Design and Control of Concrete Mixtures

An organization of cement companies to improve and extend the uses of portland cement and concrete through market development, engineering, research, education, and public affairs work.
Design and Control of Concrete Mixtures

FOURTEENTH EDITION

by Steven H. Kosmatka, Beatrix Kerkhoff, and William C. Panarese
KEYWORDS: admixtures, aggregates, air-entrained concrete, batching, cement, cold weather, curing, durability, fibers, finishing, high-performance concrete, hot weather, mixing, mixing water, mixture proportioning, placing, portland cement concrete, properties, special concrete, standards, supplementary cementing materials, tests, and volume changes.

ABSTRACT: This book presents the properties of concrete as needed in concrete construction, including strength and durability. All concrete ingredients (cementing materials, water, aggregates, admixtures, and fibers) are reviewed for their optimal use in designing and proportioning concrete mixtures. Applicable ASTM, AASHTO, and ACI standards are referred to extensively. The use of concrete from design to batching, mixing, transporting, placing, consolidating, finishing, and curing is addressed. Special concretes, including high-performance concretes, are also reviewed.


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Cover photos show ready mixed concrete being elevated by bucket and crane to the 39th floor of a high-rise building in Chicago. (69991, 70015)

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WARNING: Contact with wet (unhardened) concrete, mortar, cement, or cement mixtures can cause SKIN IRRITATION, SEVERE CHEMICAL BURNS (THIRD-DEGREE), or SERIOUS EYE DAMAGE. Frequent exposure may be associated with irritant and/or allergic contact dermatitis. Wear waterproof gloves, a long-sleeved shirt, full-length trousers, and proper eye protection when working with these materials. If you have to stand in wet concrete, use waterproof boots that are high enough to keep concrete from flowing into them. Wash wet concrete, mortar, cement, or cement mixtures from your skin immediately. Flush eyes with clean water immediately after contact. Indirect contact through clothing can be as serious as direct contact, so promptly rinse out wet concrete, mortar, cement, or cement mixtures from clothing. Seek immediate medical attention if you have persistent or severe discomfort.
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Preface and Acknowledgements

Concrete’s versatility, durability, and economy have made it the world’s most used construction material. The United States uses over 300 million cubic meters (400 million cubic yards) of ready mixed concrete each year. It is used in highways, streets, parking lots, parking garages, bridges, high-rise buildings, dams, homes, floors, sidewalks, driveways, and numerous other applications.

*Design and Control of Concrete Mixtures* has been the cement and concrete industry’s primary reference on concrete technology for over 75 years. Since the first edition was published in the early 1920s, the U.S. version has been updated 14 times to reflect advances in concrete technology and to meet the growing needs of architects, engineers, builders, concrete producers, concrete technologists, instructors, and students.

This fully revised 14th edition was written to provide a concise, current reference on concrete, including the many advances that occurred since the last edition was published in 1988. The text is backed by over 85 years of research by the Portland Cement Association. It reflects the latest information on standards, specifications, and test methods of the American Society for Testing and Materials (ASTM), the American Association of State Highway and Transportation Officials (AASHTO), and the American Concrete Institute (ACI).

Besides presenting a 50% increase in new information over the previous edition, this edition has added metric units that are currently required on most federal government projects and many state projects; AASHTO standards commonly used by many state departments of transportation are provided alongside ASTM standards; internet addresses are provided for many references for instant access; new photographs have been added to illustrate modern technology; and included are appendices on metric unit conversion, ASTM and AASHTO standards, and a listing of key concrete organizations and their web addresses. New chapters on supplementary cementing materials, fibers, and high-performance concrete have also been added.

Acknowledgements. The authors wish to acknowledge contributions made by many individuals and organizations who provided valuable assistance in the writing and publishing of the 14th edition. A special thanks to Ken Hover, Cornell University, for extensive technical recommendations; Howard “Buck” Barker, RVT Engineering Services, for photography and text edits; and Cheryl Taylor, Consultant, for months of desktop layout. Additional thanks for technical assistance, references, photography, and editorial reviews goes to: Norm MacLeod, formerly with the Canadian Portland Cement Association; Rick McGrath, Cement Association of Canada; John Bickley, John A. Bickley Associates, Ltd.; Hamid Farzam, Construction Technology Laboratories (CTL); Colin Lobo, National Ready Mixed Concrete Association; Linda Hills, CTL (SEM); Connie Field, PCA; Bill Burns, PCA; John Shaw, PCA; Basile Rabbat, PCA; Arlene Zapata, PCA (cover design); Wes Ikezoe, PCA; Richard Small, PCA; Bruce McIntosh, PCA; Susan Pepitone, PCA; Dale McFarlane, PCA; Paul Tennis, PCA; John Melander, PCA; Jamie Farny, PCA; Carmaline Spurrer, PCA; Martin McGovern, PCA; Terry Collins, PCA; Michelle Wilson, PCA; Tony Fiorato, CTL; Vagn Johansen, CTL; Wally Klemm, formerly of CTL; Peter Marlo, CTL; Ron Bard, CTL; Manoj Bharucha, CTL; Javed Bhatty, CTL; Jennifer DeStrampe, Ground Heaters, Inc.; Jim Shilstone, Shilstone Companies, Inc.; Robert E. Neal, Lehigh Portland Cement Co.; Gregory S. Barger, Ash Grove Cement Co.; Mark Luther, Holcim (US) Inc.; Fred Cohrs, Florida Rock Industries, Inc.; Phil Zacarias, Lafarge Canada, Inc.; Terry Holland, Silica Fume Association; Oscar Manz, Consultant; Jon Mullarky, FHWA; Karen Gruber, Hercules, Inc.; Mike Pistilli, Prairie Group; Sam Tyson, American Coal Ash Association; Craig Plunk, Mineral Solutions; Jim Jensen, Mineral Solutions; John Rivisto, AVR, Inc.; Charlie Misslin, County Concrete Corp.; Jamison Langdon, Cemstone; Kerry Smith, James Cape & Sons Co.; David Meyer, Lafarge North America, Inc.; Lew Kollmansberger, Mead & Hunt, Inc.; Tim Roble, Midway Concrete Corp.; George Barker, River Valley Testing Corp.; Dan Large, SI Concrete Systems; EJ Streeu, Streeu Construction; Len Swederski, Swederski Concrete Const., Inc.; Pat Bauer, W. R. Grace Co.; Darrin G. Stanke, Zenith Tech, Inc.; Scott Zignego, Zignego Ready Mix, Inc.; Peter Waismen, Trow Engineers; Mette Geiker, Technical University of Denmark; and numerous others who have provided comments and suggestions on EB001 over the past several years. Thanks also goes to ASTM, AASHTO, and ACI for the use of their material and documents referenced throughout the book.

The authors have tried to make this edition of *Design and Control of Concrete Mixtures* a concise and current reference on concrete technology. Readers are encouraged to submit comments to improve future printings and editions of this book.
Concrete is basically a mixture of two components: aggregates and paste. The paste, comprised of portland cement and water, binds the aggregates (usually sand and gravel or crushed stone) into a rocklike mass as the paste hardens because of the chemical reaction of the cement and water (Fig. 1-1). Supplementary cementitious materials and chemical admixtures may also be included in the paste.*

Aggregates are generally divided into two groups: fine and coarse. Fine aggregates consist of natural or manufactured sand with particle sizes ranging up to 9.5 mm (3/8 in.); coarse aggregates are particles retained on the 1.18 mm (No. 16) sieve and ranging up to 150 mm (6 in.) in size. The maximum size of coarse aggregate is typically 19 mm or 25 mm (3/4 in. or 1 in.). An intermediate-sized aggregate, around 9.5 mm (3/8 in.), is sometimes added to improve the overall aggregate gradation.

The paste is composed of cementitious materials, water, and entrapped air or purposely entrained air. The paste constitutes about 25% to 40% of the total volume of concrete. Fig. 1-2 shows that the absolute volume of cement is usually between 7% and 15% and the water between 14% and 21%. Air content in air-entrained concrete ranges from about 4% to 8% of the volume.

Since aggregates make up about 60% to 75% of the total volume of concrete, their selection is important. Aggregates should consist of particles with adequate strength and resistance to exposure conditions and should not contain materials that will cause deterioration of the concrete. A continuous gradation of aggregate particle sizes is desirable for efficient use of the paste. Throughout this text, it will be assumed that suitable aggregates are being used, except where otherwise noted.

The quality of the concrete depends upon the quality of the paste and aggregate, and the bond between the two. In properly made concrete, each and every particle of aggregate is completely coated with paste and all of the spaces between aggregate particles are completely filled with paste, as illustrated in Fig. 1-3.

---

* This text addresses the utilization of portland cement in the production of concrete. The term “portland cement” pertains to a calcium silicate hydraulic cement produced by heating materials containing calcium, silicon, aluminum, and iron. The term “cement” used throughout the text pertains to portland cement or blended hydraulic cement unless otherwise stated. The term “cementitious materials” means portland or blended cement, used with or without supplementary cementitious materials.
For any particular set of materials and conditions of curing, the quality of hardened concrete is strongly influenced by the amount of water used in relation to the amount of cement (Fig. 1-4). Unnecessarily high water contents dilute the cement paste (the glue of concrete). Following are some advantages of reducing water content:

- Increased compressive and flexural strength
- Lower permeability, thus lower absorption and increased watertightness
- Increased resistance to weathering
- Better bond between concrete and reinforcement
- Reduced drying shrinkage and cracking
- Less volume change from wetting and drying

The less water used, the better the quality of the concrete—provided the mixture can be consolidated properly. Smaller amounts of mixing water result in stiffer mixtures; but with vibration, stiffer mixtures can be easily placed. Thus, consolidation by vibration permits improvement in the quality of concrete.

The freshly mixed (plastic) and hardened properties of concrete may be changed by adding chemical admixtures to the concrete, usually in liquid form, during batching. Chemical admixtures are commonly used to (1) adjust setting time or hardening, (2) reduce water demand, (3) increase workability, (4) intentionally entrain air, and (5) adjust other fresh or hardened concrete properties.

After completion of proper proportioning, batching, mixing, placing, consolidating, finishing, and curing, concrete hardens into a strong, noncombustible, durable, abrasion-resistant, and watertight building material that requires little or no maintenance. Furthermore, concrete is an excellent building material because it can be formed into a wide variety of shapes, colors, and textures for use in an unlimited number of applications.

**FRESHLY MIXED CONCRETE**

Freshly mixed concrete should be plastic or semifluid and generally capable of being molded by hand. A very wet concrete mixture can be molded in the sense that it can be cast in a mold, but this is not within the definition of “plastic”—that which is pliable and capable of being molded or shaped like a lump of modeling clay.

In a plastic concrete mixture all grains of sand and pieces of gravel or stone are encased and held in suspension. The ingredients are not apt to segregate during transport; and when the concrete hardens, it becomes a homogeneous mixture of all the components. During placing, concrete of plastic consistency does not crumble but flows sluggishly without segregation.

In construction practice, thin concrete members and heavily reinforced concrete members require workable, but never soupy, mixes for ease of placement. A plastic mixture is required for strength and for maintaining homogeneity during handling and placement. While a plastic mixture is suitable for most concrete work, plasticizing admixtures may be used to make concrete more flowable in thin or heavily reinforced concrete members.
Mixing

In Fig. 1-1, the basic components of concrete are shown separately. To ensure that they are combined into a homogeneous mixture requires effort and care. The sequence of charging ingredients into a concrete mixer can play an important part in uniformity of the finished product. The sequence, however, can be varied and still produce a quality concrete. Different sequences require adjustments in the time of water addition, the total number of revolutions of the mixer drum, and the speed of revolution. Other important factors in mixing are the size of the batch in relation to the size of the mixer drum, the elapsed time between batching and mixing, and the design, configuration, and condition of the mixer drum and blades. Approved mixers, correctly operated and maintained, ensure an end-to-end exchange of materials by a rolling, folding, and kneading action of the batch over itself as concrete is mixed.

Workability

The ease of placing, consolidating, and finishing freshly mixed concrete and the degree to which it resists segregation is called workability. Concrete should be workable but the ingredients should not separate during transport and handling (Fig. 1-5).

The degree of workability required for proper placement of concrete is controlled by the placement method, type of consolidation, and type of concrete. Different types of placements require different levels of workability.

Factors that influence the workability of concrete are: (1) the method and duration of transportation; (2) quantity and characteristics of cementitious materials; (3) concrete consistency (slump); (4) grading, shape, and surface texture of fine and coarse aggregates; (5) entrained air; (6) water content; (7) concrete and ambient air temperatures; and (8) admixtures. A uniform distribution of aggregate particles and the presence of entrained air significantly help control segregation and improve workability. Fig. 1-6 illustrates the effect of casting temperature on the consistency, or slump, and potential workability of concrete mixtures.

Properties related to workability include consistency, segregation, mobility, pumpability, bleeding, and finishability. Consistency is considered a close indication of workability. Slump is used as a measure of the consistency or wetness of concrete. A low-slump concrete has a stiff consistency. If the consistency is too dry and harsh, the concrete will be difficult to place and compact and larger aggregate particles may separate from the mix. However, it should not be assumed that a wetter, more fluid mix is necessarily more workable. If the mix is too wet, segregation and honeycombing can occur. The consistency should be the driest practicable for placement using the available consolidation equipment. See Powers (1932) and Daniel (2006).

Bleeding and Settlement

Bleeding is the development of a layer of water at the top or surface of freshly placed concrete. It is caused by sedimentation (settlement) of solid particles (cement and aggregate) and the simultaneous upward migration of water (Fig. 1-7). Bleeding is normal and it should not diminish the quality of properly placed, finished, and cured concrete. Some bleeding is helpful to control plastic shrinkage cracking.
Excessive bleeding increases the water-cement ratio near the top surface; a weak top layer with poor durability may result, particularly if finishing operations take place while bleed water is present. A water pocket or void can develop under a prematurely finished surface.

After evaporation of all bleed water, the hardened surface will be slightly lower than the freshly placed surface. This decrease in height from time of placement to initial set is called settlement shrinkage.

The bleeding rate and bleeding capacity (total settlement per unit of original concrete height) increases with initial water content, concrete height, and pressure. Use of properly graded aggregate, certain chemical admixtures, air entrainment, supplementary cementitious materials, and finer cements, reduces bleeding. Concrete used to fill voids, provide support, or provide watertightness with a good bond should have low bleeding properties to avoid formation of water pockets. See Powers (1939), Steinour (1945), and Kosmatka (2006).

**Consolidation**

Vibration sets into motion the particles in freshly mixed concrete, reducing friction between them, and giving the mixture the mobile qualities of a thick fluid. The vibratory action permits use of a stiffer mixture containing a larger proportion of coarse and a smaller proportion of fine aggregate. The larger the maximum size aggregate in concrete with a well-graded aggregate, the less volume there is to fill with paste and the less aggregate surface area there is to coat with paste; thus less water and cement are needed. Concrete with an optimally graded aggregate will be easier to consolidate and place (Fig. 1-8 left). Consolidation of coarser as well as stiffer mixtures results in improved quality and economy. On the other hand, poor consolidation can result in porous, weak concrete (Fig. 1-9) with poor durability (Fig. 1-8 right).

Mechanical vibration has many advantages. Vibrators make it possible to economically place mixtures that are impractical to consolidate by hand under many conditions. As an example, Fig. 1-10 shows concrete of a stiff consistency (low slump). This concrete was mechanically vibrated in forms containing closely spaced reinforcement. With hand rodding, a much wetter consistency would have been necessary.

**Hydration, Setting Time, and Hardening**

The binding quality of portland cement paste is due to the chemical reaction between the cement and water, called hydration.

Portland cement is not a simple chemical compound, it is a mixture of many compounds. Four of these make up 90% or more of the weight of portland cement: tricalcium silicate, dicalcium silicate, tricalcium aluminate, and tetracalcium aluminoferrite. In addition to these major compounds, several others play important roles in the hydration process. Each type of portland cement contains the same four major compounds, but in different proportions.
When clinker (the kiln product that is ground to make portland cement) is examined under a microscope, most of the individual cement compounds can be identified and their amounts determined. However, the smallest grains elude visual detection. The average diameter of a typical cement particle is approximately 15 micrometers. If all cement particles were average, portland cement would contain about 300 billion particles per kilogram, but in fact there are some 16,000 billion particles per kilogram because of the broad range of particle sizes. The particles in a kilogram of portland cement have a surface area of approximately 400 square meters.

The two calcium silicates, which constitute about 75% of the weight of portland cement, react with water to form two new compounds: calcium hydroxide and calcium silicate hydrate. The latter is by far the most important cementing component in concrete. The engineering properties of concrete—setting and hardening, strength, and dimensional stability—depend primarily on calcium silicate hydrate. It is the heart of concrete.

The chemical composition of calcium silicate hydrate is somewhat variable, but it contains lime (CaO) and silicate (SiO₂) in a ratio on the order of 3 to 2. The surface area of calcium silicate hydrate is some 300 square meters per gram. In hardened cement paste, the calcium silicate hydrate forms dense, bonded aggregations between the other crystalline phases and the remaining unhydrated cement grains; they also adhere to grains of sand and to pieces of coarse aggregate, cementing everything together (Copeland and Schulz 1962).

As concrete hardens, its gross volume remains almost unchanged, but hardened concrete contains pores filled with water and air that have no strength. The strength is in the solid part of the paste, mostly in the calcium silicate hydrate and crystalline compounds.

The less porous the cement paste, the stronger the concrete. When mixing concrete, therefore, no more water than is absolutely necessary to make the concrete plastic and workable should be used. Even then, the water used is usually more than is required for complete hydration of the cement. About 0.4 grams of water per gram of cement are needed to completely hydrate cement (Powers 1948 and 1949). However, complete hydration is rare in field concrete due to a lack of moisture and the long period of time (decades) required to achieve complete hydration.

Knowledge of the amount of heat released as cement hydrates can be useful in planning construction. In winter, the heat of hydration will help protect the concrete against damage from freezing temperatures. The heat may be harmful, however, in massive structures such as dams because it may produce undesirable temperature differentials.

Knowledge of the rate of reaction between cement and water is important because it determines the rate of hardening. The initial reaction must be slow enough to allow time for the concrete to be transported and placed. Once the concrete has been placed and finished, however, rapid hardening is desirable. Gypsum, added at the cement mill when clinker is ground, acts as a regulator of the initial rate of setting of portland cement. Other factors that influence the rate of hydration include cement fineness, admixtures, amount of water added, and temperature of the materials at the time of mixing. Fig. 1-11 illustrates the setting properties of a concrete mixture at different temperatures.

### HARDENED CONCRETE

#### Curing

Increase in strength with age continues provided (1) unhydrated cement is still present, (2) the concrete remains moist or has a relative humidity above approximately 80% (Powers 1948), (3) the concrete temperature remains favorable, and (4) sufficient space is available for hydration products to form. When the relative humidity within the concrete drops to about 80%, or the temperature of the concrete drops below freezing, hydration and strength gain virtually stop. Fig. 1-12 illustrates the relationship between strength gain and moist curing, while Fig. 1-13 illustrates the relationship between strength gain and curing temperature.

If concrete is resaturated after a drying period, hydration is resumed and strength will again increase. However, it is best to moist-cure concrete continuously from the time it is placed until it has attained the desired quality; once concrete has dried out it is difficult to resaturate. Fig. 1-14 illustrates the long-term strength gain of concrete in an outdoor exposure. Outdoor exposures often continue to provide moisture through ground contact and rainfall. Indoor concretes often dry out after curing and do not continue to gain strength (Fig. 1-12).
Forexample, as mentioned, concrete must continue to hold enough moisture throughout the curing period for the cement to hydrate to the extent that desired properties are achieved. Freshly cast concrete usually has an abundance of water, but as drying progresses from the surface inward, strength gain will continue at each depth only as long as the relative humidity at that point remains above 80%.

A common illustration of this is the surface of a concrete floor that has not had sufficient moist curing. Because it has dried quickly, concrete at the surface is weak and traffic on it creates dusting. Also, when concrete dries, it shrinks as it loses water (Fig. 1-15), just as wood and clay do (though not as much). Drying shrinkage is a primary cause of cracking, and the width of cracks is a function of the degree of drying, spacing or frequency of cracks, and the age at which the cracks occur.

While the surface of a concrete element will dry quite rapidly, it takes a much longer time for concrete in the interior to dry. Fig. 1-15 (top) illustrates the rate of drying at various depths within concrete cylinders exposed to laboratory air. Field concrete elements would have different drying profiles due to environmental conditions, size effects, and concrete properties.

The moisture content of concrete depends on the concrete's constituents, original water content, drying conditions, and the size of the concrete element (Hedenblad 1997 and 1998). After several months of drying in air with a relative humidity of 50% to 90%, moisture content is about 1% to 2% by mass of the concrete. Fig. 1-15 illustrates moisture loss and resulting shrinkage.

Size and shape of a concrete member have an important bearing on the rate of drying. Concrete elements with large surface area in relation to volume (such as floor slabs)
dry faster than voluminous concrete members with relatively small surface areas (such as bridge piers).

Many other properties of hardened concrete also are affected by its moisture content; these include elasticity, creep, insulating value, fire resistance, abrasion resistance, electrical conductivity, frost resistance, scaling resistance, and resistance to alkali-aggregate reactivity.

**Strength**

Compressive strength may be defined as the measured maximum resistance of a concrete specimen to axial load-

It is generally expressed in megapascals (MPa) or pounds per square inch (psi) at an age of 28 days. One megapascal equals the force of one newton per square millimeter (N/mm²) or 1,000,000 N/m². Other test ages are also used; however, it is important to realize the relationship between the 28-day strength and other test ages. Seven-day strengths are often estimated to be about 75% of the 28-day strength and 56-day and 90-day strengths are about 10% to 15% greater than 28-day strengths as shown in Fig. 1-16. The specified compressive strength is designated by the symbol $f'_c$, and ideally is exceeded by the actual compressive strength, $f_c$.

The compressive strength that a concrete achieves, $f_c$, results from the water-cement ratio (or water-cementitious materials ratio), the extent to which hydration has progressed, the curing and environmental conditions, and the age of the concrete. The relationship between strength and water-cement ratio has been studied since the late 1800s and early 1900s (Feret 1897 and Abrams 1918). Fig. 1-17 shows compressive strengths for a wide range of concrete mixtures and water-cement ratios at an age of 28 days. Note that strengths increase as the water-cement ratios decrease. These factors also affect flexural and tensile strengths and bond of concrete to steel.

The water-cement ratio compressive strength relationships in Fig. 1-17 are for typical non-air-entrained concretes. When more precise values for concrete are required, graphs should be developed for the specific materials and mix proportions to be used on the job.

For a given workability and a given amount of cement, air-entrained concrete requires less mixing water than non-air-entrained concrete. The lower water-cement ratio possible for air-entrained concrete tends to offset the somewhat lower strengths of air-entrained concrete, particularly in lean to medium cement content mixes.
The torsional strength for concrete is related to the modulus of rupture and the dimensions of the concrete element. Hsu (1968) presents torsional strength correlations.

Shear strength–compressive strength relationships are discussed in the ACI 318 building code. The correlation between compressive strength and flexural, tensile, torsional, and shear strength varies with concrete ingredients and environment.

Modulus of elasticity, denoted by the symbol $E$, may be defined as the ratio of normal stress to corresponding strain for tensile or compressive stresses below the proportional limit of a material. For normal-weight concrete, $E$ ranges from 14,000 to 41,000 MPa (2 to 6 million psi) and can be approximated as 5,000 times the square root of the compressive strength in megapascals (57,000 times the square root of the compressive strength in pounds per square inch). Like other strength relationships, the modulus of elasticity to compressive strength relationship is ingredient specific and should be verified in a laboratory (Wood 1992).

Density

Conventional concrete, normally used in pavements, buildings, and other structures, has a density (unit weight) in the range of 2200 to 2400 kg/m³ (137 to 150 lb/ft³). The density of concrete varies, depending on the amount and density of the aggregate, the amount of air that is entrapped or purposely entrained, and the water and cement contents, which in turn are influenced by the maximum size of the aggregate. Reducing the cement paste content (increasing aggregate volume) increases density. Values of the density of fresh concrete are given in Table 1-1. In the design of reinforced concrete structures, the combination of conventional concrete and reinforcing steel is commonly assumed to weigh 2400 kg/m³ (150 lb/ft³).
The weight of dry concrete equals the weight of the freshly mixed concrete ingredients less the weight of mix water that evaporates into the air. Some of the mix water combines chemically with the cement during the hydration process, converting the cement into cement gel. Also, some of the water remains tightly held in pores and capillaries and does not evaporate under normal conditions. The amount of mix water that will evaporate from concrete exposed to ambient air at 50% relative humidity is about 1/2% to 3% of the concrete weight; the actual amount depends on initial water content of the concrete, absorption characteristics of the aggregates, and size and shape of the concrete element.

Aside from conventional concrete, there is a wide spectrum of special concretes to meet various needs; they range from lightweight insulating concretes with a density of as little as 240 kg/m³ (15 lb/ft³) to heavyweight concrete with a density of up to 6000 kg/m³ (375 lb/ft³) used for counterweights or radiation shielding.

### Permeability and Watertightness

Concrete used in water-retaining structures or exposed to weather or other severe exposure conditions must be virtually impermeable or watertight. Watertightness is often referred to as the ability of concrete to hold back or retain water without visible leakage. Permeability refers to the amount of water migration through concrete when the water is under pressure or to the ability of concrete to resist penetration by water or other substances (liquid, gas, or ions). Generally, the same properties of concrete that make it less permeable also make it more watertight.

The overall permeability of concrete to water is a function of: (1) the permeability of the paste; (2) the permeability and gradation of the aggregate; (3) the quality of the paste and aggregate transition zone; and (4) the relative proportion of paste to aggregate. Decreased permeability improves concrete’s resistance to freezing and thawing, resaturation, sulfate, and chloride-ion penetration, and other chemical attack.

The permeability of the paste is particularly important because the paste envelops all constituents in the concrete. Paste permeability is related to water-cement ratio, degree of cement hydration, and length of moist curing. A low-permeability concrete requires a low water-cement ratio and an adequate moist-curing period. Air entrainment aids watertightness but has little effect on permeability. Permeability increases with drying.

The permeability of mature hardened cement paste kept continuously moist ranges from 0.1 x 10⁻ⁱ² to 120 x 10⁻¹² cm per sec. for water-cement ratios ranging from 0.3 to 0.7 (Powers and others 1954). The permeability of rock commonly used as concrete aggregate varies from approximately 1.7 x 10⁻⁸ to 3.5 x 10⁻¹³ cm per sec. The permeability of mature, good-quality concrete is approximately 1 x 10⁻¹⁰ cm per sec.

The relationship between permeability, water-cement ratio, and initial curing for 100 x 200-mm (4 x 8-in.) cylindrical concrete specimens tested after 90 days of air drying

<table>
<thead>
<tr>
<th>Maximum size of aggregate, mm</th>
<th>Air content, percent</th>
<th>Water, kg/m³</th>
<th>Cement, kg/m³</th>
<th>Density, kg/m³**</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>2.55</td>
<td>2.60</td>
<td>2.65</td>
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<tr>
<td>19</td>
<td>6.0</td>
<td>168</td>
<td>336</td>
<td>2194</td>
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<tr>
<td>37.5</td>
<td>4.5</td>
<td>145</td>
<td>291</td>
<td>2259</td>
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<tr>
<td>75</td>
<td>3.5</td>
<td>121</td>
<td>242</td>
<td>2307</td>
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</tbody>
</table>

<table>
<thead>
<tr>
<th>Maximum size of aggregate, in.</th>
<th>Air content, percent</th>
<th>Water, lb/ft³</th>
<th>Cement, lb/ft³</th>
<th>Density, lb/ft³**</th>
</tr>
</thead>
<tbody>
<tr>
<td>¾</td>
<td>6.0</td>
<td>283</td>
<td>566</td>
<td>137</td>
</tr>
<tr>
<td>1½</td>
<td>4.5</td>
<td>245</td>
<td>490</td>
<td>141</td>
</tr>
<tr>
<td>3</td>
<td>3.5</td>
<td>204</td>
<td>408</td>
<td>144</td>
</tr>
</tbody>
</table>

* Source: Bureau of Reclamation 1981, Table 4.
** Air-entrained concrete with indicated air content.
† On saturated surface-dry basis. Multiply relative density by 1000 to obtain density of aggregate particles in kg/m³.

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### Table 1-1. Observed Average Density of Fresh Concrete (SI Units)*

<table>
<thead>
<tr>
<th>Maximum size of aggregate, mm</th>
<th>Air content, percent</th>
<th>Water, kg/m³</th>
<th>Cement, kg/m³</th>
<th>Density, kg/m³**</th>
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<td>75</td>
<td>3.5</td>
<td>121</td>
<td>242</td>
<td>2307</td>
</tr>
</tbody>
</table>

* Source: Bureau of Reclamation 1981, Table 4.
** Air-entrained concrete with indicated air content.
† On saturated surface-dry basis. Multiply specific gravity by 62.4 to obtain density of aggregate particles in lb/ft³.
greatly reduced or completely removed, producing a high volumne of air voids. Pervious concrete has been used in tennis courts, pavements, parking lots, greenhouses, and drainage structures. Pervious concrete has also been used in buildings because of its thermal insulation properties.

**Abbrasion Resistance**

Floors, pavements, and hydraulic structures are subjected to abrasion; therefore, in these applications concrete must have a high abrasion resistance. Test results indicate that abrasion resistance is closely related to the compressive strength of concrete. Strong concrete has more resistance to abrasion.
abrasion than does weak concrete. Since compressive strength depends on water-cement ratio and curing, a low water-cement ratio and adequate curing are necessary for abrasion resistance. The type of aggregate and surface finish or treatment used also have a strong influence on abrasion resistance. Hard aggregate is more wear resistant than soft aggregate and a steel-troweled surface resists abrasion better than a surface that had not been troweled.

Fig. 1-22 shows results of abrasion tests on concretes of different compressive strengths and aggregate types. Fig. 1-23 illustrates the effect hard steel troweling and surface treatments, such as metallic or mineral aggregate surface hardeners, have on abrasion resistance. Abrasion tests can be conducted by rotating steel balls, dressing wheels, or disks under pressure over the surface (ASTM C779). One type of test apparatus is pictured in Fig. 1-24. Other types of abrasion tests are also available (ASTM C418 and C944).

**Volume Stability and Crack Control**

Hardened concrete changes volume due to changes in temperature, moisture, and stress. These volume or length changes may range from about 0.01% to 0.08%. Thermal volume changes of hardened concrete are about the same as those for steel.

Concrete under stress will deform elastically. Sustained stress will result in additional deformation called creep. The rate of creep (deformation per unit of time) decreases with time.

Concrete kept continually moist will expand slightly. When permitted to dry, concrete will shrink. The primary factor influencing the amount of drying shrinkage is the water content of the freshly mixed concrete. Drying shrinkage increases directly with increases in this water content. The amount of shrinkage also depends upon several other...
factors, such as: (1) the amount of aggregate used; (2) properties of the aggregate; (3) size and shape of the concrete element; (4) relative humidity and temperature of the ambient air; (5) method of curing; (6) degree of hydration; and (7) time.

Two basic causes of cracks in concrete are: (1) stress due to applied loads and (2) stress due to drying shrinkage or temperature changes when concrete is restrained.

Drying shrinkage is an inherent, unavoidable property of concrete; therefore, properly positioned reinforcing steel is used to reduce crack widths, or joints are used to predetermine and control the location of cracks. Thermal stress due to fluctuations in ambient temperature also can cause cracking, particularly at an early age.

Concrete shrinkage cracks can occur because of restraint. When drying shrinkage occurs and there is no restraint, the concrete does not crack. Restriction comes from several sources. Drying shrinkage is always greater near the surface of concrete; the moist inner portions restrain the concrete near the surface, which can cause cracking. Other sources of restraint are reinforcing steel embedded in concrete, the interconnected parts of a concrete structure, and the friction of the subgrade on which concrete is placed.

**Joints.** Joints are the most effective method of controlling unsightly cracking. If a sizable expanse of concrete (a wall, slab, or pavement) is not provided with properly spaced joints to accommodate drying shrinkage and temperature contraction, the concrete will crack in a random manner.

Contraction (shrinkage control) joints are grooved, formed, or sawed into sidewalks, driveways, pavements, floors, and walls so that cracking will occur in these joints rather than in a random manner. Contraction joints permit movement in the plane of a slab or wall. They extend to a depth of approximately one-quarter the concrete thickness.

Isolation joints separate a concrete placement from other parts of a structure and permit horizontal and vertical movements. They should be used at the junction of floors with walls, columns, footings, and other points where restraint can occur. They extend the full depth of slabs and include a premolded joint filler.

Construction joints occur where concrete work is concluded for the day; they separate areas of concrete placed at different times. In slabs-on-ground, construction joints usually align with, and function as, control or isolation joints. They may require dowels for load transfer.

**DURABILITY**

The durability of concrete may be defined as the ability of concrete to resist weathering action, chemical attack, and abrasion while maintaining its desired engineering properties. Different concretes require different degrees of durability depending on the exposure environment and the properties desired. The concrete ingredients, proportioning of those ingredients, interactions between the ingredients, and placing and curing practices determine the ultimate durability and life of the concrete.

**Resistance to Freezing and Thawing**

Concrete used in structures and pavements is expected to have long life and low maintenance. It must have good durability to resist anticipated exposure conditions. The most potentially destructive weathering factor is freezing and thawing while the concrete is wet, particularly in the presence of deicing chemicals. Deterioration is caused by the freezing of water and subsequent expansion in the paste, the aggregate particles, or both.

With air entrainment, concrete is highly resistant to this type of deterioration as shown in Fig. 1-25. During freezing, the water displaced by ice formation in the paste is accommodated so that it is not disruptive; the microscopic air bubbles in the paste provide chambers for the water to enter and thus relieve the hydraulic pressure generated.

When freezing occurs in concrete containing saturated aggregate, disruptive hydraulic pressures can also be generated within the aggregate. Water displaced from the aggregate particles during the formation of ice cannot escape fast enough to the surrounding paste to relieve pressure. However, under most exposure conditions, a good-quality paste (low water-cement ratio) will prevent most aggregate particles from becoming saturated. Also, if the paste is air-entrained, it will accommodate the small amounts of excess water that may be expelled from aggregates, thus protecting the concrete from freeze-thaw damage.

Fig. 1-26 illustrates, for a range of water-cement ratios, that (1) air-entrained concrete is much more resistant to freeze-thaw cycles than non-air-entrained concrete, (2) concrete with a low water-cement ratio is more durable than concrete with a high water-cement ratio.
Air-entrained concrete with a low water-cement ratio and an air content of 5% to 8% will withstand a great number of cycles of freezing and thawing without distress (Woods 1956).

Freeze-thaw durability can be determined by laboratory test procedure ASTM C666, Standard Test Method for Resistance of Concrete to Rapid Freezing and Thawing (AASHTO T 161). From the test, a durability factor is calculated that reflects the number of cycles of freezing and thawing required to produce a certain amount of deterioration. Deicer-scaling resistance can be determined by ASTM C672, Standard Test Method for Scaling Resistance of Concrete Surfaces Exposed to Deicing Chemicals.

Alkali-Aggregate Reactivity

Alkali-aggregate reactivity is a type of concrete deterioration that occurs when the active mineral constituents of some aggregates react with the alkali hydroxides in the concrete. The reactivity is potentially harmful only when it produces significant expansion. Alkali-aggregate reactivity occurs in two forms—alkali-silica reaction (ASR) and alkali-carbonate reaction (ACR). Alkali-silica reaction is of more concern than alkali-carbonate reaction because the occurrence of aggregates containing reactive silica minerals is more common.

Indications of the presence of alkali-aggregate reactivity may be a network of cracks, closed or spalling joints, or displacement of different portions of a structure (Fig. 1-27). Because deterioration due to alkali-aggregate reaction is a slow process, the risk of catastrophic failure is low. Alkali-aggregate reaction can cause serviceability problems and exacerbate other deterioration mechanisms, such as those that occur in frost, deicer, or sulfate exposures.

Current practices to control alkali-silica reactivity include the use of supplementary cementitious materials or blended cements proven by testing to control the reaction. Supplementary cementitious materials include fly ash, ground granulated blast-furnace slag, silica fume, and natural pozzolans (Fig. 1-28). Blended cements also contain these materials to control alkali-silica reactivity. This practice allows the use of locally available aggregates and cementitious materials. Reduction of the alkali content of the concrete can also control the reaction.

Use of supplementary cementitious materials or blended cements does not control alkali-carbonate reaction. Fortunately this reaction is rare. If aggregate testing indicates that an aggregate is susceptible to alkali-carbonate reactivity, the reaction can be controlled through the use of selective quarrying, aggregate blending, reducing maximum aggregate size, or using special chemical compounds that inhibit the reaction.

![Fig. 1-27. Cracking, joint closing, spalling, and lateral offset are caused by severe alkali-silica reactivity in this parapet wall. (IMG12295)](image-url)
A CI 318 building code provides reinforcing steel cover requirements for different exposures.

**Chloride Resistance and Steel Corrosion**

Concrete protects embedded steel from corrosion through its highly alkaline nature. The high pH environment in concrete (usually greater than 12.5) causes a passive and noncorroding protective oxide film to form on steel. However, the presence of chloride ions from deicers or seawater can destroy or penetrate the film. Once the chloride corrosion threshold (about 0.15% water-soluble chloride by mass of cement) is reached, an electric cell is formed along the steel or between steel bars and the electrochemical process of corrosion begins. Some steel areas along the bar become the anode, discharging current in the electric cell; from there the iron goes into solution. Steel areas that receive current are the cathodes where hydroxide ions are formed. The iron and hydroxide ions form iron hydroxide, Fe\(\text{OH}\), which further oxidizes to form rust (iron oxide). Rusting is an expansive process—rust expands up to four times its original volume—which induces internal stress and eventual spalling of the concrete over reinforcing steel. The cross-sectional area of the steel can also be significantly reduced.

Once it starts, the rate of steel corrosion is influenced by the concrete’s electrical resistivity, moisture content, and the rate at which oxygen migrates through the concrete to the steel. Chloride ions alone can also penetrate the passive film.
on the reinforcement; they combine with iron ions to form a soluble iron chloride complex that carries the iron into the concrete for later oxidation (rust) (Whiting 1997, Taylor, Whiting, and Nagi 2000, and Whiting, Taylor and Nagi 2002).

The resistance of concrete to chloride is good; however, it can be increased by using a low water-cement ratio (0.40), at least seven days of moist curing, and supplementary cementitious materials, such as silica fume, to reduce permeability. Increasing the concrete cover over the steel also helps slow down the migration of chlorides.

Other methods of reducing steel corrosion include the use of corrosion inhibiting admixtures (ASTM C1582), epoxy-coated reinforcing steel (ASTM D3963 or AASHTO M 284), surface treatments, concrete overlays, and cathodic protection.

Epoxy-coated reinforcing steel works by preventing chloride ions from reaching the steel (Fig. 1-30). Surface treatments and concrete overlays attempt to stop or reduce chloride ion penetration at the concrete surface. Silanes, siloxanes, methacrylates, epoxies, and other materials are used as surface treatments.

Impermeable materials, such as most epoxies, should not be used on slabs on ground or other concrete where moisture can freeze under the coating. The freezing water can cause surface delamination under the impermeable coating. Latex-modified portland cement concrete, low-slump concrete, and concrete with silica fume are used in overlays to reduce chloride-ion ingress.

Cathodic protection methods reverse the corrosion current flow through the concrete and reinforcing steel. This is done by inserting a nonstructural anode in the concrete and forcing the steel to be the cathode by electrically charging the system. The anode is connected to the positive pole of a rectifier. Since corrosion occurs where the current leaves the steel, the steel cannot corrode if it is receiving the induced current.

Chloride present in plain concrete (not containing steel) is generally not a durability concern.

Corrosion of nonferrous metals in concrete is discussed by Kerkhoff (2007).

### Chemical Resistance

Portland cement concrete is resistant to most natural environments; however, concrete is sometimes exposed to substances that can attack and cause deterioration. Concrete in chemical manufacturing and storage facilities is especially prone to chemical attack. The effect of sulfates and chlorides is discussed in this chapter. Acids attack concrete by dissolving cement paste and calcareous aggregates. In addition to using concrete with a low permeability, surface treatments can be used to keep aggressive substances from coming in contact with concrete. Effects of Substances on Concrete and Guide to Protective Treatments (Kerkhoff 2007) discusses the effects of hundreds of chemicals on concrete and provides a list of treatments to help control chemical attack.

### Sulfates and Salt Crystallization

Excessive amounts of sulfates in soil or water can attack and destroy a concrete that is not properly designed. Sulfates (for example calcium sulfate, sodium sulfate, and magnesium sulfate) can attack concrete by reacting with hydrated compounds in the hardened cement paste. These reactions can induce sufficient pressure to disrupt the cement paste, resulting in disintegration of the concrete (loss of paste cohesion and strength). Calcium sulfate attacks calcium aluminate hydrate and forms ettringite. Sodium sulfate reacts with calcium hydroxide and calcium aluminate hydrate forming ettringite and gypsum. Magnesium sulfate attacks in a manner similar to sodium sulfate and forms ettringite, gypsum, and also brucite (magnesium hydroxide). Brucite forms primarily on the concrete surface; it consumes calcium hydroxide, lowers the pH of the pore solution, and then decomposes the calcium silicate hydrates (Santhanam and others 2001).

Thaumaisite may form during sulfate attack in moist conditions at temperatures usually between 0°C and 10°C (32°F to 50°F) and it occurs as a result of a reaction between calcium silicate hydrate, sulfate, calcium carbonate, and water (Report of the Thaumaisite Expert Group 1999). In concretes where deterioration is associated with excess thaumaisite formation, cracks can be filled with thaumaisite and haloes of white thaumaisite are present around aggregate particles. At the concrete/soil interface the surface concrete layer can be “mushy” with complete replacement of the cement paste by thaumaisite (Hobbs 2001).

Like natural rock formations such as limestone, porous concrete is susceptible to weathering caused by salt crystallization. These salts may or may not contain sulfates and they may or may not react with the hydrated compounds in concrete. Examples of salts known to cause weathering of field concrete include sodium carbonate and sodium sulfate (laboratory studies have also related saturated solutions of calcium chloride and other salts to concrete deterioration). The greatest damage occurs with drying of saturated solutions of these salts, often in an environment with specific cyclic changes in relative humidity and temperature that alter mineralogical phases. In permeable concrete exposed to drying conditions, salt solutions

---

**Fig. 1-30. Epoxy-coated reinforcing steel used in a bridge deck. (IMG12498)**
For the best defense against external sulfate attack: (1) design concrete with a low water to cementitious materials ratio (around 0.4), and (2) use cements specially formulated for sulfate environments, such as ASTM C150 (AASHTO M 85) Type II or V cements, C595 (AASHTO M 240) moderate sulfate resistant cements, or C1157 Types MS or HS. The superior sulfate resistance of ASTM C150 Type II and V cements is shown in Fig. 1-33.

Sulfate attack and salt crystallization are more severe at locations where the concrete is exposed to wetting and drying cycles, than continuously wet exposures. This is often seen in concrete posts where the concrete has deteriorated only a few centimeters above and below the soil line. The portion of concrete deep in the soil (where it is continuously wet) is in good condition (Fig. 1-31 and 1-32). However, if the sulfate exposure is severe enough, with time even the continuously moist sections can be attacked by sulfates if the concrete is not properly designed.

For the best defense against external sulfate attack:

1. Design concrete with a low water to cementitious materials ratio (around 0.4).
2. Use cements specially formulated for sulfate environments, such as ASTM C150 (AASHTO M 85) Type II or V cements, C595 (AASHTO M 240) moderate sulfate resistant cements, or C1157 Types MS or HS. The superior sulfate resistance of ASTM C150 Type II and V cements is shown in Fig. 1-33.

Fig. 1-31. Sulfate attack is often the most severe at the location of the most wetting and drying, which is usually near the soil line. Here concrete posts have been attacked by sulfates near the soil line. Also see the inset in Fig. 1-32. The concrete is in better condition deep within the soil where it is moist. (IMG12497)

Fig. 1-32. Concrete beams after seven years of exposure to sulfate-rich wet soil in a Sacramento, California, test plot. The beams in better condition have low water-cementitious materials ratios, and most have sulfate resistant cement. The inset shows two of the beams tipped on their side to expose decreasing levels of deterioration with depth and moisture level. (IMG12296, IMG12496)

Fig. 1-33. Average 16-yr ratings of concrete beams in sulfate soils for three portland cements at various water-cement ratios (Stark 2002).
Seawater Exposures

Concrete has been used in seawater exposures for decades with excellent performance. However, special care in mix design and material selection is necessary for these severe environments. A structure exposed to seawater or seawater spray is most vulnerable in the tidal or splash zone where there are repeated cycles of wetting and drying and/or freezing and thawing. Sulfates and chlorides in seawater require the use of low permeability concrete to minimize steel corrosion and sulfate attack (Fig. 1-34).

A cement resistant to moderate sulfate exposure is helpful. Portland cements with tricalcium aluminate (C₃A) contents that range from 4% to 10% have been found to provide satisfactory protection against seawater sulfate attack, as well as protection against corrosion of reinforcement by chlorides. Proper concrete cover over reinforcing steel must be provided (see ACI 318). Water-cementitious material ratios should not exceed 0.40. In northern climates, the concrete must be properly air entrained with at least 6% air. High-strength concrete should be considered where large ice formations abrade the structure. See Stark (1995 and 2001), Farny (1996), and Kerkhoff (2007).

Ettringite crystals in air voids and cracks are typically two to four micrometers in cross section and 20 to 30 micrometers long. Under conditions of extreme deterioration or decades in a moist environment, the white ettringite crystals can appear to completely fill voids or cracks. Both water and sufficient space must be present for the crystals to form. Cracks can form due to damage caused by frost action, alkali-aggregate reactivity, drying shrinkage, thermal effects, strain due to excessive stress, or other mechanisms.

Ettringite and Heat Induced Delayed Expansion

Ettringite, one form of calcium sulfoaluminate, is found in all portland cement paste. Calcium sulfate sources, such as gypsum, are added to portland cement during final grinding at the cement mill to prevent rapid setting and improve strength development. Sulfate is also present in supplementary cementitious materials and admixtures. Gypsum and other sulfate compounds react with calcium aluminate in cement to form ettringite within the first few hours after mixing with water. Most of the sulfate in cement is normally consumed to form ettringite or calcium monosulfate within 24 hours (Klemm and Miller 1997). At this stage ettringite is uniformly and discretely dispersed throughout the cement paste at a submicroscopic level (less than a micrometer in cross-section). This ettringite is often called primary ettringite.

If concrete is exposed to moisture for long periods of time (many years), the ettringite can slowly dissolve and reform in less confined locations. Upon microscopic examination, harmless white needle-like crystals of ettringite can be observed lining air voids. This reformed ettringite is usually called secondary ettringite (Fig. 1-35).

Concrete deterioration accelerates the rate at which ettringite leaves its original location in the paste to go into solution and recrystallize in larger spaces such as air voids or cracks. Both water and sufficient space must be present for the crystals to form. Cracks can form due to damage caused by frost action, alkali-aggregate reactivity, drying shrinkage, thermal effects, strain due to excessive stress, or other mechanisms.

Ettringite and Heat Induced Delayed Expansion. Heat induced delayed expansion (HIDE)—also called delayed ettringite formation (DEF)—refers to a rare condition of internal sulfate attack* in which mature concretes undergo expansion and

*Internal sulfate attack refers to deterioration mechanisms occurring in connection with sulfate that is in the concrete at the time of placement.
cracking. Only concretes of particular chemical makeup are affected when they have achieved high temperatures, usually after the first few hours of placement (between 70°C and 100°C [158°F to 212°F] depending on the concrete ingredients and the time the temperature is achieved after casting). This can occur because the high temperature decomposes any initial ettringite formed and holds the sulfate and alumina tightly in the calcium silicate hydrate (C-S-H) gel of the cement paste. The normal formation of ettringite is thus impeded.

In the presence of moisture, sulfate desorbs from the confines of the C-S-H and reacts with calcium monosulfoaluminate to form ettringite in cooled and hardened concrete. After months or years of desorption, ettringite forms in confined locations within the paste. Such ettringite can exert crystallization pressures because it forms in a limited space under supersaturation. One theory: since concrete is rigid and if there are insufficient voids to accommodate the ettringite volume increase, expansion and cracks can occur. In addition, some of the initial (primary) ettringite may be converted to monosulfoaluminate at high temperatures and upon cooling revert back to ettringite. Because ettringite takes up more space than monosulfoaluminate from which it forms, the transformation is an expansive reaction. The mechanism causing expansion in the paste is not fully understood at this time; the true influence of ettringite formation on this expansion is still being investigated. Some research indicates that there is little relationship between ettringite formation and expansion.

As a result of an increase in paste volume, separation of the paste from the aggregates is usually observed with heat induced delayed expansion. It is characterized by the development of rims of ettringite around the aggregates (Fig. 1-36). At early stages of heat induced delayed expansion, the voids between paste and aggregate are empty (no ettringite present). It should be noted that concrete can sustain a small amount of this expansion without harm. Only extreme cases result in cracking, and often heat induced delayed expansion is associated with other deterioration mechanisms, especially alkali-silica reactivity.

Only concretes in massive elements that retain the heat of hydration or elements exposed to very high temperatures at an early age are at risk of HIDE; and of these only a few have the chemical makeup or temperature profile to cause detrimental expansion. Normal sized concrete elements cast and maintained near ambient temperatures cannot experience HIDE when sound materials are used.

Fly ash and slag may help control heat induced delayed expansion, along with control over early-age temperature development. For more information, see Lerch (1945), Day (1992), Klemm and Miller (1997), Thomas (1998), and Famy (1999).


Thomas, M. D. A., *Delayed Ettringite Formation in Concrete—Recent Developments and Future Directions*, University of Toronto, 1998, 45 pages.


CHAPTER 2
Portland, Blended, and Other Hydraulic Cements

Portland cements are hydraulic cements composed primarily of hydraulic calcium silicates (Fig. 2-1). Hydraulic cements set and harden by reacting chemically with water. During this reaction, called hydration, cement combines with water to form a stonelike mass. When the paste (cement and water) is added to aggregates (sand and gravel, crushed stone, or other granular material) it acts as an adhesive and binds the aggregates together to form concrete, the world’s most versatile and most widely used construction material.

Hydration begins as soon as cement comes in contact with water. Each cement particle forms a fibrous growth on its surface that gradually spreads until it links up with the growth from other cement particles or adheres to adjacent substances. This fibrous build up results in progressive stiffening, hardening, and strength development. The stiffening of concrete can be recognized by a loss of workability that usually occurs within three hours of mixing, but is dependent upon the composition and fineness of the cement, any admixtures used, mixture proportions, and temperature conditions. Subsequently, the concrete sets and becomes hard.

Hydration continues as long as favorable moisture and temperature conditions exist (curing) and space for hydration products is available. As hydration continues, concrete becomes harder and stronger. Most of the hydration and strength development take place within the first month, but then continues, though more slowly, for a long time with adequate moisture and temperature. Continuous strength increases exceeding 30 years have been recorded (Washa and Wendt 1975 and Wood 1992).

THE BEGINNING OF AN INDUSTRY

Early builders used clay to bind stones together into a solid structure for shelter and protection. The oldest concrete discovered so far dates from around 7000 BC and was found in 1985 when a concrete floor was uncovered during the construction of a road at Yiftah El in Galilee, Israel. It consisted of a lime concrete, made from burning limestone to produce quicklime, which when mixed with water and stone, hardened to form concrete (Brown 1996 and Auburn 2000).

A cementing material was used between the stone blocks in the construction of the Great Pyramid at Giza in Ancient Egypt around 2500 BC. Some reports say it was a lime mortar while others say the cementing material was made from burnt gypsum. By 500 BC, the art of making lime-based mortar arrived in Ancient Greece. The Greeks used lime-based materials as a binder between stone and brick and as a rendering material over porous limestone commonly used in the construction of their temples and palaces.

Examples of early Roman concrete have been found dating back to 300 BC. The very word concrete is derived from the Latin word “concretus” mean-
1. Stone is first reduced to 125 mm (5 in.) size, then to 20 mm (¾ in.), and stored.

2. Raw materials are ground to powder and blended.

2. Raw materials are ground, mixed with water to form slurry, and blended.

3. Burning changes raw mix chemically into cement clinker.

4. Clinker with gypsum is ground into portland cement and shipped.

Fig. 2-3. Steps in the traditional manufacture of portland cement.
1. Stone is first reduced to 125 mm (5 in.) size, then to 20 mm (¾ in.), and stored.

2. Raw materials are ground to powder and blended.

3. Burning changes raw mix chemically into cement clinker. Note four-stage preheater, flash furnaces, and shorter kiln.

4. Clinker with gypsum is ground into portland cement and shipped.

Fig. 2-4. Steps in the modern dry-process manufacture of portland cement.
ing grown together or compounded. The Romans perfected the use of pozzolan as a cementing material. Sometime during the second century BC the Romans quarried a volcanic ash near Pozzuoli; thinking it was sand, they mixed it with lime and found the mixture to be much stronger than they had produced previously. This discovery was to have a significant effect on construction. The material was not sand, but a fine volcanic ash containing silica and alumina, which when combined chemically with lime, produced what became known as pozzolanic cement. This material was used by builders of the famous Roman walls, aqueducts and other historic structures including the Theatre at Pompeii (seating 20,000 spectators), and the Colosseum and Pantheon in Rome. Pozzolan seems to have been ignored during the Middle Ages until the quality of cementing materials deteriorated. The practice of burning lime and the use of pozzolan was not introduced again until the 1300s.

Efforts to determine why some limes possess hydraulic properties while others (those made from essentially pure limestones) do not were not made until the 18th century. John Smeaton, often referred to as the “father of civil engineering in England,” concentrated his work in this field. He found that an impure, soft limestone, containing clay minerals made the best hydraulic cement. This combined with a pozzolan, imported from Italy, was used in his project to rebuild the Eddystone Lighthouse in the English Channel, southwest of Plymouth England. The project took three years to complete and began operation in 1759; it was recognized as a significant accomplishment in the development of the cement industry. A number of discoveries followed as efforts within a growing natural cement industry were now directed to the production of a consistent quality material.

The difference between a hydraulic lime and natural cement is a function of the temperature attained during calcination. Furthermore, a hydraulic lime can hydrate in a “lump” form, whereas natural cements must be crushed and finely ground before hydration can take place. Natural cement is stronger than hydraulic lime but weaker than portland cement. Natural cement was manufactured in Rosendale, New York in the early 1800s (White 1820) and was first used to build the Erie Canal in 1818 (Snell and Snell 2000).

The development of portland cement was the result of persistent investigation by science and industry to produce a superior quality natural cement. The invention of portland cement is generally credited to Joseph Aspdin, an English mason. In 1824, he obtained a patent for his product, which he named portland cement because when set, it resembled the color of the natural limestone quarried on the Isle of Portland in the English Channel (Fig. 2-2) (Aspdin 1824). The name has endured and is used throughout the world, with many manufacturers adding their own trade or brand names.

Aspdin was the first to prescribe a formula for portland cement and the first to have his product patented. However, in 1845, L. C. Johnson, of White and Sons, Swanscombe, England, claimed to have “burned the cement raw materials with unusually strong heat until the mass was nearly vitrified,” producing a portland cement as we now know it. This cement became the popular choice during the middle of the 19th century and was exported from England to various parts of the world. Production also began in Belgium, France, and Germany about the same time and export of these products from Europe to North America began about 1865. The first recorded shipment of portland cement to the United States was in 1868. The first portland cement manufactured in the United States was produced at a plant in Coplay, Pennsylvania, in 1871.

MANUFACTURE OF PORTLAND CEMENT

Portland cement is produced by pulverizing clinker which consists primarily of hydraulic calcium silicates. Clinker also contains some calcium aluminates and calcium aluminoferrites and one or more forms of calcium sulfate (gypsum) is interground with the clinker to make the finished product.

Materials used in the manufacture of portland cement must contain appropriate amounts of calcium, silica, alumina, and iron components. During manufacture, chemical analyses of all materials are made frequently to ensure a uniformly high quality cement.

Steps in the manufacture of cement are illustrated in the flow charts in Figs. 2-3 and 2-4. While the operations of all cement plants are basically the same, no flow diagram can adequately illustrate all plants. There is no typical portland cement manufacturing facility; every plant has significant differences in layout, equipment, or general appearance (Fig. 2-5).

Selected raw materials (Table 2-1) are transported from the quarry (Fig. 2-6), crushed (Fig. 2-7), milled, and proportioned so that the resulting mixture has the desired chemical composition. The raw materials are generally a mixture...

Fig. 2-5. Aerial view of a cement plant. (IMG12442)
the kiln at a rate controlled by the slope and rotational speed of the kiln. Burning fuel (powdered coal, new or recycled oil, natural gas, rubber tires, and by-product fuel) is forced into the lower end of the kiln where temperatures of 1400°C to 1550°C change the raw material chemically into cement clinker, grayish-black pellets predominantly the size of marbles (Fig. 2-9). Fig. 2-10 illustrates important technological developments that can improve significantly the productivity and energy efficiency of dry-process plants.

After blending, the ground raw material is fed into the upper end of a kiln (Fig. 2-8). The raw mix passes through the kiln at a rate controlled by the slope and rotational speed of the kiln. Burning fuel (powdered coal, new or recycled oil, natural gas, rubber tires, and by-product fuel) is forced into the lower end of the kiln where temperatures of 1400°C to 1550°C change the raw material chemically into cement clinker, grayish-black pellets predominantly the size of marbles (Fig. 2-9). Fig. 2-10 illustrates important technological developments that can improve significantly the productivity and energy efficiency of dry-process plants.

<table>
<thead>
<tr>
<th>Table 2-1. Sources of Raw Materials Used in Manufacture of Portland Cement</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Calcium</strong></td>
</tr>
<tr>
<td>Alkali waste</td>
</tr>
<tr>
<td>Aragonite*</td>
</tr>
<tr>
<td>Calcite*</td>
</tr>
<tr>
<td>Cement-kiln dust</td>
</tr>
<tr>
<td>Cement rock</td>
</tr>
<tr>
<td>Chalk</td>
</tr>
<tr>
<td>Clay</td>
</tr>
<tr>
<td>Fuller’s earth</td>
</tr>
<tr>
<td>Limestone*</td>
</tr>
<tr>
<td>Marble</td>
</tr>
<tr>
<td>Marl*</td>
</tr>
<tr>
<td>Seashells</td>
</tr>
<tr>
<td>Shale*</td>
</tr>
<tr>
<td>Slag</td>
</tr>
</tbody>
</table>

Note: Many industrial byproducts have potential as raw materials for the manufacture of portland cement. 
*Most common sources.
### Cross-section view of kiln

<table>
<thead>
<tr>
<th>Temperature</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>To 700°C</td>
<td>Raw materials are free-flowing powder.</td>
</tr>
<tr>
<td>700-900°C</td>
<td>Powder is still free-flowing.</td>
</tr>
<tr>
<td>1150-1200°C</td>
<td>Particles start to become “sticky”.</td>
</tr>
<tr>
<td>1200-1350°C</td>
<td>As particles start to agglomerate, they are held together by the liquid. The rotation of the kiln initiates coalescing of agglomerates and layering of particles.</td>
</tr>
<tr>
<td>1350-1450°C</td>
<td>Agglomeration and layering of particles continue as material falls on top of each other.</td>
</tr>
<tr>
<td>Cooling</td>
<td>Clinker nodules remain unchanged during cooling.</td>
</tr>
</tbody>
</table>

### Nodulization process

<table>
<thead>
<tr>
<th>Temperature</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>To 700°C</td>
<td>Particles are solid. No reaction between particles.</td>
</tr>
<tr>
<td>700-900°C</td>
<td>Particles are still solid.</td>
</tr>
<tr>
<td>1150-1200°C</td>
<td>Reactions start happening between solid particles.</td>
</tr>
<tr>
<td>1200-1350°C</td>
<td>Capillary forces of the liquid keep particles together.</td>
</tr>
<tr>
<td>1350-1450°C</td>
<td>Nodules will form with sufficient liquid. Insufficient liquid will result in dusty clinker.</td>
</tr>
<tr>
<td>Cooling</td>
<td>Clinker nodules remain unchanged during cooling.</td>
</tr>
</tbody>
</table>

### Clinkering reactions

<table>
<thead>
<tr>
<th>Temperature</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>To 700°C</td>
<td>Water is lost. Dehydrated clay recrystallizes.</td>
</tr>
<tr>
<td>700-900°C</td>
<td>Powder is still free-flowing.</td>
</tr>
<tr>
<td>1150-1200°C</td>
<td>As calcination continues, free lime increases. Reactive silica combines with CaO to begin forming C₂S.</td>
</tr>
<tr>
<td>1200-1350°C</td>
<td>When calcination is complete, temperature increases rapidly. Small belite crystals form from combination of silicates and CaO.</td>
</tr>
<tr>
<td>1350-1450°C</td>
<td>Belite crystals decrease in amount, increase in size. Alite increases in size and amount.</td>
</tr>
<tr>
<td>Cooling</td>
<td>Upon cooling, the C₃A and C₄AF crystallize in the liquid phase. Lamellar structure appears in belite crystals.</td>
</tr>
</tbody>
</table>

**Fig. 2-10.** Process of clinker production from raw feed to the final product (Hills 2000).
shows the clinker production process from raw feed to the final product.

The clinker is cooled and then pulverized. During this operation a small amount of gypsum (Fig. 2-11) is added to regulate the setting time of the cement and to improve shrinkage and strength development properties (Lerch 1946 and Tang 1992). In the grinding mill, clinker is ground so fine that nearly all of it passes through a 45 micrometer (No. 325 mesh) sieve. This extremely fine gray powder is portland cement (Fig. 2-1).

TYPES OF PORTLAND CEMENT

Different types of portland cement are manufactured to meet various normal physical and chemical requirements for specific purposes. Portland cements are manufactured to meet the specifications of ASTM C150, AASHTO M 85, or ASTM C1157.

ASTM C150, Standard Specification for Portland Cement, provides for eight types of portland cement using Roman numeral designations as follows:

- **Type I** Normal
- **Type IA** Normal, air-entraining
- **Type II** Moderate sulfate resistance
- **Type II A** Moderate sulfate resistance, air-entraining
- **Type III** High early strength
- **Type III A** High early strength, air-entraining
- **Type IV** Low heat of hydration
- **Type V** High sulfate resistance

AASHTO M 85, Specification for Portland Cement, also uses type designations I through V for portland cement. The requirements of M 85 are almost identical to ASTM C150. AASHTO specifications are used by some state departments of transportation in lieu of ASTM standards.

ASTM C1157, Performance Specification for Hydraulic Cements, provides for six types of portland cement as discussed under “Hydraulic Cements” below.

A detailed review of ASTM C150 and AASHTO M 85 cements follows.

**Type I**

Type I portland cement is a general-purpose cement suitable for all uses where the special properties of other types are not required. Its uses in concrete include pavements, floors, reinforced concrete buildings, bridges, tanks, reservoirs, pipe, masonry units, and precast concrete products (Fig. 2-12).

**Type II**

Type II portland cement is used where precaution against moderate sulfate attack is important. It is used in normal structures or elements exposed to soil or ground waters where sulfate concentrations are higher than normal but not unusually severe (see Table 2-2, and Figs. 2-13 to Fig. 2-15). Type II cement has moderate sulfate resistant properties because it contains no more than 8% tricalcium aluminate (C₃A).

Sulfates in moist soil or water may enter the concrete and react with the hydrated C₃A, resulting in expansion, scaling, and cracking of concrete. Some sulfate compounds, such as magnesium sulfate, directly attack calcium silicate hydrate.

Use of Type II cement in concrete must be accompanied by the use of a low water to cementitious materials ratio and low permeability to control sulfate attack. Fig. 2-13 (top) illustrates the improved sulfate resistance of Type II cement over Type I cement.

Concrete exposed to seawater is often made with Type II cement. Seawater contains significant amounts of sulfates and chlorides. Although sulfates in seawater are capable of attacking concrete, the presence of chlorides inhibits the expansive reaction that is characteristic of sulfate attack. Chloride competes with sulfate for the aluminate phases, after which they exist together in the
concrete. The reaction products of sulfate attack are also more soluble in a chloride solution and can leach out of the concrete. Observations from a number of sources show that the performance of concretes in seawater with portland cements having C₃A contents as high as 10%, have shown satisfactory durability, providing the permeability of the concrete is low and the reinforcing steel has adequate cover (Zhang, Bremner, and Malhotra 2003). Type II cements specially manufactured to meet the moderate heat option of ASTM C150 (AASHTO M 85) will generate heat at a slower rate than Type I or most Type II cements. The requirement of moderate heat of hydration can be specified at the option of the purchaser. A cement in which heat-of-hydration maximums are specified can be used in structures of considerable mass, such as large piers, large foundations, and thick retaining walls (Fig. 2-16). Its use will reduce temperature rise and temperature related cracking, which is especially important when concrete is placed in warm weather.

Because of its increased availability, Type II cement is sometimes used in all aspects of construction, regardless of the need for sulfate resistance or moderate heat generation. Some cements may be labeled with more than one type designation, for example Type I/II. This simply means that such a cement meets the requirements of both cement Types I and II.

**Type III**

Type III portland cement provides strength at an early period, usually a week or less. It is chemically and physically similar to Type I cement, except that its particles have been ground finer. It is used when forms need to be removed as soon as possible or when the structure must be put into service quickly. In cold weather its use permits a reduction in the length of the curing period (Fig. 2-17). Although higher cement content mixes of Type I cement can be used to gain high early strength, Type III may provide it easier and more economically.
Type IV

Type IV portland cement is used where the rate and amount of heat generated from hydration must be minimized. It develops strength at a slower rate than other cement types. Type IV cement is intended for use in massive concrete structures, such as large gravity dams, where the temperature rise resulting from heat generated during hardening must be minimized (Fig. 2-16). Type IV cement is rarely available.

Type V

Type V portland cement is used in concrete exposed to severe sulfate action—principally where soils or groundwaters have a high sulfate content (Figs. 2-13 to 2-15). It gains strength more slowly than Type I cement. Table 2-2 lists sulfate concentrations requiring the use of Type V cement. The high sulfate resistance of Type V cement is attributed to a low tricalcium aluminate content, not more than 5%. Use of a low water to cementitious materials ratio and low permeability are critical to the performance of any concrete exposed to sulfates. Even Type V cement concrete cannot withstand a severe sulfate exposure if the concrete has a high water to cementitious materials ratio (Fig. 2-15).
White Portland Cements

White portland cement is a true portland cement that differs from gray cement chiefly in color. It is made to conform to the specifications of ASTM C150, usually Type I or Type III; the manufacturing process is controlled so that the finished product will be white. White portland cement is made of selected raw materials containing negligible amounts of iron and magnesium oxides—the substances that give cement its gray color. White portland cement is used primarily for architectural purposes in structural walls, precast and glass fiber reinforced concrete (GFRC) facing panels, terrazzo surfaces, stucco, cement paint, tile grout, and decorative concrete (Fig. 2-18). Its use is recommended wherever white or colored concrete, grout, or mortar is desired. White portland cement should be specified as: white portland cement meeting the specifications of ASTM C150, Type [I, II, III, or V]. White cement is also used to manufacture white masonry cement meeting ASTM C91 and white plastic cement meeting ASTM C1328 (PCA 1999). White cement was first manufactured in the United States in York, Pennsylvania in 1907. See Farny (2001) for more information.

Air-Entraining Portland Cements

Specifications for three types of air-entraining portland cement (Types IA, IIA, and IIIA) are given in ASTM C150 and AASHTO M 85. They correspond in composition to ASTM Types I, II, and III, respectively, except that small quantities of air-entraining material are interground with the clinker during manufacture. These cements produce concrete with improved resistance to freezing and thawing. Such concrete contains minute, well-distributed, and completely separated air bubbles. Air entrainment for most concretes is achieved through the use of an air-entraining admixture, rather than through the use of air-entraining cements. Air-entraining cements are available only in certain areas.

Fig. 2-17. High early strength cements are used where early concrete strength is needed, such as in (left to right) cold weather concreting, fast track paving to minimize traffic congestion, and rapid form removal for precast concrete. (IMG12350, IMG12477, IMG12476)

Fig. 2-18. White portland cement is used in white or light-colored architectural concrete, ranging from (left to right) terrazzo for floors shown here with white cement and green granite aggregate (IMG12475), to decorative and structural precast and cast-in-place elements (68981), to building exteriors. The far right photograph shows a white precast concrete building housing the ASTM Headquarters in West Conshohocken, Pennsylvania. Photo courtesy of ASTM.
BLENDED HYDRAULIC CEMENTS

Blended cements are used in all aspects of concrete construction in the same manner as portland cements. Blended cements can be used as the only cementitious material in concrete or they can be used in combination with other supplementary cementitious materials added at the concrete plant. Blended cements are often designed to be used in combination with local pozzolans and slags. If a blended cement or portland cement is used alone or in combination with added pozzolans or slags, the concrete should be tested for strength, durability, and other properties required in project specifications (PCA 1995 and Detwiler, Bhatttay, and Bhattacharja 1996).

Blended hydraulic cements are produced by intimately and uniformly intergrinding or blending two or more types of fine materials. The primary materials are portland cement, ground granulated blast-furnace slag, fly ash, silica fume, calcined clay, other pozzolans, hydrated lime, and preblended combinations of these materials (Fig. 2-19). Blended hydraulic cements must conform to the requirements of ASTM C595 (AASHTO M 240), Specification for Blended Hydraulic Cements, or ASTM C1157, Performance Specification for Hydraulic Cement.

ASTM C595 recognizes two primary classes of blended cements as follows:

- Type IS (X) Portland blast-furnace slag cement
- Type IP (X) Portland-pozzolan cement

The letter “X” stands for the nominal percentage of the SCM included in the blended cement. For example, a cement designated as Type IS (50) contains 50% by mass of slag. Types IS and IP are general purpose cements (Fig. 2-12); these and subcategory types are reviewed in the discussion below.

AASHTO M 240 also uses class designations for blended cement. The requirements of M 240 are almost identical to those in ASTM C595.

ASTM C1157 provides for six types of blended cement as discussed under “Hydraulic Cements” below. Cements meeting the requirements of C1157 meet physical performance test requirements without prescriptive restrictions on ingredients or cement chemistry. This allows the cement manufacturer to optimize strength and durability properties through use of a variety of cementitious materials, such as portland clinker, fly ash, slag, silica fume, and calcined clay (Fig. 2-19).

A detailed review of ASTM C595 and AASHTO M 240 blended cements follows:

**Type IS**

Portland blast-furnace slag cement, Type IS, may be used in general concrete construction. Historical use of slag blended cements dates back to the beginning of the 20th century in Europe, Japan, and North America (Abrams 1925). In producing these cements, granulated blast-furnace slag is either interground with portland cement clinker, separately ground and blended with portland cement, or produced with a combination of intergrinding and blending. Portland blast-furnace slag cement with a slag content equal to or exceeding 70% by mass [IS (≥70)] is permitted to contain hydrated lime. The blast-furnace slag content of this cement is up to 95% of the mass of the finished cement. Portland blast-furnace slag cement with a slag content of less than 70% by mass, Type IS (<70), may be used in general concrete construction. This cement is manufactured by either (1) intergrinding portland cement clinker and granulated blast-furnace slag, (2) blending portland cement and finely ground granulated blast-furnace slag, or (3) a combination of intergrinding and blending. Slag is less than 70% by mass of the finished cement. Subcategory types (optional special properties) such as air-entrainment, moderate sulfate resistance, high sulfate resistance, or moderate heat of hydration may be specified by adding the suffixes A, MS, HS, or MH. For example, an air-entraining portland blast-furnace slag cement with 40% slag that has high sulfate resistance would be designated as Type IS (40)-A (HS). Portland blast-furnace slag cement with a slag content of 70% by mass or more, Type IS (≥70), is used with portland cement in making concrete or with lime in making mortars, but is not used alone in structural concrete. Type IS (≥70) is manufactured by either (1) blending ground granulated blast-furnace slag and portland cement, (2) blending ground granulated blast-furnace slag and hydrated lime, or (3) blending a combination of ground granulated blast-furnace slag, portland cement, and hydrated lime. Air-entrainment may be designated in a Type IS (≥70) cement by adding the suffix A, for example, Type IS (85)-A. See Klieger and Isberner (1967) and PCA (1995).
**Type IP**

Portland-pozzolan cements are designated as Type IP. Type IP may be used for general construction. These cements are manufactured by intergrinding portland cement clinker with a suitable pozzolan, by blending portland cement or portland blast-furnace slag cement and a pozzolan, or by a combination of intergrinding and blending. The pozzolan content of these cements is up to 40% by mass. Laboratory tests indicate that performance of concrete made with Type IP cement as a group is similar to that of Type I cement concrete. Type IP may be designated as air-entraining, moderate sulfate resistant, high sulfate resistant, low heat of hydration, or with moderate heat of hydration by adding the suffixes A, MS, HS, LH, or MH.

**HYDRAULIC CEMENTS**

Hydraulic cements set and harden by reacting chemically with water. They also stay hard and maintain their stability under water. They are used in all aspects of concrete construction. All portland and blended cements are hydraulic cements. “Hydraulic cement” is merely a broader term. See also ASTM C219 for terms relating to hydraulic cements.

The 1990s saw the creation of a performance specification for hydraulic cements—ASTM C1157, *Performance Specification for Hydraulic Cements*. This specification is designed generically for hydraulic cement to include portland cement, modified portland cement, and blended hydraulic cement. Cements meeting the requirements of C1157 meet physical performance test requirements, as opposed to prescriptive restrictions on ingredients or cement chemistry as found in other cement specifications. ASTM C1157 provides for six types of hydraulic cement as follows:

<table>
<thead>
<tr>
<th>Type</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Type GU</td>
<td>General use</td>
</tr>
<tr>
<td>Type HE</td>
<td>High early strength</td>
</tr>
<tr>
<td>Type MS</td>
<td>Moderate sulfate resistance</td>
</tr>
<tr>
<td>Type HS</td>
<td>High sulfate resistance</td>
</tr>
<tr>
<td>Type MH</td>
<td>Moderate heat of hydration</td>
</tr>
<tr>
<td>Type LH</td>
<td>Low heat of hydration</td>
</tr>
</tbody>
</table>

In addition, these cements can also have an Option R—Low Reactivity with Alkali-Reactive Aggregates—specified to help control alkali-silica reactivity. For example, Type GU-R would be a general use hydraulic cement with low reactivity with alkali-reactive aggregates.

When specifying a C1157 cement, the specifier uses the nomenclature of “hydraulic cement,” “portland cement,” “air-entraining portland cement,” “modified portland cement” or “blended hydraulic cement” along with a type designation. For example, a specification may call for a Hydraulic Cement Type GU, a Blended Hydraulic Cement Type MS, or a Portland Cement Type HS. If a type is not specified, then Type GU is assumed.

ASTM C1157 defines a blended cement as having more than 15% mineral additive and a modified portland cement as containing up to 15% mineral additive. The mineral additive usually prefixes the modified portland cement nomenclature, for example, slag-modified portland cement.

ASTM C1157 also allows a strength range to be specified from a table in the standard. If a strength range is not specified, only the minimum strengths apply. Strength ranges are rarely applied in the United States.

A detailed review of ASTM C1157 cements follows:

**Type GU**

Type GU is a general-purpose cement suitable for all uses where the special properties of other types are not required. Its uses in concrete include pavements, floors, reinforced concrete buildings, bridges, pipe, precast concrete products, and other applications where Type I is used (Fig. 2-12).

**Type HE**

Type HE cement provides higher strengths at an early age, usually a week or less. It is used in the same manner as Type III portland cement (Fig. 2-17).

**Type MS**

Type MS cement is used where precaution against moderate sulfate attack is important, as in drainage structures where sulfate concentrations in ground waters are higher than normal but not unusually severe (see Table 2-2). It is used in the same manner as Type II portland cement (Fig. 2-14). Like Type II, Type MS cement concrete must be made with a low water-cementitious materials ratio to provide sulfate resistance.

**Type HS**

Type HS cement is used in concrete exposed to severe sulfate action—principally where soils or ground waters have a high sulfate content (see Table 2-2). It is used in the same manner as Type V portland cement (Fig. 2-14).

**Type MH**

Type MH cement is used where the concrete needs to have a moderate heat of hydration and a controlled temperature rise. Type MH cement is used in the same manner as a moderate heat Type II portland cement (Fig. 2-16).
Table 2-3. Applications for Commonly Used Cements

<table>
<thead>
<tr>
<th>Cement specification</th>
<th>General purpose</th>
<th>Moderate heat of hydration</th>
<th>High early strength</th>
<th>Low heat of hydration</th>
<th>Moderate sulfate resistance</th>
<th>High sulfate resistance</th>
<th>Resistance to alkali-silica reactivity (ASR)**</th>
</tr>
</thead>
<tbody>
<tr>
<td>ASTM C150 (AASHTO M 85) portland cements</td>
<td>I</td>
<td>II (moderate heat option)</td>
<td>III</td>
<td>IV</td>
<td>II</td>
<td>V</td>
<td>Low alkali option</td>
</tr>
<tr>
<td>ASTM C595 (AASHTO M 240) blended hydraulic cements</td>
<td>IS IP IS(&lt;70)(MH) IP(MH) IP(LH) IS(&lt;70)(MS) IP(MS) IS(&lt;70)(HS) IP(HS)</td>
<td>IS IP IS(&lt;70)(MH) IP(MH) IP(LH) IS(&lt;70)(MS) IP(MS) IS(&lt;70)(HS) IP(HS)</td>
<td>Low reactivity option</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>ASTM C1157 hydraulic cements***</td>
<td>GU MH HE LH MS HS</td>
<td>Option R</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

* Check the local availability of specific cements as all cements are not available everywhere.
** The option for low reactivity with ASR susceptible aggregates can be applied to any cement type in the columns to the left.
*** For ASTM C1157 cements, the nomenclature of hydraulic cement, portland cement, air-entraining portland cement, modified portland cement, or blended hydraulic cement is used with the type designation.

Type LH

Type LH cement is used where the rate and amount of heat generated from hydration must be minimized. It develops strength at a slower rate than other cement types. Type LH cement is intended for use in massive concrete structures where the temperature rise resulting from heat generated during hardening must be minimized. It is used in the same manner as Type IV portland cement (Fig. 2-16).

Table 2-3 provides a matrix of commonly used cements and where they are used in concrete construction.

SPECIAL CEMENTS

Special cements are produced for particular applications. Table 2-4 summarizes the special cements discussed below. See Odler (2000) and Klemm (1998) for more information.

Masonry and Mortar Cements

Masonry cements and mortar cements are hydraulic cements designed for use in mortar for masonry construction (Fig. 2-20). They consist of a mixture of portland cement or blended hydraulic cement and plasticizing materials (such as limestone or hydrated or hydraulic lime), together with other materials introduced to enhance one or more properties such as setting time, workability, water retention and durability. These components are proportioned and packed at a cement plant under controlled conditions to assure uniformity of performance.

Masonry cements meet the requirements of ASTM C91, which classifies masonry cements as Type N, Type S, and Type M. White masonry cement and colored masonry cements meeting ASTM C91 are also available in some areas. Mortar cements meet the requirements of ASTM C1329, which also classifies mortar cements as Type N, Type S, and Type M. A brief description of each type follows:

Type N masonry cement and Type N mortar cement are used in ASTM C270 Type N and Type O mortars. They may also be used with portland or blended cements to produce Type S and Type M mortars.

Type S masonry cement and Type S mortar cement are used in ASTM C270 Type S mortar. They may also be used with portland or blended cements to produce Type M mortar.

Type M masonry cement and Type M mortar cement are used in ASTM C270 Type M mortar without the addition of other cements or hydrated lime.

Types N, S, and M generally have increasing levels of portland cement and higher strength, Type M having the highest strength. Type N is used most commonly.

The increased use of masonry in demanding structural applications, such as high seismic areas, resulted in the recent development of mortar cement. Mortar cement is similar to masonry cement in that it is a factory-prepared cement primarily used to produce masonry mortar.
However, ASTM C1329 places lower maximum air content limits on mortar cement than permitted for masonry cements; also, ASTM C1329 is the only ASTM masonry material specification that includes bond strength performance criteria.

The workability, strength, and color of masonry cements and mortar cements stay at a high level of uniformity because of manufacturing controls. In addition to mortar for masonry construction, masonry cements and mortar cements are used for parging. Masonry cements are also used in portland-cement based plaster or stucco (Fig. 2-21) construction (see ASTM C926). Masonry cement and mortar cement should not be used for making concrete.

### Plastic Cements

Plastic cement is a hydraulic cement that meets the requirements of ASTM C1328. It is used to make portland cement-based plaster or stucco (ASTM C926), usually in the southwest and west coast of the United States (Fig. 2-21). Plastic cements consist of a mixture of portland and blended hydraulic cement and plasticizing materials (such as limestone, hydrated or hydraulic lime), together with materials introduced to enhance one or more properties such as setting time, workability, water retention, and durability.

ASTM C1328 defines separate requirements for a Type M and a Type S plastic cement with Type M having higher strength requirements. The International Building Code (IBC) does not classify plastic cement into different types,
but specifies that it must meet the requirements of ASTM C1328 plastic cement. When plastic cement is used, no lime or other plasticizer may be added to the plaster at the time of mixing.

The term “plastic” in plastic cement does not refer to the inclusion of any organic compounds in the cement; rather, “plastic” refers to the ability of the cement to impart to the plaster a high degree of plasticity or workability. Plaster made from this cement must remain workable for a long enough time for it to be reworked to obtain the desired densification and texture. Plastic cement should not be used to make concrete. For more information on the use of plastic cement and plaster, see Melander and Isberner (1996).

**Finely-Ground Cements (Ultrafine Cements)**

Finely-ground cements, also called ultrafine cements, are hydraulic cements that are ground very fine for use in grouting into fine soil or thin rock fissures (Fig. 2-22). The cement particles are less than 10 µm in diameter with 50% of particles less than 5 µm. Blaine fineness often exceeds 800 m²/kg. These very fine cements consist of portland cement, ground granulated blast-furnace slag, and other mineral additives.

**Expansive Cements**

Expansive cement is a hydraulic cement that expands slightly during the early hardening period after setting. It must meet the requirements of ASTM C845 in which it is designated as Type E-1. Currently, three varieties of expansive cement are recognized and have been designated as K, M, and S, which are added as a suffix to the type. Type E-1(K) contains portland cement, tetracalcium trialuminate, calcium sulfate, and uncombined calcium oxide (lime). Type E-1(M) contains portland cement, calcium aluminate cement, and calcium sulfate. Type E-1(S) contains portland cement with a high tricalcium aluminate content and calcium sulfate. Type E-1(K) is the most readily available expansive cement in North America.

Expansive cement may also be made of formulations other than those mentioned. The expansive properties of each type can be varied over a considerable range.

When expansion is restrained, for example by reinforcing steel, expansive cement concrete (also called shrinkage compensating concrete) can be used to (1) compensate for the volume decrease due to drying shrinkage, (2) induce tensile stress in reinforcement (post-tensioning), and (3) stabilize the long-term dimensions of post-tensioned concrete structures with respect to original design.

One of the major advantages of using expansive cement in concrete is noted in (1) above; when you can compensate for volume change due to drying shrinkage you can control and reduce drying shrinkage cracks. Fig. 2-23 illustrates the length change (early expansion and drying shrinkage) history of shrinkage-compensating concrete and conventional

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Fig. 2-21. Masonry cement and plastic cement are used to make plaster or stucco for commercial, institutional, and residential buildings. Shown are a church and home with stucco exteriors. Inset shows a typical stucco texture. (IMG12470, IMG12469, 68805)

Fig. 2-22. (left) A slurry of finely ground cement and water can be injected into the ground, as shown here, to stabilize in-place materials, to provide strength for foundations, or to chemically retain contaminants in soil. (IMG12468) Illustration (right) of grout penetration in soil.
Gajda (1996) studied a functional addition to control alkali-silica reactivity.

**Water-Repellent Cements**

Water-repellent cements, sometimes called waterproofed cements, are usually made by adding a small amount of water-repellent additive such as stearate (sodium, aluminum, or other) to cement clinker during final grinding (Lea 1971). Manufactured in either white or gray color, water-repellent cements reduce capillary water transmission when there is little to no pressure but do not stop water-vapor transmission. They are used in tile grouts, paint, and stucco finish coats.

**Regulated-Set Cements**

Regulated-set cement is a calcium fluoroaluminate hydraulic cement that can be formulated and controlled to produce concrete with setting times from a few minutes to an hour and with corresponding rapid early strength development (Greening and others 1971). It is a portland-based cement with functional additions that can be manufactured in the same kiln used to manufacture conventional portland cement. Regulated-set cement incorporates set control and early-strength-development components. Final physical properties of the resulting concrete are in most respects similar to comparable concretes made with portland cement.

**Geopolymer Cements**

Geopolymer cements are inorganic hydraulic cements that are based on the polymerization of minerals (Davidovits, Davidovits, and James 1999). The term more specifically refers to alkali-activated aluminosilicate cements, also called zeolitic or polysialate cements. They have been used in general construction, high-early strength applications, and waste stabilization. These cements do not contain organic polymers or plastics.

**Ettringite Cements**

Ettringite cements are calcium sulfoaluminate cements that are specially formulated for particular uses, such as the stabilization of waste materials (Klemm 1998). They can be formulated to form large amounts of ettringite to stabilize particular metallic ions within the ettringite structure. Ettringite cements have also been used in rapid setting applications, including use in coal mines. Also see “Expansive Cements” above.

**Rapid Hardening Cements**

Rapid hardening, high-early strength, hydraulic cement is used in construction applications, such as fast-track paving, where fast strength development is needed (design or load carrying strength in about four hours). They are classified by ASTM C1600 as Type URH – ultra rapid hard-
SELECTING AND SPECIFYING CEMENTS

When specifying cements for a project, be sure to check the availability of cement types as discussed below; specifications should allow flexibility in cement selection. Limiting a project to only one cement type, one brand, or one standard cement specification can result in project delays and it may not allow for the best use of local materials. Cements with special properties should not be required unless special characteristics are necessary. In addition, the use of supplementary cementing materials should not inhibit the use of any particular portland or blended cement. The project specifications should focus on the needs of the concrete structure and allow use of a variety of materials to accomplish those needs. A typical specification may call for portland cements meeting ASTM C150 (AASHTO M 85) or C1157, or for blended cements meeting ASTM C595 (AASHTO M 240) or C1157.

If no special properties, (such as low-heat generation or sulfate resistance) are required, all general use cements should be allowed, such as Types I, GU, IS, and IP. Also, it should be noted that some cement types meet the requirements of other types as well; for example, all Type II cements meet the requirements of Type I, but not all Type I cements meet the requirements of Type II. See Tables 2-3 and 2-4 for guidance on using different cements.

Availability of Cements

Some types of cement may not be readily available in all areas of the United States. Before specifying a type of cement, its availability should be determined.

ASTM C150 Type I portland cement is usually carried in stock and is furnished when a type of cement is not specified. Type II cement is usually available, especially in areas of the country where moderate sulfate resistance is needed. Cement Types I and II represent about 75% of the cement shipped from plants in the United States. Some cements are designated as both Type I and II (Type I/II), meaning that they meet specification requirements for both types. Type III cement and white cement are usually available in larger metropolitan areas and represent about 4% of cement produced. Type IV cement is manufactured only when specified for particular projects (massive structures like dams) and therefore is usually not readily available. Type V cement is available only in particular regions of the country where it is needed to resist high sulfate environments. Air-entraining cements are sometimes difficult to obtain. Their use has decreased as the popularity of air-entraining admixtures has increased. Masonry cement is available in most areas. See PCA (2007) for statistics on cement use.

If a given cement type is not available, comparable results frequently can be obtained with one that is available. For example, high-early-strength concrete can be made by using mixtures with higher cement contents and lower water-cement ratios. Also, the effects of heat of hydration can be minimized by using lean mixtures, smaller placing lifts, artificial cooling, or by adding a

Magnesium Phosphate Cements

Magnesium phosphate cement is a rapid setting, early strength gain cement. It is usually used for special applications, such as repair of pavements and concrete structures, or for resistance to certain aggressive chemicals. It does not contain portland cement.

Sulfur Cements

Sulfur cement is used with conventional aggregates to make sulfur cement concrete for repairs and chemically resistant applications. Sulfur cement melts at temperatures between 113°C and 121°C (235°F and 250°F). Sulfur concrete is maintained at temperatures around 130°C (270°F) during mixing and placing. The material gains strength quickly as it cools and is resistant to acids and aggressive chemicals. Sulfur cement does not contain portland or hydraulic cement.
Canadian and European Cement Specifications

In some instances, projects in the United States designed by engineering firms from other countries refer to cement standards other than those in ASTM or AASHTO. For example, the European cement standard, EN 197, sometimes appears on project specifications. EN 197 cement Types CEM I, II, III, IV, and V do not correspond to the cement types in ASTM C150, nor can ASTM cements be substituted for EN specified cement without the designer’s approval. CEM I is a portland cement and CEM II through V are blended cements. EN 197 also has strength classes and ranges (32.5, 42.5, and 52.5 MPa).

There is no direct equivalency between ASTM and other cement standards of the world because of differences in test methods and limits on required properties. EN 197 cements are usually not available in the United States; therefore, the best approach is to inform the designer as to what cements are locally available and ask for a change in the project specifications to allow use of an ASTM or AASHTO cement.

Old Canadian CSA A5 Types 10, 20, 30, 40, and 50 were essentially the same as ASTM C150 (AASHTO M 85) Types I through V.

In 2004, Canadian Standard CSA A3001-03 introduced two-letter descriptive type designations for portland cements similar to ASTM C150 with requirements similar to ASTM C150:

- GU: General use hydraulic cement (former CSA A5 Type 10)
- MS: Moderate sulphate-resistant hydraulic cement (former CSA A5 Type 20)
- MH: Moderate heat of hydration hydraulic cement (former CSA A5 Type 20)
- HE: High early-strength hydraulic cement (former CSA A5 Type 30)
- LH: Low heat of hydration hydraulic cement (former CSA A5 Type 40)
- HS: High sulphate-resistant hydraulic cement (former CSA A5 Type 50)

The nomenclature for blended hydraulic cements is a three-letter descriptive designation (GUb, MSb, MHb, HEb, LHb, and HSb) to address its equivalent performance to portland cements with up to three supplementary cementing materials. Upon request, the designations for blended cements can also provide information on the composition of blended hydraulic cements. The designations then follow the form: BHb-Axx/Byy/Czz, where BHb is the blended hydraulic cement type, xx, yy, and zz are the supplementary materials used in the cement in proportions A, B, and C respectively.

Covered supplementary cementitious materials include:

- Ground granulated blast furnace slag (S),
- Silica fume (SF),
- Natural pozzolans (N), and
- Fly ash (Classes F, C, and CH). Class F, C, and CH fly ashes are low (less than 8% CaO by mass), medium
(between 8% and 20% CaO by mass), and high calcium oxide (more than 20% CaO by mass) contents, respectively.

Examples:
- MS—portland cement (with no supplementary cementitious materials) for use when moderate sulfate resistance is required.
- GUb-30F/5SF—general use blended cement containing 30% by mass Class F fly ash (F) and 5% silica fume (SF).

CHEMICAL COMPOUNDS AND HYDRATION OF PORTLAND CEMENT

During the burning operation in the manufacture of portland cement clinker, calcium combines with the other components of the raw mix to form four principal compounds that make up 90% of cement by mass. Gypsum (4% to 6%), or other calcium sulfate source, and grinding aids are also added during grinding. Cement chemists use the following chemical shorthand (abbreviations) to describe compounds:

- A = Al₂O₃, C = CaO, F = Fe₂O₃, H = H₂O, M = MgO, S = SiO₂,
- and S = SO₃.

The term “phase” rather than “compounds” can also be used to describe the components of clinker. Following are the four primary compounds in portland cement, their approximate chemical formulas, and abbreviations:

- Tricalcium silicate: 3CaO•SiO₂ = C₃S
- Dicalcium silicate: 2CaO•SiO₂ = C₂S
- Tricalcium aluminate: 3CaO•Al₂O₃ = C₃A
- Tetracalcium aluminoferite: 4CaO•Al₂O₃•Fe₂O₃ = C₄AF

Following are the forms of calcium sulfate, their chemical formulas, and abbreviations:

- Anhydrous calcium sulfate: CaSO₄ = CaO•SO₃ = C₅
- Calcium sulfate dihydrate (gypsum): CaSO₄•2H₂O = CaO•SO₃•2H₂O = C₅H₂
- Calcium sulfate hemihydrate: CaSO₄•½ H₂O = CaO•SO₃•½ H₂O = C₅H₁/₂

Gypsum, calcium sulfate dihydrate, is the predominant source of sulfate used in cement.

C₃S and C₂S in clinker are also referred to as alite and belite, respectively. Alite constitutes 50% to 70% of the clinker, whereas belite accounts for only 15% to 30%. Aluminate compounds constitute about 5% to 10% of the clinker and ferrite compounds 5% to 15% (Taylor 1997). These and other compounds may be observed and analyzed through the use of microscopical techniques (see Fig. 2-25, ASTM C1356, and Campbell 1999).

In the presence of water, these compounds hydrate (chemically combined with water) to form new compounds that are the infrastructure of hardened cement paste in concrete (Fig. 2-26). The calcium silicates, C₃S and C₂S, hydrate to form the compounds calcium hydroxide and calcium silicate hydrate (archaically called tobermorite gel). Hydrated portland cement contains 15% to 25% calcium hydroxide and about 50% calcium silicate hydrate by mass. The strength and other properties of hydrated cement are due primarily to calcium silicate hydrate (Fig. 2-27). C₃A reacts with water and calcium hydroxide to form tetracalcium aluminate hydrates. C₄AF reacts with water to form calcium aluminoferrite hydrates. C₃A, sulfates (gypsum, anhydrite, or other sulfate source), and water combine to form ettringite (calcium trisulfoaluminate hydrate), calcium monosulfoaluminate, and other related compounds. These basic compound transformations are shown in Table 2-5.

Brunauer (1957), Copeland and others (1960), Lea (1971), Powers and Brownyard (1947), Powers (1961), and Taylor (1997) addressed the pore structure and chemistry of cement paste. Fig. 2-28 shows estimates of the relative volumes of the compounds in hydrated portland cement pastes.


The approximate percentage of each compound can be calculated from a chemical oxide analysis (ASTM C114 or AASHTO T 105) of the unhydrated cement per ASTM C150...
Fig. 2-26. Electron micrographs of (left) dicalcium silicate hydrate, (middle) tricalcium silicate hydrate, and (right) hydrated normal portland cement. Note the fibrous nature of the calcium silicate hydrates. Broken fragments of angular calcium hydroxide crystallites are also present (right). The aggregation of fibers and the adhesion of the hydration particles is responsible for the strength development of portland cement paste. Reference (left and middle) Brunauer 1962 and (right) Copeland and Schulz 1962. (IMG12464, IMG12463, IMG12462)

### Table 2-5. Portland Cement Compound Hydration Reactions (Oxide Notation)

<table>
<thead>
<tr>
<th>Reaction</th>
<th>Products</th>
<th>Equations</th>
</tr>
</thead>
<tbody>
<tr>
<td>2 (3CaO•SiO$_2$) + 11 H$_2$O = 3CaO•2SiO$_2$•6H$_2$O + 3 (CaO•H$_2$O)</td>
<td>Calcium silicate hydrate (C-S-H)</td>
<td>+ 3 (CaO•H$_2$O) Calcium hydroxide</td>
</tr>
<tr>
<td>2 (2CaO•SiO$_2$) + 9 H$_2$O = 3CaO•2SiO$_2$•8H$_2$O + CaO•H$_2$O</td>
<td>Calcium silicate hydrate (C-S-H)</td>
<td>+ CaO•H$_2$O Calcium hydroxide</td>
</tr>
<tr>
<td>3CaO•Al$_2$O$_3$ + 3 (CaO•SO$_3$•2H$_2$O) + 26 H$_2$O = 6CaO•Al$_2$O$_3$•3SO$_3$•32H$_2$O</td>
<td>Tricalcium aluminate</td>
<td>Ettringite</td>
</tr>
<tr>
<td>2 (3CaO•Al$_2$O$_3$) + 6 CaO•Al$_2$O$_3$•3SO$_3$•32H$_2$O + 4 H$_2$O = 3 (4CaO•Al$_2$O$_3$•SO$_3$•12H$_2$O)</td>
<td>Tricalcium aluminate</td>
<td>Calcium monosulfoaluminate</td>
</tr>
<tr>
<td>3CaO•Al$_2$O$_3$ + CaO•H$_2$O + 12 H$_2$O = 4CaO•Al$_2$O$_3$•13H$_2$O</td>
<td>Tricalcium aluminate</td>
<td>Tetracalcium aluminate hydrate</td>
</tr>
<tr>
<td>4CaO•Al$_2$O$_3$•Fe$_2$O$_3$ + 10 H$_2$O = 6CaO•Al$_2$O$_3$•Fe$_2$O$_3$•12H$_2$O</td>
<td>Tetracalcium aluminoferrite</td>
<td>Calcium aluminoferrite hydrate</td>
</tr>
</tbody>
</table>

Note: This table illustrates only primary transformations and not several minor transformations. The composition of calcium silicate hydrate (C-S-H) is not stoichiometric (Tennis and Jennings 2000).
Due to the inaccuracies of Bogue calculations, X-ray diffraction techniques can be used to more accurately determine compound percentages (ASTM C1365). Table 2-6 shows typical elemental and compound composition and fineness for each of the principal types of portland cement.

Although elements are reported as simple oxides for standard consistency, they are usually not found in that oxide form in the cement. For example, sulfur from the gypsum is reported as SO$_3$ (sulfur trioxide), however, cement does not contain any sulfur trioxide. The amount of calcium, silica, and alumina establish the amounts of the primary compounds in the cement and effectively the properties of hydrated cement. Sulfate is present to control setting, as well as drying shrinkage and strength gain (Tang 1992). Minor and trace elements and their effect on cement properties are discussed by Bhatty (1995) and PCA (1992).

Present knowledge of cement chemistry indicates that the primary cement compounds have the following properties:

**Tricalcium Silicate**, C$_3$S, hydrates and hardens rapidly and is largely responsible for initial set and early strength (Fig. 2-29). In general, the early strength of portland cement concrete is higher with increased percentages of C$_3$S.
Tetracalcium Aluminoferite, $C_4AF$, is the product resulting from the use of iron and aluminum raw materials to reduce the clinkering temperature during cement manufacture. It contributes little to strength. Most color effects that make cement gray are due to $C_4AF$ and its hydrates.

Calcium Sulfate, as anhydrite (anhydrous calcium sulfate), gypsum (calcium sulfate dihydrate), or hemihydrate, often called plaster of Paris or bassanite (calcium sulfate hemihydrate) is added to cement during final grinding to provide sulfate to react with $C_3A$ to form ettringite (calcium trisulfaloaluminate). This controls the hydration of $C_3A$.

Without sulfate, a cement would set rapidly. In addition to controlling setting and early strength gain, the sulfate also helps control drying shrinkage and can influence strength through 28 days (Lerch 1946).

In addition to the above primary cement compounds, numerous other lesser compound formulations also exist (PCA 1997, Taylor 1997, and Tennis and Jennings 2000).

Water (Evaporable and Nonevaporable)

Water is a key ingredient of pastes, mortars, and concretes, as the phases in portland cement must chemically react with water to develop strength. The amount of water added to a mixture controls the durability as well. The space initially taken up by water in a cementitious mixture is partially or completely replaced over time as the hydration reactions proceed (Table 2-5). If more than about 35% water by mass of cement—a water to cement ratio of 0.35—is used, then porosity in the hardened material will remain,
Even after complete hydration. This is called capillary porosity. Fig. 2-30 shows that cement pastes with high and low water to cement ratios have equal masses after drying (evaporable water was removed). The cement consumed the same amount of water in both pastes resulting in more bulk volume in the higher water-cement ratio paste. As the water to cement ratio increases, the capillary porosity increases, and the strength decreases. Also, transport properties such as permeability and diffusivity are increased, allowing detrimental chemicals to more readily attack the concrete or reinforcing steel.

Water is found in cementitious materials in several forms. Free water is mixing water that has not reacted with the cement phases. Bound water is chemically combined in the solid phases or physically bound to the solid surfaces. A reliable separation of the chemically combined from the physically adsorbed water is not possible. Therefore Powers (1949) distinguished between evaporable and nonevaporable water. The nonevaporable water is the amount retained by a sample after it has been subjected to a drying procedure intended to remove all the free water (traditionally, by heating to 105°C). Evaporable water was originally considered to be free water, but it is now recognized that some bound water is also lost upon heating to this temperature. All nonevaporable water is bound water, but the opposite is not true.

For complete hydration of portland cement, only about 40% water (a water-to-cement ratio of 0.40) is needed. If a water-to-cement ratio greater than about 0.40 is used, the excess water not needed for cement hydration remains in the capillary pores or evaporates. If a water-to-cement ratio less than about 0.40 is used, some cement will remain unhydrated.

To estimate the degree of hydration of a hydrated material the nonevaporable water content is often used. To convert the measured nonevaporable water into degrees of hydration, it is necessary to know the value of nonevaporable water-to-cement ratio \( w_{n/c} \) at complete hydration. Experimentally, this can be determined by preparing a high water-to-cement ratio cement paste (for example, 1.0) and continuously grinding while it hydrates in a roller mill. In this procedure, complete hydration of the cement will typically be achieved after 28 days.

Alternatively, an estimate of the value of nonevaporable water-to-cement ratio \( w_{n/c} \) at complete hydration can be obtained from the potential Bogue composition of the cement. Nonevaporable water contents for the major compounds of portland cement are provided in Table 2-7. For a typical Type I cement, these coefficients will generally result in a calculated \( w_{n/c} \) for completely hydrated cement somewhere between 0.22 and 0.25.

**PHYSICAL PROPERTIES OF CEMENT**

Specifications for cement place limits on both its physical properties and often chemical composition. An understanding of the significance of some of the physical properties is helpful in interpreting results of cement tests. Tests of the physical properties of the cements should be used to evaluate the properties of the cement, rather than the concrete. Cement specifications limit the properties with respect to the type of cement. Cement should be sampled in accordance with ASTM C183 (AASHTO T 127). During manufacture, cement is continuously monitored for its chemistry and the following properties:

**Particle Size and Fineness**

Portland cement consists of individual angular particles with a range of sizes, the result of pulverizing clinker in the grinding mill (Fig. 2-31 left). Approximately 95% of cement particles are smaller than 45 \( \mu \)m, with the average particle around 15 \( \mu \)m. Fig. 2-31 (right) illustrates the particle size distribution for a portland cement. The overall particle size distribution of cement is called “fineness.” The fineness of cement affects heat released and the rate of hydration. Greater cement fineness (smaller particle size) increases the rate at which cement hydrates and thus accelerates strength.

### Table 2-7. Nonevaporable Water Contents for Fully Hydrated Major Compounds of Cement

<table>
<thead>
<tr>
<th>Hydrated cement compound</th>
<th>Nonevaporable (combined) water content (g water/g cement compound)</th>
</tr>
</thead>
<tbody>
<tr>
<td>C₃S hydrate</td>
<td>0.24</td>
</tr>
<tr>
<td>C₂S hydrate</td>
<td>0.21</td>
</tr>
<tr>
<td>C₃A hydrate</td>
<td>0.40</td>
</tr>
<tr>
<td>C₄AF hydrate</td>
<td>0.37</td>
</tr>
<tr>
<td>Free lime (CaO)</td>
<td>0.33</td>
</tr>
</tbody>
</table>

![Image of cement paste cylinders](IMG12302)
development. The effects of greater fineness on paste strength are manifested principally during the first seven days.

In the early 1900s, cement fineness was expressed as the mass of cement per fractional size (percent weight retained on specific sieve sizes). Today, fineness is usually measured by the Blaine air-permeability test (ASTM C204 or AASHTO T 153—Fig. 2-32) that indirectly measures the surface area of the cement particles per unit mass. Cements with finer particles have more surface area in square meters per kilogram of cement. The older use of square centimeters per gram for reporting fineness is now considered archaic. Most cement standards only have minimum limits on fineness. ASTM C150 and AASHTO M 85 have maximum limits on fineness for Type II and Type IV cements of 420 m²/kg and 430 m²/kg respectively. The Wagner turbidimeter test (ASTM C115 or AASHTO T 98—Fig. 2-32), the 45-micrometer (No. 325) sieve (ASTM C430 or AASHTO T 192—Fig. 2-33) and the electronic (x-ray or laser) particle size analyzer (Fig. 2-34) can also be used to test fineness. Blaine fineness data are given in Table 2-6.

**Soundness**

Soundness refers to the ability of a hardened paste to retain its volume. Lack of soundness or delayed destructive expansion can be caused by excessive amounts of hard-burned free lime or magnesia. Most specifications for portland cement limit the magnesia (periclase) content and the maximum expansion as measured by the autoclave-expansion test. Since adoption of the autoclave-expansion test (ASTM C151 or AASHTO T 107) in 1943, there have been exceedingly few cases of abnormal expansion attributed to unsound cement (Fig. 2-35) (Gonnerman, Lerch, and Whiteside 1953).
Consistency

Consistency refers to the relative mobility of a freshly mixed cement paste or mortar or to its ability to flow. During cement testing, pastes are mixed to normal consistency as defined by a penetration of 10 ± 1 mm of the Vicat plunger (see ASTM C187 or AASHTO T 129 and Fig. 2-36). Mortars are mixed to obtain either a fixed water-cement ratio or to yield a flow within a prescribed range. The flow is determined on a flow table as described in ASTM C230 (AASHTO M 152) and ASTM C1437 (Fig. 2-37). Both the normal consistency method and the flow test are used to regulate water contents of pastes and mortars, respectively, to be used in subsequent tests; both allow comparing dissimilar ingredients with the same penetrability or flow.

Setting Time

The object of the setting time test is to determine (1) the time that elapses from the moment water is added until the paste ceases to be fluid and plastic (called initial set) and (2) the time required for the paste to acquire a certain degree of hardness (called final set).

To determine if a cement sets according to the time limits specified in cement specifications, tests are performed using either the Vicat apparatus (ASTM C191 or AASHTO T 131—Fig. 2-38) or the Gillmore needle (ASTM C266 or AASHTO T 154—Fig. 2-39).

The Vicat test governs if no test method is specified by the purchaser. Initial set of cement paste must not occur too early and final set must not occur too late. The setting times indicate that a paste is or is not undergoing normal hydra-
tion reactions. Sulfate (from gypsum or other sources) in the cement regulates setting time, but setting time is also affected by cement fineness, water-cement ratio, and any admixtures that may be used. Setting times of concretes do not correlate directly with setting times of pastes because of water loss to the air or substrate, presence of aggregate, and because of temperature differences in the field (as contrasted with the controlled temperature in a testing lab). Fig. 2-40 illustrates mean set times for portland cements.

dehydrates in the cement mill to form plaster. Stiffening is caused by the rapid crystallization of interlocking needle-like secondary gypsum. Additional mixing without added water breaks up these crystals to restore workability. Ettringite precipitation can also contribute to false set.

Flash set or quick set is evidenced by a rapid and early loss of workability in paste, mortar, or concrete. It is usually accompanied by the evolution of considerable heat resulting primarily from the rapid reaction of aluminates. If the proper amount or form of calcium sulfate is not available to control the calcium aluminate hydration, stiffening becomes apparent. Flash set cannot be dispelled, nor can the plasticity be regained by further mixing without the addition of water.

Proper stiffening results from the careful balance of the sulfate and aluminate compounds, as well as the temperature and fineness of the materials (which control the hydration and dissolution rates). The amount of sulfate formed into plaster has a significant effect. For example, with one particular cement, 2% plaster allowed a 5 hour set time, while 1% plaster caused flash set to occur, and 3% allowed false set to occur (Helmuth and others 1995).

Cements are tested for early stiffening using ASTM C451 (paste method) (AASHTO T 186), and ASTM C359 (mortar method) (AASHTO T 185), which use the penetration techniques of the Vicat apparatus. However, these tests do not address all the mixing, placing, temperature, and field conditions that can cause early stiffening. They also do not address early stiffening caused by interactions with other concrete ingredients. For example, concretes mixed for very short periods, less than a minute, tend to be more susceptible to early stiffening (ACI 225).
Compressive Strength

Compressive strength as specified by ASTM cement standards is that obtained from tests of 50-mm (2-in.) mortar cubes tested in accordance with ASTM C109 (AASHTO T 106) (Fig. 2-41). These cubes are made and cured in a prescribed manner using a standard sand.

Compressive strength is influenced by the cement type, or more precisely, the compound composition and fineness of the cement. ASTM C1157 has both minimum and maximum strength requirements while ASTM C150 and C95 (AASHTO M 85 and M 240) set only a minimum strength requirement. Minimum strength requirements in cement specifications are exceeded comfortably by most manufacturers. Therefore, it should not be assumed that two types of cement meeting the same minimum requirements will produce the same strength of mortar or concrete without modification of mix proportions.

In general, cement strengths (based on mortar-cube tests) cannot be used to predict concrete strengths with a great degree of accuracy because of the many variables in aggregate characteristics, concrete mixtures, construction procedures, and environmental conditions in the field (Weaver, Isabelle and Williamson 1970 and DeHayes 1990). Figs. 2-42 and 2-43 illustrate the strength development for standard mortars made with various types of portland cement. Wood (1992) provides long-term strength properties of mortars and concretes made with portland and blended cements. The strength uniformity of a cement from a single source may be determined by following the procedures outlined in ASTM C917.

Heat of Hydration

Heat of hydration is the heat generated when cement and water react. The amount of heat generated is dependent chiefly upon the chemical composition of the cement, with C₃A and C₅S being the compounds primarily responsible for high heat evolution. The water-cement ratio, fineness of the cement, and temperature of curing also are factors. An increase in the fineness, cement content, and curing temperature increases the heat of hydration. Although portland cement can evolve heat for many years, the rate of heat generation is greatest at early ages. A large amount of heat evolves within the first three days with the greatest rate of heat liberation usually occurring within the first 24 hours (Copeland and others 1960). The heat of hydration is tested in accordance with ASTM C186 or by conduction calorimetry (Figs. 2-44).

For most concrete elements, such as slabs, heat generation is not a concern because the heat is quickly dissipated into the environment. However, in structures of considerable mass, greater than a meter (yard) thick, the rate and amount of heat generated are important. If this heat is not rapidly dissipated, a significant rise in concrete temperature can occur. This may be undesirable since, after hardening at an elevated temperature, nonuniform cooling of the concrete mass to ambient temperature may create undesirable tensile stresses. On the other hand, a rise in concrete temperature caused by heat of hydration is often beneficial in cold weather since it helps maintain favorable curing temperatures.

Table 2-8 provides heat of hydration values for a variety of portland cements. This limited data show that Type III cement has a higher heat of hydration than other types while Type IV has the lowest. Also note the difference in heat generation between regular Type II cements and moderate heat Type II cements.
Fig. 2-43. Strength development of portland cement mortar cubes from combined statistics. The dashed line represents the mean value and the shaded area the range of values (adapted from Gebhardt 1995).

Fig. 2-44. Heat of hydration can be determined by (left) ASTM C186 or by (right) a conduction calorimeter. (IMG12447, IMG12448)

Table 2-8. ASTM C186 Heat of Hydration for Selected Portland Cements from the 1990s, kJ/kg*

<table>
<thead>
<tr>
<th></th>
<th>Type I cement</th>
<th>Type II cement</th>
<th>Type II moderate cement</th>
<th>Type III cement</th>
<th>Type IV cement</th>
<th>Type V cement</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>7 day</td>
<td>28 day</td>
<td>7 day</td>
<td>28 day</td>
<td>7 day</td>
<td>28 day</td>
</tr>
<tr>
<td>No. of samples</td>
<td>15</td>
<td>7</td>
<td>16</td>
<td>7</td>
<td>2</td>
<td>2</td>
</tr>
<tr>
<td>Average</td>
<td>349</td>
<td>400</td>
<td>344</td>
<td>398</td>
<td>263</td>
<td>283</td>
</tr>
<tr>
<td>Maximum</td>
<td>372</td>
<td>444</td>
<td>371</td>
<td>424</td>
<td>372</td>
<td>414</td>
</tr>
<tr>
<td>Minimum</td>
<td>320</td>
<td>377</td>
<td>308</td>
<td>372</td>
<td>227</td>
<td>368</td>
</tr>
<tr>
<td>% of Type I (7 day)</td>
<td>100</td>
<td>99</td>
<td>75</td>
<td>106</td>
<td>67</td>
<td>89</td>
</tr>
</tbody>
</table>

*This table is based on very limited data.
1 cal/g = 4.184 kJ/kg. PCA (1997).
Cements do not generate heat at a constant rate. The heat output during hydration of a typical Type I portland cement is illustrated in Fig. 2-45. The first peak shown in the heat profile is due to heat given off by the initial hydration reactions of cement compounds such as tricalcium aluminate. Sometimes called the heat of wetting, this first heat peak is followed by a period of slow activity known as the induction period. After several hours, a broad second heat peak attributed to tricalcium silicate hydration emerges, signaling the onset of the paste hardening process. Finally, there is the third peak due to the renewed activity of tricalcium aluminate; its intensity and location normally dependent on the amount of tricalcium aluminate and sulfate in the cement.

In calorimetry testing, the first heat measurements are obtained about 7 minutes after mixing the paste; as a result only the down slope of the first peak is observed (Stage 1, Fig. 2-45). The second peak (C₃S peak) often occurs between 6 and 12 hours. The third peak (renewed C₃A peak on conversion of AFT to AFm) occurs between 12 and 90 hours. This information can be helpful when trying to control temperature rise in mass concrete (Tang 1992).

When heat generation must be minimized in concrete, designers should choose a lower heat cement, such as an ASTM C150 (AASHTO M 85) Type II portland cement with the optional moderate heat of hydration requirements. Not all Type II cements are made for a moderate level of heat development, however, so the moderate heat option must be specifically requested. Type IV cement can also be used to control temperature rise, but it is rarely available. Moderate-heat and low-heat cements are also available in ASTM C595 (AASHTO M 240) and C1157 specifications. Supplementary cementitious materials can also be used to reduce temperature rise.

ASTM C150 (AASHTO M 85) has both a chemical approach and a physical approach to control heat of hydration. Either approach can be specified, but not both. ASTM C595 (AASHTO M 240) and ASTM C1157 use physical limits. See Poole (2007) for more information.

**Loss on Ignition**

Loss on ignition (LOI) of portland cement is determined by heating a cement sample of known weight to between 900°C and 1000°C until a constant weight is obtained. The weight loss of the sample is then determined. Normally, a high loss on ignition is an indication of prehydration and carbonation, which may be caused by improper or prolonged storage, or adulteration during transport. The test for loss on ignition is performed in accordance with ASTM C114 (AASHTO T 105) (Fig. 2-46). LOI values range from 0 to 3%.

**Density and Relative Density**

(Density and Specific Gravity)

The density of cement is defined as the mass of a unit volume of the solids or particles, excluding air between particles. It is reported as megagrams per cubic meter or grams per cubic centimeter (the numeric value is the same for both units). The particle density of portland cement ranges from 3.10 to 3.25, averaging 3.15 Mg/m³. Portland-blast-furnace-slag and portland-pozzolan cements have densities ranging from 2.90 to 3.15, averaging 3.05 Mg/m³. The density of a cement, determined by ASTM C188 (AASHTO T 133) (Fig. 2-47), is not an indication of the cement’s quality; its principal use is in mixture proportioning calculations.

For mixture proportioning, it may be more useful to express the density as relative density (also called specific gravity). The relative density is a dimensionless number determined by dividing the cement density by the density of water at 4°C. The density of water at 4°C is 1.0 Mg/m³ (1.0 g/cm³ or 1000 kg/m³).

---

**Fig. 2-45. Heat evolution as a function of time for cement paste. Stage 1 is heat of wetting or initial hydrolysis (C₃A and C₃S hydration). Stage 2 is a dormant period related to initial set. Stage 3 is an accelerated reaction of the hydration products that determines rate of hardening and final set. Stage 4 decelerates formation of hydration products and determines the rate of early strength gain. Stage 5 is a slow, steady formation of hydration products establishing the rate of later strength gain.**

**Fig. 2-46. Loss on ignition test of cement. (IMG12305)**
THERMAL ANALYSIS

Thermal analysis techniques have been available for years to analyze hydraulic reactions and the interactions of cement with both mineral and chemical admixtures (Figs. 2-45 and 2-49). Traditionally, thermal analysis was not part of a routine test program for cement. However, thermal analysis has gained recent popularity in analyzing chemical and physical properties of cementitious materials and raw materials for cement manufacture (Bhatty 1993, Shkolnik and Miller 1996, and Tennis 1997).

Thermal analysis involves heating a small sample at a controlled rate to high temperatures (up to 1000°C or more). As compounds react or decompose, changes that occur as a function of time or temperature are recorded. As the sample’s temperature changes, various reactions occur that cause a change in the sample’s weight, temperature, energy, or state. A solid can melt, vaporize, decompose to a gas with a residual solid, or react with a gas (at elevated temperatures) to form a different solid or a different solid and another gas.

Typical uses for thermal analysis include:

- identifying which hydration products have formed and in what quantities
- solving early stiffening problems
- identifying impurities in raw materials
- determining the amount of weathering in clinker or cement
- estimating the reactivity of pozzolans and slags for use in blended cements
- identifying the organic matter content and variations in a quarry
- quantifying the amount of carbonation that has taken place in an exposed sample
- analyzing durability problems with concrete.

Specific thermal analysis techniques are discussed below.

Bulk Density

The bulk density of cement is defined as the mass of cement particles plus air between particles per unit volume. The bulk density of cement can vary considerably depending on how it is handled and stored. Portland cement that is fluffed up may weigh only 830 kg/m³ (52 pcf), whereas when consolidated by vibration, the same cement can weigh as much as 1650 kg/m³ (103 pcf) (Toler 1963). For this reason, good practice dictates that cement must be weighed for each batch of concrete produced, as opposed to use of a volumetric measure (Fig. 2-48).

A relative density of 3.15 is assumed for portland cement in volumetric computations of concrete mix proportioning. However, as mix proportions list quantities of concrete ingredients in kilograms or pounds, the relative density must be multiplied by the density of water at 4°C stated as 1000 kg/m³ (62.4 lb/ft³) to determine the particle density in kg/m³ or lb/ft³. This product is then divided into the mass or weight of cement to determine the absolute volume of cement in cubic meters or cubic feet.
Thermogravimetric Analysis (TGA)

Thermogravimetric analysis (TGA) is a technique that measures the mass (weight) of a sample as it is being heated (or cooled) at a controlled rate. The weight change of a sample depends on the composition of the sample, the temperature, the heating rate, and the type of gas in the furnace (air, oxygen, nitrogen, argon, or other gas). A change in mass within a specific temperature range identifies the presence of a particular chemical compound. The magnitude of the weight change indicates the amount of the compound in the sample.

A sample’s free water will evaporate, lowering the mass of the sample, as the temperature is raised from room temperature to about 100°C. The sample will also lose some of the water bound in hydration products, particularly some of the water in the aluminates hydrates. Between about 100°C and 400°C, water bound in the hydration products, primarily the C-S-H gel, will be lost, together with the remainder of water in the aluminates hydrates. Between about 400°C and 500°C, calcium hydroxide provides a relatively distinct weight loss as it decomposes to calcium oxide (a solid) and water vapor. The amount of this weight loss can be used to determine how much calcium hydroxide was originally present in the sample. Above 500°C, a small amount of additional water may be lost from the hydration products. Carbonated phases lose carbon dioxide around 800°C.

By quantifying the amount of calcium hydroxide, a TGA gives an indication of the degree of hydration that has taken place in a sample. The reactivity of pozzolans can be assessed by tracking the disappearance of calcium hydroxide due to pozzolanic reaction.

Differential Thermal Analysis (DTA)

Differential thermal analysis (DTA) is an analytical method in which the difference in temperature between a sample and a control is measured as the samples are heated. The control is usually an inert material, like powdered alumina, that does not react over the temperature range being studied. If a sample undergoes a reaction at a certain temperature, then its temperature will increase or decrease relative to the (inert) control as the reaction gives off heat (exothermic) or adsorbs heat (endothermic). A thermocouple measures the temperature of each material, allowing the difference in temperature to be recorded. DTA is ideal for monitoring the transformation of cement compounds during hydration. DTA can be performed simultaneously with TGA.

Differential Scanning Calorimetry (DSC)

In differential scanning calorimetry (DSC), the heat absorbed or released is measured directly as a function of temperature or time and compared to a reference. An advantage of DTA and DSC methods is that no mass change is required; thus if a sample melts without vaporizing, measurements can still be made.

Like DTA, DSC can be used to determine what compounds are present at different stages of hydration. Fig. 2-50 shows two differential scanning calorimetry curves or thermograms. The upper curve (a) shows a portland cement paste after 15 minutes of reaction. The peaks in the curve between 100°C and 200°C are due to the endothermic (heat adsorbing) decomposition of gypsum and ettringite, while the peak around 270°C is due to syngenite (potassium calcium sulfate hydrate). Near 450°C, a smaller peak can be seen, due to calcium hydroxide.

The lower curve (b) in Fig. 2-50 shows the same cement paste after 24 hours of hydration. Note the disappearance of the peak due to gypsum, the reduction in the size of the peak due to syngenite and the growth of those due to ettringite and calcium hydroxide. The size of the areas under the curves are related to the quantity of the material in the sample.

Virtual Cement Testing

Computer technology now allows simulation of cement compounds (Fig. 2-51), hydration, microstructure development, and physical properties. Combinations of materials, cement compounds, or particle size distributions can be examined to predict cement performance. Just some of the properties that can be simulated and predicted include heat of hydration, adiabatic heat signature, compressive strength, setting time, rheology (yield stress and viscosity),
percolation, porosity, diffusivity, thermal conductivity, electrical conductivity, carbonation, elastic properties, drying profiles, susceptibility to degradation mechanisms, autogenous shrinkage, and volumes of hydration reactants and products as a function of time. The effects of varying sulfate and alkali contents can be examined, along with interaction with supplementary cementitious materials and chemical admixtures. Computer modeling predicts performance without the expense and time requirements of physical testing (NIST 2007).

TRANSPORTATION AND PACKAGING

Around 100 million metric tons of cement are shipped per year throughout the United States. Most portland cements are shipped in bulk by rail, truck, barge, or ship (Fig. 2-52). Pneumatic (pressurized air) loading and unloading of the transport vehicle is the most popular means of handling bulk cement. However, the recent introduction of large sacks holding one to twelve tons of cement provides a new alternative to bulk cement handling. Bulk cement is measured by the metric ton (1000 kg) or short ton (2000 lb).

Cement is also bagged for convenient use at construction sites (Fig. 2-53) and for small jobs. In the United States a bag of portland cement has a mass of 42 kg (92.6 lb) and a volume of 28 liters. Traditionally a U.S. bag contains 94 lb (42.6 kg) and has a volume of one cubic foot. The mass of masonry cement and mortar cement by the bag is 36 kg for Type M, 34 kg for Type S, and 32 kg for Type N. Plastic cement has a mass of 42 kg for Type M and 35 kg for Type S. All bag weights are based on a 28 liter volume standard. Information for quantities of blended and specialty cements can be found on the bag. Partial bag sizes are also available in some areas.

In Canada, a bag of portland cement has a mass of 40 kg, and in other countries a bag of cement can have a mass of 25 kg or 50 kg.

Because of the variability of bag sizes and the presence of supplementary cementitious materials, cement bag factor terminology, such as “a six bag mix” should not be used to describe the cement content of concrete mixtures.

In the 1800s and early 1900s, cement was shipped in barrels; one barrel contained 4 bags (171 kg or 376 lbs or 4 cu ft) of cement. Use of the barrel unit is archaic and should not be used.

STORAGE OF CEMENT

Cement is a moisture-sensitive material; if kept dry, it will retain its quality indefinitely. Cement stored in contact with damp air or moisture sets more slowly and has less strength than cement that is kept dry. At the cement plant, and at ready mixed concrete facilities, bulk cement is stored in silos. The relative humidity in a warehouse or shed used to store bagged cement should be as low as possible. All cracks and openings in walls and roofs should be closed. Cement bags should not be stored on damp floors but should rest on pallets. Bags should be stacked close together to reduce air circulation but should never be
Fig. 2-53. A small amount of cement is shipped in bags, primarily for mortar applications and for small projects. (IMG12439)

Fig. 2-54. When stored on the job, cement should be protected from moisture. (IMG12438)

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CHAPTER 3

Fly Ash, Slag, Silica Fume, and Natural Pozzolans

Fly ash, ground granulated blast-furnace slag, silica fume, and natural pozzolans, such as calcined shale, calcined clay or metakaolin, are materials that, when used in conjunction with portland or blended cement, contribute to the properties of the hardened concrete through hydraulic or pozzolanic activity or both (Fig. 3-1). A pozzolan is a siliceous or aluminosiliceous material that, in finely divided form and in the presence of moisture, chemically reacts with the calcium hydroxide released by the hydration of portland cement to form calcium silicate hydrate and other cementitious compounds. Pozzolans and slags are generally categorized as supplementary cementitious materials or mineral admixtures. Table 3-1 lists the applicable specifications these materials meet. The use of these materials in blended cements is discussed in Chapter 2 and by Detwiler, Bhatti, and Bhattacharya (1996).

The practice of using supplementary cementitious materials in concrete mixtures has been growing in North America since the 1970s. There are similarities between many of these materials in that most are byproducts of other industrial processes; their judicious use is desirable not only from the national environmental and energy conservation standpoint but also for the technical benefits they provide concrete.

Supplementary cementitious materials are added to concrete as part of the total cementitious system. They may be used in addition to or as a partial replacement of portland cement or blended cement in concrete, depending on the properties of the materials and the desired effect on concrete.

Supplementary cementitious materials are used to improve a particular concrete property, such as resistance to alkali-aggregate reactivity. The optimum amount to use should be established by testing to determine (1) whether the material is indeed improving the property, and (2) the correct dosage rate, as an overdose or underdose can be harmful or not achieve the desired effect. Supplementary cementitious materials also react differently with different cements.

Traditionally, fly ash, slag, calcined clay, calcined shale, and silica fume were used in concrete individually. Today, due to improved access to these materials, concrete producers can combine two or more of these materials to optimize concrete properties. Mixtures using three cementitious materials, called ternary mixtures, are becoming more

Fig. 3-1. Supplementary cementitious materials. From left to right, fly ash (Class C), metakaolin (calcined clay), silica fume, fly ash (Class F), slag, and calcined shale. (IMG12187)

Fig. 3-2. Scanning electron microscope (SEM) micrograph of fly ash particles at 1000X. Although most fly ash spheres are solid, some particles, called cenospheres, are hollow (as shown in the micrograph). (IMG12309)
FLY ASH

Fly ash, the most widely used supplementary cementitious material in concrete, is a byproduct of the combustion of pulverized coal in electric power generating plants. Upon ignition in the furnace, most of the volatile matter and carbon in the coal are burned off. During combustion, the coal’s mineral impurities (such as clay, feldspar, quartz, and shale) fuse in suspension and are carried away from the combustion chamber by the exhaust gases. In the process, the fused material cools and solidifies into spherical glassy particles called fly ash (Fig. 3-2). The fly ash is then collected from the exhaust gases by electrostatic precipitators or bag filters. Fly ash is a finely divided powder resembling portland cement (Fig. 3-3).

Most of the fly ash particles are solid spheres and some are hollow cenospheres. Also present are plerospheres, which are spheres containing smaller spheres. Ground materials, such as portland cement, have solid angular particles. The particle sizes in fly ash vary from less than 1 μm (micrometer) to more than 100 μm with the typical particle size measuring under 20 μm. Only 10% to 30% of the particles by mass are larger than 45 μm. The surface area is typically 300 to 500 m²/kg, although some fly ashes can have surface areas as low as 200 m²/kg and as high as 700 m²/kg. For fly ash without close compaction, the bulk density (mass per unit volume including air between particles) can vary from 540 to 860 kg/m³ (34 to 54 lb/ft³), whereas with close packed storage or vibration, the range can be 1120 to 1500 kg/m³ (70 to 94 lb/ft³).

Fly ash is primarily silicate glass containing silica, alumina, iron, and calcium. Minor constituents are magnesium, sulfur, sodium, potassium, and carbon. Crystalline compounds are present in small amounts. The relative density (specific gravity) of fly ash generally ranges between 1.9 and 2.8 and the color is generally gray or tan.

ASTM C618 (AASHTO M 295) Class F and Class C fly ashes are commonly used as pozzolanic admixtures for general purpose concrete (Fig. 3-4). Class F materials are generally low-calcium (less than 10% CaO) fly ashes with

<table>
<thead>
<tr>
<th>Table 3-1. Specifications and Classes of Supplementary Cementitious Materials</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ground granulated iron blast-furnace slags—ASTM C989 (AASHTO M 302)</td>
</tr>
<tr>
<td>Grade 80</td>
</tr>
<tr>
<td>Slags with a low activity index</td>
</tr>
<tr>
<td>Grade 100</td>
</tr>
<tr>
<td>Slags with a moderate activity index</td>
</tr>
<tr>
<td>Grade 120</td>
</tr>
<tr>
<td>Slags with a high activity index</td>
</tr>
<tr>
<td>Fly ash and natural pozzolans—ASTM C618 (AASHTO M 295)</td>
</tr>
<tr>
<td>Class N</td>
</tr>
<tr>
<td>Raw or calcined natural pozzolans including:</td>
</tr>
<tr>
<td>Diatomaceous earths</td>
</tr>
<tr>
<td>Opaline cherts and shales</td>
</tr>
<tr>
<td>Tuffs and volcanic ashes or pumicites</td>
</tr>
<tr>
<td>Calcined clays, including metakaolin, and shales</td>
</tr>
<tr>
<td>Class F</td>
</tr>
<tr>
<td>Fly ash with pozzolanic properties</td>
</tr>
<tr>
<td>Class C</td>
</tr>
<tr>
<td>Fly ash with pozzolanic and cementitious properties</td>
</tr>
<tr>
<td>Silica fume—ASTM C1240</td>
</tr>
</tbody>
</table>

Fig. 3-3. Fly ash, a powder resembling cement, has been used in concrete since the 1930s. (IMG12190)

Fig. 3-4. Fly ash, slag, and calcined clay or calcined shale are used in general purpose construction, such as (left to right) walls for residential buildings, pavements, high-rise towers, and dams. (IMG12431, IMG12310, IMG12430, IMG12429)
carbon contents usually less than 5%, but some may be as high as 10%. Class C materials are often high-calcium (10% to 30% CaO) fly ashes with carbon contents less than 2%. Many Class C ashes when exposed to water will hydrate and harden in less than 45 minutes. Some fly ashes meet both Class F and Class C classifications.

Fly ash is used in about 50% of ready mixed concrete (PCA 2000). Class F fly ash is often used at dosages of 15% to 25% by mass of cementitious material and Class C fly ash is used at dosages of 15% to 40% by mass of cementitious material. Dosage varies with the reactivity of the ash and the desired effects on the concrete (Thomas 2007; Helmuth 1987; and ACI 232 2004). Refer to Thomas (2007) for more information on optimizing the use of fly ash in concrete.

SLAG

Ground granulated blast-furnace slag (Fig. 3-5), also called slag cement, is made from iron blast-furnace slag; it is a nonmetallic hydraulic cement consisting essentially of silicates and aluminosilicates of calcium developed in a molten condition simultaneously with iron in a blast furnace. The molten slag at a temperature of about 1500°C (2730°F) is rapidly chilled by quenching in water to form a glassy sandlike granulated material. The granulated material, which is ground to less than 45 microns, has a surface area fineness of about 400 to 600 m²/kg Blaine. The relative density (specific gravity) for ground granulated blast-furnace slag is in the range of 2.85 to 2.95. The bulk density varies from 1050 to 1375 kg/m³ (66 to 86 lb/ft³).

The rough and angular-shaped ground slag (Fig. 3-6) in the presence of water and an activator, NaOH or CaOH, both supplied by portland cement, hydrates and sets in a manner similar to portland cement. However, air-cooled slag does not have the hydraulic properties of water-cooled slag.

Granulated blast furnace slag was first developed in Germany in 1853 (Malhotra 1996). Ground slag has been used as a cementitious material in concrete since the beginning of the 1900s (Abrams 1925). Ground granulated blast-furnace slag, when used in general purpose concrete in North America, commonly constitutes between 30% and 45% of the cementing material in the mix (Fig. 3-4) (PCA 2000). Some slag concretes have a slag component of 70% or more of the cementitious material. ASTM C989 (AASHTO M 302) classifies slag by its increasing level of reactivity as Grade 80, 100, or 120 (Table 3-1). ASTM C1073
Natural pozzolans have been used for centuries. The term "pozzolan" comes from a volcanic ash mined at Pozzuoli, a village near Naples, Italy, following the 79 AD eruption of Mount Vesuvius. However, the use of volcanic ash and calcined clay dates back to 2000 BC and earlier in other cultures. Many of the Roman, Greek, Indian, and Egyptian pozzolan concrete structures can still be seen today, attesting to the durability of these materials.

The North American experience with natural pozzolans dates back to early 20th century public works projects, such as dams, where they were used to control temperature rise in mass concrete and provide cementitious material. In addition to controlling heat rise, natural pozzolans were used to improve resistance to sulfate attack and were among the first materials to be found to mitigate alkali-silica reaction.

The most common natural pozzolans used today are processed materials, which are heat treated in a kiln and then ground to a fine powder (Figs. 3-10, 3-11, and 3-12); they include calcined clay, calcined shale, and metakaolin.
Calcined clays are used in general purpose concrete construction much the same as other pozzolans (Fig. 3-4). They can be used as a partial replacement for the cement, typically in the range of 15% to 35%, and to enhance resistance to sulfate attack, control alkali-silica reactivity, and reduce permeability. Calcined clays have a relative density of between 2.40 and 2.61 with Blaine fineness ranging from 650 m²/kg to 1350 m²/kg. Calcined shale may contain on the order of 5% to 10% calcium, which results in the material having some cementing or hydraulic properties on its own. Because of the amount of residual calcite that is not fully calcined, and the bound water molecules in the clay minerals, calcined shale will have a loss on ignition (LOI) of perhaps 1% to 5%. The LOI value for calcined shale is not a measure or indication of carbon content as would be the case in fly ash.

Metakaolin, a special calcined clay, is produced by low-temperature calcination of high purity kaolin clay. The product is ground to an average particle size of about 1 to 2 micrometers. Metakaolin is used in special applications where very low permeability or very high strength is required. In these applications, metakaolin is used more as an additive to the concrete rather than a replacement of cement; typical additions are around 10% of the cement mass.

Natural pozzolans are classified by ASTM C618 (AASHTO M 295) as Class N pozzolans (Table 3-1). ACI 232 (2000) provides a review of natural pozzolans. Table 3-2 illustrates typical chemical analysis and selected properties of pozzolans.

### EFFECTS ON FRESHLY MIXED CONCRETE

This section provides a brief understanding of the freshly mixed concrete properties that supplementary cementitious materials affect and their degree of influence. First it should be noted that these materials vary considerably in their effect on concrete mixtures. The attributes of these materials when added separately to a concrete mixture can also be found in blended cements using supplementary cementitious materials.

### Water Requirements

Concrete mixtures containing fly ash generally require less water (about 1% to 10% less at normal dosages) for a given slump than concrete containing only portland cement. Higher dosages can result in greater water reduction (Table 3-3). However, some fly ashes can in-

---

**Table 3-2. Chemical Analysis and Selected Properties of Typical Fly Ash, Slag, Silica Fume, Calcined Clay, Calcined Shale, and Metakaolin**

<table>
<thead>
<tr>
<th></th>
<th>Class F</th>
<th>Class C</th>
<th>Ground slag</th>
<th>Silica fume</th>
<th>Calcined clay</th>
<th>Calcined shale</th>
<th>Metakaolin</th>
</tr>
</thead>
<tbody>
<tr>
<td>SiO₂, %</td>
<td>52</td>
<td>35</td>
<td>35</td>
<td>90</td>
<td>58</td>
<td>50</td>
<td>53</td>
</tr>
<tr>
<td>Al₂O₃, %</td>
<td>23</td>
<td>18</td>
<td>12</td>
<td>0.4</td>
<td>29</td>
<td>20</td>
<td>43</td>
</tr>
<tr>
<td>Fe₂O₃, %</td>
<td>11</td>
<td>6</td>
<td>1</td>
<td>0.4</td>
<td>4</td>
<td>8</td>
<td>0.5</td>
</tr>
<tr>
<td>CaO, %</td>
<td>5</td>
<td>21</td>
<td>40</td>
<td>1.6</td>
<td>1</td>
<td>8</td>
<td>0.1</td>
</tr>
<tr>
<td>SO₃, %</td>
<td>0.8</td>
<td>4.1</td>
<td>2</td>
<td>0.4</td>
<td>0.5</td>
<td>0.4</td>
<td>0.1</td>
</tr>
<tr>
<td>Na₂O, %</td>
<td>1.0</td>
<td>5.8</td>
<td>0.3</td>
<td>0.5</td>
<td>0.2</td>
<td>—</td>
<td>0.05</td>
</tr>
<tr>
<td>K₂O, %</td>
<td>2.0</td>
<td>0.7</td>
<td>0.4</td>
<td>2.2</td>
<td>2</td>
<td>—</td>
<td>0.4</td>
</tr>
<tr>
<td>Total Na eq. alk, %</td>
<td>2.2</td>
<td>6.3</td>
<td>0.6</td>
<td>1.9</td>
<td>1.5</td>
<td>—</td>
<td>0.3</td>
</tr>
<tr>
<td>Loss on ignition, %</td>
<td>2.8</td>
<td>0.5</td>
<td>1.0</td>
<td>3.0</td>
<td>1.5</td>
<td>3.0</td>
<td>0.7</td>
</tr>
<tr>
<td>Fineness, m²/kg</td>
<td>420</td>
<td>420</td>
<td>400</td>
<td>20,000</td>
<td>990</td>
<td>730</td>
<td>17,000</td>
</tr>
<tr>
<td>Blaine Nitrogen absorp.</td>
<td>2.38</td>
<td>2.65</td>
<td>2.94</td>
<td>2.40</td>
<td>2.50</td>
<td>2.63</td>
<td>2.50</td>
</tr>
<tr>
<td>Relative density</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

---

![Fig. 3-11. Metakaolin, a calcined clay.](IMG12193)

![Fig. 3-12. Scanning electron microscope micrograph of calcined clay particles at 2000X.](IMG12432)
bleed water is primarily due to the reduced water demand in fly ash concretes. Gebler and Klieger (1986) correlate reduced bleeding of concrete to the reduced water requirement of fly ash mortar. Concretes containing ground slags of comparable fineness to that of cement tend to show an increased rate and amount of bleeding than plain concretes, but this appears to have no adverse effect on segregation. Slags ground finer than cement reduce bleeding.

Silica fume is very effective in reducing both bleeding and segregation; as a result, higher slumps may be used. Calcined clays, calcined shales, and metakaolin have little effect on bleeding.

**Air Content**

The amount of air-entraining admixture required to obtain a specified air content is normally greater when fly ash is used. Class C ash requires less air-entraining admixture than Class F ash and tends to lose less air during mixing (Table 3-5). Ground slags have variable effects on the required dosage rate of air-entraining admixtures. Silica fume has a marked influence on the air-entraining admixture requirement, which in most cases rapidly increases with an increase in the amount of silica fume used in the concrete. The inclusion of both fly ash and silica fume in non-air-entrained concrete will generally reduce the amount of entrapped air.

**Workability**

Fly ash, slag, and calcined clay and shale generally improve the workability of concretes of equal slump. Silica fume may contribute to stickiness of a concrete mixture; adjustments, including the use of high-range water reducers, may be required to maintain workability and permit proper compaction and finishing.

**Bleeding and Segregation**

Concretes using fly ash generally exhibit less bleeding and segregation than plain concretes (Table 3-4). This effect makes the use of fly ash particularly valuable in concrete mixtures made with aggregates that are deficient in fines. The reduction in water demand up to 5% (Gebler and Klieger 1986). Fly ash reduces water demand in a manner similar to liquid chemical water reducers (Helmuth 1987). Ground slag usually decreases water demand by 1% to 10%, depending on dosage.

The water demand of concrete containing silica fume increases with increasing amounts of silica fume, unless a water reducer or plasticizer is used. Some lean mixes may not experience an increase in water demand when only a small amount (less than 5%) of silica fume is present.

Calcined clays and calcined shales generally have little effect on water demand at normal dosages; however, other natural pozzolans can significantly increase or decrease water demand.

---

**Table 3-3. Effect of Fly Ash on Mixing Water Requirements for Air-Entrained Concrete**

<table>
<thead>
<tr>
<th>Fly ash mix identification</th>
<th>Class of fly ash</th>
<th>Fly ash content, % by mass of cementitious material</th>
<th>Change in mixing water requirement compared to control, %</th>
</tr>
</thead>
<tbody>
<tr>
<td>C1A</td>
<td>C</td>
<td>25</td>
<td>-6</td>
</tr>
<tr>
<td>C1D</td>
<td>F</td>
<td>25</td>
<td>-2</td>
</tr>
<tr>
<td>C1E</td>
<td>F</td>
<td>25</td>
<td>-6</td>
</tr>
<tr>
<td>C1F</td>
<td>C</td>
<td>25</td>
<td>-8</td>
</tr>
<tr>
<td>C1G</td>
<td>C</td>
<td>25</td>
<td>-6</td>
</tr>
<tr>
<td>C1J</td>
<td>F</td>
<td>25</td>
<td>-6</td>
</tr>
<tr>
<td>C2A</td>
<td>C</td>
<td>50</td>
<td>-18</td>
</tr>
<tr>
<td>C2D</td>
<td>F</td>
<td>50</td>
<td>-6</td>
</tr>
<tr>
<td>C2E</td>
<td>F</td>
<td>50</td>
<td>-14</td>
</tr>
<tr>
<td>C2F</td>
<td>C</td>
<td>50</td>
<td>-16</td>
</tr>
<tr>
<td>C2G</td>
<td>C</td>
<td>50</td>
<td>-12</td>
</tr>
<tr>
<td>C2J</td>
<td>F</td>
<td>50</td>
<td>-10</td>
</tr>
</tbody>
</table>

All mixtures had cementitious materials contents of 335 kg/m^3 (564 lb/yd^3), a slump of 125 ± 25 mm (5 ± 1 in.), and an air content of 6 ± 1%. Water to cement plus fly ash ratios varied from 0.40 to 0.48 (Whiting 1989).

---

**Table 3-4. Effect of Fly Ash on Bleeding of Concrete (ASTM C232, AASHTO T 158)**

<table>
<thead>
<tr>
<th>Fly ash mix identification</th>
<th>Class of fly ash</th>
<th>Change in mixing water requirement compared to control, %</th>
</tr>
</thead>
<tbody>
<tr>
<td>C1A</td>
<td>C</td>
<td>-6</td>
</tr>
<tr>
<td>C1D</td>
<td>F</td>
<td>-2</td>
</tr>
<tr>
<td>C1E</td>
<td>F</td>
<td>-6</td>
</tr>
<tr>
<td>C1F</td>
<td>C</td>
<td>-8</td>
</tr>
<tr>
<td>C1G</td>
<td>C</td>
<td>-6</td>
</tr>
<tr>
<td>C1J</td>
<td>F</td>
<td>-6</td>
</tr>
<tr>
<td>C2A</td>
<td>C</td>
<td>-18</td>
</tr>
<tr>
<td>C2D</td>
<td>F</td>
<td>-6</td>
</tr>
<tr>
<td>C2E</td>
<td>F</td>
<td>-14</td>
</tr>
<tr>
<td>C2F</td>
<td>C</td>
<td>-16</td>
</tr>
<tr>
<td>C2G</td>
<td>C</td>
<td>-12</td>
</tr>
<tr>
<td>C2J</td>
<td>F</td>
<td>-10</td>
</tr>
</tbody>
</table>

**Table 3-4. Effect of Fly Ash on Bleeding of Concrete (ASTM C232, AASHTO T 158)**

<table>
<thead>
<tr>
<th>Fly ash mix identification</th>
<th>Class of fly ash</th>
<th>Change in mixing water requirement compared to control, %</th>
</tr>
</thead>
<tbody>
<tr>
<td>C1A</td>
<td>C</td>
<td>-6</td>
</tr>
<tr>
<td>C1D</td>
<td>F</td>
<td>-2</td>
</tr>
<tr>
<td>C1E</td>
<td>F</td>
<td>-6</td>
</tr>
<tr>
<td>C1F</td>
<td>C</td>
<td>-8</td>
</tr>
<tr>
<td>C1G</td>
<td>C</td>
<td>-6</td>
</tr>
<tr>
<td>C1J</td>
<td>F</td>
<td>-6</td>
</tr>
<tr>
<td>C2A</td>
<td>C</td>
<td>-18</td>
</tr>
<tr>
<td>C2D</td>
<td>F</td>
<td>-6</td>
</tr>
<tr>
<td>C2E</td>
<td>F</td>
<td>-14</td>
</tr>
<tr>
<td>C2F</td>
<td>C</td>
<td>-16</td>
</tr>
<tr>
<td>C2G</td>
<td>C</td>
<td>-12</td>
</tr>
<tr>
<td>C2J</td>
<td>F</td>
<td>-10</td>
</tr>
</tbody>
</table>

All mixtures had cementitious materials contents of 307 kg/m^3 (517 lb/yd^3), a slump of 75 ± 25 mm (3 ± 1 in.), and an air content of 6 ± 1%. Fly ash mixtures contained 25% ash by mass of cementitious material (Gebler and Klieger 1986).

**Volume of bleed water per surface area.**
The amount of air-entraining admixture required for a certain air content in the concrete is a function of the fineness, carbon content, alkali content, organic material content, loss on ignition, and presence of impurities in the fly ash. Increases in alkalies decrease air-entraining agent dosages, while increases in the other properties increase dosage requirements. The Foam Index test provides an indication of the required dosage of air-entraining admixture for various fly ashes relative to non-ash mixtures; it can also be used to anticipate the need to increase or decrease the dosage based on changes in the foam index (Gebler and Klieger 1983).

### Heat of Hydration

Fly ash, natural pozzolans, and ground slag have a lower heat of hydration than portland cement; consequently their use will reduce the amount of heat built up in a concrete structure (Fig. 3-13). Calcined clay imparts a heat of hydration similar to moderate heat cement (Barger and others 1997). Some pozzolans have a heat of hydration of only 40% that of Type I portland cement. This reduction in temperature rise is especially beneficial in concrete used for massive structures. Silica fume may or may not reduce the heat of hydration. Detwiler, Bhatty and Bhattacharja (1996) provide a review of the effect of pozzolans and slags on heat generation.

### Setting Time

The use of fly ash and ground granulated blast-furnace slag will generally retard the setting time of concrete (Table 3-6). The degree of set retardation depends on factors such as water-to-cement ratio, temperature, and the type of fly ash used.

### Table 3-5. Effect of Fly Ash on Air-Entraining Admixture Dosage and Air Retention

<table>
<thead>
<tr>
<th>Fly ash mixtures</th>
<th>Percent of air-entraining admixture relative to control</th>
<th>Air content, %</th>
<th>Minutes after initial mixing</th>
</tr>
</thead>
<tbody>
<tr>
<td>Identification</td>
<td>Class of fly ash</td>
<td>%</td>
<td>0</td>
</tr>
<tr>
<td>A</td>
<td>C</td>
<td>126</td>
<td>7.2</td>
</tr>
<tr>
<td>B</td>
<td>F</td>
<td>209</td>
<td>5.3</td>
</tr>
<tr>
<td>C</td>
<td>F</td>
<td>553</td>
<td>7.0</td>
</tr>
<tr>
<td>D</td>
<td>F</td>
<td>239</td>
<td>6.6</td>
</tr>
<tr>
<td>E</td>
<td>F</td>
<td>190</td>
<td>5.6</td>
</tr>
<tr>
<td>F</td>
<td>C</td>
<td>173</td>
<td>6.8</td>
</tr>
<tr>
<td>G</td>
<td>C</td>
<td>158</td>
<td>5.5</td>
</tr>
<tr>
<td>H</td>
<td>F</td>
<td>170</td>
<td>7.6</td>
</tr>
<tr>
<td>I</td>
<td>C</td>
<td>149</td>
<td>6.6</td>
</tr>
<tr>
<td>J</td>
<td>F</td>
<td>434</td>
<td>5.5</td>
</tr>
<tr>
<td>Control mixture</td>
<td></td>
<td>100</td>
<td>6.6</td>
</tr>
</tbody>
</table>

Concretes had a cementitious materials content of 307 kg/m³ (517 lb/yd³) and a slump of 75 ± 25 mm (3 ± 1 in.). Fly ash mixtures contained 25% ash by mass of cementitious material (Gebler and Klieger 1983).

### Table 3-6. Effect of Fly Ash on Setting Time of Concrete

<table>
<thead>
<tr>
<th>Fly ash test mixtures</th>
<th>Setting time, hr:min</th>
<th>Retardation relative to control, hr:min</th>
</tr>
</thead>
<tbody>
<tr>
<td>Identification</td>
<td>Class of fly ash</td>
<td>Initial</td>
</tr>
<tr>
<td>A</td>
<td>C</td>
<td>4:30</td>
</tr>
<tr>
<td>B</td>
<td>F</td>
<td>4:40</td>
</tr>
<tr>
<td>C</td>
<td>F</td>
<td>4:25</td>
</tr>
<tr>
<td>D</td>
<td>F</td>
<td>5:05</td>
</tr>
<tr>
<td>E</td>
<td>F</td>
<td>4:25</td>
</tr>
<tr>
<td>F</td>
<td>C</td>
<td>4:25</td>
</tr>
<tr>
<td>G</td>
<td>C</td>
<td>4:55</td>
</tr>
<tr>
<td>H</td>
<td>F</td>
<td>5:10</td>
</tr>
<tr>
<td>I</td>
<td>C</td>
<td>5:00</td>
</tr>
<tr>
<td>J</td>
<td>F</td>
<td>5:10</td>
</tr>
<tr>
<td>Average of:</td>
<td>Class C</td>
<td>4:40</td>
</tr>
<tr>
<td></td>
<td>Class F</td>
<td>4:50</td>
</tr>
<tr>
<td>Control mixture</td>
<td></td>
<td>4:15</td>
</tr>
</tbody>
</table>

Concretes had a cementitious materials content of 307 kg/m³ (517 lb/yd³). Fly ash mixtures contained 25% ash by mass of cementitious material. Water to cement plus fly ash ratio = 0.40 to 0.45. Tested at 23°C (73°F) (Gebler and Klieger 1986).
as the amount of portland cement, water requirement, the type and reactivity of the slag or pozzolan dosage, and the temperature of the concrete. Set retardation is an advantage during hot weather, allowing more time to place and finish the concrete. However, during cold weather, pronounced retardation can occur with some materials, significantly delaying finishing operations. Accelerating admixtures can be used to decrease the setting time. Calcined shale and clay have little effect on setting time.

**Finishability**
Concrete containing supplementary cementing materials will generally have equal or improved finishability compared to similar concrete mixes without them. Mixes that contain high dosages of cementitious materials—and especially silica fume—can be sticky and difficult to finish.

**Pumpability**
The use of supplementary cementing materials generally aids the pumpability of concrete. Silica fume is the most effective, especially in lean mixtures.

**Plastic Shrinkage Cracking**
Because of its low bleeding characteristics, concrete containing silica fume may exhibit an increase in plastic shrinkage cracking. The problem may be avoided by ensuring that such concrete is protected against drying, both during and after finishing. Other pozzolans and slag generally have little effect on plastic shrinkage cracking. Supplementary cementing materials that significantly increase set time can increase the risk of plastic shrinkage cracking.

**Curing**
The effects of temperature and moisture conditions on setting properties and strength development of concretes containing supplementary cementing materials are similar to the effects on concrete made with only portland cement; however, the curing time may need to be longer for certain materials with slow-early-strength gain.

High dosages of silica fume can make concrete highly cohesive with very little aggregate segregation or bleeding. With little or no bleed water available at the concrete surface for evaporation, plastic cracking can readily develop, especially on hot, windy days if special precautions are not taken.

Proper curing of all concrete, especially concrete containing supplementary cementing materials, should commence immediately after finishing. A seven-day moist cure or membrane cure should be adequate for concretes with normal dosages of most supplementary cementitious materials. As with portland cement concrete, low curing temperatures can reduce early-strength gain (Gebler and Klieber 1986).

**EFFECTS ON HARDENED CONCRETE**

**Strength**
Fly ash, ground granulated blast-furnace slag, calcined clay, metakaolin, calcined shale, and silica fume contribute to the strength gain of concrete. However, the strength of concrete containing these materials can be higher or lower than the strength of concrete using portland cement as the only cementing material. Fig. 3-14 illustrates this for various fly ashes. Tensile, flexural, torsional, and bond strength are affected in the same manner as compressive strength.

![Fig. 3-13. Effect of a slag on heat of hydration at 20°C (68°F) compared to a Type I cement.](image)

![Fig. 3-14. Compressive strength development at 1, 3, 7, 28, and 90 days of concrete mixtures containing 307 kg/m³ (517 lb/yd³) of cementitious materials with a fly ash dosage of 25% of the cementitious materials (Whiting 1989).](image)
Because of the slow pozzolanic reaction of some supplementary cementing materials, continuous wet curing and favorable curing temperatures may need to be provided for longer periods than normally required. However, concrete containing silica fume is less affected by this and generally equals or exceeds the one-day strength of a cement-only control mixture. Silica fume contributes to strength development primarily between 3 and 28 days, during which time a silica fume concrete exceeds the strength of a cement-only control concrete. Silica fume also aids the early strength gain of fly ash-cement concretes.

The strength development of concrete with fly ash, ground slag, calcined clay, or calcined shale, is similar to normal concrete when cured around 23°C (73°F). Fig. 3-15 illustrates that the rate of strength gain of concrete with fly ash, relative to its 28-day strength, is similar to concrete without fly ash. Concretes made with certain highly reactive fly ashes (especially high-calcium Class C ashes) or ground slags can equal or exceed the control strength in 1 to 28 days. Some fly ashes and natural pozzolans require 28 to 90 days to exceed a 28-day control strength, depending on the mixture proportions. Concretes containing Class C ashes generally develop higher early-age strength than concretes with Class F ashes.

Strength gain can be increased by: (1) increasing the amount of cementitious material in the concrete; (2) adding high-early strength cementitious materials; (3) decreasing the water-cementing materials ratio; (4) increasing the curing temperature; or (5) using an accelerating admixture.

Fig. 3-16 illustrates the benefit of using fly ash as an addition instead of a cement replacement to improve strength development in cold weather. Mass concrete design often takes advantage of the delayed strength gain of pozzolans as these structures are often not put into full service immediately. Slow early strength gain resulting from the use of some supplementary cementitious materials is an advantage in hot weather construction as it allows more time to place and finish the concrete. With appropriate mixture adjustments, all supplementary cementitious materials can be used in all seasons.

Supplementary cementing materials are often essential to the production of high-strength concrete. Fly ash, especially, has been used in production of concrete with strengths up to 100 MPa (15,000 psi). With silica fume, ready mix producers now have the ability to make concrete with strengths up to 140 MPa (20,000 psi), when used with high-range water reducers and appropriate aggregates (Burg and Ost 1994).

Impact and Abrasion Resistance
The impact resistance and abrasion resistance of concrete are related to compressive strength and aggregate type. Supplementary cementing materials generally do not affect these properties beyond their influence on strength. Concretes containing fly ash are just as abrasion resistant as portland cement concretes without fly ash (Gebler and Klieger 1986). Fig. 3-17 illustrates that abrasion resistance of fly ash concrete is related to strength.

Freeze-Thaw Resistance
It is imperative for development of resistance to deterioration from cycles of freezing and thawing that a concrete have adequate strength and entrained air. For concrete containing supplementary cementing materials to provide the same resistance to freezing and thawing cycles as a concrete made using only portland cement as a binder, four conditions for both concretes must be met:
4. They must be air-dried for one month prior to exposure to saturated freezing conditions.

Table 3-7 shows equal frost resistance for concrete with and without fly ash. Fig. 3-18 illustrates the long term durability of concretes with fly ash, slag, or shale.

**Deicer-Scaling Resistance**

Decades of field experience have demonstrated that air-entrained concretes containing normal dosages of fly ash, slag, silica fume, calcined clay, or calcined shale are resistant to scaling caused by the application of deicing salts in a freeze-thaw environment. Laboratory tests also indicate that the deicer-scaling resistance of concrete made with supplementary cementing materials is often equal to concrete made without supplementary cementing materials.

Scaling resistance can decrease as the amount of certain supplementary cementing materials increases. However, concretes that are properly designed, placed, and cured, have demonstrated good scaling resistance even when made with high dosages of some of these materials.

Deicer-scaling resistance of all concrete is significantly improved with the use of a low water-cement ratio, a moderate portland cement content, adequate air entrainment, proper finishing and curing, and an air-drying period prior to exposure of the concrete to salts and freezing temperatures. Lean concrete with only about 240 kg/m³ (405 lb/yd³) or less of cementitious material can be especially vulnerable to deicer scaling. A minimum of 335 kg/m³ (564 lb/yd³) of cementitious material and air entrainment is required.

1. They must have the same compressive strength.
2. They must have an adequate entrained air content with proper air-void characteristics.
3. They must be properly cured.

![Fig. 3-17. Comparison of abrasion resistance and compressive strength of various concretes with 25% fly ash. Abrasion resistance increases with strength (Gebler and Klieger 1986).](image)

![Fig. 3-18. View of concrete slabs in PCA outdoor test plot (Skokie, Illinois) containing (A) fly ash, (B) slag, (C) calcined shale, and (D) portland cement after 30 years of deicer and frost exposure. These samples demonstrate the durability of concretes containing various cementitious materials. Source: RX 157, LTS Cement No. 51, slabs with 335 kg/m³ (564 lb/yd³) of cementing material and air entrainment. (IMG12336, IMG12334, IMG12335, IMG12333)](image)
(564 lb/yd$^3$) of cementitious material and a maximum water to cementitious materials ratio of 0.45 is recommended. A satisfactory air-void system is also critical.

The importance of using a low water-cement ratio for scale resistance is demonstrated in Fig. 3-19. The effect of high fly ash dosages and low cementing material contents is demonstrated in Fig. 3-20. The performance of scale-resistant concretes containing fly ash at a dosage of 25% of cementing material by mass is presented in Table 3-7. The table demonstrates that well designed, placed and cured concretes with and without fly ash can be equally resistant to deicer scaling.

The ACI 318 building code states that the maximum dosage of fly ash, slag, and silica fume should be 25%, 50%, and 10% by mass of cementing materials, respectively for deicer exposures. Total supplementary cementing materials should not exceed 50% of the cementitious material. Dosages less than or higher than these limits have been shown to be durable in some cases and not in others. Different materials respond differently in different environments. The selection of materials and dosages should be based on local experience and the durability should be demonstrated by field or laboratory performance.

### Drying Shrinkage and Creep

When used in low to moderate amounts, the effect of fly ash, ground granulated blast-furnace slag, calcined clay, calcined shale, and silica fume on the drying shrinkage and creep of concrete is generally small and of little practical significance. Some studies indicate that silica fume may reduce creep (Burg and Ost 1994).
Alkali-Ag gregate Reactivity

Alkali-silica reactivity can be controlled through the use of certain supplementary cementitious materials. Silica fume, fly ash, ground granulated blast-furnace slag, calcined clay, and other pozzolans have been reported to significantly reduce alkali-silica reactivity (Figs. 3-21 and 3-22). Low calcium Class F ashes have reduced reactivity

Permeability and Absorption

With adequate curing, fly ash, ground slag, and natural pozzolans generally reduce the permeability and absorption of concrete. Silica fume and metakaolin are especially effective in this regard. Silica fume and calcined clay can provide concrete with a chloride resistance of under 1000 coulombs using the ASTM C1202 rapid chloride permeability test (Barger and others 1997). Tests show that the permeability of concrete decreases as the quantity of hydrated cementing materials increases and the water-cementitious materials ratio decreases. The absorption of fly-ash concrete is about the same as concrete without ash, although some ashes can reduce absorption by 20% or more.

Alkali-Ag gregate Reactivity

Alkali-silica reactivity can be controlled through the use of certain supplementary cementitious materials. Silica fume, fly ash, ground granulated blast-furnace slag, calcined clay, calcined shale, and other pozzolans have been reported to significantly reduce alkali-silica reactivity (Figs. 3-21 and 3-22). Low calcium Class F ashes have reduced reactivity

Fig. 3-20. Relationship between deicer-scaling resistance and dosage of fly ash for air-entrained concretes made with moderate to high water-cementitious materials ratios. Replacement of portland cement with fly ash: (left) 25% and (right) 50%. A scale rating of 0 is no scaling and 5 is severe scaling (Whiting 1989).

Fig. 3-21. Effect of different fly ashes and slag on alkali-silica reactivity. Note that some ashes are more effective than others in controlling the reaction and that dosage of the ash or slag is critical. A highly reactive natural aggregate was used in this test. A less reactive aggregate would require less ash or slag to control the reaction. A common limit for evaluating the effectiveness of pozzolans or slags is 0.10% expansion using this rapid mortar bar test (Detwiler 2002).

Fig. 3-22. Reduction of alkali-silica reactivity by calcined clay and calcined shale (Lerch 1950).
expansion up to 70% or more in some cases. At optimum dosage, Class C ashes can also reduce reactivity but to a lesser degree than most Class F ashes. Supplementary cementing materials provide additional calcium silicate hydrate to chemically tie up the alkalis in the concrete (Bhatt 1985, and Bhatt and Greening 1986). Determination of optimum supplementary cementing materials dosage is important to maximize the reduction in reactivity and to avoid dosages and materials that can aggravate reactivity. Dosage rates should be verified by tests, such as ASTM C1567 or ASTM C1293.

Supplementary cementing materials that reduce alkali-silica reactions will not reduce alkali-carbonate reactions, a type of reaction involving cement alkalis and certain dolomitic limestones.

Descriptions of aggregate testing and preventive measures to be taken to prevent deleterious alkali-aggregate reaction are discussed in Farny and Kerkhoff (2007), PCA (1998), and AASHTO (2000).

**Sulfate Resistance**

With proper proportioning and material selection, silica fume, fly ash, calcined shale, and ground slag can improve the resistance of concrete to sulfate or seawater attack. This is done primarily by reducing permeability and by reducing the amount of reactive elements needed for expansive sulfate reactions. For improved sulfate resistance of lean concrete, one study showed that for a particular Class F ash, an adequate amount was approximately 20% of the cement plus fly ash; this illustrates the need to determine optimum ash contents, as higher ash contents were detrimental (Stark 1989).

The sulfate resistance of high-cement-content, low water-cement ratio concrete made with a sulfate resistant cement is so great that fly ash has little opportunity to improve resistance. Concretes with Class F ashes are generally more sulfate resistant than those made with Class C ashes. Some Class C ashes have been shown to reduce sulfate resistance at normal dosage rates.

Ground slag is generally considered beneficial in sulfate environments. However, one long-term study in a very severe environment showed a slight reduction in sulfate resistance in concrete containing ground slag compared to concrete containing only portland cement as the cementing material (Stark 1982 and 1989). One reason for decreased performance with slag in this study is that the mixtures may not have been optimized for sulfate resistance.

Other studies indicate that concrete with ground slag has a sulfate resistance equal to or greater than concrete made with Type V sulfate-resistant portland cement (ACI 233 and Detwiler, Bhatt, and Bhatta 1996). Calcined clay has been demonstrated to provide sulfate resistance greater than high-sulfate resistant Type V cement (Barger and others 1997).

**Corrosion of Embedded Steel**

Some supplementary cementing materials reduce steel corrosion by reducing the permeability of properly cured concrete to water, air, and chloride ions. Fly ash can significantly reduce chloride-ion ingress. Silica fume greatly decreases permeability and chloride-ion ingress and also significantly increases electrical resistivity, thereby reducing the electrochemical reaction of corrosion. Concrete containing silica fume or metakaolin is often used in overlays and full-depth slab placements on bridges and parking garages; these structures are particularly vulnerable to corrosion due to chloride-ion ingress.

**Carbonation**

Carbonation of concrete is a process by which carbon dioxide from the air penetrates the concrete and reacts with the hydroxides, such as calcium hydroxide, to form carbonates. In the reaction with calcium hydroxide, calcium carbonate is formed. Carbonation lowers the alkalinity of concrete. High alkalinity is needed to protect embedded steel from corrosion; consequently, concrete should be resistant to carbonation to help prevent steel corrosion.

The amount of carbonation is significantly increased in concretes with a high water-cementing materials ratio, low cement content, short curing period, low strength, and a highly permeable or porous paste. The depth of carbonation of good quality concrete is generally of little practical significance. At normal dosages, fly ash is reported to slightly increase carbonation, but usually not to a significant amount in concrete with short (normal) moist-curing periods (Campbell, Sturm, and Kosmatka 1991).

**Chemical Resistance**

Supplementary cementing materials often reduce chemical attack by reducing the permeability of concrete. Although many of these materials may improve chemical resistance, they do not make concrete totally immune to attack. Concrete in severe chemical exposure should be protected using barrier systems. Kerkhoff (2007) provides a discussion of methods and materials to protect concrete from aggressive chemicals and exposures.

**Soundness**

Normal dosages of fly ash, slag, silica fume and natural pozzolans do not affect the soundness of concrete. Concrete soundness is protected by soundness requirements, such as autoclave expansion limits, on the materials. Dosages in concrete should not exceed dosages deemed safe in the autoclave test.

**Concrete Color**

Supplementary cementitious materials may slightly alter the color of hardened concrete. Color effects are related to
the color and amount of the material used in concrete. Many supplementary cementing materials resemble the color of portland cement and therefore have little effect on color of the hardened concrete. Some silica fumes may give concrete a slightly bluish or dark gray tint and tan fly ash may impart a tan color to concrete when used in large quantities. Ground slag and metakaolin can make concrete whiter. Ground slag can initially impart a bluish or greenish undertone.

**CONCRETE MIX PROPORTIONS**

The optimum amounts of supplementary cementing materials used with portland cement or blended cement are determined by testing, by the relative cost and availability of the materials, and by the specified properties of the concrete.

Several test mixtures are required to determine the optimum amount of pozzolan or slag. These mixtures should cover a range of blends to establish the relationship between strength and water to cementing materials ratio. These mixtures should be established according to ACI Standard 211.1 or 211.2, taking into account the relative densities of the supplementary cementing materials. These are usually different from the relative density of portland cement. The results of these tests will be a family of strength curves for each age at which the concrete is required to meet certain specified requirements. The dosage of a cementitious material is usually stated as a mass percentage of all the cementitious materials in a concrete mixture.

Typical practice in the United States uses fly ash, slag, silica fume, calcined clay, or calcined shale as an addition to portland cement or as a partial replacement for some of the portland cement. Blended cements, which already contain pozzolans or slag, are designed to be used with or without additional supplementary cementitious materials. Concrete mixtures with more than one supplementary cementitious material are also used. For example, a concrete mixture may contain portland cement, fly ash, and silica fume. Such mixes are called ternary concretes. When fly ash, slag, silica fume, or natural pozzolans are used in combination with portland or blended cement, the proportioned concrete mixture should be tested to demonstrate that it meets the required concrete properties for the project.

**AVAILABILITY**

All supplementary cementitious materials may not be available in all areas. Consult local material suppliers on available materials. Class F fly ashes are usually available in the Eastern United States, while Class C ashes are available in the Midwest and West. Silica fume is available in most locations because only small dosages are used. Calcined clays and shales are available in select areas. Ground granulated slags are available in most regions.

**STORAGE**

In most cases, moisture will not affect the physical performance of supplementary cementing materials. These materials should, however, be kept dry to avoid difficulties in handling and discharge. Class C fly ash and calcined shale must be kept dry as they will set and harden when exposed to moisture. Equipment for handling and storing these materials is similar to that required for cement. Additional modifications may be required where using silica fume, which does not have the same flowing characteristics as other supplementary cementing materials (and may be supplied as a liquid).

These materials are usually kept in bulk storage facilities or silos, although some products are available in bags. Because the materials may resemble portland cement in color and fineness, the storage facilities should be clearly marked to avoid the possibility of misuse and contamination with other materials at the batch plant. All valves and piping should also be clearly marked and properly sealed to avoid leakage and contamination. Fly ash, slag, and natural pozzolans should be weighed after the portland or blended cement in the batching sequence to avoid overdosing in case valves stick.

**REFERENCES**


ACI, *Concrete Durability, Katherine and Bryant Mather International Conference*, SP100, American Concrete Institute, Farmington Hills, Michigan, 1987.


American Concrete Institute, Farmington Hills, Michigan, May-June 1988, pages 148 to 157.


Klieger, Paul, and Perenchio, William F., Laboratory Studies of Blended Cement: Portland-Pozzolan Cements, Research and


CHAPTER 4
Mixing Water for Concrete

Almost any natural water that is drinkable and has no pronounced taste or odor, also called potable water, can be used as mixing water for making concrete (Fig. 4-1). However, some waters that are not fit for drinking may be suitable for use in concrete.

**SOURCES OF MIXING WATER**

Sources of mixing water include: (1) Batch water discharged into the mixer from municipal water supply, municipal reclaimed water, or water from concrete production operations. This is the main but not only source of mixing water in concrete. (2) During hot-weather concreting, ice might by used as part of the mixing water (see Chapter 13). The ice should be completely melted by the time mixing is completed. (3) Water might also be added by the truck operator at the jobsite. ASTM C94 (AASHTO M 157) allows the addition of water on site if the slump of the concrete is less than specified, provided the maximum allowable water-cement ratio is not exceeded and several other conditions are met (see Chapter 10). (4) Free moisture on aggregate can represent a substantial portion of the total mixing water, because aggregates usually contain surface moisture. It is important that the water brought in by the aggregate is free from harmful materials. (5) Water contained in admixtures has to be considered part of the mixing water if the admixture’s water content is sufficient to affect the water-cementitious materials ratio by 0.01 or more.

Six typical analyses of city water supplies and seawater are shown in Table 4-1. These waters approximate the composition of domestic water supplies for most of the

<table>
<thead>
<tr>
<th>Chemicals</th>
<th>Analysis No.</th>
<th>1</th>
<th>2</th>
<th>3</th>
<th>4</th>
<th>5</th>
<th>6</th>
<th>Seawater*</th>
</tr>
</thead>
<tbody>
<tr>
<td>Silica (SiO₂)</td>
<td>2.4</td>
<td>0.0</td>
<td>6.5</td>
<td>9.4</td>
<td>22.0</td>
<td>3.0</td>
<td>—</td>
<td>50 to 480</td>
</tr>
<tr>
<td>Iron (Fe)</td>
<td>0.1</td>
<td>0.0</td>
<td>0.0</td>
<td>0.2</td>
<td>0.1</td>
<td>0.0</td>
<td>—</td>
<td>260 to 1410</td>
</tr>
<tr>
<td>Calcium (Ca)</td>
<td>5.8</td>
<td>15.3</td>
<td>29.5</td>
<td>96.0</td>
<td>3.0</td>
<td>1.3</td>
<td>50 to 480</td>
<td></td>
</tr>
<tr>
<td>Magnesium (Mg)</td>
<td>1.4</td>
<td>5.5</td>
<td>7.6</td>
<td>27.0</td>
<td>2.4</td>
<td>0.3</td>
<td>260 to 1410</td>
<td></td>
</tr>
<tr>
<td>Sodium (Na)</td>
<td>1.7</td>
<td>16.1</td>
<td>2.3</td>
<td>183.0</td>
<td>215.0</td>
<td>1.4</td>
<td>2190 to 12,200</td>
<td></td>
</tr>
<tr>
<td>Potassium (K)</td>
<td>0.7</td>
<td>0.0</td>
<td>1.6</td>
<td>18.0</td>
<td>9.8</td>
<td>0.2</td>
<td>70 to 550</td>
<td></td>
</tr>
<tr>
<td>Bicarbonate (HCO₃⁻)</td>
<td>14.0</td>
<td>35.8</td>
<td>122.0</td>
<td>334.0</td>
<td>549.0</td>
<td>4.1</td>
<td>—</td>
<td></td>
</tr>
<tr>
<td>Sulfate (SO₄²⁻)</td>
<td>9.7</td>
<td>59.9</td>
<td>5.3</td>
<td>121.0</td>
<td>11.0</td>
<td>2.6</td>
<td>580 to 2810</td>
<td></td>
</tr>
<tr>
<td>Chloride (Cl)</td>
<td>2.0</td>
<td>3.0</td>
<td>1.4</td>
<td>280.0</td>
<td>22.0</td>
<td>1.0</td>
<td>3960 to 20,000</td>
<td></td>
</tr>
<tr>
<td>Nitrate (NO₃⁻)</td>
<td>0.5</td>
<td>0.0</td>
<td>0.2</td>
<td>0.5</td>
<td>0.0</td>
<td>—</td>
<td>35,000</td>
<td></td>
</tr>
<tr>
<td>Total dissolved solids</td>
<td>31.0</td>
<td>250.0</td>
<td>125.0</td>
<td>983.0</td>
<td>564.0</td>
<td>19.0</td>
<td>—</td>
<td></td>
</tr>
</tbody>
</table>

* Different seas contain different amounts of dissolved salts.
impurities in the mixing water do not adversely shorten or extend the setting time of the cement. Acceptance criteria for water to be used in concrete are given in ASTM C1602 (see Tables 4-2 and 4-3).

Excessive impurities in mixing water not only may affect setting time and concrete strength, but also may cause efflorescence, staining, corrosion of reinforcement, volume instability, and reduced durability. Therefore, certain optional limits may be set on chlorides, sulfates, alkalies, and solids in the mixing water or appropriate tests can be performed to determine the effect the impurity has on various properties. Some impurities may have little effect on strength and setting time, yet they can adversely affect durability and other properties.

Water containing less than 2000 parts per million (ppm) of total dissolved solids can generally be used satisfactorily for making concrete. Water containing more than 2000 ppm of dissolved solids should be tested for its effect on strength and time of set. Additional information on the effects of mix water impurities can be found in Steinour (1960) and Abrams (1924). Over 100 different compounds and ions are discussed.

Following is a synopsis of the effects of certain impurities in mixing water on the quality of normal concrete:

### ALKALI CARBONATE AND BICARBONATE

Carbonates and bicarbonates of sodium and potassium have different effects on the setting times of different

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**Table 4-2. Acceptance Criteria for Questionable Water Supplies (ASTM C1602)**

<table>
<thead>
<tr>
<th>Limits</th>
<th>Test method</th>
</tr>
</thead>
<tbody>
<tr>
<td>Compressive strength, minimum percentage of control at 7 days</td>
<td>90</td>
</tr>
<tr>
<td>Time of set, deviation from control, hr:min. from 1:00 earlier to 1:30 later</td>
<td>C403</td>
</tr>
</tbody>
</table>

*Comparisons should be based on fixed proportions of a concrete mix design representative of questionable water supply and a control mix using 100% potable water.

**Table 4-3. Chemical Limits for Wash Water used as Mixing Water (ASTM C1602)**

<table>
<thead>
<tr>
<th>Chemical or type of construction</th>
<th>Maximum concentration, ppm</th>
<th>Test method</th>
</tr>
</thead>
<tbody>
<tr>
<td>Chloride, as Cl</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Prestressed concrete or concrete in bridge decks</td>
<td>500**</td>
<td>ASTM C114</td>
</tr>
<tr>
<td>Other reinforced concrete in moist environments or containing aluminum embedments or dissimilar metals or with stay-in-place galvanized metal forms</td>
<td>1,000**</td>
<td>ASTM C114</td>
</tr>
<tr>
<td>Sulfate, as SO₄</td>
<td>3,000</td>
<td>ASTM C114</td>
</tr>
<tr>
<td>Alkalies, as (Na₂O + 0.658 K₂O)</td>
<td>600</td>
<td>ASTM C114</td>
</tr>
<tr>
<td>Total solids by mass</td>
<td>50,000</td>
<td>ASTM C1603</td>
</tr>
</tbody>
</table>

*ppm is the abbreviation for parts per million.

** The requirements for concrete in ACI 318 shall govern when the manufacturer can demonstrate that these limits for mixing water can be exceeded. For conditions allowing the use of calcium chloride (CaCl₂) accelerator as an admixture, the chloride limitation is permitted to be waived by the purchaser.
cements. Sodium carbonate can cause very rapid setting, bicarbonates can either accelerate or retard the set. In large concentrations, these salts can materially reduce concrete strength. When the sum of the dissolved salts exceeds 1000 ppm, tests for their effect on setting time and 28-day strength should be made. The possibility of aggravated alkali-aggregate reactions should also be considered.

**CHLORIDE**

Concern over a high chloride content in mixing water is chiefly due to the possible adverse effect of chloride ions on the corrosion of reinforcing steel or prestressing strands. Chloride ions attack the protective oxide film formed on the steel by the highly alkaline (pH greater than 12.5) chemical environment present in concrete. The acid-soluble chloride ion level at which steel reinforcement corrosion begins in concrete is about 0.2% to 0.4% by mass of cement (0.15% to 0.3% water soluble). Of the total chloride-ion content in concrete, only about 50% to 85% is water soluble; the rest becomes chemically combined in cement reactions (Whiting 1997, Whiting, Taylor, and Nagi 2002, and Taylor, Whiting, and Nagi 2000).

Chlorides can be introduced into concrete with the separate mixture ingredients—admixtures, aggregates, cementitious materials, and mixing water—or through exposure to deicing salts, seawater, or salt-laden air in coastal environments. Placing an acceptable limit on chloride content for any one ingredient, such as mixing water, is difficult considering the several possible sources of chloride ions in concrete. An acceptable limit in the concrete depends primarily upon the type of structure and the environment to which it is exposed during its service life.

A high dissolved solids content of a natural water is sometimes due to a high content of sodium chloride or sodium sulfate. Both can be tolerated in rather large quantities. Concentrations of 20,000 ppm of sodium chloride are generally tolerable in concrete that will be dry in service and has low potential for corrosive reactions. Water used in prestressed concrete or in concrete that is to have aluminum embedments should not contain deleterious amounts of chloride ion. The contribution of chlorides from ingredients other than water should also be considered. Calcium chloride admixtures should be avoided in steel reinforced concrete.

The ACI 318 building code limits water soluble chloride ion content in reinforced concrete to the following percentages by mass of cement:

<table>
<thead>
<tr>
<th>Concrete Type</th>
<th>Chloride Content</th>
</tr>
</thead>
<tbody>
<tr>
<td>Prestressed concrete (Classes C0, C1, C2)</td>
<td>0.06%</td>
</tr>
<tr>
<td>Reinforced concrete exposed to chloride in service (Class C2)</td>
<td>0.15%</td>
</tr>
<tr>
<td>Reinforced concrete that will be dry or protected from moisture in service (Class C0)</td>
<td>1.00%</td>
</tr>
<tr>
<td>Other reinforced concrete construction (Class C1)</td>
<td>0.30%</td>
</tr>
</tbody>
</table>

ACI 318 does not limit the amount of chlorides in plain concrete, that is concrete not containing steel. Additional information on limits and tests can be found in ACI 222, *Corrosion of Metals in Concrete*. The acid-soluble and water-soluble chloride content of concrete can be determined by using ASTM C1152 and C1218.

**SULFATE**

Concern over a high sulfate content in mix water is due to possible expansive reactions and deterioration by sulfate attack, especially in areas where the concrete will be exposed to high sulfate soils or water. Although mixing waters containing 10,000 ppm of sodium sulfate have been used satisfactorily, the limit in Table 4-3 should be considered unless special precautions are taken.

**OTHER COMMON SALTS**

Carbonates of calcium and magnesium are not very soluble in water and are seldom found in sufficient concentration to affect the strength of concrete. Bicarbonates of calcium and magnesium are present in some municipal waters. Concentrations up to 400 ppm of bicarbonate in these forms are not considered harmful.

Magnesium sulfate and magnesium chloride can be present in high concentrations without harmful effects on strength. Good strengths have been obtained using water with concentrations up to 40,000 ppm of magnesium chloride. Concentrations of magnesium sulfate should be less than 25,000 ppm.

**IRON SALTS**

Natural ground waters seldom contain more than 20 to 30 ppm of iron; however, acid mine waters may carry rather large quantities. Iron salts in concentrations up to 40,000 ppm do not usually affect concrete strengths adversely.

**MISCELLANEOUS INORGANIC SALTS**

Salts of manganese, tin, zinc, copper, and lead in mixing water can cause a significant reduction in strength and large variations in setting time. Of these, salts of zinc, copper, and lead are the most active. Salts that are especially active as retarders include sodium iodate, sodium phosphate, sodium arsenate, and sodium borate. All can greatly retard both set and strength development when present in concentrations of a few tenths percent by mass of the cement. Generally, concentrations of these salts up to 500 ppm can be tolerated in mixing water.

Another salt that may be detrimental to concrete is sodium sulfide; even the presence of 100 ppm warrants
testing. Additional information on the effects of other salts can be found in the references.

SEAWATER

Seawater containing up to 35,000 ppm of dissolved salts is generally suitable as mixing water for concrete not containing steel. About 78% of the salt is sodium chloride, and 15% is chloride and sulfate of magnesium. Although concrete made with seawater may have higher early strength than normal concrete, strengths at later ages (after 28 days) may be lower. This strength reduction can be compensated for by reducing the water-cement ratio.

Seawater is not suitable for use in making steel reinforced concrete and it should not be used in prestressed concrete due to the risk of corrosion of the reinforcement, particularly in warm and humid environments. If seawater is used in plain concrete (no steel) in marine applications, moderate sulfate resistant cements, Types II or MS, should be used along with a low water-cement ratio.

Sodium or potassium in salts present in seawater used for mix water can aggravate alkali-aggregate reactivity. Thus, seawater should not be used as mix water for concrete with potentially alkali-reactive aggregates.

Seawater used for mix water also tends to cause efflorescence and dampness on concrete surfaces exposed to air and water (Steinour 1960). Marine-dredged aggregates are discussed in Chapter 5.

ACID WATERS

Acceptance of acid mixing water should be based on the concentration (in parts per million) of acids in the water. Occasionally, acceptance is based on the pH, which is a measure of the hydrogen-ion concentration on a log scale. The pH value is an intensity index and is not the best measure of potential acid or base reactivity. The pH of neutral water is 7.0; values below 7.0 indicate acidity and those above 7.0 alkalinity (a base).

Generally, mixing waters containing hydrochloric, sulfuric, and other common inorganic acids in concentrations as high as 10,000 ppm have no adverse effect on strength. Acid waters with pH values less than 3.0 may create handling problems and should be avoided if possible. Organic acids, such as tannic acid, can have a significant effect on strength at higher concentrations (Fig. 4-3).

Fig. 4-3. Effect of tannic acid on the strength of concrete (Abrams 1920).

ALKALINE WATERS

Waters with sodium hydroxide concentrations of 0.5% by mass of cement do not greatly affect concrete strength provided quick set is not induced. Higher concentrations, however, may reduce concrete strength.

Potassium hydroxide in concentrations up to 1.2% by mass of cement has little effect on the concrete strength developed by some cements, but the same concentration when used with other cements may substantially reduce the 28-day strength.

The possibility for increased alkali-aggregate reactivity should be considered.

WATER FROM CONCRETE PRODUCTION (WASH WATER)

The U.S. Environmental Protection Agency and state agencies forbid discharging into the nation’s waterways untreated water from concrete production operations. This is water recovered from processes of concrete production that includes wash water from mixers or that was a part of a concrete mixture; water collected in a basin as a result of storm water runoff at a concrete production facility; or water that contains quantities of concrete ingredients. In most situations, the recycled water is passed through settling ponds where the solids settle out, leaving clarified water (Fig. 4-4). In some cases, the recycled water from a reclaimer unit is kept agitated to maintain the solids in suspension for reuse as a portion of the batch water in concrete. Solid content in recycled water varies from 2.5%
to 10%. ASTM C1602 and AASHTO M 157 (with purchaser’s approval) permit the use of water from concrete production operations as mixing water in concrete, provided it meets the limits in Table 4-2 and 4-3.

INDUSTRIAL WASTEWATER

Most waters carrying industrial wastes have less than 4000 ppm of total solids. When such water is used as mixing water in concrete, the reduction in compressive strength is generally not greater than about 10%-15%. Wastewaters such as those from tanneries, paint factories, coke plants, and chemical and galvanizing plants may contain harmful impurities. It is best to test any wastewater that contains even a few hundred parts per million of unusual solids.

WATERS CARRYING SANITARY SEWAGE

A typical sewage may contain about 400 ppm of organic matter. After the sewage is diluted in a good disposal system, the concentration is reduced to about 20 ppm or less. This amount is too low to have any significant effect on concrete strength.

ORGANIC IMPURITIES

The effect of organic substances on the setting time of portland cement or the ultimate strength of concrete is a problem of considerable complexity. Such substances, like surface loams, can be found in natural waters. Highly colored waters, waters with a noticeable odor, or those in which green or brown algae are visible should be regarded with suspicion and tested accordingly. Organic impurities are often of a humus nature containing tannates or tannic acid (Fig. 4-3).

SUGAR

Small amounts of sucrose, as little as 0.03% to 0.15% by mass of cement, usually retard the setting of cement. The upper limit of this range varies with different cements. The 7-day strength may be reduced while the 28-day strength may be improved. Sugar in quantities of 0.25% or more by mass of cement may cause rapid setting and a substantial reduction in 28-day strength. Each type of sugar influences setting time and strength differently.

Less than 500 ppm of sugar in mix water generally has no adverse effect on strength, but if the concentration exceeds this amount, tests for setting time and strength should be made.

SILT OR SUSPENDED PARTICLES

About 2000 ppm of suspended clay or fine rock particles can be tolerated in mixing water. Higher amounts might not affect strength but may influence other properties of some concrete mixtures. Before use, muddy water should be passed through settling basins or otherwise clarified to reduce the amount of silt and clay added to the mixture by way of the mix water. When cement fines are returned to the concrete in reused wash water, 50,000 ppm can be tolerated.

OILS

Various kinds of oil are occasionally present in mixing water. Mineral oil (petroleum) not mixed with animal or vegetable oils probably has less effect on strength development than other oils. However, mineral oil in concentrations greater than 2.5% by mass of cement may reduce strength by more than 20%.

ALGAE

Water containing algae is unsuited for making concrete because the algae can cause an excessive reduction in strength. Algae in water leads to lower strengths either by influencing cement hydration or by causing a large amount of air to be entrained in the concrete. Algae may also be present on aggregates, in which case the bond between the aggregate and cement paste is reduced. A maximum algae content of 1000 ppm is recommended.

INTERACTION WITH ADMIXTURES

When evaluating waters for their effect on concrete properties, it is important to also test the water with chemical admixtures that will be used in the job concrete. Certain
compounds in water can influence the performance and efficiency of certain admixtures. For example, the dosage of air-entraining admixture may need to be increased when used with hard waters containing high concentrations of certain compounds or minerals.

REFERENCES


ACI Committee 201, *Guide to Durable Concrete*, ACI 201.2R-01, American Concrete Institute, Farmington Hills, Michigan, 2001, 41 pages.

ACI Committee 222, *Corrosion of Metals in Concrete*, ACI 222R-01, American Concrete Institute, Farmington Hills, Michigan, 2001, 41 pages.


The importance of using the right type and quality of aggregates cannot be overemphasized. The fine and coarse aggregates generally occupy 60% to 75% of the concrete volume (70% to 85% by mass) and strongly influence the concrete’s freshly mixed and hardened properties, mixture proportions, and economy. Fine aggregates (Fig. 5-1) generally consist of natural sand or crushed stone with most particles smaller than 5 mm (0.2 in.). Coarse aggregates (Fig. 5-2) consist of one or a combination of gravels or crushed stone with particles predominantly larger than 5 mm (0.2 in.) and generally between 9.5 mm and 37.5 mm (⅜ in. and 1½ in.). Some natural aggregate deposits, called pit-run gravel, consist of gravel and sand that can be readily used in concrete after minimal processing. Natural gravel and sand are usually dug or dredged from a pit, river, lake, or seabed. Crushed stone is produced by crushing quarry rock, boulders, cobbles, or large-size gravel. Crushed air-cooled blast-furnace slag is also used as fine or coarse aggregate.

The aggregates are usually washed and graded at the pit or plant. Some variation in the type, quality, cleanliness, grading, moisture content, and other properties is expected. Close to half of the coarse aggregates used in portland cement concrete in North America are gravels; most of the remainder are crushed stones.

Naturally occurring concrete aggregates are a mixture of rocks and minerals (see Table 5-1). A mineral is a naturally occurring solid substance with an orderly internal structure and a chemical composition that ranges within narrow limits. Rocks, which are classified as igneous, sedimentary, or metamorphic, depending on origin, are generally composed of several minerals. For example, granite contains quartz, feldspar, mica, and a few other minerals; most limestones consist of calcite, dolomite, and minor amounts of quartz, feldspar, and clay. Weathering and erosion of rocks produce particles of stone, gravel, sand, silt, and clay.

Recycled concrete, or crushed waste concrete, is a feasible source of aggregates and an economic reality, especially where good aggregates are scarce. Conventional stone crushing equipment can be used, and new equipment has been developed to reduce noise and dust.

Aggregates must conform to certain standards for optimum engineering use: they must be clean, hard, strong, durable particles free of absorbed chemicals, coatings of clay, and other fine materials in amounts that could affect hydration and bond of the cement paste. Aggregate particles that are friable or capable of being split are undesirable. Aggregates containing any appreciable amounts of shale or other shaly rocks, soft and porous materials, should be avoided; certain types of chert should be espe-
cially avoided since they have low resistance to weathering and can cause surface defects such as popouts.

Identification of the constituents of an aggregate cannot alone provide a basis for predicting the behavior of aggregates in service. Visual inspection will often disclose weaknesses in coarse aggregates. Service records are invaluable in evaluating aggregates. In the absence of a performance record, the aggregates should be tested before they are used in concrete. The most commonly used aggregates—sand, gravel, crushed stone, and air-cooled blast-furnace slag—produce freshly mixed normal-weight concrete with a density (unit weight) of 2200 to 2400 kg/m³ (140 to 150 lb/ft³). Aggregates of expanded shale, clay, slate, and slag (Fig. 5-3) are used to produce structural lightweight concrete with a freshly mixed density ranging from about 1350 to 1850 kg/m³ (90 to 120 lb/ft³). Other lightweight materials such as pumice, scoria, perlite, vermiculite, and diatomite are used to produce insulating lightweight concretes ranging in density from about 250 to 1450 kg/m³ (15 to 90 lb/ft³). Heavyweight materials such as barite, limonite, magnetite, ilmenite, hematite, iron, and steel punchings or shot are used to produce heavyweight concrete and radiation-shielding concrete (ASTM C637 and C638). Only normal-weight aggregates are discussed in this chapter. See Chapter 18 for special types of aggregates and concretes.

Normal-weight aggregates should meet the requirements of ASTM C33 or AASHTO M 6/M 80. These specifications limit the permissible amounts of deleterious substances and provide requirements for aggregate characteristics. Compliance is determined by using one or more of the several standard tests cited in the following sections and tables. However, the fact that aggregates satisfy ASTM C33 or AASHTO M 6/M 80 requirements does not necessarily assure defect-free concrete.

For adequate consolidation of concrete, the desirable amount of air, water, cement, and fine aggregate (that is, the mortar fraction) should be about 50% to 65% by absolute volume (45% to 60% by mass). Rounded aggregate, such as gravel, requires slightly lower values, while crushed aggregate requires slightly higher values. Fine aggregate content is usually 35% to 45% by mass or volume of the total aggregate content.

For brief descriptions, see “Standard Descriptive Nomenclature of Constituents of Natural Mineral Aggregates” (ASTM C294).

**CHARACTERISTICS OF AGGREGATES**

The important characteristics of aggregates for concrete are listed in Table 5-2 and most are discussed in the following section:

**Grading**

Grading is the particle-size distribution of an aggregate as determined by a sieve analysis (ASTM C136 or AASHTO T 27). The range of particle sizes in aggregate is illustrated in Fig. 5-4. The aggregate particle size is determined by using wire-mesh sieves with square openings. The seven standard ASTM C33 (AASHTO M 6/M 80) sieves for fine
aggregate have openings ranging from 150 µm to 9.5 mm (No. 100 sieve to ⅜ in.). The 13 standard sieves for coarse aggregate have openings ranging from 1.18 mm to 100 mm (0.046 in. to 4 in.). Tolerances for the dimensions of openings in sieves are listed in ASTM E11 (AASHTO M 92).

Size numbers (grading sizes) for coarse aggregates apply to the amounts of aggregate (by mass) in percentages that pass through an assortment of sieves (Fig. 5-5). For highway construction, ASTM D448 (AASHTO M 43) lists the same 13 size numbers as in ASTM C33

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**Table 5.2. Characteristics and Tests of Aggregate**

<table>
<thead>
<tr>
<th>Characteristic</th>
<th>Significance</th>
<th>Test designation*</th>
<th>Requirement or item reported</th>
</tr>
</thead>
<tbody>
<tr>
<td>Resistance to abrasion and degradation</td>
<td>Index of aggregate quality; wear resistance of floors and pavements</td>
<td>ASTM C131 (AASHTO T 96)</td>
<td>Maximum percentage of weight loss. Depth of wear and time</td>
</tr>
<tr>
<td></td>
<td></td>
<td>ASTM C535</td>
<td></td>
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<tr>
<td></td>
<td></td>
<td>ASTM C779</td>
<td></td>
</tr>
<tr>
<td>Resistance to freezing and thawing</td>
<td>Surface scaling, roughness, loss of section, and aesthetics</td>
<td>ASTM C666 (AASHTO T 161)</td>
<td>Maximum number of cycles or period of frost immunity; durability factor</td>
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<td></td>
<td></td>
<td>ASTM C682</td>
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<td></td>
<td></td>
<td>AASHTO T 103</td>
<td></td>
</tr>
<tr>
<td>Resistance to disintegration by sulfates</td>
<td>Soundness against weathering action</td>
<td>ASTM C88 (AASHTO T 104)</td>
<td>Weight loss, particles exhibiting distress</td>
</tr>
<tr>
<td>Particle shape and surface texture</td>
<td>Workability of fresh concrete</td>
<td>ASTM C295 (AASHTO T 11)</td>
<td>Maximum percentage of flat and elongated particles</td>
</tr>
<tr>
<td></td>
<td></td>
<td>ASTM D3398</td>
<td></td>
</tr>
<tr>
<td>Grading</td>
<td>Workability of fresh concrete; economy</td>
<td>ASTM C117 (AASHTO T 27)</td>
<td>Minimum and maximum percentage passing standard sieves</td>
</tr>
<tr>
<td></td>
<td></td>
<td>ASTM C136 (AASHTO T 11)</td>
<td></td>
</tr>
<tr>
<td>Fine aggregate degradation</td>
<td>Index of aggregate quality; Resistance to degradation during mixing</td>
<td>ASTM C1137 (AASHTO T 11)</td>
<td>Change in grading</td>
</tr>
<tr>
<td>Uncompacted void content of fine aggregate</td>
<td>Workability of fresh concrete</td>
<td>ASTM C1252 (AASHTO T 304)</td>
<td>Uncompacted voids and specific gravity values</td>
</tr>
<tr>
<td>Bulk density (unit weight)</td>
<td>Mix design calculations; classification</td>
<td>ASTM C29 (AASHTO T 19)</td>
<td>Compact weight and loose weight</td>
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<tr>
<td>Relative density (specific gravity)</td>
<td>Mix design calculations</td>
<td>ASTM C127 (AASHTO T 85)</td>
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<td></td>
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<td>ASTM C128 (AASHTO T 84)</td>
<td>—</td>
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<tr>
<td>initiator</td>
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<td>ASTM C127 (AASHTO T 22)</td>
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<tr>
<td>Absorption and surface moisture</td>
<td>Control of concrete quality (water-cement ratio)</td>
<td>ASTM C70 (AASHTO T 85)</td>
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<td></td>
<td></td>
<td>ASTM C127 (AASHTO T 84)</td>
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<td>ASTM C566 (AASHTO T 255)</td>
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<tr>
<td>Compressive and flexural strength</td>
<td>Acceptability of fine aggregate failing other tests</td>
<td>ASTM C39 (AASHTO T 97)</td>
<td>Strength to exceed 95% of strength achieved with purified sand</td>
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<td></td>
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<td>ASTM C78 (AASHTO T 22)</td>
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<tr>
<td>Definitions of constituents</td>
<td>Clear understanding and communication</td>
<td>ASTM C125 (AASHTO T 96)</td>
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<td></td>
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<td>ASTM C294 (AASHTO T 96)</td>
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<tr>
<td>Aggregate constituents</td>
<td>Determine amount of deleterious and organic materials</td>
<td>ASTM C40 (AASHTO T 21)</td>
<td>Maximum percentage allowed of individual constituents</td>
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<td></td>
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<td>ASTM C87 (AASHTO T 21)</td>
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<td>ASTM C117 (AASHTO T 71)</td>
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<td>ASTM C123 (AASHTO T 11)</td>
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<td>ASTM C142 (AASHTO T 113)</td>
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<td></td>
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<td>ASTM C295 (AASHTO T 112)</td>
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<tr>
<td>Resistance to alkali reactivity and volume change</td>
<td>Soundness against volume change</td>
<td>ASTM C227 (AASHTO T 11)</td>
<td>Maximum length change, constituents and amount of silica, and alkalinity</td>
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<tr>
<td></td>
<td></td>
<td>ASTM C289 (AASHTO T 11)</td>
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<td>ASTM C295 (AASHTO T 11)</td>
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<td>ASTM C342 (AASHTO T 303)</td>
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<td>ASTM C586 (AASHTO T 303)</td>
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<td></td>
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<td>ASTM C1260 (AASHTO T 303)</td>
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<td></td>
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<td>ASTM C1293 (AASHTO T 303)</td>
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<td></td>
<td></td>
<td>ASTM C1567 (AASHTO T 303)</td>
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</tr>
</tbody>
</table>

* The majority of the tests and characteristics listed are referenced in ASTM C33 (AASHTO M 6/M 80). ACI 221R-96 presents additional test methods and properties of concrete influenced by aggregate characteristics.
aggregate proportions as well as cement and water requirements, workability, pumpability, economy, porosity, shrinkage, and durability of concrete. Variations in grading can seriously affect the uniformity of concrete from batch to batch. Very fine sands are often uneconomical; very coarse sands and coarse aggregate can produce harsh, unworkable mixtures. In general, aggregates that do not have a large deficiency or excess of any size and give a smooth grading curve will produce the most satisfactory results.

The effect of a collection of various sizes in reducing the total volume of voids between aggregates is illustrated by the simple method shown in Fig. 5-7. The beaker on the left is filled with large aggregate particles of uniform size aggregate proportions as well as cement and water requirements, workability, pumpability, economy, porosity, shrinkage, and durability of concrete. Variations in grading can seriously affect the uniformity of concrete from batch to batch. Very fine sands are often uneconomical; very coarse sands and coarse aggregate can produce harsh, unworkable mixtures. In general, aggregates that do not have a large deficiency or excess of any size and give a smooth grading curve will produce the most satisfactory results.

The effect of a collection of various sizes in reducing the total volume of voids between aggregates is illustrated by the simple method shown in Fig. 5-7. The beaker on the left is filled with large aggregate particles of uniform size
and shape; the middle beaker is filled with an equal volume of small aggregate particles of uniform size and shape; and the beaker on the right is filled with particles of both sizes. Below each beaker is a graduate with the amount of water required to fill the voids in that beaker. Note that when the beakers are filled with one particle size of equal volume, the void content is constant, regardless of the particle size. When the two aggregate sizes are combined, the void content is decreased. If this operation were repeated with several additional sizes, a further reduction in voids would occur. The cement paste requirement for concrete is related to the void content of the combined aggregates.

During the early years of concrete technology it was sometimes assumed that the smallest percentage of voids (greatest density of aggregates) was the most suitable for concrete. At the same time, limits were placed on the amount and size of the smallest particles. It is now known that, even on this restricted basis, this is not the best target for the mix designer. However, production of satisfactory, economical concrete requires aggregates of low void content, but not the lowest. Voids in aggregates can be tested according to ASTM C29 or AASHTO T 19.

In reality, the amount of cement paste required in concrete is greater than the volume of voids between the aggregates. This is illustrated in Fig. 5-8. Sketch A represents large aggregates alone, with all particles in contact. Sketch B represents the dispersal of aggregates in a matrix of paste. The amount of paste is necessarily greater than the void content of sketch A in order to provide workability to the concrete; the actual amount is influenced by the workability and cohesiveness of the paste.

**Fine-Aggregate Grading**

Requirements of ASTM C33 or AASHTO M 6/M 43 permit a relatively wide range in fine-aggregate gradation, but specifications by other organizations are sometimes more restrictive. The most desirable fine-aggregate grading depends on the type of work, the richness of the mixture, and the maximum size of coarse aggregate. In leaner mixtures, or when small-size coarse aggregates are used, a grading that approaches the maximum recommended percentage passing each sieve is desirable for workability. In general, if the water-cement ratio is kept constant and the ratio of fine-to-coarse aggregate is chosen correctly, a wide range in grading can be used without measurable effect on strength. However, the best economy will sometimes be achieved by adjusting the concrete mixture to suit the gradation of the local aggregates.

Fine-aggregate grading within the limits of ASTM C33 (AASHTO M 6) is generally satisfactory for most concretes. The ASTM C33 (AASHTO M 6) limits with respect to sieve size are shown in Table 5-3.

**Table 5-3. Fine-Aggregate Grading Limits (ASTM C33/AASHTO M 6)**

<table>
<thead>
<tr>
<th>Sieve size</th>
<th>Percent passing by mass</th>
</tr>
</thead>
<tbody>
<tr>
<td>9.5 mm (% in.)</td>
<td>100</td>
</tr>
<tr>
<td>4.75 mm (No. 4)</td>
<td>95 to 100</td>
</tr>
<tr>
<td>2.36 mm (No. 8)</td>
<td>80 to 100</td>
</tr>
<tr>
<td>1.18 mm (No. 16)</td>
<td>50 to 85</td>
</tr>
<tr>
<td>600 µm (No. 30)</td>
<td>25 to 60</td>
</tr>
<tr>
<td>300 µm (No. 50)</td>
<td>5 to 30 (AASHTO 10 to 30)</td>
</tr>
<tr>
<td>150 µm (No. 100)</td>
<td>0 to 10 (AASHTO 2 to 10)</td>
</tr>
</tbody>
</table>

The AASHTO specifications permit the minimum percentages (by mass) of material passing the 300 µm (No. 50) and 150 µm (No. 100) sieves to be reduced to 5% and 0% respectively, provided:

1. The aggregate is used in air-entrained concrete containing more than 237 kilograms of cement per cubic meter (400 lb of cement per cubic yard) and having an air content of more than 3%.
2. The aggregate is used in concrete containing more than 297 kilograms of cement per cubic meter (500 lb of cement per cubic yard) when the concrete is not air-entrained.
3. An approved supplementary cementitious material is used to supply the deficiency in material passing these two sieves.

Other requirements of ASTM C33 (AASHTO M 6) are:

1. The fine aggregate must not have more than 45% retained between any two consecutive standard sieves.
2. The fineness modulus must be not less than 2.3 nor more than 3.1, nor vary more than 0.2 from the typical value of the aggregate source. If this value is ex-

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![Fig. 5-8. Illustration of the dispersion of aggregates in cohesive concrete mixtures.](image-url)
The amounts of fine aggregate passing the 300 μm (No. 50) and 150, μm (No. 100) sieves affect workability, surface texture, air content, and bleeding of concrete. Most specifications allow 5% to 30% to pass the 300 μm (No. 50) sieve. The lower limit may be sufficient for easy placing conditions or where concrete is mechanically finished, such as in pavements. However, for hand-finished concrete floors, or where a smooth surface texture is desired, fine aggregate with at least 15% passing the 300 μm (No. 50) sieve and 3% or more passing the 150 μm (No. 100) sieve should be used.

### Fineness Modulus

The fineness modulus (FM) of either fine or coarse aggregate according to ASTM C125 is calculated by adding the cumulative percentages by mass retained on each of a specified series of sieves and dividing the sum by 100. The specified sieves for determining FM are: 150 μm (No. 100), 300 μm (No. 50), 600 μm (No. 30), 1.18 mm (No. 16), 2.36 mm (No. 8), 4.75 mm (No. 4), 9.5 mm (% in.), 19.0 mm (¾ in.), 37.5 mm (½ in.), 75 mm (3 in.) and, 150 mm (6 in.).

FM is an index of the fineness of an aggregate—the higher the FM, the coarser the aggregate. Different aggregate grading may have the same FM. FM of fine aggregate is useful in estimating proportions of fine and coarse aggregates in concrete mixtures. An example of how the FM of a fine aggregate is determined (with an assumed sieve analysis) is shown in Table 5-4.

Degradation of fine aggregate due to friction and abrasion will decrease the FM and increase the amount of materials finer than the 75 μm (No. 200) sieve.

### Coarse-Aggregate Grading

The coarse aggregate grading requirements of ASTM C33 (AASHTO M 80) permit a wide range in grading and a variety of grading sizes (see Table 5-5). The grading for a given maximum-size coarse aggregate can be varied over a moderate range without appreciable effect on cement and water requirement of a mixture if the proportion of fine aggregate to total aggregate produces concrete of good workability. Mixture proportions should be changed to produce workable concrete if wide variations occur in the coarse-aggregate grading. Since variations are difficult to anticipate, it is often more economical to maintain uniformity in manufacturing and handling coarse aggregate than to reduce variations in gradation.

The maximum size of coarse aggregate used in concrete has a bearing on the economy of concrete. Usually more water and cement is required for small-size aggregates than for large sizes, due to an increase in total aggregate surface area. The water and cement required for a slump of approximately 75 mm (3 in.) is shown in Fig. 5-9 for a wide range of coarse-aggregate sizes. Fig. 5-9 shows that, for a given water-cement ratio, the amount of cement required decreases as the maximum size of coarse aggregate increases. The increased cost of obtaining and/or

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### Table 5-4. Determination of Fineness Modulus of Fine Aggregates

<table>
<thead>
<tr>
<th>Sieve size</th>
<th>Percentage of individual fraction retained, by mass</th>
<th>Percentage passing, by mass</th>
<th>Cumulative percentage retained, by mass</th>
</tr>
</thead>
<tbody>
<tr>
<td>9.5 mm (% in.)</td>
<td>0</td>
<td>100</td>
<td>0</td>
</tr>
<tr>
<td>4.75 mm (No. 4)</td>
<td>2</td>
<td>98</td>
<td>2</td>
</tr>
<tr>
<td>2.36 mm (No. 8)</td>
<td>13</td>
<td>85</td>
<td>15</td>
</tr>
<tr>
<td>1.18 mm (No. 16)</td>
<td>20</td>
<td>65</td>
<td>35</td>
</tr>
<tr>
<td>600 μm (No. 30)</td>
<td>20</td>
<td>45</td>
<td>55</td>
</tr>
<tr>
<td>300 μm (No. 50)</td>
<td>24</td>
<td>21</td>
<td>79</td>
</tr>
<tr>
<td>150 μm (No. 100)</td>
<td>18</td>
<td>3</td>
<td>97</td>
</tr>
<tr>
<td>Pan</td>
<td>3</td>
<td>0</td>
<td>—</td>
</tr>
<tr>
<td>Total</td>
<td>100</td>
<td>283</td>
<td>283</td>
</tr>
</tbody>
</table>

Fineness modulus = 283 ÷ 100 = 2.83
handling aggregates much larger than 50 mm (2 in.) may offset the savings in using less cement. Furthermore, aggregates of different maximum sizes may give slightly different concrete strengths for the same water-cement ratio. In some instances, at the same water-cement ratio, concrete with a smaller maximum-size aggregate could have higher compressive strength. This is especially true for high-strength concrete. The optimum maximum size of coarse aggregate for higher strength depends on factors such as relative strength of the cement paste, cement-aggregate bond, and strength of the aggregate particles.

The terminology used to specify size of coarse aggregate must be chosen carefully. Particle size is determined by size of sieve and applies to the aggregate passing that sieve and not passing the next smaller sieve. When speaking of an assortment of particle sizes, the size number (or grading size) of the gradation is used. The size number applies to the collective amount of aggregate that passes through an assortment of sieves. As shown in Table 5-5, the amount of aggregate passing the respective sieves is given in percentages; it is called a sieve analysis.

Because of past usage, there is sometimes confusion about what is meant by the maximum size of aggregate. ASTM C125 and ACI 116 define this term and distinguish it from nominal maximum size of aggregate. The maximum size of an aggregate is the smallest sieve that all of a particular aggregate must pass through. The nominal maximum size of an aggregate is the smallest sieve size through which the major portion of the aggregate must pass. The nominal maximum-size sieve may retain 5% to 15% of the aggregate depending on the size number. For example, aggregate size number 67 has a maximum size of 25 mm (1 in.) and a nominal maximum size of 19 mm (% in.). Ninety to one hundred percent of this aggregate must pass the 19-mm (%-in.) sieve and all of the particles must pass the 25-mm (1-in.) sieve.

The maximum size of aggregate that can be used generally depends on the size and shape of the concrete member and the amount and distribution of reinforcing steel. The maximum size of aggregate particles generally should not exceed:

<table>
<thead>
<tr>
<th>Diameter (in)</th>
<th>3/8</th>
<th>9/16</th>
<th>1/2</th>
<th>3/4</th>
<th>1</th>
<th>1 1/2</th>
<th>2</th>
<th>3 1/2</th>
</tr>
</thead>
<tbody>
<tr>
<td>37.5 mm</td>
<td>300</td>
<td>200</td>
<td>100</td>
<td>55</td>
<td>35</td>
<td>25</td>
<td>19.0</td>
<td>12.5</td>
</tr>
<tr>
<td>25.0 mm</td>
<td>250</td>
<td>200</td>
<td>100</td>
<td>55</td>
<td>35</td>
<td>25</td>
<td>19.0</td>
<td>12.5</td>
</tr>
<tr>
<td>19.0 mm</td>
<td>37.5</td>
<td>37.5</td>
<td>37.5</td>
<td>37.5</td>
<td>37.5</td>
<td>37.5</td>
<td>37.5</td>
<td>37.5</td>
</tr>
<tr>
<td>12.5 mm</td>
<td>25</td>
<td>25</td>
<td>25</td>
<td>25</td>
<td>25</td>
<td>25</td>
<td>25</td>
<td>25</td>
</tr>
<tr>
<td>9.5 mm</td>
<td>50</td>
<td>50</td>
<td>50</td>
<td>50</td>
<td>50</td>
<td>50</td>
<td>50</td>
<td>50</td>
</tr>
<tr>
<td>4.75 mm</td>
<td>200</td>
<td>200</td>
<td>200</td>
<td>200</td>
<td>200</td>
<td>200</td>
<td>200</td>
<td>200</td>
</tr>
<tr>
<td>3.7 mm</td>
<td>400</td>
<td>400</td>
<td>400</td>
<td>400</td>
<td>400</td>
<td>400</td>
<td>400</td>
<td>400</td>
</tr>
<tr>
<td>3.2 mm</td>
<td>400</td>
<td>400</td>
<td>400</td>
<td>400</td>
<td>400</td>
<td>400</td>
<td>400</td>
<td>400</td>
</tr>
<tr>
<td>2.36 mm</td>
<td>300</td>
<td>300</td>
<td>300</td>
<td>300</td>
<td>300</td>
<td>300</td>
<td>300</td>
<td>300</td>
</tr>
<tr>
<td>1.18 mm</td>
<td>200</td>
<td>200</td>
<td>200</td>
<td>200</td>
<td>200</td>
<td>200</td>
<td>200</td>
<td>200</td>
</tr>
</tbody>
</table>

Fig. 5-9. Cement and water contents in relation to maximum size of aggregate for air-entrained and non-air-entrained concrete. Less cement and water are required in mixtures having large coarse aggregate (Bureau of Reclamation 1981).
1. One-fifth the narrowest dimension of a concrete member
2. Three-quarters the clear spacing between reinforcing bars and between the reinforcing bars and forms
3. One-third the depth of slabs

These requirements may be waived if, in the judgment of the engineer, the mixture possesses sufficient workability that the concrete can be properly placed without honeycomb or voids.

Combined Aggregate Grading

Aggregate is sometimes analyzed using the combined grading of fine and coarse aggregate together, as they exist in a concrete mixture. This provides a more thorough analysis of how the aggregates will perform in concrete. Sometimes mid-sized aggregate, around the 9.5 mm (% in.) size, is lacking in an aggregate supply, resulting in a concrete with high shrinkage properties, high water demand, poor workability, poor pumpability, and poor placeability. Strength and durability may also be affected.

Fig. 5-10 illustrates an ideal gradation; however, a perfect gradation does not exist in the field—but we can try to approach it. If problems develop due to a poor gradation, alternative aggregates, blending, or special screening of existing aggregates, should be considered. Refer to Shilstone (1990) for options on obtaining optimal grading of aggregate.

The combined gradation can be used to better control workability, pumpability, shrinkage, and other properties of concrete. Abrams (1918) and Shilstone (1990) demonstrate the benefits of a combined aggregate analysis:

- With constant cement content and constant consistency, there is an optimum for every combination of aggregates that will produce the most effective water to cement ratio and highest strength.

- The optimum mixture has the least particle interference and responds best to a high frequency, high amplitude vibrator.

However, this optimum mixture cannot be used for all construction due to variations in placing and finishing needs and availability. Crouch (2000) found in his studies on air-entrained concrete that the water-cement ratio could be reduced by over 8% using combined aggregategradation. Shilstone (1990) also analyzes aggregate gradation by coarseness and workability factors to improve aggregate gradation.

Gap-Graded Aggregates

In gap-graded aggregates certain particle sizes are intentionally omitted. For cast-in-place concrete, typical gap-graded aggregates consist of only one size of coarse aggregate with all the particles of fine aggregate able to pass through the voids in the compacted coarse aggregate. Gap-graded mixes are used in architectural concrete to obtain uniform textures in exposed-aggregate finishes. They can also be used in normal structural concrete because of possible improvements in some concrete properties, and to permit the use of local aggregate gradations (Houston 1962 and Litvin and Pfeifer 1965).

For an aggregate of 19-mm (% in.) maximum size, the 4.75 mm to 9.5 mm (No. 4 to % in.) particles can be omitted without making the concrete unduly harsh or subject to segregation. In the case of 37.5 mm (1½ in.) aggregate, usually the 4.75 mm to 19 mm (No. 4 to ¾ in.) sizes are omitted.

Care must be taken in choosing the percentage of fine aggregate in a gap-graded mixture. A wrong choice can result in concrete that is likely to segregate or honeycomb because of an excess of coarse aggregate. Also, concrete with an excess of fine aggregate could have a high water demand resulting in a low-density concrete. Fine aggregate is usually 25% to 35% by volume of the total aggregate. The lower percentage is used with rounded aggregates and the higher with crushed material. For a smooth off-the-form finish, a somewhat higher percentage of fine aggregate to total aggregate may be used than for an exposed-aggregate finish, but both use a lower fine aggregate content than continuously graded mixtures. Fine aggregate content also depends upon cement content, type of aggregate, and workability.

Air entrainment is usually required for workability since low-slump, gap-graded mixes use a low fine aggregate percentage and produce harsh mixes without entrained air.

Segregation of gap-graded mixes must be prevented by restricting the slump to the lowest value consistent with good consolidation. This may vary from zero to 75 mm (to 3 in.) depending on the thickness of the section, amount of reinforcement, and height of casting. Close control of grading and water content is also required because variations might cause segregation. If a stiff mixture is required, gap-
graded aggregates may produce higher strengths than normal aggregates used with comparable cement contents. Because of their low fine-aggregate volumes and low water-cement ratios, gap-graded mixtures might be considered unworkable for some cast-in-place construction. When properly proportioned, however, these concretes are readily consolidated with vibration.

**Particle Shape and Surface Texture**

The particle shape and surface texture of an aggregate influence the properties of freshly mixed concrete more than the properties of hardened concrete. Rough-textured, angular, elongated particles require more water to produce workable concrete than do smooth, rounded, compact aggregates. Hence, aggregate particles that are angular require more cement to maintain the same water-cement ratio. However, with satisfactory gradation, both crushed and noncrushed aggregates (of the same rock types) generally give essentially the same strength for the same cement factor. Angular or poorly graded aggregates can also be more difficult to pump.

The bond between cement paste and a given aggregate generally increases as particles change from smooth and rounded to rough and angular. This increase in bond is a consideration in selecting aggregates for concrete where flexural strength is important or where high compressive strength is needed.

Void contents of compacted fine or coarse aggregate can be used as an index of differences in the shape and texture of aggregates of the same grading. The mixing water and cement requirement tend to increase as aggregate void content increases. Voids between aggregate particles increase with aggregate angularity.

Aggregates should be relatively free of flat and elongated particles. A particle is called flat and elongated when the ratio of length to thickness exceeds a specified value. See ASTM D4791 for determination of flat, and/or elongated particles. ASTM D3398 provides an indirect method of establishing a particle index as an overall measure of particle shape or texture, while ASTM C295 provides procedures for the petrographic examination of aggregate.

Flat and elongated aggregate particles should be avoided or at least limited to about 15% by mass of the total aggregate. This requirement is equally important for coarse and for crushed fine aggregate, since fine aggregate made by crushing stone often contains flat and elongated particles. Such aggregate particles require an increase in mixing water and thus may affect the strength of concrete, particularly in flexure, if the water-cement ratio is not adjusted.

A number of automated test machines are available for rapid determination of the particle size distribution of aggregate. Designed to provide a faster alternative to the standard sieve analysis test, these machines capture and analyze digital images of the aggregate particles to determine gradation. Fig. 5-11 shows a videograder that measures size and shape of an aggregate by using line-scan cameras wherein two-dimensional images are constructed from a series of line images. Other machines use matrix-scan cameras to capture two-dimensional snapshots of the falling aggregate. Maerz and Lusher (2001) developed a dynamic prototype imaging system that provides particle size and shape information by using a miniconveyor system to parade individual fragments past two orthogonally oriented, synchronized cameras.

**Bulk Density (Unit Weight) and Voids**

The bulk density or unit weight of an aggregate is the mass or weight of the aggregate required to fill a container of a specified unit volume. The volume referred to here is that occupied by both aggregates and the voids between aggregate particles.

The approximate bulk density of aggregate commonly used in normal-weight concrete ranges from about 1200 to 1750 kg/m³ (75 to 110 lb/ft³). The void content between particles affects paste requirements in mix design (see preceding sections, “Particle Shape and Surface Texture” and “Grading”). Void contents range from about 30% to 45% for coarse aggregates to about 40% to 50% for fine aggregate. Angularity increases void content while larger sizes of well-graded aggregate and improved grading decreases void content (Fig. 5-7). Methods of determining the bulk density of aggregates and void content are given in ASTM C29 (AASHTO T 19). In these standards, three methods are described for consolidating the aggregate in the container depending on the maximum size of the aggregate: rodgging, jiggging, and shoveling. The measurement of loose uncompacted void content of fine aggregate is described in ASTM C1252.

**Relative Density (Specific Gravity)**

The relative density (specific gravity) of an aggregate is the ratio of its mass to the mass of an equal absolute volume of
water. It is used in certain computations for mixture proportioning and control, such as the volume occupied by the aggregate in the absolute volume method of mix design. It is not generally used as a measure of aggregate quality, though some porous aggregates that exhibit accelerated freeze-thaw deterioration do have low specific gravities. Most natural aggregates have relative densities between 2.4 and 2.9 with corresponding particle (mass) densities of 2400 and 2900 kg/m³ (150 and 181 lb/ft³).

Test methods for determining relative densities for coarse and fine aggregates are described in ASTM C127 (AASHTO T 85) and ASTM C128 (AASHTO T 84), respectively. The relative density of an aggregate may be determined on an oven-dry basis or a saturated surface-dry (SSD) basis. Both the oven-dry and saturated surface-dry relative densities may be used in concrete mixture proportioning calculations. Oven-dry aggregates do not contain any absorbed or free water. They are dried in an oven to constant weight. Saturated surface-dry aggregates are those in which the pores in each aggregate particle are filled with water but there is no excess water on the particle surface.

Density

The density of aggregate particles used in mixture proportioning computations (not including voids between particles) is determined by multiplying the relative density (specific gravity) of the aggregate times the density of water. An approximate value of 1000 kg/m³ (62.4 lb/ft³) is often used for the density of water. The density of aggregate, along with more accurate values for water density, are provided in ASTM C127 (AASHTO T 85) and ASTM C128 (AASHTO T 84). Most natural aggregates have particle densities of between 2400 and 2900 kg/m³ (150 and 181 lb/ft³).

Absorption and Surface Moisture

The absorption and surface moisture of aggregates should be determined according to ASTM C70, C127, C128, and C566 (AASHTO T 255) so that the total water content of the concrete can be controlled and correct batch weights determined. The internal structure of an aggregate particle is made up of solid matter and voids that may or may not contain water.

The moisture conditions of aggregates are shown in Fig. 5-12. They are designated as:

1. Oven-dry—fully absorbent
2. Air dry—dry at the particle surface but containing some interior moisture, thus still somewhat absorbent
3. Saturated surface dry (SSD)—neither absorbing water from nor contributing water to the concrete mixture
4. Damp or wet—containing an excess of moisture on the surface (free water)

The amount of water added at the concrete batch plant must be adjusted for the moisture conditions of the aggregates in order to accurately meet the water requirement of the mix design. If the water content of the concrete mixture is not kept constant, the water-cement ratio will vary from batch to batch causing other properties, such as the compressive strength and workability to vary from batch to batch.

Coarse and fine aggregate will generally have absorption levels (moisture contents at SSD) in the range of 0.2% to 4% and 0.2% to 2%, respectively. Free-water contents will usually range from 0.5% to 2% for coarse aggregate and 2% to 6% for fine aggregate. The maximum water content of drained coarse aggregate is usually less than that of fine aggregate. Most fine aggregates can maintain a maximum drained moisture content of about 3% to 8% whereas coarse aggregates can maintain only about 1% to 6%.

**Bulking.** Bulking is the increase in total volume of moist fine aggregate over the same mass dry. Surface tension in the moisture holds the particles apart, causing an increase in volume. Bulking of a fine aggregate (such as sand) occurs when it is shoveled or otherwise moved in a damp condition, even though it may have been fully consoli-

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**Fig. 5-12. Moisture conditions of aggregate.**

**Fig. 5-13. Surface moisture on fine aggregate can cause considerable bulking; the amount varies with the amount of moisture and the aggregate grading (PCA Major Series 172 and PCA ST20).**
dated beforehand. Fig. 5-13 illustrates how the amount of bulking of fine aggregate varies with moisture content and grading; fine gradings bulk more than coarse gradings for a given amount of moisture. Fig. 5-14 shows similar information in terms of weight for a particular fine aggregate. Since most fine aggregates are delivered in a damp condition, wide variations can occur in batch quantities if batching is done by volume. For this reason, good practice has long favored weighing the aggregate and adjusting for moisture content when proportioning concrete.

**Resistance to Freezing and Thawing**

The frost resistance of an aggregate, an important characteristic for exterior concrete, is related to its porosity, absorption, permeability, and pore structure. An aggregate particle may absorb so much water (to critical saturation) that it cannot accommodate the expansion and hydraulic pressure that occurs during the freezing of water. If enough of the offending particles are present, the result can be expansion of the aggregate and possible disintegration of the concrete. If a single problem particle is near the surface of the concrete, it can cause a popout. Popouts generally appear as conical fragments that break out of the concrete surface. The offending aggregate particle or a part of it is usually found at the bottom of the void. Generally it is coarse rather than fine aggregate particles with higher porosity values and medium-sized pores (0.1 to 5 µm) that are easily saturated and cause concrete deterioration and popouts. Larger pores do not usually become saturated or cause concrete distress, and water in very fine pores may not freeze readily.

At any freezing rate, there may be a critical particle size above which a particle will fail if frozen when critically saturated. This critical size is dependent upon the rate of freezing and the porosity, permeability, and tensile strength of the particle. For fine-grained aggregates with low permeability (cherts for example), the critical particle size may be within the range of normal aggregate sizes. It is higher for coarse-grained materials or those with capillary systems interrupted by numerous macro pores (voids too large to hold moisture by capillary action). For these aggregates the critical particle size may be sufficiently large to be of no consequence, even though the absorption may be high. If potentially vulnerable aggregates are used in concrete subjected to periodic drying while in service, they may never become sufficiently saturated to cause failure.

Cracking of concrete pavements caused by the freeze-thaw deterioration of the aggregate within concrete is called D-cracking. This type of cracking has been observed in some pavements after three or more years of service. D-cracked concrete resembles frost-damaged concrete caused by paste deterioration. D-cracks are closely spaced crack formations parallel to transverse and longitudinal joints that later multiply outward from the joints toward the center of the pavement panel (Fig. 5-15). D-cracking is a function of the pore properties of certain types of aggregate particles and the environment in which the pavement is placed. Due to the natural accumulation of water under pavements in the base and subbase layers, the aggregate may eventually become saturated. Then with freezing and

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**Fig. 5-14.** Bulk density is compared with the moisture content for a particular sand (PCA Major Series 172).

**Fig. 5-15.** D-cracking along a transverse joint caused by failure of carbonate coarse aggregate (Stark 1976). (IMG12314)
thawing cycles, cracking of the concrete starts in the saturated aggregate (Fig. 5-16) at the bottom of the slab and progresses upward until it reaches the wearing surface. This problem can be reduced either by selecting aggregates that perform better in freeze-thaw cycles or, where marginal aggregates must be used, by reducing the maximum particle size. Also, installation of permeable bases or effective drainage systems for carrying free water out from under the pavement are helpful (Harrigan 2002).

The performance of aggregates under exposure to freezing and thawing can be evaluated in two ways: (1) past performance in the field, and (2) laboratory freeze-thaw tests of concrete specimens, ASTM C1646. If aggregates from the same source have previously given satisfactory service when used in concrete, they might be considered suitable. Aggregates not having a service record can be considered acceptable if they perform satisfactorily in air-entrained concretes subjected to freeze-thaw tests according to ASTM C1646. In these tests concrete specimens made with the aggregate in question are subjected to alternate cycles of freezing and thawing in water according to ASTM C666 (AASHO T 161). Deterioration is measured by (1) the reduction in the dynamic modulus of elasticity, (2) linear expansion, and (3) weight loss of the specimens. An expansion failure criterion of 0.035% in 350 freeze-thaw cycles or less is used by a number of state highway departments to help indicate whether or not an aggregate is susceptible to D-cracking. Different aggregate types may influence the criteria levels and empirical correlations between laboratory freeze-thaw tests. Field service records should be made to select the proper criterion (Vogler and Grove 1989).

Specifications may require that resistance to weathering be demonstrated by a sodium sulfate or magnesium sulfate test (ASTM C88 or AASHTO T 104). The test consists of a number of immersion cycles for a sample of the aggregate in a sulfate solution; this creates a pressure through salt-crystal growth in the aggregate pores similar to that produced by freezing water. The sample is then ovendried and the percentage of weight loss calculated. Unfortunately, this test is sometimes misleading. Aggregates behaving satisfactorily in the test might produce concrete with low freeze-thaw resistance; conversely, aggregates performing poorly might produce concrete with adequate resistance. This is attributed, at least in part, to the fact that the aggregates in the test are not confined by cement paste (as they would be in concrete) and the mechanisms of attack are not the same as in freezing and thawing. The test is most reliable for stratified rocks with porous layers or weak bedding planes.

An additional test that can be used to evaluate aggregates for potential D-cracking is the rapid pressure release method. An aggregate is placed in a pressurized chamber and the pressure is rapidly released causing the aggregate with a questionable pore system to fracture (Janssen and Snyder 1994). The amount of fracturing relates to the potential for D-cracking.

Wetting and Drying Properties

Weathering due to wetting and drying can also affect the durability of aggregates. The expansion and contraction coefficients of rocks vary with temperature and moisture content. If alternate wetting and drying occurs, severe strain can develop in some aggregates, and with certain types of rock this can cause a permanent increase in volume of the concrete and eventual breakdown. Clay lumps and other friable particles can degrade rapidly with repeated wetting and drying. Popouts can also develop due to the moisture-swelling characteristics of certain aggregates, especially clay balls and shales. While no specific tests are available to determine this tendency, an experienced petrographer can often be of assistance in determining this potential for distress.

Abrasion and Skid Resistance

The abrasion resistance of an aggregate is often used as a general index of its quality. Abrasion resistance is essential when the aggregate is to be used in concrete subject to abrasion, as in heavy-duty floors or pavements. Low abrasion resistance of an aggregate may increase the quantity of fines in the concrete during mixing; consequently, this may increase the water requirement and require an adjustment in the water-cement ratio.

The most common test for abrasion resistance is the Los Angeles abrasion test (rattler method) performed in accordance with ASTM C131 (AASHTO T 96) or ASTM C535. In this test a specified quantity of aggregate is placed in a steel drum containing steel balls, the drum is rotated, and the percentage of material worn away is measured. Specifications often set an upper limit on this mass loss. However, a comparison of the results of aggregate abrasion tests with the abrasion resistance of concrete made with the same aggregate do not generally show a

Fig. 5-16. Fractured carbonate aggregate particle as a source of distress in D-cracking (magnification 2.5X) (Stark 1976). (IMG12315)
clear correlation. Mass loss due to impact in the rattler is often as much as that due to abrasion. The wear resistance of concrete is determined more accurately by abrasion tests of the concrete itself (see Chapter 1).

To provide good skid resistance on pavements, the siliceous particle content of the fine aggregate should be at least 25%. For specification purposes, the siliceous particle content is considered equal to the insoluble residue content after treatment in hydrochloric acid under standardized conditions (ASTM D3042). Certain manufactured sands produce slippery pavement surfaces and should be investigated for acceptance before use.

**Strength and Shrinkage**

The strength of an aggregate is rarely tested and generally does not influence the strength of conventional concrete as much as the strength of the paste and the paste-aggregate bond. However, aggregate strength does become important in high-strength concrete. Aggregate stress levels in concrete are often much higher than the average stress over the entire cross section of the concrete. Aggregate tensile strengths range from 2 to 15 MPa (300 to 2300 psi) and compressive strengths from 65 to 270 MPa (10,000 to 40,000 psi). Strength can be tested according to ASTM C170.

Different aggregate types have different compressibility, modulus of elasticity, and moisture-related shrinkage characteristics that can influence the same properties in concrete. Aggregates with high absorption may have high shrinkage on drying. Quartz and feldspar aggregates, along with limestone, dolomite, and granite, are considered low shrinkage aggregates; while aggregates with sandstone, shale, slate, hornblende, and graywacke are often associated with high shrinkage in concrete (Fig. 5-17).

**Resistance to Acid and Other Corrosive Substances**

Portland cement concrete is durable in most natural environments; however, concrete in service is occasionally exposed to substances that will attack it.

Most acidic solutions will slowly or rapidly disintegrate portland cement concrete depending on the type and concentration of acid. Certain acids, such as oxalic acid, are harmless. Weak solutions of some acids have insignificant effects. Although acids generally attack and leach away the calcium compounds of the cement paste, they may not readily attack certain aggregates, such as siliceous aggregates. Calcereous aggregates often react readily with acids. However, the sacrificial effect of calcereous aggregates is often a benefit over siliceous aggregate in mild acid exposures or in areas where water is not flowing. With calcereous aggregate, the acid attacks the entire exposed concrete surface uniformly, reducing the rate of attack on the paste and preventing loss of aggregate particles at the surface. Calcereous aggregates also tend to neutralize the acid, especially in stagnant locations. Acids can also discolor concrete. Siliceous aggregate should be avoided when strong solutions of sodium hydroxide are present, as these solutions attack this type of aggregate.

Acid rain (often with a pH of 4 to 4.5) can slightly etch concrete surfaces, usually without affecting the performance of exposed concrete structures. Extreme acid rain or strong acid water conditions may warrant special concrete designs or precautions, especially in submerged areas. Continuous replenishment in acid with a pH of less than 4 is considered highly aggressive to buried concrete, such as pipe (Scanlon 1987). Concrete continuously exposed to liquid with a pH of less than 3 should be protected in a manner similar to concrete exposed to dilute acid solutions (ACI 515.1R).

Natural waters usually have a pH of more than 7 and seldom less than 6. Waters with a pH greater than 6.5 may be aggressive if they contain bicarbonates. Carbonic acid solutions with concentrations between 0.9 and 3 parts per million are considered to be destructive to concrete (ACI 515.1R and Kerkhoff 2007).

A low water-cement ratio, low permeability, and low-to-moderate cement content can increase the acid or corrosion resistance of concrete. A low permeability resulting from a low water-cement ratio or the use of silica fume or other pozzolans, helps keep the corrosive agent from penetrating into the concrete. Low-to-moderate cement contents result in less available paste to attack. The use of sacrificial calcereous aggregates should be considered where indicated.

Certain acids, gases, salts, and other substances that are not mentioned here also can disintegrate concrete. Acids and other chemicals that severely attack portland cement concrete should be prevented from coming in contact with the concrete by using protective coatings (Kerkhoff 2007).
Fire Resistance and Thermal Properties

The fire resistance and thermal properties of concrete—conductivity, diffusivity, and coefficient of thermal expansion—depend to some extent on the mineral constituents of the aggregates used. Manufactured and some naturally occurring lightweight aggregates are more fire resistant than normal-weight aggregates due to their insulating properties and high-temperature stability. Concrete containing a calcareous coarse aggregate performs better under fire exposure than a concrete containing quartz or siliceous aggregate such as granite or quartzite. At about 590°C (1060°F), quartz expands 0.85% causing disruptive expansion (ACI 216 and ACI 221). The coefficient of thermal expansion of aggregates ranges from 0.55 x 10^-6 per °C to 5 x 10^-6 per °C (1 x 10^-6 per °F to 9 x 10^-6 per °F). For more information refer to Chapter 15 for temperature-induced volume changes and to Chapter 18 for thermal conductivity and mass concrete considerations.

POTENTIALLY HARMFUL MATERIALS

Harmful substances that may be present in aggregates include organic impurities, silt, clay, shale, iron oxide, coal, lignite, and certain lightweight and soft particles (Table 5-6). In addition, rocks and minerals such as some cherts, strained quartz (Buck and Mather 1984), and certain dolomitic limestones are alkali reactive (see Table 5-7). Gypsum and anhydrite may cause sulfate attack. Certain aggregates, such as some shales, will cause popouts by swelling (simply by absorbing water) or by freezing of absorbed water (Fig. 5-18). Most specifications limit the permissible amounts of these substances. The performance history of an aggregate should be a determining factor in setting the limits for harmful substances. Test methods for detecting harmful substances qualitatively or quantitatively are listed in Table 5-6.

Aggregates are potentially harmful if they contain compounds known to react chemically with portland cement concrete and produce any of the following: (1) significant volume changes of the paste, aggregates, or both; (2) interference with the normal hydration of cement; and (3) otherwise harmful byproducts.

Organic impurities may delay setting and hardening of concrete, may reduce strength gain, and in unusual cases may cause deterioration. Organic impurities such as peat, humus, and organic loam may not be as detrimental but should be avoided.

Materials finer than the 75-µm (No. 200) sieve, especially silt and clay, may be present as loose dust and may form a coating on the aggregate particles. Even thin coatings of silt or clay on gravel particles can be harmful because they may weaken the bond between the cement paste and aggregate. If certain types of silt or clay are present in excessive amounts, water requirements may increase significantly.

### Table 5-6. Harmful Materials in Aggregates

<table>
<thead>
<tr>
<th>Substances</th>
<th>Effect on concrete</th>
<th>Test designation</th>
</tr>
</thead>
<tbody>
<tr>
<td>Organic impurities</td>
<td>Affects setting and hardening, may cause deterioration</td>
<td>ASTM C40 (AASHTO T 21)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>ASTM C87 (AASHTO T 71)</td>
</tr>
<tr>
<td>Materials finer than the 75-µm (No. 200) sieve</td>
<td>Affects bond, increases water requirement</td>
<td>ASTM C117 (AASHTO T 11)</td>
</tr>
<tr>
<td>Coal, lignite, or other lightweight materials</td>
<td>Affects durability, may cause stains and popouts</td>
<td>ASTM C123 (AASHTO T 113)</td>
</tr>
<tr>
<td>Soft particles</td>
<td>Affects durability</td>
<td>ASTM C235</td>
</tr>
<tr>
<td>Clay lumps and friable particles</td>
<td>Affects workability and durability, may cause popouts</td>
<td>ASTM C142 (AASHTO T 112)</td>
</tr>
<tr>
<td>Chert of less than 2.40 relative density</td>
<td>Affects durability, may cause popouts</td>
<td>ASTM C123 (AASHTO T 113)</td>
</tr>
<tr>
<td>Alkali-reactive aggregates</td>
<td>Causes abnormal expansion, map cracking, and popouts</td>
<td>ASTM C227</td>
</tr>
<tr>
<td></td>
<td></td>
<td>ASTM C289</td>
</tr>
<tr>
<td></td>
<td></td>
<td>ASTM C295</td>
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<tr>
<td></td>
<td></td>
<td>ASTM C342</td>
</tr>
<tr>
<td></td>
<td></td>
<td>ASTM C586</td>
</tr>
<tr>
<td></td>
<td></td>
<td>ASTM C1260 (AASHTO T 303)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>ASTM C1293</td>
</tr>
<tr>
<td></td>
<td></td>
<td>ASTM C1567</td>
</tr>
</tbody>
</table>

### Table 5-7. Some Potentially Harmful Reactive Minerals, Rock, and Synthetic Materials

<table>
<thead>
<tr>
<th>Alkali-silica reactive substances*</th>
<th>Alkali-carbonate reactive substances**</th>
</tr>
</thead>
<tbody>
<tr>
<td>Andesites</td>
<td>Opal</td>
</tr>
<tr>
<td>Argillites</td>
<td>Opaline shales</td>
</tr>
<tr>
<td>Certain siliceous</td>
<td>Phylites</td>
</tr>
<tr>
<td>limestones and dolomites</td>
<td>Quartzites</td>
</tr>
<tr>
<td>Chaledonic cherts</td>
<td>Quartzozes</td>
</tr>
<tr>
<td>Chaledony</td>
<td>Cherts</td>
</tr>
<tr>
<td>Cristobalite</td>
<td>Rhylolites</td>
</tr>
<tr>
<td>Dacites</td>
<td>Schists</td>
</tr>
<tr>
<td>Glassy or cryptocrystalline</td>
<td>Siliceous shales</td>
</tr>
<tr>
<td>volcanics</td>
<td>Strained quartz and certain other forms of quartz</td>
</tr>
<tr>
<td>Granite gneiss</td>
<td>Synthetic and natural silicious glass</td>
</tr>
<tr>
<td>Graywackes</td>
<td>Tridymite</td>
</tr>
<tr>
<td>Metagraywackes</td>
<td></td>
</tr>
</tbody>
</table>

* Several of the rocks listed (granite gneiss and certain quartz formations for example) react very slowly and may not show evidence of any harmful degree of reactivity until the concrete is over 20 years old.

** Only certain sources of these materials have shown reactivity.
There is a tendency for some fine aggregates to degrade from the grinding action in a concrete mixer; this effect, which is measured using ASTM C1137, may alter mixing water, entrained air and slump requirements. Coal or lignite, or other low-density materials such as wood or fibrous materials, in excessive amounts will affect the durability of concrete. If these impurities occur at or near the surface, they might disintegrate, pop out, or cause stains. Potentially harmful chert in coarse aggregate can be identified by using ASTM C123 (AASHTO T 113).

Soft particles in coarse aggregate are especially objectionable because they cause popouts and can affect durability and wear resistance of concrete. If friable, they could break up during mixing and thereby increase the amount of water required. Where abrasion resistance is critical, such as in heavy-duty industrial floors, testing may indicate that further investigation or another aggregate source is warranted.

Clay lumps present in concrete may absorb some of the mixing water, cause popouts in hardened concrete, and affect durability and wear resistance. They can also break up during mixing and thereby increase the mixing-water demand.

Aggregates can occasionally contain particles of iron oxide and iron sulfide that result in unsightly stains on exposed concrete surfaces (Fig. 5-19). The aggregate should meet the staining requirements of ASTM C330 (AASHTO M 195) when tested according to ASTM C641; the quarry face and aggregate stockpiles should not show evidence of staining.

As an additional aid in identifying staining particles, the aggregate can be immersed in a lime slurry. If staining particles are present, a blue-green gelatinous precipitate will form within 5 to 10 minutes; this will rapidly change to a brown color on exposure to air and light. The reaction should be complete within 30 minutes. If no brown gelatinous precipitate is formed when a suspect aggregate is placed in the lime slurry, there is little likelihood of any reaction taking place in concrete. These tests should be required when aggregates with no record of successful prior use are used in architectural concrete.

**ALKALI-AGGREGATE REACTIVITY**

Aggregates containing certain constituents can react with alkali hydroxides in concrete. The reactivity is potentially harmful only when it produces significant expansion (Mather 1975). This alkali-aggregate reactivity (AAR) has two forms—alkali-silica reaction (ASR) and alkali-carbonate reaction (ACR). ASR is of more concern than ACR because the occurrence of aggregates containing reactive silica minerals is more common. Alkali-reactive carbonate aggregates have a specific composition that is not very common.

Alkali-silica reactivity has been recognized as a potential source of distress in concrete since the late 1930s (Stanton 1940 and PCA 1940). Even though potentially reactive aggregates exist throughout North America, ASR distress in structural concrete is not common. There are a number of reasons for this:

- Most aggregates are chemically stable in hydraulic-cement concrete.
- Aggregates with good service records are abundant in many areas.
- Most concrete in service is dry enough to inhibit ASR.
- Use of certain pozzolans or slags can control ASR.
- In many concrete mixtures, the alkali content of the concrete is low enough to control harmful ASR.
- Some forms of ASR do not produce significant deleterious expansion.

To reduce ASR potential requires understanding the ASR mechanism; properly using tests to identify potentially reactive aggregates; and, if needed, taking steps to minimize the potential for expansion and related cracking.

**Alkali-Silica Reaction**

**Visual Symptoms of Expansive ASR.** Typical indicators of ASR might be any of the following: a network of cracks (Fig. 5-20); closed or spalled joints; relative displacements of different parts of a structure; or fragments breaking out of the surface of the concrete (popouts) (Fig. 5-21). Because ASR deterioration is slow, the risk of catastrophic failure is low. However, ASR can cause serviceability problems and can exacerbate other deterioration mechanisms such as those that occur in frost, deicer, or sulfate exposures.
Mechanism of ASR. The alkali-silica reaction forms a gel that swells as it draws water from the surrounding cement paste. Reaction products from ASR have a great affinity for moisture. In absorbing water, these gels can induce pressure, expansion, and cracking of the aggregate and surrounding paste. The reaction can be visualized as a two-step process:

1. Alkali hydroxide + reactive silica gel → reaction product (alkali-silica gel)
2. Gel reaction product + moisture → expansion

The amount of gel formed in the concrete depends on the amount and type of silica and alkali hydroxide concentration. The presence of gel does not always coincide with distress, and thus, gel presence does not necessarily indicate destructive ASR.

Factors Affecting ASR. For alkali-silica reaction to occur, the following three conditions must be present:

1. reactive forms of silica in the aggregate,
2. high-alkali (pH) pore solution, and
3. sufficient moisture.

If one of these conditions is absent, ASR cannot occur.

Test Methods for Identifying ASR Distress. It is important to distinguish between the reaction and damage resulting from the reaction. In the diagnosis of concrete deterioration, it is most likely that a gel product will be identified. But, in some cases significant amounts of gel are formed without causing damage to concrete. To pinpoint ASR as the cause of damage, the presence of deleterious ASR gel must be verified. A site of expansive reaction can be defined as an aggregate particle that is recognizably reactive or potentially reactive and is at least partially replaced by gel. Gel can be present in cracks and voids and may also be present in a ring surrounding an aggregate particle at its edges. A network of internal cracks connecting reacted aggregate particles is an almost certain indication that ASR is responsible for cracking. A petrographic examination (ASTM C856) is the most positive method for identifying ASR gel in concrete (Powers 1999). Petrography, when used to study a known reacted concrete, can confirm the presence of reaction products and verify ASR as an underlying cause of deterioration (Fig. 5-22).

Control of ASR in New Concrete. The best way to avoid ASR is to take appropriate precautions before concrete is placed. Standard concrete specifications may require modifications to address ASR. These modifications should be carefully tailored to avoid limiting the concrete producer’s options. This permits careful analysis of cementitious materials and aggregates and choosing a control strategy that optimizes effectiveness and the economic selection of materials. If the aggregate is not reactive by historical identification or testing, no special requirements are needed.
Identification of Potentially Reactive Aggregates. Field performance history is the best method of evaluating the susceptibility of an aggregate to ASR. For the most definitive evaluation, the existing concrete should have been in service for at least 15 years. Comparisons should be made between the existing and proposed concrete’s mix proportions, ingredients, and service environments. This process should tell whether special requirements are needed, are not needed, or whether testing of the aggregate or job concrete is required. The use of newer, faster test methods can be utilized for initial screening. Where uncertainties arise, lengthier tests can be used to confirm results. Table 5-8 describes different test methods used to evaluate potential alkali-silica reactivity. These tests should not be used to disqualify use of potentially reactive aggregates, as reactive aggregates can be safely used with the careful selection of cementitious materials as discussed below.

Materials and Methods to Control ASR. The most effective way of controlling expansion due to ASR is to design mixtures specifically to control ASR, preferably using locally available materials. The following options are not listed in priority order and, although usually not necessary, they can be used in combination with one another.

In North America, current practices include the use of a supplementary cementitious material or blended cement proven by testing to control ASR or limiting the alkali content of the concrete. Supplementary cementitious materials include fly ash, ground granulated blast-furnace slag, silica fume, and natural pozzolans. Blended cements use slag, fly ash, silica fume, and natural pozzolans to control ASR. Low-alkali portland cement (ASTM C150) with an alkali content of not more than 0.60% (equivalent sodium oxide) can be used to control ASR. Its use has been successful with slightly reactive to moderately reactive aggregates. However, low-alkali cements are not available in all areas. Thus, the use of locally available cements in combination with pozzolans, slags, or blended cements is preferable for controlling ASR. When pozzolans, slags, or blended cements are used to control ASR, their effectiveness must be determined by tests such as ASTM C1567 or C1293. Where applicable, different amounts of pozzolan or slag should be tested to determine the optimum dosage. Expansion usually decreases as the dosage of the pozzolan or slag increases (see Fig. 5-23). Lithium-based admixtures are also available to control ASR. Limestone sweetening (the popular term for replacing approximately 30% of the reactive sand-gravel aggregate with crushed limestone) is effective in controlling deterioration in some sand-gravel aggregate concretes. See AASHTO (2001), Farny and Kerkhoff (2007), and PCA (2007) for more information on tests to demonstrate the effectiveness of the above control measures.

Alkali-Carbonate Reaction

Mechanism of ACR. Reactions observed with certain dolomitic rocks are associated with alkali-carbonate reaction (ACR). Reactive rocks usually contain large crystals of dolomite scattered in and surrounded by a fine-grained matrix of calcite and clay. Calcite is one of the mineral forms of calcium carbonate; dolomite is the common name for calcium-magnesium carbonate. ACR is relatively rare because aggregates susceptible to this reaction are usually unsuitable for use in concrete for other reasons, such as strength potential. Argillaceous dolomitic limestone contains calcite and dolomite with appreciable amounts of clay and can contain small amounts of reactive silica. Alkali reactivity of carbonate rocks is not usually dependent upon its clay mineral composition (Hadley 1961). Aggregates have potential for expansive ACR if the following lithological characteristics exist (Ozol 2006 and Swenson 1967):
### Table 5-8. Test Methods for Alkali-Silica Reactivity (adapted from Farny and Kerkhoff 2007)

<table>
<thead>
<tr>
<th>Test name</th>
<th>Purpose</th>
<th>Type of test</th>
<th>Type of sample</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>ASTM C227</strong>, Potential Alkali-Reactivity of Cement-Aggregate Combinations (Mortar-Bar Method)</td>
<td>To test the susceptibility of cement-aggregate combinations to expansive reactions involving alcalies</td>
<td>Mortar bars stored over water at 37.8°C (100°F) and high relative humidity</td>
<td>At least 4 mortar bars; standard dimensions: 25 x 25 x 285 mm (1 x 1 x 11 1/4 in.)</td>
</tr>
<tr>
<td><strong>ASTM C289</strong>, Potential Alkali-Silica Reactivity of Aggregates (Chemical Method)</td>
<td>To determine potential reactivity of siliceous aggregates</td>
<td>Sample reacted with alkaline solution at 80°C (176°F)</td>
<td>Three 25 gram samples of crushed and sieved aggregate</td>
</tr>
<tr>
<td><strong>ASTM C294</strong>, Constituents of Natural Mineral Aggregates</td>
<td>To give descriptive nomenclature for the more common or important natural minerals—a good starting point to predict behavior</td>
<td>Visual identification</td>
<td>Varies, but should be representative of entire source</td>
</tr>
<tr>
<td><strong>ASTM C295</strong>, Petrographic Examination of Aggregates for Concrete</td>
<td>To evaluate possible aggregate reactivity through petrographic examination</td>
<td>Visual and microscopic examination of prepared samples—sieve analysis, microscopy, scratch or acid tests</td>
<td>Varies with knowledge of quarry: cores 53 to 100 mm in diameter (2% to 4 in.), 45 kg (100 lb) or 300 pieces, or 2 kg (4 lb)</td>
</tr>
<tr>
<td><strong>ASTM C342</strong>, Potential Volume Change of Cement-Aggregate Combinations</td>
<td>To determine the potential ASR expansion of cement-aggregate combinations</td>
<td>Mortar bars stored in water at 23°C (73.4°F)</td>
<td>3 mortar bars per cement-aggregate combination, standard dimensions 25 x 25 x 285 mm (1 x 1 x 11 1/4 in.)</td>
</tr>
<tr>
<td><strong>ASTM C441</strong>, Effectiveness of Pozzolans or GBFS in Preventing Excessive Expansion of Concrete Due to Alkali-Silica Reaction</td>
<td>To determine effectiveness of pozzolans or slag in controlling expansion from ASR</td>
<td>Mortar bars—using Pyrex glass as aggregate—stored over water at 37.8°C (100°F) and high relative humidity</td>
<td>At least 3 mortar bars and also 3 mortar bars of control mixture</td>
</tr>
<tr>
<td><strong>ASTM C856</strong>, Petrographic Examination of Hardened Concrete</td>
<td>To outline petrographic examination procedures of hardened concrete—useful in determining condition or performance</td>
<td>Visual (unmagnified) and microscopic examination of prepared samples</td>
<td>At least one core 150 mm diameter by 300 mm long (6 in. diameter by 12-in. long)</td>
</tr>
<tr>
<td><strong>ASTM C856 Annex (AASHTO T 299)</strong>, Uranyl-Acetate Treatment Procedure</td>
<td>To identify products of ASR in hardened concrete</td>
<td>Staining of a freshly exposed concrete surface and viewing under UV light</td>
<td>Varies: core with lapped surface, core with broken surface</td>
</tr>
<tr>
<td>Los Alamos staining method (Powers 1999)</td>
<td>To identify products of ASR in hardened concrete</td>
<td>Staining of a freshly-exposed concrete surface with two different reagents</td>
<td>Varies: core with lapped surface, core with broken surface</td>
</tr>
<tr>
<td><strong>ASTM C1260</strong> (AASHTO T 303), Potential Alkali-Reactivity of Aggregates (Mortar-Bar Method)</td>
<td>To test the potential for deleterious alkali-silica reaction of aggregate in mortar bars</td>
<td>Immersion of mortar bars in alkaline solution at 80°C (176°F)</td>
<td>At least 3 mortar bars</td>
</tr>
<tr>
<td><strong>ASTM C1293</strong>, Determination of Length Change of Concrete Due to Alkali-Silica Reaction (Concrete Prism Text)</td>
<td>To determine the potential ASR expansion of cement-aggregate combinations</td>
<td>Concrete prisms stored over water at 38°C (100.4°F)</td>
<td>3 prisms per cement-aggregate combination, standard dimensions: 75 x 75 x 285 mm (1 x 1 x 11 1/4 in.)</td>
</tr>
<tr>
<td><strong>ASTM C1567</strong>, Potential Alkali-Silica Reactivity of Combinations of Cementitious Materials and Aggregate (Accelerated)</td>
<td>To test the potential for deleterious alkali-silica reaction of cementitious materials and aggregate combinations in mortar bars</td>
<td>Immersion of mortar bars in alkaline solution at 80°C (176°F)</td>
<td>At least 3 mortar bars for each cementitious materials and aggregate combination</td>
</tr>
</tbody>
</table>
### Chapter 5 • Aggregates for Concrete

<table>
<thead>
<tr>
<th>Duration of test</th>
<th>Measurement</th>
<th>Criteria</th>
<th>Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td>Varies: first measurement at 14 days, then 1,2,3,4,5, 6,9, and 12 months and every 6 months after that as necessary</td>
<td>Length change</td>
<td>Per ASTM C33, maximum 0.10% expansion at 6 months, or if not available for a 6-month period maximum of 0.05% at 3 months</td>
<td>Test may not produce significant expansion, especially for carbonate aggregate. Long test duration. Expansions may not be from AAR.</td>
</tr>
<tr>
<td>24 hours</td>
<td>Drop in alkalinity and amount of silica solubilized</td>
<td>Point plotted on graph falls in deleterious or potentially deleterious area</td>
<td>Quick results. Some aggregates give low expansions even though they have high silica content—Not reliable.</td>
</tr>
<tr>
<td>Short duration—as long as it takes to visually examine the sample</td>
<td>Description of type and proportion of minerals in aggregate</td>
<td>Not applicable</td>
<td>These descriptions are used to characterize naturally-occurring minerals that make up common aggregate sources.</td>
</tr>
<tr>
<td>Short duration—visual examination does not involve long test periods</td>
<td>Particle characteristics, like shape, size, texture, color, mineral composition, physical condition</td>
<td>Not applicable</td>
<td>Usually includes optical microscopy; may include XRD analysis, differential thermal analysis, or infrared spectroscopy—see C294 for descriptive nomenclature.</td>
</tr>
<tr>
<td>52 weeks</td>
<td>Length change</td>
<td>Per ASTM C33, unsatisfactory aggregate if expansion equals or exceeds 0.200% at 1 year</td>
<td>Primarily used for aggregates from Oklahoma, Kansas, Nebraska, and Iowa.</td>
</tr>
<tr>
<td>Varies: first measurement at 14 days, then 1,2,3,4,5, 6,9, and 12 months and every 6 months after that as necessary</td>
<td>Length change</td>
<td>Per ASTM C989, minimum 75% reduction in expansion or 0.020% maximum expansion or per C618, comparison against low-alkali control</td>
<td>Highly reactive artificial aggregate may not represent real aggregate conditions. Pyrex contains alkalies.</td>
</tr>
<tr>
<td>Short duration—including preparation of samples and visual and microscopic examination</td>
<td>Is the aggregate known to be reactive? Orientation and geometry of cracks Is there any gel present?</td>
<td>See measurement—this examination determines if ASR reactions have taken place and their effects upon the concrete. Used in conjunction with other tests.</td>
<td>Specimens can be examined with stereomicroscopes, polarizing microscopes, metallographic microscopes, and SEM.</td>
</tr>
<tr>
<td>Immediate results</td>
<td>Intensity of fluorescence</td>
<td>Lack of fluorescence</td>
<td>Identifies small amounts of ASR gel whether they cause expansion or not. Opal, a natural aggregate, and carbonated paste can glow—interpret results accordingly. Text must be supplemented by petrographic examination and physical tests for determining concrete expansion.</td>
</tr>
<tr>
<td>Immediate results</td>
<td>Color of stain</td>
<td>Dark pink stain corresponds to ASR gel and indicates an advanced state of degradation</td>
<td></td>
</tr>
<tr>
<td>16 days</td>
<td>Length change</td>
<td>If greater than 0.10%, go to supplementary test procedures; if greater than 0.28%, indicative of potential deleterious expansion</td>
<td>Very fast alternative to C227. Useful for slowly reacting aggregates or ones that produce expansion late in the reaction.</td>
</tr>
<tr>
<td>Varies: first measurement at 7 days, then 28 and 56 days, then 3, 6, 9, and 12 months and every 6 months after that as necessary</td>
<td>Length</td>
<td>Per Appendix X1: potentially deleteriously reactive if expansion equals or exceeds 0.04% at 1 year (or 0.04% at 2 years if SCMs are to be evaluated)</td>
<td>Preferred method of assessment. Best represents the field. Long test duration for meaningful results. Use as a supplement to C227, C295, C289, C1260, and C1567. Similar to CSA A23.2-14A.</td>
</tr>
<tr>
<td>16 days</td>
<td>Length change</td>
<td>If greater than 0.10%, indicative of potential deleterious expansion; use C1293 (2-year test) to confirm</td>
<td>Very fast alternative to C1293. Allows for evaluation of supplementary cementing materials and determination of optimum dosage (Mortar-Bar Method).</td>
</tr>
</tbody>
</table>
minerals and water proportioned to have a relative density (specific gravity) less than that of the desirable aggregate particles but greater than that of the deleterious particles. The heavier particles sink to the bottom while the lighter particles float to the surface. This process can be used when acceptable and harmful particles have distinguishable relative densities.

Jigging separates particles with small differences in density by pulsating water current. Upward pulsations of water through a jig (a box with a perforated bottom) move the lighter material into a layer on top of the heavier material; the top layer is then removed.

Rising-current classification separates particles with large differences in density. Light materials, such as wood and lignite, are floated away in a rapidly upward moving stream of water.

Crushing is also used to remove soft and friable particles from coarse aggregates. This process is sometimes the only means of making material suitable for use. Unfortunately, with any process some acceptable material is always lost and removal of all harmful particles may be difficult or expensive.

HANDLING AND STORING AGGREGATES

Aggregates should be handled and stored in a way that minimizes segregation and degradation and prevents contamination by deleterious substances (Fig. 5-24). Stockpiles should be built up in thin layers of uniform thickness to minimize segregation. The most economical and acceptable method of forming aggregate stockpiles is the truck-dump method, which discharges the loads in a way that keeps them tightly joined. The aggregate is then reclaimed with a front-end loader. The loader should remove slices from the edges of the pile from bottom to top so that every slice will contain a portion of each horizontal layer.

AGGREGATE BENEFICIATION

Aggregate processing consists of: (1) basic processing—crushing, screening, and washing—to obtain proper gradation and cleanliness; and (2) beneficiation—upgrading quality by processing methods such as heavy media separation, jiggling, rising-current classification, and crushing.

In heavy media separation, aggregates are passed through a heavy liquid comprised of finely ground heavy minerals and water proportioned to have a relative density (specific gravity) less than that of the desirable aggregate particles but greater than that of the deleterious particles. The heavier particles sink to the bottom while the lighter particles float to the surface. This process can be used when acceptable and harmful particles have distinguishable relative densities.

Jigging separates particles with small differences in density by pulsating water current. Upward pulsations of water through a jig (a box with a perforated bottom) move the lighter material into a layer on top of the heavier material; the top layer is then removed.

Rising-current classification separates particles with large differences in density. Light materials, such as wood and lignite, are floated away in a rapidly upward moving stream of water.

Crushing is also used to remove soft and friable particles from coarse aggregates. This process is sometimes the only means of making material suitable for use. Unfortunately, with any process some acceptable material is always lost and removal of all harmful particles may be difficult or expensive.

Test Methods for Identifying ACR Distress. The three test methods commonly used to identify potentially alkali-carbonate reactive aggregate are:

- petrographic examination (ASTM C295);
- rock cylinder method (ASTM C586); and
- concrete prism test (ASTM C1105).

Materials and Methods to Control ACR. ACR-susceptible aggregate has a specific composition that is readily identified by petrographic testing. If a rock indicates ACR-susceptibility, one of the following preventive measures should be taken:

- selective quarrying to completely avoid reactive aggregate;
- blend aggregate according to Appendix in ASTM C1105; or
- limit aggregate size to smallest practical.

Low-alkaline cement and pozzolans are generally not very effective in controlling expansive ACR.
When aggregates are not delivered by truck, acceptable and inexpensive results can be obtained by forming the stockpile in layers with a clamshell bucket (cast-and-spread method); in the case of aggregates not subject to degradation, spreading the aggregates with a rubber-tire dozer and reclaiming with a front-end loader can be used. By spreading the material in thin layers, segregation is minimized. Whether aggregates are handled by truck, bucket loader, clamshell, or conveyor belt, stockpiles should not be built up in high, cone-shaped piles since this results in segregation. However, if circumstances necessitate construction of a conical pile, or if a stockpile has segregated, gradation variations can be minimized when the pile is reclaimed; in such cases aggregates should be loaded by continually moving around the circumference of the pile to blend sizes rather than by starting on one side and working straight through the pile.

Crushed aggregates segregate less than rounded (gravel) aggregates and larger-size aggregates segregate more than smaller sizes. To avoid segregation of coarse aggregates, size fractions can be stockpiled and batched separately. Proper stockpiling procedures, however, should eliminate the need for this. Specifications provide a range in the amount of material permitted in any size fraction partly because of segregation in stockpiling and batching operations.

Washed aggregates should be stockpiled in sufficient time before use so that they can drain to a uniform moisture content. Damp fine material has less tendency to segregate than dry material. When dry fine aggregate is dropped from buckets or conveyors, wind can blow away the fines; this should be avoided if possible.

Bulkheads or dividers should be used to avoid contamination of aggregate stockpiles. Partitions between stockpiles should be high enough to prevent intermingling of materials. Storage bins should be circular or nearly square. Their bottoms should slope not less than 50 degrees from the horizontal on all sides to a center outlet. When loading the bin, the material should fall vertically over the outlet into the bin. Chuting the material into a bin at an angle and against the bin sides will cause segregation. Baffle plates or dividers will help minimize segregation. Bins should be kept as full as possible since this reduces breakage of aggregate particles and the tendency to segregate. Recommended methods of handling and storing aggregates are discussed at length in Matthews (1965 to 1967), NCHRP (1967), and Bureau of Reclamation (1981).

Seashells may be present in the aggregate source. These shells are a hard material that can produce good quality concrete, however, a higher cement content may be required. Also, due to the angularity of the shells, additional cement paste is required to obtain the desired workability. Aggregate containing complete shells (uncrushed) should be avoided as their presence may result in voids in the concrete and lower the compressive strength.

Marine-dredged aggregates often contain salt from the seawater. The primary salts are sodium chloride and magnesium sulfate and the amount of salt on the aggregate is often not more than about 1% of the mass of the mixing water. The highest salt content occurs in sands located just above the high-tide level. Use of these aggregates with drinkable mix water often contributes less salt to the mixture than the use of seawater (as mix water) with salt-free aggregates.

Marine aggregates can be an appreciable source of chlorides. The presence of these chlorides may affect the concrete by (1) altering the time of set, (2) increasing drying shrinkage, (3) significantly increasing the risk of corrosion of steel reinforcement, and (4) causing efflorescence. Generally, marine aggregates containing large amounts of chloride should not be used in reinforced concrete.

Marine-dredged aggregates can be washed with fresh water to reduce the salt content. There is no maximum limit on the salt content of coarse or fine aggregate; however, the chloride limits presented in Chapter 9 should be followed.

RECYCLED-CONCRETE AGGREGATE

The concept of using old concrete pavements, buildings, and other structures as a source of aggregate has been demonstrated on several projects, resulting in both material and energy savings (ECCO 1999). The procedure involves (1) breaking up and removing the old concrete, (2) crushing in primary and secondary crushers (Fig. 5-25), (3) removing reinforcing steel and other embedded items,
(4) grading and washing, and (5) finally stockpiling the resulting coarse and fine aggregate (Fig. 5-26) (ACI 555). Dirt, gypsum board, wood, and other foreign materials should be prevented from contaminating the final product.

Recycled concrete is simply old concrete that has been crushed to produce aggregate. Recycled-concrete aggregate is primarily used in pavement reconstruction. It has been satisfactorily used as an aggregate in granular subbases, lean-concrete subbases, soil-cement, and in new concrete as the only source of aggregate or as a partial replacement of new aggregate.

Recycled-concrete aggregate generally has a higher absorption and a lower specific gravity than conventional aggregate (ACI 555). This results from the high absorption of porous mortar and hardened cement paste within the recycled concrete aggregate. Absorption values typically range from 3% to 10%, depending on the concrete being recycled; this absorption lies between those values for natural and lightweight aggregate. The values increase as coarse particle size decreases (Fig. 5-27). The high absorption of the recycled aggregate makes it necessary to add more water to achieve the same workability and slump than for concrete with conventional aggregates. Dry recycled aggregate absorbs water during and after mixing. To avoid this, recycled aggregate should be prewetted or stockpiles should be kept moist.

The particle shape of recycled-concrete aggregate is similar to crushed rock as shown in Fig. 5-28. The relative density decreases progressively as particle size decreases. The sulfate content of recycled-concrete aggregate should be determined to assess the possibility of deleterious sulfate reactivity. The chloride content should also be determined where applicable.

New concrete made from recycled-concrete aggregate generally has good durability. Carbonation, permeability, and resistance to freeze-thaw action have been found to be the same or even better than concrete with conventional aggregates. Concrete made with recycled coarse aggregates and conventional fine aggregate can obtain an adequate compressive strength. The use of recycled fine aggregate can result in minor compressive strength reductions. However, drying shrinkage and creep of concrete made with recycled aggregates is up to 100% higher than concrete with a corresponding conventional aggregate. This is due to the large amount of old cement paste and mortar especially in the fine aggregate. Therefore, considerably lower values of drying shrinkage can be achieved using recycled coarse aggregate with natural sand (Kerkhoff and Siebel 2001). As with any new aggregate source, recycled-concrete aggregate should be tested for durability, gradation, and other properties.

Recycled concrete used as coarse aggregate in new concrete possesses some potential for alkali-silica-reaction if the old concrete contained alkali-reactive aggregate. The
alkali content of the cement used in the old concrete has little effect on expansion due to alkali-silica-reaction. For highly reactive aggregates made from recycled concrete, special measures discussed under “Alkali–Silica Reaction” should be used to control ASR. Also, even if expansive ASR did not develop in the original concrete, it can not be assumed that it will not develop in the new concrete if special control measures are not taken. Petrographic examination and expansion tests are recommended to make this judgment (Stark 1996).

Concrete trial mixtures should be made to check the new concrete’s quality and to determine the proper mixture proportions. One potential problem with using recycled concrete is the variability in the properties of the old concrete that may in turn affect the properties of the new concrete. This can partially be avoided by frequent monitoring of the properties of the old concrete that is being recycled. Adjustments in the mixture proportions may then be needed.

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Admixtures are those ingredients in concrete other than portland cement, supplementary cementitious materials, water, and aggregates that are added to the mixture immediately before or during mixing (Fig. 6-1). Admixtures can be classified by function as follows:

1. Air-entraining admixtures
2. Water-reducing admixtures
3. Plasticizers
4. Accelerating admixtures
5. Retarding admixtures
6. Hydration-control admixtures
7. Corrosion inhibitors
8. Shrinkage reducers
9. Alkali-silica reactivity inhibitors
10. Coloring admixtures
11. Miscellaneous admixtures such as workability, bonding, dampproofing, permeability reducing, grouting, gas-forming, antiwashout, foaming, and pumping admixtures

Table 6-1 provides a much more extensive classification of admixtures.

Concrete should be workable, finishable, strong, durable, watertight, and wear resistant. These qualities can often be obtained easily and economically by the selection of suitable materials rather than by resorting to admixtures (except air-entraining admixtures when needed).

The major reasons for using admixtures are:

1. To reduce the cost of concrete construction
2. To achieve certain properties in concrete more effectively than by other means
3. To maintain the quality of concrete during the stages of mixing, transporting, placing, and curing in adverse weather conditions
4. To overcome certain emergencies during concreting operations

Despite these considerations, it should be borne in mind that no admixture of any type or amount can be considered a substitute for good concreting practice.

The effectiveness of an admixture depends upon factors such as type, brand, and amount of cementing materials; water content; aggregate shape, gradation, and proportions; mixing time; slump; and temperature of the concrete.

Admixtures being considered for use in concrete should meet applicable specifications as presented in Table 6-1. Trial mixtures should be made with the admixture and the job materials at temperatures and humidities anticipated on the job. In this way the compatibility of the admixture with other admixtures and job materials, as well as the effects of the admixture on the properties of the fresh and hardened concrete, can be observed. The amount of admixture recommended by the manufacturer or the optimum amount determined by laboratory tests should be used.
Table 6-1. Concrete Admixtures by Classification

<table>
<thead>
<tr>
<th>Type of admixture</th>
<th>Desired effect</th>
<th>Material</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Accelerators</strong> (ASTM C494 and AASHTO M 194, Type C)</td>
<td>Accelerate setting and early-strength development</td>
<td>Calcium chloride (ASTM D98 and AASHTO M 144)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Triethanolamine, sodium thiocyanate, calcium formate, calcium nitrite,</td>
</tr>
<tr>
<td></td>
<td></td>
<td>calcium nitrate</td>
</tr>
<tr>
<td><strong>Air detainers</strong></td>
<td>Decrease air content</td>
<td>Tributyl phosphate, dibutyl phthalate, octyl alcohol, water-insoluble</td>
</tr>
<tr>
<td></td>
<td></td>
<td>esters of carboxic and boric acid, silicones</td>
</tr>
<tr>
<td><strong>Air-entraining admixtures</strong> (ASTM C260 and AASHTO M</td>
<td>Improve durability in freeze-thaw, deicer, sulfate, and</td>
<td>Salts of wood resins (Vinsol resin), some synthetic detergents, salts</td>
</tr>
<tr>
<td>154)</td>
<td>alkali-reactive environments</td>
<td>of sulfonated lignin, salts of petroleum acids, salts of proteinaceous</td>
</tr>
<tr>
<td></td>
<td>Improve workability</td>
<td>material, fatty and resinous acids and their salts, alkyl/benzene</td>
</tr>
<tr>
<td></td>
<td></td>
<td>sulfonates, salts of sulfonated hydrocarbons</td>
</tr>
<tr>
<td><strong>Alkali-aggregate reactivity inhibitors</strong></td>
<td>Reduce alkali-aggregate reactivity expansion</td>
<td>Barium salts, lithium nitrate,</td>
</tr>
<tr>
<td></td>
<td></td>
<td>lithium carbonate, lithium hydroxide</td>
</tr>
<tr>
<td><strong>Antiwashout admixtures</strong></td>
<td>Cohesive concrete for underwater placements</td>
<td>Cellulose, acrylic polymer</td>
</tr>
<tr>
<td><strong>Bonding admixtures</strong></td>
<td>Increase bond strength</td>
<td>Polyvinyl chloride, polyvinyl acetate, acrylcs,</td>
</tr>
<tr>
<td></td>
<td></td>
<td>butadiene-styrene copolymers</td>
</tr>
<tr>
<td><strong>Coloring admixtures</strong> (ASTM C979)</td>
<td>Colored concrete</td>
<td>Modified carbon black, iron oxide, phthalocyanine, umber, chromium</td>
</tr>
<tr>
<td></td>
<td></td>
<td>oxide, titanium oxide, cobalt blue</td>
</tr>
<tr>
<td><strong>Corrosion inhibitors</strong> (ASTM C1582)</td>
<td>Reduce steel corrosion activity in a chloride-laden</td>
<td>Calcium nitrite, sodium nitrite, sodium benzoate, certain phosphates</td>
</tr>
<tr>
<td></td>
<td>environment</td>
<td>or fluosilicates, fluoaluminates, ester amines</td>
</tr>
<tr>
<td><strong>Dampproofing admixtures</strong></td>
<td>Retard moisture penetration into dry concrete</td>
<td>Soaps of calcium or ammonium stearate or oleate</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Butyl stearate</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Petroleum products</td>
</tr>
<tr>
<td><strong>Foaming agents</strong></td>
<td>Produce lightweight, foamed concrete with low density</td>
<td>Cationic and anionic surfactants</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Hydrolized protein</td>
</tr>
<tr>
<td><strong>Fungicides, germicides, and insecticides</strong></td>
<td>Inhibit or control bacterial and fungal growth</td>
<td>Polyhalogenated phenols</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Dieldrine emulsions</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Copper compounds</td>
</tr>
<tr>
<td><strong>Gas formers</strong></td>
<td>Cause expansion before setting</td>
<td>Aluminum powder</td>
</tr>
<tr>
<td><strong>Grouting admixtures</strong></td>
<td>Adjust grout properties for specific applications</td>
<td>See Air-entraining admixtures, Accelerators, Retarders, and Water</td>
</tr>
<tr>
<td></td>
<td></td>
<td>reducers</td>
</tr>
<tr>
<td><strong>Hydration control admixtures</strong></td>
<td>Suspend and reactivate cement hydration with stabilizer</td>
<td>Carboxylic acids</td>
</tr>
<tr>
<td></td>
<td>and activator</td>
<td>Phosphorus-containing organic acid salts</td>
</tr>
<tr>
<td><strong>Permeability reducers</strong></td>
<td>Decrease permeability</td>
<td>Latex</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Calcium stearate</td>
</tr>
<tr>
<td><strong>Pumping aids</strong></td>
<td>Improve pumpability</td>
<td>Organic and synthetic polymers</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Organic flocculents</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Organic emulsions of paraffin, coal tar, asphalt, acrylics</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Bentonite and pyrogenic silicas</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Hydrated lime (ASTM C141)</td>
</tr>
<tr>
<td><strong>Retarding admixtures</strong> (ASTM C494 and AASHTO M 194,</td>
<td>Retard setting time</td>
<td>Lignin</td>
</tr>
<tr>
<td>Type B)</td>
<td></td>
<td>Borax</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Sugars</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Tartaric acid and salts</td>
</tr>
<tr>
<td><strong>Shrinkage reducers</strong></td>
<td>Reduce drying shrinkage</td>
<td>Polyoxyalkylene alkyl ether</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Propylene glycol</td>
</tr>
<tr>
<td><strong>Superplasticizers</strong> <em>(ASTM C1017, Type 1)</em></td>
<td>Increase flowability of concrete</td>
<td>Sulfonated melamine formaldehyde condensates</td>
</tr>
<tr>
<td></td>
<td>Reduce water-cement ratio</td>
<td>Sulfonated napthalene formaldehyde condensates</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Lignosulfonates</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Polycarboxylates</td>
</tr>
</tbody>
</table>
AIR-ENTRAINING ADMIXTURES

Air-entraining admixtures are used to purposely introduce and stabilize microscopic air bubbles in concrete. Air-entrainment will dramatically improve the durability of concrete exposed to cycles of freezing and thawing (Fig. 6-2). Entrained air greatly improves concrete’s resistance to surface scaling caused by chemical deicers (Fig. 6-3). Furthermore, the workability of fresh concrete is improved significantly, and segregation and bleeding are reduced or eliminated.

Air-entrained concrete contains minute air bubbles that are distributed uniformly throughout the cement paste. Entrained air can be produced in concrete by use of an air-entraining cement, by introduction of an air-entraining admixture, or by a combination of both methods. An air-entraining cement is a portland cement with an air-entraining addition interground with the clinker during manufacture. An air-entraining admixture, on the other hand, is added directly to the concrete materials either before or during mixing.

The primary ingredients used in air-entraining admixtures are listed in Table 6-1. Specifications and methods of testing air-entraining admixtures are given in ASTM C260 and C233 (AASHTO M 154 and T 157). Air-entraining additions for use in the manufacture of air-entraining cements must meet requirements of ASTM C226. Applicable requirements for air-entraining cements are given in ASTM C150 and AASHTO M 85. See Chapter 8, Air-Entrained Concrete, Klieger (1966), and Whiting and Nagi (1998) for more information.

WATER-REDUCING ADMIXTURES

Water-reducing admixtures are used to reduce the quantity of mixing water required to produce concrete of a certain slump, reduce water-cement ratio, reduce cement content, or increase slump. Typical water reducers reduce the water content by approximately 5% to 10%. Adding a water-reducing admixture to concrete without reducing the water content can produce a mixture with a higher slump. The rate of slump loss, however, is not reduced and in most cases is increased (Fig. 6-4). Rapid slump loss results in reduced workability and less time to place concrete.

An increase in strength is generally obtained with water-reducing admixtures as the water-cement ratio is

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Table 6-1. Concrete Admixtures by Classification (Continued)

<table>
<thead>
<tr>
<th>Type of admixture</th>
<th>Desired effect</th>
<th>Material</th>
</tr>
</thead>
<tbody>
<tr>
<td>Superplasticizer* and retarder (ASTM C1017, Type 2)</td>
<td>Increase flowability with retarded set Reduce water–cement ratio</td>
<td>See superplasticizers and also water reducers</td>
</tr>
<tr>
<td>Water reducer (ASTM C494 and AASHTO M 194, Type A)</td>
<td>Reduce water content at least 5%</td>
<td>Lignosulfonates Hydroxylated carboxylic acids Carbohydrates (Also tend to retard set so accelerator is often added)</td>
</tr>
<tr>
<td>Water reducer and accelerator (ASTM C494 and AASHTO M 194, Type E)</td>
<td>Reduce water content (minimum 5%) and accelerate set</td>
<td>See water reducer, Type A (accelerator is added)</td>
</tr>
<tr>
<td>Water reducer and retarder (ASTM C494 and AASHTO M 194, Type D)</td>
<td>Reduce water content (minimum 5%) and retard set</td>
<td>See water reducer, Type A (retarder is added)</td>
</tr>
<tr>
<td>Water reducer—high range (ASTM C494 and AASHTO M 194, Type F)</td>
<td>Reduce water content (minimum 12%)</td>
<td>See superplasticizers</td>
</tr>
<tr>
<td>Water reducer—high range—and retarder (ASTM C494 and AASHTO M 194, Type G)</td>
<td>Reduce water content (minimum 12%) and retard set</td>
<td>See superplasticizers and also water reducers</td>
</tr>
<tr>
<td>Water reducer—mid range</td>
<td>Reduce water content (between 6 and 12%) without retarding</td>
<td>Lignosulfonates Polycarboxylates</td>
</tr>
</tbody>
</table>

* Superplasticizers are also referred to as high-range water reducers or plasticizers. These admixtures often meet both ASTM C494 (AASHTO M 194) and ASTM C1017 specifications.
cracks in concrete. Using a water reducer to reduce the cement and water content of a concrete mixture—while maintaining a constant water-cement ratio—can result in equal or reduced compressive strength, and can increase slump loss by a factor of two or more (Whiting and Dziedzic 1992).

Water reducers decrease, increase, or have no effect on bleeding, depending on the chemical composition of the admixture. A reduction of bleeding can result in finishing difficulties on flat surfaces when rapid drying conditions are present. Water reducers can be modified to give varying degrees of retardation while others do not signifi-

reduced. For concretes of equal cement content, air content, and slump, the 28-day strength of a water-reduced concrete containing a water reducer can be 10% to 25% greater than concrete without the admixture. Despite reduction in water content, water-reducing admixtures may cause increases in drying shrinkage. Usually the effect of the water reducer on drying shrinkage is small compared to other more significant factors that cause shrinkage.

Fig. 6-2. Frost damage (crumbling) at joints of a pavement (top), frost induced cracking near joints (bottom), and enlarged view of cracks (inset). (IMG12417, IMG12416, IMG12415)

Fig. 6-3. Scaled concrete surface resulting from lack of air entrainment, use of deicers, and poor finishing and curing practices. (IMG12414)

Fig. 6-4. Slump loss at 23°C (73°F) in concretes containing conventional water reducers (ASTM C494 and AASHTO M 194 Type D) compared with a control mixture (Whiting and Dziedzic 1992).
cantly affect the setting time. ASTM C494 (AASHTO M 194) Type A water reducers can have little effect on setting, while Type D admixtures provide water reduction with retardation, and Type E admixtures provide water reduction with accelerated setting. Type D water-reducing admixtures usually retard the setting time of concrete by one to three hours (Fig. 6-5). Some water-reducing admixtures may also entrain some air in concrete. Lignin-based admixtures can increase air contents by 1 to 2 percentage points. Concretes with water reducers generally have good air retention (Table 6-2).

The effectiveness of water reducers on concrete is a function of their chemical composition, concrete temperature, cement composition and fineness, cement content, and the presence of other admixtures. The classifications and components of water reducers are listed in Table 6-1. See Whiting and Dziedzic (1992) for more information on the effects of water reducers on concrete properties.

### MID-RANGE WATER REDUCING ADMIXTURES

Mid-range water reducers were first introduced in 1984. These admixtures provide significant water reduction (between 6 and 12%) for concretes with slumps of 125 to 200 mm (5 to 8 in.) without the retardation associated with high dosages of conventional (normal) water reducers. Normal water reducers are intended for concretes with slumps of 100 to 125 mm (4 to 5 in.). Mid-range water reducers can be used to reduce stickiness and improve finishability, pumpability, and placeability of concretes containing silica fume and other supplementary cementing materials. Some can also entrain air and be used in low slump concretes (Nmai, Schlagbaum, and Violetta 1998).

### HIGH-RANGE WATER REDUCING ADMIXTURES

High-range water reducers, ASTM C494 (AASHTO M 194) Types F (water reducing) and G (water reducing and retarding), can be used to impart properties induced by regular water reducers, only much more efficiently. They can greatly reduce water demand and cement contents and make low water-cement ratio, high-strength concrete with normal or enhanced workability. A water reduction of 12% to 30% can be obtained through the use of these admixtures. The reduced water content and water-cement ratio can produce concretes with (1) ultimate compressive strengths in excess of 70 MPa (10,000 psi), (2) increased early strength gain, (3) reduced chloride-ion penetration, and (4) other beneficial properties associated with low water-cement ratio concrete (Fig. 6-6).

---

**Fig. 6-5. Retardation of set in cement-reduced mixtures relative to control mixture. Concretes L and H contain conventional water reducer, concretes N, M, B, and X contain high-range water reducer (Whiting and Dziedzic 1992).**

**Fig. 6-6. Low water to cement ratio concrete with low chloride permeability—easily made with high-range water reducers—is ideal for bridge decks. (IMG12413)**
High-range water reducers are generally more effective than regular water-reducing admixtures in producing workable concrete. A significant reduction of bleeding can result with large reductions of water content; this can result in finishing difficulties on flat surfaces when rapid drying conditions are present. Some of these admixtures can cause significant slump loss (Fig. 6-7). Significant retardation is also possible, but can aggravate plastic shrinkage cracking without proper protection and curing (Fig. 6-5). Drying shrinkage, chloride permeability, air retention (Table 6-2), and strength development of concretes with high-range water reducers are comparable to concretes without them when compared at constant water-cement ratios (reduced cement and water contents) (Fig. 6-8).

Concretes with high-range water reducers can have larger entrained air voids and higher void-spacing factors than normal air-entrained concrete. This would generally indicate a reduced resistance to freezing and thawing; however, laboratory tests have shown that concretes with a moderate slump using high-range water reducers have good freeze-thaw durability, even with slightly higher void-spacing factors. This may be the result of lower water-cement ratios often associated with these concretes.

When the same chemicals used for high-range water reducers are used to make flowing concrete, they are often called plasticizers or superplasticizers (see discussion below).

**PLASTICIZERS FOR FLOWING CONCRETE**

Plasticizers, often called superplasticizers, are essentially high-range water reducers meeting ASTM C1017; these admixtures are added to concrete with a low-to-normal slump and water-cement ratio to make high-slump flowing concrete (Fig. 6-9). Flowing concrete is a highly fluid but workable concrete that can be placed with little or no vibration or compaction while still remaining essentially

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**Table 6-2. Loss of Air from Cement Reduced Concrete Mixtures**

<table>
<thead>
<tr>
<th>Mixture</th>
<th>Initial air content, %*</th>
<th>Final air content, %†</th>
<th>Percent air retained</th>
<th>Rate of air loss, %/minute</th>
</tr>
</thead>
<tbody>
<tr>
<td>C</td>
<td>Control</td>
<td>5.4</td>
<td>3.0</td>
<td>56</td>
</tr>
<tr>
<td>L</td>
<td>Water reducer</td>
<td>7.0</td>
<td>4.7</td>
<td>67</td>
</tr>
<tr>
<td>H</td>
<td>High-range water reducer</td>
<td>6.2</td>
<td>4.6</td>
<td>74</td>
</tr>
<tr>
<td>N</td>
<td>High-range water reducer</td>
<td>6.8</td>
<td>4.8</td>
<td>71</td>
</tr>
<tr>
<td>M</td>
<td>High-range water reducer</td>
<td>6.4</td>
<td>3.8</td>
<td>59</td>
</tr>
<tr>
<td>B</td>
<td>High-range water reducer</td>
<td>6.8</td>
<td>5.6</td>
<td>82</td>
</tr>
<tr>
<td>X</td>
<td>High-range water reducer</td>
<td>6.6</td>
<td>5.0</td>
<td>76</td>
</tr>
</tbody>
</table>

* Represents air content measured after addition of admixture.
† Represents air content taken at point where slump falls below 25 mm (1 in.).
(3-in.) slump concrete can easily produce a concrete with a 230-mm (9-in.) slump. Flowing concrete is defined by ASTM C1017 as a concrete having a slump greater than 190 mm (7 1⁄2 in.), yet maintaining cohesive properties.

ASTM C1017 has provisions for two types of admixtures: Type 1—plasticizing, and Type 2—plasticizing and retarding. Plasticizers are generally more effective than regular or mid-range water-reducing admixtures in producing flowing concrete. The effect of certain plasticizers in increasing workability or making flowing concrete is short-lived, 30 to 60 minutes; this period is followed by a rapid loss in workability or slump loss (Fig. 6-11). High temperatures can also aggravate slump loss. Due to their propensity for slump loss, these admixtures are some-

free of excessive bleeding or segregation. Following are a few of the applications where flowing concrete is used: (1) thin-section placements (Fig. 6-10), (2) areas of closely spaced and congested reinforcing steel, (3) tremie pipe (underwater) placements, (4) pumped concrete to reduce pump pressure, thereby increasing lift and distance capacity, (5) areas where conventional consolidation methods are impractical or can not be used, and (6) for reducing handling costs. The addition of a plasticizer to a 75-mm

![Fig. 6-9. Flowable concrete with a high slump (top) is easily placed (middle), even in areas of heavy reinforcing steel congestion (bottom). (IMG12324, IMG12325, IMG12326)](image1)

![Fig. 6-10. Plasticized, flowing concrete is easily placed in thin sections such as this bonded overlay that is not much thicker than 1 1⁄2 diameters of a quarter. (IMG12207)](image2)

![Fig. 6-11. Slump loss at 32°C (90°F) in flowing concretes (TN, TM, TB, and TX) compared with control mixture (TC) (Whiting and Dziedzic 1992).](image3)
times added to the concrete mixer at the jobsite. They are available in liquid and powder form. Extended-slump-life plasticizers added at the batch plant help reduce slump-loss problems. Setting time may be accelerated or retarded based on the admixture’s chemistry, dosage rate, and interaction with other admixtures and cementing materials in the concrete mixture. Some plasticizers can retard final set by one to almost four hours (Fig. 6-12). Strength development of flowing concrete is comparable to normal concrete (Fig. 6-13).

While it was previously noted that flowing concretes are essentially free of excessive bleeding, tests have shown that some plasticized concretes bleed more than control concretes of equal water-cement ratio (Fig. 6-14); but plasticized concretes bleed significantly less than control concretes of equally high slump and higher water content. High-slump, low-water-content, plasticized concrete has less drying shrinkage than a high-slump, high-water-content conventional concrete; however this concrete has similar or higher drying shrinkage than conventional low-slump, low-water-content concrete (Whiting 1979, Gebler 1982, and Whiting and Dziedzic 1992).

The effectiveness of the plasticizer is increased with an increasing amount of cement and fines in the concrete. It is also affected by the initial slump of the concrete.

Plasticized flowing concrete can have larger entrained air voids and greater void-spacing factors than conventional concrete. Air loss can also be significant. Some research has indicated poor frost- and deicer-scaling resistance for some flowing concretes when exposed to a continuously moist environment without the benefit of a drying period (Whiting and Dziedzic 1992). However, field performance of flowing concretes with low water to portland cement ratios has been good in most frost environments.
Table 6-1 lists the primary components and specifications for plasticizing (superplasticizer) admixtures.

## RETARDING ADMIXTURES

Retarding admixtures are used to delay the rate of setting of concrete. High temperatures of fresh concrete (30°C [86°F]) are often the cause of an increased rate of hardening that makes placing and finishing difficult. One of the most practical methods of counteracting this effect is to reduce the temperature of the concrete by cooling the mixing water and/or the aggregates. Retarders do not decrease the initial temperature of concrete. The bleeding rate and bleeding capacity of concrete is increased with retarders.

Retarding admixtures are useful in extending the setting time of concrete, but they are often also used in attempts to decrease slump loss and extend workability, especially prior to placement at elevated temperatures. The fallacy of this approach is shown in Fig. 6-15, where the addition of a retarder resulted in an increased rate of slump loss compared to the control mixtures (Whiting and Dziedzic 1992).

Retarders are sometimes used to: (1) offset the accelerating effect of hot weather on the setting of concrete; (2) delay the initial set of concrete or grout when difficult or unusual conditions of placement occur, such as placing concrete in large piers and foundations, cementing oil wells, or pumping grout or concrete over considerable distances; or (3) delay the set for special finishing techniques, such as an exposed aggregate surface.

The amount of water reduction for an ASTM C494 (AASHTO M 194) Type B retarding admixture is normally less than that obtained with a Type A water reducer. Type D admixtures are designated to provide both water reduction and retardation.

In general, some reduction in strength at early ages (one to three days) accompanies the use of retarders. The effects of these materials on the other properties of concrete, such as shrinkage, may not be predictable. Therefore, acceptance tests of retarders should be made with actual job materials under anticipated job conditions. The classifications and components of retarders are listed in Table 6-1.

## HYDRATION-CONTROL ADMIXTURES

Hydration controlling admixtures became available in the late 1980s. They consist of a two-part chemical system: (1) a stabilizer or retarder that essentially stops the hydration of cementing materials, and (2) an activator that reestablishes normal hydration and setting when added to the stabilized concrete. The stabilizer can suspend hydration for 72 hours and the activator is added to the mixture just before the concrete is used. These admixtures make it possible to reuse concrete returned in a ready-mix truck by suspending setting overnight. The admixture is also useful in maintaining concrete in a stabilized non-hardened state during long hauls. The concrete is reactivated when it arrives at the project. This admixture presently does not have a standard specification (Kinney 1989).

## ACCELERATING ADMIXTURES

An accelerating admixture is used to accelerate the rate of hydration (setting) and strength development of concrete at an early age. The strength development of concrete can also be accelerated by other methods: (1) using Type III or Type HE high-early-strength cement, (2) lowering the water-cement ratio by adding 60 to 120 kg/m³ (100 to 200 lb/yd³) of additional cement to the concrete, (3) using a water reducer, or (4) curing at higher temperatures.

Accelerators are designated as Type C admixtures under ASTM C494 (AASHTO M 194).

Calcium chloride (CaCl₂) is the chemical most commonly used in accelerating admixtures, especially for non-reinforced concrete. It should conform to the requirements of ASTM D98 (AASHTO M 144) and should be sampled and tested in accordance with ASTM D345.

The widespread use of calcium chloride as an accelerating admixture has provided much data and experience on the effect of this chemical on the properties of concrete. Besides accelerating strength gain, calcium chloride causes an increase in drying shrinkage, potential reinforcement corrosion, discoloration (a darkening of concrete), and an increase in the potential for scaling.

_Calcium chloride is not an antifreeze agent._ When used in allowable amounts, it will not reduce the freezing point of concrete by more than a few degrees. Attempts to protect...
concrete from freezing by this method are foolhardy. Instead, proven reliable precautions should be taken during cold weather (see Chapter 14, Cold-Weather Concreting).

When used, calcium chloride should be added to the concrete mixture in solution form as part of the mixing water. If added to the concrete in dry flake form, all of the dry particles may not be completely dissolved during mixing. Undissolved lumps in the mix can cause popouts or dark spots in hardened concrete.

The amount of calcium chloride added to concrete should be no more than is necessary to produce the desired results and in no case exceed 2% by mass of cementing material. When calculating the chloride content of commercially available calcium chloride, it can be assumed that:

1. Regular flake contains a minimum of 77% CaCl₂
2. Concentrated flake, pellet, or granular forms contain a minimum of 94% CaCl₂

An overdose can result in placement problems and can be detrimental to concrete. It may cause: rapid stiffening, a large increase in drying shrinkage, corrosion of reinforcement, and loss of strength at later ages (Abrams 1924 and Lackey 1992).

Applications where calcium chloride should be used with caution:

1. Concrete subjected to steam curing
2. Concrete containing embedded dissimilar metals, especially if electrically connected to steel reinforcement
3. Concrete slabs supported on permanent galvanized-steel forms
4. Colored concrete

Calcium chloride or admixtures containing soluble chlorides should not be used in the following:

1. Construction of parking garages
2. Prestressed concrete because of possible steel corrosion hazards
3. Concrete containing embedded aluminum (for example, conduit) since serious corrosion of the aluminum can result, especially if the aluminum is in contact with embedded steel and the concrete is in a humid environment
4. Concrete containing aggregates that, under standard test conditions, have been shown to be potentially deleteriously reactive
5. Concrete exposed to soil or water containing sulfates
6. Floor slabs intended to receive dry-shake metallic finishes
7. Hot weather generally
8. Massive concrete placements

The maximum chloride-ion content for corrosion protection of prestressed and reinforced concrete as recommended by the ACI 318 building code is presented in Table 6-3. Resistance to the corrosion of embedded steel is further improved with an increase in the depth of concrete cover over reinforcing steel, and a lower water-cement ratio. Stark (1989) demonstrated that concretes made with 1% CaCl₂ • 2H₂O by mass of cement developed active steel corrosion when stored continuously in fog. When 2% CaCl₂ • 2H₂O was used, active corrosion was detected in concrete stored in a fog room at 100% relative humidity. Risk of corrosion was greatly reduced at lower relative humidities (50%). Gaynor (1998) demonstrates how to calculate the chloride content of fresh concrete and compare it with recommended limits.

Several nonchloride, noncorrosive accelerators are available for use in concrete where chlorides are not recommended (Table 6-1). However, some nonchloride accelerators are not as effective as calcium chloride. Certain nonchloride accelerators are specially formulated for use in cold weather applications with ambient temperatures down to -7°C (20°F).

### CORROSION INHIBITORS

Corrosion inhibitors are used in concrete for parking structures, marine structures, and bridges where chloride salts are present. The chlorides can cause corrosion of steel reinforcement in concrete (Fig. 6-16). Ferrous oxide and ferric oxide form on the surface of reinforcing steel in concrete. Ferrous oxide, though stable in concrete’s alkaline environment, reacts with chlorides to form complexes that move away from the steel to form rust. The chloride ions continue to attack the steel until the passivating oxide layer is destroyed. Corrosion-inhibiting admixtures chemically arrest the corrosion reaction.

Commercially available corrosion inhibitors include: calcium nitrite, sodium nitrite, dimethyl ethanolamine, amines, phosphates, and ester amines. Anodic inhibitors, such as nitrites, block the corrosion reaction of the chloride-ions by chemically reinforcing and stabilizing the passive protective film on the steel; this ferric oxide film is created by the high pH environment in concrete. The

![Table 6-3. Maximum Chloride-Ion Content for Corrosion Protection of Reinforcement](image)
Nitrite-ions cause the ferric oxide to become more stable. In effect, the chloride-ions are prevented from penetrating the passive film and making contact with the steel.

A certain amount of nitrite can stop corrosion up to some level of chloride-ion. Therefore, increased chloride levels require increased levels of nitrite to stop corrosion.

Cathodic inhibitors react with the steel surface to interfere with the reduction of oxygen. The reduction of oxygen is the principal cathodic reaction in alkaline environments (Berke and Weil 1994). Corrosion inhibitors should conform to ASTM C1582.

**SHRINKAGE-REDUCING ADMIXTURES**

Shrinkage-reducing admixtures, introduced in the 1980s, have potential uses in bridge decks, critical floor slabs, and buildings where cracks and curling must be minimized for durability or aesthetic reasons (Fig. 6-17). Propylene glycol and polyoxyalkylene alkyl ether have been used as shrinkage reducers. Drying shrinkage reductions of between 25% and 50% have been demonstrated in laboratory tests. These admixtures have negligible effects on slump and air loss, but can delay setting. They are generally compatible with other admixtures (Nmai, Tomita, Hondo and Buffenbarger 1998 and Shah, Weiss and Yang 1998).

**CHEMICAL ADMIXTURES TO REDUCE ALKALI-AGGREGATE REACTIVITY (ASR INHIBITORS)**

Chemical admixtures to control alkali-silica reactivity (alkali-aggregate expansion) were introduced in the 1990s (Fig. 6-18). Lithium nitrate, lithium carbonate, lithium hydroxide, lithium aluminum silicate (decrepitated spodumene), and barium salts have shown reductions of alkali-silica reaction (ASR) in laboratory tests (Thomas and Stokes 1999 and AASHTO 2001). Some of these materials have potential for use as an additive to cement (Gajda 1996). There is little long-term field experience available on the effectiveness of these materials.

**COLORING ADMIXTURES (PIGMENTS)**

Natural and synthetic materials are used to color concrete for aesthetic and safety reasons (Fig. 6-19). Red concrete is used around buried electrical or gas lines as a warning to anyone near these facilities. Yellow concrete safety curbs are used in paving applications. Generally, the amount of pigments used in concrete should not exceed 10% by weight of the cement. Pigments used in amounts less than 6% generally do not affect concrete properties.
and reduce the water-cement ratio to less than 0.5. Most admixtures that reduce water-cement ratio consequently reduce permeability.

Some supplementary cementing materials, especially silica fume, reduce permeability through the hydration and pozzolanic-reaction process. Other admixtures that act to block the capillaries in concrete have been shown to be effective in reducing concrete corrosion in chemically aggressive environments. Such admixtures, designed for use in high-cement content/low-water-cement ratio concretes, contain aliphatic fatty acid and an aqueous emulsion of polymeric and aromatic globules (Aldred 1988).

PUMPING AIDS

Pumping aids are added to concrete mixtures to improve pumpability. Pumping aids cannot cure all unpumpable concrete problems; they are best used to make marginally pumpable concrete more pumpable. These admixtures increase viscosity or cohesion in concrete to reduce dewatering of the paste while under pressure from the pump.

Some pumping aids may increase water demand, reduce compressive strength, cause air entrainment, or retard setting time. These side effects can be corrected by adjusting the mix proportions or adding another admixture to offset the side effect.

A partial list of materials used in pumping aids is given in Table 6-1. Some admixtures that serve other primary purposes but also improve pumpability are air-entraining agents, and some water-reducing and retarding admixtures.

BONDING ADMIXTURES AND BONDING AGENTS

Bonding admixtures are usually water emulsions of organic materials including rubber, polyvinyl chloride, polyvinyl acetate, acrylics, styrene butadiene copolymers, and other polymers. They are added to portland cement mixtures to increase the bond strength between old and new concrete. Flexural strength and resistance to chloride-ion ingress are also improved. They are added in proportions equivalent to 5% to 20% by mass of the cementing materials; the actual quantity depending on job conditions and type of admixture used. Some bonding admixtures may increase the air content of mixtures. Nonreemulsifiable types are resistant to water, better suited to exterior application, and used in places where moisture is present.

The ultimate result obtained with a bonding admixture will be only as good as the surface to which the concrete is applied. The surface must be dry, clean, sound, free of dirt, dust, paint, and grease, and at the proper temperature. Organic or polymer modified concretes are acceptable for patching and thin-bonded overlayer, particularly where feather-edged patches are desired.

Unmodified carbon black substantially reduces air content. Most carbon black for coloring concrete contains an admixture to offset this effect on air. Before a coloring admixture is used on a project, it should be tested for color fastness in sunlight and autoclaving, chemical stability in cement, and effects on concrete properties. Calcium chloride should not be used with pigments to avoid color distortions. Pigments should conform to ASTM C979.

DAMPPROOFING ADMIXTURES

The passage of water through concrete can usually be traced to the existence of cracks or areas of incomplete consolidation. Sound, dense concrete made with a water-cement ratio of less than 0.50 by mass will be watertight if it is properly placed and cured.

Admixtures known as dampproofing agents include certain soaps, stearates, and petroleum products. They may, but generally do not, reduce the permeability of concretes that have low cement contents, high water-cement ratios, or a deficiency of fines in the aggregate. Their use in well-proportioned mixes, may increase the mixing water required and actually result in increased rather than reduced permeability.

Dampproofing admixtures are sometimes used to reduce the transmission of moisture through concrete that is in contact with water or damp earth. Many so-called dampproofers are not effective, especially when used in concretes that are in contact with water under pressure.

PERMEABILITY-REDUCING ADMIXTURES

Permeability-reducing admixtures reduce the rate at which water under pressure is transmitted through concrete. One of the best methods of decreasing permeability in concrete is to increase the moist-curing period and reduce the water-cement ratio to less than 0.5. Most admixtures that reduce water-cement ratio consequently reduce permeability.

Some supplementary cementing materials, especially silica fume, reduce permeability through the hydration and pozzolanic-reaction process. Other admixtures that act to block the capillaries in concrete have been shown to be effective in reducing concrete corrosion in chemically aggressive environments. Such admixtures, designed for use in high-cement content/low-water-cement ratio concretes, contain aliphatic fatty acid and an aqueous emulsion of polymeric and aromatic globules (Aldred 1988).
Bonding agents should not be confused with bonding admixtures. Admixtures are an ingredient in the concrete; bonding agents are applied to existing concrete surfaces immediately before the new concrete is placed. Bonding agents help “glue” the existing and the new materials together. Bonding agents are often used in restoration and repair work; they consist of portland cement or latex-modified portland cement grout or polymers such as epoxy resins (ASTM C881 or AASHTO M 235) or latex (ASTM C1059).

**GROUTING ADMIXTURES**

Portland cement grouts are used for a variety of purposes: to stabilize foundations, set machine bases, fill cracks and joints in concrete work, cement oil wells, fill cores of masonry walls, grout prestressing tendons and anchor bolts, and fill the voids in preplaced aggregate concrete. To alter the properties of grout for specific applications, various air-entraining admixtures, accelerators, retarders, and nonshrink admixtures are often used.

**GAS-FORMING ADMIXTURES**

Aluminum powder and other gas-forming materials are sometimes added to concrete and grout in very small quantities to cause a slight expansion of the mixture prior to hardening. This may be of benefit where the complete grouting of a confined space is essential, such as under machine bases or in post-tensioning ducts of prestressed concrete. These materials are also used in larger quantities to produce autoclaved cellular concretes. The amount of expansion that occurs is dependent upon the amount of gas-forming material used, the temperature of the fresh mixture, the alkali content of the cement, and other variables. Where the amount of expansion is critical, careful control of mixtures and temperatures must be exercised. Gas-forming agents will not overcome shrinkage after hardening caused by drying or carbonation.

**AIR DETRAINERS**

Air-detraining admixtures reduce the air content in concrete. They are used when the air content cannot be reduced by adjusting the mix proportions or by changing the dosage of the air-entraining agent and other admixtures. However, air-detainers are rarely used and their effectiveness and dosage rate should be established on trial mixes prior to use on actual job mixes. Materials used in air-detaining agents are listed in Table 6-1.

**FUNGICIDAL, GERMICIDAL, AND INSECTICIDAL ADMIXTURES**

Bacteria and fungal growth on or in hardened concrete may be partially controlled through the use of fungicidal, germicidal, and insecticidal admixtures. The most effective materials are polyhalogenated phenols, dieldrin emulsions, and copper compounds. The effectiveness of these materials is generally temporary, and in high dosages they may reduce the compressive strength of concrete.

**ANTIWASHOUT ADMIXTURES**

Antiwashout admixtures increase the cohesiveness of concrete to a level that allows limited exposure to water with little loss of cement. This allows placement of concrete in water and under water without the use of tremies. The admixtures increase the viscosity of water in the mixture resulting in a mix with increased thixotropy and resistance to segregation. They usually consist of water soluble cellulose ether or acrylic polymers.

**COMPATIBILITY OF ADMIXTURES AND CEMENTITIOUS MATERIALS**

Fresh concrete problems of varying severity are encountered due to cement-admixture incompatibility and incompatibility between admixtures. Incompatibility between supplementary cementing materials and admixtures or cements can also occur. Slump loss, air loss, early stiffening, and other factors affecting fresh concrete properties can result from incompatibilities. While these problems primarily affect the plastic-state performance of concrete, long-term hardened concrete performance may also be adversely affected. For example, early stiffening can cause difficulties with consolidation of concrete, therefore compromising strength.

Taylor et al. (2006 and 2006a) developed protocols for testing materials for potential incompatibility in a particular combination. This preconstruction testing can reduce the likelihood of performance problems in the field. However, reliable test methods are not available to adequately address all incompatibility issues due to variations in materials, mixing equipment, mixing time, and environmental factors. When incompatibility is discovered in the field, a common solution is to simply change admixtures or cementing materials (Helmuth, Hills, Whiting, and Bhattacharja 1995, Tagni-Hamou and Aïtcin 1993, and Tang and Bhattacharja 1997).

**STORING AND DISPENSING CHEMICAL ADMIXTURES**

Liquid admixtures can be stored in barrels or bulk tankers. Powdered admixtures can be placed in special storage bins and some are available in premeasured plastic bags. Admixtures added to a truck mixer at the jobsite are often in plastic jugs or bags. Powdered admixtures, such as cer-
tain plasticizers, or a barrel of admixture may be stored at the project site.

Dispenser tanks at concrete plants should be properly labeled for specific admixtures to avoid contamination and avoid dosing the wrong admixture. Most liquid chemical admixtures should not be allowed to freeze; therefore, they should be stored in heated environments. Consult the admixture manufacturer for proper storage temperatures. Powdered admixtures are usually less sensitive to temperature restrictions, but may be sensitive to moisture.

Liquid chemical admixtures are usually dispensed individually in the batch water by volumetric means (Fig. 6-20). Liquid and powdered admixtures can be measured by mass, but powdered admixtures should not be measured by volume. Care should be taken to not combine certain admixtures prior to their dispensing into the batch as some combinations may neutralize the desired effect of the admixtures. Consult the admixture manufacturer concerning compatible admixture combinations or perform laboratory tests to document performance.

Fig. 6-20. Liquid admixture dispenser at a ready mix plant provides accurate volumetric measurement of admixtures. (IMG12323)

Hester, Weston T., Superplasticizers in Ready Mixed Concrete (A Practical Treatment for Everyday Operations), National Ready Mixed Concrete Association, Publication No. 158, Silver Spring, Maryland, 1979.

Kinney, F. D., “Reuse of Returned Concrete by Hydration Control: Characterization of a New Concept,” Superplasticizers and Other Chemical Admixtures in Concrete, SP119, American Concrete Institute, Farmington Hills, Michigan, 1989, pages 19 to 40.


Nmai, Charles K.; Schlagbaum, Tony; and Violetta, Brad, “A History of Mid-Range Water-Reducing Admixtures,” Concrete International, American Concrete Institute, Farmington Hills, Michigan, April 1998, pages 45 to 50.

Nmai, Charles K.; Tomita, Rokuro; Hondo, Fumiaki; and Buffenbarger, Julie, “Shrinkage-Reducing Admixtures,” Concrete International, American Concrete Institute, Farmington Hills, Michigan, April 1998, pages 31 to 37.


Shah, Surendra P.; Weiss, J. Jason; and Yang, Wei, “Shrinkage Cracking—Can it be Prevented?,” Concrete International, American Concrete Institute, Farmington Hills, Michigan, April 1998, pages 51 to 55.


Fibers have been used in construction materials for many centuries. The last three decades have seen a growing interest in the use of fibers in ready-mixed concrete, pre-cast concrete, and shotcrete. Fibers made from steel, plastic, glass, and natural materials (such as wood cellulose) are available in a variety of shapes, sizes, and thicknesses; they may be round, flat, crimped, and deformed with typical lengths of 6 mm to 150 mm (0.25 in. to 6 in.) and thicknesses ranging from 0.005 mm to 0.75 mm (0.0002 in. to 0.03 in.) (Fig. 7-1). They are added to concrete during mixing. The main factors that control the performance of the composite material are:

1. Physical properties of fibers and matrix
2. Strength of bond between fibers and matrix

Although the basic governing principles are the same, there are several characteristic differences between conventional reinforcement and fiber systems:

1. Fibers are generally distributed throughout a given cross section whereas reinforcing bars or wires are placed only where required
2. Most fibers are relatively short and closely spaced as compared with continuous reinforcing bars or wires
3. It is generally not possible to achieve the same area of reinforcement to area of concrete using fibers as compared to using a network of reinforcing bars or wires

Fibers are typically added to concrete in low volume dosages (often less than 1%), and have been shown to be effective in reducing plastic shrinkage cracking.

Fibers typically do not significantly alter free shrinkage of concrete, however at high enough dosages they can increase the resistance to cracking and decrease crack width (Shah, Weiss, and Yang 1998).

ADVANTAGES AND DISADVANTAGES OF USING FIBERS

Fibers are generally distributed throughout the concrete cross section. Therefore, many fibers are inefficiently located for resisting tensile stresses resulting from applied loads. Depending on fabrication method, random orientation of fibers may be either two-dimensional (2-D) or three-dimensional (3-D). Typically, the spray-up fabrication method has a 2-D random fiber orientation whereas the premix (or batch) fabrication method typically has a 3-D random fiber orientation. Also, many fibers are observed to extend across cracks at angles other than 90º or may have less than the required embedment length for development of adequate bond. Therefore, only a small percentage of the fiber content may be efficient in resisting tensile or flexural stresses. “Efficiency factors” can be as low as 0.4 for 2-D random orientation and 0.25 for 3-D random orientation. The efficiency factor depends on fiber length and critical embedment length. From a conceptual point of view, reinforcing with fibers is not a highly efficient method of obtaining composite strength.

Fiber concretes are best suited for thin section shapes where correct placement of conventional reinforcement would be extremely difficult. In addition, spraying of fiber concrete accommodates the fabrication of irregularly shaped products. A substantial weight savings can be realized using relatively thin fiber concrete sections having
the equivalent strength of thicker conventionally reinforced concrete sections.

**TYPES AND PROPERTIES OF FIBERS AND THEIR EFFECT ON CONCRETE**

**Steel Fibers**

Steel fibers are short, discrete lengths of steel with an aspect ratio (ratio of length to diameter) from about 20 to 100, and with any of several cross sections. Some steel fibers have hooked ends to improve resistance to pullout from a cement-based matrix (Fig. 7-2).

ASTM A820 classifies four different types based on their manufacture. Type I – Cold-drawn wire fibers are the most commercially available, manufactured from drawn steel wire. Type II – Cut sheet fibers are manufactured as the name implies: steel fibers are laterally sheared off steel sheets. Type III – Melt-extracted fibers are manufactured with a relatively complicated technique where a rotating wheel is used to lift liquid metal from a molten metal surface by capillary action. The extracted molten metal is then rapidly frozen into fibers and thrown off the wheel by centrifugal force. The resulting fibers have a crescent-shaped cross section. Type IV – Other fibers. For tolerances for length, diameter, and aspect ratio, as well as minimum tensile strength and bending requirement, see ASTM A820.

Steel-fiber volumes used in concrete typically range from 0.25% to 2%. Volumes of more than 2% generally reduce workability and fiber dispersion and require special mix design or concrete placement techniques.

The compressive strength of concrete is only slightly affected by the presence of fibers. The addition of 1.5% by volume of steel fibers can increase the direct tensile strength by up to 40% and the flexural strength up to 150%.

Steel fibers do not affect free shrinkage. Steel fibers delay the fracture of restrained concrete during shrinkage and they improve stress relaxation by creep mechanisms (Altoubat and Lange 2001).

The durability of steel-fiber concrete is contingent on the same factors as conventional concrete. Freeze-thaw durability is not diminished by the addition of steel fibers provided the mix is adjusted to accommodate the fibers, the concrete is properly consolidated during placement, and is air-entrained. With properly designed and placed concrete, little or no corrosion of the fibers occurs. Any surface corrosion of fibers is cosmetic as opposed to a structural condition.

Steel fibers have a relatively high modulus of elasticity (Table 7-1). Their bond to the cement matrix can be enhanced by mechanical anchorage or surface roughness and they are protected from corrosion by the alkaline environment in the cement matrix (ACI 544.1R-96).

Steel fibers are most commonly used in airport pavements and runway/taxi overlays. They are also used in bridge decks (Fig. 7-3), industrial floors, and highway pavements. Structures exposed to high-velocity water flow have been shown to last about three times longer than conventional concrete alternatives. Steel fiber concrete is also used for many precast concrete applications that make use of the improved impact resistance or toughness imparted by the fibers. In utility boxes and septic tanks, steel fibers replace conventional reinforcement.

Steel fibers are also widely used with shotcrete in thin-layer applications, especially rock-slope stabilization and tunnel linings. Silica fume and accelerators have enabled shotcrete to be placed in thicker layers. Silica fume also reduces the permeability of the shotcrete material (Morgan 1987). Steel-fiber shotcrete has been successfully applied with fiber volumes up to 2%.

Slurry-infiltrated concrete (SIFCON) with fiber volumes up to 20% has been used since the late 1970s. Slurry-
### Table 7-1. Properties of Selected Fiber Types

<table>
<thead>
<tr>
<th>Fiber type</th>
<th>Relative density (specific gravity)</th>
<th>Diameter, µm (0.001 in.)</th>
<th>Tensile strength, MPa (ksi)</th>
<th>Modulus of elasticity, MPa (ksi)</th>
<th>Strain at failure, %</th>
</tr>
</thead>
<tbody>
<tr>
<td>Steel</td>
<td>7.80</td>
<td>100-1000 (4-40)</td>
<td>500-2600 (70-380)</td>
<td>210,000 (30,000)</td>
<td>0.5-3.5</td>
</tr>
<tr>
<td>Glass E</td>
<td>2.54</td>
<td>8-15 (0.3-0.6)</td>
<td>2000-4000 (290-580)</td>
<td>72,000 (10,400)</td>
<td>3.0-4.8</td>
</tr>
<tr>
<td>Glass AR</td>
<td>2.70</td>
<td>12-20 (0.5-0.8)</td>
<td>1500-3700 (220-540)</td>
<td>80,000 (11,600)</td>
<td>2.5-3.6</td>
</tr>
<tr>
<td>Synthetic</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Acrylic</td>
<td>1.18</td>
<td>5-17 (0.2-0.7)</td>
<td>200-1000 (30-145)</td>
<td>17,000-19,000 (2,500-2,800)</td>
<td>28-50</td>
</tr>
<tr>
<td>Aramid</td>
<td>1.44</td>
<td>10-12 (0.4-0.47)</td>
<td>2000-3100 (300-450)</td>
<td>62,000-120,000 (9,000-17,000)</td>
<td>2-3.5</td>
</tr>
<tr>
<td>Carbon</td>
<td>1.90</td>
<td>8-9 (0.3-0.35)</td>
<td>1800-2600 (260-380)</td>
<td>230,000-380,000 (33,400-55,100)</td>
<td>0.5-1.5</td>
</tr>
<tr>
<td>Nylon</td>
<td>1.14</td>
<td>23 (0.9)</td>
<td>1000 (140)</td>
<td>5,200 (750)</td>
<td>20</td>
</tr>
<tr>
<td>Polyester</td>
<td>1.38</td>
<td>10-80 (0.4-3.0)</td>
<td>280-1200 (40-170)</td>
<td>10,000-18,000 (1,500-2,500)</td>
<td>10-50</td>
</tr>
<tr>
<td>Polyethylene</td>
<td>0.96</td>
<td>25-1000 (1-40)</td>
<td>80-600 (11-85)</td>
<td>5,000 (725)</td>
<td>12-100</td>
</tr>
<tr>
<td>Polypropylene</td>
<td>0.90</td>
<td>20-200 (0.8-8)</td>
<td>450-700 (65-100)</td>
<td>3,500-5,200 (500-750)</td>
<td>6-15</td>
</tr>
<tr>
<td>Natural</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Wood cellulose</td>
<td>1.50</td>
<td>25-125 (1-5)</td>
<td>350-2000 (51-290)</td>
<td>10,000-40,000 (1,500-5,800)</td>
<td></td>
</tr>
<tr>
<td>Sisal</td>
<td>1.50</td>
<td>280-600 (40-85)</td>
<td>13,000-25,000 (1,900-3,800)</td>
<td>3.5</td>
<td></td>
</tr>
<tr>
<td>Coconut</td>
<td>1.12-1.15</td>
<td>100-400 (4-16)</td>
<td>120-200 (17-29)</td>
<td>19,000-25,000 (2,800-3,800)</td>
<td>10-25</td>
</tr>
<tr>
<td>Bamboo</td>
<td>1.50</td>
<td>50-400 (2-16)</td>
<td>350-500 (51-73)</td>
<td>33,000-40,000 (4,800-5,800)</td>
<td></td>
</tr>
<tr>
<td>Jute</td>
<td>1.02-1.04</td>
<td>100-200 (4-8)</td>
<td>250-350 (36-51)</td>
<td>25,000-32,000 (3,800-4,600)</td>
<td>1.5-1.9</td>
</tr>
<tr>
<td>Elephant grass</td>
<td>1.50</td>
<td>425 (17)</td>
<td>180 (26)</td>
<td>4,900 (710)</td>
<td>3.6</td>
</tr>
</tbody>
</table>

Adapted from PCA (1991) and ACI 544.1R-96.

Infiltrated concrete can be used to produce a component or structure with strength and ductility that far exceeds that of conventionally mixed or sprayed fiber concrete. SIFCON is not inexpensive and needs fine-tuning, but it holds potential for applications exposed to severe conditions and requiring very high strength and toughness. These applications include impact and blast-resistant structures, refractories, protective revetment, and taxiway and pavement repairs (Fig. 7-4). Table 7-2 shows a SIFCON mix design.

### Table 7-2. SIFCON Mix Design.

<table>
<thead>
<tr>
<th>Component</th>
<th>Amount</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cement</td>
<td>1000 kg/m³ (1686 lb/yd³)</td>
</tr>
<tr>
<td>Water</td>
<td>330 kg/m³ (556 lb/yd³)</td>
</tr>
<tr>
<td>Siliceous Sand ≤ 0.7 mm (≤ 0.028 in.)</td>
<td>860 kg/m³ (1450 lb/yd³)</td>
</tr>
<tr>
<td>Silica Slurry</td>
<td>13 kg/m³ (1.3 lb/yd³)</td>
</tr>
<tr>
<td>High-Range Water Reducer</td>
<td>35 kg/m³ (3.7 lb/yd³)</td>
</tr>
<tr>
<td>Steel Fibers (about 10 Vol.-%)</td>
<td>800 kg/m³ (84 lb/yd³)</td>
</tr>
</tbody>
</table>
Fiber modifications to improve long-term durability involve (1) specially formulated chemical coatings to help combat hydration-induced embrittlement, and (2) employment of a dispersed microsilica slurry to adequately fill fiber voids, thereby reducing potential for calcium hydroxide infiltration.

A low-alkaline cement has been developed in Japan that produces no calcium hydroxide during hydration. Accelerated tests with the cement in alkali-resistant-glass fiber-reinforced concrete samples have shown greater long-term durability than previously achieved.

Metaalkalin can be used in glass-fiber-reinforced concrete without significantly affecting flexural strength, strain, modulus of elasticity, and toughness. (Marikunte, Aldea, Shah 1997).

The single largest application of glass-fiber concrete has been the manufacture of exterior building façade panels (Fig. 7-5). Other applications are listed in PCA (1991).

Glass Fibers

The first research on glass fibers in the early 1960s used conventional borosilicate glass (E-glass) (Table 7-1) and soda-lime-silica glass fibers (A-glass). The test results showed that alkali reactivity between the E-glass fibers and the cement-paste reduced the strength of the concrete. Continued research resulted in alkali-resistant glass fibers (AR-glass) (Table 7-1), that improved long-term durability, but sources of other strength-loss trends were observed. One acknowledged source was fiber embrittlement stemming from infiltration of calcium hydroxide particles, by-products of cement hydration, into fiber bundles. Alkali reactivity and cement hydration are the basis for the following two widely held theories explaining strength and ductility loss, particularly in exterior glass fiber concrete:

- Alkali attack on glass-fiber surfaces reduces fiber tensile strength and, subsequently, lowers compressive strength.
- Ongoing cement hydration causes calcium hydroxide particle penetration of fiber bundles, thereby increasing fiber-to-matrix bond strength and embrittlement; the latter lowers tensile strength by inhibiting fiber pullout.

Fig. 7-4. Tightly bunched steel fibers are placed in a form, before cement slurry is poured into this application of slurry-infiltrated steel-fiber concrete (SIFCON). (IMG12409)

Fig. 7-5. (top) Glass-fiber-reinforced concrete panels are light and strong enough to reduce this building’s structural requirements. (bottom) Spray-up fabrication made it easy to create their contoured profiles. (IMG12404, IMG12327)
Synthetic Fibers

Synthetic fibers are man-made fibers resulting from research and development in the petrochemical and textile industries. Fiber types that are used in Portland cement concrete are: acrylic, aramid, carbon, nylon, polyester, polyethylene, and polypropylene. Table 7-1 summarizes the range of physical properties of these fibers.

Synthetic fibers can reduce plastic shrinkage and subsidence cracking and may help concrete after it is fractured. Ultra-thin whitetopping often uses synthetic fibers for potential containment properties to delay pothole development. Problems associated with synthetic fibers include: (1) low fiber-to-matrix bonding; (2) inconclusive performance testing for low fiber-volume usage with polypropylene, polyethylene, polyester and nylon; (3) a low modulus of elasticity for polypropylene and polyethylene; and (4) the high cost of carbon and aramid fibers.

Polypropylene fibers (Fig. 7-6), the most popular of the synthetics, are chemically inert, hydrophobic, and lightweight. They are produced as continuous cylindrical monofilaments that can be chopped to specified lengths or cut as films and tapes and formed into fine fibrils of rectangular cross section (Fig. 7-7).

Used at a rate of at least 0.1 percent by volume of concrete, polypropylene fibers reduce plastic shrinkage cracking and subsidence cracking over steel reinforcement (Suprenant and Malisch 1999). The presence of polypropylene fibers in concrete may reduce settlement of aggregate particles, thus reducing capillary bleed channels. Polypropylene fibers can help reduce spalling of high-strength, low-permeability concrete exposed to fire in a moist condition.

New developments show that monofilament fibers are able to fibrillate during mixing if produced with both, polypropylene and polyethylene resins. The two polymers are incompatible and tend to separate when manipulated. Therefore, during the mixing process each fiber turns into a unit with several fibrils at its end. The fibrils provide better mechanical bonding than conventional monofilaments. The high number of fine fibrils also reduces plastic shrinkage cracking and may increase the ductility and toughness of the concrete (Trottier and Mahoney 2001).

Acrylic fibers have been found to be the most promising replacement for asbestos fibers. They are used in cement board and roof-shingle production, where fiber volumes of up to 3% can produce a composite with mechanical properties similar to that of an asbestos-cement composite. Acrylic-fiber concrete composites exhibit high postcracking toughness and ductility. Although lower than that of asbestos-cement composites, acrylic-fiber-reinforced concrete’s flexural strength is ample for many building applications.

Aramid fibers have high tensile strength and a high tensile modulus. Aramid fibers are two and a half times as strong as E-glass fibers and five times as strong as steel fibers. A comparison of mechanical properties of different aramid fibers is provided in PCA (1991). In addition to excellent strength characteristics, aramid fibers also have excellent strength retention up to 160°C (320°F), dimensional stability up to 200°C (392°F), static and dynamic fatigue resistance, and creep resistance. Aramid strand is available in a wide range of diameters.

Carbon fibers were developed primarily for their high strength and elastic modulus and stiffness properties for applications within the aerospace industry. Compared with most other synthetic fibers, the manufacture of carbon fibers is expensive and this has limited commercial development. Carbon fibers have high tensile strength and modulus of elasticity (Table 7-1). They are also inert to...
Natural fibers were used as a form of reinforcement long before the advent of conventional reinforced concrete. Mud bricks reinforced with straw and mortars reinforced with horsehair are just a few examples of how natural fibers were used long ago as a form of reinforcement. Many natural reinforcing materials can be obtained at low levels of cost and energy using locally available manpower and technical know-how. Such fibers are used in the manufacture of low-fiber-content concrete and occasionally have been used in thin-sheet concrete with high-fiber content. For typical properties of natural fibers see Table 7-1.

Unprocessed Natural Fibers. In the late 1960s, research on the engineering properties of natural fibers, and concrete made with these fibers was undertaken; the result was these fibers can be used successfully to make thin sheets for walls and roofs. Products were made with portland cement and unprocessed natural fibers such as coconut coir, sisal, bamboo, jute, wood, and vegetable fibers. Although the concretes made with unprocessed natural fibers show good mechanical properties, they have some deficiencies in durability. Many of the natural fibers are highly susceptible to volume changes due to variations in fiber moisture content. Fiber volumetric changes that accompany variations in fiber moisture content can drastically affect the bond strength between the fiber and cement matrix.

Wood Fibers (Processed Natural Fibers). The properties of wood cellulose fibers are greatly influenced by the method by which the fibers are extracted and the refining processes involved. The process by which wood is reduced to a fibrous mass is called pulping. The kraft process is the one most commonly used for producing wood cellulose fibers. This process involves cooking wood chips in a solution of sodium hydroxide, sodium carbonate, and sodium sulfide. Wood cellulose fibers have relatively good mechanical properties compared to many manmade fibers such as polypropylene, polyethylene, polyester, and acrylic. Delignified cellulose fibers (lignin removed) can be produced with a tensile strength of up to approximately 2000 MPa (290 ksi) for selected grades of wood and pulping processes. Fiber tensile strength of approximately 500 MPa (73 ksi) can be routinely achieved using a chemical pulping process and the more common, less expensive grades of wood.

MULTIPLE FIBER SYSTEMS

For a multiple fiber system, two or more fibers are blended into one system. The hybrid-fiber concrete combines macro- and microsteel fibers. A common macrofibre blended with a newly developed microfiber, which is less than 10 mm (0.4 in.) long and less than 100 micrometer (0.004 in.) in diameter, leads to a closer fiber-to-fiber spacing, which reduces microcracking and increases tensile strength. The intended applications include thin repairs and patching (Banthia and Bindiganavile 2001). A blend of steel and polypropylene fibers has also been used for some applications. This system is supposed to combine the toughness and impact-resistance of steel fiber concrete with the reduced plastic cracking of polypropylene fiber concrete. For a project in the Chicago area (Wójtyśiak, Borden and Harrison, 2001), a blend of 30 kg/m³ (50 lb/yd³) of steel fibers and 0.9 kg/m³ (1½ lb/yd³) of fib-
rillated polypropylene fibers were used for slabs on grade. The concrete with blended fibers had a lower slump compared to plain concrete but seemed to have enhanced elastic and post-elastic strength.

REFERENCES


Banthia, Nemkumar, and Bindiganavile, Vivek, “Repairing with Hybrid-Fiber-Reinforced Concrete,” *Concrete International*, American Concrete Institute, Farmington Hills, Michigan, June 2001, pages 29 to 32.


Shah, S. P.; Weiss, W. J.; and Yang, W., “Shrinkage Cracking – Can it be prevented?,” *Concrete International*, American Concrete Institute, Farmington Hills, Michigan, April 1998, pages 51 to 55.


CHAPTER 8
Air-Entrained Concrete

One of the greatest advances in concrete technology was the development of air-entrained concrete in the mid-1930s. Today air entrainment is recommended for nearly all concretes, principally to improve freeze-thaw resistance when exposed to water and deicing chemicals. However, there are other important benefits of entrained air in both freshly mixed and hardened concrete.

Air-entrained concrete is produced by using either an air-entraining cement or adding an air-entraining admixture during batching. The air-entraining admixture stabilizes bubbles formed during the mixing process, enhances the incorporation of bubbles of various sizes by lowering the surface tension of the mixing water, impedes bubble coalescence, and anchors bubbles to cement and aggregate particles.

Anionic air-entraining admixtures are hydrophobic (repel water) and are electrically charged (nonionic or no-charge admixtures are also available). The negative electric charge is attracted to positively charged cement grains, which aids in stabilizing bubbles. The air-entraining admixture forms a tough, water-repelling film, similar to a soap film, with sufficient strength and elasticity to contain and stabilize the air bubbles and prevent them from coalescing. The hydrophobic film also keeps water out of the bubbles. The stirring and kneading action of mechanical mixing disperses the air bubbles. The fine aggregate particles also act as a three-dimensional grid to help hold the bubbles in the mixture.

Entrained air bubbles are not like entrapped air voids, which occur in all concretes as a result of mixing, handling, and placing and are largely a function of aggregate characteristics. Intentionally entrained air bubbles are extremely small in size, between 10 to 1000 µm in diameter, while entrapped voids are usually 1000 µm (1 mm) or larger. The majority of the entrained air voids in normal concrete are between 10 µm and 100 µm in diameter. As shown in Fig. 8-1, the bubbles are not interconnected; they are well dispersed and randomly distributed. Non-air-entrained concrete with a 25-mm (1-in.) maximum-size aggregate has an air content of approximately 1 1⁄2%. This same mixture air entrained for severe frost exposure would require a total air content of about 6%, made up of both the coarser “entrapped” air voids and the finer “entrained” air voids.

PROPERTIES OF AIR-ENTRAINED CONCRETE

The primary concrete properties influenced by air entrainment are presented in the following sections. A brief summary of other properties not discussed below is presented in Table 8-1.

Freeze-Thaw Resistance

The resistance of hardened concrete to freezing and thawing in a moist condition is significantly improved by the use of intentionally entrained air, even when various deicers are involved. Convincing proof of the improve-
Table 8-1. Effect of Entrained Air on Concrete Properties

<table>
<thead>
<tr>
<th>Properties</th>
<th>Effect</th>
</tr>
</thead>
<tbody>
<tr>
<td>Abrasion</td>
<td>Little effect; increased strength increases abrasion resistance</td>
</tr>
<tr>
<td>Absorption</td>
<td>Little effect</td>
</tr>
<tr>
<td>Alkali-silica reactivity</td>
<td>Expansion decreases with increased air</td>
</tr>
<tr>
<td>Bleeding</td>
<td>Reduced significantly</td>
</tr>
<tr>
<td>Bond to steel</td>
<td>Decreased</td>
</tr>
<tr>
<td>Compressive strength</td>
<td>Reduced approximately 2% to 6% per percentage point increase in air; harsh or lean mixes may gain strength</td>
</tr>
<tr>
<td>Creep</td>
<td>Little effect</td>
</tr>
<tr>
<td>Deicer scaling</td>
<td>Significantly reduced</td>
</tr>
<tr>
<td>Density</td>
<td>Decreases with increased air</td>
</tr>
<tr>
<td>Fatigue</td>
<td>Little effect</td>
</tr>
<tr>
<td>Finishability</td>
<td>Reduced due to increased cohesion (stickiness)</td>
</tr>
<tr>
<td>Flexural strength</td>
<td>Reduced approximately 2% to 4% per percentage point increase in air</td>
</tr>
<tr>
<td>Freeze-thaw resistance</td>
<td>Significantly improved resistance to water-saturated freeze-thaw deterioration</td>
</tr>
<tr>
<td>Heat of hydration</td>
<td>No significant effect</td>
</tr>
<tr>
<td>Modulus of elasticity</td>
<td>Decreases with increased air approximately 720 to 1380 MPa (105,000 to 200,000 psi) per percentage point of air</td>
</tr>
<tr>
<td>Permeability</td>
<td>Little effect; reduced water-cement ratio reduces permeability</td>
</tr>
<tr>
<td>Scaling</td>
<td>Significantly reduced</td>
</tr>
<tr>
<td>Shrinkage (drying)</td>
<td>Little effect</td>
</tr>
<tr>
<td>Slump</td>
<td>Increases with increased air approximately 25 mm (1 in.) per ½ to 1 percentage point increase in air</td>
</tr>
<tr>
<td>Specific heat</td>
<td>No effect</td>
</tr>
<tr>
<td>Sulfate resistance</td>
<td>Significantly improved</td>
</tr>
<tr>
<td>Stickiness</td>
<td>Increased cohesion—harder to finish</td>
</tr>
<tr>
<td>Temperature of wet concrete</td>
<td>No effect</td>
</tr>
<tr>
<td>Thermal conductivity</td>
<td>Decreases 1% to 3% per percentage point increase in air</td>
</tr>
<tr>
<td>Thermal diffusivity</td>
<td>Decreases about 1.6% per percentage point increase in air</td>
</tr>
<tr>
<td>Water demand of wet concrete for equal slump</td>
<td>Decreases with increased air; approximately 3 to 6 kg/m³ (5 to 10 lb/yd³) per percentage point of air</td>
</tr>
<tr>
<td>Watertightness</td>
<td>Increases slightly; reduced water-cement ratio increases watertightness</td>
</tr>
<tr>
<td>Workability</td>
<td>Increases with increased air</td>
</tr>
</tbody>
</table>

Note: The table information may not apply to all situations.

Fig. 8-2. Effect of entrained air on the resistance of concrete to freezing and thawing in laboratory tests. Concretes were made with cements of different fineness and composition and with various cement contents and water-cement ratios (Bates and others 1952, and Lerch 1960).

As the water in moist concrete freezes, it produces osmotic and hydraulic pressures in the capillaries and pores of the cement paste and aggregate. If the pressure exceeds the tensile strength of the paste or aggregate, the cavity will dilate and rupture. The accumulative effect of successive freeze-thaw cycles and disruption of paste and aggregate eventually cause significant expansion and deterioration of the concrete. Deterioration is visible in the form of cracking, scaling, and crumbling (Fig. 8-3). Powers (1965) and Pigeon and Pleau (1995) extensively review the mechanisms of frost action.

Hydraulic pressures are caused by the 9% expansion of water upon freezing; in this process growing ice crystals displace unfrozen water. If a capillary is above critical saturation (91.7% filled with water), hydraulic pressures result as freezing progresses. At lower water contents, no hydraulic pressure should exist.

Osmotic pressures develop from differential concentrations of alkali solutions in the paste (Powers 1965a). As pure water freezes, the alkali concentration increases in the adjacent unfrozen water. A high-alkali solution, through the mechanism of osmosis, draws water from lower-alkali solutions in the pores. This osmotic transfer of water continues until equilibrium in the fluids’ alkali concentration is achieved. Osmotic pressure is considered a minor factor, if present at all, in aggregate frost action, whereas it may be dominant in certain cement pastes. Osmotic pressures, as described above, are considered to be a major factor in “salt scaling.”

Capillary ice (or any ice in large voids or cracks) draws water from pores to advance its growth. Also, since most pores in cement paste and some aggregates are too
small for ice crystals to form, water attempts to migrate to locations where it can freeze.

Entrained air voids act as empty chambers in the paste where freezing and migrating water can enter, thus relieving the pressures described above and preventing damage to the concrete. Upon thawing, most of the water returns to the capillaries due to capillary action and pressure from air compressed in the bubbles. Thus the bubbles are ready to protect the concrete from the next cycle of freezing and thawing (Powers 1955, Lerch 1960, and Powers 1965).

The pressure developed by water as it expands during freezing depends largely upon the distance the water must travel to the nearest air void for relief. Therefore, the voids must be spaced close enough to reduce the pressure below that which would exceed the tensile strength of the concrete. The amount of hydraulic pressure is also related to the rate of freezing and the permeability of the paste.

The spacing and size of air voids are important factors contributing to the effectiveness of air entrainment in concrete. ASTM C457 describes a method of evaluating the air-void system in hardened concrete. Most authorities consider the following air-void characteristics as representative of a system with adequate freeze-thaw resistance (Powers 1949, Klieger 1952, Klieger 1956, Mielenz and others 1958, Powers 1965, Klieger 1966, Whiting and Nagi 1998, and Pinto and Hover 2001):

1. Calculated spacing factor, \( \bar{L} \), (an index related to the distance between bubbles but not the actual average spacing in the system)—less than 0.200 mm (0.008 in.) (Powers 1954 and 1965)
2. Specific surface, \( \alpha \) (surface area of the air voids)—24 square millimeters per cubic millimeter (600 sq in. per cubic inch) of air-void volume, or greater.

Current U.S. field quality control practice usually involves only the measurement of total air volume in freshly mixed concrete; this does not distinguish air-void size in any way. In addition to total air volume, Canadian practice also requires attainment of specific spacing factors. Fig. 8-4 illustrates the relationship between spacing factor and total air content. Measurement of air volume alone does not permit full evaluation of the important characteristics of the air-void system; however, air-entrainment is generally considered effective for freeze-thaw resistance when the volume of air in the mortar fraction of the concrete—material passing the 4.75-mm (No. 4) sieve—is about 9 ± 1% (Klieger 1952) or about 18% by paste volume. For equal admixture dosage rates per unit of cement, the air content of ASTM C185 (AASHTO T 137) mortar would be about 19% due to the standard aggregate’s properties.

Fig. 8-3. Effect of weathering on boxes and slabs on ground at the Long-Time Study outdoor test plot, Project 10, PCA, Skokie, Illinois. Specimens at top are air-entrained, specimens at bottom exhibiting severe crumbling and scaling are non-air-entrained. All concretes were made with 335 kg (564 lb) of Type I portland cement per cubic meter (cubic yard). Periodically, calcium chloride deicer was applied to the slabs. Specimens were 40 years old when photographed (see Klieger 1963 for concrete mixture information). (IMG12408, IMG12406, IMG12407, IMG12405)
the effect of water-cement ratio on the durability of non-air-entrained concrete.

Concrete elements should be properly drained and kept as dry as possible as greater degrees of saturation increase the likelihood of distress due to freeze-thaw cycles. Concrete that is dry or contains only a small amount of moisture in service is essentially not affected by even a large number of cycles of freezing and thawing. Refer to the sections on “Deicer-Scaling Resistance” and “Recommended Air Contents” in this chapter and to Chapter 9 for mixture design considerations.

Deicer-Scaling Resistance

Deicing chemicals used for snow and ice removal can cause and aggravate surface scaling. The damage is primarily a physical action. Deicer scaling of inadequately air-entrained or non-air-entrained concrete during freezing is believed to be caused by a buildup of osmotic and hydraulic pressures in excess of the normal hydraulic pressures produced when water in concrete freezes. These pressures become critical and result in scaling unless entrained air voids are present at the surface and throughout the sample to relieve the pressure. The hygroscopic (moisture absorbing) properties of deicing salts also attract water and keep the concrete more saturated, increasing the potential for freeze-thaw deterioration. However, properly designed and placed air-entrained concrete will withstand deicers for many years.

Studies have also shown that, in absence of freezing, the formation of salt crystals in concrete (from external sources of chloride, sulfate, and other salts) may contribute to concrete scaling and deterioration similar to the crumbling of rocks by salt weathering. The entrained air voids in concrete allow space for salt crystals to grow; this relieves internal stress similar to the way the voids relieve stress from freezing water in concrete (ASCE 1982 and Sayward 1984).

Deicers can have many effects on concrete and the immediate environment. All deicers can aggravate surface scaling of concrete that is not properly air entrained. Sodium chloride (rock salt) (ASTM D632 or AASHTO M 143), and calcium chloride (ASTM D98 or AASHTO M 144) are the most frequently used deicers. In the absence of freezing, sodium chloride has little to no chemical effect on concrete but can damage plants and corrode metal. Calcium chloride in weak solutions generally has little chemical effect on concrete and vegetation but does corrode metal.

Studies have shown that concentrated calcium chloride solutions can chemically attack concrete (Brown and Cady 1975). While there is disagreement in the laboratory studies about the effect of magnesium chloride on concrete deterioration (Leiser and Dombrowski 1967; Cody at al. 2006; Lee at al. 2000; and Kozikowski et al. 2007), there has not been any observed damage of field concrete that can be attributed to magnesium chloride use (NCHRP 2007). Urea does not chemically damage concrete, vegetation, or metal. Nonchloride deicers are used to minimize corrosion of rein-

The air content of concrete with 19-mm (3/4-in.) maximum-size aggregate would be about 6% for effective freeze-thaw resistance.

The relationship between air content of standard mortar and concrete is illustrated by Taylor (1948). Pinto and Hover (2001) address paste air content versus frost resistance. The total required concrete air content for durability increases as the maximum-size aggregate is reduced (due to greater paste volume) and the exposure conditions become more severe (see “Recommended Air Contents” later in this chapter).

Freeze-thaw resistance is also significantly increased with the use of the following: (1) a good quality aggregate, (2) a low water to cementing materials ratio (maximum 0.45), (3) a minimum cementitious materials content of 335 kg/m³ (564 lb/ yd³), (4) proper finishing and curing techniques, and (5) a compressive strength of 28 MPa (4,000 psi) when exposed to repeated freeze-thaw cycles. Even non-air-entrained concretes will be more freeze-thaw resistant with a low water-cement ratio. Fig. 8-5 illustrates the relationship between air content and frost resistance.

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Fig. 8-4. Spacing factor as a function of total air content in concrete (Pinto and Hover 2001).

Fig. 8-5. Durability factors vs. number of freeze-thaw cycles for selected non-air-entrained concretes (Pinto and Hover 2001).
forcing steel and minimize groundwater chloride contamination. The use of deicers containing ammonium nitrate and ammonium sulfate should be strictly prohibited as they rapidly attack and disintegrate concrete.

The extent of scaling depends upon the amount of deicer used and the frequency of application. Relatively low concentrations of deicer (on the order of 2% to 4% by mass) produce more surface scaling than higher concentrations or the absence of deicer (Verbeck and Klieger 1956).

Deicers can reach concrete surfaces in ways other than direct application, such as splashing by vehicles and dripping from the undersides of vehicles. Scaling is more severe in poorly drained areas because more of the deicer solution remains on the concrete surface during freezing and thawing. Air entrainment is effective in preventing surface scaling and is recommended for all concretes that may come in contact with deicing chemicals (Fig. 8-6).

A good air-void system with a low spacing factor (maximum of 200 micrometers) is perhaps more important to deicer environments than saturated frost environments without deicers. The relationship between spacing factor and deicer scaling is illustrated in Fig. 8-7. A low water to portland cement ratio helps minimize scaling, but is not sufficient to control scaling at normal water-cement ratios.

Fig. 8-6. Cumulative mass loss for mixtures with a water to cement ratio of 0.45 and on-time finishing (Pinto and Hover 2001).

Fig. 8-7. Visual rating as a function of spacing factor, for a concrete mixture with a water to cement ratio of 0.45 (Pinto and Hover 2001).

Fig. 8-8 illustrates the overriding impact of air content over water-cement ratio in controlling scaling.

To provide adequate durability and scale resistance in severe exposures with deicers present, air-entrained concrete should be composed of durable materials and have the following: (1) a low water to cementitious materials ratio (maximum 0.45), (2) a slump of 100 mm (4 in.) or less unless a plasticizer is used, (3) a cementitious materials content of 335 kg/m³ (564 lb/yd³), (4) proper finishing after bleed water has evaporated from the surface, (5) adequate drainage, (6) a minimum of 7 days moist curing at or above 10°C (50°F), (7) a compressive strength of 28 MPa (4000 psi) when exposed to repeated freeze-thaw cycling, and (8) a minimum 30-day drying period after moist curing if concrete will be exposed to freeze-thaw cycles and deicers when saturated. Target air contents are discussed in “Recommended Air Contents” at the end of this chapter.

Normal dosages of supplementary cementitious materials should not effect scaling resistance of properly designed, placed, and cured concrete (Table 8-2). The ACI 318 building code allows up to 10% silica fume, 25% fly ash, and 50% slag as part of the cementitious materials for deicer exposures. However, abuse of these materials along

<table>
<thead>
<tr>
<th>Table 8-2. Deicer Scaling Resistance (Visual Ratings) of Concrete with Selected Supplementary Cementing Materials</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Mass replacement</strong></td>
</tr>
<tr>
<td>of cement, %</td>
</tr>
<tr>
<td>Scale rating at 25 cycles</td>
</tr>
<tr>
<td>Scale rating at 50 cycles</td>
</tr>
</tbody>
</table>

Concrete had 335 kg of cementing materials per cubic meter (565 lb/yd³), a Type I portland cement, a water to cementing materials ratio of 0.50, a nominal slump of 75 mm (3 in.), and a nominal air content of 6%. Test method: ASTM C672. Results are for specific materials tested in 2000 and may not be representative of other materials. Scale rating: 1 = very slight scaling (3 mm depth maximum) with no coarse aggregate visible, 2 = slight to moderate scaling.
The effect of mix design, surface treatment, curing, or other variables on resistance to surface scaling can be evaluated by ASTM C672.

**Sulfate Resistance**

Sulfate resistance of concrete is improved by air entrainment, as shown in Figs. 8-9 and 8-10, when advantage is taken of the reduction in water-cement ratio possible with air entrainment. Air-entrained concrete made with a low water-cement ratio, an adequate cement content and a sulfate-resistant cement will be resistant to attack from sulfate soils and waters.

**Resistance to Alkali-Silica Reactivity**

The expansive disruption caused by alkali-silica reactivity is reduced through the use of air-entrainment (Kretzinger 1949). Alkali hydroxides react with the silica of reactive aggregates to form expansive reaction products, causing the concrete to expand. Excessive expansion will disrupt and deteriorate concrete. As shown in Fig. 8-11, the expansion of mortar bars made with reactive materials is reduced as the air content is increased.

**Strength**

When the air content is maintained constant, strength varies inversely with the water-cement ratio. Fig. 8-12 shows a typical relationship between 28-day compressive strength and water-cement ratio for concrete that has the recommended percentages of entrained air. As air content is increased, a given strength generally can be maintained by holding the voids (air + water) to cement ratio constant; this may, however, necessitate some increase in cement content.

Air-entrained as well as non-air-entrained concrete can readily be proportioned to provide similar moderate strengths. Both generally must contain the same amount

![Fig. 8-8. Measured mass loss of concrete after 40 cycles of deicer and frost exposure at various water to cement ratios (Pinto and Hover 2001).](image)

![Fig. 8-9. Effect of entrained air and cement content (Type II) on performance of concrete specimens exposed to a sulfate soil. Without entrained air the specimens made with lesser amounts of cement deteriorated badly. Specimens made with the most cement and the lowest water-cement ratio were further improved by air entrainment. Specimens were 5 years old when photographed (Stanton 1948 and Lerch 1960).](image)
Chapter 8 • Air-Entrained Concrete

Air-entrained concrete
Cement: Type I
Age: 28 days

Fig. 8-10. Performance of a variety of air-entrained and non-air-entrained concretes exposed to sulfate soil. Sulfate resistance is increased with the use of Types II and V cements, a higher cement content, lower water-cement ratio, and air entrainment (Stark 1984).

Fig. 8-11. Effect of air content on the reduction of expansion due to alkali-silica reaction (Kretsinger 1949).

of coarse aggregate. When the cement content and slump are held constant, air entrainment reduces the sand and water requirements as illustrated in Fig. 8-13. Thus, air-entrained concretes can have lower water-cement ratios than non-air-entrained concretes; this minimizes the reductions in strength that generally accompany air entrainment. At constant water-cement ratios, increases in air will proportionally reduce strength (Fig. 8-14). Pinto and Hover (2001) found that for a decrease in strength of 10 MPa (1450 psi), resulting from a four percentage point decrease in air, the water-cement ratio had to be decreased by 0.14 to maintain strength. Some reductions in strength may be tolerable in view of other benefits of air, such as improved workability. Reductions in strength become more significant in higher-strength mixes, as illustrated in Fig. 8-15. In lower-cement-content, harsh mixes, strength is generally increased by entrainment of air in proper amounts due to the reduced water-cement ratio and improved workability. For moderate to high-strength concrete, each percentile of increase in entrained air reduces the compressive strength about 2% to 9% (Cordon 1946, Klieger 1952, Klieger 1956, Whiting and Nagi 1998, and Pinto and Hover 2001). Actual strength varies and is affected by the cement source, admixtures, and other concrete ingredients.

Attainment of high strength with air-entrained concrete may be difficult at times. Even though a reduction in mixing water is associated with air entrainment, mixtures with high cement contents require more mixing water than lower-cement-content mixtures; hence, the increase in strength expected from the additional cement is offset somewhat by the additional water. Water reducers can offset this effect.
Workability

Entrained air improves the workability of concrete. It is particularly effective in lean (low cement content) mixes that otherwise might be harsh and difficult to work. In one study, an air-entrained mixture made with natural aggregate, 3% air, and a 37-mm (1 1⁄2-in.) slump had about the same workability as a non-air-entrained concrete with 1% air and a 75-mm (3-in.) slump, even though less cement was required for the air-entrained mix (Cordon 1946).

Because of improved workability with entrained air, water and sand content can be reduced significantly (Fig. 8-13). A volume of air-entrained concrete requires less water than an equal volume of non-air-entrained concrete of the same consistency and maximum size aggregate. Freshly mixed concrete containing entrained air is cohesive, looks and feels fatty or workable, and can usually be handled with ease; on the other hand, high air contents can make a mixture sticky and more difficult to finish. Entrained air also reduces segregation and bleeding in freshly mixed and placed concrete.

AIR-ENTRAINING MATERIALS

The entrainment of air in concrete can be accomplished by adding an air-entraining admixture at the mixer, by using an air-entraining cement, or by a combination of these
methods. Regardless of the method used, adequate control and monitoring is required to ensure the proper air content at all times.

Numerous commercial air-entraining admixtures, manufactured from a variety of materials, are available. Most air-entraining admixtures consist of one or more of the following materials: wood resin (Vinsol resin), sulfonated hydrocarbons, fatty and resinous acids, and synthetic materials. Chemical descriptions and performance characteristics of common air-entraining agents are shown in Table 8-3. Air-entraining admixtures are usually liquids and should not be allowed to freeze. Admixtures added at the mixer should conform to ASTM C260 (AASHTO M 154).

Air-entraining cements comply with ASTM C150 and C595 (AASHTO M 85 and M 240). To produce such cements, air-entraining additions conforming to ASTM C226 are interground with the cement clinker during manufacture. Air-entraining cements generally provide an adequate amount of entrained air to meet most job conditions; however, a specified air content may not necessarily be obtained in the concrete. If an insufficient volume of air is entrained, it may also be necessary to add an air-entraining admixture at the mixer.

Each method of entraining air has certain advantages. On jobs where careful control is not practical, air-entraining cements are especially useful to ensure that a significant portion of the required air content will always be obtained. They eliminate the possibility of human or mechanical error that can occur when adding an admixture during batching. With air-entraining admixtures, the volume of entrained air can be readily adjusted to meet job conditions by changing the amount of admixture added at the mixer. Variations in air content can be expected with variations in aggregate proportions and gradation, mixing time, temperature, and slump. The order of batching and mixing concrete ingredients when using an air-entraining admixture has a significant influence on the amount of air entrained; therefore, consistency in batching is needed to maintain adequate control.

When entrained air is excessive, it can be reduced by using one of the following defoaming (air-detraining) agents: tributyl phosphate, dibutyl phthalate, octyl alcohol, water-insoluble esters of carbonic acid and boric acid, and silicones. Only the smallest possible dosage of defoaming agent should be used to reduce the air content to the specified limit. An excessive amount might have adverse effects on concrete properties (Whiting and Stark 1983).

**FACTORS AFFECTING AIR CONTENT**

**Cement**

As cement content increases, the air content decreases for a set dosage of air-entraining admixture per unit of cement within the normal range of cement contents (Fig. 8-16). In going from 240 to 360 kilogram of cement per cubic meter (400 to 600 lb of cement per cubic yard), the dosage rate may have to be doubled to maintain a constant air content. However, studies indicate that when this is done the air-void spacing factor generally decreases with an increase in

<table>
<thead>
<tr>
<th>Classification</th>
<th>Chemical description</th>
<th>Notes and performance characteristics</th>
</tr>
</thead>
<tbody>
<tr>
<td>Wood derived acid salts</td>
<td>Alkali or alkanolamine salt of: A mixture of tricyclic acids, phenolics, and terpenes.</td>
<td>Quick air generation. Minor air gain with initial mixing. Air loss with prolonged mixing. Mid-sized air bubbles formed. Compatible with most other admixtures.</td>
</tr>
<tr>
<td>Vinsol® resin</td>
<td>Tricyclic acids-major component. Tricyclic acids-minor component.</td>
<td>Same as above.</td>
</tr>
<tr>
<td>Wood rosin</td>
<td>Tricyclic acids-major component. Fatty acids-major component. Tricyclic acids-minor component.</td>
<td>Slower air generation. Air may increase with prolonged mixing. Smallest air bubbles of all agents. Compatible with most other admixtures.</td>
</tr>
<tr>
<td>Tall oil</td>
<td>Coconut fatty acids, alkanolamine salt.</td>
<td>Slower air generation than wood rosins. Moderate air loss with mixing. Coarser air bubbles relative to wood rosins. Compatible with most other admixtures.</td>
</tr>
<tr>
<td>Vegetable oil acids</td>
<td>Alkyl-arylsulfonates and sulfates (e.g., sodium dodecylbenzenesulfonate).</td>
<td>Quick air generation. Minor air loss with mixing. Coarser bubbles. May be incompatible with some HRWR. Also applicable to cellular concretes.</td>
</tr>
<tr>
<td>Synthetic detergents</td>
<td>All these are rarely used as concrete air-entraining agents in current practice.</td>
<td></td>
</tr>
<tr>
<td>Synthetic workability aids</td>
<td>All these are rarely used as concrete air-entraining agents in current practice.</td>
<td></td>
</tr>
</tbody>
</table>

---

**Table 8-3. Classification and Performance Characteristics of Common Air-Entraining Admixtures**

(Whiting and Nagi 1998)
the amount of fine aggregate causes more air to be entrained for a given amount of air-entraining cement or admixture (more air is also entrapped in non-air-entrained concrete).

Fine aggregate particles passing the 600 µm to 150 µm (No. 30 to No. 100) sieves entrap more air than either very fine or coarser particles. Appreciable amounts of material passing the 150 µm (No. 100) sieve will result in a significant reduction of entrained air.

Fine aggregates from different sources may entrap different amounts of air even though they have identical gradations. This may be due to differences in shape and surface texture or contamination by organic materials.

Mixing Water and Slump

An increase in the mixing water makes more water available for the generation of air bubbles, thereby increasing the air content as slumps increase up to about 150 or 175 mm (6 or 7 inches). An increase in the water-cement ratio from 0.4 to 1.0 can increase the air content by four percentage points. A portion of the air increase is due to the relationship between slump and air content. Air content increases with slump even when the water-cement ratio is held constant. The spacing factor, $L$, of the air-void system also increases, that is, the voids become coarser at higher water-cement ratios, thereby reducing concrete freeze-thaw durability (Stark 1986).

The addition of 5 kg of water per cubic meter (8.4 lbs of water per cubic yard) of concrete can increase the slump by 25 mm (one inch). A 25-mm (1-in.) increase in slump increases the air content by approximately one-half to one cement content; and for a given air content the specific surface increases, thus improving durability.

An increase in cement fineness will result in a decrease in the amount of air entrained. Type III cement, a very finely ground material, may require twice as much air-entraining agent as a Type I cement of normal fineness.

High-alkali cements may entrain more air than low alkali cements with the same amount of air-entraining material. A low-alkali cement may require 20% to 40% (occasionally up to 70%) more air-entraining agent than a high-alkali cement to achieve an equivalent air content. Precautions are therefore necessary when using more than one cement source in a batch plant to ensure that proper admixture requirements are determined for each cement (Greening 1967).

Coarse Aggregate

The size of coarse aggregate has a pronounced effect on the air content of both air-entrained and non-air-entrained concrete, as shown in Fig. 8-16. There is little change in air content when the size of aggregate is increased above 37.5 mm (1½ in.).

Fine Aggregate

The fine-aggregate content of a mixture affects the percentage of entrained air. As shown in Fig. 8-17, increasing the amount of fine aggregate causes more air to be entrained for a given amount of air-entraining cement or admixture (more air is also entrapped in non-air-entrained concrete).

Fine-aggregate particles passing the 600 µm to 150 µm (No. 30 to No. 100) sieves entrap more air than either very fine or coarser particles. Appreciable amounts of material passing the 150 µm (No. 100) sieve will result in a significant reduction of entrained air.

Fine aggregates from different sources may entrap different amounts of air even though they have identical gradations. This may be due to differences in shape and surface texture or contamination by organic materials.
percentage point for concretes with a low-to-moderate slump and constant air-entraining admixture dosage. However, this approximation is greatly affected by concrete temperature, slump, and the type and amount of cement and admixtures present in the concrete. A low-slump concrete with a high dosage of water-reducing and air-entraining admixtures can undergo large increases in slump and air content with a small addition of water. On the other hand, a very fluid concrete mixture with a 200 to 250-mm (8 to 10-in.) slump may lose air with the addition of water. Refer to Tables 8-4 and 8-5 for more information.

The mixing water used may also affect air content. Algae-contaminated water increases air content. Highly alkaline wash water from truck mixers can affect air con-

### Table 8-4. Effect of Mixture Design and Concrete Constituents on Control of Air Content in Concrete

<table>
<thead>
<tr>
<th>Characteristic/Material</th>
<th>Effects</th>
<th>Guidance</th>
</tr>
</thead>
<tbody>
<tr>
<td>Alkali content</td>
<td>Air content increases with increase in cement alkali level. Less air-entraining agent dosage needed for high-alkali cements. Air-void system may be more unstable with some combinations of alkali level and air-entraining agent used.</td>
<td>Changes in alkali content or cement source require that air-entraining agent dosage be adjusted. Decrease dosage as much as 40% for high-alkali cements.</td>
</tr>
<tr>
<td>Fineness</td>
<td>Decrease in air content with increased fineness of cement.</td>
<td>Use up to 100% more air-entraining admixture for very fine (Type III) cements. Adjust admixture if cement source or fineness changes.</td>
</tr>
<tr>
<td>Cement content in mixture</td>
<td>Decrease in air content with increase in cement content. Smaller and greater number of voids with increased cement content.</td>
<td>Increase air-entraining admixture dosage rate as cement content increases.</td>
</tr>
<tr>
<td>Contaminants</td>
<td>Air content may be altered by contamination of cement with finish mill oil.</td>
<td>Verify that cement meets ASTM C150 (AASHTO M 85) requirements on air content of test mortar.</td>
</tr>
<tr>
<td>Fly ash</td>
<td>Air content decreases with increase in loss on ignition (carbon content). Air-void system may be more unstable with some combinations of fly ash/cement/air-entraining agents.</td>
<td>Changes in LOI or fly ash source require that air-entraining admixture dosage be adjusted. Perform “foam index” test to estimate increase in dosage. Prepare trial mixes and evaluate air-void systems.</td>
</tr>
<tr>
<td>Ground granulated blast-furnace slag</td>
<td>Decrease in air content with increased fineness of GGBFS.</td>
<td>Use up to 100% more air-entraining admixture for finely ground slags.</td>
</tr>
<tr>
<td>Silica fume</td>
<td>Decrease in air content with increase in silica fume content.</td>
<td>Increase air-entraining admixture dosage up to 100% for fume contents up to 10%.</td>
</tr>
<tr>
<td>Metakaolin</td>
<td>No apparent effect.</td>
<td>Adjust air-entraining admixture dosage if needed.</td>
</tr>
<tr>
<td>Water reducers</td>
<td>Air content increases with increases in dosage of lignin-based materials. Spacing factors may increase when water-reducers used.</td>
<td>Reduce dosage of air-entraining admixture. Select formulations containing air-detraining agents. Prepare trial mixes and evaluate air-void systems.</td>
</tr>
<tr>
<td>Retarders</td>
<td>Effects similar to water-reducers.</td>
<td>Adjust air-entraining admixture dosage.</td>
</tr>
<tr>
<td>Accelerators</td>
<td>Minor effects on air content.</td>
<td>No adjustments normally needed.</td>
</tr>
<tr>
<td>High-range water reducers (Plasticizers)</td>
<td>Moderate increase in air content when formulated with lignosulfonate. Spacing factors increase.</td>
<td>Only slight adjustments needed. No significant effect on durability.</td>
</tr>
<tr>
<td>Maximum size</td>
<td>Air content requirement decreases with increase in maximum size. Little increase over 37.5 mm (1½ in.) maximum size aggregate.</td>
<td>Decrease air content.</td>
</tr>
<tr>
<td>Sand-to-total aggregate ratio</td>
<td>Air content increases with increased sand content.</td>
<td>Decrease air-entraining admixture dosage for mixtures having higher sand contents.</td>
</tr>
<tr>
<td>Sand grading</td>
<td>Middle fractions of sand promote air-entrainment.</td>
<td>Monitor gradation and adjust air-entraining admixture dosage accordingly.</td>
</tr>
</tbody>
</table>
### Table 8-4. Effect of Mixture Design and Concrete Constituents on Control of Air Content in Concrete (Continued)

<table>
<thead>
<tr>
<th>Characteristic/Material</th>
<th>Effects</th>
<th>Guidance</th>
</tr>
</thead>
<tbody>
<tr>
<td>Water chemistry</td>
<td>Very hard water reduces air content. Batching of admixture into concrete wash water decreases air. Algae growth may increase air.</td>
<td>Increase air entrainer dosage. Avoid batching into wash water.</td>
</tr>
<tr>
<td>Water-to-cement ratio</td>
<td>Air content increases with increased water to cement ratio.</td>
<td>Decrease air-entraining admixture dosage as water to cement ratio increases.</td>
</tr>
<tr>
<td>Slump</td>
<td>Air increases with slumps up to about 150 mm (6 in.). Air decreases with very high slumps. Difficult to entrain air in low-slump concretes.</td>
<td>Adjust air-entraining admixture dosages for slump. Avoid addition of water to achieve high-slump concrete. Use additional air-entraining admixture; up to ten times normal dosage.</td>
</tr>
</tbody>
</table>

### Table 8-5. Effect of Production Procedures, Construction Practices, and Environment on Control of Air Content in Concrete

<table>
<thead>
<tr>
<th>Procedure/Variable</th>
<th>Effects</th>
<th>Guidance</th>
</tr>
</thead>
<tbody>
<tr>
<td>Batching sequence</td>
<td>Simultaneous batching lowers air content. Cement-first raises air content.</td>
<td>Add air-entraining admixture with initial water or sand.</td>
</tr>
<tr>
<td>Mixer capacity</td>
<td>Air increases as capacity is approached.</td>
<td>Run mixer close to full capacity. Avoid overloading.</td>
</tr>
<tr>
<td>Mixing time</td>
<td>Central mixers: air content increases up to 90 sec. of mixing. Truck mixers: air content increases with mixing. Short mixing periods (30 seconds) reduce air content and adversely affect air-void system.</td>
<td>Establish optimum mixing time for particular mixer. Avoid overmixing. Establish optimum mixing time (about 60 seconds).</td>
</tr>
<tr>
<td>Mixing speed</td>
<td>Air content gradually increases up to approx. 20 rpm. Air may decrease at higher mixing speeds.</td>
<td>Follow truck mixer manufacturer recommendations. Maintain blades and clean truck mixer.</td>
</tr>
<tr>
<td>Admixture metering</td>
<td>Accuracy and reliability of metering system will affect uniformity of air content.</td>
<td>Avoid manual-dispensing or gravity-feed systems and timers. Positive-displacement pumps interlocked with batching system are preferred.</td>
</tr>
<tr>
<td>Transport and delivery</td>
<td>Some air (1% to 2%) normally lost during transport. Loss of air in nonagitating equipment is slightly higher.</td>
<td>Normal retempering with water to restore slump will restore air. If necessary, retemper with air-entraining admixture to restore air. Dramatic loss in air may be due to factors other than transport.</td>
</tr>
<tr>
<td>Haul time and agitation</td>
<td>Long hauls, even without agitation, reduce air, especially in hot weather.</td>
<td>Optimize delivery schedules. Maintain concrete temperature in recommended range.</td>
</tr>
<tr>
<td>Retempering</td>
<td>Regains some of the lost air. Does not usually affect the air-void system. Retempering with air-entraining admixtures restores the air-void system.</td>
<td>Retemper only enough to restore workability. Avoid addition of excess water. Higher admixture dosage is needed for jobsite admixture additions.</td>
</tr>
</tbody>
</table>
The effect of water hardness in most municipal water supplies is generally insignificant; however, very hard water from wells, as used in rural communities, may decrease the air content in concrete.

**Slump and Vibration**

The effect of slump and vibration on the air content of concrete is shown in Fig. 8-18. For a constant amount of air-entraining admixture, air content increases as slump increases up to about 150 or 175 mm (6 or 7 inches); then it begins to decrease with further increases in slump. At all slumps, however, even 15 seconds of vibration (the ACI 309 limit) will cause a considerable reduction in air content. Prolonged vibration of concrete should be avoided.

The greater the slump, air content, and vibration time, the larger the percentage of reduction in air content during vibration (Fig. 8-18). However, if vibration is properly applied, little of the intentionally entrained air is lost. The air lost during handling and moderate vibration consists mostly of the larger bubbles that are usually undesirable from the standpoint of strength. While the average size of the air voids is reduced, the air-void spacing factor remains relatively constant.

Internal vibrators reduce air content more than external vibrators. The air loss due to vibration increases as the volume of concrete is reduced or the vibration frequency is significantly increased. Lower vibration frequencies (8000 vpm) have less effect on spacing factors.
and air content than high vibration frequencies (14,000 vpm). High frequencies can significantly increase spacing factors and decrease air content after 20 seconds of vibration (Brewster 1949 and Stark 1986).

For pavements, specified air contents and uniform air void distributions can be achieved by operating within paving machine speeds of 1.22 to 1.88 meters per minute (4 to 6 feet per minute) and with vibrator frequencies of 5,000 to 8,000 vibrations per minute. The most uniform distribution of air voids throughout the depth of concrete, in and out of the vibrator trails, is obtained with the combination of a vibrator frequency of approximately 5,000 vibrations per minute and a slipform paving machine forward track speeds of 1.22 meters per minute (4 feet per minute). Higher frequencies of speeds singularly or in combination can result in discontinuities and lack of required air content in the upper portion of the concrete pavement. This in turn provides a greater opportunity for water and salt to enter the pavement and reduce the durability and life of the pavement (Cable, McDaniel, Schlorholtz, Redmond, and Rabe 2000).

**Concrete Temperature**

Temperature of the concrete affects air content, as shown in Fig. 8-19. Less air is entrained as the temperature of the concrete increases, particularly as slump is increased. This effect is especially important during hot-weather concreting when the concrete might be quite warm. A decrease in air content can be offset when necessary by increasing the quantity of air-entraining admixture.

In cold-weather concreting, the air-entraining admixture may lose some of its effectiveness if hot mix water is used during batching. To offset this loss, such admixtures should be added to the batch after the temperature of the concrete ingredients have equalized.

Although increased concrete temperature during mixing generally reduces air volume, the spacing factor and specific surface are only slightly affected.

**Supplementary Cementitious Materials**

The effect of fly ash on the required dosage of air-entraining admixtures can range from no effect to an increase in dosage of up to five times the normal amount (Geblor and Klieger 1986). Class C ash typically requires less air-entraining admixture than Class F ash and tends to lose less air during mixing (Thomas 2007). Ground slags have variable effects on the required dosage rate of air-entraining admixtures. Silica fume has a marked influence on the air-entraining admixture requirement, which in most cases rapidly increases with an increase in the amount of silica fume used in the concrete. Large quantities of slag and silica fume can double the dosage of air-entraining admixtures required (Whiting and Nagi 1998). The inclusion of both fly ash and silica fume in non-air-entrained concrete will generally reduce the amount of entrapped air.

**Admixtures and Coloring Agents**

Coloring agents such as carbon black usually decrease the amount of air entrained for a given amount of admixture. This is especially true for coloring materials with increasing percentages of carbon (Taylor 1948).

Water-reducing and set-retarding admixtures generally increase the efficiency of air-entraining admixtures by 50% to 100%; therefore, when these are used, less air-entraining admixture will usually give the desired air content. Also, the time of addition of these admixtures into the mix affects the amount of entrained air; delayed additions generally increasing air content.

Set retarders may increase the air-void spacing in concrete. Some water-reducing or set-retarding admixtures are not compatible with some air-entraining admixtures. If they are added together to the mixing water before being dispensed into the mixer, a precipitate may form. This will settle out and result in large reductions in entrained air. The fact that some individual admixtures interact in this manner does not mean that they will not be fully effective if dispensed separately into a batch of concrete.

Superplasticizers (high-range water reducers) may increase or decrease the air content of a concrete mixture based on the admixture’s chemical formulation and the
slump of the concrete. Naphthalene-based superplasticizers tend to increase the air content while melamine-based materials may decrease or have little effect on air content. The normal air loss in flowing concrete during mixing and transport is about 2 to 4 percentage points (Whiting and Dziedzic 1992).

Superplasticizers also affect the air-void system of hardened concrete by increasing the size of the entrained air voids. This results in a higher-than-normal spacing factor, occasionally higher than what may be considered desirable for freeze-thaw durability. However, tests on superplasticized concrete with slightly higher spacing factors have indicated that superplasticized concretes have good freeze-thaw durability. This may be due to the reduced water-cement ratio often associated with superplasticized concretes.

A small quantity of calcium chloride is sometimes used in cold weather to accelerate the hardening of concrete. It can be used successfully with air-entraining admixtures if it is added separately in solution form to the mix water. Calcium chloride will slightly increase air content. However, if calcium chloride comes in direct contact with some air-entraining admixtures, a chemical reaction can take place that makes the admixture less effective. Nonchloride accelerators may increase or decrease air content, depending upon the individual chemistry of the admixture, but they generally have little effect on air content.

### Mixing Action

Mixing action is one of the most important factors in the production of entrained air in concrete. Uniform distribution of entrained air voids is essential to produce scale-resistant concrete; nonuniformity might result from inadequate dispersion of the entrained air during short mixing periods. In production of ready mixed concrete, it is especially important that adequate and consistent mixing be maintained at all times.

The amount of entrained air varies with the type and condition of the mixer, the amount of concrete being mixed, and the rate and duration of mixing. The amount of air entrained in a given mixture will decrease appreciably as the mixer blades become worn, or if hardened concrete is allowed to accumulate in the drum or on the blades. Because of differences in mixing action and time, concretes made in a stationary mixer and those made in a transit mixer may differ significantly in amounts of air entrained. The air content may increase or decrease when the size of the batch departs significantly from the rated capacity of the mixer. Little air is entrained in very small batches in a large mixer; however, the air content increases as the mixer capacity is approached.

Fig. 8-20 shows the effect of mixing speed and duration of mixing on the air content of freshly mixed concretes made in a transit mixer. Generally, more air is entrained as the speed of mixing is increased up to about 20 rpm, beyond which air entrainment decreases. In the tests from which the data in Fig. 8-20 were derived, the air content reached an upper limit during mixing and a gradual decrease in air content occurred with prolonged mixing. Mixing time and speed will have different effects on the air content of different mixes. Significant amounts of air can be lost during mixing with certain mixtures and types of mixing equipment.

Fig. 8-21 shows the effect of continued mixer agitation on air content. The changes in air content with prolonged agitation can be explained by the relationship between slump and air content. For high-slump concretes, the air content increases with continued agitation as the slump decreases to about 150 or 175 mm (6 or 7 in.). Prolonged agitation will decrease slump further and decrease air content. For initial slumps lower than 150 mm (6 in.), both the air content and slump decrease with continued agitation. When concrete is retempered (the addition of water and remixing to restore original slump), the air content is increased; however, after 4 hours, retempering is ineffective in increasing air content. Prolonged mixing or agitation of concrete is accompanied by a progressive reduction in slump.
Transporting and Handling

Generally, some air—approximately 1 to 2 percentage points—is lost during transportation of concrete from the mixer to the jobsite. The stability of the air content during transport is influenced by several variables including concrete ingredients, haul time, amount of agitation or vibration during transport, temperature, slump, and amount of retempering.

Once at the jobsite, the concrete air content remains essentially constant during handling by chute discharge, wheelbarrow, power buggy, and shovel. However, concrete pumping, crane and bucket, and conveyor-belt handling can cause some loss of air, especially with high-air-content mixtures. Pumping concrete can cause a loss of up to 3 percentage points of air (Whiting and Nagi 1998).

Finishing

Proper screeding, floating, and general finishing practices should not affect the air content. McNeal and Gay (1996) and Falconi (1996) demonstrated that the sequence and timing of finishing and curing operations are critical to surface durability. Overfinishing (excessive finishing) may reduce the amount of entrained air in the surface region of slabs—thus making the concrete surface vulnerable to scaling. However, as shown in Fig. 8-22, early finishing does not necessarily affect scale resistance unless bleed water is present (Pinto and Hover 2001). Concrete to be exposed to deicers should not be steel troweled.

TESTS FOR AIR CONTENT

Four methods for determining the air content of freshly mixed concrete are available. Although they measure only total air volume and not air-void characteristics, it has been shown by laboratory tests that these methods are indicative of the adequacy of the air-void system.

Acceptance tests for air content of freshly mixed concrete should be made regularly for routine control purposes. Samples should be obtained and tested in accordance with ASTM C172 (AASHTO T 141).

Following are methods for determining the air content of freshly mixed concrete:

1. Pressure method (ASTM C231 or AASHTO T 152, Standard Test Method for Air Content of Freshly Mixed Concrete by the Pressure Method)—practical for field-testing all concretes except those made with highly porous and lightweight aggregates.

2. Volumetric method (ASTM C173 or AASHTO T 196, Standard Test Method for Air Content of Freshly Mixed Concrete by the Volumetric Method)—practical for field-testing all concretes, but particularly useful for concretes made with lightweight and porous aggregates.

When aggregates larger than 50 mm (2 in.) are used, they should be removed by hand and the effect of their removal calculated in arriving at the total air content.

3. Gravimetric method (ASTM C138 or AASHTO T 121, Standard Test Method for Unit Weight, Yield, and Air Content [Gravimetric] of Concrete)—requires accurate knowledge of relative density and absolute volumes of concrete ingredients.

4. Chace air indicator (AASHTO T 199, Standard Method of Test for Air Content of Freshly Mixed Concrete by the Chace Indicator)—a very simple and inexpensive way to check the approximate air content of freshly mixed concrete. This pocket-size device tests a mortar sample from the concrete. This test is not a substitute, however, for the more accurate pressure, volumetric, and gravimetric methods.

The foam-index test can be used to measure the relative air-entraining admixture requirement for concretes containing fly ash-cement combinations (Gebler and Klieger 1983).

![Fig. 8-22. Effect of early finishing—magnesium floating 20 minutes after casting—on scale resistance for: (top) 6% air-entrained concrete; (bottom) non-air-entrained concrete.](image-url)
The air-void characteristics of hardened concrete can be determined by ASTM C457 methods. This test is used to determine void spacing factor, specific surface of voids, and number of voids per length of traverse.

**Air-Void Analysis of Fresh Concrete**

The conventional methods for analyzing air in fresh concrete, such as the pressure method noted above, measure the total air content only; consequently, they provide no information about the parameters that determine the quality of the air-void system. These parameters—the size and number of voids and spacing between them—can be measured on polished samples of hardened concrete (ASTM C457); but the result of such analysis will only be available several days after the concrete has hardened. Therefore, test equipment called an air-void analyzer (AVA) has been developed to determine the standard ASTM C457 air-void parameters in fresh samples of air-entrained concrete (Fig. 8-23). The test apparatus determines the volume and size distributions of entrained air voids; thus an estimation of the spacing factor, specific surface, and total amount of entrained air can be made.

In this test method, air bubbles from a sample of fresh concrete rise through a viscous liquid, enter a column of water above it, then rise through the water and collect under a submerged buoyancy recorder (Fig. 8-24). The viscous liquid retains the original bubble sizes. Large bubbles rise faster than small ones through the liquids. The change in buoyancy is recorded as a function of time and can be related to the number of bubbles of different size.

**RECOMMENDED AIR CONTENTS**

The amount of air to be used in air-entrained concrete depends on a number of factors: (1) type of structure, (2) climatic conditions, (3) number of freeze-thaw cycles, (4) extent of exposure to deicers, and (5) the design life of the structure.

The ACI 318 building code (2008) contains four different exposure classes in the Exposure Category “F-Freezing and Thawing:”

1. F0 (Not applicable)—Concrete not exposed to freezing and thawing cycles
2. F1 (Moderate)—Concrete exposed to freezing and thawing cycles and occasional exposure to moisture
3. F2 (Severe)—Concrete exposed to freezing and thawing cycles and in continuous contact with moisture
4. F3 (Very Severe)—Concrete exposed to freezing and thawing cycles that will be in continuous contact with moisture and exposure to deicing chemicals

Concrete in exposure categories F1, F2, and F3 has to be air entrained with the air contents listed in Table 8-6 for...
moderate and severe exposure. Furthermore, the water to cementitious materials ratio of concrete in these three categories should not exceed 0.45 and have a 28-day compressive strength of at least 28 MPa (4500 psi). ACI 318 allows a one-percentage point reduction in target air contents for concretes with strengths over 34 MPa (5000 psi), and presumably such concretes would have low water to cementitious materials ratios. Fig. 8-25 illustrates how deicer-scaling resistance is impacted by air content and low water to portland cement ratios (strengths ranging from 40 to 59 MPa [5,800 to 8,600 psi]). This illustrates that concretes with very low water to portland cement ratios are more frost and deicer resistant; hence, they may allow use of lower air contents. This relationship (Fig. 8-25) was not established for concretes containing supplementary cementitious materials as they were not studied (Pinto and Hover 2001).

ACI 318 limits the amounts of pozzolans and slag—10% for silica fume, 25% for fly ash, 50% for slag—as part of the cementitious material for deicer exposures. However, mix designers should consult local practices as to allowable dosages to prevent frost and deicer damage. Combinations of materials without historical record can be analyzed using ASTM C 666 (AASHTO T 161) and ASTM C672. Pinto and Hover (2001) evaluate the applicability of the ACI 318 requirements for frost resistance of portland cement concrete mixtures with water to cement ratios from 0.25 to 0.50. Fig. 8-26 illustrates the effect of increased air content with respect to aggregate size on reducing expansion due to saturated freezing and thawing; it emphasizes the need to follow the requirements of Table 8-6 for severe exposure.

When entrained air is not required for protection against freeze-thaw cycles or deicers, the target air contents for mild exposure given in Table 8-6 can be used. Higher air contents can also be used as long as the design strength is achieved. As noted earlier, entrained air helps to reduce bleeding and segregation and can improve the workability of concrete.

More information on air-entrained concrete can be found in Whiting and Nagi (1998).

Table 8-6. Recommended Total Target Air Content for Concrete

<table>
<thead>
<tr>
<th>Nominal maximum size aggregate, mm (in.)</th>
<th>Air content, percent*</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Severe exposure**</td>
</tr>
<tr>
<td></td>
<td>(Class F2 and F3)</td>
</tr>
<tr>
<td>&lt;9.5 (%)</td>
<td>9</td>
</tr>
<tr>
<td>9.5 (%)</td>
<td>7½</td>
</tr>
<tr>
<td>12.5 (%)</td>
<td>7</td>
</tr>
<tr>
<td>19.0 (%)</td>
<td>6</td>
</tr>
<tr>
<td>25.0 (1)</td>
<td>6</td>
</tr>
<tr>
<td>37.5 (1½)</td>
<td>5½</td>
</tr>
<tr>
<td>50 (2)††</td>
<td>5</td>
</tr>
<tr>
<td>75 (3)‡‡</td>
<td>4½</td>
</tr>
</tbody>
</table>

* Project specifications often allow the air content of the concrete to be within -1 to +2 percentage points of the table target values.

** Concrete exposed to wet-freeze-thaw conditions, deicers, or other aggressive agents (ACI 318 Exposure Class F2 and F3).

† Concrete exposed to freezing but not continually moist, and not in contact with deicers or aggressive chemicals (ACI 318 Exposure Class F1).

‡‡ Concrete not exposed to freezing conditions, deicers, or aggressive agents.

† These air contents apply to the total mix, as for the preceding aggregate sizes. When testing these concretes, however, aggregate larger than 37.5 mm (1½ in.) is removed by handpicking or sieving and air content is determined on the minus 37.5 mm (1½ in.) fraction of mix. (Tolerance on air content as delivered applies to this value.)

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### Fig. 8-25. Measured mass loss after 40 cycles of deicers and freeze-thaw exposures at various air contents (Pinto and Hover 2001).

### Fig. 8-26. Relationship between air content and expansion of concrete test specimens during 300 cycles of freezing and thawing for various maximum aggregate sizes. Smaller aggregate sizes require more air (Klieger 1952).
REFERENCES


ACI Committee 201, Guide to Durable Concrete, ACI 201.2R-01, reapproved 1997, ACI Committee 201 Report, American Concrete Institute, Farmington Hills, Michigan, 2001, 41 pages.

ACI Committee 309, Guide for Consolidation of Concrete, ACI 309R-96, ACI Committee 309 Report, American Concrete Institute, Farmington Hills, Michigan, 1996.

ACI Committee 318, Building Code Requirements for Structural Concrete and Commentary, ACI 318-88, ACI Committee 318 Report, American Concrete Institute, Farmington Hills, Michigan, 2008.

ACPA, Scale-Resistant Concrete Pavements, IS117, American Concrete Pavement Association, Skokie, Illinois, 1996.


Wang, Kejin; Monteiro, Paulo J. M.; Rubinsky, Boris; and Arav, Amir, “Microscopic Study of Ice Propagation in Concrete,” *ACI Materials Journal*, July-August 1996, American Concrete Institute, Farmington Hills, Michigan, pages 370 to 376.


The process of determining required and specifiable characteristics of a concrete mixture is called mix design. Characteristics can include: (1) fresh concrete properties; (2) required mechanical properties of hardened concrete such as strength and durability requirements; and (3) the inclusion, exclusion, or limits on specific ingredients. Mix design leads to the development of a concrete specification.

Mixture proportioning refers to the process of determining the quantities of concrete ingredients, using local materials, to achieve the specified characteristics of the concrete. A properly proportioned concrete mix should possess these qualities:

1. Acceptable workability of the freshly mixed concrete
2. Durability, strength, and uniform appearance of the hardened concrete
3. Economy

Understanding the basic principles of mixture design is as important as the actual calculations used to establish mix proportions. Only with proper selection of materials and mixture characteristics can the above qualities be obtained in concrete construction (Fig. 9-1) (Abrams 1918, Hover 1998, and Shilstone 1990).

SELECTING MIX CHARACTERISTICS

Before a concrete mixture can be proportioned, mixture characteristics are selected based on the intended use of the concrete, the exposure conditions, the size and shape of building elements, and the physical properties of the concrete (such as frost resistance and strength) required for the structure. The characteristics should reflect the needs of the structure; for example, resistance to chloride ions should be verifiable and the appropriate test methods specified.

Once the characteristics are selected, the mixture can be proportioned from field or laboratory data. Since most of the desirable properties of hardened concrete depend primarily upon the quality of the cementitious paste, the first step in proportioning a concrete mixture is the selection of the appropriate water-cementing materials ratio for the durability and strength needed. Concrete mixtures should be kept as simple as possible, as an excessive number of ingredients often make a concrete mixture difficult to control. The concrete technologist should not, however, overlook the opportunities provided by modern concrete technology.

Water-Cementing Materials Ratio and Strength Relationship

Strength (compressive or flexural) is the most universally used measure for concrete quality. Although it is an important characteristic, other properties such as durability, permeability, and wear resistance are now recognized as being equal and in some cases more important, especially when considering life-cycle design of structures.

Within the normal range of strengths used in concrete construction, the compressive strength is inversely related to the water-cement ratio or water-cementing materials ratio. For fully compacted concrete made with clean, sound aggregates, the strength and other desirable prop-

Fig. 9-1. Trial batching (inset) verifies that a concrete mixture meets design requirements prior to use in construction. (IMG12399, IMG12400).
The specified compressive strength, \( f_c \), at 28 days is the strength that is expected to be equal to or exceeded by the average of any set of three consecutive strength tests. ACI 318 requires for \( f_c \) to be at least 17.5 MPa (2500 psi). No individual test (average of two cylinders) can be more than 3.5 MPa (500 psi) below the specified strength. Specimens must be cured under laboratory conditions for an individual class of concrete (ACI 318). Some specifications allow alternative ranges.

The average strength should equal the specified strength plus an allowance to account for variations in materials; variations in methods of mixing, transporting, and placing the concrete; and variations in making, curing, and testing concrete cylinder specimens. The average strength, which is greater than \( f_c \), is called \( \bar{f} \); it is the strength required in the mix design. Requirements for \( \bar{f} \) are discussed in detail under “Proportioning” later.

### Table 9-1. Maximum Water-Cementitious Material Ratios and Minimum Design Strengths for Various Exposure Conditions

<table>
<thead>
<tr>
<th>Exposure category</th>
<th>Exposure condition</th>
<th>Maximum water-cementitious material ratio by mass for concrete</th>
<th>Minimum design compressive strength, ( f_c ), MPa (psi)</th>
</tr>
</thead>
<tbody>
<tr>
<td>F0, S0, P0, C0</td>
<td>Concrete protected from exposure to freezing and thawing, application of deicing chemicals, or aggressive substances</td>
<td>Select water-cementitious ratio on basis of strength, workability, and finishing needs</td>
<td>Select strength based on structural requirements</td>
</tr>
<tr>
<td>P1</td>
<td>Concrete intended to have low permeability when exposed to water</td>
<td>0.50</td>
<td>28 (4000)</td>
</tr>
<tr>
<td>F1, F2, F3</td>
<td>Concrete exposed to freezing and thawing in a moist condition or deicers</td>
<td>0.45</td>
<td>31 (4500)</td>
</tr>
<tr>
<td>C2</td>
<td>For corrosion protection for reinforced concrete exposed to chlorides from deicing salts, salt water, brackish water, seawater, or spray from these sources</td>
<td>0.40</td>
<td>35 (5000)</td>
</tr>
</tbody>
</table>

Adapted from ACI 318 (2002). ACI 318-08 introduced the following four exposure categories that determine durability requirements for concrete: (1) F – Freezing and Thawing; (2) S – Sulfates; (3) P – Permeability; and (4) C – Corrosion. Increasing numerical values represent increasingly severe exposure conditions.

### Table 9-2. Types of Cement Required for Concrete Exposed to Sulfates in Soil or Water

<table>
<thead>
<tr>
<th>Sulfate exposure class</th>
<th>Sulfate (SO₄) in soil, % by mass</th>
<th>Sulfate (SO₄) in water, ppm</th>
<th>Cement type*</th>
<th>Maximum water-cementitious material ratio, by mass</th>
</tr>
</thead>
<tbody>
<tr>
<td>S0 Negligible</td>
<td>Less than 0.10</td>
<td>Less than 150</td>
<td>No special type required</td>
<td>—</td>
</tr>
<tr>
<td>S1 Moderate**</td>
<td>0.10 to 0.20</td>
<td>150 to 1500</td>
<td>II (IP(MS), IS(&lt;70)(MS))</td>
<td>MS 0.50</td>
</tr>
<tr>
<td>S2 Severe</td>
<td>0.20 to 2.00</td>
<td>1500 to 10,000</td>
<td>V (IP(HS), IS(&lt;70)(HS))</td>
<td>HS 0.45</td>
</tr>
<tr>
<td>S3 Very severe</td>
<td>Over 2.00</td>
<td>Over 10,000</td>
<td>V (IP(HS), IS(&lt;70)(HS))</td>
<td>HS 0.40</td>
</tr>
</tbody>
</table>

* Pozzolans and slag that have been determined by testing according to ASTM C1012 or by service record to improve sulfate resistance may also be used. Maximum expansions when using ASTM C1012: Moderate exposure—0.10% at 6 months; Severe exposure—0.05% at 6 months or 0.10% at 12 months; Very Severe exposure—0.10% at 18 months. Sulfate resistance of individual pozzolans or slags needs to be established by demonstrating a maximum expansion of 0.10% at 1 year, before a 6 months test duration is acceptable. Refer to ACI 201 (2001) for more guidance.

** Includes seawater.


Source: Adapted from Bureau of Reclamation 1981, ACI 201, and ACI 318.
in this chapter. Tables 9-1 and 9-2 show strength requirements for various exposure conditions.

Flexural strength is sometimes used on paving projects instead of compressive strength; however, flexural strength is avoided due to its greater variability. For more information on flexural strength, see “Strength” in Chapter 1 and “Strength Specimens” in Chapter 16.

**Water-Cementitious Material Ratio**

The water-cementitious material ratio is simply the mass of water divided by the mass of cementitious material (Portland cement, blended cement, fly ash, slag, silica fume, and natural pozzolans). The water-cementitious material ratio selected for mix design must be the lowest value required to meet anticipated exposure conditions. Tables 9-1 and 9-2 show requirements for various exposure conditions.

When durability does not control, the water-cementitious materials ratio should be selected on the basis of concrete compressive strength. In such cases the water-cementitious materials ratio and mixture proportions for the required strength should be based on adequate field data or trial mixes made with actual job materials to determine the relationship between the ratio and strength. Fig. 9-2 or Table 9-3 can be used to select a water-cementitious materials ratio with respect to the required average strength, $f_{cm}$, for trial mixes when no other data are available.

In mix design, the water to cementitious materials ratio, $W/CM$, is often used synonymously with water to cement ratio ($W/C$); however, some specifications differentiate between the two ratios. Traditionally, the water to cement ratio referred to the ratio of water to Portland cement or water to blended cement.

**Aggregates**

Two characteristics of aggregates have an important influence on proportioning concrete mixtures because they affect the workability of the fresh concrete. They are:

1. Grading (particle size and distribution)
2. Nature of particles (shape, porosity, surface texture)

Grading is important for attaining an economical mixture because it affects the amount of concrete that can be made with a given amount of cementitious materials and water. Coarse aggregates should be graded up to the largest size practical under job conditions. The maximum size that can be used depends on factors such as the size and shape of the concrete member to be cast, the amount and distribution of reinforcing steel in the member, and the thickness of slabs. Grading also influences the workability and placeability of the concrete. Sometimes mid-sized aggregate, around the 9.5 mm (% in.) size, is lacking.

**Table 9-3 (Metric). Relationship Between Water to Cementitious Material Ratio and Compressive Strength of Concrete**

<table>
<thead>
<tr>
<th>Compressive strength at 28 days, MPa</th>
<th>Water-cementitious materials ratio by mass</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Non-air-entrained concrete</td>
</tr>
<tr>
<td>45</td>
<td>0.38</td>
</tr>
<tr>
<td>40</td>
<td>0.42</td>
</tr>
<tr>
<td>35</td>
<td>0.47</td>
</tr>
<tr>
<td>30</td>
<td>0.54</td>
</tr>
<tr>
<td>25</td>
<td>0.61</td>
</tr>
<tr>
<td>20</td>
<td>0.69</td>
</tr>
<tr>
<td>15</td>
<td>0.79</td>
</tr>
</tbody>
</table>

Strength is based on cylinders moist-cured 28 days in accordance with ASTM C31 (AASHTO T 23). Relationship assumes nominal maximum size aggregate of about 19 to 25 mm. Adapted from ACI 211.1 and ACI 211.3.

**Table 9-3 (Inch-Pound Units). Relationship Between Water to Cementitious Material Ratio and Compressive Strength of Concrete**

<table>
<thead>
<tr>
<th>Compressive strength at 28 days, psi</th>
<th>Water-cementitious materials ratio by mass</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Non-air-entrained concrete</td>
</tr>
<tr>
<td>7000</td>
<td>0.33</td>
</tr>
<tr>
<td>6000</td>
<td>0.41</td>
</tr>
<tr>
<td>5000</td>
<td>0.48</td>
</tr>
<tr>
<td>4000</td>
<td>0.57</td>
</tr>
<tr>
<td>3000</td>
<td>0.68</td>
</tr>
<tr>
<td>2000</td>
<td>0.82</td>
</tr>
</tbody>
</table>

Strength is based on cylinders moist-cured 28 days in accordance with ASTM C31 (AASHTO T 23). Relationship assumes nominal maximum size aggregate of about ¾ in. to 1 in. Adapted from ACI 211.1 and ACI 211.3.
in an aggregate supply; this can result in a concrete with high shrinkage properties, high water demand, and poor workability and placeability. Durability may also be affected. Various options are available for obtaining optimal grading of aggregate (Shilstone 1990).

The maximum size of coarse aggregate should not exceed one-fifth the narrowest dimension between sides of forms nor three-fourths the clear space between individual reinforcing bars or wire, bundles of bars, or prestressing tendons or ducts. It is also good practice to limit aggregate size to not more than three-fourths the clear space between reinforcement and the forms. For unreinforced slabs on ground, the maximum size should not exceed one third the slab thickness. Smaller sizes can be used when availability or economic consideration require them.

The amount of mixing water required to produce a unit volume of concrete of a given slump is dependent on the shape and the maximum size and amount of coarse aggregate. Larger sizes minimize the water requirement and thus allow the cement content to be reduced. Also, rounded aggregate requires less mixing water than a crushed aggregate in concretes of equal slump (see “Water Content”).

The maximum size of coarse aggregate that will produce concrete of maximum strength for a given cement content depends upon the aggregate source as well as its shape and grading. For high compressive-strength concrete (greater than 70 MPa or 10,000 psi), the maximum size is about 19 mm (3⁄4 in.). Higher strengths can also sometimes be achieved through the use of crushed stone aggregate rather than rounded-gravel aggregate.

The most desirable fine-aggregate grading will depend upon the type of work, the paste content of the mixture, and the size of the coarse aggregate. For leaner mixtures, a fine grading (lower fineness modulus) is desirable for workability. For richer mixtures, a coarse grading (higher fineness modulus) is used for greater economy.

In some areas, the chemically bound chloride in aggregate may make it difficult for concrete to pass chloride limits set by ACI 318 or other specifications. However, some or all of the chloride in the aggregate may not be available for participation in corrosion of reinforcing steel, thus that chloride may be ignored. ASTM C1524, Soxhlet extracted chloride test, can be used to evaluate the amount of chloride available from aggregate. ACI 222.1 also provides guidance.

The bulk volume of coarse aggregate can be determined from Fig. 9-3 or Table 9-4. These bulk volumes are based on aggregates in a dry-rodded condition as described in ASTM C29 (AASHTO T 19); they are selected from empirical relationships to produce concrete with a degree of workability suitable for general reinforced concrete construction. For less workable concrete, such as required for concrete pavement construction, they may be increased about 10%. For more workable concrete, such as may be required when placement is by pump, they may be reduced up to 10%.

### Air Content

Entrained air must be used in all concrete that will be exposed to freezing and thawing and deicing chemicals and can be used to improve workability even where not required.

### Table 9-4. Bulk Volume of Coarse Aggregate Per Unit Volume of Concrete

<table>
<thead>
<tr>
<th>Nominal maximum size of aggregate, mm (in.)</th>
<th>Bulk volume of dry-rodded coarse aggregate per unit volume of concrete for different fineness moduli of fine aggregate*</th>
</tr>
</thead>
<tbody>
<tr>
<td>9.5 (%)</td>
<td>0.50 0.48 0.46 0.44</td>
</tr>
<tr>
<td>12.5 (%)</td>
<td>0.59 0.57 0.55 0.53</td>
</tr>
<tr>
<td>19 (%)</td>
<td>0.66 0.64 0.62 0.60</td>
</tr>
<tr>
<td>25 (1)</td>
<td>0.71 0.69 0.67 0.65</td>
</tr>
<tr>
<td>37.5 (1/4)</td>
<td>0.75 0.73 0.71 0.69</td>
</tr>
<tr>
<td>50 (2)</td>
<td>0.78 0.76 0.74 0.72</td>
</tr>
<tr>
<td>75 (3)</td>
<td>0.82 0.80 0.78 0.76</td>
</tr>
<tr>
<td>150 (6)</td>
<td>0.87 0.85 0.83 0.81</td>
</tr>
</tbody>
</table>

*Bulk volumes are based on aggregates in a dry-rodded condition as described in ASTM C29 (AASHTO T 19). Adapted from ACI 211.1.
Air entrainment is accomplished by using an air-entraining portland cement or by adding an air-entraining admixture at the mixer. The amount of admixture should be adjusted to meet variations in concrete ingredients and job conditions. The amount recommended by the admixture manufacturer will, in most cases, produce the desired air content.

Recommended target air contents for air-entrained concrete are shown in Fig. 9-4 and Table 9-5. Note that the amount of air required to provide adequate freeze-thaw resistance is dependent upon the nominal maximum size of aggregate and the level of exposure. In properly proportioned mixes, the mortar content decreases as maximum aggregate size increases, thus decreasing the required concrete air content. This is evident in Fig. 9-4. The levels of exposure are defined by ACI 211.1 as follows:

**Mild Exposure.** This exposure includes indoor or outdoor service in a climate where concrete will not be exposed to freezing or deicing agents. When air entrainment is desired for a beneficial effect other than durability, such as to improve workability or cohesion or in low cement content concrete to improve strength, air contents lower than those needed for durability can be used.

**Moderate Exposure.** Service in a climate where freezing is expected but where the concrete will not be continually exposed to moisture or free water for long periods prior to freezing and will not be exposed to deicing or other aggressive chemicals. Examples include exterior beams, columns, walls, girders, or slabs that are not in contact with wet soil and are so located that they will not receive direct applications of deicing chemicals.

**Severe Exposure.** Concrete that is exposed to deicing or other aggressive chemicals or where the concrete may become highly saturated by continual contact with moisture or free water prior to freezing. Examples include pavements, bridge decks, curbs, gutters, sidewalks, canal linings, or exterior water tanks or sumps.

When mixing water is held constant, the entrainment of air will increase slump. When cement content and slump are held constant, the entrainment of air results in the need for less mixing water, particularly in leaner concrete mixtures. In batch adjustments, in order to maintain a constant slump while changing the air content, the water should be decreased by about 3 kg/m³ (5 lb/yd³) for each percentage point increase in air content or increased 3 kg/m³ (5 lb/yd³) for each percentage point decrease.

A specific air content may not be readily or repeatedly achieved because of the many variables affecting air content; therefore, a permissible range of air contents around a target value must be provided. Although a range of ±1% of the Fig. 9-4 or Table 9-5 values is often used in project specifications, it is sometimes an impractically tight limit. The solution is to use a wider range, such as –1 to +2 percentage points of the target values. For example, for a target value of 6% air, the specified range for the concrete delivered to the jobsite could be 5% to 8%.

**Slump**

Concrete must always be made with a workability, consistency, and plasticity suitable for job conditions. Workability is a measure of how easy or difficult it is to place, consolidate, and finish concrete. Consistency is the ability of freshly mixed concrete to flow. Plasticity determines concrete’s ease of molding. If more aggregate is used in a concrete mixture, or if less water is added, the mixture becomes stiff (less plastic and less workable) and difficult to mold. Neither very dry, crumbly mixtures nor very watery, fluid mixtures can be regarded as having plasticity.

The slump test is used to measure concrete consistency. For a given proportion of cement and aggregate without admixtures, the higher the slump, the wetter the mixture. Slump is indicative of workability when assessing similar mixtures. However, slump should not be used to compare mixtures of totally different proportions. When used with different batches of the same mix design, a change in slump indicates a change in consistency and in the characteristics of materials, mixture proportions, water content, mixing, time of test, or the testing itself.
### Table 9-5 (Metric). Approximate Mixing Water and Target Air Content Requirements for Different Slumps and Nominal Maximum Sizes of Aggregate

<table>
<thead>
<tr>
<th>Slump, mm</th>
<th>Water, kilograms per cubic meter of concrete, for indicated sizes of aggregate*</th>
<th>9.5 mm</th>
<th>12.5 mm</th>
<th>19 mm</th>
<th>25 mm</th>
<th>37.5 mm</th>
<th>50 mm**</th>
<th>75 mm**</th>
<th>150 mm**</th>
</tr>
</thead>
<tbody>
<tr>
<td>25 to 50</td>
<td>Non-air-entrained concrete</td>
<td>207</td>
<td>199</td>
<td>190</td>
<td>179</td>
<td>166</td>
<td>154</td>
<td>130</td>
<td>113</td>
</tr>
<tr>
<td>75 to 100</td>
<td></td>
<td>228</td>
<td>216</td>
<td>205</td>
<td>193</td>
<td>181</td>
<td>169</td>
<td>145</td>
<td>124</td>
</tr>
<tr>
<td>150 to 175</td>
<td></td>
<td>243</td>
<td>228</td>
<td>216</td>
<td>202</td>
<td>190</td>
<td>178</td>
<td>160</td>
<td>—</td>
</tr>
<tr>
<td>Approximate amount of entrapped air in non-air-entrained concrete, percent</td>
<td></td>
<td>3</td>
<td>2.5</td>
<td>2</td>
<td>1.5</td>
<td>1</td>
<td>0.5</td>
<td>0.3</td>
<td>0.2</td>
</tr>
<tr>
<td>25 to 50</td>
<td>Air-entrained concrete</td>
<td>181</td>
<td>175</td>
<td>168</td>
<td>160</td>
<td>150</td>
<td>142</td>
<td>122</td>
<td>107</td>
</tr>
<tr>
<td>75 to 100</td>
<td></td>
<td>202</td>
<td>193</td>
<td>184</td>
<td>175</td>
<td>165</td>
<td>157</td>
<td>133</td>
<td>119</td>
</tr>
<tr>
<td>150 to 175</td>
<td></td>
<td>216</td>
<td>205</td>
<td>197</td>
<td>184</td>
<td>174</td>
<td>166</td>
<td>154</td>
<td>—</td>
</tr>
<tr>
<td>Recommended average total air content, percent, for level of exposure:†</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Mild exposure</td>
<td></td>
<td>4.5</td>
<td>4.0</td>
<td>3.5</td>
<td>3.0</td>
<td>2.5</td>
<td>2.0</td>
<td>1.5</td>
<td>1.0</td>
</tr>
<tr>
<td>Moderate exposure (Class F1)</td>
<td></td>
<td>6.0</td>
<td>5.5</td>
<td>5.0</td>
<td>4.5</td>
<td>4.5</td>
<td>4.0</td>
<td>3.5</td>
<td>3.0</td>
</tr>
<tr>
<td>Severe exposure (Class F2 and F3)</td>
<td></td>
<td>7.5</td>
<td>7.0</td>
<td>6.0</td>
<td>6.0</td>
<td>5.5</td>
<td>5.0</td>
<td>4.5</td>
<td>4.0</td>
</tr>
</tbody>
</table>

* These quantities of mixing water are for use in computing cementitious material contents for trial batches. They are maximums for reasonably well-shaped angular coarse aggregates graded within limits of accepted specifications.

** The slump values for concrete containing aggregates larger than 37.5 mm are based on slump tests made after removal of particles larger than 37.5 mm by wet screening.

† The air content in job specifications should be specified to be delivered within –1 to +2 percentage points of the table target value for moderate and severe exposures.

Adapted from ACI 211.1 and ACI 318. Hover (1995) presents this information in graphical form.

### Table 9-5 (Inch-Pound Units). Approximate Mixing Water and Target Air Content Requirements for Different Slumps and Nominal Maximum Sizes of Aggregate

<table>
<thead>
<tr>
<th>Slump, in.</th>
<th>Water, pounds per cubic yard of concrete, for indicated sizes of aggregate*</th>
<th>% in.</th>
<th>% in.</th>
<th>% in.</th>
<th>1 in.</th>
<th>1 1/2 in.</th>
<th>2 in.**</th>
<th>3 in.**</th>
<th>6 in.**</th>
</tr>
</thead>
<tbody>
<tr>
<td>1 to 2</td>
<td>Non-air-entrained concrete</td>
<td>350</td>
<td>335</td>
<td>315</td>
<td>300</td>
<td>275</td>
<td>260</td>
<td>220</td>
<td>190</td>
</tr>
<tr>
<td>3 to 4</td>
<td></td>
<td>385</td>
<td>365</td>
<td>340</td>
<td>325</td>
<td>300</td>
<td>285</td>
<td>245</td>
<td>210</td>
</tr>
<tr>
<td>6 to 7</td>
<td></td>
<td>410</td>
<td>385</td>
<td>360</td>
<td>340</td>
<td>315</td>
<td>300</td>
<td>270</td>
<td>—</td>
</tr>
<tr>
<td>Approximate amount of entrapped air in non-air-entrained concrete, percent</td>
<td></td>
<td>3</td>
<td>2.5</td>
<td>2</td>
<td>1.5</td>
<td>1</td>
<td>0.5</td>
<td>0.3</td>
<td>0.2</td>
</tr>
<tr>
<td>1 to 2</td>
<td>Air-entrained concrete</td>
<td>305</td>
<td>295</td>
<td>280</td>
<td>270</td>
<td>250</td>
<td>240</td>
<td>205</td>
<td>180</td>
</tr>
<tr>
<td>3 to 4</td>
<td></td>
<td>340</td>
<td>325</td>
<td>305</td>
<td>295</td>
<td>275</td>
<td>265</td>
<td>225</td>
<td>200</td>
</tr>
<tr>
<td>6 to 7</td>
<td></td>
<td>365</td>
<td>345</td>
<td>325</td>
<td>310</td>
<td>290</td>
<td>280</td>
<td>260</td>
<td>—</td>
</tr>
<tr>
<td>Recommended average total air content, percent, for level of exposure:†</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Mild exposure</td>
<td></td>
<td>4.5</td>
<td>4.0</td>
<td>3.5</td>
<td>3.0</td>
<td>2.5</td>
<td>2.0</td>
<td>1.5</td>
<td>1.0</td>
</tr>
<tr>
<td>Moderate exposure (Class F1)</td>
<td></td>
<td>6.0</td>
<td>5.5</td>
<td>5.0</td>
<td>4.5</td>
<td>4.5</td>
<td>4.0</td>
<td>3.5</td>
<td>3.0</td>
</tr>
<tr>
<td>Severe exposure (Class F2 and F3)</td>
<td></td>
<td>7.5</td>
<td>7.0</td>
<td>6.0</td>
<td>6.0</td>
<td>5.5</td>
<td>5.0</td>
<td>4.5</td>
<td>4.0</td>
</tr>
</tbody>
</table>

* These quantities of mixing water are for use in computing cement factors for trial batches. They are maximums for reasonably well-shaped angular coarse aggregates graded within limits of accepted specifications.

** The slump values for concrete containing aggregates larger than 1 1/2 in. are based on slump tests made after removal of particles larger than 1 1/2 in. by wet screening.

† The air content in job specifications should be specified to be delivered within –1 to +2 percentage points of the table target value for moderate and severe exposures.

Adapted from ACI 211.1. Hover (1995) presents this information in graphical form.
Different slumps are needed for various types of concrete construction. Slump is usually indicated in the job specifications as a range, such as 50 to 100 mm (2 to 4 in.), or as a maximum value not to be exceeded. ASTM C94 addresses slump tolerances in detail. When slump is not specified, an approximate value can be selected from Table 9-6 for concrete consolidated by mechanical vibration. For batch adjustments, the slump can be increased by about 10 mm by adding 2 kilograms of water per cubic meter of concrete (1 in. by adding 10 lb of water per cubic yard of concrete).

Water Content

The water content of concrete is influenced by a number of factors: aggregate size, aggregate shape, aggregate texture, slump, water to cementing materials ratio, air content, cementing materials type and content, admixtures, and environmental conditions. An increase in air content and aggregate size, a reduction in water-cementing materials ratio and slump, and the use of rounded aggregates, water-reducing admixtures, or fly ash will reduce water demand. On the other hand, increased temperatures, cement contents, slump, water-cement ratio, aggregate angularity, and a decrease in the proportion of coarse aggregate to fine aggregate will increase water demand.

The approximate water contents in Table 9-5 and Fig. 9-5, used in proportioning, are for angular coarse aggregates (crushed stone). For some concretes and aggregates, the water estimates in Table 9-5 and Fig. 9-5 can be reduced by approximately 10 kg (20 lb) for subangular aggregate, 20 kg (35 lb) for gravel with some crushed particles, and 25 kg (45 lb) for a rounded gravel to produce the slumps shown. This illustrates the need for trial batch testing of local materials, as each aggregate source is different and can influence concrete properties differently.

### Table 9-6. Recommended Slumps for Various Types of Construction

<table>
<thead>
<tr>
<th>Concrete construction</th>
<th>Slump, mm (in.)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Maximum*</td>
</tr>
<tr>
<td>Reinforced foundation walls and footings</td>
<td>75 (3)</td>
</tr>
<tr>
<td>Plain footings, caissons, and substructure walls</td>
<td>75 (3)</td>
</tr>
<tr>
<td>Beams and reinforced walls</td>
<td>100 (4)</td>
</tr>
<tr>
<td>Building columns</td>
<td>100 (4)</td>
</tr>
<tr>
<td>Pavements and slabs</td>
<td>75 (3)</td>
</tr>
<tr>
<td>Mass concrete</td>
<td>75 (3)</td>
</tr>
</tbody>
</table>

*May be increased 25 mm (1 in.) for consolidation by hand methods, such as rodding and spading. Plasticizers can safely provide higher slumps.

Adapted from ACI 211.1.
It should be kept in mind that changing the amount of any single ingredient in a concrete mixture normally effects the proportions of other ingredients as well as alter the properties of the mixture. For example, the addition of 2 kg of water per cubic meter will increase the slump by approximately 10 mm (10 lb of water per cubic yard will increase the slump by approximately 1 in.); it will also increase the air content and paste volume, decrease the aggregate volume, and lower the density of the concrete. In mixture adjustments, for the same slump, a decrease in air content by 1 percentage point will increase the water demand by about 3 kg per cubic meter of concrete (5 lb per cu yd of concrete).

**Cementing Materials Content and Type**

The cementing materials content is usually determined from the selected water-cementing materials ratio and water content, although a minimum cement content frequently is included in specifications in addition to a maximum water-cementing materials ratio. Minimum cement content requirements serve to ensure satisfactory durability and finishability, to improve wear resistance of slabs, and to guarantee a suitable appearance of vertical surfaces. This is important even though strength requirements may be met at lower cementing materials contents. However, excessively large amounts of cementing materials should be avoided to maintain economy in the mixture and to not adversely affect workability and other properties.

For severe freeze-thaw, deicer, and sulfate exposures, it is desirable to specify: (1) a minimum cementing materials content of 335 kg per cubic meter (564 lb per cubic yard) of concrete, and (2) only enough mixing water to achieve the desired consistency without exceeding the maximum water-cementing materials ratios shown in Tables 9-1 and 9-2. For placing concrete underwater, usually not less than 390 kg of cementing materials per cubic meter (650 lb of cementing materials per cubic yard) of concrete should be used with a water to cementing materials ratio not exceeding 0.45. For workability, finishability, abrasion resistance, and durability in flatwork, the quantity of cementing materials to be used should be not less than shown in Table 9-7.

To obtain economy, proportioning should minimize the amount of cement required without sacrificing concrete quality. Since quality depends primarily on water-cementing materials ratio, the water content should be held to a minimum to reduce the cement requirement. Steps to minimize water and cement requirements include use of (1) the stiffest practical mixture, (2) the largest practical maximum size of aggregate, and (3) the optimum ratio of fine-to-coarse aggregate.

Concrete that will be exposed to sulfate conditions should be made with the type of cement shown in Table 9-2.

Seawater contains significant amounts of sulfates and chlorides. Although sulfates in seawater are capable of attacking concrete, the presence of chlorides in seawater inhibits the expansive reaction that is characteristic of sulfate attack. This is the major factor explaining observations from a number of sources that the performance of concretes in seawater have shown satisfactory durability; this is despite the fact these concretes were made with portland cements having tricalcium aluminate (C₃A) contents as high as 10%, and sometimes greater. However, the permeability of these concretes was low, and the reinforcing steel had adequate cover. Portland cements meeting a C₃A requirement of not more than 10% or less than 4% (to ensure durability of reinforcement) are acceptable (ACI 357R).

Supplementary cementitious materials have varied effects on water demand and air contents. The addition of fly ash will generally reduce water demand and decrease the air content if no adjustment in the amount of air-entraining admixture is made. Silica fume increases water demand and decreases air content. Slag and metakaolin have a minimal effect at normal dosages.

**Table 9-7. Minimum Requirements of Cementing Materials for Concrete Used in Flatwork**

<table>
<thead>
<tr>
<th>Nominal maximum size of aggregate, mm (in.)</th>
<th>Cementing materials, kg/m³ (lb/yd³)*</th>
</tr>
</thead>
<tbody>
<tr>
<td>37.5 (1%)</td>
<td>280 (470)</td>
</tr>
<tr>
<td>25 (1)</td>
<td>310 (520)</td>
</tr>
<tr>
<td>19 (%)</td>
<td>320 (540)</td>
</tr>
<tr>
<td>12.5 (%)</td>
<td>350 (590)</td>
</tr>
<tr>
<td>9.5 (%)</td>
<td>360 (610)</td>
</tr>
</tbody>
</table>

* Cementing materials quantities may need to be greater for severe exposure. For example, for deicer exposures, concrete should contain at least 335 kg/m³ (564 lb/yard³) of cementing materials. Adapted from ACI 302.

<table>
<thead>
<tr>
<th>Cementitious materials*</th>
<th>Maximum percent of total cementitious materials by mass**</th>
</tr>
</thead>
<tbody>
<tr>
<td>Fly ash and natural pozzolans</td>
<td>25</td>
</tr>
<tr>
<td>Slag</td>
<td>50</td>
</tr>
<tr>
<td>Silica fume</td>
<td>10</td>
</tr>
<tr>
<td>Total of fly ash, slag, silica fume and natural pozzolans</td>
<td>50†</td>
</tr>
<tr>
<td>Total of natural pozzolans and silica fume</td>
<td>35†</td>
</tr>
</tbody>
</table>

* Includes portion of supplementary cementing materials in blended cements.

** Total cementitious materials include the summation of portland cements, blended cements, fly ash, slag, silica fume and other pozzolans.

† Silica fume should not constitute more than 10% of total cementitious materials and fly ash or other pozzolans shall not constitute more than 25% of cementitious materials. Adapted from ACI 318.
Table 9-8 shows limits on the amount of supplementary cementing materials in concrete to be exposed to deicers. Local practices should be consulted as dosages smaller or larger than those shown in Table 9-8 can be used without jeopardizing scale-resistance, depending on the exposure severity.

**Admixtures**

Water-reducing admixtures are added to concrete to reduce the water-cementing materials ratio, reduce cementing materials content, reduce water content, reduce paste content, or to improve the workability of a concrete without changing the water-cementing materials ratio. Water reducers will usually decrease water contents by 5% to 10% and some will also increase air contents by $\frac{1}{2}$ to 1 percentage point. Retarders may also increase the air content.

High-range water reducers (plasticizers) reduce water contents between 12% and 30% and some can simultaneously increase the air content up to 1 percentage point; others can reduce or not affect the air content.

Calcium chloride-based admixtures reduce water contents by about 3% and increase the air content by about $\frac{1}{2}$ percentage point.

When using a chloride-based admixture, the risks of reinforcing steel corrosion should be considered. Table 9-9 provides recommended limits on the water-soluble chloride-ion content in reinforced and prestressed concrete for various conditions.

When using more than one admixture in concrete, the compatibility of intermixing admixtures should be assured by the admixture manufacturer or the combination of admixtures should be tested in trial batches. The water contained in admixtures should be considered part of the mixing water if the admixture’s water content is sufficient to affect the water-cementing materials ratio by 0.01 or more.

An excessive use of multiple admixtures should be minimized to allow better control of the concrete mixture in production and to reduce the risk of admixture incompatibility.

**PROPORTIONING**

The design of concrete mixtures involves the following: (1) the establishment of specific concrete characteristics, and (2) the selection of proportions of available materials to produce concrete of required properties, with the greatest economy. Proportioning methods have evolved from the arbitrary volumetric method (1:2:3—cement:sand: coarse aggregate) of the early 1900s (Abrams 1918) to the present-day weight and absolute-volume methods described in ACI’s Committee 211 Standard Practice for Selecting Proportions for Normal, Heavyweight and Mass Concrete (ACI 211.1).

Weight proportioning methods are fairly simple and quick for estimating mixture proportions using an assumed or known weight of concrete per unit volume. A more accurate method, absolute volume, involves use of relative density (specific gravity) values for all the ingredients to calculate the absolute volume each will occupy in a unit volume of concrete. The absolute volume method will be illustrated. A concrete mixture also can be proportioned from field experience (statistical data) or from trial mixtures.

Other valuable documents to help proportion concrete mixtures include the Standard Practice for Selecting Proportions for Structural Lightweight Concrete (ACI 211.2); Guide for Selecting Proportions for No-Slump Concrete (ACI 211.3); Guide for Selecting Proportions for High-Strength Concrete with Portland Cement and Fly Ash (ACI 211.4R); and Guide for Submittal of Concrete Proportions (ACI 211.5). Hover (1995 and 1998) provides a graphical process for designing concrete mixtures in accordance with ACI 211.1.

**Proportioning from Field Data**

A presently or previously used concrete mixture design can be used for a new project if strength-test data and standard deviations show that the mixture is acceptable. Durability aspects previously presented must also be met. Standard deviation computations are outlined in ACI 318. The statistical data should essentially represent the same materials, proportions, and concreting conditions to be used in the new project. The data used for proportioning should also be from a concrete with an $f'_c$ that is within 7 MPa (1000 psi) of the strength required for the proposed work. Also, the data should represent at least 30 consecutive tests or two groups of consecutive tests totaling at least 30 tests (one test is the average strength of two cylinders from the same sample). If only 15 to 29 consecutive tests are available, an adjusted standard deviation can be

---

**Table 9-9. Maximum Chloride-Ion Content for Corrosion Protection**

<table>
<thead>
<tr>
<th>Type of member</th>
<th>Maximum water-soluble chloride ion (Cl⁻) in concrete, percent by mass of cement*</th>
</tr>
</thead>
<tbody>
<tr>
<td>Prestressed concrete (Classes C0, C1, C2)</td>
<td>0.06</td>
</tr>
<tr>
<td>Reinforced concrete exposed to chloride in service (Class C2)</td>
<td>0.15</td>
</tr>
<tr>
<td>Reinforced concrete that will be dry or protected from moisture in service (Class C0)</td>
<td>1.00</td>
</tr>
<tr>
<td>Other reinforced concrete construction (Class C1)</td>
<td>0.30</td>
</tr>
</tbody>
</table>

*ASTM C1218. Adapted from ACI 318.
obtained by multiplying the standard deviation \( S \) for the 15 to 29 tests and a modification factor from Table 9-10. The data must represent 45 or more days of tests.

The standard or modified deviation is then used in Equations 9-1 to 9-3. The average compressive strength from the test record must equal or exceed the ACI 318 required average compressive strength, \( f'_{cr} \), in order for the concrete proportions to be acceptable. The \( f'_{cr} \) for the selected mixture proportions is equal to the larger of Equations 9-1 and 9-2 (for \( f'_c \leq 35 \) MPa [5000 psi]) or Equations 9-1 and 9-3 (for \( f'_c > 35 \) MPa [5000 psi]).

\[
\begin{align*}
\frac{f'_c}{S} &= \frac{f'_c}{S} + 3.45 \\

f'_{cr} &= f'_c + 2.33S - 3.45 \text{ (MPa)} \\

f'_{cr} &= f'_c + 2.33S - 500 \text{ (psi)} \\

f'_{cr} &= 0.90 f'_c + 2.33S \\

\end{align*}
\]

where

\( f'_{cr} \) = required average compressive strength of concrete used as the basis for selection of concrete proportions, MPa (psi)

\( f'_c \) = specified compressive strength of concrete, MPa (psi)

\( S \) = standard deviation, MPa (psi)

When field strength test records do not meet the previously discussed requirements, \( f'_{cr} \) can be obtained from Table 9-11. A field strength record, several strength test records, or tests from trial mixtures must be used for documentation showing that the average strength of the mixture is equal to or greater than \( f'_{cr} \).

If less than 30, but not less than 10 tests are available, the tests may be used for average strength documentation if the time period is not less than 45 days. Mixture proportions may also be established by interpolating between two or more test records if each meets the above and project requirements. If a significant difference exists between the mixtures that are used in the interpolation, a trial mixture should be considered to check strength gain. If the test records meet the above requirements and limitations of ACI 318, the proportions for the mixture may then be considered acceptable for the proposed work.

If the average strength of the mixtures with the statistical data is less than \( f'_{cr} \), or statistical data or test records are insufficient or not available, the mixture should be proportioned by the trial-mixture method. The approved mixture must have a compressive strength that meets or exceeds \( f'_{cr} \). Three trial mixtures using three different water to cementing materials ratios or cementing materials contents should be tested. A water to cementing materials ratio to strength curve (similar to Fig. 9-2) can then be plotted and the proportions interpolated from the data. It is also good practice to test the properties of the newly proportioned mixture in a trial batch.

ACI 214 provides statistical analysis methods for monitoring the strength of the concrete in the field to ensure that the mix properly meets or exceeds the design strength, \( f'_c \).

### Table 9-10. Modification Factor for Standard Deviation When Less Than 30 Tests Are Available

<table>
<thead>
<tr>
<th>Number of tests*</th>
<th>Modification factor for standard deviation**</th>
</tr>
</thead>
<tbody>
<tr>
<td>Less than 15</td>
<td>Use Table 9-11</td>
</tr>
<tr>
<td>15</td>
<td>1.16</td>
</tr>
<tr>
<td>20</td>
<td>1.08</td>
</tr>
<tr>
<td>25</td>
<td>1.03</td>
</tr>
<tr>
<td>30 or more</td>
<td>1.00</td>
</tr>
</tbody>
</table>

* Interpolate for intermediate numbers of tests.

** Modified standard deviation to be used to determine required average strength, \( f'_{cr} \).

Adapted from ACI 318.

### Table 9-11 (Metric). Required Average Compressive Strength When Data Are Not Available to Establish a Standard Deviation

<table>
<thead>
<tr>
<th>Specified compressive strength, ( f'_c ), MPa</th>
<th>Required average compressive strength, ( f'_{cr} ), MPa</th>
</tr>
</thead>
<tbody>
<tr>
<td>Less than 21</td>
<td>( f'_c + 7.0 )</td>
</tr>
<tr>
<td>21 to 35</td>
<td>( f'_c + 8.5 )</td>
</tr>
<tr>
<td>Over 35</td>
<td>( 1.10 f'_c + 5.0 )</td>
</tr>
</tbody>
</table>

Adapted from ACI 318.

### Table 9-11 (Inch-Pound Units). Required Average Compressive Strength When Data Are Not Available to Establish a Standard Deviation

<table>
<thead>
<tr>
<th>Specified compressive strength, ( f'_c ), psi</th>
<th>Required average compressive strength, ( f'_{cr} ), psi</th>
</tr>
</thead>
<tbody>
<tr>
<td>Less than 3000</td>
<td>( f'_c + 1000 )</td>
</tr>
<tr>
<td>3000 to 5000</td>
<td>( f'_c + 1200 )</td>
</tr>
<tr>
<td>Over 5000</td>
<td>( 1.10 f'_c + 700 )</td>
</tr>
</tbody>
</table>

Adapted from ACI 318.
the concrete should be determined by testing the cylinders in compression. The test results should be plotted to produce a strength versus water-cementing materials ratio curve (similar to Fig. 9-2) that is used to proportion a mixture.

A number of different methods of proportioning concrete ingredients have been used at one time or another, including:

- Arbitrary assignment (1:2:3), volumetric
- Void ratio
- Fineness modulus
- Surface area of aggregates
- Cement content

Any one of these methods can produce approximately the same final mixture after adjustments are made in the field. The best approach, however, is to select proportions based on past experience and reliable test data with an established relationship between strength and water to cementing materials ratio for the materials to be used in the concrete. The trial mixtures can be relatively small batches made with laboratory precision or job-size batches made during the course of normal concrete production. Use of both is often necessary to reach a satisfactory job mixture.

The following parameters must be selected first:
1. required strength,
2. minimum cementing materials content or maximum water-cementing materials ratio,
3. nominal maximum size of aggregate,
4. air content,
5. desired slump.

Trial batches are then made varying the relative amounts of fine and coarse aggregates as well as other ingredients. Based on considerations of workability and economy, the proper mixture proportions are selected.

When the quality of the concrete mixture is specified by water-cementitious material ratio, the trial-batch procedure consists essentially of combining a paste (water, cementing materials, and, generally, a chemical admixture) of the correct proportions with the necessary amounts of fine and coarse aggregates to produce the required slump and workability. Representative samples of the cementing materials, water, aggregates, and admixtures must be used.

Quantities per cubic meter (cubic yard) are then calculated. To simplify calculations and eliminate error caused by variations in aggregate moisture content, the aggregates should be prewetted then dried to a saturated surface-dry (SSD) condition; place the aggregates in covered containers to keep them in this SSD condition until they are used. The moisture content of the aggregates should be determined and the batch weights corrected accordingly.

The size of the trial batch is dependent on the equipment available and on the number and size of test specimens to be made. Larger batches will produce more accurate data. Machine mixing is recommended since it more nearly represents job conditions; it is mandatory if the concrete is to contain entrained air. The mixing procedures outlined in ASTM C192 (AASHTO T 126) should be used.

### Measurements and Calculations

Tests for slump, air content, and temperature should be made on the trial mixture, and the following measurements and calculations should also be performed:

**Density (Unit Weight) and Yield.** The density (unit weight) of freshly mixed concrete is expressed in kilograms per cubic meter (pounds per cubic foot). The yield is the volume of fresh concrete produced in a batch, usually expressed in cubic meters (cubic feet). The yield is calculated by dividing the total mass of the materials batched by the density of the freshly mixed concrete. Density and yield are determined in accordance with ASTM C138.

**Absolute Volume.** The absolute volume of a granular material (such as cement and aggregates) is the volume of the solid matter in the particles; it does not include the volume of air spaces between particles. The volume (yield) of freshly mixed concrete is equal to the sum of the absolute volumes of the concrete ingredients—cementing materials, water (exclusive of that absorbed in the aggregate), aggregates, admixtures when applicable, and air. The absolute volume is computed from a material’s mass and relative density (specific gravity) as follows:

\[
\text{Absolute volume} = \frac{\text{mass of loose material}}{\text{relative density of a material} \times \text{density of water}}
\]

A value of 3.15 can be used for the relative density (specific gravity) of portland cement. Blended cements have relative densities ranging from 2.90 to 3.15. The relative density of fly ash varies from 1.9 to 2.8, slag from 2.85 to 2.95, and silica fume from 2.20 to 2.25. The relative density of water is 1.0 and the density of water is

<table>
<thead>
<tr>
<th>Temperature, °C</th>
<th>Density, kg/m³</th>
<th>Temperature, °F</th>
<th>Density, lb/ft³</th>
</tr>
</thead>
<tbody>
<tr>
<td>16</td>
<td>998.93</td>
<td>60</td>
<td>62.368</td>
</tr>
<tr>
<td>18</td>
<td>998.58</td>
<td>65</td>
<td>62.337</td>
</tr>
<tr>
<td>20</td>
<td>998.19</td>
<td>70</td>
<td>62.302</td>
</tr>
<tr>
<td>22</td>
<td>997.75</td>
<td>75</td>
<td>62.261</td>
</tr>
<tr>
<td>24</td>
<td>997.27</td>
<td>80</td>
<td>62.216</td>
</tr>
<tr>
<td>26</td>
<td>996.75</td>
<td>85</td>
<td>62.166</td>
</tr>
<tr>
<td>28</td>
<td>996.20</td>
<td></td>
<td></td>
</tr>
<tr>
<td>30</td>
<td>995.61</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Table 9-12. Density of Water Versus Temperature
1000 kg/m³ (62.4 lb/ft³) at 4°C (39°F)—accurate enough for mix calculations at room temperature. More accurate water density values are given in Table 9-12. Relative density of normal aggregate usually ranges between 2.4 and 2.9.

The relative density of aggregate as used in mix-design calculations is the relative density of either saturated surface-dry (SSD) material or oven-dry material. Relative densities of admixtures, such as water reducers, can also be considered if needed. Absolute volume is usually expressed in cubic meters (cubic feet).

The absolute volume of air in concrete, expressed as cubic meters per cubic meter (cubic feet per cubic yard), is equal to the total air content in percent divided by 100 (for example, 7% ÷ 100) and then multiplied by the volume of the concrete batch.

The volume of concrete in a batch can be determined by either of two methods: (1) if the relative densities of the aggregates and cementing materials are known, these can be used to calculate concrete volume; or (2) if relative densities are unknown, or they vary, the volume can be computed by dividing the total mass of materials in the mixer by the density of concrete. In some cases, both determinations are made, one serving as a check on the other.

EXAMPLES OF MIXTURE PROPORTIONING

Example 1. Absolute Volume Method (Metric)

Conditions and Specifications. Concrete is required for a pavement that will be exposed to moisture in a severe freeze-thaw environment. A specified compressive strength, $f'_c$, of 35 MPa is required at 28 days. Air entrainment is required. Slump should be between 25 mm and 75 mm. A nominal maximum size aggregate of 25 mm is required. No statistical data on previous mixes are available. The materials available are as follows:

**Cement:** Type GU (ASTM C1157) with a relative density of 3.0.

**Coarse aggregate:** Well-graded, 25-mm nominal maximum-size rounded gravel (ASTM C33 or AASHTO M 80) with an oven-dry relative density of 2.68, absorption of 0.5% (moisture content at SSD condition) and oven-dry rodded bulk density (unit weight) of 1600 kg/m³. The laboratory sample for trial batching has a moisture content of 2%.

**Fine aggregate:** Natural sand (ASTM C33 or AASHTO M 6) with an oven-dry relative density of 2.64 and absorption of 0.7%. The laboratory sample moisture content is 6%. The fineness modulus is 2.80.

Air-entraining admixture: Wood-resin type (ASTM C260 or AASHTO M 154).

Water reducer: ASTM C494 (AASHTO M 194). This particular admixture is known to reduce water demand by 10% when used at a dosage rate of 3 g (or 3 mL) per kg of cement. Assume that the chemical admixtures have a density close to that of water, meaning that 1 mL of admixture has a mass of 1 g.

From this information, the task is to proportion a trial mixture that will meet the above conditions and specifications.

**Strength.** The design strength of 35 MPa is greater than the 31 MPa required in Table 9-1 for the exposure condition. Since no statistical data is available, $f'_c$ (required compressive strength for proportioning) from Table 9-11 is equal to $f'_c + 8.5$. Therefore, $f'_c = 35 + 8.5 = 43.5$ MPa.

**Water to Cement Ratio.** For an environment with moist freezing and thawing, the maximum water to cementitious material ratio should be 0.45. The recommended water to cementitious material ratio for an $f'_c$ of 43.5 MPa is 0.31 from Fig. 9-2 or interpolated from Table 9-3

\[
\frac{(45 - 43.5)(0.34 - 0.30)}{45 - 40} + 0.30 = 0.31
\]

Since the lower water to cement ratio governs, the mix must be designed for 0.31. If a plot from trial batches or field tests had been available, the water to cement ratio could have been extrapolated from that data.

**Air Content.** For a severe freeze-thaw exposure, Table 9-5 recommends a target air content of 6.0% for a 25-mm aggregate. Therefore, design the mix for 5% to 8% air and use 8% (or the maximum allowable) for batch proportions. The trial-batch air content must be within ±0.5 percentage points of the maximum allowable air content.

**Slump.** The slump is specified at 25 mm to 75 mm. Use 75 mm ± 20 mm for proportioning purposes.

**Water Content.** Table 9-5 and Fig. 9-5 recommend that a 75-mm slump, air-entrained concrete made with 25-mm nominal maximum-size aggregate should have a water content of about 175 kg/m³. However, rounded gravel should reduce the water content of the table value by about 25 kg/m³. Therefore, the water content can be estimated to be about 150 kg/m³ (175 kg/m³ minus 25 kg/m³). In addition, the water reducer will reduce water demand by 10% resulting in an estimated water demand of 135 kg/m³.

**Cement Content.** The cement content is based on the maximum water-cement ratio and the water content. Therefore, 135 kg/m³ of water divided by a water-cement ratio of 0.31 requires a cement content of 435 kg/m³; this is greater than the 335 kg/m³ required for frost resistance (Table 9-7).
Coarse-Aggregate Content. The quantity of 25-mm nominal maximum-size coarse aggregate can be estimated from Fig. 9-3 or Table 9-4. The bulk volume of coarse aggregate recommended when using sand with a fineness modulus of 2.80 is 0.67. Since it has a bulk density of 1600 kg/m³, the oven-dry mass of coarse aggregate for a cubic meter of concrete is
\[
1600 \times 0.67 = 1072 \text{ kg}
\]

Admixture Content. For an 8% air content, the air-entraining admixture manufacturer recommends a dosage rate of 0.5 g per kg of cement. From this information, the amount of air-entraining admixture per cubic meter of concrete is
\[
0.5 \times 435 = 218 \text{ g or } 0.218 \text{ kg}
\]
The water reducer dosage rate of 3 g per kg of cement results in
\[
3 \times 435 = 1305 \text{ g or } 1.305 \text{ kg}
\]
Fine-Aggregate Content. At this point, the amounts of all ingredients except the fine aggregate are known. In the absolute volume method, the volume of fine aggregate is determined by subtracting the absolute volumes of the known ingredients from 1 cubic meter. The absolute volume of the water, cement, admixtures and coarse aggregate is calculated by dividing the known mass of each by the product of their relative density and the density of water. Volume computations are as follows:

\[
\begin{align*}
\text{Water} & = \frac{135}{1 \times 1000} = 0.135 \text{ m}^3 \\
\text{Cement} & = \frac{435}{3.0 \times 1000} = 0.145 \text{ m}^3 \\
\text{Air} & = \frac{8.0}{100} = 0.080 \text{ m}^3 \\
\text{Coarse aggregate} & = \frac{1072}{2.68 \times 1000} = 0.400 \text{ m}^3 \\
\text{Total volume of known ingredients} & = 0.760 \text{ m}^3 \\
\end{align*}
\]
The calculated absolute volume of fine aggregate is then
\[
1 - 0.76 = 0.24 \text{ m}^3
\]
The mass of dry fine aggregate is
\[
0.24 \times 2.64 \times 1000 = 634 \text{ kg}
\]
The mixture then has the following proportions before trial mixing for one cubic meter of concrete:

\[
\begin{align*}
\text{Water} & = 135 \text{ kg} \\
\text{Cement} & = 435 \text{ kg} \\
\text{Coarse aggregate (dry)} & = 1072 \text{ kg} \\
\text{Fine aggregate (dry)} & = 634 \text{ kg} \\
\text{Total mass} & = 2276 \text{ kg} \\
\text{Air-entraining admixture} & = 0.218 \text{ kg} \\
\text{Water reducer} & = 1.305 \text{ kg} \\
\end{align*}
\]
Slump 75 mm (±20 mm for trial batch)

Estimated concrete 8% (±0.5% for trial batch)

Air content 8% (±0.5% for trial batch)

Estimated concrete density (using SSD aggregate) = 2286 kg/m³

The liquid admixture volume is generally too insignificant to include in the water calculations. However, certain admixtures, such as shrinkage reducers, plasticizers, and corrosion inhibitors are exceptions due to their relatively large dosage rates; their volumes should be included.

Moisture. Corrections are needed to compensate for moisture in and on the aggregates. In practice, aggregates will contain some measurable amount of moisture. The dry-batch weights of aggregates, therefore, have to be increased to compensate for the moisture that is absorbed in and contained on the surface of each particle and between particles. The mixing water added to the batch must be reduced by the amount of free moisture contributed by the aggregates. Tests indicate that for this example, coarse-aggregate moisture content is 2% and fine-aggregate moisture content is 6%.

With the aggregate moisture contents (MC) indicated, the trial batch aggregate proportions become

Coarse aggregate (2% MC) = 1072 x 1.02 = 1093 kg

Fine aggregate (6% MC) = 634 x 1.06 = 672 kg

Water absorbed by the aggregates does not become part of the mixing water and must be excluded from the water adjustment. Surface moisture contributed by the coarse aggregate amounts to 2% - 0.5% = 1.5%; that contributed by the fine aggregate is 6% – 0.7% = 5.3%. The estimated requirement for added water becomes
\[
135 - (1072 x 0.015) - (634 x 0.053) = 85 \text{ kg}
\]
The estimated batch weights for one cubic meter of concrete are revised to include aggregate moisture as follows:

\[
\begin{align*}
\text{Water (to be added)} & = 85 \text{ kg} \\
\text{Cement} & = 435 \text{ kg} \\
\text{Coarse aggregate (2% MC, wet)} & = 1093 \text{ kg} \\
\text{Fine aggregate (6% MC, wet)} & = 672 \text{ kg} \\
\text{Total} & = 2285 \text{ kg} \\
\text{Air-entraining admixture} & = 0.218 \text{ kg} \\
\text{Water reducer} & = 1.305 \text{ kg} \\
\end{align*}
\]

Trial Batch. At this stage, the estimated batch weights should be checked by means of trial batches or by full-size field batches. Enough concrete must be mixed for appropriate air and slump tests and for casting the three cylinders required for 28-day compressive-strength tests, plus beams for flexural tests if necessary. For a laboratory trial batch it is convenient, in this case, to scale down the weights to produce 0.1 m³ of concrete as follows:

\[
\begin{align*}
\text{Water (to be added)} & = 135 + 435 + (1072 x 1.005^*) \\
\text{Cement} & = + (634 x 1.007^*) \\
\text{SSD aggregate} & = 2286 \text{ kg/m}^3 \\
\end{align*}
\]

*(0.5% absorption ÷ 100) + 1 = 1.005

*(0.7% absorption ÷ 100) + 1 = 1.007

Chapter 9 • Designing and Proportioning Normal Concrete Mixtures
Batch and reduce the water content by 2 kg/m³ for each 10 mm reduction in slump. The adjusted mixture water for the reduced slump and air content is

\[(3 \text{ kg water} \times 1 \text{ percentage point difference for air}) – (2 \text{ kg water} \times 25/10 \text{ for slump change}) + 129 = 127 \text{ kg of water}\]

With less mixing water needed in the trial batch, less cement also is needed to maintain the desired water-cement ratio of 0.31. The new cement content is

\[\frac{127}{0.31} = 410 \text{ kg}\]

The amount of coarse aggregate remains unchanged because workability is satisfactory. The new adjusted batch weights based on the new cement and water contents are calculated after the following volume computations:

<table>
<thead>
<tr>
<th>Material</th>
<th>Weight (kg)</th>
<th>Density (kg/m³)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Water</td>
<td>127</td>
<td>1 x 1000</td>
</tr>
<tr>
<td>Cement</td>
<td>410</td>
<td>3.0 x 1000</td>
</tr>
<tr>
<td>Coarse aggregate (dry)</td>
<td>1072</td>
<td>2.68 x 1000</td>
</tr>
<tr>
<td>Fine aggregate (dry)</td>
<td>676</td>
<td>1.06 x 1000</td>
</tr>
<tr>
<td>Air</td>
<td>8</td>
<td>100</td>
</tr>
<tr>
<td>Total</td>
<td>2285</td>
<td></td>
</tr>
</tbody>
</table>

The mixing water required for a cubic meter of the same slump concrete as the trial batch is

\[\frac{12.97}{0.10026} = 129 \text{ kg}\]

Batch Adjustments. The measured 100-mm slump of the trial batch is unacceptable (above 75 mm ±20 mm max.), the yield was slightly high, and the 9.0% air content as measured in this example is also too high (more than 0.5% above 8.5% max.). Adjust the yield and reestimate the amount of air-entraining admixture required for an 8% air content and adjust the water to obtain a 75-mm slump. Increase the mixing water content by 3 kg/m³ for each 1% by which the air content is decreased from that of the trial batch. The above concrete, when mixed, had a measured slump of 100 mm, an air content of 9%, and a density of 2274 kg per cubic meter. During mixing, some of the pre-measured water may remain unused or additional water may be added to approach the required slump. In this example, although 8.5 kg of water was calculated to be added, the trial batch actually used only 8.0 kg. The mixture excluding admixtures therefore becomes

<table>
<thead>
<tr>
<th>Material</th>
<th>Weight (kg)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Water</td>
<td>8.0</td>
</tr>
<tr>
<td>Cement</td>
<td>43.5</td>
</tr>
<tr>
<td>Coarse aggregate (2% MC)</td>
<td>109.3</td>
</tr>
<tr>
<td>Fine aggregate (6% MC)</td>
<td>67.2</td>
</tr>
<tr>
<td>Total</td>
<td>228.0</td>
</tr>
</tbody>
</table>

The yield of the trial batch is

\[\frac{228.0 \text{ kg}}{2274 \text{ kg/m}³} = 0.10026 \text{ m}³\]

The mixing water content is determined from the added water plus the free water on the aggregates and is calculated as follows:

- Water added: 8.0 kg
- Free water on coarse aggregate: \[\frac{109.3}{1.02} \times 0.015 = 1.61 \text{ kg}\]
- Free water on fine aggregate: \[\frac{67.2}{1.06} \times 0.053 = 3.36 \text{ kg}\]
- Total water: 12.97 kg

The mixing water required for a cubic meter of the same slump concrete as the trial batch is

\[\frac{12.97}{0.10026} = 129 \text{ kg}\]

*(2% MC – 0.5% absorption) ÷ 100 = 0.015
(6% MC – 0.7% absorption) ÷ 100 = 0.053*
ally, the proportion of fine to coarse aggregate is kept constant in adjusting the batch weights to maintain workability or other properties obtained in the first trial batch. After adjustments to the cementitious materials, water, and air content have been made, the volume remaining for aggregate is appropriately proportioned between the fine and coarse aggregates.

Additional trial concrete mixtures with water-cement ratios above and below 0.31 should also be tested to develop a strength to water-cement ratio relationship. From that data, a new more economical mixture with a compressive strength closer to $f'_c$, and a lower cement content can be proportioned and tested. The final mixture would probably look similar to the above mixture with a slump range of 25 mm to 75 mm and an air content of 5% to 8%. The amount of air-entraining admixture must be adjusted to field conditions to maintain the specified air content.

**Example 2. Absolute Volume Method (Inch-Pound Units)**

**Conditions and Specifications.** Concrete is required for a building foundation. A specified compressive strength, $f'_c$, of 3500 psi is required at 28 days using a Type I cement. The design calls for a minimum of 3 in. of concrete cover over the reinforcing steel. The minimum distance between reinforcing bars is 4 in. The only admixture allowed is for air entrainment. No statistical data on previous mixes are available. The materials available are as follows:

- **Cement:** Type I, ASTM C150, with a relative density of 3.15.
- **Coarse aggregate:** Well-graded ¾-in. maximum-size gravel containing some crushed particles (ASTM C33) with an oven-dry relative density (specific gravity) of 2.68, absorption of 0.5% (moisture content at SSD condition) and oven-dry rodded bulk density (unit weight) of 100 lb per cu ft. The laboratory sample for trial batching has a moisture content of 2%.
- **Fine aggregate:** Natural sand (ASTM C33) with an oven-dry relative density (specific gravity) of 2.64 and absorption of 0.7%. The laboratory sample moisture content is 6%. The fineness modulus is 2.80.
- **Air-entraining admixture:** Wood-resin type, ASTM C260.

From this information, the task is to proportion a trial mixture that will meet the above conditions and specifications.

**Strength.** Since no statistical data is available, $f'_c$ (required compressive strength for proportioning) from Table 9-11 is equal to $f'_c + 1200$. Therefore, $f'_c = 3500 + 1200 = 4700$ psi.

**Water to Cement Ratio.** Table 9-1 requires no maximum water to cement ratio. The recommended water to cement ratio for an $f'_c$ of 4700 psi is 0.42 interpolated from Fig. 9-2 or Table 9-3 [water to cement ratio = [(5000 – 4700)(0.48 – 0.40)/(5000 – 4000)] + 0.40 = 0.42].

**Coarse-Aggregate Size.** From the specified information, a ¾-in. nominal maximum-size aggregate is adequate as it is less than ¾ of the distance between reinforcing bars and between the rebars and forms (cover).

**Air Content.** A target air content of 6.0% is specified in this instance not for exposure conditions but to improve workability and reduce bleeding. Therefore, design the mix for 6% ±1.0% air and use 7% (or the maximum allowable) for batch proportions. The trial batch air content must be within ±0.5 percentage points of the maximum allowable air content.

**Slump.** As no slump was specified, a slump of 1 to 3 in. would be adequate as indicated by Table 9-6. Use 3 in. for proportioning purposes, the maximum recommended for foundations.

**Water Content.** Fig. 9-5 and Table 9-5 recommend that a 3-in. slump, air-entrained concrete made with ¾-in. nominal maximum-size aggregate should have a water content of about 305 lb per cu yd. However, gravel with some crushed particles should reduce the water content of the table value by about 35 lb. Therefore, the water content can be estimated to be about 305 lb minus 35 lb, which is 270 lb.

**Cement Content.** The cement content is based on the maximum water-cement ratio and the water content. Therefore, 270 lb of water divided by a water-cement ratio of 0.42 requires a cement content of 643 lb.

**Coarse-Aggregate Content.** The quantity of ¾-in. nominal maximum-size coarse aggregate can be estimated from Fig. 9-3 or Table 9-4. The bulk volume of coarse aggregate recommended when using sand with a fineness modulus of 2.80 is 0.62. Since it weighs 100 lb per cu ft, the oven-dry weight of coarse aggregate for a cubic yard of concrete (27 cu ft) is

$$100 \times 27 \times 0.62 = 1674 \text{ lb per cu yd of concrete}$$

**Admixture Content.** For a 7% air content, the air-entraining admixture manufacturer recommends a dosage rate of 0.9 fl oz per 100 lb of cement. From this information, the amount of air-entraining admixture is

$$0.9 \times \frac{643}{100} = 5.8 \text{ fl oz per cu yd}$$

**Fine-Aggregate Content.** At this point, the amount of all ingredients except the fine aggregate are known. In the absolute volume method, the volume of fine aggregate is determined by subtracting the absolute volumes of the known ingredients from 27 cu ft (1 cu yd). The absolute volume of the water, cement, and coarse aggregate is cal-
culated by dividing the known weight of each by the product of their relative density (specific gravity) and the density of water. Volume computations are as follows:

- **Water**: 
  \[
  \frac{270}{1 \times 62.4} = 4.33 \text{ cu ft}
  \]

- **Cement**: 
  \[
  \frac{643}{3.15 \times 62.4} = 3.27 \text{ cu ft}
  \]

- **Air**: 
  \[
  \frac{7.0}{100} \times 27 = 1.89 \text{ cu ft}
  \]

- **Coarse aggregate**: 
  \[
  \frac{1674}{2.68 \times 62.4} = 10.01 \text{ cu ft}
  \]

- **Total volume of known ingredients**: 
  \[
  19.50 \text{ cu ft}
  \]

The liquid admixture volume is generally too insignificant to include in these calculations. However, certain admixtures such as shrinkage reducers, plasticizers, and corrosion inhibitors are exceptions due to their relatively large dosage rates; their volumes should be included.

The calculated absolute volume of fine aggregate is then 
\[
27 - 19.50 = 7.50 \text{ cu ft}
\]

The weight of dry fine aggregate is 
\[
7.50 \times 2.64 \times 62.4 = 1236 \text{ lb}
\]

The mixture then has the following proportions before trial mixing for one cubic yard of concrete:

- **Water**: 270 lb
- **Cement**: 643 lb
- **Coarse aggregate (dry)**: 1674 lb
- **Fine aggregate (dry)**: 1236 lb
- **Total weight**: 3823 lb
- **Air-entraining admixture**: 5.8 fl oz
- **Slump**: 3 in. (±¾ in. for trial batch)
- **Air content**: 7% (±0.5% for trial batch)
- **Estimated density**: \[
\frac{[270 + 643 + (1674 \times 1.005*) + (1236 \times 1.007*)]}{27} = 142.22 \text{ lb per cubic foot}
\]

**Moisture.** Corrections are needed to compensate for moisture in the aggregates. In practice, aggregates will contain some measurable amount of moisture. The dry-batch weights of aggregates, therefore, have to be increased to compensate for the moisture that is absorbed in and contained on the surface of each particle and between particles. The mixing water added to the batch must be reduced by the amount of free moisture contributed by the aggregates. Tests indicate that for this example, coarse-aggregate moisture content is 2% and fine-aggregate moisture content is 6%.

With the aggregate moisture contents (MC) indicated, the trial batch aggregate proportions become

- **Coarse aggregate (2% MC)**: \(1674 \times 1.02 = 1707 \text{ lb}\)
- **Fine aggregate (6% MC)**: \(1236 \times 1.06 = 1310 \text{ lb}\)

Water absorbed by the aggregates does not become part of the mixing water and must be excluded from the water adjustment. Surface moisture contributed by the coarse aggregate amounts to 2% – 0.5% = 1.5%; that contributed by the fine aggregate is 6% – 0.7% = 5.3%. The estimated requirement for added water becomes 
\[
270 - (1674 \times 0.015) - (1236 \times 0.053) = 179 \text{ lb}
\]

The estimated batch weights for one cubic yard of concrete are revised to include aggregate moisture as follows:

- **Water (to be added)**: 179 lb
- **Cement**: 643 lb
- **Coarse aggregate (2% MC, wet)**: 1707 lb
- **Fine aggregate (6% MC, wet)**: 1310 lb
- **Total**: 3839 lb
- **Air-entraining admixture**: 5.8 fl oz

**Trial Batch.** At this stage, the estimated batch weights should be checked by means of trial batches or by full-size field batches. Enough concrete must be mixed for appropriate air and slump tests and for casting the three cylinders required for compressive-strength tests at 28 days. For a laboratory trial batch it is convenient, in this case, to scale down the weights to produce 2.0 cu ft of concrete or \(\frac{2}{27}\) cu yd.

- **Water**: \(270 \times \frac{2}{27} = 13.26 \text{ lb}\)
- **Cement**: \(643 \times \frac{2}{27} = 47.63 \text{ lb}\)
- **Coarse aggregate (wet)**: \(1707 \times \frac{2}{27} = 126.44 \text{ lb}\)
- **Fine aggregate (wet)**: \(1310 \times \frac{2}{27} = 97.04 \text{ lb}\)
- **Total**: 284.37 lb

**Air-entraining admixture**: 5.8 fl oz

[Laboratories often convert fluid ounces to milliliters by multiplying fluid ounces by 29.57353 to improve measurement accuracy. Also, most laboratory pipets used for measuring fluids are graduated in milliliter units]

The above concrete, when mixed, had a measured slump of 4 in., an air content of 8%, and a density (unit weight) of 141.49 lb per cubic foot. During mixing, some of the premeasured water may remain unused or additional water may be added to approach the required slump. In this example, although 13.26 lb of water was calculated to be added, the trial batch actually used only 13.12 lb. The mixture excluding admixture therefore becomes:
The yield of the trial batch is

\[
\frac{284.23}{141.49} = 2.009 \text{ cu ft}
\]

The mixing water content is determined from the added water plus the free water on the aggregates and is calculated as follows:

- Water added = 13.12 lb
- Free water on coarse aggregate = \(\frac{126.44}{1.02} \times 0.015\) = 1.86 lb
- Free water on fine aggregate = \(\frac{97.04}{1.06} \times 0.053\) = 4.85 lb
- Total = 19.83 lb

The mixing water required for a cubic yard of the same slump concrete as the trial batch is

\[
19.83 \times 27 = 267 \text{ lb}
\]

**Batch Adjustments.** The measured 4-in. slump of the trial batch is unacceptable (more than 0.75 in. above 3-in. max.), the yield was slightly high, and the 8.0% air content as measured in this example is also too high (more than 0.5% above 7% max.). Adjust the yield, reestimate the amount of air-entraining admixture required for a 7% air content, and adjust the water to obtain a 3-in. slump. Increase the mixing water content by 5 lb for each 1% by which the air content is decreased from that of the trial batch and reduce the water content by 10 lb for each 1-in. reduction in slump. The adjusted mixture water for the reduced slump and air content is

\[
(5 \times 1) - (10 \times 1) + 267 = 262 \text{ lb per cu yd}
\]

With less mixing water needed in the trial batch, less cement also is needed to maintain the desired water-cement ratio of 0.42. The new cement content is

\[
\frac{262}{0.42} = 624 \text{ lb per cu yd}
\]

The amount of coarse aggregate remains unchanged because workability is satisfactory. The new adjusted batch weights based on the new cement and water contents are calculated after the following volume computations:

- Water = \(\frac{262}{1 \times 62.4}\) = 4.20 cu ft
- Cement = \(\frac{624}{3.15 \times 62.4}\) = 3.17 cu ft
- Coarse aggregate = \(\frac{1674}{2.68 \times 62.4}\) = 10.01 cu ft
- Air = \(\frac{7.0}{100} \times 27\) = 1.89 cu ft
- Total = 19.27 cu ft

The weight of dry fine aggregate required is

\[
7.73 \times 2.64 \times 62.4 = 1273 \text{ lb}
\]

An air-entraining admixture dosage of 0.8 fluid ounces per 100 pounds of cement is expected to achieve the 7% air content in this example. Therefore, the amount of air-entraining admixture required is:

\[
\frac{0.8 \times 624}{100} = 5.0 \text{ fl oz}
\]

Adjusted batch weights per cubic yard of concrete are

- Water = 262 lb
- Cement = 624 lb
- Coarse aggregate = 1674 lb (dry)
- Fine aggregate = 1273 lb (dry)
- Air-entraining admixture = 5.0 fl oz

Estimated concrete density (unit weight) with the aggregates at SSD:

\[
\frac{[262 + 624 + (1674 \times 1.005) + (1273 \times 1.007)]}{27} = 142.60 \text{ lb per cu ft}
\]

Upon completion of checking these adjusted proportions in a trial batch, it was found that the proportions were adequate for the desired slump, air content, and yield. The 28-day test cylinders had an average compressive strength of 4900 psi, which exceeds the \(f_{cr}\) of 4700 psi. Due to fluctuations in moisture content, absorption rates, and specific gravity of the aggregate, the density determined by volume calculations may not always equal the unit weight determined by ASTM C138 (AASHTO T 121). Occasionally, the proportion of fine to coarse aggregate is kept constant in adjusting the batch weights to maintain workability or other properties obtained in the first trial batch. After adjustments to the cement, water, and air content have been made, the volume remaining for aggregate is appropriately proportioned between the fine and coarse aggregates.

Additional trial concrete mixtures with water-cement ratios above and below 0.42 should also be tested to develop a strength curve. From the curve, a new more economical mixture with a compressive strength closer to \(f_{cr}\) can be proportioned and tested. The final mixture would probably look similar to the above mixture with a slump range of 1 in. to 3 in. and an air content of 5% to 7%. The

\*1 + (2% MC/100) = 1.02; 1 + (6% MC/100) = 1.06;
** (2% MC – 0.5% absorption)/100 = 0.015; (6% MC – 0.7% absorption)/100 = 0.053
amount of air-entraining admixture must be adjusted to field conditions to maintain the specified air content.

**Water Reducers.** Water reducers are used to increase workability without the addition of water or to reduce the water-cement ratio of a concrete mixture to improve permeability or other properties.

Using the final mixture developed in the last example, assume that the project engineer approves the use of a water reducer to increase the slump to 5 in. to improve workability for a difficult placement area. Assuming that the water reducer has a manufacturer’s recommended dosage rate of 4 oz per 100 lb of cement to increase slump 2 in., the admixture amount becomes

\[
\frac{624}{100} \times 4 = 25.0 \text{ oz per cu yd}
\]

The amount of air-entraining agent may also need to be reduced (up to 50%), as many water reducers also entrain air. If a water reducer was used to reduce the water-cement ratio, the water and sand content would also need adjustment.

**Pozzolans and Slag.** Pozzolans and slag are sometimes added in addition to or as a partial replacement of cement to aid in workability and resistance to sulfate attack and alkali reactivity. If a pozzolan or slag were required for the above example mixture, it would have been entered in the first volume calculation used in determining fine aggregate content. For example:

Assume that 75 lb of fly ash with a relative density (specific gravity) of 2.5 were to be used in addition to the originally derived cement content. The ash volume would be

\[
\frac{75}{2.5 \times 62.4} = 0.48 \text{ cu ft}
\]

The water to cementing materials ratio would be

\[
\frac{W}{C + P} = \frac{270}{643 + 75} = 0.38\text{ by weight}
\]

The water to portland cement only ratio would still be

\[
\frac{W}{C} = \frac{270}{643} = 0.42\text{ by weight}
\]

The fine aggregate volume would have to be reduced by 0.48 cu ft to allow for the volume of ash.

The pozzolan amount and volume computation could also have been derived in conjunction with the first cement content calculation using a water to cementing materials ratio of 0.42 (or equivalent). For example, assume 15% of the cementitious material is specified to be a pozzolan and

\[
\frac{W/CM}{W/(C + P)} = 0.42.
\]

Then with

\[
W = 270 \text{ lb and } C + P = 643 \text{ lb,}
\]

\[
P = 643 \times \frac{15}{100} = 96 \text{ lb}
\]

and

\[
C = 643 - 96 = 547 \text{ lb}
\]

Appropriate proportioning computations for these and other mix ingredients would follow.

Example 3. Laboratory Trial Mixture

Using the PCA Water-Cement Ratio Method (Metric)

With the following method, the mix designer develops the concrete proportions directly from the laboratory trial batch rather than the absolute volume of the constituent ingredients.

**Conditions and Specifications.** Concrete is required for a plain concrete pavement to be constructed in North Dakota. The pavement specified compressive strength is 35 MPa at 28 days. The standard deviation of the concrete producer is 2.0 MPa. Type IP cement and 19-mm nominal maximum-size coarse aggregate is locally available. Proportion a concrete mixture for these conditions and check it by trial batch. Enter all data in the blank spaces on a trial mixture data sheet (Fig. 9-6).

**Durability Requirements.** The pavement will be exposed to freezing, thawing, and deicers and therefore should have a maximum water to cementitious material ratio of 0.45 (Table 9-1) and at least 335 kg of cement per cubic meter of concrete.

**Strength Requirements.** For a standard deviation of 2.0 MPa, the \( f'_{ct} \) (required compressive strength for proportioning) must be the larger of

\[
f'_{ct} = f'_{c} + 1.34S = 35 + 1.34(2.0) = 37.7 \text{ MPa}
\]

or

\[
f'_{ct} = f'_{c} + 2.33S - 3.45 = 35 + 2.33(2.0) - 3.45 = 36.2 \text{ MPa}
\]

Therefore the required average compressive strength = 37.7 MPa.

**Aggregate Size.** The 19-mm maximum-size coarse aggregate and the fine aggregate are in saturated-surface dry condition for the trial mixtures.

**Air Content.** The target air content should be 6% (Table 9-5) and the range is set at 5% to 8%.

**Slump.** The specified target slump for this project is 40 (±20) mm.

**Batch Quantities.** For convenience, a batch containing 10 kg of cement is to be made. The quantity of mixing water required is 10 x 0.45 = 4.5 kg. Representative samples of fine and coarse aggregates are measured in suitable containers. The values are entered as initial mass in Column 2 of the trial-batch data sheet (Fig. 9-6).

All of the measured quantities of cement, water, and air-entraining admixture are used and added to the mixer. Fine and coarse aggregates, previously brought to a saturated, surface-dry condition, are added until a workable concrete mixture with a slump deemed adequate for placement is produced. The relative proportions of fine and coarse aggregate for workability can readily be judged by an experienced concrete technician or engineer.

**Workability.** Results of tests for slump, air content, density, and a description of the appearance and workability are noted in the data sheet and Table 9-13.
The amounts of fine and coarse aggregates not used are recorded on the data sheet in Column 3, and mass of aggregates used (Column 2 minus Column 3) are noted in Column 4. If the slump when tested had been greater than that required, additional fine or coarse aggregates (or both) would have been added to reduce slump. Had the slump been less than required, water and cement in the appropriate ratio (0.45) would have been added to increase slump. It is important that any additional quantities be measured accurately and recorded on the data sheet.

**Mixture Proportions.**

Mixture proportions for a cubic meter of concrete are calculated in Column 5 of Fig. 9-6 by using the batch yield (volume) and density (unit weight). For example, the number of kilograms of cement per cubic meter is determined by dividing one cubic meter by the volume of concrete in the batch and multiplying the result by the number of kilograms of cement in the batch. The percentage of fine aggregate by mass of total aggregate is also calculated. In this trial batch, the cement content was 341 kg/m³ and the fine aggregate made up 38% of the total aggregate by mass. The air content and slump were acceptable. The 28-day strength was 39.1 MPa, greater than $f_{ct}$.
Example 4. Laboratory Trial Mixture Using the PCA Water-Cement Ratio Method (Inch-Pound Units)

With the following method, the mix designer develops the concrete proportions directly from a laboratory trial batch, rather than the absolute volume of the constituent ingredients as in Example 2.

**Conditions and Specifications.** Air-entrained concrete is required for a foundation wall that will be exposed to moderate sulfate soils. A compressive strength, \( f'_{c} \), of 4000 psi at 28 days using Type II cement is specified. Minimum thickness of the wall is 10 in. and concrete cover over \( \frac{1}{2} \)-in.-diameter reinforcing bars is 3 in. The clear distance between reinforcing bars is 3 in. The water-cement ratio versus compressive strength relationship based on field and previous laboratory data for the example ingredients is illustrated by Fig. 9-7. Based on the test records of the materials to be used, the standard deviation is 300 psi. Proportion and evaluate by trial batch a mixture meeting the above conditions and specifications. Enter all data in the appropriate blanks on a trial-mixture data sheet (Fig. 9-8).

**Water-Cement Ratio.** For these exposure conditions, Table 9-2 indicates that concrete with a maximum water-cement ratio of 0.50 should be used and the minimum design strength should be 4000 psi.

The water-cement ratio for strength is selected from a graph plotted to show the relationship between the water-cement ratio and compressive strength for these specific concrete materials (Fig. 9-7).

For a standard deviation of 300 psi, \( f'_{c} \) must be the larger of

\[
f'_{c} = f'_{c} + 1.34S = 4000 + 1.34(300) = 4402 \text{ psi}
\]

or

\[
f'_{c} = f'_{c} + 2.33S - 500 = 4000 + 2.33(300) - 500 = 4199 \text{ psi}
\]

Therefore, \( f'_{c} = 4400 \text{ psi} \)

From Fig. 9-7, the water-cement ratio for air-entrained concrete is 0.55 for an \( f'_{c} \) of 4400 psi. This is greater than the 0.50 permitted for the exposure conditions; therefore, the exposure requirements govern. A water-cement ratio of 0.50 must be used, even though this may produce strengths higher than needed to satisfy structural requirements.

**Aggregate Size.** Assuming it is economically available, \( 1\frac{1}{2} \)-in. maximum-size aggregate is satisfactory; it is less than \( \frac{1}{5} \) the wall thickness and less than \( \frac{3}{4} \) the clear distance between reinforcing bars and between reinforcing bars and the form. If this size were not available, the next smaller available size would be used. Aggregates are to be in a saturated surface-dry condition for these trial mixtures.

**Air Content.** Because of the exposure conditions and to improve workability, a moderate level of entrained air is needed. From Table 9-5, the target air content for concrete with \( 1\frac{1}{2} \)-in. aggregate in a moderate exposure is 4.5%. Therefore, proportion the mixture with an air content range of 4.5% ±1% and aim for 5.5% ±0.5% in the trial batch.

**Slump.** The recommended slump range for placing a reinforced concrete foundation wall is 1 in. to 3 in., assuming that the concrete will be consolidated by vibration (Table 9-6). Batch for 3 in. ±0.75 in.

**Batch Quantities.** For convenience, a batch containing 20 lb of cement is to be made. The quantity of mixing water required is \( 20 \times 0.50 = 10 \text{ lb} \). Representative samples of fine and coarse aggregates are weighed into suitable containers. The values are entered as initial weights in Column 2 of the trial-batch data sheet (Fig. 9-8).

All of the measured quantities of cement, water, and air-entraining admixture are used and added to the mixer. Fine and coarse aggregates, previously brought to a saturated surface-dry condition, are added in proportions similar to those used in mixes from which Fig. 9-7 was developed. Mixing continues until a workable concrete with a 3-in. slump deemed adequate for placement is produced. The relative proportions of fine and coarse aggregate for workability can readily be judged by an experienced concrete technician or engineer.

**Workability.** Results of tests for slump, air content, unit weight, and a description of the appearance and workability (“Good” for this example) are noted on the data sheet.

The amounts of fine and coarse aggregates not used are recorded on the data sheet in Column 3, and masses of aggregates used (Column 2 minus Column 3) are noted in Column 4. If the slump when tested had been greater than that required, additional fine or coarse aggregates (or...
both) would have been added to reduce slump. Had the slump been less than required, water and cement in the appropriate ratio (0.50) would have been added to increase slump. It is important that any additional quantities be measured accurately and recorded on the data sheet.

**Mixture Proportions.** Mixture proportions for a cubic yard of concrete are calculated in Column 5 of Fig. 9-8 by using the batch yield (volume) and density (unit weight). For example, the number of pounds of cement per cubic yard is determined by dividing 27 cu ft (1 cu yd) by the volume of concrete in the batch and multiplying the result by the number of pounds of cement in the batch. The percentage of fine aggregate by weight of total aggregate is also calculated. In this trial batch, the cement content was 539 lb per cubic yard and the fine aggregate made up 33.5% of the total aggregate by weight. The air content and slump were acceptable. The 28-day strength was 4950 psi (greater than $f_{cu}$). The mixture in Column 5, along with slump and air content limits of 1 in. to 3 in. and 3.5% to 5.5%, respectively, is now ready for submission to the project engineer.

**Mixture Adjustments.** To determine the most workable and economical proportions, additional trial batches could be made varying the percentage of fine aggregate. In each batch the water-cement ratio, aggregate gradation, air content, and slump should remain about the same. Results of four such trial batches are summarized in Table 9-14.

![Figure 9-8. Trial mixture data sheet (inch-pound units).](image)

**Table 9-14. Example of Results of Laboratory Trial Mixtures (Inch-Pound Units)**

<table>
<thead>
<tr>
<th>Batch no.</th>
<th>Slump, in.</th>
<th>Air content, percent</th>
<th>Density, lb/cu ft$^3$</th>
<th>Cement content, lb/cu yd$^3$</th>
<th>Fine aggregate, percent of total aggregate</th>
<th>Workability</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>3</td>
<td>5.4</td>
<td>144</td>
<td>539</td>
<td>33.5</td>
<td>Good</td>
</tr>
<tr>
<td>2</td>
<td>2½</td>
<td>4.9</td>
<td>144</td>
<td>555</td>
<td>27.4</td>
<td>Harsh</td>
</tr>
<tr>
<td>3</td>
<td>2½</td>
<td>5.1</td>
<td>144</td>
<td>549</td>
<td>35.5</td>
<td>Excellent</td>
</tr>
<tr>
<td>4</td>
<td>3</td>
<td>4.7</td>
<td>145</td>
<td>540</td>
<td>30.5</td>
<td>Excellent</td>
</tr>
</tbody>
</table>

*Water-cement ratio was 0.50.*
Table 9-15 illustrates the change in mix proportions for various types of concrete mixtures using a particular aggregate source. Information for concrete mixtures using particular ingredients can be plotted in several ways to illustrate the relationship between ingredients and properties. This is especially useful when optimizing concrete mixtures for best economy or to adjust to specification or material changes (Fig. 9-9).

![Graphical relationship](image)

Fig. 9-9. Example graphical relationship for a particular aggregate source demonstrating the relationship between slump, aggregate size, water to cement ratio, and cement content (Hover 1995).
### Table 9-15 (Metric). Example Trial Mixtures for Air-Entrained Concrete of Medium Consistency, 75-mm to 100-mm slump

<table>
<thead>
<tr>
<th>Water-cement ratio, kg per kg</th>
<th>Nominal maximum size of aggregate, mm</th>
<th>Air content, percent</th>
<th>Water, kg per cu meter of concrete</th>
<th>Cement, kg per cu meter of concrete</th>
<th>With fine sand, fineness modulus = 2.50</th>
<th>With coarse sand, fineness modulus = 2.90</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>Fine aggregate, percent of total</td>
<td>Fine aggregate, kg per cu meter of concrete</td>
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<td>0.40</td>
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<td>7.5</td>
<td>202</td>
<td>505</td>
<td>50</td>
<td>744</td>
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<tr>
<td></td>
<td>12.5</td>
<td>7.5</td>
<td>194</td>
<td>485</td>
<td>41</td>
<td>630</td>
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<tr>
<td></td>
<td>19.0</td>
<td>7.5</td>
<td>178</td>
<td>446</td>
<td>35</td>
<td>577</td>
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<td></td>
<td>25.0</td>
<td>6</td>
<td>169</td>
<td>424</td>
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<td>534</td>
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<tr>
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<td>37.5</td>
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<td>395</td>
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<td>7.5</td>
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<td>744</td>
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<td>7.5</td>
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<td>9.5</td>
<td>7.5</td>
<td>202</td>
<td>336</td>
<td>55</td>
<td>928</td>
</tr>
<tr>
<td></td>
<td>12.5</td>
<td>7.5</td>
<td>194</td>
<td>321</td>
<td>47</td>
<td>809</td>
</tr>
<tr>
<td></td>
<td>19.0</td>
<td>7.5</td>
<td>178</td>
<td>298</td>
<td>40</td>
<td>738</td>
</tr>
<tr>
<td></td>
<td>25.0</td>
<td>6</td>
<td>169</td>
<td>282</td>
<td>37</td>
<td>670</td>
</tr>
<tr>
<td></td>
<td>37.5</td>
<td>5</td>
<td>158</td>
<td>262</td>
<td>34</td>
<td>660</td>
</tr>
<tr>
<td>0.70</td>
<td>9.5</td>
<td>7.5</td>
<td>202</td>
<td>288</td>
<td>55</td>
<td>928</td>
</tr>
<tr>
<td></td>
<td>12.5</td>
<td>7.5</td>
<td>194</td>
<td>277</td>
<td>47</td>
<td>809</td>
</tr>
<tr>
<td></td>
<td>19.0</td>
<td>7.5</td>
<td>178</td>
<td>256</td>
<td>41</td>
<td>738</td>
</tr>
<tr>
<td></td>
<td>25.0</td>
<td>6</td>
<td>169</td>
<td>240</td>
<td>37</td>
<td>688</td>
</tr>
<tr>
<td></td>
<td>37.5</td>
<td>5</td>
<td>158</td>
<td>226</td>
<td>34</td>
<td>660</td>
</tr>
</tbody>
</table>

### Table 9-15 (Inch-Pound Units). Example Trial Mixtures for Air-Entrained Concrete of Medium Consistency, 3-in. to 4-in. slump

<table>
<thead>
<tr>
<th>Water-cement ratio, lb per lb</th>
<th>Nominal maximum size of aggregate, in.</th>
<th>Air content, percent</th>
<th>Water, lb per cu yd of concrete</th>
<th>Cement, lb per cu yd of concrete</th>
<th>With fine sand, fineness modulus = 2.50</th>
<th>With coarse sand, fineness modulus = 2.90</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>Fine aggregate, percent of total</td>
<td>Fine aggregate, lb per cu yd of concrete</td>
</tr>
<tr>
<td>0.40</td>
<td>%</td>
<td>7.5</td>
<td>340</td>
<td>850</td>
<td>50</td>
<td>1250</td>
</tr>
<tr>
<td></td>
<td>%</td>
<td>7.5</td>
<td>325</td>
<td>815</td>
<td>41</td>
<td>1060</td>
</tr>
<tr>
<td></td>
<td>%</td>
<td>6</td>
<td>300</td>
<td>750</td>
<td>35</td>
<td>970</td>
</tr>
<tr>
<td></td>
<td>%</td>
<td>6</td>
<td>285</td>
<td>715</td>
<td>32</td>
<td>900</td>
</tr>
<tr>
<td>0.45</td>
<td>%</td>
<td>7.5</td>
<td>340</td>
<td>755</td>
<td>51</td>
<td>1330</td>
</tr>
<tr>
<td></td>
<td>%</td>
<td>7.5</td>
<td>325</td>
<td>720</td>
<td>43</td>
<td>1140</td>
</tr>
<tr>
<td></td>
<td>%</td>
<td>6</td>
<td>300</td>
<td>665</td>
<td>37</td>
<td>1040</td>
</tr>
<tr>
<td></td>
<td>%</td>
<td>6</td>
<td>285</td>
<td>635</td>
<td>33</td>
<td>970</td>
</tr>
<tr>
<td></td>
<td>%</td>
<td>5</td>
<td>265</td>
<td>590</td>
<td>31</td>
<td>930</td>
</tr>
</tbody>
</table>
Table 9-15 (Inch-Pound Units). Example Trial Mixtures for Air-Entrained Concrete of Medium Consistency, 3-in. to 4-in. slump (Continued)

<table>
<thead>
<tr>
<th>Water-cement ratio, lb per lb</th>
<th>Nominal maximum size of aggregate, in.</th>
<th>Air content, percent</th>
<th>Water, lb per cu yd of concrete</th>
<th>Cement, lb per cu yd of concrete</th>
<th>With fine sand, fineness modulus = 2.50</th>
<th>With coarse sand, fineness modulus = 2.90</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>Fine aggregate, percent of total aggregate</td>
<td>Fine aggregate, lb per cu yd of concrete</td>
</tr>
<tr>
<td>0.50</td>
<td>%</td>
<td></td>
<td></td>
<td></td>
<td>53</td>
<td>1400</td>
</tr>
<tr>
<td></td>
<td>%</td>
<td></td>
<td></td>
<td></td>
<td>44</td>
<td>1200</td>
</tr>
<tr>
<td></td>
<td>%</td>
<td></td>
<td></td>
<td></td>
<td>38</td>
<td>1100</td>
</tr>
<tr>
<td></td>
<td>1</td>
<td></td>
<td></td>
<td></td>
<td>34</td>
<td>1020</td>
</tr>
<tr>
<td></td>
<td>1½</td>
<td></td>
<td></td>
<td></td>
<td>32</td>
<td>980</td>
</tr>
<tr>
<td>0.55</td>
<td>%</td>
<td></td>
<td></td>
<td></td>
<td>54</td>
<td>1450</td>
</tr>
<tr>
<td></td>
<td>%</td>
<td></td>
<td></td>
<td></td>
<td>45</td>
<td>1250</td>
</tr>
<tr>
<td></td>
<td>%</td>
<td></td>
<td></td>
<td></td>
<td>39</td>
<td>1140</td>
</tr>
<tr>
<td></td>
<td>1</td>
<td></td>
<td></td>
<td></td>
<td>35</td>
<td>1060</td>
</tr>
<tr>
<td></td>
<td>1½</td>
<td></td>
<td></td>
<td></td>
<td>33</td>
<td>1030</td>
</tr>
<tr>
<td>0.60</td>
<td>%</td>
<td></td>
<td></td>
<td></td>
<td>54</td>
<td>1490</td>
</tr>
<tr>
<td></td>
<td>%</td>
<td></td>
<td></td>
<td></td>
<td>46</td>
<td>1290</td>
</tr>
<tr>
<td></td>
<td>%</td>
<td></td>
<td></td>
<td></td>
<td>40</td>
<td>1180</td>
</tr>
<tr>
<td></td>
<td>1</td>
<td></td>
<td></td>
<td></td>
<td>36</td>
<td>1100</td>
</tr>
<tr>
<td></td>
<td>1½</td>
<td></td>
<td></td>
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<td>33</td>
<td>1060</td>
</tr>
<tr>
<td>0.65</td>
<td>%</td>
<td></td>
<td></td>
<td></td>
<td>55</td>
<td>1530</td>
</tr>
<tr>
<td></td>
<td>%</td>
<td></td>
<td></td>
<td></td>
<td>47</td>
<td>1330</td>
</tr>
<tr>
<td></td>
<td>%</td>
<td></td>
<td></td>
<td></td>
<td>40</td>
<td>1210</td>
</tr>
<tr>
<td></td>
<td>1</td>
<td></td>
<td></td>
<td></td>
<td>37</td>
<td>1130</td>
</tr>
<tr>
<td></td>
<td>1½</td>
<td></td>
<td></td>
<td></td>
<td>34</td>
<td>1090</td>
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<tr>
<td>0.70</td>
<td>%</td>
<td></td>
<td></td>
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<td>55</td>
<td>1560</td>
</tr>
<tr>
<td></td>
<td>%</td>
<td></td>
<td></td>
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<td>47</td>
<td>1360</td>
</tr>
<tr>
<td></td>
<td>%</td>
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<td></td>
<td></td>
<td>41</td>
<td>1240</td>
</tr>
<tr>
<td></td>
<td>1</td>
<td></td>
<td></td>
<td></td>
<td>37</td>
<td>1160</td>
</tr>
<tr>
<td></td>
<td>1½</td>
<td></td>
<td></td>
<td></td>
<td>34</td>
<td>1110</td>
</tr>
</tbody>
</table>

Example 5. Absolute Volume Method Using Multiple Cementing Materials and Admixtures (Metric)

The following example illustrates how to develop a mix using the absolute volume method when more than one cementing material and admixture are used.

**Conditions and Specifications.** Concrete with a structural design strength of 40 MPa is required for a bridge to be exposed to freezing and thawing, deicers, and very severe sulfate soils. A coulomb value not exceeding 1500 is required to minimize permeability to chlorides. Water reducers, air entrainers, and plasticizers are allowed. A shrinkage reducer is requested to keep shrinkage under 300 millionths. Some structural elements exceed a thickness of 1 meter, requiring control of heat development. The concrete producer has a standard deviation of 2 MPa for similar mixes to that required here. For difficult placement areas, a slump of 200 mm to 250 mm is required. The following materials are available:

- **Cement:** Type HS, silica fume modified portland cement, ASTM C1157. Relative density of 3.14. Silica fume content of 5%.
- **Fly ash:** Class F, ASTM C618 (AASHTO M 295). Relative density of 2.60.
- **Slag:** Grade 120, ASTM C989 (AASHTO M 302). Relative density of 2.90.
- **Coarse aggregate:** Well-graded 19-mm nominal maximum-size crushed rock (ASTM C33 or AASHTO M 80) with an oven-dry relative density of 2.68, absorption of 0.5%, and oven-dry density of 1600 kg/m³. The laboratory sample has a moisture content of 2.0%. This aggregate has a history of alkali-silica reactivity in the field.
- **Fine aggregate:** Natural sand with some crushed particles (ASTM C33 or AASHTO M 6) with an oven-dry relative density of 2.64 and an absorption of 0.7%. The laboratory sample has a moisture content of 6%. The fineness modulus is 2.80.
- **Air entrainer:** Synthetic, ASTM C260 (AASHTO M 154).
- **Retarding water reducer:** Type D, ASTM C494 (AASHTO M 194). Dosage of 3 g per kg of cementing materials.
Plasticizer: Type 1, ASTM C1017. Dosage of 30 g per kg of cementing materials.

Shrinkage reducer: Dosage of 15 g per kg of cementing materials.

**Strength.** For a standard deviation of 2.0 MPa, the $f_{ct}$ must be the greater of

\[
f_{ct} = f'_c + 1.34S = 40 + 1.34(2) = 42.7
\]

or

\[
f_{ct} = 0.9 f'_c + 2.33S = 36 + 2.33(2) = 40.7
\]

therefore $f_{ct} = 42.7$

**Water to Cementing Materials Ratio.** Past field records using these materials indicate that a water to cementing materials ratio of 0.35 is required to provide a strength level of 42.7 MPa.

For a deicer environment and to protect embedded steel from corrosion, Table 9-1 requires a maximum water to cementing materials ratio of 0.40 and a strength of at least 35 MPa. For a severe sulfate environment, Table 9-2 requires a maximum water to cementing materials ratio of 0.40 and a strength of at least 35 MPa. Both the water to cementing materials ratio requirements and strength requirements are met and exceeded using the above determined 0.35 water to cementing materials ratio and 40 MPa design strength.

**Air Content.** For a severe exposure, Fig. 9-4 suggests a target air content of 6% for 19-mm aggregate. Therefore, design the mix for 5% to 8% and use 8% for batch proportions. The trial batch air content must be within ±0.5 percentage points of the maximum allowable air content.

**Slump.** Assume a slump of 50 mm without the plasticizer and a maximum of 200 mm to 250 mm after the plasticizer is added. Use 250 ± 20 mm for proportioning purposes.

**Water Content.** Fig. 9-5 recommends that a 50-mm slump, air-entrained concrete with 19-mm aggregate should have a water content of about 168 kg/m³. Assume the retarding water reducer and plasticizer will jointly reduce water demand by 15% in this case, resulting in an estimated water demand of 143 kg per cubic meter, while achieving the 250-mm slump.

**Cementing Materials Content.** The amount of cementing materials is based on the maximum water-cementing materials ratio and water content. Therefore, 143 kg of water divided by a water-cementing materials ratio of 0.35 requires a cement content of 409 kg. Fly ash and slag will be used to help control alkali-silica reactivity and control temperature rise. Local use has shown that a fly ash dosage of 15% and a slag dosage of 30% by mass of cementing materials are adequate. Therefore, the suggested cementing materials for one cubic meter of concrete are as follows:

<table>
<thead>
<tr>
<th>Ingredient</th>
<th>Quantity (kg)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cement</td>
<td>225</td>
</tr>
<tr>
<td>Fly ash</td>
<td>61</td>
</tr>
<tr>
<td>Slag</td>
<td>123</td>
</tr>
<tr>
<td>Air</td>
<td>8.0</td>
</tr>
<tr>
<td>Coarse aggregate</td>
<td>992</td>
</tr>
</tbody>
</table>

These dosages meet the requirements of Table 9-8 (2.8% silica fume from the cement + 15% fly ash + 30% slag = 47.8% which is less than the 50% maximum allowed).

**Coarse-Aggregate Content.** The quantity of 19-mm nominal maximum-size coarse aggregate can be estimated from Fig. 9-3. The bulk volume of coarse aggregate is estimated using sand with a fineness modulus of 2.80 is 0.62. Since the coarse aggregate has a bulk density of 1600 kg/m³, the oven-dry mass of coarse aggregate for a cubic meter of concrete is

\[
1600 \times 0.62 = 992 \text{ kg/m}^3
\]

**Admixture Content.** For an 8% air content, the air-entraining admixture manufacturer recommends a dosage of 0.5 g per kg of cementing materials. The amount of air entrainer is then

\[
0.5 \times 409 = 205 \text{ g} = 0.205 \text{ kg}
\]

The retarding water reducer dosage rate is 3 g per kg of cementing materials. This results in

\[
3 \times 409 = 1227 \text{ g or 1.227 kg of water reducer per cubic meter of concrete.}
\]

The plasticizer dosage rate is 30 g per kg of cementing materials. This results in

\[
30 \times 409 = 12,270 \text{ g or 12.270 kg of plasticizer per cubic meter of concrete.}
\]

The shrinkage reducer dosage rate is 15 g per kg of cementing materials. This results in

\[
15 \times 409 = 6135 \text{ g or 6.135 kg of shrinkage reducer per cubic meter of concrete.}
\]

**Fine-Aggregate Content.** At this point, the amounts of all ingredients except the fine aggregate are known. The volume of fine aggregate is determined by subtracting the absolute volumes of all known ingredients from 1 cubic meter. The absolute volumes of the ingredients is calculated by dividing the known mass of each by the product of their relative density and the density of water. Assume a relative density of 1.0 for the chemical admixtures. Assume a density of water of 997.75 kg/m³ as all materials in the laboratory are maintained at a room temperature of 22°C (Table 9-12). Volumetric computations are as follows:

<table>
<thead>
<tr>
<th>Ingredient</th>
<th>Volume (m³)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Water (including chemical admixtures)</td>
<td>$\frac{143}{1.0 \times 997.75}$ = 0.143 m³</td>
</tr>
<tr>
<td>Cement</td>
<td>$\frac{225}{3.14 \times 997.75}$ = 0.072 m³</td>
</tr>
<tr>
<td>Fly ash</td>
<td>$\frac{61}{2.60 \times 997.75}$ = 0.024 m³</td>
</tr>
<tr>
<td>Slag</td>
<td>$\frac{123}{2.90 \times 997.75}$ = 0.043 m³</td>
</tr>
<tr>
<td>Air</td>
<td>$\frac{8.0}{100}$ = 0.080 m³</td>
</tr>
<tr>
<td>Coarse aggregate</td>
<td>$\frac{992}{2.68 \times 997.75}$ = 0.371 m³</td>
</tr>
<tr>
<td>Total</td>
<td>$\frac{173}{2.68 \times 997.75}$ = 0.733 m³</td>
</tr>
</tbody>
</table>
The batch quantities for one cubic meter of concrete are revised to include aggregate moisture as follows:

<table>
<thead>
<tr>
<th>Material</th>
<th>Quantity</th>
</tr>
</thead>
<tbody>
<tr>
<td>Water (to be added)</td>
<td>71 kg</td>
</tr>
<tr>
<td>Cement</td>
<td>225 kg</td>
</tr>
<tr>
<td>Fly ash</td>
<td>61 kg</td>
</tr>
<tr>
<td>Slag</td>
<td>123 kg</td>
</tr>
<tr>
<td>Coarse aggregate (2% MC)</td>
<td>1012 kg</td>
</tr>
<tr>
<td>Fine aggregate (6% MC)</td>
<td>745 kg</td>
</tr>
<tr>
<td>Air entrainer</td>
<td>0.205 kg</td>
</tr>
<tr>
<td>Water reducer</td>
<td>1.227 kg</td>
</tr>
<tr>
<td>Plasticizer</td>
<td>12.27 kg</td>
</tr>
<tr>
<td>Shrinkage reducer</td>
<td>6.14 kg</td>
</tr>
</tbody>
</table>

**Trial Batch.** The above mixture is tested in a 0.1 m$^3$ batch in the laboratory (multiply above quantities by 0.1 to obtain batch quantities). The mixture had an air content of 7.8%, a slump of 240 mm, a density of 2257 kg/m$^3$, a yield of 0.1 m$^3$, and a compressive strength of 44 MPa. Rapid chloride testing resulted in a coulomb value of 990 (ASTM C1202 or AASHTO T 277). ASTM C1567 was used to evaluate the potential of the mix for alkali-silica reactivity, resulting in an acceptable expansion of 0.02%. Temperature rise was acceptable and shrinkage was within specifications. The water-soluble chloride content was 0.06%, meeting the requirements of Table 9-9. The following mix proportions meet all applicable requirements and are ready for submission to the project engineer for approval:

<table>
<thead>
<tr>
<th>Material</th>
<th>Quantity</th>
</tr>
</thead>
<tbody>
<tr>
<td>Water added</td>
<td>123 kg</td>
</tr>
<tr>
<td>Cement, Type HS</td>
<td>225 kg</td>
</tr>
<tr>
<td>Fly ash, Class F</td>
<td>61 kg</td>
</tr>
<tr>
<td>Slag, Grade 120</td>
<td>123 kg</td>
</tr>
<tr>
<td>Coarse aggregate (dry)</td>
<td>992 kg</td>
</tr>
<tr>
<td>Fine aggregate (dry)</td>
<td>703 kg</td>
</tr>
<tr>
<td>Air entrainer</td>
<td>0.205 kg</td>
</tr>
<tr>
<td>Water reducer</td>
<td>1.227 kg</td>
</tr>
<tr>
<td>Plasticizer</td>
<td>12.27 kg</td>
</tr>
<tr>
<td>Shrinkage reducer</td>
<td>6.14 kg</td>
</tr>
</tbody>
</table>

**Moisture.** The dry batch weights of aggregates have to be increased to compensate for the moisture on and in the aggregates and the mixing water reduced accordingly. The coarse aggregate and fine aggregate have moisture contents of 2% and 6%, respectively. With the moisture contents indicated, the trial batch aggregate proportions become

- Coarse aggregate (2% MC) = 992 x 1.02 = 1012 kg
- Fine aggregate (6% MC) = 703 x 1.06 = 745 kg

Absorbed water does not become part of the mixing water and must be excluded from the water adjustment. Surface moisture contributed by the coarse aggregate amounts to 2% – 0.5% = 1.5% and that contributed by the fine aggregate, 6% – 0.7% = 5.3%. The estimated added water becomes

$$123 - (992 \times 0.015) - (703 \times 0.053) = 71 \text{ kg}$$

**CONCRETE FOR SMALL JOBS**

Although well-established ready mixed concrete mixtures are used for most construction, ready mix is not always practical for small jobs, especially those requiring one cubic meter (yard) or less. Small batches of concrete mixed at the site are required for such jobs.
Table 9-16 (Metric). Proportions by Mass to Make One Tenth Cubic Meter of Concrete for Small Jobs

<table>
<thead>
<tr>
<th>Nominal maximum size coarse aggregate, mm</th>
<th>Air-entrained concrete</th>
<th>Non-air-entrained concrete</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Cement, kg</td>
<td>Wet fine aggregate, kg</td>
</tr>
<tr>
<td>9.5</td>
<td>46</td>
<td>85</td>
</tr>
<tr>
<td>12.5</td>
<td>43</td>
<td>74</td>
</tr>
<tr>
<td>19.0</td>
<td>40</td>
<td>67</td>
</tr>
<tr>
<td>25.0</td>
<td>38</td>
<td>62</td>
</tr>
<tr>
<td>37.5</td>
<td>37</td>
<td>61</td>
</tr>
</tbody>
</table>

*If crushed stone is used, decrease coarse aggregate by 5 kg and increase fine aggregate by 5 kg.

Table 9-16 (Inch-Pound). Proportions by Mass to Make One Cubic Foot of Concrete for Small Jobs

<table>
<thead>
<tr>
<th>Nominal maximum size coarse aggregate, in.</th>
<th>Air-entrained concrete</th>
<th>Non-air-entrained concrete</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Cement, lb</td>
<td>Wet fine aggregate, lb</td>
</tr>
<tr>
<td>3⁄8</td>
<td>29</td>
<td>53</td>
</tr>
<tr>
<td>1⁄2</td>
<td>27</td>
<td>46</td>
</tr>
<tr>
<td>3⁄4</td>
<td>25</td>
<td>42</td>
</tr>
<tr>
<td>1</td>
<td>24</td>
<td>39</td>
</tr>
<tr>
<td>1 1⁄2</td>
<td>23</td>
<td>38</td>
</tr>
</tbody>
</table>

*If crushed stone is used, decrease coarse aggregate by 3 lb and increase fine aggregate by 3 lb.

Table 9-17. Proportions by Bulk Volume* of Concrete for Small Jobs

<table>
<thead>
<tr>
<th>Nominal maximum size coarse aggregate, mm (in.)</th>
<th>Air-entrained concrete</th>
<th>Non-air-entrained concrete</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Cement</td>
<td>Wet fine aggregate</td>
</tr>
<tr>
<td>9.5 (%</td>
<td>1</td>
<td>2⁄4</td>
</tr>
<tr>
<td>12.5 (%)</td>
<td>1</td>
<td>2⁄4</td>
</tr>
<tr>
<td>19.0 (%)</td>
<td>1</td>
<td>2⁄4</td>
</tr>
<tr>
<td>25.0 (1)</td>
<td>1</td>
<td>2⁄4</td>
</tr>
<tr>
<td>37.5 (1½)</td>
<td>1</td>
<td>2⁄4</td>
</tr>
</tbody>
</table>

*The combined volume is approximately ⅔ of the sum of the original bulk volumes.

If mixture proportions or mixture specifications are not available, Tables 9-16 and 9-17 can be used to select proportions for concrete for small jobs. Recommendations with respect to exposure conditions discussed earlier should be followed.

The proportions in Tables 9-16 and 9-17 are only a guide and may need adjustments to obtain a workable mix with locally available aggregates (PCA 1988). Packaged, combined, dry concrete ingredients (ASTM C387) are also available.

**DESIGN REVIEW**

In practice, concrete mixture proportions will be governed by the limits of data available on the properties of materials, the degree of control exercised over the production of concrete at the plant, and the amount of supervision at the jobsite. It should not be expected that field results will be an exact duplicate of laboratory trial batches. An adjustment of the selected trial mixture is usually necessary on the job.

The mixture design and proportioning procedures presented here and summarized in Fig. 9-10 are applicable to normal-weight concrete. For concrete requiring some special property, using special admixtures or materials—lightweight aggregates, for example—different proportioning principles may be involved.

Internet web sites also provide assistance with designing and proportioning concrete mixtures (Bentz 2001). Many of these web sites are internationally oriented and assume principles not used in North America. Therefore, appropriate cautions should be taken when using the internet to design concrete mixtures.
Concrete production facility has field strength test records for the specified class of concrete or within 7 MPa (1000 psi) of the specified class.

- **Yes**
  - 30 consecutive tests
    - Calculate $S$
    - Required average strength from Eq. (9-1) or (9-2) or (9-3)

- **No**
  - Two groups of consecutive tests (total ≥ 30)
    - Calculate average $S$
    - Required average strength from Eq. (9-1) or (9-2) or (9-3)
  - 15 to 29 consecutive tests
    - Calculate and increase using Table 9-10
    - Required average strength from Table 9-11

- **Field record of at least ten consecutive test results using similar materials and under similar conditions is available**

- **No**
  - Results represent one mixture
    - Average ≥ required average
    - Submit for approval
  - Results represent two or more mixtures
    - Plot average strength versus proportions and interpolate for required average strength
    - Submit for approval

- **Make trial mixtures using at least three different w/cm ratios or cementing materials contents**
  - 15 to 29 consecutive tests
    - Calculate and increase using Table 9-10
    - Required average strength from Table 9-11

**Fig. 9-10. Flowchart for selection and documentation of concrete proportions.**
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CHAPTER 10
Batching, Mixing, Transporting, and Handling Concrete

The specification, production, and delivery of concrete are achieved in different ways. The basic processes and common techniques are explained here. ASTM C94 provides standard specifications for the manufacture and delivery of freshly mixed concrete. Standards of the Concrete Plant Manufacturers Bureau, Truck Mixer Manufacturers Bureau, and Volumetric Mixer Manufacturers Bureau can be found on the National Ready Mixed Concrete Association’s website at http://www.nrmca.org.

Three options for ordering or specifying concrete are described in ASTM C94:

1. Option A is performance based. It requires the purchaser to specify the compressive strength only, while the concrete producer selects the mixture proportions needed to obtain the required compressive strength.
2. Option B is prescription based. The purchaser specifies mixture proportions, including cement, water and admixture contents.
3. Option C is a mixed option. It requires the concrete producer to select the mixture proportions with the minimum allowable cement content and compressive strength specified by the purchaser.

**Batching**

Batching is the process of measuring concrete mix ingredients by either mass or volume and introducing them into the mixer. To produce concrete of uniform quality, the ingredients must be measured accurately for each batch. Most specifications require that batching be done by mass rather than by volume (ASTM C94 or AASHTO M 157). Water and liquid admixtures can be measured accurately by either volume or mass. Volumetric batching (ASTM C685 or AASHTO M 241) is used for concrete mixed in continuous mixers.

Specifications generally require that materials be measured for individual batches within the following percentages of accuracy: cementitious material ±1%, aggregates ±2%, water ±1%, and admixtures ±3%.

Equipment should be capable of measuring quantities within these tolerances for the smallest batch regularly used as well as for larger batches (Fig. 10-1). The accuracy of scales and batching equipment should be checked periodically and adjusted when necessary.

Liquid chemical admixtures should be charged into the mixture as aqueous solutions. The volume of liquid, if significant, should be subtracted from the batched quantity of mixing water. Admixtures that cannot be added in solution can be either batched by mass or volume as directed by the manufacturer. Admixture dispensers should be checked frequently since errors in dispensing admixtures, particularly overdoses, can lead to serious problems in both fresh and hardened concrete.

![Fig. 10-1. Control room for batching equipment in a typical ready mixed concrete plant. (IMG12398)](image-url)
MIXING CONCRETE

All concrete should be mixed thoroughly until it is uniform in appearance, with all ingredients evenly distributed. Mixers should not be loaded above their rated capacities and should be operated at the mixing speed recommended by the manufacturer. Increased output should be obtained by using a larger mixer or additional mixers, rather than by speeding up or overloading the equipment on hand. If the blades of a mixer become worn or coated with hardened concrete, mixing action will be less efficient. These conditions should be corrected.

If concrete has been adequately mixed, samples taken from different portions of a batch will have essentially the same density, air content, slump, and coarse-aggregate content. Maximum allowable differences to evaluate mixing uniformity within a batch of ready mixed concrete are given in ASTM C94 (AASHTO M 157).

Structural low-density concrete can be mixed the same way as normal-density concrete when the aggregates have less than 10% total absorption by mass or when the absorption is less than 2% by mass during the first hour after immersion in water. For aggregates not meeting these limits, mixing procedures are described in PCA (1986).

Stationary Mixing

Concrete is sometimes mixed at the jobsite in a stationary mixer or a paving mixer (Fig. 10-2). Stationary mixers include both onsite mixers and central mixers in ready mix plants. They are available in sizes up to 9.0 m³ (12 yd³) and can be of the tilting or nonlifting type or the open-top revolving blade or paddle type. All types may be equipped with loading skips and some are equipped with a swinging discharge chute. Many stationary mixers have timing devices, some of which can be set for a given mixing time and locked so that the batch cannot be discharged until the designated mixing time has elapsed.

Careful attention should be paid to the required mixing time. Many specifications require a minimum mixing time of one minute plus 15 seconds for every cubic meter (yard), unless mixer performance tests demonstrate that shorter periods are acceptable and will provide a uniform concrete mixture. Short mixing times can result in nonhomogenous mixtures, poor distribution of air voids (resulting in poor frost resistance), poor strength gain, and early stiffening problems. The mixing period should be measured from the time all cement and aggregates are in the mixer drum, provided all the water is added before one-fourth of the mixing time has elapsed (ACI 304R).

Under usual conditions, up to about 10% of the mixing water should be placed in the drum before the solid materials are added. Water then should be added uniformly with the solid materials, leaving about 10% to be added after all other materials are in the drum. When heated water is used in cold weather, this order of charging may require some modification to prevent possible rapid stiffening when hot water contacts the cement. In this case, addition of the cementitious materials should be delayed until most of the aggregate and water have intermingled in the drum. Where the mixer is charged directly from a batch plant, the materials should be added simultaneously at such rates that the charging time is about the same for all materials. If supplementary cementing materials are used, they should be added after the cement.

If retarding or water-reducing admixtures are used, they should be added in the same sequence in the charging cycle each time. If not, significant variations in the time of initial setting and percentage of entrained air may result. Addition of the admixture should be completed not later than one minute after addition of water to the cement has been completed or prior to the start of the last three-fourths of the mixing cycle, whichever occurs first. If two or more admixtures are used in the same batch of concrete, they should be added separately; this is intended to avoid any interaction that might interfere with the efficiency of any of the admixtures and adversely affect the concrete properties. In addition, the sequence in which they are added to the mix can be important too.

Ready Mixed Concrete

Ready mixed concrete is proportioned and mixed off the project site and is delivered to the construction area in a freshly mixed and unhardened state. It can be manufactured by any of the following methods:
1. Central-mixed concrete is mixed completely in a stationary mixer (Fig. 10-3) and is delivered either in a truck agitator (Fig. 10-4 bottom), a truck mixer operating at agitating speed (Fig. 10-3), or a nonagitating truck (Fig. 10-4 top). Fig. 10-5 illustrates a central mix ready mix plant.

2. Shrink-mixed concrete is mixed partially in a stationary mixer and completed in a truck mixer.

3. Truck-mixed concrete is mixed completely in a truck mixer (Fig. 10-6).

ASTM C94 (AASHTO M 157) notes that when a truck mixer is used for complete mixing, 70 to 100 revolutions of the drum or blades at the rate of rotation designated by the manufacturer as mixing speed are usually required to produce the specified uniformity of concrete. All revolutions after 100 should be at a rate of rotation designated by the manufacturer as agitating speed. Agitating speed is usually about 2 to 6 rpm, and mixing speed is generally about 6 to 18 rpm. Mixing at high speeds for long periods of time, about 1 or more hours, can result in concrete strength loss, temperature rise, excessive loss of entrained air, and accelerated slump loss.

When truck mixers are used, ASTM C94 (AASHTO M 157) also limits the time between batching and complete discharge of the concrete at the job site; this time is 1½ hours or before the drum has revolved 300 times after introduction of water to the cement and aggregates or the cement to the aggregates. Mixers and agitators should always be operated within the limits for volume and speed of rotation designated by the equipment manufacturer.
Fig. 10-5. Schematic of a ready mix plant.
Mobile Batcher Mixed Concrete (Continuous Mixer)

Mobile volumetric mixers are special trucks (Fig. 10-7) that batch by volume and continuously mix concrete as the dry concrete ingredients, water, and admixtures are continuously fed into a mixing trough, typically an auger system. The concrete must conform to ASTM C685 (AASHTO M 241) specifications and is proportioned and mixed at the jobsite in the quantities needed. The concrete mixture is also easily adjusted for project placement and weather conditions.

Remixing Concrete

Fresh concrete that is left to agitate in the mixer drum tends to stiffen before initial set develops. Such concrete may be used if upon remixing it becomes sufficiently plastic to be compacted in the forms. ASTM C94 (AASHTO M 157) allows water to be added to remix the concrete when the truck arrives on the jobsite and the slump is less than specified providing the following conditions are met: (1) maximum allowable water-cement ratio is not exceeded as calculated including surface water on aggregates as well as batch water and water added on site; (2) maximum allowable slump is not exceeded; (3) maximum allowable mixing and agitating time (or drum revolutions) are not exceeded; and (4) concrete is remixed for a minimum of 30 revolutions at mixing speed or until the uniformity of the concrete is within the limits described in ASTM C94 (AASHTO M 157). Water should not be added to a partial load. If early setting becomes a persistent problem, a retarder may be used to control early hydration, especially in high-cement-content mixes. Mixture adjustments at the jobsite for air entrainment, and the addition of other admixtures, is permitted, followed by sufficient mixing.

Indiscriminate addition of water to make concrete more fluid should not be allowed because this lowers the quality of concrete. The later addition of water and remixing to retemper the mixture can result in marked strength reduction.

TRANSPORTING AND HANDLING CONCRETE

Good advanced planning can help choose the appropriate handling method for an application. Consider the following three occurrences that, should they occur during handling and placing, could seriously affect the quality of the finished work:

Delays. The objective in planning any work schedule is to produce the fastest work with the best labor force and the proper equipment for the work at hand. Machines for transporting and handling concrete are being improved all the time. The greatest productivity will be achieved if the work is planned to get the most out of personnel and equipment and if the equipment is selected to reduce the delay time during concrete placement.

Early Stiffening and Drying Out. Concrete begins to stiffen as soon as the cementitious materials and water are mixed, but the degree of stiffening that occurs in the first 30 minutes is not usually a problem; concrete that is kept agitated generally can be placed and compacted within 1/2 hours after mixing unless hot concrete temperatures or
high cement contents speed up hydration excessively. Planning should eliminate or minimize any variables that would allow the concrete to stiffen to the extent that full consolidation is not achieved and finishing becomes difficult. Less time is available during conditions that hasten the stiffening process, such as hot and dry weather, use of accelerators, and use of heated concrete.

**Segregation.** Segregation is the tendency for coarse aggregate to separate from the sand-cement mortar. This results in part of the batch having too little coarse aggregate and the remainder having too much. The former is likely to shrink more and crack and have poor resistance to abrasion. The latter may be too harsh for full consolidation and finishing and is a frequent cause of honeycombing. The method and equipment used to transport and handle the concrete must not result in segregation of the concrete materials.

### Methods and Equipment for Transporting and Handling Concrete

Table 10-1 summarizes the most common methods and equipment for moving concrete to the point where it is needed.

**Table 10-1. Methods and Equipment for Transporting and Handling Concrete**

<table>
<thead>
<tr>
<th>Equipment</th>
<th>Type and range of work for which equipment is best suited</th>
<th>Advantages</th>
<th>Points to watch for</th>
</tr>
</thead>
<tbody>
<tr>
<td>Belt conveyors</td>
<td>For conveying concrete horizontally or to a higher or lower level. Usually positioned between main discharge point and secondary discharge point.</td>
<td>Belt conveyors have adjustable reach, traveling diverter, and variable speed both forward and reverse. Can place large volumes of concrete quickly when access is limited.</td>
<td>End-discharge arrangements needed to prevent segregation and leave no mortar on return belt. In adverse weather (hot, windy) long reaches of belt need cover.</td>
</tr>
<tr>
<td>Belt conveyors mounted on truck mixers</td>
<td>For conveying concrete to a lower, horizontal, or higher level.</td>
<td>Conveying equipment arrives with the concrete. Adjustable reach and variable speed.</td>
<td>End-discharge arrangements needed to prevent segregation and leave no mortar on return belt.</td>
</tr>
<tr>
<td>Buckets</td>
<td>Used with cranes, cableways, and helicopters for construction of buildings and dams. Convey concrete directly from central discharge point to formwork or to secondary discharge point.</td>
<td>Enables full versatility of cranes, cableways, and helicopters to be exploited. Clean discharge. Wide range of capacities.</td>
<td>Select bucket capacity to conform to size of the concrete batch and capacity of placing equipment. Discharge should be controllable.</td>
</tr>
<tr>
<td>Chutes on truck mixers</td>
<td>For conveying concrete to a lower level, usually below ground level, on all types of concrete construction.</td>
<td>Low cost and easy to maneuver. No power required; gravity does most of the work.</td>
<td>Slopes should range between 1 to 2 and 1 to 3 and chutes must be adequately supported in all positions. End-discharge arrangements (downpipe) needed to prevent segregation.</td>
</tr>
<tr>
<td>Cranes and buckets</td>
<td>The right equipment for work above ground level.</td>
<td>Can handle concrete, reinforcing steel, formwork, and sundry items in bridges and concrete-framed buildings.</td>
<td>Has only one hook. Careful scheduling between trades and operations is needed to keep crane busy.</td>
</tr>
<tr>
<td>Dropchutes</td>
<td>Used for placing concrete in vertical forms of all kinds. Some chutes are one piece tubes made of flexible rubberized canvas or plastic, others are assembled from articulated metal cylinders (elephant trunks). Dropchutes direct concrete into formwork and carry it to bottom of forms without segregation. Their use avoids spillage of grout and concrete on reinforcing steel and form sides, which is harmful when off-the-form surfaces are specified. They also will prevent segregation of coarse particles.</td>
<td>Dropchutes should have sufficiently large, splayed-top openings into which concrete can be discharged without spillage. The cross section of drop chute should be chosen to permit inserting into the formwork without interfering with reinforcing steel.</td>
<td></td>
</tr>
<tr>
<td>Mobile batcher mixers</td>
<td>Used for intermittent production of concrete at jobsite, or where only small quantities are required. A combined materials transporter and mobile batching and mixing system for quick, precise proportioning of specified concrete. One-man operation.</td>
<td>Trouble-free operation requires good preventive maintenance program on equipment. Materials must be identical to those in original mix design.</td>
<td></td>
</tr>
</tbody>
</table>
### Table 10-1. Methods and Equipment for Transporting and Handling Concrete (Continued)

<table>
<thead>
<tr>
<th>Equipment</th>
<th>Type and range of work for which equipment is best suited</th>
<th>Advantages</th>
<th>Points to watch for</th>
</tr>
</thead>
<tbody>
<tr>
<td>Nonagitating trucks</td>
<td>Used to transport concrete on short hauls over smooth roadways.</td>
<td>Capital cost of nonagitating equipment is lower than that of truck agitators or mixers.</td>
<td>Concrete slump should be limited. Possibility of segregation. Height is needed for high lift of truck body upon discharge.</td>
</tr>
<tr>
<td>Pneumatic guns (shotcrete)</td>
<td>Used where concrete is to be placed in difficult locations and where thin sections and large areas are needed.</td>
<td>Ideal for placing concrete in freeform shapes, for repairing structures, for protective coatings, thin linings, and building walls with one-sided forms.</td>
<td>Quality of work depends on skill of those using equipment. Only experienced nozzlemen should be employed.</td>
</tr>
<tr>
<td>Pumps</td>
<td>Used to convey concrete directly from central discharge point at jobsite to formwork or to secondary discharge point.</td>
<td>Pipelines take up little space and can be readily extended. Delivers concrete in continuous stream. Pump can move concrete both vertically and horizontally. Truck-mounted pumps can be delivered when necessary to small or large projects. Tower-crane mounted pump booms provide continuous concrete for tall building construction.</td>
<td>Constant supply of freshly-mixed concrete is needed with average consistency and without any tendency to segregate. Care must be taken in operating pipeline to ensure an even flow and to clean out at conclusion of each operation. Pumping vertically, around bends, and through flexible hose will considerably reduce the maximum pumping distance.</td>
</tr>
<tr>
<td>Screw spreaders</td>
<td>Used for spreading concrete over large flat areas, such as in pavements and bridge decks.</td>
<td>With a screw spreader a batch of concrete discharged from a bucket or truck can be quickly spread over a wide area to a uniform depth. The spread concrete has good uniformity of compaction before vibration is used for final compaction.</td>
<td>Screw spreaders are normally used as part of a paving train. They should be used for spreading before vibration is applied.</td>
</tr>
<tr>
<td>Tremies</td>
<td>For placing concrete underwater.</td>
<td>Can be used to funnel concrete down through the water into the foundation or other part of the structure being cast.</td>
<td>Precautions are needed to ensure that the tremie discharge end is always buried in fresh concrete, so that a seal is preserved between water and concrete mass. Diameter should be 250 to 300 mm (10 to 12 in.) unless pressure is available. Concrete mixture needs more cement, 390 kg/m³ (658 lb/yd³), and greater slump, 150 to 230 mm (6 to 9 in.), because concrete must flow and consolidate without any vibration.</td>
</tr>
<tr>
<td>Truck agitators</td>
<td>Used to transport concrete for all uses in pavements, structures, and buildings. Haul distances must allow discharge of concrete within 1½ hours, but limit may be waived under certain circumstances.</td>
<td>Truck agitators usually operate from central mixing plants where quality concrete is produced under controlled conditions. Discharge from agitators is well controlled. There is uniformity and homogeneity of concrete on discharge.</td>
<td>Timing of deliveries should suit job organization. Concrete crew and equipment must be ready onsite to handle concrete.</td>
</tr>
<tr>
<td>Truck mixers</td>
<td>Used to transport concrete for uses in pavements, structures, and buildings. Haul distances must allow discharge of concrete within 1½ hours, but limit may be waived under certain circumstances.</td>
<td>No central mixing plant needed, only a batching plant, since concrete is completely mixed in truck mixer. Discharge is same as for truck agitator.</td>
<td>Timing of deliveries should suit job organization. Concrete crew and equipment must be ready onsite to handle concrete. Control of concrete quality is not as good as with central mixing.</td>
</tr>
<tr>
<td>Wheelbarrows and buggies</td>
<td>For short flat hauls on all types of onsite concrete construction, especially where accessibility to work area is restricted.</td>
<td>Very versatile and therefore ideal inside and on jobsites where placing conditions are constantly changing.</td>
<td>Slow and labor intensive.</td>
</tr>
</tbody>
</table>
There have been few, if any, major changes in the principles of conveying concrete during the last 75 years. What has changed is the technology that led to development of better machinery to do the work more efficiently. The wheelbarrow and buggy, although still used, have advanced to become the power buggy (Fig. 10-8); the bucket hauled over a pulley wheel has become the bucket and crane (Fig. 10-9); and the horse-drawn wagon is now the ready mixed concrete truck (Figs. 10-10 and 10-11).

Fig. 10-8. Versatile power buggy can move all types of concrete over short distances. (IMG12249)

Fig. 10-9. Concrete is easily lifted to its final location by bucket and crane. (IMG12230)

Fig. 10-10. Ready mixed concrete can often be placed in its final location by direct chute discharge from a truck mixer. (IMG12243)

Fig. 10-11. In comparison to conventional rear-discharge trucks, front-discharge truck mixers provide the driver with more mobility and control for direct discharge into place. (IMG12397)

Years ago concrete was placed in reinforced concrete buildings by means of a tower and long chutes. This was a guyed tower centrally placed on the site with a hopper at the top to which concrete was hauled by winch. A series of chutes suspended from the tower allowed the concrete to flow by gravity directly to the point required. As concrete-framed buildings became taller, the need to hoist reinforcement and formwork as well as concrete to higher levels led to the development of the tower crane—a familiar sight on the building skyline today (Fig. 10-12). It is fast and versatile, but the fact that it has only one hook must be considered when planning a job.

The conveyor belt is old in concept and much changed over the years (Fig. 10-13). Recently, truck-mixer-mounted conveyor belts have come into use (Fig. 10-14). The pneumatic process for shotcreting was patented in 1911 and is
literally unchanged (see Chapter 18). The first mechanical concrete pump was developed and used in the 1930s and the hydraulic pump was developed in the 1950s. The advanced mobile pump with hydraulic placing boom (Fig. 10-15) is probably the single most important innovation in concrete handling equipment. It is economical to use in placing both large and small quantities of concrete, depending on jobsite conditions. For small to medium size projects, a combination of truck mixer and boom pump can be used to transport and place concrete. The screw spreader (Fig. 10-16) has been very effective in placing and distributing concrete for pavements. Screw spreaders can place a uniform depth of concrete quickly and efficiently.

Fig. 10-12. The tower crane and bucket can easily handle concrete for tall-building construction. (IMG12396)

Fig. 10-13. The conveyor belt is an efficient, portable method of handling concrete. A drop chute prevents concrete from segregating as it leaves the belt; a scraper prevents loss of mortar. Conveyor belts can be operated in series and on extendable booms of hydraulic cranes. (IMG12395)

Fig. 10-14. A conveyor belt mounted on a truck mixer places concrete up to about 12 meters (40 feet) without the need for additional handling equipment. (IMG12244)

Fig. 10-15. (top) A truck-mounted pump and boom can conveniently move concrete vertically or horizontally to the desired location. (bottom) View of concrete discharging from flexible hose connected to rigid pipeline leading from the pump. Rigid pipe is used in pump booms and in pipelines to move concrete over relatively long distances. Up to 8 m (25 ft) of flexible hose may be attached to the end of a rigid line to increase placement mobility. (IMG12394, IMG12393)
Choosing the Best Method

The first thing to look at is the type of job, its physical size, the total amount of concrete to be placed, and the time schedule. Studying the job details further will tell how much of the work is below, at, or above ground level. This aids in choosing the concrete handling equipment necessary for placing concrete at the required levels.

Concrete must be moved from the mixer to the point of placement as rapidly as possible without segregation or loss of ingredients. The transporting and handling equipment must have the capacity to move sufficient concrete so that cold joints are eliminated.

Work Above Ground Level

Conveyor belt, crane and bucket, hoist, pump, or the ultimate sky-hook, the helicopter, can be used for lifting concrete to locations above ground level (Fig. 10-17). The tower crane (Fig. 10-12) and pumping boom (Fig. 10-18) are the right tools for tall buildings. The volume of concrete needed per floor as well as boom placement and length affect the use of a pump; large volumes minimize pipeline movement in relation to output.

The specifications and performance of transporting and handling equipment are being continuously improved. The best results and lowest costs will be realized if the work is planned to get the most out of the equipment and if the equipment is flexibly employed to reduce total job cost. Any method is expensive if it does not get the job done. Panarese (1987) is very helpful in deciding which method to use based on capacity and range information for various methods and equipment.
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Fig. 10-17. For work aboveground or in inaccessible sites, a concrete bucket can be lifted by helicopter. (Source: Paschal)

Fig. 10-18. A pump boom mounted on a mast and located near the center of a structure can frequently reach all points of placement. It is especially applicable to tall buildings where tower cranes cannot be tied up with placing concrete. Concrete is supplied to the boom through a pipeline from a ground-level pump. Concrete can be pumped hundreds of meters (feet) vertically with these pumping methods. (IMG12245)
CHAPTER 11
Placing and Finishing Concrete

PREPARATION BEFORE PLACING

Preparation prior to placing concrete for pavements or slabs on grade includes compacting, trimming, and moistening the subgrade (Figs. 11-1, 11-2, and 11-3); erecting the forms; and setting the reinforcing steel and other embedded items securely in place. Moistening the subgrade is important, especially in hot, dry weather to keep the dry subgrade from drawing too much water from the concrete; it also increases the immediate air-moisture level thereby decreasing the amount of evaporation from the concrete surface. The strength or bearing capacity of the subgrade should be adequate to support anticipated structural loads.

In cold weather, concrete must not be placed on a frozen subgrade. Snow, ice, and other debris must be removed from within the forms before concrete is placed. Where concrete is to be deposited on rock or hardened concrete, all loose material must be removed, and cut faces should be nearly vertical or horizontal rather than sloping.

Recently placed concrete requiring an overlay is usually roughened shortly after hardening to produce a better

Fig. 11-1. A base course foundation for concrete pavement is shaped by an auto-trimmer to design grades, cross section and alignment by automatic sensors that follow string lines. (IMG12388)

Fig. 11-2. Water trucks with spray-bars are used to moisten subgrades and base course layers to achieve adequate compaction and to reduce the amount of water drawn out of concrete as it’s placed. (IMG12371)

Fig. 11-3. (top) Adequate compaction of a base course foundation for concrete pavement can be achieved by using a vibratory roller. (bottom) Vibratory plate compactors are also used to prepare subgrades under slabs. (IMG12370, IMG12369)
bond with the next placement. As long as no laitance (a weak layer of concrete), dirt, or loose particles are present, newly hardened concrete requires little preparation prior to placing freshly mixed concrete on it. When in service for a period of time, old hardened concrete usually requires mechanical cleaning and roughening prior to placement of new concrete. The subject of placing freshly mixed concrete on hardened concrete is discussed in more detail under the sections entitled “Placing on Hardened Concrete” and “Construction Joints.”

Forms should be accurately set, clean, tight, adequately braced, and constructed of or lined with materials that will impart the desired off-the-form finish to the hardened concrete. Wood forms, unless oiled or otherwise treated with a form-release agent, should be moistened before placing concrete, otherwise they will absorb water from the concrete and swell. Forms should be made for removal with minimum damage to the concrete. With wood forms, use of too large or too many nails should be avoided to facilitate removal and reduce damage. For architectural concrete, the form-release agent should be a nonstaining material. See Hurd (1979) and ACI Committee 347 (2004) for more information on formwork.

Reinforcing steel should be clean and free of loose rust or mill scale when concrete is placed. Unlike subgrades, reinforcing steel can be colder than 0°C (32°F) with special considerations. See “Concreting Aboveground” in Chapter 14 for more details. Mortar splattered on reinforcing bars from previous placements need not be removed from steel and other embedded items if the next lift is to be completed within a few hours; loose, dried mortar, however, must be removed from items that will be encased by later lifts of concrete.

All equipment used to place concrete must be clean and in good working condition. Standby equipment should be available in the event of a breakdown.

**DEPOSITING THE CONCRETE**

Concrete should be deposited continuously as near as possible to its final position without objectionable segregation (Figs. 11-4, 11-5, 11-6, 11-7, and 11-8). In slab construction, placing should be started along the perimeter at one end of the work with each batch discharged against previously placed concrete. The concrete should not be dumped in separate piles and then leveled and worked together; nor should the concrete be deposited in large piles and moved horizontally into final position. Such practices result in segregation because mortar tends to flow ahead of the coarser material.

In general, concrete should be placed in walls, thick slabs, or foundations in horizontal layers of uniform thickness; each layer should be thoroughly consolidated before the next is placed. The rate of placement should be rapid enough so that previously placed concrete has not yet set when the next layer of concrete is placed upon it. Timely placement and adequate consolidation will prevent flow lines, seams, and planes of weakness (cold joints) that result...
from placing freshly mixed concrete on concrete past initial set. Layers should be about 150 to 500 mm (6 to 20 in.) thick for reinforced members and 380 to 500 mm (15 to 20 in.) thick for mass work; the thickness will depend on the width between forms and the amount of reinforcement.

To avoid segregation, concrete should not be moved horizontally over too long a distance as it is being placed in forms or slabs. In some work, such as placing concrete in sloping wingwalls or beneath window openings in walls, it is necessary to move the concrete horizontally within the forms, but this should be kept to a minimum.

Where standing water is present, concrete should be placed in a manner that displaces the water ahead of the concrete but does not allow water to be mixed in with the concrete; to do so will reduce the quality of the concrete. In all cases, water should be prevented from collecting at the ends, in corners, and along faces of forms. Care should be taken to avoid disturbing saturated subgrade soils so they maintain sufficient bearing capacity to support structural loads.

Chutes and dropchutes are used to move concrete to lower elevations without segregation and spattering of mortar on reinforcement and forms. Properly designed concrete has been allowed to drop by free fall into caissons. Results of a field test to determine if concrete could be dropped vertically 15 meters (50 ft) into a caisson without segregation proved that there was no significant difference in aggregate gradation between control samples as delivered and free-fall samples taken from the bottom of the caisson (Turner 1970). More recent field studies indicate that free fall of concrete from heights of up to 46 m (150 ft)—directly over reinforcing steel or at a high slump—does not result in segregation of the concrete ingredients nor reduce compressive strength (Suprenant 2001). However, if a baffle is not used to control the flow of concrete onto sloped surfaces at the end of an inclined chute, segregation can occur.

Concrete is sometimes placed through openings, called windows, in the sides of tall, narrow forms. When a chute discharges directly through the opening without controlling concrete flow at the end of the chute there is danger of segregation. A collecting hopper should be used outside the opening to permit the concrete to flow more smoothly through the opening; this will decrease the tendency to segregate.

When concrete is placed in tall forms at a fairly rapid rate, some bleed water may collect on the top surface, especially with non-air-entrained concrete. Bleeding can be reduced by placing more slowly and by using concrete of a stiffer consistency, particularly in the lower portion of the form. When practical, concrete should be placed to a level 300 mm to 400 mm (about a foot) below the top of tall forms and an hour or so allowed for the concrete to partially set. Placing should resume before the surface hardens to avoid formation of a cold joint. If practical to work around vertical reinforcing steel, it is good practice to overfill the form by 25 mm (an inch) or so and cut off the excess concrete after it has stiffened and bleeding has ceased.

In monolithic placement of deep beams, walls, or columns, to avoid cracks between structural elements, concrete placement should stop (usually about 1 hr) to allow settlement of the deep element before concreting is continued in any slabs, beams, or girders framing into them. The delay should be short enough to allow the next layer of concrete to knit with the previous layer by vibration, thus preventing cold joints and honeycombing (ACI Committee 304 2000). Haunches and column capitals are considered part of the floor or roof slab and should be placed integrally with them.

**PLACING CONCRETE UNDERWATER**

Concrete should be placed in the air rather than underwater whenever possible. When it must be placed underwater, the work should be done under experienced
supervision. The basic principles for normal concrete work in the dry apply, with common sense, to underwater concreting. The following special points, however, should be observed:

The slump of the concrete should be specified at 150 to 230 mm (6 to 9 in.) and the mixture should have a maximum water-cementitious materials ratio of 0.45. Generally, the cementitious materials content will be 390 kg/m³ (600 lb/yd³) or more.

It is important that the concrete flow without segregation; therefore, the aim in proportioning should be to obtain a cohesive mixture with high workability. Anti-washout admixtures can be used to make concrete cohesive enough to be placed in limited depths of water, even without tremies. Using rounded aggregates, a higher percentage of fines, and entrained air should help to obtain the desired consistency.

The current in the water through which the concrete is deposited should not exceed 3 m (10 ft) per minute.

Methods for placing concrete underwater include the following: tremie, concrete pump, bottom-dump buckets, grouting preplaced aggregate, toggle bags, bagwork, and the diving bell.

A tremie is a smooth, straight pipe long enough to reach the lowest point to be concreted from a working platform above the water. The diameter of the tremie pipe should be at least 8 times the diameter of the maximum size of aggregate. A hopper to receive the concrete is attached to the top of the pipe. The lower end of the tremie should be kept buried in the fresh concrete to maintain a seal below the rising top surface and to force the concrete to flow in beneath it by pressure. Placing should be continuous with as little disturbance to the previously placed concrete as possible. The top surface should be kept as level as possible. See ACI Committee 304 (2000) for additional information.

Mobile concrete pumps with a variable radius boom makes easy work of placing concrete underwater. Because the flexible hose on a concrete pump is similar to a tremie, the same placement techniques apply.

With the grouting preplaced aggregate method, the forms are first filled with clean coarse aggregate, then the voids in the coarse aggregate are filled with a grout to produce concrete. Grouting preplaced aggregate has advantages when placing concrete in flowing water. Concrete can be placed more quickly and economically than by conventional placement methods. However, the method is very specialized and should only be performed by qualified experienced personnel.

Sand bags half full of plastic concrete can be used for small jobs, filling gaps, or temporary work. The tied end should face away from the outside.

**SPECIAL PLACING TECHNIQUES**

Concrete may be placed by methods other than the usual cast-in-place method. These methods, such as shotcreting, are described in Chapter 18. No matter what method is used, the basics of mixing, placing, consolidating, and curing apply to all portland cement concretes.

**CONSOLIDATING CONCRETE**

Consolidation is the process of compacting fresh concrete; to mold it within the forms and around embedded items and reinforcement; and to eliminate stone pockets, honeycomb, and entrapped air (Fig. 11-9). It should not remove significant amounts of intentionally entrained air in air-entrained concrete.

Consolidation is accomplished by hand or by mechanical methods. The method chosen depends on the consistency of the mixture and the placing conditions, such as complexity of the formwork and amount and spacing of reinforcement. Generally, mechanical methods using either internal or external vibration are the preferred methods of consolidation.

Workable, flowing mixtures can be consolidated by hand rodding, that is, thrusting a tamper rod or other suitable tool repeatedly into the concrete. The rod should be long enough to reach the bottom of the form or lift and thin enough to easily pass between the reinforcing steel and the forms. Low-slump concrete can be transformed into flowing concrete for easier consolidation through the use of superplasticizers without the addition of water to the concrete mixture.

Spading can be used to improve the appearance of formed surfaces. A flat, spadelike tool should be repeatedly inserted and withdrawn adjacent to the form. This forces the larger coarse aggregates away from the forms and assists entrapped air voids in their upward movement toward the top surface where they can escape. A mixture designed to be readily consolidated by hand methods should not be consolidated by mechanical methods; otherwise, the concrete is likely to segregate under intense mechanical action.

![Fig. 11-9. Honeycomb and rock pockets are the results of inadequate consolidation. (IMG12262)](image-url)
Even in highly reinforced elements, proper mechanical consolidation makes possible the placement of stiff mixtures with the low water-cementitious materials ratios and high coarse-aggregate contents associated with high-quality concrete (Fig. 11-10). Among the mechanical methods are centrifugation, used to consolidate moderate-to-high-slump concrete in making pipes, poles, and piles; shock or drop tables, used to compact very stiff low-slump concrete in the manufacture of architectural precast units; and vibration—internal and external.

**Vibration**

Vibration, either internal or external, is the most widely used method for consolidating concrete. When concrete is vibrated, the internal friction between the aggregate particles is temporarily destroyed and the concrete behaves like a liquid; it settles in the forms under the action of gravity and the large entrapped air voids rise more easily to the surface. Internal friction is reestablished as soon as vibration stops.

Vibrators, whether internal or external, are usually characterized by their frequency of vibration, expressed as the number of vibrations per second (Hertz), or vibrations per minute (vpm); they are also designated by the amplitude of vibration, which is the deviation in millimeters (inches) from the point of rest. The frequency of vibration can be measured using a vibrating reed tachometer.

When vibration is used to consolidate concrete, a standby vibrator should be on hand at all times in the event of a mechanical breakdown.

**Internal Vibration.** Internal or immersion-type vibrators, often called spud or poker vibrators (Figs. 11-10 and 11-11), are commonly used to consolidate concrete in walls, columns, beams, and slabs. Flexible-shaft vibrators consist of a vibrating head connected to a driving motor by a flexible shaft. Inside the head, an unbalanced weight connected to the shaft rotates at high speed, causing the head to revolve in a circular orbit. The motor can be powered by electricity, gasoline, or air. The vibrating head is usually cylindrical with a diameter ranging from 20 to 180 mm (¾ to 7 in.). Some vibrators have an electric motor built right into the head, which is generally at least 50 mm (2 in.) in diameter. The dimensions of the vibrator head as well as its frequency and amplitude in conjunction with the workability of the mixture affect the performance of a vibrator.

Small-diameter vibrators have high frequencies ranging from 160 to 250 Hz (10,000 to 15,000 vpm) and low amplitudes ranging between 0.4 and 0.8 mm (0.016 and 0.03 in.). As the diameter of the head increases, the frequency decreases and the amplitude increases. The effective radius of action of a vibrator increases with increasing diameter. Vibrators with a diameter of 20 to 40 mm (¾ to 1½ in.) have a radius of action in freshly mixed concrete ranging between 75 and 150 mm (3 and 6 in.), whereas the radius of action for vibrators of 50- to 80-mm (2- to 3-in.) diameter ranges between 180 and 350 mm (7 and 14 in.). Table 11-1 shows the range of characteristics and applications for internal vibrators for various applications.

Proper use of internal vibrators is important for best results. Vibrators should not be used to move concrete horizontally since this causes segregation. Whenever possible, the vibrator should be lowered vertically into the concrete at regularly spaced intervals and allowed to descend by gravity. It should penetrate to the bottom of the layer being placed and at least 150 mm (6 in.) into any previously placed layer. The height of each layer or lift should be about the length of the vibrator head or generally a maximum of 500 mm (20 in.) in regular formwork.

In thin slabs, the vibrator should be inserted at an angle or horizontally in order to keep the vibrator head completely immersed. However, the vibrator should not be dragged around randomly in the slab. For slabs on grade, the vibrator should not make contact with the sub-
Table 11-1. Range of Characteristics, Performance, and Applications of Internal* Vibrators

<table>
<thead>
<tr>
<th>Group</th>
<th>Diameter of head, mm (in.)</th>
<th>Recommended frequency, vibrations per minute**</th>
<th>Suggested values of</th>
<th>Approximate values of</th>
<th>Application</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
<td>Eccentric moment, mm-kg in.-lb (10^-5)</td>
<td>Average amplitude, mm (in.)</td>
<td>Centrifugal force, kg (lb)</td>
</tr>
<tr>
<td>1</td>
<td>20-40 (¾-1½)</td>
<td>9000-15,000</td>
<td>3.5-12 (0.03-0.10)</td>
<td>0.4-0.8 (0.015-0.03)</td>
<td>45-180 (100-400)</td>
</tr>
<tr>
<td>2</td>
<td>30-60 (1¼-2½)</td>
<td>8500-12,500</td>
<td>9-29 (0.08-0.25)</td>
<td>0.5-1.0 (0.02-0.04)</td>
<td>140-400 (300-900)</td>
</tr>
<tr>
<td>3</td>
<td>50-90 (2-3½)</td>
<td>8000-12,000</td>
<td>23-81 (0.20-0.70)</td>
<td>0.6-1.3 (0.025-0.05)</td>
<td>320-900 (700-2000)</td>
</tr>
<tr>
<td>4</td>
<td>80-150 (3-6)</td>
<td>7000-10,500</td>
<td>8-290 (0.70-2.5)</td>
<td>0.8-1.5 (0.03-0.06)</td>
<td>680-1800 (1500-4000)</td>
</tr>
<tr>
<td>5</td>
<td>130-150 (5-6)</td>
<td>5500-8500</td>
<td>260-400 (2.25-3.50)</td>
<td>1.0-2.0 (0.04-0.08)</td>
<td>1100-2700 (2500-6000)</td>
</tr>
</tbody>
</table>

* Generally, extremely dry or very stiff concrete does not respond well to internal vibrators.
** While vibrator is operating in concrete.
† Distance over which concrete is fully consolidated.
‡ Assumes the insertion spacing is 1½ times the radius of action, and that vibrator operates two-thirds of time concrete is being placed. These ranges reflect not only the capability of the vibrator but also differences in workability of the mix, degree of deaeration desired, and other conditions experienced in construction.
Adapted from ACI 309.

The distance between insertions should be about 1½ times the radius of action so that the area visibly affected by the vibrator overlaps the adjacent previously vibrated area by a few centimeters (inches).

The vibrator should be held stationary until adequate consolidation is attained and then slowly withdrawn. An insertion time of 5 to 15 seconds will usually provide adequate consolidation. The concrete should move to fill the hole left by the vibrator on withdrawal. If the hole does not refill, reinsertion of the vibrator at a nearby point should solve the problem.

Adequacy of internal vibration is judged by experience and by changes in the surface appearance of the concrete. Changes to watch for are the embedment of large aggregate particles, general batch leveling, the appearance of a thin film of mortar on the top surface, and the cessation of large bubbles of entrapped air escaping at the surface. Internal vibration may significantly affect the entrained-air-void system in concrete (Stark 1986, and Hover 2001). Detailed guidance for proper vibration should be followed (ACI Committee 309).

Allowing a vibrator to remain immersed in concrete after paste accumulates over the head is bad practice and can result in nonuniformity. The length of time that a vibrator should be left in the concrete will depend on the workability of the concrete, the power of the vibrator, and the nature of the section being consolidated.

In heavily-reinforced sections where an internal vibrator cannot be inserted, it is sometimes helpful to vibrate the reinforcing bars by attaching a form vibrator to
their exposed portions. This practice eliminates air and water trapped under the reinforcing bars and increases bond between the bars and surrounding concrete; use this method only if the concrete is still workable under the action of vibration. Internal vibrators should not be attached to reinforcing bars for this purpose because the vibrators may be damaged.

Revibration of previously compacted concrete can be done to both fresh concrete as well as any underlying layer that has partially hardened. Revibration is used to improve bond between concrete and reinforcing steel, release water trapped under horizontal reinforcing bars, and remove additional entrapped air voids. In general, if concrete becomes workable under revibration, the practice is not harmful and may be beneficial.

**External Vibration.** External vibrators can be form vibrators, vibrating tables, or surface vibrators such as vibratory screeds, plate vibrators, vibratory roller screeds, or vibratory hand floats or trowels. Form vibrators, designed to be securely attached to the outside of the forms, are especially useful (1) for consolidating concrete in members that are very thin or congested with reinforcement, (2) to supplement internal vibration, and (3) for stiff mixes where internal vibrators cannot be used.

Attaching a form vibrator directly to the form generally is unsatisfactory. Rather, the vibrator should be attached to a steel plate that in turn is attached to steel I-beams or channels passing through the form stiffeners themselves in a continuous run. Loose attachments can result in significant vibration energy losses and inadequate consolidation.

Form vibrators can be either electrically or pneumatically operated. They should be spaced to distribute the intensity of vibration uniformly over the form; optimum spacing is best found by experimentation. Sometimes it may be necessary to operate some of the form vibrators at a different frequency for better results; therefore, it is recommended that form vibrators be equipped with controls to regulate their frequency and amplitude. Duration of external vibration is considerably longer than for internal vibration—generally between 1 and 2 minutes. A reed tachometer can not only determine frequency of vibration, but also give a rough estimate of amplitude of vibration by noting the oscillation of the reed at various points along the forms. This will assist in identifying dead spots or weak areas of vibration. A vibrograph could be used if more reliable measurements of frequency and amplitude are needed.

Form vibrators should not be applied within the top meter (yard) of vertical forms. Vibration of the top of the form, particularly if the form is thin or inadequately stiffened, causes an in-and-out movement that can create a gap between the concrete and the form. Internal vibrators are recommended for use in this area of vertical forms.

Vibrating tables are used in precasting plants. They should be equipped with controls so that the frequency and amplitude can be varied according to the size of the element to be cast and the consistency of the concrete. Stiffer mixtures generally require lower frequencies (below 6000 vpm) and higher amplitudes (over 0.13 mm [0.005 in.]) than more workable mixtures. Increasing the frequency and decreasing the amplitude as vibration progresses will improve consolidation.

Surface vibrators, such as vibratory screeds (Figs. 11-12, 11-13, and 11-14), are used to consolidate concrete in floors and other flatwork. Vibratory screeds give positive control of the strikeoff operation and save a great deal of labor. When using this equipment, concrete need not have slumps in excess of 75 mm (3 in.). For greater than 75 mm slumps, care should be taken because surface vibration of such concrete will result in an excessive accumulation of mortar and fine material on the surface; this may reduce wear resistance. For the same reason, surface vibrators...
Segregation from striking reinforcing steel without adequate vibration may also contribute to streaking.

Cold joints are a discontinuity resulting from a delay in placement that allowed one layer to harden before the adjacent concrete was placed. The discontinuity can reduce the structural integrity of a concrete member if the successive lifts did not properly bond together. The concrete can be kept alive by revibrating it every 15 minutes or less depending on job conditions. However, once the time of initial setting approaches, vibration should be discontinued and the surface should be suitably prepared for the additional concrete.

Placement lines or “pour” lines are dark lines between adjacent placements of concrete batches. They may occur if, while vibrating the overlying layer, the vibrator did not penetrate the underlying layer enough to knit the layers together.

Subsidence cracking may occur at or near the initial setting time as concrete settles over reinforcing steel in relatively deep elements that have not been adequately vibrated. Revibration at the latest time that the vibrator will sink into the concrete under its own weight may eliminate these cracks.

Defects from overvibration include: (1) segregation as vibration and gravity causes heavier aggregates to settle while lighter aggregates rise; (2) sand streaks; (3) loss of entrained air in air-entrained concrete; (4) excessive form deflections or form damage; and (5) form failure caused by excessive pressure from vibrating the same location too long and/or placing concrete more quickly than the designed rate of pour.

Undervibration is more often a problem than overvibration.

CONCRETE SLABS

Concrete slabs can be finished in many ways, depending on the intended service use. Various colors and textures, such as exposed-aggregate or a pattern-stamped surface, may be called for. Some surfaces may require only strikeoff and screeding to proper contour and elevation, while for other surfaces a broomed, floated, or troweled finish may be specified. Details are given in ACI Committee 309.

Consequences of Improper Vibration. Following are some of the worst defects caused by undervibration: (1) honeycomb; (2) excessive amount of entrapped air voids, often called bugholes; (3) sand streaks; (4) cold joints; (5) placement lines; and (6) subsidence cracking.

Honeycomb results when the spaces between coarse aggregate particles do not become filled with mortar. Faulty equipment, improper placement procedures, a concrete mix containing too much coarse aggregate, or congested reinforcement can cause honeycomb.

Excessive entrapped air voids are similar to, but not as severe as honeycomb. Vibratory equipment and operating procedures are the primary causes of excessive entrapped air voids, but the other causes of honeycomb apply too.

Sand streaks result when heavy bleeding washes mortar out from along the form. A wet, harsh mixture that lacks workability because of an insufficient amount of mortar or fine aggregate may cause sand streaking.

Segregation from striking reinforcing steel without adequate vibration should not be operated after the concrete has been adequately consolidated.

Because surface vibration of concrete slabs is least effective along the edges, a spud or poker-type vibrator should be used along the edge forms immediately before the vibratory screed is applied.

Vibratory screeds are used for consolidating slabs up to 250 mm (10 in.) thick, provided such slabs are nonreinforced or only lightly reinforced (welded-wire fabric). Internal vibration or a combination of internal and surface vibration is recommended for reinforced slabs. More detailed information regarding internal and external vibration of concrete can be obtained from ACI Committee 309.

Fig. 11-14. A laser level stimulating the sensors on this screed guides the operator as he strikes off the concrete. Screed poles and chairs are not needed and fewer workers are required to place concrete. Laser screeds interfaced with total station surveying equipment can also strike off sloped concrete surfaces. (IMG12385)
ditions on the setting time of the concrete and the size of
the placement to be completed.

Subgrade Preparation

Cracks, slab settlement, and structural failure can often be
traced to an inadequately prepared and poorly compacted
subgrade. The subgrade on which a slab on ground is to
be placed should be well drained, of uniform bearing
capacity, level or properly sloped, and free of sod, organic
matter, and frost. The three major causes of nonuniform
support are: (1) the presence of soft unstable saturated
soils or hard rocky soils, (2) backfilling without adequate
compaction, and (3) expansive soils. Uniform support
cannot be achieved by merely dumping granular material
on a soft area. To prevent bridging and settlement
cracking, soft or saturated soil areas and hard spots (rocks)
should be dug out and filled with soil similar to the sur-
rounding subgrade or if a similar soil is not available, with
granular material such as sand, gravel, or crushed stone.
All fill materials must be compacted to provide the same
uniform support as the rest of the subgrade. Proof rolling
the subgrade using a fully-loaded dump truck or similar
heavy equipment is commonly used to identify areas of
unstable soils that need additional attention.

During subgrade preparation, it should be remem-
bered that undisturbed soil is generally superior to com-
pacted material for supporting concrete slabs. Expansive,
compressible, and potentially troublesome soils should be
evaluated by a geotechnical engineer; a special slab design
may be required.

The subgrade should be moistened with water in
advance of placing concrete, but should not contain pud-
dles or wet, soft, muddy spots when the concrete is placed.

Subbase

A satisfactory slab on ground can be built without a sub-
base. However, a subbase is frequently placed on the sub-
grade as a leveling course to equalize minor surface
irregularities, enhance uniformity of support, bring the
site to the desired grade, and serve as a capillary break
between the slab and the subgrade.

Where a subbase is used, the contractor should place
and compact to near maximum density a 100-mm (4-in.)
thickness of granular material such as sand, gravel,
crushed stone, or slag. If a thicker subbase is needed for
achieving the desired grade, the material should be com-
pacted in thin layers about 100 mm (4 in.) deep unless tests
determine compaction of thicker lifts is possible (Fig. 11-15).
Subgrades and subbases can be compacted with small plate
vibrators, vibratory rollers, or hand tampers. Unless the sub-
base is well compacted, it is better not to use a subbase;
simply leave the subgrade uncovered and undisturbed.

Vapor Retarders and
Moisture-Problem Prevention

Many of the moisture problems associated with enclosed
slabs on ground (floors) can be minimized or eliminated
by (1) sloping the landscape away from buildings, (2) us-
ing a 100-mm (4-in.) thick granular subbase to form a
capillary break between the soil and the slab, (3) pro-
viding drainage for the granular subbase to prevent
water from collecting under the slab, (4) installing foun-
dation drain tile, and (5) installing a vapor retarder, often
polyethylene sheeting.

For years vapor retarders have been mistakenly called
vapor barriers. A vapor retarder slows the movement of
water vapor by use of a 0.15 to 0.25 mm (6 to 10 mil) poly-
ethylene film that is overlapped approximately 150 mm
(6 in.) at the edges. A vapor retarder does not stop 100% of
vapor migration; a vapor barrier does. Vapor barriers are
thick, rugged multiple-ply-reinforced membranes that are
sealed at the edges. Vapor retarders are discussed in this
text because they are more commonly used; but many of
the following principles apply to vapor barriers as well.

A vapor retarder should be placed under all concrete
floors on ground that are likely to receive an impermeable
floor covering such as sheet vinyl tile or be used for any
purpose where the passage of water vapor through the
floor might damage moisture-sensitive equipment or
materials in contact with the floor. However, a few project
sites with deep groundwater tables and sandy soils con-
taining very low silt or clay contents may not require the
use of a vapor retarder under concrete slabs.

Vapor retarders placed directly under concrete slabs
may increase the time delay before final finishing due to
longer bleeding times, particularly in cold weather. To
minimize this effect, a minimum 75-mm (3-in.) thick layer
of approved granular, self-draining compactible subbase material should be placed over the vapor barrier (or insulation if present) (ACI Committee 302). Some contractors find only 75 mm of granular sand over polyethylene sheeting to be slippery, somewhat dangerous, and difficult to keep in place while concreting. A 150 to 200 mm (6 to 8 in.)-thick subbase will alleviate this problem. The subbase over a vapor retarder must be kept from getting saturated by rain or construction activities to prevent excessive vapor migration after the concrete slab is placed.

If concrete is placed directly on a vapor retarder, the water-cementitious materials ratio should be kept low (0.45 or less) because excess mix water can only escape to the surface as bleed water. Because of a longer bleeding period, settlement cracking over reinforcement and shrinkage cracking is more likely. For more information see ACI (2001) and ACI Committee 302.

Good quality, well-consolidated concrete at least 100-mm (4-in.) thick is practically impermeable to the passage of liquid water unless the water is under considerable pressure; however, such concrete—even concrete several times as thick—is not impermeable to the passage of water vapor.

Water vapor that passes through a concrete slab evaporates at the surface if it is not sealed. Floor coverings such as linoleum, vinyl tile, carpeting, wood, and synthetic surfacing effectively seal the moisture within the slab; eventually this moisture may deteriorate latex adhesives causing the floor covering to loosen, buckle, or blister.

To prevent problems with floor covering materials caused by moisture within the concrete, the following steps should be taken: (1) use a low water-cement ratio concrete, (2) moist-cure the slab for 7 days, (3) allow the slab a 2-or-more-month drying period (Hedenblad 1997 and 1998), and (4) test the slab moisture condition before installing the floor covering.

In one commonly used test (ASTM F1869), the moisture vapor emission rate from a concrete slab is determined by taping a domed plastic vapor barrier with a desiccant under it to the floor. After about 72 hours the desiccant is weighed and the moisture vapor emission rate is calculated. The slab is considered dry enough for placing a flooring material if the moisture vapor emission rate is below either 1.4 or 2.3 kg/1000 m²/24 hr (3 or 5 lbs/1000 ft²/24 hr) depending on the type of floor covering to be installed. Flooring-material manufacturers often have their own recommended test and specified moisture limits for installing their product. For more information and additional tests for water vapor transmission, see “Moisture Testing” in Chapter 16, Kosmatka (1985), and PCA (2000).

Insulation is sometimes installed over the vapor barrier to assist in keeping the temperature of a concrete floor above the dew point; this helps prevent moisture in the air from condensing on the slab surface. This practice also creates a warm floor for thermal comfort. Codes and specifications often require insulation at the perimeter of a floor slab. Placing insulation under the entire slab on ground for energy conservation alone usually cannot be justified economically. For more details, see PCA (1985).

Forms

Edge forms and intermediate screeds should be set accurately and firmly to the specified elevation and contour for the finished surface. Slab edge forms are usually metal or wood braced firmly with wood or steel stakes to keep them in horizontal and vertical alignment. The forms should be straight and free from warping and have sufficient strength to resist concrete pressure without bulging. They should also be strong enough to support any mechanical placing and finishing equipment used.

Rain Protection

Prior to commencing placing of concrete, the owner and contractor should be aware of procedures to be followed in the event of rain during the placing operation. Protective coverings such as polyethylene sheets or tarpaulins should be available and onsite at all times. When rain occurs, all batching and placing operations should stop and the fresh concrete should be covered to the extent that the rain does not indent the surface of the concrete or wash away the cement paste. When rain ceases, the covering should be removed and remedial measures taken such as surface retexturing or reworking in-place plastic concrete, before concrete placing resumes.

Placing and Spreading

Placement should start at the far point of a slab and proceed toward the concrete supply source. The concrete, which should be placed as close as possible to its final position, should slightly overfill the forms and be roughly leveled with square ended shovels or concrete rakes. Large voids trapped in the concrete during placing should be removed by consolidation.

Screeding (Strikeoff)

Screeding or strikeoff is the process of cutting off excess concrete to bring the top surface of a slab to proper grade. The template used in the manual method is called a straightedge, although the lower edge may be straight or slightly curved, depending on the surface specified. It should be moved across the concrete with a sawing motion while advancing forward a short distance with each movement. There should be a surplus (surcharge) of concrete against the front face of the straightedge to fill in low areas as the straightedge passes over the slab. A 150-mm (6-in.) slab needs a surcharge of about 25 mm (1 in.). Straightedges are sometimes equipped with vibrators that
consolidate the concrete and assist in reducing the strikeoff work. This combination of straightedge and vibrator is called a vibratory screed (Fig. 11-12). Vibratory screeds are discussed earlier in this chapter under “Consolidating Concrete.” Screeding, consolidation, and bullfloating must be completed before excess bleed water collects on the surface.

**Bullfloating or Darbying**

To eliminate high and low spots and to embed large aggregate particles, a bullfloat or darby (Fig. 11-16 top) should be used immediately after strikeoff. The long-handle bullfloat (Fig. 11-16 bottom) is used on areas too large to reach with a short-handle darby. Highway straightedges are often used to obtain very flat surfaces (Fig. 11-17). For non-air-entrained concrete, these tools can be made of wood; for air-entrained concrete they should be of aluminum or magnesium alloy.

Bullfloating or darbying must be completed before bleed water accumulates on the surface. Care must be taken not to overwork the concrete as this could result in a less durable surface.

The preceding operations should level, shape, and smooth the surface and work up a slight amount of cement paste. Although sometimes no further finishing is required, on most slabs bullfloating or darbying is followed by one or more of the following finishing operations: edging, jointing, floating, troweling, and brooming. A slight hardening of the concrete is necessary before the start of any of these finishing operations. When the bleed-water sheen has evaporated and the concrete will sustain foot pressure with only about 6-mm (1⁄4-in.) indentation, the surface is ready for continued finishing operations (Fig. 11-18).
**Warning:** One of the principal causes of surface defects in concrete slabs is finishing while bleed water is on the surface. If bleed water is worked into the surface, the water-cement ratio is significantly increased which reduces strength, entrained-air content, and watertightness of the surface. Any finishing operation performed on the surface of a concrete slab while bleed water is present can cause crazing, dusting, or scaling (PCA 2001). Floating and troweling the concrete (discussed later) before the bleeding process is completed may also trap bleed water under the finished surface producing a weakened zone or void under the finished surface; this occasionally results in delaminations. The use of low-slump concrete with an adequate cement content and properly graded fine aggregate will minimize bleeding and help ensure maintenance-free slabs. For exterior slabs, air entrainment also reduces bleeding. ACI Committee 302 and Tarr and Farny (2008) present the placing and finishing techniques in more detail and PCA (2001) discusses defects.

**Edging and Jointing**

Edging is required along all edge forms and isolation and construction joints in floors and outdoor slabs such as walks, drives, and patios. Edging densifies and compacts concrete next to the form where floating and troweling are less effective, making it more durable and less vulnerable to scaling, chipping, and popouts.

In the edging operation, the concrete should be cut away from the forms to a depth of 25 mm (1 in.) using a pointed mason trowel or a margin trowel. Then an edger should be held almost flat on the surface and run with the front of the tool slightly raised to prevent the edger from leaving too deep an impression. Edging may be required after each subsequent finishing operation for interior slabs.

Proper jointing practices can eliminate unsightly random cracks. Contraction joints, also called control joints, are made with a hand groover or by inserting strips of plastic, wood, metal, or preformed joint material into the unhardened concrete. When hand methods are used to form control joints in exterior concrete slabs, mark the forms to accurately locate the joints. Prior to bullfloating, the edge of a thin strip of wood or metal may be used to knock down the coarse aggregate where the joint will be hand tooled. The slab should then be jointed immediately after bullfloating or in conjunction with the edging operation. Control joints also can be made in hardened concrete by sawing. Jointing is discussed further under the heading “Making Joints in Floors and Walls” later in this chapter.

**Floating**

After the concrete has been hand-edged and hand-jointed, it should be floated with a hand float or with a finishing machine using float blades (Fig. 11-18).

The purpose of floating is threefold: (1) to embed aggregate particles just beneath the surface; (2) to remove slight imperfections, humps, and voids; and (3) to compact the mortar at the surface in preparation for additional finishing operations. The concrete should not be overworked as this may bring an excess of water and fine material to the surface and result in subsequent surface defects.

Hand floats usually are made of fiberglass, magnesium, or wood. The metal float reduces the amount of work required because drag is reduced as the float slides more readily over the concrete surface. A magnesium float is essential for hand-floating air-entrained concrete because a wood float tends to stick to and tear the concrete surface. The light metal float also forms a smoother surface than the wood float.

The hand float should be held flat on the concrete surface and moved with a slight sawing motion in a sweeping arc to fill in holes, cut off lumps, and smooth ridges. When finishing large slabs, power floats can be used to reduce finishing time.

Floating produces a relatively even (but not smooth) texture that has good slip resistance and is often used as a final finish, especially for exterior slabs. Where a float finish is the desired final finish, it may be necessary to float the surface a second time after it has hardened a little more.

Marks left by hand edgers and groovers are normally removed during floating unless the marks are desired for decorative purposes; in such cases the edger and groover should be used again after final floating.

**Troweling**

Where a smooth, hard, dense surface is desired, floating should be followed by steel troweling (Fig. 11-19). Troweling should not be done on a surface that has not been floated; troweling after only bullfloating or darbying is not an adequate finishing procedure.

It is customary when hand-finishing large slabs to float and immediately trowel an area before moving the

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**Fig. 11-19.** Hand floating (right hand) the surface with a hand float held flat on the concrete surface and moved in a sweeping arc with a slight sawing motion. Troweling (left hand) with blade tilted is performed before moving the kneeboards. (IMG12378)
kneeboards. These operations should be delayed until the concrete has hardened sufficiently so that water and fine material are not brought to the surface. Too long a delay, of course, will result in a surface that is too hard to float and trowel. The tendency, however, is to float and trowel the surface too soon. Premature floating and troweling can cause scaling, crazing, or dusting and a surface with reduced wear resistance.

Spreading dry cement on a wet surface to take up excess water is a bad practice and can cause crazing. Such wet spots should be avoided, if possible, by adjustments in aggregate gradation, mix proportions, and consistency. When wet spots do occur, finishing operations should be delayed until the water either evaporates or is removed with a rubber floor squeegee or by dragging a soft rubber garden hose. If a squeegee or hose is used, care must be taken so that excess cement paste is not removed with the water.

The first troweling may produce the desired surface free of defects. However, surface smoothness, density, and wear resistance can all be improved by additional trowelings. There should be a lapse of time between successive trowelings to permit the concrete to become harder. As the surface stiffens, each successive troweling should be made with smaller trowels, using progressively more tilt and pressure on the trowel blade. The final pass should make a ringing sound as the trowel moves over the hardening surface.

A power trowel is similar to a power float, except that the machine is fitted with smaller, individual steel trowel blades that are adjustable for tilt and pressure on the concrete surface. When the first troweling is done by machine, at least one additional troweling by hand should be done to remove small irregularities. If necessary, tooled edges and joints should be rerun after troweling to maintain uniformity and true lines.

Exterior concrete should not be troweled for several reasons: (1) because it can lead to a loss of entrained air caused by overworking the surface, and (2) troweled surfaces can be slippery when wet. Floating and brooming should be sufficient for outdoor concrete.

**Brooming**

Brooming should be performed before the concrete has thoroughly hardened, but it should be sufficiently hard to retain the scoring impression to produce a slip-resistant surface (Fig. 11-20). Rough scoring can be achieved with a rake, a steel-wire broom, or a stiff, coarse, fiber broom; such coarse-textured brooming usually follows floating. If a finer texture is desired, the concrete should be floated to a smooth surface and then brushed with a soft-bristled broom. Interior concrete could also be troweled before brooming. Best results are obtained with brooms that are specially made for texturing concrete. Slabs are usually broomed transversely to the main direction of traffic.

Highway pavements are textured by “tining” the surface with stiff wires; this improves traction and reduces hydroplaning (Fig. 11-21).

**Curing and Protection**

All newly placed and finished concrete slabs should be cured and protected from drying, from extreme changes in temperature, and from damage by subsequent construction and traffic.
Curing should begin immediately after finishing (Fig. 11-22). Curing is needed to ensure continued hydration of the cement, strength gain of the concrete, and a minimum of early drying shrinkage.

Special precautions are necessary when concrete work continues during periods of adverse weather. In cold weather, arrangements should be made in advance for heating, covering, insulating, or enclosing the concrete. Hot-weather work may require special precautions against rapid evaporation and drying and high temperatures.

Fig. 11-22. An excellent method of wet curing is to completely cover the surface with wet burlap and keep it continuously wet during the curing period. (IMG12374)

PREPARING HARDENED CONCRETE

Bonded Construction Joints in Structural Concrete

A bonded construction joint is needed between two structural concrete placements. When freshly mixed concrete is placed in contact with existing hardened concrete, a high-quality bond and watertight joint are required. Poorly bonded construction joints are usually the result of (1) lack of bond between old and new concrete, or (2) a weak porous layer in the hardened concrete at the joint. The quality of a bonded joint therefore depends on the quality of the hardened concrete and preparation of its surface.

In columns and walls, the concrete near the top surface of a lift is often of inferior quality to the concrete below. This may be due to poor consolidation or use of badly proportioned or high-slump mixtures that cause excessive laitance, bleeding, and segregation. Even in well-proportioned and carefully consolidated mixtures, some aggregate particle settlement and water gain (bleeding) at the top surface is unavoidable; this is partic-ularly true with high rates of placement. Also, the encasing formwork prevents the escape of moisture from the fresh concrete. While formwork provides adequate curing as long as it remains in place, the top surface where there is no encasing formwork may dry out too rapidly; this may result in a weak porous layer unless protection and curing are provided.

Preparing Hardened Concrete

When freshly mixed concrete is placed on recently hardened concrete, certain precautions must be taken to secure a well-bonded, watertight joint. The hardened concrete must be clean, sound, and reasonably rough with some coarse aggregate particles exposed. Any laitance, soft mortar, dirt, wood chips, form oil, or other foreign materials must be removed since they could interfere with proper bonding of the subsequent placement.

The surface of old concrete upon which fresh concrete is to be placed must be thoroughly roughened and cleaned of all dust, surface films, deposits, loose particles, grease, oil, and other foreign material. In most cases it will be necessary to remove the entire surface down to sound concrete. Roughening and cleaning with lightweight chipping hammers, waterblasting, scarifiers, sandblasting (Fig. 11-23), shotblasting and hydrojetting are some of the satisfactory methods for exposing sound concrete. Care must be taken to avoid contamination of the clean surface before a bonding grout and overlay concrete are placed.

Partially set or recently hardened concrete may only require stiff-wire brushing. In some types of construction

Fig. 11-23. Sandblasting can clean any size or shape surface – horizontal, vertical or overhead. Consult local environmental regulations regarding sandblasting. (IMG12252)
such as dams, the surface of each concrete lift is cut with a high-velocity air-water jet to expose clean, sound concrete just before final set. This is usually done 4 to 12 hours after placing. The surface must then be protected and continuously cured until concreting is resumed for the next lift.

For two-course floors, the top surface of the base slab can be roughened just before it sets with a steel or stiff fiber broom. The surface should be level, heavily scored, and free of laitance; then it should be protected until it is thoroughly cleaned just before the grout coat and topping mix are placed. When placing a bonded topping on a floor slab, the base slab should be cleaned of all laitance, dust, debris, grease or other foreign substances by using one of the following methods:

a. Wet- or dry-grit sandblasting
b. High-pressure water blasting
c. Mechanical removal by scabblers, or grinding wheels
d. Power brooming and vacuuming

Hardened concrete may be left dry or be moistened before new concrete is placed on it; however, the surface should not be wet or have any free-standing water. Laboratory studies indicate a slightly better bond is obtained on a dry surface than on a damp surface; however, the increased moisture level in the hardened concrete and in the environment around the concrete reduces water loss from the concrete mixture. This can be very beneficial, especially on hot, dry days.

For making a horizontal construction joint in reinforced concrete wall construction, good results have been obtained by constructing the forms to the level of the joint, overfilling the forms a few centimeters (inches), and then removing the excess concrete just before hardening occurs; the top surface then can be manually roughened with stiff brushes. The procedure is illustrated in Fig. 11-24.

In the case of vertical construction joints cast against a bulkhead, the concrete surface generally is too smooth to permit proper bonding. So, particular care should be given to removal of the smooth surface finish before reerecting the forms for placing freshly mixed concrete against the joint. Stiff-wire brushing may be sufficient if the concrete is less than three days old; otherwise, bushhammering or sandblasting may be needed, followed by washing with clean water to remove all dust and loose particles.

**Bonding New to Previously Hardened Concrete**

Care must be used when making horizontal construction joints in wall sections where freshly-mixed concrete is to be placed on hardened concrete. A good bond can be obtained by placing a rich concrete (higher cement and sand content than normal) in the bottom 150 mm (6 in.) of the new lift and thoroughly vibrating the joint interface. Alternatively, a cement-sand grout can be scrubbed into a clean surface immediately ahead of concreting.

A topping concrete mix for slabs can be bonded to the previously prepared base slab by one of the following procedures:

1. **Portland cement-sand grouting**: A 1 to 1 cement-sand grout having a water-cement ratio of not greater than 0.45, mixed to a creamlike consistency, is scrubbed into the prepared dry or damp (no free water) base slab surface.
2. **Latex.** A latex-bonding agent is added to the cement-sand grout and is spread in accordance with the latex manufacturer’s direction.

3. **Epoxy.** An approved epoxy-bonding agent placed on the base concrete, prepared in accordance with the epoxy manufacturer’s direction.

The bonding procedure should produce tensile bond strength with the base concrete in excess of 1.0 MPa (150 psi).

Grout is placed just a short distance ahead of the overlay or top-course concrete (Fig. 11-25). This method may also be applicable to horizontal joints in walls. The grout should not be allowed to dry out prior to the overlay placement; otherwise, the dry grout may act as a poor surface for bonding. The surface of the base slab should have been prepared by one of the methods discussed previously. Overlays are discussed further under “Patching, Cleaning, and Finishing” later in this chapter.

**MAKING JOINTS IN FLOORS AND WALLS**

The following three types of joints are common in concrete construction: isolation joints, contraction joints, and construction joints.

**Isolation Joints**

Isolation joints (Fig. 11-26) permit both horizontal and vertical differential movements at adjoining parts of a structure. They are used, for example, around the perimeter of a floor on ground, around columns, and around machine foundations to separate the slab from the more rigid parts of the structure.

Isolation-joint material (often called expansion-joint material) can be as thin as 6 mm (1/4 in.) or less, but 13-mm (1/2-in.) material is commonly used. Care should be taken to ensure that all the edges for the full depth of the slab are isolated from adjoining construction; otherwise cracking can occur.

Columns on separate footings are isolated from the floor slab either with a circular or square-shaped isolation joint. The square shape should be rotated to align its corners with control and construction joints.

**Contraction Joints**

Contraction joints (Fig. 11-27) provide for movement in the plane of a slab or wall and induce controlled cracking caused by drying and thermal shrinkage at preselected locations. Contraction joints (also sometimes called contract-
contraction joints) should be constructed to permit transfer of loads perpendicular to the plane of a slab or wall. If no contraction joints are used, or if they are too widely spaced in slabs on ground or in lightly reinforced walls, random cracks may occur; cracks are most likely when drying and thermal shrinkage produce tensile stresses in excess of the concrete’s tensile strength.

Contraction joints in slabs on ground can be made in several ways. One of the most common methods is to saw a continuous straight slot in the top of the slab (Fig. 11-28). This creates a plane of weakness in which a crack will form. Vertical loads are transmitted across a contraction joint by aggregate interlock between the opposite faces of the crack providing the crack is not too wide and the spacing between joints is not too great. Crack widths at saw-cut contraction joints that exceed 0.9 mm (0.035 in.) do not reliably transfer loads. The effectiveness of load transfer by aggregate interlock depends on more than crack width. Other factors include: slab thickness, subgrade support, load magnitude, repetitions of load, and aggregate angularity. Steel dowels (Figs. 11-6 and 11-29b) may be used to increase load transfer at contraction joints when heavy wheel loads are anticipated. Sizes and spacing of dowels, which are placed at the center of the slab depth, are shown in Tarr and Fanny (2008). See ACI Committee 302 and PCA (1982) for further discussions on doweled joints.

Sawing must be coordinated with the setting time of the concrete. It should be started as soon as the concrete has hardened sufficiently to prevent aggregates from being dislodged by the saw (usually within 4 to 12 hours after the concrete hardens); sawing should be completed before drying shrinkage stresses become large enough to produce cracking. The timing depends on factors such as mix proportions, ambient conditions, and type and hardness of aggregates. New dry-cut sawing techniques allow saw cutting to take place shortly after final finishing is completed. Generally, the slab should be cut before the concrete cools, when the concrete sets enough to prevent raveling or tearing while saw cutting, and before drying-shrinkage cracks start to develop.

Contraction joints also can be formed in the fresh concrete with hand groovers or by placing strips of wood, metal, or preformed joint material at the joint locations. The top of the strips should be flush with the concrete surface. Contraction joints, whether sawed, grooved, or preformed, should extend into the slab to a depth of at least one-fourth the slab thickness or a minimum of 25 mm (1 in.) deep. It is recommended that the joint depth not exceed one-third the slab thickness if load transfer from aggregate interlock is important.

Contraction joints in walls are also planes of weakness that permit differential movements in the plane of the wall. The thickness of the wall at a contraction joint should be reduced by 25%, preferably 30%. Under the guidance of the design engineer, in lightly reinforced walls, half of the horizontal steel rebars should be cut at the joint. Care must be taken to cut alternate bars precisely at the joint. At the corners of openings in walls where contraction joints are located, extra diagonal or vertical and horizontal reinforcement should be provided to control cracking. Contraction joints in walls should be spaced not more than about 6 meters (20 ft) apart. In addition, contraction joints should be placed where abrupt changes in wall thickness or height occur, and near corners—if possible, within 3 to 4 meters (10 to 15 ft). Depending on the structure, these joints may need to be caulked to prevent the passage of water through the wall. Instead of caulking, a waterstop (or both) can be used to prevent water from leaking through the crack that occurs in the joint.

The spacing of contraction joints in floors on ground depends on (1) slab thickness, (2) shrinkage potential of the concrete, (3) subgrade friction, (4) environment, and (5) the absence or presence of steel reinforcement. Unless reliable data indicate that more widely spaced joints are feasible, the suggested intervals given in Table 11-2 should be used for well-proportioned concrete with aggregates having normal shrinkage characteristics. Joint spacing should be decreased for concrete suspected of having high shrinkage characteristics. The panels created by contraction joints should be approximately square. Panels with excessive length-to-width ratio (more than 1½ to 1) are likely to crack at an intermediate location. In joint layout design it is also important to remember that contraction (control) joints should only terminate at a free edge or at an isolation joint. Contraction joints should never terminate at another contraction joint as cracking will be induced from the end of the terminated joint into the adjacent panel. This is sometimes referred to as sympathetic cracking. Refer to Fig. 11-31, which illustrates one possible joint layout solution to eliminate the potential for induced sympathetic cracking.
Construction Joints

Construction joints (Fig. 11-29) are stopping places in the process of construction. A true construction joint should bond new concrete to existing concrete and permit no movement. Deformed tiebars are often used in construction joints to restrict movement. Because extra care is needed to make a true construction joint, they are usually designed and built to function as contraction or isolation joints. For example, in a floor on ground the construction joints align with columns and function as contraction joints and therefore are purposely made unbonded. The structural designer of suspended slabs should decide the location of construction joints. Oils, form-release agents, and paints are used as debonding materials. In thick, heavily-loaded floors, unbonded doweled construction joints are commonly used. For thin slabs, the flat-faced butt-type joint will suffice.

On most structures it is desirable to have wall joints that will not detract from appearance. When properly made, wall joints can become inconspicuous or hidden by rustication strips. They thus can become an architectural as well as a functional feature of the structure. However, if rustication strips are used in structures that may be exposed to deicing salts, such as bridge columns and abutments, care should be taken to ensure that the reinforcing steel has the required depth of concrete cover to prevent corrosion.

Horizontal joints in walls should be made straight, exactly horizontal, and should be placed at suitable locations. A straight horizontal construction joint can be made by nailing a 25 mm (1 in.) wood strip to the inside face of the form near the top (see Fig. 11-24). Concrete should then be placed to a level slightly above the bottom of the strip. After the concrete has settled and before it becomes too hard, any laitance that has formed on the top surface

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**Table 11-2 (Metric). Spacing of Contraction Joints in Meters**

<table>
<thead>
<tr>
<th>Slab thickness, mm</th>
<th>Maximum-size aggregate less than 19 mm</th>
<th>Maximum-size aggregate 19 mm and larger</th>
</tr>
</thead>
<tbody>
<tr>
<td>100</td>
<td>2.4</td>
<td>3.0</td>
</tr>
<tr>
<td>125</td>
<td>3.0</td>
<td>3.75</td>
</tr>
<tr>
<td>150</td>
<td>3.75</td>
<td>4.5</td>
</tr>
<tr>
<td>175</td>
<td>4.25</td>
<td>5.25**</td>
</tr>
<tr>
<td>200</td>
<td>5.0**</td>
<td>6.0**</td>
</tr>
<tr>
<td>225</td>
<td>5.5**</td>
<td>6.75**</td>
</tr>
<tr>
<td>250</td>
<td>6.0**</td>
<td>7.5**</td>
</tr>
</tbody>
</table>

* Spacings are appropriate for slumps between 100 mm and 150 mm. If concrete cools at an early age, shorter spacings may be needed to control random cracking. (A temperature difference of only 6°C may be critical.) For slumps less than 100 mm, joint spacing can be increased by 20%.

** When spacings exceed 4.5 m, load transfer by aggregate interlock decreases markedly.

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**Table 11-2 (Inch-Pound Units). Spacing of Contraction Joints in Feet**

<table>
<thead>
<tr>
<th>Slab thickness, in.</th>
<th>Maximum-size aggregate less than ¾ in.</th>
<th>Maximum-size aggregate ¾ in. and larger</th>
</tr>
</thead>
<tbody>
<tr>
<td>4</td>
<td>8</td>
<td>10</td>
</tr>
<tr>
<td>5</td>
<td>10</td>
<td>13</td>
</tr>
<tr>
<td>6</td>
<td>12</td>
<td>15</td>
</tr>
<tr>
<td>7</td>
<td>14</td>
<td>18**</td>
</tr>
<tr>
<td>8</td>
<td>16**</td>
<td>20**</td>
</tr>
<tr>
<td>9</td>
<td>18**</td>
<td>23**</td>
</tr>
<tr>
<td>10</td>
<td>20**</td>
<td>25**</td>
</tr>
</tbody>
</table>

* Spacings are appropriate for slumps between 4 in. and 6 in. If concrete cools at an early age, shorter spacings may be needed to control random cracking. (A temperature difference of only 10°F may be critical.) For slumps less than 4 in., joint spacing can be increased by 20%.

** When spacings exceed 15 ft, load transfer by aggregate interlock decreases markedly.
A typical joint layout for all three joint types—isolation, contraction, and construction—is illustrated in Fig. 11-31. Isolation joints are provided around the perimeter of the floor where it abuts the walls and around all fixed elements that may restrain movement of the slab. This includes columns and machinery bases that penetrate the floor slab. With the slab isolated from other building elements, the remaining task is to locate and correctly space contraction joints to eliminate random cracking. Construction joint locations are coordinated with the floor contractor to accommodate work schedules and crew size. Unbonded construction joints should coincide with the contraction joint pattern and act as contraction joints. Construction joints should be planned to provide long-strips for each placement rather than a checker-board pattern. Contraction joints are then placed to divide the long-strips into relatively square panels, with panel length not exceeding 1.5 times the width. Contraction joints should stop at free edges or isolation joints. For more information on joints, see ACI Committee 302 (2004), PCA (1982), and Tarr and Farny (2008). For joints in walls, see PCA (1982), PCA (1982a), PCA (1984), PCA (1984a), and PCA (1984b).

**FILLING FLOOR JOINTS**

There are three options for treating joints: they can be filled, sealed, or left open. The movement at contraction joints in a floor is generally very small. For some industrial and commercial uses, these joints can be left unfilled or unsealed. Where there are wet conditions, hygienic and dust-control requirements, or considerable traffic by small, hard-wheel vehicles, joint filling is necessary.

The difference between a filler and a sealer is the hardness of the material; fillers are more rigid than sealers and provide support to joint edges. In many places where traffic loading is light, a resilient material such as a polyurethane elastomer sealant is satisfactory. However, heavy-traffic areas require support for joint edges to prevent spalling at saw-cuts; in such cases a good quality, semi-rigid epoxy or polyurea filler with a Shore Hardness of A-80 or D-50 (ASTM D2240) should be used. The mate-
rial should be installed full depth in the saw cut, without a backer rod, and flush with the floor surface.

Isolation joints are intended to accommodate movement; thus a flexible, elastomeric sealant should be used to keep foreign materials out of the joint.

UNJOINTED FLOORS

An unjointed floor, or one with a limited number of joints, can be constructed when joints are unacceptable. Three unjointed floor methods are suggested:

1. A prestressed floor can be built through the use of post-tensioning. With this method, steel strands in ducts are tensioned after the concrete hardens to produce compressive stress in the concrete. This compressive stress will counteract the development of tensile stresses in the concrete and provide a crack-free floor. Large areas, 1000 m² (10,000 ft²) and more, can be constructed in this manner without intermediate joints.

2. Large areas—a single day of slab placement, usually 800 to 1000 m² (8000 to 10,000 ft²)—can be cast without contraction joints when the amount of distributed steel in the floor is about one-half of one percent of the cross-sectional area of the slab. Special effort should be made to reduce subgrade friction in floors without contraction joints. Tarr and Farny (2008) discusses use of distributed steel in floors.

3. Concrete made with expansive cement can be used to offset the amount of drying shrinkage to be anticipated after curing. Contraction joints are not needed when construction joints are used at intervals of 10 to 35 meters (40 to 120 ft). Large areas, up to 2000 m² (20,000 ft²), have been cast in this manner without joints. Steel reinforcement is needed in order to produce compressive stresses during and after the expansion period—this is a form of prestressing.

REMOVING FORMS

It is advantageous to leave forms in place as long as possible to continue the curing period. However, there are times when it is necessary to remove forms as soon as possible. For example, where a rubbed finish is specified, forms must be removed early to permit the first rubbing before the concrete becomes too hard. Furthermore, it is often necessary to remove forms quickly to permit their immediate reuse.

In any case, shoring should not be removed until the concrete is strong enough to satisfactorily carry the stresses from both the dead load of the structure and any imposed construction loads. The concrete should be hard enough so that the surfaces will not be injured in any way when reasonable care is used in removing forms. In general, for concrete temperatures above 10°C (50°F), the side forms of reasonably thick, supported sections can be removed 24 hours after concreting. Beam and floor slab forms and supports (shoring) may be removed between 3 and 21 days, depending on the size of the member and the strength gain of the concrete. For most conditions, it is better to rely on the strength of the concrete as determined by in situ or field-cured specimen testing rather than arbitrarily selecting an age at which forms may be removed. Advice on shoring is provided by ACI Committee 347.

For form removal, the designer should specify the minimum strength requirements for various members. The age-strength relationship should be determined from representative samples of concrete used in the structure and field-cured under job conditions. It should be remembered, however, that strengths are affected by the materials used, temperature, and other conditions. The time required for form removal, therefore, will vary from job to job.

A pinch bar or other metal tool should not be placed against the concrete to wedge forms loose. If it is necessary to wedge between the concrete and the form, only wooden wedges should be used. Stripping should be started some distance away from and move toward a projection. This relieves pressure against projecting corners and reduces the chance of edges breaking off.

Recessed forms require special attention. Wooden wedges should be gradually driven behind the form and the form should be tapped lightly to break it away from the concrete. Forms should not be pulled off rapidly after wedging has been started at one end; this is almost certain to break the edges of the concrete.

PATCHING, CLEANING, AND FINISHING

After forms are removed, all bulges, fins, and small projections can be removed by chipping or tooling. Undesired bolts, nails, ties, or other embedded metal can be removed or cut back to a depth of 13 mm (½ in.) from the concrete surface. When required, the surface can be rubbed or ground to provide a uniform appearance. Any cavities such as tiered holes should be filled unless they are intended for decorative purposes. Honeycombed areas must be repaired and stains removed to present a concrete surface that is uniform in color. All of these operations can be minimized by exercising care in constructing the formwork and placing the concrete. In general, repairs are easier to make and more successful if they are made as soon as practical, preferably as soon as the forms are removed. However, the procedures discussed below apply to both new and old hardened concrete.

Holes, Defects, and Overlays

Patches usually appear darker than the surrounding concrete; therefore, some white cement should be used in mortar or concrete for patching where appearance is
Important. Samples should be applied and cured in an inconspicuous location, perhaps a basement wall, several days in advance of patching operations to determine the most suitable proportions of white and gray cements. Steel troweling should be avoided since this darkens the patch.

Bolt holes, tierod holes, and other cavities that are small in area but relatively deep should be filled with a dry-pack mortar. The mortar should be mixed as stiff as is practical: use 1 part cement, 2 1/2 parts sand passing a 1.25 mm (No. 16) sieve, and just enough water to form a ball when the mortar is squeezed gently in the hand. The cavity should be clean with no oil or loose material and kept damp with water for several hours. A neat-cement paste should be scrubbed onto the void surfaces, but not allowed to dry before the mortar is placed. The mortar should be tamped into place in layers about 13 mm (1/2 in.) thick. Vigorous tamping and adequate curing will ensure good bond and minimum shrinkage of the patch.

Concrete used to fill large patches and thin-bonded overlays should have a low water-cement ratio, often with a cement content equal to or greater than the concrete to be repaired. Cement contents range from 360 to 500 kg per cubic meter (600 to 850 lb per cubic yard) and the water-cement ratio is usually 0.45 or less. The aggregate size should be no more than 1/3 the patch or overlay thickness. A 9.5-mm (3/8-in.) nominal maximum size coarse aggregate is commonly used. The sand proportion can be higher than usual, often equal to the amount of coarse aggregate, depending on the desired properties and application.

Before the patching concrete is applied, the surrounding concrete should be clean and sound (Fig. 11-32). Abrasive methods of cleaning (sandblasting, hydrojetting, waterblasting, scarification, or shotblasting) are usually required. For overlays, a cement-sand grout, a cement-sand-latex grout, or an epoxy bonding agent should be applied to the prepared surface with a brush or broom (see the earlier section “Bonding New to Previously Hardened Concrete”). Typical grout mix proportions are 1 part cement and 1 part fine sand and latex or epoxy admixtures. The grout should be applied immediately before the new concrete is placed. The grout should not be allowed to dry before the freshly mixed concrete is placed; otherwise bond may be impaired. The concrete may be dry or damp when the grout is applied but not wet with free-standing water. The minimum thickness for most patches and overlays is 20 mm (3/4 in.). Some structures, like bridge decks, should have a minimum repair thickness of 40 mm (1 1/2 in.). A superplasticizer is one of many admixtures often added to overlay or repair concrete to reduce the water-cement ratio and to improve workability and ease of consolidation (Kosmatka 1985a).

Honeycombed and other defective concrete should be cut out to expose sound material. If defective concrete is left adjacent to a patch, moisture may get into the voids; in time, weathering action will cause the patch to spall. The edges of the defective area should be cut or chipped straight and at right angles to the surface, or slightly undercut to provide a key at the edge of the patch. No featheredges should be permitted (Fig. 11-33). Based on the size of the patch, either a mortar or a concrete patching mixture should be used.

Shallow patches can be filled with a dry-pack mortar as described earlier. This should be placed in layers not more than 13 mm (1/2 in.) thick, with each layer given a scratch finish to improve bond with the subsequent layer. The final layer can be finished to match the surrounding concrete by floating, rubbing, or tooling, or on formed surfaces by pressing a section of form material against the patch while still plastic.

Deep patches can be filled with concrete held in place by forms. Such patches should be reinforced and doweled to the hardened concrete (Concrete Manual 1981). Large,
shallow vertical or overhead repairs may best be accomplished by shotcreting. Several proprietary low-shrinkage cementitious repair products are also available.

**Curing Patches**

Following patching, good curing is essential (Fig. 11-34). Curing should be started immediately to avoid early drying. Wet burlap, wet sand, plastic sheets, curing paper, tarpaulins, or a combination of these can be used. In locations where it is difficult to hold these materials in place, an application of two coats of membrane-curing compound is often the most convenient method.

**Cleaning Concrete Surfaces**

Concrete surfaces are not always uniform in color when forms are removed; they may have a somewhat blotchy appearance and there may be a slight film of form-release agent in certain areas. There may be mortar stains from leaky forms or there may be rust stains. Flatwork can also become discolored during construction. Where appearance is important, all surfaces should be cleaned after construction has progressed to the stage where there will be no discoloration from subsequent construction activities.

There are three techniques for cleaning concrete surfaces: water, chemical, and mechanical (abrasion). Water dissolves dirt and rinses it from the surface. Chemical cleaners, usually mixed with water, react with dirt to separate it from the surface, and then the dirt and chemicals are rinsed off with clean water. Mechanical methods—sandblasting is most common—remove dirt by abrasion.

Before selecting a cleaning method, it should be tested on an inconspicuous area to be certain it will be helpful and not harmful. If possible, identify the characteristics of the discoloration because some treatments are more effective than others in removing certain materials.

**Water cleaning** methods include low-pressure washes, moderate-to-high-pressure waterblasting, and steam. Low-pressure washing is the simplest, requiring only that water run gently down the concrete surface for a day or two. The softened dirt then is flushed off with a slightly higher pressure rinse. Stubborn areas can be scrubbed with a nonmetallic-bristle brush and rinsed again. High-pressure waterblasting is used effectively by experienced operators. Steam cleaning must be performed by skilled operators using special equipment. Water methods are the least harmful to concrete, but they are not without potential problems. Serious damage may occur if the concrete surface is subjected to freezing temperatures while it is still wet; and water can bring solublesalts to the surface, forming a chalky, white deposit called efflorescence.

**Chemical cleaning** is usually done with water-based mixtures formulated for specific materials such as brick, stone, and concrete. An organic compound called a surfactant (surface-active agent), which acts as a detergent to wet the surface more readily, is included in most chemical cleaners. A small amount of acid or alkali is included to separate the dirt from the surface. For example, hydrochloric (muriatic) acid is commonly used to clean masonry walls and remove efflorescence. There can be problems related to the use of chemical cleaners. Their acid or alkaline properties can lead to reaction between cleaner and concrete as well as mortar, painted surfaces, glass, metals, and other building materials. Since chemical cleaners are used in the form of water-diluted solutions, they too can liberate soluble salts from within the concrete to form efflorescence. Some chemicals can also expose the aggregate in concrete. Chemicals commonly used to clean concrete surfaces and remove discoloration include weak solutions (1% to 10% concentration) of hydrochloric, acetic, or phosphoric acid. Diammonium citrate (20% to 30% water solution) is especially useful in removing discoloration stains and efflorescence on formed and flatwork surfaces. Chemical cleaners should be used by skilled operators taking suitable safety precautions. See Greening (1966) and PCA (1988) for more information.

**Mechanical cleaning** includes sandblasting, shotblasting, scarification, power chipping, and grinding. These methods wear the dirt off the surface rather than separate it from the surface. They, in fact, wear away both the dirt and some of the concrete surface; it is inevitable that there will be some loss of decorative detail, increased surface roughness, and rounding of sharp corners. Abrasive methods may also reveal defects (voids) hidden just beneath the formed surface.

Chemical and mechanical cleaning can each have an abrading effect on the concrete surface that may change the appearance of a surface compared to that of an adjacent uncleaned surface.
Finishing Formed Surfaces

Many off-the-form concrete surfaces require little or no additional treatment when they are carefully constructed with the proper forming materials. These surfaces are divided into two general classes: smooth and textured or patterned. Smooth surfaces are produced with plastic-coated forms, steel forms, fiberglass-reinforced plastic forms, formica forms, or tempered-hardboard forms. Textured or patterned surfaces are achieved with form liners, rough-sawn lumber, special grades and textures of plywood, or by fracturing the projections of a striated surface.

Rough-form finishes require patching of all tieholes and defects, unless tieholes are left open for architectural effect. Otherwise, these surfaces need no further work since texture and finish are imparted by the forms.

For a smooth off-the-form finish, it is important to arrange the smooth-facing forming material and tierods in a symmetrical pattern. Studs and wales that are capable of preventing excessive deflections must support smooth-finish forms that are somewhat lightweight.

A smooth, rubbed finish is produced on a newly hardened concrete surface no later than the day following form removal. The forms are removed and necessary patching completed as soon as possible. The surface is then wet and rubbed with a carborundum brick or other abrasive until a satisfactory uniform color and texture are produced.

A sand-floated finish can also be produced on newly hardened concrete surfaces. No later than 5 to 6 hours following form removal, the surface should be thoroughly wetted and rubbed with a wood float in a circular motion, working fine sand into the surface until the resulting finish is even and uniform in texture and color.

A grout cleandown (sack-rubbed finish) can be used to impart a uniform color and appearance to a smooth surface. After defects have been repaired, the surface should be saturated thoroughly with water and kept wet at least one hour before finishing operations begin. Next a grout of 1 part cement, 1 1⁄2 to 2 parts of fines and passing a 600 μm (No. 30) sieve, and sufficient water for a thick, creamy consistency should be prepared. It should be preshrunk by mixing at least one hour before use and then remixed without the addition of water and applied uniformly by brush, plasterer’s trowel, or rubber float to completely fill all air bubbles and holes.

The surface should be vigorously floated with a wood, sponge rubber, or cork float immediately after applying the grout to fill any small air holes (bugholes) that are left; any remaining excess grout should be scraped off with a sponge-rubber float. If the float pulls grout from the holes, a sawing motion of the tool should correct the difficulty; any grout remaining on the surface should be allowed to stand undisturbed until it loses some of its plasticity but not its damp appearance. Then the surface should be rubbed with clean, dry burlap to remove all excess grout. All air holes should remain filled, but no visible film of grout should remain after the rubbing. Any section being cleaned with grout must be completed in one day, since grout remaining on the surface overnight is difficult to remove.

If possible, work should be done in the shade and preferably during cool, damp weather. During hot or dry weather, the concrete can be kept moist with a fine fog spray.

The completed surface should be moist-cured by keeping the area wet for 36 hours following the clean down. When completely dry, the surface should have a uniform color and texture.

SPECIAL SURFACE FINISHES

Patterns and Textures

A variety of patterns and textures can be used to produce decorative finishes. Patterns can be formed with divider strips or by scoring or stamping the surface just before the concrete hardens. Textures can be produced with little effort and expense with floats, trowels, and brooms; more elaborate textures can be achieved with special techniques (Fig. 11-35). See Kosmatka and Collins (2004).

Exposed-Aggregate Concrete

An exposed-aggregate finish provides a rugged, attractive surface in a wide range of textures and colors. Select aggregates are carefully chosen to avoid deleterious substances; they are usually of uniform size such as 9.5 to 12.5 mm (% to 1⁄2 in.) or larger. They should be washed thoroughly before use to assure satisfactory bond. Flat or elongated aggregate particles should not be used since they are easily dislodged when the aggregate is exposed. Caution should be exercised when using crushed stone; it not only has a greater tendency to stack during the seeding operation (requiring more labor), but it also may be undesirable in some applications (pool decks, for example).

The aggregate should be evenly distributed or seeded in one layer onto the concrete surface immediately after the slab has been bullfloated or dribbled. The particles must be completely embedded in the concrete. This can be done by lightly tapping with a wooden hand float, a darby, or the broad side of a piece of lumber; then, when the concrete can support a finisher on kneeboards, the surface should be hand-floated with a magnesium float or darby until the mortar completely surrounds and slightly covers all the aggregate particles.

Methods of exposing the aggregate usually include washing and brushing, using retarders, and scrubbing. When the concrete has hardened sufficiently, simultaneously brushing and flushing with water should expose the aggregate. In washing and brushing, the surface layer of mortar should be carefully washed away with a light spray of water and brushed until the desired exposure is achieved.
Waterblasting can also be used to texture the surface of hardened concrete, especially where local ordinances prohibit the use of sandblasting for environmental reasons. High-pressure water jets are used on surfaces that have or have not been treated with retarders. In tooling or bushhammering, a layer of hardened concrete is removed and the aggregate is fractured at the surface. The surfaces attained can vary from a light scaling to a deep, bold texture obtained by jackhammering with a single-pointed chisel. Combs and multiple points can be used to produce finishes similar to some finishes used on cut stone.

Grinding and polishing will produce an exposed-aggregate concrete such as terrazzo, which is primarily used indoors. This technique is done in several successive steps using either a stone grinder or diamond-disk grinder. Each successive step uses finer grit than the preceding one. A polishing compound and buffer can then be used for a honed finish.

Regardless of the method employed, it is wise for the contractor to make a preconstruction mock-up (field sample) for each finish to determine the timing and steps involved; in addition, the mock-up is used to obtain aesthetic approval from the architect and owner. For more information see Kosmatka and Collins (2004), PCA (1972), and PCA (1995).
Colored Finishes

Colored concrete finishes for decorative effects in both interior and exterior applications can be achieved by four different methods: (1) the one-course or integral method, (2) the two-course method, (3) the dry-shake method, and (4) stains and paints (discussed below).

Color pigments added to the concrete in the mixer to produce a uniform color is the basis for the one-course method. Both natural and synthetic pigments are satisfactory if they are: (1) insoluble in water, (2) free from soluble salts and acids, (3) fast to sunlight, (4) fast to alkali and weak acids, (5) limited to small amounts of calcium sulphate, and (6) ground fine enough so that 90% passes a 45 micron screen. Use only the minimum amount necessary to produce the desired color and not more than 10% by weight of the cement.

In the two-course method, a base slab is placed and left with a rough texture to bond better to a colored topping layer. As soon as the base slab can support a cement mason’s weight, the topping course can be placed. If the base slab has hardened, prepare a bonding grout for the base slab prior to placing the topping mix. The topping mix is normally 13 mm (1/2 in.) to 25 mm (1 in.) thick, with a ratio of cement to sand of 1:3 or 1:4. The mix is floated and troweled in the prescribed manner. The two-course method is more commonly used because it is more economical than the one-course method.

In the dry-shake method, a prepackaged dry-color material is cast onto the surface of a concrete slab. The dry-shake material is applied after the concrete has been screeded and dabbled or bullfloated, excess moisture has evaporated from the surface, and preliminary floating has been done. Two-thirds of the dry material is shaken evenly by hand over the surface and thoroughly floated into the surface in a manner that evenly distributes the material. Immediately, the rest of the material is cast onto the surface and floated as before. The surface can then be troweled at the same time as a typical slab. For exterior surfaces that will be exposed to freezing and thawing, little or no troweling followed by brooming with a soft bristle concrete broom is usually sufficient.

Stains, Paints, and Clear Coatings

Many types of stains, paints and clear coatings can be applied to concrete surfaces. Among the principal paints used are portland cement base, latex-modified portland cement, and latex (acrylic and polyvinyl acetate) paints (PCA 1992). However, stains and paints are used only when it is necessary to color existing concrete. It is difficult to obtain a uniform color with dyes or stains; therefore, the manufacturer’s directions should be closely followed.

Portland cement based paints can be used on either interior or exterior exposures. The surface of the concrete should be damp at the time of application and each coat should be dampened as soon as possible without disturbing the paint. Damp curing of conventional portland cement paint is essential. On open-textured surfaces, such as concrete masonry, the paint should be applied with stiff-bristle brushes (scrub brushes). Paint should be worked well into the surface. For concrete with a smooth or sandy finish, whitewash or Dutch-type calcimine brushes are best.

The latex materials used in latex-modified portland cement paints retard evaporation, thereby retaining the necessary water for hydration of the portland cement. When using latex-modified paints, moist curing is not required.

Most latex paints are resistant to alkali and can be applied to new concrete after 10 days of good drying weather. The preferred method of application is by long fiber, tapered nylon brushes 100 to 150 mm (4 to 6 in.) wide; however, roller or spray methods can also be used. The paints may be applied to damp, but not wet surfaces. If the surface is moderately porous, or if extremely dry conditions prevail, prewetting the surface is advisable.

Clear coatings are frequently used on concrete surfaces to (1) prevent soiling or discoloration of the concrete by air pollution, (2) to facilitate cleaning the surface if it does become dirty, (3) to brighten the color of the aggregates, and (4) to render the surface water-repellent and thus prevent color change due to rain and water absorption. The better coatings often consist of methyl methacrylate forms of acrylic resin, as indicated by a laboratory evaluation of commercial clear coatings (Litvin 1968). The methyl methacrylate coatings should have a higher viscosity and solids content when used on smooth concrete, since the original appearance of smooth concrete is more difficult to maintain than the original appearance of exposed-aggregate concrete.

Other materials, such as silane and siloxane penetrating sealers, are commonly used as water repellents for many exterior concrete applications.

PRECAUTIONS

Protect Your Head and Eyes. Construction equipment and tools represent constant potential hazards to busy construction personnel. That's why hard hats are required on construction projects. It is therefore recommended that some sort of head protection, such as a hard hat or safety hat, be worn when working any construction job, large or small.

Proper eye protection is essential when working with cement or concrete. Eyes are particularly vulnerable to blowing dust, splattering concrete, and other foreign objects. On some jobs it may be advisable to wear full-cover goggles or safety glasses with side shields. Actions that cause dust to become airborne should be avoided. Local or general ventilation can control exposures below applicable exposure limits; respirators may be used in poorly ventilated areas, where exposure limits are exceeded, or when dust causes discomfort or irritation.
Protect Your Back. All materials used to make concrete—Portland cement, sand, coarse aggregate, and water—can be quite heavy, even in small quantities. When lifting heavy materials, the back should be straight, legs bent, and the weight between the legs as close to the body as possible. Mechanical equipment should be used to place concrete as close as possible to its final position. After the concrete is deposited in the desired area by chute, pump, or wheelbarrow, it should be pushed—not lifted—into final position with a shovel; a short-handled, square-end shovel is an effective tool for spreading concrete, but special concrete rakes or come-alongs also can be used. Excessive horizontal movement of the concrete should be avoided; it not only requires extra effort, but may also lead to segregation of the concrete ingredients.

Protect Your Skin. When working with fresh concrete, care should be taken to avoid skin irritation or chemical burns (see warning statement in the box). Prolonged contact between fresh concrete and skin surfaces, eyes, and clothing may result in burns that are quite severe, including third-degree burns. Eyes and skin that come in contact with fresh concrete should be flushed thoroughly with clean water. If irritation persists, consult a physician. For deep burns or large affected skin areas, seek medical attention immediately.

The A-B-Cs of fresh concrete’s effect on skin are:

- **Abrasiven** Sand contained in fresh concrete is abrasive to bare skin.
- **Basic & Caustic** Portland cement is alkaline in nature, so wet concrete and other cement mixtures are strongly basic (pH of 12 to 13). Strong bases—like strong acids—are harmful, or caustic to skin.
- **Drying** Portland cement is hygroscopic—it absorbs water. In fact, portland cement needs water to harden. It will draw water away from any material it contacts—including skin.

Clothing worn as protection from fresh concrete should not be allowed to become saturated with moisture from fresh concrete because saturated clothing can transmit alkaline or hygroscopic effects to the skin. Clothing that becomes saturated from contact with fresh concrete should be rinsed out promptly with clear water to prevent continued contact with skin surfaces. Waterproof gloves, a long-sleeved shirt, and long pants should be worn. If you must stand in fresh concrete while it is being placed, screeded, or floated, wear rubber boots high enough to prevent concrete from getting into them (PCA 1998).

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Curing is the maintenance of a satisfactory moisture content and temperature in concrete for a period of time immediately following placing and finishing so that the desired properties may develop (Fig. 12-1). The need for adequate curing of concrete cannot be overemphasized. Curing has a strong influence on the properties of hardened concrete; proper curing will increase durability, strength, watertightness, abrasion resistance, volume stability, and resistance to freezing and thawing and deicers. Exposed slab surfaces are especially sensitive to curing as strength development and freeze-thaw resistance of the top surface of a slab can be reduced significantly when curing is defective.

When portland cement is mixed with water, a chemical reaction called hydration takes place. The extent to which this reaction is completed influences the strength and durability of the concrete. Freshly mixed concrete normally contains more water than is required for hydration of the cement; however, excessive loss of water by evaporation can delay or prevent adequate hydration. The surface is particularly susceptible to insufficient hydration because it dries first. If temperatures are favorable, hydration is relatively rapid the first few days after concrete is placed; however, it is important for water to be retained in the concrete during this period, that is, for evaporation to be prevented or substantially reduced.

With proper curing, concrete becomes stronger, more impermeable, and more resistant to stress, abrasion, and freezing and thawing. The improvement is rapid at early ages but continues more slowly thereafter for an indefinite period. Fig. 12-2 shows the strength gain of concrete with age for different moist curing periods and Fig. 12-3 shows the relative strength gain of concrete cured at different temperatures.

Fig. 12-1. Curing should begin as soon as the concrete stiffens enough to prevent marring or erosion of the surface. Burlap sprayed with water is an effective method for moist curing. (IMG12363)

Fig. 12-2. Effect of moist curing time on strength gain of concrete (Gonnerman and Shuman 1928).
The most effective method for curing concrete depends on the materials used, method of construction, and the intended use of the hardened concrete. For most jobs, curing generally involves applying curing compounds, or covering the freshly placed and finished concrete with impermeable sheets or wet burlap. In some cases, such as in hot and cold weather, special care using other precautions is needed.

Concrete mixtures with high cement contents and low water-cement ratios (less than 0.40) may require special curing needs. As cement hydrates (chemically combining with water) the internal relative humidity decreases causing the paste to self-desiccate (dry out) if no external water is provided. The paste can self-desiccate to a level where hydration stops. This may influence desired concrete properties, especially if the internal relative humidity drops below 80% within the first seven days. In view of this, membrane-forming curing compounds may not retain enough water in the concrete. Therefore, fogging and wet curing become necessary to maximize hydration (Copeland and Bragg 1955). Fogging during and after placing and finishing also helps minimize plastic cracking in concretes with very low water-cement ratios (especially around 0.30 or less).

When moist curing is interrupted, the development of strength continues for a short period and then stops after the concrete’s internal relative humidity drops to about 80%. However, if moist curing is resumed, strength development will be reactivated, but the original potential strength may not be achieved. Although it can be done in a laboratory, it is difficult to resaturate concrete in the field. Thus, it is best to moist-cure the concrete continuously from the time it is placed and finished until it has gained sufficient strength, impermeability, and durability.

Loss of water will also cause the concrete to shrink, thus creating tensile stresses within the concrete. If these stresses develop before the concrete has attained adequate tensile strength, surface cracking can result. All exposed surfaces, including exposed edges and joints, must be protected against moisture evaporation.

Hydration proceeds at a much slower rate when the concrete temperature is low. Temperatures below 10°C (50°F) are unfavorable for the development of early strength; below 4°C (40°F) the development of early strength is greatly retarded; and at or below freezing temperatures, down to -10°C (14°F), little or no strength develops.

In recent years, a maturity concept has been introduced to evaluate the development of strength when there is variation in the curing temperature of the concrete. Maturity is the product of the age of the concrete and its average curing temperature above a certain base temperature. Refer to Chapter 14 for more information on the maturity concept. It follows that concrete should be protected so that its temperature remains favorable for hydration and moisture is not lost during the early hardening period.

**CURING METHODS AND MATERIALS**

Concrete can be kept moist (and in some cases at a favorable temperature) by three curing methods:

1. **Methods that maintain the presence of mixing water in the concrete during the early hardening period.** These include ponding or immersion, spraying or fogging, and saturated wet coverings. These methods afford some cooling through evaporation, which is beneficial in hot weather.

2. **Methods that reduce the loss of mixing water from the surface of the concrete.** This can be done by covering the concrete with impervious paper or plastic sheets, or by applying membrane-forming curing compounds.
3. Methods that accelerate strength gain by supplying heat and additional moisture to the concrete. This is usually accomplished with live steam, heating coils, or electrically heated forms or pads.

The method or combination of methods chosen depends on factors such as availability of curing materials, size, shape, and age of concrete, production facilities (in place or in a plant), esthetic appearance, and economics. As a result, curing often involves a series of procedures used at a particular time as the concrete ages. For example, fog spraying or plastic covered wet burlap can precede application of a curing compound. The timing of each procedure depends on the degree of hardening of the concrete needed to prevent the particular procedure from damaging the concrete surface (ACI 308 2001).

**Ponding and Immersion**

On flat surfaces, such as pavements and floors, concrete can be cured by ponding. Earth or sand dikes around the perimeter of the concrete surface can retain a pond of water. Ponding is an ideal method for preventing loss of moisture from the concrete; it is also effective for maintaining a uniform temperature in the concrete. The curing water should not be more than about 11°C (20°F) cooler than the concrete to prevent thermal stresses that could result in cracking. Since ponding requires considerable labor and supervision, the method is generally used only for small jobs.

The most thorough method of curing with water consists of total immersion of the finished concrete element. This method is commonly used in the laboratory for curing concrete test specimens. Where appearance of the concrete is important, the water used for curing by ponding or immersion must be free of substances that will stain or discolor the concrete. The material used for dikes may also discolor the concrete.

**Fogging and Sprinkling**

Fogging (Fig. 12-4) and sprinkling with water are excellent methods of curing when the ambient temperature is well above freezing and the humidity is low. A fine fog mist is frequently applied through a system of nozzles or sprayers to raise the relative humidity of the air over flatwork, thus slowing evaporation from the surface. Fogging is applied to minimize plastic shrinkage cracking until finishing operations are complete. Once the concrete has set sufficiently to prevent water erosion, ordinary lawn sprinklers are effective if good coverage is provided and water runoff is of no concern. Soaker hoses are useful on surfaces that are vertical or nearly so.

The cost of sprinkling may be a disadvantage. The method requires an ample water supply and careful supervision. If sprinkling is done at intervals, the concrete must be prevented from drying between applications of water by using burlap or similar materials; otherwise alternate cycles of wetting and drying can cause surface crazing or cracking.

**Wet Coverings**

Fabric coverings saturated with water, such as burlap, cotton mats, rugs, or other moisture-retaining fabrics, are commonly used for curing (Fig. 12-5). Treated burlaps that reflect light and are resistant to rot and fire are available. The requirements for burlap are described in the Specification for Burlap Cloths Made from Jute or Kenaf (AASHTO M 182), and those for white burlap-polyethylene sheeting are described in ASTM C171 (AASHTO M 171).
Burlap must be free of any substance that is harmful to concrete or causes discoloration. New burlap should be thoroughly rinsed in water to remove soluble substances and to make the burlap more absorbent.

Wet, moisture-retaining fabric coverings should be placed as soon as the concrete has hardened sufficiently to prevent surface damage. During the waiting period other curing methods are used, such as fogging or the use of membrane forming finishing aids. Care should be taken to cover the entire surface with wet fabric, including the edges of slabs. The coverings should be kept continuously moist so that a film of water remains on the concrete surface throughout the curing period. Use of polyethylene film over wet burlap is a good practice; it will eliminate the need for continuous watering of the covering. Periodically rewetting the fabric under the plastic before it dries out should be sufficient. Alternate cycles of wetting and drying during the early curing period may cause crazing of the surface.

Wet coverings of earth, sand, or sawdust are effective for curing and are often useful on small jobs. Sawdust from most woods is acceptable, but oak and other woods that contain tannic acid should not be used since deterioration of the concrete may occur. A layer about 50 mm (2 in.) thick should be evenly distributed over the previously moistened surface of the concrete and kept continuously wet.

Wet hay or straw can be used to cure flat surfaces. If used, it should be placed in a layer at least 150 mm (6 in.) thick and held down with wire screen, burlap, or tarpaulins to prevent its being blown off by wind.

A major disadvantage of moist earth, sand, sawdust, hay, or straw coverings is the possibility of discoloring the concrete.

### Impervious Paper

Impervious paper for curing concrete consists of two sheets of kraft paper cemented together by a bituminous adhesive with fiber reinforcement. Such paper, conforming to ASTM C171 (AASHTO M 171), is an efficient means of curing horizontal surfaces and structural concrete of relatively simple shapes. An important advantage of this method is that periodic additions of water are not required. Curing with impervious paper enhances the hydration of cement by preventing loss of moisture from the concrete (Fig. 12-6).

As soon as the concrete has hardened sufficiently to prevent surface damage, it should be thoroughly wetted and the widest paper available applied. Edges of adjacent sheets should be overlapped about 150 mm (6 in.) and tightly sealed with sand, wood planks, pressure-sensitive tape, mastic, or glue. The sheets must be weighted to maintain close contact with the concrete surface during the entire curing period.

Impervious paper can be reused if it effectively retains moisture. Tears and holes can easily be repaired with curing-paper patches. When the condition of the paper is questionable, additional use can be obtained by using it in double thickness.

In addition to curing, impervious paper provides some protection to the concrete against damage from subsequent construction activity as well as protection from the direct sun. It should be light in color and nonstaining to the concrete. Paper with a white upper surface is preferable for curing exterior concrete during hot weather.

### Plastic Sheets

Plastic sheets materials, such as polyethylene film, can be used to cure concrete (Fig. 12-7). Polyethylene film is a lightweight, effective moisture retarder and is easily applied to complex as well as simple shapes. Its application is the same as described for impervious paper.

Curing with polyethylene film (or impervious paper) can cause patchy discoloration, especially if the concrete contains calcium chloride and has been finished by hard-steel troweling. This discoloration is more pronounced when the film becomes wrinkled, but it is difficult and time consuming on a large project to place sheet materials without wrinkles. Flooding the surface under the covering may prevent discoloration, but other means of curing should be used when uniform color is important.

Polyethylene film should conform to ASTM C171 (AASHTO M 171), which specifies a 0.10-mm (4-mil) thickness for curing concrete, but lists only clear and white opaque film. However, black film is available and is satisfactory under some conditions. White film should be used for curing exterior concrete during hot weather to reflect the sun’s rays. Black film can be used during cool weather
Membrane-Forming Curing Compounds

Liquid membrane-forming compounds consisting of waxes, resins, chlorinated rubber, and other materials can be used to retard or reduce evaporation of moisture from concrete. They are the most practical and most widely used method for curing not only freshly placed concrete but also for extending curing of concrete after removal of forms or after initial moist curing. However, the most effective methods of curing concrete are wet coverings or water spraying that keeps the concrete continually damp. Curing compounds should be able to maintain the relative humidity of the concrete surface above 80% for seven days to sustain cement hydration.

Membrane-forming curing compounds are of two general types: clear, or translucent; and white pigmented. Clear or translucent compounds may contain a fugitive dye that makes it easier to check visually for complete coverage of the concrete surface when the compound is applied. The dye fades away soon after application. On hot, sunny days, use of white-pigmented compounds are recommended; they reduce solar-heat gain, thus reducing the concrete temperature. Pigmented compounds should be kept agitated in the container to prevent pigment from settling out.

Curing compounds should be applied by hand-operated or power-driven spray equipment immediately after final finishing of the concrete (Fig. 12-8). The concrete surface should be damp when the coating is applied. On dry, windy days, or during periods when adverse weather conditions could result in plastic shrinkage cracking, application of a curing compound immediately after final finishing and before all free water on the surface has evaporated will help prevent the formation of cracks. Power-driven spray equipment is recommended for uniform application of curing compounds on large paving projects. Spray nozzles and windshields on such equipment should be arranged to prevent wind-blown loss of curing compound.

Normally only one smooth, even coat is applied at a typical rate of 3 to 4 m² per liter (150 to 200 sq ft per gallon); but products may vary, so manufacturer’s recommended application rates should be followed. If two coats are necessary to ensure complete coverage, for effective protection the second coat should be applied at right angles to the first. Complete coverage of the surface must be attained because even small pinholes in the membrane will increase the evaporation of moisture from the concrete.

Curing compounds might prevent bonding between hardened concrete and a freshly placed concrete overlay. And, most curing compounds are not compatible with adhesives used with floor covering materials. Consequently, they should either be tested for compatibility, or not used when bonding of overlying materials is necessary. For example, a curing compound should not be applied to the
base slab of a two-course floor. Similarly, some curing compounds may affect the adhesion of paint to concrete floors. Curing compound manufacturers should be consulted to determine if their product is suitable for the intended application.

Curing compounds should be uniform and easy to maintain in a thoroughly mixed solution. They should not sag, run off peaks, or collect in grooves. They should form a tough film to withstand early construction traffic without damage, be nontoxic, and have good moisture-retention properties.

Caution is necessary when using curing compounds containing solvents of high volatility in confined spaces or near sensitive occupied spaces such as hospitals because evaporating volatiles may cause respiratory problems. Applicable local environmental laws concerning volatile organic compound (VOC) emissions should be followed.

Curing compounds should conform to ASTM C309 (AASHTO M 148). A method for determining the efficiency of curing compounds, waterproof paper, and plastic sheets is described in ASTM C156 (AASHTO T 155). ASTM C1151, discontinued in 2000, also evaluates the effectiveness of curing compounds. Curing compounds with sealing properties are specified under ASTM C1315.

Internal Moist Curing

Internal moist curing refers to methods of providing moisture from within the concrete as opposed to outside the concrete. This water should not affect the initial water to cement ratio of the fresh concrete. Lightweight (low-density) fine aggregate or absorbent polymer particles with an ability to retain a significant amount of water may provide additional moisture for concretes prone to self-desiccation. When more complete hydration is needed for concretes with low water to cement ratios (around 0.30 or less), 60 kg/m³ to 180 kg/m³ (100 lb/yd³ to 300 lb/yd³) of saturated lightweight fine aggregate can provide additional moisture to extend hydration, resulting in increased strength and durability. All of the fine aggregate in a mixture can be replaced with saturated lightweight fine aggregate to maximize internal moist curing. Internal moist curing must be accompanied by external curing methods.

Forms Left in Place

Forms provide satisfactory protection against loss of moisture if the top exposed concrete surfaces are kept wet. A soaker hose is excellent for this. The forms should be left on the concrete as long as practical.

Wood forms left in place should be kept moist by sprinkling, especially during hot, dry weather. If this cannot be done, they should be removed as soon as practical and another curing method started without delay. Color variations may occur from formwork and uneven water curing of walls.

Steam Curing

Steam curing is advantageous where early strength gain in concrete is important or where additional heat is required to accomplish hydration, as in cold weather.

Two methods of steam curing are used: live steam at atmospheric pressure (for enclosed cast-in-place structures and large precast concrete units) and high-pressure steam in autoclaves (for small manufactured units). Only live steam at atmospheric pressure will be discussed here.

A typical steam-curing cycle consists of (1) an initial delay prior to steaming, (2) a period for increasing the temperature, (3) a period for holding the maximum temperature constant, and (4) a period for decreasing the temperature. A typical atmospheric steam-curing cycle is shown in Fig. 12-9.

Steam curing at atmospheric pressure is generally done in an enclosure to minimize moisture and heat losses. Tarpaulins are frequently used to form the enclosure. Application of steam to the enclosure should be delayed until initial set occurs or delayed at least 3 hours after final placement of concrete to allow for some hardening of the concrete. However, a 3- to 5-hour delay period prior to steaming will achieve maximum early strength, as shown in Fig. 12-10.

Steam temperature in the enclosure should be kept at about 60°C (140°F) until the desired concrete strength has developed. Strength will not increase significantly if the maximum steam temperature is raised from 60°C to 70°C (140°F to 160°F). Steam-curing temperatures above 70°C (160°F) should be avoided; they are uneconomical and may result in damage. It is recommended that the internal

![Fig. 12-9. A typical atmospheric steam-curing cycle.](image-url)
temperature of concrete not exceed 70°C (160°F) to avoid heat induced delayed expansion and undue reduction in ultimate strength. Use of concrete temperatures above 70°C (160°F) should be demonstrated to be safe by test or historic field data.

Concrete temperatures are commonly monitored at the exposed ends of the concrete element. Monitoring air temperatures alone is not sufficient because the heat of hydration may cause the internal temperature of the concrete to exceed 70°C (160°F). Besides early strength gain, there are other advantages of curing concrete at temperatures of around 60°C (140°F); for example, there is reduced drying shrinkage and creep as compared to concrete cured at 23°C (73°F) for 28 days (Klieger 1960 and Tepponen and Eriksson 1987).

Excessive rates of heating and cooling should be avoided to prevent damaging volume changes. Temperatures in the enclosure surrounding the concrete should not be increased or decreased more than 22°C to 33°C (40°F to 60°F) per hour depending on the size and shape of the concrete element.

The curing temperature in the enclosure should be held until the concrete has reached the desired strength. The time required will depend on the concrete mixture and steam temperature in the enclosure (ACI Committee 517).

**Insulating Blankets or Covers**

Layers of dry, porous material such as straw or hay can be used to provide insulation against freezing of concrete when temperatures fall below 0°C (32°F).

Formwork can be economically insulated with commercial blanket or batt insulation that has a tough moistureproof covering. Suitable insulating blankets are manufactured of fiberglass, sponge rubber, cellulose fibers, mineral wool, vinyl foam, and open-cell polyurethane foam. When insulated formwork is used, care should be taken to ensure that concrete temperatures do not become excessive.

Framed enclosures of canvas tarpaulins, reinforced polyethylene film, or other materials can be placed around the structure and heated by space heaters or steam. Portable hydronic heaters are used to thaw subgrades as well as heat concrete without the use of an enclosure.

Curing concrete in cold weather should follow the recommendations in Chapter 14 and ACI 306 (1997), *Cold-Weather Concreting.* Recommendations for curing concrete in hot weather can be found in Chapter 13 and ACI 305, *Hot-Weather Concreting.*

**Electrical, Oil, Microwave, and Infrared Curing**

Electrical, hot oil, microwave and infrared curing methods have been available for accelerated and normal curing of concrete for many years. Electrical curing methods include a variety of techniques: (1) use of the concrete itself as the electrical conductor, (2) use of reinforcing steel as the heating element, (3) use of a special wire as the heating element, (4) electric blankets, and (5) the use of electrically heated steel forms (presently the most popular method). Electrical heating is especially useful in cold-weather concreting. Hot oil may be circulated through steel forms to heat the concrete. Infrared rays and microwave have had limited use in accelerated curing of concrete. Concrete that is cured by infrared methods is usually under a covering or enclosed in steel forms. Electrical, oil, and infrared curing methods are used primarily in the precast concrete industry.

**CURING PERIOD AND TEMPERATURE**

The period of time that concrete should be protected from freezing, abnormally high temperatures, and against loss of moisture depends upon a number of factors: the type of cementing materials used; mixture proportions; required strength, size and shape of the concrete member; ambient weather; and future exposure conditions. The curing period may be 3 weeks or longer for lean concrete mixtures used in massive structures such as dams; conversely, it may be only a few days for rich mixes, especially if Type III or HE cement is used. Steam-curing periods are nor-
mally much shorter, ranging from a few hours to 3 days; but generally 24-hour cycles are used. Since all the desirable properties of concrete are improved by curing, the curing period should be as long as necessary.

For concrete slabs on ground (floors, pavements, canal linings, parking lots, driveways, sidewalks) and for structural concrete (cast-in-place walls, columns, slabs, beams, small footings, piers, retaining walls, bridge decks), the length of the curing period for ambient temperatures above 5°C (40°F) should be a minimum of 7 days; additional time may be needed to attain 70% of the specified compressive or flexural strength. When the daily mean ambient temperature is 5°C (40°F) or lower, ACI Committee 306 recommendations for curing should be followed to prevent damage by freezing.

A higher curing temperature provides earlier strength gain in concrete than a lower temperature but it may decrease 28-day strength as shown in Fig. 12-11. If strength tests are made to establish the time when curing can cease or forms can be removed, representative concrete test cylinders or beams should be fabricated in the field, kept adjacent to the structure or pavement they represent, and cured by the same methods. Equipment is available that can monitor internal concrete temperatures and match that temperature in the concrete cylinder curing box; this is the most accurate means of representing in-place concrete strengths. Cores, cast-in-place removable cylinders, and nondestructive testing methods may also be used to determine the strength of a concrete member.

Since the rate of hydration is influenced by cement type and the presence of supplementary cementing materials, the curing period should be prolonged for concretes made with cementing materials possessing slow-strength-gain characteristics. For mass concrete (large piers, locks, abutments, dams, heavy footings, and massive columns and transfer girders) in which no pozzolan is used as part of the cementitious material, curing of unreinforced sections should continue for at least 2 weeks. If the mass concrete contains a pozzolan, minimum curing time for unreinforced sections should be extended to 3 weeks. Heavily-reinforced mass concrete sections should be cured for a minimum of 7 days.

During cold weather, additional heat is often required to maintain favorable curing temperatures of 10°C to 20°C (50°F to 70°F). Vented gas or oil-fired heaters, heating coils, portable hydronic heaters, or live steam can be used to supply the required heat. In all cases, care must be taken to avoid loss of moisture from the concrete. Exposure of fresh concrete to heater or engine exhaust gases must be avoided as this can result in surface deterioration and dusting (rapid carbonation).

High-early-strength concrete can be used in cold weather to speed-up setting time and strength development. This can reduce the curing period, but a minimum temperature of 10°C (50°F) must be maintained.

For adequate deicer scale resistance of concrete, the minimum curing period generally corresponds to the time required to develop the design strength of the concrete at the surface. A period of air-drying after curing will enhance resistance to scaling. This drying period should be at least 1 month of relatively dry weather before the application of deicing salts.

**SEALING COMPOUNDS**

Sealing compounds (sealers) are liquids applied to the surface of hardened concrete to reduce the penetration of liquids or gases such as water, deicing solutions, and carbon dioxide that cause freeze-thaw damage, corrosion of reinforcing steel, and acid attack. In addition, sealing compounds used on interior floor slabs reduce dusting and the absorption of spills while making the surface easier to clean.

Sealing compounds differ in purpose from curing compounds; they should not be confused as being the same. The primary purpose of a curing compound is to retard the loss of water from newly placed concrete and it is applied immediately after finishing. Surface sealing compounds on the other hand retard the penetration of harmful substances into hardened concrete and are typi-
cally not applied until the concrete is 28 days old. Surface sealers are generally classified as either film-forming or penetrating.

Sealing exterior concrete is an optional procedure generally performed to help protect concrete from freeze-thaw damage and chloride penetration from deicers. Curing is not optional when using a sealer; curing is necessary to produce properties needed for concrete to perform adequately for its intended purpose. Satisfactory performance of exterior concrete still primarily depends on an adequate air-void system, sufficient strength, and the use of proper placing, finishing and curing techniques. However, not all concrete placed meets those criteria; surface sealers can help improve the durability of these concretes.

Film-forming sealing compounds remain mostly on the surface with only a slight amount of the material penetrating the concrete. The relatively large molecular structure of these compounds limits their ability to penetrate the surface. Thinning them with solvents will not improve their penetrating capability. These materials not only reduce the penetration of water, they also protect against mild chemicals; furthermore, they prevent the absorption of grease and oil as well as reduce dusting under pedestrian traffic.

Surface sealers consist of acrylic resins, chlorinated rubber, urethanes, epoxies, and alpha methyl styrene. The effectiveness of film-forming sealers depends on the continuity of the layer formed. Abrasive grit and heavy traffic can damage the layer requiring the reapplication of the material. Consult manufacturers’ application recommendations because some of these materials are intended for interior use only and may yellow and deteriorate under exposure to ultraviolet light.

The penetrating sealer that has been used most extensively for many years is a mixture of 50 percent boiled linseed oil and 50 percent mineral spirits (AASHTO M 233). Although this mixture is an effective sealer, it has two main disadvantages: it darkens the concrete, and periodic reapplication is necessary for long-term protection.

A new generation of water-repellent penetrating sealers have a very small molecular size that allows penetration and saturation of the concrete as deep as 3 mm (¼ in.). The two most common are silane and siloxane, compounds which are derived from the silicone family. These sealers allow the concrete to breath, thus preventing a buildup of vapor pressure between the concrete and sealer that can occur with some film-forming materials. Because the sealer is embedded within the concrete, making it more durable to abrasive forces or ultraviolet deterioration, it can provide longer lasting protection than film-forming sealers. However, periodic retreatment is recommended. In northern states and coastal areas silanes and siloxanes are popular for protecting bridge decks and other exterior structures from corrosion of reinforcing steel caused by chloride infiltration from deicing chemicals or sea spray (Fig 12-12).

Application of any sealer should only be done on concrete that is clean and allowed to dry for at least 24 hours at temperatures above 16°C (60°F). At least 28 days should be allowed to elapse before applying sealers to new concrete. Penetrating sealers cannot fill surface voids if they are filled with water. Some surface preparation may be necessary if the concrete is old and dirty. Concrete placed in the late fall should not be sealed until spring because the sealer may cause the concrete to retain water that may exacerbate freeze-thaw damage.

The precautions outlined earlier regarding volatile solvents in curing compounds also apply to sealing compounds. The effectiveness of water-based surface sealers is still being determined. The scale resistance provided by concrete sealers should be evaluated based on criteria established in ASTM C 672. For more information on surface sealing compounds, see AASHTO M 224, ACI Committee 330 and ACI Committee 362.

Fig 12-12. Penetrating sealers help protect reinforcing steel in bridge decks from corrosion due to chloride infiltration without reducing surface friction. (IMG12358)
REFERENCES

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CHAPTER 13
Hot-Weather Concreting

Weather conditions at a jobsite—hot or cold, windy or calm, dry or humid—may be vastly different from the optimum conditions assumed at the time a concrete mix is specified, designed, or selected, or from laboratory conditions in which concrete specimens are stored and tested. Hot weather conditions adversely influence concrete quality primarily by accelerating the rate of moisture loss and rate of cement hydration that occur at higher temperatures. Detrimental hot weather conditions include:

- high ambient temperature
- high concrete temperature
- low relative humidity
- high wind speed
- solar radiation

Hot weather conditions can create difficulties in fresh concrete, such as:

- increased water demand
- accelerated slump loss leading to the addition of water on the jobsite
- increased rate of setting resulting in placing and finishing difficulties
- increased tendency for plastic cracking
- critical need for prompt early curing
- difficulties in controlling entrained air
- increased concrete temperature resulting in long-term strength loss
- increased potential for thermal cracking

Adding water to the concrete at the jobsite can adversely affect properties and serviceability of the hardened concrete, resulting in:

- decreased strength from higher water to cement ratio
- decreased durability due to cracking
- increased permeability
- nonuniform surface appearance
- increased tendency for drying shrinkage
- reduced abrasion resistance from tendency to sprinkle water during finishing

Only by taking precautions to alleviate these difficulties in anticipation of hot-weather conditions can concrete work proceed smoothly. For more information on the above topics, see ACI Committee 305 (1999).

WHEN TO TAKE PRECAUTIONS

During hot weather the most favorable temperature for achieving high quality freshly mixed concrete is usually lower than can be obtained without artificial cooling. A concrete temperature of 10°C to 15°C (50°F to 60°F) is desirable to maximize beneficial mix properties, but such temperature are not always practical. Many specifications require only that concrete when placed should have a temperature of less than 29°C to 32°C (85°F to 90°F). The ASTM C94 (AASHTO M 157) specifications for ready

Fig. 13-1. Liquid nitrogen added directly into a truck mixer at the ready mix plant is an effective method of reducing concrete temperature for mass concrete placements or during hot-weather concreting. (IMG12357)
mixed concrete notes in some situations difficulty may be encountered when concrete temperatures approach 32°C (90°F). However, this specification does not mandate a maximum concrete temperature unless heated aggregates or heated water are used.

Precautions should be planned in advance to counter the effects of a high concrete temperature when the concrete placed is somewhere between 25°C and 35°C (77°F and 95°F). Last-minute attempts to prevent hot-weather damage are rarely performed soon enough. If acceptable field data is not available, the maximum temperature limit should be established for conditions at the jobsite; this should be based on trial-batch tests at the temperature and for the typical concrete section thickness anticipated, rather than on ideal temperatures of 20°C to 30°C (68°F to 86°F) cited in ASTM C192 (AASHTO T 126). If possible, large batches should be made to measure mix properties at time intervals to establish the relationship for the property of interest as a function of time at various batch temperatures of interest. This process will establish the maximum allowable time to discharge concrete after batching for various concrete temperatures.

More than controlling the maximum temperature is required to determine when to employ precautions to produce concrete with the required strength and durability. For most work it is too complex to simply limit only the maximum temperature of concrete as placed; circumstances and concrete requirements vary too widely. For example, a temperature limit that would serve successfully at one jobsite could be highly restrictive at another. Atmospheric conditions, including air temperature, relative humidity and wind speed, in conjunction with site conditions influence the precautions needed. For example, flatwork done under a roof that blocks solar radiation with exterior walls in place that screen the wind could be completed using a high temperature concrete; this concrete would cause difficulty if placed outdoors on the same day where it would be exposed to direct sun and wind.

Which precautions to use and when to use them will depend on: the type of construction; characteristics of the materials being used; and the experience of the placing and finishing crew in dealing with the atmospheric conditions on the site. The following list of precautions will reduce or avoid the potential problems of hot-weather concreting:

- use materials and mix proportions that have a good record in hot-weather conditions
- cool the concrete or one or more of its ingredients (Fig. 13-1)
- use a concrete consistency that allows rapid placement and consolidation
- reduce the time of transport, placing and finishing as much as possible
- schedule concrete placements to limit exposure to atmospheric conditions, such as at night or during favorable weather conditions
- consider methods to limit moisture loss during placing and finishing, such as sunshades, wind-screens, fogging, or spraying
- apply temporary moisture-retaining films after screeding
- organize a preconstruction conference to discuss the precautions required for the project

The above precautions are discussed in further detail throughout this chapter.

**EFFECTS OF HIGH CONCRETE TEMPERATURES**

As concrete temperature increases there is a loss in slump that is often unadvisedly compensated for by adding water to the concrete at the jobsite. At higher temperatures a greater amount of water is required to hold slump constant than is needed at lower temperatures. Adding water without adding cement results in a higher water-cement ratio, thereby lowering the strength at all ages and adversely affecting other desirable properties of the hardened concrete. This is in addition to the adverse effect on strength at later ages due to the higher temperature, even without the addition of water. Adding cement to compensate for the use of additional mix water may not be enough to achieve the desired concrete properties because additional cement will further increase the concrete temperature and water demand.

As shown in Fig. 13-2, if the temperature of freshly mixed concrete is increased from 10°C to 38°C (50°F to 100°F), about 20 kg/m³ (33 lb/ yd³) of additional water is needed to maintain the same 75-mm (3-in.) slump. This additional water could reduce strength by 12% to 15% and...
produce a compressive strength cylinder test result that may not comply with specifications.

High temperatures of freshly mixed concrete increase the rate of setting and shorten the length of time within which the concrete can be transported, placed, and finished. Setting time can be reduced by 2 or more hours with a 10°C (18°F) increase in concrete temperature (Fig. 13-3). Concrete should remain plastic long enough so that each layer can be placed without development of cold joints or discontinuities in the concrete. Retarding admixtures, ASTM C494 (AASHTO M 194) Type B, and hydration control admixtures can be beneficial in offsetting the accelerating effects of high temperature.

In hot weather, there is an increased tendency for cracks to form both before and after hardening. Rapid evaporation of water from freshly placed concrete can cause plastic-shrinkage cracks before the surface has hardened (discussed in more detail later in this chapter). Cracks may also develop in the hardened concrete because of increased drying shrinkage due to higher water contents or thermal volume changes as the concrete cools.

Air entrainment is also affected in hot weather. At elevated temperatures, an increase in the amount of air-entraining admixture is required to produce a given air content.

Fig. 13-4 shows the effect of high initial concrete temperatures on compressive strength. The concrete temperatures at the time of mixing, casting, and curing were 23°C (73°F), 32°C (90°F), 41°C (105°F), and 49°C (120°F). After 28 days, the specimens were all moist-cured at 23°C (73°F) until the 90-day and one-year test ages. The tests, using identical concretes of the same water-cement ratio, show that while higher concrete temperatures give higher early strength than concrete at 23°C (73°F), at later ages concrete strengths are lower. If the water content had been increased to maintain the same slump (without increasing cement content), the reduction in strength would have been even greater than shown.

The proper fabrication, curing, and testing of compression test specimens during hot weather is critical. Steps should be taken to make sure ASTM C31 (AASHTO T 23) procedures are followed regarding initial curing of strength specimens for acceptance or quality control testing at 16°C to 27°C (60°F to 80°F). If the initial 24 hour curing is at 38°C (100°F), the 28-day compressive strength of the test specimens may be 10% to 15% lower than if cured at the required ASTM C31 (AASHTO T 23) curing temperatures (Gaynor 1985).

Because of the detrimental effects of high concrete temperatures, all operations in hot weather should be directed toward keeping the concrete as cool as possible.

**COOLING CONCRETE MATERIALS**

The usual method of cooling concrete is to lower the temperature of the concrete materials before mixing. One or more of the ingredients should be cooled. In hot weather the aggregates and mixing water should be kept as cool as practicable; these materials have a greater influence on concrete temperature after mixing than other ingredients.

The contribution of each ingredient in a concrete mixture to the temperature of the freshly mixed concrete is related to the temperature, specific heat, and quantity of each material. Fig. 13-5 shows graphically the effect of temperature of materials on the temperature of fresh concrete. It is evident that although concrete temperature is

**Fig. 13-3. Effect of concrete temperature on setting time (Burg 1996).**

**Fig. 13-4. Effect of high concrete temperatures on compressive strength at various ages (Klieger 1958).**
primarily dependent upon the aggregate temperature, cooling the mixing water can be effective.

The approximate temperature of concrete can be calculated from the temperatures of its ingredients by using the following equation (NRMCA 1962):

$$T = \frac{0.22(T_a M_a + T_c M_c + T_w M_w + T_{wa} M_{wa})}{0.22(M_c + M_c + M_w + M_{wa})}$$

where

$T = \text{temperature of the freshly mixed concrete, °C (°F)}$

$T_a, T_c, T_w, \text{and } T_{wa} = \text{temperature in °C (°F) of aggregates, cement, added mixing water, and free water on aggregates, respectively}$

$M_a, M_c, M_w, \text{and } M_{wa} = \text{mass, kg (lb), of aggregates, cementing materials, added mixing water, and free water on aggregates, respectively}$

Example calculations for initial concrete temperature are shown in Table 13-1A.

Of all the materials in concrete, water is the easiest to cool. Even though it is used in smaller quantities than the other ingredients, cold water will produce a moderate reduction in the concrete temperature. Mixing water from a cool source should be used. It should be stored in tanks that are not exposed to the direct rays of the sun. Tanks and pipelines carrying mixing water should be buried, insulated, shaded, or painted white to keep water as cool as practical. Water can be cooled by refrigeration, liquid nitrogen, or ice. Cooling the mix water temperature 2.0°C to 2.2°C (3.5°F to 4°F) will usually lower the concrete temperature about 0.5°C (1°F). However, because mix water is such a small percentage of the total mixture, it is difficult to lower concrete temperatures more than about 4.5°C (8°F) by cooling the water alone.

Ice can be used as part of the mixing water provided it is completely melted by the time mixing is completed.

### Table 13-1A (Metric). Effect of Temperature of Materials on Initial Concrete Temperatures

<table>
<thead>
<tr>
<th>Material</th>
<th>Mass, $M$, kg</th>
<th>Specific heat kJ/kg • K</th>
<th>Joules to vary temperature, °C</th>
<th>Initial temperature of material, $T$, °C</th>
<th>Total joules in material</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>(1)</td>
<td>(2)</td>
<td>Col.1 x Col.2</td>
<td>(4)</td>
<td>(5) Col. 3 x Col. 4</td>
</tr>
<tr>
<td>Cement</td>
<td>335 ($M_c$)</td>
<td>0.92</td>
<td>308</td>
<td>66 ($T_c$)</td>
<td>20,328</td>
</tr>
<tr>
<td>Water</td>
<td>123 ($M_w$)</td>
<td>4.184</td>
<td>1692</td>
<td>27 ($T_w$)</td>
<td>45,684</td>
</tr>
<tr>
<td>Total aggregate</td>
<td>1839 ($M_a$)</td>
<td>0.92</td>
<td>2515</td>
<td>27 ($T_a$)</td>
<td>79,917</td>
</tr>
</tbody>
</table>

Initial concrete temperature = \(\frac{79,917}{2515} = 31.8°C\)

To achieve 1°C reduction in initial concrete temperature:

- Cement temperature must be lowered = \(\frac{2515}{308} = 8.2°F\)
- Or water temperature dropped = \(\frac{2515}{515} = 4.9°F\)
- Or aggregate temperature cooled = \(\frac{2515}{1692} = 1.5°F\)

### Table 13-1A (Inch-Pound Units). Effect of Temperature of Materials on Initial Concrete Temperatures

<table>
<thead>
<tr>
<th>Material</th>
<th>Mass, $M$, lb</th>
<th>Specific heat Btu/°F • lb</th>
<th>Btu to vary temperature, °F</th>
<th>Initial temperature of material, $T$, °F</th>
<th>Total Btu’s in material</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>(1)</td>
<td>(2)</td>
<td>Col.1 x Col.2</td>
<td>(4)</td>
<td>(5) Col. 3 x Col. 4</td>
</tr>
<tr>
<td>Cement</td>
<td>564 ($M_c$)</td>
<td>0.22</td>
<td>124</td>
<td>150 ($T_c$)</td>
<td>18,600</td>
</tr>
<tr>
<td>Water</td>
<td>282 ($M_w$)</td>
<td>1.00</td>
<td>282</td>
<td>80 ($T_w$)</td>
<td>22,560</td>
</tr>
<tr>
<td>Total aggregate</td>
<td>3100 ($M_a$)</td>
<td>0.22</td>
<td>682</td>
<td>80 ($T_a$)</td>
<td>54,560</td>
</tr>
</tbody>
</table>

Initial concrete temperature = \(\frac{95,720}{1088} = 88.0°F\)

To achieve 1°F reduction in initial concrete temperature:

- Cement temperature must be lowered = \(\frac{1088}{124} = 8.8°F\)
- Or water temperature dropped = \(\frac{1088}{282} = 3.9°F\)
- Or aggregate temperature cooled = \(\frac{1088}{682} = 1.6°F\)
When using crushed ice, care must be taken to store it at a temperature that will prevent the formation of lumps.

When ice is added as part of the mixing water, the effect of the heat of fusion of the ice must be considered; so the equation for temperature of fresh concrete is modified as follows:

$$T (°C) = \frac{0.22(T_a M_a + T_c M_c) + T_w M_w + T_{wa} M_{wa} - 80 M_i}{0.22(M_a + M_c) + M_w + M_{wa} + M_i}$$

$$T (°F) = \frac{0.22(T_a M_a + T_c M_c) + T_w M_w + T_{wa} M_{wa} - 112 M_i}{0.22(M_a + M_c) + M_w + M_{wa} + M_i}$$

where $M_i$ is the mass in kg (lb) of ice (NRMCA 1962 and Mindess and Young 1981).

The heat of fusion of ice in metric units is 335 kJ per kg (in British thermal units, 144 Btu per pound). Calculations in Table 13-1B show the effect of 44 kg (75 lb) of ice in reducing the temperature of concrete. The amount of water and ice must not exceed the total mixing-water requirements.

Fig. 13-6 shows crushed ice being charged into a truck mixer prior to the addition of other materials. Mixing time should be long enough to completely melt the ice. The volume of ice should not replace more than approximately 75% of the total batch water. The maximum temperature reduction from the use of ice is limited to about 11°C (20°F).

If a greater temperature reduction is required, the injection of liquid nitrogen into the mixer may be the best alternative method.

The liquid nitrogen can be added directly into a central mixer drum or the drum of a truck mixer to lower concrete temperature. Fig. 13-1 shows liquid nitrogen added directly into a truck mixer near a ready mix plant. Care should be taken to prevent the liquid nitrogen from contacting the metal drum; the super cold liquid nitrogen may crack the drum. The addition of liquid nitrogen does not in itself influence the amount of mix water required except that lowering the concrete temperature can reduce water demand.

Aggregates have a pronounced effect on the fresh concrete temperature because they represent 70% to 85% of the total mass of concrete. To lower the temperature of concrete 0.5°C (1°F) requires only a 0.8°C to 1.1°C (1.5°F to 2°F) reduction in the temperature of the coarse aggregate.

There are several simple methods of keeping aggregates cool. Stockpiles should be shaded from the sun and

<table>
<thead>
<tr>
<th>Table 13-1B (Metric). Effect of Ice (44 kg) on Temperature of Concrete</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Material</strong></td>
</tr>
<tr>
<td></td>
</tr>
<tr>
<td>Cement</td>
</tr>
<tr>
<td>Water</td>
</tr>
<tr>
<td>Total aggregate</td>
</tr>
<tr>
<td>Ice</td>
</tr>
<tr>
<td>minus</td>
</tr>
</tbody>
</table>

Concrete temperature = $\frac{65,177}{2699} = 24.1°C$

<table>
<thead>
<tr>
<th>Table 13-1B (Inch-Pound Units). Effect of Ice (75 lb) on Temperature of Concrete</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Material</strong></td>
</tr>
<tr>
<td></td>
</tr>
<tr>
<td>Cement</td>
</tr>
<tr>
<td>Water</td>
</tr>
<tr>
<td>Total aggregate</td>
</tr>
<tr>
<td>Ice*</td>
</tr>
<tr>
<td>minus</td>
</tr>
</tbody>
</table>

Concrete temperature = $\frac{81,320}{1088} = 74.7°F$

*32 $M_i = -112 M_i$
kept moist by sprinkling. Do not spray salt water on aggregate stockpiles. Since evaporation is a cooling process, sprinkling provides effective cooling, especially when the relative humidity is low.

Sprinkling of coarse aggregates should be adjusted to avoid producing excessive variations in the surface moisture content and thereby causing a loss of slump uniformity. Refrigeration is another method of cooling materials. Aggregates can be immersed in cold-water tanks, or cooled air can be circulated through storage bins. Vacuum cooling can reduce aggregate temperatures to as low as 1°C (34°F).

Cement temperature has only a minor effect on the temperature of the freshly mixed concrete because of cement’s low specific heat and the relatively small amount of cement in a concrete mixture. A cement temperature change of 5°C (9°F) generally will change the concrete temperature by only 0.5°C (1°F). Because cement loses heat slowly during storage, it may still be warm when delivered. (This heat is produced in grinding the cement clinker during manufacture.) Since the temperature of cement does affect the temperature of the fresh concrete to some extent, some specifications place a limit on its temperature at the time of use. This limit varies from 66°C to 82°C (150°F to 180°F) (ACI Committee 305). However, it is preferable to specify a maximum temperature for freshly mixed concrete rather than place a temperature limit on individual ingredients (Lerch 1955).

SUPPLEMENTARY CEMENTITIOUS MATERIALS

Many concrete producers consider the use of supplementary cementitious materials to be essential in hot weather conditions. The materials of choice are fly ash and other pozzolans (ASTM C618 or AASHTO M 295) and ground granulated blast-furnace slag (ASTM C989 or AASHTO M 302). These materials generally slow both the rate of setting as well as the rate of slump loss. However, some caution regarding finishing is needed; because the rate of bleeding can be slower than the rate of evaporation, plastic shrinkage cracking or crazing may result. This is discussed in greater detail under “Plastic Shrinkage Cracking” below.

PREPARATION BEFORE CONCRETING

Before concrete is placed, certain precautions should be taken during hot weather to maintain or reduce concrete temperature. Mixers, chutes, conveyor belts, hoppers, pump lines, and other equipment for handling concrete should be shaded, painted white, or covered with wet burlap to reduce solar heat.

Forms, reinforcing steel, and subgrade should be fogged or sprinkled with cool water just before the con-
crete is placed. Fogging the area during placing and finishing operations not only cools the contact surfaces and surrounding air but also increases its relative humidity. This reduces the temperature rise of the concrete and minimizes the rate of evaporation of water from the concrete after placement. For slabs on ground, it is a good practice to moisten the subgrade the evening before con creting. There should be no standing water or puddles on forms or subgrade at the time concrete is placed.

During extremely hot periods, improved results can be obtained by restricting concrete placement to early morning, evening, or nighttime hours, especially in arid climates. This practice has resulted in less thermal shrinkage and cracking of thick slabs and pavements.

**TRANSPORTING, PLACING, FINISHING**

Transporting and placing concrete should be done as quickly as practical during hot weather. Delays contribute to loss of slump and an increase in concrete temperature. Sufficient labor and equipment must be available at the jobsite to handle and place concrete immediately upon delivery.

Prolonged mixing, even at agitating speed, should be avoided. If delays occur, stopping the mixer and then agitating intermittently can minimize the heat generated by mixing. ASTM C94 (AASHTO M 157) requires that discharge of concrete be completed within 1½ hours or before the drum has revolved 300 times, whichever occurs first. During hot weather the time limit can be reasonably reduced to 1 hour or even 45 minutes. If specific time limitations on the completion of discharge of the concrete are desired, they should be included in the project specifications. It is also reasonable to obtain test data from a trial batch simulating the time, mixing, and anticipated concrete temperature to document, if necessary, a reduction in the time limit.

Since the setting of concrete is more rapid in hot weather, extra care must be taken with placement techniques to avoid cold joints. For placement of walls, shallower layers can be specified to assure enough time for consolidation with the previous lift. Temporary sunshades and windbreaks help to minimize cold joints.

Floating of slabs should be done promptly after the water sheen disappears from the surface or when the concrete can support the weight of a finisher with no more than a 5-mm (¼-in.) indentation. Finishing on dry and windy days requires extra care. Rapid drying of the concrete at the surface may cause plastic shrinkage cracking.

**PLASTIC SHRINKAGE CRACKING**

Plastic shrinkage cracks sometimes occur in the surface of freshly mixed concrete soon after it has been placed, while it is being finished or shortly thereafter (Fig. 13-7). These cracks which appear mostly on horizontal surfaces can be substantially eliminated if preventive measures are taken.

Plastic shrinkage cracking is usually associated with hot-weather concreting; however, it can occur any time ambient conditions produce rapid evaporation of moisture from the concrete surface. These cracks occur when water evaporates from the surface faster than it can travel to the surface during the bleeding process. This creates rapid drying shrinkage and tensile stresses in the surface that often result in short, irregular cracks. The following conditions, singly or collectively, increase evaporation of surface moisture and increase the possibility of plastic shrinkage cracking:

1. High air temperature
2. High concrete temperature
3. Low humidity
4. High wind speed

The crack length is generally 50 to 1000 mm (a few inches to 3 ft) in length and they are usually spaced in an irregular pattern from 50 to 700 mm (a few inches to 2 ft) apart. Fig. 13-8 is useful for determining when precautionary measures should be taken. There is no way to predict with certainty when plastic shrinkage cracking will occur.

When the rate of evaporation exceeds 1 kg/m² (0.2 lb/ft²) per hour, precautionary measures such as wind-screens are almost mandatory. With some concrete mixtures, such as those containing pozzolans, cracking is possible if the rate of evaporation exceeds 0.5 kg/m² (0.1 lb/ft²) per hour. Concrete containing silica fume is particularly prone to plastic shrinkage because bleeding rates are commonly only 0.25 kg/m² (0.05 lb/ft²) per hour. Therefore, protection from premature drying is essential at lower evaporation rates. At some point in the process of setting, bleeding goes to zero and the surface begins to dry at evaporation rates much lower than the typically specified 1.0 kg/m² (0.2 lb/ft²) per hour; in such cases, further protection becomes necessary regardless of the type of concrete mixture.

![Fig. 13-7. Typical plastic shrinkage cracks.](IMG12267)
To use these charts:

1. Enter with air temperature, move **up** to relative humidity.
2. Move **right** to concrete temperature.
3. Move **down** to wind velocity.
4. Move **left**: read approximate rate of evaporation.

One or more of the precautions listed below can minimize the occurrence of plastic shrinkage cracking. They should be considered while planning for hot-weather concrete construction or while dealing with the problem after construction has started. They are listed in the order in which they should be done during construction.

1. Moisten concrete aggregates that are dry and absorptive.
2. Keep the concrete temperature low by cooling aggregates and mixing water.
3. Dampen the subgrade (Fig. 13-9) and fog forms prior to placing concrete.
4. Erect temporary windbreaks to reduce wind velocity over the concrete surface.
5. Erect temporary sunshades to reduce concrete surface temperatures.

Fig. 13-8. Effect of concrete and air temperatures, relative humidity, and wind velocity on rate of evaporation of surface moisture from concrete. Wind speed is the average horizontal air or wind speed in km/h (mph) measured at 500 mm (20 in.) above the evaporating surface. Air temperature and relative humidity should be measured at a level approximately 1.2 to 1.8 m (4 to 6 ft) above the evaporating surface and on the windward side shielded from the sun’s rays (Menzel 1954).
6. Protect the concrete with temporary coverings, such as polyethylene sheeting, during any appreciable delay between placing and finishing.
7. Fog the slab immediately after placing and before finishing, taking care to prevent the accumulation of water that may reduce the quality of the cement paste in the slab surface.
8. Add plastic fibers to the concrete mixture to help reduce plastic shrinkage crack formation.

Fogging the concrete before and after final finishing is the most effective way to minimize evaporation and reduce plastic shrinkage cracking. Use of a fog spray will raise the relative humidity of the ambient air over the slab, thus reducing evaporation from the concrete. Fog nozzles atomize water using air pressure (Figs. 13-10 and 13-11) to create a fog blanket. They should not be confused with garden-hose nozzles, which leave an excess amount of water on the slab. Fogging should be continued until a suitable curing material such as a curing compound, wet burlap, or curing paper can be applied.

Other methods to prevent the rapid loss of moisture from the concrete surface include:

- Spray application of temporary moisture-retaining films (usually polymers); these compounds can be applied immediately after screeding to reduce water evaporation before final finishing operations and curing commence. These materials are floated and troweled into the surface during finishing and should have no adverse effect on the concrete or inhibit the adhesion of membrane-curing compounds.
- Reduction of time between placing and the start of curing by eliminating delays during construction.

If plastic shrinkage cracks should appear during finishing, striking each side of the crack with a float and refinishing can close the cracks. However, the cracking may reoccur unless the causes are corrected.

**CURING AND PROTECTION**

Curing and protection are more critical in hot weather than in temperate periods. Retaining forms in place cannot be considered a satisfactory substitute for curing in hot weather; they should be loosened as soon as practical without damage to the concrete. Water should then be applied at the top exposed concrete surfaces—for example,
with a soil-soaker hose—and allowed to run down inside
the forms. On hardened concrete and on flat concrete sur-
faces in particular, curing water should not be more than
about 11°C (20°F) cooler than the concrete. This will mini-
mize cracking caused by thermal stresses due to tempera-
ture differentials between the concrete and curing water.

The need for moist curing is greatest during the first
few hours after finishing. To prevent the drying of exposed
concrete surfaces, moist curing should commence as soon
as the surfaces are finished and continue for at least 24
hours. In hot weather, continuous moist curing for the
entire curing period is preferred. However, if moist curing
cannot be continued beyond 24 hours, while the surfaces
are still damp, the concrete should be protected from
drying with curing paper, heat-reflecting plastic sheets, or
membrane-forming curing compounds.

White-pigmented curing compounds can be used on
horizontal surfaces. Application of a curing compound
during hot weather should be preceded by 24 hours of
moist curing. If this is not practical, the compound should
be applied immediately after final finishing. The concrete
surfaces should be moist.

Moist-cured surfaces should dry out slowly after the
curing period to reduce the possibility of surface crazing
and cracking. Crazing, a network pattern of fine cracks
that do not penetrate much below the surface, is caused by
minor surface shrinkage. Crazing cracks are very fine and
barely visible except when the concrete is drying after the
surface has been wet. The cracks encompass small con-
crete areas less than 50 mm (2 in.) in dimension, forming a
chicken-wire like pattern.

ADMIXTURES

For unusual cases in hot weather and where careful in-
spection is maintained, a retarding admixture may be ben-
eficial in delaying the setting time, despite the somewhat
increased rate of slump loss resulting from their use. A
hydration control admixture can be used to stop cement
hydration and setting. Hydration is resumed, when de-
sired, with the addition of a special accelerator (reactivator).

Retarding admixtures should conform to the require-
ments of ASTM C494 (AASHTO M 194) Type B. Ad-
mixtures should be tested with job materials under job
conditions before construction begins; this will determine
their compatibility with the basic concrete ingredients and
their ability under the particular conditions to produce the
desired results.

HEAT OF HYDRATION

Heat generated during cement hydration raises the tem-
perature of concrete to a greater or lesser extent de-
pending on the size of the concrete placement, its
surrounding environment, and the amount of cement in
the concrete. As a general rule a 5°C to 9°C (10°F to 15°F)
temperature rise per 45 kg (100 lb) of portland cement can
be expected from the heat of hydration (ACI Committee
211 1991). There may be instances in hot-weather-concrete
work and massive concrete placements when measures
must be taken to cope with the generation of heat from
cement hydration and attendant thermal volume changes
to control cracking (see Chapters 15 and 18).

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CHAPTER 14
Cold-Weather Concreting

Concrete can be placed safely without damage from freezing throughout the winter months in cold climates if certain precautions are taken. Cold weather is defined by ACI Committee 306 as a period when for more than 3 successive days the average daily air temperature drops below 5°C (40°F) and stays below 10°C (50°F) for more than one-half of any 24 hour period. Under these circumstances, all materials and equipment needed for adequate protection and curing must be on hand and ready for use before concrete placement is started. Normal concreting practices can be resumed once the ambient temperature is above 10°C (50°F) for more than half a day.

During cold weather, the concrete mixture and its temperature should be adapted to the construction procedure and ambient weather conditions. Preparations should be made to protect the concrete; enclosures, windbreaks, portable heaters, insulated forms, and blankets should be ready to maintain the concrete temperature (Fig. 14-1).

Forms, reinforcing steel, and embedded fixtures must be clear of snow and ice at the time concrete is placed. Thermometers and proper storage facilities for test cylinders should be available to verify that precautions are adequate.

**EFFECT OF FREEZING FRESH CONCRETE**

Concrete gains very little strength at low temperatures. Freshly mixed concrete must be protected against the disruptive effects of freezing (Fig. 14-2) until the degree of saturation of the concrete has been sufficiently reduced by the process of hydration. The time at which this reduction is accomplished corresponds roughly to the time required for the concrete to attain a compressive strength of 3.5 MPa (500 psi) (Powers 1962). At normal temperatures and water-cement ratios less than 0.60, this occurs within the first 24 hours after placement. Significant ultimate strength reductions, up to about 50%, can occur if concrete is frozen within a few hours after placement or before it attains a compressive strength of 3.5 MPa (500 psi) (McNeese 1952). Concrete to be exposed to deicers should attain a strength of 28 MPa (4,000 psi) prior to repeated cycles of freezing and thawing (Klieger 1957).

Concrete that has been frozen just once at an early age can be restored to nearly normal strength by providing favorable subsequent curing conditions. Such concrete, however, will not be as resistant to weathering nor as impermeable as concrete that had not been frozen. The critical period after which concrete is not seriously damaged by one

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Fig. 14-1. When suitable preparations to build enclosures and insulate equipment have been made, cold weather is no obstacle to concrete construction. (IMG12272, IMG12271)
or two freezing cycles is dependent upon the concrete ingredients and conditions of mixing, placing, curing, and subsequent drying. For example, air-entrained concrete is less susceptible to damage by early freezing than non-air-entrained concrete. See Chapter 8, “Air-Entrained Concrete,” for more information.

### STRENGTH GAIN OF CONCRETE AT LOW TEMPERATURES

Temperature affects the rate at which hydration of cement occurs—low temperatures retard hydration and consequently retard the hardening and strength gain of concrete.

If concrete is frozen and kept frozen above about minus 10°C (14°F), it will gain strength slowly. Below that temperature, cement hydration and concrete strength gain cease. Fig. 14-3 illustrates the effect of cool temperatures on setting time. Fig. 14-4 illustrates the effects of casting temperature on slump. Figs. 14-5 and 14-6 show the age-compressive strength relationship for concrete that has been cast and cured at various temperatures. Note in Fig. 14-6 that concrete cast and cured at 4°C (40°F) and 13°C (55°F) had relatively low strengths for the first week; but after 28 days—when all specimens were moist-cured at 23°C (73°F)—strengths for the 4°C (40°F) and 13°C (55°F) concretes grew faster than the 23°C (73°F) concrete and at one year they were slightly higher.
Higher-early strengths can be achieved through use of Type III high-early-strength cement as illustrated in Fig. 14-7. Principal advantages occur during the first 7 days. At a 4°C (40°F) curing temperature, the advantages of Type III cement are more pronounced and persist longer than at the higher temperature.

HEAT OF HYDRATION

Concrete generates heat during hardening as a result of the chemical process by which cement reacts with water to form a hard, stable paste. The heat generated is called heat of hydration; it varies in amount and rate for different cements. Dimensions of the concrete placement, ambient air temperature, initial concrete temperature, water-cement ratio, admixtures, and the composition, fineness, and amount of cementitious material all affect heat generation and buildup.

Heat of hydration is useful in winter concreting as it contributes to the heat needed to provide a satisfactory curing temperature; often without other temporary heat sources, particularly in more massive elements.

Concrete must be delivered at the proper temperature and account must be taken of the temperature of forms, reinforcing steel, the ground, or other concrete on which the fresh concrete is cast. Concrete should not be cast on frozen concrete or on frozen ground.

Fig. 14-8 shows a concrete pedestal being covered with a tarpaulin just after the concrete was placed. Tarpaulins and insulated blankets are often necessary to retain the heat of hydration more efficiently and keep the concrete as warm as possible. Thermometer readings of
the concrete’s temperature will tell whether the covering is adequate. The heat liberated during hydration will offset to a considerable degree the loss of heat during placing, finishing, and early curing operations. As the heat of hydration slows down, the need to cover the concrete becomes more important.

SPECIAL CONCRETE MIXTURES

High strength at an early age is desirable in winter construction to reduce the length of time temporary protection is required. The additional cost of high-early-strength concrete is often offset by earlier reuse of forms and shores, savings in the shorter duration of temporary heating, earlier setting times that allows the finishing of flatwork to begin sooner, and earlier use of the structure. High-early-strength concrete can be obtained by using one or a combination of the following:

1. Type III or HE high-early-strength cement
2. Additional portland cement (60 to 120 kg/m³ or 100 to 200 lb/yd³)
3. Chemical accelerators

Small amounts of an accelerating admixture can be used to accelerate the setting and early-age strength development of concrete in cold weather. Accelerators containing chlorides should not be used where there is an in-service potential for corrosion, such as in concrete members containing steel reinforcement or where aluminum or galvanized inserts will be used. Chlorides are not recommended for concretes exposed to soil or water containing sulfates or for concretes susceptible to alkali-aggregate reaction.

Accelerators must not be used as a substitute for proper curing and frost protection. Specially designed accelerating admixtures allow concrete to be placed at temperatures down to -7°C (20°F). The purpose of these admixtures is to reduce the time of initial setting, but not necessarily to speed up strength gain. Covering concrete to keep out moisture and to retain heat of hydration is still necessary. Furthermore, traditional antifreeze solutions, as used in automobiles, should never be used; the quantity of these materials needed to appreciably lower the freezing point of concrete is so great that strength and other properties can be seriously affected.

Since the goal of using special mixtures during cold weather concreting is to reduce the time of setting, a low water-cement ratio, low-slump concrete is particularly desirable, especially for cold-weather flatwork; concrete mixtures with higher slumps usually take longer to set. In addition, evaporation is minimized so that finishing can be accomplished quicker (Fig. 14-9).

AIR-ENTRAINED CONCRETE

Entrained air is particularly desirable in any concrete that will be exposed to freezing weather. Concrete that is not air entrained can suffer strength loss and internal as well as surface damage as a result of freezing and thawing (Fig. 14-10). Air entrainment provides the capacity to absorb stresses due to ice formation within the concrete. See Chapter 8, “Air-Entrained Concrete.”
Air entrainment should always be used for construction during the freezing months. The exception is concrete work done under roof where there is no chance that rain, snow, or water from other sources can saturate the concrete and where there is no chance of freezing.

The likelihood of water saturating a concrete floor during construction is very real. Fig. 14-11 shows conditions in the upper story of an apartment building during winter construction. Snow fell on the top deck. When heaters were used below to warm the deck, the snow melted. Water ran through floor openings down to a level that was not being heated. The water-saturated concrete froze, which caused a strength loss, particularly at the floor surface. This could also result in greater deflection of the floor and a surface that is less wear-resistant than it might have been.

**TEMPERATURE OF CONCRETE**

**Temperature of Concrete as Mixed**

The temperature of fresh concrete as mixed should not be less than shown in Lines 1, 2, or 3 of Table 14-1 for the respective thickness of section. Note that lower concrete temperatures are recommended for more massive concrete sections because heat generated during hydration is dissipated less rapidly in heavier sections. Also note that at lower ambient air temperatures more heat is lost from concrete during transporting and placing; hence, the recommended concrete temperatures as mixed are higher for colder weather.

### Table 14-1. Recommended Concrete Temperature for Cold-Weather Construction—Air-Entrained Concrete*

<table>
<thead>
<tr>
<th>Line</th>
<th>Condition</th>
<th>Thickness of sections, mm (in.)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>Less than 300 (12)</td>
</tr>
<tr>
<td>1</td>
<td>Minimum temperature of fresh concrete as mixed for weather indicated.</td>
<td>Above -1°C (30°F)</td>
</tr>
<tr>
<td>2</td>
<td>-18°C to -1°C (0°F to 30°F)</td>
<td>18°C (65°F)</td>
</tr>
<tr>
<td>3</td>
<td>Below -18°C (0°F)</td>
<td>21°C (70°F)</td>
</tr>
<tr>
<td>4</td>
<td>Minimum temperature of fresh concrete as placed and maintained.**</td>
<td>13°C (55°F)</td>
</tr>
</tbody>
</table>

* Adapted from Table 3.1 of ACI 306R-88.
** Placement temperatures listed are for normal-weight concrete. Lower temperatures can be used for lightweight concrete if justified by tests. For recommended duration of temperatures in Line 4, see Table 14-3.
Design and Control of Concrete Mixtures • EB001

There is little advantage in using fresh concrete at a temperature much above 21°C (70°F). Higher concrete temperatures do not afford proportionately longer protection from freezing because the rate of heat loss is greater. Also, high concrete temperatures are undesirable since they increase thermal shrinkage after hardening, require more mixing water for the same slump, and contribute to the possibility of plastic-shrinkage cracking (caused by rapid moisture loss through evaporation). Therefore, the temperature of the concrete as mixed should not be more than 5°C (10°F) above the minimums recommended in Table 14-1.

**Aggregate Temperature.** The temperature of aggregates varies with weather and type of storage. Aggregates usually contain frozen lumps and ice when the temperature is below freezing. Frozen aggregates must be thawed to avoid pockets of aggregate in the concrete after batching, mixing, and placing. If thawing takes place in the mixer, excessively high water contents in conjunction with the cooling effect due to the ice melting must be avoided.

At temperatures above freezing it is seldom necessary to heat aggregates, the desired concrete temperature can usually be obtained by heating only the mixing water. At temperatures below freezing, in addition to heating the mixing water, often only the fine aggregate needs to be heated to produce concrete of the required temperature, provided the coarse aggregate is free of frozen lumps.

Three of the most common methods for heating aggregates are: (1) storing in bins or weigh hoppers heated by steam coils or live steam; (2) storing in silos heated by hot air or steam coils; and (3) stockpiling over heated slabs, stem vents or pipes. Although heating aggregates stored in bins or weigh hoppers is most commonly used, the volume of aggregate that can be heated is often limited and quickly consumed during production. Circulating steam through pipes over which aggregates are stockpiled is a recommended method for heating aggregates. Stockpiles can be covered with tarpaulins to retain and distribute heat and to prevent formation of ice. Live steam, preferably at pressures of 500 to 900 kPa (75 to 125 psi), can be injected directly into the aggregate pile to heat it, but the resultant variable moisture content in aggregates might result in erratic mixing-water control.

On small jobs aggregates can be heated by stockpiling over metal culvert pipes in which fires are maintained. Care should be taken to prevent scorching the aggregates.

**Mixing-Water Temperature.** Of the ingredients used to make concrete, mixing water is the easiest and most practical to heat. The mass of aggregates and cement in concrete is much greater than the mass of water; however, water can store about five times as much heat as can cement and aggregate of the same weight. For cement and aggregates, the average specific heat (that is, heat units required to raise the temperature 1°C (1°F) per kg (lb) of material) can be assumed as 0.925 kJ (0.22 Btu) compared to 4.187 kJ (1.0 Btu) for water.

Fig. 14-12 shows the effect of temperature of materials on temperature of fresh concrete. The chart is based on the equation

\[
T = \frac{0.22 (T_a M_a + T_c M_c) + T_w M_w + T_{wa} M_{wa}}{0.22 (M_a + M_c) + M_w + M_{wa}}
\]

where

\[
T = \text{temperature in degrees Celsius (Fahrenheit) of the fresh concrete}
\]

\[
T_a, T_c, T_w, \text{ and } T_{wa} = \text{temperature in degrees Celsius (Fahrenheit) of the aggregates, cement, added mixing water, and free moisture on aggregates, respectively; generally } T_a = T_{wa}
\]

\[
M_a, M_c, M_w, \text{ and } M_{wa} = \text{mass in kilograms (pounds) of the aggregates, cement, free moisture on aggregates, and mixing water, respectively}
\]

If the weighted average temperature of aggregates and cement is above 0°C (32°F), the proper mixing-water temperature for the required concrete temperature can be selected from Fig. 14-12. The range of concrete temperatures in the chart corresponds with the recommended values given in Lines 1, 2, and 3 of Table 14-1.

![Fig. 14-12. Temperature of mixing water needed to produce heated concrete of required temperature. Temperatures are based on the mixture shown but are reasonably accurate for other typical mixtures.](image-url)
To avoid the possibility of a quick or flash set of the concrete when either water or aggregates are heated to above 38°C (100°F), they should be combined in the mixer first before the cement is added. If this mixer-loading sequence is followed, water temperatures up to the boiling point can be used, provided the aggregates are cold enough to reduce the final temperature of the aggregates and water mixture to appreciably less than 38°C (100°F).

Fluctuations in mixing-water temperature from batch to batch should be avoided. The temperature of the mixing water can be adjusted by blending hot and cold water.

**Temperature of Concrete as Placed and Maintained**

There will be some temperature loss after mixing while the truck mixer is traveling to the construction site and waiting to discharge its load. The concrete should be placed in the forms before its temperature drops below that given on Line 4 of Table 14-1; that concrete temperature should be maintained for the duration of the protection period given in Chapter 12 under “Curing Period and Temperature.”

**Cooling After Protection**

To avoid cracking of the concrete due to sudden temperature change near the end of the curing period, ACI Committee 306 requires that the source of heat and cover protection be slowly removed. The maximum allowable temperature drop during the first 24 hours after the end of the protection is given in Table 14-2. The temperature drops apply to surface temperatures. Notice that the cooling rates for surfaces of mass concrete (thick sections) are lower than for thinner members.

**CONTROL TESTS**

Thermometers are needed to check the concrete temperatures as delivered, as placed, and as maintained. An inexpensive pocket thermometer is shown in Fig. 14-13.

After the concrete has hardened, temperatures can be checked with special surface thermometers or with an ordinary thermometer that is kept covered with insulating blankets. A simple way to check temperature below the concrete surface is shown in Fig. 14-14. Instead of filling the hole shown in Fig. 14-14 with a fluid, it can be fitted with insulation except at the bulb.

Concrete test cylinders must be maintained at a temperature between 16°C (60°F) and 27°C (80°F) at the jobsite for up to 48 hours until they are taken to a laboratory for curing (ASTM C31 or AASHTO T 23). For concrete mixtures with a specified strength of 40 MPa (6,000 psi) or greater, the initial curing temperature shall be between 20°C and 26°C (68°F and 78°F). During this period, cylinders should be kept in a curing box and covered with a nonabsorptive, nonreactive plate or impervious plastic bag; the temperature in the box should be accurately con-

<table>
<thead>
<tr>
<th>Section size, minimum dimensions, mm (in.)</th>
<th>Less than 300 (12)</th>
<th>300 to 900 (12 to 36)</th>
<th>900 to 1800 (36 to 72)</th>
<th>Over 1800 (72)</th>
</tr>
</thead>
<tbody>
<tr>
<td>28°C (50°F)</td>
<td>22°C (40°F)</td>
<td>17°C (30°F)</td>
<td>11°C (20°F)</td>
<td></td>
</tr>
</tbody>
</table>

* Adapted from Table 5.5 of ACI 306R-88.
Concreting on ground during cold weather involves some extra effort and expense, but many contractors find that it more than pays for itself. In winter, the site around the structure may be frozen rather than a morass of mud. The concrete will furnish some if not all of the heat needed for proper curing. Internal concrete temperatures should be monitored. Insulated blankets or simple enclosures are easily provided. Embankments are frozen and require less bracing. With a good start during the winter months, construction gets above the ground before warmer weather arrives.

Placing concrete on the ground involves different procedures from those used at an upper level: (1) the ground must be thawed before placing concrete; (2) cement hydration will furnish some of the curing heat; (3) construction of enclosures is much simpler and use of insulating blankets may be sufficient; (4) in the case of a floor slab, a vented heater is required if the area is enclosed; and (5) hydronic heaters can be used to thaw subgrades using insulated blankets or to heat enclosures without concern for carbonation. For more on hydronic heaters, see “Heaters” later in this chapter.

Once cast, footings should be backfilled as soon as possible with unfrozen fill. Concrete should never be placed on a frozen subgrade or backfilled with frozen fill; otherwise once they thaw, uneven settlements may occur causing cracking.

ACI Committee 306 requires that concrete not be placed on any surface that would lower the temperature of the concrete in place below the minimum values shown on Line 4 in Table 14-1. In addition, concrete placement temperatures should not be higher than these minimum values by more than 11°C (20°F) to reduce rapid moisture loss and the potential development of plastic shrinkage cracks.

When the subgrade is frozen to a depth of approximately 80 mm (3 inches), the surface region can be thawed by (1) steaming; (2) spreading a layer of hot sand, gravel, or other granular material where the grade elevations allow it; (3) removing and replacing with unfrozen fill; (4) covering the subgrade with insulation for a few days; or (5) using hydronic heaters under insulated blankets which can thaw frozen ground at a rate of 0.3 m (1 ft) per 24 hours to a depth up to 3 m (10 ft) (Grochoski 2000). Placing concrete for floor slabs and exposed footings should be delayed until the ground thaws and warms sufficiently to ensure that it will not freeze again during the protection and curing period.

Slabs can be cast on ground at ambient temperatures as low as 2°C (35°F) as long as the minimum concrete temperature as placed is not less than shown on Line 4 of Table 14-1. Although surface temperatures need not be higher than a few degrees above freezing, they also should preferably not be more than 5°C (10°F) higher than the minimum placement temperature either. The duration of curing should not be less than that described in Chapter 12 for the

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**Fig. 14-15. Insulated curing box with thermostat for curing test cylinders. Heat is supplied by electric rubber heating mats on the bottom. A wide variety of designs are possible for curing boxes. (IMG12268)**

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trolled by a thermostat (Fig. 14-15). When stored in an insulated curing box outdoors, cylinders are less likely to be jostled by vibrations than if left on the floor of a trailer. If kept in a trailer where the heat may be turned off at night or over a weekend or holiday, the cylinders would not be at the prescribed curing temperatures during this critical period.

In addition to laboratory-cured cylinders, it is useful to field-cure some test cylinders in order to monitor actual curing conditions on the job in cold weather. It is sometimes difficult to find the right locations for field curing. Differences in the surface to volume ratios between cylinders and the structure, in conjunction with differences in mass, make correlating field-cured cylinder strengths to in-place strengths difficult. A preferred location is in a boxout in a floor slab or wall with thermal insulation for cover. When placed on a formwork ledge just below a heated, suspended floor, possible high temperatures there will not duplicate the average temperature in the slab, nor the lowest temperature on top of the slab. Still, field-cured cylinders are more indicative of actual concrete strength than laboratory-cured cylinders. Particular care should be taken to protect compressive strength test cylinders from freezing; their small mass may not generate enough heat of hydration to protect them.

Molds stripped from cylinders after the first 24 ± 8 hours must be wrapped tightly in plastic bags or laboratory curing started immediately. When cylinders are picked up for delivery to the laboratory, they must be maintained at a temperature of 16°C (60°F) to 27°C (80°F) until they are placed in the laboratory curing room.

Cast-in-place cylinders (ASTM C873) and non-destructive testing methods discussed in Chapter 16, as well as maturity techniques discussed later in this chapter, are helpful in monitoring in-place concrete strength.

**CONCRETING ON GROUND**
appropriate exposure classification. Because of the risk of surface imperfections that might occur on exterior concrete placed in late fall and winter, many concrete contractors choose to delay concrete placement until spring. By waiting until spring, temperatures will be more favorable for cement hydration; this will help generate adequate strengths along with sufficient drying so the concrete can resist freeze-thaw damage.

CONCRETING ABOVEGROUND

Working aboveground in cold weather usually involves several different approaches compared to working at ground level:

1. The concrete mixture need not be changed to generate more heat because portable heaters can be used to heat the undersides of floor and roof slabs. Nevertheless, there are advantages to having a mix that will produce a high strength at an early age; for example, artificial heat can be cut off sooner (see Table 14-3), and forms can be recycled faster.
2. Enclosures must be constructed to retain the heat under floor and roof slabs.
3. Portable heaters used to warm the underside of formed concrete can be direct-fired heating units (without venting).

Before placing concrete, the heaters under a formed deck should be turned on to preheat the forms and melt any snow or ice remaining on top. Temperature requirements for surfaces in contact with fresh concrete are the same as those outlined in the previous section “Concreting on Ground.” Metallic embedments at temperatures below the freezing point may result in local freezing that decreases the bond between concrete and steel reinforcement. ACI Committee 306 suggests that a reinforcing bar having a cross-sectional area of about 650 mm² (1 in.²) should have a temperature of at least -12°C (10°F) immediately before being surrounded by fresh concrete at a temperature of at least 13°C (55°F). Caution and additional study are required before definitive recommendations can be formulated. See ACI 306 for additional information.

When slab finishing is completed, insulating blankets or other insulation must be placed on top of the slab to ensure that proper curing temperatures are maintained. The insulation value (R) necessary to maintain the concrete surface temperature of walls and slabs above-ground at 10°C (50°F) or above for 7 days may be estimated from Fig. 14-16. To maintain a temperature for longer periods, more insulation is required. ACI 306 has additional graphs and tables for slabs placed on ground at

Table 14-3.

A. Recommended Duration of Concrete Temperature in Cold Weather–Air-Entrained Concrete*

<table>
<thead>
<tr>
<th>Service category</th>
<th>Protection from early-age freezing</th>
<th>For safe stripping strength</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Conventional concrete,** days</td>
<td>High-early strength concrete,† days</td>
</tr>
<tr>
<td>No load, not exposed;†</td>
<td>2</td>
<td>1</td>
</tr>
<tr>
<td>No load, exposed, but later has favorable moist-curing</td>
<td>3</td>
<td>2</td>
</tr>
<tr>
<td>Partial load, exposed</td>
<td>6</td>
<td>4</td>
</tr>
<tr>
<td>Fully stressed, exposed</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Table 14-3.

B. Recommended Duration of Concrete Temperature for Fully Stressed, Exposed, Air-Entrained Concrete

<table>
<thead>
<tr>
<th>Required percentage of standard-cured 28-day strength</th>
<th>Days at 10°C (50°F)</th>
<th>Days at 21°C (70°F)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>I or GU</td>
<td>II or MS</td>
</tr>
<tr>
<td>50</td>
<td>6 9 3</td>
<td></td>
</tr>
<tr>
<td>65</td>
<td>11 14 5</td>
<td></td>
</tr>
<tr>
<td>85</td>
<td>21 28 16</td>
<td></td>
</tr>
<tr>
<td>95</td>
<td>29 35 26</td>
<td></td>
</tr>
</tbody>
</table>

* Adapted from Tables 5.1 and 5.3 of ACI 306. Cold weather is defined as that in which the average daily temperature is less than 4°C (40°F) for 3 successive days except that if temperatures above 10°C (50°F) occur during at least 12 hours in any day, the concrete should no longer be regarded as winter concrete and normal curing practice should apply. For recommended concrete temperatures, see Table 14-1. For concrete that is not air entrained, ACI Committee 306 states that protection for durability should be at least twice the number of days listed in Table A.

Part B was adapted from Table 6.8 of ACI 306R-88. The values shown are approximations and will vary according to the thickness of concrete, mix proportions, etc. They are intended to represent the ages at which supporting forms can be removed. For recommended concrete temperatures, see Table 14-1.

** Made with ASTM Type I, II, GU, or MS portland cement.
† Made with ASTM Type III or HE cement, or an accelerator, or an extra 60 kg/m³ (100 lb/yard³) of cement.
‡ “Exposed” means subject to freezing and thawing.
Corners and edges are particularly vulnerable during cold weather. As a result, the thickness of insulation for these areas, especially on columns, should be about three times the thickness that is required to maintain the same for walls or slabs. On the other hand, if the ambient temperature rises much above the temperature assumed in selecting insulation values, the temperature of the concrete may become excessive. This increases the probability of thermal shock and cracking when forms are removed. Temperature readings of insulated concrete should therefore be taken at regular intervals and should not vary from ambient air temperatures by more than the values given in ACI 306. In addition, insulated concrete temperatures should not be allowed to rise much above 27°C (80°F). In case of a sudden increase in concrete temperature, up to say 35°C (95°F), it may be necessary to remove some of the insulation or loosen the formwork. The maximum temperature differential between the concrete interior and the concrete surface should be about 20°C (35°F) to minimize cracking. The weather forecast

Fig. 14-16. Thermal resistance ($R$) of insulation required to maintain the concrete surface temperature of walls and slabs aboveground at 10°C (50°F) or above for 7 days. Concrete temperature as placed: 10°C (50°F). Maximum wind velocity: 24 km/h (15 mph). Note that in order to maintain a certain minimum temperature for a longer period of time, more insulation or a higher $R$ value is required (adapted from ACI 306).
should be checked and appropriate action taken for expected temperature changes.

Columns and walls should not be cast on foundations at temperatures below 0°C (32°F) because chilling of concrete in the bottom of the column or wall will retard strength development. Concrete should not be placed on any surface that would lower the temperature of the as-placed concrete below the minimum values shown on Line 4 in Table 14-1.

**ENCLOSURES**

Heated enclosures are very effective for protecting concrete in cold weather, but are probably the most expensive too (Fig. 14-17). Enclosures can be of wood, canvas tarpaulins, or polyethylene film (Fig. 14-18). Prefabricated, rigid-plastic enclosures are also available. Plastic enclo-

---

**Table 14-4. Insulation Values of Various Materials**

<table>
<thead>
<tr>
<th>Material</th>
<th>Density kg/m³ (lb/ft³)</th>
<th>Thermal resistance, R, for 10-mm (1-in.) thickness of material,* (m² · K)/W ([°F · hr · ft²]/Btu)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Expanded polyurethane</td>
<td>24 (1.5)</td>
<td>0.438 (6.25)</td>
</tr>
<tr>
<td>Expanded polystyrene, extruded smooth-skin surface</td>
<td>29 to 56 (1.8 to 3.5)</td>
<td>0.347 (5.0)</td>
</tr>
<tr>
<td>Expanded polystyrene, extruded cut-cell surface</td>
<td>29 (1.8)</td>
<td>0.377 (4.4)</td>
</tr>
<tr>
<td>Glass fiber, organic bonded</td>
<td>64 to 144 (4 to 9)</td>
<td>0.277 (4.0)</td>
</tr>
<tr>
<td>Expanded polystyrene, molded beads</td>
<td>16 (1)</td>
<td>0.247 (3.85)</td>
</tr>
<tr>
<td>Mineral fiber with resin binder</td>
<td>240 (15)</td>
<td>0.239 (3.45)</td>
</tr>
<tr>
<td>Mineral fiberboard, wet felted</td>
<td>256 to 272 (16 to 17)</td>
<td>0.204 (2.94)</td>
</tr>
<tr>
<td>Vegetable fiberboard sheathing</td>
<td>288 (18)</td>
<td>0.182 (2.64)</td>
</tr>
<tr>
<td>Cellular glass</td>
<td>136 (8.5)</td>
<td>0.201 (2.86)</td>
</tr>
<tr>
<td>Laminated paperboard</td>
<td>480 (30)</td>
<td>0.193 (2.00)</td>
</tr>
<tr>
<td>Particle board (low density)</td>
<td>590 (37)</td>
<td>0.128 (1.85)</td>
</tr>
<tr>
<td>Plywood</td>
<td>545 (34)</td>
<td>0.087 (1.24)</td>
</tr>
<tr>
<td><strong>Loose fill</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Wood fiber, soft woods</td>
<td>32 to 56 (2.0 to 3.5)</td>
<td>0.231 (3.33)</td>
</tr>
<tr>
<td>Perlite (expanded)</td>
<td>80 to 128 (5.0 to 8.0)</td>
<td>0.187 (2.70)</td>
</tr>
<tr>
<td>Vermiculite (exfoliated)</td>
<td>64 to 96 (4.0 to 6.0)</td>
<td>0.157 (2.27)</td>
</tr>
<tr>
<td>Vermiculite (exfoliated)</td>
<td>112 to 131 (7.0 to 8.2)</td>
<td>0.148 (2.13)</td>
</tr>
<tr>
<td>Sawdust or shavings</td>
<td>128 to 240 (8.0 to 15.0)</td>
<td>0.154 (2.22)</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Material</th>
<th>Material thickness mm (in.)</th>
<th>Thermal resistance, R, for thickness of material,* (m² · K)/W ([°F · hr · ft²]/Btu)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mineral fiber blanket, fibrous form (rock, slag, or glass)</td>
<td>5 to 32 kg/m³ (0.3 to 2 lb/ft³)</td>
<td>1.23 (7)</td>
</tr>
<tr>
<td>50 to 70 (2 to 2.75)</td>
<td></td>
<td>1.90 (11)</td>
</tr>
<tr>
<td>75 to 85 (3 to 3.5)</td>
<td></td>
<td>3.34 (19)</td>
</tr>
<tr>
<td>90 to 165 (5.5 to 6.5)</td>
<td></td>
<td>1.90 (11)</td>
</tr>
<tr>
<td>Mineral fiber loose fill (rock, slag, or glass)</td>
<td>10 to 32 kg/m³ (0.6 to 2 lb/ft³)</td>
<td>3.34 (19)</td>
</tr>
<tr>
<td>95 to 125 (3.75 to 5)</td>
<td></td>
<td>3.87 (22)</td>
</tr>
<tr>
<td>165 to 220 (6.5 to 8.75)</td>
<td></td>
<td>5.28 (30)</td>
</tr>
<tr>
<td>190 to 250 (7.5 to 10)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>260 to 350 (10.25 to 13.75)</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>


R values are the reciprocal of U values (conductivity).
the minimum required on Line 4 in Table 14-1, additional insulating material, or material with a higher \( R \) value, should be applied. Corners and edges of concrete are most vulnerable to freezing. In view of this, temperatures at these locations should be checked often.

The thermal resistance (\( R \)) values for common insulating materials are given in Table 14-4. For maximum efficiency, insulating materials should be kept dry and in close contact with concrete or formwork.

Concrete pavements can be protected from cold weather by spreading 300 mm (1 ft) or more of dry straw or hay on the surface for insulation. Tarpaulins, polyethylene film, or waterproof paper should be used as a protective cover over the straw or hay to make the insulation more effective and prevent it from blowing away. The straw or hay should be kept dry or its insulation value will drop considerably.

Stay-in-place insulating concrete forms became popular for cold-weather construction in the 1990s (Fig. 14-19).
Forms built for repeated use of tent can be economically insulated with commercial blanket or batt insulation. The insulation should have a tough moisture-proof covering to withstand handling abuse and exposure to the weather. Rigid insulation can also be used (Fig. 14-21).

Insulating blankets for construction are made of fiberglass, sponge rubber, open-cell polyurethane foam, vinyl foam, mineral wool, or cellulose fibers. The outer covers are made of canvas, woven polyethylene, or other tough fabrics that will withstand rough handling. The R value for a typical insulating blanket is about 1.0 m²·°C/W (5.6°F·hr·ft²) /Btu. (IMG12274)

Fig. 14-21. With air temperatures down to -23°C (-10°F), concrete was cast in this insulated column form made of 19-mm (¾-in.) high-density plywood inside, 25-mm (1-in.) rigid polystyrene in the middle, and 13-mm (½-in.) rough plywood outside. R value: 1.0 m²·°C/W (5.6°F·hr·ft²) /Btu. (IMG12274)

14-20). Forms built for repeated use often can be economically insulated with commercial blanket or batt insulation. The insulation should have a tough moisture-proof covering to withstand handling abuse and exposure to the weather. Rigid insulation can also be used (Fig. 14-21).

Insulating blankets for construction are made of fiberglass, sponge rubber, open-cell polyurethane foam, vinyl foam, mineral wool, or cellulose fibers. The outer covers are made of canvas, woven polyethylene, or other tough fabrics that will withstand rough handling. The R value for a typical insulating blanket is about 1.2 m²·°C/W for 50 to 70 mm thickness, 7°F·hr·ft²)/Btu, but since R values are not marked on the blankets, their effectiveness should be checked with a thermometer. If necessary, they can be used in two or three layers to attain the desired insulation.

HEATERS

Three types of heaters are used in cold-weather concrete construction: direct fired, indirect fired, and hydronic systems (Figs. 14-22 to 14-25). Indirect-fired heaters are vented to remove the products of combustion. Where heat is to be supplied to the top of fresh concrete— for example, a floor slab—vented heaters are required. Carbon dioxide

Fig. 14-22. Two types of air heaters.

Fig. 14-23. An indirect-fired heater. Notice vent pipe that carries combustion gases outside the enclosure. (IMG12275)
(CO₂) in the exhaust must be vented to the outside and prevented from reacting with the fresh concrete (Fig. 14-23). Direct-fired units can be used to heat the enclosed space beneath concrete placed for a floor or a roof deck (Fig. 14-24).

Hydronic systems transfer heat by circulating a glycol/water solution in a closed system of pipes or hoses (see Fig. 14-25). These systems transfer heat more efficiently than forced air systems without the negative effects of exhaust gases and drying of the concrete from air movement. The specific heat of water/glycol solutions is more than six times greater than air. As a result, hydronic heaters can deliver very large quantities of heat at low temperature differentials of 5°C (10°F) or less between the heat transfer hose and the concrete. Cracking and curling induced by temperature gradients within the concrete are almost eliminated along with the danger of accidentally overheating the concrete and damaging long-term strength gain.

Typical applications for hydronic systems include thawing and preheating subgrades. They are also used to cure elevated and on-grade slabs, walls, foundations, and columns. To heat a concrete element, hydronic heating hoses are usually laid on or hung adjacent to the structure and covered with insulated blankets and sometimes plastic sheets. Usually, construction of temporary enclosures is not necessary. Hydronic systems can be used over areas much larger than would be practical to enclose. If a heated enclosure is necessary for other work, hydronic hoses can be sacrificed (left under a slab on grade) to make the slab a radiant heater for the structure built above (Grochoski 2000).

Any heater burning a fossil fuel produces carbon dioxide (CO₂); this gas will combine with calcium hydroxide on the surface of fresh concrete to form a weak layer of calcium carbonate that interferes with cement hydration (Kauer and Freeman 1955). The result is a soft, chalky surface that will dust under traffic. Depth and degree of carbonation depend on concentration of CO₂, curing temperature, humidity, porosity of the concrete, length of exposure, and method of curing. Direct-fired heaters, therefore, should not be permitted to heat the air over concreting operations—at least until 24 hours have elapsed. In addition, the use of gasoline-powered construction equipment should be restricted in enclosures during that time. If unvented heaters are used, immediate wet curing or the use of a curing compound will minimize carbonation.

Carbon monoxide (CO), another product of combustion, is not usually a problem unless the heater is using recirculated air. Four hours of exposure to 200 parts per million of CO will produce headaches and nausea. Three hours of exposure to 600 ppm can be fatal. The American National Standard Safety Requirements for Temporary and Portable Space Heating Devices and Equipment Used in the Construction Industry (ANSI A10.10) limits concentrations of CO to 50 ppm at worker breathing levels. The standard also establishes safety rules for ventilation and the stability, operation, fueling, and maintenance of heaters.
A salamander is an inexpensive combustion heater without a fan that discharges its combustion products into the surrounding air; heating is accomplished by radiation from its metal casing. Salamanders are fueled by coke, oil, wood, or liquid propane. They are but one form of a direct-fired heater. A primary disadvantage of salamanders is the high temperature of their metal casing, a definite fire hazard. Salamanders should be placed so that they will not overheat formwork or enclosure materials. When placed on floor slabs, they should be elevated to avoid scorching the concrete.

Some heaters burn more than one type of fuel. The approximate heat values of fuels are as follows:

<table>
<thead>
<tr>
<th>Fuel Type</th>
<th>Heat Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>No. 1 fuel oil</td>
<td>37,700 kJ/L (135,000 Btu/gal)</td>
</tr>
<tr>
<td>Kerosene</td>
<td>37,400 kJ/L (134,000 Btu/gal)</td>
</tr>
<tr>
<td>Gasoline</td>
<td>35,725 kJ/L (128,000 Btu/gal)</td>
</tr>
<tr>
<td>Liquid-propane gas</td>
<td>25,500 kJ/L (91,500 Btu/gal)</td>
</tr>
<tr>
<td>Natural gas</td>
<td>37,200 kJ/m³ (1,000 Btu/ft³)</td>
</tr>
</tbody>
</table>

The output rating of a portable heater is usually the heat content of the fuel consumed per hour. A rule of thumb is that about 134,000 kJ are required for each 100 m³ (36,000 Btu for 10,000 ft³) of air to develop a 10°C (20°F) temperature rise.

Electricity can also be used to cure concrete in winter. The use of large electric blankets equipped with thermostats is one method. The blankets can also be used to thaw subgrades or concrete foundations.

Use of electrical resistance wires that are cast into the concrete is another method. The power supplied is under 50 volts, and from 7.0 to 23.5 MJ (1.5 to 5 kilowatt-hours) of electricity per cubic meter (cubic yard) of concrete is required, depending on the circumstances. The method has been used in the Montreal, Quebec, area for many years. Where electrical resistance wires are used, insulation should be included during the initial setting period. If insulation is removed before the recommended time, the concrete should be covered with an impervious sheet and the power continued for the required time.

Steam is another source of heat for winter concreting. Live steam can be piped into an enclosure or supplied through radiant heating units. In choosing a heat source, it must be remembered that the concrete itself supplies heat through hydration of cement; this is often enough for curing needs if the heat can be retained within the concrete with insulation.

**DURATION OF HEATING**

After concrete is in place, it should be protected and kept at the recommended temperatures listed on Line 4 of Table 14-1. These curing temperatures should be maintained until sufficient strength is gained to withstand exposure to low temperatures, anticipated environment, and construction and service loads. The length of protection required to accomplish this will depend on the cement type and amount, whether accelerating admixtures were used, and the loads that must be carried. Recommended minimum periods of protection are given in Table 14-3. The duration of heating structural concrete that requires the attainment of full service loading before forms and shores are removed should be based on the adequacy of in-place compressive strengths rather than an arbitrary time period. If no data are available, a conservative estimate of the length of time for heating and protection can be made using Table 14-3.

**Moist Curing**

Strength gain stops when moisture required for curing is no longer available. Concrete retained in forms or covered with insulation seldom loses enough moisture at 5°C to 15°C (40°F to 55°F) to impair curing. However, a positive means of providing moist curing is needed to offset drying from low wintertime humidities and heaters used in enclosures during cold weather.

Live steam exhausted into an enclosure around the concrete is an excellent method of curing because it provides both heat and moisture. Steam is especially practical in extremely cold weather because the moisture provided offsets the rapid drying that occurs when very cold air is heated.

Liquid membrane-forming compounds can be used for early curing of concrete surfaces within heated enclosures.

**Terminating the Heating Period**

Rapid cooling of concrete at the end of the heating period should be avoided. Sudden cooling of the concrete surface while the interior is still warm may cause thermal cracking, especially in massive sections such as bridge piers, abutments, dams, and large structural members; thus cooling should be gradual. A safe temperature differential between a concrete wall and the ambient air temperature can be obtained from ACI 306R-88. The maximum uniform drop in temperature throughout the first 24 hours after the end of protection should not be more than the amounts given in Table 14-2. Gradual cooling can be accomplished by lowering the heat or by simply shutting off the heat and allowing the enclosure to cool to outside ambient air temperature.

**FORM REMOVAL AND RESHORING**

It is good practice in cold weather to leave forms in place as long as possible. Even within heated enclosures, forms serve to distribute heat more evenly and help prevent drying and local overheating.

If the curing temperatures listed on Line 4 of Table 14-1 are maintained, Table 14-3A can be used to determine the minimum time in days that vertical support for forms
should be left in place. Before shores and forms are removed, fully stressed structural concrete should be tested to determine if in-place strengths are adequate, rather than waiting an arbitrary time period. In-place strengths can be monitored using one of the following: (1) field-cured cylinders (ASTM C31 or AASHTO T 23); (2) probe penetration tests (ASTM C803); (3) cast-in-place cylinders (ASTM C873); (4) pullout testing (ASTM C900); or (5) maturity testing (ASTM C1074). Many of these tests are indirect methods of measuring compressive strength; they require correlation in advance with standard cylinders before estimates of in-place strengths can be made.

If in-place compressive strengths are not documented, Table 14-3B lists conservative time periods in days to achieve various percentages of the standard laboratory-cured 28-day strength. The engineer issuing project drawings and specifications in cooperation with the formwork contractor must determine what percentage of the design strength is required (see ACI Committee 306). Side forms can be removed sooner than shoring and temporary falsework (ACI Committee 347).

MATURITY CONCEPT

The maturity concept is based on the principle that strength gain in concrete is a function of curing time and temperature. The maturity concept, as described in ACI 306R-88 and ASTM C1074 can be used to evaluate strength development. Two maturity methods to estimate the in-place concrete strength are shown in Table 14-5. The first method is based on the Nurse-Saul function, also called Time-Temperature Factor method. This method is simple and very popular. However, it doesn’t recognize that maturity increases disproportionately at elevated temperatures and that this increase depends on the type(s) of cementitious materials used and the water-to-cementitious materials ratio. Therefore, the Time-Temperature Factor method typically underestimates the strength development at elevated temperatures. The “Equivalent Age” maturity function is based on the Arrhenius equation; this function presents maturity in terms of equivalent age of curing at a specified temperature.

The Time-Temperature Factor method presents maturity in terms of °C·hr. Most maturity equipment uses a datum temperature of 0°C, which further simplifies the calculation. Given this simplification, maturity is typically calculated using metric units.

To monitor the strength development of concrete in place using the maturity concept, the following information must be available:

1. The strength-maturity relationship of the concrete used in the structure. The results of compressive strength tests at various ages on a series of cylinders made of a concrete similar to that used in the structure; this must be done to develop a strength-maturity curve.

These cylinders are cured in a laboratory at 23°C ± 2°C (73°F ± 3°F).

2. A time-temperature record of the concrete in place. Temperature readings are obtained by placing expendable thermistors or thermocouples at varying depths in the concrete. The location giving the lowest values provides the series of temperature readings to be used in the computation (Fig. 14-26).

See Fig. 14-27 for an example using the maturity concept.

Before construction begins, a calibration curve is drawn plotting the relationship between compressive strength and the maturity factor for a series of test cylinders (of the particular concrete mixture proportions) cured in a laboratory and tested for strength at successive ages.

The maturity concept is not precise and has some limitations. But, the concept is useful in checking the curing of concrete and estimating strength in relation to time and temperature. It assumes that all other factors affecting concrete strength have been properly controlled. With these limitations in mind, the maturity concept has gained greater acceptance for representing the compressive strength of the concrete for removal of shoring or opening a pavement to traffic; but it is no substitute for quality control and proper concreting practices (Gajda 2007; Malhotra 1974; and ACI Committee 347).

<table>
<thead>
<tr>
<th>Table 14-5. Time-Temperature Factor and Equivalent Age Maturity Equations</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Time-Temperature Factor</strong></td>
</tr>
<tr>
<td>$M = \sum_{0}^{t} \frac{(T - T_0) \Delta t}{\Delta t}$</td>
</tr>
<tr>
<td>$T$ = average concrete temperature, °C, during the time interval $\Delta t$</td>
</tr>
<tr>
<td>$T_0$ = datum temperature (usually taken to be 0°C)</td>
</tr>
<tr>
<td>$t$ = elapsed time, hours</td>
</tr>
<tr>
<td>$\Delta t$ = time intervals, hours</td>
</tr>
<tr>
<td><strong>Equivalent Age</strong></td>
</tr>
<tr>
<td>$t_e = \sum_{0}^{t} \frac{E}{R(T - T_r)} \Delta t$</td>
</tr>
<tr>
<td>$E$ = apparent activation energy, J/mol (see ASTM C1074 for typical values)</td>
</tr>
<tr>
<td>$R$ = universal gas constant, 8.314 J/mol-K</td>
</tr>
<tr>
<td>$T$ = average concrete temperature, Kelvin, during the time interval $\Delta t$</td>
</tr>
<tr>
<td>$T_r$ = absolute reference temperature, Kelvin</td>
</tr>
<tr>
<td>$\Delta t$ = time intervals, hours</td>
</tr>
</tbody>
</table>
REFERENCES

ACI Committee 306, Cold-Weather Concreting, ACI 306R-88, reapproved 2002, American Concrete Institute, Farmington Hills, Michigan, 1988, 23 pages.

ACI Committee 347, Guide to Formwork for Concrete, ACI 347-04, American Concrete Institute, Farmington Hills, Michigan, 2004, 32 pages.


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McNeese, D. C., “Early Freezing of Non-Air-Entrained Concrete,” Journal of the American Concrete Institute Proceedings, vol. 49, American Concrete Institute, Farmington Hills, Michigan, December 1952, pages 293 to 300.

NRMCA, Cold Weather Ready Mixed Concrete, Publication No. 130, National Ready Mixed Concrete Association, Silver Spring, Maryland, 1968.


Concrete changes slightly in volume for various reasons, and understanding the nature of these changes is useful in planning or analyzing concrete work. If concrete were free of any restraints to deform, normal volume changes would be of little consequence; but since concrete in service is usually restrained by foundations, subgrades, reinforcement, or connecting members, significant stresses can develop. This is particularly true of tensile stresses.

Cracks develop because concrete is relatively weak in tension but quite strong in compression. Controlling the variables that affect volume changes can minimize high stresses and cracking. Tolerable crack widths should be considered in the structural design.

Volume change is defined merely as an increase or decrease in volume. Most commonly, the subject of concrete volume changes deals with linear expansion and contraction due to temperature and moisture cycles. But chemical effects such as carbonation shrinkage, sulfate attack, and the disruptive expansion of alkali-aggregate reactions also cause volume changes. Also, creep is a volume change or deformation caused by sustained stress or load. Equally important is the elastic or inelastic change in dimensions or shape that occurs instantaneously under applied load.

For convenience, the magnitude of volume changes is generally stated in linear rather than volumetric units. Changes in length are often expressed as a coefficient of length in parts per million, or simply as millionths. It is applicable to any length unit (for example, m/m or ft/ft); one millionth is 0.000001 m/m (0.000001 in./in.) and 600 millionths is 0.000600 m/m (0.000600 in./in.). Change of length can also be expressed as a percentage; thus 0.06% is the same as 0.000600, which incidentally is approximately the same as 6 mm per 10 m (¾ in. per 100 ft). The volume changes that ordinarily occur in concrete are small, ranging in length change from perhaps 10 millionths up to about 1000 millionths.

**EARLY AGE VOLUME CHANGES**

The volume of concrete begins to change shortly after it is cast. Early volume changes, within 24 hours, can influence the volume changes (such as drying shrinkage) and crack formation in hardened concrete, especially for low water to cement ratio concrete. Following are discussions on various forms of early volume change:

**Chemical Shrinkage**

Chemical shrinkage refers to the reduction in absolute volume of solids and liquids in paste resulting from cement hydration. The absolute volume of hydrated cement products is less than the absolute volume of cement and water before hydration. This change in volume of cement paste during the plastic state is illustrated by the first two bars in Fig. 15-1. This does not include air bubbles from mixing. Chemical shrinkage continues to occur at a microscopic scale as long as cement hydrates. After initial set, the paste cannot deform as much as when it was in a plastic state.
Therefore, further hydration and chemical shrinkage is compensated by the formation of voids in the microstructure (Fig. 15-1). Most of this volume change is internal and does not significantly change the visible external dimensions of a concrete element.

The amount of volume change due to chemical shrinkage can be estimated from the hydrated cement phases and their crystal densities or it can be determined by physical test as illustrated in Fig. 15-2. The Japan Concrete Institute has a test method for chemical shrinkage of cement paste (Tazawa 1999). An example of long-term chemical shrinkage for portland cement paste is illustrated in Fig. 15-3. Early researchers sometimes referred to chemical shrinkage as the absorption of water during hydration (Powers 1935). Le Chatelier (1900) was the first to study chemical shrinkage of cement pastes.

**Autogenous Shrinkage**

Autogenous shrinkage is the macroscopic volume reduction (visible dimensional change) of cement paste, mortar, or concrete caused by cement hydration. The macroscopic volume reduction of autogenous shrinkage is much less than the absolute volume reduction of chemical shrinkage because of the rigidity of the hardened paste structure. Chemical shrinkage is the driving force behind autogenous shrinkage. The relationship between autogenous shrinkage and chemical shrinkage is illustrated in Figs. 15-1, 15-4, and 15-5. Some researchers and organizations consider that

![Fig. 15-2. Test for chemical shrinkage of cement paste showing flask for cement paste and pipet for absorbed water measurement.](image1)

![Fig. 15-3. Chemical shrinkage of cement paste (Tazawa 1999).](image2)

![Fig. 15-4. Relationship between autogenous shrinkage and chemical shrinkage of cement paste at early ages (Hammer 1999).](image3)

![Fig. 15-5. Volumetric relationship between subsidence, bleed water, chemical shrinkage, and autogenous shrinkage. Only autogenous shrinkage after initial set is shown. Not to scale.](image4)
autogenous shrinkage starts at initial set while others evaluate autogenous shrinkage from time of placement.

When external water is available, autogenous shrinkage cannot occur. When external water is not available, cement hydration consumes pore water resulting in self desiccation of the paste and a uniform reduction of volume (Copeland and Bragg 1955). Autogenous shrinkage increases with a decrease in water to cement ratio and with an increase in the amount of cement paste. Normal concrete has negligible autogenous shrinkage; however, autogenous shrinkage is most prominent in concrete with a water to cement ratio under 0.42 (Holt 2001). High-strength, low water to cement ratio (0.30) concrete can experience 200 to 400 millionths of autogenous shrinkage. Autogenous shrinkage can be half that of drying shrinkage for concretes with a water to cement ratio of 0.30.

Recent use of high performance, low water to cement ratio concrete in bridges and other structures has renewed interest in autogenous shrinkage to control crack development. Concretes susceptible to large amounts of autogenous shrinkage should be cured with external water for at least 7 days to help control crack development. Fogging should be provided as soon as the concrete is cast. The hydration of supplementary cementing materials also contributes to autogenous shrinkage, although at different levels than portland cement. In addition to adjusting paste content and water to cement ratios, autogenous shrinkage can be reduced by using shrinkage reducing admixtures or internal curing techniques. Some cementitious systems may experience autogenous expansion. Tazawa (1999) and Holt (2001) review techniques to control autogenous shrinkage.

Test methods for autogenous shrinkage and expansion of cement paste, mortar, and concrete and tests for autogenous shrinkage stress of concrete are presented by Tazawa (1999).

**Subsidence**

Subsidence refers to the vertical shrinkage of fresh cementitious materials before initial set. It is caused by bleeding (settlement of solids relative to liquids), air voids rising to the surface, and chemical shrinkage. Subsidence is also called settlement shrinkage. Subsidence of well-consolidated concrete with minimal bleed water is insignificant. The relationship between subsidence and other shrinkage mechanisms is illustrated in Fig. 15-5. Excessive subsidence is often caused by a lack of consolidation of fresh concrete. Excessive subsidence over embedded items, such as supported steel reinforcement, can result in cracking over embedded items. Concretes made with air entrainment, sufficient fine materials, and low water contents will minimize subsidence cracking. Also, plastic fibers have been reported to reduce subsidence cracking (Suprenant and Malisch 1999).

**Plastic Shrinkage**

Plastic shrinkage refers to volume change occurring while the concrete is still fresh, before hardening. It is usually observed in the form of plastic shrinkage cracks occurring before or during finishing (Fig. 15-6). The cracks often resemble tears in the surface. Plastic shrinkage results from a combination of chemical and autogenous shrinkage and rapid evaporation of moisture from the surface that exceeds the bleeding rate. Plastic shrinkage cracking can be controlled by minimizing surface evaporation through use of fogging, wind breaks, shading, plastic sheet covers, wet burlap, spray-on finishing aids (evaporation retarders), and plastic fibers.

**Swelling**

Concrete, mortar, and cement paste swell in the presence of external water. When water drained from capillaries by chemical shrinkage is replaced by external water, the volume of the concrete mass increases. As there is no self desiccation, there is no autogenous shrinkage. External water can come from wet curing or submersion. Swelling occurs due to a combination of crystal growth, absorption of water, and osmotic pressure. The swelling is not large, only about 50 millionths at early ages (Fig. 15-7). When the

![Fig. 15-6. Plastic shrinkage cracks resemble tears in fresh concrete. (IMG12283)](image)

![Fig. 15-7. Early age swelling of 100 x 100 x 375-mm (4 x 4 x 15-in.) concrete specimens cured under water (Alticin 1999).](image)
external water source is removed, autogenous shrinkage and drying shrinkage reverse the volume change.

**Early Thermal Expansion**

As cement hydrates, the exothermic reaction provides a significant amount of heat. In large elements the heat is retained, rather than dissipated as happens with thin elements. This temperature rise, occurring over the first few hours and days, can induce a small amount of expansion that counteracts autogenous and chemical shrinkage (Holt 2001).

**MOISTURE CHANGES (DRYING SHRINKAGE) OF HARDENED CONCRETE**

Hardened concrete expands slightly with a gain in moisture and contracts with a loss in moisture. The effects of these moisture cycles are illustrated schematically in Fig. 15-8. Specimen A represents concrete stored continuously in water from time of casting. Specimen B represents the same concrete exposed first to drying in air and then to alternate cycles of wetting and drying. For comparative purposes, it should be noted that the swelling that occurs during continuous wet storage over a period of several years is usually less than 150 millionths; this is about one-fourth of the shrinkage of air-dried concrete for the same period. Fig. 15-9 illustrates swelling of concretes wet cured for 7 days following by shrinkage when sealed or exposed to air drying. Autogenous shrinkage reduces the volume of the sealed concretes to a level about equal to the amount of swelling at 7 days. Note that the concretes wet cured for 7 days had less shrinkage due to drying and autogenous effects than the concrete that had no water curing. This illustrates the importance of early, wet curing to minimize shrinkage (Aïtcin 1999).

Tests indicate that the drying shrinkage of small, plain concrete specimens (without reinforcement) ranges from about 400 to 800 millionths when exposed to air at 50% humidity. Concrete with a unit drying shrinkage of 550 millionths shortens about the same amount as the thermal contraction caused by a decrease in temperature of 55°C (100°F). Preplaced aggregate concrete has a drying shrinkage of 200 to 400 millionths; this is considerably less than normal concrete due to point-to-point contact of aggregate particles in preplaced aggregate concrete. The drying shrinkage of structural lightweight concrete ranges from slightly less than to 30 percent more than that of normal-density concrete, depending on the type of aggregate used.

The drying shrinkage of reinforced concrete is less than that for plain concrete, the difference depending on the amount of reinforcement. Steel reinforcement restricts but does not prevent drying shrinkage. In reinforced concrete structures with normal amounts of reinforcement, drying shrinkage is assumed to be 200 to 300 millionths. Similar values are found for slabs on ground restrained by subgrade.

For many outdoor applications, concrete reaches its maximum moisture content in winter; so in winter the volume changes due to increase in moisture content and the decrease in average temperature tend to offset each other.

The amount of moisture in concrete is affected by the relative humidity of the ambient air. The free moisture content of concrete elements after drying in air at relative
humidities of 50% to 90% for several months is about 1% to 2% by weight of the concrete; the actual amount depends on the concrete’s constituents, original water content, drying conditions, and the size and shape of the concrete element.

After concrete has dried to a constant moisture content at one relative humidity condition, a decrease in humidity causes it to lose moisture while an increase causes it to gain moisture. The concrete shrinks or swells with each such change in moisture content due primarily to responses of the cement paste to moisture changes. Most aggregates show little response to changes in moisture content, although there are a few aggregates that swell or shrink in response to such changes.

As drying takes place, concrete shrinks. Where there is no restraint, movement occurs freely and no stresses or cracks develop (Fig. 15-10a top). If the tensile stress that results from restrained drying shrinkage exceeds the tensile strength of the concrete, cracks can develop (Fig. 15-10a bottom). Random cracks may develop if joints are not properly provided and the concrete element is restrained from shortening (Fig. 15-10b). Contraction joints for slabs on ground should be spaced at distances of 24 to 36 times the slab thickness to control random cracks (Fig. 15-10c). Joints in walls are equally important for crack control (Fig. 15-10d). Fig. 15-11 illustrates the relationship between drying rate at different depths, drying shrinkage, and mass loss for normal-density concrete (Hanson 1968).

Shrinkage may continue for a number of years, depending on the size and shape of the concrete mass. The rate and ultimate amount of shrinkage are usually smaller for large masses of concrete than for small masses; on the other hand, shrinkage continues longer for large masses. Higher volume-to-surface ratios (larger elements) experience lower shrinkage as shown in Fig. 15-12.

The rate and amount of drying shrinkage for small concrete specimens made with various cements are shown in Fig. 15-13. Specimens were initially moist-cured for 14 days at 21°C (70°F), then stored for 38 months in air at the same temperature and 50% relative humidity. Shrinkage recorded at the age of 38 months ranged from 600 to 790 millionths. An average of 34% of this shrinkage occurred within the first month. At the end of 11 months an average of 90% of the 38-month shrinkage had taken place.

Fig. 15-10. (a) Illustration showing no crack development in concrete that is free to shrink (slab on rollers); however, in reality a slab on ground is restrained by the subbase (or other elements) creating tensile stresses and cracks. (b) Typical shrinkage cracks in a slab on ground. (c) A properly functioning contraction joint controls the location of shrinkage cracking. (d) Contraction joints in the slabs and walls shown will minimize the formation of cracks. (IMG12279, IMG12281, IMG12282)
Design and Control of Concrete Mixtures

Fig. 15-11. Relative humidity distribution at various depths, drying shrinkage, and mass loss of 150 x 300-mm (6 x 12-in.) cylinders moist-cured for 7 days followed by drying in laboratory air at 23°C (73°F) and 50% RH (Hanson 1968).

Normal-density concrete

Cement content: 270 kg/m³ (454 lb/yd³)
w/c ratio: 0.66

75 mm (3 in.) depth
45 (1/4)
20 (1/4)
6 (1/4)

Shrinkage, millionths

Time of drying, days

Normal-density concrete

Mass loss, kg

Time of drying, days

Normal-density concrete

Fig. 15-12. Drying shrinkage of various sizes of cylindrical specimens made of Elgin, Illinois gravel concrete (Hansen and Mattock 1966).

Fig. 15-13. Results of long-term drying shrinkage tests by the U.S. Bureau of Reclamation. Shrinkage ranged from 600 to 790 millionths after 38 months of drying. The shrinkage of concretes made with air-entraining cements was similar to that for non-air-entrained concretes in this study (Bureau of Reclamation 1947 and Jackson 1955).
Effect of Concrete Ingredients on Drying Shrinkage

The most important controllable factor affecting drying shrinkage is the amount of water per unit volume of concrete. The results of tests illustrating the water content to shrinkage relationship are shown in Fig. 15-14. Shrinkage can be minimized by keeping the water content of concrete as low as possible. This is achieved by keeping the total coarse aggregate content of the concrete as high as possible (minimizing paste content). Use of low slumps and placing methods that minimize water requirements are thus major factors in controlling concrete shrinkage. Any practice that increases the water requirement of the cement paste, such as the use of high slumps (without superplasticizers), excessively high freshly mixed concrete temperatures, high fine-aggregate contents, or use of small-size coarse aggregate, will increase shrinkage. A small amount of water can be added to ready mixed concrete at the jobsite without affecting drying shrinkage properties as long as the additions are within mix specifications (Suprenant and Malisch 2000).

The general uniformity of shrinkage of concretes with different types of cement at different ages is illustrated in Fig. 15-13. However, this does not mean that all cements or cementing materials have similar shrinkage.

Supplementary cementing materials usually have little effect on shrinkage at normal dosages. Fig. 15-15 shows that concretes with normal dosages of selected fly ashes performed similar to the control concrete made with only portland cement as the cementing material.

Aggregates in concrete, especially coarse aggregate, physically restrain the shrinkage of hydrating cement paste. Paste content affects the drying shrinkage of mortar more than that of concrete. Drying shrinkage is also dependent on the type of aggregate. Hard, rigid aggregates are difficult to compress and provide more restraint to shrinkage than softer, less rigid aggregates. As an extreme example, if steel balls were substituted for ordinary coarse aggregate, shrinkage would be reduced 30% or more. Drying shrinkage can also be reduced by avoiding aggregates that have high drying shrinkage properties and aggregates containing excessive amounts of clay. Quartz, granite, feldspar, limestone, and dolomite aggregates generally produce concretes with low drying shrinkages (ACI Committee 224). Steam curing will also reduce drying shrinkage.

Most chemical admixtures have little effect on shrinkage. The use of accelerators such as calcium chloride will increase drying shrinkage of concrete. Despite reductions in water content, some water-reducing admixtures can increase drying shrinkage, particularly those that contain an accelerator to counteract the retarding effect of the admixture. Air entrainment has little or no effect on drying shrinkage. High-range water reducers usually have little effect on drying shrinkage (Fig. 15-16). Drying shrinkage can be evaluated in accordance with ASTM C157 (AASHTO T 160).

Fig. 15-14. Relationship between total water content and drying shrinkage. A large number of mixtures with various proportions is represented within the shaded area of the curves. Drying shrinkage increases with increasing water contents.

Fig. 15-15. Drying shrinkage of fly ash concretes compared to a control mixture. The graphs represent the average of four Class C ashes and six Class F ashes, with the range in drying shrinkage rarely exceeding 0.01 percentage points. Fly ash dosage was 25% of the cementing material (Gebler and Klieger 1986).
Steam curing will also reduce drying shrinkage. Computer software is available to predict the effect of curing and environmental conditions on shrinkage and cracking (FHWA and Transtec 2001). Hedenblad (1997) provides tools to predict the drying of concrete as effected by different curing methods and type of construction.

**TEMPERATURE CHANGES OF HARDENED CONCRETE**

Concrete expands slightly as temperature rises and contracts as temperature falls, although it can expand slightly as free water in the concrete freezes. Temperature changes may be caused by environmental conditions or by cement hydration. An average value for the coefficient of thermal expansion of concrete is about 10 millionths per degree Celsius (5.5 millionths per degree Fahrenheit), although values ranging from 6 to 13 millionths per degree Celsius (3.2 to 7.0 millionths per degree Fahrenheit) have been observed. This amounts to a length change of 5 mm for 10 m of concrete (2/3 in. for 100 ft of concrete) subjected to a rise or fall of 50°C (100°F). The coefficient of thermal expansion for structural low-density (lightweight) concrete varies from 7 to 11 millionths per degree Celsius (3.6 to 6.1 millionths per degree Fahrenheit). The coefficient of thermal expansion of concrete can be determined by AASHTO TP 60.

Thermal expansion and contraction of concrete varies with factors such as aggregate type, cement content, water-cement ratio, temperature range, concrete age, and relative humidity. Of these, aggregate type has the greatest influence.

Table 15-1 shows some experimental values of the thermal coefficient of expansion of concretes made with aggregates of various types. These data were obtained from tests on small concrete specimens in which all factors were the same except aggregate type. In each case, the fine aggregate was of the same material as the coarse aggregate.

The thermal coefficient of expansion for steel is about 12 millionths per degree Celsius (6.5 millionths per degree Fahrenheit), which is comparable to that for concrete. The coefficient for reinforced concrete can be assumed as 11 millionths per degree Celsius (6 millionths per degree Fahrenheit), the average for concrete and steel.

Temperature changes that result in shortening can crack concrete members that are highly restrained by another part of the structure or by ground friction. Consider a long restrained concrete member cast without joints that, after moist curing, is allowed to drop in temperature. As the temperature drops, the concrete wants to shorten, but cannot because it is restrained longitudinally. The resulting tensile stresses cause the concrete to crack. Tensile strength and modulus of elasticity of concrete both may be assumed proportional to the square root of concrete compressive strength. And calculations show that a large enough temp-
Low Temperatures

Concrete continues to contract as the temperature is reduced below freezing. The amount of volume change at subfreezing temperatures is greatly influenced by the moisture content, behavior of the water (physical state—ice or liquid), and type of aggregate in the concrete. In one study, the coefficient of thermal expansion for a temperature range of 24°C to -155°C (75°F to –250°F) varied from $6 \times 10^{-6}$ per °C (3.3 × 10^{-6} per °F) for a low density (light-weight) aggregate concrete to 8.2 × 10^{-6} per °C (4.5 × 10^{-6} per °F) for a sand and gravel mixture. Subfreezing temperatures can significantly increase the compressive and tensile strength and modulus of elasticity of moist concrete. Dry concrete properties are not as affected by low temperatures. In the same study, moist concrete with an original compressive strength of 35 MPa at 24°C (5000 psi at 75°F) achieved over 117 MPa (17,000 psi) at -100°C (-150°F). The same concrete tested oven-dry or at a 50% internal relative humidity had strength increases of only about 20%. The modulus of elasticity for sand and gravel concrete with 50% relative humidity was only 8% higher at -150°C than at 24°C (-250°F than at 75°F), whereas the moist concrete had a 50% increase in modulus of elasticity. Going from 24°C to -170°C (75°F to –250°F), the thermal conductivity of normal-weight concrete also increased, especially for moist concrete. The thermal conductivity of lightweight aggregate concrete is little affected (Monfore and Lentz 1962 and Lentz and Monfore 1966).

High Temperatures

Temperatures greater than 95°C (200°F) that are sustained for several months or even several hours can have significant effects on concrete. The total amount of volume change of concrete is the sum of volume changes of the cement paste and aggregate. At high temperatures, the paste shrinks due to dehydration while the aggregate expands. For normal-aggregate concrete, the expansion of the aggregate exceeds the paste shrinkage resulting in an overall expansion of the concrete. Some aggregates such as expanded shale, andesite, or pumice with low coefficients of expansion can produce a very volume-stable concrete in high-temperature environments (Fig. 15-19). On the other hand, some aggregates undergo extensive and abrupt volume changes at a particular temperature, causing disruption in the concrete. For example, in one study a dolomitic limestone aggregate contained an iron sulfide impurity caused severe expansion, cracking, and disintegration in concrete exposed to a temperature of 150°C (302°F) for four months; at temperatures above and below 150°C (302°F) there was no detrimental expansion (Carette, Painter, and Malhotra 1982). The coefficient of thermal expansion tends to increase with temperature rise.

Besides volume change, sustained high temperatures can also have other, usually irreversible, effects such as a

<table>
<thead>
<tr>
<th>Aggregate type (from one source)</th>
<th>Coefficient of expansion, millionths per °C</th>
<th>Coefficient of expansion, millionths per °F</th>
</tr>
</thead>
<tbody>
<tr>
<td>Quartz</td>
<td>11.9</td>
<td>6.6</td>
</tr>
<tr>
<td>Sandstone</td>
<td>11.7</td>
<td>6.5</td>
</tr>
<tr>
<td>Gravel</td>
<td>10.8</td>
<td>6.0</td>
</tr>
<tr>
<td>Granite</td>
<td>9.5</td>
<td>5.3</td>
</tr>
<tr>
<td>Basalt</td>
<td>8.6</td>
<td>4.8</td>
</tr>
<tr>
<td>Limestone</td>
<td>6.8</td>
<td>3.8</td>
</tr>
</tbody>
</table>

Coefficients of concretes made with aggregates from different sources may vary widely from these values, especially those for gravels, granites, and limestones (Davis 1930).
If stable aggregates are used and strength reduction and the effects on other properties are accounted for in the mix design, high quality concrete can be exposed to temperatures of 90°C to 200°C (200°F to 400°F) for long periods. Some concrete elements have been exposed to temperatures up to 250°C (500°F) for long periods of time; however, special steps should be taken or special materials (such as heat-resistant calcium aluminate cement) should be considered for exposure temperatures greater than 200°C (400°F). Before any structural concrete is exposed to high temperatures (greater than 90°C or 200°F), laboratory testing should be performed to determine the particular concrete’s thermal properties. This will avoid any unexpected distress.

**CURLING (WARPING)**

In addition to horizontal movement caused by changes in moisture and temperature, curling of slabs on ground can be a problem; this is caused by differences in moisture content and temperature between the top and bottom of slabs (Fig. 15-21).

The edges of slabs at the joints tend to curl upward when the surface of a slab is drier or cooler than the bottom. A slab will assume a reverse curl when the surface is wetter or warmer than the bottom. However, enclosed slabs, such as floors on ground, curl only upward. When the edges of an industrial floor slab are curled upward they lose support from the subbase and become a cantilever. Lift-truck traffic passing over joints causes a repetitive vertical deflection that creates a great potential for fatigue cracking in the slab. The amount of vertical upward curl (curling) is small for a short, thick slab.
Curling can be reduced or eliminated by using design and construction techniques that minimize shrinkage differentials and by using techniques described earlier to reduce temperature and moisture-related volume changes. Thickened edges, shorter joint spacings, permanent vapor-impermeable sealers, and large amounts of reinforcing steel placed 50 mm (2 in.) below the surface all help reduce curling (Ytterberg 1987).

ELASTIC AND INELASTIC DEFORMATION

Compression Strain

The series of curves in Fig. 15-22 illustrate the amount of compressive stress and strain that results instantaneously due to loading of unreinforced concrete. With water-cement ratios of 0.50 or less and strains up to 1500 millionths, the upper three curves show that strain is closely proportional to stress; in other words, the concrete is almost elastic. The upper portions of the curves and beyond show that the concrete is inelastic. The curves for high-strength concrete have sharp peaks, whereas those for lower-strength concretes have long and relatively flat peaks. Fig. 15-22 also shows the sudden failure characteristics of higher strength, low water to cement ratio, concrete cylinders.

When load is removed from concrete in the inelastic zone, the recovery line usually is not parallel to the original line for the first load application. Therefore, the amount of permanent set may differ from the amount of inelastic deformation (Fig. 15-23).

The term “elastic” is not favored for general discussion of concrete behavior because frequently the strain may be in the inelastic range. For this reason, the term “instantaneous strain” is often used.

Modulus of Elasticity

The ratio of stress to strain in the elastic range of a stress-strain curve for concrete defines the modulus of elasticity (E) of that concrete (Fig. 15-23). Normal-density concrete has a modulus of elasticity of 14,000 to 41,000 MPa.
(2,000,000 psi to 6,000,000 psi), depending on factors such as compressive strength and aggregate type. For normal-density concrete with compressive strengths \( f_c \) between 20 MPa and 35 MPa (3000 psi and 5000 psi), the modulus of elasticity can be estimated as 5000 times the square root of \( f_c \) (57,000 times the square root of \( f_c \) in psi). The modulus of elasticity for structural lightweight concrete is between 7000 MPa and 17,000 MPa (1,000,000 psi and 2,500,000 psi). \( E \) for any particular concrete can be determined in accordance with ASTM C469.

**Deflection**

Deflection of concrete beams and slabs is one of the more common and obvious building movements. The deflections are the result of flexural strains that develop under dead and live loads and that may result in cracking in the tensile zone of concrete members. Reinforced concrete structural design anticipates these tension cracks. Concrete members are often cambered, that is, built with an upward bow, to compensate for the expected later deflection.

**Poisson’s Ratio**

When a block of concrete is loaded in uniaxial compression, as in Fig. 15-24, it will shorten and at the same time develop a lateral strain or bulging. The ratio of lateral to axial strain is called Poisson’s ratio, \( \mu \). A common value used is 0.20 to 0.21, but the value may vary from 0.15 to 0.25 depending upon the aggregate, moisture content, concrete age, and compressive strength. Poisson’s ratio (ASTM C469) is generally of no concern to the structural designer; it is used in advanced structural analysis of flat-plate floors, shell roofs, arch dams, and mat foundations.

**Shear Strain**

Concrete, like other materials, deforms under shear forces. The shear strain produced is important in determining the load paths or distribution of forces in indeterminate structures—for example where shear-walls and columns both participate in resisting horizontal forces in a concrete building frame. The amount of movement, while not large, is significant in short, stubby members; in larger members it is overshadowed by flexural strains. Calculation of the shear modulus (modulus of rigidity), \( G \), is shown in Fig. 15-25; \( G \) varies with the strength and temperature of the concrete.

**Torsional Strain**

Plain rectangular concrete members can also fail in torsion, that is, a twisting action caused by bending about an axis parallel to the wider face and inclined at an angle of about 45 degrees to the longitudinal axis of a member. Microcracks develop at low torque; however, concrete behaves reasonably elastic up to the maximum limit of the elastic torque (Hsu 1968).

**CREEP**

When concrete is loaded, the deformation caused by the load can be divided into two parts: a deformation that occurs immediately (elastic strain) and a time-dependent deformation that begins immediately but continues at a decreasing rate for as long as the concrete is loaded. This latter deformation is called creep.

The amount of creep is dependent upon (1) the magnitude of stress, (2) the age and strength of the concrete.
when stress is applied, and (3) the length of time the concrete is stressed. It is also affected by other factors related to the quality of the concrete and conditions of exposure, such as: (1) type, amount, and maximum size of aggregate; (2) type of cementing materials; (3) amount of cement paste; (4) size and shape of the concrete element; (5) volume to surface ratio of the concrete element; (6) amount of steel reinforcement; (7) prior curing conditions; and (8) the ambient temperature and humidity.

Within normal stress ranges, creep is proportional to stress. In relatively young concrete, the change in volume or length due to creep is largely unrecoverable; in older or drier concrete it is largely recoverable.

The creep curves shown in Fig. 15-26 are based on tests conducted under laboratory conditions in accordance with ASTM C512. Cylinders were loaded to almost 40% of their compressive strength. Companion cylinders not subject to load were used to measure drying shrinkage; this was then deducted from the total deformation of the loaded specimens to determine creep. Cylinders were allowed to dry while under load except for those marked “sealed.” The two 28-day curves for each concrete strength in Fig. 15-26 show that creep of concrete loaded under drying conditions is greater than creep of concrete sealed against drying. Concrete specimens loaded at a late age will creep less than those loaded at an early age. It can be seen that as concrete strength decreases, creep increases. Fig. 15-27 illustrates recovery from the elastic and creep strains after load removal.

A combination of strains occurring in a reinforced column is illustrated in Fig. 15-28. The curves represent deformations and volume changes in a 14th-story column of a 76-story reinforced concrete building while under construction. The 400 x 1200-mm (16 x 48-in.) column contained 2.08% vertical reinforcement and was designed for 60-MPa (9000-psi) concrete.

**Fig. 15-26.** Relationship of time and age of loading to creep of two different strength concretes. Specimens were allowed to dry during loading, except for those labeled as sealed (Russell and Corley 1977).

**Fig. 15-27.** Combined curve of elastic and creep strains showing amount of recovery. Specimens (cylinders) were loaded at 8 days immediately after removal from fog curing room and then stored at 21°C (70°F) and 50% RH. The applied stress was 25% of the compressive strength at 8 days (Hansen and Mattock 1966).

**Fig. 15-28.** Summation of strains in a reinforced concrete column during construction of a tall building (Russell and Corley 1977).
in-place concrete is insignificant and does not have to be considered in engineering practice. Carbonation of paste proceeds slowly and produces little direct shrinkage at relative humidities of 100% and 25%. Maximum carbonation and carbonation shrinkage occurs at about 50% relative humidity. Irreversible shrinkage and weight gain occurs during carbonation. And the carbonated product may show improved volume stability to subsequent moisture change and reduced permeability (Verbeck 1958).

During manufacture some concrete masonry units are deliberately exposed to carbon dioxide after reaching 80% of their rated strength; this introduction to carbonation shrinkage makes the units more dimensionally stable. Future drying shrinkage is reduced 30% or more (Toennies and Shideler 1963).

One of the causes of surface crazing of concrete is the shrinkage that accompanies natural air carbonation of young concrete. More research is needed on the effect of early carbonation on deicer scaling resistance.

Carbonation of another kind also can occur in freshly placed, unhardened concrete. This carbonation causes a soft, chalky surface called dusting; it usually takes place during cold-weather concreting when there is an unusual amount of carbon dioxide in the air due to unvented heaters or gasoline-powered equipment operating in an enclosure.

**CHEMICAL CHANGES AND EFFECTS**

Some volume changes of concrete result from chemical reactions; these may take place shortly after placing and finishing or later due to reactions within the hardened concrete in the presence of water or moisture.

**Carbonation**

Hardened concrete containing some moisture reacts with carbon dioxide present in air, a reaction that results in a slight shrinkage of the surface paste of the concrete. The effect, known as carbonation, is not destructive but actually increases the chemical stability and strength of the concrete. However, carbonation also reduces the pH of concrete. If steel is present in the carbonated area, steel corrosion can occur due to the absence of the protective oxide film provided by concrete’s high pH. Rust is an expansive reaction and results in cracking and spalling of the concrete. The depth of carbonation is very shallow in dense, high-quality concrete, but can penetrate deeply in porous, poor-quality concrete. Because so little of a concrete element carbonates, carbonation shrinkage of cast-in-place concrete is insignificant and does not have to be considered in engineering practice.

Carbonation of paste proceeds slowly and produces little direct shrinkage at relative humidities of 100% and 25%. Maximum carbonation and carbonation shrinkage occurs at about 50% relative humidity. Irreversible shrinkage and weight gain occurs during carbonation. And the carbonated product may show improved volume stability to subsequent moisture change and reduced permeability (Verbeck 1958).

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**Sulfate Attack**

Sulfate attack of concrete can occur where soil and groundwater have a high sulfate content and measures to reduce sulfate attack, such as use of a low water to cementing materials ratio, have not been taken. The attack is greater in concrete that is exposed to wetting and drying, such as foundation walls and posts. Sulfate attack usually results in an expansion of the concrete because of the formation of solids from the chemical action or salt crystallization. The amount of expansion in severe circumstances has been significantly higher than 0.1%, and the disruptive effect within the concrete can result in extensive cracking and disintegration. The amount of expansion cannot be accurately predicted.

**Alkali-Aggregate Reactions**

Certain aggregates can react with alkali hydroxides in concrete, causing expansion and cracking over a period of years. The reaction is greater in those parts of a structure exposed to moisture. A knowledge of the characteristics of local aggregates is essential. There are two types of alkali-reactive aggregates, siliceous and carbonate. Alkali-aggregate reaction expansion may exceed 0.5% in concrete and can cause the concrete to fracture and break apart.

Structural design techniques cannot counter the effects of alkali-aggregate expansion, nor can the expan-
sion be controlled by jointing. In areas where deleteriously reactive aggregates are known to exist, special measures must be taken to prevent the occurrence of alkali-aggregate reaction.

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Satisfactory concrete construction and performance requires concrete with specific properties. To assure that these properties are obtained, quality control and acceptance testing are indispensable parts of the construction process. Test results provide important feedback used to base decisions regarding mix adjustments. However, past experience and sound judgment must be relied on in evaluating tests and assessing their significance in controlling the design, batching and placement processes that influence the ultimate performance of the concrete.

Specifiers are moving toward performance-based specifications (also called end-result or end-property specifications) that require the final performance of concrete be achieved independent of the process used to achieve the performance. Physical tests and concrete properties are used to measure acceptance. Such specifications may not have acceptance limits for process control tests such as slump or limits on the quantities of concrete ingredients as do prescriptive specifications. The end result of compressive strength, low permeability, documented durability, and a minimal number of cracks, for example, would be the primary measure of acceptance. Of course, even though process control tests may not be specified, the wise concrete producer would use them to guide the product to a successful end result. However, most specifications today are still a combination of prescriptive and performance requirements (Parry 2000).

**CLASSES OF TESTS**

Project specifications may affect (1) characteristics of the mixture, such as maximum size of aggregate, aggregate proportions, or minimum cement content; (2) characteristics of the cement, water, aggregates, and admixtures; and (3) characteristics of the freshly mixed and hardened concrete, such as temperature, slump, air content, and compressive or flexural strengths.

Cementitious materials are tested for their compliance with ASTM or AASHTO standards to avoid any abnormal performance such as early stiffening, delayed setting, or low strengths in concrete. More details regarding cementitious materials can be found in Chapters 2 and 3.

Tests of aggregates have two major purposes: (1) to determine the suitability of the material itself for use in concrete, including tests for abrasion, soundness against saturated freeze-thaw cycles, harmful materials by petrographic examination, and potential alkali-aggregate reactivity; and (2) to assure uniformity, such as tests for moisture control, relative density (specific gravity), and gradation of aggregates. Some tests are used for both purposes. Testing aggregates to determine their potential alkali-aggregate reactivity is discussed in Chapter 5, “Aggregates for Concrete.” Tests of concrete to evaluate the performance of available materials, to establish mixture proportions, and to control concrete quality during construction include slump, air content, temperature, density (unit weight), and strength. Slump, air content, and strength tests are usually required in project specifications for concrete quality control, whereas density is more useful in mixture proportioning. Nevertheless, ASTM C94 (AASHTO M 157) specifies that slump, air-content, density, and temperature tests should be made when strength specimens are made.

Following is a discussion of frequency of testing and descriptions of the major control tests to ensure uniformity of materials, desired properties of freshly mixed concrete, and required strength of hardened concrete. Special tests are also described.


**Computational Software.** In order to make computation of test data easier, NRMCA (2001) provides a CD with spreadsheets for a variety of concrete and aggregate tests.

**FREQUENCY OF TESTING**

Frequency of testing is a significant factor in the effectiveness of quality control of concrete. Specified test frequencies are intended for acceptance of the material or one of its components at a random location within the quantity or time period represented by the test. Such frequencies may not occur often enough to control the material within
specified limits during production. Process control tests are nonrandom tests performed more often than specified to document trends that allow adjustments to be made before acceptance tests are performed.

The frequency of testing aggregates and concrete for typical batch-plant procedures depends largely upon the uniformity of materials, including the moisture content of aggregates, and the production process. Initially it is advisable to make process control tests several times a day, but as work progresses and the material becomes more predictable, the frequency often can be reduced. ASTM C1451 provides a standard practice for determining the uniformity of cementitious materials, aggregates, and chemical admixtures used in concrete.

Usually, aggregate moisture tests are made once or twice a day. The first batch of fine aggregate in the morning is often overly wet since moisture will migrate overnight to the bottom of the storage bin. As fine aggregate is drawn from the bottom of the bin and additional aggregate is added, the moisture content should stabilize at a lower level and the first moisture test can be made. Obtaining moisture samples representative of the aggregates being batched is important; a one percent change in moisture content of fine aggregate will change the amount of mix water needed by approximately eight kilograms per cubic meter (13 lb/yd³).

Slump tests should be made for the first batch of concrete each day, whenever consistency of concrete appears to vary, and whenever strength-test cylinders are made at the jobsite.

Air-content tests should be made often enough at the point of delivery to ensure proper air content, particularly if temperature and aggregate grading change. An air-content test should be performed for each sample of concrete from which cylinders are made; a record of the temperature of each sample of concrete should also be kept.

The number of strength tests made will depend on the job specifications and the occurrence of variations. The ACI 318 building code and ASTM C94 require that strength tests of each class of concrete placed each day should be taken not less than once a day, nor less than once for each 115 cubic meters (150 cu yd) of concrete. In addition, ACI 318 recommends not less than once for each 500 square meters (5000 sq ft) of surface area for slabs or walls placed each day. The average strength of two 28-day cylinders is required for each test used for evaluating concrete. A 7-day test cylinder, along with the two 28-day test cylinders, is often made and tested to provide an early indication of strength development. As a rule of thumb, the 7-day strength is about 60% to 75% of the 28-day compressive strength, depending upon the type and amount of cementitious materials, water-cement ratio, curing temperature, and other variables. Additional specimens may be required when high-strength concrete is involved or where structural requirements are critical. Specimens should be laboratory cured when tested for acceptance or ultimate performance of the concrete. However, laboratory-cured specimens should not be used as an indication of in-place concrete strengths (ACI 318).

In-place concrete strengths are typically documented by casting specimens that are field-cured (as nearly as practical) in the same manner as concrete in the structure. Field-cured specimens are commonly used to decide when forms and shores might be removed under a structural slab or to determine when traffic will be allowed on new pavement. ASTM C31 (AASHTO T 23) contains additional instructions regarding the handling of field-cured cylinders. Although field-cured specimens may be tested at any age, 7-day tests are often made for comparison with laboratory tests at the same age; these are useful to judge if curing and protection during cold weather concreting is adequate.

TESTING AGGREGATES

Sampling Aggregates

Methods for obtaining representative samples of aggregates are given in ASTM D75 (AASHTO T 2). Accurate sampling is important. The location in the production process where samples will be obtained must be carefully planned. Sampling from a conveyor belt, stockpile, or aggregate bin may require special sampling equipment. Caution must be exercised to obtain a sample away from stockpile segregation and the sample must be large enough to meet ASTM minimum sample size requirements. In addition, samples obtained for moisture content testing should be placed in a sealed container or plastic bag as soon as possible to retain moisture until testing.

Reducing large field samples to small quantities for individual tests must be done with care so that the final samples will be truly representative ASTM C702 (AASHTO T 248). For coarse aggregate, this is done by the quartering method: The sample, thoroughly mixed, is spread on a piece of canvas in an even layer 75 or 100 mm (3 or 4 in.) thick. It is divided into four equal parts. Two opposite parts are then discarded. This process is repeated until the desired size of sample remains. A similar procedure is sometimes used for moist, fine aggregate. Sample splitters are desirable for dry aggregate (Fig. 16-1) but should not be used for samples that are more than saturated surface dry.

Organic Impurities

Organic impurities in fine aggregate should be determined in accordance with ASTM C40 (AASHTO T 21). A sample of fine aggregate is placed in a sodium hydroxide solution and shaken. The following day the color of the sodium hydroxide solution is compared with a glass color.
standard or standard color solution. If the color of the solution containing the sample is darker than the standard color solution or Organic Glass Plate No. 3, the fine aggregate should not be used for important concrete work without further investigation.

Some fine aggregates contain small quantities of coal or lignite that give the liquid a dark color. The quantity may be insufficient to reduce the strength of the concrete appreciably. If surface appearance of the concrete is not important, ASTM C33 (AASHTO M 6) states that fine aggregate is acceptable if the amount of coal and lignite does not exceed 1.0% of the total fine aggregate mass. A fine aggregate failing this ASTM C33 (AASHTO M 6) limit may be used if, when tested in accordance with ASTM C87 (AASHTO T 71), the 7-day strengths of mortar cubes made with the sand (ASTM C109 or AASHTO T 106) are at least 95% of the 7-day strengths of mortar made with the same sand, but washed in a 3% solution of sodium hydroxide and then thoroughly rinsed in water. It should be realized that appreciable quantities of coal or lignite in aggregates can cause popouts and staining of the concrete and can reduce durability when concrete is exposed to weathering. Local experience is often the best indication of the durability of concrete made with such aggregates.

**Objectionable Fine Material**

Large amounts of clay and silt in aggregates can adversely affect durability, increase water requirements, and increase shrinkage. Specifications usually limit the amount of material passing the 75 μm (No. 200) sieve to 2% or 3% in fine aggregate and to 1% or less in coarse aggregate. Testing for material finer than the 75 μm (No. 200) sieve should be done in accordance with ASTM C117 (AASHTO T 11). Testing for clay lumps should be in accordance with ASTM C142 (AASHTO T 112).

**Grading**

Gradation of aggregates significantly affects concrete mixture proportioning and workability. Hence, gradation tests are an important element in the assurance of concrete quality. The grading or particle size distribution of an aggregate is determined by a sieve analysis test in which the particles are divided into their various sizes by standard sieves. The analysis should be made in accordance with ASTM C136 (AASHTO T 27).

Results of sieve analyses are used in three ways: (1) to determine whether or not the materials meet specifications; (2) to select the most suitable material if several aggregates are available; and (3) to detect variations in grading that are sufficient to warrant blending selected sizes or an adjustment of concrete mix proportions.

The grading requirements for concrete aggregate are shown in Chapter 5 and ASTM C33 (AASHTO M 6/M 80). Materials containing too much or too little of any one size should be avoided. Some specifications require that mixture proportions be adjusted if the average fineness modulus of fine aggregate changes by more than 0.20. Other specifications require an adjustment in mixture proportions if the amount retained on any two consecutive sieves changes by more than 10% by mass of the total fine-aggregate sample. A small quantity of clean particles that pass a 150 μm (No. 100) sieve but are retained on a 75 μm (No. 200) sieve is desirable for workability. For this reason, most specifications permit up to 10% of this material in fine aggregate.

Well-graded aggregates contain particles on each sieve size. Well-graded aggregates enhance numerous factors that result in greater workability and durability. The more well-graded an aggregate is, the more it will pack together efficiently, thus reducing the volume between aggregate particles that must be filled by paste. On the other hand, gap-graded aggregates—those having either a large quantity or a deficiency of one or more sieve sizes—can result in reduced workability during mixing, pumping, placing, consolidation and finishing. Durability can suffer too as a result of using more fine aggregate and water to produce a workable mix. See Chapter 5 and Graves (2006) for additional information on aggregate grading.

**Moisture Content of Aggregates**

Several methods are used for determining the amount of moisture in aggregate samples. The total moisture content for fine or coarse aggregate can be tested in accordance with ASTM C566 (AASHTO T 255). In this method a measured sample of damp aggregate is dried either in a ventilated conventional oven, microwave oven, or over an electric or gas hotplate. From the mass before and after drying, the total moisture content can be calculated as follows:
When drying equipment is not available a field or plant determination of surface (free) moisture in fine aggregate can be made in accordance with ASTM C70. The same procedure can be used for coarse aggregate with appropriate changes in the size of sample and dimensions of the container. This test depends on displacement of water by a known mass of moist aggregate; therefore, the relative density (specific gravity) of the aggregate must be known accurately.

Electric moisture meters are used in many concrete batching plants primarily to monitor the moisture content of fine aggregates, but some plants also use them to check coarse aggregates. They operate on the principle that the electrical resistance of damp aggregate decreases as moisture content increases, within the range of dampness normally encountered. The meters measure the electrical resistance of the aggregate between electrodes protruding into the batch hopper or bin. Moisture meters using the microwave-absorption method are gaining popularity because they are more accurate than the electric meters. However, both methods measure moisture contents accurately and rapidly, but only at the level of the probes. These meters require frequent calibration and must be maintained properly. The variable nature of moisture contents in aggregates cause difficulty in obtaining representative samples for comparison to electric moisture meters. Several oven-dried moisture content tests should be performed to verify the calibration of these meters before trends in accuracy can be established.

Table 16-1 illustrates a method of adjusting batch weights for moisture in aggregates.

### Table 16-1. Example of Adjustment in Batch Weights for Moisture in Aggregates

<table>
<thead>
<tr>
<th>Aggregate data</th>
<th>Absorption, %</th>
<th>Moisture content, %</th>
</tr>
</thead>
<tbody>
<tr>
<td>Fine aggregate</td>
<td>1.2</td>
<td>5.8</td>
</tr>
<tr>
<td></td>
<td>0.4</td>
<td>0.8</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Concrete ingredients</th>
<th>Mix design mass (aggregates in dry BOD condition), kg/m³ (lb/yd³)</th>
<th>Aggregate mass (SSD condition), kg/m³ (lb/yd³) (Absorbed %)</th>
<th>Aggregate mass (in moist condition), kg/m³ (lb/yd³) (Moisture %)</th>
<th>Mix water correction for surface moisture in aggregates, kg/m³ (lb/yd³)</th>
<th>Adjusted batch weight, kg/m³ (lb/yd³)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cement</td>
<td>355 (598)</td>
<td>703 (1185)</td>
<td>735 (1239)</td>
<td>32 (54)</td>
<td>355 (598)</td>
</tr>
<tr>
<td>Fine aggregate</td>
<td>695 (1171)</td>
<td>1064 (1793)</td>
<td>1068 (1800)</td>
<td>4 (7)</td>
<td>735 (1239)</td>
</tr>
<tr>
<td>Coarse aggregate</td>
<td>1060 (1787)</td>
<td>1068 (1800)</td>
<td></td>
<td></td>
<td>1068 (1800)</td>
</tr>
<tr>
<td>Water</td>
<td>200 (337)</td>
<td></td>
<td></td>
<td></td>
<td>164 (276)</td>
</tr>
<tr>
<td>Total</td>
<td>2310 (3893)</td>
<td></td>
<td></td>
<td></td>
<td>2322 (3913)†</td>
</tr>
</tbody>
</table>

* An aggregate in a bulk-oven dry (BOD) condition is one with its permeable voids completely dry so that it is fully absorbent.
** An aggregate in a saturated, surface-dry (SSD) condition is one with its permeable voids filled with water and with no surface moisture on it. Concrete suppliers often request mix design proportions on a SSD basis because of batching software requirements.
† Total adjusted batch weight is higher than total mix design weight by the amount of water absorbed in the aggregate.
TESTING FRESHLY MIXED CONCRETE

Sampling Freshly Mixed Concrete

The importance of obtaining truly representative samples of freshly mixed concrete for control tests must be emphasized. Unless the sample is representative, test results will be misleading. Samples should be obtained and handled in accordance with ASTM C172 (AASHTO T 141). Except for routine slump and air-content tests performed for process control, ASTM C172 (AASHTO T 141) requires that sample size used for acceptance purposes be at least 28 liter (1 cu ft) and be obtained within 15 minutes between the first and final portions of the sample. The composite sample, made of two or more portions, should not be taken from the very first or last portion of the batch discharge. The sample should be protected from sunlight, wind, and other sources of rapid evaporation during sampling and testing.

Consistency

The slump test, ASTM C143 (AASHTO T 119), is the most generally accepted method used to measure the consistency of concrete (Fig. 16-2). The test equipment consists of a slump cone (a metal conical mold 300 mm [12 in.] high, with a 200-mm [8-in.] diameter base and 100-mm [4-in.] diameter top) and a steel rod 16 mm (5⁄8 in.) in diameter and 600 mm (24 in.) long with a hemispherically shaped tip. The dampened slump cone, placed upright on a flat, nonabsorbent rigid surface, should be filled in three layers of approximately equal volume. Therefore, the cone should be filled to a depth of about 70 mm (2 1⁄2 in.) for the first layer, a depth of about 160 mm (6 in.) for the second layer, and overfilled for the third layer. Each layer is rodded 25 times. Following rodding, the last layer is struck off and the cone is slowly raised vertically 300 mm (12 in.) in 5 ± 2 seconds. As the concrete subsides or settles to a new height, the empty slump cone is then inverted and gently placed next to the settled concrete. The slump is the vertical distance the concrete settles, measured to the nearest 5 mm (¼ in.); a ruler is used to measure from the top of the slump cone (mold) to the displaced original center of the subsided concrete (see Fig. 16-2).

A higher slump value is indicative of a more fluid concrete. The entire test through removal of the cone should be completed in 2 1⁄2 minutes, as concrete loses slump with time. If a falling away or shearing off occurs from a portion of the concrete, another test should be run on a different portion of the sample.

Another test method for flow of fresh concrete involves the use of the K-Slump Tester (ASTM C1362). This is a probe-type instrument that is thrust into the concrete in any location where there is a minimum depth of 175 mm (7 in.) of concrete a 75-mm (3-in.) radius of concrete around the tester. The amount of mortar flowing into openings in the tester is reported as a measure of flow.

Additional consistency tests include: the FHWA vibrating slope apparatus (Wong and others 2001 and Saucier 1966); British compacting factor test (BS 1881); Powers remolding test (Powers 1932); German flow table test (DIN 1048-1); Vebe consistometer for roller-compacted concrete (ASTM C1170); Kelly ball penetration test (ASTM C360-92 now discontinued); Thaulow tester; the inverted slump cone for fiber-reinforced concrete (ASTM C995); Powers and Wiler plastometer (Powers and Wiler 1941);

![Fig. 16-2. Slump test for consistency of concrete. Figure A illustrates a lower slump, Figure B a higher slump. (IMG12179, IMG12180)](IMG12179, IMG12180)
Tattersall (1971) workability device; Colebrand test; BML viscometer (Wallevik 1996); BTRHEOM rheometer for fluid concrete (de Larrard, Szitkar, Hu, and Joly 1993); free-orifice rheometer (Bartos 1978); delivery chute torque meter (US patent 4,332,158 [1982]); delivery-chute vane (US patent 4,578,989 [1986]); Angles flow box (Angles 1974); ring penetration test (Teranishs and others 1994); and the Wigmore (1948) consistometer. The Vebe test and the Thaulow test are especially applicable to stiff and extremely dry mixes while the flow table is especially applicable to flowing concrete (Daniel 2006).

Temperature Measurement

Because of the important influence concrete temperature has on the properties of freshly mixed and hardened concrete, many specifications place limits on the temperature of fresh concrete. Glass or armored thermometers are available (Figs. 16-3). The thermometer should be accurate to plus or minus 0.5°C (±1°F) and should remain in a representative sample of concrete for a minimum of 2 minutes or until the reading stabilizes. A minimum of 75 mm (3 in.) of concrete should surround the sensing portion of the thermometer. Electronic temperature meters with precise digital readouts are also available. The temperature measurement (ASTM C1064 or AASHTO T 309) should be completed within 5 minutes after obtaining the sample.

Density and Yield

The density (unit weight) and yield of freshly mixed concrete (Fig. 16-4) are determined in accordance with ASTM C138 (AASHTO T 121). The results should be sufficiently accurate to determine the volumetric quantity (yield) of concrete produced per batch (see Chapter 9). The test also can give indications of air content provided the relative densities of the ingredients are known. A balance or scale sensitive to 0.3% of the anticipated mass of the sample and container is required. For example, a 7-liter (0.25-ft³) density container requires a scale sensitive to 50 g (0.1 lb). The size of the container used to determine density and yield varies with the size of aggregate; if in good condition, the 7-liter (0.25-ft³) airmeter container is commonly used with aggregates up to 25 mm (1 in.); a 14-liter (0.5-ft³) container is used with aggregates up to 50 mm (2 in.). The container should be calibrated at least annually (ASTM C1077). Care is needed to consolidate the concrete adequately by either rodding or internal vibration. Strike off the top surface using a flat plate so that the container is filled to a flat smooth finish. The density is expressed in kilograms per cubic meter (pounds per cubic foot) and the yield in cubic meters (cubic feet).

The density of unhardened as well as hardened concrete can also be determined by nuclear methods, ASTM C1040 (AASHTO T 271).

Air Content

A number of methods for measuring air content of freshly mixed concrete can be used. ASTM standards include the pressure method (C231) (AASHTO T 152), the volumetric method (C173) (AASHTO T 196), and the gravimetric method (C138) (AASHTO T 121).

The pressure method (Fig. 16-5) is based on Boyle’s law, which relates pressure to volume. Many commercial air meters of this type are calibrated to read air content directly when a predetermined pressure is applied. The applied pressure compresses the air within the concrete sample, including the air in the pores of aggregates. For this reason, tests by this method are not suitable for determining the air content of concretes made with some lightweight aggregates or other very porous materials. Aggregate correction factors that compensate for air trapped in...
normal-weight aggregates are relatively constant and, though small, should be subtracted from the pressure meter gauge reading to obtain the correct air content. The instrument should be calibrated for various elevations above sea level if it is to be used in localities having considerable differences in elevation. Some meters utilize change in pressure of a known volume of air and are not affected by changes in elevation. Pressure meters are widely used because the mix proportions and specific gravities of the concrete ingredients need not be known. Also, a test can be conducted in less time than is required for other methods.

The volumetric method (Fig. 16-6) outlined in ASTM C173 (AASHTO T 196) requires removal of air from a known volume of concrete by agitating the concrete in an excess of water. This method can be used for concrete containing any type of aggregate, including lightweight or porous materials. An aggregate correction factor is not necessary with this test. The volumetric test is not affected by atmospheric pressure, and specific gravity of the concrete ingredients need not be known. Care must be taken to agitate the sample sufficiently to remove all air. The addition of 500 mL (1 pt) of alcohol accelerates the removal of air, thus shortening test times; it also dispels most of the foam and increases the precision of the test, including those performed on high-air-content or high-cement-content concretes.

The gravimetric method utilizes the same test equipment used for determining the density (unit weight) of concrete. The measured density of concrete is subtracted from the theoretical density as determined from the absolute volumes of the ingredients, assuming no air is present (see ASTM C138 or AASHTO T 121). This difference, expressed as a percentage of the theoretical density, is the air content. Mixture proportions and specific gravities of the ingredients must be accurately known; otherwise results may be in error. Consequently, this method is suitable only where laboratory-type control is exercised. Significant changes in density can be a convenient way to detect variability in air content.

A pocket-size air indicator (AASHTO T 199) can be used as a quick check for the presence of low, medium, or high levels of air in concrete, but it is not a substitute for the other more accurate methods. A representative sample of mortar from the concrete is placed in the container. The container is then filled with alcohol and rolled with the thumb over the open end to remove the air from the mortar. The indicated air content is determined by comparing the drop in the level of the alcohol with a calibration chart. The test can be performed in a few minutes. It is especially useful in checking for the presence of air in concrete near the surface that may have suffered reductions in air because of faulty finishing procedures.

With any of the above methods, air-content tests should be started within 5 minutes after the final portion of the composite sample has been obtained.

Studies into the effect of fly ash on the air-void stability of concrete have resulted in the development of the foam-index test. The test can be used to measure the relative air-entraining admixture requirements for concrete mixtures containing fly ash. The fly ash to be tested is placed in a wide mouth jar along with the air-entraining admixture and shaken vigorously. Following a waiting period of 45 seconds, a visual determination of the stability of the foam or bubbles is made (Gebler and Klieger 1983).
Strength Specimens

Specimens molded for strength tests should be made and cured in accordance with ASTM C31 or AASHTO T 23 (field-molded specimens) or ASTM C192 or AASHTO T 126 (laboratory-molded specimens). Molding of strength specimens should be started within 15 minutes after the composite sample is obtained.

The standard test specimen for compressive strength of concrete with a maximum aggregate size of 50 mm (2 in.) or smaller is a cylinder 150 mm (6 in.) in diameter by 300 mm (12 in.) high (Fig. 16-7). For larger aggregates, the diameter of the cylinder should be at least three times the maximum-size aggregate, and the height should be twice the diameter. While rigid metal molds are preferred, parafined cardboard, plastic, or other types of disposable molds conforming to ASTM C470 can be used. They should be placed on a smooth, level, rigid surface and filled carefully to avoid distortion of their shape.

A 100-mm (4-in.) diameter by 200-mm (8-in.) high cylinder mold has been commonly used with high strength concrete containing up to 19 mm (3/4 in.) maximum-size aggregate (Burg and Ost 1994, Forstie and Schnormeier 1981, and Date and Schnormeier 1984). The 100 x 200-mm cylinder is easier to cast, requires less sample, weighs considerably less than a 150 x 300-mm (6 x 12-in.) concrete cylinder; it is therefore easier to handle and requires less moist-curing storage space. In addition, the smaller cross-sectional area allows higher compressive strengths to be reached by a testing machine that has a smaller load capacity. The difference in indicated strength between the two cylinder sizes is insignificant as illustrated in Fig. 16-8. The standard deviation and coefficient of variation of 100-mm cylinders is slightly higher or similar to that for 150-mm cylinders (Burg and others 1999 and Pistilli and Willems 1993). The predominant size used in Canada is the 100-mm diameter cylinder. Consult job specifications for allowable cylinder sizes.

Beams for the flexural strength test should be 150 x 150 mm (6 x 6 in.) in cross section for aggregates up to 50 mm (2 in.). For larger aggregates, the minimum cross-sectional dimension should be not less than three times the maximum size of aggregate. The length of beams should be at least three times the depth of the beam plus 50 mm (2 in.), or a total length of not less than 500 mm (20 in.) for a 6 x 6-in. beam.

Test cylinders to be rodded (slump of 25 mm [1 in.] or more) should be filled in three approximately equal layers with each...
layer rodded 25 times for 150-mm (6-in.) diameter cylinders; beam specimens up to 200 mm (8 in.) deep should be filled in two equal layers with each layer rodded once with a 16-mm (5/8-in.) rod for each 1400 mm² (2 in.²) of the specimen’s top surface area. If the rodding leaves holes, the sides of the mold should be lightly tapped with a mallet or open hand. Cylinders to be vibrated should be filled in two layers with one insertion per layer for 100-mm (4-in.) diameter cylinders and two insertions per layer for 150-mm (6-in.) cylinders.

Beams over 200 mm (8 in.) deep and cylinders 300 to 450 mm (12 to 18 in.) deep to be vibrated (slump of 75 mm [3 in.] or less) should be filled in two layers; beams 150 to 200 mm (6 to 8 in.) deep to be vibrated can be filled in one layer. Internal vibrators should have a maximum width of no more than 1/3 the width of beams or 1/4 the diameter of cylinders. Immediately after casting, the tops of the specimens should be (1) covered with an oiled glass or steel plate, (2) sealed with a plastic bag, or (3) sealed with a plastic cap.

The strength of a test specimen can be greatly affected by jostling, changes in temperature, and exposure to drying, particularly within the first 24 hours after casting. Thus, test specimens should be cast in locations where subsequent movement is unnecessary and where protection is possible. Cylinders and test beams should be protected from rough handling at all ages. Remember to identify specimens on the exterior of the mold to prevent confusion and errors in reporting. Do not etch identification numbers into the surface of fresh concrete test specimens. Use tape or identification tags that do not damage the sample.

Standard testing procedures require that specimens be cured under controlled conditions, either in the laboratory (Fig. 16-9) or in the field. Controlled laboratory curing in a moist room or in a limewater storage tank gives an accurate indication of the quality of the concrete as delivered. Limewater must be saturated with hydrated lime, not agricultural lime, in accordance with ASTM C511 (AASHTO M 201) to prevent leaching of lime from concrete specimens.

Specimens cured in the field in the same manner as the structure more closely represent the actual strength of concrete in the structure at the time of testing; however, they give little indication of whether a deficiency is due to the quality of the concrete as delivered or to improper handling and curing. On some projects, field-cured specimens are made in addition to those destined for controlled laboratory curing; these are especially useful when the weather is unfavorable, to determine when forms can be removed, or when the structure can be put into use. For more information see “Strength Tests of Hardened Concrete” in this chapter and ASTM (2000).

In-place concrete strength development can also be evaluated by maturity testing (ACI Committee 306 and ASTM C1074), which was discussed in Chapter 14.

**Time of Setting**

Test method ASTM C403 (AASHTO T 197) is used to determine the time of setting of concrete by means of penetration resistance measurements made at regular time intervals on mortar sieved from the concrete mixture (Fig. 16-9).
Accelerated Compression Tests to Project Later-Age Strength

The need to assess the quality of concrete at early ages in comparison to traditional 28-day tests has received much attention due to the quickening pace of today’s construction. ASTM has developed two methods for estimating later-age strengths of concrete specimens based upon early-age tests.

ASTM C684 uses accelerated strength tests to expedite quality control of concrete. Cylinder strength tests are accelerated using one of four curing procedures: warm water at $35\pm 3{^\circ}C (95\pm 5{^\circ}F)$, boiling water, autogenous (insulated), and high temperature at $150\pm 3{^\circ}C (300\pm 5{^\circ}F)$. Accelerated strength tests are performed at ages ranging between 5 and 49 hours, depending on the curing procedure used. Later-age strengths are estimated using previously established relationships between accelerated strength and standard 28-day compressive strength tests.

ASTM C918 uses the maturity method of monitoring temperature of cylinders cured in accordance with standard methods outlined in ASTM C31 (AASHTO T 23). Cylinders are tested at early ages beyond 24 hours, and the concrete temperature history is used to compute the maturity index at the time of test. Using historic data, a prediction equation is developed to project the strength at later ages based on the maturity index and early-age strength tests. See Carino (2006).

Chloride Content

The chloride content of concrete and its ingredients should be checked to make sure it is below the limit necessary to avoid corrosion of reinforcing steel. An approximation of the water-soluble chloride content of freshly mixed concrete, aggregates, and admixtures can be made using a method initiated by the National Ready Mixed Concrete Association (NRMCA 1986). A determination of the total chloride content of freshly mixed concrete may be made by summing up the chloride contents of all of the individual constituents of the mix. The NRMCA method gives a quick approximation and should not be used to determine compliance. See Chapter 9 for chloride-ion limitations for concrete.

Portland Cement Content, Water Content, and Water-Cement Ratio

Test methods are available for estimating the portland cement and water content of freshly mixed concrete. These test results can assist in determining the strength and durability potential of concrete prior to setting and hardening and can indicate whether or not the desired cement and water contents were obtained. ASTM test methods C1078-87 and C1079-87 (discontinued in 1998), based on the Kelly-Vail method, can be used to determine cement content and water content, respectively. Experimental methods using microwave absorption have been developed to estimate water to cement ratio. The disadvantage of these test methods is they require sophisticated equipment and special operator skills, which may not be readily available.

Other tests for determining cement or water contents can be classified into four categories: chemical determination, separation by settling and decanting, nuclear related, and electrical. The Rapid Analysis Machine and nuclear cement gage have been used to measure cement contents (Forester, Black, and Lees 1974 and PCA 1983). The microwave oven (AASHTO TP 23) and neutron-scattering methods have been used to measure water contents. For an overview of these and other tests from all four categories, see Hime (1994). A combination of these tests can be run independently of each other to determine either cement content or water content to calculate the water-cement ratio.

Supplementary Cementitious Materials Content

Standard test methods are not available for determining the supplementary cementitious materials content of freshly mixed concrete. However, the presence of certain supplementary cementitious materials, such as fly ash, can be determined by washing a sample of the concrete’s mortar over a 45 $\mu$m (No. 325) sieve and viewing the residue retained with a stereo microscope (150 to 250X) (Fig. 16-11). Fly ash particles appear as spheres of various colors. Sieving the mortar through a 150 or 75 $\mu$m (No. 100 or 200) sieve is helpful in removing sand grains.

Bleeding of Concrete

The bleeding properties of fresh concrete can be determined by two methods described in ASTM C232 (AASHTO T 158). One method consolidates the specimen by tamping without further disturbance; the other method consolidates the specimen by vibration after which the specimen is vibrated intermittently throughout the test. The amount of bleed water at the surface is expressed as the volume of bleed water per unit area of exposed concrete, or as a percentage of the net mixing water in the test specimen. Typical values range from 0.01 to 0.08
Strength Tests of Hardened Concrete

Strength tests of hardened concrete can be performed on the following: (1) cured specimens molded in accordance with ASTM C31 or C192 (AASHTO T 23 and T 126) from samples of freshly mixed concrete; (2) in-situ specimens cored or sawed from hardened concrete in accordance with ASTM C42 (AASHTO T 24); or (3) specimens made from cast-in-place cylinder molds, ASTM C873 (Fig. 16-13).

Cast-in-place cylinders can be used in concrete that is 125 to 300 mm (5 to 12 in.) in depth. The mold is filled in the normal course of concrete placement. The specimen is then cured in place and in the same manner as the rest of the concrete section. The specimen is removed from the concrete and mold immediately prior to testing to determine the in-place concrete strength. This method is particularly applicable in cold-weather concreting, post-tensioning work, slabs, or any concrete work where a minimum in-place strength must be achieved before construction can continue.

For all methods, cylindrical samples should have a diameter at least three times the maximum size of coarse aggregate in the concrete and a length as close to twice the diameter as possible. Correction factors are available in ASTM C42 (AASHTO T 24) for samples with lengths of 1 to 2 times the diameter. Cores and cylinders with a height of less than 95% of the diameter before or after capping should not be tested. Use of a minimum core diameter of 95 mm (3.75 in.) is suggested where a length to diameter (L/D) ratio greater than one is possible.

Drilled cores should not be taken until the concrete can be sampled without disturbing the bond between the mortar and the coarse aggregate. For horizontal surfaces, cores should be taken vertically and not near formed joints or edges. For vertical or sloped faces, cores should be taken perpendicular to the central portion of the concrete element. Although diamond-studded coring bits can cut through reinforcing steel, it should be avoided if possible.

mL/cm² or 0.1% to 2.5% of mix water. The bleeding test is rarely used in the field (Fig. 16-12). Bleeding was also discussed in Chapter 1.

TESTING HARDENED CONCRETE

Premolded specimens described in the previous section “Strength Specimens” (ASTM C31 [AASHTO T 23], ASTM C192 [AASHTO T 126], or ASTM C873), or samples of hardened concrete obtained from construction or laboratory work (ASTM C42 [AASHTO T 24], ASTM C823, or ASTM C873) can be used in tests on hardened concrete. Separate specimens should be obtained for each test performed because specimen preconditioning for certain tests can make the specimen unusable for other tests.
A cap is placed on one or both ends of the cylinder; the specimen is then tested in accordance with ASTM C39 (AASHTO T 22) with the added step to stop the test at 10% of the anticipated load to check that the axis of the cylinder is vertical within a tolerance of 0.5°. If either the perpendicularity of the cylinder end, or the vertical alignment during loading are not met, the load applied to the cylinder may be concentrated on one side of the specimen. This can cause a short shear fracture in which the failure plane intersects the end of the cylinder. This type of fracture usually indicates the cylinder failed prematurely, yielding results lower than the actual strength of the concrete. If perpendicularity requirements are not met, the cylinder can be saw-cut, ground, or capped with a sulfur mortar compound in accordance with ASTM C617 (AASHTO T 231).

Short shear fractures can also be reduced by: dusting the pad and end of cylinder with corn starch or talcum powder, preventing excess water from cylinders or burlap from draining into the retainer and below the pad, and checking bearing surfaces of retainers for planeness and indentations. In addition, annually clean and lubricate the spherically seated block and adjacent socket on the compression machine.

Testing of specimens (Fig. 16-15) should be done in accordance with (1) ASTM C39 (AASHTO T 22) for compressive strength, (2) ASTM C78 (AASHTO T 97) for flexural strength using third-point loading, (3) ASTM C293 (AASHTO T 177) for flexural strength using center-point loading, and (4) ASTM C496 (AASHTO T 198) for splitting tensile strength. Fig. 16-16 shows the correlation between compressive strength and flexural strength test results.

For both pavement thickness design and pavement mixture proportioning, the modulus of rupture (flexural strength) should be determined by the third-point loading test (ASTM C78 or AASHTO T 97). However, modulus of rupture by center-point loading (ASTM C293 or AASHTO T 177) or cantilever loading can be used for job control if empirical relationships to third-point loading test results are determined before construction starts.
The moisture content of the specimen has considerable effect on the resulting strength (Fig. 16-15). Beams for flexural tests are especially vulnerable to moisture gradient effects. A saturated specimen will show lower compressive strength and higher flexural strength than those for companion specimens tested dry. This is important to consider when cores taken from hardened concrete in service are compared with molded specimens tested as taken from the moist-curing room or water storage tank. Cylinders used for acceptance testing for a specified strength must be cured in accordance with ASTM C31 and C511 (AASHTO T 23 and M 201) to accurately represent the quality of the concrete. However, cores are subject to workmanship, variable environmental site conditions, and variable conditioning after extraction. Cores are tested in either a dry or moist condition, but rarely in a saturated condition similar to lab-cured cylinders. Because cores and cylinders are handled in very different ways, they cannot be expected to yield the same results.

The amount of variation in compressive-strength testing is far less than for flexural-strength testing. To avoid the extreme care needed in field flexural-strength testing, compressive-strength tests can be used to monitor concrete quality; however, a laboratory-determined empirical relationship (Fig. 16-16) must be developed between the compressive and flexural strength of the concrete used (Kosmatka 1985a). Because of this empirical relationship and the economics of testing cylinders instead of beams, most state departments of transportation are now utilizing compression tests of cylinders to monitor concrete quality for their pavement and bridge projects.

**Evaluation of Compression Test Results.** The ACI 318 building code states that the compressive strength of concrete can be considered satisfactory if the following conditions are met: the averages of all sets of three consecutive strength tests equal or exceed the specified 28-day strength $f_{cu}$ and no individual strength test (average of two cylinders) is more than 3.5 MPa (500 psi) below the specified strength. If the results of the cylinder tests do not meet this criteria, the strength of the in-place concrete should be evaluated by drilled cores.

In addition to the two 28-day cylinders, job specifications often require one or two 7-day cylinders and one or more "hold" cylinders. The 7-day cylinders monitor early strength gain. Hold cylinders are commonly used to provide additional information in case the 28-day cylinders are damaged or do not meet the required compressive strength. For low 28-day test results, the hold cylinders are typically tested at 56 days in age.

Protection and curing procedures should also be evaluated to judge if they are adequate when field-cured cylinders have a strength of less than 85% of that of companion laboratory-cured cylinders. The 85% requirement may be waived if the field-cured strength exceeds $f_{cu}$ by more than 3.5 MPa (500 psi).

When necessary, the in-place concrete strength should be determined by testing three cores for each strength test taken in the portion of the structure where the laboratory-cured cylinders did not meet acceptance criteria. Moisture conditioning of cores prior to compression testing should follow the guidelines in ASTM C42 (AASHTO T 24) or ACI 318.

Nondestructive test methods are not a substitute for core tests (ASTM C42 or AASHTO T 24). If the average strength of three cores is at least 85% of $f_{cu}$, and if no single core is less than 75% of $f_{cu}$, the concrete in the area represented by the cores is considered structurally adequate. If the results of properly made core tests are so low as to leave structural integrity in doubt, load tests as outlined in Chapter 20 of ACI 318 may be performed. Refer to Chapter 9 and NRMCA (1979), ACI Committee 214 (2002), and ACI Committee 318 (2008).

**Air Content**

The air-content and air-void-system parameters of hardened concrete can be determined by ASTM C457. The hardened air-content test is performed to assure that the air-void system is adequate to resist damage from a freeze-thaw environment. The test is also used to determine the effect different admixtures and methods of placement and consolidation have on the air-void system. The test can be performed on premolded specimens or samples removed from the structure. Using a polished section of a concrete sample, the air-void system is documented by making measurements using a microscope. The information obtained from this test includes the volume of entrained and entrapped air, its specific surface (surface area of the air voids), the spacing factor, and the number of voids per
can render the specimens useless for certain additional tests, especially strength tests. The density can be obtained by multiplying the relative density (specific gravity) by the density of water (1000 kg/m³ or 62.4 lb/ft³).

Saturated, surface-dry density (SSD) is often required for specimens to be used in other tests. In this case, the density can be determined by soaking the specimen in water for 48 hours and then determining its weight in air (when SSD) and immersed in water. The SSD density is then calculated as follows:

\[ D_{SSD} = \frac{M_1 \rho}{M_1 - M_2} \]

where

- \( D_{SSD} \) is density in the SSD condition
- \( M_1 \) is the SSD mass in air, kg (lb)
- \( M_2 \) is the apparent mass immersed in water, kg (lb)
- \( \rho \) is the density of water, 1000 kg/m³ (62.4 lb/ft³)

The SSD density provides a close indication of the freshly mixed density of concrete. The density of hardened concrete can also be determined by nuclear methods (ASTM C1040 or AASHTO T 271).

The rate of absorption (sorptivity) of water by hardened concrete can be determined using ASTM C1585.

### Table 16-2. Permeability and Absorption of Concretes Moist Cured 7 Days and Tested After 90 Days.

<table>
<thead>
<tr>
<th>Mix No.</th>
<th>Cement, kg/m³ (lb/yd³)</th>
<th>w/cm</th>
<th>Compressive strength at 90 days, MPa (psi)</th>
<th>RCPT, coulombs</th>
<th>90 days ponding, % CI</th>
<th>Water, m/s**</th>
<th>Air, m/s**</th>
<th>Porosity, %‡</th>
<th>Vol. of permeable voids, %</th>
<th>Absorption after immersion, %</th>
<th>Absorption after immersion and boiling, %</th>
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</thead>
<tbody>
<tr>
<td>1</td>
<td>445 (750)</td>
<td>0.26*</td>
<td>ASTM C39 (AASHTO T 22)</td>
<td>104.1 (15100)</td>
<td>65</td>
<td>0.013</td>
<td>—</td>
<td>2.81 x 10⁻¹⁰</td>
<td>7.5</td>
<td>6.2</td>
<td>2.43</td>
</tr>
<tr>
<td>2</td>
<td>445 (750)</td>
<td>0.29*</td>
<td>ASTM C1202 (AASHTO T 277)</td>
<td>76.7 (11130)</td>
<td>852</td>
<td>0.022</td>
<td>—</td>
<td>3.19 x 10⁻¹⁰</td>
<td>8.8</td>
<td>8.0</td>
<td>3.13</td>
</tr>
<tr>
<td>3</td>
<td>381 (642)</td>
<td>0.40*</td>
<td>ASTM C1543 (AASHTO T 259)</td>
<td>46.1 (6690)</td>
<td>3242</td>
<td>0.058</td>
<td>—</td>
<td>2.61 x 10⁻¹²</td>
<td>11.3</td>
<td>12.2</td>
<td>4.96</td>
</tr>
<tr>
<td>4</td>
<td>327 (550)</td>
<td>0.50</td>
<td>API RP 27</td>
<td>38.2 (5540)</td>
<td>4315</td>
<td>0.076</td>
<td>1.94 x 10⁻¹²</td>
<td>1.65 x 10⁻⁸</td>
<td>12.5</td>
<td>12.7</td>
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<td>5</td>
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<td>0.60</td>
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<td>39.0 (5660)</td>
<td>4526</td>
<td>0.077</td>
<td>2.23 x 10⁻¹²</td>
<td>1.45 x 10⁻⁸</td>
<td>12.7</td>
<td>12.5</td>
<td>5.37</td>
</tr>
<tr>
<td>6</td>
<td>245 (413)</td>
<td>0.75</td>
<td>ASTM C642</td>
<td>28.4 (4120)</td>
<td>5915</td>
<td>0.085</td>
<td>8.32 x 10⁻¹²</td>
<td>1.45 x 10⁻⁸</td>
<td>13.0</td>
<td>13.3</td>
<td>5.81</td>
</tr>
</tbody>
</table>

* Admixtures: 59.4 kg/m³ (100 lb/yd³) silica fume and 25.4 ml/kg of cement (30 fl.oz/cwt) HRWR (Mix 1); 13.0 ml/kg (20 fl.oz/cwt) HRWR (Mix 2); 2.2 ml/kg (3.4 fl. oz/cwt) WR (Mix 3).

** To convert from m/s to Darcy, multiply by 1.03 x 10⁵, from m/s to m², multiply by 1.02 x 10⁻⁷.

† Measured with helium porosimetric.

Adapted from Whiting (1988).
**Portland Cement Content**

The portland cement content of hardened concrete can be determined by ASTM C1084 (AASHTO T 178) standard methods. Although not frequently performed, the cement content tests are valuable in determining the cause of lack of strength gain or poor durability of concrete. Aggregate content can also be determined by these tests. However, the user of these test methods should be aware of certain admixtures and aggregate types that can alter test results. The presence of supplementary cementitious materials would be reflected in the test results.

**Supplementary Cementitious Material and Organic Admixture Content**

The presence and amount of certain supplementary cementitious materials, such as fly ash, can be determined by petrographic techniques (ASTM C856). A sample of the supplementary cementitious material used in the concrete is usually necessary as a reference to determine the type and amount of the supplementary cementitious material present. The presence and possibly the amount of an organic admixture (such as a water reducer) can be determined by infrared spectrophotometry (Hime, Mivelaz, and Connolly 1966).

**Chloride Content**

Concern with chloride-induced corrosion of reinforcing steel has led to the need to monitor and limit the chloride content of reinforced concrete. Limits on the water-soluble chloride ion content of hardened reinforced concrete are given in ACI 318. The water-soluble chloride ion content of hardened concrete can be determined in accordance with procedures outlined in ASTM C1218. In addition, ASTM C1152 can be used to determine the acid-soluble chloride content of concrete which in most cases is equivalent to total chloride.

Many of the above tests for chloride ion content also extract chloride ions from the fine and coarse aggregates that generally do not contribute to corrosion of reinforcing steel. ASTM C1524 is a standard for the analysis of aggregate for water-extractable chloride (Soxhlet method). It is used when chloride contents have been found to be significantly high in aggregates, concretes, or mortars when tested by either ASTM C1152 or C1218. Because ASTM C1524 does not pulverize the aggregates as other tests do, it theoretically measures more closely the chloride ions available for corrosion. ACI 222.1 is also a Soxhlet procedure that tests chunks of concrete for water-extractable chloride. The true meaning of results from the Soxhlet procedures is still being debated.

**Petrographic Analysis**

Petrographic analysis uses microscopic techniques described in ASTM C856 to determine the constituents of concrete, concrete quality, and the causes of inferior performance, distress, or deterioration. Estimating future performance and structural safety of concrete elements can be facilitated. Some of the items that can be reviewed by a petrographic examination include paste, aggregate, fly ash, and air content; frost and sulfate attack; alkali-aggregate reactivity; degree of hydration and carbonation; water-cement ratio; bleeding characteristics; fire damage; scaling; popouts; effect of admixture; and several other aspects. Almost any kind of concrete failure can be analyzed by petrography (St. John, Poole, and Sims 1998). However, a standard petrographic analysis is sometimes accompanied by “wet” chemical analyses, infrared spectroscopy, X-ray diffractometry, scanning electron microscopy with attendant elemental analysis, differential thermal analysis, and other analytical tools.

The Annex to ASTM C856 (AASHTO T 299) describes a technique for field and laboratory detection of alkali-silica gel. Using this method, a uranyl-acetate solution is applied to a broken or roughened concrete surface that has been dampened with distilled or deionized water. After one minute, the solution is rinsed off and the treated surface is viewed under ultraviolet light. Areas of gel fluoresce bright yellow-green. It must be recognized, however, that several materials not related to ASR in concrete can fluoresce and interfere with an accurate indication of ASR gel. Materials that fluoresce like gel include: naturally fluorescent minerals, carbonated paste, opal, and some other rock ingredients, and reactions from fly ash, silica fume, and other pozzolans. ASTM C856 includes a prescreening procedure that gives a visual impression to compensate for the effects of these materials. However, this test is considered ancillary to more definitive petrographic examinations and other tests. In addition, the toxicity and radioactivity of uranyl acetate warrants special handling and disposal procedures regarding the solution and treated concrete. Caution regarding potential eye damage from ultraviolet light also merits attention.

The Los Alamos method is a staining technique that does not require ultraviolet light or uranyl-acetate solution. Instead, solutions of sodium cobaltinitrite and rhodamine B are used to condition the specimen and produce a dark pink stain that corresponds to calcium-rich ASR gel. It should be noted that these methods can produce evidence of ASR gel without causing damage to concrete. ASR gel can be present when other mechanisms such as freeze-thaw action, sulfate attack, and other deterioration mechanisms have caused the damage. These rapid methods for detecting the presence of ASR gel are useful but their limitations must be understood. Neither of the rapid procedures is a viable substitute for petrographic examination coupled with proper field inspection (Powers 1999).
**Volume and Length Change**

Volume or length change limits are sometimes specified for certain concrete applications. Volume change is also of concern when a new ingredient is added to concrete to make sure there are no significant adverse effects. ASTM C157 (AASHTO T 160) (water and air storage methods) determines length change in concrete due to drying shrinkage, chemical reactivity, and forces other than those intentionally applied. Determination of early volume change of concrete before hardening can be performed using ASTM C827. Creep can be determined in accordance with ASTM C512. The static modulus of elasticity and Poisson’s ratio of concrete in compression can be determined by methods outlined in ASTM C469 and dynamic values of these parameters can be determined by using ASTM C215. Age of cracking due to both drying shrinkage and deformations caused by autogenous shrinkage and heat of hydration can be determined in accordance with ASTM C1581.

**Durability**

Durability refers to the ability of concrete to resist deterioration from the environment or from the service in which it is placed. Properly designed concrete should endure without significant distress throughout its service life. In addition to tests for air content and chloride content described previously, the following tests are used to measure the durability of concrete:

**Frost Resistance.** The freeze-thaw resistance of concrete is usually determined in accordance with ASTM C666 (AASHTO T 161). Samples are monitored for changes in dynamic modulus, mass, and volume over a period of 300 or more cycles of freezing and thawing. ASTM C682 is also available to evaluate frost resistance. Concrete that will be exposed to deiers as well as saturated freezing should be tested according to ASTM C672 for deicer-scaling resistance. Although ASTM C672 requires that only surface scaling be monitored, many practitioners also measure mass loss, as is done in Canada (Fig. 16-18). Concrete mixtures that perform well in ASTM C666 (AASHTO T 161) do not always perform well in ASTM C672. ASTM C666 (AASHTO T 161) and ASTM C672 are often used to evaluate innovative mix designs, or new materials such as chemical admixtures, supplementary cementsitious materials, and aggregates to determine their effect on frost and deicer resistance.

**Sulfate Resistance.** The sulfate resistance of concrete materials can be evaluated by using a saturated mortar bar test, ASTM C1012. This test is valuable in assessing the sulfate resistance of concrete that will be continuously wet, but it does not evaluate the more aggressive wet-dry cycling environment. The test can be modified to include wet-dry cycling or the U.S. Bureau of Reclamation’s wet-dry concrete prism test for sulfate attack can be used. ASTM D516 (AASHTO T 290), ASTM C1580 or the Bureau’s method (U.S. Bureau of Reclamation 1975) can be used to test soil and water for sulfate ion content to determine the severity of the sulfate exposure.

**Alkali-Silica Reactivity.** Alkali-silica reaction is best controlled at the design stage when selecting materials for use in a specific concrete mixture. Aggregate can be tested for potential alkali-silica reactivity by using the ASTM C227 mortar bar test (for moderate to highly reactive aggregate), ASTM C289 chemical method (for highly reactive aggregate), ASTM C295 (petrographic analysis), ASTM C1260 (AASHTO T 303) rapid mortar bar, and ASTM C1293 concrete prism test. Materials, such as fly ash and slag, are often used to control alkali-silica reaction; they should be evaluated by tests such as ASTM C227, ASTM C441, C1567, or C1293 to determine their effectiveness. An alternate to testing aggregate separately for potential reactivity is to test the concrete mixture to be used on the job; here one would use ASTM C1567 or C1293. A rapid 13-week version of ASTM C1293 is being developed by the University of Texas at Austin through the International Center for Aggregate Research (Touma, Fowler, Folliard, and Nelson 2001). Existing concrete structures can be evaluated for alkali-silica reaction using ASTM C856.

**Alkali-Carbonate Reactivity.** Alkali-carbonate reactivity is more rare than alkali-silica reactivity. Potential reactivity of aggregates can be evaluated by using ASTM C295, ASTM C586 rock cylinder, and ASTM C1105. Existing concrete structures can be evaluated for alkali-carbonate reactivity using ASTM C856.

**Corrosion Resistance.** The corrosion resistance of reinforced concrete is rarely tested unless unusual materials are used, concrete will be used in a very severe environment, or there is a need to evaluate the potential for inplace corrosion. Corrosion activity can be evaluated using ASTM C876.

**Abrasion Resistance.** Abrasion resistance can be determined by using ASTM C418 (sandblasting), ASTM C779 (revolving disk, dressing wheel, and ball bearing methods), ASTM C944 (rotating cutter), and ASTM C1138 (underwater test).
**Moisture Testing**

The in-place moisture content, water vapor emission rate, and relative humidity of hardened concrete are useful indicators in determining if concrete is dry enough for application of floor-covering materials and coatings. The moisture content of concrete should be low enough to avoid spalling when exposed to temperatures above the boiling point of water. Moisture related test methods fall into two general categories: either qualitative or quantitative. Qualitative tests provide an indication of the presence or absence of moisture while quantitative tests measure the amount of moisture. Qualitative tests may give a strong indication that excessive moisture is present and the floor is not ready for floor-covering materials. Quantitative tests are performed to assure that the floor is dry enough for these materials.

Qualitative moisture tests include: plastic sheet, mat bond, electrical resistance, electrical impedance, and nuclear moisture gauge tests. The plastic sheet test (ASTM D4263) uses a square sheet of clear plastic film that is taped to the slab surface and left for 24 hours to see if moisture develops under it. The plastic sheet test is unreliable. In the mat bond test, a 1-m² (9-ft²) sheet of floor covering is glued to the floor with the edges taped to the concrete for 72 hours. The force needed to remove the flooring is an indication of the slab moisture condition. Electrical resistance is measured using a moisture meter through two probes placed in contact with the concrete. Electrical impedance uses an electronic signal that is influenced by the moisture in the concrete. Nuclear moisture gauges contain high-speed neutrons that are slowed by the hydrogen atoms in water. The affect of these encounters is a measure of the moisture content of the concrete. Although the last three tests each yield a numeric test result, their value is quite limited. Experience and skill are needed to judge the trustworthiness of the devices and the test results produced by them.

Quantitative test methods include: gravimetric moisture content, moisture vapor emission rate, and relative humidity probe tests. The most direct method for determining moisture content is to dry cut a specimen from the concrete element in question, place it in a moisture proof container, and transport it to a laboratory for testing. After obtaining the specimen’s initial mass, dry the specimen in an oven at about 105°C (220°F) for 24 hours or until constant mass is achieved. The difference between the two masses divided by the dry mass, times 100, is the moisture content in percent. The moisture vapor emission rate (ASTM F1869) is the most commonly used test in the United States for measuring the readiness of concrete for application of floor coverings. The emission rate is expressed as kilograms (pounds) of moisture emitted from 93 m² (1000 ft²) in 24 hours. See Kosmatka (1985) and Kanare (2005) for more information.

Relative humidity tests are used in several countries outside the United States for measuring moisture in concrete slabs. Two British standards, BS 5325: 1996 and BS 8203: 1996 use a hygrometer or relative humidity probe sealed under an insulated, impermeable box to trap moisture in an air pocket above the floor. The probe is allowed to equilibrate for at least 72 hours or until two consecutive readings at 24 hours intervals are within the precision of the instrument (typically ± 3% RH). Acceptable relative humidity limits for the installation of floor coverings range from a maximum of 60% to 90%. It can require several months of air-drying to achieve the desired relative humidity. A method for estimating drying time to reach a specified relative humidity based on water-cement ratio, thickness of structure, number of exposed sides, relative humidity, temperature and curing conditions can be found in Hedenblad (1997), Hedenblad (1998), and Tarr and Farny (2008).

**Carbonation**

The depth or degree of carbonation can be determined by petrographic techniques (ASTM C856) through the observation of calcium carbonate—the primary chemical product of carbonation. In addition, a phenolphthalein color test can be used to estimate the depth of carbonation by testing the pH of concrete (carbonation reduces pH). Upon application of the phenolphthalein solution to a freshly fractured or freshly cut surface of concrete, noncarbonated areas turn red or purple while carbonated areas remain colorless. (Fig. 16-19). The phenolphthalein indicator when observed against hardened paste changes color at a pH of 9.0 to 9.5. The pH of good quality noncarbonated concrete without admixtures is usually greater than 12.5. For more information, see “pH Testing Methods” below, and see Verbeck (1958), Steinour (1964), and Campbell, Sturm, and Kosmatka (1991).
Each method has limitations and caution should be exercised against acceptance of nondestructive test results as having a constant correlation to the traditional compression test; for example, empirical correlations must be developed prior to use (Malhotra 1976, NRMCA 1979, Malhotra 1984, Clifton 1985, Malhotra and Carino 1991).

An NDT program may be undertaken for a variety of purposes regarding the strength or condition of hardened concrete, including:

- Determination of in-place concrete strength
- Monitoring rate of concrete strength gain
- Location of nonhomogeneity, such as voids or honeycombing in concrete
- Determination of relative strength of comparable members
- Evaluation of concrete cracking and delaminations
- Evaluation of damage from mechanical or chemical forces
- Steel reinforcement location, size, and corrosion activity
- Member dimensions

Irrespective of the type of NDT test used, adequate and reliable correlation data with standard 28-day compressive strength data is usually necessary to evaluate the accuracy of the NDT method. In addition, correlation to in-place compressive strengths using drilled cores from one or two locations can provide guidance in interpreting NDT test results; these can then be used to survey larger portions of the structure. Care should be taken to consider the influence that varying sizes and locations of structural elements can have on the NDT test being used.

Rebound Hammer Tests. The Schmidt rebound hammer (Fig. 16-20) is essentially a surface-hardness tester that provides a quick, simple means of checking concrete uniformity. It measures the rebound of a spring-loaded...
plunger after it has struck a smooth concrete surface. The rebound number reading gives an indication of the compressive strength and stiffness of the concrete. Two different concrete mixtures having the same strength but different stiffnesses will yield different readings. In view of this, an understanding of the factors influencing the accuracy of the test is required.

The results of a Schmidt rebound hammer test (ASTM C805) are affected by surface smoothness, size, shape, and rigidity of the specimen; age and moisture condition of the concrete; type of coarse aggregate; and degree of carbonation of the concrete surface. When these limitations are recognized; and the hammer is calibrated for the particular materials used in the concrete (Fig. 16-21) by comparison with cores or cast specimens, then this instrument can be useful for determining the relative compressive strength and uniformity of concrete in the structure.

**Penetration Tests.** The Windsor probe (ASTM C803), like the rebound hammer, is basically a hardness tester that provides a quick means of determining the relative strength of the concrete. The equipment consists of a powder-actuated gun that drives a hardened alloy probe into the concrete (Fig. 16-22). The exposed length of the probe is measured and related by a calibration table to the compressive strength of the concrete.

The results of the Windsor-probe test will be influenced by surface smoothness of the concrete and the type and hardness of aggregate used. Therefore, to improve accuracy, a calibration table or curve for the particular concrete to be tested should be made, usually from cores or cast specimens.

Both the rebound hammer and the probe damage the concrete surface to some extent. The rebound hammer leaves a small indentation on the surface; the probe leaves a small hole and may cause minor cracking and small craters similar to popouts.

**Maturity Tests.** The maturity principle is that strength gain is a function of time and temperature. ASTM C1074 generates a maturity index that is based on temperature and time factors. The estimated strength depends on properly determining the strength-maturity function for a particular concrete mixture. The device uses thermocouples or thermistors placed in the concrete and connected to strip-chart recorders or digital data-loggers that record concrete temperature as a function of time. The temperature in relation to time data is correlated to compression tests performed on cylindrical specimens to generate a temperature-time versus strength curve that is used to estimate in-place concrete strength.

**Pullout Tests.** A pullout test (ASTM C900) involves casting the enlarged end of a steel rod in the concrete to be tested and then measuring the force required to pull it out (Fig. 16-23). The test measures the direct shear strength of the concrete. This in turn is correlated with the compressive strength; thus a measurement of the in-place compressive strength is made.

**Break-Off Tests.** The break-off test (ASTM C1150) determines the in-place strength of the concrete by breaking off an in situ cylindrical concrete specimen at a failure plane parallel to the finished surface of the concrete element. A break-off number is generated and assessed in relation to the strength of the concrete. Similar to pullout tests, the relationship between break-off test numbers and compres-
C1040 (AASHTO T 271) procedures use gamma radiation to determine the density of unhardened and hardened concrete in place.

Battery-operated magnetic detection devices, like the pachometer or covermeter, are available to measure the depth of reinforcement in concrete and to detect the position of rebars. Electrical-resistivity equipment is being developed to estimate the thickness of concrete pavement slabs.

A microwave-absorption method has been developed to determine the moisture content of porous building materials such as concrete. Acoustic-emission techniques show promise for studying load levels in structures and for locating the origin of cracking.

Ground-penetrating (short-pulse) radar is a rapid technique for nondestructive detection of delaminations and other types of defects in overlaid reinforced concrete decks. It also shows potential for monitoring strength development in concrete, measuring the thickness of concrete members, and locating reinforcement.

Infrared thermographic techniques are used to detect and show, both large and small, internal voids, delaminations and cracks in bridges, highway pavements, garage decks, buildings and other structural elements exposed to direct sunlight. For more information, see Malhotra and Carino (1991). ACI Committee 228 presents additional information on these and other nondestructive test methods.

Finally, acoustic impact methods also employ simple hammer and chain drag soundings that are low-cost accurate tests used to identify delaminated areas of concrete. Hammer soundings can be used on either vertical or horizontal surfaces, but are usually limited to small areas of delaminations. These areas are identified by striking the surface of the concrete with a hammer while listening for either a ringing or hollow sound. Dragging either a single chain, in small areas, or for larger areas, a T-bar with or without wheels having four or more chains attached are also used to identify delaminated concrete (ASTM D4580). Approximately one meter (3 ft) of chain is in contact with the concrete during chain drag soundings. The sound emitted indicates whether the concrete is delaminated or not. Chain drag soundings are usually limited to horizontal surfaces that have a relatively rough texture. Smooth concrete may not bounce the chain links enough to generate adequate sound to detect delaminated areas. Note that corrosion of reinforcing bars in the area of delaminated concrete will probably extend beyond the boundary identified as delaminated.

Table 16-3 lists several nondestructive test methods along with main applications.
## Table 16-3. Nondestructive Test Methods for Concrete

<table>
<thead>
<tr>
<th>Concrete properties</th>
<th>Recommended NDT methods</th>
<th>Possible NDT methods</th>
</tr>
</thead>
<tbody>
<tr>
<td>Strength</td>
<td>Penetration probe, Rebound hammer, Pullout methods, Break off</td>
<td>Ultrasound pulse echo, Visual examination</td>
</tr>
<tr>
<td>General quality and uniformity</td>
<td>Penetration probe, Rebound hammer, Ultrasound pulse velocity, Gamma radiography</td>
<td>Radar, Gamma radiography, Ultrasound pulse echo</td>
</tr>
<tr>
<td>Thickness</td>
<td>Ultrasonic pulse velocity</td>
<td></td>
</tr>
<tr>
<td>Stiffness</td>
<td>Ultrasonic pulse velocity</td>
<td>Proof loading (load-deflection)</td>
</tr>
<tr>
<td>Density</td>
<td>Gamma radiography, Ultrasonic pulse velocity</td>
<td>Neutron density gage</td>
</tr>
<tr>
<td>Rebar size and location</td>
<td>Covertmeter (pachometer), Gamma radiography</td>
<td>X-ray radiography, Ultrasonic pulse echo, Radar</td>
</tr>
<tr>
<td>Corrosion state of reinforcing steel</td>
<td>Electrical potential measurement</td>
<td></td>
</tr>
<tr>
<td>Presence of subsurface voids</td>
<td>Acoustic impact, Gamma radiography, Ultrasonic pulse velocity</td>
<td>Infrared thermography, X-ray radiography, Ultrasonic pulse echo, Radar, Resonant frequency testing</td>
</tr>
<tr>
<td>Structural integrity of concrete</td>
<td>Proof loading (load-deflection)</td>
<td>Proof testing using acoustic emission</td>
</tr>
</tbody>
</table>

Adapted from ACI Subcommittee 364 (1994) and Clifton (1985).

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CHAPTER 17
High-Performance Concrete

High-performance concrete (HPC) exceeds the properties and constructability of normal concrete. Normal and special materials are used to make these specially designed concretes that must meet a combination of performance requirements. Special mixing, placing, and curing practices may be needed to produce and handle high-performance concrete. Extensive performance tests are usually required to demonstrate compliance with specific project needs (ASCE 1993, Russell 1999, and Bickley and Mitchell 2001). High-performance concrete has been primarily used in tunnels, bridges, and tall buildings for its strength, durability, and high modulus of elasticity (Fig. 17-1). It has also been used in shotcrete repair, poles, parking garages, and agricultural applications.

High-performance concrete characteristics are developed for particular applications and environments; some of the properties that may be required include:

- High strength
- High early strength
- High modulus of elasticity
- High abrasion resistance
- High durability and long life in severe environments
- Low permeability and diffusion
- Resistance to chemical attack
- High resistance to frost and deicer scaling damage
- Toughness and impact resistance
- Volume stability
- Ease of placement
- Compaction without segregation
- Inhibition of bacterial and mold growth

High-performance concretes are made with carefully selected high-quality ingredients and optimized mixture designs; these are batched, mixed, placed, compacted and cured to the highest industry standards. Typically, such concretes will have a low water-cementing materials ratio of 0.20 to 0.45. Plasticizers are usually used to make these concretes fluid and workable.

High-performance concrete almost always has a higher strength than normal concrete. However, strength is not always the primary required property. For example, a normal strength concrete with very high durability and very low permeability is considered to have high-performance properties. Bickley and Fung (2001) demonstrated that 40 MPa (6,000 psi) high-performance concrete for bridges could be economically made while...
Table 17-1. Materials Used in High-Performance Concrete

<table>
<thead>
<tr>
<th>Material</th>
<th>Primary contribution/Desired property</th>
</tr>
</thead>
<tbody>
<tr>
<td>Portland cement</td>
<td>Cementing material/durability</td>
</tr>
<tr>
<td>Blended cement</td>
<td>Cementing material/durability/high strength</td>
</tr>
<tr>
<td>Fly ash</td>
<td>Cementing material/durability/high strength</td>
</tr>
<tr>
<td>Slag</td>
<td>Cementing material/durability/high strength</td>
</tr>
<tr>
<td>Silica fume</td>
<td>Cementing material/durability/high strength</td>
</tr>
<tr>
<td>Slag</td>
<td>Cementing material/durability/high strength</td>
</tr>
<tr>
<td>Metakaolin</td>
<td>Cementing material/durability/high strength</td>
</tr>
<tr>
<td>Calcined shale</td>
<td>Cementing material/durability/high strength</td>
</tr>
<tr>
<td>Superplasticizers</td>
<td>Flowability</td>
</tr>
<tr>
<td>High-range water reducers</td>
<td>Reduce water to cement ratio</td>
</tr>
<tr>
<td>Hydration control admixtures</td>
<td>Control setting</td>
</tr>
<tr>
<td>Retarders</td>
<td>Control setting</td>
</tr>
<tr>
<td>Accelerators</td>
<td>Accelerate setting</td>
</tr>
<tr>
<td>Corrosion inhibitors</td>
<td>Control steel corrosion</td>
</tr>
<tr>
<td>Water reducers</td>
<td>Reduce cement and water content</td>
</tr>
<tr>
<td>Shrinkage reducers</td>
<td>Reduce shrinkage</td>
</tr>
<tr>
<td>ASR inhibitors</td>
<td>Control alkali-silica reactivity</td>
</tr>
<tr>
<td>Polymer/latex modifiers</td>
<td>Durability</td>
</tr>
<tr>
<td>Optimally graded aggregate</td>
<td>Improve workability and reduce paste demand</td>
</tr>
</tbody>
</table>

Table 17-2. Selected Properties of High-Performance Concrete

<table>
<thead>
<tr>
<th>Property</th>
<th>Test method</th>
<th>Criteria that may be specified</th>
</tr>
</thead>
<tbody>
<tr>
<td>High strength</td>
<td>ASTM C39 (AASHTO T 22)</td>
<td>70 to 140 MPa (10,000 to 20,000 psi) at 28 to 91 days</td>
</tr>
<tr>
<td>High-early compressive strength</td>
<td>ASTM C39 (AASHTO T 22)</td>
<td>20 to 28 MPa (3000 to 4000 psi) at 3 to 12 hours or 1 to 3 days</td>
</tr>
<tr>
<td>High-early flexural strength</td>
<td>ASTM C78 (AASHTO T 97)</td>
<td>2 to 4 MPa (300 to 600 psi) at 3 to 12 hours or 1 to 3 days</td>
</tr>
<tr>
<td>Abrasion resistance</td>
<td>ASTM C944</td>
<td>0 to 1 mm depth of wear</td>
</tr>
<tr>
<td>Low permeability</td>
<td>ASTM C1202 (AASHTO T 277)</td>
<td>500 to 2000 coulombs</td>
</tr>
<tr>
<td>Chloride penetration</td>
<td>ASTM C1543 (AASHTO T 259 &amp; T 260)</td>
<td>Less than 0.07% Cl at 6 months</td>
</tr>
<tr>
<td>High resistivity</td>
<td>ASTM G59</td>
<td></td>
</tr>
<tr>
<td>Low absorption</td>
<td>ASTM C642</td>
<td>2% to 5%</td>
</tr>
<tr>
<td>Low diffusion coefficient</td>
<td>ASTM C1556</td>
<td>1000 x 10^{-13} m/s</td>
</tr>
<tr>
<td>Resistance to chemical attack</td>
<td>Expose concrete to saturated solution in wet/dry environment</td>
<td>No deterioration after 1 year</td>
</tr>
<tr>
<td>Sulfate attack</td>
<td>ASTM C1012</td>
<td>0.10% max. expansion at 6 months for moderate sulfate exposures or 0.5% max. expansion at 6 months for severe sulfate exposure</td>
</tr>
<tr>
<td>High modulus of elasticity</td>
<td>ASTM C469</td>
<td>More than 40 GPa (5.8 million psi) (Aïtcin 1998)</td>
</tr>
<tr>
<td>High resistance to freezing and thawing damage</td>
<td>ASTM C666, Procedure A</td>
<td>Durability factor of 95 to 100 at 300 to 1000 cycles (max. mass loss or expansion can also be specified)</td>
</tr>
<tr>
<td>High resistance to deicer scaling</td>
<td>ASTM C672</td>
<td>Scale rating of 0 to 1 or mass loss of 0 to 0.5 kg/m² after 50 to 300 cycles</td>
</tr>
<tr>
<td>Low shrinkage</td>
<td>ASTM C157</td>
<td>Less than 400 millionths (Aïtcin 1998)</td>
</tr>
<tr>
<td>Low creep</td>
<td>ASTM C512</td>
<td>Less than normal concrete</td>
</tr>
</tbody>
</table>

High-early-strength concrete, also called fast-track concrete, achieves its specified strength at an earlier age than normal concrete. The time period in which a specified strength should be achieved may range from a few
hours (or even minutes) to several days. High-early-strength can be attained by using traditional concrete ingredients and concreting practices, although sometimes special materials or techniques are needed.

High-early-strength can be obtained by using one or a combination of the following, depending on the age at which the specified strength must be achieved and on job conditions:

1. Type III or HE high-early-strength cement
2. High cement content (400 to 600 kg/m³ or 675 to 1000 lb/yd³)

Table 17-3 (Metric). Typical High-Performance Concretes Used in Structures

<table>
<thead>
<tr>
<th>Mixture number</th>
<th>1</th>
<th>2</th>
<th>3</th>
<th>4</th>
<th>5</th>
<th>6</th>
</tr>
</thead>
<tbody>
<tr>
<td>Water, kg/m³</td>
<td>151</td>
<td>145</td>
<td>135</td>
<td>145</td>
<td>130</td>
<td>130</td>
</tr>
<tr>
<td>Cement, kg/m³</td>
<td>311</td>
<td>398</td>
<td>500</td>
<td>335</td>
<td>513</td>
<td>315</td>
</tr>
<tr>
<td>Fly ash, kg/m³</td>
<td>31</td>
<td>45</td>
<td>—</td>
<td>—</td>
<td>—</td>
<td>40</td>
</tr>
<tr>
<td>Slag, kg/m³</td>
<td>47</td>
<td>—</td>
<td>—</td>
<td>125</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td>Silica fume, kg/m³</td>
<td>16</td>
<td>32</td>
<td>30</td>
<td>40</td>
<td>43</td>
<td>23</td>
</tr>
<tr>
<td>Coarse aggregate, kg/m³</td>
<td>1068</td>
<td>1030</td>
<td>1100</td>
<td>1130</td>
<td>1080</td>
<td>1140</td>
</tr>
<tr>
<td>Fine aggregate, kg/m³</td>
<td>676</td>
<td>705</td>
<td>700</td>
<td>695</td>
<td>685</td>
<td>710</td>
</tr>
<tr>
<td>Water reducer, L/m³</td>
<td>1.6</td>
<td>1.7</td>
<td>—</td>
<td>1.0</td>
<td>—</td>
<td>1.5</td>
</tr>
<tr>
<td>Retarder, L/m³</td>
<td>—</td>
<td>—</td>
<td>1.8</td>
<td>—</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td>Air, %</td>
<td>7 ± 1.5</td>
<td>5 – 8</td>
<td>—</td>
<td>—</td>
<td>—</td>
<td>5.5</td>
</tr>
<tr>
<td>HRWR or plasticizer, L/m³</td>
<td>2.1</td>
<td>3</td>
<td>14</td>
<td>6.5</td>
<td>15.7</td>
<td>5.0</td>
</tr>
<tr>
<td>Water to cementing materials ratio</td>
<td>0.37</td>
<td>0.30</td>
<td>0.27</td>
<td>0.29</td>
<td>0.25</td>
<td>0.34</td>
</tr>
<tr>
<td>Comp. strength at 28 days, MPa</td>
<td>59</td>
<td>—</td>
<td>93</td>
<td>99</td>
<td>119</td>
<td>—</td>
</tr>
<tr>
<td>Comp. strength at 91 days, MPa</td>
<td>—</td>
<td>60</td>
<td>107</td>
<td>104</td>
<td>145</td>
<td>—</td>
</tr>
</tbody>
</table>

4. BCE Place Phase 2, Toronto, 1993.

* Originally used a blended cement containing silica fume. Portland cement and silica fume quantities have been separated for comparison purposes.

Table 17-3 (Inch-Pound Units). Typical High-Performance Concretes Used in Structures

<table>
<thead>
<tr>
<th>Mixture number</th>
<th>1</th>
<th>2</th>
<th>3</th>
<th>4</th>
<th>5</th>
<th>6</th>
</tr>
</thead>
<tbody>
<tr>
<td>Water, lb/yd³</td>
<td>254</td>
<td>244</td>
<td>227</td>
<td>244</td>
<td>219</td>
<td>219</td>
</tr>
<tr>
<td>Cement, lb/yd³</td>
<td>525</td>
<td>671*</td>
<td>843</td>
<td>565*</td>
<td>865</td>
<td>531</td>
</tr>
<tr>
<td>Fly ash, lb/yd³</td>
<td>53</td>
<td>76</td>
<td>—</td>
<td>—</td>
<td>—</td>
<td>67</td>
</tr>
<tr>
<td>Slag, lb/yd³</td>
<td>79</td>
<td>—</td>
<td>—</td>
<td>211</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td>Silica fume, lb/yd³</td>
<td>27</td>
<td>54*</td>
<td>51</td>
<td>67*</td>
<td>72</td>
<td>39</td>
</tr>
<tr>
<td>Coarse aggregate, lb/yd³</td>
<td>1800</td>
<td>1736</td>
<td>1854</td>
<td>1905</td>
<td>1820</td>
<td>1921</td>
</tr>
<tr>
<td>Fine aggregate, lb/yd³</td>
<td>1140</td>
<td>1188</td>
<td>1180</td>
<td>1171</td>
<td>1155</td>
<td>1197</td>
</tr>
<tr>
<td>Water reducer, oz/yd³</td>
<td>41</td>
<td>47</td>
<td>—</td>
<td>27</td>
<td>—</td>
<td>38</td>
</tr>
<tr>
<td>Retarder, oz/yd³</td>
<td>—</td>
<td>—</td>
<td>48</td>
<td>—</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td>Air, %</td>
<td>7 ± 1.5</td>
<td>5 – 8</td>
<td>—</td>
<td>—</td>
<td>—</td>
<td>5.5</td>
</tr>
<tr>
<td>HRWR or plasticizer, oz/yd³</td>
<td>55</td>
<td>83</td>
<td>375</td>
<td>175</td>
<td>420</td>
<td>131</td>
</tr>
<tr>
<td>Water to cementing materials ratio</td>
<td>0.37</td>
<td>0.30</td>
<td>0.27</td>
<td>0.29</td>
<td>0.25</td>
<td>0.34</td>
</tr>
<tr>
<td>Comp. strength at 28 days, psi</td>
<td>8,590</td>
<td>—</td>
<td>13,500</td>
<td>14,360</td>
<td>17,250</td>
<td>—</td>
</tr>
<tr>
<td>Comp. strength at 91 days, psi</td>
<td>—</td>
<td>8700</td>
<td>15,300</td>
<td>15,080</td>
<td>21,000</td>
<td>—</td>
</tr>
</tbody>
</table>

4. BCE Place Phase 2, Toronto, 1993.

* Originally used a blended cement containing silica fume. Portland cement and silica fume quantities have been separated for comparison purposes.
3. Low water-cementing materials ratio (0.20 to 0.45 by mass)
4. Higher freshly mixed concrete temperature
5. Higher curing temperature
6. Chemical admixtures
7. Silica fume (or other supplementary cementing materials)
8. Steam or autoclave curing
9. Insulation to retain heat of hydration
10. Special rapid hardening cements.

High-early-strength concrete is used for prestressed concrete to allow for early stressing; precast concrete for rapid production of elements; high-speed cast-in-place construction; rapid form reuse; cold-weather construction; rapid repair of pavements to reduce traffic downtime; fast-track paving; and several other uses.

In fast-track paving, use of high-early-strength mixtures allows traffic to open within a few hours after concrete is placed. An example of a fast-track concrete mixture used for a bonded concrete highway overlay consisted of 380 kg (640 lb) of Type III cement, 42 kg (70 lb) of Type C fly ash, 6½% air, a water reducer, and a water-to-cementing materials ratio of 0.4. Strength data for this 40-mm (1½-in.) slump concrete are given in Table 17-4. Figs. 17-2 and 17-3 illustrate early strength development of concretes designed to open to traffic within 4 hours after placement. Fig. 17-4 illustrates the benefits of blanket curing to develop early strength for patching or fast-track applications.

When designing early-strength mixtures, strength development is not the only criteria that should be evaluated; durability, early stiffening, autogenous shrinkage, drying shrinkage, temperature rise, and other properties also should be evaluated for compatibility with the project. Special curing procedures, such as fogging, may be needed to control plastic shrinkage cracking.

### Table 17-4. Strength Data for Fast-Track Bonded Overlay

<table>
<thead>
<tr>
<th>Age</th>
<th>Compressive strength, MPa (psi)</th>
<th>Flexural strength, MPa (psi)</th>
<th>Bond strength, MPa (psi)</th>
</tr>
</thead>
<tbody>
<tr>
<td>4 hours</td>
<td>1.7 (252)</td>
<td>0.9 (126)</td>
<td>0.9 (120)</td>
</tr>
<tr>
<td>6 hours</td>
<td>7.0 (1020)</td>
<td>2.0 (287)</td>
<td>1.1 (160)</td>
</tr>
<tr>
<td>8 hours</td>
<td>13.0 (1883)</td>
<td>2.7 (393)</td>
<td>1.4 (200)</td>
</tr>
<tr>
<td>12 hours</td>
<td>17.6 (2546)</td>
<td>3.4 (494)</td>
<td>1.6 (225)</td>
</tr>
<tr>
<td>18 hours</td>
<td>20.1 (2920)</td>
<td>4.0 (574)</td>
<td>1.7 (250)</td>
</tr>
<tr>
<td>24 hours</td>
<td>23.9 (3467)</td>
<td>4.2 (604)</td>
<td>2.1 (302)</td>
</tr>
<tr>
<td>7 days</td>
<td>34.2 (4960)</td>
<td>5.0 (722)</td>
<td>2.1 (309)</td>
</tr>
<tr>
<td>14 days</td>
<td>36.5 (5295)</td>
<td>5.7 (825)</td>
<td>2.3 (328)</td>
</tr>
<tr>
<td>28 days</td>
<td>40.7 (5900)</td>
<td>5.7 (830)</td>
<td>2.5 (359)</td>
</tr>
</tbody>
</table>

Adapted from Knutson and Riley 1987
HIGH-STRENGTH CONCRETE

The definition of high strength changes over the years as concrete strength used in the field increases. This publication considers high-strength concrete (HSC) to have a strength significantly beyond what is used in normal practice. For example, today about 90% of ready mixed concrete has a 28-day specified compressive strength ranging from 20 MPa (3000 psi) to 40 MPa (6000 psi), with most of it between 28 MPa (4000 psi) and 35 MPa (5000 psi). Therefore, HSC considered here has a design strength of at least 70 MPa (10,000 psi).

Most high-strength concrete applications are designed for compressive strengths of 70 MPa (10,000 psi) or greater as shown in Tables 17-3 and 17-5. For strengths of 70 MPa (10,000 psi) and higher, stringent application of the best practices is required. Compliance with the guidelines and

Table 17-5 (Metric). Mixture Proportions and Properties of Commercially Available High-Strength Concrete (Burg and Ost 1994)

<table>
<thead>
<tr>
<th>Units per m³</th>
<th>Mix number</th>
<th>1</th>
<th>2</th>
<th>3</th>
<th>4</th>
<th>5</th>
<th>6</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cement, Type I, kg</td>
<td>564</td>
<td>475</td>
<td>487</td>
<td>564</td>
<td>475</td>
<td>327</td>
<td></td>
</tr>
<tr>
<td>Silica fume, kg</td>
<td>—</td>
<td>24</td>
<td>47</td>
<td>89</td>
<td>74</td>
<td>27</td>
<td></td>
</tr>
<tr>
<td>Fly ash, kg</td>
<td>—</td>
<td>59</td>
<td>—</td>
<td>—</td>
<td>104</td>
<td>87</td>
<td></td>
</tr>
<tr>
<td>Coarse aggregate SSD (12.5 mm crushed limestone), kg</td>
<td>1068</td>
<td>1068</td>
<td>1068</td>
<td>1068</td>
<td>1068</td>
<td>1121</td>
<td></td>
</tr>
<tr>
<td>Fine aggregate SSD, kg</td>
<td>647</td>
<td>659</td>
<td>676</td>
<td>593</td>
<td>593</td>
<td>742</td>
<td></td>
</tr>
<tr>
<td>HRWR Type F, liters</td>
<td>11.6</td>
<td>11.6</td>
<td>11.22</td>
<td>20.11</td>
<td>16.44</td>
<td>6.3</td>
<td></td>
</tr>
<tr>
<td>HRWR Type G, liters</td>
<td>—</td>
<td>—</td>
<td>—</td>
<td>—</td>
<td>—</td>
<td>3.24</td>
<td></td>
</tr>
<tr>
<td>Retarder, Type D, liters</td>
<td>1.12</td>
<td>1.05</td>
<td>0.97</td>
<td>1.46</td>
<td>1.5</td>
<td>—</td>
<td></td>
</tr>
<tr>
<td>Water to cementing materials ratio</td>
<td>0.28</td>
<td>0.29</td>
<td>0.29</td>
<td>0.22</td>
<td>0.23</td>
<td>0.32</td>
<td></td>
</tr>
</tbody>
</table>

Fresh concrete properties

| Slump, mm | 197 | 248 | 216 | 254 | 235 | 203 |
| Density, kg/ m³ | 2451 | 2453 | 2433 | 2486 | 2459 | 2454 |
| Air content, % | 1.6 | 0.7 | 1.3 | 1.1 | 1.4 | 1.2 |
| Concrete temp., °C | 24 | 24 | 18 | 17 | 17 | 23 |

Compressive strength, 100 x 200-mm moist-cured cylinders

| 3 days, MPa | 57 | 54 | 55 | 72 | 53 | 43 |
| 7 days, MPa | 67 | 71 | 71 | 92 | 77 | 63 |
| 28 days, MPa | 79 | 92 | 90 | 117 | 100 | 85 |
| 56 days, MPa | 84 | 94 | 95 | 122 | 116 | — |
| 91 days, MPa | 88 | 105 | 96 | 124 | 120 | 92 |
| 182 days, MPa | 97 | 105 | 97 | 128 | 120 | — |
| 426 days, MPa | 103 | 118 | 100 | 133 | 119 | — |
| 1085 days, MPa | 115 | 122 | 115 | 150 | 132 | — |

Modulus of elasticity in compression, 100 x 200-mm moist-cured cylinders

| 91 days, GPa | 50.6 | 49.9 | 50.1 | 56.5 | 53.4 | 47.9 |

Drying shrinkage, 75 by 75 x 285-mm prisms

| 7 days, millioths | 193 | 123 | 100 | 87 | 137 | — |
| 28 days, millioths | 400 | 287 | 240 | 203 | 233 | — |
| 90 days, millioths | 573 | 447 | 383 | 320 | 340 | — |
| 369 days, millioths | 690 | 577 | 520 | 453 | 467 | — |
| 1075 days, millioths | 753 | 677 | 603 | 527 | 523 | — |
Cementing materials are usually incorporated into the concrete mixture. This produces additional benefits in the form of reduced heat generation during hydration.

With use of low-slump or no-slump mixes, high-compressive-strength concrete is produced routinely under careful control in precast and prestressed concrete plants. These stiff mixes are placed in ruggedly-built forms and consolidated by prolonged vibration or shock methods. However, cast-in-place concrete uses more fragile forms that do not permit the same compaction procedures, hence more workable concretes are necessary to achieve the required compaction and to avoid segregation.

**Table 17-5 (Inch-Pound Units). Mixture Proportions and Properties of Commercially Available High-Strength Concrete (Burgand Ost 1994)**

<table>
<thead>
<tr>
<th>Units per yd³</th>
<th>Mix number</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>1</td>
</tr>
<tr>
<td>Cement, Type I, lb</td>
<td>950</td>
</tr>
<tr>
<td>Silica fume, lb</td>
<td>—</td>
</tr>
<tr>
<td>Fly ash, lb</td>
<td>—</td>
</tr>
<tr>
<td>Coarse aggregate SSD (% in. crushed limestone), lb</td>
<td>1800</td>
</tr>
<tr>
<td>Fine aggregate SSD, lb</td>
<td>1090</td>
</tr>
<tr>
<td>HRWR Type F, fl oz</td>
<td>300</td>
</tr>
<tr>
<td>HRWR Type G, fl oz</td>
<td>—</td>
</tr>
<tr>
<td>Retarder, Type D, fl oz</td>
<td>29</td>
</tr>
<tr>
<td>Water to cementing materials ratio</td>
<td>0.28</td>
</tr>
</tbody>
</table>

**Fresh concrete properties**

<table>
<thead>
<tr>
<th></th>
<th>7%</th>
<th>9%</th>
<th>8½</th>
<th>10</th>
<th>9%</th>
<th>8%</th>
</tr>
</thead>
<tbody>
<tr>
<td>Slump, in.</td>
<td>7¾</td>
<td>9¾</td>
<td>8½</td>
<td>10</td>
<td>9½</td>
<td>8%</td>
</tr>
<tr>
<td>Density, lb/ft³</td>
<td>153.0</td>
<td>153.1</td>
<td>151.9</td>
<td>155.2</td>
<td>153.5</td>
<td>153.2</td>
</tr>
<tr>
<td>Air content, %</td>
<td>1.6</td>
<td>0.7</td>
<td>1.3</td>
<td>1.1</td>
<td>1.4</td>
<td>1.2</td>
</tr>
<tr>
<td>Concrete temp., °F</td>
<td>75</td>
<td>75</td>
<td>65</td>
<td>63</td>
<td>62</td>
<td>74</td>
</tr>
</tbody>
</table>

**Compressive strength, 4 x 8-in. moist-cured cylinders**

<table>
<thead>
<tr>
<th></th>
<th>3 days, psi</th>
<th>7 days, psi</th>
<th>28 days, psi</th>
<th>56 days, psi</th>
<th>91 days, psi</th>
<th>182 days, psi</th>
<th>426 days, psi</th>
<th>1085 days, psi</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>8,220</td>
<td>7,900</td>
<td>7,970</td>
<td>10,430</td>
<td>7,630</td>
<td>6,170</td>
<td>14,910</td>
<td>16,720</td>
</tr>
<tr>
<td></td>
<td>9,660</td>
<td>10,230</td>
<td>10,360</td>
<td>13,280</td>
<td>11,150</td>
<td>9,170</td>
<td>17,100</td>
<td>17,730</td>
</tr>
<tr>
<td></td>
<td>11,460</td>
<td>13,300</td>
<td>13,070</td>
<td>17,000</td>
<td>14,530</td>
<td>12,270</td>
<td>19,900</td>
<td>21,170</td>
</tr>
<tr>
<td></td>
<td>12,230</td>
<td>13,660</td>
<td>13,840</td>
<td>17,630</td>
<td>16,760</td>
<td>—</td>
<td>17,760</td>
<td>19,190</td>
</tr>
<tr>
<td></td>
<td>12,800</td>
<td>15,170</td>
<td>13,950</td>
<td>18,030</td>
<td>17,350</td>
<td>13,310</td>
<td>21,230</td>
<td>22,850</td>
</tr>
<tr>
<td></td>
<td>14,110</td>
<td>15,160</td>
<td>14,140</td>
<td>18,590</td>
<td>17,400</td>
<td>—</td>
<td>19,440</td>
<td>21,200</td>
</tr>
<tr>
<td></td>
<td>14,910</td>
<td>17,100</td>
<td>14,560</td>
<td>19,230</td>
<td>17,290</td>
<td>—</td>
<td>20,140</td>
<td>21,950</td>
</tr>
<tr>
<td></td>
<td>16,720</td>
<td>17,730</td>
<td>16,650</td>
<td>21,750</td>
<td>19,190</td>
<td>—</td>
<td>22,490</td>
<td>24,200</td>
</tr>
</tbody>
</table>

**Modulus of elasticity in compression, 4 x 8-in. moist-cured cylinders**

<table>
<thead>
<tr>
<th></th>
<th>91 days, million psi</th>
<th>7.34</th>
<th>7.24</th>
<th>7.27</th>
<th>8.20</th>
<th>7.75</th>
<th>6.95</th>
</tr>
</thead>
</table>

**Drying shrinkage, 3 x 3 by 11.5-in. prisms**

<table>
<thead>
<tr>
<th></th>
<th>7 days, millionths</th>
<th>193</th>
<th>123</th>
<th>100</th>
<th>87</th>
<th>137</th>
<th>—</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>28 days, millionths</td>
<td>400</td>
<td>287</td>
<td>240</td>
<td>203</td>
<td>233</td>
<td>—</td>
</tr>
<tr>
<td></td>
<td>90 days, millionths</td>
<td>573</td>
<td>447</td>
<td>383</td>
<td>320</td>
<td>340</td>
<td>—</td>
</tr>
<tr>
<td></td>
<td>369 days, millionths</td>
<td>690</td>
<td>577</td>
<td>520</td>
<td>453</td>
<td>467</td>
<td>—</td>
</tr>
<tr>
<td></td>
<td>1075 days, millionths</td>
<td>753</td>
<td>677</td>
<td>603</td>
<td>527</td>
<td>523</td>
<td>—</td>
</tr>
</tbody>
</table>

With use of low-slump or no-slump mixes, high-compressive-strength concrete is produced routinely under careful control in precast and prestressed concrete plants. These stiff mixes are placed in ruggedly-built forms and consolidated by prolonged vibration or shock methods. However, cast-in-place concrete uses more fragile forms that do not permit the same compaction procedures, hence more workable concretes are necessary to achieve the required compaction and to avoid segrega-
tion and honeycomb. Superplasticizing admixtures are invariably added to HPC mixtures to produce workable and often flowable mixtures.

Production of high-strength concrete may or may not require the purchase of special materials. The producer must know the factors affecting compressive strength and know how to vary those factors for best results. Each variable should be analyzed separately in developing a mix design. When an optimum or near optimum is established for each variable, it should be incorporated as the remaining variables are studied. An optimum mix design is then developed keeping in mind the economic advantages of using locally available materials. Many of the items discussed below also apply to most high-performance concretes.

Cement

Selection of cement for high-strength concrete should not be based only on mortar-cube tests but should also include tests of comparative strengths of concrete at 28, 56, and 91 days. A cement that yields the highest concrete compressive strength at extended ages (91 days) is preferable. For high-strength concrete, a cement should produce a minimum 7-day mortar-cube strength of approximately 30 MPa (4350 psi).

Trial mixtures with cement contents between 400 and 550 kg/m³ (675 to 930 lb/yd³) should be made for each cement being considered for the project. Amounts will vary depending on target strengths. Other than decreases in sand content as cement content increases, the trial mixtures should be as nearly identical as possible.

Supplementary Cementing Materials

Fly ash, silica fume, or slag are often mandatory in the production of high-strength concrete; the strength gain obtained with these supplementary cementing materials cannot be attained by using additional cement alone. These supplementary cementing materials are usually added at dosage rates of 5% to 20% or higher by mass of cementing material. Some specifications only permit use of up to 10% silica fume, unless evidence is available indicating that concrete produced with a larger dosage rate will have satisfactory strength, durability, and volume stability. The water-to-cementing materials ratio should be adjusted so that equal workability becomes the basis of comparison between trial mixtures. For each set of materials, there will be an optimum cement-plus-supplementary cementing materials content at which strength does not continue to increase with greater amounts and the mixture becomes too sticky to handle properly. Blended cements containing fly ash, silica fume, slag, or calcined clay can be used to make high-strength concrete with or without the addition of supplementary cementing materials.

Aggregates

In high-strength concrete, careful attention must be given to aggregate size, shape, surface texture, mineralogy, and cleanliness. For each source of aggregate and concrete strength level there is an optimum-size aggregate that will yield the most compressive strength per unit of cement. To find the optimum size, trial batches should be made with 19 mm (3/4 in.) and smaller coarse aggregates and varying cement contents. Many studies have found that 9.5 mm to 12.5 mm (3/8 in. to 1/2 in.) nominal maximum-size aggregates give optimum strength.

In high-strength concretes, the strength of the aggregate itself and the bond or adhesion between the paste and aggregate become important factors. Tests have shown that crushed-stone aggregates produce higher compressive strength in concrete than gravel aggregate using the same size aggregate and the same cementing materials content; this is probably due to a superior aggregate-to-paste bond when using rough, angular, crushed material. For specified concrete strengths of 70 MPa (10,000 psi) or higher, the potential of the aggregates to meet design requirements must be established prior to use.

Coarse aggregates used in high-strength concrete should be clean, that is, free from detrimental coatings of dust and clay. Removing dust is important since it may affect the quantity of fines and consequently the water demand of a concrete mix. Clay may affect the aggregate-
paste of the aggregate and also by the mixture proportions (Baalbaki and others 1991). If an aggregate has the ability and suggested formulas for high-strength concrete. There are code formulas for normal-strength concrete. The modulus of elasticity is not necessarily proportional to the compressive strength of a concrete. There are code formulas for normal-strength concrete and suggested formulas for high-strength concrete. The modulus achievable is affected significantly by the properties of the aggregate and also by the mixture proportions (Baalbaki and others 1991). If an aggregate has the ability to produce a high modulus, then the optimum modulus in concrete can be obtained by using as much of this aggregate as practical, while still meeting workability and cohesiveness requirements. If the coarse aggregate being used is a crushed rock, and manufactured fine aggregate of good quality is available from the same source, then a combination of the two can be used to obtain the highest possible modulus.

Due to the high amount of cementitious material in high-strength concrete, the role of the fine aggregate (sand) in providing workability and good finishing characteristics is not as crucial as in conventional strength mixes. Sand with a fineness modulus (FM) of about 3.0—considered a coarse sand—has been found to be satisfactory for producing good workability and high compressive strength. For specified strengths of 70 MPa (10,000 psi) or greater, FM should be between 2.8 and 3.2 and not vary by more than 0.10 from the FM selected for the duration of the project. Finer sand, say with a FM of between 2.5 and 2.7, may produce lower-strength, sticky mixtures.

**Admixtures**

The use of chemical admixtures such as water reducers, retarders, high-range water reducers or superplasticizers is necessary. They make more efficient use of the large amount of cementitious material in high-strength concrete and help to obtain the lowest practical water to cementing materials ratio. Chemical admixture efficiency must be evaluated by comparing strengths of trial batches. Also, compatibility between cement and supplementary cementing materials, as well as water-reducing and other admixtures, must be investigated by trial batches. From these trial batches, it will be possible to determine the workability, setting time, and amount of water reduction for given admixture dosage rates and times of addition.

The use of air-entraining admixtures is not necessary or desirable in high-strength concrete that is protected from the weather, such as interior columns and shearwalls of high-rise buildings. However, for bridges, concrete piles, piers, or parking structures, where durability in a freeze-thaw environment is required, entrained air is mandatory. Because air entrainment decreases concrete strength of rich mixtures, testing to establish optimum air contents and spacing factors may be required. Certain high-strength concretes may not need as much air as normal-strength concrete to be frost resistant. Pinto and Hover (2001) found that non-air-entrained, high-strength concretes had good frost and deicer-scaling resistance at a water to portland cement ratio of 0.25. Burg and Ost (1996) found good frost resistance with non-air-entrained concrete containing silica fume at a water to cementing materials ratio of 0.22 (Mix No. 4 in Table 17-5); however, this was not the case with other mixtures, including a portland-only mixture with a water to cement ratio of 0.28.

**Proportioning**

The trial mixture approach is best for selecting proportions for high-strength concrete. To obtain high strength, it is necessary to use a low water to cementing materials ratio and a high portland cement content. The unit strength obtained for each unit of cement used in a cubic meter (yard) of concrete can be plotted as strength efficiency to assist with mix designs.

The water requirement of concrete increases as the fine aggregate content is increased for any given size of coarse aggregate. Because of the high cementing materials content of these concretes, the fine aggregate content can be kept low. However, even with well-graded aggregates, a low water-cementing materials ratio may result in concrete that is not sufficiently workable for the job. If a superplasticizer is not already being used, this may be the time to consider one. A slump of around 200 mm (8 in.) will provide adequate workability for most applications. ACI Committee 211 (1993), Farny and Panarese (1994), and Nawy (2001) provide additional guidance on proportioning.

**Mixing**

High-strength concrete has been successfully mixed in transit mixers and central mixers; however, many of these concretes tend to be sticky and cause build-up in these mixers. Where dry, uncompacted silica fume has been batching into a mix, “balling” of the mix has occurred and mixing has been less than complete. In such instances it has been found necessary to experiment with the sequence in which solids and liquids are added, and the
percentage of each material added at each step in the batching procedure. Batching and mixing sequences should be optimized during the trial mix phase. Where truck mixing is unavoidable, the best practice is to reduce loads to 90% of the rated capacity of the trucks.

Where there is no recent history of HSC or HPC mixtures that meet specified requirements, it is essential to first make laboratory trial mixes to establish optimum proportions. At this stage, the properties of the mix, such as workability, air content, density, strength, and modulus of elasticity can be determined. Once laboratory mixture proportions have been determined, field trials using full loads of concrete are essential; they should be delivered to the site or to a mock-up to establish and confirm the suitability of the batching, mixing, transporting and placing systems to be used.

Prequalification of concrete suppliers for high-strength concrete projects is recommended (Bickley 1993). In a prequalification procedure, one or more loads of the proposed mixture is cast into a trial mock-up. The fresh concrete is tested for slump, air content, temperature, and density. Casting the mock-up provides the opportunity to assess the suitability of the mix for placing and compaction. The mock-up can be instrumented to record temperatures and temperature gradients; it can also be cored and tested to provide correlation with standard cylinder test results. The cores can be tested to provide the designer with in-place strength and modulus values for reference during construction. The heat characteristics of the mixture can also be determined using a computer interactive program, and the data used to determine how curing technology should be applied to the project.

Placing, Consolidation, and Curing

Close liaison between the contractor and the concrete producer allows concrete to be discharged rapidly after arrival at the jobsite. Final adjustment of the concrete should be supervised by the concrete producer’s technicians at the site, by a concrete laboratory, or by a consultant familiar with the performance and use of high-strength concrete.

Delays in delivery and placing must be eliminated; sometimes it may be necessary to reduce batch sizes if placing procedures are slower than anticipated. Rigid surveillance must be exercised at the jobsite to prevent any addition of retempering water. Increases in workability should only be achieved by the addition of a superplasticizer. This should be done by the supplier’s technician. The contractor must be prepared to receive the concrete and understand the consequences of exceeding the specified slump and water-cementitious materials ratio.

Consolidation is very important in achieving the potential strengths of high-strength concrete. Concrete must be vibrated as quickly as possible after placement in the forms. High-frequency vibrators should be small enough to allow sufficient clearance between the vibrating head and reinforcing steel. Over-vibration of workable normal-strength concrete often results in segregation, loss of entrained air, or both. On the other hand, high-strength concrete without a superplasticizer, will be relatively stiff and contain little air. Consequently, inspectors should be more concerned with under-vibration rather than over-vibration. Most high-strength concrete, particularly very high-strength-concrete, is placed at slumps of 180 mm to 220 mm (7 in. to 9 in.). Even at these slumps, some vibration is required to ensure compaction. The amount of compaction should be determined by onsite trials.

High-strength concrete is often difficult to finish because of its sticky nature. High cementitious materials contents, large dosages of admixtures, low water contents, and air entrainment all contribute to the difficulty of finishing these concretes. Because the concrete sticks to the trowels and other finishing equipment, finishing activities should be minimized. The finishing sequence should be modified from that used for normal concrete.

Curing of high-strength concrete is even more important than curing normal-strength concrete. Providing adequate moisture and favorable temperature conditions is recommended for a prolonged period, particularly when 56- or 91-day concrete strengths are specified.

Additional curing considerations apply with HSC and HPC. Where very low water-cement ratios are used in flatwork (slabs and overlays), and particularly where silica fume is used in the mixture, there will be little if any bleeding before or after finishing. In these situations it is imperative that fog curing or evaporation retarders be applied to the concrete immediately after the surface has been struck off. This is necessary to avoid plastic shrinkage cracking of horizontal surfaces and to minimize curling. Fog curing, followed by 7 days of wet curing, has proven to be very effective.

It is inevitable that some vertical surfaces, such as columns, may be difficult to cure effectively. Where projects are fast-tracked, columns are often stripped at an early age to allow raising of self-climbing form systems. Concrete is thus exposed to early drying, sometimes within eleven hours after casting. Because of limited access, providing further curing is difficult and impractical.

Tests were conducted on column concrete to determine if such early exposure and lack of curing have any harmful effects. The tests showed that for a portland cement-slag-silica fume mixture with a specified strength of 70 MPa (10,000 psi), the matrix was sound and a very high degree of impermeability to water and chloride ions had been achieved (Bickley and others 1994). Nevertheless, the best curing possible is recommended for all HPC.

The temperature history of HPC is an integral part of its curing process. Advantage should also be taken of recent developments in curing technology. Temperature
increases and gradients that will occur in a concrete placement can be predicted by procedures that provide data for this purpose. With this technique, measures to heat, cool, or insulate a concrete placement can be determined and applied to significantly reduce both micro- and macro-cracking of the structure and assure durability. The increasing use of these techniques will be required in most structures using HPC to assure that the cover concrete provides long term protection to the steel, and results in the intended service life of the structure.

Quality Control

A comprehensive quality-control program is required at both the concrete plant and onsite to guarantee consistent production and placement of high-strength concrete. Inspection of concreting operations from stockpiling of aggregates through completion of curing is important. Closer production control than is normally obtained on most projects is necessary. Also, routine sampling and testing of all materials is particularly necessary to control uniformity of the concrete.

While tests on concrete should always be made in strict accordance with standard procedures, some additional requirements are recommended, especially where specified strengths are 70 MPa (10,000 psi) or higher. In testing high-strength concrete, some changes and more attention to detail are required. For example, cardboard cylinder molds, which can cause lower strength-test results, should be replaced with reusable steel or plastic molds. Capping of cylinders must be done with great care using appropriate capping compounds. Lapping (grinding) the cylinder ends is an alternative to capping. For specified strengths of 70 MPa (10,000 psi) or greater, end grinding to a flatness tolerance of 0.04 mm is recommended.

The physical characteristics of a testing machine can have a major impact on the result of a compression test. It is recommended that testing machines be extremely stiff, both longitudinally and laterally.

The quality control necessary for the production of high compressive strength concrete will, in most cases, lead to low variance in test results. Strict vigilance in all aspects of quality control on the part of the producer and quality testing on the part of the laboratory are necessary on high-strength concrete projects. For concretes with specified strengths of 70 MPa (10,000 psi), or greater, the coefficient of variation is the preferred measure of quality control.

HIGH-DURABILITY CONCRETE

Most of the attention in the 1970s and 1980s was directed toward high strength HPC; today the focus is more on concretes with high durability in severe environments resulting in structures with long life. For example, the Confederation Bridge across the Northumberland Strait between Prince Edward Island and New Brunswick has a 100-year design life (see Mix No. 2 in Table 17-3). This bridge contains HPC designed to efficiently protect the embedded reinforcement. The concrete had a diffusion coefficient of $4.8 \times 10^{-13}$ at six months (a value 10 to 30 times lower than that of conventional concrete). The electrical resistivity was measured at 470 to 530 ohm-m, compared to 50 for conventional concrete. The design required that the concrete be rated at less than 1000 coulombs. The high concrete resistivity in itself will result in a rate of corrosion that is potentially less than 10 percent of the corrosion rate for conventional concrete (Dunaszegi 1999). The following sections review durability issues that high-performance concrete can address.

Abrasion Resistance

Abrasion resistance is directly related to the strength of concrete. This makes high strength HPC ideal for abrasive environments. The abrasion resistance of HPC incorporating silica fume is especially high. This makes silica-fume concrete particularly useful for spillways and stilling basins, and concrete pavements or concrete pavement overlays subjected to heavy or abrasive traffic.

Holland and others (1986) describe how severe abrasion-erosion had occurred in the stilling basin of a dam; repairs using fiber-reinforced concrete had not proven to be durable. The new HPC mix used to repair the structure the second time contained 386 kg/m$^3$ (650 lb/yd$^3$) of cement, 70 kg/m$^3$ (118 lb/yd$^3$) of silica fume, admixtures, and had a water to cementing materials ratio of 0.28, and a 90-day compressive strength exceeding 103 MPa (15,000 psi).

Berra, Ferrara, and Tavano (1989) studied the addition of fibers to silica fume mortars to optimize abrasion resistance. The best results were obtained with a mix using slag cement, steel fibers, and silica fume. Mortar strengths ranged from 75 MPa to 100 MPa (11,000 psi to 14,500 psi). In addition to better erosion resistance, less drying shrinkage, high freeze-thaw resistance, and good bond to the substrate were achieved.

In Norway steel studs are allowed in tires; this causes severe abrasion wear on pavement surfaces, with resurfacing required within one to two years. Tests using an accelerated road-wear simulator showed that in the range of 100 MPa to 120 MPa (14,500 psi to 17,000 psi), concrete had the same abrasion resistance as granite (Helland 1990). Abrasion-resistant highway mixes usually contain between 320 and 450 kg/m$^3$ (539 and 758 lb/yd$^3$) of cement, plus silica fume or fly ash. They have water to cementing materials ratios of 0.22 to 0.36 and compressive strengths in the range of 85 to 130 MPa (12,000 to 19,000 psi). Applications have included new pavements and overlays to existing pavements.
**Blast Resistance**

High-performance concrete can be designed to have excellent blast resistance properties. These concretes often have a compressive strength exceeding 120 MPa (14,500 psi) and contain steel fibers. Blast-resistant concretes are often used in bank vaults and military applications.

**Permeability**

The durability and service life of concrete exposed to weather is related to the permeability of the cover concrete protecting the reinforcement. HPC typically has very low permeability to air, water, and chloride ions. Low permeability is often specified through the use of a coulomb value, such as a maximum of 1000 coulombs.

Test results obtained on specimens from a concrete column specified to be 70 MPa (10,000 psi) at 91 days and which had not been subjected to any wet curing were as follows (Bickley and others):

- **Water permeability of vacuum-saturated specimens:**
  - Age at test: 7 years
  - Applied water pressure: 0.69 MPa
  - Permeability: $7.6 \times 10^{-13} \text{ cm/s}$

- **Rapid chloride permeability (ASTM C1202):**
  - Age at test, years: 1, 2, 7
  - Coulombs: 303, 258, 417

The dense pore structure of high-performance concrete, which makes it so impermeable, gives it characteristics that make it eminently suitable for uses where a high quality concrete would not normally be considered. Latex-modified HPC is able to achieve these same low levels of permeability at normal strength levels without the use of supplementary cementing materials.

A large amount of concrete is used in farm structures. It typically is of low quality and often porous and with a rough surface, either when placed or after attack by farmyard wastes.

Gagne, Chagnon, and Parizeau (1994) provided a case history of the successful application of high performance concrete for agricultural purposes. In one case a farmer raising pigs on a large scale was losing about 1 kg per pig through diarrhea. This problem was resolved by reconstructing the pig pens with high performance concrete. Cited as beneficial properties in this application were:

- Surface smoothness that is compatible with the sensitive skin of a piglet
- Non-slip surface
- Good thermal conductivity resulting in uniform distribution of heat
- Impermeable surface to resist the growth of bacteria and viruses
- Easy to place

- Resistant to attack by acid (farmyard) wastes
- Price effective

The mix used had a water to portland cement ratio of 0.33 and had a 7-day compressive strength of 50 MPa (7,250 psi).

**Diffusion**

Aggressive ions, such as chloride, in contact with the surface of concrete will diffuse through the concrete until a state of equilibrium in ion concentration is achieved. If the concentration of ions at the surface is high, diffusion may result in corrosion-inducing concentrations at the level of the reinforcement.

The lower the water-cementing materials ratio the lower the diffusion coefficient will be for any given set of materials. Supplementary cementing materials, particularly silica fume, further reduce the diffusion coefficient. Typical values for diffusion for HPC are as follows:

<table>
<thead>
<tr>
<th>Type of Concrete</th>
<th>Diffusion Coefficient (m$^2$/s)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Portland cement-fly-ash mix:</td>
<td>$1000 \times 10^{-15}$</td>
</tr>
<tr>
<td>Portland cement-fly ash mix:</td>
<td>$1600 \times 10^{-15}$</td>
</tr>
</tbody>
</table>

**Carbonation**

HPC has a very good resistance to carbonation due to its low permeability. It was determined that after 17 years the concrete in the CN Tower in Toronto had carbonated to an average depth of 6 mm (0.24 in.) (Bickley, Sarkar, and Langlois 1992). The concrete mixture in the CN Tower had a water-cement ratio of 0.42. For a cover to the reinforcement of 35 mm (1.4 in.), this concrete would provide corrosion protection for 500 years. For the lower water-cementing materials ratios common to HPC, significantly longer times to corrosion would result, assuming a crack free structure. In practical terms, uncracked HPC cover concrete is immune to carbonation to a depth that would cause corrosion.

**Temperature Control**

The quality, strength, and durability of HPC is highly dependent on its temperature history from the time of delivery to the completion of curing. In principle, favorable construction and placing methods will enable: (1) a low temperature at the time of delivery; (2) the smallest possible maximum temperature after placing; (3) minimum temperature gradients after placing; and (4) a gradual reduction to ambient temperature after maximum temperature is reached. Excessively high temperatures and gradients can cause excessively fast hydration and micro- and macro-cracking of the concrete.

It has been a practice on major high-rise structures incorporating concretes with specified strengths of 70 MPa.
to 85 MPa (10,000 psi to 12,000 psi) to specify a maximum delivery temperature of 18°C (64°F) (Ryell and Bickley 1987). In summertime it is possible that this limit could only be met by using liquid nitrogen to cool the concrete. Experience with very-high-strength concrete suggests that a delivery temperature of no more than 25°C (77°F), preferably 20°C (68°F), should be allowed. In addition to liquid nitrogen, measures to cool HPC in the summer may involve using ice or chilled water as part of the mix water. The specifier should state the required delivery temperature.

In HPC applications such as high-rise buildings, column sizes are large enough to be classed as mass concrete. Normally, excessive heat generation in mass concrete is controlled by using a low cement content. When high-cement-content HPC mixes are used under these conditions, other methods of controlling maximum concrete temperature must be employed. Burg and Ost (1994) recorded temperature rise for 1220-mm (4-ft) concrete cubes using the mixtures in Table 17-5. A maximum temperature rise of 9.4°C to 11.7°C for every 100 kg of cement per cubic meter of concrete (10°F to 12.5°F for every 100 lb of cement per cubic yard of concrete) was measured. Burg and Fiorato (1999) monitored temperature rise in high-strength concrete caissons; they determined that in-place strength was not affected by temperature rise due to heat of hydration.

Freeze-Thaw Resistance

Because of its very low water-cementing materials ratio (less than 0.25), it is widely believed that HPC should be highly resistant to both scaling and physical breakup due to freezing and thawing. There is ample evidence that properly air-entrained high performance concretes are highly resistant to freezing and thawing and to scaling. Gagne, Pigeon, and Aïtcin (1990) tested 27 mixes using cement and silica fume with water-cementing materials ratios of 0.30, 0.26, and 0.23 and a wide range of quality in air-voids systems. All specimens performed exceptionally well in salt-scaling tests, confirming the durability of high-performance concrete, and suggesting that air-entrainment is not needed. Tachitana and others (1990) conducted ASTM C666 (Procedure A) tests on non-air-entrained high performance concrete with water-cementing materials ratios between 0.22 and 0.31. All were found to be extremely resistant to freeze-thaw damage and again it was suggested that air-entrainment is not needed.

Pinto and Hover (2001) found that non-air-entrained concrete with a water to portland cement ratio of 0.25 was deicer-scaling resistant with no supplementary cementing materials present. They found that higher strength portland cement concretes needed less air than normal concrete to be frost and scale resistant.

Burg and Ost (1994) found that of the six mixes tested in Table 17-5 using ASTM C666, only the silica fume concrete (Mix 4) with a water to cementing materials ratio of 0.22 was frost resistant.

Sidewalks constructed in Chicago in the 1920s used 25-mm (1-in.) thick toppings made of no-slump dry-pack mortar that had to be rammed into place. The concrete contained no air entrainment. Many of these sidewalks are still in use today; they are in good condition (minus some surface paste exposing fine aggregate) after 60 years of exposure to frost and deicers. No documentation exists on the water to cement ratio; however, it can be assumed that the water to cement ratio was comparable to that of modern HPCs.

While the above experiences prove the excellent durability of certain high-performance concretes to freeze-thaw damage and salt scaling, it is considered prudent to use air-entrainment. No well-documented field experiments have been made to prove that air-entrainment is not needed. Until such data are available, current practice for air-entrainment should be followed. It has been shown that the prime requirement of an air-void system for HPC is a preponderance of air bubbles of 200 µm size and smaller. If the correct air bubble size and spacing can be assured, then a moderate air content will ensure durability and minimize strength loss. The best measure of air-entrainment is the spacing factor.

Chemical Attack

For resistance to chemical attack on most structures, HPC offers a much improved performance. Resistance to various sulfates is achieved primarily by the use of a dense, strong concrete of very low permeability and low water-to-cementing materials ratio; these are all characteristics of HPC. Similarly, as discussed by Gagne and others (1994), resistance to acid from wastes is also much improved.

Alkali-Silica Reactivity

Reactivity between certain siliceous aggregates and alkali hydroxides can affect the long-term performance of concrete. Two characteristics of HPC that help combat alkali-silica reactivity are:

(1) HPC concretes at very low water to cement ratios can self desiccate (dry out) to a level that does not allow ASR to occur (relative humidity less than 80%). Burg and Ost (1994) observed relative humidity values ranging from 62% to 72% for their six mixes in Table 17-5. The low permeability of HPC also minimizes external moisture from entering the concrete.

(2) HPC concretes can use significant amounts of supplementary cementing materials that may have the ability to control alkali-silica reactivity. However, this must be demonstrated by test. HPC concretes can also use ASR inhibiting admixtures to control ASR.

HPC concretes are not immune to alkali-silica reactivity and appropriate precautions must be taken.
Resistivity

HPC, particularly that formulated with silica fume, has very high resistivity, up to 20 to 25 times that of normal concrete. This increases resistance to the flow of electrical current and reduces corrosion rates. Particularly if dry, HPC acts as an effective dielectric. Where cracking occurs in HPC, the corrosion is localized and minor; this is due to the high resistivity of the concrete which suppresses the development of a macro corrosion cell.

SELF-CONSOLIDATING CONCRETE

Self-consolidating concrete (SCC), also referred to as self-compacting concrete, is able to flow and consolidate under its own weight. At the same time it is cohesive enough to fill spaces of almost any size and shape without segregation or bleeding. This makes SCC particularly useful wherever placing is difficult, such as in heavily-reinforced concrete members or in complicated formwork.

This technology, developed in Japan in the 1980s, is based on increasing the amount of fine material, for example fly ash or limestone filler, without changing the water content compared to common concrete. This changes the rheological behavior of the concrete. SCC has to have a low yield value to ensure high flowability; a low water content ensures high viscosity, so the coarse aggregate can float in the mortar without segregating. To achieve a balance between deformability and stability, the total content of particles finer than the 150 µm (No. 100) sieve is typically high, usually about 520 to 560 kg/m³ (880 to 950 lb/yd³). Generally, the higher the required flowability of the SCC mix, the higher the amount of fine material needed to produce a stable mixture. However, in some cases, a viscosity-modifying admixture (VMA) can be used instead of, or in combination with, an increased fine content to stabilize the concrete mixture. High-range water reducers based on polycarboxylate ethers are typically used to plasticize the mixture. Fig. 17-6 shows an example of mix proportions used in self-consolidating concrete as compared to a regular concrete mix.

Since SCC is characterized by special fresh concrete properties, many new tests have been developed to measure flowability, viscosity, blocking tendency, self-leveling, and stability of the mixture (Skarendahl and Peterson 1999 and Ludwig and others 2001). The slump flow test (ASTM C1611) is performed to measure filling ability and stability. The test is performed similarly to the conventional slump test (ASTM C143) using the Abrams cone. However, instead of measuring the slumping distance vertically, the mean spread of the resulting concrete paty is measured horizontally. This number is recorded as the slump flow. The J-Ring test (ASTM C1621) measures SCC’s passing ability. The J-Ring consists of a ring of reinforcing bar such that it will fit around the base of a standard slump cone (Fig. 17-7). The slump flow with and without J-Ring is measured, and the difference calculated. The column segregation test (ASTM C1610) evaluates static stability of a concrete mixture by quantifying aggregate segregation. A column is filled with concrete and allowed to sit for awhile after placement. The column is then separated into three pieces. Each section is removed individually and the concrete from that section is washed over a 4.75 mm (No. 4) sieve and the retained aggregate weighed. A non-segregating mixture will have a consistent aggregate mass distribution in each section. A segregating mix will have higher concentrations of aggregate in the lower sections.

Strength and durability of well-designed SCC are almost similar to conventional concrete. Without proper curing, SCC tends to have higher plastic shrinkage cracking than conventional concrete (Grube and Rickert 2001). Research indicates greater tensile creep for SCC, resulting in a reduced tendency to crack (Bickley and Mitchell 2001). The use of fly ash as a filler seems to be

Fig. 17-6. Examples of materials used in regular concrete and self-consolidating concrete by absolute volume.
advantageous compared to limestone filler; it results in higher strength and higher chloride resistance (Bouzoubaa and Lachemi 2001 and Ludwig and others 2001).

The production of SCC is more expensive than regular concrete and it is difficult to keep SCC in the desired consistency over a long period of time. However, construction time is shorter and production of SCC is environmentally friendly (no noise, no vibration). Furthermore, SCC produces a good surface finish. These advantages make SCC particularly interesting for use in precasting plants. SCC has been successfully used in a number of rehabilitation projects in Canada (Bickley and Mitchell 2001). Refer to ACI Committee 237 and ACBM (2007) for more information on SCC.

REACTIVE-POWDER CONCRETE

Reactive-powder concrete (RPC) was first patented by a French construction company in 1994. It is characterized by high strength and very low porosity, which is obtained by optimized particle packing and low water content.

The properties of RPC are achieved by: (1) eliminating the coarse aggregates; just very fine powders are used such as sand, crushed quartz, and silica fume, all with particle sizes between 0.02 and 300 µm; (2) optimizing the grain size distribution to densify the mixture; (3) post-set heat-treatment to improve the microstructure; (4) addition of steel and synthetic fibers (about 2% by volume); and (5) use of superplasticizers to decrease the water to cement ratio—usually to less than 0.2—while improving the rheology of the paste. See Fig. 17-8 for a typical fresh RPC.

The compressive strength of reactive-powder concrete is typically around 200 MPa (29,000 psi), but can be produced with compressive strengths up to 810 MPa (118,000 psi) (Semioli 2001). However, the low comparative tensile strength requires prestressing reinforcement in severe structural service. Table 17-6 compares hardened concrete properties of RPC with those of an 80-MPa (11,600-psi) concrete.

![Fig. 17-8. Freshly-mixed reactive-powder concrete.](image)

RPC has found some applications in pedestrian bridges (Fig. 17-9) (Bickley and Mitchell 2001 and Semioli 2001). Also, the low porosity of RPC gives excellent durability and transport properties, which makes it a suitable material for the storage of nuclear waste (Matte and Moranville 1999). A low-heat type of reactive-powder concrete has been developed to meet needs for mass concrete pours for nuclear reactor foundation mats and underground containment of nuclear wastes (Gray and Shelton 1998).

REFERENCES


<table>
<thead>
<tr>
<th>Property</th>
<th>Unit</th>
<th>80 MPa</th>
<th>RPC</th>
</tr>
</thead>
<tbody>
<tr>
<td>Compressive strength</td>
<td>MPa (psi)</td>
<td>80 (11,600)</td>
<td>200 (29,000)</td>
</tr>
<tr>
<td>Flexural strength</td>
<td>MPa (psi)</td>
<td>7 (1000)</td>
<td>40 (5800)</td>
</tr>
<tr>
<td>Tensile strength</td>
<td>MPa (psi)</td>
<td>8 (1160)</td>
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<tr>
<td>Modulus of Elasticity</td>
<td>GPa (psi)</td>
<td>40 (5.8 \times 10^6)</td>
<td>60 (8.7 \times 10^6)</td>
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<td>Fracture Toughness</td>
<td>10^3 J/m²</td>
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<td>30</td>
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<td>Freeze-thaw, ASTM C666</td>
<td>RDF</td>
<td>90</td>
<td>100</td>
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<tr>
<td>Carbonation depth: 36 days in CO₂</td>
<td>mm</td>
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<td>0</td>
</tr>
<tr>
<td>Abrasion</td>
<td>10^{-12} m²/s</td>
<td>275</td>
<td>1.2</td>
</tr>
</tbody>
</table>

Table 17-6. Typical Mechanical Properties of Reactive Powder Concrete (RPC) Compared to an 80-MPa Concrete (Perry 1998)
ACI Committee 211, Guide for Selecting Proportions for High-Performance Concrete with Portland Cement and Fly Ash, ACI 211.4-93, reapproved 1998, American Concrete Institute, Farmington Hills, Michigan, 1993, 13 pages.

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Bickley, J. A.; Sarkar, S.; and Langlois, M., The CN Tower, Concrete International, American Concrete Institute, Farmington Hills, Michigan, August 1992, pages 51 to 55.

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Ferraris, Chiara F., and Lobo, Colin L., “Processing of HPC,” Concrete International, American Concrete Institute, Farmington Hills, Michigan, April 1998, pages 61 to 64.


Holland, T. C.; Krysa, Anton; Luther, Mark D.; and Liu, Tony C., “Use of Silica-Fume Concrete to Repair Abrasion-Erosion Damage in the Kinzua Dam Stillling Basin,” Proceedings of the Second International Conference on Fly Ash, Silica Fume, Slag and Natural Pozzolans in Concrete, SP-91, American Concrete Institute, Farmington Hills, Michigan, 1986, pages 841 to 864.


CHAPTER 18
Special Types of Concrete

Special types of concrete are those with out-of-the-ordinary properties or those produced by unusual techniques. Concrete is by definition a composite material consisting essentially of a binding medium and aggregate particles, and it can take many forms. Table 18-1 lists many special types of concrete made with portland cement and some made with binders other than portland cement. In many cases the terminology of the listing describes the use, property, or condition of the concrete. Brand names are not given. Some of the more common concretes are discussed in this chapter.

<table>
<thead>
<tr>
<th>Special types of concrete made with portland cement</th>
<th>Special types of concrete not using portland cement</th>
</tr>
</thead>
<tbody>
<tr>
<td>Architectural concrete</td>
<td>Recycled concrete</td>
</tr>
<tr>
<td>Autoclaved cellular concrete</td>
<td>Roller-compacted concrete</td>
</tr>
<tr>
<td>Centrifugally cast concrete</td>
<td>High-early-strength concrete</td>
</tr>
<tr>
<td>Colloidal concrete</td>
<td>High-strength concrete</td>
</tr>
<tr>
<td>Colored concrete</td>
<td>Insulating concrete</td>
</tr>
<tr>
<td>Controlled-density fill</td>
<td>Latex-modified concrete</td>
</tr>
<tr>
<td>Cyclopean (rubble) concrete</td>
<td>Low-density concrete</td>
</tr>
<tr>
<td>Dry-packed concrete</td>
<td>Mass concrete</td>
</tr>
<tr>
<td>Epoxy-modified concrete</td>
<td>Moderate-strength lightweight concrete</td>
</tr>
<tr>
<td>Exposed-aggregate concrete</td>
<td>Nailable concrete</td>
</tr>
<tr>
<td>Ferrocement</td>
<td>No-slump concrete</td>
</tr>
<tr>
<td>Fiber concrete</td>
<td>Polymer-modified concrete</td>
</tr>
<tr>
<td>Fill concrete</td>
<td>Pervious (porous) concrete</td>
</tr>
<tr>
<td>Flowable fill</td>
<td>Pozzolan concrete</td>
</tr>
<tr>
<td>Flowing concrete</td>
<td>Precast concrete</td>
</tr>
<tr>
<td>Fly-ash concrete</td>
<td>Prepacked concrete</td>
</tr>
<tr>
<td>Gap-graded concrete</td>
<td>Preplaced aggregate concrete</td>
</tr>
<tr>
<td>Geopolymer concrete</td>
<td>Reactive-powder concrete</td>
</tr>
<tr>
<td></td>
<td></td>
</tr>
<tr>
<td>Acrylic concrete</td>
<td>Furan concrete</td>
</tr>
<tr>
<td>Aluminum phosphate concrete</td>
<td>Gypsum concrete</td>
</tr>
<tr>
<td>Asphalt concrete</td>
<td>Latex concrete</td>
</tr>
<tr>
<td>Calcium aluminate concrete</td>
<td>Magnesium phosphate concrete</td>
</tr>
<tr>
<td>Epoxy concrete</td>
<td>Methyl methacrylate (MMA) concrete</td>
</tr>
</tbody>
</table>

Most of the definitions of these types of concrete appear in Cement and Concrete Terminology, ACI 116.

STRUCTURAL LIGHTWEIGHT CONCRETE

Structural lightweight concrete is similar to normal-weight concrete except that it has a lower density. It is made with lightweight aggregates (all-lightweight concrete) or with a combination of lightweight and normal-weight aggregates. The term “sand lightweight” refers to lightweight concrete made with coarse lightweight aggregate and natural sand.

Structural lightweight concrete has an air-dry density in the range of 1350 to 1850 kg/m³ (85 to 115 pcf) and a
28-day compressive strength in excess of 17 MPa (2500 psi). Some job specifications allow air-dry densities up to 1920 kg/m³ (120 pcf). For comparison, normal-weight concrete containing regular sand, gravel, or crushed stone has a dry density in the range of 2080 to 2480 kg/m³ (130 to 155 pcf). ASTM C567 provides a test for density of structural lightweight concrete. Structural lightweight concrete is used primarily to reduce the dead-load weight in concrete members, such as floors in high-rise buildings.

**Structural Lightweight Aggregates**

Structural lightweight aggregates are usually classified according to their production process because various processes produce aggregates with somewhat different properties. Processed structural lightweight aggregates should meet the requirements of ASTM C330, which includes:

- Rotary kiln expanded clays (Fig. 18-1), shales, and slates
- Sintering grate expanded shales and slates
- Pelletized or extruded fly ash
- Expanded slags

Structural lightweight aggregates can also be produced by processing other types of material, such as naturally occurring pumice and scoria.

Structural lightweight aggregates have densities significantly lower than normal-weight aggregates, ranging from 560 to 1120 kg/m³ (35 to 70 pcf), compared to 1200 to 1760 kg/m³ (75 to 110 pcf) for normal-weight aggregates. These aggregates may absorb 5% to 20% water by weight of dry material. To control the uniformity of structural lightweight concrete mixtures, the aggregates are pre-wetted (but not saturated) prior to batching.

**Compressive Strength**

The compressive strength of structural lightweight concrete is usually related to the cement content at a given slump and air content, rather than to a water-to-cement ratio. This is due to the difficulty in determining how much of the total mix water is absorbed into the aggregate and thus not available for reaction with the cement. ACI 211.2 provides guidance on the relationship between compressive strength and cement content. Typical compressive strengths range from 20 to 35 MPa (3000 to 5000 psi). High-strength concrete can also be made with structural lightweight aggregates.

In well-proportioned mixtures, the cement content and strength relationship is fairly constant for a particular source of lightweight aggregate. However, the relationship will vary from one aggregate source or type to another. When information on this relationship is not available from the aggregate manufacturer, trial mixtures with varying cement contents are required to develop a range of compressive strengths, including the strength specified. Fig. 18-2 shows the relationship between cement content and compressive strength. An example of a 28-MPa (4000-psi) structural lightweight concrete mixture with an air-dry density of about 1800 kg/m³ (112 pcf), a combination of natural sand and gravel, and a lightweight rotary kiln expanded clay coarse aggregate follows:
• 356 kg (600 lb) Type I portland cement
• 534 kg (900 lb) sand, oven-dry
• 320 kg (540 lb) gravel (12.5 to 2.36 mm [1/2 in. to #8]), oven-dry
• 356 kg (600 lb) lightweight aggregate (9.5 mm to 600 µm [3/8 in. to #30]), oven-dry
• 172 kg (290 lb) mix water added
• 0.7 L (20 oz) water-reducing admixture
• 0.09L (2.5 oz) air-entraining admixture
• 1m³ (1yd³) yield
• Slump—75 mm (3 in.)
• Air content—6%

Material proportions vary significantly for different materials and strength requirements.

**Entrained Air**

As with normal-weight concrete, entrained air in structural lightweight concrete ensures resistance to freezing and thawing and to deicer applications. It also improves workability, reduces bleeding and segregation, and may compensate for minor grading deficiencies in the aggregate.

The amount of entrained air should be sufficient to provide good workability to the plastic concrete and adequate freeze-thaw resistance to the hardened concrete. Air contents are generally between 5% and 8%, depending on the maximum size of coarse aggregate (paste content) used and the exposure conditions. Testing for air content should be performed by the volumetric method (ASTM C173 or AASHTO T 196). The freeze-thaw durability is also significantly improved if structural lightweight concrete is allowed to dry before exposure to a freeze-thaw environment.

**Specifications**

Many suppliers of lightweight aggregates for use in structural lightweight concrete have information on suggested specifications and mixture proportions pertaining to their product. The usual specifications for structural concrete state a minimum compressive strength, a maximum density, a maximum slump, and an acceptable range in air content.

The contractor should also be concerned with the bleeding, workability, and finishing properties of structural lightweight concrete.

**Mixing**

In general, mixing procedures for structural lightweight concrete are similar to those for normal-density concrete; however, some of the more absorptive aggregates may require prewetting before use. Water added at the batching plant should be sufficient to produce the specified slump at the jobsite. Measured slump at the batch plant will generally be appreciably higher than the slump at the site. Pumping can especially aggravate slump loss.

**Workability and Finishability**

Structural lightweight concrete mixtures can be proportioned to have the same workability, finishability, and general appearance as a properly proportioned normal-density concrete mixture. Sufficient cement paste must be present to coat each particle, and coarse-aggregate particles should not separate from the mortar. Enough fine aggregate is needed to keep the freshly mixed concrete cohesive. If aggregate is deficient in minus 600 µm (No. 30) sieve material, finishability may be improved by using a portion of natural sand, by increasing cement content, or by using satisfactory mineral fines. Since entrained air improves workability, it should be used regardless of exposure.

**Slump**

Due to lower aggregate density, structural lightweight concrete does not slump as much as normal-weight concrete with the same workability. A lightweight air-entrained mixture with a slump of 50 to 75 mm (2 to 3 in.) can be placed under conditions that would require a slump of 75 to 125 mm (3 to 5 in.) for normal-weight concrete. It is seldom necessary to exceed slumps of 125 mm (5 in.) for normal placement of structural lightweight concrete. With higher slumps, the large aggregate particles tend to float to the surface, making finishing difficult.

**Vibration**

As with normal-weight concrete, vibration can be used effectively to consolidate lightweight concrete; the same frequencies commonly used for normal-density concrete are recommended. The length of time for proper consolidation varies, depending on mix characteristics. Excessive vibration causes segregation by forcing large aggregate particles to the surface.

**Placing, Finishing, and Curing**

Structural lightweight concrete is generally easier to handle and place than normal-weight concrete. A slump of 50 to 100 mm (2 to 4 in.) produces the best results for finishing. Greater slumps may cause segregation, delay finishing operations, and result in rough, uneven surfaces.

If pumped concrete is being considered, the specifier, suppliers, and contractor should all be consulted about performing a field trial using the pump and mixture planned for the project. Adjustments to the mixture may be necessary; pumping pressure causes the aggregate to absorb more water, thus reducing the slump and increasing the density of the concrete.

Finishing operations should be started earlier than for comparable normal-weight concrete, but finishing too early may be harmful. A minimum amount of floating
and troweling should be done; magnesium finishing tools are preferred.

The same curing practices should be used for lightweight concrete as for normal-weight concrete. The two methods commonly used in the field are water curing (ponding, sprinkling, or using wet coverings) and preventing loss of moisture from the exposed surfaces (covering with waterproof paper, plastic sheets, or sealing with liquid membrane-forming compounds). Generally, 7 days of curing are adequate for ambient air temperatures above 10°C (50°F).

**INSULATING AND MODERATE-STRENGTH LIGHTWEIGHT CONCRETES**

Insulating concrete is a lightweight concrete with an ovendry density of 800 kg/m³ (50 pcf) or less. It is made with cementing materials, water, air, and with or without aggregate and chemical admixtures. The ovendry density ranges from 240 to 800 kg/m³ (15 to 50 pcf) and the 28-day compressive strength is generally between 0.7 and 7 MPa (100 and 1000 psi). Cast-in-place insulating concrete is used primarily for thermal and sound insulation, roof decks, fill for slab-on-grade subbases, leveling courses for floors or roofs, firewalls, and underground thermal conduct linings.

Moderate-strength lightweight concrete has a density of 800 to 1900 kg/m³ (50 to 120 pcf) ovendry and has a compressive strength of approximately 7 to 17 MPa (1000 to 2500 psi). It is made with cementing materials, water, air, and with or without aggregate and chemical admixtures. At lower densities, it is used as fill for thermal and sound insulation of floors, walls, and roofs and is referred to as fill concrete. At higher densities it is used in cast-in-place walls, floors and roofs, and precast wall and floor panels. See ACI documents for more information.

For discussion purposes, insulating and moderate-strength lightweight concretes can be grouped as follows:

**Group I** is made with expanded aggregates such as perlite, vermiculite, or expanded polystyrene beads. Ovendry concrete densities using these aggregates generally range between 240 and 800 kg/m³ (15 and 50 pcf). This group is used primarily in insulating concrete. Some moderate-strength concretes can also be made from aggregates in this group.

**Group II** is made with aggregates manufactured by expanding, calcining, or sintering materials such as blast-furnace slag, clay, diatomite, fly ash, shale, or slate, or by processing natural materials such as pumice, scoria, or tuff. Ovendry concrete densities using these aggregates can range between 720 and 1440 kg/m³ (45 and 90 pcf). Aggregates in this group are used in moderate-strength lightweight concrete and some of these materials (expanded slag, clay, fly ash, shale, and slate) are also used in both moderate-strength and structural lightweight concrete (up to about 1900 kg/m³ or 120 pcf air-dry).

**Group III** concretes are made by incorporating into a cement paste or cement-sand mortar a uniform cellular structure of air voids that is obtained with preformed foam (ASTM C869), formed-in-place foam, or special foaming agents. Ovendry densities ranging between 240 to 1900 kg/m³ (15 to 120 pcf) are obtained by substitution of air voids for some or all of the aggregate particles; air voids can consist of up to 80% of the volume. Cellular concrete can be made to meet the requirements of both insulating and moderate strength lightweight concrete.

Aggregates used in Groups I and II should meet the requirements of ASTM C332, *Standard Specification for Lightweight Aggregates for Insulating Concrete*. These aggregates have dry densities in the range of from 96 to 1120 kg/m³ (6 to 70 pcf) down to 16 kg/m³ (1 pcf) for expanded polystyrene beads.

**Mixture Proportions**

Examples of mixture proportions for Group I and III concretes appear in Table 18-2. In Group I, air contents may be as high as 25% to 35%. The air-entraining agent can be prepackaged with the aggregate or added at the mixer. Because of the absorptive nature of the aggregate, the volumetric method (ASTM C173 or AASHTO T 196) should be used to measure air content.

Water requirements for insulating and fill concretes vary considerably, depending on aggregate characteristics, entrained air, and mixture proportions. An effort should be made to avoid excessive amounts of water in insulating concrete used in roof fills. Excessive water causes high drying shrinkage and cracks that may damage the waterproofing membrane. Accelerators containing calcium chloride should not be used where galvanized steel will remain in permanent contact with the concrete because of possible corrosion problems.

Mixture proportions for Group II concretes usually are based on volumes of dry, loose materials, even when aggregates are moist as batched. Satisfactory proportions can vary considerably for different aggregates or combinations of aggregates. Mixture proportions ranging from 0.24 to 0.90 cubic meters of aggregate per 100 kg (4 to 14 cu ft per 100 lb) of cement can be used in lightweight concretes that are made with pumice, expanded shale, and expanded slag. Some mixtures, such as those for no-fines concretes, are made without fine aggregate but with total void contents of 20% to 35%. Cement contents for Group II concretes range between 120 to 360 kg per cubic meter (200 and 600 lb per cubic yard) depending on air content, aggregate gradation, and mixture proportions.

No-fines concretes containing pumice, expanded slag, or expanded shale can be made with 150 to 170 kg of water per cubic meter (250 to 290 lb of water per cubic yard), total
Workability

Because of their high air content, lightweight concretes weighing less than 800 kg/m³ (50 pcf) generally have excellent workability. Slumps of up to 250 mm (10 in.) usually are satisfactory for Group I and Group III concretes; appearance of the mix, however, may be a more reliable indication of consistency. Cellular concretes are handled as liquids; they are poured or pumped into place without further consolidation.

Mixing and Placing

All concrete should be mechanically mixed to produce a uniform distribution of materials of proper consistency and required density. In batch-mixing operations, various sequences can be used for introducing the ingredients; the preferred sequence is to first introduce the required amount of water into the mixer, then add the cement, air-entraining or foaming agent, aggregate, preformed foam, and any other ingredients.

Excessive mixing and handling should be avoided because they tend to break up aggregate particles, thereby changing density and consistency. Segregation is not usually a problem (though it could be for Group II) because of the relatively large amounts of entrained air in these mixtures.

Pumping is the most common method of placement, but other methods can be used. Finishing operations should be kept to a minimum; smoothing with a darby or bullfloat is usually sufficient. Placement of insulating concretes should be done by workers experienced with these special concretes.

Periodic wet-density tests (ASTM C138 or AASHTO T 121) at the jobsite can be performed to check the uniformity of the concrete. Variations in density generally should not exceed plus or minus 32 kg/m³ (2 pcf). A close approximation of the oven-dry density can be determined from the freshly mixed density.

<table>
<thead>
<tr>
<th>Type of concrete</th>
<th>Ratio: portland cement to aggregate by volume</th>
<th>Ovendry density, kg/m³ (pcf)</th>
<th>Type I portland cement, kg/m³ (lb/yard³)</th>
<th>Water-cement ratio, by mass</th>
<th>28-day compressive strength, MPa (psi)</th>
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</thead>
<tbody>
<tr>
<td>Perlite*</td>
<td>1:4</td>
<td>480 to 608 (30 to 38)</td>
<td>362 (610)</td>
<td>0.94</td>
<td>2.75 (400)</td>
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<tr>
<td></td>
<td>1:5</td>
<td>416 to 576 (26 to 36)</td>
<td>306 (516)</td>
<td>1.12</td>
<td>2.24 (325)</td>
</tr>
<tr>
<td></td>
<td>1:6</td>
<td>352 to 545 (22 to 34)</td>
<td>245 (414)</td>
<td>1.24</td>
<td>1.52 (220)</td>
</tr>
<tr>
<td></td>
<td>1:8</td>
<td>320 to 512 (20 to 32)</td>
<td>234 (395)</td>
<td>1.72</td>
<td>1.38 (200)</td>
</tr>
<tr>
<td>Vermiculite*</td>
<td>1:4</td>
<td>496 to 593 (31 to 37)</td>
<td>380 (640)</td>
<td>0.98</td>
<td>2.07 (300)</td>
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<tr>
<td></td>
<td>1:5</td>
<td>448 to 496 (28 to 31)</td>
<td>295 (498)</td>
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<td>1.17 (170)</td>
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<td></td>
<td>1:6</td>
<td>368 to 464 (23 to 29)</td>
<td>245 (414)</td>
<td>1.60</td>
<td>0.90 (130)</td>
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<tr>
<td></td>
<td>1:8</td>
<td>320 to 336 (20 to 21)</td>
<td>178 (300)</td>
<td>2.08</td>
<td>0.55 (80)</td>
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<tr>
<td>Polystyrene:**</td>
<td>sand:</td>
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<td></td>
<td></td>
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</tr>
<tr>
<td></td>
<td>0 kg/m³ (0 lb/yd³)</td>
<td>1:3.4</td>
<td>545 (34)††</td>
<td>445 (750)</td>
<td>0.40††</td>
</tr>
<tr>
<td></td>
<td>73 kg/m³ (124 lb/yd³)</td>
<td>1:3.1</td>
<td>625 (39)††</td>
<td>445 (750)</td>
<td>0.40††</td>
</tr>
<tr>
<td></td>
<td>154 kg/m³ (261 lb/yd³)</td>
<td>1:2.9</td>
<td>725 (44)††</td>
<td>445 (750)</td>
<td>0.40††</td>
</tr>
<tr>
<td></td>
<td>200 kg/m³ (338 lb/yd³)</td>
<td>1:2.5</td>
<td>769 (48)††</td>
<td>474 (800)</td>
<td>0.40††</td>
</tr>
<tr>
<td>Cellular*</td>
<td>(neat cement)</td>
<td>—</td>
<td>625 (39)††</td>
<td>524 (884)</td>
<td>0.57††</td>
</tr>
<tr>
<td></td>
<td>—</td>
<td>545 (34)††</td>
<td>468 (790)</td>
<td>0.56††</td>
<td>1.45 (210)</td>
</tr>
<tr>
<td></td>
<td>—</td>
<td>448 (28)††</td>
<td>396 (668)</td>
<td>0.57††</td>
<td>0.90 (130)</td>
</tr>
<tr>
<td></td>
<td>—</td>
<td>368 (23)††</td>
<td>317 (535)</td>
<td>0.65††</td>
<td>0.34 (50)</td>
</tr>
<tr>
<td>Cellular†</td>
<td>(sanded)††</td>
<td>1:1</td>
<td>929 (58)††</td>
<td>429 (724)</td>
<td>0.40††</td>
</tr>
<tr>
<td></td>
<td>1:2</td>
<td>1250 (78)††</td>
<td>373 (630)</td>
<td>0.41††</td>
<td>5.66 (820)</td>
</tr>
<tr>
<td></td>
<td>1:3</td>
<td>1602 (100)††</td>
<td>360 (602)</td>
<td>0.51††</td>
<td>15.10 (2190)</td>
</tr>
</tbody>
</table>

* Reichard (1971).
** Source: Hanna (1978). The mix also included air entrainment and a water-reducing agent.
† Source: Gustafsson (1970).
†† Dry-rodded sand with a bulk density of 1600 kg/m³ (100 pcf).
‡ Air-dry density at 28 days, 50% relative humidity.
**Thermal Resistance**

ASTM C177, Test Method for Steady-State Heat Flux Measurements and Thermal Transmission Properties by Means of the Guarded-Hot-Plate Apparatus, is used to determine values of thermal conductivity. Fig. 18-3 shows an approximate relationship between thermal resistance and density. The thermal conductivity of concrete increases with an increase in moisture content and density. See Brewer (1967) for additional density and conductivity relationships.

![Graph of Thermal Resistance vs Density](image)

**Fig. 18-3. Thermal resistance of concrete versus density (PCA 1980).**

**Strength**

Strength requirements depend on the intended use of the concrete. For example, a compressive strength of 0.7 MPa (100 psi), or even less, may be satisfactory for insulation of underground steam lines. Roof-fill insulation requires sufficient early strength to withstand foot traffic. Compressive strengths of 0.7 to 1.5 MPa (100 to 200 psi) are usually adequate for roof fills, but strengths up to 3.5 MPa (500 psi) are sometimes specified. In general, the strength of insulating concrete is of minor importance. Compressive strength of lightweight insulating concrete should be determined by the methods specified in ASTM C495 or C513.

Table 18-2 and Fig. 18-4 give examples of the relationship between density and strength for lightweight insulating concretes. Fig. 18-5 shows examples for cellular concrete containing sand. Mixtures with strengths outside the ranges shown can be made by varying the mixture proportions. Strengths comparable to those at 28 days

![Graph of Compressive Strength vs Density](image)

**Fig. 18-5. Plastic density versus compressive strength for sanded cellular concretes. Compressive strength was determined with 150 x 300-mm (6 x 12-in.) cylinders that were cured for 21 days in a 100% relative humidity moist room followed by 7 days in air at 50% RH (McCormick 1967 and ACI 523.3R).**
would be obtained at 7 days with high-early-strength cement. The relationships shown do not apply to autoclaved products.

**Resistance to Freezing and Thawing**

Insulating and moderate-strength lightweight concretes normally are not required to withstand freeze-thaw exposure in a saturated condition. In service they are normally protected from the weather, thus little research has been done on their resistance to freezing and thawing.

**Drying Shrinkage**

The drying shrinkage of insulating or moderate-strength lightweight concrete is not usually critical when it is used for insulation or fill; however, excessive shrinkage can cause curling. In structural use, shrinkage should be considered. Moist-cured cellular concretes made without aggregates have high drying shrinkage. Moist-cured cellular concretes made with sand may shrink from 0.1% to 0.6%, depending on the amount of sand used. Autoclaved cellular concretes shrink very little on drying. Insulating concretes made with perlite or pumice aggregates may shrink 0.1% to 0.3% in six months of drying in air at 50% relative humidity; vermiculite concretes may shrink 0.2% to 0.45% during the same period. Drying shrinkage of insulating concretes made with expanded slag or expanded shale ranges from about 0.06% to 0.1% in six months.

**Expansion Joints**

Where insulating concrete is used on roof decks, a 25-mm (1-in.) expansion joint at the parapets and all roof projections is often specified. Its purpose is to accommodate expansion caused by the heat of the sun so that the insulating concrete can expand independently of the roof deck. Transverse expansion joints should be placed at a maximum of 30 m (100 ft) in any direction for a thermal expansion of 1 mm per meter (1 in. per 100 lin ft). A fiberglass material that will compress to one-half its thickness under a stress of 0.17 MPa (25 psi) is generally used to form these joints.

**AUTOCLAVED CELLULAR CONCRETE**

Autoclaved cellular concrete (also called autoclaved aerated concrete) is a special type of lightweight building material. It is manufactured from a mortar consisting of pulverized siliceous material (sand, slag, or fly ash), cement and/or lime, and water; to this a gas forming admixture, for example aluminum powder, is added. The chemical reaction of aluminum with the alkaline water forms hydrogen, which expands the mortar as macro pores with a diameter of 0.5 mm to 1.5 mm (0.02 in. to 0.06 in.) form. The material is then pressure steam cured (autoclaved) over a period of 6 to 12 hours using a temperature of 190°C (374°F) and a pressure of 1.2 MPa (174 psi). This forms a hardened mortar matrix, which essentially consists of calcium silicate hydrates.

This porous mineral building material has densities between 300 and 1000 kg/m³ (19 and 63 lb/ft³) and compressive strengths between 2.5 and 10 MPa (300 and 1500 lb/in²). Due to the high macropore content—up to 80 percent by volume—autoclaved cellular concrete has a thermal conductivity of only 0.15 to 0.20 W/(m·K) (1 to 1.4 Btu·in. /[h·ft²·ºF]).

Autoclaved cellular concrete is produced in block or panel form for construction of residential or commercial buildings (Fig. 18-6).

Additional information can be found in ACI 523.2R, Guide for Precast Cellular Concrete.

**HIGH-DENSITY CONCRETE**

High-density (heavyweight) concrete and has a density of up to about 6400 kg/m³ (400 pcf). Heavyweight concrete is used principally for radiation shielding but is also used for counterweights and other applications where high-density is important. As a shielding material, heavyweight concrete protects against the harmful effects of X-rays, gamma rays, and neutron radiation. Selection of concrete for radiation shielding is based on space requirements and on the type and intensity of radiation. Where space requirements are not important, normal-weight concrete will generally produce the most economical shield; where space is limited, heavyweight concrete will allow for reductions in shield thickness without sacrificing shielding effectiveness.
Type and intensity of radiation usually determine the requirements for density and water content of shielding concrete. Effectiveness of a concrete shield against gamma rays is approximately proportional to the density of the concrete; the heavier the concrete, the more effective the shield. On the other hand, an effective shield against neutron radiation requires both heavy and light elements. The hydrogen in water provides an effective light element in concrete shields. Some aggregates contain crystallized water, called fixed water, as part of their structure. For this reason, heavyweight aggregates with high fixed-water contents often are used if both gamma rays and neutron radiation are to be attenuated. Boron glass (boron frit) is also added to attenuate neutrons.

**High-Density Aggregates**

High-density aggregates such as barite, ferrophosphorus, goethite, hematite, ilmenite, limonite, magnetite, and degreased steel punchings and shot are used to produce high-density concrete. Where high fixed-water content is desirable, serpentine (which is slightly heavier than normal-weight aggregate) or bauxite can be used (see ASTM C637 and C638).

Table 18-3 gives typical bulk density, relative density (specific gravity), and percentage of fixed water for some of these materials. The values are a compilation of data from a wide variety of tests or projects reported in the literature. Steel punchings and shot are used where concrete with a density of more than 4800 kg/m³ (300 pcf) is required.

In general, selection of an aggregate is determined by physical properties, availability, and cost. Heavyweight aggregates should be reasonably free of fine material, oil, and foreign substances that may affect either the bond of paste to aggregate particle or the hydration of cement. For good workability, maximum density, and economy, aggregates should be roughly cubical in shape and free of excessive flat or elongated particles.

**Additions**

Boron additions such as colemanite, boron frits, and boro-calcite are sometimes used to improve the neutron shielding properties of concrete. However, they may adversely affect setting and early strength of concrete; therefore, trial mixes should be made with the addition under field conditions to determine suitability. Admixtures such as pressure-hydrated lime can be used with coarse-sand sizes to minimize any retarding effect.

**Properties of High-Density Concrete**

The properties of high-density concrete in both the freshly mixed and hardened states can be tailored to meet job conditions and shielding requirements by proper selection of materials and mixture proportions.

Except for density, the physical properties of heavy-weight concrete are similar to those of normal-weight concrete. Strength is a function of water-cement ratio; thus, for any particular set of materials, strengths comparable to those of normal-weight concretes can be achieved. Typical densities of concretes made with some commonly used high-density aggregates are shown in Table 18-3. Because each radiation shield has special requirements, trial mixes should be made with job materials and under job conditions to determine suitable mixture proportions.

**Proportioning, Mixing, and Placing**

The procedures for selecting mix proportions for heavyweight concrete are the same as those for normal-weight concrete. However, additional mixture information and sample calculations are given in ACI 211.1. Following are the most common methods of mixing and placing high-density concrete:

*Conventional methods of mixing and placing often are used, but care must be taken to avoid overloading the mixer, especially with very heavy aggregates such as steel.

### Table 18-3. Physical Properties of Typical High-Density Aggregates and Concrete

<table>
<thead>
<tr>
<th>Type of aggregate</th>
<th>Fixed-water,* percent by weight</th>
<th>Aggregate relative density</th>
<th>Aggregate bulk density, kg/m³ (pcf)</th>
<th>Concrete density, kg/m³ (pcf)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Goethite</td>
<td>10–11</td>
<td>3.4–3.7</td>
<td>2080–2240 (130–140)</td>
<td>2880–3200 (180–200)</td>
</tr>
<tr>
<td>Limonite**</td>
<td>8–9</td>
<td>3.4–4.0</td>
<td>2080–2400 (130–150)</td>
<td>2880–3360 (180–210)</td>
</tr>
<tr>
<td>Barite</td>
<td>0</td>
<td>4.0–4.6</td>
<td>2320–2560 (145–160)</td>
<td>3360–3680 (210–230)</td>
</tr>
<tr>
<td>Ilmenite†</td>
<td>4.3–4.8</td>
<td></td>
<td>2560–2700 (160–170)</td>
<td>3520–3850 (220–240)</td>
</tr>
<tr>
<td>Hematite†</td>
<td>4.9–5.3</td>
<td></td>
<td>2880–3200 (180–200)</td>
<td>3850–4170 (240–260)</td>
</tr>
<tr>
<td>Magnetite†</td>
<td>4.2–5.2</td>
<td></td>
<td>2400–3040 (150–190)</td>
<td>3360–4170 (210–260)</td>
</tr>
<tr>
<td>Ferrophosphorus</td>
<td>0</td>
<td>5.8–6.8</td>
<td>3200–4160 (200–260)</td>
<td>4080–5290 (255–330)</td>
</tr>
<tr>
<td>Steel punchings or shot</td>
<td>0</td>
<td>6.2–7.8</td>
<td>3860–4650 (230–290)</td>
<td>4650–6090 (290–380)</td>
</tr>
</tbody>
</table>

* Water retained or chemically bound in aggregates.
** Test data not available.
† Aggregates may be combined with limonite to produce fixed-water contents varying from about ½% to 5%.
punchings. Batch sizes should be reduced to about 50% of the rated mixer capacity. Because some heavy aggregates are quite friable, excessive mixing should be avoided to prevent aggregate breakup with resultant detrimental effects on workability and bleeding.

Preplaced aggregate methods can be used for placing normal and high-density concrete in confined areas and around embedded items; this will minimize segregation of coarse aggregate, especially steel punchings or shot. The method also reduces drying shrinkage and produces concrete of uniform density and composition. With this method, the coarse aggregates are preplaced in the forms and grout made of cement, sand, and water is then pumped through pipes to fill the voids in the aggregate.

Pumping of heavyweight concrete through pipelines may be advantageous in locations where space is limited. Heavyweight concretes cannot be pumped as far as normal-weight concretes because of their higher densities.

Puddling is a method whereby a 50-mm (2-in.) layer or more of mortar is placed in the forms and then covered with a layer of coarse aggregate that is rodded or internally vibrated into the mortar. Care must be taken to ensure uniform distribution of aggregate throughout the concrete.

MASS CONCRETE

Mass concrete is defined by ACI Committee 116 as “Any large volume of cast-in-place concrete with dimensions large enough to require that measures be taken to cope with the generation of heat and attendant volume change to minimize cracking.” Mass concrete includes not only low-cement-content concrete used in dams and other massive structures but also moderate- to high-cement-content concrete in structural members of bridges and buildings (Fig. 18-7). Mass concrete placements require special considerations to reduce heat of hydration and the resulting temperature rise to avoid damaging the concrete through excessive temperatures and temperature differences that can result in thermal cracking (Gajda and VanGeem, 2002).

In mass concrete, temperature rise (Fig. 18-8) results from the heat of hydration of cementitious materials. As the interior concrete increases in temperature and expands, the surface concrete may be cooling and contracting. This causes tensile stresses that may result in thermal cracks at the surface if the temperature differen-
tial between the surface and center is too great. The width and depth of cracks depends upon the temperature differential, physical properties of the concrete, and the reinforcing steel.

A definite member size beyond which a concrete structure should be classified as mass concrete is not readily available. Many large structural elements may be massive enough that heat generation should be considered; this is particularly critical when the minimum cross-sectional dimensions of a solid concrete member approach or exceed 1 meter (3 feet) or when cement contents exceed 355 kg/m³ (600 lb per cubic yard). Temperature rise in mass concrete is related to the initial concrete temperature (Fig. 18-9), ambient temperature, size of the concrete element (volume to surface ratio and minimum dimension), and type and quantity of cementitious materials. Smaller concrete members less than 0.3 meters (1 ft) thick with moderate amounts of cementitious materials are typically of little concern as the generated heat is rapidly dissipated.

To avoid cracking, the internal concrete temperature for dams and other nonreinforced mass concrete structures of relatively low compressive strength should not be allowed to rise more than 11°C to 14°C (20°F to 25°F) above the mean annual ambient temperature (ACI 308). Internal concrete temperature gain can be controlled a number of ways: (1) a low cement content—120 to 270 kg/m³ (200 to 450 lb per cubic yard); (2) large aggregate size—75 to 150 mm (3 to 6 in.); (3) high coarse aggregate content—up to 80% of total aggregate; (4) low-heat-of-hydration cement; (5) pozzolans—where heat of hydration of a pozzolan can be 25% to 75% that of cement; (6) reductions in the initial concrete temperature to be cooling the concrete ingredients; (7) cooling the concrete through the use of embedded cooling pipes; (8) steel forms for rapid heat dissipation; (9) water curing; and (10) low lifts—1.5 m (5 ft) or less during placement. In massive structures of high volume-to-surface ratio, an estimate of the adiabatic temperature rise can be made using equations in Gajda (2007).

Massive structural reinforced concrete members with high cement contents (300 to 600 kg per cubic meter or 500 to 1000 lb per cu yard) cannot use many of the placing techniques and controlling factors mentioned above to maintain low temperatures to control cracking. For these concretes (often used in bridges, foundations, and power plants), a good technique is to (1) avoid external restraint from adjacent concrete elements, (2) reduce the size of the member by placing the concrete in multiple smaller pours, or (3) control internal differential thermal strains by preventing the concrete from experiencing an excessive temperature differential between the surface and the center. The latter is done by properly designing the concrete and either keeping the concrete surface warm through use of insulation or reducing the internal concrete temperature by precooling of the concrete or postcooling with internal cooling pipes.

Studies and experience have shown that by limiting the maximum temperature differential between the interior and exterior surface of the concrete to less than about 20°C (36°F), surface cracking can be minimized or avoided (FitzGibbon 1977 and Fintel and Ghosh 1978). Some sources indicate that the maximum temperature differential (MTD) for concrete containing granite or limestone (low-thermal-coefficient aggregates) should be 25°C and 31°C (45°F and 56°F), respectively (Bamforth 1981). The actual MTD for a particular mass concrete placement and concrete mix design can be determined using equations in ACI 207 (2007).

In general, an MTD of 20°C (36°F) should be assumed unless a demonstration or calculations based on physical properties of the actual concrete mix the geometry of the concrete member show that higher MTD values are allowable. By limiting the temperature differential to 20°C (36°F) or less, the concrete will cool slowly to ambient temperature with little or no surface cracking; however, this is true only if the member is not restrained by continuous reinforcement crossing the interface of adjacent or opposite sections of hardened concrete. Restrained concrete will tend to crack due to eventual thermal contraction after the cool down. Unrestrained concrete should not crack if proper procedures are followed and the temperature differential is monitored and controlled. If there is any concern over excess temperature differentials in a concrete member, the element should be considered as mass concrete and appropriate precautions taken.

Fig. 18-10 illustrates the relationship between temperature rise, cooling, and temperature differentials for a section of mass concrete. As can be observed, if the forms (which are providing adequate insulation in this case) are removed too early, cracking will occur once the difference between interior and surface concrete temperatures exceeds the critical temperature differential of 20°C (36°F). If higher temperature differentials are permissible, the forms can be removed sooner. For large concrete placements, surface insulation may be need for an extended period of time of up to several weeks or longer.
The maximum temperature rise can be estimated by approximation, if the concrete contains 300 to 600 kg of cement per cubic meter (500 to 1,000 lb of Type I/II cement per cubic yard) and the least dimension of the member is 1.8 m (6 ft). This approximation (under normal, not adiabatic conditions) would be 12°C for every 100 kg of cement per cubic meter (12.8°F for every 100 lb of cement per cubic yard). For example, the maximum temperature of such an element made with concrete having 535 kg of Type I/II cement per cubic meter (900 lb of cement per cubic yard) and cast at 16°C (60°F) would be about

\[16°C + (12°C \times 535/100)\text{ or } 80°C\]

\[(60°F + [12.8°F \times 900/100])\text{ or } 175°F\]

Temperatures and temperature differences in mass concrete can also be calculated by a method in ACI 207 (2005).

The slow rate of heat exchange between concrete and its surroundings is due to the concrete’s heat capacity. Heat escapes from concrete at a rate that is inversely proportional to the square of its least dimension. A 150-mm (6-in.) thick wall cooling from both sides will take approximately 1½ hours to dissipate 95% of its developed heat. A 1.5-m (5-ft) thick wall would take an entire week to dissipate the same amount of heat (ACI 207). Inexpensive thermocouples can be used to monitor concrete temperature.

### PREPLACED AGGREGATE CONCRETE

Preplaced aggregate concrete is produced by first placing coarse aggregate in a form and later injecting a cement-sand grout, usually with admixtures, to fill the voids. Properties of the resulting concrete are similar to those of comparable concrete placed by conventional methods; however, considerably less thermal and drying shrinkage can be expected because of the point-to-point contact of aggregate particles.

Coarse aggregates should meet requirements of ASTM C33 (AASHTO M 80). In addition, most specifications limit both the maximum and minimum sizes; for example, 75-mm (3-in.) maximum and 12.5-mm (½-in.) minimum. Aggregates are generally graded to produce a void content of 35% to 40%. Fine aggregate used in the grout is generally graded to a fineness modulus of between 1.2 and 2.0, with nearly all of the material passing a 1.25-mm (No. 16) sieve.

Although the preplaced aggregate method has been used principally for restoration work and in the construction of reactor shields, bridge piers, and underwater structures, it has also been used in buildings to produce unusual architectural effects. Since the forms are completely filled with coarse aggregate prior to grouting, a dense and uniform exposed-aggregate facing is obtained when the surface is sandblasted, tooled, or retarded and wire-brushed at an early age.

Tests for preplaced aggregate concrete are given in ASTM C937 through C943. Preplaced aggregate concrete is discussed in more detail in ACI 304-00, Guide for Measuring, Transporting, and Placing Concrete.

### NO-SLUMP CONCRETE

No-slump concrete is defined as concrete with a consistency corresponding to a slump of 6 mm (¼ in.) or less. Such concrete, while very dry, must be sufficiently workable to be placed and consolidated with the equipment to be used on the job. The methods referred to here do not necessarily apply to mixtures for concrete masonry units or for compaction by spinning techniques.

Many of the basic laws governing the properties of higher-slump concretes are applicable to no-slump concrete. For example, the properties of hardened concrete depend principally on the ratio of water to cement, provided the mix is properly consolidated.

Measurement of the consistency of no-slump concrete differs from that for higher-slump concrete because the slump cone is impractical for use with the drier consistencies. ACI 211.3, Guide for Selecting Proportions for No-Slump Concrete, describes three methods for measuring the consistency of no-slump concrete: (1) the Vebe apparatus; (2) the compacting-factor test; and (3) the Thaulow drop table. In the absence of the above test equipment, workability can be adequately judged by a trial mixture that is placed and compacted with the equipment and methods to be used on the job.

Intentionally entrained air is recommended for no-slump concrete where durability is required. The amount of air-entraining admixture usually recommended for higher-slump concretes will not produce air contents in no-slump concretes that are as high as those in the higher-slump concretes. The lower volume of entrained air, however, generally provides adequate durability for no-slump concretes; while the volume of entrained air is not there, sufficient small air voids are present. This departure from the usual methods of designing and controlling entrained air is necessary for no-slump concretes.

For a discussion of water requirements and computation of trial mixtures, see ACI 211.3.
ROLLER-COMPACTED CONCRETE

Roller-compacted concrete (RCC) is a lean, no-slump, almost dry concrete that is compacted in place by vibratory roller or plate compaction equipment (Fig. 18-11). RCC is a mixture of aggregate, cement, and water; supplementary cementing materials such as fly ash also have been used. Cement contents range from 60 to 360 kg per cubic meter (100 to 600 lb per cubic yard). Mixing is done with conventional batch mixers, continuous mixers, or in some instances tilting-drum truck mixers.

Applications for RCC fall into two distinct categories—water control structures (dams) and pavements. While the same term is used to describe both types of concrete use, the design and construction processes are different.

Water Control Structures

RCC can be used for the entire dam structure, or as an overtopping protection on the upper section and on the downstream face. The nominal maximum aggregate size can range up to 150 mm (6 in.). The zero slump mix is produced in a high-capacity central-mixing plant near the site and delivered by truck or by conveyor belt. Cement content is usually lower than that used in a conventional concrete mix, but similar to that of mass concrete. Compressive strengths ranging from 7 to 30 MPa (1000 to 4500 psi) have been obtained from roller-compacted concrete in dam projects. The RCC mix is transported by trucks and conveyor belts and spread by grader or bulldozer, followed by rolling with vibratory compactors. No forms are used. On some projects the upstream face is surfaced with higher strength conventional air-entrained concrete or precast panels for improved durability.

RCC dams have the advantage of allowing much steeper slopes on both faces than an earth fill dam. In addition to the advantage of using less material, the dam is completed and placed in service earlier, usually at a significant savings in overall cost compared to an earth fill structure.

Other water control RCC applications include use as an emergency spillway or overtopping protection for embankment dams, low permeable liner for settling ponds, bank protection, and grade control structure for channels and riverbeds.

Pavements

The uses for RCC paving range from pavements as thick as one meter (one yard) for the mining industry to city streets, paved surfaces for composting operations, logging, truck staging areas, and warehouse floors. The procedures for construction of an RCC pavement require tighter control than for dam construction (Arnold and Zamensky 2000). Cement content is in the same range as conventional concrete, 300 to 360 kg/m$^3$ (500 to 600 lb/yd$^3$), and compressive strength is of the same order, 30 to 40 MPa (4000 to 6000 psi). The nominal maximum aggregate size is limited to 19 mm (¾ in.) to provide a smooth, dense surface. For even better surface textures, a 16 mm (5⁄8 in.) maximum size aggregate is recommended.

The zero slump mix is usually produced in a continuous flow pugmill mixer at production rates as high as 400 tons per hour. It is possible to mix RCC in a central batch plant, but the plant must be dedicated to RCC production exclusively, because the material tends to stick to the inside drums. Specifications usually require that the mix be transported, placed, and compacted within 60 minutes of the start of mixing; although ambient weather conditions may increase or decrease that time window.

RCC is typically placed in layers 125 to 250 mm (5 to 10 in.) in thickness using an asphalt-type paving machine. High-density paving equipment is preferred for layers thicker than 150 mm (6 in.) since the need for subsequent compaction by rollers is reduced. Where a design calls for pavement thickness greater than 250 mm (10 in.), the RCC should be placed in multiple layers. In this type of construction, it is important that there be a minimum time delay in placing subsequent layers so that good bond is assured. Following placement by a paver, RCC can be compacted with a combination of vibratory steel-wheeled rollers and rubber-tired equipment.

Curing is vitally important in RCC pavement construction. The very low water content at the initial mixing stage means that an RCC mix will dry out very quickly once it is in place. Continuous water curing is the recommended method, although sprayed on asphaltic emulsion, plastic sheeting, and concrete curing compounds have been used in some cases. Pavement projects have had design compressive strengths of about 35 MPa (5000 psi) with field strengths in the range of 35 to 70 MPa (5000 to 10,000 psi) (Hansen 1987).

High-performance roller compacted concrete for areas subjected to high impact and abrasive loading were developed in the mid-1990’s. These mixes are based on obtaining the optimum packing of the various sizes of aggregate particles, and the addition of silica fume to the mix (Marchand and others 1997 and Reid and others 1998).

**SOIL-CEMENT**

Soil-cement is a mixture of pulverized soil or granular material, cement, and water. Some other terms applied to soil-cement are “cement-treated base or subbase,” “cement stabilization,” “cement-modified soil,” and “cement-treated aggregate.” The mixture is compacted to a high density, and as the cement hydrates the material becomes hard and durable.

Soil-cement is primarily used as pavement base course for roads, streets, airports, and parking areas. A bituminous or portland cement concrete wearing course is usually placed over the base. Soil-cement is also used as slope protection for earth dams and embankments, reservoir and ditch linings, deep-soil mixing, and foundation stabilization (Fig. 18-12).

The soil material in soil-cement can be almost any combination of sand, silt, clay, and gravel or crushed stone. Local granular materials (such as slag, caliche, limestone, and scoria) plus a wide variety of waste materials (such as cinders, ash, and screenings from quarries and gravel pits) can be used to make soil-cement. Also, old granular-base roads, with or without their bituminous surfaces, can be recycled to make soil-cement.

Soil-cement should contain sufficient portland cement to resist deterioration from freeze-thaw and wet-dry cycling and sufficient moisture for maximum compaction. Cement contents range from 80 to 255 kg per cubic meter (130 to 430 lb per cubic yard).

There are four steps in soil-cement construction: spreading cement, mixing, compaction, and curing. The proper quantity of cement must be spread on the in-place soil; the cement and soil material, and the necessary amount of water are mixed thoroughly using any of several types of mixing machines; and finally, the mixture is compacted with conventional road-building equipment to 96% to 100% of maximum density. See ASTM D558 (AASHTO T 134) and PCA 1992.

A light coat of bituminous material is commonly used to prevent moisture loss; it also helps bond and forms part of the bituminous surface. A common type of wearing surface for light traffic is a surface treatment of bituminous material and chips 13 to 19 mm (½ to ¾ in.) thick. For heavy-duty use and in severe climates a 38-mm (11/2-in.) asphalt mat is used.

Depending on the soil used, the 7-day compressive strengths of saturated specimens at the minimum cement content meeting soil-cement criteria are generally between 2 to 5 MPa (300 and 800 psi). Like concrete, soil-cement continues to gain strength with age; compressive strengths in excess of 17 MPa (2500 psi) have been obtained after many years of service.

See ACI Committee 230 (1990) and PCA (1995) for detailed information on soil-cement construction.

**SHOTCRETE**

Shotcrete is mortar or small-aggregate concrete that is pneumatically projected onto a surface at high velocity (Fig. 18-13). Also known as “gunite” and “sprayed concrete,” shotcrete was developed in 1911 and its concept is essentially unchanged even in today’s use. The relatively
Shrinkage-compensating concrete is used in concrete slabs, pavements, structures, and repair work to minimize drying shrinkage cracks. Expansion of concrete made with shrinkage-compensating cement should be determined by the method specified in ASTM C878.

Reinforcing steel in the structure restrains the concrete and goes into tension as the shrinkage compensating concrete expands. Upon shrinking due to drying contraction caused by moisture loss in hardened concrete, the tension in the steel is relieved; as long as the resulting tension in the concrete does not exceed the tensile strength of the concrete, no cracking should result. Shrinkage-compensating concrete can be proportioned, batched, placed, and cured similarly to normal concrete with some precautions; for example, it is necessary to assure the expected expansion by using additional curing. More information can be found in Chapter 2 and in ACI 223-98, Standard Practice for the Use of Shrinkage-Compensating Concrete.

PERVIOUS CONCRETE

Pervious (porous or no-fines) concrete contains a narrowly graded coarse aggregate, little to no fine aggregate, and insufficient cement paste to fill voids in the coarse aggregate. This low water-cement ratio, low-slump concrete resembling popcorn is primarily held together by cement paste at the contact points of the coarse aggregate particles; this produces a concrete with a high volume of voids (20% to 35%) and a high permeability that allows water to flow through it easily (Tennis, Leming, and Akers 2004).

Pervious concrete is used in hydraulic structures as drainage media, and in parking lots, pavements, and airport runways to reduce stormwater run off (Fig. 18-14). It also recharges the local groundwater supply by allowing
water to penetrate the concrete to the ground below. Pervious concretes have also been used in tennis courts and greenhouses. Leming, Malcom, and Tennis (2007) describe the fundamental hydrologic behavior of pervious concrete pavement systems.

As a paving material, porous concrete is raked or slip-formed into place with conventional paving equipment and then roller compacted. Vibratory screeds or hand rollers can be used for smaller jobs. In order to maintain its porous properties, the surfaces of pervious concrete should not be closed up or sealed; therefore, troweling and finishing are not desired. The compressive strength of different mixes can range from 3.5 to 27.5 MPa (500 to 4000 psi). Drainage rates commonly range from 100 to 900 liters per minute per square meter (2 to 18 gallons per minute per square foot).

No-fines concrete is used in building construction (particularly walls) for its thermal insulating properties. For example, a 250-mm (10-in.) thick porous-concrete wall can have an R value of 0.9 (5 using inch-pound units) compared to 0.125 (0.75) for normal concrete. No-fines concrete is also lightweight, 1600 to 1900 kg/m³ (100 to 120 pcf), and has low drying shrinkage properties (Malhotra 1976 and Concrete Construction 1983).

**WHITE AND COLORED CONCRETE**

**White Concrete**

White portland cement is used to produce white concrete, a widely used architectural material (Fig. 18-15). It is also used in mortar, plaster, stucco, terrazzo, and portland cement paint. White portland cement is manufactured from raw materials of low iron content; it conforms to ASTM C150 (AASHTO M 85) even though these specifications do not specifically mention white portland cement.

White concrete is made with aggregates and water that contain no materials that will discolor the concrete. White or light-colored aggregates can be used. Oil that could stain concrete should not be used on the forms. Care must be taken to avoid rust stains from tools and equipment. Curing materials that could cause stains must be avoided. Refer to Farny (2001) and http://www.cement.org/decorative for more information.

**Colored Concrete**

Colored concrete can be produced by using colored aggregates or by adding color pigments (ASTM C979) or both. When colored aggregates are used, they should be exposed at the surface of the concrete. This can be done several ways; for example, casting against a form that has been treated with a retarder. Unhydrated paste at the surface is later brushed or washed away. Other methods involve removing the surface mortar by sandblasting, waterblasting, bushhammering, grinding, or acid washing. If surfaces are to be washed with acid, a delay of approximately two weeks after casting is necessary. Colored aggregates may be natural rock such as quartz, marble, and granite, or they may be ceramic materials.

Pigments for coloring concrete should be pure mineral oxides ground finer than cement; they should be insoluble in water, free of soluble salts and acids, colorfast in sunlight, resistant to alkalies and weak acids, and virtually free of calcium sulfate. Mineral oxides occur in nature and are also produced synthetically; synthetic pigments generally give more uniform results.

The amount of color pigments added to a concrete mixture should not be more than 10% of the mass of the cement. The amount required depends on the type of pigment and the color desired. For example, a dose of pigment equal to 1.5% by mass of cement may produce a pleasing pastel color, but 7% may be needed to produce a deep color. Use of white portland cement with a pigment will produce cleaner, brighter colors and is recommended in preference to gray cement, except for black or dark gray colors (Fig. 18-16).
To maintain uniform color, do not use calcium chloride, and batch all materials carefully by mass. To prevent streaking, the dry cement and color pigment must be thoroughly blended before they are added to the mixer. Mixing time should be longer than normal to ensure uniformity.

In air-entrained concrete, the addition of pigment may require an adjustment in the amount of air-entraining admixture to maintain the desired air content.

**Dry-Shake Method.** Slabs or precast panels that are cast horizontally can be colored by the dry-shake method. Prepackaged, dry coloring materials consisting of mineral oxide pigment, white portland cement, and specially graded silica sand or other fine aggregate are marketed ready for use by various manufacturers.

After the slab has been bull floated once, two-thirds of the dry coloring material should be broadcast evenly by hand over the surface. The required amount of coloring material can usually be determined from previously cast sections. After the material has absorbed water from the fresh concrete, it should be floated into the surface. Then the rest of the material should be applied immediately at right angles to the initial application, so that a uniform color is obtained. The slab should again be floated to work the remaining material into the surface.

Other finishing operations may follow depending on the type of finish desired. Curing should begin immediately after finishing; take precautions to prevent discoloring the surface. See Kosmatka and Collins (2004) for more information.

**POLYMER-PORTLAND CEMENT CONCRETE**

Polymer-portland cement concrete (PPCC), also called polymer-modified concrete, is basically normal portland cement concrete to which a polymer or monomer has been added during mixing to improve durability and adhesion. Thermoplastic and elastomeric latexes are the most commonly used polymers in PPCC, but epoxies and other polymers are also used. In general, latex improves ductility, durability, adhesive properties, resistance to chloride-ion ingress, shear bond, and tensile and flexural strength of concrete and mortar. Latex-modified concretes (LMC) also have excellent freeze-thaw, abrasion, and impact resistance. Some LMC materials can also resist certain acids, alkalies, and organic solvents. Polymer-portland cement concrete is primarily used in concrete patching and overlays, especially bridge decks. See ACI 548.3R for more information on polymer-modified concrete and ACI 548.4 for LMC overlays.

**FERROCEMENT**

Ferrocement is a special type of reinforced concrete composed of closely spaced layers of continuous relatively thin metallic or nonmetallic mesh or wire embedded in mortar. It is constructed by hand plastering, shotcreting, laminating (forcing the mesh into fresh mortar), or a combination of these methods.

The mortar mixture generally has a sand-cement ratio of 1.5 to 2.5 and a water-cement ratio of 0.35 to 0.50. Reinforcement makes up about 5% to 6% of the ferrocement volume. Fibers and admixtures may also be used to modify the mortar properties. Polymers or cement-based coatings are often applied to the finished surface to reduce porosity.

Ferrocement is considered easy to produce in a variety of shapes and sizes; however, it is labor intensive. Ferrocement is used to construct thin shell roofs, swimming pools, tunnel linings, silos, tanks, prefabricated houses, barges, boats, sculptures, and thin panels or sections usually less than 25 mm (1 in.) thick (ACI 549R and ACI 549.1R).

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Appendix

GLOSSARY
The intent of this Glossary is to clarify terminology used in concrete construction, with special emphasis on those terms used in Design and Control of Concrete Mixtures. Additional terminology that may not be in this book is included in the Glossary for the convenience of our readers. Other sources for terms include ACI Committee 116, and ASTM standards.

Absorption—see Water absorption.

Accelerating admixture—admixture that speeds the rate of hydration of hydraulic cement, shortens the normal time of setting, or increases the rate of hardening, of strength development, or both, of portland cement, concrete, mortar, grout, or plaster.

Addition—substance that is interground or blended in limited amounts into a hydraulic cement during manufacture—not at the jobsite—either as a “processing addition” to aid in manufacture and handling of the cement or as a “functional addition” to modify the useful properties of the cement.

Admixture—material, other than water, aggregate, and hydraulic cement, used as an ingredient of concrete, mortar, grout, or plaster and added to the batch immediately before or during mixing.

Aggregate—granular mineral material such as natural sand, manufactured sand, gravel, crushed stone, air-cooled blast-furnace slag, vermiculite, or perlite.

Air content—total volume of air voids, both entrained and entrapped, in cement paste, mortar, or concrete. Entrained air adds to the durability of hardened mortar or concrete and the workability of fresh mixtures.

Air entrainment—intentional introduction of air in the form of minute, disconnected bubbles (generally smaller than 1 mm) during mixing of portland cement concrete, mortar, grout, or plaster to improve desirable characteristics such as cohesion, workability, and durability.

Air-entraining admixture—admixture for concrete, mortar, or grout that will cause air to be incorporated into the mixture in the form of minute bubbles during mixing, usually to increase the material's workability and frost resistance.

Air-entraining portland cement—Portland cement containing an air-entraining addition added during its manufacture.

Air void—entrapped air pocket or an entrained air bubble in concrete, mortar, or grout. Entrapped air voids usually are larger than 1 mm in diameter; entrained air voids are smaller. Most of the entrapped air voids should be removed with internal vibration, power screeding, or rodding.

Alkali-aggregate reactivity—production of expansive gel caused by a reaction between aggregates containing certain forms of silica or carbonates and alkali hydroxides in concrete.

Architectural concrete—concrete that will be permanently exposed to view and which therefore requires special care in selection of concrete ingredients, forming, placing, consolidating, and finishing to obtain the desired architectural appearance.

Autoclaved cellular concrete—concrete containing very high air content resulting in low density, and cured at high temperature and pressure in an autoclave.

Batching—process of weighing or volumetrically measuring and introducing into the mixer the ingredients for a batch of concrete, mortar, grout, or plaster.

Blast-furnace slag—nonmetallic byproduct of steel manufacturing, consisting essentially of silicates and aluminum silicates of calcium that are developed in a molten condition simultaneously with iron in a blast furnace.

Bleeding—flow of mixing water from a newly placed concrete mixture caused by the settlement of the solid materials in the mixture.
**Blended hydraulic cement**—cement containing combinations of portland cement, pozzolans, slag, and/or other hydraulic cement.

**Bulking**—increase in volume of a quantity of sand when in a moist condition compared to its volume when in a dry state.

**Calcined clay**—clay heated to high temperature to alter its physical properties for use as a pozzolan or cementing material in concrete.

**Calcined shale**—shale heated to high temperature to alter its physical properties for use as a pozzolan or cementing material in concrete.

**Carbonation**—reaction between carbon dioxide and a hydroxide or oxide to form a carbonate.

**Cellular concrete**—high air content or high void ratio concrete resulting in low density.

**Cement**—see Portland cement and Hydraulic cement.

**Cement paste**—constituent of concrete, mortar, grout, and plaster consisting of cement and water.

**Cementitious material (cementing material)**—any material having cementing properties or contributing to the formation of hydrated calcium silicate compounds. When proportioning concrete, the following are considered cementitious materials: portland cement, blended hydraulic cement, fly ash, ground granulated blast-furnace slag, silica fume, calcined clay, metakaolin, calcined shale, and rice husk ash.

**Chemical admixture**—see Admixture.

**Chemical bond**—bond between materials resulting from cohesion and adhesion developed by chemical reaction.

**Clinker**—end product of a portland cement kiln; raw cementitious material prior to grinding

**Chloride (attack)**—chemical compounds containing chloride ions, which promote the corrosion of steel reinforcement. Chloride deicing chemicals are primary sources.

**Coarse aggregate**—natural gravel, crushed stone, or iron blast-furnace slag, usually larger than 5 mm (0.2 in.) and commonly ranging in size between 9.5 mm and 37.5 mm (⅜ in. to 1½ in.).

**Cohesion**—mutual attraction by which elements of a substance are held together.

**Colored concrete**—concrete containing white cement and/or mineral oxide pigments to produce colors other than the normal gray hue of traditional gray cement concrete.

**Compaction**—process of inducing a closer arrangement of the solid particles in freshly mixed and placed concrete, mortar, or grout by reduction of voids, usually by vibration, tamping, rodging, puddling, or a combination of these techniques. Also called consolidation.

**Compressive strength**—maximum resistance that a concrete, mortar, or grout specimen will sustain when loaded axially in compression in a testing machine at a specified rate; usually expressed as force per unit of cross sectional area, such as megapascals (MPa) or pounds per square inch (psi).

**Concrete**—mixture of binding materials and coarse and fine aggregates. Portland cement and water are commonly used as the binding medium for normal concrete mixtures, but may also contain pozzolans, slag, and/or chemical admixtures.

**Consistency**—relative mobility or ability of freshly mixed concrete, mortar, or grout to flow. (See also Slump and Workability.)

**Construction joint**—a stopping place in the process of construction. A true construction joint allows for bond between new concrete and existing concrete and permits no movement. In structural applications their location must be determined by the structural engineer. In slab on grade applications, construction joints are often located at contraction (control) joint locations and are constructed to allow movement and perform as contraction joints.

**Contraction joint**—weakened plane to control cracking due to volume change in a concrete structure. Joint may be grooved, sawed, or formed. Also known as a “Control joint.”

**Corrosion**—deterioration of metal by chemical, electrochemical, or electrolytic reaction.

**Creep**—time-dependent deformation of concrete, or of any material, due to a sustained load.

**Curing**—process of maintaining freshly placed concrete mortar, grout, or plaster moist and at a favorable temperature for a suitable period of time during its early stages so that the desired properties of the material can develop. Curing assures satisfactory hydration and hardening of the cementitious materials.

**Dampproofing**—treatment of concrete, mortar, grout, or plaster to retard the passage or absorption of water, or water vapor.

**Density**—mass per unit volume; the weight per unit volume in air, expressed, for example, in kg/m³ (lb/ft³).

**Durability**—ability of portland cement concrete, mortar, grout, or plaster to resist weathering action and other conditions of service, such as chemical attack, freezing and thawing, and abrasion.

**Early stiffening**—rapidly developing rigidity in freshly mixed hydraulic cement paste, mortar, grout, plaster, or concrete.

**Entrapped air**—irregularly shaped, unintentional air voids in fresh or hardened concrete 1 mm or larger in size.
**Entrained air**—spherical microscopic air bubbles—usually 10 µm to 1000 µm in diameter—intentionally incorporated into concrete to provide freezing and thawing resistance and/or improve workability.

**Epoxy resin**—class of organic chemical bonding systems used in the preparation of special coatings or adhesives for concrete or masonry or as binders in epoxy-resin mortars and concretes.

**Ettringite**—needle like crystalline compound produced by the reaction of C₃A, gypsum, and water within a portland cement concrete.

**Expansion joint**—a separation provided between adjoining parts of a structure to allow movement.

**Ferrocement**—one or more layers of steel or wire reinforcement encased in portland cement mortar creating a thin-section composite material.

**Fibers**—thread or thread like material ranging from 0.05 to 4 mm (0.002 to 0.16 in.) in diameter and from 10 to 150 mm (0.5 to 6 in.) in length and made of steel, glass, synthetic (plastic), carbon, or natural materials.

**Fiber concrete**—concrete containing randomly oriented fibers in 2 or 3 dimensions through out the concrete matrix.

**Fine aggregate**—aggregate that passes the 9.5-mm (%)-in.) sieve, almost entirely passes the 4.75-mm (No. 4) sieve, and is predominantly retained on the 75-µm (No. 200) sieve.

**Fineness modulus (FM)**—factor obtained by adding the cumulative percentages of material in a sample of aggregate retained on each of a specified series of sieves and dividing the sum by 100.

**Finishing**—mechanical operations like screeding, consolidating, floating, troweling, or texturing that establish the final appearance of any concrete surface.

**Fire resistance**—that property of a building material, element, or assembly to withstand fire or give protection from fire; it is characterized by the ability to confine a fire or to continue to perform a given structural function during a fire, or both.

**Flexural strength**—ability of solids to resist bending.

**Fly ash**—residue from coal combustion, which is carried in flue gases, and is used as a pozzolan or cementing material in concrete.

**Forms**—temporary supports for keeping fresh concrete in place until it has hardened to such a degree as to be self supporting (when the structure is able to support its dead load).

**Freeze-thaw resistance**—ability of concrete to withstand cycles of freezing and thawing. (See also Air entrainment and Air-entraining admixture.)

**Fresh concrete**—concrete that has been recently mixed and is still workable and plastic.

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**Grading**—size distribution of aggregate particles, determined by separation with standard screen sieves.

**Grout**—mixture of cementitious material with or without aggregate or admixtures to which sufficient water is added to produce a pouring or pumping consistency without segregation of the constituent materials.

**Hardened concrete**—concrete that is in a solid state and has developed a certain strength.

**High-density concrete (heavyweight concrete)**—concrete of very high density; normally designed by the use of heavyweight aggregates.

**High-strength concrete**—concrete with a design strength of at least 70 MPa (10,000 psi).

**Honeycomb**—term that describes the failure of mortar to completely surround coarse aggregates in concrete, leaving empty spaces (voids) between them.

**Hydrated lime**—dry powder obtained by treating quicklime with sufficient water to satisfy its chemical affinity for water; consists essentially of calcium hydroxide or a mixture of calcium hydroxide and magnesium oxide or magnesium hydroxide, or both.

**Hydration**—in concrete, mortar, grout, and plaster, the chemical reaction between hydraulic cement and water in which new compounds with strength-producing properties are formed.

**Hydraulic cement**—cement that sets and hardens by chemical reaction with water, and is capable of doing so under water. (See also Portland cement.)

**Inch-pound units**—units of length, area, volume, weight, and temperature commonly used in the United States during the 18th to 20th centuries. These include, but are not limited to: (1) length— inches, feet, yards, and miles; (2) area—square inches, square feet, square yards, and square miles; (3) volume—cubic inches, cubic feet, cubic yards, gallons, and ounces; (4) weight— pounds and ounces; and (5) temperature—degrees Fahrenheit.

**Isolation Joint**—separation that allows adjoining parts of a structure to move freely to one another, both horizontally and vertically.

**Joint**—see Construction joint, Contraction joint, Isolation joint, and Expansion joint.

**Kiln**—rotary furnace used in cement manufacture to heat and chemically combine raw inorganic materials, such as limestone, sand and clay, into calcium silicate clinker.
**Lightweight aggregate**—low-density aggregate used to produce lightweight (low-density) concrete. Could be expanded or sintered clay, slate, diatomaceous shale, perlite, vermiculite, or slag; natural pumice, scoria, volcanic cinders, tuff, or diatomite; sintered fly ash or industrial cinders.

**Lightweight concrete**—low-density concrete compared to normal-density concrete.

**Lime**—general term that includes the various chemical and physical forms of quicklime, hydrated lime, and hydraulic lime. It may be high-calcium, magnesian, or dolomitic.

**Masonry**—concrete masonry units, clay brick, structural clay tile, stone, terra cotta, and the like, or combinations thereof, bonded with mortar, dry-stacked, or anchored with metal connectors to form walls, building elements, pavements, and other structures.

**Masonry cement**—hydraulic cement, primarily used in masonry and plastering construction, consisting of a mixture of portland or blended hydraulic cement and plasticizing materials (such as limestone, hydrated or hydraulic lime) together with other materials introduced to enhance one or more properties such as setting time, workability, water retention, and durability.

**Mass concrete**—cast-in-place concrete in volume large enough to require measures to compensate for volume change caused by temperature rise from heat of hydration in order to keep cracking to a minimum.

**Metakaolin**—highly reactive pozzolan made from kaolin clays.

**Metric units**—also called System International (SI) Units. System of units adopted by most of the world by the 21st Century. These include but are not limited to: (1) length—millimeters, meters, and kilometers; (2) area—square millimeters and square meters; (3) volume—cubic meters and liters; (4) mass—milligrams, grams, kilograms, and megagrams; and (5) degrees Celsius.

**Mineral admixtures**—see Supplementary cementitious materials.

**Modulus of elasticity**—ratio of normal stress to corresponding strain for tensile or compressive stress below the proportional limit of the material; also referred to as elastic modulus, Young’s modulus, and Young’s modulus of elasticity; denoted by the symbol \( E \).

**Moist-air curing**—curing with moist air (no less than 95% relative humidity) at atmospheric pressure and a temperature of about 23°C (73°F).

**Mortar**—mixture of cementitious materials, fine aggregate, and water, which may contain admixtures, and is usually used to bond masonry units.

**Mortar cement**—hydraulic cement, primarily used in masonry construction, consisting of a mixture of portland or blended hydraulic cement and plasticizing materials (such as limestone, hydrated or hydraulic lime) together with other materials introduced to enhance one or more properties such as setting time, workability, water retention, and durability. Mortar cement and masonry cement are similar in use and function. However, specifications for mortar cement usually require lower air contents and they include a flexural bond strength requirement.

**Normal weight concrete**—class of concrete made with normal density aggregates, usually crushed stone or gravel, having a density of approximately 2400 kg/m³ (150 lb/ft³). (See also Lightweight concrete and High-density concrete.)

**No-slump concrete**—concrete having a slump of less than 6 mm (¼ in.).

**Overlay**—layer of concrete or mortar placed on or bonded to the surface of an existing pavement or slab. Normally done to repair a worn or cracked surface. Overlays are seldom less than 25 mm (1 in.) thick.

**Pavement (concrete)**—highway, road, street, path, or parking lot surfaced with concrete. Although typically applied to surfaces that are used for travel, the term also applies to storage areas and playgrounds.

**Permeability**—property of allowing passage of fluids or gases.

**Pervious concrete (no-fines or porous concrete)**—concrete containing insufficient fines or no fines to fill the voids between aggregate particles in a concrete mixture. The coarse aggregate particles are coated with a cement and water paste to bond the particles at their contact points. The resulting concrete contains an interconnected pore system allowing storm water to drain through the concrete to the subbase below.

**pH**—chemical symbol for the logarithm of the reciprocal of hydrogen ion concentration in gram atoms per liter, used to express the acidity or alkalinity (base) of a solution on a scale of 0 to 14, where less than 7 represents acidity, and more than 7 alkalinity

**Plastic cement**—special hydraulic cement product manufactured for plaster and stucco application. One or more inorganic plasticizing agents are interground or blended with the cement to increase the workability and molding characteristics of the resultant mortar, plaster, or stucco.

**Plasticity**—that property of freshly mixed cement paste, concrete, mortar, grout, or plaster that determines its workability, resistance to deformation, or ease of molding.

**Plasticizer**—admixture that increases the plasticity of portland cement concrete, mortar, grout, or plaster.
Polymer-portland cement concrete—fresh portland cement concrete to which a polymer is added for improved durability and adhesion characteristics, often used in overlays for bridge decks; also referred to as polymer-modified concrete and latex-modified concrete.

Popout—shallow depression in a concrete surface resulting from the breaking away of pieces of concrete due to internal pressure.

Portland blast-furnace slag cement—hydraulic cement consisting of: (1) an intimately interground mixture of portland cement clinker and granulated blast-furnace slag; (2) an intimate and uniform blend of portland cement and fine granulated blast-furnace slag; or (3) finely ground blast-furnace slag with or without additions.

Portland cement—Calcium silicate hydraulic cement produced by pulverizing portland cement clinker, and usually containing calcium sulfate and other compounds. (See also Hydraulic cement.)

Portland cement plaster—a combination of portland cement-based cementitious material(s) and aggregate mixed with a suitable amount of water to form a plastic mass that will adhere to a surface and harden, preserving any form and texture imposed on it while plastic. See also Stucco.

Portland-pozzolan cement—hydraulic cement consisting of an intimate and uniform blend of portland cement or portland blast-furnace slag cement and fine pozzolan produced by intergrinding portland cement clinker and pozzolan, by blending portland cement or portland blast-furnace slag cement and finely divided pozzolan, or a combination of intergrinding and blending, in which the amount of the pozzolan constituent is within specified limits.

Pozzolan—siliceous or siliceous and aluminous materials, like fly ash or silica fume, which in itself possess little or no cementitious value but which will, in finely divided form and in the presence of moisture, chemically react with calcium hydroxide at ordinary temperatures to form compounds possessing cementitious properties.

Precast concrete—concrete cast in forms in a controlled environment and allowed to achieve a specified strength prior to placement on location.

Prestressed concrete—concrete in which compressive stresses are induced by high-strength steel tendons or bars in a concrete element before loads are applied to the element which will balance the tensile stresses imposed in the element during service. This may be accomplished by the following: Post-tensioning—a method of prestressing in which the tendons/bars are tensioned after the concrete has hardened; or Pre-tensioning—a method of prestressing in which the tendons are tensioned before the concrete is placed.

Polymer-portland cement concrete

Quality control—actions taken by a producer or contractor to provide control over what is being done and what is being provided so that applicable standards of good practice for the work are followed.

Reactive-powder concrete—high-strength, low-water and low-porosity concrete with high silica content and aggregate particle sizes of less than 0.3 mm.

Ready-mixed concrete—concrete manufactured for delivery to a location in a fresh state

Recycled concrete—hardened concrete that has been processed for reuse, usually as an aggregate.

Reinforced concrete—concrete to which tensile bearing materials such as steel rods or metal wires are added for tensile strength.

Relative density—a ratio relating the mass of a volume of material to that of water; also called specific gravity.

Relative humidity—The ratio of the quantity of water vapor actually present in the atmosphere to the amount of water vapor present in a saturated atmosphere at a given temperature, expressed as a percentage.

Retarder—an admixture that delays the setting and hardening of concrete.

Roller-compacted concrete (RCC)—a zero slump mix of aggregates, cementitious materials and water that is consolidated by rolling with vibratory compactors; typically used in the construction of dams, industrial pavements, storage and composting areas, and as a component of composite pavements for highways and streets.

Scaling—disintegration and flaking of a hardened concrete surface, frequently due to repeated freeze-thaw cycles and application of deicing chemicals.

Segregation—separation of the components (aggregates and mortar) of fresh concrete, resulting in a nonuniform mixture.

Self-compacting concrete—concrete of high workability that require little or no vibration or other mechanical means of consolidation.

Set—the degree to which fresh concrete has lost its plasticity and hardened.

Silica fume—very fine noncrystalline silica which is a byproduct from the production of silicon and ferrosilicon alloys in an electric arc furnace; used as a pozzolan in concrete.

Shotcrete—mortar or small-aggregate concrete that is conveyed by compressed air through a hose and applied at high velocity to a surface. Also known as gunite and sprayed concrete.

Shrinkage—decrease in either length or volume of a material resulting from changes in moisture content, temperature, or chemical changes.
Shrinkage-compensating concrete—concrete containing expansive cement, or an admixture, which produces expansion during hardening and thereby offsets the contraction occurring later during drying (drying shrinkage).

Slag cement—hydraulic cement consisting mostly of an intimate and uniform blend of ground, granulated blast-furnace slag with or without portland cement or hydrated lime.

Slump—measure of the consistency of freshly mixed concrete, equal to the immediate subsidence of a specimen molded with a standard slump cone.

Slurry—thin mixture of an insoluble substance, such as portland cement, slag, or clay, with a liquid, such as water.

Soil cement—mixture of soil and measured amounts of portland cement and water compacted to a high density; primarily used as a base material under pavements; also called cement-stabilized soil.

Specific gravity—see Relative density.

Stucco—portland cement plaster and stucco are the same material. The term “stucco” is widely used to describe the cement plaster used for coating exterior surfaces of buildings. However, in some geographical areas, “stucco” refers only to the factory-prepared finish coat mixtures. (See also Portland cement plaster.)

Sulfate attack—most common form of chemical attack on concrete caused by sulfates in the groundwater or soil manifested by expansion and disintegration of the concrete.

Superplasticizer (plasticizer)—admixture that increases the flowability of a fresh concrete mixture.

Supplementary cementitious (cementing) materials—Cementitious material other than portland cement or blended cement. See also Cementitious material.

Tensile strength—stress up to which concrete is able to resist cracking under axial tensile loading.

U


Unit weight—density of fresh concrete or aggregate, normally determined by weighing a known volume of concrete or aggregate (bulk density of aggregates includes voids between particles).

V

Vibration—high-frequency agitation of freshly mixed concrete through mechanical devices, for the purpose of consolidation.

Volume change—Either an increase or a decrease in volume due to any cause, such as moisture changes, temperature changes, or chemical changes. (See also Creep.)
### ASTM Standards

American Society for Testing and Materials (ASTM) documents related to aggregates, cement, and concrete that are relevant to or referred to in the text are listed as follows and can be obtained at [www.astm.org](http://www.astm.org).

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<tr>
<td>C1150-96</td>
<td>Standard Test Method for The Break-Off Number of Concrete (Discontinued 2002)</td>
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<td>C1151-91</td>
<td>Standard Test Method for Evaluating the Effectiveness of Materials for Curing Concrete (Discontinued 2000)</td>
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<td>Standard Test Method for Electrical Indication of Concrete’s Ability to Resist Chloride Ion Penetration</td>
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<td>C1218</td>
<td>Standard Test Method for Water-Soluble Chloride in Mortar and Concrete</td>
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<td>Standard Practice for Use of Unbonded Caps in Determination of Compressive Strength of Hardened Concrete Cylinders</td>
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<td>C1240</td>
<td>Standard Specification for Use of Silica Fume for Use as a Mineral Admixture in Hydraulic-Cement Concrete, Mortar, and Grout</td>
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<td>Standard Test Method for Potential Alkali Reactivity of Aggregates (Mortar-Bar Method)</td>
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<td>Standard Test Method for Thermal Integrity of Flexible Water Vapor Retarders</td>
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<td>C1293</td>
<td>Standard Test Method for Determination of Length Change of Concrete Due to Alkali-Silica Reaction</td>
</tr>
<tr>
<td>C1315</td>
<td>Standard Specification for Liquid Membrane-Forming Compounds Having Special Properties for Curing and Sealing Concrete</td>
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<td>C1328</td>
<td>Standard Specification for Plastic (Stucco) Cement</td>
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<td>C1329</td>
<td>Standard Specification for Mortar Cement</td>
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<td>C1362</td>
<td>Standard Test Method for Flow of Freshly Mixed Hydraulic Cement Concrete</td>
</tr>
<tr>
<td>Code</td>
<td>Description</td>
</tr>
<tr>
<td>------</td>
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<tr>
<td>C1383</td>
<td>Standard Test Method for Measuring the P-Wave Speed and the Thickness of Concrete Plates Using the Impact-Echo Method</td>
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<tr>
<td>C1435</td>
<td>Standard Practice for Molding Roller-Compacted Concrete in Cylinder Molds Using a Vibrating Hammer</td>
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<tr>
<td>C1436</td>
<td>Standard Specification for Materials for Shotcrete</td>
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<td>C1438</td>
<td>Standard Specification for Latex and Powder Polymer Modifiers for Hydraulic Cement Concrete and Mortar</td>
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<td>C1439</td>
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<tr>
<td>C1451</td>
<td>Standard Practice for Determining Uniformity of Ingredients of Concrete From a Single Source</td>
</tr>
<tr>
<td>C1480</td>
<td>Standard Specification for Packaged, Pre-Blended, Dry, Combined Materials for Use in Wet or Dry Shotcrete Application</td>
</tr>
<tr>
<td>C1524</td>
<td>Standard Test Method for Water-Extractable Chloride in Aggregate (Soxhlet Method)</td>
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<tr>
<td>C1542</td>
<td>Standard Test Method for Measuring Length of Concrete Cores</td>
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<tr>
<td>C1543</td>
<td>Standard Test Method for Determining the Penetration of Chloride Ion into Concrete by Ponding</td>
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<tr>
<td>C1556</td>
<td>Standard Test Method for Determining the Apparent Chloride Diffusion Coefficient of Cementitious Mixtures by Bulk Diffusion</td>
</tr>
<tr>
<td>C1580</td>
<td>Standard Test Method for Water-Soluble Sulfate in Soil</td>
</tr>
<tr>
<td>C1581</td>
<td>Standard Test Method for Determining Age at Cracking and Induced Tensile Stress Characteristics of Mortar and Concrete under Restrained Shrinkage</td>
</tr>
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<td>C1582</td>
<td>Standard Specification for Admixtures to Inhibit Chloride-Induced Corrosion of Reinforcing Steel in Concrete</td>
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<tr>
<td>C1585</td>
<td>Standard Test Method for Measurement of Rate of Absorption of Water by Hydraulic-Cement Concretes</td>
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<tr>
<td>C1600</td>
<td>Standard Specification for Rapid Hardening Hydraulic Cement</td>
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<tr>
<td>C1602</td>
<td>Standard Specification for Mixing Water Used in the Production of Hydraulic Cement Concrete</td>
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<td>Standard Test Method for Measurement of Solids in Water</td>
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<td>Standard Test Method for Obtaining and Testing Drilled Cores of Shotcrete</td>
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<tr>
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<td>Standard Test Method for Chemical Shrinkage of Hydraulic Cement Paste</td>
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<td>Standard Test Method for Static Segregation of Self-Consolidating Concrete Using Column Technique</td>
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<tr>
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<td>C1621</td>
<td>Standard Test Method for Passing Ability of Self-Consolidating Concrete by J-Ring</td>
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<td>C1622</td>
<td>Standard Specification for Cold-Weather Admixture Systems</td>
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<tr>
<td>C1646</td>
<td>Standard Practice for Making and Curing Test Specimens for Evaluating Frost Resistance of Coarse Aggregate in Air-Entrained Concrete by Rapid Freezing and Thawing</td>
</tr>
<tr>
<td>D75</td>
<td>Standard Practice for Sampling Aggregates</td>
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<tr>
<td>D98</td>
<td>Standard Specification for Calcium Chloride</td>
</tr>
<tr>
<td>D345</td>
<td>Standard Test Method for Sampling and Testing Calcium Chloride for Roads and Structural Applications</td>
</tr>
<tr>
<td>D448</td>
<td>Standard Classification for Sizes of Aggregate for Road and Bridge Construction</td>
</tr>
<tr>
<td>D512</td>
<td>Standard Test Methods for Chloride Ion in Water</td>
</tr>
<tr>
<td>D516</td>
<td>Standard Test Method for Sulfate Ion in Water</td>
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<tr>
<td>D558</td>
<td>Standard Test Methods for Moisture-Density Relations of Soil-Cement Mixtures</td>
</tr>
<tr>
<td>D632</td>
<td>Standard Specification for Sodium Chloride</td>
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<tr>
<td>D2240</td>
<td>Standard Test Method for Rubber Property-Durometer Hardness</td>
</tr>
<tr>
<td>D3042</td>
<td>Standard Test Method for Insoluble Residue in Carbonate Aggregates</td>
</tr>
<tr>
<td>D3398</td>
<td>Standard Test Method for Index of Aggregate Particle Shape and Texture</td>
</tr>
<tr>
<td>D3963</td>
<td>Standard Specification for Fabrication and Jobsite Handling of Epoxy-Coated Steel Reinforcing Bars</td>
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<tr>
<td>D4263</td>
<td>Standard Test Method for Indicating Moisture in Concrete by the Plastic Sheet Method</td>
</tr>
<tr>
<td>D4580</td>
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</tr>
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<td>D4791</td>
<td>Standard Test Method for Flat Particles, Elongated Particles, or Flat and Elongated Particles in Coarse Aggregate</td>
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<td>E11</td>
<td>Standard Specification for Wire Cloth and Sieves for Testing Purposes</td>
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<tr>
<td>F1869</td>
<td>Standard Test Method for Measuring Moisture Vapor Emission Rate of Concrete Subfloor Using Anhydrous Calcium Chloride</td>
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<tr>
<td>G59</td>
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AASHTO STANDARDS

American Association of State Highway and Transportation Officials (AASHTO) documents related to aggregates, cement, and concrete that are relevant to or referred to in the text are listed as follows and can be obtained at www.aashto.org.

M 6  Fine Aggregate for Portland Cement Concrete
M 43  Sizes of Aggregate for Road and Bridge Construction
M 80  Coarse Aggregate for Portland Cement Concrete
M 85  Portland Cement
M 92  Wire-Cloth Sieves for Testing Purposes
M 143  Sodium Chloride
M 144  Calcium Chloride
M 148  Liquid Membrane-Forming Compounds for Curing Concrete
M 152  Flow Table for Use in Test of Hydraulic Cements
M 154  Air-Entraining Admixtures for Concrete
M 157  Ready-Mixed Concrete
M 171  Sheet Materials for Curing Concrete
M 182  Burlap Cloth Made From Jute or Kenaf
M 194  Chemical Admixtures For Concrete
M 195  Lightweight Aggregate for Structural Concrete
M 200  Epoxy Protective Coatings
M 201  Moist Cabinets, Moist Rooms, and Water Storage Tanks Used in the Testing of Hydraulic Cements and Concretes
M 205  Molds for Forming Concrete Test Cylinders Vertically
M 210  Use of Apparatus for the Determination of Length Change of Hardened Cement Paste, Mortar, and Concrete
M 224  Use of Protective Sealers for Portland Cement Concrete
M 231  Weighing Devices Used in the Testing of Materials
M 233  Boiled Linseed Oil Mixture for Treatment of Portland Cement Concrete
M 235  Epoxy Resin Adhesives
M 240  Blended Hydraulic Cements
M 241  Concrete Made by Volumetric Batching and Continuous Mixing
M 284  Epoxy Coated Reinforcing Bars
M 295  Coal Fly Ash and Raw and Calcined Natural Pozzolan for use as a Mineral Admixture in Concrete
M 302  Ground Granulated Blast-Furnace Slag for Use in Concrete and Mortars
M 307  Microsilica for Use in Concrete and Mortar
M 321  High-Reactivity Pozzolans for Use in Hydraulic-Cement Concrete, Mortar, and Grout
T 2  Sampling of Aggregates
T 11  Materials Finer than 75-µm (No. 200) Sieve in Mineral Aggregates by Washing
T 19  Bulk Density (“Unit Weight”) and Voids in Aggregate
T 21  Organic Impurities in Fine Aggregate for Concrete
T 22  Compressive Strength of Cylindrical Concrete Specimens
T 23  Making and Curing Concrete Test Specimens
T 24  Obtaining and Testing Drilled Cores and Sawed Beams of Concrete
T 26  Quality of Water to be Used in Concrete
T 27  Sieve Analysis of Fine and Coarse Aggregate
T 71  Effect of Organic Impurities in Fine Aggregate on Strength of Mortar
T 84  Specific Gravity and Absorption of Fine Aggregate
T 85  Specific Gravity and Absorption of Coarse Aggregate
T 96  Resistance to Degradation of Small-Size Coarse Aggregate by Abrasion and Impact in the Los Angeles Machine
T 97  Flexural Strength of Concrete (Using Simple Beam with Third Point Loading)
T 98  Fineness of Portland Cement by the Turbidimeter
T 103  Soundness of Aggregates by Freezing and Thawing
T 104  Soundness of Aggregate by Use of Sodium Sulfate or Magnesium Sulfate
T 105  Chemical Analysis of Hydraulic Cement
T 106  Compressive Strength of Hydraulic Cement Mortar (Using 50-mm or 2-in. Cube Specimens)
T 107  Autoclave Expansion of Portland Cement
T 112  Clay Lumps and Friable Particles in Aggregate
T 113  Lightweight Pieces in Aggregate
T 119  Slump of Hydraulic Cement Concrete
T 121  Mass per Cubic Meter (Cubic Foot), Yield, and Air Content (Gravimetric) of Concrete
T 126  Making and Curing Concrete Test Specimens in the Laboratory
T 127  Sampling and Amount of Testing by Hydraulic Cement
T 128  Fineness of Hydraulic Cement by the 150-µm (No. 100) and 75-µm (No. 200) Sieves (discontinued)
T 129  Normal Consistency of Hydraulic Cement
T 131  Time of Setting of Hydraulic Cement by Vicat Needle
T 133  Density of Hydraulic Cement
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<td>TP 60</td>
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# Design and Control of Concrete Mixtures

## METRIC CONVERSION FACTORS

The following list provides the conversion relationship between U.S. customary units and SI (International System) units. The proper conversion procedure is to multiply the specified value on the left (primarily U.S. customary values) by the conversion factor exactly as given below and then round to the appropriate number of significant digits desired. For example, to convert 11.4 ft to meters: 11.4 x 0.3048 = 3.47472, which rounds to 3.47 meters. Do not round either value before performing the multiplication, as accuracy would be reduced. A complete guide to the SI system and its use can be found in IEEE/ASTM SI-10, Metric Practice.

### To convert from to multiply by

#### Length

<table>
<thead>
<tr>
<th>Unit (symbol)</th>
<th>Conversion Factor</th>
<th>Note</th>
</tr>
</thead>
<tbody>
<tr>
<td>inch (in.)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>micrometer (µm)</td>
<td>25,400 E*</td>
<td></td>
</tr>
<tr>
<td>millimeter (mm)</td>
<td>25.4 E</td>
<td></td>
</tr>
<tr>
<td>meter (m)</td>
<td>0.0254 E</td>
<td></td>
</tr>
<tr>
<td>foot (ft)</td>
<td>0.3048 E</td>
<td></td>
</tr>
<tr>
<td>yard (yd)</td>
<td>0.9144</td>
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</tr>
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#### Area

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<tr>
<th>Unit (symbol)</th>
<th>Conversion Factor</th>
<th>Note</th>
</tr>
</thead>
<tbody>
<tr>
<td>square foot (ft²)</td>
<td>0.09290304E</td>
<td></td>
</tr>
<tr>
<td>square inch (in.²)</td>
<td>0.00694444E</td>
<td></td>
</tr>
<tr>
<td>square yard (yd²)</td>
<td>0.8361274</td>
<td></td>
</tr>
<tr>
<td>square meter (m²)</td>
<td></td>
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</tbody>
</table>

#### Volume

<table>
<thead>
<tr>
<th>Unit (symbol)</th>
<th>Conversion Factor</th>
<th>Note</th>
</tr>
</thead>
<tbody>
<tr>
<td>cubic inch (in.³)</td>
<td>16.387064</td>
<td></td>
</tr>
<tr>
<td>cubic foot (ft³)</td>
<td>0.02831685</td>
<td></td>
</tr>
<tr>
<td>cubic yard (yd³)</td>
<td>0.7645549</td>
<td></td>
</tr>
<tr>
<td>gallon (gal)</td>
<td>3.7854118</td>
<td></td>
</tr>
<tr>
<td>fluid ounce (fl oz)</td>
<td>29.57353</td>
<td></td>
</tr>
<tr>
<td>cubic meter (m³)</td>
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<td></td>
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</tbody>
</table>

#### Force

<table>
<thead>
<tr>
<th>Unit (symbol)</th>
<th>Conversion Factor</th>
<th>Note</th>
</tr>
</thead>
<tbody>
<tr>
<td>kip (1000 lb)</td>
<td>453.6</td>
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</tr>
<tr>
<td>pound (lb)</td>
<td>0.4535924</td>
<td></td>
</tr>
<tr>
<td>pound (lb)</td>
<td>4,448.222</td>
<td></td>
</tr>
<tr>
<td>newton (N)</td>
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</table>

#### Pressure or stress

<table>
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<tr>
<th>Unit (symbol)</th>
<th>Conversion Factor</th>
<th>Note</th>
</tr>
</thead>
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<tr>
<td>pound per square foot (psf)</td>
<td>0.000020482</td>
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</tr>
<tr>
<td>pound per square foot (psf)</td>
<td>47.88</td>
<td></td>
</tr>
<tr>
<td>pound per square inch (psi)</td>
<td>0.00694444E</td>
<td></td>
</tr>
<tr>
<td>pound per square inch (psi)</td>
<td>6,894.757</td>
<td></td>
</tr>
<tr>
<td>pound per square inch (psi)</td>
<td>0.000020482</td>
<td></td>
</tr>
<tr>
<td>kilogram per square meter (kg/m²)</td>
<td>4.8824</td>
<td></td>
</tr>
<tr>
<td>kilogram per square meter (kg/m²)</td>
<td>0.000020482</td>
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<tr>
<td>pascal (Pa)</td>
<td>1.4503773</td>
<td></td>
</tr>
<tr>
<td>pascal (Pa)</td>
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<tr>
<td>pascal (Pa)</td>
<td>4,133.752</td>
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<tr>
<td>pascal (Pa)</td>
<td>47.88</td>
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</tr>
<tr>
<td>pascal (Pa)</td>
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</tr>
<tr>
<td>megapascal (MPa)</td>
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</tr>
<tr>
<td>megapascal (MPa)</td>
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#### Mass (weight)

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<tr>
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<th>Conversion Factor</th>
<th>Note</th>
</tr>
</thead>
<tbody>
<tr>
<td>pound (lb)</td>
<td>0.4535924</td>
<td></td>
</tr>
<tr>
<td>ton, 2000 lb</td>
<td>907.1848</td>
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<tr>
<td>kilogram (kg)</td>
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### To convert from to multiply by

#### Mass (weight) per length

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<th>Conversion Factor</th>
<th>Note</th>
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<tr>
<td>kip per linear foot (klf)</td>
<td>0.001488</td>
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<tr>
<td>pound per linear foot (plf)</td>
<td>1.488</td>
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</table>

#### Mass per volume (density)

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<th>Conversion Factor</th>
<th>Note</th>
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</thead>
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<tr>
<td>pound per cubic foot (lb/ft³)</td>
<td>0.07031</td>
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<tr>
<td>pound per cubic yard (lb/yd³)</td>
<td>0.5933</td>
<td></td>
</tr>
<tr>
<td>kilogram per cubic meter (kg/m³)</td>
<td>16.01846</td>
<td></td>
</tr>
<tr>
<td>kilogram per cubic meter (kg/m³)</td>
<td>0.07031</td>
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</tr>
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### Temperature

<table>
<thead>
<tr>
<th>Unit (symbol)</th>
<th>Conversion Factor</th>
<th>Note</th>
</tr>
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<tbody>
<tr>
<td>degree Fahrenheit (°F)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>degree Celsius (°C)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>degree Kelvin (°K)</td>
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</table>

#### Energy and heat

<table>
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<tr>
<th>Unit (symbol)</th>
<th>Conversion Factor</th>
<th>Note</th>
</tr>
</thead>
<tbody>
<tr>
<td>British thermal unit (Btu)</td>
<td>1055.056</td>
<td></td>
</tr>
<tr>
<td>calorie (cal)</td>
<td>4.1868 E</td>
<td></td>
</tr>
<tr>
<td>kilowatt-hour (kwh)</td>
<td>3,600,000 E</td>
<td></td>
</tr>
<tr>
<td>British thermal unit per pound (Btu/lb)</td>
<td>0.5556</td>
<td></td>
</tr>
<tr>
<td>watt (W)</td>
<td>0.2930711</td>
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</tr>
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</table>

### Permeability

<table>
<thead>
<tr>
<th>Unit (symbol)</th>
<th>Conversion Factor</th>
<th>Note</th>
</tr>
</thead>
<tbody>
<tr>
<td>darcy</td>
<td>2.4909631</td>
<td></td>
</tr>
<tr>
<td>feet per day (ft/day)</td>
<td>0.000352</td>
<td></td>
</tr>
</tbody>
</table>

* E indicates that the factor given is exact.
** One U.S. gallon equals 0.8327 Canadian gallon.
† A pascal equals 1.000 newton per square meter.

Note:
One U.S. gallon of water weighs 8.34 pounds (U.S.) at 60°F.
One cubic foot of water weighs 62.4 pounds (U.S.).
One milliliter of water has a mass of 1 gram and has a volume of one cubic centimeter.
One U.S. bag of cement weighs 94 lb.

The prefixes and symbols listed below are commonly used to form names and symbols of the decimal multiples and submultiples of the SI units.

### Multiplication Factor

<table>
<thead>
<tr>
<th>Unit (symbol)</th>
<th>Prefix</th>
<th>Symbol</th>
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<tbody>
<tr>
<td>1,000,000,000 = 10⁹</td>
<td>giga</td>
<td>G</td>
</tr>
<tr>
<td>1,000,000 = 10⁶</td>
<td>mega</td>
<td>M</td>
</tr>
<tr>
<td>1,000 = 10³</td>
<td>kilo</td>
<td>k</td>
</tr>
<tr>
<td>1 = 1</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td>0.01 = 10⁻²</td>
<td>centi</td>
<td>c</td>
</tr>
<tr>
<td>0.001 = 10⁻³</td>
<td>milli</td>
<td>m</td>
</tr>
<tr>
<td>0.000001 = 10⁻⁶</td>
<td>micro</td>
<td>µ</td>
</tr>
<tr>
<td>0.000000001 = 10⁻⁹</td>
<td>nano</td>
<td>n</td>
</tr>
</tbody>
</table>
## CEMENT AND CONCRETE RESOURCES

<table>
<thead>
<tr>
<th>Acronym</th>
<th>Description</th>
<th>Website</th>
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</thead>
<tbody>
<tr>
<td>AASHTO</td>
<td>American Association of State Highway and Transportation Officials</td>
<td><a href="http://www.transportation.org">http://www.transportation.org</a></td>
</tr>
<tr>
<td>ACA</td>
<td>American Association of State Highway and Transportation Officials</td>
<td><a href="http://www.aashto.org">http://www.aashto.org</a></td>
</tr>
<tr>
<td>ACA</td>
<td>American Concrete Association</td>
<td><a href="http://www.aca-usa.org">http://www.aca-usa.org</a></td>
</tr>
<tr>
<td>ACBM</td>
<td>Center for Advanced Cement Based Materials (Northwestern University)</td>
<td><a href="http://www.civil.northwestern.edu/ACBM">http://www.civil.northwestern.edu/ACBM</a></td>
</tr>
<tr>
<td>ACI</td>
<td>American Concrete Institute</td>
<td><a href="http://www.aci-int.org">http://www.aci-int.org</a></td>
</tr>
<tr>
<td>ACPA</td>
<td>American Concrete Pavement Association</td>
<td><a href="http://www.pavement.com">http://www.pavement.com</a></td>
</tr>
<tr>
<td>ACPA</td>
<td>American Concrete Pipe Association</td>
<td><a href="http://www.concrete-pipe.org">http://www.concrete-pipe.org</a></td>
</tr>
<tr>
<td>ACPA</td>
<td>American Concrete Pumping Association</td>
<td><a href="http://www.concretepumpers.com/">http://www.concretepumpers.com/</a></td>
</tr>
<tr>
<td>ACPPA</td>
<td>American Concrete Pressure Pipe Association</td>
<td><a href="http://www.acppa.org">http://www.acppa.org</a></td>
</tr>
<tr>
<td>ACS</td>
<td>American Ceramic Society</td>
<td><a href="http://www.acers.org">http://www.acers.org</a></td>
</tr>
<tr>
<td>AIT</td>
<td>International Ferrocement Society</td>
<td><a href="http://www.ferrocement.org">http://www.ferrocement.org</a></td>
</tr>
<tr>
<td>APA</td>
<td>Architectural Precast Association</td>
<td><a href="http://www.archprecast.org">http://www.archprecast.org</a></td>
</tr>
<tr>
<td>API</td>
<td>American Petroleum Institute</td>
<td><a href="http://www.api.org">http://www.api.org</a></td>
</tr>
<tr>
<td>ANSI</td>
<td>American National Standards Institute</td>
<td><a href="http://www.ansi.org">http://www.ansi.org</a></td>
</tr>
<tr>
<td>ASCC</td>
<td>American Society of Concrete Contractors</td>
<td><a href="http://www.ascconline.org">http://www.ascconline.org</a></td>
</tr>
<tr>
<td>ASCE</td>
<td>American Society of Civil Engineers</td>
<td><a href="http://www.asce.org">http://www.asce.org</a></td>
</tr>
<tr>
<td>ASI</td>
<td>American Shotcrete Association</td>
<td><a href="http://www.shotcrete.org">http://www.shotcrete.org</a></td>
</tr>
<tr>
<td>BCA</td>
<td>British Cement Association</td>
<td><a href="http://www.cementindustry.co.uk">http://www.cementindustry.co.uk</a></td>
</tr>
<tr>
<td>BDZ</td>
<td>Bundesverband der Deutschen Zementindustrie e. V.</td>
<td><a href="http://www.bdzentrum.de">http://www.bdzentrum.de</a></td>
</tr>
<tr>
<td>BFRL</td>
<td>Building and Fire Research Laboratory (NIST)</td>
<td><a href="http://www.bfrl.nist.gov">http://www.bfrl.nist.gov</a></td>
</tr>
<tr>
<td>BRE</td>
<td>Building Research Establishment (UK)</td>
<td><a href="http://www.bre.co.uk">http://www.bre.co.uk</a></td>
</tr>
<tr>
<td>BSI</td>
<td>British Standards Institution</td>
<td><a href="http://www.bsi-global.com">http://www.bsi-global.com</a></td>
</tr>
<tr>
<td>CAC</td>
<td>Cement Association of Canada</td>
<td><a href="http://www.cement.ca">http://www.cement.ca</a></td>
</tr>
<tr>
<td>CC</td>
<td>Cement Careers</td>
<td><a href="http://www.cementcareers.org">http://www.cementcareers.org</a></td>
</tr>
<tr>
<td>CCAA</td>
<td>Cement and Concrete Association of Australia</td>
<td><a href="http://www.concrete.net.au">http://www.concrete.net.au</a></td>
</tr>
<tr>
<td>CCRI</td>
<td>Concrete Corrosion Inhibitors Foundation</td>
<td><a href="http://www.corrosioninhibitors.org/">http://www.corrosioninhibitors.org/</a></td>
</tr>
<tr>
<td>CEMBUREAU</td>
<td>European Cement (Industry) Association</td>
<td><a href="http://www.cembureau.be">http://www.cembureau.be</a></td>
</tr>
<tr>
<td>CEN</td>
<td>European Committee for Standardization</td>
<td><a href="http://www.cen.eu">http://www.cen.eu</a></td>
</tr>
<tr>
<td>CERF</td>
<td>Civil Engineering Research Foundation</td>
<td><a href="http://www.cerf.org">http://www.cerf.org</a></td>
</tr>
<tr>
<td>CFA</td>
<td>Concrete Foundations Association</td>
<td><a href="http://www.cfaana.org">http://www.cfaana.org</a></td>
</tr>
<tr>
<td>CH</td>
<td>Concrete Homes</td>
<td><a href="http://www.concretehomes.com">http://www.concretehomes.com</a></td>
</tr>
<tr>
<td>CP Tech</td>
<td>National Concrete Pavement Technology Center</td>
<td><a href="http://www.cpcenter.org">http://www.cpcenter.org</a></td>
</tr>
<tr>
<td>CSI</td>
<td>Construction Specifications Institute</td>
<td><a href="http://www.csinet.org">http://www.csinet.org</a></td>
</tr>
<tr>
<td>CSI</td>
<td>Cast Stone Institute</td>
<td><a href="http://www.caststone.org">http://www.caststone.org</a></td>
</tr>
<tr>
<td>DIN</td>
<td>Deutsches Institut für Normung e.V. (German Standards Institution)</td>
<td><a href="http://www.din.de">http://www.din.de</a></td>
</tr>
<tr>
<td>ECCO</td>
<td>Environmental Council of Concrete Organizations</td>
<td><a href="http://www.ecco.org">http://www.ecco.org</a></td>
</tr>
<tr>
<td>ERMCO</td>
<td>European Ready Mixed Concrete Association</td>
<td><a href="http://www.ermco.eu">http://www.ermco.eu</a></td>
</tr>
<tr>
<td>ESCSI</td>
<td>Expanded Shale, Clay and Slate Institute</td>
<td><a href="http://www.escsi.org">http://www.escsi.org</a></td>
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An organization of cement companies to improve and extend the uses of portland cement and concrete through market development, engineering, research, education, and public affairs work.