

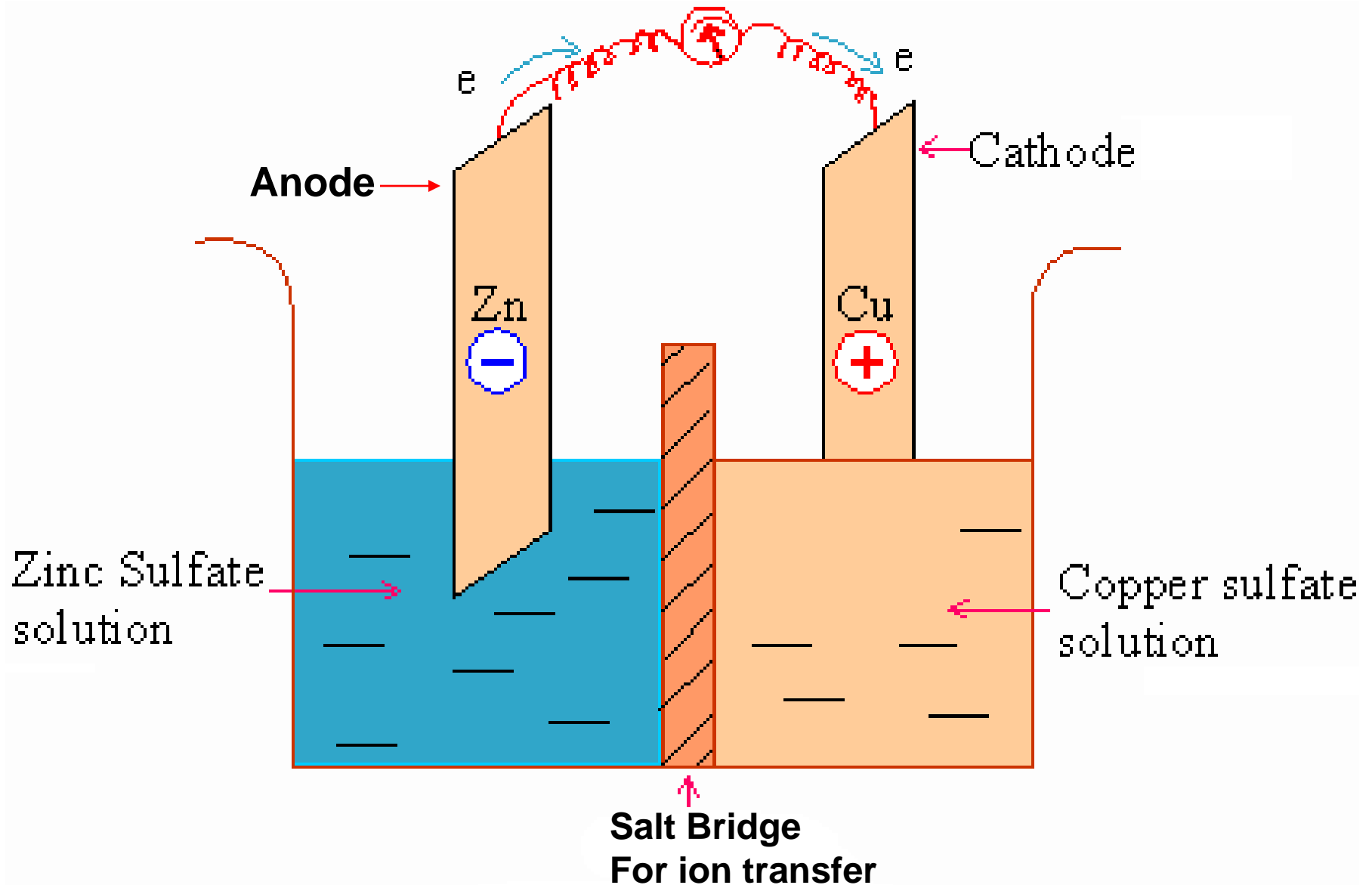
CHEM.5140 Advanced Analytical Chemistry

Electroanalytical Methods

Two general categories:

- 1) Potentiometric Systems – measure voltage (i.e., potential) of a galvanic cell (produces electricity spontaneously)
- 2) Voltammetric Systems – control potential & usually measure current in an electrolytic cell (consumes power to cause an electrochemical reaction to occur)

Electrochemical Cell



Potentiometry

- Determine concentrations by measuring the potential (i.e., voltage) of an electrochemical cell (**galvanic cell**)
- Two electrodes are required (2 half cells)
 - 1) Indicator Electrode – potential responds to activity of species of interest
 - 2) Reference Electrode – chosen so that its potential is independent of solution composition.

$$E_{\text{cell}} = E_{\text{ind}} - E_{\text{ref}} (+ E_{\text{J}})$$

Nernst Equation

$$E = E^{\circ} - \frac{RT}{nF} \ln \frac{[\text{Red}]}{[\text{Ox}]}$$

Where

- R = gas constant
- T = absolute temperature
- n = number of electrons in reaction
- F = Faraday's constant
- E = potential
- E° = standard potential
- [Red] = molar concentration of reduced form of species
- [Ox] = molar concentration of oxidized form of species

Reference Electrodes

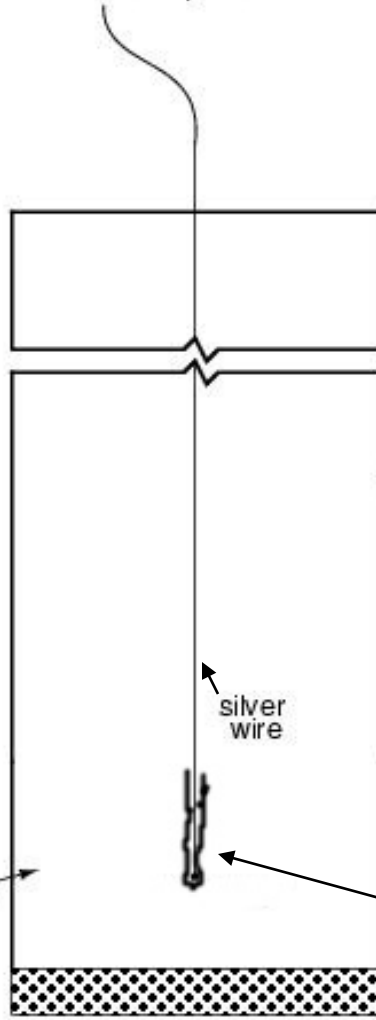
- Normal Hydrogen Electrode (NHE)
- $2\text{H}^+ + 2\text{e}^- \leftrightarrow \text{H}_2 \quad E^\circ = 0.000 \text{ v}$

- Saturated Calomel Electrode (SCE)
- $\text{Hg}_2\text{Cl}_2 + 2 \text{e}^- \leftrightarrow 2 \text{Hg} + 2\text{Cl}^- \quad E^\circ = 0.268 \text{ v}$

- Silver/Silver Chloride Electrode (AgCl)
- $\text{AgCl} + \text{e}^- \leftrightarrow \text{Ag} + \text{Cl}^- \quad E^\circ = 0.222 \text{ v}$

Silver/Silver Chloride Reference Electrode

wire connection point



- Reaction is $\text{AgCl} + \text{e}^- \leftrightarrow \text{Ag} + \text{Cl}^-$
- $E^\circ = 0.222 \text{ v}$
- $E = 0.228$ for 1 M KCl
- Easily made in the lab
- Reliable
- Non-toxic vs Calomel

filled with potassium chloride solution

silver wire

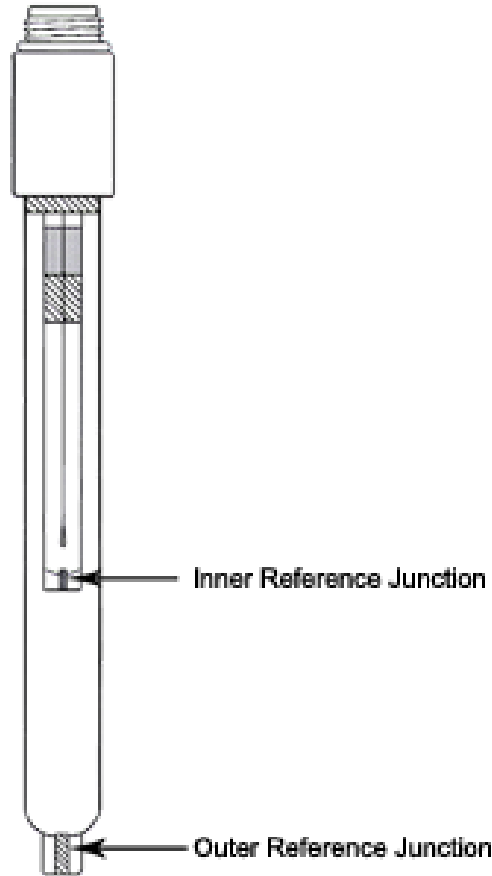
glass or plastic body

silver chloride coating on silver wire

porous junction

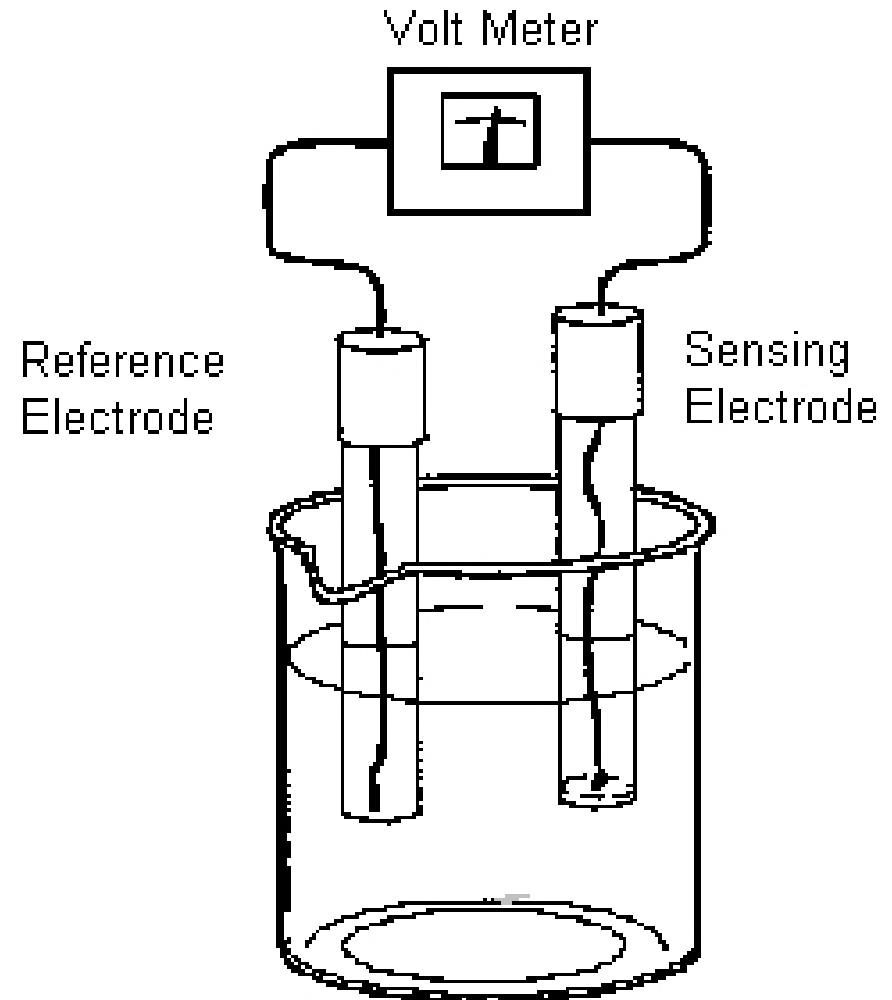
Double Junction Reference

Double Junction Reference Electrode



Indicator Electrodes

- Potential “**indicates**” **activity** of species
- Terms Working Electrode or Sensing Electrode are sometimes used
- Coupled to reference and meter as usual



Indicator Electrodes

Electrode of the First Kind

– Metallic Indicator Electrodes

1) Active metals (e.g., Ag, Cu, Hg, Pb, Cd)
can serve as indicators for their own ions



Nernst Equation

$$E = E^{\circ} - \frac{RT}{nF} \ln \frac{[\text{Red}]}{[\text{Ox}]}$$

At 25 °C this becomes

$$E = E^{\circ} - \frac{0.0591}{n} \log \frac{[\text{Red}]}{[\text{Ox}]}$$

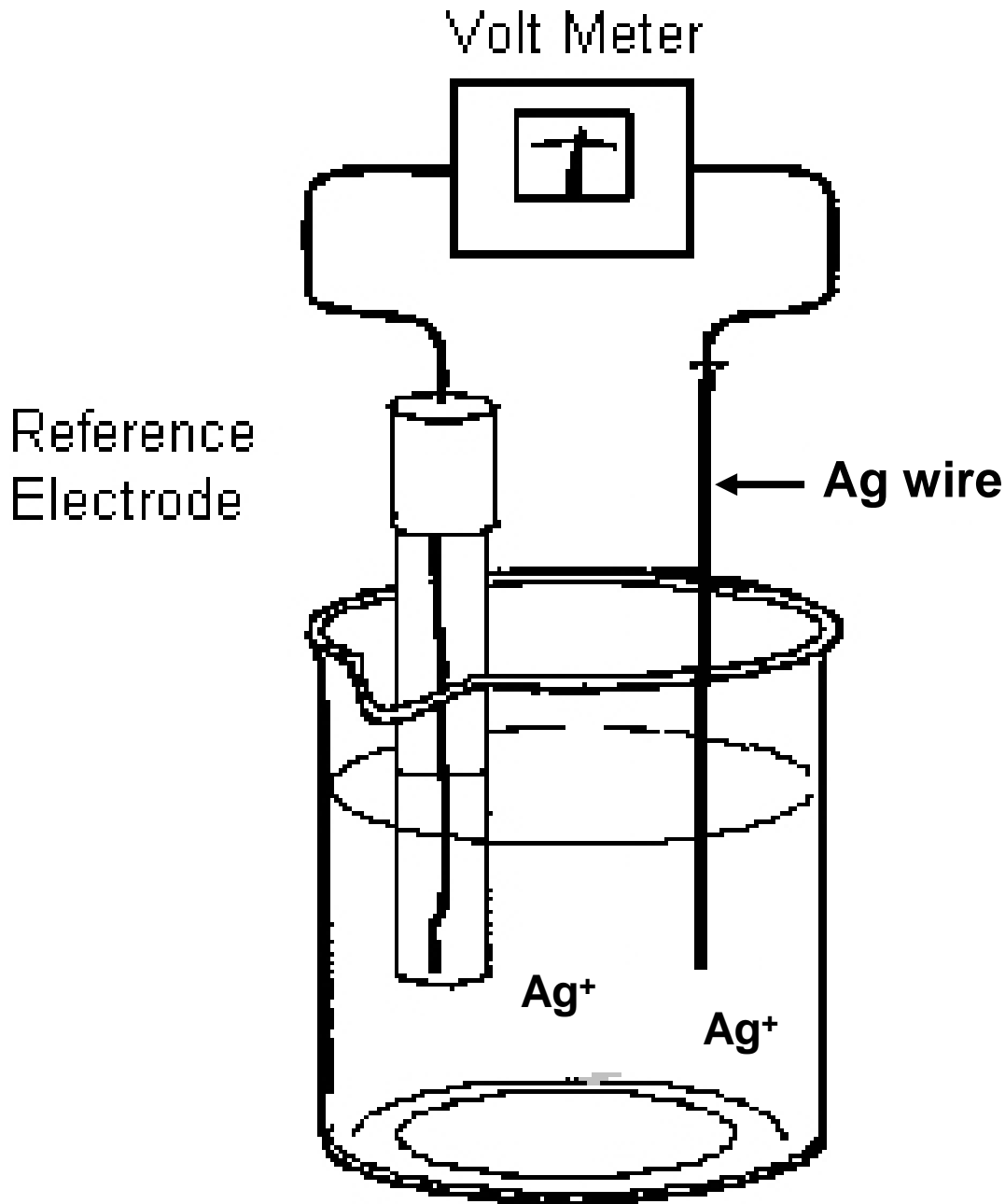
For the Silver/Silver ion system



this becomes

$$E = 0.799 - \frac{0.0591}{1} \log \frac{1}{[\text{Ag}^+]}$$

Electrode potential is inversely proportional to the concentration (activity) of silver ions, even for a simple electrode made only of **Ag metal**



**Silver metallic
Electrode to
Measure
 Ag^+ ions**



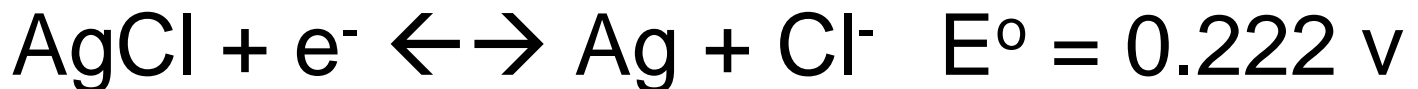
Indicator Electrodes

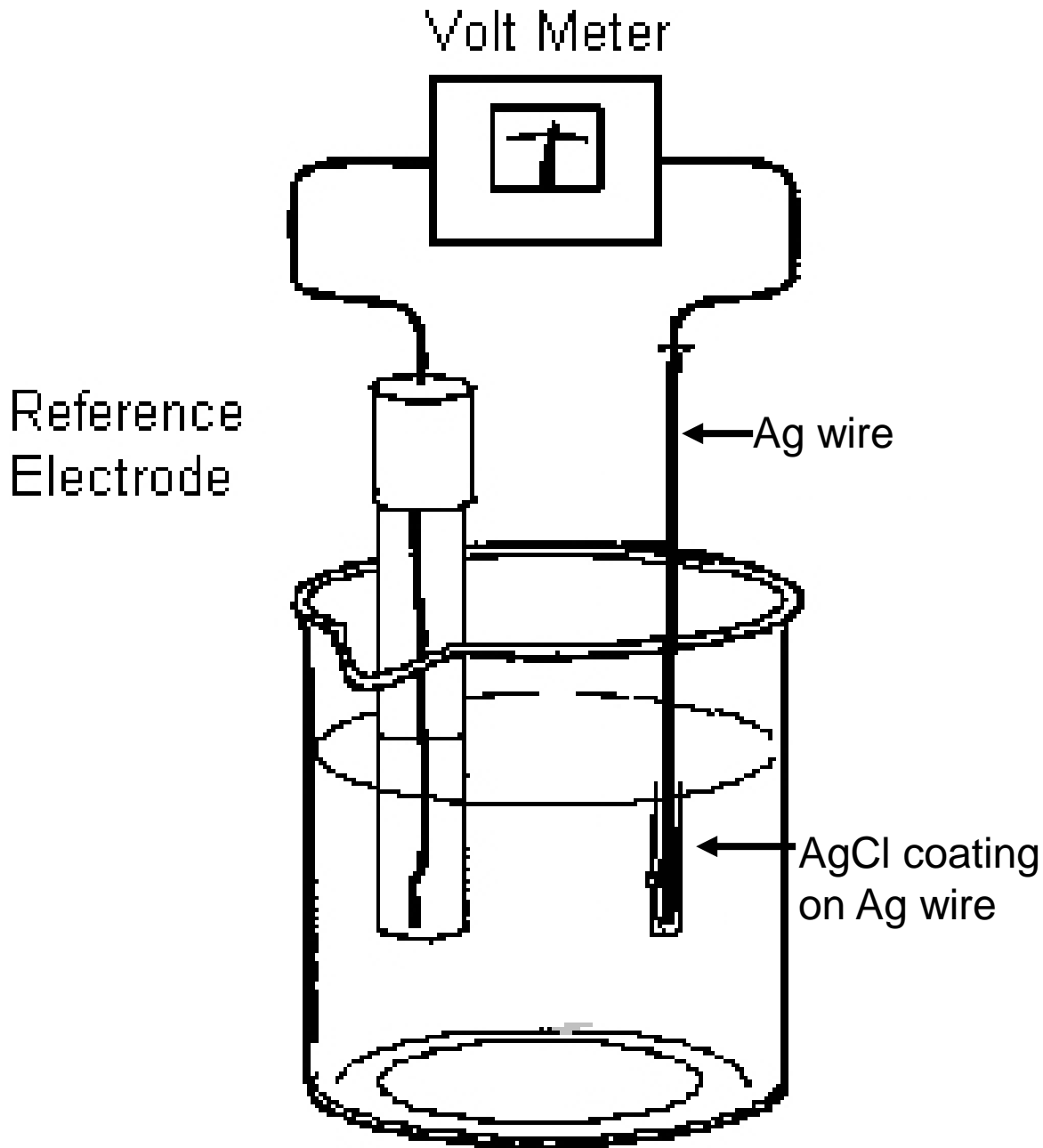
– Metallic Indicator Electrodes

2) Active metal in contact with slightly soluble precipitate involving the metal cation
–responds to anion concentration

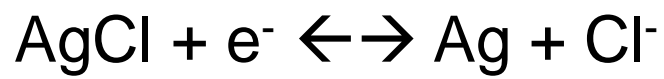
This is an **Electrode of the Second Kind**

e.g., Silver/Silver Chloride Electrode (AgCl)





Silver/Silver Chloride Electrode to measure Cl⁻ ions



For the Silver/Silver Chloride system



The Nernst equation becomes

$$E = 0.222 - \frac{0.0591}{1} \log \frac{[\text{Cl}^-]}{1}$$

Electrode potential is **directly** proportional to the concentration (activity) of the chloride ions for this Electrode of the Second Kind

Indicator Electrodes

– Metallic Indicator Electrodes

3) Inert Electrodes e.g., Pt or Au (noble metal)

Don't participate in the reaction, but serve only as a source or sink for electrons.

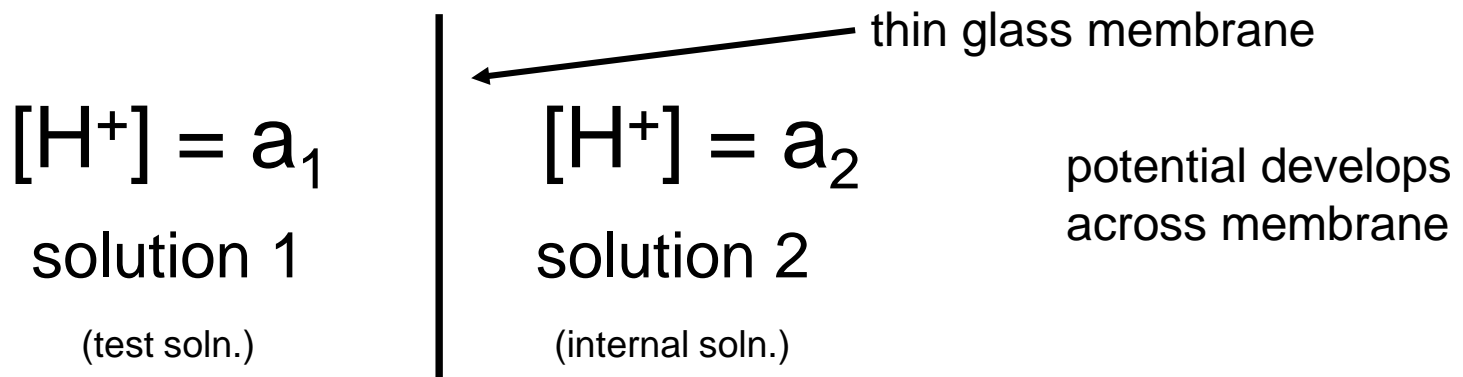
Respond to the ratio of [Ox] to [Red] when both oxidized and reduced species are soluble e.g., could be used to follow a redox titration like Fe^{2+} with MnO_4^-

Summary of Metallic Electrodes

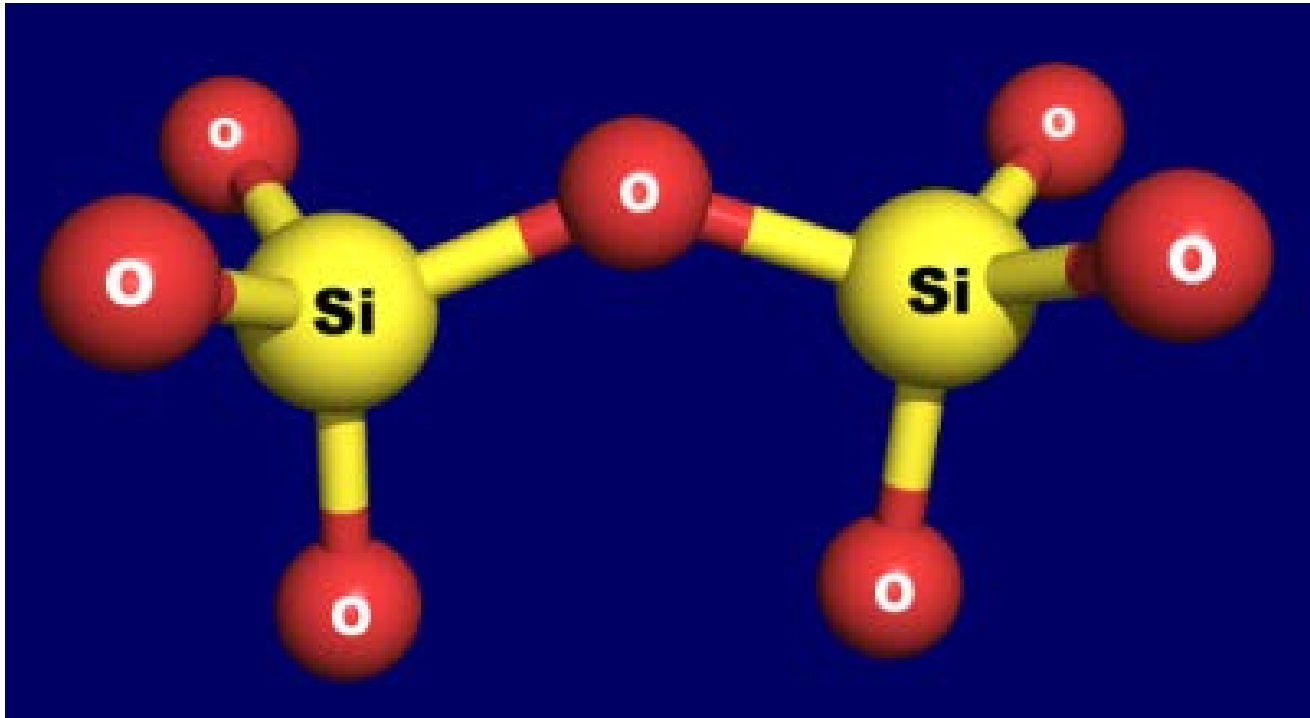
- Important historically for development of potentiometric sensors
- Not very practical in modern electrochem because of poor detection limits
- Inert electrodes are used in redox titrations
- Better systems now available for Cl^- , etc.
- Electrode of the Third Kind can be made that senses an ion that reacts with Cl^- and changes the activity of Cl^- detected

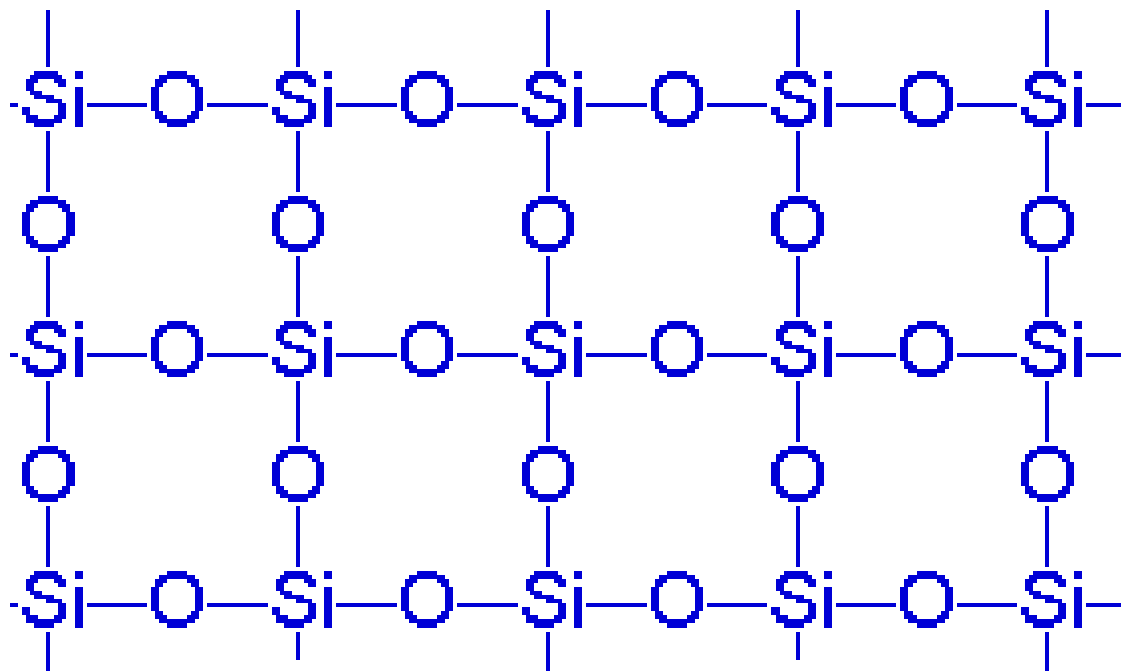
Membrane Electrodes

- Several types – Glass membrane electrode
 - Solid State “ “
 - Liquid Junction “ “
 - Permeable “ “
- Most important is glass electrode for pH



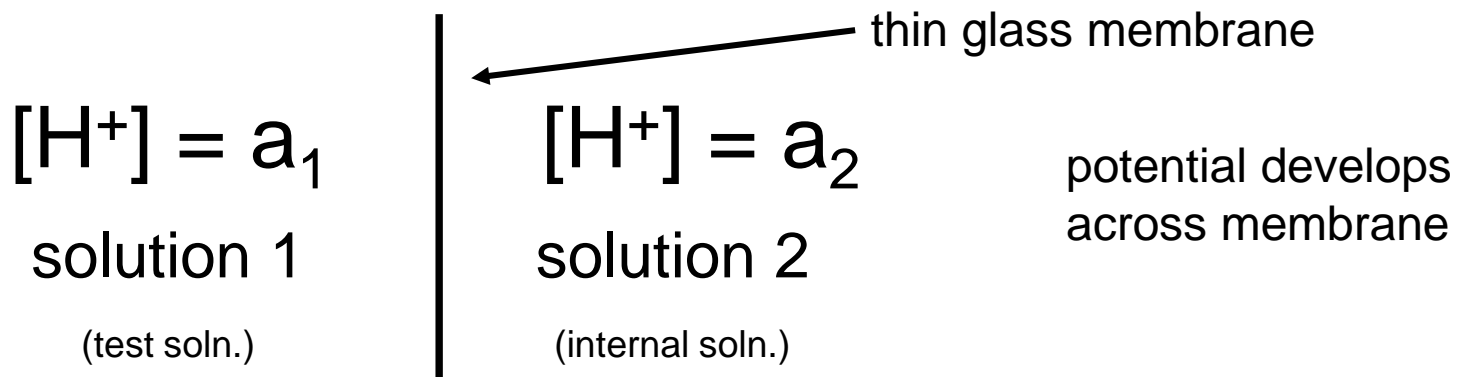
Glass is SiO_2





Membrane Electrodes

- Several types – Glass membrane electrode
 - Solid State “ “
 - Liquid Junction “ “
 - Permeable “ “
- Most important is glass electrode for pH



Glass Membrane Electrode for pH

$$E = K + 0.0591 \log \frac{a_1}{a_2}$$

where K is constant at fixed temp

In Glass Electrode, a_2 is fixed at 0.1 M HCl

$$E = K' + 0.0591 \log a_1$$

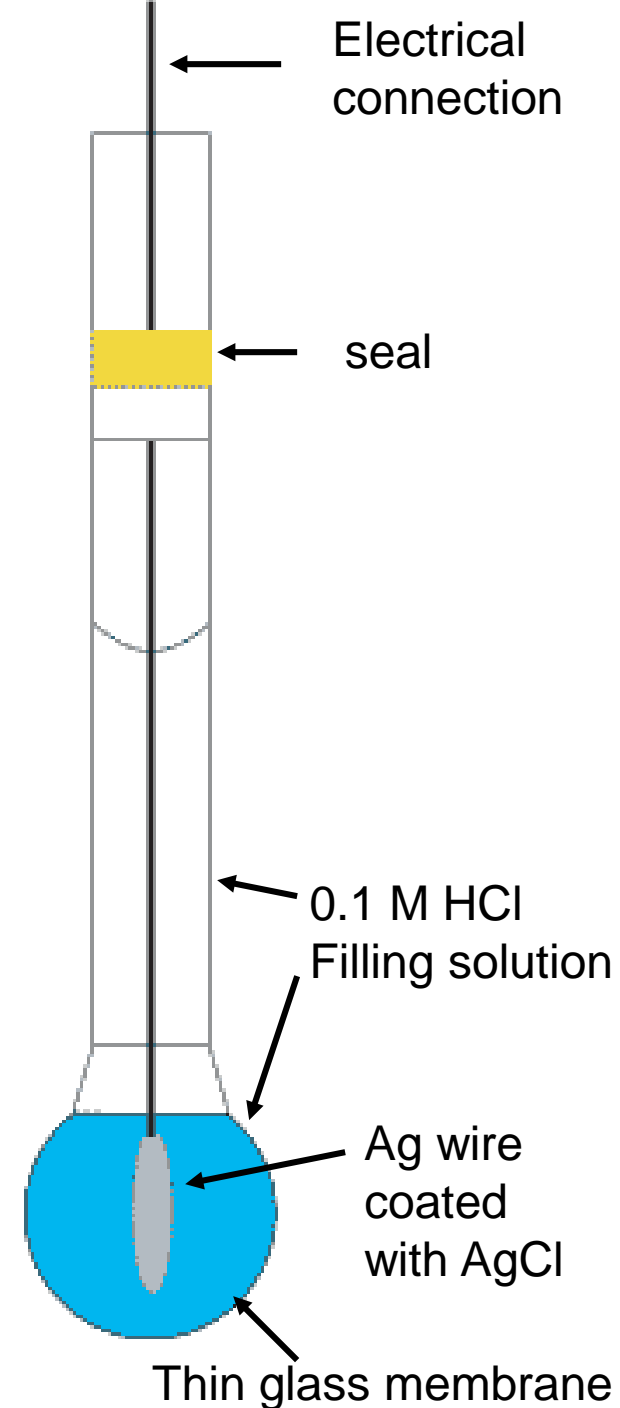
K' new constant that includes a_2

$$E = K' - 0.0591 \text{ pH}$$

since $\text{pH} = -\log a_{\text{H}^+}$

Glass pH Electrode

- $E = K' - 0.0591 \text{ pH}$
- Combine with reference electrode and meter
- Half cell voltage proportional to pH
- Nernstian slope
- Intercept is K' , no E°
- Calibrate with buffers



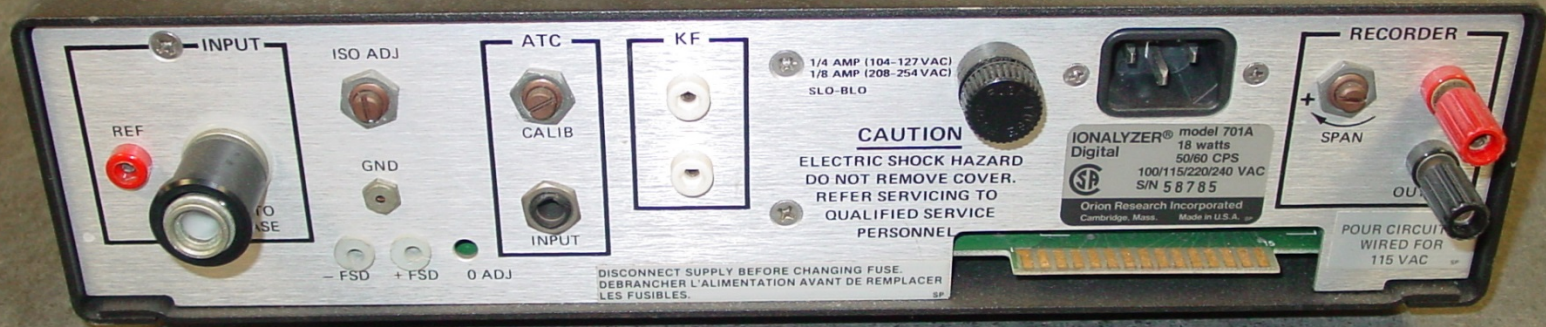
6.888

REL MV MV STD BY pH/.01 pH/.001

CALIB

TEMP °C 0 20 40 60 80 100 % SLOPE

ORION RESEARCH model 701A / digital IONALYZER



INPUT

REF

ISO ADJ

GND

- FSD

+ FSD

0 ADJ

ATC

CALIB

INPUT

KF

1/4 AMP (104-127 VAC)
1/8 AMP (208-254 VAC)
SLO-BLO

CAUTION
ELECTRIC SHOCK HAZARD
DO NOT REMOVE COVER.
REFER SERVICING TO
QUALIFIED SERVICE
PERSONNEL

IONALYZER® model 701A
Digital
18 watts
50/60 CPS
100/115/220/240 VAC
S/N 58785
Orion Research Incorporated
Cambridge, Mass. Made in U.S.A.

RECORDER

+

SPAN

OUT

POUR CIRCUIT
WIRED FOR
115 VAC

DISCONNECT SUPPLY BEFORE CHANGING FUSE
DEBRANCHER L'ALIMENTATION AVANT DE REMPLACER
LES FUSIBLES.

inoLab

pH Level 1



wtw



M



CAL



AR



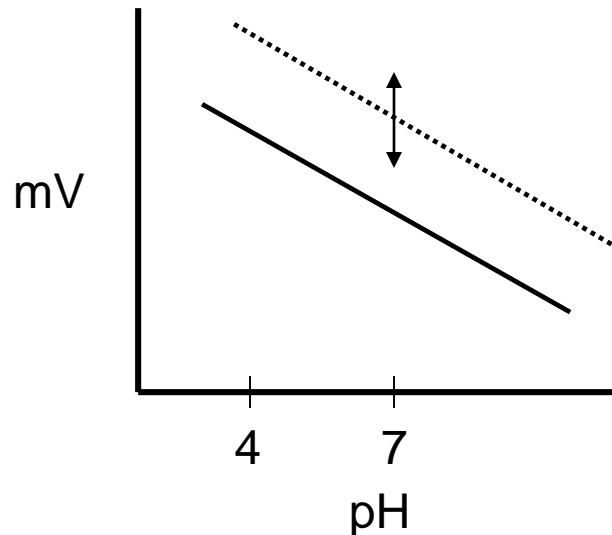
RUN
ENTER





Proper pH Calibration

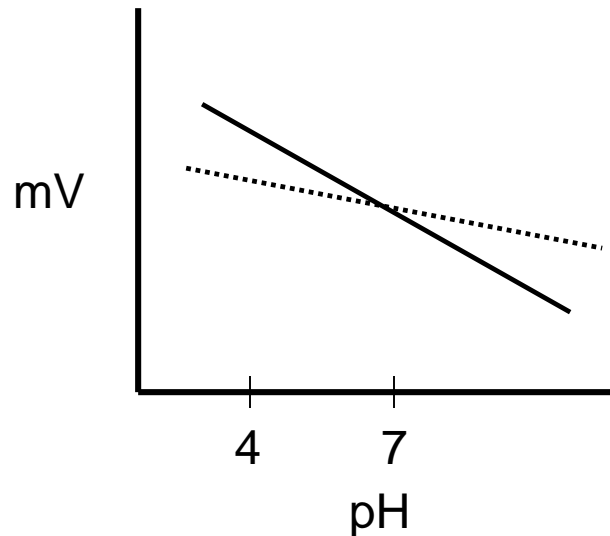
- $E = K' - 0.0591 \text{ pH}$
- Meter measures E vs pH – must calibrate both slope & intercept on meter with buffers
- Meter has two controls – calibrate & slope
- 1st use $\text{pH } 7.00$ buffer to adjust calibrate knob



Calibrate knob raises and lowers the line without changing slope

Proper pH Calibration (cont.)

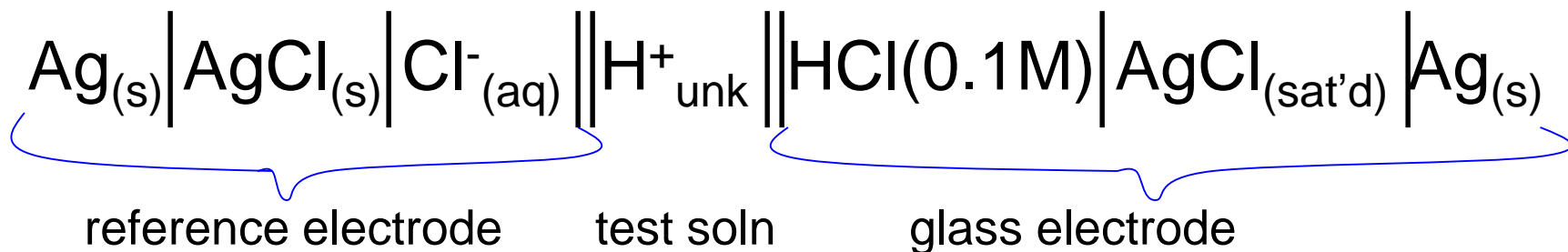
- 2nd step is to use any other pH buffer
- Adjust slope/temp control to correct pH value
- This will pivot the calibration line around the isopotential which is set to 7.00 in all meters



Slope/temp control pivots line around isopotential without changing it

- Slope comes from RT/nF in Nernst Equation
- Slope is temperature sensitive
- Other factors influence slope including
 - Impurities in glass membrane
 - Overall quality of electrode construction
- Many electrodes exhibit “full Nernstian response” while others may give only 90%

Cell for pH measurement (shorthand notation)



Errors in pH Measurement 1

- pH measurements are only as good as the buffers used to calibrate
 - Accuracy good to ± 0.01 units*
 - Precision may be good to ± 0.001 units
 - Junction potential dependent on ionic strength of solution – E_j may be a significant error if test solution has different ionic strength than buffers
- * Unless using special buffers, temp. control & a Faraday cage

Errors in pH Measurement 2

- Asymmetry potential is another non-ideal potential that arises possibly from strain in the glass. When both internal & external H^+ solutions are the same activity, potential should be 0 but it's not

$$E_{\text{cell}} = E_{\text{ind}} - E_{\text{ref}} + E_j + E_a$$

- Temperature of electrodes, calibration buffers and sample solutions must be the same primarily because of T in Nernst Eq. ATC probes are available for many meters

Errors in pH Measurement 3

- Alkaline Error or Sodium Error occurs when pH is very high (e.g., 12) because Na^+ concentration is high (from NaOH used to raise pH) and H^+ is very low. Electrode responds slightly to Na^+ & gives a lower reading than actual pH. This is related to the concept of selectivity coefficients where the electrode responds to many ions but is most selective for H^+ . Problem occurs because Na^+ is 10 orders of magnitude higher than H^+ in the solution.