CHEM.5140 Advanced Analytical Chemistry

Electroanalytical Methods Two general categories:

- Potentiometric Systems measure voltage (i.e., potential) of a galvanic cell (produces electricity spontaneously)
- 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)

   Indicator Electrode potential responds to activity of species of interest
   Reference Electrode chosen so that its potential is independent of solution composition.

#### $\mathsf{E}_{\mathsf{cell}} = \mathsf{E}_{\mathsf{ind}} - \mathsf{E}_{\mathsf{ref}} \ (+ \mathsf{E}_{\mathsf{J}})$

#### **Nernst Equation**



Where R = gas constant

- T = absolute temperature
- n = number of electrons in reaction
- F = Faraday's constant
- E = potential
- E<sup>o</sup> = standard potential

[Red] = molar concentration of reduced form of species

[Ox] = molar concentration of oxidized form of species

### **Reference Electrodes**

- Normal Hydrogen Electrode (NHE)
- $2H^+ + 2e^- \leftrightarrow H_2$   $E^\circ = 0.000$  v
- Saturated Calomel Electrode (SCE)
- $Hg_2CI_2 + 2 e^- \leftrightarrow 2 Hg + 2CI^- E^\circ = 0.268 v$
- Silver/Silver Chloride Electrode (AgCI)
- AgCl +  $e^- \leftrightarrow Ag$  + Cl<sup>-</sup>  $E^\circ = 0.222 v$

#### Silver/Silver Chloride Reference Electrode



## **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

$$Ag^+ + e^- \leftrightarrow Ag \qquad E^\circ = 0.799 v$$

#### **Nernst Equation**



#### At 25 °C this becomes

For the Silver/Silver ion system

 $Ag^+ + e^- \leftrightarrow Ag \qquad E^\circ = 0.799 v$ 

this becomes

 $\begin{array}{ccc} 0.0591 & 1 \\ E = 0.799 - ---- & \log & ----- \\ 1 & & [Ag^+] \end{array}$ 

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<sup>+</sup> ions

 $Ag^+ + e^- \leftrightarrow Ag$ 

## 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) AgCl +  $e^- \leftarrow \rightarrow$  Ag + Cl<sup>-</sup>  $E^\circ = 0.222 v$



For the Silver/Silver Chloride system

 $AgCI + e^{-} \leftrightarrow Ag + CI^{-} E^{\circ} = 0.222 v$ 

The Nernst equation becomes

0.0591 [Cl<sup>-</sup>] E = 0.222 - ----- log ------1 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<sup>2+</sup> with MnO<sub>4</sub><sup>-</sup>

# 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<sup>-</sup>, etc.
- Electrode of the Third Kind can be made that senses an ion that reacts with Cl<sup>-</sup> and changes the activity of Cl<sup>-</sup> detected

## Membrane Electrodes

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



# Glass is SiO<sub>2</sub>





## Membrane Electrodes

- Several types Glass membrane electrode
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#### Glass Membrane Electrode for pH $a_1$ $E = K + 0.0591 \log ------a_2$ $a_2$

#### where K is constant at fixed temp

In Glass Electrode, a<sub>2</sub> 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 pH since pH = -log  $a_{H+}$

### Glass pH Electrode

- E = K' 0.0591 pH
- Combine with reference electrode and meter
- Half cell voltage proportional to pH
- Nernstian slope
- Intercept is K', no E<sup>o</sup>
- Calibrate with buffers











# Proper pH Calibration

- E = K' 0.0591 pH
- Meter measures E vs pH must calibrate both slope & intercept on meter with buffers
- Meter has two controls calibrate & slope
- 1<sup>st</sup> use pH 7.00 buffer to adjust calibrate knob



Calibrate knob raises and lowers the line without changing slope

# Proper pH Calibration (cont.)

- 2<sup>nd</sup> 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)

$$Ag_{(s)} |AgCI_{(s)}| CI_{(aq)} ||H^+_{unk}|| HCI(0.1M) |AgCI_{(sat'd)}| Ag_{(s)}$$

reference electrode

# 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  $\pm 0.001$  units

- Junction potential dependent on ionic strength of solution – E<sub>j</sub> 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<sup>+</sup> solutions are the same activity, potential should be 0 but it's not

$$\mathsf{E}_{\mathsf{cell}} = \mathsf{E}_{\mathsf{ind}} - \mathsf{E}_{\mathsf{ref}} + \mathsf{E}_{\mathsf{j}} + \mathsf{E}_{\mathsf{a}}$$

 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<sup>+</sup> concentration is high (from NaOH used to raise pH) and H<sup>+</sup> is very low. Electrode responds slightly to Na<sup>+</sup> & 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<sup>+</sup>. Problem occurs because Na<sup>+</sup> is 10 orders of magnitude higher than H<sup>+</sup> in the solution.