PHOENIX ELECTRODE COMPANY FLUOROBORATE ION ELECTRODES INSTRUCTION MANUAL

GENERAL INSTRUCTIONS

Introduction

The pHoenix Fluoroborate Ion Electrodes are used to quickly, simply , accurately, and economically measure fluoroborate in aqueous solutions.

Required Equipment

1.

A pH/mV meter or an ion meter, either line operated or portable.

2. A magnetic stirrer.

3.

Semi-logarithmic 4-cycle graph paper for preparing c alibration curves when using the meter in the mV mode.

4.

Plastic beakers and storage bottles for all fluoroborate standards and samples. The hydrofluoric acid present in fluo roborate solutions will etch glass.

5.

The pHoenix Fluoroborate Ion Electrode, Cat. No. BF41501, (reference electrode necessary), the pHoenix Fluoroborate Gl ass Combination Electrode, Cat. No. BF41502, or the Fluorobora te Epoxy Combination Electrode, Cat. No. BF41503.

6.

The pHoenix Double Junction Reference Electrode, Cat. No . 5731429 (for use with the BF41501) with pHoenix Filling Solu tion, Cat No. R001011, in the inner junction and with pHoenix Filling Solution, Cat. No. R001044, in the outer junction.

Required Solutions

1.

Deionized or distilled water for solution and standard p reparation.

2.

pHoenix Ionic Strength Adjuster (ISA), 2M $(NH_4)_2SO_4$, Cat. No. BF4IS01. To prepare this solution from your own laborator y stock, half fill a 1000 ml volumetric flask with distilled w

ater and add 264 grams of reagent-grade ammonium sulfate, $(NH_4)_2SO_4$. Swirl the flask to dissolve the solid. Fill the flask to the mark with distilled water, cap, and invert the flask se veral times to mix the contents. ISA is added at the rate of 2 ml of ISA to each 100 ml of standard or sample to adjust the ionic strength to about 0.12M.

3.

pHoenix Electrode Filling Solution, $0.1M (NH_4)_2SO_4/0.01M$ KCl, Cat. No. R001045, for the BF41503 epoxy combination elect rode, or pHoenix Electrode Filling Solution, $0.1M (NH_4)_2SO_4$, Ca t. No. R001044, for the 5731429 double junction electrode or B F41502 glass combination electrode.

4.

pHoenix Fluoroborate Standard, 0.1M NaBF₄, Cat. No. B F4AS01. To prepare this solution, add 10.98 grams of reagentgrade NaBF₄ to about 500 ml of distilled water and dissolve th e solid. Filter quantitatively. Add the filtrate to a 1 lite r volumetric flask and fill to the mark with distilled water. This solution should be stored in polyethylene bottles and d iscarded after one week to avoid errors introduced by hydrol ysis of fluoroborate.

5.

pHoenix Fluoroborate Standard, 1000 ppm BF_4^{-1} , Cat. No.BF4 AS02. To prepare this solution, add 1.27 grams of reagent-gra de NaBF₄ to about 500 ml of distilled water and dissolve the s olid. Filter quantitatively. Add the filtrate to a 1 liter v olumetric flask and fill to the mark with distilled water. Th is solution should be stored in polyethylene bottles and disca rded after one week to avoid errors introduced to avoid errors introduced by hydrolysis of fluoroborate.

GENERAL PREPARATION

Electrode Preparation

Remove the rubber cap(s) covering the electrode tip(s) and any rubb er inserts covering the filling hole(s) of the reference electrode. Fill the combination electrode or the reference electrode with th e filling solution shipped with the electrode to a level just below the fill hole. No preparation is required for a sealed reference electrode. Gently shake the electrode downward in the same manner as a clinical thermometer to remove any air bubbles which may be tr apped behind the fluoroborate membrane. Prior to first usage, or a fter long-term storage, immerse the fluoroborate membrane in fluoro borate standard for thirty minutes. The electrode is now ready for use.

Connect the electrode(s) to the proper terminal(s) of the meter as

recommended by the meter manufacturer.

Electrode Slope Check (with pH/mV meter) (check electrodes each day)

1.

To a 150 ml plastic beaker, add 100 ml of distilled wate r and 2 ml of ISA. Place the beaker on a magnetic stirrer and begin stirring at a constant rate. After assuring that the m eter is in the mV mode, lower the electrode tip(s) into the solution.

2.

Using a pipet, add 1 ml of 0.1M or 1000 ppm fluoroborate standard to the beaker. When the reading has stabilized, rec ord the millivolt reading.

3.

Using a pipet, add 10 ml of the same fluoroborate s tandard used above to the beaker. When the reading has sta bilized, record the millivolt reading.

4.

Determine the difference between the two readings. A d ifference of 56 " 2 mV indicates correct electrode operati on, assuming the solution temperature is between 20° and 25°C. See the **TROUBLESHOOTING** section if the potential change is not within this range.

<u>Slope</u> is defined as the change in potential observed when the conce ntration changes by a factor of 10.

Electrode Slope Check (with ion meter) (check electrodes each day)

1.

Prepare standard fluoroborate solutions whose concentrations vary by a factor of ten. Use either the 0.1M or 1000 ppm fluoroborate standard. Use the serial dilution meth od for this preparation.

2.

To a 150 ml plastic beaker, add 100 ml of the lower valu e standard and 2 ml of ISA. Place the beaker on the magnetic stirrer and begin stirring at a constant rate. Lower the elect rode tip(s) into the solution. Assure that the meter is in th e concentration mode.

3.

Adjust the meter to the concentration of the standard an d fix the value in the memory according to the meter manufactu rer's instructions.

4. Rinse the electrode(s) and blot dry.

5.

To a 150 ml plastic beaker, add 100 ml of the higher value standard and 2 ml of ISA. Place the beaker on the magne tic stirrer and begin stirring at a constant rate. Lower the electrode tip(s) into the solution.

6.

Adjust the meter to the concentration of the standard an d fix the value in the memory.

7.

Read the electrode slope according to the meter m anufacturer's instructions. Correct electrode operation is in dicated by a slope of 90-100%. See the **<u>TROUBLESHOOTING</u>** secti on if the slope is not within this range.

MEASUREMENT

Measuring Hints

The sensing membrane is normally subject to water uptake and might appear milky. This has no effect on performance.

All samples and standards should be at the same temperature for pre cise measurement, preferably ambient temperature.

Constant, but not violent, stirring is necessary for accurate measu rement. Magnetic stirrers can generate sufficient heat to change t he solution temperature. To counteract this effect, place a piece of insulating material, such as styrofoam sheet, between the stirre r and beaker.

Always rinse the electrode(s) with distilled water and blot dry bet ween measurements. Use a clean, dry tissue to prevent cross-contam ination.

For samples with high ionic strength, prepare standards whose compo sition is similar to the sample.

A slow responding electrode may be caused by interferences to the e lectrode. To restore proper performance, soak the electrode in dis tilled water for about 5 minutes to clean the membrane, rinse, and soak in standard solution for about 5 minutes.

Always check to see that the membrane is free from air bubbles afte r immersion into standard or sample.

Sample Requirements

All samples must be aqueous and not contain organics which can diss olve in the membrane or extract out the liquid ion exchanger.

The temperature of the standard solutions and of the sample solutions should be the same and below 40° C. About a 2% error will be introduced for a 1°C difference in temperature.

Interferences should be absent. If they are present, use the proce dures found in the **Interferences** section to remove them.

The pH range for the fluoroborate ion electrode is 2.5-11. Neutral ize samples outside this range with acid or base to bring them in r ange. Fluoroborate sample with either high or low pH must be analy zed immediately after preparation or hydrolysis of fluoroborate to BF_3OH^{-1} , $BF_2(OH)_2^{-1}$, and $BF(OH)_3^{-1}$ will occur. Convert boric acid or b orate ion to fluoroborate by addition of HF.

Units of Measurement

Fluoroborate concentrations are measured in units of ppm as boron, ppm as fluoroborate, moles per liter, or any other convenient conce ntration unit. Table 1 indicates some of the concentration units.

TABLE 1: Concentration Unit Conversion Factors

ppm B	$\underline{ppm BF_4}^{-1}$	moles/liter
108.9	868.0	1.0×10^{-2}
10.8	86.8	1.0×10^{-3}
1.1	8.7	1.0×10^{-4}

MEASUREMENT PROCEDURE

Direct Measurement

A simple procedure for measuring a large number of samples. A sing le meter reading is all that is required for each sample. The ioni c strength of samples and standards should be made the same by adju stment with ISA for all fluoroborate solutions. The temperature of both sample solution and standard solutions should be the same.

Direct Measurement of Fluoroborate (using a pH/mV meter)

1.

By serial dilution of the 0.1 M or 1000 ppm standard, pr epare 10^{-2} , 10^{-3} , and 10^{-4} M or 100 and 10 ppm fluoroborate stand ards. Add 2 ml of ISA per 100 ml of standard. Prepare standa rds with a composition similar to the samples if the samples have an ionic strength above 0.1M. Use plastic lab ware.

2.

Place the most dilute solution $(10^{-4}M \text{ or } 10 \text{ ppm})$ on the m agnetic stirrer and begin stirring at a constant rate After a ssuring that the meter is in the mV mode, lower the electrode tip(s) into the solution. When the reading has stabilized, re cord the mV reading.

3.

Place the mid-range solution $(10^{-3}M \text{ or } 100 \text{ ppm})$ on the m agnetic stirrer and begin stirring. After rinsing the electr ode(s) in distilled water, blot dry and immerse the electrode tip(s) in the solution. When the reading has stabilized, record the mV reading.

4.

Place the most concentrated solution $(10^{-2}M \text{ or } 1000 \text{ ppm})$ on the magnetic stirrer and begin stirring. After rinsing the electrode(s) in distilled water, blot dry, and immerse the el ectrode tip(s) in the solution. When the reading has stabiliz ed, record the mV reading.

5.

Using the semi-logarithmic graph paper, plot the mV read ing (linear axis) against the concentration (log axis). A typ ical calibration curve can be found in Figure 1.

A calibration curve is constructed on semi-logarithmic paper when u sing the pH/mV meter in the millivolt mode. The measured elec trode potential in mV (linear axis) is plotted against the st andard concentration (log axis). In the linear region of the curve, only three standards are necessary to determine a cali bration curve. In the non-linear region, additional points m ust be measured. The direct measurement procedures given are for the linear portion of the curve. The non-linear portion of the curve requires the use of low level procedures.

Extrapolate the calibration curve down to about 1.0 x $10^{-5}M$ (0.1 ppm B).

To a clean, dry 150 ml plastic beaker, add 100 ml of the sample and 2 ml of ISA. Place the beaker on the magnetic sti rrer and begin stirring at a constant rate. After rinsing th e electrode tip(s) with distilled water and blotting dry, lowe r the electrode tip(s) into the solution. When the reading ha s stabilized, record the mV reading. Determine the concentrat ion directly from the calibration curve.

7.

The calibration should be checked every two hours. Assu ming no change in ambient temperature, place the electrode tip (s) in the mid-range standard. After the reading has stabiliz ed, compare it to the original reading recorded in Step 3 abov e. A reading differing by more than 0.5 mV or a change in the ambient temperature will necessitate the repetition of Steps 2-5 above. A new calibration curve should be prepared daily.

Direct Measurement of Fluoroborate (using an ion meter)

1.

By serial dilution of the 0.1M or 1000 ppm fluoroborate standard, prepare two fluoroborate standards whose concentrati on is near the expected sample concentration. Measure out 100 ml of each standard into individual 150 ml plastic beakers an d add 2 ml of ISA to each.

2.

Place the most dilute solution on the magnetic stirrer a nd begin stirring at a constant rate. Assure that the meter i s in the concentration mode. Lower the electrode tip(s) into the solution.

3.

Adjust the meter to the concentration of the fluoroborat e standard and fix the value in the memory according to the me termanufacturer's instructions after stabilization of the reading.

4. Rinse the electrode(s) with distilled water and blot dry.

5.

Place the most concentrated solution on the magnetic stirrer and begin stirring at a constant rate. Lower the electro de tip(s) into the solution.

6.

Adjust the meter to the concentration of the fluoroborat e standard and fix the value in the memory according to the me termanufacturer's instructions after stabilization of the reading.

6.

7.

For low level measurements, place the rinsed, dried elec trode(s) into a solution containing 100 ml of distilled water and 2 ml of ISA. After stabilization, fix the blank value in the meter according to the meter manufacturer's instructions.

8.

Place 100 ml of the sample and 2 ml of ISA in a 150 ml p lastic beaker. Place the beaker on the magnetic stirrer and b egin stirring.

9.

Immerse the electrode tip(s) in the solution and wait fo r the reading to stabilize. Read the concentration directly f rom the meter display.

10.

The calibration should be checked every two hours. Assu ming no change in ambient temperature, place the electrode tip (s) in the first fluoroborate standard. After the reading ha s stabilized, compare it to the original reading in Step 3 ab ove. A reading differing by more than 0.5 mV or a change in ambient temperature will necessitate the repetition of Steps 2 -6 above. The meter should be re-calibrated daily.

Low Level Fluoroborate Determination (using a pH/mV meter)

This procedure is recommended for solutions with ionic strengths le ss than 1.0×10^{-2} M. If the solution is high in ionic strength, but low in fluoroborate, use the same procedure, but prepare a calibra tion solution with a composition similar to the sample.

1.

Using 20 ml of standard ISA, dilute to 100 ml with disti lled water. This low level ISA $[0.4M (NH_4)_2SO_4]$ is added at th e rate of 1 ml low level ISA to each 100 ml of sample. The b ackground ionic strength will be $4.0 \times 10^{-3}M$.

2.

Dilute 1 ml of 0.1M standard to 100 ml to prepare a 1.0 x $10^{-3}M$ BF₄⁻¹ solution for measurements in moles per liter. Use the 1000 ppm standard for preparing a 100 ppm BF₄⁻¹ standard b y diluting 10 ml of the 1000 ppm standard to 100 ml. Standard s should be prepared fresh daily.

3.

Add 1 ml of the low level ISA to a 100 ml volumetric f lask and fill to the mark with distilled water. Pour this sol ution into a 150 ml plastic beaker and place the beaker on the magnetic stirrer. Begin stirring at a constant rate. Place the electrode tip(s) in the solution. Assure that the meter is in the mV mode.

5.

4.

Add increments of the 1.0 x $10^{-3} \mbox{M}$ or 100 ppm standards as given in Table 2.

6.

After the reading has stabilized, record the mV reading after each addition.

TABLE 2: Step-wise Calibration for Low Level Fluoroborate Measurements

Step	Pipet	Added <u>Volume (ml)</u>	Concentra <u>M BF₄⁻¹</u>	tion <u>ppm BF_4^{-1}</u>
1	А	0.1	1.0×10^{-6}	0.1
2	A	0.1	2.0×10^{-6}	0.2
3	A	0.2	4.0×10^{-6}	0.4
4	A	0.2	6.0×10^{-6}	0.6
5	A	0.4	9.9 x 10^{-6}	1.0
б	В	2.0	2.9×10^{-5}	2.9
7	В	2.0	4.8×10^{-5}	4.8

Pipet A = 1 ml graduated pipet Pipet B = 2 ml pipet Solutions: additions of 1.0×10^{-3} M or 100 ppm standard to 100 ml of ISA as prepared in Step 3 above.

7.

On semi-logarithmic graph paper, plot the millivolt r eading (linear axis) against the concentration (log axis) as i n Figure 1.

8. Rinse the electrode(s) and blot dry.

9.

Measure out 100 ml of the sample into a 150 ml plastic beaker, add 1 ml of low level ISA, and place the beaker on the magnetic stirrer. Begin stirring. Lower the electrode tip(s) into the solution. After the reading has stabilized, record the mV reading and determine the concentration from the low le vel calibration curve. 10.

Prepare a new low level calibration curve daily. Check the calibration curve every two hours by repeating Steps 2-7.

Low Level Fluoroborate Determination (using an ion meter)

Follow the procedure given for normal fluoroborate determinations u sing an ion meter and the blank correction procedure.

ELECTRODE CHARACTERISTICS

Reproducibility

Electrode measurements reproducible to "2% can be obtained if the e lectrode is calibrated every hour. Factors such as temperature flu ctuations, drift, and noise limit reproducibility. Reproducibility is independent of concentration within the electrode's operating range.

Interferences

Certain anions are electrode interferences and will cause electrode malfunction, drift or measurement errors if present in high enough levels. The level of interfering common anions that will cause a 10% error at three levels of fluoroborate is given in Table 3.

<u>TABLE 3</u>: Concentration of Possible Interferences Causing a 10% Error at Various Levels of Fluoroborate; Background Ionic Strength of 0.12M (NH₄)₂SO₄.

Interferer (moles/lit	$\frac{10^{-2}M}{10^{-2}M}$	$10^{-3}M$	$10^{-4}M$ BF ₄ ⁻¹
PO_4^{-3}	8.0×10^{-1}	8.0x10 ⁻²	8.0x10 ⁻³
$\mathrm{HPO_4}^{-2}$	8.0x10 ⁻¹	8.0x10 ⁻²	8.0x10 ⁻³
$H_2PO_4^{-1}$	8.0x10 ⁻¹	8.0x10 ⁻²	8.0x10 ⁻³
Cl^{-1}	5.0×10^{-1}	5.0×10^{-2}	5.0x10 ⁻³
CO_{3}^{-2}	3.0×10^{-1}	3.0×10^{-2}	3.0×10^{-3}
HCO_3^{-1}	3.0×10^{-1}	3.0x10 ⁻²	3.0×10^{-3}
NO_3^{-1}	5.0×10^{-2}	5.0x10 ⁻³	5.0×10^{-4}
NO_2^{-1}	1.0×10^{-2}	1.0×10^{-3}	1.0×10^{-4}

Br^{-1}	1.0×10^{-2}	1.0×10^{-3}	1.0×10^{-4}
CN^{-1}	5.0×10^{-3}	5.0×10^{-4}	5.0×10^{-5}
ClO_3^{-1}	5.0×10^{-4}	5.0×10^{-5}	5.0×10^{-6}
I^{-1} ClO_4^{-1}	5.0x10 ⁻⁵ 5.0x10 ⁻⁶	5.0×10^{-6} 5.0×10^{-7}	5.0×10^{-7} 5.0×10^{-8}

Interferences (ppm)	100 ppm	10 ppm 1	ppm
PO_4^{-3}	7.0×10^{4}	7.0×10^{3}	7.0×10^{2}
HPO_4^{-2}	7.1×10^{4}	7.1×10^{3}	7.1×10^{2}
$H_2PO_4^{-1}$	7.2×10^{4}	7.2×10^{3}	7.2×10^{2}
Cl ⁻¹	1.6×10^{4}	1.6x10 ³	1.6×10^{2}
CO_{3}^{-2}	1.7x10 ⁴	1.7x10 ³	1.7×10^{2}
HCO3 ⁻¹	1.7x10 ⁴	1.7x10 ³	1.7×10^{2}
NO ₃ ⁻¹	2.9×10^{3}	2.9×10^{2}	2.9×10^{1}
NO_2^{-1}	4.2×10^{2}	4.2x10 ¹	4.2
Br^{-1}	7.4×10^{2}	7.4x10 ¹	7.4
CN^{-1}	1.2×10^{2}	1.2x10 ¹	1.2
ClO_3^{-1}	4.0x10 ¹	4.0	$4.0 x 10^{-1}$
I ⁻¹	6.0	6.0x10 ⁻¹	6.0x10 ⁻²
ClO_4^{-1}	5.0×10^{-1}	5.0×10^{-2}	5.0x10 ⁻³

Interferences such as phosphate, bromide, iodide and cyanide can be removed by precipitation with 0.5 grams of silver sulfate added to 100 ml of sample. Interferences such as carbonate and bicarbonate can be removed by conversion to carbon dioxide through acidificati on of the sample to pH 4.5 with sulfuric acid. Nitrite interferenc e can be removed by adding 0.3 grams of sulfamic acid to 100 ml of sample.

The above interference removal procedures require similar treatment of standards as well as samples.

If the electrode is exposed to high levels of interfering ions whic h cannot be removed, the electrode reading may drift and the respon se may become sluggish. Restore performance by soaking in distille d water for 30 minutes followed by soaking in fluoroborate standard for 30 minutes.

Temperature Influences

Samples and standards should be at the same temperature, since elec trode potentials are influenced by changes in temperature. A 1°C di fference in temperature results in a 2% error at the 10^{-3} M level. B ecause of the solubility equilibria on which the electrode depends, the absolute potential of the reference electrode changes slowly w ith temperature. The slope of the fluoroborate electrode, as indic ated by the factor "S" in the Nernst equation, also varies with tem perature. Table 4 gives values for the "S" factor in the Nernst eq uation for the fluoroborate ion.

The operating range of the fluoroborate ion electrode is $0^{\circ}-40^{\circ}C$, pr ovided that temperature equilibrium has occurred.

If the temperature varies substantially from room temperature, equi librium times up to one hour are recommended.

TABLE 4: Temperature vs Values for the Electrode Slope

Temp (°C)	"S"
0	54.20
10	56.18
20	58.18
25	59.16
30	60.15
40	62.13
50	64.11

Electrode Response

Plotting the mV potential against the fluoroborate concentration on semi-logarithmic paper results in a straight line with a slope of about 56 mV per decade. (Refer to Figure 1.)

The time needed to reach 99% of the stable electrode potential read ing, the electrode response time, varies from one minute or less in highly concentrated solutions to several minutes near the detectio n limit. (Refer to Figure 2.)

Limits of Detection

The upper limit of detection in pure sodium fluoroborate solutions is 1M. In the presence of other ions, the upper limit of detection is above 10⁻¹M fluoroborate, but two factors influence this upper l imit. Both the possibility of a liquid junction potential developi ng at the reference electrode and the salt extraction effect influe nce this upper limit. Some salts may infuse into the electrode mem brane at high salt concentrations, causing deviation from the theor etical response. Either dilute samples between 1M and 10⁻¹M or cali brate the electrode at 4 or 5 intermediate points.

The lower limit of detection is influenced by the slight water solu bility of the ion exchanger used in the sensing portion of the elec trode. Refer to Figure 1 for a comparison of the theoretical response to the actual response at low levels of fluoroborate. Fluorobo rate measurements below $10^{-5}M$ BF₄⁻¹ (0.87 ppm as BF₄⁻¹ or 0.11 ppm as B) should employ low level procedures.

Electrode Storage

The fluoroborate electrodes may be stored for short periods of time in 10^{-2} M fluoroborate solution. For longer storage (longer than tw o weeks), rinse and dry the fluoroborate membrane and cover the tip with any protective cap shipped with the electrode(s). The refere nce portion of the combination electrode (or the outer chamber of t he reference electrode) should be drained of filling solution, if r efillable and the rubber insert placed over the filling hole.

pH Effects

The operating range of the fluoroborate electrode is from pH 2.5 to pH 11.

Electrode Life

The fluoroborate electrode will last six months in normal laborator y use. On-line measurement might shorten operational lifetime to s everal months. In time, the response time will increase and the ca libration slope will decrease to the point calibration is difficult and electrode replacement is required.

ELECTRODE THEORY

Electrode Operation

The fluoroborate electrode consists of an electrode body containing a liquid internal filling solution in contact with a gelled organo philic membrane containing a fluoroborate ion exchanger. When the membrane is in contact with a solution containing free fluoroborate ions, an electrode potential develops across the membrane. This e lectrode potential is measured against a constant reference potenti al, using a standard pH/mV meter or an ion meter. The level of flu oroborate ions, corresponding to the measured potential, is describ ed by the Nernst equation:

$$E = E$$
 - S log X

where: E = measured electrode potential E\$= reference potential (a constant) S = electrode slope (-56 mV) X = level of fluoroborate ions in solution

The activity, X, represents the effective concentration of the ions in solution. The total fluoroborate ion concentration, C_t , is the sum of free fluoroborate ion, C_f , and complexed or bound fluoroborate te ion, C_b . The electrode is able to respond to only the free ions, whose concentration is :

 $C_f = C_t - C_b$

Since fluoroborate ions form very few stable complexes, the free io n concentration may be equated to the total ion concentration.

The activity is related to the free ion concentration, C_f , by the a ctivity coefficient, a, by:

 $X = a C_f$

Activity coefficients vary, depending on total ions strength, I, de fined as:

I = $1/2 \ 3C_xZ_x$? where: C_x = concentration of ion X Z_x = charge of ion X 3 = sum of all of the types of ions in the solution

In the case of high and constant ionic strength relative to the sen sed ion concentration, the activity coefficient, a, is constant and the activity, X, is directly proportional to the concentration. To adjust the background ionic strength to a high and constant valu e, ionic strength adjuster (ISA) is added to samples and standards. The recommended ISA for fluoroborate is $(NH_4)_2SO_4$. Solutions other than this may be used as ionic strength adjusters as long as ions that they contain do not interfere with the electrode's response to fluoroborate ions.

The reference electrode must also be considered. When two solution s of different composition are brought into contact with one anothe r, liquid junction potentials arise. Millivolt potentials occur fr om the inter-diffusion of ions in the two solutions. Electrode cha rge will be carried unequally across the solution boundary resultin g in a potential difference between the two solutions, since ions d iffuse at different rates. When making measurements, it is importa nt to remember that this potential be the same when the reference i s in the standardizing solution as well as in the sample solution o r the change in liquid junction potential will appear as an error i n the measured electrode potential.

The composition of the liquid junction filling solution in the refe rence electrode is most important. The speed with which the positi ve and negative ions in the filling solutions diffuse into the samp les should be as nearly equal as possible, that is, the filling sol ution should be equitransferent. No junction potential can result if the rate at which positive and negative charge carried into the sample is equal.

Strongly acidic (pH = 0-2) and strongly basic (pH = 12-14) solution s are particularly troublesome to measure. The high mobility of hy drogen and hydroxide ions in samples make it impossible to mask the ir effect on the junction potential with any concentration of equit ransferent salt. One must either calibrate the electrodes in the s ame pH range as the sample or use a known incremental method for io n measurement.

TROUBLESHOOTING GUIDE

The goal of troubleshooting is the isolation of a problem through c hecking each of the system components in turn: the meter, the plast ic ware, the electrode(s), the standards & reagents, the sample, an d the technique.

Meter

The meter may be checked by following the check-out procedure in th e instrument instruction manual.

Plastic ware

Clean plastic ware is essential for good measurement. Be sure to w ash the plastic ware well with a mild detergent and rinse very well with distilled or deionized water.

Electrodes

The electrodes may be checked by using the procedure found in the s ections entitled **Electrode Slope Check**.

1.

Be sure to use distilled or deionized water when following the procedures given in **Electrode Slope Check**.

2.

If the electrode fails to respond as expected, see the s ection **Measuring Hints**. Repeat the slope check.

3.

If the electrodes still fail to respond as expected, su bstitute another fluoroborate ion electrode that is known to b e in good working order for the questionable electrode. If th e problem persists and you are using an electrode pair, try th e same routine with a working reference electrode.

4.

If the problem persists, the standards or reagent may be of poor quality, interferences in the sample may be present or the technique may be faulty. See **Standards & Reagents**, **Sampl** e, and **Technique** sections below.

5.

If another electrode is not available for test purposes, or if the electrode in use is suspect, review the instruction manual and be sure to:

- Clean and rinse the electrode(s) thoroughly.

- Prepare the electrode(s) properly.

- Use the proper filling solution.

- Adjust the pH and the ionic strength of the solution by the use of the proper ISA.

- Measure correctly and accurately.

- Review TROUBLESHOOTING HINTS.

Standards & Reagents

Whenever problems arise with the measuring procedure that has been used successfully in the past, be sure to check the standard and re agent solutions. If in doubt about the credibility of any of the s olutions, prepare them again. Errors may result from contamination of the ISA, incorrect dilution of standards, poor quality distille d/deionized water, or a simple mathematical miscalculation.

Sample

Look for possible interferences, complexing agents, or substances w hich could affect the response or physically damage the sensing ele ctrode (or the reference electrode) if the electrode(s) work perfec tly in the standard, but not in the sample.

Try to determine the composition of the samples prior to testing to eliminate a problem before it starts. (See Measuring Hints, Sampl e Requirements, and Interferences.)

Technique

Be sure that the electrode's limit of detection has not been exceed ed. Be sure that the analysis method is clearly understood and tha t Good Laboratory Practice has been followed. Refer to the instruc tion manual again. Reread <u>GENERAL</u> <u>PREPARATION</u> and <u>ELECTRODE</u> <u>CHARAC</u> <u>TERISTICS</u>.

If trouble still persists, call pHoenix Electrode Company at 1-800-522-7920 and ask for the Technical Services Department.

TROUBLESHOOTING HINTS

Symptom	Possible Causes	Next Step
Out of Range Reading	defective meter (i manual)	check meter with shorting strap see meter .nstruction
	defective electrode	check electrode
plugge	electrode(s) not ed in properly and	<pre>unplug electrode(s) reseat</pre>
	reference electrode not filled	be sure reference electrode is filled
	air bubble on membrane	remove bubble by re-dipping electrode

	electrodes not in solution	<pre>put electrode(s) in solution</pre>
Noisy or Unstable Readings (reading	defective meter s	check meter with shorting strap
rapidly changing)	air bubble on membrane	remove bubble by re-dipping electrode
	defective electrode	replace electrode
	ISA not used	use recommended ISA
	meter or stirrer not grounded	ground meter or stirrer
	electrode exposed to interferences standard	soak electrode in fluoroborate
	outer filling solution level too low	fill electrode to level just below the fill hole
Drift (reading slowly changing e direction) t	samples and stand at different emperatures m	dards allow solutions to come to room temper-in on ature before easurement
	electrode exposed to interferences standard	soak electrode in fluoroborate
	incorrect reference filling solution	use recommended filling solution
Low Slope or ope ate	standards contam d or incorrectly s made	in- prepare fresh No Sl standards
	ISA not used I	use recommended SA
	standard used as IS.	A use ISA
	electrode exposed to interferences standard	soak electrode in fluoroborate

defective electrode check electrode operation air bubble on remove bubble by re-dipping probe membrane "Incorrect Answer" incorrect scaling plot millivolts on (but calibration of semi-log paper the linear axis. On cur the log axis, be ve is good) sure concentration numbers within each decade are increa sing with increasing concentration. incorrect sign be sure to note sign of millivolt number correctly incorrect standards prepare fresh standards wrong units used apply correct conversion factor: $10^{-3}M = 10.8 \text{ ppm as}$ BF_4^{-1} Boron = 86.8 ppm as complexing agents use de-complexing in sample procedure sample carryover rinse electrodes thoroughly between samples

SPECIFICATIONS

Concentrati	on Range:	1M to 7 x 10^{-6} M (1.1 x 10^4 to 9.0 x 10^{-2} ppm as B)
pH Range:		2.5 to 11
Temperature	Range:	0° to 40° C
Resistance:		100 megaohms
Reproducibi	lity:	" 2%
Samples:		aqueous solutions only no organic solvents
Size:		110 mm in length 12 mm in diameter 1 m cable length
Storage:		Store in dilute fluoroborate solution
ORDERING IN P/N	FORMATION DESCRIPTION	N
BF41501	Fluorobo electrode	rate Electrode, mono (reference necessary), PVC body
BF41502	Fluorobo	rate Electrode, combination, glass body
BF41503	Fluorobo	rate Electrode, combination, epoxy body
5731429 bo	Referenc ody, for us	e Electrode, double junction, sleeve, epox se with the BF41501
BF4AS01	Fluorobo	rate Standard, 0.1M $NaBF_4$
BF4AS02	Fluorobo	rate Standard. 1000 ppm BF_4^{-1}
BF4IS01	Fluorobo 2 M (NH ₄) ₂ S	rate ISA (Ionic Strength Adjustor), SO_4
R001011	5731429 H 4 M KCl (w	Reference Electrode Inner Filling Solution ith Ag+)
R001044	5731429 H Solution & Solution,	Reference Electrode Outer Filling BF41502 Combination Electrode Filling 0.1M $(NH_4)_2SO_4$

epoxy

BF41501 Combination Electrode Filling Solution, 0.1M $(\rm NH_4)_2SO_4/0.01M$ KCl (with Ag+) R001045

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