2-10-1994

Solubility Studies of Iron(III) Oxides and Hydroxides

Victor L. Menting
Portland State University

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THESIS APPROVAL

The abstract and thesis of Victor L. Menting for the Master of Science in Chemistry was presented February 10, 1994, and accepted by the thesis committee and the department.

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**************************************************************************  

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ABSTRACT


Title: Solubility Studies Of Iron(III) Oxides And Hydroxides.

The hydrolysis of iron(III) ions in aqueous solution forms a series of soluble hydroxide complexes with associated equilibrium constants. The solubility of iron(III) is controlled by the various soluble hydroxide complexes, and can, in theory, be calculated from the pH and equilibrium constants. Experimental verification of the calculated solubility has proven difficult due to the lack of sensitive analytical techniques and the presence of colloidal ferric hydroxide interferences.

Recently, electrochemical methods for the determination of low levels of iron(III) have been developed using adsorptive cathodic stripping voltammetry which relies on the interfacial accumulation of the chelate of iron with Solochrome Violet RS on a hanging mercury drop electrode.

The purpose of this investigation was to experimentally verify the calculated solubility of iron(III) in the pH 4-12 region using adsorptive cathodic stripping voltammetry.

The ubiquitous nature of iron requires background
levels of iron be reduced below the experimental concentrations to be determined. Attempts to lower the background levels of iron were ineffective as concentrations below about $10^{-8}$ M iron could not be attained.

Verification of the calculated solubility of iron(III) was unsuccessful as background concentrations of iron(III) and the presence of colloidal ferric hydroxide hindered the experimental results. The dissolution of the ferric hydroxide colloids coupled with the background levels of iron resulted in the determination of experimental concentrations which exceeded theoretical values by two to four orders of magnitude.
SOLUBILITY STUDIES OF IRON(III) OXIDES AND HYDROXIDES

by

VICTOR L. MENTING

A thesis submitted in partial fulfillment of the requirements for the degree of

MASTER OF SCIENCE

in

CHEMISTRY

Portland State University

1994
DEDICATION

This work is dedicated to the two people I admire and respect most, my parents, James R. and Judith A. Menting. Their love and support has been a constant source of inspiration.
ACKNOWLEDGEMENTS

The author would like to extend his deepest gratitude and sincerest appreciation to Dr. Dennis Barnum for sharing his wealth of knowledge. Dr. Barnum's insights greatly aided the author in the completion of this investigation.
# TABLE OF CONTENTS

<table>
<thead>
<tr>
<th>DEDICATION</th>
<th>ii</th>
</tr>
</thead>
<tbody>
<tr>
<td>ACKNOWLEDGEMENTS</td>
<td>iii</td>
</tr>
<tr>
<td>LIST OF TABLES</td>
<td>v</td>
</tr>
<tr>
<td>LIST OF FIGURES</td>
<td>vi</td>
</tr>
<tr>
<td>CHAPTER</td>
<td></td>
</tr>
<tr>
<td>I  INTRODUCTION</td>
<td>1</td>
</tr>
<tr>
<td>Theory</td>
<td>1</td>
</tr>
<tr>
<td>II  EXPERIMENTAL</td>
<td>10</td>
</tr>
<tr>
<td>Apparatus</td>
<td>10</td>
</tr>
<tr>
<td>Possible Sources Of Iron Contamination</td>
<td>19</td>
</tr>
<tr>
<td>III RESULTS</td>
<td>33</td>
</tr>
<tr>
<td>Differential Pulse Polarography</td>
<td>33</td>
</tr>
<tr>
<td>Fast Scan D.C. Polarography</td>
<td>42</td>
</tr>
<tr>
<td>IV  DISCUSSION</td>
<td>46</td>
</tr>
<tr>
<td>Conclusions</td>
<td>46</td>
</tr>
<tr>
<td>Future Work</td>
<td>53</td>
</tr>
<tr>
<td>V  REFERENCES</td>
<td>54</td>
</tr>
<tr>
<td>Literature Cited</td>
<td>54</td>
</tr>
<tr>
<td>Selected Bibliography</td>
<td>56</td>
</tr>
</tbody>
</table>
# LIST OF TABLES

<table>
<thead>
<tr>
<th>TABLE</th>
<th>PAGE</th>
</tr>
</thead>
<tbody>
<tr>
<td>I</td>
<td>2</td>
</tr>
<tr>
<td>II</td>
<td>4</td>
</tr>
<tr>
<td>III</td>
<td>8</td>
</tr>
<tr>
<td>IV</td>
<td>14</td>
</tr>
<tr>
<td>V</td>
<td>20</td>
</tr>
<tr>
<td>VI</td>
<td>22</td>
</tr>
<tr>
<td>VII</td>
<td>31</td>
</tr>
<tr>
<td>VIII</td>
<td>38</td>
</tr>
</tbody>
</table>

**TABLE I**  
Equilibrium Reactions And Equilibrium Constants For The Iron(III) Hydroxy Complexes From Baes And Mesmer [1976]  

**TABLE II**  
Literature Equilibrium Constants  

**TABLE III**  
Solubility Product Constants Of The Different Crystalline Forms Of Ferric Hydroxide $K_{sp} = [Fe^{3+}][OH^-]^3$ From Lindsay [1985]  

**TABLE IV**  
Preparation Of Buffers With Actual pH's, Ionic Strengths, And Buffer System  

**TABLE V**  
Concentration Of Citrate (ng/mL) Added And The Resulting Peak Current Increases  

**TABLE VI**  
Change In The Calculated Iron Concentration For Buffers With And Without Filtration  

**TABLE VII**  
Increases In Current With Time At Various Conditions Involving Solution Stirring And Nitrogen Purging  

**TABLE VIII**  
Concentration of Total Dissolved Iron(III) Calculated From Aqueous Ferric Perchlorate, Colloidal Ferric Hydroxide, And Goethite At Various pH's
## LIST OF FIGURES

<table>
<thead>
<tr>
<th>FIGURE</th>
<th>PAGE</th>
</tr>
</thead>
<tbody>
<tr>
<td>2. Comparison Of The Calculated Solubility Of Fe(OH)$_3$(s) With Experimental Solubility Measurements From The Literature</td>
<td>5</td>
</tr>
<tr>
<td>3. Structure Of Solochrome Violet RS</td>
<td>17</td>
</tr>
<tr>
<td>4. Concentration Versus Time Showing The Dissolution Of Colloidal Ferric Hydroxide</td>
<td>24</td>
</tr>
<tr>
<td>5. Concentration Versus Total Iron Added As Colloidal Ferric Hydroxide</td>
<td>26</td>
</tr>
<tr>
<td>6. Differential Pulse Voltammogram Of Solution Containing Supporting electrolyte/SVRS And Iron, Spiked With Al(III)</td>
<td>28</td>
</tr>
<tr>
<td>7. Approximate Concentration Of Iron As A Function Of Time To Determine The Extent Of Iron Contamination Resulting From Iron Diffusion Out Of The Electrode Frit</td>
<td>29</td>
</tr>
</tbody>
</table>
8. Voltammograms Obtained In The Differential Pulse Mode For SVRS Containing Solutions In The Absence And Presence Of Added Iron 34

9. Voltammograms Of Current Versus Added Iron In The Differential Pulse Mode 35

10. Analysis Of Background Iron By Standard Addition For Differential Pulse Polarography 36

11. Comparison Of Calculated Solubility With Experimental Concentrations Of Iron In Solutions Equilibrated With Ferric Perchlorate 39

12. Comparison Of Calculated Solubility With Experimental Concentrations Of Iron In Solutions Equilibrated With Colloidal Ferric Hydroxide 40

13. Comparison Of Calculated Solubility With Experimental Concentrations Of Iron In Solutions Equilibrated With Goethite 41

14. Voltammogram Of Fast Scan D.C. Polarography 43

15. Analysis Of Background Iron By Standard Addition For Fast Scan D.C. Polarography 45

16. Comparison Of The Solubility of Fe(OH)$_{2.35}X_{0.65}$ Calculated Using The Solubility Product Proposed By Fox [1988] 47
17. Comparison Of The Solubility of Fe(OH)$_{2.35X_{0.65}}$
Calculated Using The Solubility Product
Proposed By Fox [1988], With The
Experimentally Measured Solubility Found In
This Study . . . . . . . . . . . . . . . . . 48

18. Graphical Representation Of U{M(OH)$_y$} Versus
y For Literature Values Of The
Equilibrium Constants Of The Iron Hydroxy
Complexes . . . . . . . . . . . . . . . . . 51
CHAPTER I

INTRODUCTION

THEORY

Iron(III) ions in aqueous solution are surrounded by a primary hydration sphere of six water molecules, $\text{Fe(H}_2\text{O)}_6^{3+}$. As the pH increases, $\text{H}^+$ ions are removed from the hydration sphere to form a series of soluble hydroxide complexes according to the reaction (water molecules are omitted for brevity):

$$x \text{Fe}^{3+} + y \text{H}_2\text{O} \rightleftharpoons \text{Fe}_x(\text{OH})_{y}^{(3x-y)^+} + y \text{H}^+ \quad (1)$$

The complexes formed are the mononuclear $\text{Fe(OH)}_2^{2+}$, $\text{Fe(OH)}_2^+$, $\text{Fe(OH)}_3^-$, and $\text{Fe(OH)}_4^-$; and the polynuclear complexes $\text{Fe}_2(\text{OH})_4^{4+}$ and $\text{Fe}_3(\text{OH})_4^{5+}$. (The superscript "o" in $\text{Fe(OH)}_3^-$ will be used to distinguish the neutral water soluble complex from the $\text{Fe(OH)}_3$ precipitate.) For simplicity, the general form, $K_{xy}$, will be used for the equilibrium constant of the complex, $\text{Fe}_x(\text{OH})_{y}^{3x-y}$. Equilibrium constants are given in Table I.

The following equation can be written to describe the solubility of iron(III) in aqueous solution:

$$[\text{Fe}^{3+}]_T = [\text{Fe}^{3+}] + [\text{Fe(OH)}_2^{2+}] + [\text{Fe(OH)}_2^+] + [\text{Fe(OH)}_3^-] + [\text{Fe(OH)}_4^-] + 2[\text{Fe}_2(\text{OH})_2^{4+}] + 3[\text{Fe}_3(\text{OH})_4^{5+}] \quad (2)$$

where $[\text{Fe}^{3+}]_T$ is the total iron concentration, i.e. the
solubility.

TABLE I

EQUILIBRIUM REACTIONS AND EQUILIBRIUM CONSTANTS FOR THE IRON(III) HYDROXY COMPLEXES
FROM BAES AND MESMER [1976]

\[
\begin{align*}
\text{Equilibrium Reaction} & & \log K_{xy} \\
\text{Fe}^{3+} + H_2O & \leftrightarrow \text{FeOH}^{2+} + H^+ & -3.05 \\
\text{Fe}^{3+} + 2H_2O & \leftrightarrow \text{Fe(OH)}_2^{3+} + 2H^+ & -6.31 \\
\text{Fe}^{3+} + 3H_2O & \leftrightarrow \text{Fe(OH)}_3^- + 3H^+ & < -12 \\
\text{Fe}^{3+} + 4H_2O & \leftrightarrow \text{Fe(OH)}_4^{4+} + 4H^+ & -21.6 \\
2\text{Fe}^{3+} + 2H_2O & \leftrightarrow \text{Fe}_2(\text{OH})_4^{2+} + 2H^+ & -2.96 \\
3\text{Fe}^{3+} + 4H_2O & \leftrightarrow \text{Fe}_3(\text{OH})_4^{4+} + 4H^+ & -5.77
\end{align*}
\]

Substitution of the equilibrium expressions for the complexes into Equation 2 yields:

\[
[\text{Fe}^{3+}]_T = \frac{K_{sp}(K_{11}[H^+]^{-1} + K_{12}[H^+]^{-2} + K_{13}[H^+]^{-3} + K_{14}[H^+]^{-4} + 2K_{22}[H^+]^{-2} + 3(K_{34}[H^+]^{-4})}{2K_{22}[H^+]^{-2} + 3(K_{34}[H^+]^{-4})}
\]

where \( K_{xy} \) is the equilibrium constant for the \( \text{Fe}_x(\text{OH})_y^{3x-y} \) complex. From Equation 3 it can be seen that the solubility depends on the pH and can, in principle, be calculated from known equilibrium constants. The concentration of soluble iron in a solution saturated with \( \text{Fe(OH)}_3(s) \) is graphically represented in Figure 1 as a function of pH. It can be seen that the solubility of \( \text{Fe(OH)}_3(s) \) is controlled by the various soluble hydroxide complexes. [The polynuclear complexes -- \( \text{Fe}_2(\text{OH})_2^{4+} \) and \( \text{Fe}_3(\text{OH})_4^{5+} \) -- will be ignored, as such complexes are important only in fairly concentrated solutions (above approximately 0.01M).]

Experimental verification of the calculated solubility

\(^1\)Determined at 25°C with I = 1.5
Figure 1. Calculated solubility of Fe(OH)₃ (s) using the equilibrium constants presented by Stumm and Morgan [1981]. Note that Stumm and Morgan omitted Fe(OH)₆ from their calculations. (-) = [Fe³⁺]ₕ (log $K_{sp}$ = -39.1, log $K_{11}$ = -2.19, log $K_{12}$ = -5.67, and log $K_{14}$ = -21.6)
curve as seen in Figure 1, has proven to be difficult. Figure 2 compares the calculated solubility of Fe(OH)$_3$(s) with experimental measurements taken from the literature. Clearly, agreement is poor, although it might be made to appear better depending on the values selected for $K_{13}$ and $K_{sp}$. Furthermore data at high pH values are lacking.

In the pH 6-10 range the solubility might be controlled by the soluble Fe(OH)$_3^-$, however the value of $K_{13}$ is not well established. So little is known and understood about this equilibrium constant that some authors [Stumm and Morgan, 1981; and Martell and Smith, 1982] exclude $K_{13}$ from their solubility calculations as shown in Figure 1. Table II shows different values of $K_{13}$ taken from the literature. An initial objective of this research was to obtain a more reliable value.

**TABLE II**

<table>
<thead>
<tr>
<th>REFERENCE</th>
<th>$K_{13}$</th>
</tr>
</thead>
<tbody>
<tr>
<td>Gayer and Woontner, 1956</td>
<td>2.9<em>10$^{-9}$ to 2.9</em>10$^{-13}$</td>
</tr>
<tr>
<td>Byrne and Kester, 1976</td>
<td>5.3*10$^{-13}$</td>
</tr>
<tr>
<td>Baes and Mesmer, 1976</td>
<td>&lt; 1*10$^{-12}$</td>
</tr>
<tr>
<td>Zafiriou and True, 1980</td>
<td>&lt;&lt;2.4*10$^{-14}$</td>
</tr>
<tr>
<td>Barnum, 1983</td>
<td>1.3*10$^{-12}$</td>
</tr>
</tbody>
</table>

One problem in the experimental verification of $K_{13}$ is the lack of sensitive analytical techniques which determine sub-nanomolar concentrations of iron. Colloidal ferric hydroxide particles are also a problem as they can pass through all but the finest pore size filters and dialysis
Figure 2. Comparison of the calculated solubility (−) of Fe(OH)₃(s) with experimental solubility measurements from the literature. (X)Kuma, et al. [1992], (◊)Kuma, et al. [1993], and (+)Byrne and Kester [1976].
tubing, interfering with the determination of iron.

Another problem is the nature of the ferric hydroxide solid phase. Different solid forms of ferric hydroxide are known: fresh ("active") and aged ("inactive") amorphous ferric hydroxide (Fe(OH)₃·nH₂O), a hypothesized amorphous non-stoichiometric Fe(OH)₂₋ₓₓₒₓₓ₋ₓₓ, goethite (α-FeOOH), akaganeite (β-FeOOH), lepidocrocite (γ-FeOOH), feroxyhyte (δ-FeOOH), haematite (α-Fe₂O₃), and maghemite (γ-Fe₂O₃).

**Amorphous Ferric Hydroxide, Fe(OH)₃·nH₂O**

Amorphous ferric hydroxide, also known as hydrous ferric oxide, changes slowly with age. The "active" form is freshly precipitated, while the "inactive" form is aged. Both crystalline forms are produced by the addition of base to aqueous iron(III). Still further aging, over a period of months or years, results in a slow transformation to goethite and haematite; in some cases akaganeite and lepidocrocite forms on aging [Kolthoff, et al., 1969].

**Non-Stoichiometric Fe(OH)₂₋ₓₓₒₓₓ₋ₓₓ**

Fox [1988] suggested that amorphous ferric hydroxide is non-stoichiometric, as Fe(OH)₂₋ₓₓₒₓₓ₋ₓₓ.

**Goethite, α-FeOOH**

Goethite, a crystalline form of iron(III) hydroxide, is formed by slow hydrolysis of iron(III) salts -- although FeCl₃ results in akaganeite -- or by aging of ferrihydrite
(Fe₅O₃(OH)₉) [Schwertmann and Taylor, 1977]. Goethite can also be formed by the conversion of lepidocrocite in alkali hydroxide or ferrous sulfate solutions [Schwertmann and Taylor, 1972].

**Akaganeite, β-FeO(OH)**

Crystalline akaganeite formation occurs by the hydrolysis of iron(III) in solutions containing chloride or fluoride ions [Sherman, et al., 1982].

**Lepidocrocite, γ-FeOOH**

Lepidocrocite, a crystalline solid, is formed by oxidation of precipitated Fe(OH)₂ (amakinite) at lower pH and temperature and in the absence of iron(III) ions [Kabata-Pendias and Pendias, 1984].

**Feroxyhyte, δ-FeOOH**

Feroxyhyte forms a crystalline solid by violent oxidation of Fe(OH)₂ (amakinite) in very basic solutions [Schwertmann and Taylor, 1977].

**Haematite, α-Fe₂O₃**

Crystalline haematite formation results from the transformation of ferrihydrite or by dehydration of goethite at elevated temperatures [Sherman, et al., 1982].

**Maghemite, γ-Fe₂O₃**

Crystalline maghemite typically is formed by the
oxidation of magnetite (Fe$_3$O$_4$), but thermal dehydration of lepidocrocite can occur [Sherman, et al., 1982].

The various solid forms of the iron(III) minerals have different solubility products as seen in Table III. The literature values for the solubility product of ferric hydroxide vary from $10^{-35.5}$ to $10^{-44}$ [Stumm and Lee, 1960]. Most of the literature values make no mention as to the solid form of iron(III) or the age of the precipitate used. Since the solid forms, and therefore the solubilities, form an intricate network of interconversions, it is important to select the appropriate solubility product for calculations.

**TABLE III**

<table>
<thead>
<tr>
<th>Crystal Form</th>
<th>$K_{sp}$</th>
</tr>
</thead>
<tbody>
<tr>
<td>Amorphous, Fe(OH)$_3$ $\cdot$ nH$_2$O</td>
<td>$3.5 \times 10^{-29}$</td>
</tr>
<tr>
<td>Non-Stoichiometric, Fe(OH)$<em>{2.35}X</em>{0.65}$</td>
<td>$2 \times 10^{-32}$</td>
</tr>
<tr>
<td>Goethite, $\alpha$-FeO(OH)</td>
<td>$9.6 \times 10^{-43}$</td>
</tr>
<tr>
<td>Lepidocrocite, $\gamma$-FeO(OH)</td>
<td>$2.5 \times 10^{-41}$</td>
</tr>
<tr>
<td>Haematite, $\alpha$-Fe$_2$O$_3$</td>
<td>$1.2 \times 10^{-42}$</td>
</tr>
<tr>
<td>Maghemite, $\gamma$-Fe$_2$O$_3$</td>
<td>$3.9 \times 10^{-41}$</td>
</tr>
</tbody>
</table>

Recently, electrochemical methods for the determination of low concentrations of iron(III) in aqueous solutions have been developed. Levels of iron as low as $10^{-9}$-$10^{-10}$M can be determined using adsorptive cathodic stripping voltammetry.

\[ K_{sp} = [Fe^{3+}][OH^-]^{2.35} \]  
\[ X \] is an anion belonging to the solvent and can be emitted from the expression as its concentration is constant.

\[ 2 \text{From Fox [1988]; with } K_{sp} = [Fe^{3+}][OH^-]^{2.35} \]
(CSV). The procedure relies on the interfacial accumulation of the chelate of iron with Solochrome Violet RS (SVRS) on a hanging mercury drop electrode. Iron(III) forms a complex with SVRS at pH values between 4 and 5, while iron(II) does not [Latimer, 1968]. This method offers several advantages such as: detection of iron(III) in the presence of other metal ions, including iron(II), and a detection limit of 0.04µg/L (7*10^{-10}M) iron(III) (calculated from 3 times the noise) [Wang and Mahmoud, 1987].

The purpose of this investigation was to experimentally verify the calculated solubility of iron(III) in the pH 4-12 region using adsorptive CSV.
CHAPTER II

EXPERIMENTAL

APPARATUS

Polarograph

The polarographic equipment used was a PAR 264A Voltammetric Analyzer connected to a PAR303A Static Mercury Drop Electrode. Purge time, deposition time, equilibration time, and scan time were controlled by the potentiostat. The drop size used was "medium", which gave an approximate surface area of 0.015cm$^2$. Solutions were stirred using a teflon coated stirring bar. Borosilicate glass, plastic, or siliconized glass voltammetric cells (EG&G PARC) were used.

The nitrogen line used for purging solutions was equipped with an activated carbon column, a zeolite column, and a 0.20µm filter.

The mercury was examined by atomic emission spectrographic analysis and proved to have no detectible impurities.

Clean Box

To provide a dust free working environment, experiments were carried out in a dry box fitted with a fan to circulate air through an electrostatically charged, high performance
Hammock clean air filter (Filtrete by 3M). The gloves were removed and the holes covered with two layers of the clean air filter material.

**Cleaning Protocol**

All glassware was cleaned by soaking in hydrochloric acid via the following protocol. The glassware was washed in warm tap water with standard laboratory soap (Labetone or Sparkleen), rinsed five times with tap water, five times with deionized water, and placed in a hydrochloric acid bath. Smaller glassware was soaked in 6M HCl for a minimum of four hours; while the larger glassware was soaked in 1.5M HCl for a minimum of 12 hours. After the acid bath, glassware was rinsed five times with deionized water and placed inverted in the clean box to air dry.

**Water**

"Nanopure" Water. Deionized water was used to prepare all solutions and wash the equipment and glassware. Tap water was distilled and passed through a Barnstead NANOpure II Water System, consisting of an organic removal cartridge, two ion exchange cartridges, a submicron filter, and a bacterial removal filter.

Quartz Distilled Water. The preparation of quartz distilled water (Q-water) was accomplished by first distilling deionized water over alkaline KMnO₄ to oxidize any organics, then a second distillation, and finally a sub-
boiling point distillation. The entire apparatus was constructed of quartz.

**Sodium Hydroxide And Hydrochloric Acid**

A 1.075M solution of sodium hydroxide was prepared by diluting 19M (50%) sodium hydroxide (J.T. Baker Chemical Co.) in nanopure water. The solution was standardized by titration with primary standard potassium acid phthalate to the phenolphthalein end point and stored in a plastic container equipped with a soda lime tube to prevent the absorption of carbon dioxide.

A 1.124M solution of hydrochloric acid was prepared by diluting 12M (35%) hydrochloric acid (J.T. Baker Chemical Co.) with nanopure water. The solution was standardized by titration with the standardized sodium hydroxide to the phenolphthalein end point.

**Colloidal Ferric Hydroxide**

A suspension of colloidal, hydrated ferric hydroxide was prepared by the procedure outlined by Brescia, et al. A 2.5M solution of ferric chloride (Mallinckrodt Analytical Reagent) was prepared by dissolving the solid in nanopure water. Two milliliters of the concentrated ferric chloride solution was added to 150mL of nanopure water and boiled to produce the colloidal suspension. The colloidal suspension was dialyzed until there was no longer residual chloride in the rinse solution when tested with a silver nitrate
solution. The concentration of the resulting colloidal suspension was approximately $3.3 \times 10^{-2}$ M in total iron.

**Goethite ($\alpha$-FeO(OH))**

A sample of natural goethite (Ward's Natural Science) was ground to less than 100 mesh and suspended in nanopure water to remove the fines and possible impurities. After 30 seconds, the water was decanted off and the process repeated four more times. The remaining sediments were dried in a 110°C oven for 2 hours.

An X-ray powder diffraction pattern was taken of the sample using a Philips XRG-3000 X-ray diffractometer to confirm the identity of the mineral sample. The diffraction pattern confirmed the identity of the sample as goethite, and also indicated the possible presence of trace amounts of amakinite [($Fe_{0.72}Mg_{0.22}Mn_{0.05})$ $(OH)_2$].

**pH 4-12 Buffers**

A series of buffers covering the pH range 4 through 12 in 0.5 pH increments was prepared. Solutions chosen were acetic acid (pKa = 4.76), PIPES [Piperazine-N,N'-bis(2-ethanesulfonic acid)] (pKa = 6.80), TRIS [Tris(hydroxymethyl)aminomethane] (pKa = 8.30), and CAPS [3-(cyclohexylamino)propanesulfonic acid] (pKa = 10.4). These buffers were chosen to minimize complexation with iron. The acetic acid was obtained from J.T. Baker Chemical Co. and the PIPES, TRIS, and CAPS were obtained from Sigma Chemical
Co. The 1M solutions of hydrochloric acid and sodium hydroxide were used in conjunction with these compounds to prepare the desired buffer solutions. All buffers were prepared and diluted with nanopure water, and filtered through 0.2µm filters before use to remove bacteria. Table IV lists the detailed preparation of each buffer with the resulting pH and ionic strength. The pH values were measured using a Lazar Digital pH meter with a Corning general purpose combination electrode calibrated against pH 4.01, 7.00 and 10.00 standard buffers.

**TABLE IV**

**PREPARATION OF BUFFERS WITH ACTUAL pH's, IONIC STRENGTHS, AND BUFFER SYSTEM**

<table>
<thead>
<tr>
<th>pH</th>
<th>Moles Acid</th>
<th>Moles Base</th>
<th>Actual pH</th>
<th>Ionic Strength</th>
<th>Buffer</th>
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</thead>
<tbody>
<tr>
<td>4</td>
<td>6.622E-2</td>
<td>1.074E-2</td>
<td>3.93</td>
<td>1.074E-1</td>
<td>HOAc</td>
</tr>
<tr>
<td>4.5</td>
<td>2.777E-2</td>
<td>1.074E-2</td>
<td>4.45</td>
<td>1.074E-1</td>
<td>HOAc</td>
</tr>
<tr>
<td>5</td>
<td>1.563E-2</td>
<td>1.074E-2</td>
<td>4.99</td>
<td>1.074E-1</td>
<td>HOAc</td>
</tr>
<tr>
<td>5.5</td>
<td>1.178E-2</td>
<td>1.074E-2</td>
<td>5.49</td>
<td>1.074E-1</td>
<td>HOAc</td>
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<tr>
<td>6</td>
<td>8.428E-3</td>
<td>1.003E-2</td>
<td>6.13</td>
<td>8.428E-2</td>
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<td>6.5</td>
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<td>1.002E-2</td>
<td>6.55</td>
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<td>PIPES</td>
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<td>9.977E-3</td>
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<td>3.813E-2</td>
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<td>1.002E-2</td>
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<td>1.000E-2</td>
<td>8.56</td>
<td>3.736E-2</td>
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<tr>
<td>9</td>
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<td>1.003E-2</td>
<td>8.94</td>
<td>1.648E-2</td>
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<tr>
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<td>8.958E-3</td>
<td>9.999E-4</td>
<td>9.68</td>
<td>9.999E-3</td>
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</tr>
<tr>
<td>10</td>
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<tr>
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<td>9.999E-4</td>
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<td>9.999E-3</td>
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<td>CAPS</td>
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<tr>
<td>12</td>
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<td>9.999E-4</td>
<td>12.12</td>
<td>1.022E-2</td>
<td>NaOH</td>
</tr>
</tbody>
</table>

Supporting Electrolyte

The supporting electrolyte used in the polarographic
analysis was acetate buffer (pH 5.1), prepared by isothermal distillation from acetic acid (Allied Chemical) and ammonium hydroxide. The ionic strength of the supporting electrolyte solution was 0.03.

**Solochrome Violet RS**

The Solochrome Violet RS was purchased from ICN Biomedicals and used without further purification. A stock solution, $1.5 \times 10^{-4}$ M, was prepared fresh daily.

**Standard Iron Solution**

A stock iron solution was prepared by dissolving a weighed quantity of Hach Iron Powder (99.9+%) in perchloric acid (Merck & Co.) with gentle heating and oxidizing to ferric iron with hydrogen peroxide. To produce the standard ferric solution used for the experiment, the stock solution was diluted and sodium citrate (Mallinckrodt Analytical Reagent) added to prevent the precipitation of iron. The concentrations of the iron and citrate in the standard solution were $4.000 \times 10^{-6}$ M and $3.002 \times 10^{-5}$ M, respectively.

**Solubility Measurements**

Three series of solubility measurements were made; one with colloidal iron(III) hydroxide, one with goethite, and one in which aqueous iron(III) perchlorate was added and the precipitate formed in situ. For each equilibration, 30.0 mL of buffer was placed in a 50mL polypropylene centrifuge tube
("Oakridge" type, Nalgene) under sterile conditions. To one
series of buffers was added 1mL of 0.04000M aqueous ferric
perchlorate; to the next was added 1mL of colloidal ferric
hydroxide suspension which contained a total of 0.032 moles
of Fe$^{3+}$ per liter; and to the third was added 0.1g of solid
goethite. At the pH with the highest theoretical iron
solubility, the total ferric iron placed into the tubes
exceeded the total expected concentration of dissolved iron
by approximately one hundred.

The solutions were equilibrated by constant agitation
on a vertical rotating wheel at 14 RPM in a constant
temperature compartment (25°C) for 3 months. This reaction
time is more than adequate to reach equilibrium [Biederman
and Schindler, 1957].

Iron Analysis

Willard and Dean [1950] reported that the single
reductive voltammetric wave of Solochrome Violet RS (Figure
3) was split into two waves in the presence of aluminum
ions. The height of the second wave, which was 0.2V more
negative that the first, was proportional to the aluminum
concentration. The reduction of the aluminum complex of
SVRS occurs at a more negative potential than the free dye,
because the complex is more stable and requires a greater
voltage to bring about reduction. Later studies showed that
SVRS can be used to determine several metal ions, including
iron(III).
The mechanism for the reduction of the SVRS compound at the mercury electrode has been determined with reasonable certainty [Florence and Belew, 1969]. The reduction involves a potential determining 2-electron step giving an unstable hydrazo intermediate, which rapidly disproportionates. Disproportionation yields two amines and the original SVRS azo compound.

\[-\text{N}=\text{N}^- + 2\text{H}^+ + 2\text{e}^- \leftrightarrow -\text{NH}_2\text{NH}_2^-\]  
\[2 -\text{NH}_2\text{NH}_2^- \rightarrow -\text{N}=\text{N}^- + 2 -\text{NH}_2\]  

The reduction of SVRS has a working range of at least pH 3 through 11, however, the reduction potential becomes more negative with increasing pH (i.e. \(E_h=-0.278\text{V}\) at pH 4.30 and \(E_h=-0.573\text{V}\) at pH 9.20) [Florence and Belew, 1969]. While the mechanism for the reduction of the SVRS is well known, the mechanism for the reduction of the iron/SVRS complex is still under investigation.

Figure 3. Structure of Solochrome Violet RS (5-sulfo-2-hydroxybenzene-azo-2-naphthol, C.A.#[2092-55-9]).
This method of adsorptive CSV in the presence of SVRS involves the formation of a surface-active complex of Fe(III)/SVRS, and its interfacial accumulation onto the hanging mercury drop electrode. The complex is then quantified by reductive voltammetric stripping of the adsorbed complex. The current produced by reduction of the surface concentration of the ion complex is proportional to the solution concentration. The complexation and interfacial accumulation of the Fe(III)/SVRS system can be described by the following equations [Wang, et al., 1987]:

\[
\text{Fe}^{3+} + \text{SVRS} \leftrightarrow [\text{FeSVRS}]^{3+} \quad (6)
\]

\[
[\text{FeSVRS}]^{3+} \leftrightarrow [\text{FeSVRS}]^{3+}_{\text{adsorbed}} \quad (7)
\]

Analytical Procedure. The analytical procedure used to determine iron(III) concentrations is similar to that used by Wang and Mahmoud [1987].

Ten milliliters of the supporting electrolyte solution (pH 5.1 acetate buffer) containing 1.5*10^{-6} M of SVRS was plastic tip pipetted into the electrolytic cell, and deaerated by bubbling with nitrogen for 8 minutes. Deaeration was stopped and the nitrogen was passed over the solution surface while the preconcentration potential (usually -0.10V) was applied to a fresh mercury drop with the solution being stirred. Following the 30 second preconcentration period, stirring was stopped. After a 15 second equilibration time, a voltammogram was recorded using differential pulse polarographic stripping, by applying a
negative going linear scan from -0.10V to -1.10V at a scan rate of 10mV/sec. After the background voltammogram was obtained, aliquots of the standard iron(III) solution were added and voltammograms obtained. All analyses employed the standard addition method and all voltammograms were recorded at room temperature.

POSSIBLE SOURCES OF IRON CONTAMINATION

With decreasing concentration of iron(III) to be determined, contamination can prevail to such an extent that the analytical results can be incorrect by orders of magnitude. In the determination of extremely low concentrations of iron(III) the various sources of contamination were investigated.

Water

Because of the low concentrations of iron to be determined, it was important to obtain water which had a lower concentration of iron than that to be determined. The best water that could be obtained contained approximately 10⁻⁸M iron. Both Nanopure water and Q-water gave this result. Even at this low level, adsorptive CSV gives a large iron peak.

While water in this iron concentration range may not give extremely accurate results, accurate results in ultralow concentration determinations of heavy metals are extremely difficult to obtain [Boutron, 1990].
Although any iron impurities in the SVRS would have been subtracted out with the background, it was still considered as a possible source of iron contamination. The amount of iron that would have to be in the SVRS to create the concentrations seen was calculated to be approximately 1% w/w of the SVRS used. This large quantity of impurity seemed unreasonable, therefore the SVRS was not purified further. Wang and Mahmoud [1987] also found that commercial SVRS did not require further recrystallization.

Citrate

Voltammograms were recorded with successive standard additions of $3.4 \times 10^{-5} \text{M} \ (63.88 \text{ng/L})$ sodium citrate to determine if the citrate contained iron impurities or would interfere with the electrochemical method. The data obtained are listed in Table V.

<table>
<thead>
<tr>
<th>Citrate Added (ng/mL)</th>
<th>Current (nA)</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>153.125</td>
</tr>
<tr>
<td>6.12</td>
<td>181.250</td>
</tr>
<tr>
<td>12.24</td>
<td>196.875</td>
</tr>
<tr>
<td>18.36</td>
<td>212.500</td>
</tr>
<tr>
<td>24.48</td>
<td>225.000</td>
</tr>
</tbody>
</table>

The data indicate a linear increase with a slope of $2.86 \text{nA}/(\text{ng/mL})$ and a correlation coefficient of 0.99. The slope indicates the presence of iron in the citrate.
solution. The iron in the citrate solution can be attributed, in part, to the water used to prepare the solution. The increase in peak current is insignificant as small quantities of citrate are used during the analytical procedure.

**Pipet Tips**

Although the pipet tips were acid washed, iron that had not been removed in the acid washing process, can desorb during use. This hypothesis was not tested, as the literature [Mart, 1979; and Gretzinger, et al., 1982] suggests that this is not a problem for the small volumes of dilute samples pipetted.

**Syringes And 0.20μm Filters**

Aliquots of the original buffer solutions were analyzed with and without filtering; any change in the calculated concentration between the filtered and non-filtered aliquots could be due to the adsorption or desorption of iron from either the syringes or the filters. Table VI presents the data collected from the buffers analyzed, including the change in concentration due to filtration. A positive value indicates an increase in iron concentration due to filtration and a negative indicates a decrease. While changes in the iron concentration are apparent, there is not a significant increase or decrease in the total concentration.
TABLE VI

CHANGE IN THE CALCULATED IRON CONCENTRATION FOR BUFFERS WITH AND WITHOUT FILTRATION

<table>
<thead>
<tr>
<th>Buffer pH</th>
<th>([\text{Fe}^{3+}]_0 \text{(M)})</th>
</tr>
</thead>
<tbody>
<tr>
<td>3.93</td>
<td>+ 6.661*10^{-10}</td>
</tr>
<tr>
<td>8.15</td>
<td>+ 5.750*10^{-9}</td>
</tr>
<tr>
<td>12.12</td>
<td>- 2.137*10^{-9}</td>
</tr>
</tbody>
</table>

The changes in concentration of iron due to filtration are very sporadic and no conclusion, except the fact that the syringes and filters have a small impact on the concentration, can be drawn.

Colloids

The 0.2 µm filters are probably too coarse to collect all colloidal iron particles. These colloidal particles could then dissolve in the pH 5.1 acetate buffer, increasing the concentration of dissolved iron. Kennedy, et al. [1974] presented a study on the effects of filter pore-size on the analysis of some metal ions, including iron, in water samples. The study found that approximately 10.5-17.5 µg/L (1.9*10^{-7} to 3.1*10^{-7} M) colloidal iron can pass through a 0.22 µm filter, but are retained on a 0.10 µm filter. The study mentioned that very small amounts of particulates can pass through 0.10 µm filters.

To determine the effects of the dissolution of colloidal iron hydroxide on the analysis, an aliquot of a colloidal ferric hydroxide suspension was placed into an
electrolytic cell containing the supporting electrolyte/SVRS solution. Voltammograms were obtained on the solution as a function of time to determine if the dissolution of the colloidal material was affecting the peak current. Figure 4 is a plot of approximate concentration versus time. The resulting plot shows an approximate linear relationship with a slope of 5.62nA/min (approximately $3.0 \times 10^{-8}$ (mol/L)/nA) and a correlation coefficient of 0.99. This indicates that there may be a significant increase in peak current due to colloidal particles that pass through the 0.20µm filter.

The high pH equilibration solutions may be more affected by the dissolution of colloids. Lengweiler, et al. [1961b] demonstrated that at lower pH's few colloidal particles were found; large size colloidal particles were formed near pH 7 with particle size decreasing through the higher pH range. The pH effect on particle size was evident by the lack of particulate matter retained on the 0.2µm filters of the lower pH range solutions. Large amounts of particulate matter was retained on the filters of the mid-pH range solutions with decreasing amounts retained on the filters of the higher pH solutions.

The presence of colloids may also affect the peak current by interfering with the equilibrium between aqueous Fe$^{3+}$ and SVRS in the solution, or by decreasing the concentration of SVRS below the optimum requirement. If active sites on the colloidal surface were to complex with
Figure 4. Concentration versus time showing the dissolution of colloidal ferric hydroxide. The slope is 5.62 nA/min (about $10^{-7}$ (µg/mL)/nA) with a correlation coefficient of 0.99.
SVRS in the same manner as the aqueous iron, the colloid/SVRS complex could adsorb on the mercury drop increasing the peak current as the complex is reduced. On the other hand, if several SVRS molecules were to react with each colloid particle, reduction of the adsorbed complex would decrease the peak current.

The possible effects of colloid/SVRS complexation were examined by adding increasing increments of colloidal ferric hydroxide suspension, in excess relative to iron solubility, into two series of vessels containing 12mL of the pH 5.1 acetate buffer. One series of vessels was allowed to equilibrate for 2 days, the other for 7 days. When equilibrium is reached, the concentration of dissolved iron should be equal; whereas the amount of colloidal material in the vessels will be increasing with increasing colloidal suspension added. Upon determination of the concentration of iron in the samples, an increase in the concentration can be attributed to the reduction of the colloid/SVRS complex. Figure 5 represents iron concentration determined versus total moles of iron per liter for both the 2 and 7 day equilibrations. The concentrations remained fairly constant with increasing colloidal concentration. This indicates that the peak current is not affected by the presence of colloidal material in the solution.

**Metal Ions**

Wang and Mahmoud [1987] tested several metal ions at
Figure 5. Concentration versus total iron added as colloidal ferric hydroxide. (□) Two day and (+) Seven day equilibration.
50µg/L and found that they do not interfere with the determination of iron(III). Under the conditions of their experiment, they concluded that 50µg/L Al³⁺ created a minor current increase to the iron/SVRS peak. To determine if Al(III) will interfere with the determination of iron concentration, a sample of the supporting electrolyte/SVRS containing iron was spiked with 100µL of 10⁻⁴M Al(NO₃)₃, and a voltammogram obtained (Figure 6). The addition of Al(III) to the sample resulted in the appearance of an additional peak at -0.76V, however, there was no increase in the iron/SVRS peak. From this demonstration it was determined that Al(III) will not interfere with the determination of iron concentrations.

Electrode Frit

When the Ag/AgCl reference electrode is in contact with iron in solution, some of the iron may diffuse into the electrode frit. When a solution of lower iron concentration comes in contact with the frit, the iron can diffuse back out of the frit at a rate which may increase the iron concentration in solution.

A solution containing the normal concentration of iron (1.9*10⁻⁷M) was placed in contact with the electrode for a period of 1 hour. The solution was then removed and the electrodes thoroughly washed with 1% nitric acid and deionized water. A solution of supporting electrolyte/SVRS containing 7.8*10⁻⁸M iron (to ensure visualization of the
iron/SVRS peak) was analyzed as a function of time to determine if iron was diffusing out of the electrode frit.

Figure 6. Differential pulse voltammogram of solution containing supporting electrolyte/SVRS and iron, spiked with Al(III). The addition of Al(III) results in a peak at -0.76V (see current scale in upper right).

Figure 7 shows the resulting graph of the data. The data show a linear increase in current with time. The slope of
Figure 7. Approximate concentration of iron as a function of time to determine the extent of contamination resulting from iron diffusion out of the electrode frit.
this line is 0.2414nA/min with a correlation coefficient of 1.00. While there is an increase in current with time, the increase would not significantly alter the peak current of iron in solution. If iron can diffuse out of the frit, perhaps some of the Ag⁺ can diffuse out of the frit as well. Wang and Mahmoud [1987] did not consider this in their analysis of the effects of other metal ions on the iron/SVRS peak. A sample solution was spiked with 5×10⁻⁷M Ag⁺ to determine if it would interfere with the iron/SVRS peak. The addition of the silver to the sample showed no increase in the peak current, indicating that silver was not responsible for an increase in peak current with time.

**Purge Gas**

To determine the effects on the peak current due to nitrogen gas, voltammograms were obtained with and without solution stirring and nitrogen purging. When the solution was not purged with nitrogen, nitrogen was still passed over the solution to prevent oxygen from entering the system. The data obtained from the voltammograms was plotted, and the resulting slopes obtained indicate the increases in current with time. Table VII lists the various conditions and the resulting increases in current. The data suggest that the nitrogen gas has a significant effect on the peak current, even when the solution is not purged it appears that iron is entering the system possibly a result of the nitrogen gas passing over the solution. To ensure these
results were indeed an effect of the nitrogen, two separate test solutions were analyzed by recording the peak current increase as a function of varied purge times. The results found the current increases to be 1.977 and 1.948 nA/min, indicating the nitrogen gas may be a possible source of iron contamination.

TABLE VII

INCREASES IN CURRENT WITH TIME AT VARIOUS CONDITIONS INVOLVING SOLUTION STIRRING AND NITROGEN PURGING

<table>
<thead>
<tr>
<th>Conditions</th>
<th>Current Increase</th>
</tr>
</thead>
<tbody>
<tr>
<td>Stir/Purge</td>
<td>0.8984 nA/min</td>
</tr>
<tr>
<td>Stir/ --</td>
<td>0.4911</td>
</tr>
<tr>
<td>-- / Purge</td>
<td>0.9961</td>
</tr>
<tr>
<td>-- / --</td>
<td>0.3906</td>
</tr>
</tbody>
</table>

To rule out the possible effects of the stir bar causing the release of iron from the cell surface by abrasive action, voltammograms were obtained without the presence of a stir bar in the solution. When the solution was analyzed without nitrogen purging or passing over the surface, there was no increase in the iron/SVRS peak current. This ruled out any effects of iron increase due to abrasive effects from the stir bar.

It was speculated that the increase in peak current due to purge gas could be a result of dust particles from the activated carbon or zeolite columns in the gas train, which pass through the 0.20 µm filter or from gaseous forms of iron in the nitrogen. While the existence of gaseous forms of
iron(III) are rare, ferric chloride (FeCl₃) and ferric carbonyl (Fe(CO)₅) can exist in the gaseous phase. The effects of gaseous forms of iron is pure speculation, and the most probable effect is the existence of dust particles entering the solution.
CHAPTER III

RESULTS

DIFFERENTIAL PULSE POLAROGRAPHY

Instrumental Characteristics

Prior to the collection of analytical results, voltammograms were obtained to ensure the system was functioning as previously reported. Figure 8 shows voltammograms for SVRS in acetate buffer in the absence (a) and presence (b,c) of iron(III).

When the SVRS-containing solution was stirred for 30 seconds with the electrode held at -0.1V prior to the scan, a large cathodic peak associated with the reduction of the adsorbed SVRS is observed at -0.46V (a). A smaller peak is observed at -0.36V, 0.10V more positive than the SVRS reduction peak. Florence and Belew [1969] attribute this peak to a 1-electron reduction process. When the same experiment is conducted in the presence of iron(III) (b), an additional peak associated with the reduction of the adsorbed Fe(III)/SVRS complex is observed at -0.60V. The peak height of the iron/SVRS complex increases if the deposition potential is changed to -0.50V (c). The increased peak height can be explained by the reduced competition by the uncomplexed SVRS for sites on the hanging
mercury drop. To increase analytical response, all voltammograms for the determination of ultratrace levels of iron were obtained using a -0.50V preconcentration potential.

\[ -0.46V \]

\[ -0.36V - 0.60V \]

\[ 100nA \]

\[ -0.10V - 1.10V \]

\[ -0.50V - 0.70V \]

\[ -0.60V \]

Figure 8. Voltammograms obtained in the differential pulse mode for SVRS containing solutions in the absence (a) and presence (b,c) of added iron.

To ensure that the instrumentation would give a linear response over the concentration range required, voltammograms were recorded with successive standard additions of 2.11\( \mu \)g/L (3.773*10^{-4}M) iron. Figure 9 shows linear scan voltammograms in the differential pulse mode for
iron solutions of increasing concentration, 2.11-8.44µg/L. Figure 10 depicts the peak current versus concentration of iron added. The resulting plot gives a straight line with a correlation coefficient of 0.99.

These data indicate a linear relationship between the response (peak current) and the concentration of added iron over the required concentration range.

The parameters used in the study include: the use of a
Figure 10. Analysis of background iron by standard addition for differential pulse polarography. Extrapolation determined background iron to be $2.465 \times 10^{-5}$ M, with a correlation coefficient of 0.99.
pH 5.1 acetate buffer as the supporting electrolyte; SVRS concentration of $1.5 \times 10^{-6}$ M; "medium" mercury drop size; and a 30 second deposition time. These parameters were thoroughly examined by Dean and Bryan [1957]; Florence and Belew [1969]; Wang and Mahmoud [1987]; Wang, et al. [1987]; and van den Berg, et al. [1991].

**Analysis Of Samples**

Portions of the equilibrated iron/buffer solutions were removed from the plastic centrifuge tubes and filtered through 0.2µm syringe filters (Rainin Instrument Co.) to remove any colloidal material. An aliquot of the filtered sample was placed into the electrolytic cell containing the buffer/SVRS solution. Voltammograms were obtained for the solution and successive standard additions of iron. The resulting data were plotted as current versus concentration of iron added. The concentration of iron in the sample was determined from the x-intercept of the graph.

Due to the ubiquitous nature of iron, it was important to accurately determine the background concentration of iron present in the buffer/SVRS system. Prior to each analysis, background voltammograms were obtained and the background iron concentration calculated. The background concentration was then subtracted from the concentration determined for the equilibrated pH 4-12 buffers.
Analytical Results. The concentrations of dissolved iron in the buffer solutions equilibrated with a) excess aqueous ferric perchlorate, b) colloidal ferric hydroxide, and c) goethite are tabulated in Table VIII, and the data are plotted in Figures 11, 12, and 13. For comparison, these plots also show the iron solubility as calculated from equilibrium theory.

<table>
<thead>
<tr>
<th>pH</th>
<th>(\log [\text{Fe}^{3+}]_{\text{aq}}) (moles/L)</th>
<th>(\text{Fe(ClO}_4)^{3-}) (aq)</th>
<th>Colloidal</th>
<th>Goethite</th>
</tr>
</thead>
<tbody>
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<td>3.93</td>
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<td>-6.20</td>
<td>-5.85</td>
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</tr>
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</tr>
<tr>
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<td>-5.88</td>
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</tr>
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</tr>
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<td>-6.61</td>
</tr>
<tr>
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<tr>
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<td>-5.73</td>
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<td>N/A</td>
</tr>
<tr>
<td>12.12</td>
<td>-5.92</td>
<td>-5.60</td>
<td>-6.16</td>
<td>-6.16</td>
</tr>
</tbody>
</table>
Figure 11. Comparison of calculated solubility (-) with experimental concentrations (+) of iron in solutions equilibrated with ferric perchlorate. (log $K_{sp} = -39.1$, log $K_{11} = -2.19$, log $K_{12} = -5.67$, log $K_{13} = -12.3$, and log $K_{14} = -21.6$)
Figure 12. Comparison of calculated solubility (-) with experimental concentrations (◊) of iron in solutions equilibrated with colloidal ferric hydroxide. (log $K_\text{sp} = -39.1$, log $K_{11} = -2.19$, log $K_{12} = -5.67$, log $K_{13} = -12.3$, and log $K_{14} = -21.6$)
Figure 13. Comparison of calculated solubility (-) with experimental concentrations (Δ) of iron in solutions equilibrated with goethite. (log $K_{sp} = -39.1$, log $K_{11} = -2.19$, log $K_{12} = -5.67$, log $K_{13} = -12.3$, and log $K_{14} = -21.6$)
Wang and Mahmoud [1987] stated that the sensitivity of the procedure can be increased by using fast scan d.c. polarographic adsorptive CSV.

In an effort to improve the determination of iron concentrations in the pH 6-10 range several methodological procedures were altered to reduce the possibility of iron contamination. To prevent any adsorptive/desorptive losses of iron to glass surfaces, glassware was strictly avoided; except for the electrolytic cell, which was silanized with siliconizing fluid (EG&G PARC) to pacify the surface. Also, all solutions, including the pH 5.1 buffer and SVRS solutions, were prepared using Q-water in acid washed plastic containers.

Instrumental Characteristics

When a SVRS containing solution was analyzed, it was noticed that the cathodic peak associated with the reduction of the adsorbed SVRS was observed at -0.52V. The same experiment carried out in the presence of iron produced a peak at -0.81V, corresponding to the reduction of the iron/SVRS complex (Figure 14).

The shift to more negative potentials could not be explained, however, spiking the solution with iron or SVRS resulted in increases in their respective peaks. As stated earlier, changing the initial potential increases the peak
current as a result of reduced competition for sites on the mercury drop. The initial potential for the polarographic analysis was begun at -0.48 V and terminated at -1.10 V with a scan rate of 100 mV/sec. Analysis of the optimal deposition time proved 45 seconds gave the optimal response.

With the new parameters, voltammograms were obtained to determine the background levels of iron in the water.

Figure 14. Voltammogram of fast scan (100 mV/sec) d.c. polarography.
Figure 15 is a graphical representation of the peak current versus concentration of added iron. The data points indicate a linear response with a correlation coefficient of 0.98. Extrapolation of this line to the x-intercept results in a background iron concentration of $5.916 \times 10^{-8}$ M. Determinations of background iron in Q-water by this method were repeated several times, all resulting in similar concentrations of iron. The concentration of background iron had not improved by this method; therefore, determination of iron concentrations in the pH 6-10 range was not affected.

**Summary.** Attempts to lower the background levels of iron were not successful as concentrations below about $10^{-8}$ M iron could not be attained. This is approximately the level shown by Wang and Mahmoud [1987] and van den Berg, et al. [1991]. Also, it is approximately the same level determined by Barnum in purified water using a kinetic analysis in which iron(III) catalyzed the oxidation of N,N-Dimethyl-p-phenylenediamine by hydrogen peroxide [Hirayama and Unohara, 1988].

The signal/noise ratio in both the adsorptive CSV and fast scan differential pulse experiments for the iron peak is still quite high, even at $10^{-8}$ M iron.

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1. Personal communication.
Figure 15. Analysis of background iron by standard addition for fast scan d.c. polarography. Extrapolation determined background iron to be $5.916 \times 10^{-8}$ M, with a correlation coefficient of 0.98.
CHAPTER IV

DISCUSSION

CONCLUSIONS

Fox [1988] discussed the existence of a non-stoichiometric \( \text{Fe(OH)}_{2.35}(X)_{0.65} \). Fox determined the solubility of colloidal ferric hydroxide by dialysis followed by atomic absorption. The results were combined with literature values and plotted as \( p_a \text{Fe(III)} \) versus pH, where \( a \) represents activity. The plot resulted in a straight line \( (r = 0.998) \) with the least squares equation:

\[
-\log a_{\text{Fe(III)}} = 2.35pH - 1.17
\]  

(8)

Rearrangement using \( K_w = 10^{-14} \) yielded:

\[
a_{\text{Fe(III)}}/a_{\text{OH}^-}^{-2.35} = K_{sp} = 10^{-31.7}
\]  

(9)

This equation implies a solid phase with the formula \( \text{Fe(OH)}_{2.35}X_{0.65} \).

Figure 16 shows the calculated solubility of \( \text{Fe(OH)}_{2.35}X_{0.65} \) versus pH using Fox's solubility product, \( K_{sp} = [\text{Fe}^{3+}][\text{OH}^-]^{2.35} = 10^{-31.7} \). The plot predicts iron concentrations much larger than those expressed in the literature as well as those found in this work. These data are compared in Figure 17. The experimental values from this study are approximately two to four orders of magnitude
Figure 16. Comparison of the solubility of Fe(OH)$_{2}$.35X$_{0.65}$ calculated (−) using the solubility product proposed by Fox [1988]. The calculated solubility predicts Fe(OH)$_{2}$.35X$_{0.65}$ will dissolve in excess base, which is not observed.
Figure 17. Comparison of the solubility of Fe(OH)$_{2.35}$X$_{0.65}$ calculated (-) using the solubility product proposed by Fox [1988], with the experimentally measured solubility found in this study. (□) Aqueous ferric perchlorate, (+) Colloidal ferric hydroxide, and (◇) Goethite.
lower than those postulated by Fox in the higher pH region. For this reason it was determined that the results expressed by Fox would not be pursued further.

Barnum [1983] presented an empirical correlation which enables the prediction of equilibrium constants and standard free energies of formation for metal mononuclear and polynuclear hydroxy complexes. Barnum found the following empirical correlation between the standard free energy of formation for hydroxy complexes and the number of hydroxy groups:

$$\Delta G^\circ_f \{M(OH)_y\} = \Delta G^\circ_f \{M\} + By + Cy^2 + D/y$$  \hspace{1cm} (10)

For purposes of interpolating or extrapolating to unknown equilibrium constants this equation was converted into a linear form by combining with the standard free energy of the equation:

$$x\text{ M}^{n+} + y\text{ H}_2\text{O} \leftrightarrow M_x(\text{OH})_{y-x}^{-y} + y\text{ H}^+$$  \hspace{1cm} (11)

$$\Delta G^\circ = \Delta G^\circ_f \{M(\text{OH})_y\} - \Delta G^\circ_f \{M\} - y\Delta G^\circ_f \{\text{H}_2\text{O}\}$$  \hspace{1cm} (12)

Combining these with $\Delta G = -RT\ln K$ gives:

$$U\{M(\text{OH})_y\} = \Delta G^\circ_f \{\text{H}_2\text{O}\} - D/y^2 - (2.303RT\ln K_y)/y$$  \hspace{1cm} (13)

where $y$ is the number of hydroxyl groups in the complex and $D = 8.37\text{ kJ/mole}$ for divalent and trivalent metal ions. When $U\{M(\text{OH})_y\}$ is graphed as a function of $y$ for metal ions with step-wise hydrolysis, a straight line is produced. These calculations work well for 24 of the metal ions investigated. In the four cases of metal ions that do not fit the empirical correlation, all but scandium(III) can be
traced to variations in the structure of the hydroxide complex. Barnum's estimated equilibrium constant of Fe(OH)$_3$ is $K_{13} = 10^{-11.89}$ which agrees with the value ($<10^{-12}$) presented by Baes and Mesmer [1976].

The suggestion in recent literature that the value of $K_{13}$ is significantly lower than previously reported does not fit Barnum's empirical correlation. The values for the equilibrium constants of Fe(OH)$_{2+}$, Fe(OH)$_3^+$, and Fe(OH)$_4^-$ have been thoroughly investigated in the literature and the values presented do not vary significantly. Figure 18 shows several literature values of the equilibrium constants graphed as $U\{M(OH)_y\}$ versus $y$. The correlation coefficient for the data is 0.96, and removal of the literature values for $K_{13}$ produced a correlation coefficient of 1.00.

The disagreement between Barnum's correlation and the more recent literature values of $K_{13}$ could be the result of possible structural changes in one or more of the iron(III) hydroxy complexes, which would affect the free energies of formation. Possible changes that could occur are: a shift from high spin to low spin, a change from octahedral to tetrahedral coordination, or the formation of an oxo complex such as O=Fe(OH)(H$_2$O)$_4$ (octahedral) or O=Fe(OH)(H$_2$O)$_2$ (tetrahedral).

The idea that Fe$^{3+}$, a d$^5$ metal ion, could exist in a low spin state is highly unlikely as it has a large exchange stabilization energy for its half-filled d subshell [Huheey,
Figure 18. Graphical representation of $U\{M(OH)\}$ versus $y$ for literature values of the equilibrium constants of the iron(III) hydroxy complexes.
Iron(III) is high spin in nearly all its complexes, except those with extremely strong ligands, such as [Fe(CN)_4]^{3-}, [Fe(bipy)_3]^{3+}, and [Fe(phen)_3]^{3+}. While octahedral iron(III) complexes can exhibit spin crossover behavior -- from the high spin configuration, \(^6\text{A}_{1g}\), to the low spin configuration, \(^2\text{T}_{2g}\) -- most complexes have a trigonally distorted octahedral configuration with six sulfur atoms [Cotton and Wilkinson, 1988], which the complexes in question do not have.

An ion in an octahedral hole is always at least as stable, and usually more stable, than the same ion in an equivalent tetrahedral field. The increased stability in octahedral fields indicates that iron(III) hydroxy complexes are unlikely to undergo structural changes from an octahedral to a tetrahedral structure.

Iron(III) complexes have the ability to form oxo complexes, although they typically form polymeric complexes such as the \(\mu\)-oxo dimer:

\[
2[\text{Fe(OH)(H}_2\text{O)}_5]^{2+} \leftrightarrow [(\text{H}_2\text{O})_5\text{FeOFe(H}_2\text{O)}_5]^{4+} + \text{H}_2\text{O} \quad (14)
\]

Although several possible structural changes can occur, none of them appear to be reasonable explanations for the disagreement between recent literature values of \(K_{13}\) and Barnum's empirical correlation.
FUTURE WORK

Future work in this area should begin with identifying the source of iron in the water supply and reducing background levels to enable more precise measurements of "ultratrace" concentrations of iron(III).

While the method in this study has a detection limit of $7 \times 10^{-10} \text{M}$, perhaps another method with a similar, or lower, detection limit could be employed. Another possible method is the kinetic method presented by Hirayama and Unohara [1988]. Regardless of the method, colloidal ferric hydroxide should be strictly avoided. Methods such as the use of smaller sized filters or dialysis would be useful to obtain the desired results.
CHAPTER V

REFERENCES

LITERATURE CITED


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