Mossbauer Spectroscopy, X-ray Diffraction and IR Spectroscopy of Oxide Precipitates Formed from FeS04 Solution

Musić, Svetozar; Czako-Nagy, I.; Popović, Stanko; Vertes, A.; Tonković, M.

Source / Izvornik: **Croatica Chemica Acta, 1986, 59, 833 - 851**

Journal article, Published version Rad u časopisu, Objavljena verzija rada (izdavačev PDF)

Permanent link / Trajna poveznica: <https://urn.nsk.hr/urn:nbn:hr:217:416049>

Rights / Prava: [In copyright](http://rightsstatements.org/vocab/InC/1.0/) / [Zaštićeno autorskim pravom.](http://rightsstatements.org/vocab/InC/1.0/)

Download date / Datum preuzimanja: **2024-05-04**

Repository / Repozitorij:

[Repository of the Faculty of Science - University o](https://repozitorij.pmf.unizg.hr)f [Zagreb](https://repozitorij.pmf.unizg.hr)

CCA-1691

YU ISSN 0011-1643 UDC 543.42 Original Scientific Paper

Mossbauer Spectroscopy, X-ray Diffraction and IR Spectroscopy of Oxide Precipitates Formed from FeS04 Solution

S. Music, I. Czak6-Nagy, S. Popovic, A. Vertes*, and M. Tonkovic*

Ruder Boskovic Institute, P. 0. Box 1016, 41001 Zagreb, Croatia, Yugoslavia

and

*Laboratory of Nuclear Chemistry, Lorand-Eotvos University, 1088-Budapest, Hungary

Received February 19, 1986

Iron oxyhydroxides and oxides were precipitated from FeS04 solution at low oxygen content. The composition and structure, stoichiometry, particle size and nuclear magnetism of the precipitates were studied using Mössbauer spectroscopy, X-ray diffraction and IR spectroscopy. The standard iron oxyhydroxides and oxides were also characterized using the same instrumental techniques. The results have indicated a strong dependence of the chemical composition and structure of the precipitates on the $[FeSO₄] / [NaOH]$ concentration ratio. α -FeOOH of poor crystallinity was isolated at low pH values. α -Fe₂O₃ was formed by internal crystallization of α -FeOOH. At 90° C, a mixture of $Fe(OH)_2/Fe(OH)_3$ was transformed, with the time of heating, to nonstoichiometric Fe₃O₄, α -FeOOH and further to α -Fe₂O₃. Nonstoichiometric Fe_9O_4 was the final precipitation product in su-spension with the $[Fe^{2+}] / [NaOH]$ stoichiometric ratio. The importance of these results for the corrosion science of steel in the pre- sence of sulfates has been discussed.

INTRODUCTION

Iron hydroxides, oxyhydroxides and oxides, as well as the ferrite oxides are typical components of the rust formed during the corrosion of iron and its alloys. The chemical and physical properties of iron oxides as corrosion products are strongly dependent on the physico-chemical conditions during the corrosion process. The formation of iron oxide rust creates a great engineering problem in many industries. Therefore, it is not surprising that corrosion engineers have shown a continuous interest in the achievements of iron oxide chemistry.

In the last 50 years a great number of papers on the precipitation of iron oxides have been published (corrosion, soil science, pigments, magnetic tapes, etc.). Also, several review papers with a large body of literature citations have appeared. ¹-s

The chemical composition and structure of precipitated iron oxides greatly depend on the iron salt concentration, the nature of anion, pH, temperature, time of aging and other topochemical parameters. The precipitation of iron oxides from Fe(III)-salt solutions has been studied to a much greater extent than the precipitation from Fe(II)-salt solutions. Iron oxides can be precipitated by adding a base to Fe(III)-salt solution or by forced hydrolysis at elevated temperatures. Fe³⁺ ions are characterized by high hydrolyzability. Precipitation of the solid phase without alkali addition can be easily achieved even at room temperature after a proper time of aging. The hydrolysis of Fe3+ ions with nitrate, chloride or perchlorate anions can be described as a sequence of stages for the formation of hydroxy complexes, hydroxy polymers and hydrous oxides as the end product. The nature of the anion present in the Fe3+ solution affects strongly the composition and structural properties of the precipitate formed.

The precipitation of iron oxides from $Fe(II)$ -salt solutions is strongly dependent on the oxidation/reduction conditions in a solution. Iron(Il) hydroxide will precipitate, upon the addition of alkali to a $Fe²⁺$ solution, when the solubility product constant of $Fe(OH)_2$ is exceeded. In the presence of oxidative agents (O_2 , atmospheric O_2 , H_2O_2), mixed Fe^{2+}/Fe^{3+} hydroxides, and different oxyhydroxides and oxides are formed. Studies on the precipitation of oxides from Fe(II)-salt solutions can be related to some natural processes (corrosion of iron, soil genesis, etc.). The usefulness of these studies increases as the conditions in laboratory experiments approach those in nature.

In this work the precipitation of iron oxides from $FeSO₄$ solution was investigated. Since the precipitates obtained were of a complex nature, their phase analysis was performed using Mossbauer spectroscopy, X-ray diffraction and IR spectroscopy. The combination of these techniques made it possible to follow the changes in the composition and structure, stoichiometry, particle size and nuclear magnetism of the precipitate formed. On the basis of the results obtained in this study and those of other researchers the mechanism of the formation of iron oxides from FeS04 solution and their structural transformations are discussed. The contribution of such investigations to the corrosion science is also pointed out. The sulfate salt of iron(II) was chosen for our experiments, since many corrosion processes involve a sulfate-rich medium. $9-16$ The influence of sulfate ions on the composition and structure of corrosion products was observed.

EXPERIMENTAL

Standard Iron Oxide Samples

The following iron oxyhydroxides and oxides were used to record their standard Mössbauer spectra, diffraction patterns and IR spectra:

- a -FeOOH · nH₂O, precipitated from FeSO₄ solution and after isolation additionally heated,
- β -FeOOH, precipitated by slow hydrolysis of FeCl₃ solution at room temperature,
- $-$ y-FeOOH, precipitated from FeCl₂ solution,
- a -Fe₂O₃, produced by PFIZER Co.,
- $-y$ -Fe₂O₃, prepared by oxidation of magnetite at elevated temperature,
- $-$ Fe₃O₄ (nonstoichiometric), produced by PFIZER Co., and
- $-$ Fe₃O₄ (stoichiometric), formed as a corrosion product.

Precipitation of Iron Oxides from FeS04 Solution

AnalaR grade FeSO₄ · 7H₂O, NaOH, H₂SO₄, ethanol and bidistilled water were used. Fine grained FeSO₄ · 7H₂O was kept in oxygen free atmosphere to prevent the Fe²⁺ oxidation into Fe³⁺. 0.1 M or 0.2 M NaOH solutions $(f = 1.0000)$ were used. The solutions were prepared by mixing given volumes of NaOH (or $H₂SO₄$) solution and bidistilled water (not deaerated). After addition of a proper amount of $F \in SO_4$.
 $T H_2O$ salt, the flasks were vigorously shaken. Then the flasks were closed with glass stoppers and heated at 90 °C for different tim solutions used for precipitation of iron oxides is shown in Table I. In the systems

Chemical Composition of the Solution used for Precipitation of Iron Oxides

* To each solution 5.5610 g of $FESO₄ \cdot 7H₂O$ was added.

with no NaOH addition or with a small quantity of NaOH, the yields of the precipitate were small in relation to the total quantity of Fe(II). Yields of the preci-
pitate are increased with the increase of NaOH in the solution.

Instrumentation

Mössbauer spectra were recorded using a constant-acceleration spectrometer of standard design combined with a multichannel analyzer. The spectrometer was calibrated using α -Fe foil, and a ⁵⁷Co/Pd source was used. The spectra were recorded at room temperature or liquid nitrogen temperature in transmission geometry. All isomer shifts are given with respect to metallic iron.

The X-ray powder diffraction measurements were carried out at room tem-
perature using a counter diffractometer with monochromatized $CuKa$ radiation. Iron oxides and oxyhydroxides were easily and indisputably identified according to the data in the JCPDS Powder Diffraction File (card no. $13-534$ for α -Fe₂O₃, 4-755 for γ -Fe₂O₃, 19-629 for Fe₃O₄, 17-536 for α -FeOOH and 8-98 for γ -FeOOH). The infrared spectra were recorded using a Model 580B Perkin-Elmer spectrophoto- meter. The specimens were pressed in KBr disks.

836 S. MUSIC ET AL.

EXPERIMENTAL RESULTS

Mossbauer Spectroscopy

The knowledge of standard Mössbauer spectra and the corresponding Mössbauer parameters is very important in the analysis of different iron oxides. However, the shape of a measured spectrum and its parameters can sometimes differ significantly from that known as the standard spectrum. Mössbauer spectrum is highly influenced by poor crystallinity, nonstoichiometry, crystal defects, etc.

In this work the standard Mössbauer spectra of Fe oxyhydroxides and oxides were also measured and the corresponding Mössbauer parameters obtained by computer evaluation are shown in Table II. The origin of the samples is given in the experimental section.

TABLE II

s1Fe Mossbauer Parameters of Some Fe Oxyhydroxides and Oxides Obtained at Room Temperature

* Isomer shifts are given relative to alpha iron.

** Errors: \pm 0.005 mm/s and \pm 1 kOe.

*** Q_1 , due to the presence of FeOOH.

[•] In the presence of magnetic hyperfine splitting $\Delta = \Delta_{12} - \Delta_{56}$.

At room temperature, goethite $(\alpha$ -FeOOH) should be characterized with a sextet of lines. However, many samples of α -FeOOH generate spectra of a poor quality with shapes that deviate considerably from the Lorentzian. Also, partially relaxed Mössbauer spectra of the α -FeOOH with intermediate particle size have asymmetrically broadened lines and the values of the internal magnetic field are decreased.

Lepidocrocite (y-FeOOH) indicates its paramagnetic behavior at room temperature, and also at 77 K.

The Mössbauer spectrum of hematite $(\alpha - Fe_2O_3)$ is characterized by a well shaped sextet with $H_{5/2} = 523$ kOe at room temperature. This sextet may

collapse into a doublet at room temperature, if α -Fe₂O₃ consists only of very fine particles (superparamagnetism phenomenon). The superparamagnetic behavior is also characteristic of other Fe oxyhydroxides and oxides.

Maghemite (γ -Fe₂O₃) prepared by the oxidation of Fe²⁺ in Fe₃O₄ is characterized by a sextet of lines. The line-width of γ -Fe₂O₃ ($\Gamma = 0.458$ mm/s) is greater than that of α -Fe₂O₃ ($\Gamma = 0.288$ mm/s), since the Fe³⁺ ions in γ -Fe₂O₃ are distributed between tetrahedral and octahedral sites.

Nonstoichiometric and stoichiometric magnetite ($Fe₃O₄$) are characterized by two sextets at room temperature. The outer sextet of $Fe₃O₄$ corresponds to the tetrahedral sites of Fe^{3+} , and the inner sextet to octahedral sites of (Fe^{2+}/Fe^{3+}) . The nonstoichiometry of $Fe₃O₄$ results in an increased intensity of the peaks of the outer sextet. Table II. shows an increase of the internal magnetic fields in the order:

$$
\text{Fe}_3\text{O}_4 \text{ (stoich.)} \rightarrow \text{Fe}_3\text{O}_4 \text{ (nonstoich.)} \rightarrow \gamma \text{-Fe}_2\text{O}_3
$$
\n
$$
M_1, M_2 \qquad M_1, M_2 \qquad M_1
$$

In the interpretation of the Mossbauer spectra of oxides precipitated from FeS04 solutions, the results and observations of other researchers were also $used.^2,6,17-18$

The Mössbauer spectroscopic results obtained for iron oxide precipitates, prepared on the basis of experimental data shown in Table I., can be summarized as follows.

Samples S-1 and S-2 were precipitated from 0.1 M $\text{FeSO}_4 + 0.02$ M H_2SO_4 solution, and the corresponding Mössbauer spectra recorded at room temperature are shown in Figure 1. The Mössbauer spectrum shown in Figure 1a (sample S-1) is characterized by the central quadrupole doublet. For the same sample the sextet of lines with an internal magnetic field of \sim 490 kOe is measured at liquid nitrogen temperature. In the Mossbauer spectrum of sample S-2 (Figure lb), the appearance of a very poor sextet can be observed. The Mossbauer spectroscopic results indicate the presence of superparamagnetic goethite of poor crystallinity. The chemical composition of these samples can be approximately describes as α -FeOOH \cdot nH₂O.

Sample S-3 was prepared by precipitation from 0.1 M FeSO₄ solution without any NaOH or H_2SO_4 addition. The time of heating (at 90° C) was 48 hours. An internal magnetic field of 382 kOe was measured at room temperature. The Mossbauer spectrum shown in Figure 2 has the form of an asymmetric sextet. More precisely, an asymmetric sextet at room temperature can be considered as a superposition of several sextets. At liquid nitrogen temperature a single sextet was measured with Mossbauer parameters corresponding to α -FeOOH. For shorter times of heating (3 and 24 hours), a broad asymmetric sextet with a central quadrupole doublet was obtained. The central quadrupole doublet disappeared at liquid nitrogen temperature, due to superparamagnetism. After 24 hours of heating, the total area under the peaks of the central quadrupole doublet decreased up to $2^0/6$. Also, a tendency of increase of the internal magnetic field (RT) with time of heating is observed:

363 kOe \rightarrow 378 kOe \rightarrow 382 kOe

(3 hrs.) (24 hrs.) (48 hrs.)

Figure 1. Mössbauer spectra of samples S-1 (Figure 1a) and S-2 (Figure 1b) recorded at room temperature.

Figure 2. Mössbauer spectrum of sample S-3 recorded at room temperature.

Samples S-4 and S-5 were precipitated from 0.1 M $FeSO₄ + 0.002$ M NaOH solution. The Mössbauer spectra of samples S-4 and S-5 have shown that these precipitates contain only α -FeOOH. The central quadrupole doublet was not observed in the spectrum recorded at room temperature. Sample S-5 is characterized by $H_{5/2} = 376$ kOe at room temperature and $H_{5/2} = 513$ kOe at liquid nitrogen temperature.

Samples S-6 and S-7 were precipitated from 0.1 M $FeSO₄ + 0.004$ M NaOH solution. The Mossbauer spectra of these samples were characterized by two magnetic splitting components, M_1 and M_2 . The room temperature Mössbauer spectrum of sample S-7 is shown in Figure 3. For this sample, M_1 (α -Fe₂O₃) = $= 521$ kOe and M_2 (α -FeOOH) = 382 kOe were measured at room temperature. Sample S-6 contained $\sim 8\%$ α -Fe₂O₃ and with prolonged time of heating (48) hours at 90 °C) this value was increased to 20% α -Fe₂O₃.

Figure 3. Mossbauer spectrum of sample S-7 recorded at room temperature.

Samples S-8, S-9 and S-10 were prepared by precipitation from 0.1 M $F \in SO_4 + 0.075$ M NaOH solution. The corresponding times of heating at 90 °C were 4, 24 and 48 hours, respectively. The Mössbauer spectra of these samples are shown in Figure 4. The spectrum shown in Figure 4a can be ascribed to nonstoichiometric $Fe₃O₄$ with a significant loss of $Fe²⁺$ ions.

Sample S-9 is a mixture of nostoichiometric $Fe₃O₄$ (36%) and α -FeOOH (64⁰/₀). In sample S-10, a mixture of 90⁹/₀ α -FeOOH and 10⁰/₀ α -Fe₂O₃ was determined.

Samples S-11 and S-12 were precipitated from 0.1 M $\text{FeSO}_4 + 0.015$ M NaOH solution. At room temperature, the Mossbauer spectrum of sample S-11 consisted of three sextets and a central quadrupole doublet. The areas under the spectrum components corresponded to $Fe₃O₄$ (40⁰/o), α -FeOOH (40⁰/o) and to the central quadrupole doublet (20%). This doublet ($\delta = 0.36$ mm/s, $\Delta = 0.65$ mm/s) was also present ($\delta = 0.48$ mm/s, $\Delta = 0.70$ mm/s) at liquid N₂ temperature. The Mossbauer spectrum did not make it possible to distinguish

Figure 4. Mössbauer spectra of samples S-8 (Figure 4a), S-9 (Figure 4b) and S-10 (Figure 4c) recorded at room temperature.

between γ -FeOOH and the superparamagnetic α -FeOOH. In sample S-11 the presence of γ -FeOOH was detected by IR spectroscopy. With prolonged time of heating (48 hours), the central quadrupole doublet disappeared (sample S-12). Also, that amount of $Fe₃O₄$ decreased to 20% and the corresponding amount of α -FeOOH increased to 80%.

Samples S-13 and S-14 were precipitated from 0.1 M $FeSO₄ + 0.05$ M NaOH solution. The situation with samples S-13 and S-14 is close to that for samples S-11 and S-12. The Mössbauer spectra of sample S-13 recorded at room temperature and liquid N_2 temperature are shown in Figure 5. Sample

S-13 shows the presence of 50% Fe₃O₄, 15% α -FeOOH and 35% of a component, which can be ascribed to γ -FeOOH or the superparamagnetic α -FeOOH.

Samples S-15 and S-16 were precipitated from 0.1 M $\text{FeSO}_4 + 0.2$ M NaOH solution. Mössbauer spectroscopy showed the presence of γ -Fe₂O₃ or a very nonstoichiometric Fe₃O₄.

X-ray Diffraction

Results of the phase analysis are given in Table III (standard samples) and Table IV. The molar fractions of the components in mixtures are estimated

TABLE III

X-ray Diffraction Phase Analysis of Standard Iron Oxides and Oxyhydroxides

| As-declared | Found |
|---|---|
| α -Fe ₂ O ₃ | α -Fe ₂ O ₃ |
| v -Fe ₂ O ₃ | γ -Fe ₂ O ₃ + α -Fe ₂ O ₃ (< 10 ⁰ / ₀) |
| stoichiometric Fe ₃ O ₄ | stoichiometric Fe ₃ O ₄ |
| α -FeOOH | a -FeOOH |
| v -FeOOH | ν -FEOOH + α -FeOOH ($\leq 5\%$) |

TABLE IV

X-ray Diffraction Phase Analysis of Samples Precipitated From FeS04 Solution

by means of a semiquantitative phase analysis. The characteristic parts of X-ray powder diffraction patterns are shown in Figures 6 (standard samples of oxyhydroxides), 7 (standard samples of oxides) and 8 (samples S-13, S-9, S-2 and S-3). Particular attention was paid to the accurate measurement of the Bragg angles in order to distinguish stoichiometric $Fe₃O₄$ from nonstoichiometric $Fe₃O₄ \cdot \gamma - Fe₂O₃$ and $Fe₃O₄$ have a very similar unit-cell parameter, but different Bravais lattice and space group: $P2_13$ (or $P4_23$) for γ -Fe₂O₃ and Fd3m for $Fe₃O₄$. Therefore, these two oxides can be easily distinguished from each other, as γ -Fe₂O₃ shows an excess of faint diffraction lines due to the Bravais lattice P. Nonstoichiometric $Fe₃O₄$ also possesses the Bravais lattice F, but a smaller unit-cell parameter than the stoichiometric $Fe₃O₄$. The values for

Figure 5. Mössbauer spectra of sample S-13 recorded at room temperature (Figure 5a) and at liquid N_2 temperature (Figure 5b).

Figure 6. Characteristic parts of the X-ray diffraction powder patterns of standard samples of iron oxyhydroxides (radiation CuKa).

Figure 7. Characteristic parts of the X-ray diffraction powder patterns of standard samples of iron oxides (radiation $CuKa$).

the unit-cell parameter of these oxides obtained in the present work are as follows:

IR Spectroscopy

The results of IR spectroscopic measurements arc summarized in Tables V and VI. The characteristic positions of the IR bands of the standard oxyhydroxides and oxides, as well as literature data $19-22$ were used for the identification of oxide components in the samples investigated. However, in some cases the IR data were not sufficient for a reliable identification of oxide components.

In previous studies²³⁻²⁵ on the adsorption of sulfate ions by iron hydroxides and oxides, it has been found that the bridged binuclear sulfato complex shows IR bands at 1150-1250, 1100-1140, 1030-1060 and 900-970 cm⁻¹ with the highest intensity at 1100-1140 cm⁻¹ The IR band at 1130 cm⁻¹ is sometimes of a higher intensity, due to the overlap with peak of goethite.

Figure 8. Characteristic parts of the X-ray diffraction powder patterns of samples precipitated from $FeSO₄$ solution (radiation CuKa).

Sulfato complexes were detected in samples S-3, S-13, S-14, S-15 and S-16, which contain $Fe₃O₄$ or a mixture of $Fe₃O₄$ and α -FeOOH. Two IR bands of very weak intensities at 2930 and 2850 cm⁻¹ were ascribed to C_2H_5OH traces, which probably remained during the process of isolation and cleaning of the precipitates. In all samples and some standards small amounts of water were detected.

DISCUSSION

In the experiments presented, the $Fe³⁺$ ions needed for the precipitation of Fe(III) oxyhydroxides and oxides were generated by the oxidation of $Fe²⁺$ ions with atmospheric oxygen dissolved in an aqueous solution and by the $Fe(OH)_2$ decomposition. Also, in all precipitation systems investigated, the oxygen content was kept low in relation to the total concentration of $Fe(II)$. The mechanism of the oxidation of $Fe²⁺$ ions by dissolved oxygen, under different physico-chemical conditions, has already been studied by other researchers.²⁶⁻³⁰ They have established that the oxidation of Fe^{2+} ions is affected by several factors, i.e. the concentrations of Fe^{2+} and O_2 , the solution pH, the kinds of coexisting anions and the amount of Fe(III) oxyhydroxide. According to Gato and coworkers,²⁶ the oxidation of Fe^{2+} ions can be described by a sequence of several reactions. The rate-determining step is the reaction between $[Fe(H₂O)₆]²⁺$ and hydrated oxygen in the form of $O₂ \cdot HO⁻$ complex. The exi-

Characteristic Infrared Positions of Standard Iron Oxyhydroxides and Oxides

Descriptions:

 $s =$ strong, vs = very strong, ms = medium strong, sh = sharp, w = weak, vw = very weak, b = broad.

Characteristic Infrared Band Positions of Oxide Precipitates Formed From FeSO4 Solution

stence of the $O_2 \cdot HO^-$ complex is assumed to be a consequence of the dissociation equilibrium of dissolved $O₂$ in water:

$$
O_2\cdot H_2O \rightleftharpoons O_2\cdot HO^- + H^*
$$

The hydroxyl group of the $O_2 \cdot HO^-$ complex substitutes one H₂O molecule in the $[Fe(H₂O)₆]²⁺$ complex and one electron is transferred to the oxygen molecule by the bridge mechanism. The result of these reactions is the formation of the $[Fe(H₂O)₃OH]²⁺$ complex ion and of the $O₂^-$ radical. The chief difference, in comparison with the precipitation from $Fe_2(SQ_4)$ solution, lies in the fact that the sulfate ions in the FeS04 solution are not included in the formation of Fe(III)-hydroxy complexes. In the case of Fe^{3+} hydrolysis in $Fe^{(SO_4)}$ solution, the SO₄² ion occupies the first ligand sphere of Fe³⁺ in the early stage of the reaction, which leads further to the formation of basic iron(III) sulfates. In a previous study,³¹ a strong influence of SO_4^2 ions on the precipitation process from $Fe₂(SO₄)₃$ or $(NH₄)FeSO₄$ solution has been shown. Also, sulfate ions have a dominant influence on the mechanism of the thermal decomposition of basic iron(III) sulfates.32

Fe(III)-hydroxy complexes can be considered precursors of »amorphous« hydroxide and oxyhydroxide in the precipitation process from FeS04 solution in slightly acidic and neutral pH regions. The hydrolysis of Fe(III) is followed by the nucleation of structures which are responsible for further development of γ -FeOOH, »amorphous« Fe(III)-hydroxide and α -FeOOH. »Amorphous« $Fe(III)$ -hydroxide may be formed during the oxidation of $FeSO₄$ solution by air flow in neutral and weakly alkaline solutions.³³ Generally, the $Fe³⁺$ ions in »amorphous« Fe(III)-hydroxide are octahedrally coordinated by a mixture of $O²$, OH⁻ and OH₂. On the basis of X-ray diffraction and Mössbauer spectroscopic measurements, $34-38$ it has been suggested that oxygen ions form a hexagonal close-packed lattice in »amorphous« Fe(III)-hydroxide. Also, on the basis of X-ray diffraction data and theoretical considerations, it has been stated that the structure of »ferric hydroxide gel« is close to that of γ -FeOOH.³⁹

In this work the processes of phase transformations are forced by heating of the precipitation systems at 90° C. The results of Mössbauer spectroscopic analysis of iron oxides precipitated from FeS04 solution indicate a strong dependence of their chemical composition and structure on the $[FeSO_4]/[NaOH]$ ratio at the beginning of the precipitation process. The phase analysis of isolated precipitates did not show the presence of an »amorphous« Fe(III) hydroxide. Also, this compound was not detected in precipitates isolated after 3 hours of heating of the FeS04 solution in weakly acidic pH-medium. The absence of »amorphous« $Fe(III)$ -hydroxide is probably due to the high reduction conditions (low oxygen content) and longer times of heating at 90° C.

The formation of γ -FeOOH from FeSO_; solution in weakly acidic pH medium is strongly a function of the oxidation/reduction conditions.13 Low $[Fe^{2+}]/[Fe^{3+}]$ ratios in sulfate solutions lead to γ -FeOOH, whereas high $[Fe^{2+}]/$ /[Fe³⁺] ratios favor α -FeOOH formation. The mechanism of the conversion of γ -FeOOH to α -FeOOH in sulfate solutions can be explained in terms of γ -FeOOH dissolution and recrystallization. The presence of γ -FeOOH was observed in the course of the hydrolysis of $\text{Fe(ClO}_4)$ solution under special circumstances (temperature below 37° C and partial neutralization of solution).⁴⁰⁻⁴² α -FeOOH was the dominant hydrolytical product in Fe(ClO₄)₃ solution. γ -FeOOH is a typical constituent of the rust formed during the atmospheric corrosion of steel^{13,16,43-44} and during the corrosion of steel in aerated water.⁴⁵⁻⁴⁹ This oxyhydroxide forms in early stages of the rusting of steel and transforms with time to α -FeOOH, γ -Fe₂O₃ and Fe₃O₄.

In the present work, the final pH values of suspensions were in the 1.93-3.83 pH range, depending on the NaOH addition at the beginning of the precipitation process. These pH values are comparable with the pH conditions during the actual corrosion of steel in water medium. Also, in the case of the atmospheric corrosion of steel, which can be regarded as wet corrosion in the thin water film on steel surface, pH values are close to pH 4. The appearance of acidic rains decreases the pH value of the thin water film on steel surface.

The Mössbauer spectra of samples S-1 to S-5, isolated from suspensions in the 1.93-2.64 pH range, have shown the presence of goethite. The measured tendency of increase of the internal magnetic field (RT) with the time of heating can be correlated with the ordering of the goethite crystal structure. A very good agreement between Mössbauer spectroscopy and X-ray diffraction identification is obtained. The superparamagnetic goethite of poor crystallinity (samples S-1 and S-2) is characterized by very broad X-ray diffraction lines. IR spectroscopy has also shown that goethite is the main component in these samples. However, on the basis of the IR spectra, the presence of small amounts of α -Fe₂O₃ or Fe₃O₄ could be also ascribed. Since the IR bands of the standard oxyhydroxides and oxides were well defined, it can be concluded that the nature of goethite precipitated from FeSO_4 solution is responsible for the above mentioned finding.

With a further increase in NaOH concentration in the precipitation systems, additional Fe-oxide components in the precipitate appear. In samples S-6 and S-7, hematite (α -Fe₂O₃) was also detected. For the interpretation of this finding the transformation of α -FeOOH to α -Fe₂O₃ by internal crystallization can be suggested. This mechanism has already been proposed in a previous study³¹ on the hydrolysis of $Fe(NO₃)$ ₃ solution at 90 °C. Preliminary experiments with electron diffraction of the isolated particles have shown the development of α -Fe₂O₃ structure inside the α -FeOOH particle.

The results of the phase analysis of samples S-8, S-9 and S-10 make it possible to follow the complex transformation of Fe-oxide precipitates. At the beginning of the precipitation process, a green-brownish suspension, indicating a mixture of $Fe(OH)_2$ and $Fe(OH)_3$, was formed. After 4 hours of heating, the precipitate was composed of nonstoichiometric $Fe₃O₄$. In the precipitate we detected $\sim 60\%$ α -FeOOH + $\sim 40\%$ Fe₃O₄ after 24 hours of heating at 90 °C. The Mössbauer spectrum (Figure 4c) shows outer peaks of small intensities, which can be ascribed to a small amount of α -Fe₂O₃; the other peaks correspond to α -FeOOH.

The Mössbauer spectra recorded at RT and 77 K were not sufficient for a precise phase analysis of samples S-11 and S-13. The presence of a central quadrupole doublet at both temperatures and the corresponding Mossbauer parameters can be ascribed to γ -FeOOH or the superparamagnetic α -FeOOH. This problem was solved by additional measurements using X-ray diffraction and IR spectroscopy. Samples S-15 and S-16 were precipitated from FeSO₄ solution with a stoichiometric ratio of $[Fe^{2+}]/[OH^-]$ which is necessary for the $Fe(OH)_2$ precipitation. The analysis of these samples has shown the presence of nonstoichiometric Fe₃O₄. In sample S-15 hematite traces were detected by X-ray diffraction.

Magnetite $(Fe₃O₄)$ is a typical corrosion product of steel in contact with water molecules. The stoichiometry, particle size and magnetic properties of $Fe₃O₄$ are very sensitive to environmental conditions during the corrosion of steel. The stoichiometry of $Fe₃O₄$ formed during corrosion can be followed by Mössbauer spectroscopy by observing the changes in the intensity of M_1 and M_2 components. An increase in the intensity of the M_1 component (outer sextet) and a corresponding decrease in the M_2 component (inner sextet) are indications of $Fe₃O₄$ nonstoichiometry. The stoichiometry of $Fe₃O₄$ is better in the adherent layer of the rust, due to the more reductive conditions there. In nonadherent layer of the rust nonstoichiometric $Fe₃O₄$ is generally present. The internal magnetic field of the M_1 component slightly increases with the increase in $Fe₃O₄$ nonstoichiometry and approaches the $H_{5/2}$ value for γ -Fe₂O₃. This result is expected, since total oxidation of Fe²⁺ ions in octahedral sites will produce γ -Fe₂O₃. The difference in Fe₃O₄ stoichiometry of the adherent and nonadherent layers can be explained by a slow penetration of O_2 through rust layers. Mössbauer spectroscopy identified the crevice corrosion products formed at 288 °C as mainly stoichiometric magnetite.50

In this study a particular attention was paid to the Mössbauer spectroscopy and X-ray diffraction of γ -Fe₂O₃, nonstoichiometric Fe₃O₄ and stoichiometric $Fe₃O₄$. The experimental results presented show that a proper use of the capabilities of Mossbauer spectroscopy and X-ray diffraction makes it possible to distinguish between γ -Fe₂O₃, nonstoichiometric Fe₃O₄ and stoichiometric $Fe₃O₄$ in Fe-oxide precipitates (or rust samples) of a complex chemical composition and structure. On the basis of a precise phase analysis of rust samples it is possible to discuss correctly the mechanism of corrosion of steel in contact with water. Naturally, in the discussion of the mechanism of corrosion, electrochemical and colloid-chemical data should also be taken into consideration.

REFERENCES

- 1. A. L. Mac k a y, Some Aspects of the Topochemistry of the Iron Oxides and Hydroxides in Reactivity of Solids, Proc. of the 4-th lnt. Symp. on the Reactivity of Solids (J. H. de Boer et al., Eds.), Elsevier, Amsterdam 1960, p. 571.
- 2. G. W. Simmons and H. Leidheiser, Jr., Corrosion and Interfacial Reactions in Application of Mössbauer Spectroscopy, Vol. I, Bell Telephone Laboratories, Inc. 1976, p. 85.
- 3. E. Matijević, Progr. Colloid Polym. Sci. 61 (1976) 24.
- 4. U. Schwertmann, and R. M. Taylor, Iron Oxides in Minerals in Soil
Environments, Soil Sci. Soc. of America, Inc., Madison WI. 1977, p. 145.
- 5. F. J. Berry, Trans. Met. Chem. 4 (1979) 209.
- 6. L. H. Bowen, Mössbauer Spectroscopy of Ferric Oxides and Hydroxides in Mössbauer Effect Data Index-1979, Plenum, New York 1979, p. 76.
7. N. C. Datta, J. Sci. Industr. Res. 40 (1981) 571.
8. C. M. Flynn, Jr., Chem. Rev
-
-
-
- 10. L. G. Johansson, and N. G. Vannerberg, Werkstoffe Korros. 32 (1981) 265.
- 11. K. Kaneko and K. Inou ye, Corrosion Sci. 27 (1981) 639.
- 12. B. Bavarian, A. Moccari, and D. D. Macdonald, Corrosion 38 (1982) 104.
- 13. H. Leidheiser, Jr., and S. Musić, Corrosion Sci. 22 (1982) 1089.
- 14. T. M. De ^v ine, Corrosion of Iron-Base Alloys in Treatise on Materials Science
- and Technology, Vol. 25., Academic Press. Inc. 1983, p. 201.
15. W. Bogaerts, P. Vanslembrouck, and A. Van Haute, Influence
of HCO_3^- , PO_4^{-3} and SO_4^{2} on localized corrosion phenomena in hot water systems in Industrial Water Treatment and Conditioning, 36-th Int. Conf. CEBE DEAU - Liege, 25-27 May 1983, p. 123.
- 16. H. Leidheiser, Jr., and I. Czakó-Nagy, Corrosion Sci. 24 (1984) 569. 17. W. Meisel, Kémia Közlemények 48 (1977) 41.
- 18. I. Dézsi and M. Fodor, Phys. Stat. Sol. 15 (1966) 247.
-
- 19. T. Misawa, J. Japan Soc. Colour Mater. 54 (1981) 309.
20. C. Morterra, C. Mirra, and E. Borello, J. Chem. Soc., Chem. Commun. 1983, p. 767.
- 21. E. Mendelovici, R. Villalba, and A. Sagarzazu, Mater. Res. Bull. 17 (1982) 241.
- 22. Sh. Yariv and E. Mendelovici, Appl. Spectr. 33 (1979) 410.
- 23. N. C. Datta, A. B. Ghatak, A. K. Cha k raborty, an S. P. Sen, Fertilizer Technol. 19 (1982) 134.
- 24. J. B. Harrison and V. E. Berkheiser, Clays Clay Miner. 30 (1982) 97.
- 25. R. L. Parfitt and R. St. C. Smart, J. Chem. Soc. Faraday 73 (1977) 796.
- 26. K. Go to, H. T a mu r a, and M. N a gay a m a, Denki Kagaku 39 (1971) 690.
- 27. H. Tamura, S. Kawamura, and M. Nagayama, Corrosion Sci. 20 (1980) 963.
- 28. H. Tamura, K. Goto, and M. Nagayama, Corrosion Sci. 16 (1976) 197.
- 29. E. J. Roekens and R. E. Van Grieken, Marine Chem. 13 (1983) 195.
- 30. T. Chmielewski and W. A. Charewicz, Hydrometallurgy 12 (1984) 21.
- 31. S. Musić, A. Vertes, G. W. Simmons, I. Czakó-Nagy, and H. Leidheiser, Jr., J. Colloid Interface Sci. 85 (1982) 256.
- 32. S. Musić, A. Vertes, G. W. Simmons, I. Czakó-Nagy, and H. Leid he is er, J r., Radiochem. Radioanal. Letters 49 (1981) 315.
- 33. A. Šolcova, J. Šubrt, J. Vinš, F. Hanousek, V. Zapletal, and
J. Tlaskal, Coll. Czech. Chem. Commun. 46 (1981) 3049.
- 34. A. A. Van der Giessen, Chemical and Physical Properties of Iron(III)--Oxide Hydrate, Ph. D. Thesis, Technical University, Eindhoven 1968.
- 35. S. Okamoto, H. Sekizawa, and S. I. Okamoto, Proc. 7-th Int. Symp. on the Reactivity of Solids (J. S. And erson et al., Eds.), Chapman and Hall Ltd., London 1972, p. 341.
- 36. S. 0 k am o to, IEEE Trans. Magn. MAG-10 (1974) 923.
- 37. J. M. D. Coey, and P. W. Readman, Earth Planet. Sci. Letters 21 (1973) 45.
- 38. E. Murad and U. Schwertmann, Am. Mineral. 65 (1980) 1044.
- 39. M. Kobayashi, and M. Uda, J. Non-Crystall. Solids 29 (1978) 419.
- 40. W. Feitknecht, and W. Michaelis, Helv. Chim. Acta 45 (1962) 212.
- 41. P. J. Murphy, A. M. Posner, and J. P. Quirk, Aust. J. Soil Res. 13 (1975) 189.
- 42. M. K. Wang and P. H. Hsu, Soil Sci. Soc. Am. J. 44 (1980) 1089.
- 43. T. Morozumi, M. Otsuka, and H. Ohashi, Boshoku Gijutsu 28 (1979) 617.
- 44. I. Kina, R. Avotina, O. Kukurs, and Z. Konstants, Izv. Akad. Nauk Latv. SSR, Ser. Khim., 1983, p. 281.
- 45. Y. Ujihira and K. Nomura, Bunseki Kagaku 27 (1978) 782.
- 46. D. C. Smith and B. McEnaney, Corrosion Sci. 19 (1979) 379.
- 47. J. Kassim, T. Baird, and J. R. Fryer, Corrosion Sci. 22 (1982) 147. 48. P. C. Bhat, M. P. Sathyavathiamma, N. G. Puttaswamy, and
- R. M. Ma 11 y a, Corrosion Sci. 23 (1983) 733.
- 49. H. Leidheiser, Jr., S. Musić, and J. F. McIntyre, Corrosion Sci. 24 (1984) 197.
- 50. H. Leid he is er, Jr., R. D. Gran at a, G. W. Simmons, S. Mu sic, and H. L. Vedage, Metal Cation Inhibitors for Controlling Denting Corrosion in Steam Generators, The Report: EPRI-NP-2655, Palo Alto, California 1982.

SAŽETAK

Miissbauerova spektroskopija, difrakcija X-zrake i IC spektroskopija oksidnih taloga nastalih iz otopine FeS04

s. Music, I. Czak6-Nagy, *s.* Popovic, *A.* Vertes i **M.** Tonkovic

Željezni oksihidroksidi i oksidi taloženi su iz otopine FeSO₄ pri malom sadrfaju kisika. Sastav i struktura, stehiometrija, velicina cestica i nuklearni magnetizam proučavani su s pomoću Mössbauerove spektroskopije, difrakcije X-zraka i IC spektroskopije. lstim instrumentalnim tehnikama karakterizirani su standardni zeljezni oksihidroksidi i oksidi. Eksperimentalni rezultati pokazali su jaku ovisnost sastava i strukture taloga o koncentracijskom odnosu [FeSO4] / [NaOH]. Pri niskim pH-vrijednostima izoliran je slabo kristalni α -FeOOH. α -Fe₂O₃ je dobiven internom kristalizacijom α -FeOOH. S vremenom zagrijavanja kod 90 °C, smjesa Fe(OH)₂/Fe(OH)₃ transformirala se u nestehiometr $a-Fe₂O₃$. Nestehiometrijski Fe₃O₄ bio je konačni taložni produkt u suspenziji sa stehiometrijskim odnosom [FeSO₄]/[NaOH] potrebnim za taloženje Fe(OH)₂. Razmotrena je važnost dobivenih rezultata za znanost o koroziji čelika u prisutnosti sulfata.