Vacancy infilling during the crystallization of Fe-deficient hematite: An in situ synchrotron X-ray diffraction study of non-classical crystal growth

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ABSTRACT

The crystallization of hematite from precursor ferrihydrite was studied using time-resolved, angle-dispersive synchrotron X-ray diffraction in aqueous solutions at pH 10 and 11 and at temperatures ranging from 80 to 170 °C. Rietveld analyses revealed a nonclassical crystallization pathway involving vacancy infilling by Fe as defective hematite nanocrystals evolved. At 90 °C and pH 11, incipient hematite particles exhibited an Fe site occupancy as low as 0.68(2), and after 30 min, Fe occupancy plateaued at 0.84(1), achieving a metastable steady state with a composition corresponding to “hydrohematite”. During crystal growth, unit-cell volume increased with an increase in Fe occupancy. The increase in Fe occupancy in hydrohematite was accomplished by deprotonation, resulting in a shortening of the long Fe-O(H) bonds and decreased distortion of the octahedral sites. Once the occupancy stabilized, the unit-cell volume contracted following further nanoparticle growth. Our study documented a variety of synthetic routes to the formation of “hydrohematite” with an Fe vacancy of 10-20 mol% in the final product.

The structure refined for synthetic hydrohematite at 90 °C and pH 11 closely matched that of natural hydrohematite from Salisbury, CT, with a refined Fe occupancy of 0.83(2). Dry heating this natural hydrohematite generated anhydrous, stoichiometric hematite, again by continuous infilling of vacancies. The transformation initiated at 150 °C and was complete at 700 °C, and it was accompanied by the formation of a minor amorphous phase that served as a reservoir for Fe during the inoculation of the defective crystalline phase.
INTRODUCTION

Hematite occurs as a naturally abundant coating on sediments and soils, often controlling the sorption and redox states of dissolved metals and organic pollutants (Barrón and Torrent 2013; Colombo et al. 2017). Hematite also has been used as a paleoclimate proxy for environments thousands to millions of years in the past, via hematite:goethite ratios (Ji et al. 2004; Hyland et al. 2015) and also as the primary carrier of natural remanent magnetization in redbeds, enabling plate tectonic reconstruction (e.g., Walker et al. 1981; Løvlie et al. 1984; Szaniawski et al. 2012). In addition, hematite is an important industrial material that is widely used in pigments, catalysts, batteries, sensors, photoelectrochemical water splitting devices, heavy-metal decontaminants, and many other applications (Paterson 1999; Walter 2006; Lu et al. 2014; McBriarty et al. 2018; Guo et al. 2020).

The presence of vacancies in hematite greatly enhances its efficiency as a photocatalyst (Guo et al. 2020), as an adsorbent of radiotoxic elements (Wang et al. 2015; McBriarty et al. 2018), as a charge-storage capacitor (Hahn et al. 2011; Lu et al. 2014), and as an electrical conductor (Yang et al. 2013). Many studies have focused on the effects of oxygen vacancies and surface iron deficiencies in hematite (Yang et al. 2013; Hu et al. 2016; Kim 2020). In this paper, we explore hematite that exhibits bulk Fe cation deficiencies.

The hematite structure (space group $R$3$c$) contains sheets of oxygen anions that are hexagonal closest packed, and Fe fills 2/3 of the octahedral sites (Blake et al. 1966). The substitution of $3\text{H}^+ \leftrightarrow 1\text{Fe}^{3+}$ yields the following general formula for hydrohematite:
Fe$_{2-x/3}$(OH)$_x$O$_{3-x}$ (Fig. 1). Wolska and Schwertmann (1989) designated the term 
“hydrohematite” for the compositional range $0.5 \geq x > 0$, and “protohematite” for $1 \geq x > 0.5$.
“Protohematite” with Fe vacancies up to 10 mol% has been observed as an intermediate during the heating of goethite (Gualtieri and Venturelli 1999; Burgina et al. 2000). In their study of the transformation of akageneite to hematite, however, Peterson et al. (2015, 2016) reported transient hematite-like precursors with Fe vacancy concentrations greatly exceeding 10 mol%. Despite the importance of hematite for commercial, technological, and paleoenvironmental applications, little is known about its tolerance for Fe deficiencies and the range of Fe occupancies in natural and synthetic hematite minerals.

Figure 1 represents the Fe stoichiometries along a join connecting compositions extending from “ferrihydrite” (simplified to Fe(OH)$_3$) to endmember hematite (Fe$_2$O$_3$), all having the $R\bar{3}c$ symmetry of stoichiometric hematite. When only $\frac{3}{4}$ of the octahedral sites of hematite are filled (i.e, Fe$_{occ} = 0.75$), then the ratio of Fe to O is $2/3 \times 3/4 = 1/2$, yielding a formula of FeOOH, with H serving to charge balance. To distinguish this phase from FeOOH-goethite with space group (S.G.) \textit{Pnma}, we will describe "$R\bar{3}c$ hematite-type FeOOH" as "Hm-FeOOH" hereafter in this paper.

Despite the modern definition of “hydrohematite” introduced by Wolska and Schwertmann (1989), the mineral name actually was first proposed in the 1840s. Hermann (1844) and Breithaupt (1847) reported the discovery of hematite-like minerals with chemistries exactly halfway between the compositions of goethite and hematite, such that Fe$_{occ} = 0.875$. Hermann (1844) called this mineral “turgite” (a term now employed by collectors to describe iridescent goethite), and Breitmann (1847) christened...
this phase “hydrohematite”. In terms of crystallographic Fe occupancy, the “hydrohematite” of Wolska and Schwertmann (1989) includes compositions for which $1 > F_{occ} > 0.917$, whereas “protohematite” includes phases for which $0.917 > F_{occ} > 0.833$. Therefore, Breitmann’s “hydrohematite” would be classified by Wolska and Schwertmann (1989) as “protohematite” rather than “hydrohematite”.

Our analyses of Fe-deficient hematite-like phases from historical collections (Chen et al. 2021b) revealed that Fe occupancies are highly variable in natural specimens. Despite the groundbreaking character of the studies by Wolska and co-workers (Wolska 1981, 1988; Wolska and Schwertmann 1989), we regard their revised nomenclature as historically problematic. In this article, we use the term “superhydrous hematite” to represent $\bar{R}3c$ structures over the range of stoichiometries between Fe(OH)$_3$ and Fe$_2$O$_3$ (Fig. 1). Although the International Mineralogical Association has not approved the name “hydrohematite” for any usage, we will employ hydrohematite (without quotations) to refer to compositions that lie halfway between those of Fe$_2$O$_3$ and FeOOH – formally, near Fe$_7$O$_9$(OH)$_3$ (Fig. 1).

We have performed a detailed atomic-scale study of the transformation of two-line ferrihydrite to hydrohematite, and hydrohematite to endmember hematite, by time-resolved synchrotron X-ray powder diffraction (TRXRD). Using freshly prepared ferrihydrite in alkaline (pH 10, 11) solutions as the starting material, we synthesized metastable Fe-deficient hydrohematite over a range of temperatures and monitored Fe occupancy parameters as nanocrystalline hematite evolved. To further explore the metastability of hydrohematite, we dry-heated natural hydrohematite and documented its conversion to stoichiometric hematite. Our study highlights a non-classical
crystallization mechanism involving vacancy inoculation with Fe during crystal growth, and it demonstrates that hematite can accommodate high concentrations of Fe vacancies and maintain structural integrity. The exceptionally Fe-deficient hydrohematite phases that we observed have the potential to tolerate high levels of dopants in their structures and to serve as candidate materials for adsorbents and photoelectrochemical energy conversion devices. In addition, they will exhibit magnetic properties that markedly depart from those observed in endmember hematite (Hill et al. 2008; Jiang et al. 2022).

EXPERIMENTAL METHODS

Ferrihydrite preparation

Ferrihydrite was the starting material for hydrohematite crystallization. Fresh ferrihydrite was prepared at room temperature using the method described by Schwertmann and Cornell (2003). We dissolved 4.039 g iron nitrate nonahydrate (Fe(NO₃)₃ • 9 H₂O) in 10 mL deionized water to produce a 1 M solution of Fe(NO₃)₃. This solution was titrated with 5 M KOH drop by drop until the pH held steady for >10 mins at pH 10 and 11. The drop-by-drop titration was completed about 40 minutes. Two-line ferrihydrite, as identified by synchrotron X-ray diffraction, precipitated as a gel (containing ~ 0.5 M ferrihydrite) immediately upon the reaction of the Fe(NO₃)₃ and KOH. We used this fresh ferrihydrite (aged no longer than 4 hours) as the reaction precursor. When the ferrihydrite aged longer than 24 hours at the room temperature, X-ray diffraction (XRD) patterns still revealed two-line ferrihydrite, but the aged ferrihydrite failed to transform to either hematite or goethite upon heating at 80 to 170 °C.
for at least 12 hours. Therefore, we utilized freshly prepared ferrihydrite for all of our TXRD experiments.

Using a 1 mL syringe, we injected ferrihydrite gel into a thin-walled quartz glass capillary (4.0 cm long and 1.0 mm outer-diameter; Charles Supper Company). UV-cured epoxy (OG142-87, EPO-TEK) then was injected to seal the capillary. The epoxy caps were cured using a full-spectrum UV lamp (EXFO X-Cite Series 120). The ferrihydrite gels were masked during this step to minimize interaction of UV light with the gels, and each capillary was rotated during curing to ensure even curing of the epoxy. The volumes of the ferrihydrite gels (~30 µL), the aging times of the gels (within 4 hr), and the amount of headspace in the capillary (1 mm away from the top of the gel surface) were maintained as similar as possible to minimize sample variations. Capillaries were inserted into a standard Cu brass mounting pin for loading in an X-ray goniometer head. The capillary was secured within the brass pin by a small amount of clay.

**Time-resolved synchrotron X-ray diffraction (TRXRD)**

TRXRD patterns were collected at the GeoSoilEnviroCARS (GSECARS) 13-BM-C beamline at the Advanced Photon Source (APS), Argonne National Laboratory (ANL). The monochromatic X-ray wavelength was refined as 0.8291(5) Å using a LaB₆ standard. The distance from the sample to the camera was refined to be 108.45(6) mm. The partially focused X-ray beam measured approximately 0.3 mm in height and 0.4 mm in width, and it was directed to the center of the capillary. Capillary orientation was maintained perpendicular to the forced-gas heater (described below). Capillaries were rotated 30° about their long axes at 1° per second to minimize preferred orientation.
effects in the XRD patterns. Each pattern was collected with an exposure time of 30 seconds using either a MAR 165 CCD detector or a Dectris PILATUS 1M pixel array detector. Full-circle images were integrated into linear intensity-versus-2θ XRD datasets using the program Dioptas (Prescher and Prakapenka 2015).

Electrically resistive forced helium heaters

A low-temperature forced-gas heater was fabricated at GSECARS for heating experiments below 250 °C, similar to the one described by Heaney et al. (2020). This low-temperature heater consisted of wound Ni coils around an inner ceramic tube, with an applied DC voltage using a Sorensen 33 V x 33 A (Model XHR 33-33) power supply to achieve resistive heating. This assembly was encased in an outer ceramic sleeve, and He gas was forced through the interior of the heated cylinder.

A second forced-gas heater was fabricated for high temperature heating experiments from room temperature to 1000 °C. The design of this high-temperature heater was nearly identical to that of the low-temperature heater but used freestanding W coils instead of Ni wound around a central ceramic support.

Both heaters were placed below the capillary with the gas flowing out of a 6 mm nozzle directed perpendicular to the capillary’s long dimension and centered on the region where XRD measurements were made. The temperature was measured by placing the tip of a type-K thermocouple into the center of the He gas flow approximately 3 millimeters from the heater’s exit and a few millimeters from the capillary. The temperature was monitored with a Keithley 2700 Multimeter. We heated samples from room temperature to the target temperature using proportional-integral-derivative (PID)
software. The low-temperature furnace heated at a rate of 14 °C/s, such that the target temperatures were achieved in few seconds before the start of the second pattern, and these temperatures stabilized at ± 0.5 °C.

Temperature was calibrated using phase transitions of RbNO$_3$ (Alfa Aesar, 99.8%, metals basis) and the melting point of metallic silver for both heaters. We monitored the transformation of RbNO$_3$ loaded in a 1.0 mm quartz glass capillary and heated from 25.7 to 400 °C. Based on this standardization, we estimate that the temperature measured by the thermocouple was within ± 1.5 °C of the actual temperature inside the capillary.

**Heating temperatures**

For experiments involving the low-temperature hydrothermal transformation of ferrihydrite to hydrohematite, runs were conducted using the low-temperature heater at the following constant temperatures: 80, 90, 100, 110, 130, 150, and 170 °C, all at pH 10, and 90 °C at pH 11. We terminated data collection after reaction progress ceased, with run times of 9, 2, 1.5, 1, 1, 0.5, 2, and 1.6 h respectively. Individual 2-dimensional diffraction patterns were collected for 30 s, and we allowed 8 seconds for software integration of the 2-dimensional ring patterns into 1-dimensional intensity-versus-2θ patterns, for a total data collection time of 38 s per XRD pattern.

To analyze the transformation of hydrohematite to hematite, we used the high-temperature furnace to heat dry natural hydrohematite powders from Salisbury, CT (Genth 255-3, Chen et al. 2021b) according to the following schedule: 1) Initial heating up to 90 °C at 13 °C/min; 2) Ramping from 90 to 1000 °C at 2 °C/min; and 3) Sustained
heating at 1000 °C for 0.5 h, to match the protocol of Gualtieri and Venturelli (1999). We loaded the natural hydrohematite powders from Salisbury in a 0.7 mm thin-walled quartz capillary with one end unsealed to allow for the release of water vapor and other gases during heating. The diameter of quartz capillary in high-temperature dry heating experiment was smaller from our low-temperature hydrothermal experiments, 0.7 versus 1.0 mm, because of the significant higher X-ray absorption among the well-packed dry powders.

Rietveld structure refinement

Rietveld structure refinements were performed using the EXPGUI interface of the General Structure Analysis System (GSAS) program (Larson and Dreele 2000; Toby and Von Dreele 2013). The starting structural parameters for hematite (S.G. \( R\bar{3}c \)), goethite (S.G. \( Pnma \)), and magnetite (S.G. \( Fd\bar{3}m \)) came from Blake et al. (1966), Gualtieri and Venturelli (1999), and Haavik et al. (2000), respectively. The Gaussian peak shape coefficients \( GU \), \( GV \), and \( GW \) determined by refinement of a LaB\(_6\) standard analyzed at the start of our data collection and fixed for the remainder of the refinements. First, we refined background, weight fraction, sample displacement, and unit-cell parameters. Backgrounds were best fit for all patterns using a shifted Chebyshev polynomial with 12 terms. Peak profiles were modeled using a pseudo-Voigt function described by Thompson et al. (1987). Second, the profile parameters sensitive to crystal size broadening (\( LX \)), strain broadening (\( LY \)), anisotropic crystal size broadening (\( ptec \)), and anisotropic microstrain broadening (\( S_{hkl} \)) were refined. After the above parameters had converged, atomic positions, and finally Fe occupancies in hematite were allowed to
refine. When the refinement of isotropic temperature factors ($U_{iso}$) generated negative values, values for $U_{iso}$ were fixed to 0.006 for Fe and 0.012 for O. Refinements were performed over a d-spacing range from 1.36 - 5.94 Å (35.50° to 8.00° 2θ). Goodness-of-fit parameters indicated high-quality refinements (Fig. S1), with $\chi^2$ ranging from 0.1 to 1.2 (± 0.03 to 0.05), $R_{wp}$ from 0.001 to 0.005 (± 0.001 to 0.002), and $R_{Bragg}$ from 0.010 to 0.014 (± 0.001 to 0.002).

Crystallite size determination

We estimated the crystallite size using the Scherrer (1918) equation:

$$\tau = \frac{K\lambda}{\beta \cos \theta}$$  \hspace{1cm} (1)

where $\tau$ is the mean particle size; $K$ is the crystal shape factor, in this case, 0.9 (Bagheri et al. 2013); $\beta$ is the full width at half maximum of a Bragg peak (FWHM, corrected for instrumental broadening by a LaB$_6$ standard); $\lambda$ is the X-ray wavelength; and $\theta$ is the Bragg angle. In order to account for instrumental broadening, the FWHM of the (211) diffraction peak of a LaB$_6$ standard was measured using the MDI JADE 2010 software (Materials Data, Inc. Livermore, CA), and that value was subtracted from the FWHM of the (116) diffraction peak of the emergent hydrohematite.

RESULTS

Transformation pathway

Two-line ferrihydrite transformed to hydrohematite and goethite in all of our experiments. Despite prior reports that this reaction sequence involves a maghemite-like
phase as an intermediate (Barrón et al., 2003; Cao et al., 2017), diffraction peaks for maghemite were never detected in any of our experiments. A representative TRXRD sequence (90 °C and pH 11) is shown in Figure 2 and a complimentary stacked 2D plot is shown in Supporting Information (Fig. S2). Precursor two-line ferrihydrite is indicated by two broad peaks at 2.54 and 1.46 Å. Within a data collection time of 38 s, the strongest diffraction peak for hydrohematite (014) typically appeared shortly before (within 2 min of) the appearance of the strongest goethite peak (101). One should note, however, that hematite diffracts more strongly than does goethite; the reference intensity ratio (RIR) for hematite is 3.18 whereas that for goethite is 2.63 (Gates-Rector and Blanton 2019). Moreover, the peak-to-background ratios for these emergent phases are extremely low. Despite these uncertainties, our analyses from the suite of experiments consistently suggested that hydrohematite precipitates from ferrihydrite slightly sooner than goethite.

The refined weight fractions and normalized scale factors as a function of time for 90 °C and pH 11 are plotted in Figure 3. Scale factors directly reflect the mass of each phase as detected by the X-ray beam within the sampling window. In order to compare the behaviors of hematite and goethite, the scale factors were normalized to their final refined values for each phase at the end of the experiment. As seen in Figure 3A, both hydrohematite and goethite continuously increased in mass abundance until attaining a plateau when ferrihydrite was depleted (~3,500 s in Fig. 3A). Despite the first appearance by hematite, the weight fraction (Fig. 3B) of goethite dominated in the earliest stages of each run but was overtaken by hematite. At 90 °C and pH 11, the experimental systems achieved a steady state with ~75 wt% hematite and ~25 wt%
goethite. As seen in Figure 3B, this hematite-to-goethite ratio was maintained despite continued co-precipitation of goethite and hematite, as reflected in the refined scale factors beyond 1000 s in Figures 3A and 3B. We attribute the constancy of these phase proportions to a uniformity in the kinetics of precipitation, which depend strongly on pH and temperature (Chen, 2021a). In agreement with Das et al. (2011), goethite growth was more favorable at low temperatures (Table 1).

Fe occupancy in hydrohematite

As shown in Figure 3, when ferrihydrite at pH 11 was heated to 90 °C, the Fe occupancy initially refined to 0.68(2) at 17 min and increased to 0.84(1) at 30 min. For the next 60 min, Fe_{occ} stabilized at 0.84. Although peak-to-background was low and errors were accordingly high for structure refinements of the first-formed crystals, our analyses revealed that Fe occupancies of incipient nuclei ranged between 0.60(2) and 0.75(2) – below that of Hm-FeOOH. Estimated standard deviations (ESDs) for refined occupancies as calculated during Rietveld refinements typically are underestimated (Post and Bish 1989). Experiments at other temperatures, however, exhibited similar behaviors (Fig. S3), such that hydrohematite started with iron occupancies below 0.75(2), and then evolved to a metastable hydrohematite composition by Fe infilling during crystal growth.

The refined Fe occupancies for the final metastable hydrohematite structures ranged from 0.80-0.90 in our experiments (Table 2, Fig. 1). Though we were time-limited during our synchrotron runs, our experiments demonstrated that highly Fe-deficient hydrohematite remained unchanged for more than 4 hours when temperatures were lower than 150 °C. Prior researchers have observed Fe-deficient hematite-like
phases as unstable, intermediate precursors during stoichiometric hematite formation 
(Gualtieri and Venturelli 1999; Peterson et al. 2018). For the first time, we have shown 
that at pH 10 and 11, ferrihydrite gels will generate hydrohematite with Fe occupancies 
ranging from 0.80-0.90 as a steady-state product (Figs. 3, 4, and S3).

In Figure 4, we have plotted the Fe occupancy of hematite as a function of 
reaction time and temperature for all runs at pH 10 from 80-170 °C. This plot shows the 
Fe concentration of hydrohematite at different time stamps at various temperatures. A 
2D projection of Figure 4 is shown in the Supplementary Information (Fig. S4). At pH 
10, higher temperatures favor hydrohematite with fewer Fe vacancies, suggesting that 
highly Fe-deficient hematite, with \( F_{\text{occ}} = 0.8-0.9 \), is evidence of precipitation at 
temperatures less than 200 °C. In a separate study, we showed that stoichiometric 
hematite forms in acidic solutions (pH ≤ 5) at high temperatures (T ≥ 180 °C) (Chen et 
al., 2022).

Changes in unit-cell parameters and crystal size during hydrohematite growth

We monitored structural changes during the transformation from ferrihydrite to 
hydrohematite by combining synchrotron TRXRD and Rietveld analysis. The growth of 
hydrohematite nanoparticles was accompanied by an initially rapid expansion of the unit 
cell, followed by a decrease in the magnitudes of \( a \) and \( c \), and therefore, of unit-cell 
volume (Fig. 5). The initial unit-cell expansion strongly correlated with an increase in Fe 
occupancy. We hypothesize that with higher occupancies of Fe, the Fe-Fe repulsion 
between adjacent face-sharing octahedra increased, resulting in the observed unit-cell 
expansion. The \( c \)-axis, which is normal to the plane of closest packed oxygen atoms,
exhibited a more dramatic extension than did the $a$-axis. For example, at 90 °C and pH 11, as the Fe occupancy increased from 0.68(2) to 0.84(1) in the first 25 min, the $c$-axis expanded 0.050% while the $a$-axis expanded 0.023%, and unit-cell volume increased 0.083%. As described in more detail below, the greater expansion along $c$ relative to $a$ was associated with a larger change in the Fe-O-Fe angle of face-sharing octahedra along the $c$-axis than in the edge-sharing octahedra perpendicular to $c$. This unit-cell expansion continued until the crystallite sizes attained ~36 nm, as ascertained by Scherrer analysis.

Once the Fe occupancy achieved a steady state of ~0.85, mean crystallite sizes continued to increase, and the unit-cell volume contracted. During this second stage, the $a$-axis contracted relatively more than the $c$-axis. Specifically, when hydrohematite nanoparticles grew from ~37 to ~50 nm at 90 °C and pH 11, the Fe occupancy remained at 0.84(1), the $a$-axis contracted 0.065% but the $c$-axis contracted only 0.025%. The decrease in lattice parameters during hydrohematite nanocrystal growth is consistent with the crystallization behavior of many other ionic nanoparticles (Perebeinos et al. 2002; Fischer et al. 2008; Diehm et al. 2012), although the cause of this inverse relationship remains controversial (Heaney et al. 2020).

Unit-cell volume, crystallite size, and scale factors plateaued at the same time in our experimental runs (~3300 s in Figs. 3 and 5), indicating that depletion of ferrihydrite induced a cessation in crystal growth. The attainment of a steady-state value for Fe occupancy occurred much earlier than the termination of crystal growth. This result suggests that the generation of Fe-deficient hydrohematite was favored relative to stoichiometric hematite in our experiments. Chen et al. (2021b) reported a crystal structure for natural hydrohematite from Salisbury, CT. The final structure refined for...
synthetic hydrohematite at pH 11, 90 °C (Table 2) was nearly identical to that of the
natural material, which exhibited a refined Fe occupancy of 0.83(2), \(a\)-axis of 5.040(1) Å, and \(c\)-axis of 13.797(1) Å.

Geometrical evolution of octahedral sites

For the end-product hydrohematite synthesized at pH 11 and 90 °C, the three longer Fe-O bonds refined as 2.164(4) Å, and the three shorter Fe-O bonds refined as 1.916(3) Å (Fig. 6). Blake et al. (1966), however, report that in stoichiometric hematite, the three longer Fe-O bonds are 2.119(2) Å and the three shorter Fe-O bonds are 1.941(2) Å. The duality of Fe-O bond lengths in the FeO\(_6\) octahedra of hematite is attributed to Fe-Fe repulsion in face-sharing octahedra. In Fe-poor hydrohematite, the distortion associated with Fe-Fe repulsion therefore is diminished, but the octahedral site geometries also are influenced by OH\(^-\) groups. The long Fe-O bond lengths that we refined for hydrohematite reflect a 15% contribution of octahedral vacancy sites as well as a reduced effective charge for O in OH\(^-\). The average Fe-O bond lengths refined here for hydrohematite (2.040(2) Å) are not significantly different from the stoichiometric hematite (2.030(2) Å) (Blake et al. 1966). We provide a full cif file of our representative synthetic hydrohematite structure at pH 11 and 90 °C in the Supplementary Materials, but we acknowledge that our neutron diffraction refinements in a separate study will present the magnetic structure and hydrogen positions of natural hydrohematite (Chen, 2021a).

Figure 7 offers some insight into the evolution of Fe-O octahedra during hydrohematite crystal growth. As Fe occupancy increased from 0.68(2) to 0.84(1), the Fe-O-Fe angle of the face-sharing octahedra increased 16.6% (from 71.4(5) to 83.25(24));
the Fe-O-Fe angle of edge-sharing octahedra increased 5.4% (from 88.4(3) to 93.13(7));
and the Fe-O-Fe angle for corner-sharing octahedra decreased 14.7% (from 142.0(9) to
123.75(32)). These trends of Fe-O-Fe angles versus Fe occupancies are consistent with
those described in Gualtieri and Venturelli (1999) for the transition of hydrohematite to
stoichiometric hematite during the dry heating of goethite powders.

The greater expansion in the Fe-O-Fe angle oriented along \( c \) for face-sharing
octahedra relative to the Fe-O-Fe angle between edge-sharing octahedra would be
expected as Fe infills sites and Fe-Fe repulsion increases. Consequently, expansion along
the \( c \)-axis is more significant relative to \( a \), as described above. An animation of the
structural changes observed during the evolution of hydrohematite at 90 °C and pH 11 is
accessible in the Supplementary Materials. This movie reveals that as the Fe occupancy
for hydrohematite increases from ~0.68 to ~0.84, the FeO\(_6\) octahedra develop from
distorted to highly symmetrical coordination polyhedra when viewed along \( c \).

**Thermal stability of hydrohematite**

We have reported natural occurrences of hydrohematite that likely persisted at
low temperatures for hundreds of millions of years (Chen et al. 2021b). Can
hydrohematite diagenetically transform to stoichiometric hematite? To test the stability
of hydrohematite, we dry-heated some hydrohematite samples from room temperature to
1000 °C using TRXRD (Figs. 8, 9). Because our synthetic hydrohematite contained low
concentrations of goethite, we employed phase pure natural hydrohematite from
Salisbury, CT (Penn State University, Genth Collection #255.3). A complete structural
refinement of Salisbury hydrohematite has been reported in Chen et al (2021b).
Rietveld refinement allowed us to explore the changes in structure and Fe
occupancy with elevated temperatures. The refined Fe occupancy of Salisbury (CT)
hydrohematite before heating was 0.81(1) (Fig. 9), within error of the value reported in
Chen et al. (2021b) of 0.83(2). The Fe occupancy increased very slightly from room
temperature to ~150 °C, beyond which the rate of Fe infilling was notably more rapid
(Fig. 10). Above 200 °C, the refined Fe occupancy increased linearly with temperature
until Fe_{occ} achieved unity at 700 °C (Fig. 10G). Magnetite appeared at 400 °C due to the
reduction of Fe^{3+} to Fe^{2+}, but only in small quantities, representing less than 2 wt% of the
mixture at 700 °C.

The transformation from hydrohematite to stoichiometric hematite was
accompanied by measurable structural changes. Consistent with our synthetic
hydrohematite, within each FeO_{6} octahedron in Salisbury hydrohematite, two types of
Fe-O bonds exist, with refined distances of 2.154(2) Å and 1.923(2) Å at room
temperature. Between room temperature and 150 °C, the overall thermal expansion
induced a stretching of all Fe-O bond lengths and an increase in unit-cell volume with
scarcely any change in Fe occupancy (Fig. 10). When the temperature exceeded 150 °C,
however, hydrohematite started to lose hydroxyls with a concomitant increase in Fe
occupancy. At the onset of this dehydration, we observed a decrease in refined crystallite
size and a rapid increase in the Fe-Fe bond length. Despite the overall thermal expansion
that accompanied heating, the long Fe-O bond contracted in response to deprotonation.
From 150 °C to 400 °C, the dehydration process dominated the thermal expansion such
that the unit-cell expansion slowed and, even more dramatically, c contracted between
230 °C and 330 °C (Fig. 11). These crystallographic changes between 150 and 400 °C
are consistent with a significant water loss, as further supported by thermogravimetric analysis-mass spectroscopy (TGA-MS) analysis (Chen et al. 2021b). Moreover, crystal annealing was revealed by the sustained increase in crystallite size from 300 to 700 °C. From 400-600 °C, the refined Fe occupancy increased from ~0.91 to ~0.96, and all lattice parameters continued to expand. The long Fe-O bond length was constant within error, but the short Fe-O bond increased due to thermal expansion. The most tightly bonded hydroxyls escaped at ~600 °C and were completely depleted at 700 °C, again consistent with TGA-MS data (Chen et al. 2021b). The long Fe-O bond length again decreased between 600 and 700 °C, whereas Fe-Fe distances hardly changed. The final decrease in the long Fe-O bond length (Fig. 10) may be related to the loss of magnetic coupling around the Néel temperature \( T_N \) at ~690 °C (Lu and Meng 2010).

The overall increase in Fe occupancies during the transformation from hydrohematite to stoichiometric hematite suggests that some hydrohematite steadily disintegrated and supplied the residual crystalline material with Fe. The breakdown of some hydrohematite into an amorphous phase was evidenced by a continual decrease in scale factor during the heating experiment (Fig. 10A). Moreover, we observed a slight increase in background between 2\( \theta \) of 8.3° and 13.8° (d-spacings of 5.7 to 3.4 Å) (Fig. 9).
DISCUSSION

Conditions for hydrohematite formation

Past researchers have reported hydrohematite as an unstable intermediate phase in the transformation of Fe-deficient to stoichiometric hematite (Wolska 1981; Wolska and Schwertmann 1989; Kustova et al. 1992; Gualtieri and Venturelli 1999; Liu et al. 2013; Peterson et al. 2015, 2016, 2018). It was not clear whether, under certain conditions, hydrohematite could be generated as a steady-state metastable product in preference to stoichiometric hematite. In the present work, we have successfully synthesized metastable hydrohematite from ferrihydrite at pH 10 and 11 as an end-stage material. Peterson et al. (2018) also described an increase in refined Fe occupancies from 0.6 to 1.0 during the synthesis of hematite (Fig. 12). However, that study employed a different reactant (akageneite) at very low starting pH (~1.0). We used fresh two-line ferrihydrite as the starting material because ferrihydrite is the most common precursor to hematite and goethite in soils (Fischer and Schwertmann 1975; Schwertmann and Murad 1983; Paterson 1999; Waychunas et al. 2005; Colombo et al. 2014; Lagroix et al. 2016). We argue that high pH may be necessary to crystallize Fe-deficient hydrohematite as a steady-state product, at least when temperatures are above 25 °C. High OH⁻ concentrations in alkaline solutions may stabilize hydrous hydrohematite relative to anhydrous hematite. Thus, our results suggest hydrohematite preferentially precipitates in moderately alkaline, low-temperature fluids.

Structural mechanism of metastable hydrohematite crystallization
Classical crystal growth theories assume atom-by-atom or molecule-by-molecule attachment to the surface of a crystal growing in solution (Karthika et al. 2016). Schwertmann and Murad (1983) first proposed a departure from this simple mechanism by asserting that hematite forms through ferrihydrite aggregation and “internal structural re-arrangement”. However, the nature of this structural change was not explained. We propose that the initially Fe-deficient structure of hydrohematite is inherited from precursor ferrihydrite through a rapid aggregation of ferrihydrite nanoparticles, consistent with the models of Soltis and Penn (2016) and Soltis et al. (2016). These authors report that trace amounts of defect-rich hematite nanoparticles already were detectable using low-temperature SQuID magnetometry in as-prepared two-line ferrihydrite, even though hematite peaks were not visible by synchrotron XRD. Hematite growth occurred by “a particle-mediated growth mechanism”, possibly involving oriented aggregation of hematite nanoparticles.

As these nanocrystals grow, we infer that Fe from dissolving or transforming two-line ferrihydrite infills the defective hematite nanocrystals by vacancy diffusion, leading to an increase in Fe occupancy in the hematite. Since X-rays average over domains that are roughly 10 times the lattice repeat, it is useful to note that the low Fe occupancies that we observed represent at least nanoscale-sized domains; aggregates of defect-free nanocrystals separated by Fe-deficient grain boundaries produced during oriented aggregation would not have yielded the XRD patterns that we observed. As more Fe infills the vacancies, stronger Fe-Fe repulsion leads to an increase in the Fe-Fe distances of face-sharing and edge-sharing octahedra, and therefore an expansion in the lattice parameters. As the Fe occupancy reaches steady-state equilibrium ($Fe_{occ} \sim 0.85$), the Fe...
occupancy ceases to change but the crystals can continue to grow. Thus, we suggest that hydrohematite forms through a non-classical process in which nanocrystal-mediated growth is accompanied by the infilling of vacancies with Fe.

Hydrogen Positions

Due to the low X-ray scattering factor for H⁺, the locations of the hydroxyl groups cannot be directly detected by XRD. However, our present study may offer some constraints with respect to the H positions in the hydrohematite structure: 1) the H atoms likely are bonded to the three longer Fe-O bonds; 2) the continuous loss of H on heating to 700 °C suggests that hydrohematite contains a variety of hydroxyl bonding environments; and 3) a high activation energy is required to completely deprotonate the structure. Indeed, prior studies have reported that temperatures >1000 °C were required to remove all hydroxyls from synthetic hydrohematite (Wolska 1981; Wolska and Schwertmann 1989).

Figure 13 shows the relationship between Fe-O bond length and Fe occupancy when our hydrothermal and dry heating data sets are combined. Although mixing of these datasets does not account for the different thermal regimes of the wet and dry heating experiments, the results do suggest that decreasing Fe occupancies -- and greater hydroxyl contents -- lead to longer “long Fe-O bonds” and shorter “short Fe-O bonds” (Fig. 13). Thus, our heating data are consistent with trends described in Gualtieri and Venturelli (1999). As we ponder the peculiar metastability of hydrohematite when Fe_{occ} = 0.85, it is interesting to note the near constancy in the long Fe-O distance as Fe_{occ} increases from 0.85 to 1.00.
XRD patterns of the hydrohematite produced from ferrihydrite in this study did not exhibit extra reflections or split peaks. Indeed, despite the proposed existence of monoclinic “clinohematite” when akageneite transforms at low pH (Peterson et al. 2015), none of the synthetic hematite in the current study supported a departure from $R\bar{3}c$ symmetry. Thus, the incorporation of considerable concentrations of H in the hematite structure does not require a violation of the $R\bar{3}c$ symmetry. One may interpret the absence of superstructures as evidence that the vacancies, and also the H positions, lack long-range order. On the other hand, neutron scattering analyses do offer evidence for partial positional ordering of H, with approximately half of the H ordered and half disordered (Chen, 2021a). TGA-MS analyses of hydrohematite, as described in Chen et al. (2021b), reveal a thermally continuous depletion of H as seen with TRXRD of heated samples and consistent with locally ordered OH groups with different bonding environments.

**IMPLICATIONS**

Our crystallographic analysis alters the conventional view of the crystallization mechanism for hematite, and perhaps other metal hydroxides. Our XRD results reflect changes in the state of the bulk structure as hematite nanoparticles nucleated and grew. Importantly, the TRXRD data were insensitive to surface structures. Consequently, the observation of non-unitary occupancy factors for Fe reflects deficiencies within the inner volumes of the nanoparticles. Thus, these experiments demonstrate that the hydrothermal transformation of ferrihydrite first generates hematite nanoparticles that are a crystallographic “Swiss cheese”. The overall space group symmetry and atomic structure
conform to those of hematite, but the vacancy concentrations of the first nanocrystals were so high that the compositions fell between Hm-Fe(OH)$_3$ and Hm-FeOOH (Fig. 1). The evolution of these highly Fe-deficient phases towards Fe$_2$O$_3$ must involve rapid diffusion of Fe into interior sites. This study demonstrates the reach of in situ synchrotron X-ray diffraction coupled with Rietveld analysis to unravel transformation mechanisms of minerals in hydrothermal systems.

Chen et al. (2021b) report the widespread occurrence of natural hydrohematite with Fe occupancies near 0.85. Here, we provide a recipe to synthesize Fe-deficient hematite as a metastable product. Combining the natural discoveries of hydrohematite (Chen et al. 2021b) and our laboratory synthesis of hydrohematite, we interpret our results to indicate that hydrohematite may form in alkaline and low-temperature solutions in sedimentary and hydrothermal deposits. Moreover, the Fe occupancy in hydrohematite and the hematite-to-goethite ratio are sensitive to pH and temperature, thus serving as a possible indicator of paleoclimatic conditions. Since the concentration of Fe vacancies in hematite can significantly modify its physical and chemical properties, such as color, magnetic properties, reactivity, and electrochemistry (Paterson 1999; Walter 2006; Lu et al. 2014; McBriarty et al. 2018; Guo et al. 2020), this study provides direction for the synthesis of hydrohematite with tailored characteristics. By extension, these results offer technological as well as natural implications, as they suggest that hydrohematite rather than hematite can form through water-saturated corrosion reactions at moderately alkaline pH, such as in the passivation layers on steel reinforcement in concrete (Leek 1991; Angst et al. 2017).
Acknowledgments

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Table 1. The weight fraction of hydrohematite (Hyhm) and goethite (Gt) and Rietveld fitting parameters at pH 10 from 80-170 °C, pH 11 and 90 °C.

<table>
<thead>
<tr>
<th>Temp (°C)</th>
<th>pH</th>
<th>Heating time (min)</th>
<th>Wt, Fraction</th>
<th>Goodness of Fit (GOF)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
<td>Hyhm</td>
<td>Gt</td>
</tr>
<tr>
<td>80</td>
<td>10</td>
<td>50</td>
<td>0.67(1)</td>
<td>0.33(1)</td>
</tr>
<tr>
<td>90</td>
<td>10</td>
<td>18</td>
<td>0.78(1)</td>
<td>0.22(1)</td>
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<tr>
<td>110</td>
<td>10</td>
<td>44</td>
<td>0.88(1)</td>
<td>0.12(1)</td>
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<td>10</td>
<td>30</td>
<td>0.79(1)</td>
<td>0.21(1)</td>
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<tr>
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<td>7</td>
<td>0.84(1)</td>
<td>0.16(1)</td>
</tr>
<tr>
<td>170</td>
<td>10</td>
<td>7</td>
<td>0.90(1)</td>
<td>0.10(1)</td>
</tr>
<tr>
<td>90</td>
<td>11</td>
<td>90</td>
<td>0.71(1)</td>
<td>0.29(1)</td>
</tr>
</tbody>
</table>

1 Weight fractions were constrained to sum to 1.0.
Table 2. Rietveld refinement lattice parameters of metastable hydrohematite and goethite synthesized at pH 10, 11 in the present study.

<table>
<thead>
<tr>
<th>Temp (°C)</th>
<th>pH</th>
<th>Heating time (min)</th>
<th>Hydrohematite</th>
<th>Goethite</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
<td>a (Å)</td>
<td>c (Å)</td>
</tr>
<tr>
<td>80</td>
<td>10</td>
<td>50</td>
<td>5.032(1)</td>
<td>13.748(1)</td>
</tr>
<tr>
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<td>18</td>
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<td>13.747(1)</td>
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<td>44</td>
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<tr>
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<td>10</td>
<td>7</td>
<td>5.039(1)</td>
<td>13.766(1)</td>
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<tr>
<td>170</td>
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<td>7</td>
<td>5.038(1)</td>
<td>13.763(1)</td>
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<tr>
<td>90</td>
<td>11</td>
<td>90</td>
<td>5.041(1)</td>
<td>13.777(1)</td>
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</table>
Figure Captions

Figure 1  The compositional relationships among ferrihydrite, hydrohematite, and stoichiometric hematite. Synthetic hydrohematite products in this study exhibited refined Fe occupancies in the range of 0.80 to 0.90.

Figure 2  Stacked TRXRD patterns showing the crystallization of hydrohematite (Hyhm, red peaks) and goethite (Gt, yellow peaks) from 2-line ferrihydrite (Fh) at 90 °C pH 11. Zoom-in plot showed that the ratio of starred peaks at (124) (left) and (030) (right) in hydrohematite is sensitive to time and Fe occupancies.

Figure 3.  (A) A representative refinement at 90 °C and pH 11 showed that hydrohematite (Hyhm, black dots) and goethite (Gt, blue squares) precipitated from ferrihydrite with time. (B) Normalized scale factors and (C) crystallite sizes achieved a plateau at ~3300 s, indicating that ferrihydrite had depleted and crystal growth ceased. (D) Refined Fe occupancy in hematite (Chen et al., 2021b) increased from 0.68(2) to 0.84(1) after 1550 s, and then stabilized at 0.84(1), suggesting metastable hydrohematite formed. Rietveld refined standard deviations were smaller than symbols.

Figure 4  The dependence of Fe occupancy on transformation time and temperature (80-170 °C) as revealed by a best fit surface to our refined TRXRD data at pH 10. Our refined Fe occupancy data are indicated by black dots. Initial structures for hydrohematite were always highly Fe-deficient and then stabilized.
at a refined Fe occupancy of 0.8-0.9. Higher temperatures produced metastable hydrohematite with fewer Fe vacancies.

**Figure 5** Evolution of unit-cell parameters $a$ (A), $c$ (B), and volume (C) during hydrohematite growth at 90 °C pH 11. Changes in Fe occupancy are included (D, Chen et al., 2021b) for comparison. Refinement errors, as reported by GSAS, are smaller than symbols.

**Figure 6** The crystal structure of hydrohematite synthesized at pH 11 and heated at 90 °C for 1.5 hr, with a refined Fe occupancy of 0.84(1). The hydrohematite structure exhibits sheets of oxygen (O) anions that are hexagonal closest packed, and iron (Fe) resides within octahedral sites. Two types of Fe-O bonds refined as 2.164(4) and 1.916(3) Å. The Fe-O-Fe angles of face-sharing, edge-sharing, and corner-sharing octahedra are 83.25°(24), 93.13°(7), and 123.75°(32) respectively.

**Figure 7** Changes in refined bond angles during hydrohematite growth at 90 °C pH 11. (A) The Fe-O-Fe angle reflects octahedra that are adjacent along the $c$-axis. These face-sharing octahedra exhibited the greatest expansion (16.65%) of all Fe-O-Fe and O-Fe-O angles. (B) The Fe-O-Fe angle represent adjacent octahedra within the (001) plane. These edge-sharing octahedra exhibited a smaller expansion of 5.34%. (C) The Fe-O-Fe angle of corner-sharing
octahedra contracted 14.77%. Standard deviations of refined values are smaller than symbols.

Figure 8  Stacked TRXRD patterns showing the transformation of hydrohematite to hematite when dry heated from room temperature to 700 °C.

Figure 9  XRD patterns with varied Fe occupancies during the transformation of hydrohematite to hematite. The ratio of starred peaks at (12\bar{4}) and (030) is sensitive to Fe occupancies.

Figure 10  Crystallographic evolution in the transformation of hydrohematite to hematite with increased temperatures. Standard deviations of refined values are smaller than symbols.

Figure 11  Refined lattice parameters in the dehydration of hydrohematite to hematite with increased temperatures.

Figure 12  A comparison of the formation of hydrohematite from ferrihydrite in our study (Chen et al., 2021b) and hematite formation from akageneite in Peterson et al. (2018). The variation in Fe occupancy in hematite is shown as a function of final phase abundance (as represented by final refined scale factors). Akageneite powders in solution were heated to 175 °C at pH 1.4 (red squares), whereas the ferrihydrite gels were heated in this study to 90 °C at pH 11.
Figure 13  Fe-O bond length dependence on Fe occupancy, combining the data from the transformation of ferrihydrite to hydrohematite at 90 °C and pH11, hydrohematite to hematite from room temperature to 700 °C, and goethite to hematite from 200 to 800 °C (Gualtieri and Venturelli, 1999). With an increased Fe occupancy, the three longer Fe-O bonds shorten, and the three shorter Fe-O bonds lengthen.
**Fe**$_{1.5}$**O**$_{2.0}$ or **Fe**$_2$**O**$_3$

**Hematite**

- Water content (%): 7.80 0.0018.42 12.67

**Fe**$_{1.0}$**(OH)**$_{3.0}$ or **Fe**(OH)$_3$

"**Ferrihydrite**"

- **Fe**$_{2-\frac{x}{3}}$**O**$_{3-x}$(**OH**)$_x$

- **Fe**$_{1.5}$**(OH)**$_{1.5}$ or **FeOOH

"**Goethite**"

- **Fe**$_{1.75}$**(OH)**$_{0.75}$ or **Fe$_7$O$_9$(**OH**)$_3$

"**Hydrohematite**"

- **Fe**$_{1.5}$**(OH)**$_{2.0}$ or **Fe$_2$O$_3$

**Hematite**

**Figure 1**

- Synthetic Hyhm at pH 10
- Natural Hyhm (Salisbury, CT)
Figure 3
Figure 4
Figure 5

A

B

C

D

Volume (Å³)

Fe_{occ}

Time (s)
Figure 6
Figure 7

(A) Face sharing Fe-O-Fe (°) over time.
(B) Edge sharing Fe-O-Fe (°) over time.
(C) Corner sharing Fe-O-Fe (°) over time.
(D) Fe_{occ} over time with error bars.
Figure 8
Figure 9
Figure 10
Figure 11
Figure 12

Iron occupancy vs. Scale factor

Akag → Hyhm → Hm

Fh → Hyhm

- Our data at 90°C, pH11
- Peterson et al., 2018