Oxidation-assisted alkaline precipitation: the effect of H$_2$O$_2$ on the size of CuO and FeOOH nanoparticles†

Sam G. F. Eggermont, a,b Ana Rua-Ibarz, c Kristof Tirez, c Xochitl Dominguez-Benetton d, b,c and Jan Fransaer* a

H$_2$O$_2$ was demonstrated to narrow the size distribution and decrease the size of CuO and hydrous FeOOH (2-line ferrihydrite) nanoparticles under conditions of high supersaturation. We introduce oxidation-assisted alkaline precipitation (Ox-AP) and compare it to traditional alkaline precipitation (AP). While for AP, a metal salt solution (e.g., CuCl$_2$) is mixed only with an alkali (e.g., NaOH), for Ox-AP, the more reduced form of that metal salt solution (e.g., CuCl) is simultaneously mixed with that alkali and an oxidant (e.g., H$_2$O$_2$). The resulting precipitates were characterized with SEM, XRD, DLS and single particle ICP-MS and shown to be nanoparticles (NPs). Ox-AP CuO NPs were up to 3 times smaller than AP NPs. Ox-AP FeOOH NPs were up to 22.5% smaller than AP NPs. We discuss and propose a possible mechanism of Ox-AP through careful consideration of the known reaction chemistry of iron and copper. We propose that an increased monomer formation rate enhances the nucleation rate, which ultimately results in smaller particles with a more narrow distribution. The more distinct effect of Ox-AP on copper, was attributed to the fast formation of the stable CuO monomer, compared to AP, where the Cu(OH)$_2$ and/or Cu$_2$(OH)$_3$Cl monomers are more likely formed. Although, the exact mechanism of Ox-AP needs experimental confirmation, our results nicely demonstrate the potential of using Ox-AP to produce smaller NPs with a more narrow distribution in comparison to using AP.

Introduction

We introduce oxidation-assisted alkaline precipitation (Ox-AP) as an alternative method to traditional alkaline precipitation (AP). With AP, a metal salt (e.g., CuCl$_2$) is mixed only with an alkali (e.g., NaOH) to achieve the supersaturation and precipitation of a sparingly soluble metal (oxy)(hydr)oxide (e.g., CuO). With Ox-AP the metal salt of the more reduced metal ion (e.g., CuCl) is mixed with both that alkali and an oxidant (e.g., H$_2$O$_2$). A careful comparison is only possible if the metal ion has at least two relatively stable oxidation states in solution. For this reason, we chose aqueous copper- and iron chloride solutions. The comparison of Ox-AP to AP is relevant given that AP is frequently cited for the production of CuO and FeOOH nanoparticles. Our original hypothesis was that, for the same supersaturation values, AP and Ox-AP would yield nanoparticles with the same characteristics (i.e., composition, size, morphology, etc.).

According to classical nucleation theory (CNT), the particle formation process via hydrolysis proceeds in four steps: (1) formation of zero-charge monomers in the form of [M(OH)$_3$(H$_2$O)$_{n-3}$]$^-$, with metal ion charge z and coordination N, (2) creation of nuclei via the olation and/or oxolation of zero-charge precursors, (3) growth of the nuclei via addition of matter via olation and/or oxolation and (4) aging of the primary particles via Ostwald ripening and/or aggregation. Recently, steps (3) and (4) have been scrutinized for several systems, where meta-stable primary particles (i.e., pre-nucleation clusters, primary nanoparticles, etc.), can also serve as building blocks for growth of nanoparticles, a mechanism not included in CNT. These mechanisms are coined nonclassical and most of the recent findings are summarized in the review of De Yoreo and coworkers.

CuO and FeOOH particles are easily produced via AP. For example, mixing a CuCl$_2$ or FeCl$_3$ solution with a highly alkaline solution (e.g., with NaOH) yields Cu(OH)$_2$ or Fe(OH)$_3$, respectively, which gradually (either fast or slow) form CuO or FeOOH via oxolation. The overall reaction scheme simplifies to:

\[
\text{Cu}^{2+} + 2\text{OH}^- \rightarrow \text{Cu(OH)}_2 \rightarrow \text{CuO} + \text{H}_2\text{O} \quad (1)
\]

\[
\text{Fe}^{3+} + 3\text{OH}^- \rightarrow \text{Fe(OH)}_3 \rightarrow \text{FeOOH} + \text{H}_2\text{O} \quad (2)
\]
where the coordinated water ligands of Cu$^{2+}$ and Fe$^{3+}$ are omitted for clarity. The corresponding reaction quotients $Q_{\text{Cu(OH)$_2$}}$ and $Q_{\text{Fe(OH)$_3$}}$ are:

$$Q_{\text{Cu(OH)$_2$}} = a_{\text{Cu$_2$}} \hat{\gamma}_{\text{Cu$_2$}} \hat{\gamma}_{\text{OH}} \left[\text{Cu$_2$}\hat{\gamma}\text{[OH$^{-}$]}\right]^2$$  \hspace{1cm} (3)

$$Q_{\text{Fe(OH)$_3$}} = a_{\text{Fe$_3$}} \hat{\gamma}_{\text{Fe$_3$}} \hat{\gamma}_{\text{OH}} \left[\text{Fe$_3$}\hat{\gamma}\text{[OH$^{-}$]}\right]^3$$  \hspace{1cm} (4)

with $a_i$ the activities and $\gamma_i$ the activity coefficients of a given species $i$, and the concentrations of Cu$^{2+}$, Fe$^{3+}$ and OH$^{-}$ [mol L$^{-1}$] given in between square brackets.

CNT couples the chemical reactions of the monomer formation to the physical process of phase separation (i.e., from liquid to liquid + solid) via the concept of supersaturation. For reaction precipitation, the supersaturation $S$ is defined as the ratio between the reaction quotient $Q$ and solubility product $K_{\text{sp}}$ as exemplified for Cu(OH)$_2$:

$$S_{\text{Cu(OH)$_2$}} = \frac{Q_{\text{Cu(OH)$_2$}}}{K_{\text{sp,Cu(OH)$_2$}}}$$  \hspace{1cm} (5)

with $K_{\text{sp,Cu(OH)$_2$}}$:

$$K_{\text{sp,Cu(OH)$_2$}} = a_{\text{Cu$_2$},eq} \hat{\gamma}_{\text{Cu$_2$}} \hat{\gamma}_{\text{OH}} \cdot \text{eq}^2$$  \hspace{1cm} (6)

and $a_{i,eq}$ the equilibrium activities of species $i$.

If $S > 1$, the amount of reactants is in excess for equilibrium and precipitation can occur. Often $S$ needs to be many times higher than 1 before precipitation occurs, depending on other limiting factors (e.g., surface energy of the precipitate, temperature, etc.). If $S < 1$, the amount of reactants is limiting for precipitation and hence the reactants stay in solution. According to CNT, a higher supersaturation level increases the nucleation rate which results in smaller particles.$^{46-8}$

The (homogeneous) nucleation rate $J$ [s$^{-1}$] depends on supersaturation via a power law, which in the simplest form is written as:

$$J = 10^{-4} [\log S]^2$$  \hspace{1cm} (7)

with $J_{\text{max}}$ [s$^{-1}$] the nucleation rate at infinitely high supersaturation and $A$ [—] a collection of variables:$^{56-8}$

$$A = \frac{4\beta_p^3 \gamma^3 V^2}{27 \beta_p^2 (k_B T \ln 10)}$$  \hspace{1cm} (8)

with $\beta_p$ [—] the form factor of the particle surface, $\gamma$ [J m$^{-2}$] the surface free energy per unit area, $V$ [m$^3$] the molecular volume, $\beta_p$ [—] the form factor of the particle volume, $k_B$ [m$^2$ kg s$^{-2}$ K$^{-1}$] the Boltzmann constant and $T$ [K] the temperature.

A more elaborate approach, including for example heterogeneous nucleation and other important phenomena during precipitation, is found in scientific CNT literature.$^{46-8,11}$ Although CNT demonstrated its usefulness in the past, recent studies increasingly demonstrate the limitations of CNT. These studies discovered different nonclassical crystallization routes and the importance of chemical kinetics.$^{10,12}$

In this study we refute our original hypothesis, by demonstrating for two cases that Ox-AP yielded smaller nanoparticles with a more narrow size distribution than AP. Ox-AP achieves these feats with as little as possible alteration of the supersaturation. Contrary to recent discoveries of nonclassical crystallization routes, which are related to steps (3) and (4) of the precipitation process, we believe that the effect of Ox-AP is related to steps (1) and (2) of the precipitation process. We discuss the possible nature of Ox-AP of CuO and FeOOH through careful consideration of the known reaction chemistry of iron and copper.

### Experimental

Complete experimental details are described in the ESL.$^\dagger$

#### Preparation of the reagent solutions

A 10 mM metal chloride solution, supplemented with 0.5 M of NaCl, was prepared for each of the metal ions discussed above (i.e., CuCl$_2$, CuCl, FeCl$_3$, and FeCl$_2$, respectively). The NaCl was a necessity in the case of CuCl to keep Cu$^+$ in solution (for complexing between Cl$^-$ and Cu$^+$).$^{13}$ In the other cases, NaCl was added to keep composition consistency between all cases. To further isolate the effect of H$_2$O$_2$, we tweaked the individual NaOH solutions, with or without 10 mM H$_2$O$_2$ (100% excess), in function of their matching metal chloride solution, to always have the same theoretical leftover end-concentration of OH$^-$ (0.01 M, i.e., pH 12). This was deemed important, because Jolivet and coworkers demonstrated the importance of pH on the size of metal oxide nanoparticles.$^{14}$

#### Preparation of the nanoparticles

To have a consistent reaction front and consistent mixing, the solutions were mixed in a Y-junction by pumping them separately at high equal volumetric rates, after which the mixture was collected in a glass beaker. A summary of the electrolytes and a schematic drawing of the Y-junction tubular mixing-reactor are shown in Fig. 1.
Material characterization techniques

The steady state pH after mixing was measured with a pH probe. Scanning electron microscopy (SEM) was used for the determination of size and morphology of the nanoparticles. X-ray powder diffraction (XRD) was used for the determination of the composition of the nanoparticles. Single particle inductively coupled plasma mass spectrometry (sp ICP-MS) and dynamic light scattering (DLS) were used for the determination of the size of the nanoparticles.

Preparation of the samples for material characterization

Every effluent was divided into three samples. Two samples for wet analysis (i.e., sp ICP-MS and DLS). The other samples for dry analysis (i.e., XRD and SEM). All samples were treated according to a washing procedure. The samples for wet analysis required dilution steps specific to each of the applied techniques. The dry analysis samples required a drying step.

Results

Visual observations of the nanoparticle suspension

Fig. 2 demonstrates the dispersed precipitates just after mixing, stored in plastic storage tubes. When H₂O₂ was used, some gas evolution was observed, in the form of bubbles, suggesting the decomposition of excess H₂O₂ to oxygen gas and water (see insets of Fig. 2b and d). Table 1 shows the pH measured after mixing, which was lower than the expected value of 12. This is most likely due to the sodium error at 0.5 M NaCl. The pH difference between the copper case and the iron case was approximately 1 mM OH⁻, and thus considered insignificant for this study, because it is less than 1% of the amount of the total OH⁻ added.

The copper samples had a distinct color difference (Fig. 2). Initially, the Cu²⁺ sample was distinctly blue, but developed into a dark brown color by the end of the washing process. The Cu⁺ (with H₂O₂) sample was dark greenish brown for a very short instance, but even before the washing step started its color changed to dark brown. Both the Fe³⁺ and the Fe²⁺ (with H₂O₂) formed in the cases with H₂O₂.

| Solution                  | pH measured after mixing |
|---------------------------|--------------------------|
| CuCl₂                    | 11.67 ± 0.02             |
| CuCl + H₂O₂               | 11.72 ± 0.04             |
| FeCl₃                     | 11.49 ± 0.05             |
| FeCl₂ + H₂O₂              | 11.54 ± 0.06             |

Table 1 pH resulting after mixing the different solutions (the theoretical pH value anticipated for all instances is pH 12)

Sample were bright orange, with little observable change over time.

Size of the nanoparticles

Single particle ICP-MS and DLS were used to determine the size of the nanoparticles. In all cases, spherical shape was assumed, and thus, the particle spherical equivalent diameter is reported. For rod-shaped particles of similar length over width ratio, comparison of particles with different sizes remains valid. Fig. 3 demonstrates the particle size distributions calculated from the sp ICP-MS measurements for the different cases. In the case of copper, the size of the particles shows a remarkable difference, contradicting our initial hypothesis of size consistency between AP and Ox-AP for the same supersaturation value. In the case of iron, the size difference follows a similar trend, although much less pronounced. Fig. 4a and b respectively compare the DLS and sp ICP-MS size measurements. Both measurements clearly demonstrate that the Ox-AP yields smaller particles than the traditional AP. In the case of copper, the average size of the oxidation-induced precipitates is 2.8 nm smaller (i.e., 30 nm instead of 90 nm) according to sp ICP-MS, and 2.2 times smaller (i.e., 100 nm instead of 250 nm) according to DLS. In the case of iron the results are less pronounced, yet inconsistent with our original hypothesis: the average size of the oxidation-induced precipitates is 12.5% smaller (i.e., 28 nm instead of 32 nm) than with the traditional AP-obtained precipitates according to sp ICP-MS, and 22.5% smaller (i.e., 117 nm instead of 151 nm) according to DLS. The difference in the sizes obtained by both techniques, sp ICP-
MS and DLS, can mainly be attributed to two factors. Firstly, there exists a chance that agglomerates remain in our washed, but unstabilized, sample, resulting in increased apparent particle diameters. Secondly, because of anisotropy, the particles size (z-average) of DLS measurements can be strongly distorted. Table 2 shows the polydispersity index (PDI) for both single-particle ICP-MS and DLS, respectively. A lower PDI value corresponds to a more narrow distribution. Either technique demonstrates a more narrow distribution for Ox-AP than for traditional AP and this for both the copper case and the iron case.

Composition of the nanoparticles

The XRD-patterns for the copper and iron cases are shown in Fig. 5a and b, respectively. CuO is the main phase for the copper case. The CuCl case shows two additional peaks, indicating an additional Cu₂(OH)₃Cl phase. This indicates that at least part of the precipitated phase is the Cu₂(OH)₃Cl instead of the Cu(OH)₂ and besides the reactions described in eqn (1), following overall reactions are likely:

\[
2\text{Cu}^{2+} + 3\text{OH}^- + \text{Cl}^- \rightarrow \text{Cu}_3(\text{OH})_2\text{Cl} \quad (9)
\]

\[
\text{Cu}_2(\text{OH})_3\text{Cl} + \text{OH}^- \rightarrow 2\text{CuO} + 2\text{H}_2\text{O} + \text{Cl}^- \quad (10)
\]

The mechanism in eqn (10) would be similar to the conversion mechanism of Cu(OH)₂ to CuO at higher pH. Both iron cases show highly amorphous XRD patterns of 2-line ferrihydrite (hydrus FeOOH), although only the main (110) peak (one of two peaks in 2-line ferrihydrite) is visible. It is known that, with time ferrihydrite can convert into more crystalline products such as goethite and hematite.

Morphology of the nanoparticles

Fig. 6 shows the SEM images for the different precipitates. For the copper case, separate particles are easily distinguished for AP and Ox-AP. Ellipsoidal needle-shaped NPs are observable in either case, but the Ox-AP NPs are clearly smaller than the AP NPs. For the iron case, even at higher magnification than for the copper case, separate particles are hardly distinguishable for AP and Ox-AP (Fig. 6c and d). The nanoparticles form large aggregates of smaller nanoparticles. With digital magnification, it was possible to identify a few separate nanoparticles (Fig. 1 ESI†).
However, these do not provide conclusive evidence on the size differences between AP and Ox-AP for the iron case.

Discussion

The observations can be summarized as follows: (1) the experiments with iron result in smaller nanoparticles with more narrow size distributions than the experiments with copper, both for Ox-AP and AP; (2) Ox-AP results in smaller nanoparticles with more narrow size distributions than AP, both for the copper and the iron cases; (3) the effect of Ox-AP is much more pronounced in the case of copper than in the case of iron.

The observations can be discussed in the light of the particle formation process. As discussed above, the particle formation process via hydrolysis proceeds in four steps: (1) formation of zero-charge precursors, (2) creation of nuclei via the olation and/or oxolation of zero-charge precursors, (3) growth of the nuclei via addition of matter via olation and/or oxolation and (4) aging of the primary particles via Ostwald ripening and/or aggregation.\(^9\) Step (1) depends on the solubility of the species and the concentration of the reagents and the available ligands (i.e., especially on the pH). Moreover, it is also limited by the lability of the water ligand in the first coordination shell of the metal ion. Step (2) depends on the concentration of the zero-charge species and the water lability as well, because the lability codetermines the rates at which olation and oxolation can occur. Step (3) and step (4) are still intensively debated in current literature and strongly depend on step (1) and step (2), and any of the other process parameters.\(^9\)

In the AP process, OH\(^-\) ions compete with H\(_2\)O and Cl\(^-\) ions to enter the first hydration sphere of the metal aquo complex. The strong nucleophilic character of OH\(^-\) allows it to form more stable complexes with the metal ion, and at the right OH\(^-\) concentration the most stable form is a sparingly soluble zero-charge monomer. This monomer can form polymers via olation and oxolation until a critical size is reached and they are stable in solution. The critical size strongly depends on the inherent chemical nature of the monomer (i.e., the metal ion, the ligands, etc.), the olation and oxolation kinetics to which it is subjected, and also on the level of supersaturation (i.e., the concentration of the precursor ions).

Supersaturation S is much higher for the iron case than for the copper case (Table 3, more detailed calculations are given in the ESI†), and in accordance to CNT, the NPs in the iron case should be smaller than in the copper case.

Since the end pH, background electrolyte concentration, metal ion concentration and process operating conditions were the same for the copper and iron cases, the inherent differences between aqueous behaviour of Cu\(^{2+}\) and Fe\(^{3+}\) (i.e., the case-specific values of \(J_{\text{max}}\) and \(A\) from eqn (7)) likely cause the observed size difference between the CuO and hydrous FeOOH nanoparticles. The values of \(J_{\text{max}}\) and \(A\) depend on the chemical nature of the monomers and the olation and oxolation kinetics of those monomers. This is inherently different for copper and iron. Discussion of these differences would lead us to far afield.

More interesting is to compare the Ox-AP cases of copper and iron with the AP cases of copper and iron, respectively. Our original hypothesis was that, for the same supersaturation values, AP and Ox-AP would yield nanoparticles with the same characteristics (i.e., composition, size, morphology, etc.). Imagine infinitely fast oxidation of the metal ions in the Ox-AP case. This would cause complete oxidation prior to any other reaction (i.e., any of the four steps in the particle formation process), OH\(^-\) would be formed according to following equations:

\[
2\text{Cu}^+ + \text{H}_2\text{O}_2 + 2\text{OH}^- \rightarrow 2\text{Cu}^{2+} + 2\text{OHCu}^{2+} + 2\text{OH}^- + \text{2H}_2\text{O}
\]

(11)

\[
2\text{Fe}^{2+} + \text{H}_2\text{O}_2 + 4\text{OH}^- \rightarrow 2\text{Fe}^{3+} + 2\text{OHFe}^{2+} + 4\text{OH}^- + \text{2FeOOH} + \text{2H}_2\text{O}
\]

(12)

where the Cu\(^{2+}\) and Fe\(^{3+}\) substrresses the fact that these OH\(^-\) ions are formed by the redox reaction of the metal ion with H\(_2\)O\(_2\). The metal ion concentration and the amount of OH\(^-\) would be the same for AP and Ox-AP. Therefore, the overall supersaturation would be the same and CNT would thus predict the same precipitates with the same sizes. Interestingly, our experimental results contradict this: compared to AP, Ox-AP results in smaller nanoparticles with a more narrow size distribution. This refutes the hypothesis. Therefore H\(_2\)O\(_2\) likely interferes on the mechanisms of steps (1) to (4), rather than cause complete oxidation prior to those steps. The question remains, where does H\(_2\)O\(_2\) interfere in the particle formation process?

Because Ox-AP results in smaller particles with more narrow size-distributions, it is unlikely that in our process H\(_2\)O\(_2\) would interfere significantly in the ageing process (i.e., step (4)), because although ageing tends to narrow the distribution size, it generally increases the average particle size. This is not in agreement with our observations. Therefore, prior to ageing, the particles should already have a smaller sizes and more narrow distributions.

The later stages of growth and ageing are often suppressed using large molecules that block the surface to avoid coalescence or by using certain ligands that suppress the redissolution of smaller particles in favour of the larger particles (i.e., Ostwald ripening). It is also unlikely that hydrogen peroxide would interfere in these stages, especially because most of the hydrogen peroxide reacts away in an oxidation reaction with the metal, and the excess quickly decomposes (i.e., bubbles are visible in the final solution). Therefore the interference of H\(_2\)O\(_2\) is most likely occurring during step (1) and/or step (2) of the precipitation process. The olation and oxolation reactions in

| Precursor     | \(K_{\text{sp}}\) | \(S\) for end-pH 12 |
|---------------|------------------|---------------------|
| Cu(OH)\(_2\)  | \(2.20 \times 10^{-20}\) mol\(^3\) L\(^{-3}\) | 9.1 \times 10\(^{13}\) |
| Cu\(_2\)(OH)\(_3\)Cl | \(2.34 \times 10^{-35}\) mol\(^4\) L\(^{-6}\) | 5.2 \times 10\(^{24}\) |
| Fe(OH)\(_3\) | \(2.79 \times 10^{-39}\) mol\(^4\) L\(^{-6}\) | 2.8 \times 10\(^{31}\) |
step (2) depend strongly on the characteristics of the zero-charge monomers formed in step 1. Therefore, we argue that H$_2$O$_2$ takes an important role in the formation of the monomers and the subsequent olation and/or oxolation reactions.

In Ox-AP, both OH$^-$ and HO$_2^-$ (i.e., the deprotonated form of H$_2$O$_2$ at high p$\text{H}$ values$^{28}$) act as strong nucleophiles (i.e., HO$_2^-$ even more so than OH$^-$) and compete with water and Cl$^-$ to enter the first hydration sphere of the metal ion. There are many possible reaction pathways for HO$_2^-$ (i.e., H$_2$O$_2$) with the metal ion, known in scientific literature as Fenton(-like) reactions.$^{28,29}$ The exact pathways are still subject of much research and heavily system dependent. Here, we propose, both for copper and iron, only one of the possible pathways. For clarity, we do not take into account the many possible side-reactions and intermediaries.

Without hydrogen peroxide, the expected zero-charge monomer of Cu$^+$ is CuOH at elevated p$\text{H}$, with a subsequent oxolation reaction to Cu$_2$O:

$$2\text{Cu}^+ + 2\text{OH}^- \rightarrow 2\text{CuOH} \rightarrow \text{Cu}_2\text{O} + \text{H}_2\text{O}$$

(13)

With hydrogen peroxide, the CuOOH is another possible monomer:

$$\text{Cu}^+ + \text{HO}_2^- \rightarrow \text{CuOOH}$$

(14)

Subsequently, the oxidative nature of the HO$_2^-$ ligand could result in a Fenton reaction:$^{29}$

$$\text{CuOOH} \rightarrow \text{CuO} + \text{OH}^-$$

(15)

The proposed reactions require little ligand exchanges and result directly in the CuO monomer, rather than the Cu(OH)$_2$ monomer or the Cu$_2$(OH)$_3$Cl dimer. The OH$^-$ radical is a non-selective oxidant which can oxidise neighbouring species, for example the Cu$^+$ ion or the CuOH monomer, at high rates$^{30}$

$$\text{Cu}^+ + \text{OH}^- \rightarrow \text{CuO} + \text{H}^+$$

(16)

$$\text{CuOOH} + \text{OH}^- \rightarrow \text{CuO} + \text{H}_2\text{O}$$

(17)

In any of the proposed reactions, the CuO is the end product. This might explain why, the conversion to the brown CuO precipitate is much faster for Ox-AP than for AP. The rapid formation of the stable CuO monomer, is a likely explanation of the smaller nanoparticles. The rapid depletion of monomers during the nucleation phase would then explain the more narrow size distribution. The lack of available monomers also slows down the subsequent growth. Although no evidence is currently available, we hope that the exact mechanism can be determined experimentally in the near future.

In the case of iron, the effect of Ox-AP is less obvious. A possible reason follows from our proposed reaction mechanism. Without hydrogen peroxide, the expected zero-charge monomer is Fe(H$_2$O)$_6$(OH)$_3$. The complexed water results from the fact that Fe$^{2+}$ and Fe$^{3+}$ easily form hexaqua complexes.$^{31}$ The Fe(H$_2$O)$_3$(OH)$_3$ monomer, subsequently transforms into FeOOH(H$_2$O)$_3$ and with time (i.e., with ageing) crystallizes into FeOOH, with different possible crystal structures depending on the solution and process characteristics. Ageing was not allowed in our experiments, which limited crystallization. With hydrogen peroxide, we can draft a reaction similar to eqn (14), taking into account that Fe$^{2+}$ easily forms the hexaqua complex:

$$[\text{Fe}(\text{H}_2\text{O})_6^{2+}] + \text{HO}_2^- \rightarrow [\text{Fe}(\text{H}_2\text{O})_6\text{OOH}]^+ + \text{H}_2\text{O}$$

(18)

$$[\text{Fe}(\text{H}_2\text{O})_6\text{OOH}]^+ + \text{OH}^- + \text{H}_2\text{O}$$

(19)

Unlike with copper, with iron the reaction does not directly result in a zero-charge monomer. Moreover, the product itself (i.e., $[\text{Fe}(\text{H}_2\text{O})_6(\text{OH})_3]^+$) is an intermediary for ferrihydrite precipitation in AP.$^{32,33}$ The ferrihydrite results from a second step with OH$^-$:

$$[\text{Fe}(\text{H}_2\text{O})_6(\text{OH})_3]^+ + \text{OH}^- \rightarrow \text{Fe}(\text{OH})_3(\text{H}_2\text{O})_3 + \text{H}_2\text{O}$$

(20)

Remark that the sequence of these reactions is not fixed. The hydrolysis reaction with OH$^-$ might also occur first in the sequence, after which HO$_2^-$ reacts with the $[\text{Fe}(\text{H}_2\text{O})_6(\text{OH})]^+$ complex to form the zero-charge $[\text{Fe}(\text{H}_2\text{O})_6\text{OOH}]$ monomer. This monomer would then further oxidise to form the Fe(OH)$_3$(H$_2$O)$_3$ monomer.

In the above reaction mechanisms, the reason might be found for the lesser effect of Ox-AP in the case of iron. The reaction with HO$_2^-$ does not immediately give rise to a zero-charge monomer, requiring an extra step. Additionally, the resulting monomer is the same as the one expected for AP of iron, rather than directly resulting in a more stable form, as with copper. This is in line with the visibly observed alikeness of the precipitates (i.e., no transformation is obvious like with copper). One possible reason that an effect is still observable, could be that with Fe$^{2+}$, the zero-charge monomer forms with only two ligand substitutions rather than three in the AP case of Fe$^{3+}$. Additionally, the water exchange kinetics of Fe$^{2+}$ are much faster than those of Fe$^{3+}$ (4 orders of magnitude$^{34}$), which might cause faster nucleation rates and the related observed, but small, effects.

### Conclusion

We demonstrated that Ox-AP yields smaller nanoparticles with a more narrow size distribution than traditional AP for iron- and copper-chloride solutions in conditions of high supersaturation. For CuO, the differences between traditional AP and Ox-AP are very clear. For FeOOH, the differences are observable to a lesser extent. The proposed mechanisms of Ox-AP are the improved condensation kinetics and the faster formation of a stable zero-charge CuO monomer. For iron, we believe that the condensation kinetics improve less than for copper. We propose that the lack of a more stable monomer, in contrast to the copper case, causes this lesser effect. Although Ox-AP of iron lacks the more stable monomer, two other factors were proposed as possible causes for the observed effects. The first factor is the lower amount of necessary ligand exchanges to...
form a zero-charge monomer for Fe$^{2+}$ compared to Fe$^{3+}$. The second factor is the higher water exchange kinetics for Fe$^{3+}$ than for Fe$^{2+}$. Both factors possibly result in higher condensation kinetics for Ox-AP. In light of our observations, other systems can be prospected for the production of small nanoparticles with a narrow size distribution. Herein, a sufficiently stable precursor metal ion that can be oxidised is detrimental. Our theory is insightful for further investigations of other systems, but we hope that the exact mechanism can be confirmed experimentally in the near future to allow for better synthesis-by-design.

**Conflicts of interest**

There are no conflicts to declare.

**Acknowledgements**

This research was funded by the Strategic Initiative Materials (SIM) in Flanders, within the recyclable materials MaRes program, under grant agreement no. 150626 (Get-A-Met project). JF and XD acknowledge funding from the European Union’s Horizon 2020 research and innovation programme under grant agreement no. 654100 (CHPM2030 project).

**Notes and references**

1 Q. Zhang, K. Zhang, D. Xu, G. Yang, H. Huang, F. Nie, C. Liu and S. Yang, *Prog. Mater. Sci.*, 2014, 60, 208–337.
2 M. Mohapatra and S. Anand, *Int. J. Eng. Sci. Technol.*, 2010, 2(8), 127–146.
3 V. K. LaMer and R. H. Dinegar, *J. Am. Chem. Soc.*, 1950, 72, 4847–4854.
4 A. E. Nielsen, *Kinetik of precipitation*, Pergamon, 1964, vol. 18.
5 T. Sugimoto, *Adv. Colloid Interface Sci.*, 1987, 28, 65–108.
6 J. Dirksen and T. Ring, *Chem. Eng. Sci.*, 1991, 46, 2389–2427.
7 J. Klein and R. David, *Crystallization technology handbook*, 1995, p. 359.
8 G. Demopoulos, *Hydrometallurgy*, 2009, 96, 199–214.
9 J.-P. Jolivet, M. Henry and J. Livage, *Metal oxide chemistry and synthesis: from solution to solid state*, Wiley-Blackwell, 2000.
10 J. J. De Yoreo, P. U. Gilbert, N. A. Sommerdijk, R. L. Penn, S. Whiteham, D. Joester, H. Zhang, J. D. Rimer, A. Navrotsky, J. F. Banfield, *et al.*, *Science*, 2015, 349, aaa6760.
11 N. T. Thanh, N. Maclean and S. Mahiddine, *Chem. Rev.*, 2014, 114, 7610–7630.
12 R. Xie, Z. Li and X. Peng, *J. Am. Chem. Soc.*, 2009, 131, 15457–15466.
13 J. Fritz, *J. Phys. Chem.*, 1981, 85, 890–894.
14 J.-P. Jolivet, S. Cassaignon, C. Chanéac, D. Chiche and E. Tronc, *J. Sol-Gel Sci. Technol.*, 2008, 46, 299–305.
15 M. D. Montaño, J. W. Olesik, A. G. Barber, K. Challis and J. F. Ranville, *Anal. Bioanal. Chem.*, 2016, 408, 5053–5074.
16 A. Ortega and J. García de la Torre, *J. Chem. Phys.*, 2003, 119, 9914–9919.
17 R. C. Murdock, L. Braydich-Stolle, A. M. Schrand, J. J. Schlager and S. M. Hussain, *Toxicol. Sci.*, 2008, 101, 239–253.
18 P. A. Hassan, S. Rana and G. Verma, *Langmuir*, 2014, 31, 3–12.
19 H. B. Weiser, W. Milligan and E. Cook, *J. Am. Chem. Soc.*, 1942, 64, 503–508.
20 Y. Cudennec and A. Lecerf, *Solid State Sci.*, 2003, 5, 1471–1474.
21 H. Liu, P. Li, B. Lu, Y. Wei and Y. Sun, *J. Solid State Chem.*, 2009, 182, 1767–1771.
22 B.-S. Zhu, Y. Jia, Z. Jin, B. Sun, T. Luo, L.-T. Kong and J.-H. Liu, *RSC Adv.*, 2015, 5, 84389–84397.
23 S. Das, M. J. Hendry and J. Essiflie-Dughan, *Environ. Sci. Technol.*, 2010, 45, 268–275.
24 P. Patnaik, *Handbook of inorganic chemicals*, McGraw-Hill, New York, 2003, vol. 529.
25 J. W. Ball, D. K. Nordstrom, *et al.*, *WATEQ4F – User’s manual with revised thermodynamic data base and test cases for calculating speciation of major, trace and redox elements in natural waters*, Open-File Report, 90–129, 1991, p. 185, DOI: 10.3133/ofr90129.
26 M. Pourbaix, *Atlas of Electrochemical Equilibrium in Aqueous Solutions*, NACE, 1974, p. 307.
27 J. O. Edwards and R. G. Pearson, *J. Am. Chem. Soc.*, 1962, 84, 16–24.
28 E. Neyens and J. Baeyens, *J. Hazard. Mater.*, 2003, 98, 33–50.
29 A. D. Bokare and W. Choi, *J. Hazard. Mater.*, 2014, 275, 121–135.
30 H. L. Wiegand, C. T. Orths, K. Kerpen, H. V. Lutze and T. C. Schmidt, *Environ. Sci. Technol.*, 2017, 51, 14321–14329.
31 G. Wilkinson, R. D. Gillard and J. A. McCleverty, *Comprehensive coordination chemistry, The synthesis, reactions, properties and applications of coordination compounds, Main group and early transition elements*, vol. 3, 1987.
32 R. M. Cornell, R. Giovanoli and W. Schneider, *J. Chem. Technol. Biotechnol.*, 1989, 46, 115–134.
33 A. Stefánsson, *Environ. Sci. Technol.*, 2007, 41, 6117–6123.
34 D. T. Richens, *Chem. Rev.*, 2005, 105, 1961–2002.