Rapidly self-deoxygenating controlled radical polymerization in water via in situ disproportionation of Cu(I)†

Evelina Liarou, Yisong Han, Ana M. Sanchez, Marc Walker and David M. Haddleton *a

Rapidly self-deoxygenating Cu-RDRP in aqueous media is investigated. The disproportionation of Cu(I)/Me₆Tren in water towards Cu(II) and highly reactive Cu(0) leads to O₂-free reaction environments within the first seconds of the reaction, even when the reaction takes place in the open-air. By leveraging this significantly fast O₂-reducing activity of the disproportionation reaction, a range of well-defined water-soluble polymers with narrow dispersity are attained in a few minutes or less. This methodology provides the ability to prepare block copolymers via sequential monomer addition with little evidence for chain termination over the lifetime of the polymerization and allows for the synthesis of star-shaped polymers with the use of multi-functional initiators. The mechanism of self-deoxygenation is elucidated with the use of various characterization tools, and the species that participate in the rapid oxygen consumption is identified and discussed in detail.

Introduction

Reversible Deactivation Radical Polymerization (RDRP) methods, including atom-transfer radical polymerization (ATRP),1,4 single electron transfer-living radical polymerization (SET-LRP),4,4 reversible addition–fragmentation chain-transfer polymerization (RAFT)5,6 and nitroxide-mediated polymerization (NMP)7–9 have provided access to an increasing range of well-defined materials with sophisticated architectures, various functionalities and controlled (macro)molecular characteristics.10–12 Until recently, a notable hindrance for their development has been oxygen intolerance. Consequently, controlled radical polymerization processes conducted in the presence of oxygen are inhibited due to the ability of oxygen to react with carbon-centred radicals, leading to the formation of peroxo radicals via side reactions.13,14

Although the traditionally applied deoxygenation approaches, including freeze–pump–thaw cycles and N₂/Ar sparging are unarguably efficient for O₂ removal, they can be disadvantageous when volatile reagents are deoxygenated leading to their evaporation, and they are incompatible with new polymerization platforms (i.e. high-throughput) were the reaction scales are low.15,16 In this context, research has been focused on the replacement of these traditional methods either chemically through oxygen scavengers and reducing agents (i.e. glucose oxidase (GOx)),17–20 ascorbic acid,21–23 hydrazine,24,25 photoredox catalysts26–27 or by physically displacing oxygen through headspace elimination.28–31

In particular, the reaction medium plays an important role in the evolution of an oxygen tolerant polymerization since the concentration of dissolved O₂ varies in different solvents and can also exhibit differences depending on temperature, solution viscosity and pressure.22–24 Although oxygen tolerance and consumption have been thoroughly investigated in organic media,14,24,25,28–31,35–38 oxygen tolerant polymerization in aqueous media can be considered challenging. Apart from the high concentration of dissolved O₂ in even purified water (> 8 mg L⁻¹ at ambient temperature), there is high potential of side reactions, such as hydrolysis or elimination of the R-X or P-X bond, dissociation of the deactivating Cu(n) species when Cu-RDRP is applied,39–40 and hydrolysis of the (macro-)chain transfer agent (CTA or macro-CTA)41–43 in the case of RAFT.44,45 Consequently, oxygen tolerant polymerizations in aqueous media require the efficient removal of oxygen from the polymerization solution, and also necessitate rapid rates in order to avoid these chain termination events.

In order to avoid conventional deoxygenation in aqueous-mediated RAFT photopolymerization of N,N-dimethylacrylamide, Boyer and colleagues reported the use of a zinc porphyrin photocatalyst (ZnTPPS) with ascorbic acid as a singlet oxygen quencher.46 In a separate study, the same group demonstrated the combination of eosin Y with ascorbic acid as a reducing agent system, for the oxygen tolerant aqueous RAFT photopolymerization at low volumes.23 In the field of Cu-mediated
polymerization, among other elegant approaches.\(^47,48\) He and co-workers reported the addition of Cu(0) and ascorbic acid as reducing agents, for the surface-initiated ATRP of 2-hydroxyethyl methacrylate.\(^49,50\) Matyjaszewski and colleagues reported on the conversion of O\(_2\) to CO\(_2\) through GO\(_2\), for the aqueous ATRP of oligo(ethylene oxide) methyl ether methacrylate (OEOMA\(_{300}\)) in the presence of pyruvates\(^51\) or horseradish peroxidase.\(^22\) Percec has reported the Cu(0) wire-mediated SET-LRP of methyl acrylate in the presence of air, with the addition of hydrazine as additional reducing agent.\(^24\) Recently, Bennetti and colleagues reported an oxygen tolerant Fe(0) system for the Si-ATRP of polymer brushes where the iron acts both as a source of catalyst and as a reducing agent showing excellent cytocompatibility towards mammalian cells for preparation of biomaterials.\(^31\) Theodorou et al. have reported a versatile, oxygen tolerant photoinduced Cu-RDRP approach in emulsion for the synthesis of various protein–polymer bioconjugates, with different proteins used as macroinitiators and plastic syringes as reaction vessels, in order to efficiently eliminate headspace. Their approach was compatible with various monomers, proteins and reaction conditions, achieving well-defined bioconjugates within 2 hours.\(^34\)

Herein, we report the first instantaneously self-deoxygenating Cu-RDRP of various monomers in aqueous media, by omitting any type of conventional deoxygenation method (Scheme 1). The disproportionation of Cu(0) in the presence of Me\(_6\)Tren as a tertiary amine aliphatic sigma-donor ligand in water towards Cu(0) and Cu(II) is exploited, leading to full oxygen consumption pro reaction, a range of hydrophilic homo- and block co-polymers being achieved (Fig. 1a). Furthermore, when the PNiPAm from this non-deoxygenated reaction was compared with the N\(_2\)-sparged deoxygenation, good agreement was observed between both the two M\(_n\)SEC values and the dispersities (Table 1, Fig. S1 and S2†).

Based on previous work on Cu(0)-wire mediated-RDRP in organic solvents where it was found that elimination of the headspace was important for successful polymerization, we were interested in investigating the effect of the [O\(_2\)] concentration. The aqueous Cu-RDRP of NiPAm with targeted DP\(_n\) = 50 was tested under open to air conditions, performing both the disproportionation of Cu(0) and the polymerization reaction without sealing the vial, whilst applying a fast stirring rate (900 rpm). Although the polymerization reached near-quantitative conversion (>99%) after 12 minutes (similarly to the sealed reaction) an induction period, ascribed to continuous O\(_2\) diffusion into the reaction was observed, which was further assisted by the rapid stirring, as discussed in the mechanistic section below. Importantly, the robustness of the system was further verified when higher molecular weights were targeted, applying the same conditions i.e. low monomer concentration of 10 w/v % and lower temperature without any external deoxygenation. It has previously been reported that when DP\(_n\) ≥ 80 was targeted, the ratio [Cu(i)Br] : [Me\(_6\)Tren] = 1 : 1 resulted in inefficient deactivation, which was observed as high dispersity values.\(^26\) Therefore, for PNiPAm with targeted DP\(_n\) = 100, 200

**Results and discussion**

Initially, in order to examine the ability to carry out efficient polymerization reactions in the presence of oxygen, the aqueous Cu-RDRP of N-isopropylacrylamide (NiPAm) was conducted without any type of external deoxygenation (Scheme S1†), using only headspace elimination, following the conditions [I] : [NiPAm] : [Cu(i)Br] : [Me\(_6\)Tren] = 1 : 50 : 0.4 : 0.4. For this purpose, a sealed 8 mL glass vial was charged with Cu(i)Br, 1 mL H\(_2\)O and Me\(_6\)Tren, and was placed in ice-bath with rapid stirring (900 rpm) for 60 seconds. Upon formation of a heterogeneous blue solution (indicative of the formation [Cu(H\(_2\)O)\(_6\)]\(^{2+}\)) with a black-purple Cu(0) precipitate, an aqueous solution of NiPAm and the water-soluble initiator (2,3-dihydroxypropyl 2-bromo-2-methylpropanoate) were added to the disproportionation solution, and the polymerization was left to commence. Kinetic studies were performed to reveal rapid polymerization rates, with >99% monomer conversion only after 12 minutes in accordance with previous results,\(^35\) as well as low dispersity being achieved (D = 1.15) and good agreement between experimental and theoretical M\(_n\) values verifying the versatility to carry out the polymerization without prior removal of dissolved oxygen (Fig. 1a). Furthermore, when the PNiPAm from this non-deoxygenated reaction was compared with the N\(_2\)-sparged deoxygenation, good agreement was observed between both the two M\(_n\)SEC values and the dispersities (Table 1, Fig. S1 and S2†).

![Scheme 1 Schematic representation of the self-deoxygenating aqueous Cu-RDRP of acrylamides employing the pre-disproportionation of Cu(i)Br/Me\(_6\)Tren.](image-url)
and 400 twice as much Cu(i)Br as for DPₙ = 50 was used, following the conditions [I] : [M] : [Cu(i)Br] : [Me₆Tren] = 1 : 0.8 : 0.4. It should be noted that, although the polymerization with targeted DPₙ = 50 reached >99% conversion after only 12 minutes, for higher molar masses longer reaction times were required with DPₙ = 200 taking 60 minutes to reach >99% conversion and DPₙ = 400 taking up to 90 minutes (Table S1 and Fig. S3a†). Furthermore, when DPₙ = 400 was initially targeted with 10 w/v % monomer concentration, the polymerization was not successful while when 20 w/v % (thus higher catalyst concentration) was used, high conversion, low dispersity and controlled molecular weights were obtained. In no case, was background, uncontrolled, polymerization observed, as verified by the SEC chromatograms from dual angle light scattering and viscosity detectors across the entire calibrated region (Fig. S3b†). The versatility of this methodology was expanded with the implementation of this aqueous self-deoxygenating system to both acrylates and acrylamides (Table 2), as well as more complex polymer architectures including star-shaped polymers.

NiPAm, N-hydroxyethyl acrylamide (HEAm) (Fig. S4†), N,N-dimethylacrylamide (DMA) (Fig. S5†), poly(ethylene glycol) methyl ethyl acrylate (PEGA₄₈₀) (Fig. S6†) and N-acryloyl-oxymorpholine (NAM) (Fig. S7†) were polymerized through aqueous Cu-RDRP, without applying any type of external deoxygenation, resulting in excellent control over the obtained Mn,SEC values, with low dispersity and near-quantitative conversions over short reaction times (Table 2). Additionally, since star-shaped polymers have gained a lot of interest due to their diverse properties, the self-deoxygenating approach was applied for the synthesis of star-shaped PHEAm polymers with overall targeted DPₙ = 60, 80 and 120.

For this purpose, 3-, 4- and 8-arm multifunctional initiators were used for the synthesis of PHEAm star-shaped polymers with each arm having a targeted DPₙ = 20. As the multi-arm star initiators were relatively hydrophobic, polymerizations were

![Fig. 1](image_url)  
**Fig. 1** Kinetic studies for the self-deoxygenating aqueous Cu-RDRP of NiPAm with targeted DPₙ = 50 when the polymerization was carried out (a) in sealed vial and (b) open-to-air conditions with ¹H NMR spectra (left), plots of conversion and ln([M]₀/[M]) over time (middle) and DMF-SEC derived molecular weight distributions (right).

<table>
<thead>
<tr>
<th>Conditions</th>
<th>Conv. (%)</th>
<th>Mn,SEC (g mol⁻¹)</th>
<th>Mₙ,th (g mol⁻¹)</th>
<th>D</th>
<th>Reaction time</th>
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<tr>
<td>N₂ sparging</td>
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<td>6200</td>
<td>5900</td>
<td>1.10</td>
<td>12 min</td>
</tr>
<tr>
<td>ND/sealed</td>
<td>&gt;99</td>
<td>7200</td>
<td>5900</td>
<td>1.13</td>
<td>12 min</td>
</tr>
<tr>
<td>ND/open-air</td>
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<td>6800</td>
<td>5900</td>
<td>1.12</td>
<td>12 min</td>
</tr>
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</table>

In all polymerizations, the monomer concentration was 10 w/v % and the conditions were [I] : [DPₙ] : [Cu(i)Br] : [Me₆Tren] = 1 : 50 : 0.4 : 0.4.

Conversion was calculated via ¹H NMR in D₂O. *Determined by DMF-SEC analysis and expressed as molecular weight equivalents to PMMA narrow molecular weight standards.

<table>
<thead>
<tr>
<th>Polymer</th>
<th>Conv. (%)</th>
<th>Mn,SEC (g mol⁻¹)</th>
<th>Mₙ,th (g mol⁻¹)</th>
<th>D</th>
<th>Time (min)</th>
</tr>
</thead>
</table>
| NiPAm₅₀  
| b | >99 | 16 200 | 11 600 | 1.14 | 15 |
| PHEAm₅₀  
| b | >99 | 26 200 | 11 800 | 1.17 | 15 |
| PDMA₈₀  
| b | >99 | 11 300 | 8200 | 1.15 | 20 |
| P(PEGA₄₈₀)₉₀  
| c | 98 | 9300 | 9600 | 1.17 | 30 |
| PNAM₄₀  
| b | 98 | 7100 | 5800 | 1.09 | 240 |

The monomer concentration = 10 w/v %, [Cu(i)Br] : [Me₆Tren] = [0.8] : [0.4]. *Determined by DMF-SEC analysis and expressed as molecular weight equivalents to PMMA narrow molecular weight standards.
conducted in water – organic solvent mixtures (either methanol or DMSO), with the pre-disproportionation of Cu(i) being carried out in pure water in each case. As a result, PHEAm star-shaped polymers were obtained at near-quantitative conversions (>99%) and narrow molecular weight distributions (Đ = 1.11–1.2) in less than 2 hours (Fig. S8†). It should be noted that the higher than the theoretical \( M_n,\text{SEC} \) values for both linear and star-shaped PHEAm are attributed to interactions of the two -OH groups with the SEC-column and such deviations are observed for all the HEAm-derived polymers.

Given the non-deoxygenated environment, as well as the side reactions which are well-known to occur in aqueous media, we were interested in investigating the extent of end group fidelity in this approach. For this purpose, MALDI-ToF-MS was employed for the mass characterization of PNiPAm with targeted DPₙ = 50, synthesized in a sealed vial. The MALDI-ToF analysis revealed that there are peak distributions corresponding to polymer chains that had undergone both elimination and hydrolysis of the alkyl halide, whilst bromine capped chains were also observed (Fig. S9†). These chain termination events are also observed under conventionally deoxygenated systems originating from the excess of water where the rate of hydrolysis is lower than the rate of chain propagation, thus they cannot be correlated only with the presence or absence of oxygen. However, the analysis of the \(^1\text{H} \text{NMR} \) spectra of both PNiPAm which was analyzed through MALDI and the extreme case of PNiPAm synthesized in open air, showed no peaks corresponding to vinyl protons originating from elimination (Fig. S9b†). This observation led us to hypothesize that elimination might occur either during the MALDI process or during the sample preparation, and possibly is not occurring in the timescale of the polymerization. This hypothesis was further verified by performing in situ block copolymerizations which resulted in polyacrylamide diblock copolymers with well-defined molecular characteristics at high conversions (Fig. 2a and b). Although this rapid polymerization without loss of end group fidelity via radical- radical termination is not in accordance with classical free radical polymerization kinetics, Ballard and Asua have shown that when the probability density functions of the termination reactions are altered to allow for radical diffusion, the reduced rate of termination can explain “the ability for a seemingly impossible level of control of radical reactions”. The versatility of the methodology was further verified through low volume reactions carried out in 96-well plates which were sealed with a plate-lid. Both PNiPAm and HEAm were polymerized in 300 µL total reaction volume, exhibiting good control over the molecular weights and the dispersity at high conversion (Fig. S10†).

Rate of oxygen consumption during Cu(І) disproportionation to Cu(ІІ) and Cu(0)

The rate of the oxygen reducing activity of the Cu(І)/Me₆Tren aqueous solution was investigated via the online monitoring of the dissolved \( \text{O}_2 \) concentration with the use of an oxygen probe. Initially, the [dissolved \( \text{O}_2 \)] in the disproportionation solution used for the polymerization of NiPAm with targeted DPₙ = 50 (with [Cu(I)]:[Me₆Tren] = 1:1) was monitored, resulting in rapid (a few seconds) oxygen consumption (Fig. 3a – grey). Subsequently, since the concentration of the components able to consume oxygen affect the rate of oxygen consumption, different concentrations were examined, using 1–7 mL of H₂O (Fig. 3a).

As expected, the rate of oxygen consumption exhibited differences depending on the concentration, with the fastest oxygen consumption (~3 s) observed for the most concentrated solution (1 mL of H₂O), and the slowest rate observed for 7 mL of H₂O (~1 min). It is noted that when 1 mL of the solution was used prior to monomer addition (HEAm in this case) and initiator addition, the polymerization exhibited the highest control over the molecular weights (Fig. S11 and Table S2†). This suggests that the rate of an aqueous non-deoxygenated polymerization is dependent on the concentration of the disproportionation reagents (Cu(I)Br/Me₆Tren) and/or their products (Cu(0), Cu(ІII)), and sufficient amount of these species is important in order to facilitate the \( \text{O}_2 \) consumption process. In order to provide an insight into the open-to-air polymerization where almost perfect control over the macromolecular characteristics was observed, the [O₂] monitoring was performed in an open vial.
In this case, the oxygen consumption was again rapid (~10–60 s) (Fig. 3b), with the 1 mL solution becoming self-deoxygenated within the first 10 seconds. The trend of consumption exhibited the same profile as in the sealed vial, again depending on the concentration of the disproportionation solution. The main difference between the sealed and the open-to-air experiments was that the latter required slightly longer reaction times for total deoxygenation to occur, which could be attributed to the constant diffusion of O₂ into the solution from the air during high stirring rates (Fig. 3a and b). It should be noted that at ~0 °C (temperature applied at the disproportionation and polymerization reactions), the solubility of oxygen is higher than at ambient (i.e. 25 °C) temperature (~14 mg L⁻¹ and ~8 mg L⁻¹ respectively), while the diffusion rate of oxygen is lower.⁴⁴–⁴⁶ Hence, although there is a constant exposure of the reaction solution to air/oxygen, the reducing ability of Cu(0) (as well as Cu(I)), combined with the low-temperature conditions could further facilitate the oxygen tolerant nature of this aqueous Cu-RDRP.

Subsequently, the role of the ligand was investigated by using different amounts of Me₆Tren with respect to Cu(I)Br. Initially, a solution containing only Cu(I)Br and 3 mL H₂O was examined which, although being heterogeneous (due to the insolubility of Cu(I)Br in water), a very slow O₂ consumption was observed (from 14 mg L⁻¹ to ~13 mg L⁻¹ after 1 min). However, with the addition of low loadings of Me₆Tren ([Cu(I)Br] : [Me₆Tren] = 1 : 0.125) full O₂ consumption was observed after ~40 s (Fig. 3d – green and pink). Following this, higher concentrations of Me₆Tren resulted in accelerated consumption rates (Fig. 3d), with the fastest O₂ consumption being observed when excess of Me₆Tren was applied ([Cu(I)Br] : [Me₆Tren] = 1 : 2.5, Fig. 3d – dark cyan). This indicates that, upon formation of the Cu-complex with the amine ligand in water, the rapid disproportionation of Cu(I) is followed by rapid O₂ consumption, even when low loadings of Cu(I)Br or ligand are used. In this context, it is speculated that even small amounts of the in situ generated Cu(0) particles can “consume” oxygen. However, since the “nascent” Cu(0) particles are highly reactive, we hypothesized that their generation is followed by their oxidation in the presence of O₂ (vide infra electron microscopy and XPS).

In order to examine which of the Cu-species following disproportionation participate in the O₂ consumption, the [O₂] evolution was also monitored in different solvents including ethanol, methanol and dimethylformamide (DMF) which promote disproportionation of Cu(I),⁴⁷ as well as solvents that stabilize Cu(I) and thus, disproportionation is not favoured (i.e. acetone, acetonitrile, toluene) (Fig. 3e). Specifically, when EtOH and MeOH that facilitate the disproportionation of Cu(I) were used, the fastest O₂-consumption rates were observed. Noteworthy is that the same trend was observed when MeCN, which favours the stabilization of Cu(I), was used. Slower consumption was evidenced when acetone, H₂O and the polar

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Fig. 3 Line graphs illustrating the oxygen consumption over time with (a) different concentrations of the Cu(I)Br/Me₆Tren complex in a sealed environment (with [Cu(I)Br] : [Me₆Tren] = 1 : 1) and (b) open-to-air conditions (with [Cu(I)Br] : [Me₆Tren] = 1 : 1), (c) various concentrations of Cu(I)Br in 3 mL reaction volume, (d) various concentrations of Me₆Tren in 3 mL reaction volume, (e) in different solvents with 3 mL reaction volume and (f) upon addition of 6 mL aqueous solution of monomer and initiator in 1 mL disproportionation reaction. All measurements were conducted in glass vials with 7 mL capacity, at ~0 °C and 900 rpm stirring rate.
apo tic solvent DMF were used, while toluene where Cu(I) is insoluble, exhibited the slowest rates. Since acetonitrile stabil ises Cu(I) and does not promote disproportionation, we speculate that O₂ consumption occurs from the oxidation of Cu(I) to Cu(II). Based on this, it is possible that O₂ consumption is assisted by the oxidation of both Cu(I) and the “nascent” Cu(0), and depending on the choice of solvent, the oxidation of Cu(0) and Cu(I) are competing reactions which can both lead to rapid deoxygenation. The continuous deoxygenation profile (even after the addition of monomer and initiator solutions the [O₂] has been reduced, and the O₂-free environment is re-established) (Fig. 3f) might be attributed to the regeneration of Cu(I) upon oxidation of Cu(0), which can re-disproportionate and thus “perpetuate” the reduction of O₂ until the reagents are fully consumed. As a result, it is hypothesized that Cu(I), originating either from the initially added Cu(I)Br or from the oxidation of Cu(0), and “nascent” Cu(0) participate in the consumption of both dissolved and in gas phase O₂, and due to the long-lasting paucity of these Cu-species, the reaction solutions remain deoxygenated even open-to-air.

To better understand the effect of O₂ on the catalyst and in an attempt to identify the oxidation products, XPS and EM (TEM, SEM, ADF-STEM and EELS) were employed. Initially, the solution following disproportionation in the presence of O₂ was examined through SEM and aiming to monitor any changes in the morphology of the Cu particles over time, Cu(I)Br in water and aliquots of the disproportionation reaction were imaged (Fig. 4a–h). The Cu(I)Br sample in water consists of >1 μm size well-shaped aggregates. Upon addition of Me₆Tren and after the ~1 s second of the reaction, where there is an instantaneous observation of black/purple Cu(0) precipitate with fast stirring, the sample mainly includes three different structures, namely dendrite-like shaped aggregates which consist of particles >100 nm, small multi-sized particles, as well as faceted crystal structures (Fig. 4b). It is notable that the crystal-shaped morphologies are mainly present for the sample taken immediately after the addition of Me₆Tren (“~1 s” sample).

Considering the loss of the larger Cu(I)Br aggregates, it is hypothesized that the crystals are formed upon instant consumption of Cu(I)Br, following a clusterization process (Fig. S12 and S13†) which can be correlated with the formation of the dendrite-like shaped aggregates. After ~2 seconds of the reaction, the observed faceted structures are significantly less, whilst the dendritic-like aggregates, which we hypothesize that follow a nucleation and growth process become more evident (Fig. 4c–e). It is notable that this morphology is still observed after 60 seconds of the reaction, while after 5 minutes this dendritic pattern becomes “softer”, with the dendritic-like branches becoming less evident and more uniform (Fig. 4f). Finally, after 30 minutes, small particle-like aggregates are observed which are covered by more uniform larger aggregates (Fig. 4g) and finally, after 24 hours of the reaction, large aggregates that consist of smaller particles are observed (Fig. 4h).

It is hypothesized that the consumption of O₂ would lead to the possible formation of different copper oxides and thus, the EDX studies were focused on the presence of oxygen with respect to copper. Initially, Cu(I)Br was measured as blank sample, showing a distinctive peak at 0.9 keV corresponding to Cu, as well as a very small peak assigned to oxygen at 0.5 keV (Fig. S14†) and could be attributed to the oxidation of the Cu(I)Br powder from air over time. When the precipitate from the disproportionation was examined immediately after the addition of Me₆Tren (sample was taken after ~1 second), the peak at 0.5 keV, assigned to oxygen increased slightly (as well as after ~2–3 seconds) and became even more evident after 60 seconds of the reaction (Fig. S14†). This observation might be related to the formation of oxides as the reaction takes place in the presence of oxygen, a hypothesis that is corroborated by the full O₂ consumption within the first 60 seconds. After 30 minutes, the intensity of the oxygen peak further increased, with the highest intensity being observed after 24 hours. The presence of copper oxides and hydroxide was also verified through X-ray photoelectron spectroscopy (XPS) for both the black precipitate and the supernatant after 60 s. Examination of the Cu 2p₃/2 region revealed the presence of Cu(II) states in the precipitate (Fig. 5a), as evidenced by the shake-up features observed between 940 eV and 945 eV. Detailed fitting of this region and the peak around 934 eV using the work of Biesinger, showed

![Fig. 4](image-url)

**Fig. 4** SEM images of (a) Cu(I)Br dispersed in H₂O and (b–h) the self-deoxygenating disproportionation reaction precipitate collected at different times. The “~1 s” was taken immediately after the addition of Me₆Tren.
Fig. 5 XPS core level (a) Cu 2p3/2 and (b) O 1s spectra of the disproportionation precipitate after 60 seconds of the reaction. The features between 940 eV and 945 eV (a) are due to shake-up peaks from Cu(II) states.

Evidence for Cu(OH)$_2$, CuO and Cu(II)Br$_2$. Turning to the peak at 932.3 eV, it is not possible to distinguish between Cu(0) and Cu(n) states so one must look at alternative regions in order to understand the chemistry of the system. Fig. S15† presents the data from the Br 3d region, where two doublets were required to fit the data corresponding to Cu(II)Br and Cu(II)Br$_2$.

Next, the Cu LMM Auger emission region was analysed (Fig. S16†) and required the addition of the Cu(0) components as described by Biesinger to replicate the data.† Comparing the Auger spectra of the precipitate and the supernatant, we observe a downward shift of around 2 eV in the kinetic energy of the peak intensity when moving from the precipitate to the supernatant, suggesting a higher Cu(n) concentration in the supernatant. This hypothesis is corroborated by the Cu 2p3/2 spectrum acquired from the supernatant (Fig. S17a†), where the intensity of the shake-up features and the peak at 934.2 eV have both increased relative to the precipitate (Fig. 5a).

The O 1s data acquired also suggest the existence of both Cu(OH)$_2$ and Cu oxides in both the supernatant (Fig. S17b†) and the precipitate (Fig. 5b). Overall, based on the elemental analysis of both the precipitate and supernatant (Tables S3 and S4†), copper and oxygen are mainly predominant in the precipitate, while nitrogen originating from the ligand (Fig. S18†), as well as bromine (originating from Cu(II)Br) are mainly present in the supernatant (Fig. S15†).

High-resolution transmission electron microscopy (HR-TEM) and annular dark field-scanning transmission electron microscopy (ADF-STEM) in combination with electron energy loss spectroscopy (EELS) were used in order to provide more details on the chemical state of the Cu-species generated upon reaction with oxygen. The HR-TEM of the disproportionation precipitate (60 seconds aliquot) verified the presence of copper species with dramatically different morphology in the nanoscale, i.e. dendrite-like shaped structures (Fig. 6c and g), big particles with well-defined facets (Fig. 6d and h) and much smaller particles (Fig. 6a, b, e and f).

Based on the EELS analysis, the faceted particles exhibited a Cu–L$_2$ edge peaked at 935.9 eV (Fig. 6j, blue) but no detectable O–K edge. This suggests that these particles are metallic Cu, which has remained unaffected from the presence of oxygen within the first 60 seconds of the reaction. In contrast, the dendrite-like structures exhibited a prominent O–K edge peaked at 533.8 eV and a Cu–L$_3$ edge peaked at 936.2 eV, indicating the formation of Cu(n) materials (Fig. 6j, red).†

It is therefore speculated that these dendritic structures exist in the form of CuO which, based on their immediate presence upon addition of the ligand (after ~1 second), were rapidly formed. The Cu–L$_3$ edge from the smaller (<80 nm) particles is shifted to a lower energy loss of 933.5 eV (Fig. 6j, yellow). In combination with the energy loss features shown in the O–K edge (Fig. 6j, yellow), it is confirmed that these particles are CuO. In addition, even smaller (2–10 nm) particles were found (Fig. 6b and f) within the samples. They show a Cu–L$_3$ edge corresponding to Cu(n) and a not clearly defined O–K edge. However, the analysis of the relative intensities between the O and Cu edges revealed that there is shortage of O within the very small particles and they are in fact a mixture of Cu(n) and Cu(0).

In summary, the fast oxygen consumption rates observed from the online O$_2$-monitoring suggest that the solution following disproportionation can be considered as a rapidly self-deoxygenating system. In order to investigate the effect of the ligand and the Cu-source (Cu(n)Br) on oxygen consumption, different concentrations of these components were examined individually. In the absence of Cu(n)Br (when only Me$_6$Tren is present) no oxygen consumption was observed over the timescale of the disproportionation reaction, while a gradual
increase of the [Cu(i)Br] led to progressively faster rate of O2-consumption. In the absence of ligand (when only Cu(i)Br was present), a slightly descending trend was observed which became evident with the addition of even low loadings of Me2Tren. It is noteworthy that although the different concentrations of Cu(i)Br have a gradual effect on the consumption rate, alterations on the ligand concentrations did not show the same effect. This observation leads us to hypothesize that the rate of O2-consumption is mainly dependent on the concentration of Cu(i) which is directly related to the concentration of in situ generated highly reactive Cu(0) particles. Although the ligand cannot be considered as the primary O2-reducing agent based on these findings, its role is highlighted when negligible oxygen levels are observed even after the addition of monomer and initiator. This might be attributed to the ability of the ligand either to reduce Cu(iii) back to Cu(i), or re-initiate a new “disproportionation” cycle, thus leading to elimination of dissolved O2. To further clarify the role of the disproportionation components, the O2-consumption profile of the reaction was investigated in various solvents. It was shown that fast O2-consumption rates were observed not only for solvents that favour disproportionation of Cu(i) into Cu(0) and Cu(II), but also for solvents that stabilize Cu(i) i.e. acetoniitrile. These data indicate that both Cu(i) and Cu(0) can consume oxygen via being oxidised, and the oxidation of Cu(0) and Cu(i) are competing reactions which both lead to rapid deoxygenation.

These observations were further corroborated when electron microscopy (SEM, HR-TEM) and spectroscopic (XPS, EDX, EELS) techniques were collectively employed for the determination of the oxidized products. SEM along with EDX analyses indicated that within the first seconds of the reaction there is formation of copper oxides which becomes more evident as the reaction progresses, supporting our hypothesis that the copper species are primarily responsible for oxygen consumption. The morphological differences of the copper species facilitated further analysis through HR-TEM and EELS spectroscopy, which identified the different copper species. Characterization of both the reaction supernatant and precipitate through XPS revealed again that copper oxides are the main products of this self-deoxygenation process.

Conclusions

The rapidly self-deoxygenating Cu-RDRP in aqueous media of various linear and non-linear, homo- and block-copolymers is reported. Disproportionation of Cu(i)/Me2Tren in water towards Cu(ii) and highly reactive Cu(0), leads to O2-free reaction environments within the first seconds of the reaction, even when the reaction is taking place in the open-air. By leveraging this significantly fast O2-reducing activity of the disproportionation reaction, well-defined water-soluble polymers with very narrow dispersity are attained in a few minutes or less. Importantly, this methodology provides the ability to prepare block copolymers via sequential monomer addition with little evidence for chain termination over the lifetime of the polymerization and allows for the synthesis of star-shaped polymers with the use of multi-functional initiators. The use of a range of characterization tools provides insights into this self-deoxygenating platform and identifies the species that participate in the oxygen consumption, as well as the species generated upon exposure of the solution to O2-rich environments. The unprecedentedly fast reducing ability of the Cu(i) disproportionation can be an advantageous and mild platform for applications that require instantaneous O2-free environments for long periods.

Conflicts of interest

There are no conflicts to declare.

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References