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Insights into water coordination associated to the Cu$^{II}$/Cu$^1$ electron transfer at a biomimetic Cu centre

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The coordination properties of the biomimetic complex [Cu(TMPA)(H$_2$O)](CF$_3$SO$_2$)$_2$ (TMPA=tris(2-pyridylmethyl)amine) have been investigated by electrochemistry combined to UV-Vis and EPR spectroscopies in different non-coordinating media including imidazolium-based room-temperature ionic liquids, for different water contents. The solid-state X-ray diffraction analysis of the complex shows that the cupric centre lies in a N$_4$O coordination environment with a nearly perfect trigonal bipyramidal geometry (TBP), the water ligand being axially coordinated to Cu$^{II}$. In solution, the coordination geometry of the complex remains TBP in all media. Neither the triflate ion nor the anions of the ionic liquids were found to coordinate the copper centre. Cyclic voltammetry in all media shows that the decoordination of the water molecule occurs upon mono-electronic reduction of the Cu$^{II}$ complex. Back-coordination of the water ligand at the cuprous state can be detected by increasing the water content and/or decreasing the timescale of the experiment. Numerical simulations of the voltammograms allow the determination of kinetics and thermodynamics for the water association-dissociation mechanism. The resulting data suggest that (i) the binding/unbinding of water at the Cu$^{II}$ redox state is relatively slow and equilibrated in all media, and (ii) the binding of water at Cu$^1$ is somewhat faster in the ionic liquids than in the non-coordinating solvents, while the decoordination process is weakly sensitive to the nature of the solvents. These results suggest that ionic liquids favour water exchange without interfering with the coordination sphere of the metal centre. This makes them promising media for studying host-guest reactions with biomimetic complexes.

Introduction

Reorganizational processes coupled to electron transfer are of particular importance in many chemical and biological systems associated to the Cu$^{II}$/Cu$^1$ reaction.1 Such processes are due to crystal field forces and Jahn-Teller effects when passing from Cu$^1$ (d$^{19}$) to Cu$^{II}$ (d$^9$). They involve most of the time the coordination/decoordination of one or several donor ligands together with geometric changes. The Cu$^1$ (d$^{19}$) ion is quite “plastic” toward its environment and can readily adopt 2-, 3- or 4-coordinated geometry. Contrariwise, the Cu$^{II}$ (d$^9$) ion has a strong preference for tetragonal environments, leading to 5- and 6-coordinated cores, and Cu$^{II}$ can exert a strong electronic/geometric effect on its surroundings in the absence of sterical constraints. Many studies have been devoted to understand/control this chemical upheaval with the variation of the electronic/geometric/steric environment of the copper centre, using mainly synthetic Cu complexes as model systems.1 One of the most popular system is the mononuclear complex [Cu$^{II}$(TMPA)(L)]$^{2+}$ (L=solvent), based on the tripodal tetradentate ligand TMPA (TMPA = tris(2-pyridylmethyl)amine), which is a ligand of choice for mimicking metallo enzymes of relevance to oxygen activation, for instance. This complex and its derivatives have been studied in depth over the past years since they react at Cu$^1$ redox state with dioxygen to form transient Cu$_2$O$_2$ reactive species.2,3,4 Recent works have shown that these derivatives were able to activate dioxygen, leading to the catalytic oxidation of organic substrates as homogeneous catalysts5,6,7, or to reduce O$_2$ into H$_2$O by heterogeneous catalysis.8 Also, in acidic aqueous media, catalytic properties towards reduction of nitrite ions into nitric oxide were emphasized by voltammetric studies for [Cu$^{II}$(TMPA)(H$_2$O)]$^+$,8 In another connection, Le Mest et al. have used the nitrito Cu complex [Cu$^{II}$(TMPA)(CH$_3$CN)]$^{2+}$ for evidencing secondary-sphere effects at the redox level by comparison with a more sophisticated complex.10 The chemical modifications associated to the electronic transfer are usually detected by complementing spectroscopic studies (NMR, EPR, UV-Vis...) with electrochemical techniques (often cyclic voltammetry). In almost all cases, the electrochemical studies of [Cu$^{II}$(TMPA)(L)]$^{2+}$ were performed in coordinating solvents (DMF, CH$_3$CN, H$_2$O) and/or in presence of coordinating anion (Cl$^-$, N$_3^-$). The purpose of these studies was essentially to compare the donating effect of each ligand by correlation with the thermodynamic standard potential value $E^{0}$ in the same solvents,
the redox potential being highly influenced by electronic, steric and geometric factors. So far, none of the voltammetric studies in solution have ever intended to investigate the chemical mechanisms associated to the Cu\(\text{II}\) electron exchange. Particularly the water ligand(s) decoordination process at Cu\(^{\text{II}}\) redox state has not been explored by electrochemistry, despite its high interest. Indeed, water plays an important role in many enzymatic processes.\(^{11}\) In particular, it was shown that water could inhibit oxygen reaction in cytochrome c oxidase models.\(^{12}\) Also, it can affect the redox potential of proteins through secondary sphere effects.\(^{13}\) Recently, water molecules in the active site of Cu-nitrite reductase, were shown to specifically protect from nitrite coordination at Cu(II) and favor catalytic activation at Cu(I) by displacement of the water ligand.\(^{14}\) Water dissociation is thus of fundamental importance in the activation process. Here we propose to explore such a phenomenon with water as an exogenous ligand with a TMPA Cu complex. Herein, with the help of electrochemistry combined to EPR/UV-Vis spectroscopies, we report a detailed study of the behaviour of the [Cu\(^{\text{II}}\)(TMPA)(H\(_2\))](CF\(_3\)SO\(_2\))\(_2\) complex upon electron transfer in different non-coordinating media including dichloromethane (CH\(_2\)Cl\(_2\)), tetrahydrofuran (THF), propylene carbonate (PC) and imidazolium-based room temperature ionic liquids (RTILs) (Scheme 1). This study is aimed at investigating the chemical processes associated to the Cu\(^{\text{II}}\) electron transfer reaction, particularly the possible release of the water molecule at the Cu\(^{\text{II}}\) redox state in terms of kinetics and thermodynamics. Ionic liquids are liquid at room temperature (or close to) and are composed entirely of ions. Generally consisting in a combination of bulky organic cations and weakly coordinating anions, they have been scarcely considered as solvents for analyzing the electrochemical reactivity of coordination metal complexes.\(^{15}\) A major advantage of the ionic liquids is the ease of change of the anion/cation combination, allowing the preparation of highly “tunable” media especially designed for a specific application. As such, they nowadays cover a wide range of applications including synthetic chemistry, catalysis, (bio) analytical sensors, and material science.\(^{15,16}\) They have been widely used as solvents in fundamental electrochemical reactions.\(^{17}\) However ionic liquids have been scarcely considered as solvents for analysing the electrochemical reactivity of coordination metal complexes.\(^{18}\) While they can represent advantageous alternative to conventional solvents due to their intrinsic properties, their specific nature may also influence the coordination properties of the complexes. Thus, some studies have explored the coordinating properties of the anion from the ionic liquids towards metallic cations (Cu\(^{2+}\), Li\(^+\)).\(^{19,20}\) Nevertheless, the influence of the RTILs as secondary sphere on the coordination properties of metallic complexes associated to their redox properties has not been investigated so far.

**Results and Discussion**

![Fig. 1](image)

**Scheme 1.** Representation of the RTILs used in this study.

**Synthesis and structural properties of the [Cu\(^{\text{II}}\)(TMPA)(H\(_2\))](CF\(_3\)SO\(_2\))\(_2\) complex.**

The synthesis of the bis-triflate Cu complex [Cu\(^{\text{II}}\)(TMPA)(H\(_2\))](CF\(_3\)SO\(_2\))\(_2\) was inspired by the procedure developed for the bis-perchlorato analogous complex [Cu\(^{\text{II}}\)(TMPA)(H\(_2\))](ClO\(_4\))\(_2\). Blue single crystals were obtained and analysed by X-ray diffraction. As shown in Fig. 1, the cupric centre lies in a N\(_2\)O coordination mode with a nearly perfect trigonal bipyramidal geometry (TB) (\(\tau = 0.95\)),\(^{21}\) the O(7)-Cu-N(4) angle being almost linear (176.91°) (see Table 1). Clearly, none of both triflate anions (CF\(_3\)SO\(_2\))\(_2\) is coordinated to the Cu centre as previously reported for the bis-perchlorato analogous complex.\(^{9}\) Cu-N distances in the trigonal plane are almost equal (2.06 Å), and N-Cu-N angles are close to the ideal 120° value. In the axial part, Cu-N(4) and Cu-O(7) distances are shorter (2.01 Å and 1.95 Å respectively) depicting strong interaction of the amine and the water ligands with the metal ion. These structural features are similar to those found for the perchlorato derivative [Cu(TMPA)(H\(_2\)))](ClO\(_4\))\(_2\) (\(\tau = 0.96\)),\(^{9}\) emphasizing the minor role of the counter anions on the structural properties at the solid state.

**Table 1** Selected bond distances and angles for [Cu\(^{\text{II}}\)(TMPA)(H\(_2\))](CF\(_3\)SO\(_2\))\(_2\). See Supporting Information part for details

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<thead>
<tr>
<th>Bond</th>
<th>Distance Å</th>
<th>Angle /°</th>
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<tbody>
<tr>
<td>Cu-N(1)</td>
<td>2.060(3)</td>
<td>98.5(1)</td>
</tr>
<tr>
<td>Cu-N(2)</td>
<td>2.058(3)</td>
<td>94.8(1)</td>
</tr>
<tr>
<td>Cu-N(3)</td>
<td>2.069(3)</td>
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<tr>
<td>Cu-N(4)</td>
<td>2.011(3)</td>
<td>176.9(1)</td>
</tr>
<tr>
<td>Cu-O(7)</td>
<td>1.945(8)</td>
<td>119.9(1)</td>
</tr>
<tr>
<td>N(1)-Cu-N(2)</td>
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<td></td>
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<tr>
<td>N(1)-Cu-N(3)</td>
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<td></td>
</tr>
<tr>
<td>N(2)-Cu-N(3)</td>
<td>117.4(1)</td>
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<tr>
<td>N(2)-Cu-N(4)</td>
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<tr>
<td>N(3)-Cu-N(4)</td>
<td>82.2(1)</td>
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</tr>
</tbody>
</table>
Spectroscopic studies in solution.

The coordination properties of [Cu(TMPA)(H₂O)](CF₃SO₃)₂ were examined by Electron Paramagnetic Resonance (EPR) and UV-Visible spectroscopies in three non-coordinating solvents (CH₂Cl₂, THF and PC) and in three different ionic liquids where both anions and cations were varied ([BMIm][NTf₂], [BMIm][PF₆] and [Et₃BuN][NTf₂]) (Scheme 1). Addition of coordinating species such as H₂O and CH₃CN was performed at room temperature for a full characterization of the coordinating environment around Cu²⁺, with a particular focus on the nature of the fifth exogenous ligand. Whatever the non-coordinating media, [Cu²⁺(TMPA)(H₂O)]²⁺ displays an EPR reversed axial spectral pattern with gₓ > gᵧ ≈ 2.0 (Fig. 2A and SI). The hyperfine coupling constants Aₓ have small values in the range 100-120 G (Table 2). These features are typical of a Cu¹⁺ complex in a TBP geometry with a dz² ground state, as previously observed for the nitrilo analogue [Cu¹⁺(TMPA)(CH₃CN)]²⁺ in frozen acetonitrile. The g and A values are slightly different than those obtained in acetonitrile (gₓ = 2.19, gᵧ = 2.02, Aₓ = 113 10⁻⁴ cm⁻¹). Moreover, the addition of water at room temperature did not induce any modification of the frozen solution spectra (see SI for each solvent). Hence, taking into account that the EPR data are not affected at all by the presence of water whatever the solvent and that the donor ability of THF is much weaker compared to water, it can reasonably be concluded that H₂O is the fifth ligand in axial position in THF as well as in CH₂Cl₂ and PC, and that the solid state aqua TBP structure determined by XRD is kept in solution. The same conclusion could be drawn for the ionic liquids, as the EPR responses of the complex were almost identical to those obtained in CH₂Cl₂, THF and PC (Fig. 2A). Moreover, for the three different ionic liquids media, the recorded spectra were very similar, evidencing that neither the nature of the cation (imidazolium vs ammonium) nor that of the anion (NTf₂⁻ vs PF₆⁻) do modify the coordination properties of the Cu²⁺TMPA complex in the ionic liquids. In the case of the anions, this result deserves to be emphasized because previous studies showed that the NTf₂⁻ anion, despite its non-coordinating nature, is able to bind ligand-deficient metal complexes through various nitrogen and/or oxygen binding modes.²² As for non-coordinating solvents, addition of acetonitrile to the ionic liquids yielded a change of the EPR response, indicating the substitution of H₂O by CH₃CN in the Cu²⁺ coordination sphere. UV-Vis absorption analysis is in good concordance with EPR results. In all media, d-d transition bands in the 850-950 nm wavelength range, with a shoulder at ca. 680 nm were observed. This is indicative of a large contribution of trigonal bipyramidal TBP geometry in solution, as confirmed by the solid state spectrum of the complex (Table 2).²³ Interestingly, the addition of water induces a significant shift of the λmax value, between 10 and 50 nm in all media (see SI and Table 2). For instance, in THF, a shift of 46 nm is observed by addition of 1 % of water (Fig. 2B). This clearly shows, in agreement with EPR data, that it is not correlated to the coordination of the solvent. Possibly, this effect is due to the increased polarity of the solution and solvation of the counterion and/or water. To summarize, the spectroscopic solution studies show that H₂O remains coordinated to the cupric center in all the non-coordinating media starting from the solid-state characterized [Cu²⁺(TMPA)(H₂O)]²⁺ complex. The geometric features of the Cu complex in solution are not significantly perturbed by the change

Table 2  EPR data (150 K) and UV-Vis data for [Cu²⁺(TMPA)(H₂O)](SO₃CF₃)²⁻ in the different media.

<table>
<thead>
<tr>
<th>Medium</th>
<th>[H₂O] / ppm</th>
<th>gₓ (Aₓ / 10⁻⁴ cm⁻¹)</th>
<th>gᵧ (Aᵧ / 10⁻⁴ cm⁻¹)</th>
<th>λmax / nm (ε / M⁻¹ cm⁻¹)</th>
<th>λmax / nm (ε / M⁻¹ cm⁻¹)</th>
</tr>
</thead>
<tbody>
<tr>
<td>CH₂Cl₂ dry</td>
<td>200[^a]</td>
<td>1.99 (66)</td>
<td>2.21 (103)</td>
<td>695 (54)</td>
<td>940 (249)</td>
</tr>
<tr>
<td>CH₂Cl₂ wet</td>
<td>190[^b]</td>
<td>2.00 (66)</td>
<td>2.21 (105)</td>
<td>694 (53)</td>
<td>898 (317)</td>
</tr>
<tr>
<td>THF dry</td>
<td>50[^c]</td>
<td>2.00 (64)</td>
<td>2.20 (106)</td>
<td>677 (67)</td>
<td>892 (185)</td>
</tr>
<tr>
<td>THF wet</td>
<td>190[^d]</td>
<td>2.00 (64)</td>
<td>2.20 (106)</td>
<td>667 (120)</td>
<td>846 (215)</td>
</tr>
<tr>
<td>PC dry</td>
<td>150[^e]</td>
<td>2.00 (63)</td>
<td>2.20 (106)</td>
<td>689 (69)</td>
<td>895 (183)</td>
</tr>
<tr>
<td>PC wet</td>
<td>680[^f]</td>
<td>2.00 (68)</td>
<td>2.20 (105)</td>
<td>682 (80)</td>
<td>870 (210)</td>
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<tr>
<td>[BMIm][NTf₂] dry</td>
<td>50[^g]</td>
<td>1.99 (62)</td>
<td>2.20 (103)</td>
<td>676 (52)</td>
<td>892 (120)</td>
</tr>
<tr>
<td>[BMIm][NTf₂] wet</td>
<td>360[^h]</td>
<td>0[^j]</td>
<td>0[^j]</td>
<td>0[^j]</td>
<td>0[^j]</td>
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<tr>
<td>[BMIm][PF₆] dry</td>
<td>77.5[^i]</td>
<td>0[^j]</td>
<td>0[^j]</td>
<td>0[^j]</td>
<td>0[^j]</td>
</tr>
<tr>
<td>[Et₃BuN][NTf₂] wet</td>
<td>0[^j]</td>
<td>0[^j]</td>
<td>0[^j]</td>
<td>0[^j]</td>
<td>0[^j]</td>
</tr>
<tr>
<td>Solid state</td>
<td>0[^j]</td>
<td>0[^j]</td>
<td>0[^j]</td>
<td>0[^j]</td>
<td>0[^j]</td>
</tr>
</tbody>
</table>

Corresponding concentration in mol L⁻¹: [^a] 1.5 10⁻⁴, [^b] 1.4 10⁻⁴, [^c] 2.5 10⁻⁴, [^d] 9.5 10⁻⁴, [^e] 9.9 10⁻⁴, [^f] 4.5 10⁻⁴, [^g] 4.1 10⁻⁴, [^h] 2.9 10⁻⁴, [^i] 5.8 10⁻⁴, [^j] not determined. [^j] pseudo-axial spectrum gₓ = 2.21 (117), gᵧ = 2.17 (117); gₓ = 1.99 (72), dₓ → dₓ transition; dₓ → dₓ transition.

Fig. 2. A) EPR spectra at T = 150 K of [Cu²⁺(TMPA)(H₂O)]²⁺ (10⁻⁴ mol L⁻¹) in a) “dry” CH₂Cl₂ ([H₂O] = 1.5 10⁻³ M (200 ppm)) and b) “dry” [BMIm][NTf₂] ([H₂O] = 4 10⁻⁴ M (200 ppm)). B) UV-Vis spectra of [Cu²⁺(TMPA)(H₂O)]²⁺ (1.5 10⁻⁴ mol L⁻¹) in dry (green) and wet (blue) THF.
of medium (TBP conformation) and water content. This point indicates that all voltammetric studies will be initiated with the same (geometry, coordination) complex \([\text{Cu}^{II}(\text{TMPA})(\text{H}_2\text{O})]^2+\) and is prerequisite for a comparison of the redox behavior and associated mechanisms in the different media.

In addition to their non-coordinating nature, the solvents \([\text{BMIm}][\text{NTf}_2]\) and \([\text{BMIm}][\text{PF}_6]\) allow a comparative study of the influence of the medium on the redox behaviour of \([\text{Cu}^{II}(\text{TMPA})(\text{H}_2\text{O})]^2+\). As the concentration of water is increasing, the anodic peak disappears while the redox system at \(E^0\) tends to be more reversible, even at the lowest scan rates (Fig. 4). The electrochemical behaviour of \([\text{Cu}^{II}(\text{TMPA})(\text{H}_2\text{O})]^2+\) in \(\text{CH}_2\text{Cl}_2\) is similar to those described above but it is complicated by a reaction between the cuprous complex, formed during the electrochemical reduction of the initial \(\text{Cu}^{II}\) complex, and dichloromethane to produce \([\text{Cu}^{II}(\text{TMPA})(\text{Cl})]^+\) as previously described by Karlin and co-workers. A second reversible redox process could be then observed at \(V_{0.75}\) V, as in ionic liquids (Fig. 5). Under mild conditions, the \(\text{Cu}^{II}\) complex reacts in a stoechiometric manner with alkyl or benzyls halides to produce near quantitative yields of coupled products.

Electrochemical studies of the \([\text{Cu}^{II}(\text{TMPA})(\text{H}_2\text{O})]^2+\) complex.

Cyclic voltammetric experiments were performed in the different solvents containing \(\text{NBu}_4\text{PF}_6\) as supporting salt, as well as in \([\text{BMIm}][\text{NTf}_2]\) and in \([\text{BMIm}][\text{PF}_6]\) as typical examples of ionic liquids. In addition to their non-coordinating nature, the solvents exhibit different properties of viscosity, polarity (Table 3) for allowing a comparative study of the influence of the medium on the redox behaviour of \([\text{Cu}^{II}(\text{TMPA})(\text{H}_2\text{O})]^2+\). The water content in the solvents was varied to examine the possible effect of the presence of water on the chemical processes associated to the electron transfer reaction for this aqua complex. For sake of clarity, solvents were denominated as “dry” and “wet” according to the water concentration contained in the medium as described in Table 2.
The copper center acts as the halogen atom abstractor and the halide CuII complex [CuII(TMPA)Cl]2+ is formed. This chloro-CuII complex is then reversibly reduced at -0.75 V explaining the occurrence of the second reversible redox process in CH2Cl2. The formation of the chloro complex is fully corroborated by the addition of chloride to the solution, which leads to the formation of a reversible system at the same potential.

![Graph](image)

**Fig. 5.** Cyclic voltammetry of [CuII(TMPA)H2O]2+ (~ 10^-3 mol L^-1) at a glassy carbon disk electrode at v = 0.1 V s^-1 in “dry” CH2Cl2.

**Mass transport of [CuII(TMPA)(H2O)]2+ complex.**

For each medium, under dry conditions (Table 4), the diffusion coefficient (D) value for [CuII(TMPA)(H2O)]2+ was estimated from current peak values by using the Randles-Sevcik equation.27 The diffusion coefficient decreases as the viscosity of the medium is increasing, in agreement with the Stokes-Einstein relationship (1):

\[
D = \frac{k_B T}{\eta \theta \pi n_0}
\]

where D is the diffusion coefficient, \( \eta \) is the dynamic viscosity of the medium, \( k_B \) is the Boltzmann constant, \( T \) is the absolute temperature, \( \theta \) is the hydrodynamic radius of the species, and \( \pi n_0 \) is a numerical value being equal to 6 or 4 depending if the “stick” or the “slip” conditions apply. It can be noted that the Stokes-Einstein equation applies in ionic liquids in the case where the diffusing species are similar in size with the ions of the IL but was shown not to apply for small molecules such as dihydrogen or sulphur dioxide.28 A non-Stokesian behaviour has been even evidenced for concentrated iodide in ionic liquids.29 In more recent works, Licence and co-workers have demonstrated that the Stokes-Einstein equation describes the behaviour of electroactive species such as ferrocenemethanol and ferrocenemethylidiazole in ionic liquids but the numerical value lies somewhere between 4 and 6.30 As can also be observed, the [CuII(TMPA)(H2O)]2+ complex diffuses very slowly in the two ionic liquids, \( D = 6 \times 10^{-12} \text{ m}^2 \text{ s}^{-1} \) in [BMIm][NTf2] and \( D = 0.4 \times 10^{-12} \text{ m}^2 \text{ s}^{-1} \) in [BMIm][PF6].

For comparison, the diffusion coefficient of inorganic metal complexes such as ferrocene or ferrocenemethanol have been reported as \( D = 6.3 - 46 \times 10^{-12} \text{ m}^2 \text{ s}^{-1} \) in [EMIm][NTf2] (\( \eta \approx 34 \text{ mPa s} \)) or \( D = 25 \times 10^{-12} \text{ m}^2 \text{ s}^{-1} \) in [BMIm][NTf2] and \( D = 5 \times 10^{-12} \text{ m}^2 \text{ s}^{-1} \) in [BMIm][PF6].30b Lower diffusion coefficient values have been found for a rhennium tetratetrazolato complex\(^{30b} \ (D = 5.8 \times 10^{-12} \text{ m}^2 \text{ s}^{-1} \) in [BMIm][NTf2] and \( D = 1.3 \times 10^{-12} \text{ m}^2 \text{ s}^{-1} \) in [BMIm][PF6]) or for a large polyoxometalate ion\(^{31} \ [\text{Mo}_{9}\text{O}_{26}]^{2-} \ (D = 1.4 \times 10^{-12} \text{ m}^2 \text{ s}^{-1} \) in [BMIm][PF6]).

**Variation of the redox potentials of [CuII(TMPA)(H2O)]2+ vs the media.**

As shown in Table 4, the value of the redox potential \( E^{0}_V \) is highly dependent on the medium and on the amount of water present in that medium. Addition of water results in a large negative shift of the potential values except in CH2Cl2. In this latter medium, the “dry” medium still contains a large amount of water regarding the water saturation value (1.5 \times 10^{-3} \text{ mol L}^{-1}) in CH2Cl2, then the further addition of water does not lead to a considerable shift of the potential (20 mV). For the other media, a 120-140 mV shift of the apparent potential is observed. More surprising is the relative order of the apparent redox potential in the different media (Table 4). It varies according to \( E^{0}_V \) (THF) < \( E^{0}_V \) (PC) < \( E^{0}_V \) (CH2Cl2) < \( E^{0}_V \) ([BMIm][NTf2]) < \( E^{0}_V \) ([BMIm][PF6]). As the polarity is increased (for instance THF to PC), the stabilization should be higher for the dication (CuII) than for the monocation (CuI). As a consequence, a negative shift of the redox potential is expected.30 As shown in Table 4, the \( E^{0}_V \) values vary with the media but not in the trend expected on the basis of the dielectric constants values (PC > [BMIm][NTf2] > [BMIm][PF6] > CH2Cl2 > THF). Moreover, if we compare the “dry” THF medium with the “dry” ionic liquids medium - because those media contain a similar amount of water - it can be observed that the apparent redox potential is positively shifted in the ionic liquids media (\( \Delta E^{0} = 60 \text{ mV} \)). This effect is even stronger (140 mV) when the fifth ligand bound to Cu is a chloride ion (-0.75 V vs -0.61 V). Such a situation suggests that the CuI complex is better stabilized than the CuII cation. This observation can be related to specific solvation effect in ionic liquids media.34 Ionic liquids are organised media and the CuI complex is likely to induce much less disorder than its corresponding CuII analogue, leading to an apparent stabilization of the cupric complex (entropic factor).

**Medium effects on the kinetics of the reduction process of [CuII(TMPA)(H2O)]2+.**

A further analyse of the electrochemical behaviour of the [CuII(TMPA)(H2O)]2+ complex allows to explore in more details the dynamics of the process namely regarding its kinetics. First, we will consider the apparent heterogeneous electron-transfer kinetics of the process in “dry” and “wet” media. One problem for measuring the associated rate constant with cyclic voltammetry is the large resistance resulting from low conductive media as those used in this work and which is somewhat difficult to mitigate with the positive feedback compensation feature included in most of the commercially available potentiostat. A way to overcome this difficulty is to use electrochemical impedance spectroscopy (EIS) that permits to determine the apparent heterogeneous electron-transfer kinetic rate constant

<table>
<thead>
<tr>
<th>Medium</th>
<th>Dry</th>
<th>Wet</th>
</tr>
</thead>
<tbody>
<tr>
<td>[H2O]</td>
<td>15</td>
<td>6</td>
</tr>
<tr>
<td>THF</td>
<td>2.5</td>
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</tr>
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</tbody>
</table>

Table 4: Selected electrochemical parameters for [CuII(TMPA)(H2O)]2+ in the different media. \( E^{0}_V \) vs Fe.
from a charge transfer resistance ($R_{ct}$) measurement separated from the uncompensated solution resistance ($R_{unc}$). EIS was performed in the different media for dry and wet conditions. As depicted in Fig. 6 for [BMIm][NTf$_2$], the addition of water systematics leads to a variation of the Nyquist Plots ($Z^*$ vs $Z'$ where $Z^*$ and $Z'$ correspond to the imaginary and real part of the impedance respectively). Reasonable fits of the experimental curves were obtained on the basis of a classical equivalent circuit modelling the electrochemical cell including a constant phase element (CPE) (instead of a pure double layer capacitance) and a Warburg impedance ($W$) (related to the diffusion limiting process) (Scheme 2). The Nyquist plots fitting provides a determination of $R_{ct}$ values then of the apparent electron transfer rate constant $k_{app}$ in the different media, under “dry” and “wet” conditions. Results are gathered in Table 5 for comparison purpose. For any of the media used in this work, the overall electrochemical process appears as a very slow process since the reported $k_{app}$ values are small. The addition of water results in an enhancement of the apparent electron-transfer kinetics, indicating that the overall kinetics depends on the water concentration in the medium. This suggests that the reduction process of the $[\text{Cu}^{II}(\text{TMPA})(\text{H}_2\text{O})]^+$ complex involves chemical reactions ($\text{H}_2\text{O}$ coordination/decoordination) that are faster than the electrochemical reaction as previously described by Laviron in a square scheme mechanism.

Fig. 6. Nyquist plots ($Z^*$ vs $Z'$) from EIS measurements at a glassy carbon electrode of $[\text{Cu}^{II}(\text{TMPA})(\text{H}_2\text{O})]^+$ in [BMIm][NTf$_2$] upon addition of water ($E_{acc} = 0.01 \ \text{V}, 0.1 \ \text{Hz} < \omega < 10^4 \ \text{Hz}$). A) $[\text{H}_2\text{O}] = 4 \times 10^3$ mol L$^{-1}$ (50 ppm) ($\square$); $[\text{H}_2\text{O}] = 6 \times 10^2$ mol L$^{-1}$ (700 ppm) ($\triangleright$), and $[\text{H}_2\text{O}] = 2.9 \times 10^5$ mol L$^{-1}$ (3600 ppm) ($\bullet$); B) $[\text{H}_2\text{O}] = 2.9 \times 10^3$ mol L$^{-1}$ (3600 ppm) exp. data and simulated curve; D) $[\text{H}_2\text{O}] = 4 \times 10^3$ mol L$^{-1}$ (50 ppm) exp. data and simulated curve.

Table 5 Apparent heterogeneous electron-transfer kinetic rate constant ($k_{app}$) values in different media (dry/wet) from Nyquist plots fitting.

<table>
<thead>
<tr>
<th>Medium</th>
<th>$k_{app}$ ($\text{dry}$)</th>
<th>$k_{app}$ ($\text{wet}$)</th>
</tr>
</thead>
<tbody>
<tr>
<td>$[\text{H}_2\text{O}]$ mol L$^{-1}$</td>
<td>$k_{app}$ cm s$^{-1}$</td>
<td>$k_{app}$ cm s$^{-1}$</td>
</tr>
<tr>
<td>$\text{CH}_2\text{Cl}_2$</td>
<td>$1.5 \times 10^3$</td>
<td>$1.4 \times 10^4$</td>
</tr>
<tr>
<td>$\text{THF}$</td>
<td>$2.5 \times 10^3$</td>
<td>$9.4 \times 10^4$</td>
</tr>
<tr>
<td>$\text{PC}$</td>
<td>$9.9 \times 10^3$</td>
<td>$4.5 \times 10^5$</td>
</tr>
<tr>
<td>$[\text{BMIm}][\text{NTf}_2]$</td>
<td>$4.1 \times 10^3$</td>
<td>$2.9 \times 10^5$</td>
</tr>
<tr>
<td>$[\text{BMIm}][\text{Pf}_6]$</td>
<td>$5.8 \times 10^3$</td>
<td>$2.0 \times 10^3$</td>
</tr>
</tbody>
</table>

The comparison of $k_{app}$ values between organic solvents and the ionic liquids clearly shows (for both dry and wet conditions) a slower electron transfer process in the IL media. This effect was already reported for different electroactive species. From a kinetic point of view, IL could affect the activation energy, through solvent-solute interactions, dynamic solvent effects, as well as the entropic term because of the strong structuration of the media. In the framework of the Marcus-Hush theory, outer-sphere electron transfer kinetics could be expressed under its Arrhenius form (equation (2)):

$$k_{app} = \kappa K_p \tau_L \left[\frac{-\Delta G^*}{4\pi RT}\right]^{1/2} \exp \left[\frac{-\Delta G^*}{RT}\right]$$

where $\kappa$ is the probability of electron tunnelling (1 in case of adiabaticity), $K_p$ is the equilibrium constant for a precursor model, $\tau_L$ is the longitudinal relaxation time, $\Delta G^*$ is the activation energy.

A recent theoretical work on the use of Marcus theory to redox processes in RTILs has predicted that lower rate constants obtained in ionic liquid vs usual organic solvents could be due to solvent dynamics, i.e the $\tau_L$ parameter from the pre-exponential factor in equation (2). Indeed, screening times are supposed to be higher in ionic liquids than in organic solvents. It has been suggested that this could result from slow translational motion of the ions in the ionic liquid media whereas fast longitudinal relaxation occur in organic solvents.

Voltammetric simulations

The cyclic voltammetry experiments as described in Fig. 3 are in agreement with an electron transfer associated with a slow (at the cyclic voltammetry timescale) decoordination of the $\text{H}_2\text{O}$ axial ligand when $\text{Cu}^{II}$ is reduced to $\text{Cu}^{I}$. The anodic peak located at ca. 0.4 V vs Fc is $\text{H}_2\text{O}$-related and arises upon the reduction of $\text{Cu}^{II}$ complex to $\text{Cu}^{I}$ complex. This irreversible oxidation peak could be reasonably ascribed to the cuprous complex with no water molecule bounded to $\text{Cu}$. This is supported by the fact that the oxidation potential corresponding to the formed $[\text{Cu}^{II}(\text{TMPA})]^+$ is
hardly affected by the media since a value $E_p \sim 0.4$ V vs Fc is found in all the four media. Notice that this system remains totally irreversible for all the scan rates, up to 20 V s$^{-1}$, and its oxidation occurs at a potential around 0.7 V more positive than the aqua complex. We could derive that, contrarily to [Cu(TMPA)]$^{2+}$, the binding of water to [Cu(TMPA)]$^{2+}$ is fast and totally irreversible.

![Scheme 3](image)

Scheme 3. Formal square scheme mechanism for the exchange of water associated with electron transfer for the [Cu(TMPA)(H$_2$O)]$^{2+}$ complex.

The electrochemical process can be then described as in scheme 3: [Cu(TMPA)(H$_2$O)]$^{2+}$ is first reduced to [Cu(TMPA)(H$_2$O)]$^+$ and a slow decoordination of the H$_2$O axial ligand is following to produce a cuprous complex [Cu(TMPA)]$^+$ which is oxidized on the return scan. The 4-coordinate Cu species likely displays a pseudo-tetrahedral environment in agreement with previous studies. Voltammograms of Fig. 3 and 4 were simulated (see Fig. S1) to extract the kinetic ($k_i$, $k_f$) and thermodynamic ($K = k_f/k_i$) parameters associated to the binding/unbinding at the Cu$^{2+}$ redox state according to Scheme 3. It was possible to get a good agreement between experimental and fitted parameters considering the mechanism depicted in Scheme 3. Two criteria could be considered to extract the data. The reversibility of the reduction of [Cu(TMPA)(H$_2$O)]$^{2+}$ considerably increases with the scan rates from 0.02 to 0.5 V s$^{-1}$. Similarly, the reversibility increases with an addition of water in the media. These two observations imply that the reaction of [Cu(TMPA)]$^+$ with H$_2$O is relatively slow and equilibrated.

![Table 6](image)

Table 6. Kinetics and thermodynamic parameters of the dissociation reaction of [Cu(TMPA)(H$_2$O)]$^{2+}$

<table>
<thead>
<tr>
<th>Medium</th>
<th>$k_i$ / s$^{-1}$</th>
<th>$k_f$ / L mol$^{-1}$ s$^{-1}$</th>
<th>$K$ / L mol$^{-1}$</th>
</tr>
</thead>
<tbody>
<tr>
<td>THF</td>
<td>0.15</td>
<td>4</td>
<td>26</td>
</tr>
<tr>
<td>PC</td>
<td>0.02</td>
<td>4</td>
<td>200</td>
</tr>
<tr>
<td>[BMIm][NTf]</td>
<td>0.08</td>
<td>20</td>
<td>250</td>
</tr>
</tbody>
</table>

[a] Estimations from simulations of experimental voltammograms (See Fig. S1 and Supplementary Information part).

These conclusions are illustrated by the derived rate constants of Table 6. On the whole, there is little change with the solvent. We just notice that the $H_2O$ binding is more favoured and faster in the ionic liquid than in the two other classical solvents. On one hand, the sluggishness of the decoordination process in all media may be associated to the fact that pyridyl moieties can be very labile at

40 Cu$^2+$ redox state when an exogenous ligand is bounded, as previously reported. Indeed, the pyridyl decoordination may induce a reinforcement of the metal-water bond, thus disfavouring water departure. On the other hand, the higher kinetics for the water binding reaction at Cu$^2+$ detected for the imidazolium-based ionic liquids vs THF and PC may be ascribed to the existence of specific domains in these media which significantly favour the diffusion of water molecules despite high viscosity. This behaviour is reminiscent of channel enzymes and could be exploited in catalytic reactions involving fast water coordination/decoordination.

Conclusions

The coordination behaviour of a mononuclear complex [Cu(TMPA)(H$_2$O)]$^{2+}$, simple mimic of metallo enzymes was investigated in different non-coordinating solvents, including IL, by combining electrochemistry and spectroscopic techniques. Only a few studies report on the fundamental behaviour of inorganic metal complexes in IL. The coordination geometry of the Cu$^{2+}$ complex remains trigonal bipyramidal in all the media with the water ligand being axially coordinated to Cu$^{2+}$. In particular, the anions of the ionic liquids were not found to coordinate the copper centre. The dissociation mechanism of the water axial ligand upon electron transfer in terms of kinetics and thermodynamics was studied in different media. The overall process is highly dependent on the water concentration, with a slow electron transfer reaction, slower in ionic liquids than in the conventional solvents, as already reported for different electroactive species. The decoordination of the water molecule was detected by cyclic voltammetry from the mono-electronic reduction of the Cu$^{2+}$ complex. The increase of the water content and/or decrease of the timescale of the experiment resulted in the disappearance of the oxidation peak related to the water-free Cu$^{2+}$ complex. Numerical simulations of the experimental CVs suggest that the binding/unbinding of water at the Cu(I) redox state is relatively slow and equilibrated in all media. Moreover, the binding of water at Cu$^{2+}$ is somewhat faster in the ionic liquids than in the non-coordinating conventional solvents, while the unbinding process is weakly sensitive to the nature of the solvents.

In summary, the ionic liquids exhibit promising properties in comparison to classical solvents, to investigate redox systems which are Cu-enzymes models: (i) no coordination of the anion to the metal ion at any redox state, (ii) enhanced water flow/exchange, and (iii) tunable molecular structure (shape and bulkyness). This last property can be exploited for investigating host-guest reactions with biomimetic systems incorporating cavities such as calix[n]arenes, since adequate design of the RTILs can preclude competitive solvent hosting and thus promote better access of the substrate to the cavity.

Experimental

Materials and methods

Anhydrous “extra-dry” dichloromethane (H$_2$O < 30 ppm, Acros), tetrahydrofuran (99.85%, Acros), and propylene carbonate (99.7%, Sigma-Aldrich) were used as received and kept under N$_2$ in the glovebox. CH$_2$Cl$_2$, THF and PC were dried on activated molecular sieves in the glovebox before use. All chemicals were

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of reagent grade and were used without purification. The supporting electrolyte NBu₄PF₆ was synthesized from NBu₄OH (Fluka) and HFPE (Aldrich). It was then purified, dried under vacuum for 48 hours at 100 °C, then kept under N₂ in the glovebox. EPR spectra were run at T=150 K on a Bruker Elexys spectrometer (X-band). EPR simulation was carried on the XSophe-Sophe-Xpreview software and afforded the quantitative determination of the gₓ and gᵧ parameters as well as the related hyperfine coupling constants Aₓ and Aᵧ.45 UV-Vis-NIR spectroscopy was performed with a JASCO V-670 spectrophotometer and an integration-sphere accessory for solid-state measurements. The electrochemical studies were performed in a glovebox (Jacomex) (O₂ < 1 ppm, H₂O < 1 ppm) with a home-designed 3-electrodes cell (Working electrode: glassy carbon disk, reference electrode: Pt in Fe²⁺/Fe solution, counter electrode: Pt wire). The glassy carbon electrode was carefully polished before each voltammetry experiment with a 3 µm alumina aqueous suspension and ultrasonically rinsed in water then acetone. The concentration in Cu complex was 1 0⁻⁹ mol·L⁻¹ except in the ionic liquid where the concentration was set to 3 10⁻⁵ mol·L⁻¹, in order to get a better signal-to-noise ratio. All potential values are given against the ferrocene/ferrocenium couple. Cyclic voltammetry and impedance spectroscopy measurements were performed with an Autolab electrochemical analyzer (PGSTAT 302 potentiostat/galvanostat from Eco Chemie B.V., equipped with the GEPES/FRA software). For cyclic voltammetry, scan rates were varied between 0.02 and 5 V s⁻¹. For impedance spectroscopy, a 10 mV rms sinusoidal signal was applied along with the dc potential, which was set at the E₀ value previously determined by cyclic voltammetry. The frequency range was varied from 0.1 Hz to 10 kHz. The charge transfer resistance value (Rₓ) was extracted from the experimental Nyquist plots on the basis of the equivalent circuit. The apparent rate constant k_app was evaluated from the simplification of the Butler-Volmer equation at small overpotential (equation (3)), assuming that α = 0.5 and Cₓ = Cᵧ at the equilibrium potential. [R is the perfect gas constant, T is the absolute temperature, n is the number of electrons transferred, F is the Faraday constant, A is the electrode area, Cₓ and Cᵧ are the bulk concentrations of the oxidized and reduced species, α is the symmetry barrier coefficient and was taken as 0.5].

\[
Rₓ = \frac{RT}{n²F²A k_app Cₓ e^{(1 - α)} - α F R}
\]  (3)

Synthesis of the ligand and Cu complex

The TMPA ligand (TMPA = tris(2-pyridylmethyl)amine) was prepared according to a previously published procedure.46 The synthesis of the bis-triflate Cu complex \([\text{Cu}^{II}(\text{TMPA})(\text{H₂O})](\text{CF₃SO₃})₂\) was carried out by a method previously described47 for the synthesis of Cu(II) complexes, with a small modification: the Cu(II) solution was prepared from CuCl₂·2H₂O in water with the addition of an equimolar amount of H₂SO₄. The ligand solution was added and the mixture was stirred for 48 hours. The complex was filtered over neutral alumina and washed with acetone, then evaporated under reduced pressure. The residue was dissolved in THF and the solution was added to the Cu(II) solution. The mixture was stirred for 48 hours, filtered over neutral alumina and dried under vacuum. The complex was then purified by repeated washing with H₂O, filtered over neutral alumina and silica. Prior to each experiment, vacuum pumping carefully dried RTIL overnight and then dried over molecular sieves at least for 48 h. The amount of water was measured by Karl Fischer titration (Karl Fischer 652 Metrohm).

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Notes and references

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† Electronic Supplementary Information (ESI) available: numerical simulations of cyclic voltammetry experiments and XRD data collection (CCDC 946244). See DOI: 10.1039/b000000x/.
‡ Addition of chloride ions leads to the increase of the intensity of the redox system at -0.61 V vs Fe/C₆. #The E₉ potential was systematically determined from the cyclic voltammetric curve before ac impedance measurement by taking the mid-potential value for different water concentrations.


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Insights into water binding/unbinding associated to the Cu$^{II}$/Cu$^{I}$ electron transfer at a biomimetic Cu centre

Ana Gabriela Porras Gutiérrez, Joceline Zeitouny, Antoine Gomila, Bénédicte Douziech, Nathalie Cosquer, Françoise Conan, Olivia Reinaud, Philippe Hapiot, Yves Le Mest, Corinne Lagrost, and Nicolas Le Poul

By combining electrochemistry and spectroscopic techniques the coordination chemistry of an aqua Cu complex was investigated in non-coordinating solvents and in imidazolium-based ionic liquids. The dynamics of the unbinding-binding process upon electronic transfer was particularly studied.

Keywords: copper complexes / electron transfer / non coordinating media / ionic liquids / ligand exchange