

REVIEW

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characterization and properties towards C–H
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This review summarizes the latest discoveries in the field of C–H activation by copper monooxygenases and more particularly by their bioinspired systems. This work first describes the recent background on copper-containing enzymes along with additional interpretations about the nature of the active copper–oxygen intermediates. It then focuses on relevant examples of bioinorganic synthetic copper–oxygen intermediates according to their nuclearity (mono to polynuclear). This includes a detailed description of the spectroscopic features of these adducts as well as their reactivity towards the oxidation of recalcitrant C_{sp}³–H bonds. The last part is devoted to the significant expansion of heterogeneous catalytic systems based on copper–oxygen cores (*i.e.* within zeolite frameworks).

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Organized as a French collaborative consortium (from Grenoble, Marseille and Brest), the authors of this review carry out research on the structure–function relationships of copper–oxygen species for the activation of C–H bonds in alkanes. Two French ANR grants, the COMEBAC project (ANR-13-BS07-0018) and COSACH project (ANR-22-CE07-0032), supported their research. Based on new catalysts inspired by the structure and functioning of the copper-containing monooxygenase pMMO, the COMEBAC project aimed to design and study a mixed-valent bis(μ -oxo)Cu^{II}Cu^{III} species that was proposed to be active in the oxidation of alkanes. The subsequent COSACH project continues the objective of COMEBAC but in extending its investigations towards mono-nuclear and dinuclear copper–oxygen entities. The architecture of these new catalysts is based on recent experimental and theoretical research on bioinspired models of copper-containing monooxygenases (pMMO, LPMO and PHM). COMEBAC and COSACH cover different areas of chemistry, including the synthesis/characterization of new ligands/complexes, generation and identification of transient copper–oxygen species using for example cryo-spectroelectrochemical approaches, theoretical calculations and studies of reactivity for the oxidation of substrates.





Scheme 1 Reaction type catalyzed by copper-containing mono-oxygenases (a) DβH (Dopamine β-monooxygenase); PHM (Peptidyl-glycine α-Hydroxylating Monooxygenase); pMMO (particulate Methane Monooxygenase); LPMO (Lytic Polysaccharide Monooxygenase); FGE (Formylglycine-Generating Enzyme) and (b) Ty (Tyrosinase).

1 Introduction

Copper-oxygen species (mono to multi-copper, derived from O₂ or O-atom donors represented by Cu/O) have been widely investigated since the 1980s due to their ubiquitous involvement in various biological processes such as oxidation and oxygenation reactions. Given their ability to act on various substrates as different as methane and phenols (pMMO vs. Ty, Scheme 1) copper-containing mono-oxygenases have drawn particular attention¹ for the development of synthetic catalysts



Chart 2 Representation of the polydentate ligands reported in this review.



Chart 1 Representation of the monodentate ligands reported in this review.

for efficient C–H bond activation. The approach, which consists of going back and forth between enzymes and chemical models, has allowed us to better understand the structure/reactivity of Cu/O species in enzymes as well as the O-atom transfer (OAT) mechanisms. This has been particularly well illustrated by studies of type III copper-containing proteins (hemocyanin, catechol oxidase and tyrosinase) which now constitute an important part of several bioinorganic chemistry textbooks.

Over the past decade, some aspects of the structure and reactivity of copper-oxygen models have been reported by Stack,^{2,3} Itoh,⁴ Tolman,⁵ Karlin,⁶ and Garcia-Bosch.⁷ This review covers the literature from the year 2017 to early 2024, focusing on recent advances on the structures and reactivities of the characterized copper-oxygen species possibly involved in C–H bond activation (Scheme 1a), as models for copper containing mono-oxygenases. Many articles in the literature report the oxidation of catechol to quinone or non-catalytic oxidation of phenol to catechol and subsequently to quinone by bioinspired complexes modelling tyrosinase's activity (Scheme 1b). The majority of these articles fall outside the scope of this review. Only those which aimed at highlighting the reactivity of Cu/O entities towards X–H bonds are discussed herein. Beyond the inventory of research work accomplished in the field of structure/reactivity of copper-oxygen species, our objective is to highlight the trends towards the challenging goal of efficient C–H bond activation. Mechanisms of C–H bond activation by reactive Cu/O intermediates (mono to polynuclear) are discussed with particular emphasis on the latest understandings and key catalytic systems (including heterogeneous catalysts).



Charts 1 and 2 in this review offer a comprehensive list of the main ligands discussed here, along with their respective abbreviations, towards the formation and stabilization of both monodentate and polydentate Cu/O species.

2 General context

Recent years have seen the evolution of our knowledge on some copper-containing monooxygenases. Indeed, without mentioning the particulate Methane Monooxygenase (pMMO), which has not yet revealed all its secrets, or Lytic Polysaccharide Monooxygenase (LPMO), the activity of which is still debated (monooxygenase *vs.* peroxigenase), others like Peptidylglycine α -Hydroxylating Monooxygenase (PHM) and Dopamine β -monooxygenase (D β H), could well go from uncoupled to coupled binuclear (see Section 2.1).² Moreover, the emerging advances could allow polyethylene deconstruction with hexamerins.⁸ The first section of this review is an opportunity to update new knowledge related to copper-containing monooxygenases.

Another development observed in recent years concerns the thermodynamic and kinetic evaluation of reactions linked to proton-coupled electron transfer (PCET) processes. These are important advances for a better understanding of the intrinsic properties of the catalysts. Therefore, we have devoted a second section to these thermodynamic aspects.

2.1. Copper-containing monooxygenases

Among the copper-containing monooxygenases described so far, tyrosinases (Tys) occupy an emblematic place due to the important role played by chemical models in deciphering their oxy-state structure and their reactivity. Tyrosinases do not catalyze the oxidation of phenols through the activation of a C–H bond but more precisely *via* an $S_{RN}1$ mechanism (*vide infra*). However, as the mechanism of this transformation involves Cu/O species common to the other monooxygenases, we start this section with tyrosinase. Two recent reviews from Tuczek⁹ and Mukherjee¹⁰ deal with this close link. Tys are present in prokaryotes and eukaryotes (plants, arthropods, fungi and mammals) where they are involved in tissues pigmentation, wound healing, radiation protection and primary immune response. Tys belong to the commonly called binuclear coupled enzymes due to their active site composed of two copper ions coordinated to three histidines each and spaced from each other by 3.1–4.2 Å depending on the redox and/or ligation state. The Ty-catalyzed oxygenation of phenolic compounds consists of two consecutive reactions, the *ortho*-hydroxylation of phenol to catechol (cresolase or monophenolase activity, EC 1.14.18.1) and its subsequent oxidation into *ortho*-quinone (catecholase or diphenolase activity, EC 1.10.3.1) (Fig. 1). Beyond many applications in the medical and biotechnological fields,^{11,12} recent studies on tyrosinases from Solomon's group have revisited the mechanism of the O-atom transfer that was previously considered as an aromatic electrophilic substitution (S_EAr) on a phenolate anion bound to the μ - η^2 : η^2 peroxido dicopper(II) core (Fig. 1b).^{13,14} On the basis of new kinetic studies and QM/MM calculations, the group of Solomon concluded that the most likely mechanism would be a radical-nucleophilic aromatic substitution ($S_{RN}1$) initiated by a H-bonding of the present phenol (no deprotonation) and then proton-coupled electron transfer (PCET) to the μ - η^2 : η^2 peroxido dicopper(II) core of oxy-Ty (Fig. 1c).^{15–18}

Two other copper-containing monooxygenases have recently gained renewed interest. Dopamine β -monooxygenase (D β H, EC 1.14.17.1, Fig. 2b) and Peptidylglycine α -Hydroxylating Monooxygenase (PHM, EC 1.14.17.3, Fig. 2a) were until now considered as uncoupled binuclear copper-containing monooxygenases with two copper ions coordinated one to three histidines (Cu_H) and the second to two histidines and one methionine (Cu_M). Cu_H and Cu_M , which are spaced from each other by approximately 11 Å, were proposed to possess distinct functions: Cu_M would be involved in the hydroxylation process through a copper-superoxido species while Cu_H would act as an electronic relay (Fig. 2d).¹⁹ In this representation, the copper-superoxido species would abstract an H-atom (Hydrogen Atom Abstraction, HAA) from the substrate which would be followed by a rebound event (Scheme 2a). This proposal, which was based on initial X-ray data (Fig. 2c and d), spectroscopic and kinetic studies on the reduced and oxy-PHM, was recently revisited and could become obsolete according to the recent X-ray 3D structure of the human D β H (Fig. 2e and f).²⁰

Indeed, the D β H X-ray structure revealed that the enzyme can adopt two different conformations: (i) an open



Fig. 1 Type III copper-containing proteins, (a) reactions catalyzed by Ty and oxy-Ty 3D-structure of the active site (PDB ID: 1wx4); (b) mechanism involving an electrophilic aromatic substitution (S_EAr) and (c) mechanism involving a radical-nucleophilic aromatic substitution ($S_{RN}1$).





Fig. 2 Uncoupled binuclear copper-containing monooxygenases, PHM and DβH. (a) and (b) reactions catalyzed by PHM and DβH, respectively; (c) X-ray structure of PHM from *Rattus norvegicus* showing the two domains (Cu_M domain in green and Cu_H domain in magenta) connected by a loop (in blue) in which each domain binds a single copper atom spaced from each other by approximately 11 Å and (d) structure of the Cu_M-superoxido intermediate in PHM (PDB ID 1opm); (e) and (f) X-ray structure of the human DβM (PDB ID 4zel) showing the open (e) and the closed (f) forms. Only the Cu_M domain of the closed form contains a copper atom, the other copper ions (red sphere) are manually placed.

conformation with two distant copper ions, resembling the previously reported PHM structure (Fig. 2e) and (ii) a closed conformation, in which the two copper sites are only 4 to 5 Å apart, in what might be a coupled binuclear copper site (Fig. 2f).



Scheme 2 Recently revisited mechanism for PHM and DβH showing the mechanisms involving an HAA by (a) Cu_M-superoxido intermediate in an open dicopper active site and (b) μ-oxyl dicopper intermediate in closed dicopper active site. Abbreviation: AscOH for ascorbic acid.

Could PHM and DβH leave the uncoupled binuclear family for the coupled binuclear ones? This question is relevant since an alternative mechanism corroborated by QM/MM calculations,²¹ involving a dinuclear copper intermediate formed *via* an open to closed conformational transition managed to better account for the experimental spectroscopic data collected over decades on PHM and the DβH (Scheme 2b).^{22–25} On one hand, it is interesting to note that this proposal was already tentatively suggested in 2002, though without sufficient support from the scientific community at the time.²⁶ On the other hand, if confirmed, this new finding illustrates once more the important role of chemical models in these studies. This takes us back few decades to Réglier's proposals when: (i) the first DβH functional model involving the occurrence of a μ-η²:η² peroxido dicopper(II) species was reported^{27,28} and (ii) based on the substrate binding ligand approach^{29,30} the occurrence of a dicopper mixed-valent species such as (μ-oxido)Cu^{II}Cu^{III} ↔ (μ-oxyl)Cu^{II}Cu^{II} was proposed to explain the O-atom transfer to the ligand.³¹

The membrane-bound Methane Monooxygenase (pMMO, EC 1.14.18.3) continues to be in the headlines of discussions about the nature, localization and nuclearity of the active site, as well as on the mechanism. Methane monooxygenases (MMOs) are



crucial enzymes in bacterial methane metabolism since they catalyze the conversion of methane to methanol ($\text{CH}_4 + \text{O}_2 + 2\text{e}^- + 2\text{H}^+ \rightarrow \text{CH}_3\text{OH} + \text{H}_2\text{O}$). They represent a potential route towards the use and mitigation of methane emissions. MMOs exist in two forms, a cytoplasmic soluble form (sMMO) and a membrane-bound enzyme form (pMMO). These MMOs are completely unrelated, with different architectures, metal cofactors (iron for sMMO and copper for pMMO), and mechanisms.³² While the mechanism of sMMO is well described and agreed upon, that is not the case for pMMO. To date, crystal structures of pMMOs from five species have been determined.^{33,34} These structures show that pMMOs are heterotrimers in which each monomer is composed of three subunits, PmoA (α), PmoB (β), and PmoC (γ). The heterotrimer ($\alpha\beta\gamma$)₃ adopts a cylindrical shape composed of twelve transmembrane helices almost entirely embedded in the lipid bilayer (Fig. 3a).

Despite these structures, the identity of the copper sites remains unclear. According to several spectroscopic and biochemical analyses there would be 12–14 copper ions found within each functional pMMO. Only two sites (Cu_B and Cu_C)

have been clearly identified by X-ray crystallography in all pMMO sources (Fig. 3). The nuclearity of the Cu_B site was the subject of a long debate. In the original crystal structure of the pMMO of *M. capsulatus* (Bath), this site could be modeled with one or two copper ions. However, thanks to spectroscopic^{35–37} and computational³⁸ studies, the debate has evolved in favor of a mononuclear site. In contrast, the Cu_C site located within the transmembrane domain of the PmoC subunit has been identified as mononuclear featuring two histidines and one aspartate as ligands. Biochemical studies of pMMO activity suggest that the Cu_C site would likely be the catalytic site for O_2 activation and CH_4 oxidation.³⁴ A recent publication by Shaik attempts to address this question through a QM/MM study of dioxygen activation at the Cu_C center in the presence of the physiological reductant duroquinol (DQH2).³⁹ This simulation has converged towards the occurrence of the $\text{Cu}_\text{C}^{\text{II}}$ -oxyl species which would be responsible for the C–H activation. These results reveal the role that phenols could play in the activation of O_2 at the monocopper center.

However, two recent cryo-electron microscopy (cryo-EM) structures of the pMMO have re-opened the debate (Fig. 3). The first structure published by Chan regards pMMO from *M. capsulatus* (Bath) in the copper(I) oxidation state. This structure reveals a dinuclear copper B site, and more surprising unveils additional copper clusters at the PmoA/PmoC interface within the membrane (D site) and in the water-exposed C-terminal subdomain of the PmoB (E site).⁴⁰ The other structures have been published by Rosenzweig of the pMMOs *M. capsulatus* (Bath), *M. alcaliphilum* and *Methylocystis* sp. strain Rockwell in the copper(II) oxidation state and embedded in the lipid bilayer closer to the *in vivo* environment.⁴¹ These new structures reveal a mononuclear copper A site (only for *M. capsulatus* and *M. alcaliphilum*), a mononuclear copper B site and the presence of copper in C or D sites but never together in both sites. The different nuclearities observed for the copper B site (mono vs. binuclear) could be linked to the oxidation state of copper ions, mono for copper(II) vs. dinuclear for copper(I). This observation is perhaps not surprising, since copper tridentate complexes bearing the same ligand were shown to crystallize in dimeric and monomeric forms in the copper(I) and copper(II) oxidation states, respectively.⁴² More interesting is the proximity of the copper C and D sites which are only approximately 4 Å apart from each other. Recent QM/MM calculations⁴³ suggest that the $\text{Cu}_\text{D}^{\text{I}}$ site would be more reactive than the $\text{Cu}_\text{C}^{\text{I}}$ site in dioxygen activation for the generation of a Cu^{II} -oxyl species. Furthermore, these simulations which tend to demonstrate that the natural reductant ubiquinol (CoQH_2) prefers to bind to the $\text{Cu}_\text{D}^{\text{I}}$ site rather than to the $\text{Cu}_\text{C}^{\text{I}}$ site could support the fact that the active site of pMMO is Cu_D rather than Cu_C .⁴³ Even if the structures did not reveal the presence of copper ions in the two sites, given the short distance between them, one may raise the question of the physiological relevance of a possible binuclear Cu_C – Cu_D site for methane oxidation (Fig. 3c).

Lytic polysaccharide monooxygenases (LPMOs, EC 1.14.99.54) are copper-containing enzymes that comprise a large enzyme super-family. LPMOs hydroxylate polysaccharide substrates at either the C1 or the C4 position followed by the



Fig. 3 X-ray structure of *M. capsulatus* (Bath) pMMO (PDB ID 3rgb). (a) Heterotrimer ($\alpha\beta\gamma$)₃ adopting a cylindrical shape composed of twelve transmembrane helices; (b) 3D-view of one subunit $\alpha\beta\gamma$ showing the different location of copper ions, PmoA (α) in green, PmoB (β) in blue and PmoC (γ) in orange; (c) 3D-views of the identified copper center by cryo-EM. The Cu_B center displays a histidine brace similar to the one reported on LPMO (see Fig. 4).





Fig. 4 Lytic polysaccharide monooxygenase (LPMO). (a) Reactions catalyzed by LPMO; (b) LPMO X-ray structure from *Thermoascus aurantiacus* (PDB ID 2yet) showing the copper active site at the binding surface of the LPMO.

subsequent formation of the aldonic acids (C1 hydroxylation) or 4-keto sugars (C4 hydroxylation) at oxidized chain ends (Fig. 4a).⁴⁴ All LPMOs display a mononuclear copper active site that is located on the binding surface of the enzyme (Fig. 4b). Coordinated with two histidine residues, the copper ion exhibits a distinctive feature: it is linked to the *N*-amino-terminal histidine through its NH_2 -terminal end and with the imidazole side chain, a coordination motif that has been named “histidine brace”.⁴⁵ Since the discovery in the early 2010s of the first LPMO, these enzymes have continuously attracted interest and stimulated the curiosity of the scientific community.^{44–47} One major reason for interest in LPMOs is their key role in the deconstruction of recalcitrant polysaccharides (cellulose, chitin, *etc.*), a role of paramount commercial importance to produce advanced biofuels or bio-based chemicals.⁴⁸ Another matter of interest is that LPMOs were the first “real” mononuclear copper-containing monooxygenases easily accessible by genetic expression.

LPMOs are active in complex media, with heterogeneous substrates (cellulose, chitin, *etc.*) in the presence of numerous reducing polyphenolic compounds (degradation of lignin), and



Scheme 3 Putative mechanisms to explain LPMO catalyzed hydroxylation: (a) the monooxygenase pathway and (b) peroxxygenase pathway.

in the presence of both dioxygen and hydrogen peroxide. Therefore, the mechanism associated with their activity is difficult to study. Although LPMOs were considered at first as monooxygenases (with dioxygen as the co-substrate), recent investigations have suggested that they could instead be considered as peroxxygenase (with hydrogen peroxide as the co-substrate).⁴⁹ The exact nature of this oxygenated co-substrate (dioxygen or hydrogen peroxide) is still debated^{50–53} as well as the nature of the copper-oxygen species involved in polysaccharide hydroxylation.^{54,55} Recently, Schröder *et al.* resolved a new cryo-trapped structure with an activated dioxygen intermediate in a mixture of superoxido and hydroperoxido states.⁵⁶ Several alternative active species have been proposed to be formed within LPMOs' catalytic mechanism, including Cu^{II} -oxyl, or even the hydroxyl radical^{57,58} without having been spectroscopically characterized or observed. Scheme 3 summarizes the possible intermediates and mechanisms for LPMO catalyzed hydroxylation.

The Formylglycine-Generating Enzyme (FGE, EC 1.8.3.7), which catalyzes the post-translational activation of type I sulfatases (cysteine residues in formylglycine, Fig. 5a) in eukaryotes and aerobic microbes, was discovered at the end of the 1990s.⁵⁹ Recently, it drew renewed interest not only because of its potential in protein engineering,⁶⁰ but also because of its unique active site which differentiates it from other copper-containing monooxygenases (Fig. 5).⁶¹ Indeed, while copper ions bind mainly to histidines with tetrahedral or square-planar coordination spheres in the classical copper-containing monooxygenases (Ty, D β H, PHM, pMMO and LPMO), a single copper(I) binds two cysteines in a linear coordination configuration in FGE. Thanks to several techniques (stopped-flow absorbance spectroscopy,⁶² X-ray,⁶³ QM/MM⁶⁴), a mechanism has been proposed in which a non-coordinated dioxygen complex was





Fig. 5 Formylglycine-Generating Enzyme (FGE). (a) Reaction catalyzed by FGE; (b) the proposed mechanism and within the center the 3D view of the FGE active site from *Thermomonospora curvata* (PDB ID 6xtr) showing the copper center linked to two cysteines and the peptide substrate as well as the hydrophobic pocket consisting of a tryptophan and a serine where the dioxygen binds before to be reduced to superoxide.

suggested as a precatalytic complex. The dioxygen would not be coordinated to the copper(I) but rather bound in a hydrophobic pocket juxtaposed to the metal. Thus, the binding and activation of dioxygen by FGE would be a stepwise process conversely to many other metal-containing oxygenases. The mechanism would then involve HAA by a copper superoxido intermediate followed by the formation of an alkyl-peroxido intermediate precursor of the aldehyde. However, the mechanism is far from being fully elucidated and investigation using small copper model complexes could be highly beneficial.

In 2023, it was reported that the buccal secretion of the lepidopteran *Galleria mellonella* larvae was capable of oxidizing and depolymerizing polyethylene at room temperature in the presence of a copper salt. The cryo-EM analysis and 3D-reconstructions revealed that the lepidopteran buccal secretion is mainly composed of four hexamerins (Demetra, Cibeles, Ceres and Cora) belonging to the type III copper-containing protein family.⁸ However, despite close homologies with Hc and Ty, the type III canonical active site present in such lepidopteran hexamerins does not have the capacity to bind copper ions. Surprisingly, it consists of a combination of aromatic/hydrophobic and charged residues instead of histidine residues present in Hc/Ty. However, X-ray fluorescence

spectroscopy at the Cu-absorption edge indicated the presence of 6 copper ions per subunit in lepidopteran hexamerins and the 3D reconstructions revealed the presence of several different metal ion binding sites. The diverse location of metal sites, sometimes present at subunit interfaces, could have a structural function, but it cannot be excluded that the PE-modifying activity of these proteins is assisted by any of these metals. The results are still quite vague, and the occurrence of copper ions needs to be confirmed. However, this work must receive our full attention considering that plastic waste management is a pressing ecological, societal, and economical challenge.

In this constantly evolving context where knowledge of biological systems is refined over the years, the design of model complexes of the different active sites is necessary to contribute to this progress. From these studies, a clear emphasis on the chemistry of dioxygen and/or hydrogen peroxide at mono, bi or trinuclear copper centers en route to produce Cu^{II} -oxyl or $(\mu\text{-oxido})\text{Cu}^{\text{II}}\text{Cu}^{\text{III}} \leftrightarrow (\mu\text{-oxyl})\text{Cu}^{\text{II}}\text{Cu}^{\text{II}}$ active species is more relevant than ever. This review focuses on this subject.

2.2. Thermodynamic and kinetic aspects of C–H bond abstraction

Evaluation of the intrinsic mechanisms occurring in HAA with the afore-mentioned enzymes is obviously extremely difficult due to the transient character of the copper–oxygen adducts generated along the catalytic cycle. Nevertheless, synthetic models of copper enzymes, although generally less effective than their natural counterparts, have been extensively studied since they afford better understanding of oxygen activation through a fine characterization of the intermediates, mostly by using low-temperature conditions. In recent years, a common strategy has been targeted to evaluate reaction thermodynamics and kinetics related to Proton-Coupled Electron Transfer (PCET) processes to obtain better insights into the intrinsic properties of the catalysts. In particular, the bond dissociation free energy (BDFE) of the O–H bond of hydrogenated copper–oxygen adducts is considered as a strong indicator of the oxidizing HAA power of the catalyst, although it provides no direct information on the stepwise/concerted pathways for H-atom transfer and the related kinetics.⁶⁵

BDFEs for solution-phase PCET processes are usually determined by using a square scheme involving two parallel ET and



Scheme 4 Square-scheme mechanism for the HAA reaction: concerted (HAT/CPET) vs. stepwise (ET/PT) pathways. X is the reactive copper–oxygen species and XH stands for the hydrogenated copper–oxygen species.



PT processes, where XH denotes the hydrogenated copper-oxygen catalyst (Scheme 4).^{66,67}

According to this scheme, the BDFE of the X–H bond can be obtained by considering stepwise electron transfer (ET) and proton transfer (PT) processes. Indeed, independent determination of pK_a and one-electron reduction standard potential (E^0 in V) values related to ET–PT or PT–ET pathways yields BDFE according to eqn (1):^{66,67}

$$\text{BDFE} = 23.06 E^0 + 1.37 pK_a + C_G \quad (1)$$

where C_G denotes the free-energy for the H^+/H^\bullet reaction ($C_G = -FE^0(H^+/H^\bullet)$), which is solvent-dependent.

Given the high uncertainty on C_G , Mayer and co-workers have recently proposed another general expression for BDFE (see eqn (2)) that includes the potential of hydrogenation E^0 (V vs. H_2) formally related to the PCET conversion of X into XH involving H_2 , as well as the free enthalpy of formation of H^\bullet :^{66,68}

$$\text{BDFE} = 23.06 E^0 (\text{V vs. } H_2) + \Delta G_f^0(H^\bullet) \quad (2)$$

According to Mayer, this alternative approach is of interest since (i) E^0 (V vs. H_2) is easily accessible from open-circuit potential (OCP) measurements, and (ii) $\Delta G_f^0(H^\bullet)$ is almost constant ($52.2 \pm 0.6 \text{ kcal mol}^{-1}$) for a wide range of commonly used solvents, in contrast to C_G . As a consequence, the solution-phase measured values of E^0 (V vs. H_2) are considered as equivalent to BDFE ones. In practice, OCP measurements require the presence in solution of known concentrations of X, XH, acid/base buffer and electrolyte species.⁶⁶ Although it was shown to be fully reliable for stable X/XH systems, it is more difficult to use with highly unstable X/XH species such as those encountered with copper-oxygen adducts.

Beyond thermochemical considerations for PCET processes, supplementary information can be obtained by different kinetic analyses and/or computational studies. Various experimental approaches are possible to decipher the concerted or stepwise character of the studied reaction. According to the general definition adopted in recent years by chemists working in the field of PCET reactions, two classes of concerted HAA processes are usually considered, namely CPET (also coined as cPCET) and HAT. The difference between concerted proton–electron transfer (CPET) and H-atom transfer (HAT) relies on the locus of the electron and proton transfer, *i.e.* on the same orbital (HAT) or on different orbitals (CPET).⁶⁷ For these two mechanisms, high-energy intermediates are precluded in strong contrast to elementary ET and PT stepwise pathways. Interestingly, in-between situations which were referred to as concerted asynchronous HAA reactions have been recently proposed on the basis of computational studies.^{69–72} Hence, a basic asynchronous mechanism may develop a PT-like character in the transition-state whereas an oxidatively asynchronous process would feature an ET-like character.

The rate constant k of an HAA reaction is classically determined experimentally from the UV-Vis monitoring of either the catalyst or substrate spectroscopic signal throughout the reaction.^{65,73,74} For instance, the determination of k for various

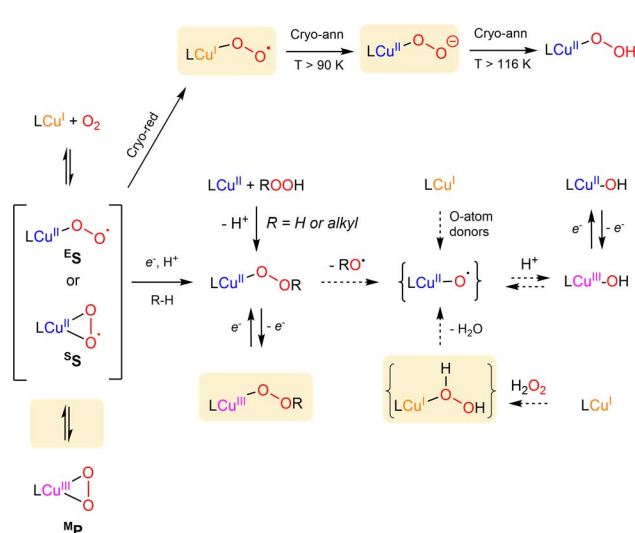
concentrations of hydrogenated and deuterated substrates allows the determination of a Kinetic Isotope Effect (KIE) value, suggesting either a stepwise (ET–PT/PT–ET) process if the KIE is close to 1, or a concerted (HAT/CPET) reaction if $KIE > 1$. Other approaches like Hammett plots ($\ln k$ vs. Hammett coefficients of substrates), Marcus plots ($\ln k$ vs. $E^0(\text{substrate})$) and Bell–Evans–Polanyi plots (Gibbs free energy of activation ΔG^\ddagger vs. BDFE(substrate)) have been also used to obtain mechanistic information.⁷³ Hence, the rate constant for a concerted PCET process is expected to increase as the BDFE of the substrate decreases, as classically found for KIE (H/D) analysis. In contrast, a stepwise PCET featuring a PT rate-determining step shows an increase of k with the BDFE of the substrate. In addition to these experimental approaches, recent computational studies have proposed to differentiate between HAT and CPET processes on the basis of the variation of the dipolar moment of the catalyst–substrate adduct bond throughout the HAA reaction.⁷²

Overall, quantification of thermodynamics and kinetics of HAA reactions involving copper-oxygen species has been extensively standardized over the last few years, allowing direct comparison between the different families of copper catalysts. With the support of computational studies, such approach gives more insights into the mechanisms and helps in the rationalization of the oxidative properties in relation to the structure of the active species.

3 Mononuclear copper–oxygen species

3.1. Cu/O₂ entities: copper-superoxido or copper-peroxido adducts

3.1.1. Structural and/or electronic data. Cu/O₂ entities are generally the first intermediates formed upon reaction of L–Cu(I) complexes with dioxygen. In most cases, they are obtained upon



Scheme 5 Summary of known formation and interconversion pathways of mononuclear copper-oxygen entities. Recently isolated species or pathways are highlighted in orange. Dotted arrows correspond to postulated conversions pathways. The nomenclature used is derived from Stack *et al.*²



reaction of Cu(I) precursors with dioxygen at low temperature.^{5,19,75–77} This procedure has led to the isolation and characterization of several mononuclear entities with different structural and electronic properties. These species can be formulated as either Cu(II)-superoxo (with the superoxido ligand bound either in the η^1 or η^2 binding mode (Scheme 5, ^ES and ^SS respectively)) or Cu(III)-peroxido entities (with the peroxido bound in the η^2 binding mode, ^MP) depending on the degree of electron transfer achieved between dioxygen and the initial Cu(I) precursor. In general, η^1 -superoxo entities display an $S = 1$ ground state arising from ferromagnetic coupling between the Cu(II) and the superoxide radical. They are characterized by intense UV-Vis transitions at 380–420 nm ($\epsilon \approx 1000\text{--}5000\text{ M}^{-1}\text{ cm}^{-1}$) and $\nu(\text{O--O}) \sim 1100\text{ cm}^{-1}$ in their resonance Raman (rR) spectra, values typical for superoxido adducts. Intermediates displaying η^2 -coordination modes generally have singlet ground states and tuning between superoxido and peroxido forms has been found to be dependent on the electron-donating power and denticity of the supporting ligands. O–O vibrations are found ranging from 950 cm^{-1} to 1100 cm^{-1} consistent with the existence of a continuum between the two above-mentioned resonant structures.

In 2017, only four Cu/O₂ entities had been structurally characterized.^{78–81} Two additional structures were since reported by the group of Betley^{82,83} using sterically encumbered dipyrin ligands ($^{\text{Ar}}\text{L}$ or $^{\text{EMind}}\text{L}$, with Ar = 5-mesityl-1,9-(2,4,6-Ph₃-C₆H₂) and EMind = 1,1,7,7-tetraethyl-1,2,3,5,6,7-hexahydro-3,3,5,5-tetramethyl-s-indacene, Chart 1). In both cases, the dioxygen derived ligand is bound in the η^2 mode. In particular, high-quality single crystals were obtained by exposing single

crystals of the ($^{\text{EMind}}\text{L}$)Cu(N₂) precursor to air. The reaction proceeded in the solid state, reflecting that replacement of N₂ by O₂ induces minimal changes in lattice parameters (Fig. 6a).⁸³ The molecular structure of ($^{\text{EMind}}\text{L}$)Cu(O₂) revealed a square-planar geometry around the copper center with relatively short Cu–O bonds of 1.824 and 1.834 Å, together with an O–O distance of 1.379 Å. The resonance Raman spectrum of the intermediate showed the presence of a $\nu(\text{O--O})$ vibration at 1003 cm^{-1} . These structural and spectroscopic data, combined with DFT calculations, led to the assignment to a predominant ^SS description of the two intermediates.

Although no crystallographic structures could be obtained, the groups of Karlin and Hoffman have recently reported on new types of copper-oxygen intermediates using tmpa and tren-derived ligands ($\text{L} = ^{\text{PV}}\text{tmpa}$, $^{\text{DMT}}\text{tmpa}$, $^{\text{TMG}^3}\text{tren}$, Chart 1).⁸⁴ The well-known ^ES [(L)Cu^{II}(O₂)]⁺ species were generated and cryo-reduced by γ -irradiation at 77 K using a ^{60}Co source. This allows the generation of energetic mobile electrons through solvent ionization processes. All primary cryo-reduced species display similar axial Q-band and X-band EPR spectra with $g_{\parallel} = 2.52$ and $g_{\perp} \approx 1.95$. Detailed spectroscopic analysis led to the proposal that [(L)Cu^I(η^1 -O₂)] species are formed upon reduction of the copper ions, which are trapped in the geometry of the parent complexes (Scheme 5).

To the best of our knowledge, this corresponds to the first isolation of such species in the literature. Warming of the frozen solution of [($^{\text{DMT}}\text{tmpa}$)Cu^I(O₂)] at 90 K for 5 minutes (a temperature below the MeTHF solvent melting point) led to the change of the EPR signature that was assigned to the formation of the [($^{\text{DMT}}\text{tmpa}$)Cu^{II}(η^1 -O₂[−])] intermediate resulting from internal electron transfer between the copper center and the superoxido ligand (Scheme 5). Although already crystallographically detected in metalloenzymes (e.g. LPMO⁸⁸), such η^1 -Cu(II)-peroxido species was never evidenced in model complexes. Finally, warming at a temperature above the solvent melting point allowed protonation of the terminal peroxido ligand to provide the η^1 -hydroperoxido [($^{\text{DMT}}\text{tmpa}$)Cu^{II}(η^1 -OOH)]⁺ intermediate.

As previously mentioned, the dioxygen-derived ligand of the primary Cu/O₂ adducts is found either in η^1 or η^2 coordination modes. Yet, using the bulky $^{\text{TIP}^3}\text{tren}$ ligand (Chart 1), the group of Itoh observed the coexistence of the two binding modes with one single ligand-system.⁸⁹ Indeed, oxygenation of the Cu(I) precursor at low temperature led to the detection of an ^ES species together with another intermediate, assigned to the ^MP adduct based on UV-Vis, Raman, NMR and DFT calculations.⁸⁹

Finally, using a redox active ligand (DHP = (2,5-bis((2-*t*-butylhydrazono)(*p*-tolyl)methyl)-pyrrole, Chart 1), the group of Anderson prepared a diamagnetic Cu^{II}-L[•] complex.⁹⁰ Upon oxygenation at low temperature, an ^ES intermediate was characterized arising from electron transfer between the ligand and dioxygen.

3.1.2. Reactivity. The reactivity of Cu/O₂ adducts has been well documented.^{5,19,75–77} Variation of the first coordination sphere, as well as ligand denticity, geometry and bulkiness has been conducted over the past 40 years to provide substantial data on the stability and reactivity of Cu/O₂ species. In the



Fig. 6 Selected crystallographic structures of recently described mononuclear copper-oxygen adducts. a) η^2 -superoxo based on the $^{\text{EMind}}\text{L}$ ligand.⁸³ Hydrogen atoms were omitted for clarity; b) The anion [(L)Cu(OOCm)][−] based on the bis(arylcarboxamido)-pyridine ligand.⁸⁵ Hydrogen atoms and the Et₄N⁺ counter ion were omitted for clarity; c) η^1 -Hydroperoxido complex in an artificial enzyme based on a streptavidin scaffold: [(biot-et-dpea)Cu^{II}(OOH)(OH₂)]⁺ c Sav;⁸⁶ d) High-valent [(L)CuOH] species based on the bis(arylcarboxamido)-pyridine ligand (W as deprotonated water ligand).⁸⁷



absence of a sterically encumbered ligand, these adducts commonly react with excess Cu(I) precursors to generate dinuclear peroxido entities (Scheme 10). Direct conversion to the dinuclear bis- μ -oxido species (**O**) has also been recently reported.⁹¹

^ES species have been shown to display promising HAA reactivity towards O–H and C–H bonds, leading to the corresponding Cu–OOH species and radical products.^{5,19,75–77} The group of England prepared bulky tmpa derived ligands (bearing aryl substituents) and successfully stabilized $[(\text{Ar}^3\text{tmpa})\text{Cu}^{\text{II}}(\eta^1\text{-O}_2)]^+$ species that are stable up to -20°C .⁹² This allowed extensive reactivity studies using a broad range of substrates bearing O–H, N–H, or C–H bonds. As already observed for other complexes, HAA from C–H bonds proved to be more challenging than O–H and N–H bonds and the intermediates were found to react only with weak C–H bonds (*ca.* 73 kcal mol^{−1}).

It has been reported that introduction of thioether ligation to the copper, modelling the coordination sphere found in PHM and D β H enzymes, enhances the oxidizing power of $\eta^1\text{-Cu}^{\text{II}}$ -superoxido adducts.⁹³ Although several complexes were prepared with thioether-containing ligands, it has to be noted that the occurrence of thioether ligation to copper was only recently experimentally attested to using EXAFS on the previously reported $[(^{\text{TMG}}\text{N}_3\text{S})\text{Cu}^{\text{II}}(\text{O}_2)]^+$ species.^{94,95}

Second coordination sphere interactions are also known to modulate the reactivity of metal–oxygen species.⁹⁶ In order to probe the effect of second coordination sphere H-bonding effects on the reactivity of η^1 cupric-superoxido species, the group of Karlin has prepared a series of tmpa-based ligands with varying H-bonding ability.^{97,98} Increasing H-bonding to the superoxido moiety by changing the nature and the number of H-bonding substituents on the tmpa ligand was shown to both decrease the amount of binuclear entities formed to the profit of ^ES species and increase the electrophilic reactivity of the intermediate towards HAA (Fig. 7). Accordingly, the bis-pivalamido substituted ligand provided the most reactive superoxido intermediate that was able to react with O–H BDEs up to 91.5 kcal mol^{−1} and C–H BDEs up to 79.7 kcal mol^{−1}, emphasizing

the importance of second-coordination sphere elements to modulate the reactivity of copper–oxygen species, as already reported for other reactive intermediates. Very recently, Borovik *et al.* demonstrated that intramolecular H-bonds, introduced by external ligands, could influence the structural and magnetic properties of the mononuclear complex.⁹⁹

The nucleophilic reactivity of the ^ES species based on the bis(arylcarboxamido)-pyridine anionic ligand $[N,N'\text{-bis}(2,6\text{-diisopropylphenyl})\text{-}2,6\text{-pyridinedicarboxamido}]^{100}$ towards the aldehyde deformylation reaction has been reported in the literature.¹⁰¹ The group of Tolman has recently re-investigated the mechanism of this reaction.¹⁰² The preparation of the intermediate was improved by the use of a cryptand (Kryptofix® 222) when reacting the Cu(II) precursor with KO₂. This allowed additional mechanistic investigations suggesting HAA rather than nucleophilic attack processes.

Following this first study, the reaction of this anionic $[(\text{L})\text{Cu}^{\text{II}}(\eta^1\text{-O}_2)]^-$ with *para*-substituted phenols ^XArOH was investigated at low temperature.¹⁰³ Interestingly, kinetic data suggested a change in mechanism across the series depending on the nature of the substituents of the phenols. In the case of phenols with low acidities (*X* = OMe, NMe₂) a CPET/HAT mechanism was proposed. The reaction rates were significantly lower than that obtained with the high-valent $[(\text{L})\text{Cu}^{\text{III}}(\text{OH})]$ core containing the same ligand.⁶⁵ The O–H BDE value for the $[(\text{L})\text{Cu}^{\text{II}}\text{OOH}]^-$ species was estimated to be ~ 3 kcal mol^{−1} lower than that for $[(\text{L})\text{Cu}^{\text{II}}(\text{OH}_2)]^{2-}$, consistent with the higher PCET/HAT rates of $[(\text{L})\text{Cu}^{\text{III}}(\text{OH})]$ species as compared to $[(\text{L})\text{Cu}^{\text{II}}(\eta^1\text{-O}_2)]^-$ (Scheme 6). In the case of electron-deficient and acidic phenols (*X* = NO₂, CF₃, Cl), the results were consistent with a rate-determining proton transfer (PT) from the phenol to the $[(\text{L})\text{Cu}^{\text{II}}(\text{O}_2)]^-$ species to yield the formally Cu^{III}–OOH adduct. The enhanced basicity of $[(\text{L})\text{Cu}^{\text{II}}(\text{O}_2)]^-$ as compared to $[(\text{L})\text{Cu}^{\text{III}}(\text{OH})]^+$ (*pK_a* ~ 19 vs. 12 respectively) likely favors PT from acidic phenols while for the least acidic ones a CPET/HAT mechanism is followed. Finally, using the ligand L^{pyim} (Chart 1), the group of Itoh has observed an unprecedented reactivity of the corresponding ^ES intermediate towards aldehydes and ketones, leading to C–C bond formation with acetone (the reaction solvent), and to the corresponding β -hydroxyketones (aldols).¹⁰⁴ Nucleophilic reactivity was supported by

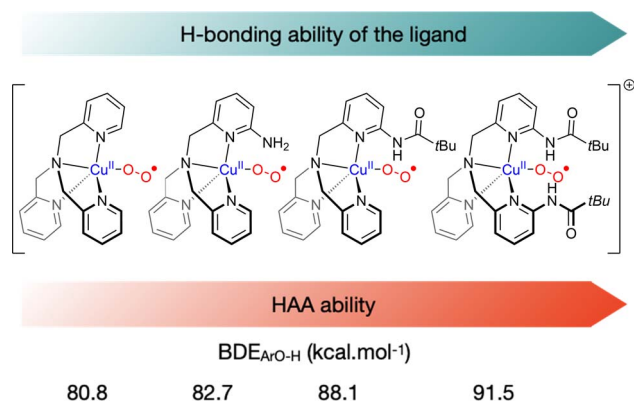


Fig. 7 Representation of selected tmpa-based ligands prepared with different H-bonding substituents to modulate their HAA reactivity.^{97,98} Highest BDE_{OH} from phenolic substrates that react with the superoxido intermediate.



Scheme 6 Comparison of the thermodynamic properties of $[(\text{L})\text{Cu}^{\text{II}}(\eta^1\text{-O}_2)]^-$ and $[(\text{L})\text{Cu}^{\text{III}}(\text{OH})]$ using the same bis(arylcarboxamido)-pyridine ligand. Adapted from ref. 103.



Scheme 7 Reaction of $[(EInd)L]Cu^{II}(O_2)$ with 1,2-diphenylhydrazine to produce azoarene and $[(EInd)L]Cu^{II}(H_2O_2)$. Adapted from ref. 83.

detailed kinetic studies and DFT calculations. A Cu-alkylperoxido species was transiently detected and characterized by rR spectroscopy, which is in agreement with nucleophilic addition of the Cu(II)-superoxido species on the carbonyl carbon of the substrate.

The reactivity of η^2 -Cu/O₂ species for HAA is less documented. The group of Betley has studied the reactivity of two η^2 -Cu/O₂ based on dipyrin ligands.^{82,83} In the case of the ^{Ar}L ligand, although thermally stable at room temperature, the intermediate reacted with 1,4-cyclohexadiene and was able to perform HAA and acid/base chemistry towards a range of phenolic substrates as well as O-atom transfer to triphenylphosphine. Although the mechanistic features remain unclear, it was proposed that an end-on ^{FS} could be formed prior to reaction with substrates. The intermediate $[(EInd)L]Cu^{II}(\eta^2-O_2)$ reacted with a stoichiometric amount of 1,2-diphenylhydrazine to provide azoarene and H₂O₂ as well as the initial $[(EInd)L]Cu^{II}(\eta^2-O_2)$ species. When the experiments were conducted under an inert atmosphere, a new diamagnetic species was detected and assigned to $[(EInd)L]Cu^I(H_2O_2)$ (Scheme 7). The same species could also be produced upon reaction of the initial $[(EInd)L]Cu^I(N_2)$ species with dry H₂O₂ ((Ph₃PO)₂(H₂O₂))¹⁰⁵ supporting the assignment as a Cu(I)-(H₂O₂) species. Such intermediate has been proposed to be pivotal in the LPMO catalytic cycle both in monooxygenase and peroxygenase pathways.⁵⁵ Further understanding of its structural and reactivity properties would therefore be of high interest.

3.2. CuOOR entities: copper-hydroperoxido or alkylperoxido adducts

3.2.1. Structural and/or electronic data. Cu^{II}-hydroperoxido intermediates are probably pivotal intermediates in enzymatic or biomimetic catalysis.^{5,19,106} In biomimetic systems, they have mainly been prepared by two different procedures (Scheme 5): (i) HAA by a superoxido $[Cu^{II}(\eta^1-O_2)]^+$ precursor on a substrate to generate the corresponding $[Cu^{II}(\eta^1-OOH)]^+$ species (many examples were reported using TEMPOH as a hydrogen atom donor) or (ii) a “peroxide-shunt” approach, *i.e.*

treatment of a Cu(II) precursor (or Cu(I)) by H₂O₂ or ROOH (in the presence or absence of a base) yielding the corresponding $[Cu^{II}(\eta^1-OOR)]^+$ species. Several new examples were recently prepared following these strategies.^{85,92,94,107–111} It has to be noted that a novel route involving cryoreduction of a superoxide $[Cu^{II}(\eta^1-O_2)]^+$ precursor followed by annealing has recently been described by the group of Karlin (Scheme 5).⁸⁴ This route has also allowed the isolation of unprecedented Cu^{II}- η^1 -peroxido species.

Copper(II)-hydro(alkyl)peroxido intermediates that have been isolated in model systems all display an η^1 coordination mode. They are characterized by intense UV-Vis features at 350 nm assigned to peroxide-to-copper LMCT as well as EPR signatures characteristic of copper(II) species. Two characteristic vibration modes are generally detected in their rRaman spectra: $\nu(O-O) \sim 850\text{ cm}^{-1}$ and $\nu(C-O) \sim 550\text{ cm}^{-1}$. In 2017, only two $[Cu^{II}(\eta^1-OOR)]^+$ species (R=H or cumyl, Cm) had been structurally characterized.^{112,113} Three additional structures were recently reported.^{85–87}

The group of Tolman has used the dianionic bis(arylcarboxamido)-pyridine ligand (Chart 1) and reacted the $[(L)Cu^{II}OH]^-$ precursor with excess ROOH (with R = ^tBu or Cm).¹² The corresponding $[(L)Cu^{II}(OOR)](Et_4N)$ complexes were successfully crystallized (Fig. 6b for R = Cm).⁸⁷ These correspond to the only additional examples of copper-alkylperoxide species structurally characterized after the first structure reported in 1993. The intermediates display O–O bond distances of 1.416 and 1.468 Å for R = ^tBu and Cm respectively, which agree well with that of the already reported Cu–OOCm intermediate (1.460 Å).¹¹² Interestingly, cyclic voltammetry measurements of the two intermediates revealed quasi-reversible processes at $E_{1/2} = -150$ and $-200\text{ mV vs. }Fc^+/Fc$, respectively. Chemical oxidations led to the isolation of one-electron oxidized species, formally Cu(III)-alkylperoxido species. These intermediates are EPR silent and display intense absorptions in the visible region (500–700 nm). Resonance Raman experiments revealed a $\nu(O-O) = 831\text{ cm}^{-1}$ sensitive to isotopic labeling with ¹⁸O. The exact electronic description (either Cu^{III}-alkylperoxido, or antiferromagnetically coupled singlet ground-state Cu^{II}-ligand radical or ferromagnetically coupled triplet Cu^{II}-ligand radical) could not be unambiguously confirmed using DFT calculations. These formally Cu^{III}-alkylperoxido species did not show any reactivity towards weak O–H or C–H bonds. They were however found to react with 4-dimethylamino-phenol (^{NMe2}ArOH) yielding a Cu(II)-phenoxy radical product, *via* possible PCET or ET/PT pathways.

Using the biotin-streptavidin strategy used to develop artificial metalloenzymes,¹¹⁴ the group of Borovik have anchored a biotinylated dpea (dipyridine ethylamine) ligand into a wild-type (WT) streptavidin (Sav) scaffold ($[(\text{biot-et-dpea})Cu^{II}(H_2O)_2]^{2+} \subset \text{Sav}$ (with an ethyl spacer between biotin and dpea)).¹¹⁵ Addition of hydrogen peroxide to a buffered solution of the Cu(II) artificial enzyme led to the formation of a $[(\text{biot-et-dpea})Cu^{II}(OOH)(OH_2)]^+ \subset \text{Sav}$ intermediate, persistent for one day at room temperature.⁸⁶ The intermediate was also generated by incubating crystals of the Cu(II) precursor with hydrogen peroxide, which allowed the obtention of crystallographic data



at 1.62 Å resolution (Fig. 6c). The copper is found in a 5-coordinated distorted trigonal bipyramidal geometry with 3 nitrogen atoms from the dpea ligand, a water molecule and an end-on hydroperoxido ligand. The Cu–O distance is found to be equal to 1.94 Å and the O–O distance is equal to 1.52 Å. The structure reveals that the streptavidin scaffold provides a H-bonding network to both the distal and the proximal O-atoms of the hydroperoxido ligand, H-bonded to asparagine residue N49 or to a water molecule stabilized by an S112 residue respectively. Remarkably, this structure represents the second crystallographic data obtained on Cu(II)–OOH species. Mutations of serine (S112) or asparagine (N89) into alanine were performed to evaluate the effect of H-bonding disruption on the stability of the intermediate. In the case of N89A, no drastic difference in stability was noted. In contrast, mutation of S122 into alanine resulted into drastic loss of stability of the intermediate. Although no crystal structure of the Cu(II)–OOH species could be obtained, the structure of the ([biot-et-dpea] Cu^{II}(H₂O)₂)²⁺ ⋅ Sav S112A precursor indicated that the occupancy of the targeted water molecule was reduced by 50%. In addition, the Cu complex was no longer located into a single position but over two positions, consistent with the importance of H-bonding interactions to govern the position of the complex within the Sav cavity. The reactivity of the different copper-hydroperoxido species confined within the native and mutant sav scaffolds was evaluated using 4-chlorobenzylamine as substrate. The copper-hydroperoxido species confined within the native and N49A Sav scaffolds were not active. In contrast, ([biot-et-dpea] Cu^{II}(H₂O)(OOH))²⁺ ⋅ Sav S112A was found to react with the substrate. These results nicely corroborate the known effects of H-bond interactions to either the proximal or distal atoms of the OOH ligand on the properties of Cu(II)–OOH species.^{116–118}

In most cases, the Cu–OOR intermediates are found in square-pyramidal or bipyramidal geometries. Interestingly, the groups of Itoh and Comba have isolated a Cu–OOCm intermediate using a rigid and bulky tridentate ligand, ^{TMG3}tach (Chart 1), that imposes an unusual tetrahedral geometry to the copper.¹¹⁰ The intermediate was prepared by reacting the methoxide precursor [(^{TMG3}tach)Cu^{II}(OMe)]⁺ with excess CmOOH. Its spectral properties were highly similar to those of alkylperoxido intermediates with different geometries.

Finally, although many Cu–OOR(H) species have been isolated and characterized, very few Cu–acylperoxido species have been studied.^{119–121} Using a ligand based on a rigid 6-membered cyclic diamine with two CH₂Py side arms (L6^{PyM2}, Chart 1), the group of Itoh has recently isolated a Cu(II)–acylperoxido intermediate upon reaction of Cu(II) complexes with mCPBA.¹²² The purple complex [(L6^{PyM2})Cu^{II}(OOCAR)]⁺ was found to be stable at 30 °C. Using similar ligands but with 7- or 8-membered rings, the adducts were unstable and converted to the corresponding *m*-chlorobenzoic acid (*m*-CBA) adducts.

3.2.2. Reactivity. Copper(II)–alkylperoxido or hydroperoxido species are considered as central intermediates in catalytic reactions and were detected in the case of several bioinspired complexes and more recently in LPMO mimics.^{42,123–125} Their reactivity can be modulated by the nature of the ligand as well as by H-bonding interactions with the distal or proximal oxygen

atoms. However, they are generally poorly efficient for HAA reactions and are often considered as precursors for more reactive species. In many cases, the intermediates were suggested to undergo O–O bond homolysis, generating Cu^{II}–O[•] and RO[•] reactive species for HAA reactions.⁵ Several studies were recently reported strengthening the hypothesis of O–O homolysis post reactivity.^{109,126} In some cases, the O–O bond cleavage of LCu^{II}–OOR complexes was proposed to occur concertedly with the C–H bond activation of external substrates such as 1,4-cyclohexadiene (1,4-CHD).^{110,127} Interestingly, a recent study from Cho and coworkers evidenced the importance of solvent in the reactivity of [(^{iPr3}tren)Cu^{II}(OOCm)]⁺.¹²⁸ In particular, non-polar solvents were suggested to enhance the reactivity towards the external C–H bonds of a substrate *via* solvation effects leading to more efficient O–O homolytic cleavage and HAA reactions. Also, the absence of C_{sp3}–H bond on the solvent was proposed to reduce the consumption of reactive species.

Using the ligand CHDAP (*N,N'*-dicyclohexyl-2,11-diaza [3,3']-(2,6)pyridinophane) the group of Cho prepared two [(CHDAP)Cu^{II}(OOR)]⁺ intermediates (with R = C(CH₃)₂Ph and ^tBu) that were characterized using UV-Vis, EPR, resonance Raman and mass spectrometry.¹⁰⁷ Self-decay of the intermediates at 25 °C followed first-order kinetics and was shown to involve O–O bond homolysis. The stability of the intermediates was not affected by the addition of thioanisole or cyclohexadiene, consistent with the fact that they are poorly/not reactive in electrophilic C–H bond activation. In contrast, reaction with aldehydes and benzoyl chlorides was observed and a mechanism involving nucleophilic attack of the alkylperoxido ligand on the carbonyl groups was proposed. A few years later, the same group showed that a copper-hydroperoxido intermediate, [(^{iPr3}tren)Cu^{II}(OOH)]⁺, was also capable of nucleophilic reactivity with acyl chlorides and with aldehydes for deformylation reactions.¹⁰⁸ In the case of aldehyde deformylation, two routes were discussed that involve a nucleophilic attack of either the proximal or the distal oxygen from the intermediate. These two studies highlight new properties of CuOOR species that will need to be further explored and understood. Aromatic hydroxylation by Cu–OOH species was also reported by the groups of Mayilmurugan and Ghosh.^{129,130}

In the catalytic cycle of LPMO, it has been suggested that the Cu(I)–(H₂O₂) intermediate might be involved (Scheme 3).⁵⁵ Although already suggested as possible intermediates in oxidation reactions,^{131,132} such species have only been recently detected in a model complex (Scheme 2).⁸³

Recently, the groups of Karlin and Solomon have provided experimental evidence that such a peroxygenase pathway can be efficient in bioinspired model chemistry.¹³³ The complex [(^{TMG3}tren)Cu^I]⁺ was treated with a sub-stoichiometric amount of dry H₂O₂ yielding the hydroxylated ^{TMG3}tren-OH ligand. Interestingly, no Cu(II) was detected at the end of the reaction suggesting a Cu(I) redox state of the complex. When hydroxyl radical scavengers were added, ligand hydroxylation was cancelled supporting the involvement of hydroxyl radicals and Fenton-like pathways. In the presence of a slight excess (3 equivalents) of H₂O₂, overoxidation occurred and the alkoxide complex [(^{TMG3}tren-O)Cu^{II}]⁺ resulting from ligand hydroxylation





Scheme 8 Reaction of $[(\text{TMG}^3\text{tren})\text{Cu}]^+$ with varying amounts of H_2O_2 . Adapted from ref. 133.

was isolated. Overall, experimental results pointed towards a “peroxygenase-like” mechanism with the involvement of controlled Fenton pathways for the hydroxylation of adequately positioned C–H bonds (Scheme 8).

Similar conclusions were obtained by the group Garcia-Bosch using bppa, a tmpa based ligand (noted LH_2 by the authors) that possesses two H-bond donors on the pyridine ligands.¹²⁴ The authors have investigated the reactivity of mononuclear Cu complexes with O_2 and H_2O_2 . $[(\text{LH}_2)\text{Cu}^{\text{I}}]^+$ was reacted with dioxygen in acetonitrile at -35°C and, in the presence of TEMPOH, full conversion to $[(\text{LH}_2)\text{Cu}^{\text{II}}(\text{OOH})]^+$ species was achieved. Addition of triflic acid led to the release of H_2O_2 together with $[(\text{LH}_2)\text{Cu}^{\text{II}}]^+$. This reactivity was discussed as relevant for the production of hydrogen peroxide from dioxygen by LPMO. Upon reaction of $[(\text{LH}_2)\text{Cu}^{\text{I}}]^+$ with H_2O_2 , $[(\text{LH}_2)\text{Cu}^{\text{II}}(\text{OH})]^+$ was produced which then evolved towards $[(\text{LH}_2)\text{Cu}^{\text{II}}(\text{OOH})]^+$ when excess H_2O_2 was used. Hydroxyl radical formation was also evidenced. The chemistry in play here seems therefore highly similar to the one described above¹³³ and could be of relevance for LPMO peroxygenase pathway.

3.3. Copper-oxyl and high-valent copper oxygen species

High-valent Cu cores have been proposed as a key species in the catalytic cycle of Cu-containing monooxygenases and in synthetic oxidation catalysts. Increasing number of formally Cu(III) complexes, mainly organometallic ones, are described in the literature, although the existence of Cu(III) and the exact redox state of the metal center is questioned.^{72,87,134–143} Most of these species are beyond the scope of this review which focuses on copper-oxygen species for C–H bond activation. Two main species will therefore be discussed here: $[\text{CuO}]^+$ and the protonated $[\text{CuOH}]^{2+}$ cores.

$[\text{CuO}]^+$ cores are often postulated in the catalytic cycles of copper-containing monooxygenases and synthetic model complexes for the activation of strong C–H bonds such as those of polysaccharides or methane.^{5,55,140,144} Calculations support the electronic description as triplet ground state Cu(II)-oxyl species ($[\text{Cu}^{\text{II}}\text{O}^\bullet]^+$). Based on the biradical character as well as the predicted weak Cu–O bond, they are expected to be highly reactive, probably explaining the lack of experimental evidence gathered on this reactive species. However, beyond numerous computational studies, the only data on the reactivity of such high-valent species came from gas-phase reactions, confirming their ability to perform C–H abstraction on alkanes, including methane.^{144–147} Attempts to isolate $[\text{CuO}]^+$ cores in solution upon reaction of a Cu(I) precursor with oxo transfer reagents such as pyridinium *N*-oxides or PhIO (Scheme 5) remained unsuccessful (with the exception of one ESI-MS data set).¹⁴⁸ Following this strategy, the group of Karlin has recently reacted several Cu(I) complexes with *para*-substituted *N,N*-dimethylaniline *N*-oxides (DMAO) as O-atom donors. The reported results pointed towards the involvement of reactive copper-oxyl species, able to hydroxylate the strong C–H bonds ($\sim 90 \text{ kcal mol}^{-1}$) of the *N,N*-dimethylaniline product generated after O-atom transfer reactions to the Cu(I) precursors.⁹⁸

The suggestions that $[\text{CuOH}]^{2+}$ cores could be of biological relevance came from the work of the group of Tolman.^{5,140} Using electron donating bis(arylcarboxamido) ligands, $[\text{CuOH}]^{2+}$ species were generated upon one-electron oxidation of the corresponding Cu(II) precursors and a Cu(III)–OH formulation was suggested based on spectroscopic data and DFT calculations. These intermediates were shown to react with strong C–H bonds (up to 99 kcal mol^{-1} for cyclohexane).⁶⁵ This strong C–H bond activation ability was correlated with the strong basicity of the Cu(II)–OH precursors compensating the weak oxidizing power of the $[\text{CuOH}]^{2+}$ core leading to a strong O–H BDE of the Cu(II)–(OH₂) species and thus to a strong thermodynamic driving force to attack energetic C–H bonds (Scheme 6).^{140,149} The group of Cramer computationally rationalized the effect of various factors on the reactivity of such high valent $[\text{CuOH}]^{2+}$ cores, including ligand modifications, counter ions, van der Waals interactions and quantum-mechanical tunneling effects.¹⁵⁰ Based on their oxidative power, it has been anticipated that such species could be involved in the catalytic cycle of LPMO, possibly stabilized by deprotonation of the terminal primary amine of the histidine brace motif.¹⁵¹ It has to be noted that deprotonation of the terminal amine of the histidine brace has been suggested to occur at high pH based on EPR and UV-Vis data.¹⁵² Interestingly, high valent $[\text{CuOH}]^{2+}$ cores were recently postulated in the course of H_2O_2 splitting by a second-coordination sphere mutant of a fungal LPMO based on DFT calculations.¹⁵³

Following previously reported work, the group of Tolman has successfully stabilized $[(\text{L})\text{Cu}(\text{OR})]$ cores (with $\text{R}=\text{H}$ or CH_2CF_3) and obtained X-ray structural data using a bis(arylcarboxamido)-pyridine ligand bearing a methoxy substituent on the pyridine moiety (Fig. 6d).⁸⁷ The two neutral intermediates were found in square-planar geometries with metal–ligand bond distances significantly shortened (average of 0.102 \AA) as



compared to that found in the Cu(II) precursors, in agreement with previously predicted contractions using EXAFS analyses and DFT calculations. Resonance Raman spectra of $[(L)Cu(OR)]$ complexes displayed a $\nu(Cu-OR)$ vibration around 635 cm^{-1} similar to the ones already reported for known high-valent Cu-OH species using similar ligands. Finally, NMR spectra were consistent with diamagnetic $S = 0$ ground-states supporting the description as $Cu^{III}-OH$ cores, in agreement with previous proposals.¹⁴⁰ This study represents the first structural characterization of a $[CuOH]^{2+}$ core as well as the first isolation of a Cu^{III} -alkoxide analog.

The groups of Tolman and Cramer have also reported on a complete mechanistic investigation of the C-H and O-H bond abstraction mechanism of three different high valent species using the bis(arylcarboxamido)-pyridine ligand:⁷² $[(L)Cu^{III}(OH)]$, $[(L)Cu^{III}(OOCm)]$ ¹⁰³ and $[(L)Cu^{III}(OCOAr)]$ a formally $Cu(III)$ -carboxylate species (Ar = *meta*-chlorophenyl) prepared for the first time in this study. It was observed that the most reactive species among the three was the $Cu(III)-OH$ species. Then the carboxylate complex was found to be more reactive than the alkylperoxido one (mostly unreactive). Detailed DFT calculations indicated that O-H or C-H bond abstraction proceeds with different synchronicity of the CPET/HAT depending on the nature of the intermediate and on the substrate. Although C-H bond abstraction from dihydroanthracene (DHA) by $[(L)Cu(OH)]$ follows a HAT mechanism, O-H bond abstraction from phenol by the same intermediate is best described as a concerted "separated" cPCET with proton and electron transfers involving different acceptor centers (the O and the Cu atoms respectively). In the case of $[(L)Cu(OCOAr)]$, both reactions were found to proceed by oxidative asynchronous cPCET, probably driven by the greater oxidizing power of the copper(III) species and the lower basicity of the carboxylate ligand.

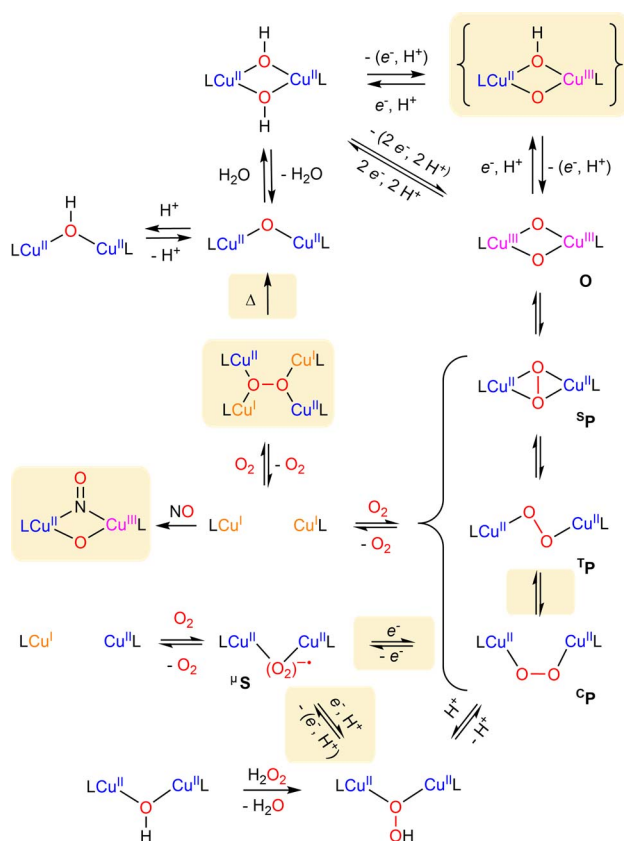
It is worth mentioning that several additional $[Cu-Y]^+$ cores have been isolated and studied using bis(arylcarboxamido)-pyridine ligands (Y = SR, halide, CN, NO₂) providing interesting tools to investigate the electronic structures and reactivity of high-valent copper cores.^{141–143,154,155} In particular, using halides, the group of Zhang has stabilized high-valent $[Cu-X]^+$ species.¹⁴² Calculations have suggested significant halide radical characters for all the complexes which are therefore best described as $[(L)Cu^II(X^{\bullet})]$, making these species analogs to copper-oxyl ones. The $[(L)CuX]$ species were able to perform HAA from DHA. In addition, upon heating of $[(L)CuX]$ species in

the presence of azobis(isobutyronitrile) (AIBN) as a carbon-centered radical precursor, formation of 2-haloisobutyronitriles was observed demonstrating the ability of these high-valent species to perform radical capture and halogenation. $[(L)CuF]$ was then employed for the direct fluorination of a variety of alkyl substrates with allylic and benzylic C-H bonds as well as α -C-H bonds of ethers in a 2 : 1 Cu : substrate stoichiometry. These findings were further developed to establish electrocatalytic C-H fluorination of various substrates.¹⁵⁵

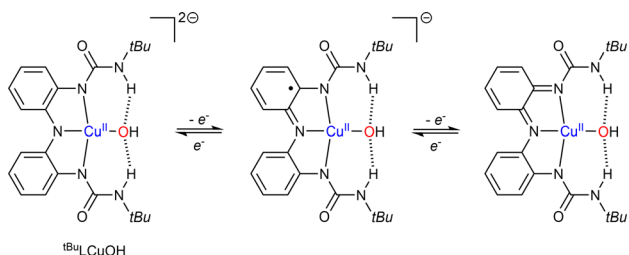
Finally, the groups of Lancaster and Garcia-Bosch have studied a copper-hydroxide complex based on tridentate redox active ligand (Scheme 9).⁷³ The starting $[(^{tBu}L)Cu^{II}-OH]^{2-}$ species was shown to undergo two reversible ligand-centered oxidation processes. The three Cu-OH cores did not show any reactivity with C-H bonds but oxidized phenols featuring relatively weak O-H bonds.

4 Dinuclear copper–oxygen species

Since the first structurally characterized *trans*-peroxido ($^T\mathbf{P}$) and μ - $\eta^2:\eta^2$ -peroxido ($^S\mathbf{P}$) dicopper(II) species (Scheme 10) by Karlin¹⁵⁶ and Kitajima¹⁵⁷ in the late eighties, numerous studies related to the characterization and reactivity of dicopper-oxygen adducts in biological and synthetic systems have been reported. Reviews related to dinuclear systems for the 2014–2017



Scheme 10 Summary of known formation and interconversion pathways of dinuclear copper–oxygen entities. Recently described species and pathways are highlighted.



Scheme 9 Characterized oxidation states reached via reduction/oxidation of $^{tBu}LCuOH$. Adapted from ref. 73.

period^{1,3,5,6,158} have remarkably highlighted various equilibria between isomeric copper-oxygen species, demonstrating how this could impact their oxidative properties. In the following section, we present recent advances in the field of dinuclear Cu₂/O₂ adducts particularly on isomer interconversion and C-H bond activation.

4.1 Peroxido dicopper(II) core

As shown in Scheme 10, peroxido dicopper(II) complexes can be prepared from the oxygenation of either mononuclear or dinuclear copper(I) complexes. As previously reported in many reviews,^{3,5,6,158,159} various types of dicopper(II) peroxides differing by their configuration can be obtained through subtle variation of controllable parameters (ligand, solvent, temperature...), as described in this section.

4.1.1 New intermediate peroxido species. Following previous works based on pyrazolato/TACN (TACN for triazacyclononane) dicopper-oxygen systems, Meyer reported in 2022 a new type of peroxide, named as 'intermediate' ([⊥]P) peroxido species (Fig. 8) with a Pyr/TACN_{1,2} ligand (Chart 2). The particularity of this adduct was to display an unusual Cu–

O–O–Cu dihedral angle φ close to 90°, yielding an intermediate geometry between *cis*-peroxides ^CP ($\varphi \approx 0^\circ$) and *trans*-peroxides ^TP ($\varphi \approx 180^\circ$).^{160,161} Comparison with previously reported analogous complexes^{160,162} indicated that the Cu...Cu distance, Cu–O–O–Cu torsion angles and spin state were significantly impacted by the spacer length. For [⊥]P, the ferromagnetic coupling ($J = 25 \text{ cm}^{-1}$) was attributed to the orthogonality of the magnetic orbitals with the bridging peroxide. Magnetic investigations associated with Frequency-Domain Fourier Transform Terahertz EPR (FD-FT THz-EPR)¹⁶³ following a methodology deployed at the BESSY II synchrotron¹⁶⁴ were successfully employed¹⁶¹ for probing the spin state of [⊥]P species^{160,161} (Fig. 8). Such species are of interest since they represent relevant proposed intermediates formed after O₂ binding on type III dicopper metalloprotein (*i.e.* hemocyanin or tyrosinase enzyme), prior to formation of the identified planar Cu₂O₂ core (^SP)¹⁵⁹ observed in oxy forms.

4.1.2 *Cis* (^CP) and *trans* (^TP) peroxido interconversion. In 2021, Lehnert, Ertem and Robinson¹⁶⁵ identified reversible interconversion of ^TP/^CP dicopper cores mediated by calcium using the well-known $\{[\text{Cu}^{\text{II}}(\text{tmpa})]_2(\mu\text{-}1,2\text{-O}_2)\}^{2+}$ system (Chart 1 for the ligand). The same year, the group of Meyer¹⁶⁶ established similar reversible interconversions induced by the binding of various alkali metal ions. The direct interactions of Ca²⁺, Li⁺, Na⁺ and K⁺ cations on the peroxido core were established by the obtention of the corresponding X-ray structures. Interestingly, ion-binding induced significant changes on the adduct's properties (electronic, magnetic, electrochemical...) depending on the nature of the bound alkali cation. These variations of properties were explained on the basis of theoretical calculations, showing modifications of the charge transfer from the copper ion to dioxygen with the alkali cation.¹⁶⁶ Such phenomena highlight the importance of taking into consideration the presence of alkali metal ions in reaction media (synthetic or biological).

4.1.3 Dinuclear Cu^{II} peroxido/superoxido interconversion. Dinuclear superoxido species have drawn attention due to their potential oxidative properties towards inert C–H bonds. As shown in Scheme 11, $\mu\text{-}1,1$ and $\mu\text{-}1,2$ superoxido-dicopper(II) complexes can be prepared either by 1e[−] oxidation of $\mu\text{-}1,2$ -



Fig. 8 Summary of the structurally characterized dicopper(II) peroxido species with pyrrolato/TACN based ligands from the Meyer group.^{160–162,167} φ represents the torsion angle Cu–O–O–Cu and θ the Cu–O–O angle.



Scheme 11 Dinuclear Cu^{II} peroxido/superoxido species interconversions.

peroxido Cu_2^{II} species or H-atom abstraction from μ -1,1 hydroperoxido adducts.

In 2019, Karlin reported the generation and characterization of two different superoxido dicopper species by chemical oxidation from a μ -1,2 peroxido complex based on the XYLO(H) dinucleating ligand (XYLO(H) = 2,6-bis{[(bis(2-pyridylethyl) amino)methyl]phenol}) (Chart 2 and Table 1).¹⁶⁸ A mixture of μ -1,2 and μ -1,1 superoxido species was detected by UV-Vis and rR spectroscopies exhibiting similar behavior to other superoxido derivatives previously described. In 2021, by adopting the ligand used for the previous structural characterization of a dinuclear copper(II) *cis* peroxido,¹⁶² Meyer reported the X-ray structures of both peroxido and superoxido species bearing the same pyrazolato/TACN_{2,2} ligand¹⁶⁷ (Chart 2 and Fig. 8). Redox investigation revealed one electron process characterized by a small reorganization energy. This was in accordance with other superoxido/peroxido interconversions reported in solution, reflecting that the redox process occurs at the peroxido-unit.^{169–171} Very recently Karlin and co-workers have completed the studies using a phenolato bridged ligand (BPMPO(H), Chart 2 and Table 1) analogous to XYLO(H).¹⁷² A peroxido-dicopper(II) complex was characterized and shown to undergo reversible oxidation to a superoxido-dicopper(II) species. The various $\text{Cu}_2^{\text{II}}(\text{O}_2^-)$ complexes (in the nature of the dinucleating ligand) were compared and discussed with respect to physical properties and reactivity.¹⁷² Based on spectroscopic, titration and electrochemical data, thermodynamic parameters ($E_{1/2}$, $\text{p}K_{\text{a}}$) were obtained, allowing a comparative analysis (Table 1). For each species described (peroxido, superoxido and hydroperoxido) despite various bridging ligands and various experimental conditions (temperature and solvent), the UV-Vis absorptions for each form are in a similar energy range. Similarly, O–O stretching frequencies obtained from rR ($\nu(\text{O}=\text{O})$) are between 798 and 815 cm^{-1} for all peroxido species (Table 1). The rR data on superoxido species range from 1070 cm^{-1} (Cu_2^{II} -

O_2^- based on the Pyr-TACN_{2,2} ligand) to 1143 cm^{-1} ($\text{Cu}_2^{\text{II}}\text{O}_2^-$ based on the XYLO[−] ligand). The decrease of the O–O stretching frequencies correlates with the weaker O–O bonds. In more detail, from various phenolato (Karlin group)^{168,172} and pyrazolato (Meyer group)^{161,167,171} bridging ligands, the particular binucleating ligand design was shown to considerably influence/control the reduction potential of the superoxido-dicopper(II) complex, the OO–H $\text{p}K_{\text{a}}$ and BDFE (see part 2.2, Table 1 and Scheme 4). The phenoxido-bridged complexes appeared to be more basic compared to the pyrazolato/TACN_{2,2} based complex (Pyr/TACN_{2,2} ligand, Chart 2 and Table 1). They were also demonstrated to be good candidates for HAA of inert C–H bonds (BDFE > 80 kcal mol^{-1}). EPR analysis for $[\text{Cu}_2^{\text{II}}(\text{Pyr}/\text{TACN}_{2,2})(\text{O}_2^-)]^{2+}$ and $[\text{Cu}_2^{\text{II}}(\text{BPMPO})(\text{O}_2^-)]^{2+}$ displayed simple mono- Cu^{II} ($S = \frac{1}{2}$) type axial spectra, thus indicating that the unpaired electron at the superoxide moiety was antiferromagnetically coupled to one of the Cu^{II} centers, leaving the remaining spin on the detected second Cu^{II} ion. In contrast $[\text{Cu}_2^{\text{II}}(\text{XYLO})(\text{O}_2^-)]^{2+}$ and its analog $[\text{Cu}_2^{\text{II}}(\text{UNO})(\text{O}_2^-)]^{2+}$ displayed at $g = \sim 2.0$ a typical organic radical single line spectra, in accordance to a localized unpaired electron on the O_2^- moiety associated with antiferromagnetic coupling occurring between the two copper(II) ions of the complexes.¹⁶⁹ This different behaviors pointed out the influence of chelate ring size on electronic-bonding properties.

These significant advances owe a lot to the recent development of cryogenic spectroscopies and analysis for unstable species identification¹⁷³ (*i.e.* low-temperature spectroelectrochemistry reviewed in 2021 by Le Poul¹⁷⁴) associated with theoretical modelling at low temperatures. Since a lot of reactive intermediates have been characterized by cryo-ESI-MS (cold-spray ionization-mass spectrometry), the association of such ESI together with tandem MS could be of interest in the field of copper as reported for iron oxido intermediates.¹⁷⁵

Table 1 Comparison of the main physical and thermodynamic properties determined for $\text{Cu}_2^{\text{II}}(\text{O}_2^{2-})$, $\text{Cu}_2^{\text{II}}(\text{O}_2^{\cdot-})$ and $\text{Cu}_2^{\text{II}}(\text{OOH})$ from Meyer^{162,167,171} and Karlin^{168,172}

Ligands (references)	Pyr/TACN _{2,2} (ref. 162, 167 and 171)	XYLO(H) (ref. 168)	BPMPO(H) (ref. 172)
Conditions of the studies	CH_3CN , $-20\text{ }^\circ\text{C}$	MeTHF, $-125\text{ }^\circ\text{C}$	MeTHF, $-90\text{ }^\circ\text{C}$
Dicopper(II) peroxido			
rR, $\nu(\text{O}=\text{O})\text{ cm}^{-1}$	798	803	815
UV-Vis	527; $\epsilon = 5000$	523; $\epsilon = 7150$	536; $\epsilon = 8500$
λ_{max} (nm); ϵ ($\text{M}^{-1}\text{ cm}^{-1}$)	648; $\epsilon = 3900$		420 620
Dicopper(II) superoxido			
rR $\nu(\text{O}=\text{O})\text{ cm}^{-1}$	1070	1143	1086
UV-Vis, λ_{max} (nm); ϵ ($\text{M}^{-1}\text{ cm}^{-1}$)	440; $\epsilon = 11\,500$	402; $\epsilon = 18\,750$ 480; $\epsilon = 4380$	405; $\epsilon = 15\,100$
$E_{1/2}$ vs. $\text{Fc}^+/\text{O}_2^{\cdot-}/\text{O}_2^{2-}$	-0.59 V	-0.53 V	-0.44 V
Dicopper(II) hydroperoxido			
UV-Vis	416; $\epsilon = 5700$	398; $\epsilon = 10\,725$	391; $\epsilon = 5800$
λ_{max} (nm); ϵ ($\text{M}^{-1}\text{ cm}^{-1}$)			630
OO–H $\text{p}K_{\text{a}}$	22.2	24	22.3
OO–H BDFE kcal mol^{-1}	69.4	80.7	80.3



4.1.4 Peroxido dicopper(II)/bis(μ -oxido)dicopper(III) conversion.

The first $[\text{Cu}_2^{\text{III}}(\mu\text{-O})_2]^{2+}$ (**O**) species was structurally described in 1995 by the group of Tolman.¹⁷⁶ The same group reported in 1996 a possible equilibrium between **O** and **^SP** species.¹⁷⁷ Another type of interconversion was described in 2014 by Karlin and co-workers between **O** and **^TP** species.¹⁷⁸ Since then, a number of subsequent studies related to the factors tuning these isomeric equilibria (ligand structure and electronics, steric effects, acid–base control,¹⁷⁹ choice of solvent¹⁸⁰ ...) have been reported and reviewed. For example, Dey,¹⁸¹ in a series of ethylenediamine-based dinuclear copper(II) complexes (Chart 1), reproduced experimental observations using DFT calculations. The strength of the nitrogen donors was correlated with the difference in energy between the corresponding peroxido and oxido complexes. The stronger the donor, the weaker the difference.

In another example, DeBeer used Valence-to-Core (VtC) X-ray emission spectroscopy (XES)¹⁸² as a sensitive probe to discriminate between **^SP** and **O** cores through O–O bond detection.¹⁸³ More generally, such synchrotron-based X-ray spectroscopy approaches have been more frequently used in these last few years for the characterization of dicopper(II)–oxygen adducts, with the support of theoretical calculations, and provide their electronic structure description.¹³⁸

4.2 Peroxido species as an active intermediate and reactivity

Iron, manganese or cobalt complexes are the most predominant dinuclear systems reported in the literature for C–H hydroxylation.¹⁸⁴ They display multiple oxidation or spin states and can generate binuclear peroxido, superoxido or oxido reactive intermediates in the presence of dioxygen. Although dinuclear copper complexes share similar features, few studies have exploited these behaviors to activate substrates.

Nevertheless, in the last few years, significant advancements were made in the use of low molecular weight dicopper complexes capable of activating inert C–H bond in a direct manner through bioinspired approaches. Such developments are presented in the following parts.

4.2.1 Peroxido dicopper(II)/bis(μ -oxido)dicopper(III). As the **^SP** core represents a key intermediate identified in oxygenated type III copper proteins,¹ many synthetic systems yielding **^SP** adducts have been developed in the last few years, with the objective of performing electrophilic aromatic substitutions. Usually, the corresponding complexes display bi- or tridentate nitrogen-based ligands affording an exchangeable position for substrate fixation. The typical reaction is the *ortho*-hydroxylation of phenolic substrates to catechols and the subsequent two-electron oxidation in the corresponding quinones in relation with tyrosinase enzyme activity (see Fig. 1a) and also with diverse aliphatic C–H bond activations.¹⁸⁵ Since the first catalytically active model,¹⁸⁶ examples of dinuclear **^SP** or **O** systems that can perform C–H activation in a catalytic manner have remained scarce. For instance, several active **^SP** or **O** intermediates were established by Tucek's group by means of various bidentate or tridentate pyrazole or triazole based ligands.^{187–189} Alternatively, Herres-Pawlis and co-workers developed hybrid

guanidine based ligands that led to the formation of dinuclear catalytic systems (Chart 1).^{190–192} Their reactivities towards various natural and non-natural phenolic substrates in the presence of molecular oxygen yielded mono- and bicyclic quinones. Besides, chiral tyrosinase biomimetic model complexes for stereoselective oxidation were reviewed in 2018 by Casella and Monzani¹⁹³ and a new dinuclear chiral complex was described in 2019 (ref. 194) by the same group.

In some cases, such side-one (**^SP**) species exhibited no phenol *ortho*-hydroxylation (tyrosinase like-activity) as reported by Company for dinuclear PyNMe-based peroxides (PyNMe = 3,6,9-trimethyl-3,6,9-triaza-1(2,6)-pyridinacyclodecaphane, Chart 1).¹⁹⁵ The presence of the **^SP** species was ascertained by a unique highly intense UV-Vis signature at 353 nm analogously to other **^SP** compounds. The steric hindrance, rigidity and the saturated coordination imposed by the tetradentate PyNMe ligand prevented phenolic substrate binding and explained that no phenol *ortho*-hydroxylation could be detected. In another example, the room-temperature stable and inert **^SP** complex based on an hexapyridine ligand¹⁹⁶ (h6m4h = 1,2-bis[2-(bis(6-methyl-2-pyridyl)methyl)-6-pyridyl]ethane ligand, Chart 2) was investigated by Kodera and Le Poul.¹⁹⁷ The two-electron reduction of the **^SP** complex resulted in the O–O bond cleavage, but no C–H or O–H bond activation of exogeneous substrates was detected. This result was attributed to the negative standard potential and weak basicity of the bis-reduced species, precluding any HAA process due to a low BDFE.

Dicopper(III) bis(μ -oxido) adducts (**O**) resulting from the conversion of **^SP** species by O–O bond breakage have also been suggested to be active species.^{198,199} Since the pioneering work of Itoh,^{200–202} various examples of such reactivity have been described and reviewed.^{203,204} Karlin, Solomon and Garcia-Bosch⁷⁴ converted a non-active **^SP** in an active **O** species *via* two possible processes following coordination of either: (i) an external substrate (sodium phenolate) that was further *ortho*-hydroxylated (tyrosinase like-activity) or (ii) a Lewis acid (one equivalent of $\text{Sc}(\text{CF}_3\text{SO}_3)_3$ or two equivalents of $\text{B}(\text{C}_6\text{F}_5)_3$ or $\text{DMF} \cdot \text{CF}_3\text{SO}_3\text{H}$) in the ligand scaffold able to enhance the C–H bond activation ability. More recently, the formation of the reactive **O** species after addition of two equivalents of a phenolic



Scheme 12 Complex LCu(I) ($\text{L} = \text{tmpa}$ (tris(2-pyridylmethyl)amine)) after reaction with O_2 according Göttlich and Schindler²⁰⁶ and proposed general catalytic cycle for toluene oxidation (adapted from ref. 206).



substrate was explored by Stack with diamine ligands (Chart 1).²⁰⁵ The reactivity studies were extended to hydroxylation associated with the dehalogenation of 2-halophenolate substrates.

An attractive one-pot approach has been recently developed by Göttlich and Schindler,²⁰⁶ involving a dicopper(II)-O₂ adduct in combination with a photochemical reducing agent. Reaction of 2 equivalents of the LCu^I complex (L = tmpa ligand) with O₂ yielded an active dinuclear intermediate (**O** or **^SP**). In toluene solution, selective catalytic oxidation of toluene into benzaldehyde at room temperature was achieved, without formation of benzyl alcohol or over-oxidized products (Scheme 12). Regeneration of the LCu^I active species was performed by using a ruthenium bipyridine photosensitizer [Ru(bpy)₃](PF₆)₂ in the presence of triethyl amine (TEA, sacrificial electron donor) and irradiation at 475 nm. Under these conditions, the tmpa complex achieved a toluene conversion of 42%. Briefly, the proposed mechanism involved a dinuclear intermediate leading to the benzyl radical PhCH₂• through a H-atom abstraction from toluene. Reaction of PhCH₂• with O₂ yielded the benzylperoxido radical (PhCH₂OO•) after additional hydrogen abstraction on toluene to the corresponding reactive hydroperoxide PhCH₂-OOH species which evolves to benzaldehyde and water.

4.3 Dicopper-μ-oxido species as active intermediate

Dicopper(II)-μ-oxido complexes are accessible from multiple routes, including the reaction of Cu(I) complexes (mono or dinuclear) with O₂, PhIO and even with NO.²⁰⁷ Already, Limberg

mentioned in 2014 that “complexes featuring Cu^{II}-O-Cu^{II} units do not have a developed chemistry”.²⁰⁷ Since then, studies have remained focused on the spectroscopic features and reactivity of such entities. Dicopper(II)-μ-oxido species have been proposed as active intermediates in homogeneous catalytic activation of inert alkanes (such methane) as well as in heterogeneous catalysts (see Section 6 below).

4.3.1 Cu₂^{II}O from Cu₄^{II}O₂ species and C-H activation. In 2021, Tuczek²⁰⁸ reported the binding of O₂ on a [Cu₂-¹(bdptz)(CH₃CN)₂]²⁺ complex (bdptz for 1,4-bis(2,2'-dipyridylmethyl)phthalazine, Chart 2) at low temperature (−80 °C), leading to the formation of a tetranuclear species (Scheme 10). Upon temperature warming, homolytic cleavage of the O–O bond resulted in the generation of two mono-μ-oxido dicopper(II) complexes (Scheme 10). The formation of the Cu₂^{II}O core was evidenced by resonance Raman (Cu–O vibration at 629 cm^{−1}) and cryo-ESI MS, both including measurement with labelled oxygen and EXAFS/XANES studies. In acetone, the Cu₂O species displayed catalytic activity in oxygenation of various hydrocarbons (xanthene, dihydroanthracene, fluorene, diphenylmethane) with BDFE ranging from 75 to 82 kcal mol^{−1}. Despite poor turnover numbers, the ability of this Cu₂O intermediate in catalytic activity was demonstrated and now opens new catalytic routes in solution.

4.3.2 Cu₂^{II,III} mixed valent μ-oxido like species and C-H activation. Dicopper(II) μ-hydroxido species have been extensively characterized and since a long time regarded as poorly reactive intermediates and even considered as ‘dead ends’ in



Fig. 9 (Top) Starting materials for the preparation of reported dicopper high valent species. (Down) Selected experimental properties (A (wet DMF),²⁰⁹ B²¹⁵ and D²¹⁶ (CH₃CN) and C²¹⁰ (DMF)), *E* in V vs. Fc⁺/Fc.

catalytic processes initiated from high-valent $\text{Cu}^{\text{II}}\text{Cu}^{\text{III}}$ bis-oxido (O) complexes (known to abstract H atoms from weak to moderate C–H bonds). On the other hand, rare examples of mixed-valent $\text{Cu}^{\text{II}}\text{Cu}^{\text{III}}\mu\text{-OH}$ (or bis- $\mu\text{-OH}$) species generated from their dicopper(II) precursors have been described (Fig. 9).^{209–213} The corresponding high values of the oxidation potentials preclude chemical oxidation with a usual oxidant.²¹⁴ To date, the only report of one chemically and electrochemically achieved $\text{Cu}^{\text{II}}\text{Cu}^{\text{III}}$ species was published by Tolman²⁰⁹ (Fig. 9A) who performed chemical oxidation of the $\text{Cu}^{\text{II}}\text{Cu}^{\text{II}}$ precursor by addition of acetylferrocenium hexafluoroantimonate, $(\text{AcFc})(\text{SbF}_6)$, at -50°C in DMF.

Although no structural report has yet been reported for such species, given their high instability, they remain of high interest since $\mu\text{-oxido}$ bridged $\text{Cu}^{\text{II}}\text{Cu}^{\text{III}}$ cores have been postulated as putative intermediates for C–H bond activation.^{215,217,218} More to this point, De Tovar recently evidenced the tunability of their electrochemical properties through the formation of intimate ion pairs with surrounding anions by non-covalent interactions, highlighting the importance of second sphere effects when designing such $[\text{Cu}_2^{\text{II}}(\mu\text{-OH})_2]^{2+}$ species.²¹⁹ Interestingly and counter-intuitively, $\text{Cu}^{\text{II}}\text{Cu}^{\text{III}}(\mu\text{-O})_x$ intermediates could be more efficient for strong C–H bond activation than their corresponding $\text{Cu}^{\text{III}}\text{Cu}^{\text{III}}(\mu\text{-O})_x$ counterparts. In 2013, Yoshizawa and Shiota postulated that HAA activation could preferably occur after protonation and reduction of an O species, leading to a mixed-valent intermediate.²¹⁷ The catalytic oxidative ability of such an intermediate was experimentally demonstrated by Belle using a naphthyridine-based spacer $[\text{Py}_4\text{Cu}^{\text{II}}\text{Cu}^{\text{II}}(\mu\text{-OH})_2]^{2+}$ complex ($\text{Py}_4 = 2,7\text{-bis}(1,1\text{-dipyridylethyl})\text{-}1,8\text{-naphthyridine}$ ligand, Chart 2 and Fig. 9, structure D). Mono-electrochemical oxidation of the complex $[\text{LCu}^{\text{II}}\text{Cu}^{\text{II}}(\mu\text{-OH})_2]^{2+}$ in acetonitrile at room temperature resulted in the transient generation of a mixed-valent $\text{LCu}^{\text{II}}\text{Cu}^{\text{III}}$ species,²¹³ that could perform catalytic HAA from strong $\text{C}_{\text{sp}^3}\text{-H}$ bonds (toluene and CH_3CN with BDFE = 87.0 and 94.2 kcal mol^{−1}, respectively) in the presence of a relatively weak base (lutidine, $\text{pK}_a = 14.16$).²¹⁶ On the other hand, Thibon-Pourret reported on an intramolecular ligand C–H bond activation using a monooxidized dicopper(II) bis($\mu\text{-OH}$) complex with the Ox_2Py_2 ligand (Chart 2) bearing a bis-oxazoline arm in place of a bis-pyridine arm on a Py_4 ligand (Fig. 9-structure D).²²⁰

Finally, in 2020, Kiebers-Emmons²¹⁵ rationalized these observations through experimental determination of thermodynamic parameters ($E_{1/2}$, pK_a , BDFEs) for tmpa-based complexes (Fig. 9, structure B). BDFEs(OH) for the HAA reactions $\text{LCu}^{\text{II}}\text{-OH-Cu}^{\text{II}}\text{L} \rightarrow \text{LCu}^{\text{II}}\text{-O-Cu}^{\text{III}}\text{L}$ and $\text{LCu}^{\text{II}}\text{-OH-Cu}^{\text{III}}\text{L} \rightarrow \text{LCu}^{\text{III}}\text{-O-Cu}^{\text{III}}\text{L}$ of 103.4 and 91.7 kcal mol^{−1} (Fig. 9), respectively indicated that the higher oxidation $\text{Cu}^{\text{III}}\text{-O-Cu}^{\text{III}}$ state had a lower HAA ability than the mixed valent $\text{Cu}^{\text{II}}\text{-O-Cu}^{\text{III}}$ state. It was suggested that geometry (specifically the Cu–O–Cu angle), solvation, magnetic, and mixed valency contributions in dicopper complexes participate in the tuning of the BDFE(OH) contrarily to mononuclear species. Altogether, the results demonstrated that such high-valent dicopper(II,III) species could be key reactive intermediates in copper-based oxidative processes.



Fig. 10 Synthesis of $[\text{R}_2\text{Cu}_2(\mu\text{-O})(\mu\text{-NO})]^{2+}$ species and influence of the spin state on HAA reactivity from ref. 73 and 74.

4.3.3 $\text{Cu}_2^{\text{II,III}}$ mixed valent $\mu\text{-oxido}$ $\mu\text{-nitrosyl}$ species: C–H activation coupled to denitrification. In 2019, Zhang reported the first structurally characterized $[\text{R}_2\text{Cu}_2(\mu\text{-O})(\mu\text{-NO})]^{2+}$ complex (R_2L with $\text{R}=\text{H}$, Chart 2) prepared from $\text{R}_2\text{LCu}^{\text{I}}\text{Cu}^{\text{I}}$ and an excess of NO (Fig. 10),²²¹ or alternatively from $\text{R}_2\text{LCu}^{\text{I}}\text{Cu}^{\text{II}}$ and NO_2 . This complex was able to perform oxygen atom transfer (OAT), HAA, and C–H hydroxylation of various substrates such as phosphines, cyclohexadienes, and isochroman, yielding phosphine oxide, benzene, and 1-isochromanone, respectively. The oxidizing entity displayed a spin-switching behavior between $S = 1/2$ and $S = 3/2$ which was shown to be temperature-dependent. Modifications on the ligand (R groups introduced in position 4 of the catechol spacer (Chart 2)) induced an unsymmetrical environment for the dicopper center $[\text{R}_2\text{Cu}_2(\mu\text{-O})(\mu\text{-NO})]^{2+}$ and allowed spin state control of the intermediate.²²² The $S = 1/2$ state appeared more reactive for HAA than the $S = 3/2$ state. Best activities were obtained with $\text{R}=\text{F}$ and NH_2 , despite possessing opposing electron-donating/withdrawing properties.

4.4 Dicopper-hydroperoxido as active intermediates

Catalytic hydroxylation of benzene to phenol considering $\text{Cu}^{\text{II}}(\mu\text{-OH})\text{Cu}^{\text{II}}$ species and H_2O_2 or Cu_2^{I} and O_2 has been mechanistically investigated by Kodera²²³ and very recently by Glaser²²⁴ starting from dinuclear species $[\text{Cu}_2^{\text{II}}(\mu\text{-OH})(6\text{-hpa})]^{3+}$ or $[\text{Cu}_2^{\text{II}}(\mu\text{-OH})(\text{susan})]^{3+}$ respectively (6-hpa and susan ligands are depicted in Chart 2). In both cases, remarkably high activities compared to phenol oxidation in the *ortho* position (tyrosinase like activity) were observed (see Section 4.2.1). The active catalytic species were examined and both a Fenton-like radical mechanism or dicopper-hydroperoxido based mechanism were shown to be feasible depending of the catalytic conditions (temperature, excess amount of H_2O_2 ...). Subsequently, Kodera reported that the same dinuclear $[\text{Cu}_2^{\text{II}}(\mu\text{-OH})(6\text{-hpa})]^{3+}$ species placed in the presence of H_2O_2 was able to oxidize methane (using a high-pressure reactor) to methanol (and formaldehyde as the minor product) with TON of 43 and 7.4 respectively.²²⁵ The copper-oxyl and peroxy radicals species

detected by cryo-ESI MS were proposed to be the active species for benzene oxidation.²²³

5 Polynuclear copper–oxygen species

5.1 Trinuclear copper–oxygen adducts and C–H bond activation

So far, very few trinuclear copper complexes have been designed to activate dioxygen and perform oxidative reaction of hydrogenated substrates. Amongst reported works, Chan and co-workers have been particularly prone to develop synthetic tricopper models of the D active site of particulate methane monooxygenases (pMMOs) (see Section 2.1). Over several studies, they designed tricopper diazepane-based complexes capable of mediating methane oxidation in the presence of hydrogen peroxide in acetonitrile, but with a low efficiency under homogeneous conditions. According to DFT calculations²²⁶ and EXAFS experiments,²²⁷ the active species was proposed to be a mixed-valent adduct $[\text{Cu}^{\text{II}}\text{Cu}^{\text{II}}(\mu\text{-O})_2\text{Cu}^{\text{III}}(\text{L})]^{1+}$ (where L denotes a trinucleating ligand, for example L = 7-*N*-Etpyz, Chart 2) generated from the reaction of tris-copper(i) complexes with O₂ (see Scheme 13). The presence of a third copper center was suggested to increase the reactivity of the putative intermediate towards methane through a combination of different effects such as (i) a favored planar conformation and lengthening of the Cu^{II}–Cu^{II} distance, (ii) an increased Lewis

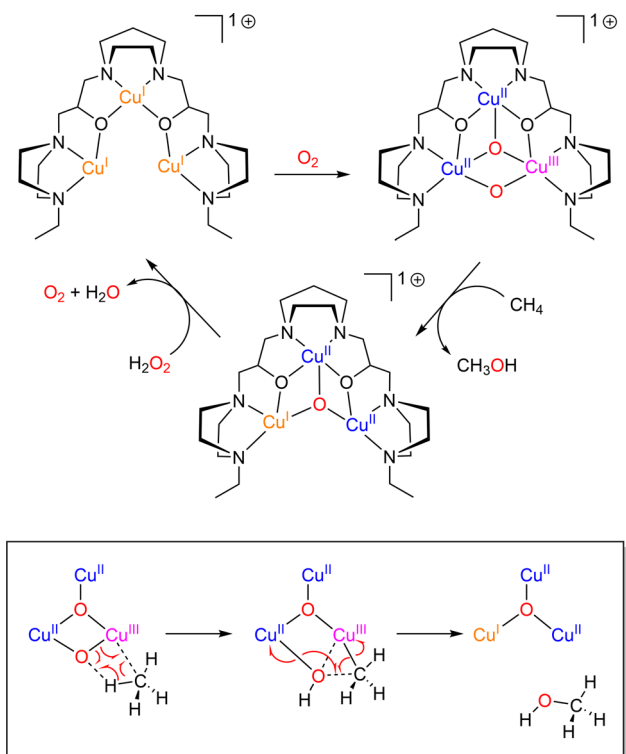
acidity and (iii) an overall singlet state of the “oxene” cluster. Interestingly, recent work by Nandi *et al.* has investigated the role of acetonitrile in the methane oxidation process mediated by the tricopper piperazine complexes from the group of Chan.²²⁸ In particular, they showed that methanol could be produced from acetonitrile in the absence of methane, but in the presence of a copper complex and hydrogen peroxide.

Significant progress in methane-to-methanol conversion was more recently obtained by Chan's group by carrying out catalysis under heterogeneous conditions.²²⁹ Experimental results demonstrated that grafting of the copper catalysts onto mesoporous silica nanoparticles significantly improved methane oxidation relative to bulk studies. One main reason for this enhancement was proposed to originate from the over-solubility of methane in the confined spaces of nanoporous materials, hence increasing the turnover frequency by raising methane concentration and avoiding the competitive direct oxidation of hydrogen peroxide by the active copper–oxygen species.

In a further paper, Chan reported the catalytic oxidation of ethane and propane using the same strategy as that used with methane.²³⁰ Indeed, the $[\text{Cu}_3^{\text{II}}(7\text{-}N\text{-Etpyz})]^{4+}$ complex (see Scheme 13) was immobilized on anionic-based mesoporous silica nanoparticles (TP-MSN, 70 nm size, 2 nm pore size) and *in situ* reduced into a tris-copper(i) species by sodium ascorbate at room temperature. In the presence of O₂ and H₂O₂, the grafted complex mediated the oxidation of ethane into ethanol with a 7-fold higher efficiency (21% conversion) compared to the same reaction under homogeneous conditions. Oxidation of propane led to the formation of both isopropanol and acetone. Notably, kinetic studies for a mixture of gases demonstrated that the catalyst was more selective for propane than for ethane or methane conversions into alcohols (30:22:1, distribution, respectively).

Catalytic methane oxidation by the tricopper-(7-*N*-Etpyz) complex was also investigated by DFT calculations.²³¹ Two mechanisms were considered for CH₄ oxidation by the reactive $[\text{Cu}_3(\mu\text{-O})_2(7\text{-}N\text{-Etpyz})]^{1+}$ complex: (i) a concerted electrophilic oxene insertion and (ii) a nucleophilic hydrogen-atom abstraction (HAA) followed by a radical rebound process. Lower activation energies were obtained for the latter, hence suggesting the occurrence of a two-step mechanism. Calculations revealed that C–O recombination was the rate-determining step in the radical rebound mechanism, hence predicting a low H/D Kinetic Isotopic Effect (KIE).

In 2020, Chan investigated the activation of the tricopper(i) complex $[\text{Cu}^{\text{I}}\text{Cu}^{\text{I}}\text{Cu}^{\text{I}}(7\text{-Dipy})]^{1+}$ (for ligand 7-Dipy see Chart 2) by H₂O₂ by using rapid-freeze-quench and cryo-ESI MS.²³² The complex was activated by H₂O₂ rather than O₂ to offer better chances in terms of kinetics to observe and characterize copper–oxygen adducts. Through an optimized methodology, the targeted $[\text{Cu}^{\text{II}}\text{Cu}^{\text{II}}(\mu\text{-O})_2\text{Cu}^{\text{III}}(7\text{-Dipy})]^{1+}$ species was detected together with other copper oxygen intermediates. A deactivation pathway involving this reactive species and a tris-Cu(i) bis-protonated complex was proposed to explain the detection of $[\text{Cu}^{\text{II}}\text{Cu}^{\text{III}}(\mu\text{-O})_2\text{Cu}^{\text{III}}(7\text{-Dipy})]^{2+}$ by high resolution mass spectrometry (Scheme 14).



Scheme 13 Top: Catalytic cycle for the oxidation of CH₄ mediated by the $[\text{Cu}^{\text{I}}\text{Cu}^{\text{I}}\text{Cu}^{\text{I}}(7\text{-}N\text{-Etpyz})]^{1+}$ complex and O₂ in the presence of hydrogen peroxide. Bottom: Proposed intermediates for methane oxidation. Adapted from ref. 227 and 229.





Scheme 14 Deactivation pathway proposed from mass-spectrometry experiments with the tricopper (7-Dipy) complex. Adapted from ref. 232.

In a more recent paper, the same group reported the development of electrocatalytic devices to perform methane-to-methanol conversion in aqueous media.²³³ As a first step, the tricopper(II)-oxido (7-*N*-Etpyz) catalyst was immobilized in mesoporous silica nanoparticles. The modified particles were then dispersed in water and deposited onto screen-printed carbon electrodes through a drop casting method, yielding a surface concentration of $0.17 \text{ nmol cm}^{-2}$. The electrocatalytic oxidation of methane was carried out by electrochemical reduction below $-0.50 \text{ V vs. Ag/AgCl}$ in potassium perchlorate ($\text{pH} = 6.5$). Production of methanol with high turnover number ($\text{TON} = \text{approx. } 2400$ after 3 h of electrolysis at $E_{\text{app}} = -0.9 \text{ V}$) was evidenced by gas chromatography from bulk electrolysis under air. However, the catalytic device suffered from the competitive oxygen reduction reaction (ORR), as well as copper leaching over time. To overcome this, sodium ascorbate and copper(II) chloride were added to the solution, showing significant improvements for the recyclability. Comparative studies with pMMO's modified electrodes emphasized the interest of using such synthetic catalysts instead of proteins in terms of efficiency over mass of catalyst.

The group of Doerrer has recently reported the generation of tricopper $[\text{Cu}_3\text{O}_2]$ cores using various monocopper(I) complexes bearing fluorinated alkoxide ligands, the main objective being to perform efficient C–H bond activation. The utilization of

perfluoropinacolate bidentate ligands, such as pin^F , afforded the formation of bis(μ_3 -oxido) tricopper(II,II,III) adducts.²³⁴ These species were characterized by UV/Vis spectroscopy at -80°C , cryo-ESI MS, and EPR spectroscopy. Reactivity studies demonstrated that the trinuclear adduct was able to catalytically oxidize *p*-hydroquinone into benzoquinone but with a limited efficiency. It also displayed OAT (Oxygen Atom Transfer) properties towards triphenylphosphines. A further step was obtained later using a fluorinated O-donor monodentate ligand (OC_4F_9).²³⁵ UV-Vis kinetic analysis suggested the transient formation of bis- μ -oxido intermediates that rapidly converted into bis- $(\mu_3\text{-O})\text{Cu}^{\text{II}}\text{Cu}^{\text{II}}\text{Cu}^{\text{III}}$ species. The latter were characterized by resonance Raman spectroscopy (the related $[\text{Cu}_3\text{O}_2]$ core associated with stretching at 679 cm^{-1} shifted to 718 cm^{-1} after isotopic labeling with ^{18}O) and UV-Vis ($\lambda_{\text{max}} = 307, 520$ and 602 nm) spectroscopy, as well as mass spectrometry. The (OC_4F_9) copper-oxygen trimer was shown to catalytically oxidize hydroquinone. Furthermore, it was also able to stoichiometrically convert 2,4-di-*tert*-butylphenolate into 3,5-di-*tert*-butyl-1,2-catechol. Such property that was not observed with the bidentate pin^F analogue was ascribed to the better flexibility of the monodentate ligand. Aiming at reproducing multicopper oxidase activity, Murray recently reported the C–H bond activation of diverse organic substrates (toluene, dihydroanthracene...) starting from a tricopper complex bearing a cyclophanate ligand (Chart 2).²³⁶ Low-temperature UV-Vis stopped-flow experiments highlighted the reaction of the tricopper complex with O_2 , leading to the generation of trinuclear copper-oxygen adducts ($\lambda_{\text{max}} = 490$ and 630 nm) with a similar spectroscopic signature to that of Chan's complex. Computational studies suggested that oxidation of exogenous substrates may occur through a concerted HAA reaction from a reactive $(\mu\text{-oxyl})(\mu_3\text{-oxido})\text{tricopper(II)}$ intermediate, followed by radical rebound and OAT. Competitive intramolecular ligand oxidation was invoked to explain the low yield obtained for substrate oxidation ($<1\%$).

5.2. Trinuclear copper complexes: related reactivities

The four-electron reduction of O_2 to water mediated by a trinuclear copper complex has been poorly investigated in the last 5 years despite its technological significance in fuel cell applications, and its relevance to the biological process in multicopper oxidases (laccase, bilirubin oxidase).

Ray reported in 2018 a comparative study of the $4e^-$, $4H^+$ reduction of dioxygen mediated by either monocopper(I) or tricopper(I) complexes bearing identical amine-based coordination spheres.²³⁷ Low-temperature UV-Vis spectroscopic measurements, supported by X-ray absorption near-edge spectroscopy (XANES), were employed to monitor the reaction of the trinuclear complex $[\text{Cu}_3^{\text{I}}(\text{L})]^{3+}$ ($\text{L} = \text{tateb} = 1,3,5\text{-tris}(3,3\text{-imino-bis}(N,N\text{-dimethylpropylamine})\text{-}2,4,6 \text{ triethylbenzene}$, Chart 2) with O_2 in the absence of reductant and proton sources. A green species assigned to a $[\text{Cu}^{\text{I}}\text{Cu}^{\text{II}}(\text{O}_2^{2-})\text{Cu}^{\text{II}}(\text{L})]^{3+}$ intermediate was hence detected at -80°C . Increasing temperature to -60°C revealed its conversion into a brownish $[\text{Cu}^{\text{II}}\text{Cu}^{\text{II}}(\text{O}_2^{3-})\text{Cu}^{\text{II}}(\text{L})]^{2+}$ species. In the presence of both reductant and proton sources at





Fig. 11 Stepwise proton-coupled reduction of the $[\text{Cu}_3^{\text{II}}(\mu\text{-O})_3(\text{LH})]^{5+}$ complex ($\text{L} = \text{tren}_4$) studied by Zhang as a model of multicopper oxidases for the reduction of dioxygen into water. Adapted from ref. 238.

-20°C , the tricopper(I) complex produced water as the sole product. Kinetic studies revealed that O_2 binding was the rate-determining step (RDS), resulting from the high instability of the trinuclear copper-oxygen adducts which underwent a very fast PCET step. In contrast, a different mechanism was found for its mononuclear analogue, generating a bis- μ -oxido Cu_2^{III} intermediate which ran the PCET reaction in a slower manner.

A supramolecular tricopper(II) complex bearing four assembled tren units was developed by Zhang as a catalytic model of multicopper oxidases for oxygen reduction into water (Fig. 11).²³⁸ The ligand was specifically designed to provide protons in the secondary sphere of the copper ions and to prevent any disassembling of the catalyst structure through the PCET reactions. Successive reduction/protonation of $[\text{Cu}_3^{\text{II}}(\mu\text{-O})_3(\text{LH})]^{5+}$

$[\text{O}_3(\text{LH})]^{5+}$ ($\text{L} = \text{tren}_4$, Chart 2) led to the generation of a tricopper(I) $(\mu\text{-OH}_2)_2$ species (Fig. 11). The different redox and protonated states were assigned thanks to voltammetric and spectroscopic characterizations. Interestingly, these reactions were shown to occur within a narrow potential range (170 mV). This effect was ascribed to favor proton relay by the ligand scaffold, as occurring in multicopper oxidases.

In 2018, Chan *et al.* reported on the efficient electrocatalytic reduction of dioxygen into water thanks to heterogeneous tricopper catalysts.²³⁹ Four different copper derivatives were investigated to account for the ligand topology and the effect of the number of copper centers on the catalytic properties was investigated. All of them were immobilized onto screen-printed carbon electrodes through a dropcasting approach. Amongst the complexes, $[\text{Cu}_3^{\text{I}}(7\text{-N-Etppz-CH}_2\text{OH})]^+$ exhibited the best performances with a 680 mV vs. RHE overpotential value at pH = 7, a turnover frequency of 12 s^{-1} , and the production of 27% H_2O_2 . Kinetic studies demonstrated that the rate-determining step was not dioxygen binding to the catalyst but merely the O-O bond cleavage through the third electron transfer to the copper cluster. Further comparative studies between dinuclear and trinuclear analogues demonstrated the key role played by the third copper center in terms of overpotential.

6 Heterogenized copper-oxygen species

6.1 Materials framework for the immobilization of copper-oxygen species

Heterogenized copper-oxygen species typically refer to Cu-containing compounds where Cu ions/clusters are



Fig. 12 Selected zeolite-based Cu-O catalyst.





Fig. 13 Selected zeolite framework types. Adopted from ref. 240.

incorporated into a heterogeneous support or matrix. These heterogenized species play a crucial role in catalysis due to copper's ability to undergo redox reactions and participate in O₂ activation and C–H bond activation, particularly in methane oxidation. This represents a crucial process with significant environmental implications. The development of selective oxidation heterogenized systems has thus received great attention over the last few years becoming a hot approach due to its advantages arising from its straight-forward implementation into industrial devices particularly focusing on methane-to-methanol (MTM) transformation.^{240,241} Within this context, understanding the nature of the true active species in such heterogenized systems has become a target towards improved performances. In the literature, there are several heterogenized systems covering the use of zeolites as coordinating materials for Cu–O species immobilization (Fig. 12) and their further MTM conversion utilization.

Zeolites are traditionally referred to as a family of open-framework aluminosilicate-based materials exhibiting distributed micropores at molecular dimensions. From here, depending on how aluminosilicate tetrahedra are connected, a different zeolite framework type can be assigned. This classification follows a three-letter code by the International Zeolite Association. Within this context, different zeolite frameworks have been preferentially studied for MTM transformation such as beta (BEA), faujasite (FAU), mordenite (MOR), mordenite framework inverted (MFI), chabazite (CHA) and omega (MAZ) (Fig. 13) among the identified 235 distinct zeolite framework types. However, despite the efforts for the characterization of such heterogenized systems, the nature of the true active

species is still a topic of debate since it depends on many factors ranging from the catalytic conditions used for the MTM conversion, the nature of such material lattices, the copper incorporation method and the signal treatment for the identification and mechanism elucidation of active species.²⁴² Additionally, many techniques and analyses have been applied for this purpose, ranging from rR spectroscopy, isotope labelling experiments and DFT calculations and machine learning.^{243,244} The aim of this section is to present the approach dealing with the parameters mentioned above towards a better understanding covering the reusability of the zeolite-containing copper-heterogenized systems.

6.2 Heterogenization of copper–oxygen species in zeolites

Cu–O species immobilized in zeolite micropores have been shown to be efficient catalysts for the selective oxidation of methane, among other substrates. Despite the lack of exact structure characterization of these Cu–O species, different structures have been proposed to exist within the Cu-exchange zeolites (Fig. 12).

6.2.1 Incorporation method, loading and distribution of copper–oxygen species. High Cu loadings contribute to the formation of oligomeric chains within the channels for both incipient wetness impregnation and for ion-exchange Cu incorporation methods, respectively. Within this context, van Bokhoven²⁴⁵ studied MOR zeolites with different Cu loadings and found that higher Cu site density requires higher temperatures for their activation.

This phenomenon is due to the relative distance between the monomeric [CuOH]⁺ species within the zeolite lattice (Fig. 14).





Fig. 14 Suggested reaction pathways of Cu–O species in Cu-MOR and their transformations during the reaction with CH₄. (a) Pathway involving [Cu₂(μ-O)]²⁺ species. (b) Pathway involving [CuOH]⁺ species, highlighting the inactivity of monomeric Cu species exhibiting dipolar spin–spin coupling at a separation distance of 4.8 Å.

These monomeric species (Fig. 14b) facilitate the formation of molecular methanol rather than the methoxy species [Cu₂(μ-OCH₃)]⁺ formed by copper dimers.

However, if these monomeric species are at a distance of *ca.* 4.8 Å, a significant dipolar spin–spin coupling between two copper sites makes them inactive (Fig. 14b). Recently, Lercher²⁴⁶ studied Cu-MOR catalysts finding that the most active Cu site was a [Cu₂Al(μ-O₃)]²⁺ cluster (Fig. 12) being able even to form an ethyl methyl ether when replacing H₂O by ethanol in the product extraction step (see Section 6.2.2). Additionally,²⁴⁷ pre-treated MFI with large cations (K⁺, Cs⁺) tunes the formation of the Cu–O species, generating a new [Cu₂(μ-O)]²⁺ site, responsible for methanol production at low copper loadings. In general, the formation of Cu-based nanoparticles or aggregates

hinders the ability of Cu-zeolites to activate methane. However, depending on their size, location and nature, they are able to perform such activation.^{248,249}

6.2.2 Catalytic setup in copper-exchanged zeolites. A specific catalytic setup is established when Cu-exchanged zeolites are used as Cu-heterogenized catalysts for C–H activation and further oxidation. The implementation of such catalytic systems usually follows a stoichiometric, “stepwise conversion” process consisting in three independent sets of reaction conditions (Fig. 15): (i) catalyst activation and treatment, (ii) methane activation and (iii) methanol extraction. Within this context, the research community has revisited these three steps during the past few years leading to significant yields and understanding improvements.

(i) Catalyst activation and treatment. The activation temperature plays a pivotal role in both the formation and nature of the copper sites prior to methane activation. In general, the oxidation of Cu-heterogenized materials requires temperatures higher than 300 °C in order to increase the population of active species. However, depending on the intrinsic kinetics of the Cu site types and the distribution of such types within a given material, this activation temperature has an impact on the overall methanol formation as well as on the selectivity.²⁵⁰ Additionally, although O₂ as oxidizing gas has been largely used for such catalyst activation, other gases like N₂O have also been employed showing a higher proportion of [Cu₂O]²⁺ species.²⁵¹ Furthermore, both [Cu₂(*trans*-μ-O₂)]²⁺ and [Cu₂(μ-O)]²⁺ species can be generated through a [Cu₂(μ-η²:η²-O₂)]²⁺ intermediate.²⁵² Additionally, although [Cu₂(μ-O)]²⁺ species are postulated to be one type of active species under these conditions, their formation ratio can be tuned by applying a self-reduction treatment, usually performed at *T* > 500 K under He, CO or mixtures of both gases.^{252,253} However, the formation

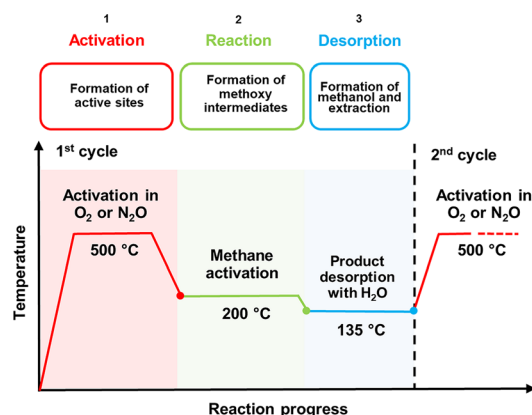


Fig. 15 Typical experimental protocol of the catalytic conversion of methane to methanol over Cu-exchanged zeolites. Adapted from ref. 246.

of Cu–O sites can be drastically affected by the presence of O₂ or water impurities.

(ii) Methane activation. After the catalyst activation, CH₄ can react with this latter in a stoichiometric way at 120 °C to 200 °C. However, the reactivity of the catalyst is sensitive to H₂O due to the intrinsic instability of the species.

(iii) Methanol extraction. Methanol extraction is usually performed by desorption of the methoxy or adsorbed methanol species with a steam flow at lower temperatures allowing the catalytic species to be regenerated.^{254,255} Recently, Cu-MOR catalysts have²⁵⁶ been found to form both CH₃ and H[•] radicals under non-thermal dielectric barrier discharge (DBD) plasma conditions facilitating methanol desorption.²⁵⁷ The authors also noticed that longer reaction times lead to the reduction of the [Cu₂(μ-O)]²⁺ species into [Cu]⁺ and [CuOH]⁺ as well as the formation of a carbon deposit. Nevertheless, the [Cu₂(μ-O)]²⁺ species can be regenerated rapidly while eliminating the carbon deposit under O₂.

6.3 Parameters affecting the catalytic activity in copper-exchanged zeolites

The reactivity of the catalytic system seems to be notably influenced by the specific type of zeolite. Despite the need to individually consider and discuss these materials, only a few factors have been identified as affecting the overall performance of the catalysts. This variation is attributed to the distinct nature of the copper sites within the zeolite framework, dependent on factors such as the Si/Al ratio, loading and distribution, pore size, lattice conformation and 2nd sphere effects.

6.3.1 Zeolite topology and lattice constraints. Zeolite topology governs the structure of active copper-oxo species, which, in turn, determines the activity of the material and the nature and distribution of the formed products.²⁵⁸ Methane overoxidation over copper-exchanged MFI takes place even at low temperature, indicating high intrinsic activity and low selectivity of the [Cu₂(μ-O)]²⁺ species, stabilized by this framework. In contrast, copper-exchanged MOR selectively converts methane into partial oxidation products such as CH₃OH but not CO₂, at temperatures below 500 K, possibly due to the low concentration of copper dimers. For Cu-BEA and Cu-FAU, reaction temperatures above 500 K are required for methanol synthesis, pointing to a lower activity for large copper-oxo clusters in comparison to dimers and monomers. Within this context, Sushkevich discovered that oxygen-activated Cu-exchange zeolites of different framework types, namely, MOR, MFI, BEA and FAU, yield different products depending on the reaction temperature. At low temperatures, preferably partial oxidation of methane occurs, resulting in the formation of different species such as adsorbed methanol and dimethyl ether. For all zeolites, the amount of such products goes through a maximum as a function of reaction temperature. However, the temperature of the maximum depends on zeolite topology and increases in the following order: Cu-MFI < Cu-MOR < Cu-BEA < Cu-FAU.

The decrease of products by partial oxidation at higher reaction temperatures is associated with their further



Fig. 16 Formation of a [Cu₃(μ₃-O)]²⁺ species and two CH₃OH molecules from the reaction of CH₄ with [Cu₃(μ-O)₃]²⁺ in MOR zeolite. Adapted from ref. 252.

transformation *via* three main pathways. First is their further oxidation by active Cu–O sites resulting in the formation of surface [CuCO]⁺ and CO₂. The second direction implies the decomposition of methoxy species to CO above 630 K. The last pathway is C–C coupling into light saturated hydrocarbons, such as ethane and propane. Additionally, the previously formed methanol can be converted into hydrocarbons because of the presence of acid sites in Cu-CHA. More to this point, the lattice of Cu-MOR allows the [Cu₃(μ-O₃)]²⁺ species to be formed as an intermediate active species able to activate and promote the oxidation of two CH₄ molecules (Fig. 16).²⁵⁹ Then, both zeolite topology and zeolite lattice constraints are found to play a significant role as they strengthen the Cu⋯Cu interaction and consequently weaken the μ–O bond destabilizing the [Cu₂(μ-O)_x]²⁺ species for CH₄ activation.²⁶⁰

6.3.2 Pore or window size. Cu sites are located within the zeolite lattice, particularly inside the pores. These pores are formed by aluminosilicate chains, which can present different Si/Al ratios (*vide supra*), and are called an X-membered ring (XMR) depending on the terminal atoms delimiting the pore entry.²⁶¹ Then, one can refer to the Cu sites on these materials by 6MR, 8MR, 10MR and 12MR (Fig. 13). In general, for CHA zeolites, Cu sites located in 6MR are the most active species but those in 8MR show an increased stability. In the former we find the [Cu₂(μ-O₂)]²⁺ species mainly with aluminate contributions whereas in the latter we find [Cu₂(μ-O₂)]²⁺ and [Cu₂(μ-O)]²⁺ species with both aluminate and silanol contributions. It is worth mentioning that, since the Si/Al ratio alters the position of aluminates within the pores (*vide supra*), the nature and location of these species within the pores lead to different activities. Due to this fact, van der Waals interactions between CH₄ and the zeolite lattice control not only the substrate adsorption but also lower the hydrogen atom abstraction (HAA) barrier.²⁶² Also, Svelle²⁴⁹ studied different Cu-exchange zeolites for ethane to ethylene transformation and observed that the pore size of the framework plays a pivotal role in the product



productivity, which increases with the pore size as follows: CHA (8MR) < MFI (10MR) < MOR (12MR).

6.3.3 Si/Al ratio, loading and distribution. Alternation of both Si and Al content in such structures has been shown to create new pores with distinct chemical and physical environments. These pores play a crucial role in MTM, providing not only insights into the distribution of copper within the lattice but also enabling better control over the formation and characterization of various Cu–O species.^{263–272} In general, three of the most active Cu–O species are $[\text{Cu}_2(\text{trans-}\mu\text{-O}_2)]^{2+}$, $[\text{Cu}_2(\mu\text{-O})]^{2+}$ and $[\text{Cu}_3(\mu\text{-O}_3)]^{2+}$,²⁵⁰ in which the copper atoms are connected to the zeolite scaffold by aluminate units.²⁷³ However, it is worth mentioning that not only dicopper and tricopper species are formed within the zeolite lattice but also monomeric species. Then, increasing the Cu/Al ratio leads to an increase of the activity of the system due to a higher density of Cu–O species and the increased probability of these species to be located at distances favorable for the formation of dinuclear and trinuclear Cu species. Furthermore, both the increase of the Cu/Al ratio and decrease of the Si/Al ratio favor the formation of polynuclear compounds such as $[\text{CuO}_x]_n$ oligomers, which present lower specific activity in MTM conversion. Within this context, Mavrikakis²⁷³ studied the effect of the Al distribution on Cu-SSZ-13 for the MTM process and found that Cu monomers, dimers and trimers can be formed depending on the different local Al distribution.

6.3.4 2nd sphere effects. As mentioned before, $[\text{Cu}_2(\mu\text{-O})]^{2+}$ species are known to be active for methane oxidation. However, depending on the lattice conformation, these species exhibit different activities.^{263,274–277} Within this context, Solomon and co-workers²⁵¹ studied both the stabilization and activities of such Cu–O species within the 8MR and 10MR for CHA and MFI zeolites, respectively unraveling that, for a given CHA zeolite, the conformation of $[\text{Cu}_2(\mu\text{-O})]^{2+}$ coordinated by two aluminates moieties drives the bridging copper oxygen atom out of the 8MR into a CHA cage, increasing both the formation of O–H bonds and methane activation. In contrast to this scenario, the positioning of aluminates within the 10MR of the MFI results in the bridging oxygen atom remaining in-plane (Fig. 17).

These phenomena where the active site is found to be located in a constricted region of the zeolite lattice are denominated as 2nd sphere effects. Here, close van der Waals contact between the substrate and the zeolite lattice in the vicinity of the active site is increased leading thus to higher activities. Recently, the authors found that the superior activity of such $[\text{Cu}_2(\mu\text{-O})]^{2+}$ species within the CHA framework is tightly related to their inner ferromagnetic character due to the π bonding ligation with their oxido bridges rather than their Cu–



Fig. 18 Proposed catalytic cycle for the $\text{CH}_4/\text{O}_2/\text{H}_2\text{O}$ reaction over Cu zeolites.

O–Cu angles as previously observed for homogeneous complexes.²⁷⁸ Within this context, when bending the Cu–O–Cu angle, each Cu^{II} nucleus binds with a different oxido p-orbital, lowering overlap and leading to orthogonal magnetic orbitals on the bridge. Also, Stepanov²⁷⁹ studied the reactivity of $[\text{Cu}_3(\mu\text{-O})_3]^{2+}$ oxido clusters on Cu-ZSM-5 after methane activation for both benzene and CO methylation finding that methoxy species at $[\text{Cu}_3(\mu\text{-O})_3]^{2+}$ oxido clusters easily interact with benzene yielding toluene.

6.3.5 Effect of temperature and water. Higher reaction temperatures not only enhance the methane conversion rate but also trigger the reactivity of Cu species.²⁸⁰ This phenomenon is attributed to higher energies required for the activation of $[\text{CuOH}]^+$ species and isolated $[\text{Cu}]^{2+}$ sites within the zeolite framework.²⁸¹ Additionally, the addition of Fe^{2+} , Zn^{2+} , La^{3+} and Mg^{2+} species favors the transformation of such monocopper species into highly active $[\text{Cu}_2(\text{trans-}\mu\text{-O}_2)]$ sites allowing even CH_4 to be oxidized to CH_3COOH .^{282,283} On the other hand, the effect of water was investigated by Chen for Cu-BEA systems²⁸⁴ unraveling that water can directly participate in the reaction through a proton transfer route over the $[\text{Cu}_2(\mu\text{-O})]^{2+}$ site, which significantly favors methanol generation and desorption and hinders carbon deposition. The effect of water was also investigated by Zhang for Cu-MOR systems,²⁸⁵ in which water can convert inactive copper species into active copper species during catalyst activation even allowing it to be recycled. Also, it is worth mentioning that Cu-zeolites show dynamic structures of Cu species with H_2O adsorption (Fig. 18).²⁸⁶ This is the case of Cu-MOR, in which Takahashi and co-workers²⁸⁷ found that H_2O adsorption is crucial for both oxidation of $[\text{CuOH}]^+$ sites and methanol production (Fig. 18). In particular, MTM reaction can only occur if methane molecules reach the active site while methanol molecules leave the material without high energetic cost for the migration. Within this context, Cu-CHA was found to exhibit a considerably higher energy barrier for methanol diffusion, which leads to the choice of higher temperature for proper methanol extraction.²⁸⁸

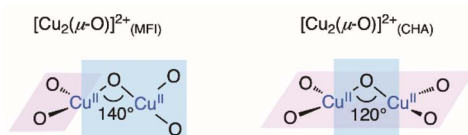


Fig. 17 $[\text{Cu}_2(\mu\text{-O})]^{2+}$ in MFI (left) and CHA (right).



Fig. 19 Incorporation of copper centers and subsequent treatment with O_2 affording Ti_8-Cu_1 and Ti_8-Cu_2 .

6.4 Other supports

6.4.1 Metal-organic frameworks (MOFs). MOFs can serve as a scaffold for immobilizing copper complexes by incorporating ligands post synthesis. Then, one can modulate the porous structure when designing MOFs to reach an enormous specific area with excellent permeability. This facilitates an enriched substrate adsorption and enhances the reactant mass transfer allowing the transformation of CH_4 to CH_3COOH .^{289–291} Additionally, Lin and co-workers²⁹² developed Ti-based MOFs (MIL-125) allowing the incorporation of $[Cu_2(\mu-O)]^{2+}$ and $[Cu_2(\mu-OH)_2]$ species leading to Ti_8-Cu_1 and Ti_8-Cu_2 catalysts, respectively (Fig. 19). Such systems have been demonstrated to exhibit a wide range of mono-oxygenation processes such as alkene epoxidation, alkane oxidation, Baeyer-Villiger and thioether oxidation activities for different substrates. Within this context, the utilization of a co-reductant (isobutyraldehyde) not only allows recycling the catalysts but also greatly accelerates the regeneration of the Cu/O species. However, a high concentration of such co-reductant competes with substrate oxidation thus decreasing the yield of monooxygenation products.

6.4.2 Al_2O_3 . Recently, Copéret²⁹³ reported the formation of well-dispersed monomeric $[Cu]^{2+}$ species supported on alumina using surface organometallic chemistry and their reactivity towards the selective and stepwise conversion of methane to methanol unraveling the active sites having a tri-coordinated $[(Al_2(\mu-O)(\mu-OH)(\mu-O)Cu)]^-$ geometry (Fig. 20).

6.4.3 Graphitic carbon nitride ($g-C_3N_4$). Zuo²⁹⁴ designed a photo-active catalyst based on Cu-MOR/ $g-C_3N_4$ able to operate under visible light. The authors found that the photo-assisted catalysis was ca. 1.85 times higher than that under thermal conditions because part of O_2 is reduced to $O_2^{\cdot-}$ by $g-C_3N_4$ under visible light accelerating the formation rate of Cu_xO_y species using $O_2^{\cdot-}$ as oxidant rather than O_2 . However, a large



Fig. 20 Strategy used to generate monomeric $[Cu]^{2+}$ sites in a pure alumina environment and the subsequent reaction with CH_4 to form hydroxyl and methoxy species.

excess of $g-C_3N_4$ blocks the pores of MOR hardening the diffusion of methane in the channel of MOR.

6.5 New strategies

6.5.1 Improvement of gas separation. Thanks to the tunable porosity when building MOFs together with the demonstrated Cu–O adduct–hydrocarbon interaction, the design of devices able to separate carbon-based gas mixtures through their designed pores is emerging as a new field of interest. Recently, He and co-workers²⁹⁵ developed a porous dicopper paddlewheel-based MOF using the pyridine-3,5-dicarboxylic acid *N*-oxide ligand resulting in a structure with two cages enriched with oxygen donor atoms. Interestingly, these O-enriched cages are able to selectively interact with C_2H_2 being able to separate the latter from $C_2H_2/CO_2/CH_4$ mixtures. Within the same context, Song²⁹⁶ studied the Cu–NaY systems for ethylene separation and found that the Cu/Al ratio dictates their adsorption ability through non-covalent interactions.

6.5.2 Overcoming the overoxidation of methane. Recently, Tang and co-workers developed a Cu-doped WO_3 photocatalyst able to overcome the overoxidation of CH_4 giving a selective production of $HCHO$.²⁹⁷ WO_3 photogenerates electrons under visible-light irradiation at room temperature, while atomically dispersed Cu^{II} acts as an effective electron acceptor (Fig. 21). The resulting ensemble synergistically leads to an efficient charge separation and transfer, resulting in a superior photocatalytic CH_4 conversion efficiency.

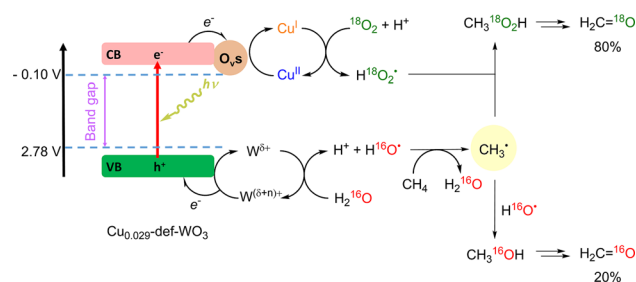


Fig. 21 Proposed mechanism for charge transfer steps during selective CH_4 oxidation over $Cu_{0.029}\text{-def-}WO_3$.

7 Concluding remarks and perspectives

Copper–oxygen species constitute a distinctive and important class of oxidation catalysts within natural systems. Although significant advancements have been achieved over the past few years in terms of fundamental and applied research still significant gaps persist in our understanding of their efficiency. This is particularly highlighted by the ongoing reconsideration of some established mechanisms in light of recent breakthroughs. The typical examples are the recent debates about the nuclearity of the reactive copper–oxygen species (mono or di) in D β H, as well as the re-evaluation of an LPMO enzyme as peroxidases rather than monooxygenases.

In the bio-inspired chemistry field, strong efforts have been particularly made during these last few years to develop synthetic copper–oxygen species able to perform efficient oxygenation of recalcitrant C–H bonds, methane being the targeted substrate.

Noteworthy, recent developments have included the exploration of copper cluster complexes incorporated within heterogeneous systems, such as zeolites. These catalysts were shown to advantageously afford the generation of highly oxidative mono- and polynuclear active sites which are hardly accessible in solution. Aside from the promising results obtained with heterogeneous systems, homogeneous copper–oxygen species still remain relevant due to valuable insights on the description of catalytic intermediates. Recent technological breakthroughs have strongly contributed to the characterization of new types of copper–oxygen transient species, such as dinuclear mixed-valent systems. In particular, *in situ* spectroscopic cryogenic techniques and synchrotron-based methods have played pivotal roles in the field in defining structure/function correlations and reaction coordinates in catalysis. Additionally, cryo-electron microscopy (cryo-EM) introduces innovative tools for elucidating the three-dimensional structure of proteins and has played a role in reopening the debate on pMMO. Altogether, these approaches have allowed easier determination of the thermodynamic properties (BDFEs) of copper–oxygen species related to their oxidative HAA power. In addition, significant progress in the field of computational studies has obviously demonstrated that these theoretical approaches have become crucial tools to improve our in-depth understanding and assist in the analysis of ambiguous experimental data.

Future work in the field of copper–oxygen species will have to consider all these new aspects/findings in order to design efficient catalysts for the activation of inert C–H bonds. The development of coupled techniques (spectroscopy, electrochemistry, spectrometry) for the *in situ* and time-resolved characterization of transient species will be of help to elucidate more deeply the different mechanisms.²⁹⁸ Alternatively, novel approaches based on photoinduced copper catalysis and radical chemistry may be applied to activate substrates under mild conditions, as recently reported.²⁹⁹

Author contributions

J. Simaan, M. Réglér, J. De Tovar, N. Le Poul and C. Belle contributed equally to writing the original draft and to reviewing and editing the manuscript. A. Thibon-Pourret, L. Wojcik, R. Leblay, Y. Wang contributed to writing, reviewing and editing the manuscript.

Conflicts of interest

There are no conflicts to declare.

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