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Unusually air-stable copper(1) complexes showing high selectivity for carbon monoxide[†]

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We report two Cu(i)-tren host molecules with unusual air-stability, as revealed by strong preference for axial CO binding over bent O₂. Spectroscopy, electrochemical, and X-ray crystal structure analyses indicate that the phenyl rotators of the capsule select for small axial ligands.

Structural features that regulate binding of diatomic molecules to metal centres are of central importance in many biochemical and chemical-industrial processes.^{1,2} As dioxygen and carbon monoxide can compete for binding at protein metal centres, probing the structural origins of selectivity between O₂ and CO at metalloproteins is of interest.³⁻⁶ Selective CO binding holds importance for sensing applications^{7,8} and also for delivery, due to its pathophysiological and therapeutic roles.9 Here we highlight copper proteins that bind dioxygen and perform vital functions such as production of norepinephrine (dopamine-βhydroxylase, DBH), mitochondrial function (cytochrome c oxidase, COX), and O2 transport in molluscs and arthropods (hemocyanin).¹⁰ These and many other copper proteins carefully regulate the Cu(I)/Cu(II) redox couple, for purposes such as electron transfer, oxidation of substrates, and O2 transport.11,12 Synthetic models for the study of cuproproteins are well-precedented13,14 and provide an opportunity to examine stereoelectronic features that control function.¹⁵ However, Cu(I) systems demonstrating CO/O2 selectivity are rare,16-18 and control of selectivity by steric factors alone has, to the best of our knowledge, never been demonstrated.

Metalloproteins can achieve ligand selectivity by varying the size and accessibility of the binding pocket.¹⁵ Many metalloproteins, including the hemocyanins (where a single dioxygen ligand coordinates two copper centres), possess deeply-buried active sites in which few guests bind.¹⁹ Inspired by this selectivity, we were interested in creating a model system with a defined metallo-cavity for ligands to enter.²⁰ However, many classes of macromolecules possessing hydrophobic cavities such as cucurbiturils,^{21–23} calixarenes,^{24–27} resorcinarenes,²⁸ and

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cyclodextrins^{29,30} generally lack defined metal-binding sites. Hemicryptophanes are a class of small-molecule cages^{31,32} that are excellent candidates for metalloprotein-inspired binding pockets, owing to their well-defined hydrophobic cavity created by a cyclotriveratrylene (CTV) north pole³³⁻³⁶ and the ease of attaching a south pole with a metal-binding site.^{31,37,38} Furthermore, hemicryptophanes and other CTV-based supramolecular structures have been used in copper-binding systems by the groups of Hardie and Dutasta.^{39–41} In designing the south pole, we were inspired by myoglobin and haemoglobin. While CO/O_2 discrimination is an important feature of these ironhaem proteins, which has been mimicked in various smallmolecule and protein models,^{42–45} such discrimination has not yet been observed in an analogous monocopper protein.

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First synthesised by Ogawa,⁴⁶ L1 (Fig. 1) is a particularly versatile hemicryptophane. In addition to selective guest⁴⁷ and anion⁴⁸ binding with protonated L1, Zn(II)-coordinated L1 catalyses the hydrolysis of activated alkyl carbonates^{46,49} and Ru(III)-L1 can catalyse the oxidation of primary alcohols.⁵⁰ Here, we sought to prepare Cu(I)-bound L1 and investigate its ligand-binding properties (Fig. 1). L1 provides an unusual example of a tripodal amine-substituted copper ligand. Many other tripodal amine- and amide-based Cu(I) systems are known, which use ligands such as tren, TMPA, and their derivatives⁵¹⁻⁵⁵ to explore the effects of alkyl groups, chelate ring size, and steric bulk on substrate binding.⁵⁶ However, with rare exceptions,⁵⁷⁻⁶⁰ most



Fig. 1 Structures of L1, L2, [CuL1]PF₆, and [CuL2]PF₆.

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notably from the Reinaud group,^{61–65} these complexes lack a well-defined cavity at the metal site.

In this work, we synthesised the copper(1) complex [CuL1]PF₆ from the previously-reported L1.^{46,47} We observed air-stability and sought to elucidate the origin of this effect, as few examples of air-stable, 4-coordinate Cu(1) complexes with an open coordination site exist due to their typically high oxidation potential.^{66–70} In similar complexes, portal size and/or steric blockage have been observed to control the size of the guests that can access the cavity.⁴⁷ Seeking to increase the portal size, we synthesised L2 (Fig. 1), a tren-hemicryptophane identical to the L1 ligand, but lacking the methoxy groups on the north pole (see ESI[†] for synthetic details). The resulting copper complex [CuL2]PF₆ maintained an unusual degree of air-stability, despite its more accessible cavity.

Interestingly, no signs of oxidation in the solid state were observed for either [CuL1]PF6 or [CuL2]PF6 upon storage as the solid powder under air for several months (Fig. S2[†]). To assess stability in the solution phase, a solution of each Cu(1) compound was prepared in DMSO-d₆ under air. Both solutions were monitored by ¹H NMR spectroscopy for two weeks, at the end of which the solutions were bubbled with air for 1 h (Fig. S3 and S4[†]). In the case of [CuL1]PF₆, no meaningful changes to the spectrum were observed after two weeks of air exposure (Fig. S3b[†]), nor was bubbling air observed to induce decomposition (Fig. S3c[†]). For [CuL2]PF₆, slight decomposition was observable after air exposure for two weeks (Fig. S4b[†]). Subsequent bubbling with air did not appear to cause significant additional decomposition (Fig. S4c†). However, repetition of these experiments with the internal standard revealed a 26% loss of compound over 14 days (Fig. S5 and S6[†]). This is presumably due to the formation of a paramagnetically broadened $Cu(\pi)$ species. While susceptible to oxidation in solution over several weeks, the compound demonstrates an unusual lack of air-sensitivity compared to other N₃Cu(1) complexes, which generally undergo oxidation on the time scale of seconds to minutes.71,72 Both Cu(I) complexes are unusually air-stable in the solution phase, but [CuL1]PF₆ appears to be marginally more stable than [CuL2]PF₆. From this, it appears that the size of the portals is important, but cannot fully explain the diminished reactivity towards dioxygen in [CuL1]PF₆.

We hypothesised that increased air stability results from a cavity that is too constrained to easily accommodate $Cu(I)-O_2$ coordination. Binding to carbon monoxide—a ligand of similar size to dioxygen—was investigated for [CuL1]PF₆ and [CuL2]PF₆



Fig. 2 IR spectra of (a) $[CuL1]PF_6$ under N_2 (grey) and CO (red) and (b) $[CuL2]PF_6$ under N_2 (grey) and CO (red).

using infrared (IR) spectroscopy in solution (Fig. 2). When purged with CO, in addition to the free CO peak at 2137 cm^{-1} , a peak was observed at 2079 cm⁻¹ for both [CuL1]PF₆ and [CuL2]PF₆, which we attribute to the formation of the Cu(1)-CO complex. Control experiments using fresh samples under dinitrogen showed neither IR peak. The 2079 cm⁻¹ stretching frequency for the bound peak is not differentiable between [CuL1]PF₆ and [CuL2]PF₆, which is expected given the similar Cu(1)-tren ligand electronic environment. These vibrational frequencies are in line with other N3CuCO complexes reported in the literature, which possess a $\nu_{\rm CO}$ of 2040–2100 cm⁻¹.^{73,74} The experiment was repeated using ¹³C-labeled CO, demonstrating the expected isotope shift (Fig. S7[†]). Binding studies were continued by comparing ¹H NMR spectra for both compounds under air, after degassing through freeze-pumpthaw cycles, and upon pressurization with CO to 20 psi (Fig. 3). The proton NMR spectra of both compounds showed changes indicative of binding behaviour in the presence of CO, eliciting peak broadening and changes in chemical shift. The degassed and under-air spectra show no differences. As expected, the most-affected NMR peaks belong to the south pole and linker regions of the molecule, nearest the copper centre. In order to remove CO and restore the ligand-free NMR spectrum for both compounds, it was necessary to degas rigorously using the



Fig. 3 (a) ¹H NMR spectra (500 MHz, CD₃CN) of [CuL1]PF₆ under air (top), after degassing (middle) and under a 20 psi pressure of carbon monoxide (bottom). (b) ¹H NMR spectra (500 MHz, CD₃CN) of [CuL2] PF₆ under air (top), after degassing (middle) and under a 20 psi pressure of carbon monoxide (bottom).

freeze-pump-thaw method. The changes to both the ¹H and ¹³C NMR spectra (Fig. S8–S14[†]) of **[CuL1]PF**₆ and **[CuL2]PF**₆ can be attributed to the presence of the CO as the guest.^{75,76} Notably, however, no signal corresponding to free or bound carbon monoxide was observed in the ¹³C NMR spectrum for either compound. Based on the evidence for a significant Cu(i)–CO bound population, we inferred that the CO off-rate is in an intermediate exchange regime resulting in extreme broadening of the CO-bound peak (Fig. S10 and S11[†]).⁷⁷ This hypothesis was confirmed by applying ¹³C-labeled CO to the sample, upon which one broad ¹³C NMR peak corresponding to bound and free CO was observed (Fig. S12 and S13[†]).

To examine carbon monoxide binding further, cyclic voltammetry (CV) experiments were performed in the absence and presence of carbon monoxide for [CuL1]PF₆ and [CuL2]PF₆ (Fig. 4). Both compounds displayed highly irreversible electrontransfer kinetics in ligand-free and ligand-bound states. We propose that the irreversible nature of the CVs is attributable to several factors, including the rigidity of the tren ligand within the capsule. In similar copper complexes, ligand rearrangement is necessary to facilitate redox changes at the metal.78,79 Considerable evidence supports the role of a "rack" or entatic state in copper electron-transfer proteins, in which small movements in one or more axial ligands assist the complex in shifting between preferred coordination geometries, i.e., tetrahedral Cu(I) and square planar/higher-coordinate Cu(II).11,80 Our system may demonstrate the opposite: the movement of ligands is constrained by the macrocycle, hindering electron transfer. This phenomenon has been seen in similar TREN-based systems displaying irreversible electron-transfer kinetics, in which the constrained ligand enforces a particular coordination geometry upon the metal centre, thus favouring the corresponding oxidation state.58 The very sluggish ET kinetics observed for these Cu model systems helps to explain their notable Cu(I) air-stability. We note that the extremely slow ET kinetics observed for both Cu(1)-CO species (Fig. 4) may be attributable to the high inner-sphere reorganisation energy associated with rapid CO ligand exchange.81

Titration of CO revealed a correlation between the CO concentration and reduction potential of both compounds. As



Fig. 4 Cyclic voltammograms for (a) [CuL1]PF₆ and (b) [CuL2]PF₆ in the presence of varying concentrations of carbon monoxide. All experiments were performed in anhydrous DMSO with 250 mM TBAPF₆ supporting electrolyte, glassy carbon working electrode, Pt mesh counter electrode, and Ag/AgNO₃ reference electrode calibrated to Fc⁺/Fc at a scan rate of 100 mV s⁻¹. Arrows show the start point of the scan (-0.457 V vs. Fc⁺/Fc) and the scan direction.

more CO was titrated during the CV measurement, both apparent $E_{1/2}$ values shifted anodically, providing additional evidence that CO is coordinating to the metal sites in [CuL1]PF₆ and [CuL2]PF₆. Consistent with observations from ¹H NMR, purging the sample with nitrogen did not revert the CV to its original state in either case. Finally, the Cu(1) oxidation potentials measured in both [CuL1]PF₆ and [CuL2]PF₆ indicate that oxidation by dioxygen should be thermodynamically favourable. This makes the relative Cu(1) air-stability all the more striking, particularly with such prominent CO binding. The larger portals in [CuL2]PF₆ barely affect CO binding or air stability.

To shed light on the preference for CO binding and against O_2 binding, we sought to obtain various crystal structures to visualize the cavity. Crystals suitable for single crystal X-ray diffraction were obtained for [**CuL1**]**PF**₆ by vapor diffusion of ether into acetonitrile (Fig. 5a). Aware of the "induced fit" model



Fig. 5 Crystal structures of (a) $[CuL1]PF_6$, (b) $[CuL1(MeCN)](OTf)_2$ and (c) $[CuL1(N_3)](OTf)_2$ (counter ions have been removed for clarity). Mapped out cavities for (d) $[CuL1]PF_6$, (e) $[CuL1(MeCN)](OTf)_2$ and (f) $[CuL1(N_3)](OTf)_2$ overlaid with the corresponding crystal structure. Guests and counter ions have been removed for clarity.

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observed for related supramolecular complexes,⁸² we suspected that a structure of the empty capsule L1 would not accurately predict the size and shape of the cavity when a ligand is present. Therefore, we also sought to crystallise Cu(1)–ligand and Cu(1)– ligand complexes that should more closely resemble the putative Cu(1)–O₂ complex. Thus, L1 was crystallised in the presence of Cu(OTf)₂.† The resulting X-ray structure contained a molecule of acetonitrile residing in the cavity and bound axially to the copper (Fig. 5b), similar to the expected binding for CO. Crystallisation of L1 was also attempted in the presence of Cu(OTf)₂ and *n*-Bu₄NN₃ in hopes of capturing a ligand coordinated to the Cu centre in a bent fashion, similar to the expected dioxygen coordination geometry.^{53,83,84} It was revealed that under these conditions an azide anion was bound to the copper(π) centre (Fig. 5c).

We mapped the surface of the ligand-accessible cavity for each of these crystal structures using Molovol.85 Cavity surfaces overlaid with the crystal structure (not including the ligands) are presented in Fig. 5d-f. For the empty structure, we observed the cavity to be elongated and narrow near the copper centre, perfectly suited to accommodate acetonitrile (Fig. 5b) or carbon monoxide when bound axially. In fact, the cavity's oblong shape does not permit a bent ligand to bind entirely inside the host molecule. Instead, upon binding to azide at a bent angle, the aryl linkers rotate, allowing the azide ligand to thread between them such that it is only partially contained within the host cavity (Fig. 5c and f). While making space for the ligand, the linker conformational change comes at an entropic cost as it restricts the rotation of the linkers. Azide binding suggests that such binding angles are attainable, but the loss of entropy needs to be compensated for by enthalpic gains. As such, we propose that O2 binding in [CuL1]PF6 and [CuL2]PF6 lacks sufficient enthalpy to fully compensate for the loss of entropy in these host molecules at rt.

A related phenomenon has been observed previously: in the Karlin group's seminal studies of Cu(I)–CO and $-O_2$ binding, the CO adduct formation rate constant is higher than that of the O_2 adduct,⁸⁶ which was attributed to a slower k_{Off} rate of CO for entropic reasons pertaining to ligand re-organisation. In comparison, the intermediate exchange that we observe in the ¹³CO NMR may be due to the more constrained nature of the ligand, in which a smaller entropic penalty would be paid for the re-ligation of the TREN ligand than for a pendant pyridyl ligand.⁷³

These observations carry implications for the study of cuproproteins and Cu-enzyme mimetics as a whole. Several model systems in the literature use CO as a redox-inactive ligand with similar binding behaviour to O_2 .^{15,87,88} Our model system, while unusual, highlights the possibility that some monocopper proteins may bind CO and discriminate against O_2 , as is well known to occur in haemoglobin. Arthropod and molluscan hemocyanins display differing selectivities for CO and O_2 ,⁸⁹ but this effect is presumably mediated by the cooperativity seen in anywhere from 6 to 160 dicopper sites.⁹⁰ The current study suggests the possibility of achieving discrimination in a monocopper protein based upon binding angle and cavity shape.

In conclusion, we have synthesised two unusually air-stable Cu(i) tren-hemicryptophane host molecules. We propose that our system is the first CO-binding Cu(i) complex for which O_2 binding/reaction is demonstrated to be electronically favourable, but is disfavoured due to steric factors alone. IR, NMR, and CV investigations yield strong evidence of discrimination for CO over O_2 . These findings shed insight on an alternate approach for stabilizing Cu(i) species, in which the coordination of a fifth ligand, as preferred by Cu(i), is made entropically unfavourable. This concept may be possible to adapt for small-molecule catalyst development and directed evolution of copper enzymes in the future, granting unusual ligand selectivity and reactivity.

Data availability

Our laboratory maintains all laboratory notebooks in a secure site, and maintains electronic copies of all spectra on secure servers. We have provided the most critical data in the MS and ESI,[†] and we will be happy to provide any additional information relevant to this paper upon request from reviewers or the broader public, after publication.

Author contributions

Conceptualization, Y. L., B. S., A. M. S., C. J. S., A. S. M., and I. J. D.; methodology, Y. L., B. S., A. M. S., S. A. D., and S. P. O.; investigation, Y. L., B. S., A. M. S., C. J. S., A. S. M., M. R. G., P. J. C.; formal analysis, B. S.; writing – original draft, A. M. S. and B. S.; writing – review and editing, A. M. S., B. S., C. J. S., A. S. M., and I. J. D.; supervision, I. J. D. and T. E. M.; project administration, I. J. D. and T. E. M.; funding acquisition, I. J. D. and T. E. M.

Conflicts of interest

There are no conflicts to declare.

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