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Cite this: Chem. Sci., 2024, 15, 5327

o All publication charges for this article

have been paid for by the Royal Society of Chemistry

Received 29th November 2023 Accepted 4th March 2024

DOI: 10.1039/d3sc06390a

rsc.li/chemical-science

Introduction

generally challenging to study.

Reversible dioxygen uptake at [Cu₄] clusters[†]

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Dioxygen binding solely through non-covalent interactions is rare. In living systems, dioxygen transport takes place via iron or copper-containing biological cofactors. Specifically, a reversible covalent interaction is established when O_2 binds to the mono or polynuclear metal center. However, O_2 stabilization in the absence of covalent bond formation is challenging and rarely observed. Here, we demonstrate a unique example of reversible non-covalent binding of dioxygen within the cavity of a well-defined synthetic all-Cu(I) tetracopper cluster.

> creating a unique pocket binding O2 solely through noncovalent interactions.

Our group has developed modular amine-based ligands Reversible O₂ binding is the cornerstone of cellular respiration.¹ serving as templates for metal cluster formation. Recently, we Hemoglobin, myoglobin, hemerythrin, and hemocyanin all reported a rigid ligand scaffold that enables the formation of serve as O₂ transporters across all living organisms.² In the square planar tetranuclear [Cu₄] clusters (Fig. 1).¹² Keeping process of biological uptake, transport, and delivery of dioxysome of the design principles employed before, we decided to gen, a covalent bond is established between O_2 and Fe^3 , or increase the degrees of freedom of the ligand scaffold as shown similarly between O2 and the Cu2 site in hemocyanin,⁴ resulting in Fig. 1. Others in the field have adopted similar measures in charge transfer to form a superoxo or peroxo moiety, either allowing or restricting the templating ligand's rigidity to respectively. Although, non-covalent interactions have been alter the cluster reactivity and composition. For example, Betley described to be operative in stabilizing ferric-superoxo intermediates in heme proteins,⁵ these nominally weak contacts are and coworkers demonstrated that increasing the template's rigidity exchanging the tame ligand backbone (tame = 1,1,1tris(aminomethyl)ethane),¹³⁻¹⁶ for the tris-amine α-α-α-1,3,5-Aside from biological cofactors, numerous studies have retris-aminocyclohexane,17-19 opens the door for substrate activaported supramolecular complexes, porous materials, and tion pathways at trinuclear species not available in the former organic cages capable of binding dioxygen. For instance, metal tame-based clusters. Similarly, C3-symmetric 1,3,5-benzene complexes formed within macrocyclic species, e.g., palladiumbound cyclodextrin,6 and Mn-supported calixarene,7 lead to substituted ligands developed by Holm to mimic iron sulfur



Fig. 1 Previous ligand architecture $*LH_8$ ($R^1 = n$ -pentyl; $R^2 = Me$, Ph, or i-Bu) compared to LH_8 ($R^3 = n$ -heptyl) reported herein with increased degrees of freedom.

built within a flexible supramolecular scaffold capable of

peroxo and superoxo moieties, respectively, stabilized inside the macrocycle. Similar occurrences are observed within metal organic frameworks (MOFs), where O2 binding to embedded

metal sites is used for O₂/N₂ separations,⁸ or bond activation

cleaving the O-O bond.9,10 Last, macrocycles alone are also

known to stabilize peroxo species.11 However, to our knowledge,

non-covalent interactions alone have not been described to stabilize neutral O₂ in biological metal cofactors or synthetic metal clusters. Here, we describe a polynuclear copper cluster

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[†] Electronic supplementary information (ESI) available: Experimental details, characterization, and spectroscopic data. CCDC 2332560 and 2333831. For ESI and crystallographic data in CIF or other electronic format see DOI: https://doi.org/10.1039/d3sc06390a

clusters,^{20,21} inspired the synthesis of oxygen-donor congeners in work reported by Agapie and coworkers towards the creation of Mn_3Ca subsite mimics of the oxygen evolving complex.²²⁻²⁴ Most recently, Suess *et al.* using a similar ligand base fragment developed a nitrogen-donor analogue capable of isolating an iron sulfur alkyl cluster which mimics elusive enzymatic intermediates.^{25,26} Other systems benefiting from ligand rigidification include those from the Murray group, whereupon limiting the degrees of freedom of their initial cryptand design,²⁷ they uncovered [M₃] clusters, M = Fe, Co, Cu, and Zn, capable of ligating and activating N_2 and CO_2 .²⁸⁻³⁴ In our case, by increasing the degrees of freedom of our ligand scaffold we have created a binding pocket within an all-Cu(1) [Cu₄] cluster that binds dioxygen through non-covalent interactions.

Results and discussion

Synthetic procedures

Our synthetic protocol provides square planar copper clusters in three steps from **1**, a precursor obtained readily in gram-scale quantities,³⁵ as shown in Scheme 1. First, a four-fold $S_N 2$ reaction on **1** by 4-methyl-3-nitrophenol in basic conditions using K_2CO_3 in DMF for 24 hours produces ligand precursor $L(NO_2)_4$ in 64% isolated yield. This tetranitro species is reduced under 60 psi of H_2 over Pd/C in refluxing THF for 48 h. The reaction is quantitative by ¹H NMR; however, after work up the isolated yield of LH_8 is 97%. *In situ* deprotonation and metalation of this tetraamine ligand with $Cu_4(Mes)_4(py)_2$ in THF forms the all-Cu(l) diamagnetic species LH_4Cu_4 in 63% yield. ¹H NMR analysis of LH_4Cu_4 reveals its ideal C_{4v} symmetry in solution (Fig. S11†).

Cluster topology

Single crystal X-ray diffraction data was collected on crystals grown by vapor diffusion of pentane into a concentrated solution of LH_4Cu_4 in THF. The molecular structure of LH_4Cu_4 confirmed the formation of a square planar arrangement of Cu atoms sitting at an average distance d_{avg} (Cu–Cu) of 2.69(2) Å (Fig. 2a). For comparison, in Cu metal the Cu–Cu distance in Cu(100) is 2.55(1) Å.³⁶ Additionally, the [Cu₄] core displays a d_{avg} (Cu–N) of 1.89(1) Å and an almost linear local Cu coordination environment with an average \angle N–Cu–N of 177.3(7) degrees. Similar square-shaped tetranuclear copper clusters mimicking Cu_z in N₂O reductase³⁷ have been previously reported by Mankad and coworkers using bridging diphosphine^{38,39} or formamidinate^{40–42} ligands resulting in cluster cores with Cu– Cu distances ranging from around 2.4 to 3.5 Å across all compounds reported therein. In fact, a recent example by the same group demonstrates the formation of a [Cu₄(O₂)] adduct, where O₂ is bound to a single metal site.⁴³

LH₄Cu₄ is closely related to our previously reported [Cu₄] clusters;^{12,44} however, we hypothesized that a larger internal cavity should be created in this newly synthesized cluster located in between the [Cu₄N₄] fragment and the resorcinarene backbone as a consequence of the axially longer LH₈ relative to *LH₈. Our hypothesis was confirmed upon analyzing the molecular crystal structure obtained when LH₄Cu₄ is exposed to MeCN (Fig. 2b). The MeCN molecule is hosted below the [Cu₄] plane establishing a [Cu₄] centroid-to-N_{MeCN} distance of 3.852 Å. Note that the structure metrics of LH₄Cu₄(MeCN) are relatively unchanged (d_{avg} (Cu–Cu) = 2.678(5) Å, d_{avg} (Cu–N) = 1.890(3) Å, \angle N–Cu–N = 177.4(6) degrees) from LH₄Cu₄.

Host-guest properties

Titration of MeCN to LH4Cu4 provided insight into the hydrogen atom resonances involved in the non-covalent bonding of MeCN in the host-guest adduct LH₄Cu₄(MeCN). As observed in Fig. 3a, the addition of MeCN in CDCl₃ leads to resonance shifts in the ¹H NMR. Particularly noteworthy is that protons a, b, and c, as labeled in Scheme 1, shift downfield as equivalents of MeCN are added (Fig. S15† contains the full spectrum), a well-known effect for H atoms involved in hydrogen bonding.45,46 The ¹H NMR spectrum additionally reveals the location of the MeCN signal at -1.76 ppm. Similar upfield shifts or shielding of the methyl resonance in acetonitrile has been observed in calixarene-based copper clusters.47 Moreover, the stepwise addition of MeCN to LH₄Cu₄ produces a new set of resonances corresponding to LH4Cu4(MeCN) while the location of those for LH4Cu4 remain unchanged, indicating a large host-guest association constant ($K_a > 10^5 \text{ M}^{-1}$ in CDCl₃)



Scheme 1 Synthetic pathway for ligand LH_8 and metalation conditions leading to LH_4Cu_4



Fig. 2 Molecular crystal structure of (a) LH₄Cu₄ (100 K) and (b) LH₄Cu₄(MeCN) (100 K). Sphere packing model is shown for LH₄Cu₄(MeCN) to showcase the congested nature of its internal cavity. The C, N, O, Cu, and H atoms are coloured grey, blue, red, orange, and white, respectively. R groups (*n*-heptyl) and most hydrogen atoms, except for select ones, are omitted for clarity. Thermal ellipsoids are set at 50% probability level.

that goes beyond the measurable range *via* NMR.⁴⁸ Variabletemperature ¹H NMR in C₆D₆ provided insight into the thermodynamics of MeCN dissociation from LH₄Cu₄(MeCN) (Fig. S16†). A van't Hoff analysis reveals a dissociation enthalpy and entropy of 12.7 kcal mol⁻¹ and 25.4 cal mol⁻¹ K⁻¹ (Fig. S17†), respectively. The calculated binding free energy of -5.14 kcal mol⁻¹ at 298 K is similar to solvent binding in cavitands.⁴⁹ Most importantly, structural fidelity of LH₄Cu₄(MeCN) in combination with the ¹H NMR spectroscopic signatures resulting from MeCN binding provides a roadmap to investigate binding of other guests to LH₄Cu₄.

Copper–dioxygen chemistry is cornerstone in living systems and has inspired the realization of many synthetic compounds seeking to replicate its structure and function.^{50,51} Seeking to probe potential ligation and activation modes of O_2 at LH₄Cu₄, an all-Cu₁ cluster, we dosed dry O_2 to an air-free solution of LH_4Cu_4 in CDCl₃. As the atmosphere is exchanged from N₂ (Fig. 3b bottom) to O_2 (Fig. 3b middle), we observe broadening in proton resonances a, b, c, and the methine at 4.85 ppm. Note that the same proton resonances a, b, and c, are affected during MeCN binding. These proton resonances sharpen back when the O₂ atmosphere is removed (Fig. 3b top). Altogether, our experiments indicate that O₂ is reversibly accommodated in the same binding pocket as MeCN. Note that $L(NO_2)_4$ and LH_8 do not show any broadening of resonances a, b, and c under the same conditions (Fig. S19 and S20[†]). It is important to highlight that the vast majority of synthetic molecular $[Cu_n^I]$ systems, with nuclearities ranging from n = 1 to 4, engage O₂ by establishing a copper-oxygen covalent bond giving rise to terminal or bridging peroxo, superoxo, or oxo complexes.52 To the best of our knowledge, this is the first formally all-Cu(I) cluster reversibly binding O2 solely via non-covalent interactions.



Fig. 3 (a) ¹H NMR spectra indicating resonances a, b, and c as MeCN is titrated into LH_4Cu_4 in $CDCl_3$ at room temperature under a dinitrogen atmosphere. (b) Sequence of ¹H NMR spectra of LH_4Cu_4 collected under N₂ (bottom), 1 atm of O₂ (middle), and last reverting back to N₂ (top). (c) Variable isotope resonance Raman spectra of LH_4Cu_4 plus O₂ collected at room temperature. The region expected for the O–O stretch is shown. Partial fits (in grey) of the orange and purple traces to Gaussian curves served to locate the maximum.



Non-covalent contact surface

Fig. 4 IGMH map displaying the non-covalent interactions between O_2 and LH₄Cu₄ (isovalue = 0.0011 a.u.).

Further examination of O₂ binding to LH₄Cu₄ was carried out by collecting resonance Raman vibrational spectra on dried thin films of LH₄Cu₄ contained within a quartz cuvette either under vacuum or dry dioxygen atmosphere. Laser excitation at 532 nm gave a band centered at 1584 cm⁻¹ assigned to the O-O stretching of dioxygen hosted within LH₄Cu₄ (Fig. 3c). Note that the symmetrical stretching band in free O2 appears at 1556.4 cm⁻¹ (Fig. S22⁺).⁵³ The same set of experiments were executed using isotopically labeled ¹⁸O₂ displaying a stretching band at 1556 cm⁻¹. The simple harmonic oscillator model predicts a shift of ~ 90 cm⁻¹, however we only observe a $\Delta^{18}O$ of 28 cm⁻¹.⁵² We hypothesize that dioxygen enters the cavity of LH₄Cu₄ likely through partial ligand dissociation or via the formation of a dilated aperture.54,55 Altogether, it appears that the non-covalent interactions serve to stabilize the O2 molecule within LH₄Cu₄ and at the same time strengthens the O-O bond by removing π^* electron density.

To further investigate the electronic properties of LH₄Cu₄ and its adduct with O2, we employed density functional theory (DFT) methods. $LH_4Cu'_4(R^3 = Me)$ was optimized at B3LYP-D3BJ/Def2-SVP+PCM(CHCl₃) level of theory. The optimized structure reproduced the experimental data with high accuracy as determined by overlaying these structures and obtaining a root mean square displacement (RMSD) of 0.3 Å and d_{avg} (Cu-Cu) of 2.67 Å (Fig. S23[†]), indicating the reliability of the selected method.

Intrigued by the non-covalent binding of O2 to LH4Cu4, the adduct $LH_4Cu'_4(O_2)$ was first optimized employing B3LYP-D3BJ/ Def2-SVP+PCM(CHCl₃) level of theory. The binding pocket within $LH_4Cu'_4$ is best visualized by applying the independent gradient model based on Hirshfeld partition of molecular density (IGMH),56 which clearly reveals the contact surface of weak non-covalent interactions involved in hosting the O₂ molecule. Note that the oxygen molecule positions itself in a way to maximize interactions with the C-H bonds from the bridging methylenes and top aromatic rings, protons a and c, respectively, in Scheme 1. The O2

molecule was placed at different starting positions during structure optimization, e.g., close to the resorcin[4]arene base, near the Cu₄ plane, and in all cases the optimum location found is that shown in Fig. 4. Overall, the IGMH isosurface for the O2 adduct is well supported experimentally as reflected in the ¹H NMR data shown in Fig. 3b.

Conclusions

Here, we designed a novel ligand architecture carrying four aniline moieties to obtain a rigidified square-planar $[Cu_4]$ topology despite the ligand's rotational degrees of freedom. The foregoing results show that, changing the resorcinarene backbone employed previously by our group engenders a cluster compound with a cavity to encapsulate small molecules, where we showcase a unique example of reversible non-covalent binding of O₂ within the [Cu₄] cluster built around a supramolecular scaffold.

Data availability

All data including experimental and analytical details are in the ESI.†

Author contributions

Manasseh K. Osei: conceptualization, investigation, data curation, writing - review & edition. Saber Mirzaei: conceptualization, investigation, data curation, formal analysis, writing review & edition. M. Saeed Mirzaei: investigation, data curation, formal analysis. Agustin Valles: investigation. Raúl Hernández Sánchez: conceptualization, project administration, resources, funding acquisition, supervision, visualization, writing - original draft, writing - review & edition.

Conflicts of interest

There are no conflicts to declare.

Acknowledgements

This research was supported by Rice University and the Robert A. Welch Foundation Young Investigator Award. Acknowledgment is made to the donors of the American Chemical Society Petroleum Research Fund for partial support of this research. The authors acknowledge the use of Raman microscope and NMR spectrometers through the Shared Equipment Authority at Rice University. Also, we thank startup funding and the Center for Research Computing from the University of Pittsburgh for their support. We thank Ying Chen for helpful discussions. We thank Dr Xu Wang for providing help with VT ¹H NMR experiments.

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