Self-Assembled Anion-Binding Cryptand for the Selective Liquid–Liquid Extraction of Phosphate Anions

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Abstract: The ligands $L^1$ and $L^2$ form trinuclear self-assembled complexes with Cu$^{2+}$ (i.e. [(L$^1$)$_2$Cu$_3$]$^{6+}$ or [(L$^2$)$_2$Cu$_3$]$^{6+}$) both of which act as a host to a variety of anions. Inclusion of long aliphatic chains on these ligands allows the assemblies to extract anions from aqueous media into organic solvents. Phosphate can be removed from water efficiently and highly selectively, even in the presence of other anions.

Modern agriculture is totally reliant on phosphate for the mass production of foodstuffs and annual demand for phosphates is growing twice as fast as the growth in human population.[1] The “peak phosphate problem” concerns the dependency on phosphate fertilizer which, unlike nitrogen based fertilizer, is produced via a finite supply located in only a few countries. It is believed that supply will outstrip demand in 20 to 30 years with depletion of reserves in the next 50 to 100 years.[2] Correspondingly, green and sustainable methodologies for phosphate use need to be developed to avoid a decrease in food production.[3] Conversely, inefficient use of phosphate results in 50 Tg yr$^{-1}$ of this fertilizer entering the environment, and oceans, causing toxic algal blooms and eutrophication.[4]

Despite their obvious medical, environmental and agricultural significance, the ability to detect and sequester anions has significantly lagged behind recognition of their cationic counterparts.[5] As a result much effort has been focused upon anion recognition in supramolecular chemistry, and the ability to synthesize receptors capable of binding anions has become increasingly more understood.[6] Anion receptors are generally organic scaffolds that contain functional groups capable of interacting with anions and often contain amine, amide and alcohol groups along with other hydrogen-bond donor units. Self-assembly is an attractive alternative to covalent synthesis for these scaffolds as it allows the construction of structurally complex architectures from relatively simple subunits, and self-assembled hosts for anion binding have attracted a lot of recent attention.[7,8]

Recently we have shown that the tripodal ligand $L^1$ (Figure 1) self-assembles with Cu$^{2+}$ ions to form the trinuclear species [(L$^1$)$_2$Cu$_3$]$^{6+}$. This complex contains a cavity which incorporates six -NH donor atoms and three Cu$^{2+}$ metal ions in an arrangement that allows all of them to interact with anions. It was demonstrated that anions are encapsulated both in the solid-state and aqueous systems and, upon encapsulation, spherical, trigonal planar, tetrahedral and octahedral anions are all precipitated from solution and can be removed by filtration. Furthermore, the cavity is selective for phosphate anions and precipitates these from water, reducing the concentration from 1000 to $<0.1$ ppm and recovering $\approx 99\%$ of the phosphate anion.[9]

Whilst this is a promising method for phosphate recovery, anion extraction by precipitation is a batch-based process and if removal/recovery of phosphate is to be practical, a method of anion sequestration using liquid-liquid extraction is essential, as this can be more readily developed into a continuous process. Whilst this would be an attractive industrial process, it is complicated by the hydrophilicity of the phosphate anion which makes it difficult to extract into organic solvents, especially in comparison to the less hydro-
philic anions (e.g. halides, nitrate etc).

As a result, the ability to selectively extract phosphate into organic solvents that partition with aqueous media is challenging but important, if resource recovery is to be an achievable goal.

In this work we show that the organic backbone of \( L^1 \) can be changed (cf. the cyclohexyl linked ligand \( L^2 \)) which gives a modified ligand that still forms a trinuclear species \([\langle L^2 \rangle_3 \text{Cu}^3\]^+) which acts as a host to anionic guests in a variety of media (Figure 1). Furthermore, ligands \( L^1a \) and \( L^2a \) are based on their parent species (\( L^1 \) and \( L^2 \)) but contain aliphatic hexyl esters which allow the formation of the tripodal complexes (e.g. \([ L_2 \text{Cu}_4 \text{SO}_4 ] \)) in organic solvents. Both of these trinuclear assemblies extract phosphate anions from water into dichloromethane and are highly selective for Na\( _2 \)HPO\(_4 \) in the presence of other common anions.

Ligand \( L^2 \) was prepared from cis, cis-1,3,5-triaminocyclohexane by reaction with benzoyl isothiocyanate, hydrolysis to the trithiourea and reaction with 2-(\( \alpha \)-bromoacetyl)pypyridine. Reaction of this tripodal ligand with Cu(ClO\(_4 \)_2 in MeNO\(_2 \) gave a pale blue solution which turned yellow upon reaction with Bu\(_4\)NH\(_2\)SO\(_4\) and deposited yellow crystals upon slow diffusion of diethyl ether. In the solid-state each of the three copper ions coordinates to a different copper metal ion, and each metal ion is coordinated by two separate ligands resulting in a host by a total of nine interactions, comprising three Cu···O NH···O hydrogen-bond interactions are substantially shorter for \([ L^2 \text{Cu}_3 \text{SO}_4 ] \]^+ (−NH···O distance 1.996–2.145 Å for \([ L^2 \text{Cu}_3 \text{SO}_4 ] \]^+ vs. 2.255–2.286 Å for \([ L^1 \text{Cu}_3 \text{SO}_4 ] \]^+). This reduction in distances is a consequence of the spacer unit, as the 1,3,5-cyclohexyl spacer of \( L^2 \) is dimensionally smaller than the trimethylamine unit of \( L^1 \), with the amine units separated by 4 bonds in \( L^2 \) as opposed to 6 bonds in \( L^1 \). The host-guest complex is also observed in the gas phase with ions in the ESI-MS at \( m/z \) 1952 and 902 corresponding to \([ [ L^2 \text{Cu}_3 \text{SO}_4 ] \text{(OTf)}_2 ]^+ \) and \([ [ L^2 \text{Cu}_3 \text{SO}_4 ] \text{(OTf)}_2 ]^+ \) respectively.

Reaction of \([ L^2 \text{Cu}^3\]^+ with Bu\(_4\)NBr in water and acetone (1:3) gave a blue solution from which blue block shaped crystals were formed upon slow evaporation. In the solid-state the trinuclear species persists and encapsulated within this core is a bromide ion, that is, the complex is \([ [ L^2 \text{Cu}_3 \text{Br} ]^+ \). As in the previous example of the sulfate complex, the halide anion interacts with all three of the Cu\(^{2+} \) ions and is supplemented by six −NH···Br hydrogen-bonding interactions (see ESI).

As can be seen in the two solid-state structures the cyclohexyl-based ligand \( L^2 \) behaves in a similar fashion to \( L^1 \); viz., it forms the trinmetallic self-assembly and incorporates anions within the cryptand. Whilst we don't have solid-state evidence for encapsulation of anions, other than sulfate and bromide, it seems highly likely that other tetrahedral oxoanions and halides would be bound in the cavities of complexes with \( L^2 \) in an analogous fashion, as has been extensively observed in \( L^1 \).

In an effort to produce compounds that could abstract anions from aqueous media into an organic solvent, ligands \( L^1a \) and \( L^2a \) were prepared, each containing three -CH\(_2\)O-CH\(_2\)CH\(_2\) ester units attached to the pyridyl units at the C\(^1\) position. The inclusion of these units should increase the solubility of the complexes in organic solvents, but are sufficiently remote from the metal binding sites not to interfere with either the self-assembly of the complex or its ability to act as a host for anions. Reaction of either \( L^1a \) and \( L^2a \) with Cu(OTf)_2 in DCM (containing 3% MeOH) resulted in dark-yellow or pale-yellow solutions, respectively (Figure 3a). Partitioning of either of these solutions with water containing one equivalent of NaH\(_2\)PO\(_4 \) resulted in a color change indicative of phosphate encapsulation (Figure 3b). UV/Vis studies show that binding of anions occurs in organic solvent as addition of one equivalent of a range of anions (as their tetraalkylammonium salts) to either \([ [ L^1 \text{Cu}_3 \text{SO}_4 ] \text{(OTf)}_2 ] \) or \([ [ L^2 \text{Cu}_3 \text{SO}_4 ] \text{(OTf)}_2 ] \) in DCM does result in a change in the UV/Vis spectrum with chloride, bromide, sulfate and phosphate but not with nitrate. Furthermore, examination of the
organic layer from the biphasic system by ESI-MS showed ions at m/z 2606 and 1228 (for L₁a) and m/z 2572 and 1211 (for L₂a) corresponding to [(L₁a)Cu₃(PO₄)₃](OTf)₆⁺ and [(L₂a)Cu₃(PO₄)₃](OTf)₆⁺ for each ligand, respectively. This indicates that the trinuclear assembly persists in DCM and that the phosphate can be transferred, via incorporation into the host assembly, to the organic phase.[11]

In an effort to ascertain how much of the phosphate anion was extracted to the organic phase a series of extraction experiments were carried out and the amount of anion remaining in the aqueous solution examined by ion chromatography (Table 1). In this experiment 3 mL of 3% MeOH in DCM containing 4.95 μmol of either [(L₁a)Cu₃]₆⁺ or [(L₂a)Cu₃]₆⁺ (1.6 mm) was exposed to water containing 4.95 μmol of Na₂HPO₄, and the biphasic system was stirred for 18 hrs. As described previously the aqueous solution was analyzed for residual anion content by ion chromatography (Table 2).

A series of competitive experiments was carried out to investigate the selectivity of the assemblies to common anions. 3 mL of 3% MeOH in DCM containing 4.95 μmol of either [(L₁a)Cu₃]₆⁺ or [(L₂a)Cu₃]₆⁺ (1.6 mm) was exposed to water containing a mixture of NaCl, NaNO₃, NaHSO₄ and NaH₂PO₄ (4.95 μmol of each) and the biphasic system was stirred for 18 hrs. As described previously the aqueous solution was analyzed for residual anion content by ion chromatography (Table 2).

For [(L₁a)Cu₃]₆⁺ no chloride or nitrate anions are removed from the aqueous layer, but removal of both NaHSO₄ and NaH₂PO₄ occurs with the former being removed more effectively. For the L₂₅ assembly there is no preference between NaHSO₄ and NaH₂PO₄ but this cage also removes chloride from the system and it shows similar affinity for both phosphate and sulfate.

The same experiment was carried out on the extraction systems but the disodium salts Na₂SO₄ and Na₂HPO₄ were used instead of their monosodium analogues (Table 3). This data shows a very different trend from the previous results with host [(L₁a)Cu₃]₆⁺ showing a preference for phosphate, removing 59% of this anion from solution compared to only 15% of sulfate. Remarkably [(L₂a)Cu₃]₆⁺ removes no nitrate and only a small amount of either chloride or sulfate anions, but it removed 64% of Na₂HPO₄ demonstrating significant selectivity for this anion. The difference in selectivity between Na₂HPO₄/NaH₂PO₄ and Na₂SO₄/NaHSO₄ can be rationalized by the acidity of the mono-anion. In these systems once encapsulation has occurred the anions are fully deprotonated regardless of their original protonated state (c.f.

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**Table 1:** Percentage remaining Na₂HPO₄ from aqueous solutions exposed to an organic solution of either [(L₁a)Cu₃]₆⁺ or [(L₂a)Cu₃]₆⁺.

| Equivalents of Host | [(L₁a)Cu₃](OTf)₆ | [(L₂a)Cu₃](OTf)₆ |
|---------------------|-----------------|-----------------|
| 1                   | 18              | 24              |
| 1.1                 | 11              | 20              |
| 1.2                 | 6               | 14              |

Concentration of the phosphate anion measured by ion chromatography.

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**Table 2:** Percentage remaining of a solution of different anions from an aqueous solution exposed to an organic solution of either [(L₁a)Cu₃]₆⁺ or [(L₂a)Cu₃]₆⁺.

| Complex                     | NaCl [%] | NaNO₃ [%] | NaHSO₄ [%] | NaH₂PO₄ [%] |
|-----------------------------|----------|-----------|------------|-------------|
| [(L₁a)Cu₃](OTf)₆            | 110      | 103       | 47         | 73          |
| [(L₂a)Cu₃](OTf)₆            | 68       | 99        | 61         | 62          |

[a] Excess chloride anions are probably due to impurities from the other salts and solvents. [b] The removal is possibly slightly higher than reported due to excess chloride from contamination. Concentration of anions measured by ion chromatography.

**Table 3:** Percentage remaining of a solution of different anions from an aqueous solution exposed to an organic solution of either [(L₁a)Cu₃]₆⁺ or [(L₂a)Cu₃]₆⁺.

| Complex                     | NaCl [%] | NaNO₃ [%] | NaHSO₄ [%] | NaH₂PO₄ [%] |
|-----------------------------|----------|-----------|------------|-------------|
| [(L₁a)Cu₃](OTf)₆            | 107      | 101       | 85         | 41          |
| [(L₂a)Cu₃](OTf)₆            | 97       | 100       | 95         | 36          |

[a] Excess chloride anions are probably due to impurities from the other salts and solvents. [b] The removal is possibly slightly higher than reported due to excess chloride from contamination. Concentration of anions measured by ion chromatography.
reaction of \([L^2]^6\) with \(\text{Bu}_2\text{NH}_2\text{O}^+\) giving \([L^2]^6\). The monoanionic \(\text{HSO}_4^-\) \((pK_a = 1.81)\) is significantly more acidic than \(\text{H}_2\text{PO}_4^-\) \((pK_a = 7.21)\) and as a result \(\text{HSO}_4^-\) is more readily deprotonated and will occupy the cavity in preference to dihydrogen phosphate. This issue does not arise with the diatomic \(\text{SO}_4^{2-}\) and the cavity is selective for the phosphate (due to the difference in anionic charge) demonstrating that the selectivity of anion encapsulation, and extraction into organic solutions, can be controlled by \(pK_a\).[2]

Molecular modelling of these systems shows that there is a thermodynamic preference for both the self-assembled species \([L^1]^6\) and \([L^2]^6\) to act as hosts for anionic guests but the former (based on \(L^1\)) gives stronger anion binding than the latter (based on \(L^2\)). The formation energies of the complexes are \(-1197\) \(\text{kJ mol}^{-1}\) \([L^1]^6\text{Cu}_2(\text{PO}_4)^{4+}\), \(-1030\) \(\text{kJ mol}^{-1}\) \([L^2]^6\text{Cu}_2(\text{PO}_4)^{4+}\), and \(-1750\) \(\text{kJ mol}^{-1}\) \([L^1]^6\text{Cu}_2(\text{SO}_4)^{4+}\), and \(-478\) \(\text{kJ mol}^{-1}\) \([L^2]^6\text{Cu}_2(\text{SO}_4)^{4+}\). Based on the calculated energetics of the systems, the ratios of phosphate to sulphate distribution are predicted to be 54:46 for \([L^1]^6\text{Cu}_2(\text{PO}_4)^{4+}\) and \([L^2]^6\text{Cu}_2(\text{PO}_4)^{4+}\) and 79:21 for \([L^1]^6\text{Cu}_2(\text{SO}_4)^{4+}\) and \([L^2]^6\text{Cu}_2(\text{SO}_4)^{4+}\). Compared to \([L^2]^6\) \(\text{Cu}_2\), \([L^1]^6\text{Cu}_2\) shows a marginal preference for phosphate and but a much greater preference for sulphate. Comparison between the energies of the phosphate and sulfate host-guest complexes with each different ligand system shows that the distribution of phosphate and sulphate in the presence of \([L^1]^6\text{Cu}_2\) is 41:59, which indicates a thermodynamic preference for this system to act as a host for sulphate compared to phosphate (e.g. \([L^1]^6\text{Cu}_2(\text{SO}_4)^{4+}\) has a more free energy of formation than \([L^1]^6\text{Cu}_2(\text{PO}_4)^{4+}\)). The distribution of phosphate and sulphate inclusion in the presence of \([L^2]^6\text{Cu}_2\) is 68:32, which indicates a thermodynamic for the formation of \([L^2]^6\text{Cu}_2(\text{PO}_4)^{4+}\) over \([L^2]^6\text{Cu}_2(\text{SO}_4)^{4+}\), the opposite selectivity of the \(L^1\) system. The modelling results are in good agreement with the experimental results, which show that the extraction of phosphate from water with \([L^1]^6\text{Cu}_2\) is approximately 10% greater than with \([L^2]^6\text{Cu}_2\). Extraction of \(\text{NaCl}, \text{NaNO}_3, \text{NaHSO}_4\) and \(\text{NaH}_2\text{PO}_4\) shows a clear preference for the extraction of sulfate using \([L^2]^6\text{Cu}_2\), but little difference between \(\text{NaHSO}_4\) and \(\text{NaH}_2\text{PO}_4\) with \([L^2]^6\text{Cu}_2\). For the extraction of \(\text{Na}_2\text{SO}_4\) and \(\text{Na}_2\text{HPO}_4\), both complexes have a preference for phosphate, but this preference is more pronounced for \([L^1]^6\text{Cu}_2\). The subtle differences in the extraction ability of the two hosts \([L^1]^6\text{Cu}_2\) and \([L^2]^6\text{Cu}_2\) is a probable consequence of the different sizes of the cavities formed. Even though \(\text{PO}_4^{3-}\) is slightly larger than \(\text{SO}_4^{2-}\) (1.54 Å vs. 1.49 Å respectively)\[3\] the rigidity and smaller size of the cyclohexyl spacer unit, which induces shorter \(\text{Cu}^-\)–anion and \(\text{NH}^-\)–anion distances, may have a better size match to the phosphate and account for the greater selectivity of this anion.

In conclusion, we have shown that self-assembled systems with differing spacer units (viz. \([L^1]^6\text{Cu}_2\) and \([L^2]^6\text{Cu}_2\)) can act as anion binding hosts exhibiting differing selectivities for different anions. Inclusion of long chain ester units on these ligands can make the complexes soluble in organic solvents and these are capable of liquid-liquid extraction of phosphate anions from water, showing excellent selectivity in the presence of other common anions. Indeed, extraction experiments on Basel growth medium (BBM, an algal growth medium)\[4\] which contains a variety of nutrients (and possible interferences) demonstrates that the phosphate anion concentration can be reduced from 168 ppm to <5 ppm but chloride (27.5 ppm) and nitrate (521 ppm) are hardly affected. The sulfate is also reduced (28.4 ppm to <1 ppm) which is unsurprising as the complex is used in slight excess (1.2 equivalents with respect to phosphate). This clearly shows that phosphate can be selectively removed from model aquatic systems, indicating a possible method for phosphate sequestration from eutrophic systems.

Acknowledgements

M.M. would like to acknowledge the EPSRC (EP/R010366/1) for funding. P.B.W. would like to acknowledge The Royal Society (RG5/R1/191154) for funding.

Conflict of interest

The authors declare no conflict of interest.

Keywords: anions · copper · macrocycles · self-assembly · structure elucidation

[1] D. Cordell, J.-O. Drangert, S. White, Global Environ. Change 2009, 19, 292 – 305; J. A. Foley, N. Ramankutty, K. A. Braunam, E. S. Cassidy, J. S. Gerber, M. Johnston, N. D. Mueller, C. O’Connell, D. K. Ray, P. C. West, C. Balzer, E. M. Bennett, S. R. Carpenter, J. Hill, C. Monfreda, S. Polasky, J. Rockström, J. Sheehan, S. Siebert, D. Tilman, D. P. M. Aks, Nature 2011, 478, 337 – 342.
[2] D. Cordell, S. White, Food Security 2015, 7, 337 – 350; D. Cordell, S. White, Annu. Rev. Environ. Res. 2014, 39, 161 – 188; J. Elser, E. Bennett, Nature 2011, 478, 29 – 31.
[3] J. Grantham, Nature 2012, 491, 303.
[4] M. Oliveira, A. V. Machado, Environ. Technol. Rev. 2013, 2, 117 – 127; V. H. Smith, G. D. Tilman, J. C. Neiola, Environ. Pollut. 1999, 100, 179 – 196; B. Grizzetti, F. Bouraoui, A. Aloe, Global Change Biol. 2012, 18, 769 – 782.
[5] J. W. Steed, Chem. Soc. Rev. 2009, 38, 506 – 519; S. O. Kang, J. M. Llaneras, V. W. Day, K. Bowman-James, Chem. Soc. Rev. 2010, 39, 3980 – 4003; P. A. Gale, E. N. Howe, X. Wu, Chem 2016, 1, 351 – 422; N. Buschhaert, S. Park, K. Baek, Y. Choi, J. Park, E. Howe, J. Hiscock, L. Karagiannidis, I. Marques, V. Felix, W. Namkung, J. L. Sessler, P. A. Gale, I. Shin, Nat. Chem. 2017, 9, 667 – 675; P. A. Gale, J. Davis, R. Quesada, Chem. Soc. Rev. 2017, 46, 2497 – 2519; P. A. Gale, C. Talgaritone, Chem. Soc. Rev. 2015, 44, 4212 – 4227; N. Buschhaert, C. Talgaritone, V. W. Cornamus, P. A. Gale, Chem. Rev. 2015, 115, 8038 – 8155; J. Hiscock, M. Sambrook, N. Wells, P. A. Gale, Chem. Sci. 2015, 6, 5680 – 5684.
[6] M. J. Langton, C. J. Serpell, P. Beer, Angew. Chem. Int. Ed. 2016, 55, 1974 – 1987; Angew. Chem. 2016, 128, 2012 – 2026; J. L. Sessler, P. A. Gale, W.-S. Cho, Anion Receptor Chemistry, RSC, Cambridge, 2006; N. H. Evans, P. D. Beer, Angew. Chem. Int. Ed. 2014, 53, 11716 – 11754; Angew. Chem. 2014, 126, 11908 – 11948; S. Kubik, Chem. Soc. Rev. 2010, 39, 3648 – 3663; S. Kubik, C. Reyheller, S. Stiuwe, J. Inclusion Phenom. Macrocyclic Chem. 2005, 52, 137 – 187; S. Otto, S. Kubik, J. Am. Chem. Soc. 2003,
[7] R. Custelcean, Chem. Soc. Rev. 2014, 43, 1813–1824; P. Ballester, Chem. Soc. Rev. 2010, 39, 3810–3830; H. Amouri, C. Desmarests, J. Moussa, Chem. Rev. 2012, 112, 2015–2041; C. R. Rice, Coord. Chem. Rev. 2006, 250, 3190–3199; D. Zhang, T. K. Ronson, J. Mosquera, A. Martinez, L. Guy, J. R. Nitschke, J. Am. Chem. Soc. 2017, 139, 6574–6577; J.-F. Ayme, J. E. Beves, C. J. Campbell, G. Gil-Ramirez, D. A. Leigh, A. J. Stephens, J. Am. Chem. Soc. 2015, 137, 9812–9815; R. Custelcean, P. V. Bonnensen, N. C. Duncan, X. Zhang, L. A. Watson, G. V. Berkel, W. B. Parson, B. P. Hay, J. Am. Chem. Soc. 2012, 134, 8525–8534; R. Custelcean, A. Bock, B. A. Moyer, J. Am. Chem. Soc. 2010, 132, 7177–7185; R. Custelcean, F. V. Sloop, Jr., A. Rajbanshi, S. Wang, B. A. Moyer, Cryst. Growth Des. 2015, 15, 517–522.

[8] R. A. Faulkner, L. P. Harding, J. Higginson, C. R. Rice, C. Slater, Angew. Chem. Int. Ed. 2014, 53, 13540–13543; Angew. Chem. 2014, 126, 13758–13761; L. P. Harding, J. C. Jeffery, T. Rii-Johannessen, C. R. Rice, Z. Zeng, Chem. Commun. 2004, 654–655; L. P. Harding, J. C. Jeffery, T. Rii-Johannessen, C. R. Rice, Z. Zeng, J. Chem. Soc. Dalton Trans. 2004, 2396–2397; R. A. Faulkner, N. J. Patmore, C. R. Rice, C. Slater, Chem. Commun. 2018, 54, 9159–9162.

[9] C. R. Rice, C. Slater, R. A. Faulkner, R. L. Allan, Angew. Chem. Int. Ed. 2018, 57, 13071–13075; Angew. Chem. 2018, 130, 13255–13259.

[10] N. J. Williams, C. A. Seipp, K. A. Garrabrant, R. Custelcean, E. Holguin, J. K. Keum, R. J. Ellis, B. A. Moyer, Chem. Commun. 2018, 54, 10048–10051; P. Sokkalingam, J. Shraberg, S. W. Rick, B. C. Gibb, J. Am. Chem. Soc. 2016, 138, 48–51; M. D. Zhang, T. K. Ronson, J. Mosquera, A. Martinez, J. R. Nitschke, Angew. Chem. Int. Ed. 2018, 57, 3717–3721; Angew. Chem. 2018, 130, 3779–3783.

[11] A UV/Vis titration to ascertain the binding constants of the cryptand shows that under the conditions measured (near 1 mM) K values are $\approx 10^3$ M$^{-1}$.

[12] For a solution of NaCl, NaNO$_3$, Na$_2$SO$_4$, and Na$_2$HPO$_4$ pH 8.85 whereas NaCl, NaNO$_3$, NaHSO$_4$, and NaH$_2$PO$_4$ pH 2.81.

[13] Bond lengths measured in [(L$_3^-$)Cu$_3$(PO$_4$)]$^{3+}$ and [(L$_3^-$)Cu$_3$(SO$_4$)]$^{3+}$ allow direct comparison (see Ref. [9]) and are in good agreement with a variety of literature sources.

[14] H. C. Bold, Bull. Torrey Bot. Club 1949, 76, 101–108.

Manuscript received: July 20, 2020
Accepted manuscript online: August 2, 2020
Version of record online: September 2, 2020