Selective Anion Extraction and Recovery Using a Fe^{II}₄L₄ Cage

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Abstract: Selective anion extraction is useful for the recovery and purification of valuable chemicals, and in the removal of pollutants from the environment. Here we report that $Fe^{II}_{4}L_{4}$ cage 1 is able to extract an equimolar amount of ReO_{4}^{-} , a high-value anion and a nonradioactive surrogate of TcO_{4}^{-} , from water to nitromethane. Importantly, the extraction was efficiently performed even in the presence of 10 other common anions in water, highlighting the high selectivity of 1 for ReO_{4}^{-} . The extracted guest could be released into water as the cage disassembled in ethyl acetate, and then 1 could be recycled by switching the solvent to acetonitrile. The versatile solubility of the cage also enabled complete extraction of ReO_{4}^{-} (as the tetrabutylammonium salt) from an organic phase into water by using the sulfate salt of 1 as the extractant.

Rhenium is among the rarest elements in the Earth's crust,^[1] but it is a key ingredient for modern industry. It is used as catalyst for petroleum refining,^[2] in the high-melting superalloys of jet engines,^[3] and in new superhard materials,^[4] to cite only three examples. The limited supply and great demand lead to a high cost, generating an economic incentive for new means to extract, separate and recycle rhenium as perrhenate (ReO4^-).^[5]

Because of its similar structure and almost identical charge density, perrhenate is also used as a nonradioactive surrogate for pertechnetate (99TcO4-),[6] which is an important radiopharmaceutical and one of the most problematic radioactive ions in nuclear waste.^[7] Significant advances have been made in designing sorbent materials for removing ReO₄⁻/TcO₄⁻ from aqueous solution by liquid-solid extraction.^[7-8] These solid materials take up anionic targets from water via anion exchange. An attractive alternative to such sorbents is the use of supramolecular receptors as liquid-phase extractants,^[9] although only a few such ReO4-/TcO4- receptors have been reported.^[10] Compared to solid-state anion exchange materials, supramolecular extractants functioning through molecular recognition offer the potential for better selectivity toward target anions. Their flexibility in solution may provide a better size and shape match in order to optimize specific interactions between receptors and substrates.^[7] Such receptors can thus help address the major challenge in supramolecular chemistry of anion recognition in water.[11]

Most supramolecular anion extractants have been robust covalent receptors^[12] as opposed to coordination cages.^[13] Such extractants must be stable in the presence of both water and organic solvents,^[14] properties that are easier to engineer for covalent systems. Nevertheless, compared to the synthesis of

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covalent cages, the preparation of self-assembled capsules usually involves less synthetic complexity. The dynamic nature of coordination bonds^[15] may also enable guest release and subsequent recycling of the extractant.^[16]

We recently reported the water-soluble sulfate salt of azaphosphatrane-based $Fe^{II}_4L_4$ tetrahedron 1 (Figure 1), which can adaptively encapsulate different anions *via* hydrogen bonding and electrostatic interactions in water.^[17] Herein, we develop 1 as an efficient and selective extractant, capable of extracting ReO_4^- in either direction between organic and aqueous phases. We also establish a simple solvent-switching procedure that allows 1 to be disassembled, releasing its anionic cargo and allowing it to be recycled.

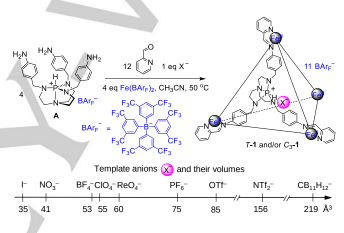


Figure 1. Subcomponent self-assembly of 1 around 1 equiv of template anion.

Non-coordinating tetrakis(3,5bis(trifluoromethyl)phenyl)borate (BAr_F⁻) was selected as the counter-anion for **1** in this work based on its lipophilicity and bulk (Figure 1). The lipophilic nature of BAr_F⁻ renders **1** soluble in water-immiscible organic solvents such as nitromethane. BAr_F⁻ is larger (968 Å³)^[18] than the cavity volume of **1** at its most expansive (253 Å³; see below), precluding competition with any of the anions discussed below.

The BAr_F⁻ salt of subcomponent **A** (Figure 1) was obtained by anion metathesis (Supporting Information section 2.1). As was observed in water,^[17] the reaction of **A** (4 equiv) with Fe(BAr_F)₂ (4 equiv) and 2-formylpyridine (12 equiv) in acetonitrile failed to give the expected cage complex **1**·[BAr_F]₁₂, which required an internal template anion (listed in Figure 1) for its formation.

In acetonitrile, template anions with volumes below 53 Å³ gave rise to both a C_3 -symmetric isomer (C_3 -1, with one azaphosphatrane ⁺P–H group oriented away from the inner cavity and the other three pointed inward) and a *T*-symmetric isomer (*T*-1, containing four inwardly-directed ⁺P–H groups) (Figure S1), whereas larger anionic templates, having volumes \geq 55 Å³, resulted in the formation of *T*-1 exclusively (Figure S2), as was observed in water.^[17] The initially obtained mixture of isomers in the former case is kinetically metastable and gradual

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interconversion between cage isomers was observed. Energy barriers of conversion in CD_3CN at 323 K were determined to be similar to the values previously obtained in water at 298 K^[17] (Figures S3-S6).

We then tested the stability of the cage, as $Tf_2N^- \subset 1 \cdot [BAr_F]_{11}$ ($Tf = CF_3SO_2$), in ethyl acetate and nitromethane, both of which are water-immiscible organic solvents suitable for liquid-liquid extraction experiments. Ca. 65% of 1 was observed to disassemble at a concentration of 1.5 mM in EtOAc after 4 h (Figure S7), with complete disassembly occurring at more dilute concentrations. In contrast, the cage was stable without any decomposition in CD_3NO_2 for at least two weeks at room temperature (Figure S9). We infer that the more polar solvent nitromethane offers a greater degree of stabilization to highly cationic 1 than does less polar ethyl acetate.^[19] Nitromethane was thus chosen as the organic solvent for liquid-liquid extractions.

Interestingly, cage reassembly was observed after evaporation of EtOAc and redissolution of **1** in CD_3CN , indicating a reversible process (Figure S8). This phenomenon provides an original means of guest release and extractant recovery, as explored further below.

Through competitive guest exchange, we were able to measure the relative binding affinities of different anions in CD₃NO₂. The following hierarchy was observed: CB₁₁H₁₂⁻ > ReO₄⁻ > TfO⁻ > PF₆⁻ > CIO₄⁻ > Tf₂N⁻ > BF₄⁻ > I⁻ > NO₃⁻ (Figures S10-S17, Table S1). This ordering differs from the one observed in water: PF₆⁻ > ReO₄⁻ > TfO⁻ > CIO₄⁻ > CB₁₁H₁₂⁻ > Tf₂N⁻ > BF₄⁻ > I⁻ > NO₃⁻ [^{17]} especially as regards the binding affinity of CB₁₁H₁₂⁻. To accommodate this largest anion, the cage framework must expand; we infer that this larger conformation in water is unfavorable because it involves greater exposure of hydrophobic surface to water. In both solvents, ReO₄⁻ binds more strongly than other common anions, indicating potential for its selective extraction.

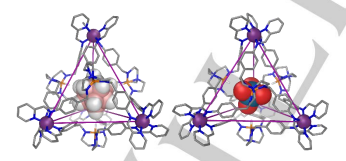


Figure 2. X-ray crystal structures of $CB_{11}H_{12}$ —C1 (left) and ReO_4 —C1 (right). Disorder, unbound counterions, non-P-bound hydrogen atoms, and solvents are omitted for clarity.

We obtained single crystals of 1 encapsulating the two most strongly bound anions in nitromethane, $CB_{11}H_{12}^-$ and ReO_4^- . X-ray diffraction analyses^[20] (Figure 2) showed a *T*-symmetric framework for both structures. The structures demonstrate the flexibility of the cage skeleton, allowing adaptation to guests of different sizes. Calculated cavity volumes of 157 Å³ and 253 Å³ were obtained for the ReO_4^- (volume 60 Å³) and $CB_{11}H_{12}^-$

(volume 219 Å³) complexes respectively (Figure S18). Cavity expansion occurs through outward motion of the azaphosphatrane faces, resulting in a more open surface having pores of ca. 2.5 Å in $CB_{11}H_{12}$ -C1, compared to ca. 1.2 Å in ReO_4 -C1.

Since Tf_2N^- is the most weakly bound among anions capable of templating 7-1 exclusively, extraction of ReO_4^- was initially investigated using $Tf_2N^-C1\cdot[BAr_F]_{11}$ as the extractant. After mixing 0.8 mM $Tf_2N^-C1\cdot[BAr_F]_{11}$ in CD_3NO_2 with 0.8 mM NaReO₄ in D₂O for 7 h, no further uptake of ReO_4^- by 1 was observed. ¹H NMR spectroscopy of the CD_3NO_2 phase revealed that 60% of the ReO_4^- from the aqueous phase had been extracted as $ReO_4^-C1\cdot[BAr_F]_{11}$, with the remainder of the 1 binding Tf_2N^- (Figure S19). After displacement by the extracted ReO₄⁻, free Tf_2N^- thus transferred from CD_3NO_2 to D₂O as the sodium salt.

We investigated the effect of the counter-ions of the Tf₂N⁻ template by adding TBANTf₂ (TBA = tetra-*n*-butylammonium), KNTf₂ or LiNTf₂ during the self-assembly, but no cation effect on the efficiency of ReO₄⁻ extraction was observed (Figure S20). Similarly, increasing the concentrations of Tf₂N⁻⊂**1**·[BAr_F]₁₁ in CD₃NO₂ and NaReO₄ in D₂O to 1.3 mM (Figure S20f) did not impact extraction efficiency.

The extraction of TfO⁻ (using NaOTf) from water under identical liquid-liquid conditions was also successful but with a lower efficiency (43%, Figure S21). Control experiments confirmed that without the cage, NaOTf did not transfer to the CD_3NO_2 phase (Figure S22).

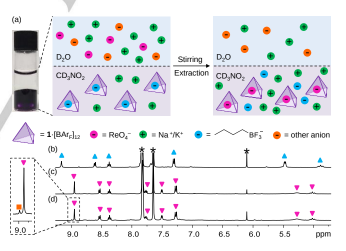


Figure 3. (a) Selective liquid-liquid extraction of ReO₄⁻ in the presence of other anions. Conditions: 0.8 mM "BuBF₃⁻ ⊂ 1·[BAr_F]₁₁ in CD₃NO₂; 0.8 mM in D₂O of each of NaReO₄, NaF, NaCl, NaBr, Nal, Na₂SO₄, KCIO₄, KNO₃, NaBF₄, NaH₂PO₄, and NaOAc; 7 hours stirring at rt; (b) - (d) Partial ¹H NMR spectra of (b) the CD₃NO₂ phase before extraction, showing only the presence of "BuBF₃⁻ ⊂ 1·[BAr_F]₁₁ (▲); (c) the CD₃NO₂ phase after extraction in the absence of competing anions, showing only the presence of ReO₄⁻ ⊂ 1·[BAr_F]₁₁ (▲); (d) the CD₃NO₂ phase after extraction in the presence of competing anions, showing the presence of 97% ReO₄⁻ ⊂ 1·[BAr_F]₁₁ (▼) and 3% CIO₄⁻ ⊂ 1·[BAr_F]₁₁ (■). The peaks of BAr_F⁻ and the trimethoxybenzene standard are denoted by asterisks.

In order to improve the extraction efficiency, we sought a more weakly bound template anion that avoided the complexity

of generating a mixture of cage diastereomers. Such an anion was found to be *n*-butyltrifluoroborate ("BuBF₃⁻). We found "BuBF₃⁻ to be able to template **7-1** exclusively (Figures S23-S28), and the resultant "BuBF₃⁻⊂**1**·[BAr_F]₁₁ to be stable in CD₃NO₂ for weeks. Moreover, 1 equiv of Tf₂N⁻ in CD₃NO₂ almost completely displaced the encapsulated "BuBF₃⁻ (Figure S29), marking "BuBF₃⁻ as the weaker binder.

When the extractant ${}^{n}BuBF_{3} \subset 1 \cdot [BAr_{F}]_{11}$ in CD₃NO₂ was mixed with an equimolar amount of NaReO₄ in D₂O, only ReO₄- $\subset 1 \cdot [BAr_{F}]_{11}$ was observed after extraction, indicating complete removal of ReO₄- from water (Figure 3c and Figure S30). Complete extraction of TfO⁻ from aqueous NaOTf was also achieved by using ${}^{n}BuBF_{3} \subset 1 \cdot [BAr_{F}]_{11}$ (Figure S31).

Encouraged by these results, we evaluated the selectivity of 1 toward ReO₄⁻ in the presence of 10 other different anions simultaneously in water: F⁻, Cl⁻, Br⁻, l⁻, SO₄²⁻, ClO₄⁻, NO₃⁻, BF₄⁻, H₂PO₄⁻, and AcO⁻ (1 equiv to ReO₄⁻ in each case). The extraction efficiency for ReO₄⁻ by "BuBF₃⁻⊂1·[BAr_F]₁₁ in the presence of this anion library was 97%, with ClO₄⁻ comprising the other 3% extracted (Figure 3).

We also developed a strategy to release and separate the extracted guest and recover the cage extractant by exploiting the instability of **1** in less polar solvents. As shown in Figure 4, after extraction, the nitromethane layer was separated and the solvent evaporated. The isolated cage was then redissolved in degassed EtOAc. As described above, the cage disassembled in this solvent. The extracted guest transferred to the water phase as KReO₄, pairing with K⁺ from "BuBF₃K, allowing its removal as the phases were separated. Regeneration of "BuBF₃⁻ \subset 1·[BAr_F]₁₁, which could be reused for further extraction experiments, was realized by evaporating the ethyl acetate and adding acetonitrile, along with "BuBF₃K (Figure S32).

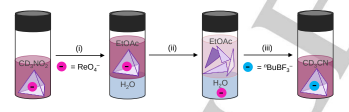


Figure 4. Illustration of cage extractant recycling: (i) After evaporation of CD₃NO₂, ReO₄⁻ ⊂ 1·[BAr_F]₁₁ was redissolved in degassed EtOAc; degassed H₂O was then added. (ii) After stirring for 4 h, the cage disassembled and ReO₄⁻ was released, transferring to the H₂O phase. (iii) After separation and evaporation of the EtOAc layer, addition of CD₃CN and "BuBF₃⁻ resulted in regeneration of the extractant "BuBF₃⁻ ⊂ 1·[BAr_F]₁₁.

Interestingly, due to the versatile solubility of **1**, either ReO₄⁻ or TfO⁻ could also be extracted from an organic phase into water, in the opposite direction to what was described above. In this case, Tf₂N⁻⊂**1**·[SO₄]_{5.5} as extractant completely removed either ReO₄⁻ or TfO⁻ from CD₃NO₂ into D₂O (Figures 5 and S33). Control experiments showed that without the cage, TBAReO₄ and TBAOTf did not transfer to D₂O (Figures S34).

In summary, we have demonstrated for the first time the feasibility of using a coordination cage for biphasic extraction. By employing BAr_{F}^{-} as counter-anion and ${}^{n}BuBF_{3}^{-}$ as template, ${}^{n}BuBF_{3}^{-}$ –1·[BAr_F]₁₁ was capable of completely extracting ReO₄⁻

from water into nitromethane. An efficiency of 97% was achieved even in the presence of 10 competing anions. A novel strategy for extractant regeneration was developed by taking advantage of the differential stability of **1** across solvents. Moreover, due to the versatile solubility of **1** when paired with different counter-anions, complete extraction of ReO_4^- (TBAReO₄) from an organic phase into water could also be accomplished by using Tf₂N⁻C1·[SO4]_{5.5}. The selective extraction properties of the cage toward perrhenate suggest great potential for recycling rhenium compounds, purification of chemicals, and for pertechnetate removal from water. Concepts developed in this study may also be generalized to enable the purification of other species using different coordination cages.

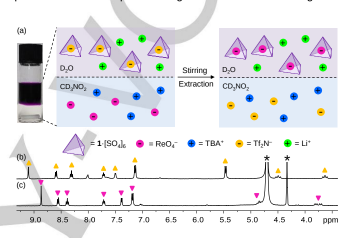


Figure 5. (a) Illustration of the liquid-liquid extraction of ReO₄⁻ from an organic phase into water. Conditions: 0.8 mM Tf₂N⁻ \subset 1·[SO₄]_{5.5} in D₂O; 0.8 mM TBAReO₄ in CD₃NO₂; 3 hours stirring. (b) – (c) Partial ¹H NMR spectra of (b) the D₂O phase before extraction, showing only Tf₂N⁻ \subset 1·[SO₄]_{5.5} (\blacktriangle), and (c) the D₂O phase after extraction, showing only ReO₄⁻ \subset 1·[SO₄]_{5.5} (\checkmark).HDO and CHD₂NO₂ peaks are represented by asterisks.

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Keywords: coordination cage • anion receptor • liquid-liquid extraction • self-assembly • supramolecular chemistry

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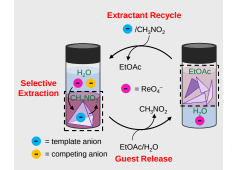
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COMMUNICATION

A Fe^{II}₄L₄ coordination cage enabled complete extraction of ReO₄⁻, a nonradioactive surrogate of TcO₄⁻, from water into an organic phase. In the presence of 10 other anions, 97% of ReO₄⁻ was selectively removed. The extracted ReO₄⁻ could be released and the cage extractant was recycled by a solvent-switching strategy. Rendering the cage watersoluble also allowed complete extraction of ReO₄⁻ from an organic phase into water.



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Page 1 – Page 4

Selective Anion Extraction and Recovery using a Fe^{II}₄L₄ Cage