Lanthanides Singing the Blues: Their Fascinating Role in the Assembly of Gigantic Molybdenum Blue Wheels

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ABSTRACT: Molybdenum blues (MBs) are a distinct class of polyoxometalates, exhibiting versatile/impressive architectures and high structural flexibility. In acidified and reduced aqueous environments, isopolyoxomolybdates generate precisely organize building blocks, which enable unique nanoscopic molecular systems (MBs) to be constructed and further fine-tuned by hetero elements such as lanthanide (Ln) ions. This Review discusses wheel-shaped MB-based structure types with strong emphasis on the ∼30 Ln-containing MBs as of August 2021, which include both organically hybridized and nonhybridized structures synthesized to date. The spotlight is thereby put on the lanthanide ions and ligand types, which are crucial for the resulting Ln-patterns and alterations in the gigantic structures. Several critical steps and reaction conditions in their synthesis are highlighted, as well as appropriate methods to investigate them both in solid state and in solution. The final section addresses the homogeneous/heterogeneous catalytic, molecular recognition and separation properties of wheel-shaped Ln-MBs, emphasizing their inimitable behavior and encouraging their application in these areas.

KEYWORDS: polyoxomolybdate, hybrid organic—inorganic, stereoselectivity, catalysis, molecular recognition, lanthanide separation

1. INTRODUCTION

Orthomolybdate ([MoO$_4$]$^{2-}$) solutions exhibit a unique chemistry as they allow for the assembly of nanosized scaffolds with the general formula $[X_{a}Y_{b}MoVI_{a}MoV_{b}O_{v}(H_{2}O)_{w}]^{z-}$ $([Mo_{x}X_{y}])$ (a = number of organic ligands; b = number of metallic hetero elements; c = degree of protonation; x and y = number of unreduced and reduced molybdenum, respectively; z = number of oxygens; v = number of coordinated water; n = resulting charge of the nanosized scaffold; w = sum of unreduced and reduced molybdenum). They are up to ∼4 nm large (for wheel-type clusters)$^1$ and exhibit unprecedented topology.$^2$ In a one-pot reaction an aqueous orthomolybdate solution is acidified (pH ≤ 4.5)$^3$ to initiate the polymerization process resulting in a number of isopolyoxomolybdates (IPOMos) (Figure 1). Generally, polymerization processes of orthomolybdate [MoO$_4$]$^{2-}$ in acidified solutions with different concentrations greater than 10$^{-8}$ M (Figure 1).$^4$ The speciation of [MoO$_4$]$^{2-}$ in acidified solutions with different concentrations and ionic strengths has been comprehensively investigated.$^5-^8$ In strongly alkaline environments, the monomeric [MoO$_4$]$^{2-}$ anion predominates, while in strongly acidic solutions the dominant species are stable mono-[MoO$_4$](OH)(H$_2$O)$_3$$^{+}$ or bi-[MoO$_4$](OH)(H$_2$O))$_2$$^{+}$ oxocations able to form inorganic—organic complexes with organic ligands (e.g., dicarboxylic acids, hydroxyaldehydes, amines, etc.).$^7$ The major species formed first upon acidification of aqueous [MoO$_4$]$^{2-}$ (Figure 1) is heptamolybdate, H$_3$[Mo$_7$O$_{24}$](6−x)$^{−}$ (Figure 1), which reaches typically its maximum concentration at pH 5 and can...
be protonated upon further acidification.\textsuperscript{4} When the environment becomes more acidic, the generation of $\beta\cdot\left[\text{Mo}_8\text{O}_{26}\right]^{4-}$ prevails over $\text{H}_x\left[\text{Mo}_8\text{O}_{26}\right]^{(6-x)-}$ ($x = 0-2$; max. conc. of $\text{H}_x\left[\text{Mo}_8\text{O}_{26}\right]^{(6-x)-}$ is reached at pH 4.5) and reaches a maximum at pH $\sim 2.5$. $\left[\text{Mo}_{36}\text{O}_{112}\left(\text{OH}_2\right)_{16}\right]^{8-}$ (Figure 1), which is a dimer of $\left[\text{Mo}_{36}\text{O}_{56}\left(\text{H}_2\text{O}\right)_{8}\right]^{14-}$ with an inversion center, is the largest known IOPOMo present in solution under nonreducing conditions below pH 2.8.\textsuperscript{15}

Upon acidification and subsequent addition of a reducing agent (e.g., $\text{N}_2\text{H}_4\cdot\text{N}_2\text{S}_2\text{O}_4$) the orthomolybdate ($\left[\text{MoO}_{6}\right]^{4-}$) solution turns blue and the virtual linker units ($\text{MoO}_6$) (Figure 2A)\textsuperscript{16} and the fundamental building block ($\text{Mo}_8$) (Figure 2A) was synthesized and structurally elucidated (Figure 5B)), whereby clusters of lower nuclearity (starting from $\text{Mo}_{138}^{20}$ (Figure 5C) to $\text{Mo}_{176}^{17}$) contain defect sites (\textsuperscript{3} = vacancies within the MBs due to the lack of $\text{Mo}_2$-units) (Figure 5C). $\text{Mo}_{176}$\textsuperscript{1} is the largest discrete wheel to date, which is constructed of two $\text{Mo}_1$, $\text{Mo}_2$, and $\text{Mo}_8$ building blocks more than $\text{Mo}_{154}$.\textsuperscript{19} The incorporation of these additional building blocks also results in the largest known nanocavity ($\text{Mo}_{154}$) exhibits a 20 Å and $\text{Mo}_{178}$ a 25 Å nanocavity; see Figure 5A and B).

Smaller fragments of suitable size, such as the IPOPOMo $\left[\text{Mo}_{36}\text{O}_{112}\left(\text{OH}_2\right)_{16}\right]^{8-}$ ($\left[\text{Mo}_{36}\right]^{15}$),\textsuperscript{15} can be templated into these wheel-shaped structures by stabilization in the central nanocavity of the wheel via supramolecular interactions, resulting in host–guest systems (e.g., $\text{Mo}_{36}\text{C}\left(\text{Mo}_{110}\right)$ with the symbol “C” representing the inclusion of a template cluster in a ring) (Figure 3B).\textsuperscript{21} This template effect was first realized by the incorporation of two nonsoluble $\left[\text{Mo}_{36}\text{O}_{56}\left(\text{H}_2\text{O}\right)_{8}\right]^{14-}$ “hubcaps” into the nanocavity of the $\text{Mo}_{176}$ nanowheel, thereby forming the “nanocookie” ($\text{Mo}_{248}$).\textsuperscript{22}

Aside from generating defects in the cluster or introducing templates, another structural modification is the coordinative fixation of organic ligands onto $\left[\text{Mo}_8\right]^{(\text{MoVII})}\left(\text{MoIII}\right)$\textsuperscript{17} units or replacement of $\text{Mo}_2$ units by metallic hetero elements of adequate size, such as lanthanide ions (Figure 2B) in the inner rings (the terms inner and outer ring rim refer to the inner and outer regions of the wheels; Figure 5). The coordinative attachment of organic ligands on MBs was achieved initially by hybridizing $\text{Mo}_{154}$ with cysteines.\textsuperscript{23} This type of MBs (= organic–inorganic hybrids) (Figure 3D) represents a fusion of inorganic, nanosized giant clusters with organic components and possess characteristics that vary from their parent cluster, such as...
decreased solubility in comparison to the parent wheel\textsuperscript{20,24} and a functionalized nanocavity (Figure 6B).

The integration of lanthanide ions into MBs was accomplished first by the introduction of Pr\textsuperscript{III} into an MB scaffold, forming the wheel \{Mo\textsubscript{12}Pr\textsubscript{6}\}\textsuperscript{155} (Figure 5D), which exhibited a "Japanese rice-ball"-shape with an outer diameter of \(\sim 31\ \text{Å}\) rather than a circular-shape as the parent wheel \{Mo\textsubscript{154}\} (Figure 5A).

The incorporation of lanthanide ions in MBs is particularly intriguing since new unique MB characteristics may arise. For example, when complexed, lanthanides can behave like typical Lewis acids and exhibit interesting catalytic properties. In several organic transformation reactions, lanthanide-containing POMs show significant chemoselectivity.\textsuperscript{25-28} Furthermore, lanthanide ions in POM-based nanostructures are also able to form inorganic–organic adducts, which can be exploited in chiral recognition.\textsuperscript{29}

This Review addresses (i) the synthesis and reaction conditions of all available structural types of MB-based wheels (Figure 3) with strong emphasis on their derivatization with lanthanide ions as the fascinating role of lanthanide ions in wheel-type MBs has not been covered yet in a review, (ii) thoroughly the geometrical changes and charge shifts to the parent \{Mo\textsubscript{154}\} cluster caused by the introduction of lanthanide ions, (iii) the inclusion of templates, and (iv) the introduction and role of (structure directing and chiral-inducing) organic ligands in MB-based architectures. As the main focus is on Ln-MB structures, techniques for confirming their intactness in solution as well as investigating their solution speciation (e.g., "blackberries"\textsuperscript{30,31}) are presented using selected examples. The final section is devoted to potential applications of wheel-shaped Ln-MB architectures as homo- and heterogeneous catalysts as well as attractive components for molecular recognition. Additionally, specific Ln-MB syntheses are described as an elegant approach in lanthanide separation.\textsuperscript{32}

2. GENERAL SYNTHESIS APPROACH OF MB/Ln-MB-BASED NANOCLUSTERS

The general approach to synthesize all types of MBs/Ln-MBs, which are shown in Figure 3, is quite straightforward. The partial reduction of molybdate(VI) solutions in an acidic environment is shared by all cluster type syntheses. Adjusted molybdate concentrations allow for the production of desirable clusters while preventing the formation of undesired by-products.\textsuperscript{22,23} A synthetic approach to force these cluster to crystallize is to destroy their hydration shell by a high electrolyte concentration.\textsuperscript{33}

The amount of reducing agent has less of an effect on MB formation as the pH of the reaction solution decreases.\textsuperscript{33} Above a pH value of \(\sim 1.5\), excessively high amounts of reducing agents (e.g., Na\textsubscript{2}S\textsubscript{2}O\textsubscript{4}) lead to brown reaction mixtures (a compound class containing the structural building block unit \(\text{O}^{6-}\text{Mo}^{\text{IV}}=\text{O}^{5-}\text{Mo}^{\text{V}}\text{O}^{\text{IV}}\)) rather than blue solutions. Typically, the pH is indirectly proportional to the nuclearity of the cluster (i.e., a decrease of the pH results in an increase of the clusters’ nuclearity)\textsuperscript{17,33-36} and the molar ratio of molybdate/additional framework-building reagents (metallic heteroions, ligands) must be optimized for successful crystallization.\textsuperscript{24,25}

Organic ligands for the synthesis of hybridized inorganic–organic MBs/Ln-MBs should have a carboxyl group in order to be able to attach to \{Mo\textsubscript{x}\} units (always depicted as red polyhedra in all the figures). Furthermore, amino groups in ligands (e.g., amino acids and peptides) are of essential importance particularly in Ln-MB syntheses when it comes to (a) stabilization of templates inside the rings,\textsuperscript{38} (b) introduction of chirality,\textsuperscript{24,39} and (c) modulation of the shape and size of Ln-MBs.\textsuperscript{39}

3. SYNTHESIS OF MB/Ln-MB BASED NANOCLUSTERS

3.1. MB Based Nanoclusters

Complete MB ring systems (Figure 4A) such as \{Mo\textsubscript{154}\} are often synthesized at room temperature, with the pH value (typically below 1.4) being the most critical factor due to its strong effect on the nuclearity of the wheel\textsuperscript{17,19}. The complete MB ring system can be transformed into templated complete MB ring systems (Figure 4B) or into lacunary MB ring systems by creating defects (i.e., vacancies caused by a lack of \{Mo\textsubscript{x}\} building blocks) (Figure 4C). The inclusion of a template was first achieved by synthesizing the unprecedented structure \{Mo\textsubscript{148}\}, which is often described as a "nanocookie"\textsuperscript{22} and the adjustment of the pH between 1.4 and 4.5 made it possible to synthesize MBs with a different number of defects.\textsuperscript{33} The obtained defect-containing MB wheels are derived from the \{Mo\textsubscript{154}\}-cluster missing a certain number of \{Mo\textsubscript{x}\}-units (one unit at pH 1.4 and eight units at pH 4.5). They are specified with \{Mo\textsubscript{154-}\textsubscript{x}\} \((x = \text{number of defect sites})\), with \{Mo\textsubscript{138}\} being the wheel bearing the largest number of defects \((x = 16)\) to date (eight \{Mo\textsubscript{x}\}-units missing in total) (Figure 5C).\textsuperscript{5}

Complete MB ring systems (Figure 3A) can also be organically hybridized to build hybridized complete MB ring systems (Figure 3D). The ligation of \{Mo\textsubscript{154}\} to eight L-histidine...
(Figure 6B) molecules represents a good example for organically functionalized MBs.24

Organic hybridization reactions can also be used to stabilize anionic templates in the center of MBs via hydrogen bonds resulting in templated and hybridized complete MB ring systems (Figure 4E). An example of such a host–guest complex is \{Mo_{36}\}⊂\{Mo_{150}\text{ornithine}_6\} (Figure 6C).38 The last representatives of lanthanide-free MB archetypes are lacunary hybridized MB ring systems, which contain defects within their scaffold as well as grafted carboxylic acids onto their remaining \{Mo\}-units (Figure 4F). Lacunary hybridized MB ring systems were initially realized with the organic-chemical hybridization of a \{Mo_{138}\} cluster leading to the inorganic−organic MB \{Mo_{138}\text{acetate}_6\} (Figure 6A).

Figure 4 gives an overview of the main synthesis factors in the assembly of all structural types of MB-based clusters.

### 3.2. Ln-MB Based Nanoclusters

#### 3.2.1. Synthesis of Purely Inorganic Ln-MBs

#### 3.2.1.1. “Japanese Rice-Ball”-, Egg-, Ellipsoid-, and Circular-Shaped Ln-MBs without Template Clusters.

The “Japanese rice-ball”-shaped wheels \{Mo_{120}\text{Ln}_6\}40 are complete Ln-MB ring systems (Figure 3G) with \text{Ln} = \text{Pr}^{III}, \text{Ce}^{III}, or \text{La}^{III} ions. The synthesis was performed at room temperature for 1–2 weeks at room temperature (see Table 2 for more details), whereby the lanthanide substitution pattern

Table 2 lists all crystal structures of lanthanide-containing molybdenum wheels as of August 2021, including information about the geometry, lanthanide pattern within the wheel, the most critical reaction parameters (such as total reagents and their stoichiometric ratios, pH (if specified), reducing agent/method, temperature, and reaction time), and applied methods to characterize the structure.

#### 3.2.1.2. Synthesis of Purely Inorganic Ln-MBs

#### 3.2.1.3. “Japanese Rice-Ball”-, Egg-, Ellipsoid-, and Circular-Shaped Ln-MBs with Template Clusters.

The “Japanese rice-ball”-shaped wheels \{Mo_{120}\text{Ln}_6\} were synthesized in a similar manner, but a significantly different synthetic protocol was used for the \text{Ln} = \text{La}^{III} and \text{Ce}^{III} ions. The synthesis was performed at a rather high temperature (90 °C for 30 min) in an acidic and reduced medium immediately after all starting materials were produced, combined and had formed a cloudy suspension.37 A \[\text{NH}_4\text{Mo}_2\text{O}_5\text{H}_2\text{O}_{25}\] solution was photochemically reduced for 2 days at room temperature to get \{Mo_{120}\text{La}_6\}.40 All three “Japanese rice-ball”-shaped Ln-MBs crystallize within 1–2 weeks at room temperature (see Table 2 for more details), whereby the lanthanide substitution pattern

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(Figure 5). Polyhedral representation (and inner and outer diameter of the respective ring/rim) of the purely inorganic complete MB and all purely inorganic complete Ln-MB ring systems as of August 2021 described in this review starting with the (A) circle-shaped \{Mo_{154}\} (simplified archetype is shown in Figure 3A), (B) circle-shaped \{Mo_{176}\} (Figure 3A), (C) circle-shaped \{Mo_{138}\} (Figure 3C), (D) “Japanese rice-ball” \{Mo_{20}\text{Ln}_6\} (Figure 3G), (E) egg-shaped \{Mo_{96}\text{La}_8\} (Figure 3G), (F) egg-shaped \{Mo_{100}\text{Ce}_6\} (Figure 3G), (G) ellipsoidal \{Mo_{150}\text{La}_2\} (Figure 3G), (H) ellipsoidal \{Mo_{150}\text{La}_2\} (Figure 3G), (I) circular-shaped \{Mo_{120}\text{La}_6\} with \text{Ln} = \text{Nd}^{III} or \text{Sm}^{III} (Figure 3G), (J) circular-shaped \{Mo_{120}\text{La}_6\} with \text{Ln} = \text{La}^{III}, \text{Ce}^{III}, or \text{Pr}^{III} (Figure 3G), and (K) ellipsoidal half closed \{Mo_{130}\text{Ce}_6\} (Figure 3H). The open side is shown on the left and the closed side on the right. Coloring code: \{MoO_6\}, yellow; \{Mo_2O_{11}\}, red; \{Mo_8O_{35}\}, blue with central \{MoO_7\}-unit in cyan; \{LaO_9\}, green.
was built up in a one-pot synthesis and is therefore less controllable.

A complete Ln-MB ring system (Figure 3G) with the shape of an “egg” was first realized through the synthesis of \{\text{Mo}_{96}\text{La}_{8}\} (Figure 5E), in which a lacunary MB with an ambiguous number of defects was first dissolved and then treated with La\text{III} ions (see Table 2 for more details).  

Elliptically shaped Ln-MB scaffolds belonging to complete Ln-MB ring systems (Figure 3G) were assembled for the first time in a one-pot approach through the use of Eu\text{III} ions, which have smaller ionic radii than those of Ce\text{III} and Pr\text{III} ions. For the assembly of \{\text{Mo}_{128}\text{Eu}_{4}\} a precipitate of Eu\text{2}(\text{MoO}_{4})_{3} was suspended in an acidic solution and then reduced with hydrazine and heated at 60−65 °C for 45 min (see Table 2 for more details).

Table 1. Prominent Representatives of Purely Inorganic Complete Ln-MB Ring Systems Arranged in Descending Symmetry to the Parent MB \{\text{Mo}_{154}\}

| cluster | lanthanide ions | symmetry | ref |
|---------|-----------------|----------|-----|
| \{\text{Mo}_{154}\} | | D_{7d} | 19 |
| \{\text{Mo}_{152}\text{Ln}_{6}\} | La\text{III}, Ce\text{III}, Pr\text{III} | D_{3} | 25, 37, 40 |
| \{\text{Mo}_{152}\text{Ln}_{6}\} | Eu\text{III} | D_{2} | 44 |
| \{\text{Mo}_{152}\text{Ln}_{6}\} | Ce\text{III} | D_{d} | 31 |
| \{\text{Mo}_{152}\text{Ln}_{6}\} | Ce\text{III} | C_{s} | 32 |
| \{\text{Mo}_{152}\text{Ln}_{6}\} | Ce\text{III} | C_{2} | 24 |
| \{\text{Mo}_{152}\text{Ln}_{6}\} | Ce\text{III} | C_{2} | 41 |
| \{\text{Mo}_{152}\text{Ln}_{6}\} | Ce\text{III} | C_{2} | 42 |
| \{\text{Mo}_{152}\text{Ln}_{6}\} | Ce\text{III} | C_{2} | 43 |
| \{\text{Mo}_{152}\text{Ln}_{6}\} | Ce\text{III} | C_{2} | 39 |
Table 2. List of All Purely Inorganic and Hybridized Inorganic–Organic Lanthanide-Containing Molybdenum Blue Wheel Crystal Structures According to the CCDC and ICSD Database (August 2021); Ln-MBs Are Listed Based on Their Shape, Starting with the “Japanese Rice-Ball”

| Formula and building blocks given below | Shape | Ln-substitution pattern† | Synthesis method, pH, reaction temperature, and molar ratio of the cluster-building reagents§ | Characterization methods applied | Ref |
|----------------------------------------|-------|--------------------------|-----------------------------------------------------------------|---------------------------------|-----|
| \([\text{Mo}_6\text{O}_{24}\text{H}_6\text{O}_{18}]^{\text{2}^-}\) \((\text{Mo}_2\text{O}_2\text{H}_2\text{O})_6\text{Pr}_3\) \((\text{Figure} \ 6)\) | | | \(\text{Pr}_2\text{(MoO}_4\text{)}_2 + \text{hydrazine, acidified with } 0.4 \text{ M HCl, } 70^\circ \text{C (45 min)}\) | Single-crystal XRD, redox titration, TGA, IR, and Raman spectroscopy, BVS, elemental analysis | 41 |
| \([\text{Mo}_{13}\text{Ce}_{11}\text{O}_{32}\text{H}_{16}\text{O}_{18}]^{\text{2}^-}\) \((\text{Mo}_6\text{O}_2\text{H}_2\text{O})_6\text{La}(\text{H}_2\text{O})_4\) \((\text{Figure} \ 3)\) | “Japanese rice-ball” | | \(\text{La}^{3+} + [\text{MoO}_4]^{2-} + \text{sodium p-toluenesulfonate, pH 10}, \text{irradiation (1 day)}, \text{room temperature}\) | Single-crystal XRD, redox titration, UV-vis-NIR, IR spectroscopy, BVS, elemental analysis (only organic) | 37 |
| \([\text{Mo}_{13}\text{O}_{24}\text{H}_{16}\text{O}_{18}]^{\text{2}^-}\) \((\text{Mo}_6\text{O}_2\text{H}_2\text{O})_6\text{La}(\text{H}_2\text{O})_4\) \((\text{Figure} \ 3)\) | | | \(\text{Ce}^{3+} + \text{MO}_{2}^{2+} + \text{hydrazine + L-arginine, acidified with } 5 \text{ M HClO}_4, 90^\circ \text{C (1 h) (CeL-Arg/Mo = 0.1:0.04:1)}\) | Single-crystal XRD, TGA, UV-vis-NIR spectroscopy, IR spectroscopy, BVS, elemental analysis | 38 |
| \([\text{Mo}_{13}\text{O}_{24}\text{H}_{16}\text{O}_{18}]^{\text{2}^-}\) \((\text{Mo}_6\text{O}_2\text{H}_2\text{O})_6\text{La}(\text{H}_2\text{O})_4\) \((\text{Figure} \ 3)\) | | | \(\text{Ce}^{3+} + \text{MO}_{2}^{2+} + \text{hydrazine + L-ornithine, acidified with } 5 \text{ M HClO}_4, 90^\circ \text{C (1 h) (CeO} \_{\text{O}}\text{rM/Mo = 0.1:0.04:1)}\) | Single-crystal XRD, TGA, UV-vis-NIR spectroscopy, IR spectroscopy, BVS, elemental analysis | 38 |
| \([\text{Mo}_{13}\text{O}_{24}\text{H}_{16}\text{O}_{18}]^{\text{2}^-}\) \((\text{Mo}_6\text{O}_2\text{H}_2\text{O})_6\text{La}(\text{H}_2\text{O})_4\) \((\text{Figure} \ 3)\) | egg-shaped | | \(\text{La}^{3+} + \text{un-crystallized intermediate generated by the irradiation of } [\text{MoO}_4]^{2-}\text{propionic acid, pH 4.} + 4^\circ \text{C}\) | Single-crystal XRD, IR spectroscopy, UV-vis-NIR spectroscopy, elemental analysis | 40 |
| \([\text{Mo}_{13}\text{O}_{24}\text{H}_{16}\text{O}_{18}]^{\text{2}^-}\) \((\text{Mo}_6\text{O}_2\text{H}_2\text{O})_6\text{La}(\text{H}_2\text{O})_4\) \((\text{Figure} \ 3)\) | | | \(\text{Ce}_3\text{(MoO}_4\text{)}_2 + \text{hydrazine, acidified with } 5 \text{ M HClO}_4, 80-85^\circ \text{C (45 min)}\) | Single-crystal and powder XRD, redox titration TGA, UV-vis-NIR spectroscopy, IR and Raman spectroscopy, ESI-MS-MS spectroscopy, BVS, elemental analysis | 44 |
| \([\text{Mo}_{13}\text{O}_{24}\text{H}_{16}\text{O}_{18}]^{\text{2}^-}\) \((\text{Mo}_6\text{O}_2\text{H}_2\text{O})_6\text{La}(\text{H}_2\text{O})_4\) \((\text{Figure} \ 3)\) | ellipsoidal | | \(\text{La}^{3+} + [\text{MoO}_4]^{2-} + \text{dipropylamine, pH 10, irradiation (2 days), room temperature}\) | Single-crystal XRD, redox titration, UV-vis-NIR spectroscopy, IR spectroscopy, BVS, elemental analysis (only organic) | 40 |
| \([\text{Mo}_{13}\text{O}_{24}\text{H}_{16}\text{O}_{18}]^{\text{2}^-}\) \((\text{Mo}_6\text{O}_2\text{H}_2\text{O})_6\text{La}(\text{H}_2\text{O})_4\) \((\text{Figure} \ 3)\) | | | \(\text{La}^{3+} + [\text{MoO}_4]^{2-} + \text{cholic acid, pH 3.} + 4^\circ \text{C}\) | Single-crystal XRD, redox titration, TGA, IR spectroscopy, ELSAFS spectroscopy, magnetic susceptibility measurements | 45 |
| \([\text{Mo}_{13}\text{O}_{24}\text{H}_{16}\text{O}_{18}]^{\text{2}^-}\) \((\text{Mo}_6\text{O}_2\text{H}_2\text{O})_6\text{La}(\text{H}_2\text{O})_4\) \((\text{Figure} \ 3)\) | | | \(\text{Ce}_3\text{(MoO}_4\text{)}_2 + \text{hydrazine + oxalic acid, acidified with } 5 \text{ M HClO}_4, 90^\circ \text{C (1 h) (oxalic acid was used for Ln-MB synthesis) (Ce}_3\text{O}_4\text{rM/Mo = 0.26:0.05)}\) | Single-crystal XRD, redox titration, TGA, UV-vis-NIR spectroscopy, IR spectroscopy, BVS, elemental analysis | 45 |
| \([\text{Mo}_{13}\text{O}_{24}\text{H}_{16}\text{O}_{18}]^{\text{2}^-}\) \((\text{Mo}_6\text{O}_2\text{H}_2\text{O})_6\text{La}(\text{H}_2\text{O})_4\) \((\text{Figure} \ 3)\) | | | \(\text{Ce}^{3+} + \text{MoO}_4^{2-} + \text{hydrazine + D-arginine, acidified with } 5 \text{ M HClO}_4, 90^\circ \text{C (1 h) (CeO}_3\text{D-Arg/Mo = 0.1:0.04:1)}\) | Single-crystal XRD, TGA, UV-vis-NIR spectroscopy, IR spectroscopy, ESI-MS-MS spectroscopy, NMR, circular dichroism, BVS, elemental analysis | 42 |
Table 2. continued

| Formula and building blocks given below | Shape | Ln-substitution pattern | Synthetic method, pH, reaction temperature, and molar ratio of the cluster-building reagents | Characterization methods applied | Ref |
|----------------------------------------|-------|------------------------|----------------------------------------------------------------------------------|-------------------------------|-----|
| $\text{A}_{1} \text{[III]} \text{[III]} \text{[III]} \text{[IV]} \text{[IV]} \text{[V]} \text{[V]}$ | circular | $\text{La}^{3+} + \text{MoO}_{4}^{2-} + \text{h} \text{HCl}$ | $\text{pH} = 2.0$ and $\text{h} \text{HCl}$ | Single-crystal XRD, TGA, UV-vis-NIR spectroscopy, IR spectroscopy, ESI-HRMS, elemental analysis | $^a$ |
| $\text{B}_{1} \text{[III]} \text{[III]} \text{[III]} \text{[IV]} \text{[IV]} \text{[V]} \text{[V]}$ | ellippoidal | $\text{La}^{3+} + \text{MoO}_{4}^{2-} + \text{h} \text{HCl}$ | $\text{pH} = 2.0$ and $\text{h} \text{HCl}$ | Single-crystal XRD, TGA, UV-vis-NIR spectroscopy, IR spectroscopy, ESI-HRMS, elemental analysis | $^a$ |
| $\text{C}_{1} \text{[III]} \text{[III]} \text{[III]} \text{[IV]} \text{[IV]} \text{[V]} \text{[V]}$ |  | $\text{La}^{3+} + \text{MoO}_{4}^{2-} + \text{h} \text{HCl}$ | $\text{pH} = 2.0$ and $\text{h} \text{HCl}$ | Single-crystal XRD, TGA, UV-vis-NIR spectroscopy, IR spectroscopy, ESI-HRMS, elemental analysis | $^a$ |

$^a$The lanthanide substitution pattern of the respective wheel-type molybdenum blue is shown graphically. The lanthanide ions on the common inner rim of the wheel are shown in one color (green and pink). Important notes regarding the synthesis and characterization of the respective wheel are also indicated. $^b$As the amount of organic ligand used has a significant influence on the final structure, the molar ratio of the main cluster-building reagents is indicated when an amino acid, peptide, or (di)carboxylic acid was used in the chemical synthesis. $^c$X-ray diffraction.
more details). The elliptical wheel \(\{\text{Mo}_{12}\text{Eu}_4\}\) (Figure 7B) led to the crystallization of the dimer \(\{\text{Mo}_{25}\text{Eu}_8\}\) after 2 weeks.\(^{44}\)

Figure 7. Polyhedral and ball-and-stick representation of the (A) dimer \(\{\text{Mo}_{134}\text{Ce}_6\}\) \(^{42}\) (Figure 3G) and (B) ellipsoidal cluster \(\{\text{Mo}_{32}\text{Eu}_4\}\) \(^{44}\) (Figure 3G) (with a side view shown in (C)). Coloring code: \(\{\text{Mo}_6\}\), yellow; \(\{\text{Mo}_{13}\}\), red; \(\{\text{Mo}_{32}\}\), blue with central \(\{\text{Mo}_3\}\)-unit in cyan; \(\{\text{Ce}_9\}\) and \(\{\text{Eu}_9\}\), green; \(\text{Mo}\), blue spheres; \(\text{Ce}\), green spheres; \(\text{O}\), red spheres.

Remarkably, the self-assembly of the complete Ln-MB ring system (Figure 3G) \(\{\text{Mo}_{25}\text{La}_{10}\}\) (Figure 5G) with the shape of an ellipsoid was obtained in a photochemical reduction (irradiation for 2 days) of an acidified (pH 1.0) \(\text{La}^{III}\), containing solution with the IPOMo \(\{\text{Mo}_{6}\}\) (Figure 1).\(^{40}\) A two-step reaction in which \(\text{Ln}^{III}\) ions react in the vacancies of preformed wheels can also be applied to synthesize ellipsoidal structures.\(^{43}\) This has been realized with the synthesis of the highly Ln-doped complete Ln-MB ring system (Figure 3G) \(\{\text{Mo}_{134}\text{La}_{10}\}\) (Figure 5H).\(^{43}\) By treating the preformed wheel \(\{\text{Mo}_{134}\}\) \(^{35}\) with \(\text{La}^{III}\) ions in a NaCl-solution without acidification at room temperature, crystals of \(\{\text{Mo}_{134}\text{La}_{10}\}\) form at 4 °C after 2 weeks.

The "Japanese rice-ball"-egg-, and ellipsoid-shaped Ln-MBs are all typically synthesized at temperatures 60-90 °C for 30–45min (in a one-pot approach). A fourth type of complete Ln-MB ring systems (Figure 3G), the circular-shaped \(\{\text{Mo}_{25}\text{Ln}_{9}\}\) (Figure 5I) and \(\{\text{Mo}_{25}\text{Ln}_{10}\}\) (Figure 5J) archetypes, can be generated when the synthesis temperature is increased to 100 °C for up to 4 days accompanied by controlled evaporation of the solvent (see Table 2 for more details). This changes the reaction control from kinetic to thermodynamic control (Figure 8), which is the key step in assembling these circular-shaped Ln-MBs.\(^{31}\)

Figure 8. Four basic shapes of Ln-MBs. The "Japanese rice-ball"- (existing in the structural types shown in Figure 3G, I, and J), egg- (existing in the structural type shown in Figure 3G), and ellipsoidal-shaped Ln-MBs (existing in the structural types shown in Figure 3G–J) are all assembled in kinetically controlled syntheses; however, the circular-shaped Ln-MBs (existing in the structural type shown in Figure 3G) are synthesized in thermodynamically controlled reactions. The lanthanides that can be incorporated in the corresponding cluster types are indicated in the center of the shapes.

3.2.2.1. Ellipsoid-Shaped Ln-MBs with Template Clusters.

The incorporation of a template cluster in an Ln-MB can be achieved without hybridizing the scaffold. This was accomplished with the assistance of oxalic acid, a dicarboxylic acid, which acts as a structure-directing ligand without attaching to \(\{\text{Mo}_{6}\}\)-units to form templated complete Ln-MB ring systems (Figure 3H). This was discovered in the synthesis of \(\{\text{Mo}_{134}\text{Ce}_6\}\) (Figure 5K), which was assembled by suspending \(\text{Ce}_2(\text{MoO}_4)_3\) in an acidified aqueous solution, adding oxalic acid and heating to 90 °C for 2 h (see Table 2 for more details).

The crucial variable for the assembly of \(\{\text{Mo}_{134}\text{Ce}_6\}\) appears to be a structural ligand (in this case oxalic acid) in the proper ratio to the molybdate, which does not appear in the final structure but more likely, decomposes during synthesis. This hypothesis is based on the fact that oxalic acid decomposes in an acidic environment at 80 °C.\(^{46}\) Single crystals suitable for X-ray diffraction were obtained at room temperature after 2 weeks.

3.2.2. Structural Description of Purely Inorganic Ln-MBs.

3.2.2.1. "Japanese Rice-Ball", Egg-, Ellipsoid-, and Circular-Shaped Ln-MBs without Template Clusters. The Japanese rice-ball-shaped Ln-MBs are composed of 12 \(\{\text{Mo}_{6}\}\), 6 \(\{\text{Mo}_{5}\}\), 12 \(\{\text{Mo}_{3}\}\), and 6 \(\{\text{Ln}\}\) building units (Table 2). Six \(\{\text{Mo}_{3}\}\) groups are replaced by smaller \(\text{Ln}^{III}\) ions (\(\text{Ln}^{III}\) = \(\text{La}^{III}\), \(\text{Ce}^{III}\), or \(\text{Pr}^{III}\)), according to reported crystal structures.\(^{3,31,32,42–44}\) The average size of the integrated \(\{\text{LnO}_9\}\) in the inner ring (O–Ln-O) is 4.6 Å, while the corner-sharing \(\{\text{Mo}_{3}\}\) units (O–Mo–O–Mo–O) are 7.3 Å, forcing the cluster into a "more contracted" structure exhibiting an irregular ring-shape and a lower symmetry (\(\text{D}_{3d}\)) compared to that of the ideal circular parent structure \(\{\text{Mo}_{154}\}\) \(^{19}\) (\(\text{D}_{3d}\) point group) (Figure 5A). All lanthanide centers on both the upper and lower surfaces of

Table 2. continued

- Thermogravimetric analyses.
- Infrared spectroscopy.
- Bond valence sum.
- Ultraviolet–visible–near-infrared spectroscopy.
- Electrospray ionization–ion mobility spectrometry–mass spectrometry.
- Extended X-ray absorption fine structure spectroscopy.
- Lanthan-nuclear magnetic resonance spectroscopy.
- Building blocks marked with an asterisk (*) indicate an insertion between two \(\{\text{Mo}_{6}\}\) building blocks.
The egg-shaped Ln-MB (\{Mo_{90}La_{10}\}) assembled in a two-step approach consists of 10 {Mo1}, 4 {Mo2}, 10 {Mo8}, 2 {Mo*}, and 6 {Ln} building units (Table 2) and exhibits an inner and outer diameter of \(\sim 14\) and \(\sim 30\) Å, respectively, with four lanthanide ions and one {Mo2} unit coordinated in each inner rim (Figure 5E).41 In comparison to the "Japanese rice-ball"-shaped scaffolds \{Mo_{120}La_{20}\} (\(La^{3+} = La^{3+}, Ce^{3+}, or Pr^{3+}\)) (Figure 5D), which were synthesized in a one-pot reaction, the "egg" \{Mo_{90}La_{10}\} is smaller and has a less uniform lanthanide pattern.42−45 

With NdIII and SmIII, which belong to the later lanthanides and therefore have a smaller radius, the scaffold \{Mo_{90}Ln_{10}\} (\(Ln^{3+} = Nd^{3+} or Sm^{3+}\)) with a charge of minus one was obtained (Figure 5I) in which one {Mo2} building block unit is present in the wheel. \{Mo_{90}Ln_{10}\} is composed of 10 {Mo1}, 1 {Mo2}, 10 {Mo8}, and 9 {Ln}-building units (Table 2). The replacement of all inner-rim {Mo2}-units could not be accomplished in \{Mo_{90}La_{10}\} as the wheel contraction would become too large (radii: SmIII < NdIII < PrIII < CeIII < LaIII < \(Mo^{3+}\)). The lanthanide substitution pattern of \{Mo_{90}La_{10}\} and \{Mo_{90}Ln_{10}\} does not induce a structural deformation compared to the original cluster \{Mo_{120}\}, in contrast to \{Mo_{90}Pr_{10}\}.45−47 \{Mo_{100Ce_{6}}\},42 \{Mo_{120}La_{6}\},41 and \{Mo_{120}Eu_{6}\} are likely to be the most versatile ones, which explain the variety.43

The replacement of all inner-rim {Mo2}-units could not be accomplished in \{Mo_{120}Ce_{6}\} as the wheel contraction would become too large (radii: SmIII < NdIII < PrIII < CeIII < LaIII < \(Mo^{3+}\)). The lanthanide substitution pattern of \{Mo_{120}La_{6}\} and \{Mo_{120}Ln_{6}\} does not induce a structural deformation compared to the original cluster \{Mo_{120}\}, in contrast to \{Mo_{90}Pr_{10}\}.45−47 \{Mo_{100Ce_{6}}\},42 \{Mo_{120}La_{6}\},41 and \{Mo_{120}Eu_{6}\} are likely to be the most versatile ones, which explain the variety.43

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relatively high number of Ce\(^{III}\) containing Ln-MBs that have been reported (20 as of August 2021; see Table 2).

Pr\(^{III}\) ions, the third-longest lanthanides, are currently only known in "Japanese rice-ball"- and circular-shaped Ln-MBs.\(^{25,31}\) The latter shape could hitherto only be assembled thermodynamically. The smaller lanthanide ions Nd\(^{III}\) and Sm\(^{III}\) have so far only been incorporated into circular-shaped Ln-MBs in thermodynamically controlled Ln-MB synthesis approaches.\(^{31}\) Due to their ion size, Nd\(^{III}\) and Sm\(^{III}\) seem reluctant to be incorporated into "Japanese rice-balls" or "eggs", which are typically formed in kinetically controlled synthesis approaches. Surprisingly, it is feasible to incorporate Eu\(^{III}\), the smallest lanthanide ion so far (which one would only expect to be incorporated thermodynamically into a circular-shaped Ln-MB ring due its size) into an ellipsoidal Ln-MB through a kinetically controlled reaction.\(^{14}\) However, the incorporation of Eu\(^{III}\) into an ellipsoidal Ln-MB prevented ring closure of the inner rim (Figure 7C), which is most likely owing to its relatively small size. An open inner rim in an Ln-MB is only known for Eu\(^{III}\) ions.

The size of a lanthanide determines whether it can replace a {Mo\(_3\)} unit in MBs. The smallest lanthanide ion integrated in the inner rim of MBs and identified by X-ray structure analysis is Eu\(^{III}\).\(^{44}\) This suggests that the slightly larger lanthanide ions Nd\(^{III}\) and Sm\(^{III}\) can likewise serve as building units in kinetically controlled self-assemblies of Ln-MBs. The lanthanide ions from Gd\(^{III}\) to Lu\(^{III}\) appear to be too small to function as building units in Ln-MBs.

When incorporated in MBs lanthanide ions generate significant deformations relative to the circular parent cluster \{Mo\(_{154}\}\), resulting in the main basic shapes ("Japanese rice-ball", "egg", and ellipsoidal forms) (Figure 8). However, if the degree of replacement of the \{Mo\(_3\}\) units by Ln\(^{III}\) ions is very high (9 or 10 Ln\(^{III}\) ions per molecule), the resulting shape of the Ln-MB is nearly identical to that of \{Mo\(_{154}\}\). In order to visualize the effect of the integration site of built-in lanthanides generating deformations and to compare them directly, the cavity skeletons of the circular-shaped \{Mo\(_{90}Ln_{10}\}\) (Figure 3G), the egg-shaped \{Mo\(_{100}Ce_{6}\}\) and \{Mo\(_{96}La_{8}\}\), as well as the ellipsoidal \{Mo\(_{124}Eu_{4}\}\) are illustrated in Figure 9. Due to the even distribution of both the Pr\(^{III}\) ions within the inner surface of \{Mo\(_{124}Pr_{6}\}\) and the Ln\(^{III}\) ions in \{Mo\(_{90}Ln_{10}\}\) as well as the antipodal location of the Eu\(^{III}\) ions in \{Mo\(_{124}Eu_{4}\}\), these wheels show a more regular structure than the remaining wheels in Figure 9. In contrast, the uneven distribution of three Ce\(^{IV}\) ions in the inner rim of \{Mo\(_{105}Ce_{5}\}\) forms a triangle in which two Ce\(^{IV}\) ions are located adjacent to each other (Ce–O–Mo–O–Ce) while the third ion is positioned at the opposite side of the cavity. Also, the uneven lanthanide ion distribution in \{Mo\(_{90}La_{8}\}\), where all four La\(^{III}\) ions are asymmetrically positioned and connected to each other via O–Mo–O bridges results in an irregular rather than regular structure. Depending on the nature of the lanthanide, a specific replacement of \{Mo\(_3\}\)-units can be accomplished, allowing for curvature tuning.

### 3.2.3. Synthesis of Hybridized Ln-MBs

As of August 2021, there are 11 crystal structures of hybridized complete Ln-MB ring systems (Figure 3I) and 7 templated and hybridized complete Ln-MB ring systems (Figure 3J) reported (Table 2).

Wheel-shaped clusters have a chaotropic character, due to the large molecular surface with a relatively low charge density.
Scheme 1. Schematic Representation of the Stereoselective Synthesis of Chiral Ln-MBs through the Use of Chiral Amino Acids

“(A) d-Histidine results in the Δ-{Mo124Ce4(d-histidine)6} enantiomer (Figure 3I), whereas n-tryptophan and n-arginine yield the Λ-{Mo124Ce4(n-tryptophan)4} and Λ-{Mo124Ce4(n-arginine)4} (Figure 3I), enantiomers.24 (B) l-Histidine results in the Λ-{Mo124Ce4(l-histidine)6} enantiomer (Figure 3I), whereas l-tryptophan and l-arginine yield the Δ-{Mo124Ce4(l-tryptophan)4} and Δ-{Mo124Ce4(l-arginine)4} (Figure 3I), enantiomers.25 (C) Without the addition of a chiral amino acid, {Mo124Ce4}24 (Figure 3G) [Reprinted from ref 24. Copyright 2018 American Chemical Society.] is obtained as a racemate. Coloring code: {MoO6}, yellow; {Mo2O11}, red; {Mo8O35}, blue with central {MoO7}-unit in cyan; {CeO9}, green. Λ and Δ represent right-handed and left-handed propeller, respectively and organic ligands in the final complex are omitted for clarity.

Of Ln-MBs prompted researchers to investigate enantiopure oligopeptides, which extended the number of hybridized complete Ln-MB ring systems (Figure 3I) by six structures.39 In a one-pot synthesis, an acidified aqueous solution containing all main starting materials (CeCl3, Na2MoO4 and the respective histidine-terminated oligopeptide) (see Table 2 for quantity ratios) was reduced and heated at 90 °C for 2 h. Crystals of {Mo124Ce4(oligopeptide)3}, {Mo124Ce4(oligopeptide)}2 and {Mo124Ce4(oligopeptide)3} were obtained at room temperature after two to 8 weeks.39 The reaction solution was occasionally heated for a longer period of time (2 h) than in the case for the hybridization of Ln-MBs with proteinogenic amino acids (1 h).

3.2.3. Ln-MBs with Template Clusters. To date, Ln-MBs with template clusters have only been assembled with the assistance of organic ligands.23,24,38 It is quite difficult to ascertain which ligand will be able to enclose a polynucleotidetate as a template. The only established requirement for the ligand (aside from the carbonyl group for grafting to the {Mo3} units) is the presence of a nitrogen-containing pendant group (e.g., histidine, ornithine). In an acidic environment, this nitrogen-containing group gets protonated and could therefore stabilize template clusters within the Ln-MB electrostatically and through hydrogen bonds. The use of chiral proteinogenic amino acids not only enables the organic hybridization of the inner rim of Ln-MBs, resulting in nanosized chiral Ln-MBs, but also the synthesis of chiral templated and hybridized complete Ln-MB ring systems (Figure 3J). For the self-assembly of Δ-{Mo8}@{Mo124Ce4(histidine)6} and Λ-{Mo8}@{Mo124Ce4(histidine)6}, an acidified aqueous solution containing all of the major starting materials (CeCl3, Na2MoO4 and l-histidine (for the Λ-enantiomer) or d-histidine (for the Δ-enantiomer) was reduced and heated at 90 °C for 1 h leading to suitable crystals for X-ray diffraction after 1 week at room temperature.24 Besides the proteinogenic amino acids histidine,24 arginine,24 and tryptophan,24 the nonproteinogenic amino acid l-ornithine (Orn)24 also has a template-directing effect in the assembly of Ln-MBs. l-Ornithine enabled the synthesis of the “Japanese rice-ball”-shaped (almost ellipsoidal shaped) α-{Mo3}@{Mo124Ce4(Orn)6} (Figure 3J), the elliptically shaped {Mo17}@{Mo150Ce4(Orn)6} (Figure 6G), and β-{PMo12}@{Mo150Ce2(Orn)6} (Figure 6H). The critical synthetic variable in the self-assembly of these three templated and hybridized complete Ln-MB ring systems (Figure 3J) is the concentration of the lanthanide ion, ligand, and molybdate. For {Mo17}@{Mo150Ce2(Orn)6} and {Mo17}@{Mo150Ce2(Orn)6}, the concentrations of the ornithine and molybdate were ≈3-fold and that of the CeIII ions ≈1.5-fold higher than for the formation of α-{Mo3}@{Mo124Ce4(Orn)6} (see Table 2 for details). In α-{Mo3}@{Mo124Ce4(Orn)6} and {Mo17}@{Mo150Ce2(Orn)6}, the templated polyoxomolybdate was formed in situ, whereby the enclosed template in β-{PMo12}@{Mo150Ce2(Orn)6} was a separately added preformed Keggin.38 The ornithine and molybdate concentrations were significantly increased most likely to enhance the ligand’s template-directing effect and to enable both the encapsulation and stabilization of a larger negatively charged polyoxomolybdate cluster.

3.2.4. Structural Description of Hybridized Ln-MBs. Remarkably, organic hybridization of {Mo3} building blocks in Ln-MB assemblies predominantly yields ellipsoidal-shaped clusters. The hybridization of {Mo3} units in the cavity of ellipsoidal Ln-MBs seems to be less restricted by steric hindrances as in “Japanese rice-balls”, “eggs”, and circular-shaped clusters.

3.2.4.1. Structural Description of Ln-MBs without Template Clusters. All Ln-MBs without template clusters obtained through the hybridization with proteinogenic amino acids are hybridized complete Ln-MB ring systems (Figure 3I) sharing the same elliptical skeletal {Mo124Ce4}24 with either four arginines24 (Figure 6D) or six tryptophans24 (Figure 6E) grafted to the cluster. The skeletal {Mo124Ce4} consists of 12 {Mo1}, 8 {Mo2}, 12 {Mo3} and 4 {Ce} building units (Table 2) and exhibits an inner and outer ring diameter of approximately 12 and 31 Å, respectively (Figure 10). The four CeIV ions built into the scaffold are symmetrically located on the upper and lower rims of the wheel, and the MB-functionactivizing amino acids arginine and tryptophan are coordinated to the organically chemically modifiable24

Figure 10. Polyhedral representation of the {Mo124Ce4}24 scaffold of the hybridized, elliptical Ln-MB frameworks (Figure 3I). Coloring code: {MoO6}, yellow; {MoO11}, red; {MoO33}, blue with central {MoO}-unit in cyan; {CeO9}, green.
{Mo₃}-type building blocks via their carboxylate groups (Figure 10).

Different enantiomers are generated depending on the concentration, type, and configuration of the used amino acid. D- and L-amino acids are capable of producing the Δ and Λ enantiomer in the pure form, the configuration of the amino acid not favoring any specific enantiomer (Scheme 1). In the absence of D- or L-amino acids, only the \{Mo₁₂₂Ce₅\} (Scheme 1) wheel is obtained as a racemate, which is structurally similar to \{Mo₁₄₂Ce₁\} \(^{34}\) (slightly compressed in contrast to \{Mo₁₃₂Ce₄\}). Incorporating histidine-terminated oligopeptides into Ln-MB systems revealed that the length and sequence of the peptides markedly influence the shape, Ln-substitution pattern, and stereochemistry of the resulting Ln-MBs. \(^{35}\) For instance, if glycine (Scheme 2A) is replaced with serine/alanine (Scheme 2B) in a histidine-based dipeptide, an ellipsoidal Δ-enantiomer in the pure form, the conformation of the amino acid not favoring any specific enantiomer (Scheme 1). In the absence of D- or L-amino acids, only the \{Mo₁₂₂Ce₅\} (Scheme 1) wheel is obtained as a racemate, which is structurally similar to \{Mo₁₄₂Ce₁\} \(^{34}\) (slightly compressed in contrast to \{Mo₁₃₂Ce₄\}). Incorporating histidine-terminated oligopeptides into Ln-MB systems revealed that the length and sequence of the peptides markedly influence the shape, Ln-substitution pattern, and stereochemistry of the resulting Ln-MBs. \(^{35}\) For instance, if glycine (Scheme 2A) is replaced with serine/alanine (Scheme 2B).

**Scheme 2. Schematic representation of the histidine-terminated oligopeptide-controlled synthesis of Ln-MBs\(^ {39a}\)**

“The oligopeptides used were (A) H-Gly-L-His-OH (GH) (Figure 3I), (B) H-L-Ala-L-His-OH (AH) (Figure 3I) and H-L-Ser-L-His-OH (SH) (Figure 3I), (C) H-Gly-Gly-L-His-OH (G₃H) (Figure 3I), and (D) H-Gly-Gly-Gly-L-His-OH (G₄H) (Figure 3I). The toroidal shapes of the wheels are displayed in blue. The lanthanide ions (Ce\(^{3+}\)) are shown in green; carbon, black; and nitrogen, purple.

The oligopeptides used were (A) H-Gly-L-His-OH (GH) (Figure 3I), (B) H-L-Ala-L-His-OH (AH) (Figure 3I) and H-L-Ser-L-His-OH (SH) (Figure 3I), (C) H-Gly-Gly-L-His-OH (G₃H) (Figure 3I), and (D) H-Gly-Gly-Gly-L-His-OH (G₄H) (Figure 3I). The toroidal shapes of the wheels are displayed in blue. The lanthanide ions (Ce\(^{3+}\)) are shown in green; carbon, black; and nitrogen, purple.

The dipeptides H-Gly-L-His-OH (GH), H-L-Ala-L-His-OH (AH), and H-L-Ser-L-His-OH (SH) are attached to the ellipsoidal \{Mo₁₃₂Ce₄\} scaffold, where GH mediates the self-assembly of the Δ-{Mo₁₃₂Ce₄} enantiomer, and AH and SH that of the Λ-{Mo₁₃₂Ce₄} enantiomer. Thus, the helicity of the chiral \{Mo₁₃₂Ce₄\}-Ln-MBs depends on the N-terminal rather than the C-terminal residue of the oligopeptide. This allows an elegant switch from the Δ to the Λ enantiomer by just altering the amino acid at the N-terminus without changing the stereochemistry of the entire peptide. The tripeptide H-Gly-Gly-L-His-OH mediates the self-assembly of the wheel-skeleton \{Mo₁₂₂Ce₅\} \(^{39}\) and binds to it three times (Scheme 2C). \{Mo₁₃₂Ce₄\} is an unprecedented Ln-MB scaffold containing two differently shaped cavities in the same Ln-MB ("Japanese rice-ball" and elliptical-shaped cavity, Figure 11) and an odd number of lanthanide ions. The pentapeptide H-Gly-Gly-Gly-Gly-L-His-OH and hexapeptide H-Gly-Gly-Gly-Gly-Gly-L-His-OH mediate racemic Ln-MB mixtures, comprising the asymmetric ellipsoidal scaffold \{Mo₁₂₆Ce₄\} \(^{39}\) (Scheme 2D). Since the oligopeptides differ in length and sequence, they form different numbers of hydrogen bonds when introduced into Ln-MBs, resulting in a variety of distinct structural characteristics.

**3.2.4.2. Structural Description of Ln-MBs with Template Clusters.** Notably, only the functionalization of \{Mo₁₂₂Ce₅\} scaffold with the proteinogenic amino acids L-/D-histidine transforms the nanocavity into a reaction tube, which can be described as a confined molecular reaction vessel, wherein the in situ assembled, chiral metal cluster \{Mo₆\} (structurally related to γ-[Mo₃O₁₆] \(^{48}\) is embedded. \(^{24}\) The inclusion of the chiral cluster \{Mo₆\} into \{Mo₁₂₂Ce₅\} scaffold results in the formation of the supramolecular guest@host assembly \{Mo₆\} @ \{Mo₁₂₂Ce₅(histidine)\} \(^{24}\) (Figure 6). The orientation of the introduced \{Mo₆\} fragment within \{Mo₁₂₂Ce₅(histidine)\} depends strongly on the spatial arrangement of the coordinating histidine ligands and their hydrogen bonds formed with \{Mo₆\}. \(^{24}\) The crystal structure analysis strongly indicates that all six histidine ligands must be protonated in order to minimize the repulsive electrostatic forces between the negatively charged assemblies (nanoring and \{Mo₆\} cluster) and therefore increase system stability. Moreover, the orientation of the introduced \{Mo₆\} fragment within \{Mo₁₂₂Ce₅(histidine)\} depends strongly on the spatial arrangement of the coordinating histidine ligands and their hydrogen bonds formed with \{Mo₆\}. \(^{24}\)

The nonproteinogenic amino acid L-ornithine coordinates to \{Mo₆\} units during the self-assembly processes and acts as a structure-directing ligand, triggering the formation of various guest@host architectures. \(^{38}\) Three cerium-containing nanoclusters were obtained using L-ornithine: the "Japanese rice-ball"-shaped α-{Mo₆}[@Mo₁₂₂Ce₅(Or₉)] \(^{38}\) (Figure 6F), the elliptically shaped \{Mo₁₄₂Ce₁[@Mo₁₃₀Ce₅(Or₉)] \(^{38}\) (Figure 6G, and the elliptically β-PMo₁₂[@Mo₁₀₂Ce₅(Or₉)] \(^{38}\) (Figure 6H). In all three nanowheels, six L-ornithine ligands are attached to the inner rim. In addition, the negatively charged polyoxomolybdate template cluster in the cavity of each nanowheel is stabilized by electrostatic forces and hydrogen bonds. \{Mo₁₂₂Ce₅(Or₉)\} encapsulates the in situ formed...
octamolybdate-type anion \(\alpha\{\text{Mo}_{12}\text{O}_{40}\}\text{)}\text{\(^{4-}\) (\(\alpha\{\text{Mo}_7\) \(\text{O}_2\})\text{\(^{1-}\) (Figure 6F), whereas the nanoring \(\{\text{Mo}_{17}\text{Ce}_{2}(\text{Orn})_6\}\) can trap either the in situ formed \(\text{Mo}_{17}\text{O}_{52}(\text{H}_2\text{O})_{10}\)\text{\(^{2-}\) (\(\text{Mo}_{17}\) \(\text{O}_2\))\text{\(^{1-}\) (Figure 6G) or the separately added Keggin-type anion \(\beta\{\text{PMo}_{12}\text{O}_{40}\}\text{\(^{3-}\) (\(\beta\{\text{PMo}_{12}\) \(\text{O}_2\))\text{\(^{2-}\) (Figure 6H).\) So far, \(\{\text{Mo}_{17}\) \(\text{O}_2\)\) has only been isolated upper and lower rim, located opposite to each other (Figure 6F), whereas the nanoring \(\{\text{Mo}_{150}\text{Ce}_{2}(\text{Orn})_6\}\) can trap either the in situ formed \(\text{Mo}_{17}\text{O}_{52}(\text{H}_2\text{O})_{10}\)\text{\(^{2-}\) (\(\text{Mo}_7\) \(\text{O}_2\))\text{\(^{1-}\) (Figure 6G) or the separately added Keggin-type anion \(\beta\{\text{PMo}_{12}\text{O}_{40}\}\text{\(^{3-}\) (\(\beta\\text{ PMo}_{12}\) \(\text{O}_2\))\text{\(^{2-}\) (Figure 6H).\) So far, \(\{\text{Mo}_{17}\) \(\text{O}_2\)\) has only been isolated upper and lower rim, located opposite to each other (Figure 6F).

4. CHARACTERIZATION OF Ln-MBs IN THE SOLID STATE

4.1. Single Crystal X-ray Diffraction

First and foremost, for new Ln-MB-based nanoclusters, it is essential to obtain suitable crystals for single crystal X-ray structure analysis to determine their three-dimensional shape and size in the solid state. The size, stability, scattering power, and quality of the crystal are critical for a single crystal X-ray structure analysis. Since Ln-MBs are composed of heavy atoms and thus produce a greater diffraction pattern than molecules containing just light atoms, the crystal to be examined can be significantly smaller than that of organic compounds. Crystals of Ln-MB-based nanoclusters are typically measured around 200 K and below rather than at room temperature. To generate Ln-MB crystals suitable for X-ray diffraction, the crystallization process may need to be slowed down (for example, by allowing the solvent to evaporate more slowly or by lowering the crystallization temperature), or the synthesis parameters have to be adjusted. The majority of Ln-MB synthesis optimizations are based on the “trial and error” concept, with the aim of finding the optimal starting component ratio.\textsuperscript{32,37}

4.2. FTIR and Raman Spectroscopy, Powder Diffractometry, and TGA

Infrared spectroscopy displays characteristic bands, which appear \(<1100\text{ cm}^{-1}\) (Mo=O vibrations are located at higher wave numbers than Mo–O–Mo) and confirms only the presence of a polyoxometalate scaffold, whereas Raman spectroscopy shows characteristic bands for wheel-type scaffolds.\textsuperscript{39} Sharp bands are typically observed in the Raman spectra of MB/Ln-MBs at both low and high wavenumbers, which can be attributed to vibrations of \(\{\text{Mo}^{\text{VII}}\}_m\text{O}\) \((m = 2,3,4)\) fragments (Figure 12) with the band at around 800 cm\(^{-1}\) being a symmetric stretching vibration of the equatorial \(\{\text{Mo}^{\text{VII}}\}_m\text{O}_{\gamma}\) \(\mu_\gamma\text{-O}\) fragment.\textsuperscript{19,50} Raman spectroscopy also allows investigations of MBs/Ln-MBs in aqueous solution, since water molecules are only weakly Raman-active.

5. STABILITY AND SPECIATION STUDIES OF Ln-MBs IN SOLUTION

The stability and speciation of Ln-MBs must first be studied when dissolved before they can be used as possible molecules for applications in solution. How can the integrity of Ln-MBs be confirmed? How are aggregates, such as the spontaneous self-assembly of homogeneous, hollow spherical “blackberry” structures,\textsuperscript{30} identified? Which spectroscopic methods are relevant for the investigation of chirality in Ln-MBs? Is it feasible to show that the organic–inorganic hybridization in solution is preserved? Is it possible to determine the charge of these Ln-MB wheels at different pH values?

Among all the various investigation techniques for stability and speciation studies in solution, such as Raman-spectroscopy, ESI-MS and ES-IMS-MS, or UV–vis-Nir, small-angle X-ray scattering (SAXS) stands out when it comes to exploring nanoscopic structures (usually 1–100 nm) in solution.\textsuperscript{51} The utilization of SAXS, a highly powerful, nondestructive technique enables the determination of size, shape, and interactions of nanoscopic POMs and additionally allows for a variety of in situ analyses. Investigations of solution-phase speciation under various conditions (pH, conc., type of counterions, aged solutions, etc.) can be conducted in order to obtain information about their morphology and assembly when dissolved (e.g., is the POM oligomerized or present as a monomer? Is an elliptically stacked monomeric arrangement or rather a shifted face-to-face dimeric configuration preferred)?\textsuperscript{52,53} Stability measurements can be performed to determine whether POMs are still intact under certain conditions (e.g., in catalytic processes).\textsuperscript{51} Fundamental understanding of reaction mechanisms\textsuperscript{51} can be provided and possible preferences for particular organic compounds and whether they accelerate the cluster formation of Ln-MBs can be examined.\textsuperscript{17} Furthermore, the feasibility of registering the start and maximum of nanosized cluster formation by using SAXS, enables the determination of critical reaction parameters for their synthesis.\textsuperscript{54} SAXS studies on Ln-MB systems have not yet been conducted.

5.1. ES-IMS-MS, Redox Titration, and UV–Vis-Nir

Electrospray ionization-ion mobility spectrometry-mass spectrometry (ESI-IMS-MS) is an excellent method for investigating the speciation of gigantic MBs in solution/gas phase. ESI-IMS-MS is superior to ESI-MS for the following reason: MB/Ln-MB building blocks generate multiple species with fairly similar mass-to-charge ratios (\(m/z\)) in very different sizes/structures, resulting in overlapping envelopes in the mass spectra.\textsuperscript{55} Figure 13A shows an ESI-IMS-MS spectrum of the dimer \(\{\text{Mo}_{100}\text{Ce}_{6}\}_2\) and the dominating monomer \(\{\text{Mo}_{100}\text{Ce}_{6}\}\).\textsuperscript{45} The spectrum of the monomer \(\{\text{Mo}_{100}\text{Ce}_{6}\}\) was very similar to that of \(\{\text{Mo}_{100}\text{Ce}_{6}\}_2\) with the significant difference that no peak for the dimer was observed. Redox titration is used to determine the number of Mo\textsuperscript{V} centers in the cluster (e.g., with a Ce\textsuperscript{III} titration).\textsuperscript{55} However,
sufficient solubility of the compound is required for a redox titration; otherwise, the Mo\(^V\) centers must be identified using bond valence sum calculations based on the crystal structure.

Solution stability of MB/Ln-MB-based nanoclusters can be monitored with UV–vis-Nir, as they show characteristic absorption bands from 600 to 1100 nm, which are attributed to the intervalence charge transfer between Mo\(^{VI}\) and Mo\(^{V}\). Absorption bands from 600 to 1100 nm, which are attributed to the intervalence charge transfer between Mo\(^{VI}\) and Mo\(^{V}\). Additional absorption bands were observed around 274 nm originating from histidine. A red shift of about 20 nm relative to unattached histidine strongly suggests coordination on the \{Mo\(_{124}\)Ce\(_4\}\) scaffold and, as a result, a restriction of the histidine’s free rotation.

\(^1\)H NMR spectroscopy is well suited for examining hybridized inorganic–organic Ln-MBs for organic attachment, provided the molecule to be examined is sufficiently soluble in deuterated solvents. The \(^1\)H NMR analysis of \{Mo\(_8\)@Mo\(_{124}\)Ce\(_4\)(histidine)\(_6\)\} (Figure 13C) displays the characteristic signals for histidine, with the difference that the signals obtained are both broader and upfield- and downfield-shifted when compared to unattached histidine. The broadening of the signals as well as their upfield and downfield shift are probably caused by the shielding effect from the 24-electron reduced \{Mo\(_{124}\)Ce\(_4\)\}.

5.3. “Blackberry”, Encapsulation Experiments, SLS, and DLS

Charged MB/Ln-MBs exhibit a unique behavior in solution, assembling into hollow, spherical blackberry-like aggregates. The interactions involved for their formation are counterion-mediated electrostatic attractions, van der Waals forces, and hydrogen bonding. A critical variable for the assembly of “blackberries” is the pH, which determines their size. Typically, the lower the pH, the smaller the overall negative charge (= weaker repulsive electrostatic interactions between adjacent anion clusters) and the larger the size of the “blackberry.” Encapsulation tests using the positively charged cetrimonium bromide (CTAB) surfactant allow for the investigation of the total Ln-MB charge. For \{Mo\(_{90}\)Ce\(_{10}\)\}, the charge could be determined at various pH levels (pH 0.5–2), indicating that \{Mo\(_{90}\)Ce\(_{10}\)\} exists in solution as a decaanionic cluster above pH 2.19 The concentration of the cations has the opposite effect, as demonstrated clearly by the increasing addition of divalent Ba\(^{II}\) cations, which exhibits enhanced attraction of macroion and counterion. The higher the concentration of Ba\(^{II}\) ions is, the larger the “blackberry” becomes, confirming that its assembly is a charge-regulated process. The first “blackberry” assembly of an Ln-MB was confirmed for \{Mo\(_{90}\)Ce\(_{10}\)\} (at pH 1 in the presence of Ba\(^{II}\)) and verified by static light scattering (SLS). The hydrodynamic radius of \{Mo\(_{90}\)Ce\(_{10}\)\} was identified by the use of dynamic light scattering (DLS). POM superstructures, as well as structural changes induced by pH, counterion concentration, etc., can be observed and studied using SLS and DLS. Both light scattering techniques are able to determine dynamic properties of small POMS, such as the Keggin archetype, and gigantic assemblies.

6. The Influence of Lanthanides on the Redox Characteristics of Ln-MB Nanoclusters

The partial reduction of Mo\(^{VI}\)-solutions results in Mo-scaffolds with a different degree of reduction (given in % in Table 3), which is calculated by dividing the number of Mo\(^V\) centers by the total number of Mo centers in the wheel (Table 3). When lanthanides are incorporated into MB scaffolds, the resulting Ln-MBs demonstrate almost the same degree of reduction as Ln-free MBs (Table 3) with Ln-MBs containing a slightly smaller number of Mo\(^V\) centers within their scaffold compared with Ln-free MBs, as they are typically made up of a smaller number of building blocks.

Mo\(^V\) centers, reflecting the scaffold’s degree of reduction, are integrated in Ln-MBs as Mo\(^V\)-O–Mo\(^{VI}\) bridges, exactly as in

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Figure 13. (A) ESI-IMS-MS of a solution containing the dimer \{Mo\(_{100}\)Ce\(_6\)\} with the dominating monomer \{Mo\(_{100}\)Ce\(_6\)\}. Reprinted from ref 42. Copyright 2014 American Chemical Society. (B) CD spectra of Δ-{Mo\(_8\)@Mo\(_{124}\)Ce\(_4\)(histidine)\(_6\)}, A-{Mo\(_8\)@Mo\(_{124}\)Ce\(_4\)(histidine)\(_6\)}, l-histidine, and D-histidine. Reprinted from ref 24. Copyright 2019, American Chemical Society. (C) \(^1\)H NMR spectra of \{Mo\(_8\)@Mo\(_{124}\)Ce\(_4\)(histidine)\(_6\)\} (black) and l-histidine (blue). Reprinted from ref 24. Copyright 2019, American Chemical Society.
MBs, and present in a considerable number within the Ln-MB scaffold. These MoV−O−MoVI structural motifs are considered to be the catalytic centers, as proposed mechanistically in the {Mo154}−catalyzed oxidation of cyclohexane to cyclohexanol by Con et al.58

A correlation has been established between the degree of reduction of gigantic Ln-free POMs and the yield of products in photoredox mediated organic coupling reactions. In the synthesis of 1,4-diphenyl substituted butane-1,4-dione, a yield of 87% was achieved with the keplerate {Mo152} (degree of reduction 45%) and a yield of 78% with the MB {Mo154} (degree of reduction 18%) as a catalyst.59 As Ln-MBs and MBs exhibit a comparable degree of reduction (Table 3), similar redox characteristics are to be expected.

7. POSSIBLE APPLICATIONS AND RELEVANCE OF Ln-MBs

7.1. Homogeneous and Heterogeneous Catalysis

The development of applications in materials using giant POMs, e.g., keplerates and [Mo154−x−] is described in the recently published book chapter by Long and Cronin.60 While several intriguing examples of MBs as homogeneous and heterogeneous catalysts have been reported,58,59,61–83 merely one report about a Ln-MB showing catalytic behavior exists.24 However, based on their structure and stability in solution, potential applications can be foreseen. In homogeneous catalysis organosoluble POMs bearing lanthanides (e.g., the Wells–Dawson archetype) can exhibit chemoselectivity in organic reactions, such as in aldol, Diels–Alder–Mannich–, and Mukaiyama-type reactions.27,28 Chiral hybridized complete Ln-MB ring systems can enclose nanosized guest molecules within their cavity,24 and as they contain lanthanides, which are Lewis acidic cations, these LnIII ions might therefore enable the performance of (selective) catalytic reactions on entrapped substrates in their "Japanese rice-ball"-, circle-, egg-, and ellipse-shaped (see Figure 8) catalytic performance and function as nanoveossels for homogeneous catalysis in organic media. For their use in organic homogeneous catalysis, the Ln-MBs must be readily soluble in organic media. To increase the hydrophobicity of Ln-MBs, appropriate surface reagents can be used, which has already been implemented for the circular-shaped, lanthanide-free MB {Mo176}.64 Using the amphiphilic complexing agents didodecyldimethylammonium (DDMA) bromide and dioctadecyl(dimethyl)ammonium (DOMA) bromide, the hydrophobic aggregates Ln-MB-DDMA/DOMA could be obtained in which the rigid, hydrophilic, inorganic nanocluster would have been encapsulated by a flexible hydrophobic outer shell.

In heterogeneous catalysis, MB nanorings have been shown to be effective oxidation catalysts65 and that they can be used as a solid solution.66 The chirality of Ln-MBs, caused with the help of LnIII ions,24 could be exploited in heterogeneous catalysis, for instance, in asymmetric oxidation reactions. A Ln-MB-based solid solution for various oxidation reactions could be obtained as follows: (1) Ln-MBs could be embedded in silica using a sol−gel process as it has already been realized for the circular-shaped lanthanide-free MB {Mo176} (Figure 14).65

Figure 14. Schematic representation of Ln-MB-silica. (A) Embedding of {Mo176} in silica yields the hybrid material {Mo176}-silica.65 (B) {Mo176}-silica hybrid material with additional micropores created by PEG.65

(2) Amorphous, optically clear objects (monoliths) of different sizes and shapes could be generated, creating a porous system that contains nanostructures with access to their inner surface (Figure 14). The synthesized Ln-MB-silica could behave like a solid solution because the incorporated POMs would be homogeneously distributed in the silica matrix. Additional micropores of different sizes could be created by adding polyethylene glycol (PEG) without causing a macroscopic phase separation or the release/decomposition of the POMs. As the pores introduced by PEG would be much smaller than the cavities of the POMs, molecules could diffuse into the POM, without the POM being released. These POM hybrids could be described as precisely defined, monodisperse reaction systems in which the amount of molybdenum remains constant regardless of the chemical process.

7.2. Molecular Recognition

Molecular recognitions are among the most significant processes of biological and supramolecular systems.67 Chiral Ln-MBs are supramolecular wheels with unique functional sites inside a well-defined cavity and, therefore, should be capable of

### Table 3. All MB/Ln-MB-Based Nanoclusters with Approximate Degree of Reduction (Number (n) of Accepted e− Reflects the Number of MoV Centers in the Entire Wheel) Determined through Redox Titration

| MB/Ln-MB scaffold | n of accepted e− | % of reduction | examples of reported applications | ref |
|--------------------|-----------------|----------------|----------------------------------|-----|
| {Mo$_{154}$}C{Mo$_{150}$} | 20 | 0 in {Mo$_{154}$} and 13 in {Mo$_{150}$} | 21 |
| {Mo$_{154}$} | 24 | 17 | 20 |
| {Mo$_{150}$} | 28 | 18 | cathode material | 19, 60 |
| {Mo$_{154}$} | 32 | 18 | 1 |
| {Mo$_{154}$} (ornithine)$_{2}$ | 32 | 18 | 35 |
| {Mo$_{150}$} | 28 | 19 | 21 |
| {Mo$_{154}$} (acetate)$_{4}$ | 28 | 20 | sensor technology | 20, 60 |
| {Mo$_{154}$}C{Mo$_{176}$} | 67 | 67 in {Mo$_{154}$} and 18 in {Mo$_{176}$} | 22 |
| {Mo$_{132}$}La$_{4}$ | 24 | 18 | 44 |
| {Mo$_{120}$}Ce$_{4}$ | 24 | 18 | 35 |
| {Mo$_{120}$}La$_{4}$ | 28 | 18 | 31 |
| {Mo$_{120}$}Ce$_{4}$ | 20 | 19 | 32 |
| {Mo$_{120}$}Pr$_{4}$ | 20 | 19 | 33 |
| {Mo$_{120}$}Ce$_{4}$ | 24 | 19 | 32 |
| {Mo$_{120}$}Ce$_{4}$ | 24 | 19 | 41 |
| {Mo$_{120}$}Ce$_{4}$ | 24 | 19 | 31 |
| {Mo$_{120}$}La$_{4}$ | 28 | 19 | 34 |
| {Mo$_{90}$}Ce$_{10}$ | 20 | 20 | 30 |

The structures are arranged in increasing degree of reduction of the MBs/LnMBs. The first half of the table contains MBs, and the second half contains Ln-MBs. The symbol “C” represents the inclusion of a template cluster within the wheel.
chiral recognition and sensing in host–guest systems. The following findings indicate that Ln-MBs can be used for chiral recognition: (1) Chiral lanthanide-containing POMs are capable of conducting chiral recognition. This has been observed for the chiral POM \[\text{Ce} (\alpha_1\text{P}_2\text{W}_{17}\text{O}_{61})(\text{H}_2\text{O})_x]^{7−}\] by chiral amino acids; the resulting stereoselective interactions were confirmed by $^{13}$P NMR spectroscopy.\(^{29}\) (2) Even achiral MBs can incorporate appropriate nanosized guests chemo-, regio-, and size-selectively\(^{67}\) (Figure 15 A), and selective recognition will most likely be enhanced by introducing chirality into the system. (3) The interaction of biphenyl dicarboxylic acid (bridging ligand), cyclodextrin (chiral species), and the lanthanide TbIII yielded polypseudorotaxane-type supramolecular assemblies, which enabled the recognition of small chiral molecules.\(^{68}\) Wheels can also form polyrotaxane-type supramolecular assemblies.\(^{69}\) The coassembly of the polymer $p$-phenylenebutadiynylene ($\text{PB}_n$) (Figure 15B) with a MB yielded a one-dimensional, tubular arrangement of cofacially connected wheels. Both the rigidity and the pendant ammonium groups of the $\text{PB}_n$ have a high affinity for the wheel surface, which lead to threading interactions with the wheel and the formation of so-called inorganic–organic polypseudorotaxanes. The threading interaction is so strong that even electrostatically fixed metalloporphyrins can be pushed out of the wheel.

Chiral Ln-MBs could also potentially serve as inorganic counterparts of cucurbit[$n$]urils, which are organic macrocyclic molecules composed of several glycoluril units and known for their outstanding recognition properties.\(^{70,71}\) In particular, the possibility to produce chiral Ln-MBs specially functionalized with different amino acids and peptide sequences makes them promising candidates for recognizing specific peptides and proteins. Thus, stable, highly scalable, and potentially more reliable synthetic receptors of inorganic–organic nature could be produced with the ability to specifically inhibit or enhance the biological activity of certain peptides or proteins. For example, smaller POM-peptide hybrids have been shown to inhibit the aggregation of $\beta$-amyloid peptides.\(^{72}\)

### 7.3. Lanthanide Separation

Lanthanides are important metals in the design of special materials such as illuminant, colorants, permanent magnets, and lasers.\(^{73−75}\) They coexist quite often isostructural in sources of occurrence due to their chemical similarities.\(^{76−78}\)
(e.g., ion radii close together, lack of redox activity, etc.). As a result, the separation of neighboring 4f-elements in particular poses a significant challenge. A selective lanthanide separation based on purely inorganic chemical recognition has been realized through the crystallization of borates. The separation was based on the binding differences of the LnIII ions. Remarkably, the thermodynamically controlled synthesis of the circular-shaped Ln-MBs \( \{\text{Mo}_{90}\text{Ln}_{10}\}\) (LnIII = LaIII, CeIII, or PrIII) and \( \{\text{Mo}_{92}\text{Ln}_{9}\}\) (LnIII = NdIII or SmIII) (both depicted in Figure 5J and I) can be used to selectively enrich lanthanides from binary lanthanide mixtures as well. This was confirmed by ICP-OES analyses of lanthanide containing solid products formed in a reaction solution which contained two lanthanide species (Figure 16 shows the binary system La/Sm, Eu, or Er).

The separation factor of various binary lanthanide mixtures is associated with the difference in ionic radius between the lanthanides; that is, the larger the difference in ionic radius, the better the separation. Therefore, lanthanides with large ionic radii are preferred for thermodynamically regulated wheel construction. Lanthanide separation is typically performed in industry using hydrometallurgical techniques; however, on a laboratory scale, the Ln-MB-based lanthanide separation method could prove to be an elegant approach.

8. CONCLUSION

The spectrum of Ln-MB wheel clusters is highly diverse, as they come in a variety of structural types, shapes (“Japanese rice-ball”, “egg”, “ellipse” and “circle”) and sizes with an inner and outer diameter of 10–22 and 26–39 Å, respectively. These unique inorganic ring systems possess a nanometer-sized cavity that allows for unprecedented and intriguing host–guest chemistry. Specific scaffolds with the tuning of the nanometer-sized cavity can be conveniently achieved as all architecturally labile \( \{\text{Mo}_x\}\)-type building blocks in Ln-MBs are both completely replaceable by lanthanide ions and organically modifiable. Not only the nanocavity but also the properties of the outer surface can be altered, specifically, by introducing organic surfactants to increase the hydrophobicity, which allows the adjustment of the cluster’s polarity. MBs with integrated lanthanide ions might be critical in applications such as catalysis, molecular recognition, and lanthanide ion separation, and as several spectroscopic techniques, including the powerful ESI-IMS-MS, have demonstrated, Ln-MBs preserve their integrity in solution, implying that they could endure robust reactions without decomposing. The archive of crystal structures of wheel-shaped Ln-MBs is still considerably expandable as many organic ligands, especially amine-containing ligands, remain to be explored. The conduction of SAXS studies of Ln-MBs in the solution phase is strongly recommended to explore their stability and speciation behavior in reactions and various media. We are confident that Ln-MBs will be used as an inorganic platform for the development of innovative molecular architectures and nanostructured materials as well as for the design of adaptive inorganic–organic chemical systems.

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Notes

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