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Structure and Electronic Properties of Large Oligomeric Heterometallic 3d/Ce^{IV} Polyoxometalates

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ABSTRACT: Merging the rich chemistry of Ce(IV) polyoxometalates (POMs) with that of 3d polyanions remains a challenge due to the strong competition between these highly oxophilic lanthanide cations and 3d metallic ions for coordination to lacunary molecular metal oxides. We report herein the characterization of an unprecedented water stable hexameric Ce^{IV}/Co^{II} POM (**Ce₁₂Co₆**) made of two {(SiW₉)₂Ce₆} units connected to a {(SiW₁₀)₂Co₆(PO₄)₂} core, which stability in aqueous media has been evidenced. In addition, the pentameric Ce^{IV}/Ni^{II} compound **Ce₆Ni₈** where two {PW₉Ni₃W} and a {PW₁₀Ni₂} fragments are grafted on a {(PW₉)₂Ce₆} moiety has been obtained. Magnetic studies of **Ce₆Ni₈** revealed ferromagnetic interactions between the Ni^{II} centers constituting the {Ni₃PW₁₀} fragments, in agreement with the geometry of such trinuclear cluster. Related insoluble barium salts of **Ce₁₂Co₆** and **Ce₆Ni₈** were also prepared, allowing their solid-state electrochemical investigations and showing in particular that in **Ce₁₂Co₆** both the cobalt, cerium and silicotungstate moieties are electroactive. Finally, photo-physical studies demonstrate the formation of long-lived reduced POMs photosensitized by [Ru(bpy)₃]²⁺, suggesting that **Ce₁₂Co₆** and **Ce₆Ni₈** could be used as an efficient reservoir of reduction equivalents for photocatalytic reactions.

INTRODUCTION

The chemistry of 3d-4f heterometallic polyoxometalates (POMs) has attracted remarkable interest over the last decade, not only because of the enormous variety of new structures and topologies of these compounds that could result, but also because of their many potential applications, for example in magnetism and catalysis.^{1,2} However, very few molecular POMs in which both 3d and 4f metal ions are incorporated *within* the polyoxometalate framework have been reported to date, the number of 1D to 3D species consisting of 3d homometallic entities connected by 4f cations and 4f homometallic molecular oxides connected by 3d metal ions also remaining limited.¹ The difficulty in obtaining a large library of heterometallic molecular POMs can be explained considering that (i) 3d and 4f cations compete for coordination with lacunary POMs, and as a result pure 3d or 4f species can be obtained concomitantly rather than the targeted 3d-4f POMs, and (ii) the combination of rare earth and 3d cations with lacunary POMs often lead to the precipitation of amorphous materials that are not easy to characterize. However, the (photo)³-(electro)⁴-catalytic⁵ properties of some of the successfully isolated compounds could be investigated, suggesting that the presence of both electron acceptor and donor metallic centers in a single POM framework can confer interesting properties. Nevertheless, most of the obtained species have been synthesized for magnetic purposes, as exemplified by the isolation of the POM-based {Dy^{III}Mn^{III}}₄⁶

and {Dy^{III}₃₀Co^{II}}₈⁷ compounds, both of which were found to be single-molecule magnets. This could explain why very few species incorporating the diamagnetic, electron acceptor Ce^{IV} ion have been characterized, while the chemistry of polyoxometalates incorporating this ion has proved to be very rich as evidenced in particular by the works of Duval and coll.. This group has indeed shown that large oligomeric homometallic POMs based on {Ce^{IV}}₃,⁸ {Ce^{IV}}₄⁹ or {Ce^{IV}}₆^{10,11} building units and with tremendously various topologies can be obtained. Very interestingly, Kortz and coll. also evidenced that large, water stable, cyclic peroxo-Ce^{IV} POM species can be isolated.¹² In contrast, the two 3d-4f heterometallic POMs obtained by Reinoso et al. in presence of this ion only incorporate one Ce^{IV} center. In the {Ce^{IV}Cu^{II}}₃ molecular compound [(Ce^{IV}(O₂CCH₃))Cu^{II}₃(H₂O)(B-α-GeW₉O₃₄)₂]¹¹, resulting from the addition of a Ce^{IV} salt to the well-known [Cu^{II}₄(H₂O)₂(B-α-GeW₉O₃₄)₂]¹² Weakley-type POM, a {Ce^{IV}Cu^{II}}₃ tetranuclear cluster is sandwiched between two {GeW₉} units, a Cu^{II} center being substituted by a Ce^{IV} one during the synthetic process.¹³ Alternatively, the {Ce^{IV}Mn^{IV}}₆ [(α-P₂W₁₆O₅₇(OH)₂)(Ce^{IV}Mn^{IV}₆O₉(CH₃COO)₈)]⁸ molecular POM, which consists in a divacant Dawson unit capped by a planar Mn₆ hexagon enclosing a cerium cation, was prepared by Fang et al. by reacting the pre-formed [Ce^{IV}Mn^{IV}₆O₉(O₂CCH₃)₉(NO₃)(H₂O)₂] heterometallic precursor with the vacant [α-P₂W₁₅O₅₆]¹² Dawson species.¹⁴ Also, we can note that an additional heterometallic Ce/Mn

compound has been characterized. However, this last compound consists in the co-crystallization of the cerium $\{[A-\beta\text{-SiW}_9\text{O}_{34}]_2\text{Ce}^{\text{IV}}_4\text{O}_2(\text{CH}_3\text{COO})_2\}$ and manganese $\{[A-\beta\text{-SiW}_9\text{O}_{34}]\text{Mn}^{\text{III}}_3\text{Mn}^{\text{IV}}\text{O}_3(\text{CH}_3\text{COO})_3\}$ tetranuclear POMs,¹⁵ highlighting the great difficulty to gather in a single molecule 3d and cerium(IV) clusters.

Herein we report for the first time on the synthesis and characterization of large $\text{Ce}^{\text{IV}}/3\text{d}$ heterometallic POM systems. The hexameric $\text{Ce}_{12}\text{Co}_6$ and pentameric Ce_6Ni_8 molecular POMs have been obtained from one-pot reactions performed under mild conditions. These two compounds have been fully characterized in the solid state and their behavior in solution investigated. Their magnetic properties have been studied. Electrochemical investigations have also been carried out, while photophysical studies have been performed to investigate potential electron transfer reaction between these heterometallic species and a photosensitizer, namely $[\text{Ru}(\text{bpy})_3]^{2+}$.

EXPERIMENTAL SECTION

Synthesis. $\text{Na}_{10}[\text{A}-\alpha\text{-SiW}_9\text{O}_{34}]\cdot 12\text{H}_2\text{O}$ and $\text{Na}_9[\text{A}-\alpha\text{-PW}_9\text{O}_{34}]\cdot 7\text{H}_2\text{O}$ have been synthesized as previously described.¹⁶ All other chemicals were used as purchased without purification.

$\text{Rb}_{24}\text{Na}_8\{[\text{SiW}_9\text{O}_{34}]\text{Ce}_3\text{O}_2(\text{OH})_2(\text{H}_2\text{O})\}_2\{\text{SiW}_{10}\text{O}_{37}\text{Co}_2(\text{OH})(\text{PO}_4)\}_2\text{Co}(\text{H}_2\text{O})_2\text{Co}(\text{H}_2\text{O})_4\cdot 85\text{H}_2\text{O}$ (**Ce₁₂Co₆**). $\text{Na}_{10}[\text{A}-\alpha\text{-SiW}_9\text{O}_{34}]\cdot 12\text{H}_2\text{O}$ (500 mg, 0.19 mmol) and $\text{CoCl}_2\cdot 6\text{H}_2\text{O}$ (38 mg, 0.16 mmol) were dissolved in 10 mL of an aqueous solution of 0.125 M NaOAc (pH 7, acidified with 1 M HCl). Then, $(\text{NH}_4)_2\text{Ce}(\text{NO}_3)_6$ (200 mg, 0.36 mmol) and $\text{Na}_3\text{PO}_4\cdot 12\text{H}_2\text{O}$ (50 mg, 0.15 mmol) were added as solids, and the pH adjusted to 6.0 with 1 M NaOH. The suspension was heated to 80°C for 20 min, affording a dark orange solution which was cooled to room temperature. After centrifugation, RbCl (150 mg, 1.24 mmol) was added and the mixture stirred at room temperature for 15 min. The obtained solid was filtrated ($m = 230$ mg), dissolved in 5 mL of 0.125 M NaOAc (pH 7) and let to slowly evaporate. Brown crystals were collected after one night and washed with a 1 M RbCl aqueous solution, ethanol and ether (44 mg, yield 7% vs. W). M.W. = 20 281 g.mol⁻¹. IR: ν (cm⁻¹) = 1050 (w), 1009 (w), 991 (w), 936 (m), 859 (s), 668 (s), 498 (s). EDX: W:Co:Ce:Rb:Na; calculated 56:6:12:24:8; experimental 56:5.4:12.6:22.0:6.8.

$\text{Ba}_{16}\{[\text{SiW}_9\text{O}_{34}]\text{Ce}_3\text{O}_2(\text{OH})_2(\text{H}_2\text{O})\}_2\{\text{SiW}_{10}\text{O}_{37}\text{Co}_2(\text{OH})(\text{PO}_4)\}_2\text{Co}(\text{H}_2\text{O})_2\text{Co}(\text{H}_2\text{O})_4\cdot 108\text{H}_2\text{O}$ (**BaCe₁₂Co₆**). Crystals of **Ce₁₂Co₆** (230 mg, 1.13 10⁻⁵ mmol) were dissolved in 20 mL of 0.125 M NaOAc (pH 6, acidified with 1M HCl) and added to a 20 mL $\text{BaCl}_2\cdot 2\text{H}_2\text{O}$ (1 g, 4.09 mmol) pH 6 aqueous solution. The obtained precipitate was filtrated and washed with water, ethanol and ether (180 mg, yield = 76%). M.W. = 20 680 g.mol⁻¹. IR: ν (cm⁻¹) = 1040 (sh), 1008 (w), 993 (w), 936 (m), 864 (s), 662 (s), 495 (s). EDX: W:Co:Ce:Ba; calculated 56:6:12:16; experimental 56:5.1:12.1:18.

$\text{Rb}_{20}\text{Na}_3\{[\text{PW}_9\text{O}_{34}\text{Ni}_3(\text{H}_2\text{O})_3(\text{OH})_3(\text{WO}_3)]_2\{[\text{PW}_{10}\text{Ni}_2(\text{H}_2\text{O})_2\text{O}_36(\text{OH})_2](\text{PW}_9\text{O}_{34})_2\text{Ce}_6(\text{OH})_4\text{O}_4\}\}\cdot 60\text{H}_2\text{O}$ (**Ce₆Ni₈**): $\text{Na}_9[\text{A}-\alpha\text{-PW}_9\text{O}_{34}]\cdot 7\text{H}_2\text{O}$ (500 mg, 0.19 mmol) and $\text{NiCl}_2\cdot 6\text{H}_2\text{O}$ (129 mg, 0.54 mmol) were dissolved in 10 mL of a 1 M NaOAc (pH = 7) solution. $(\text{NH}_4)_2\text{Ce}(\text{NO}_3)_6$ (100 mg, 0.18 mmol) was

added as a solid, and the pH adjusted to 7.0 with 1M NaOH. The suspension was heated to 80°C for 30 min, affording a yellow solution which was cooled to room temperature. RbCl (100 mg, 0.83 mmol) was then added and the mixture stirred at room temperature for 1 h. The solution was centrifuged and let to stand at room temperature. The solution was centrifuged again the day after and then let to evaporate slowly. Yellow crystals were collected one day later and washed with a 1 M RbCl aqueous solution, ethanol and ether (86 mg, yield 15% vs. W). M.W. = 16 409 g.mol⁻¹. IR: ν (cm⁻¹) = 1101 (w), 1036 (m), 935 (m), 874 (m), 810 (m), 676 (s), 484 (s). EDX: W:Ni:Ce:Rb:Na; calculated 48:8:6:20:3; experimental 48:9.3:6.6:20.1:2.4.

$\text{Ba}_{11.5}\{[\text{PW}_9\text{O}_{34}\text{Ni}_3(\text{H}_2\text{O})_3(\text{OH})_3(\text{WO}_3)]_2\{[\text{PW}_{10}\text{Ni}_2(\text{H}_2\text{O})_2\text{O}_36(\text{OH})_2](\text{PW}_9\text{O}_{34})_2\text{Ce}_6(\text{OH})_4\text{O}_4\}\}\cdot 80\text{H}_2\text{O}$ (**BaCe₆Ni₈**). **Ce₆Ni₈** (100 mg, 6.09 10⁻³ mmol) was dissolved in 5 mL of water and added to a 5 mL $\text{BaCl}_2\cdot 2\text{H}_2\text{O}$ (200 mg, 0.82 mmol) pH 6 aqueous solution. The obtained precipitate was filtrated and washed with water, ethanol and ether (80 mg, yield = 79%). M.W. = 16 570 g.mol⁻¹. IR: ν (cm⁻¹) = 1099 (w), 1043 (m), 937 (m), 876 (m), 813 (m), 672 (s), 479 (s). EDX: W:Ni:Ce:Ba; calculated 48:8:6:11.5; experimental 48:9.9:5.7:12.2.

Physical methods. Infrared (IR) spectra were recorded on a Nicolet 30 ATR 6700 FT spectrometer. Thermogravimetry analyses (TGA) were performed on a Mettler Toledo TGA/DSC 1, STARE System apparatus under air flow (50 mL min⁻¹) at a heating rate of 5°C min⁻¹ up to 700°C. UV-vis spectra were recorded on a Perkin Elmer Lambda 750 UV/Vis/NIR spectrometer using a quartz cuvette with a 1 cm long optical pathway. EDX experiments were performed with a Jeol JSM-5800LV Scanning Microscope equipped with an integrated EDX system, which allows, playing on the applied voltage, to analyze samples up to 2 μm from the surface and not only at the surface of the crystals. Magnetic data were collected using a Quantum Design MPMS XL7 SQUID magnetometer. Magnetic susceptibility has been recorded under a magnetic field of 0.1 T.

Nanosecond transient absorption measurements were performed with the setup described previously.¹⁷ Briefly, an OPO pumped by a pulsed nanosecond Nd:YAG laser (EK-SPLA) was used to excite the samples. The laser pulse width was ~ 4 ns (energy of about 1-2 mJ/pulse) and the pulse frequency was 10 Hz. A white light super continuum laser (STM-2-UV LEUKOS) was used as a probe light. The probe laser was operated at a repetition rate of 20 Hz. The probe beam was split by means of a beam splitter that directs one part of beam to the sample cell while the other is directed towards the reference, to monitor spectral energy distribution fluctuations. The transmitted light from the sample enters into the center of the optical fiber (round-to-linear) bundle that goes to a spectrograph (SPEX 270M, Jobin-Yvon) for analysis. The dispersed white light from the spectrograph is detected by an intensified charge coupled device (ICCD) camera PIMAX-4 (Princeton Instrument). Upon varying the delay time between pump and probe, the transient absorption spectra with and without pump can be recorded

using Equation: $\Delta A = \log_{10} \left(\frac{s_{ref}^p}{s_{ref}^0} \times \frac{s_{sample}^0}{s_{sample}^p} \right)$, where s_{ref}^p and s_{ref}^0 are reference spectra with and without pump

respectively while s_{sample}^p and s_{sample}^0 are sample spectra with and without pump respectively. All the measurements were performed under argon saturation using quartz cuvette of dimensions 10 mm x 4 mm.

For the electrochemical characterization in the solid state, the barium salts **BaCe₁₂Co₆**/GPO and **BaCe₆Ni₈**/GPO (GPO = graphene oxide) were integrated into carbon paste electrodes. These were prepared by ball milling of a mixture of paraffin oil (20 mg), graphite powder (80 mg), and **BaCe₁₂Co₆** or **BaCe₆Ni₈** (20 mg), respectively. The composites were introduced in a pocket electrode (0.07 cm² surface area and 4 mm depth). All electrochemical experiments were performed under ambient conditions with a Bio-Logic VMP3 multichannel potentiostat and implemented with a three-electrode configuration using carbon rod as counter electrode, Ag/AgCl (3 M KCl) as reference electrode, and the pocket working electrode. 1 M H₂SO₄ (pH 0.1) was used as working electrolyte solution. All potentials were measured vs. Ag/AgCl and converted to the reversible hydrogen electrode (RHE) reference scale using $E_{RHE} = E_{Ag/AgCl} + 0.21 + 0.059 \text{ pH (V)}$. Cyclic voltammetry (CV) experiments were carried out with 50 mV s⁻¹ scan rate.

For **Ce₆Ni₈**, X-ray Diffraction Intensity data collection was carried out with a Bruker D8 VENTURE diffractometer equipped with a PHOTON 100 CMOS bidimensional detector using a high brilliance 1μS microfocus X-ray Mo Kα monochromatized radiation ($\lambda = 0.71073 \text{ \AA}$). Despite numerous attempts, achieving good resolution factors for the structure of **Ce₁₂Co₆** using this diffractometer has proven challenging. In a final effort, crystals of **Ce₁₂Co₆** were subjected to synchrotron diffraction single-crystal data collection with a microfocused X-rays on the Proxima 2A beamline (Synchrotron SOLEIL, France, $\lambda = 0.72936 \text{ \AA}$),¹⁸ allowing to obtain a good model of the **Ce₁₂Co₆** structure. The absorption corrections were based on multiple and symmetry-equivalent reflections in the data sets using the SADABS program¹⁹ based on the method of Blessing.²⁰ The structures were solved by direct methods and refined by full-matrix least-squares using the SHELX-TL package.²¹ The hydrogen atoms were theoretically located on the basis of the conformation of the supporting atoms. The crystallographic data are gathered in Table S1. CCDC –2268425 (**Ce₆Ni₈**) and 2268426 (**Ce₁₂Co₆**) contain the supplementary crystallographic data for this paper. These data can be obtained free of charge from The Cambridge Crystallographic Data Centre via www.ccdc.cam.ac.uk/data_request/cif.

RESULTS AND DISCUSSION

Synthesis and Characterizations. **Ce₁₂Co₆** has been obtained via a one-pot reaction by mixing at 80°C in a sodium acetate aqueous solution the trivalent POM [A- α -SiW₉O₃₄]¹⁰⁻, sodium phosphate and the Co^{II} and Ce^{IV} precursors. The powder obtained after addition of rubidium chloride to the solution cooled to room temperature was recrystallized in sodium acetate (pH 7), affording single crystals. X-ray diffraction studies (see Table S1 for crystallographic data) revealed that **Ce₁₂Co₆** is a hexameric compound made of two $\{(\text{SiW}_9)_2\text{Ce}^{\text{IV}}_6\}$ units connected to a $\{(\text{SiW}_{10}\text{Co}^{\text{II}}_2)_2\text{Co}^{\text{II}}_2(\text{PO}_4)_2\}$ fragment (Figure 1). In each $[(\text{SiW}_9\text{O}_{34})_2\text{Ce}_6\text{O}_4(\text{OH})_4(\text{H}_2\text{O})_2]^{8-}$ unit, six octacoordinated Ce^{IV} centers forming an octahedron are sandwiched

between two [A- α -SiW₉O₃₄]¹⁰⁻ trivacant POMs, the cerium ions being linked via both oxo and hydroxo ligands ($d_{\text{Ce-O(H)}} = 2.10 - 2.51 \text{ \AA}$). Besides, the $\{(\text{SiW}_{10}\text{O}_{37}\text{Co}_2(\text{OH})(\text{PO}_4))_2\text{Co}(\text{H}_2\text{O})_2\text{Co}(\text{H}_2\text{O})_4\}$ fragment is made of a central Co^{II} ion located on a pseudo inversion center and connected to two $[\alpha\text{-SiW}_{10}\text{O}_{37}(\text{OH})\text{Co}_2]^{7-}$ Keggin polyoxometalates constituted of two $\{\text{W}_3\text{O}_{13}\}$ and two $\{\text{W}_2\text{CoO}_{13}\}$ units around an $\{\text{SiO}_4\}$ tetrahedron, the cobalt ions being connected via a vertex. In addition, two phosphate groups not only bind these cobalt centers but also cerium cations of the $\{(\text{SiW}_9)_2\text{Ce}^{\text{IV}}_6\}$ fragments. Finally, a sixth cobalt ion is connected both to a $\{\text{SiW}_{10}\text{Co}_2\}$ unit and a phosphato group, affording the hexameric **Ce₁₂Co₆** POM represented at the bottom of Figure 1.

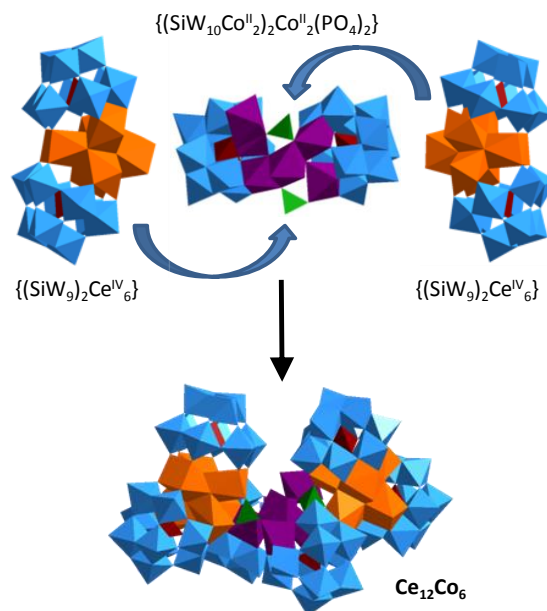


Figure 1. Polyhedral representation of **Ce₁₂Co₆** and of its constituting units; blue octahedra, WO₆; orange polyhedra, CeO₈; brown tetrahedra, SiO₄; green tetrahedra, PO₄; purple octahedra, CoO₆.

EDX experiments confirm the metallic composition of the reported heterometallic POM. Its infrared spectrum (Figure 2a) presents the characteristic W=O and W-O_b-W intense bands at wavenumbers below 1000 cm⁻¹, while the bands located at 1050 cm⁻¹ and 1009 cm⁻¹ can be attributed to P-O vibrations. All these characterizations, combined to TGA experiments (Figure 2b), allow to propose the $\text{Rb}_{24}\text{Na}_8[\{(\text{SiW}_9\text{O}_{34})\text{Ce}_3\text{O}_2(\text{OH})_2(\text{H}_2\text{O})_2\}_2\text{SiW}_{10}\text{O}_{37}\text{Co}_2(\text{OH})(\text{PO}_4))_2\text{Co}(\text{H}_2\text{O})_2\text{Co}(\text{H}_2\text{O})_4 \cdot 85\text{H}_2\text{O}]$ formula for **Ce₁₂Co₆**.

Ce₁₂Co₆ is poorly soluble in water but can be solubilized in a pH 7 0.125M sodium acetate aqueous solution. Infrared spectrum of the compound dissolved in this medium and reprecipitated with cesium chloride after 30 min at room temperature is analogous to that of pristine **Ce₁₂Co₆** crystals (Figure 2a), suggesting that this heterometallic POM is stable in aqueous solution.

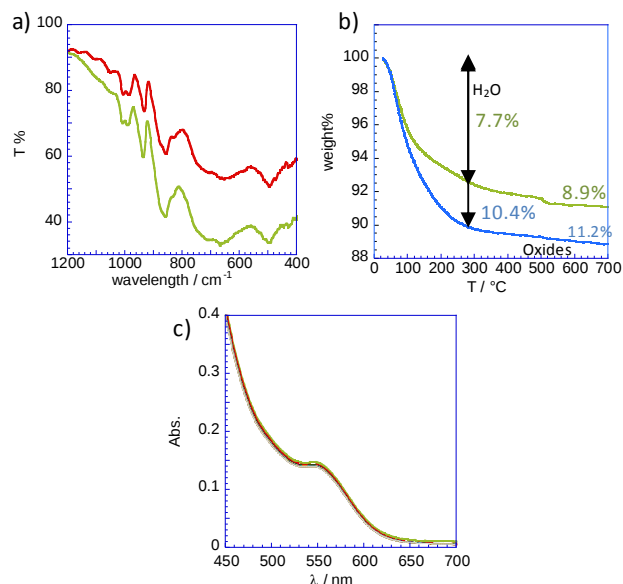


Figure 2. a) IR spectra of pristine **Ce**₁₂**Co**₆ crystals (green line) and **Ce**₁₂**Co**₆ dissolved in 0.125 M NaOAc pH 7 and re-precipitated with CsCl (red line); b) TGA measurements of **Ce**₁₂**Co**₆ (green line) and **BaCe**₁₂**Co**₆ (blue line); c) UV-Vis spectra of **Ce**₁₂**Co**₆ (0.125 M NaOAc pH 7, C = 4.2 10⁻⁴ mol.L⁻¹) recorded every hour in the 0-14 h range.

This is supported by electronic absorption spectroscopy experiments, the visible spectrum of **Ce**₁₂**Co**₆ dissolved in 0.125 M sodium acetate (pH 7) showing no evolution after 14 hours at room temperature (Figure 2c). Solid-state electronic absorption spectrum of **Ce**₁₂**Co**₆ (Figure S1) is reminiscent to that obtained in aqueous solution, this last exhibiting d-d transitions at λ = 554 nm and 541 nm, respectively, supporting again the stability of this complex in aqueous solution. Preliminary magnetic studies have been performed on a powdered sample of **Ce**₁₂**Co**₆. Related DC susceptibility curves recorded at H = 1000 and 10 000 Oe are represented in Figure S2. When the sample is cooled, the χ_MT value diminishes from χ_MT = 21.4 cm³ mol⁻¹ K at 295 K to χ_MT = 2.85 cm³ mol⁻¹ K at 2 K. The 300-30 K χ_MT behavior is characteristic of Co^{II} spin-orbit coupling phenomena, the paramagnetic ions in **Ce**₁₂**Co**₆ possessing a high-spin ground state ⁴T₁ splitted into six anisotropic Kramers doublets. However, at low temperature, the observed decrease can also be due to magnetic anisotropy and exchange interactions between the lowest Kramers doublets of the Co^{II} centers characterized by effective spins S' = 1/2, these doublets being the only ones significantly populated at low temperature (typically T < 30 K). Concerning the exchange interactions, they could in fact be both of ferromagnetic and anti-ferromagnetic natures considering that Co-O-Co angles in the 91.5-98.8 and 116.9-126.5° ranges are found in the hexanuclear cobalt cluster constituting **Ce**₁₂**Co**₆. Nevertheless, a complete treatment considering the Hamiltonian adapted to the problem would be necessary to confirm this assumption. Besides, it could be noticed that the χ_MT value determined at room temperature for **Ce**₁₂**Co**₆ is in good agreement with those previously reported for Co(II) POMs. For example, a χ_MT value of 37.5 cm³ mol⁻¹ K was found at 300 K for the decanuclear Co(II) compound [Co^{II}₁₀Co^{III}(OH)₅(H₂O)₅(W₆O₂₄)(PW₉O₃₄)₃]²²⁻ reported by

Duan et al.,²² leading to χ_MT = 3.7 cm³ mol⁻¹ K per cobalt ion vs. 3.6 cm³ mol⁻¹ K at 295 K for **Ce**₁₂**Co**₆.

The cesium salt of **Ce**₁₂**Co**₆ obtained above is slightly soluble in water. However, it is known that POM barium salts are fully insoluble in aqueous media.²³ In view to investigate the solid-state electrochemical properties of **Ce**₁₂**Co**₆ (see below), its alkaline counter-cations have been replaced by Ba²⁺ ions by simply adding BaCl₂ to a sodium acetate solution of the title compound, affording **BaCe**₁₂**Co**₆ as a fully water-insoluble powder. Infrared spectra of **Ce**₁₂**Co**₆ and **BaCe**₁₂**Co**₆ are identical (Figure S3). EDX experiments indicate that all the rubidium and sodium cations have been replaced by Ba²⁺. TGA measurement of **BaCe**₁₂**Co**₆ (Figure 2b) indicates a weight loss of 11.2% at 700°C - a temperature at which the compound is fully decomposed into oxides -, allowing to propose Ba₁₆[{(SiW₉O₃₄)Ce₃O₂(OH)₂(H₂O)}₂{SiW₁₀O₃₇Co₂(OH)(PO₄)}₂Co(H₂O)₂Co(H₂O)₄•108H₂O as formula.

Ce₆**Ni**₈ has been obtained by heating in water at pH 7 for 30 min the Ni^{II}, Ce^{IV} and [PW₉O₃₄]⁹⁻ precursors. After cooling down to room temperature, rubidium chloride was added to the yellow solution and single crystals were isolated from the slowly evaporated mother solution. X-Ray diffraction studies (Table S1) revealed that **Ce**₆**Ni**₈ is a pentameric compound (Figure 3).

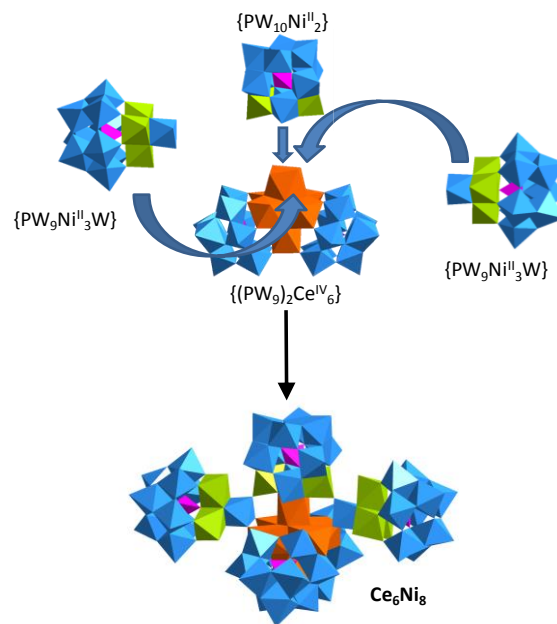


Figure 3. Polyhedral representation of **Ce**₆**Ni**₈ and of its constituting units; blue octahedra, WO₆; orange polyhedra, CeO₈; purple tetrahedra, PO₄; green octahedra, NiO₆.

Six octacoordinated Ce^{IV} centers sandwiched between two [A-α-PW₉O₃₄]⁹⁻ POMs form an octahedron highly reminiscent of that found in **Ce**₁₂**Co**₆ (see above), the cerium ions being also connected via oxo and hydroxo ligands (d_{Ce-O(H)} = 2.19 – 2.58 Å). One of the cerium ions of the {(PW₉O₃₄)₂Ce₆(OH)₄O₄} unit ensures a connection to an {α-PW₁₀Ni₂} entity in which the nickel centers belong to two adjacent {NiW₂O₁₃} trimetallic triads, the 3d ions being bridged via two {Ni-(O-W-O)₂-Ni} and a {Ni-(O-P-O)₂-Ni} fragments (d_{Ni...Ni} = 4.76 Å). The resulting trimeric unit

forms an isosceles triangle ($d_{P...P} = 8.82$ and 10.49 Å) which is sandwiched by two $\{B-\alpha-PW_9Ni_3W\}$ units, the tungsten atom capping each $\{PW_9Ni_3\}$ group ensuring connections via two $W-O-Ce$ ($d_{W...Ce} = 3.90$ and 3.91 Å) and one $W-O-Ni$ ($d_{W-O...Ni} = 3.70$ Å) bonds. Such $\{PW_9Ni_3W\}$ fragment is analogous to that found in the $[Ni_3(H_2O)_3PW_{10}O_{39}H_2O]^{7-}$ compound previously characterized.²⁴ Overall, it follows that the **Ce₆Ni₈** molecular pentameric compound adopts a pseudo trigonal bipyramidal arrangement. To the best of our knowledge, despite the tremendously huge number of POMs structurally characterized to date, such arrangement is unprecedented in polyoxometalate chemistry.

EDX measurements confirm the $W:Ce:Ni$ metallic ratio in **Ce₆Ni₈**. TGA experiments (Figure S4) indicate a weight loss of 8.1% at 700°C. Considering that the metallic oxides are entirely formed at this temperature, this leads to propose the formula $Rb_{20}Na_3[\{PW_9O_{34}Ni_3(H_2O)_3(OH)_3(WO_3)\}_2\{(PW_{10}Ni_2(H_2O)_2O_{36}(OH)_2)((PW_9O_{34})_2Ce_6(OH)_4O_4)\}]\cdot 60H_2O$ for **Ce₆Ni₈**. The calculated weight loss due to the departure of water molecules (7.5%) is in good agreement with the weight loss observed at 250°C (7.6%). In addition to the $W=O$ (935 cm^{-1}), $W-O_b-W$ (874 cm^{-1}) and $W-O_c-W$ (810 cm^{-1}) bands, the FT-IR spectrum of **Ce₆Ni₈** (Figure S5) exhibits two bands at 1101 and 1036 cm^{-1} which correspond to P-O antisymmetric stretching modes of the phosphotungstate groups.

The magnetic properties of **Ce₆Ni₈** have been investigated and interpreted considering the $\{PW_{10}Ni_2\}$ and the two $\{PW_9Ni_3W\}$ units as three isolated fragments. Also, the Ni^{II} centers in $\{PW_{10}Ni_2\}$ have been considered as magnetically non-interacting. Figure 4 evidences that the $\chi_M T$ curve exhibits a continuous increase upon cooling from 300 to 10 K, after which $\chi_M T$ rapidly decreases. This is characteristic of a ferromagnetic behavior, in agreement with $Ni-O-Ni$ angles in the 93.5 - 101.9° range in $\{PW_9Ni_3W\}$,²⁵ while the sudden decrease in $\chi_M T$ below 10 K is due to the zero-field splitting (ZFS) effect. The following Hamiltonian has thus been considered:

$$\hat{H} = -2J(\hat{S}_1\hat{S}_2 + \hat{S}_2\hat{S}_3 + \hat{S}_3\hat{S}_1 + \hat{S}'_1\hat{S}'_2 + \hat{S}'_2\hat{S}'_3 + \hat{S}'_3\hat{S}'_1) + \sum_{i=1}^8 D(\hat{S}_z^2 - \hat{S}(\hat{S} + 1))$$

with $S_1 = S_2 = S_3 = S'_1 = S'_2 = S'_3 = 1$ associated to the local spin of the Ni^{II} centers constituting the two $\{PW_9Ni_3W\}$ units, and the axial ZFS Hamiltonian being relative to the eight Ni^{II} centers present in **Ce₆Ni₈**. The $\chi_M T = f(T)$ and the variation of the magnetization M with the applied magnetic field H in the $0 - 5.5$ T range at 2, 3, 4, 6 and 8 K (Figure S6) curves were thus fitted, the best fit parameters obtained being $J = +3.8$ cm^{-1} , $g = 2.27$ and $D = +6.7$ cm^{-1} (Res = 0.019).²⁶ The determined parameters are in excellent agreement with those extracted from the magnetic data of the related $[Ni_3(H_2O)_3PW_{10}O_{39}H_2O]^{7-}$ POM ($J = +3.9$ cm^{-1} , $g = 2.33$ and $D = +5.8$ cm^{-1}), validating our model.²⁷

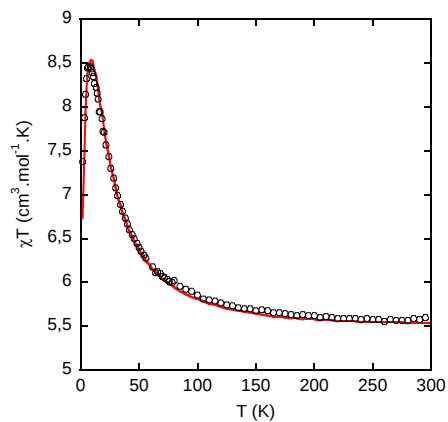


Figure 4. Temperature times magnetic susceptibility per mole of **Ce₆Ni₈** as a function of temperature between 300 and 2 K.

As some small deviations between the fits and the experimental $\chi_M T = f(T)$ and $M = f(H)$ curves are observed at low temperatures ($T < 6$ K), we have attempted to introduce a J' exchange parameter relating the magnetic interaction between the two Ni^{II} centers constituting the $\{PW_{10}Ni_2\}$ unit. The best fit parameters obtained were $g = 2.276$, $D = +6.62$ cm^{-1} , $J = +3.89$ cm^{-1} and $J' = -0.349$ cm^{-1} . However, only a slight improvement of the low temperature fit was observed (Figure S7). Also, considering the $\{Ni_3\}$ trinuclear units as isosceles triangles did not significantly improve the fit for $T < 6$ K.

The complex **Ce₆Ni₈** is well soluble in water. Infrared spectrum of the compound dissolved in water and re-precipitated with cesium chloride is analogous to that of pristine **Ce₆Ni₈** crystals (Figure S5). Both the solution (Figure S8) and solid state (Figure S9) electronic spectra evidenced two d-d centered bands in the visible range, with $\lambda = 715$ and 749 nm and $\lambda = 721$ and 767 nm, respectively. Recording UV-Vis spectra of **Ce₆Ni₈** in solution as a function of time (Figure S8) has highlighted a slight decrease of the absorbance (ca. 6.5% after 1 h) at $\lambda_{max} = 715$ nm. Nevertheless, as the observed **Ce₆Ni₈** evolving in solution is slow, a water insoluble salt of **Ce₆Ni₈** has been obtained by readily reprecipitating the alkaline salt of **Ce₆Ni₈** with Ba^{2+} , as performed for **Ce₁₂Co₆**. The IR spectrum of the obtained **BaCe₆Ni₈** is strictly analogous to that of **Ce₆Ni₈** (Figure S5). Finally, its full formula was determined after TGA (Figure S10) and EDX experiments.

Cyclic voltammetry (CV) experiments in 1 M H_2SO_4 solutions were then conducted to investigate the redox behaviors of **BaCe₁₂Co₆** and **BaCe₆Ni₈**, respectively (Figure 5).

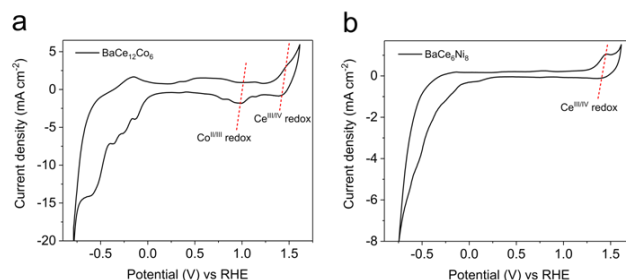


Figure 5. CV curves of (a) **BaCe₁₂Co₆**/graphene oxide (GPO) and (b) **BaCe₆Ni₈**/GPO in 1 M H_2SO_4 .

At negative potentials, both **BaCe₁₂Co₆** and **BaCe₆Ni₈** functionalized electrodes show multi-electron reduction processes, assigned to multiple electron uptake by {XW₉} (X = Si, P) moieties.^{28,29,30} The current density reaches maximum values at lowest potential suggesting the concomitant appearance of hydrogen evolution albeit, at very high overpotential. The overall process appears irreversible, since the electron oxidation waves are not observable in the scans towards positive potentials. When positive potentials over 1 V vs RHE are applied, both electrodes show a reversible Ce^{III/IV} wave. This redox wave appears at significantly high oxidation potentials when compared with aqueous Ce^{III/IV}, which must be due to electronic withdrawing effects imposed by the POM structure. The major difference between both electrodes resides in the appearance of an additional quasi-reversible Co^{III/II} redox peak in **BaCe₁₂Co₆**, before and independently of the Ce^{III/IV} event.³¹

To investigate potential electron transfer reaction between these heterometallic species and a photosensitizer (PS), nanosecond transient absorption measurements were performed in samples containing the POM, [Ru(bpy)₃]²⁺ as PS and ascorbate as electron donor by photoexcitation at 460 nm, which correspond to the absorption band of the PS. Transient absorption spectra (TAS) in the presence of **Ce₁₂Co₆** together with those of the control experiment in the absence of the POM were recorded at different delay times in pH = 7 acetate solutions (Figure 6). In the absence of POM, photoinduced electron transfer between photoexcited PS and ascorbate produced the reduced PS ([Ru(bpy)₃]^{•+}) with characteristic transient absorption band at ca. 500 nm rising until 1 μs (Figure 6a). At longer delay time, the decay of this transient band reflects the charge recombination between the reduced PS and oxidized ascorbate. The data were fitted globally using a bi-exponential function and the resulted kinetics at selected wavelengths and decays associated difference spectra (DADS) are presented in Figure 6b and Figure S11a, respectively. The first DADS has an associated lifetime of about 170 ns and exhibits a bleaching signal at 460 nm corresponding to the excited state of the PS. The second DADS associated to 38 μs lifetime exhibits absorption maxima at 500 nm and a negative signal at 460 nm, representing the reduced PS.

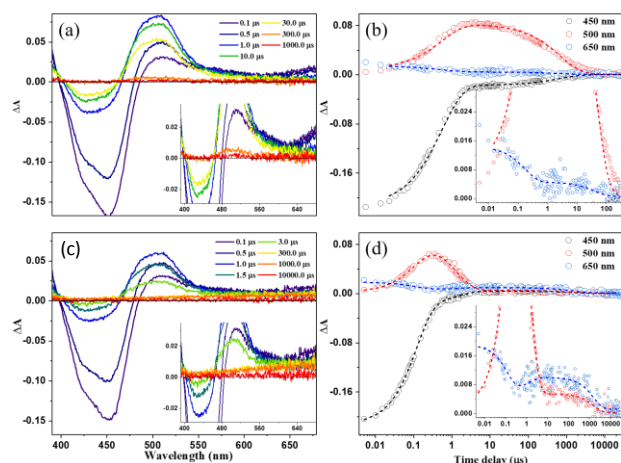


Figure 6. Nanosecond transient absorption spectra and selected kinetic measurements after excitation at 460 nm of [Ru(bpy)₃]²⁺ (51 μM) and ascorbate (100 mM) in acetate

buffer (pH = 7), (a,b) in the absence and (c, d) in the presence of **Ce₁₂Co₆** (100 μM). Inset: corresponding zoomed in Figures.

In the presence of POM, new broad absorption features, tentatively attributed to reduced POM,³² appeared at longer delay times, from hundreds μs and lives for almost 10 ms (Figure 6c). This is consistent with previous studies which suggested broad and shapeless spectra of reduced POM appearing at particularly longer time delays.^{32,33} Kinetics of bleaching signal (450 nm), absorption signals of [Ru(bpy)₃]^{•+} (500 nm) and of reduced POM (650 nm) are shown in Figure 6d. Decay of [Ru(bpy)₃]^{•+} absorption band at 500 nm is more rapid in the presence of POM, and its disappearance together with the growth of absorption bands at 650 nm of reduced POM (inset of Figure 6d) indicate an electron transfer process from reduced PS to POM. The decays were fitted globally using tri-exponential function yielding DADS shown in Figure S11b. The first and the second time components of 120 ns and 1.4 μs are attributed to decay time of excited and reduced PS, respectively. The longest time component of 1500 μs reflects the lifetime of the reduced POM, corresponding to its recombination with oxidized ascorbate. The mechanism of the formation of reduced POM in [Ru(bpy)₃]²⁺/ascorbate/POM transient absorption experiment is summarized in Figure S12.

In the similar manner, the experiments were also performed under identical conditions using **Ce₆Ni₈**, [Ru(bpy)₃]²⁺ as the photosensitizer and ascorbate as the electron donor in water (Figure S13-S14). The results showed the signature of reduced POM with a lifetime of 732 μs. The detailed discussion of the experiments with **Ce₆Ni₈** can be found in the supplementary information. Overall, these results support the possibility of POM reduction photosensitized by [Ru(bpy)₃]²⁺ for both reported heterometallic POMs.

CONCLUSIONS

We have thus shown here for the first time that it is possible to build oligomeric polyoxometalates associating in a single molecule high-nuclearity cerium(IV) and 3d-substituted POM clusters. Moreover, it has been possible to readily obtain these species under mild conditions via a one-pot synthetic strategy and starting from simple precursors. In the cobalt complex **Ce₁₂Co₆**, two {(SiW₉)₂Ce₆} units connect a {(SiW₁₀)₂Co₆(PO₄)₂} core, resulting in a hexameric compound. In contrast, a {(PW₉)₂Ce₆} entity constitutes the central unit of the pentameric species **Ce₆Ni₈**, which adopts a unique pseudo trigonal bipyramidal arrangement, the cerium POM cluster being connected to a {PW₁₀Ni₂} and two {PW₉Ni₃W} fragments. UV-vis and IR studies showed that these high-nuclearity species are stable in aqueous media. Insoluble barium salts of **Ce₁₂Co₆** and **Ce₆Ni₈** have also been prepared, and solid-state electrochemical investigations evidenced that in **Ce₁₂Co₆** the cobalt, cerium and silicotungstate moieties are electroactive. Moreover, photophysical studies demonstrate the formation of long-lived reduced POM photosensitized by [Ru(bpy)₃]²⁺ for both **Ce₁₂Co₆** and **Ce₆Ni₈**, which can thus potentially be used as efficient electron reservoirs. The photocatalytic properties of these new species are currently under study.

ASSOCIATED CONTENT

Supporting Information: Complementary crystallographic, TGA, IR, magnetic, photophysical and electrochemical data (PDF). This material is available free of charge via the Internet at <http://pubs.acs.org>.

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The manuscript was written through contributions of all authors. All authors have given approval to the final version of the manuscript.

Notes

The authors declare no competing financial interest.

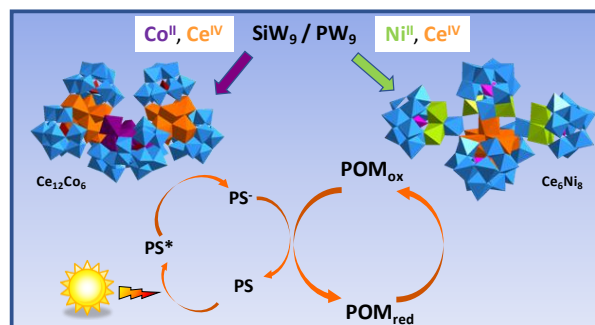
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Synopsis

Unprecedented water stable hexameric $\text{Ce}^{\text{IV}}/\text{Co}^{\text{II}}$ and pentameric $\text{Ce}^{\text{IV}}/\text{Ni}^{\text{II}}$ large polyoxometalates have been synthesized and their magnetic properties investigated. Solid-state electrochemical investigations showed that in the Ce/Co system both the cobalt, cerium and silicotungstate moieties are electroactive, while photo-physical studies demonstrate the formation of long-lived reduced POMs photosensitized by $[\text{Ru}(\text{bpy})_3]^{2+}$, suggesting that these species could be used as an efficient reservoir of reduction equivalents for photocatalytic reactions.
