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Evaluation of ultrasmall coinage metal M_{13} (dppe)₆ M = Cu, Ag, and Au clusters. Bonding, structural and optical properties from relativistic DFT calculations*

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Ultrasmall ligand-protected clusters are prototypical species for evaluating the variation at the bottom of the nanoscale range. Here we explored the ultrasmall gold-phosphine M₁₃(dppe)₆ cluster, as a prototypical framework to gain insights into the fundamental similarities and differences between Au, Ag, and Cu, in the 1-3 nm size range, via relativistic DFT calculations. Different charge states involving 8- and 10-cluster electron (ce) species with a 1S²1P⁶ and 1S²1P⁶1D² configuration, leading to structural modification in the Au species between $Au_{13}(dppm)_6^{5+}$ and $Au_{13}(dppm)_6^{3+}$, respectively. Furthermore, this structural distortion of the M_{13} core is found to occur to a lower degree for the calculated Ag and Cu counterparts. Interestingly, optical properties exhibit similar main patterns along with the series, inducing a blue-shift for silver and copper, in comparison to the gold parent cluster. For 10-ce species, the main features of 8-ce are retained with the appearance of several weak transitions in the range. The ligand-core interaction is enhanced for gold counterparts and decreased for lighter counterparts resulting in the Au > Cu > Ag trend for the interaction stabilization. Hence, the Ag and Cu counterparts of the Au₁₃(dppm)₆ cluster appear as useful alternatives, which can be further explored towards different cluster alternatives for building blocks for nanostructured materials.

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Introduction

Atomically precise gold clusters have attracted increased interest due to their unique properties relative to the bulk material, 1-9 enabling further elucidation of the nonscalable properties of ultrasmall nanoparticles. 10-12 In the last few decades, several synthetic breakthroughs^{13–24} led the characterization of ligand passivated clusters in the 1-3 nm range with both structural and electronic features ascribed to the nature of the metallic core, 25-30 usually rationalized via the divide-andprotect approach.31,32

Among the different stabilizing groups, thiolates and phosphine ligands are widespread in the literature, 25-28,33,34 ensuring strong gold-gold and gold-ligand bonds in stable structures against degradation. $^{35-42}$ Au₂₅(SR)₁₈ $^-$ (R = ligand) is one of the most noticeable ligand-protected gold clusters, composed of an atom-centered icosahedral core surrounded by six staple units, Au₂(SR)₃, 35,43-45 featuring eight cluster electrons (8-ce)^{25,46} in a 1S²1P⁶ electronic configuration. This fact accounts for its particular stability⁴⁷ in analogy to closedshell atoms, which are coined as superatoms with the central Au₁₃⁵⁺ core⁴⁸ as a recursive motif in small and ultrasmall nanoparticles. 49 It is noteworthy that Au₂₅(SR)₁₈ possesses a silver isoelectronic and isostructural analog, Ag₂₅(SR)₁₈-,^{50,51} suggesting that the Au₁₃⁵⁺ core features can also be observed in other lighter coinage metals³¹ such as Ag₁₃^{5+,50,52} and Cu₁₃^{5+,53}

Recently, ultrasmall Au₁₃(dppm)₆ has been the focus of renewed interest since its early structural characterization in 1981,¹³ depicting a fully phosphine-protected Au₁₃ core. Independently, Sun and Zhu groups showed full characterization of UV/vis and cyclic-voltammetry patterns and synthetic advantages, denoting the stability of such a cluster in several charge states, with structural rearrangements between the +5 and



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+3 charge states. 54,55 The electronic structure of $Au_{13}(dppm)_6^{5+}$ is based on a closed-shell 1S21P6 configuration, similarly to Au₂₅(SR)₁₈-, providing a 1D character of low-lying unoccupied levels, with a variable structure in Au₁₃(dppm)₆³⁺ denoting a distorted 10-ce Au₁₃ core in a 1S²1P⁶1D² configuration.

Such species provide useful templates to further evaluate the bonding and electronic characteristics, denoting the role of the ligand-shell in the stabilization and the relation between the core-structure and optical properties. Herein, we set to evaluate the plausible formation of lighter coinage-metal group counterparts to gain more insights into the century-old concern of fundamental similarities and differences between Au, Ag, and Cu, in terms of the structure and related properties, involving different charge states. In this contribution, we provide information on the overall ligand-core interaction, and optical and structural properties, for the ultrasmall M₁₃(dppm)₆ series, by using relativistic DFT methods, as a prototypical case of ligand-protected clusters.

Computational details

Computations were carried out at the relative density functional level of theory⁵⁶ by using the ADF code,⁵⁷ incorporating scalar corrections *via* the ZORA Hamiltonian.⁵⁸ The triple-ξ Slater basis set, plus two polarization functions (STO-TZ2P) for valence electrons, were employed within the generalized gradient approximation (GGA) according to the Perdew-Burke-Ernzerhof (PBE) exchange-correlation functional. 59,60 Dispersion corrections to DFT were taken into account via the pairwise Grimme3 approach. 61,62 The use of the PBE-GGA functional provides accurate results for both the structure and optical properties of gold nanoclusters with minimal ligand simplifications, 44,63-67 at an affordable computational cost, as denoted by Muniz-Miranda and coworkers. 68 Optical properties were simulated by using the Van Leeuwen and Baerends functional (LB94),69 denoting a good performance in the study of excitation in ligand-protected clusters.70-74 Solvation effects were taken into account via a continuum solvation scheme given by the 'COnductor-like Screening Model' (COSMO) module from ADF, to describe the screening effects from solvation by using dichloromethane.⁷⁵

The frozen core approximation was applied to the $[1s^2-4f^{14}]$ shells for Au, $[1s^2]$ for C, and $[2s^2]$ for P, leaving the remaining electrons to be treated variationally. Geometry optimizations were performed without any symmetry restraint via the analytical energy gradient method implemented by Versluis and Ziegler. 76 An energy convergence criterion of 10-4 Hartree, gradient convergence criteria of 10⁻³ Hartree Å⁻¹, and radial convergence criteria of 10^{-2} Å were employed for the evaluation of the relaxed structures.

Results and discussion

The characterized fully phosphine-protected Au₁₃ core in its +5 charge state, 55 exhibits a distorted atom-centered icosahedron passivated with six dppm units (Fig. 1), as the structure features

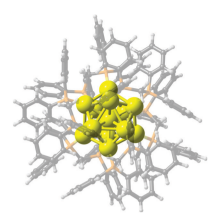


Fig. 1 Schematic structures for M₁₃(dppm)₆ clusters. Color code: Au, Ag, Cu, yellow; C: black; P: orange; H: white.

two groups of distances found for the central atom and icosahedral cage (Au_{cent}-Au_{ico}) length of 2.702 and 2.925 Å, respectively. Similarly, two groups of Auico-P distances are obtained of 2.282 and 2.346 Å, respectively, owing to a slight icosahedron ↔ cuboctahedron distortion with an elongation of certain distances (Table S1, ESI†). In its +3 charge state counterpart, the distortion of the icosahedron is increased⁵⁴ leading to several Au_{cent}-Au_{ico} distances (Tables S1 and S2, ESI†) in roughly six groups, which average to 2.819 Å, slightly shorter than 2.832 Å obtained for $[Au_{13}(dppm)_6]^{5+}$, denoting a decrease of the cage radius upon addition of two electrons.

The calculated Au_{cent} - Au_{ico} distances for $[Au_{13}(dppm)_6]^{5+}$ (1), are averaged in two groups of 2.817 and 2.921 Å, respectively, leading to an average of 2.869 Å (Table 1), in good agreement to experimental values (2.832 Å), as also obtained for Au_{ico}-P separations (2.327 and 2.341 Å). For the +3 charge state (2), similar values to experimental data are obtained, with an averaged Au_{cent}-Au_{ico} bond distance of 2.321 Å (experimental 2.314 Å).54

In order to account for structural deviations from an ideal atom-centered icosahedron (Ih), we evaluated the continuousshape-measure (CShM) developed by the group of Alvarez and coworkers. 77-79 The resulting CShM values are close to zero for structures fully coincident with the reference icosahedron and increasing values accounting for the distortion degree. For the X-ray structure of $[Au_{13}(dppm)_6]^{5+}$ (1), the Au_{13} core exhibits a CShM value of 0.568 (Calculated, 0.315), denoting the distortion of the cage given by the different groups of Au_{cent}-Au_{ico} distances (vide intra). Moreover, for [Au₁₃(dppm)₆]³⁺ (2), the CShM value increases consequently to 17.352 (calc. 17.472) owing to the different types of Au_{cent}-Au_{ico} distances.

The comparison between a perfect icosahedral structure, obtained by constraining equal Auico-Auico distances previous to relax the protecting ligand shell, and the fully optimized structure for $[Au_{13}(dppm)_6]^{5+}$ (1), reveals a small energy difference of 6.0 kcal mol⁻¹, in agreement with the small deviation accounted by the CShM value. In contrast, the larger distortion observed for $[Au_{13}(dppm)_6]^{3+}(2)$ leads to a larger energy difference of 171.5 kcal mol⁻¹ in comparison with the relaxed and perfect

Table 1 Averaged geometrical parameters in angstroms (Å), and HOMO-LUMO gap values in eV, for q = +5 and +3 species of $[M_{13}(dppm)_6]^q$. In addition, continuous-shape-measure data for the M₁₃ core is given

M	M _{cent} -M _{ico}	CShM	M _{ico} -P	H-L gap
q = +5				
Au	2.869	0.315	2.334	1.431
Au $(Exp.)^a$	2.832	0.568	2.314	
Ag	2.881	0.178	2.502	1.375
Cu	2.537	0.144	2.312	1.440
q = +3				
Âu	2.872	17.472	2.321	1.502
Au $(Exp.)^a$	2.819	17.352	2.314	
Ag	2.889	0.217	2.503	0.183
Cu	2.483	0.058	2.279	0.323

^a Experimental data from ref. 55 (q = +5) and ref. 54 (q = +3).

icosahedral Au₁₃ core structure, denoting the preference for a high distortion of the core in the latter charge state.

The electronic structure for $[Au_{13}(dppm)_6]^{5+}$ (1) is evaluated in terms of the superatom approach, 25,46,80-82 denoting an 8 cluster electron (ce) count for the Au₁₃⁵⁺ core, fulfilling a 1S²1P⁶ configuration (Fig. 2).^{25,46} The 1P shell remains the highest occupied molecular orbital (HOMO), in an almost three-fold degenerate level, whereas the low-lying unoccupied MO is a split of the 1D shell into three- and two-fold levels (Fig. 2), leading to a sizable HOMO-LUMO gap of 1.534 eV, in the range of the determined by electrochemical and UV-vis data (1.66 and

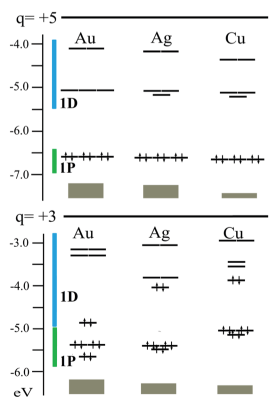


Fig. 2 Electronic structure for $M_{13}(dppm)_6$ at +5 and +3 charge states. M = Au, Ag, and Cu 1P and 1D superatomic shells are denoted by green and

1.68 eV, respectively).⁵⁵ Interestingly, for 2, the extra two-electrons reside at the $1D_{z^2}$ level, which in turn leads to a high stabilization of this shell. Now the HOMO–LUMO gap amounts to 1.502 eV, suggesting that the observed structural distortion of the Au_{13}^{3+} core enables a sizable HOMO–LUMO gap in a 10-ce $1S^21P^61D^2_{z_2}$ configuration. Isosurfaces for relevant shells are given at the ESI† (Fig. S1).

Furthermore, driven by the full characterization of $Ag_{25}(SR)_{18}^-$ with a central Ag_{13}^{5+} core, 50,51 and similarities between Au_{13}^{5+} and Cu_{13}^{5+} cores, 31 here we evaluate the hypothetical Ag and Cu counterparts of $[Au_{13}(dppm)_6]^{5+}$. Interestingly, similar electronic features are found with calculated HOMO-LUMO gaps of 1.375 and 1.440 eV, for Ag and Cu species, respectively, denoting 1P and 1D based frontier orbitals. The relaxed structures show similar averaged M_{cent}-M_{ico} distances for Ag (2.881 Å) and the Au parent (2.869 Å), with contraction to 2.537 Å for Cu species, denoting deviations from a perfect icosahedron core as given by the CShM values of 0.178 and 0.144, respectively, where values >0.1 are considered as distorted icosahedral cores.34 Moreover, this small distortion from the icosahedron led to a decreased energy difference between an ideal Ih-M13 core situation and the fully relaxed structures, of 6.1 for Ag (6.1 kcal mol⁻¹) and for Cu (4.0 kcal mol⁻¹) counterparts, which are similar to the observed for $[Au_{13}(dppm)_6]^{5+}$ (1) (6.0 kcal mol⁻¹).

However, the +3 charge state exhibits larger differences along with the series where the 10-ce 1S²1P⁶1D²_{z2} configuration exhibits a decreased HOMO-LUMO gap of 0.183 and 0.323 eV for Ag and Cu, respectively (Fig. 2). The relaxed structures for $[Ag_{13}(dppm)_6]^{3+}$ and $[Cu_{13}(dppm)_6]^{3+}$, are slightly distorted from a perfect icosahedron with CShM values of 0.217 and 0.058, respectively, in comparison to the Au parent (CShM = 17.472). Such a result denotes that the capabilities of coinage metal clusters to enable a structural rearrangement to maximize the HOMO-LUMO gap in 10-ce clusters, are different along with the triad, which is mostly enabled in gold species, whereas lighter silver and copper counterparts suggest lower structural flexibility upon 8- to 10-ce shift. Thus, such results suggest gold clusters to be more prone to reaccommodate structurally rather than Ag and Cu counterparts, resulting in a small HOMO-LUMO gap for the latter hypothetical species suggesting a lower kinetic and chemical stability. Hence, for 10-ce counterparts Ag and Cu clusters are less favored than the parent gold system, in contrast to 8-ce species which are expected to have a similar in magnitude HOMO-LUMO gap along the coinage metal group.

In order to gain more insights into the favorable stabilization given by the protecting layer of dppm ligands, the M_{13}^{5+} -dppm interaction energy is evaluated ($\Delta E_{\rm int}$, Table 2). A more favorable situation is obtained for the Au case, amounting to -1168.2 kcal ${\rm mol}^{-1}$, which decreases to -856.9 kcal ${\rm mol}^{-1}$ for Ag, and -1038.0 kcal ${\rm mol}^{-1}$ for Cu. Thus, the favorable bonding situation evolves along with the triad according to Au $> {\rm Cu} > {\rm Ag}$, as depicted for PPh₃ and other ligands. In addition, for $[{\rm Au}_{13}({\rm dppm})_6]^{3+}$ (2), the $\Delta E_{\rm int}$ amounts to -787.8 kcal ${\rm mol}^{-1}$ as a result of the two-extra electrons in the Au₁₃ core denoting a sizable decrease in the core-ligand

Table 2 Energy decomposition analysis for the core-ligand interaction. Values in kcal mol^{-1} , at different charge states, +5 and +3, respectively

Charge +5	Au		Ag		Cu	
$\Delta E_{ m Pauli} \ \Delta E_{ m elstat} \ \Delta E_{ m orb} \ \Delta E_{ m disp} \ \Delta E_{ m int}$	1917.6 -1722.1 -1254.3 -109.4 -1168.2	55.8% 40.6% 3.5%	876.8 -901.6 -745.5 -86.6 -856.9	52.0% 43.0% 5.0%	878.2 -917.9 -902.1 -96.2 -1038.0	47.9% 47.1% 5.0%
Charge +3	Au		Ag		Cu	

stabilization, which is similarly found for Ag and Cu, to -507.7 and -628.3 kcal mol^{-1} , respectively. The preference for electronegative ligands for silver protected clusters observed in the literature over electroneutral ligands, 34 account for the calculated less favorable core-ligand interaction along with the series, suggesting that electronegative ligands provide stronger interaction. However, the core-ligand interaction energy (ΔE_{int}) for silver counterpart suggest a decreased but still sizable interaction, which in addition to the sizable HOMO-LUMO gap, provides the basis for further conceivable characterization.

Moreover, the nature of the core-ligand interaction is accounted by the energy decomposition analysis (EDA) within the Morokuma–Ziegler scheme, $^{84-86}$ dissecting the $\Delta E_{\rm int}$ quantity into different chemically meaningful terms, as 85,87,88

$$\Delta E_{\text{int}} = \Delta E_{\text{Pauli}} + \Delta E_{\text{elstat}} + \Delta E_{\text{orb}} + \Delta E_{\text{disp}}$$

In this framework, the stabilizing $\Delta E_{\rm elstat}$ and $\Delta E_{\rm orb}$ terms are related to the electrostatic and covalent character of the interaction, respectively. The $\Delta E_{\rm Pauli}$ term is given by the repulsive four-electron two-orbital interactions between occupied orbitals of the different fragments. In addition, the pairwise correction of Grimme62 (DFT-D3) accounts for London dispersion interactions ($\Delta E_{\rm disp}$) of stabilizing character. To overcome the basis set superposition error (BSSE) in the fragment interaction analysis of $\Delta E_{\rm orb}$, the counterpoise method was employed.

For $[\mathrm{Au_{13}(dppm)_6}]^{5^+}$ (1), the interaction is given by a 55.8% of electrostatic character ($\Delta E_{\mathrm{elstat}}$), and 40.6% from covalent contributions (ΔE_{orb}), and a small addition from ΔE_{disp} . Thus, the core-ligand interaction is of the main electrostatic character with a sizable bonding contribution. Upon addition of two electrons, in the 10-ce $[\mathrm{Au_{13}(dppm)_6}]^{3^+}$ (2) species, the interaction is similarly of electrostatic character, denoting a sizable decrease in the bonding contribution (ΔE_{orb}) from -1254.3 in 1 to -990.0 kcal mol^{-1} in 2, and an increase in the Pauli repulsion term, owing to the shorter $\mathrm{Au_{ico}}$ -P distance required to maximize the bonding stabilization in the latter.

For silver and copper counterparts, the nature of the coreligand interaction is similar in nature to gold species, which denotes a decrease in the contributing terms. Interestingly, by **PCCP** Paper

comparing Ag and Cu species, the different energetic terms remain similar, with the exception of the covalent character, which increases sizably for Cu counterparts, resulting in the Au > Cu > Ag trend for the interaction stabilization.

Moreover, the covalent character $(\Delta E_{\rm orb})$ can be further decomposed through the Natural Orbitals for Chemical Valence⁸⁹⁻⁹¹ extension of EDA (EDA-NOCV).⁹¹ As a result, the identification of different individual orbital contributions given by their deformation densities accounts for individual chargetransfer channels between the core and the ligand layer. 92,93 The obtained results exhibit a σ -donation from the phosphine ligands towards the core, as the main source of the bonding interaction in the Au₁₃(dppm)₆ formation (Fig. S2, ESI†).

The cyclic voltammetry (CV) results for $[Au_{13}(dppm)_6]^{5+}(1)$, ⁵⁵ exhibit reversible redox couples for both first oxidation (+5 \rightarrow +6) and reduction (+5 \rightarrow +4). In addition, a redox couple above the solvent window (>-2.20 V vs. Ag/Ag^+) is expected from calculations, suggesting the plausible formation $[Au_{13}(dppm)_6]^{3+}$ (2) by electrochemical methods. The calculations for the different charge states observed from CV experiments of 1, and by the +3 state in 2, allows us to evaluate the evolution from +3 to +6 species. The structures for $[Au_{13}(dppm)_6]^q$ from q = +3 to +6 (Table S1, ESI†), suggest similar averaged Au_{cent}-Au_{ico} distances, accounting for a similar Au₁₃ radius along with the redox processes. For silver, the averaged Ag_{cent}-Ag_{ico} tend to decrease slightly from 2.889 to 2.872 Å, suggesting a small compression of the Ag₁₃ cage, which contrasts with that found for the copper case, where averaged Cucent-Cuico distance increases from 2.483 to 2.634 Å. Such results suggest a variation of the M₁₃ radii along with different charge states, depicted as a decrease in Ag, a lesser variation for Au, and in contrast, increasing for Cu species, according to the electron count from 10- to 7-ce. In comparison, for Au₂₅(PET)₁₈ species with charge states from -1 to +1, denoting 8- to 5-ce, the averaged Au_{cent}-Au_{ico} distances taken from X-ray structures, 35,94 show an increase as 2.774 < 2.783 < 2.808 Å (Table S3, ESI†), denoting that the ligand shell environment is relevant in a further relaxation of the core upon different charge states.

Lastly, optical properties were evaluated to account for the possible variation of the optical transition in the coinage metal triad. For $[Au_{13}(dppm)_6]^{5+}(1)$, 55 the optical spectrum is dominated by an absorption band at 440 nm, and weak transitions between 500 and 750 nm, which are ascribed to the main 1P \rightarrow 1D transition via time-dependent DFT calculations, which are located mainly at the Au₁₃⁵⁺ core (ESI†). Our calculations are in agreement with the previous analysis for Au₁₃, with the absorption maxima at 437 nm involving a wide 1P → Ligand, and 5d-Au block → 1D character manifold, denoting lower energy transitions between 1P-HOMO and 1D-LUMO as a small peak at 747 nm, and very weak lower energy transitions (Fig. 3). At 515 nm, a small shoulder is found, contributed by 71% of 1P \rightarrow 1D, and 29% from Ligand \rightarrow 1D transitions. Interestingly, for Ag and Cu counterparts, the same features of the absorption spectrum remain with a slight hypsochromic (blue-) shift of all the signals, with a small peak located at 728 and 713 nm, for Ag and Cu, respectively, showing an increase in the intensity of the Ligand → 1D shoulder, now

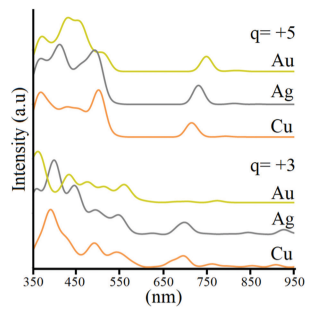


Fig. 3 Simulated optical absorption spectrum for $[M_{13}(dppm)_6]$ (M = Au, Ag, and Cu) species at +5 and +3 charge states. Absorption spectra data were interpolated by applying a Gaussian broadening with a full width of 0.03 eV.

located at 490 and 499 nm, respectively. The main transition is shifted to 409 and 366 nm, for Ag and Cu, denoting a similar pattern expected for the UV/vis spectrum, with a blue-shift of main peaks. At the +3 charge state, the spectrum retains the features shown for the +5 species, with the addition of several lower energy transitions of weak intensity owing to the appearance of $1D^2_{dz2} \rightarrow 1D^0$ excitations, allowed by structural distortions within the M13 core. Hence the main features of the UV-vis spectrum are preserved with the addition of weak lower energy peaks for the 10-ce clusters.

Conclusions

The ultrasmall Au₁₃(dppm)₆ cluster was studied in its +5 and +3 charge states, denoting structural, bonding, and optical property variations, ascribed to the respective 8- and 10-cluster electron counts. For $Au_{13}(dppm)_6^{5+}$, the Au_{13} core deviates from the ideal icosahedral structure, which is increased upon the addition of two electrons at $Au_{13}(dppm)_6^{3+}$, but retaining a similar averaged Aucore-Auico bond distance, accounting for the core radius. For Ag and Cu, the distortion from a perfect icosahedron is to a lesser extent, resulting in a reduced HOMO-LUMO gap for 10-ce (+3 charge state) species, in contrast to the sizable gap in 8-ce. Interestingly, the more significant distortion in Au₁₃(dppm)₆³⁺ enables a further stabilization of the 1Dz2 shell, retaining a sizable HOMO-LUMO gap, despite the intermediate electron count. Thus, $Cu_{13}(dppm)_6^{3+}$ and $Ag_{13}(dppm)_6^{3+}$ species are expected to be less stable owing to the decreased HOMO-LUMO gap.

Ligand-core interactions are larger for gold, followed by Cu, and lastly, Ag species, in agreement with the Au > Cu > Ag bonding energy trend observed for other ligands in monometallic complexes. From 8- to 10-ce, the ligand-core bonding decreases, owing to the two extra electrons in the M_{13} core, for all the species.

Optical properties reveal common features with a small blue-shift of the relevant peaks for lighter coinage metals, which are retained between 8- and 10-ce species, where the latter includes weak low-energy transition owing to $1D_{z^2} \rightarrow 1D$ transitions slightly allowed by the deviations from a perfect icosahedron core. Hence, lighter coinage metal counterparts of the parent $Au_{13}(dppm)_6$ appear as feasible targets for further exploratory synthesis efforts, resulting in similar properties towards alternative ligand-protected clusters as building blocks for nanostructured materials.

Conflicts of interest

There are no conflicts to declare.

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