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Introduction

[Au₁₃(AsPh₃)₈Cl₄]⁺: comparative studies on ligand effect and surface reactivity with its stibine analogue[†]

Jiu-Hong Yu,ª Zhi-Rui Yuan,ª Jing Xu,ª Jin-Gui Wang, ២ a Mohammad Azam, ^c Tian-Duo Li,ª Ying-Zhou Li 🖻 *ª and Di Sun 🖻 *b

Monoarsine-protected icosahedral cluster

Ligand shells of gold nanoclusters play important roles in regulating their molecular and electronic structures. However, the similar but distinct impacts of the homologous analogues of the protecting ligands remain elusive. The C_{2v} symmetric monoarsine-protected cluster $[Au_{13}(AsPh_3)_8Cl_4]^+$ $(Au_{13}As_8)$ was facilely prepared by direct reduction of $(Ph_3As)AuCl$ with NaBH₄. This cluster is isostructural with its previously reported stibine analogue $[Au_{13}(SbPh_3)_8Cl_4]^+$ $(Au_{13}Sb_8)$, enabling a comparative study between them. $Au_{13}As_8$ exhibits a blue-shifted electronic absorption band, and this is probably related to the stronger π -back donation interactions between the Au_{13} core and $AsPh_3$ ligands, which destabilize its superatomic 1P and 1D orbitals. In comparison to the thermodynamically less stable $Au_{13}Sb_8$, $Au_{13}As_8$ achieves a better trade-off between catalytic stability and activity, as demonstrated by its excellent catalytic performance towards the aldehyde–alkyne–amine (A^3) coupling reaction. Moreover, the ligand exchange reactions between $Au_{13}As_8$ with phosphines, as exemplified by PPh₃ and Ph₂P(CH₂)₂PPh₂, suggest that $Au_{13}As_8$ may be a good precursor cluster for further cluster preparation through the "cluster-to-cluster" route.

The icosahedral Au_{13} core is considered a fundamental building block which is commonly observed as the metal core in various structurally well-defined Au nanoclusters.^{1–17} It was theoretically found that the electronic structures of the Au_{13} unit might dictate the physicochemical properties of its assembled clusters.^{18–21} It is therefore of importance to study the influencing factors in depth on the geometric and electronic structures of the individual icosahedral Au_{13} clusters. In this context, the past 15 years have first witnessed a renewed interest in the typical phosphine-protected Au_{13} clusters involving development of new synthetic routes (*e.g.*, HCl-induced convergence),^{6,7,22,23} exploration of formation mechanisms,^{24,25} tuning of electronic properties (*e.g.*, by heterometal doping^{26–29}), and optical properties (*e.g.*, NIR emission^{6,7,30} and chiroptical activity^{31–33}), as well as further expansion of their potential applications (*e.g.*, photodynamic catalysis and nano-electronic devices).^{34,35} More recently, comparative studies on several new types of Au₁₃ analogues protected by other pnictines (*e.g.*, functionalized diphosphine,³⁶ phosphinous acid,³⁷ diarsine,³⁸ and monostibine^{39–41}) or mimics of pnictine (*N*-heterocyclic carbenes)^{42–47} have also been extensively made both experimentally and theoretically.

The large number of studies on icosahedral Au_{13} clusters showed that the stereoelectronic factors of the pnictine/NHC ligands have a non-negligible influence on their electronic structures.^{14,48-54} These ligand-based influencing factors, however, are often intertwined with other factors such as the core geometry and cluster charge state. For example, in 2021, the Konishi group synthesized a pair of dipnictine-protected clusters $[Au_{13}(dppe)_5Cl_2]^{3+}$ (dppe = Ph₂P(CH₂)₂PPh₂) and $[Au_{13}(dpap)_5Cl_2]^{3+}$ (dpap = Ph₂As(CH₂)₃AsPh₂), and demonstrated that the Au_{13} core geometry (deformation), rather than the coordinating P or As atoms, has a more pronounced effect on their absorption and photoluminescence properties.³⁸ In this sense, it is necessary to employ Au_{13} cluster analogues, with essentially the same metal cores and ligand arrangements, as the comparable cases for better understanding the pnictine

^aSchool of Chemistry and Chemical Engineering, Qilu University of Technology (Shandong Academy of Sciences), Ji'nan, 250353, P. R. China. E-mail: liyz@qlu.edu.cn ^bSchool of Chemistry and Chemical Engineering, State Key Laboratory of Crystal Materials, Shandong University, Ji'nan, 250100, P. R. China. E-mail: dsun@sdu. edu.cn

Department of Chemistry, College of Science, King Saud University, P. O. Box 2455, Riyadh 11451, Saudi Arabia

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ligand effect. The monopnictine-protected Au_{13} clusters may serve as good candidates for this purpose since the monopnictines are usually more fluxional in the coordination sphere, and the Au_{13} core geometry is expected to be less affected by the ligands due to the absence of (possibly) strained Au-L'-Austaple/bridge moieties (L' = bridging ligands). Unfortunately, few has been done with the monopnictine-protected Au_{13} clusters, largely due to the lack of comparable analogue clusters with known crystal structures.

To our knowledge, only four monophosphine-protected Au₁₃ clusters have been reported, *i.e.*, $[Au_{13}(PMe_2Ph)_{10}Cl_2]^{3+}$, $[Au_{13}(PMePh_2_8Cl_4^{\dagger}$, $[Au_{13}(POct_3)_8Cl_4^{\dagger}]^+$ ($POct_3 = trioctylphosphine$) and $[Au_{13}(L)_8Cl_4]^+$ (L = P(CH₂CH₂CO₂CH₃)₃), and the last of these could further convert to Au₂₅ nanoclusters by reacting with thiols.^{2,3,7,16} In contrast, much less is known about their heavier analogues. In 2018, we reported the monostibineprotected Au₁₃ cluster [Au₁₃(SbPh₃)₈Cl₄]⁺ (Au₁₃Sb₈),³⁹ which was found more amenable to ligand exchange and also more catalytically active than PPh3-protected Au11 clusters.40,41 Systematic studies on monoarsine-protected Au₁₃ clusters have lagged behind, despite the structurally similar cluster [Au₁₆(-AsPh₃)₈Cl₆] published long ago.⁵⁵ We are thus intrigued by further synthesis of the AsPh₃-protected Au₁₃ clusters, so as to better understand the heavier pnictine ligand effect on their structures and reactivities by comparative studies with their heavier analogue Au₁₃Sb₈.

Herein, we present the synthesis, characterization and reactivity of the monoarsine-protected cluster [Au₁₃(AsPh₃)₈- Cl_{4}^{+} (Au₁₃As₈) which is isostructural with its heavier analogue Au₁₃Sb₈, in terms of the core geometry and ligand arrangement. Comparative studies between them revealed that the blue-shift electronic absorption band of Au13As8 is probably related to the stronger π -back donation interactions between the Au₁₃ core and AsPh₃ ligands, which destabilize its superatomic 1P and 1D orbitals. Moreover, $Au_{13}As_8$ is thermodynamically more stable than its stibine counterparts due to the moderate Au-As coordination bond strength, but its surface arsine ligands are kinetically more labile than phosphines. The unique characteristic of Au₁₃As₈ enables a better trade-off between its catalytic stability and activity, and also makes it an ideal candidate for the precursor cluster in the preparation of other clusters via the "cluster-to-cluster" strategy.

Results and discussion

Synthesis and characterization

Briefly, **Au₁₃As**₈ was prepared by direct reduction of (Ph₃As)AuCl with 0.5 equivalent of NaBH₄ in dichloromethane (DCM)– ethanol mixed solvent (Scheme 1). The identity of the precursor



Scheme 1 Synthetic route to $Au_{13}As_8$

(Ph₃As)AuCl was first confirmed prior to reduction (Fig. S1†). After reduction, the crude $Au_{13}As_8$ could be easily precipitated from its DCM solution by addition of hexane and is proved to be fairly pure by comparison of its IR spectrum with that of the purer crystalline $Au_{13}As_8$, which was obtained from its DCMhexane solution by slow evaporation. In addition, the primary byproduct in this reaction, mainly found in the supernatant separated in the step of using hexane to precipitate the crude $Au_{13}As_8$, was identified as (Ph₃As)₃AuCl by X-ray diffraction analysis (Fig. S2†). This finding provides strong evidence that even minor alterations to the reaction conditions, such as the

product.55 Crystallographic data showed that Au13As8 crystallized in a monoclinic lattice with the $P2_1/n$ space group (Table S1[†]). Two crystallographically independent molecules of Au₁₃As₈, with almost the same bond parameters, are present in the unit cell (Fig. S3[†]). Each comprises an icosahedral Au₁₃ core, eight AsPh₃ ligands and four Cl⁻ ligands (Fig. 1). As is the case with its heavier analogue Au13Sb₈,³⁹ the ligands in Au13As₈ also arranged in an approximate C_{2v} symmetry. The radial Au-Au bond lengths range from 2.726(2) Å to 2.810(2) Å (2.757(2) Å on average), while the peripheral ones are in the range of 2.808(2)-2.985(2) Å (2.900(2) Å on average). The Au-Cl bond lengths fall in the range of 2.31(1)–2.37(1) Å. The Au–As bond lengths range from 2.392(4) Å to 2.414(6) Å, which is longer than that in its precursor complex (Ph₃As)AuCl (2.334(1) Å). As expected, these Au-Au and Au-Cl bond lengths in Au₁₃As₈ are also quite close to the corresponding ones in Au₁₃Sb₈ (Table S2[†]).

type of solvent or quantity of reducing agent used, can have

a considerable impact on the structure of the resulting

The mass spectrum of $Au_{13}As_8$ shows a major peak at m/z = 5151.7672, attributed to the molecular ion $[Au_{13}(AsPh_3)_8Cl_4]^+$ (calcd = 5151.7738), which matches well with its calculated one; the isotopic interval of 1.0 amu support its +1 charge (Fig. 2). Besides, the ³¹P NMR of the product shows a heptet resonance ascribable to PF_6^- (Fig. S4[†]), and its IR spectrum also shows the characteristic absorptions of PF_6^- at 839 and 557 cm⁻¹ (Fig. S5[†]). All these indicate that $Au_{13}As_8$ bears +1 charge, and the counterion is most probably PF_6^- , though it fails to be appropriately located crystallographically, probably due to its severe disorder.

The ¹H NMR spectrum of $Au_{13}As_8$ shows three doublet resonances in the lower magnetic field (*ca.* 7.10–7.50 ppm)



Fig. 1 Crystal structure of $Au_{13}As_8$ with all hydrogen atoms omitted (left: side view; right: top view). Au: orange, As: magenta, Cl: green, C: grey.



Fig. 2 ESI-MS(+) spectrum of $Au_{13}As_8$ in CH_2Cl_2 . Inset: the experimental (black trace) and calculated (magenta trace) isotopic patterns of the $Au_{13}As_8$ ion (A) $[Au_{13}(AsPh_3)_8Cl_4]^+$ and its fragment ion (B) $[Au_{13}(AsPh_3)_7Cl_4]^+$.

region in an integration ratio of 1:2:1, and with a pattern quite akin to that of $Au_{13}Sb_8$ (Fig. S6†). This suggests the presence of three types of chemically non-equivalent AsPh₃ ligands, which is consistent with its solid-state structure; this also further confirms that the icosahedral Au_{13} clusters are stereochemically more rigid than the phosphine-protected Au_{11} clusters.^{56,57}

Theoretical calculations

Since the $Au_{13}As_8$ and $Au_{13}Sb_8$ are isostructural either in the solid state or in solution, they constitute a good comparable pair for comparatively studying the heavier pnictine ligand effect on their structures and reactivities. As shown in Fig. 3, the experimental electronic absorptions for $Au_{13}As_8$ appear at 346 and 439 nm, which are blue-shifted by 6 and 15 nm, respectively, as compared to those for $Au_{13}Sb_8$ at 352 and 454 nm. Interestingly, the absorptions for the previously reported monophosphine-protected Au_{13} clusters are further blue-shifted (Fig. S7†).^{2,3,7} Given the more similar structures of $Au_{13}As_8$ and $Au_{13}Sb_8$, TD-DFT calculations for them were performed at the same level of theory, to reveal the stereoelectronic



Fig. 3 Experimental (solid lines) and TD-DFT calculated (dash lines) UV-vis spectra for $Au_{13}As_8$ (magenta traces) and $Au_{13}Sb_8$ (cyan traces) in CH_2Cl_2 .

influences of the pnictine ligands on the absorptions; the metal cores in both after structure relaxation are more of C_{2v} symmetry, probably transferred from the ligand shells and reflected by the continuous symmetry measure (CSM) value of 0.00 for both.⁵⁸ The optimized Au–Au bond lengths in Au₁₃As₈ are also slightly longer than those in Au₁₃Sb₈ (Table S3†), which is consistent with the crystallographic data. The calculated UV-vis spectra of them are well matched with their respective experimental ones, despite a small blue-shift, probably due to the overestimation of the excitation energies for the functional employed (Fig. S8 and S9†). Thus, the calculated peaks at 335 and 417 nm for Au₁₃As₈ (340 and 435 nm for Au₁₃Sb₈) should correspond to its experimental ones at 346 and 439 nm (352 and 454 nm for Au₁₃Sb₈), respectively.

Au₁₃As₈ and Au₁₃Sb₈ possess a similar electronic structure, with a HOMO-LUMO gap of 3.32 and 3.09 eV, respectively. The HOMO, HOMO - 1 and HOMO - 2 orbitals are typical of 1P superatomic orbitals while LUMO through LUMO + 4 show 1D characteristics (Tables S4 and S5[†]). The long-wavelength absorptions (417 nm for Au₁₃As₈ and 435 nm for Au₁₃Sb₈) are mainly ascribable to metal-to-metal charge transfer (MMCT) transitions from 1P to 1D shell, while the shorter-wavelength absorptions (335 nm for Au₁₃As₈ and 340 nm for Au₁₃Sb₈) are mainly attributed to ligand-to-metal charge transfer (LMCT) transitions to 1P orbitals from higher-lying occupied orbitals (HOMO - 3 to ca. HOMO - 10). In addition, the absorptions below 300 nm for both result from the even higher-energy transitions, such as HOMO - n (n > 15) to 1D, or HOMO - n(n < 5) to LUMO + n (n > 10); the phenyl rings in the ligands contributed significantly to HOMO - n (n > 15) orbitals with energy lower than -8.2 eV, and LUMO + n (n > 10) orbitals with energy higher than -2.2 eV (Fig. 4a, b and Tables S6, S7[†]).

As shown in Fig. 4c, on average, the 1P orbitals of $Au_{13}As_8$ comprise Au 5d/6sp orbitals (ca. 64%), As 4sp orbitals (ca. 13%), Cl 3sp orbitals (ca. 9%) and C 2sp orbitals (ca. 14%). The 1D orbitals are also mainly composed of Au atomic orbitals (Au: As: Cl: C = 72: 13: 3: 12). The proportions of Au and As in orbitals from HOMO - 3 to HOMO - 10 decreased to a certain extent (48% for Au and 7% for As), but those of Cl and phenyl rings increased (21% for Cl and 24% for C). This means that, for a given type of cluster $[Au_{13}(ER_3)_8Cl_4]^+$ (E = pnictogen), the R substituent of ER₃ and the inorganic Cl⁻ may less affect the longer-wavelength absorptions resulting from 1P to 1D transitions, due to their less contributions to 1P/D orbitals; in other words, the shorter-wavelength absorptions are more susceptible to the stereoelectronic property of the ER₃ ligand. Consistent with this are the experimental UV-vis spectra of $[Au_{13}(L)_8Cl_4]^+$ (L = PPh₂Me; P(CH₂CH₂CO₂CH₃)₃),^{3,16} which show almost the same absorption peaks at 432 nm in the lower-energy region but obviously different peaks at 342 nm and 333 nm, respectively, in the higher-energy region (Fig. S7[†]).

The orbital composition pattern of $Au_{13}Sb_8$ is akin to that of $Au_{13}As_8$. The 1P orbitals are composed of Au 5d/6sp (*ca*. 63%), Sb 5sp (*ca*. 13%), Cl 3sp (*ca*. 13%) and C 2sp (*ca*. 12%) orbitals. The 1D orbitals also mainly comprise Au atomic orbitals (Au : Sb : Cl : C = 67 : 17 : 4 : 11). The proportions of Au and As from HOMO - 3 to HOMO - 10 also decreased (44% for Au and 6%

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Fig. 4 Total density-of-states (TDOS) and partial density-of-states (PDOS) spectra of (a) $Au_{13}As_8$, (b) $Au_{13}Sb_8$, and (c) their selected Kohn–Sham molecular orbitals showing the populations of atomic orbitals.

for Sb), but those of Cl and phenyl rings increased (29% for Cl and 20% for C). In comparison, the superatomic 1P, 1D and the higher-lying occupied orbitals (*e.g.* HOMO – 3 to HOMO – 10) in $Au_{13}As_8$ all up-shifted relative to their counterparts in $Au_{13}Sb_8$, except for the HOMO in $Au_{13}As_8$, which is marginally lower in energy by 0.025 eV than that in $Au_{13}Sb_8$.

Previous studies with thiol/alkynyl-protected Au_{25} clusters based on the jellium model assumed that the more electronegative (usually X-type) ligands bonded to the Au_{13} core might lead to the reduction of potential volume for the confinement of Au_{13} core valence electrons, and result in the destabilization of the superatomic 1P orbitals.¹⁵ In this work, the L-type twoelectron-donor pnictine ligands are more suitable to be classified by overall electron-donating ability. AsPh₃ is a stronger σ - donating and also a better π -accepting ligand than SbPh₃;⁵⁹ it is thus envisioned that both interactions may influence the superatomic cluster orbitals. The π -back donation from the Au core to the π -acid ligand is not uncommon, for instance, Au(5d) \rightarrow CO(π^*) in the carbonyl-protected Au cluster,⁶⁰ and Au(5d) \rightarrow N=C(π^*) in the guanine-protected Au₁₃ cluster.⁶¹

To get more insight into this, the natural bond orbital (NBO) analyses were performed. The qualitative natural population analyses (NPA) show that the Au₁₃ core in Au₁₃As₈ is less negatively charged than that in $Au_{13}Sb_8$ (-0.50 vs. -1.43, Table $S9^{\dagger}$), as is also the case with the Mulliken charge analysis (-1.95) vs. -2.27, Table S10[†]). This implies that, given the stronger σ donating ability of AsPh₃, a stronger electron back donation from the Au₁₃ core to AsPh₃ may contribute to the relatively lower electron density for the Au13 core in the case of Au13As8. As expected, the second-order perturbation theory analyses in NBO basis for Au13As8 show strong interactions between the central Au lone pair orbitals and the radial Au–As σ^* orbitals, as reflected by the large second-order perturbation stabilization energies E(2) (Table S11[†]). In comparison, this type of interaction in Au₁₃Sb₈ is much weaker (Table S12[†]). This is probably associated with the fact that the As 4sp orbitals involved in the Au–As σ^* orbitals are less diffuse, lower in energy, and thus more electrophilic, compared to the Sb 5sp orbitals in Au–Sb σ^* orbitals. Since the stronger π -back donation interactions between Au and Au–As σ^* orbitals would reduce the Au–As bond orders more significantly, the Wiberg bond indices for the Au-As bonds in $Au_{13}As_8$ are indeed smaller (ca. 0.46) than those for the Au-Sb bonds in Au₁₃Sb₈ (ca. 0.54), whereas those for the innocent Au-Cl bonds in both are almost the same (ca. 0.42) (Table S13[†]). In addition, other types of π -back donation interactions between surface Au lone pairs and As/Sb-C σ^* orbitals are rather weak and comparable for both Au₁₃As₈ and Au13Sb8 (Tables S11 and S12[†]). Based on these, the upshift of 1P orbitals in Au₁₃As₈ relative to that in Au₁₃Sb₈ is tentatively accounted for by the stronger π -back donation interactions between central Au and radial Au–As σ^* orbitals, by which the better π -accepting L-type ligand AsPh₃ exerts a similar impact on the Au₁₃ core valence electrons to that for the case of the more electronegative X-type ligand.

On the other hand, the correlations of 1D orbitals with the electronegativity of ligands are rather complicated, though they shift to the same direction as 1P orbitals in most cases but lack an obvious trend in the shift amount.¹⁵ In this work, the 1D orbitals in $Au_{13}As_8$ up-shifted more (by 0.156 eV on average) than 1P orbitals (by 0.031 eV on average). That is to say, the 1D orbitals are more susceptible to the electronic factors of the pnictine ligands. In view of this, more comparative studies, including with the lighter phosphine-protected analogue clusters, may be of interest to better explain the behavior of 1D orbitals with respect to the integrated σ -donating/ π -accepting parameters of the L-type pnictine ligands.

Optical properties

The solution photoluminescence (PL) of $Au_{13}As_8$ emits at *ca*. 802 nm, which is red-shifted in wavelength and also higher in



Fig. 5 Photoluminescence spectra for Au₁₃As₈ (magenta line, $\lambda_{ex} = 460 \text{ nm}$) and Au₁₃Sb₈ (cyan line, $\lambda_{ex} = 470 \text{ nm}$) in CH₂Cl₂ (ca. 2.0 mg mL⁻¹) at room temperature.

intensity, as compared to the main emission of $Au_{13}Sb_8$ at *ca*. 740 nm (Fig. 5). This is presumably due to the heavy metal effect of Sb, which facilitates the reverse intersystem crossing (RISC) process from the T₁ state to the S₁ state and therefore leads to the blue-shift and lower intensity of the emission for $Au_{13}Sb_8$.^{62,63} The previously reported two phosphine-protected clusters $[Au_{13}(L^P)_nCl_4]^+$ (L^P = trioctylphosphine (POct₃), n = 8; L^P = Ph₂P(CH₂)₃PPh₂, n = 4) both emit at *ca*. 775 nm.⁷ Considering the likely dual emissions of $Au_{13}Sb_8$, if the shoulder at *ca*. 825 nm is of the same phosphorescence character as the emissions of $Au_{13}As_8$ and $[Au_{13}(L^P)_nCl_4]^+$, the emission wavelengths would be in the order of $Au_{13}Sb_8 >$ $Au_{13}As_8 > [Au_{13}(L^P)_nCl_4]^+$.

 $Au_{13}As_8$ in its crystalline state exhibits a PL emission over a wider range of wavelengths (from *ca*. 650 nm to *ca*. 850 nm), and the emission maximum at *ca*. 740 nm is blue-shifted by *ca*. 60 nm compared to that in solution. The PL emission of $Au_{13}Sb_8$ in the crystalline state displays a similar but also blue-shifted weak dual-emission-like pattern to that in solution, with an emission peak centered at *ca*. 805 nm and a shoulder at *ca*. 710 nm (Fig. S10†). In general, the PL blue-shift for both can be partially attributed to the rigidity enhancement of the structures in the solid state. In addition, the largely broadened and blueshifted solid PL spectrum of $Au_{13}As_8$, somewhat resembling that of $Au_{13}Sb_8$, also suggest that a similar dual-emission phenomenon to that of $Au_{13}Sb_8$ may be present in its solid/ crystalline state.

Thermal stability

The $Au_{13}As_8$ is rather stable in the solid state, but gradually decomposes with time in solution at room temperature, although such a process is much slower than for the case of $Au_{13}Sb_8$. Lower temperature could effectively inhibit the decomposition of $Au_{13}As_8$; its UV-vis monitoring spectra showed no essential decomposition even after 12 days when its DCM solution was stored at 4 °C (Fig. S11†). Thermogravimetric

analysis (TGA) of $Au_{13}As_8$ at a heating rate of 10 °C min⁻¹ under N₂ purge (60 mL min⁻¹) showed that the rapid weight loss started at *ca*. 200 °C, whereas that for $Au_{13}Sb_8$ was observed at 155 °C (Fig. S12†).³⁹ These results suggest that $Au_{13}As_8$ is more stable than $Au_{13}Sb_8$ either in solution or in the solid state but most probably still less stable than the phosphine-protected analogue clusters; this is believed to result from the moderate Au-As bond strength.

Surface reactivity-catalysis

Since the electron-rich Au₁₃ core was speculated to activate the C=O bond,^{36,64,65} the pnictine-protected Au₁₃ clusters may find uses in the aldehyde/ketone-involved catalytic reactions. Au₁₃Sb₈ was found to be able to efficiently catalyze the aldehyde-alkyne-amine (A^3) coupling reaction, but the drawback of lower catalytic stability could also not be neglected.40 To better realize the trade-off between stability and activity, the catalytic performance of Au₁₃As₈ towards the same A³ coupling reaction was tested (Table 1). It was found that, despite its relatively stronger Au-As coordination bonds, Au13As8 could also achieve a comparable conversion ratio to Au₁₃Sb₈ after 12 h, probably associated with different reaction kinetics. More importantly, Au₁₃As₈ is catalytically more stable than Au₁₃Sb₈; it can be simply recovered (and also purified) from the reaction mixture by precipitation with hexane (Fig. S13[†]), and can be reused for five cycles with only slight decrease in catalytic activity (Fig. S14 and S15[†]). This finding further underlines the important role of ligands in tuning the catalytic performance and may provide some implications for the design of more practically useful ligand-protected metal nanocluster catalysts.

Surface reactivity-ligand exchange

The exquisite "cluster-to-cluster" synthetic strategy based on ligand exchange has recently become increasingly prevailing.^{47,66,67} Taking Au_{13} clusters as examples, reactivity studies on $Au_{13}Sb_8$ or its bromo-analogue $[Au_{13}(SbPh_3)_8Br_4]^+$ had shown

Table 1 A³ coupling reactions catalyzed by Au₁₃As₈ or Au₁₃Sb₈^a Catalyst (0.5 mol%) r.t. 12 h neat reaction NMR yield (%) Catalyst^b Cycle 1 Cycle 2 Cycle 3 Cycle 4 Cycle 5 Au13As8 73 74 69 64 62 Au13Sb8 81 32

^{*a*} Benzaldehyde (1.0 mmol), piperidine (1.2 mmol), phenylacetylene (1.3 mmol), solvent-free reaction. ^{*b*} With respect to the amount of benzaldehyde. ^{*c*} Given the obvious decomposition in each cycle for Au₁₃Sb₈, no further trial was made after the second cycle due to the marked decrease in yield.



that they could readily react with glutathione (GSH) or 1-adamantanethiol (S-Adm) to give the achiral Au₂₅(SG)₁₈ or chiral Au₁₈(SAdm)₈(SbPh₃)₄Br₂, respectively.^{39,41} The Wang group also developed a new method of synthesizing thiol-protected Au₂₅ clusters through reactions of $[Au_{13}(L'')_8Cl_4]^+$ $(L'' = P(CH_2CH_2 (CO_2CH_3)_3$ with appropriate thiols.¹⁶ In analogy to these stibine or phosphine-protected Au13 clusters, Au13As8 may also possess the potential as a precursor cluster in the syntheses of other new clusters by reacting with stronger coordinating ligands. Thus, the reactions of Au₁₃As₈ with monophosphine PPh₃ or diphosphine Ph₂P(CH₂)₂PPh₂ (dppe) at room temperature were carried out as a proof-of-concept in this work. As shown in Scheme 2, two well-known clusters $[Au_{11}(PPh_3)_8Cl_2]^+$ and $[Au_{13}(dppe)_5 Cl_2$ ³⁺ were obtained in the cases of PPh₃ and dppe, respectively, and their structures were determined by comparison of their UV-vis, ¹H NMR and MS spectra with those from literature (Fig. S16-S20[†]).^{7,68} These preliminary trials confirm the potential use of Au13As8 in the "cluster-to-cluster" syntheses. In comparison, the reactions with Au₁₃As₈ proceed more slowly than with Au₁₃Sb₈, indicating that it is more likely for the former to realize more controlled derivatization or functionalization by ligand engineering.

Conclusions

In this work, a monoarsine-protected icosahedral cluster $[Au_{13}(AsPh_3)_8Cl_4]^+$ (Au₁₃As₈) was prepared. It is isostructural with its previously reported stibine analogue $[Au_{13}(SbPh_3)_8Cl_4]^+$ $(Au_{13}Sb_8)$, which provides an opportunity to better comparatively study the heavier pnictine ligand effect on their electronic structures, optical properties, and surface reactivities. The influencing mechanism of the L-type pnictine ligands on the Au₁₃ superatomic orbitals may be different from those of the Xtype ligands (e.g., organic thiol, selenol, alkynyl). The superatomic 1P and 1D orbitals of Au13As8 are up-shifted relative to those of Au₁₃As₈, which is tentatively ascribed to the stronger π back donation interactions between the Au₁₃ core and AsPh₃ ligands. In addition, compared to the thermodynamically less stable Au₁₃Sb₈, a better trade-off between stability and activity was realized with Au13As8, as reflected by its good catalytic performance towards the A³ coupling reaction. Finally, the ligand exchange reactions between Au₁₃As₈ with phosphines, as exemplified by PPh₃ and Ph₂P(CH₂)₂PPh₂, suggest that Au₁₃As₈ may be a good precursor cluster and may find more use in the preparation of other clusters via the prevailing "cluster-tocluster" strategy. Taken together, the heavier organometalloidal arsines or organometallic stibines, especially the latter, are not simply "weaker organic phosphines", and they are more likely

to exert distinct ligand effects on the structures and properties of metal nanoclusters. Since there has been only a limited number of heavier pnictine-protected metal nanoclusters, more diverse syntheses of such type of clusters, as well as explorations of their physicochemical properties and applications, are of much interest and deserve deeper scrutiny.

Data availability

All experimental and computational data associated with this article have been included in the main text and ESI. \dagger

Author contributions

Y.-Z. L. and D. S. conceived the manuscript; J. H. Y. performed the experiments; J.-H. Y., Z.-R. Y., J. X., J.-G. W., M. A., T.-D. L. and Y.-Z. L. analyzed data, prepared figures and provided conceptual contributions; J. H. Y., Y. Z. L. and D. S. wrote the manuscript with contributions from all co-authors.

Conflicts of interest

There are no conflicts to declare.

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