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Silyl- and germyl-bridged neutral square-planar Ag₄ clusters with short Ag–Ag distances exhibiting red emission†

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We report silyl- and germyl-bridged neutral square-planar Ag₄ clusters with very short Ag–Ag distances (2.695 and 2.704 Å), as revealed by single-crystal X-ray diffraction analysis. On the basis of the results of theoretical calculations, we attribute these short distances to attractive Ag–Ag interactions that reduce the optical energy gap, resulting in a red emission reaching wavelengths of up to 700 nm. These wavelengths are among the longest observed for emissions from Ag₄ clusters.

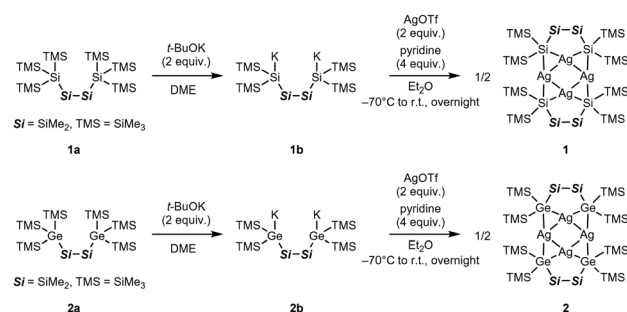
Considerable attention has been given to multinuclear transition metal molecular clusters because of their unique electronic states. These states arise from intermetallic interactions between metal centres and distinguish these multinuclear clusters from their mononuclear counterparts. Specific ligand designs can be used to produce multinuclear clusters with distinct electronic states, resulting in unique structural, optical, and reactive properties.¹ Group 11 coinage metals (Cu, Ag, and Au) in the d¹⁰ monovalent state are known to form multinuclear cluster structures with non-covalent intermetallic interactions.² Structures with d¹⁰ Ag(I) centres form various multinuclear clusters, where Ag₄ clusters are among the stable forms frequently observed in silver cluster chemistry.³ The Cambridge structural database (CSD) contains 157 structures determined by X-ray diffraction (XRD) for Ag₄ clusters with Ag–Ag interactions (Table S1, ESI†). These clusters have been studied both for their structural diversity and potential applications in chemical reactions and sensing.⁴

From a photophysical perspective, intermetallic distances within clusters play a crucial role in determining the electronic and optical properties of clusters.⁵ Short intermetallic distances

enhance orbital interactions, modifying the electronic structure and narrowing the energy gap (ΔE_g) between high-lying occupied dσ* orbitals and low-lying empty pσ/σ orbitals.⁶ Consequently, the excited states and emission properties are changed. Although d¹⁰ Cu(I) and Au(I) tetranuclear clusters readily exhibit long-wavelength red emissions,^{1a} Ag₄ clusters with emission maxima (λ_{MAX}) exceeding 600 nm are rare. This rarity is primarily due to Ag(I) complexes having larger ΔE_g s than Cu(I) and Au(I) complexes^{6,7} because ligand-centred transitions often dominate over transitions involving metal centres.

In this study, we developed two square-planar Ag₄ clusters bridged by silyl and germyl ligands. The clusters have exceptionally short Ag–Ag distances and exhibit a deep red emission around 700 nm. We performed a single-crystal XRD (SC-XRD) analysis and theoretical calculations to investigate the intermetallic interactions contributing to these distinctive structural properties and electronic states. Additionally, we examined the origin of these long-wavelength emissions by comparing them with previously reported Ag₄ clusters. We present results showing how intermetallic interactions produce red-shifted emissions in this communication.

The Ag₄ clusters **1** and **2** were synthesised as shown in Scheme 1. Dipotassium salts (**1b** and **2b**) were prepared, in a manner similar to the foundational procedure developed by Marschner and co-workers,⁸ by reacting the silyl compound **1a**



Scheme 1 Synthesis of the Ag₄ clusters **1** and **2**.

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† Electronic supplementary information (ESI) available: Summary of CSD search results for previously reported Ag₄ clusters and details of cluster synthesis, theoretical calculations, and photophysical measurements. CCDC 2400625 and 2400626. For crystallographic data in CIF or other electronic format see DOI: <https://doi.org/10.1039/d4cc06105e>



and germyl compound **2a** with 2 equiv. of *t*-BuOK in 1,2-dimethoxyethane (DME), followed by treatment with 2 equiv. of AgOTf in Et₂O under -70°C . The corresponding Ag₄ clusters were selectively obtained, as confirmed by ¹H NMR. It was necessary to add 4 equiv. of pyridine as an auxiliary ligand to improve the cluster yield (see ESI†); otherwise, only a trace quantity of **2** was obtained. Recrystallisation from Et₂O produced both **1** and **2** as orange crystals in yields of 52% and 36%, respectively. Both clusters were fully characterised by SC-XRD and elemental analysis, as well as by ¹H, ¹³C, and ²⁹Si NMR (Fig. S3–S10, ESI†).

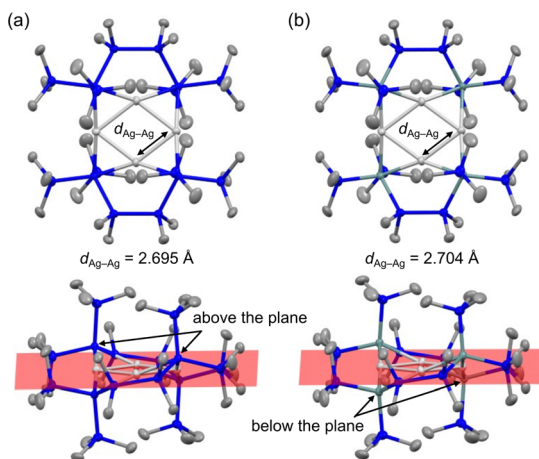


Fig. 1 ORTEP drawings (50% probability level) of (a) **1** and (b) **2** from a top-down view. The lower images show the corresponding side profiles with the Ag₄ plane highlighted in red. The blue, pale teal, silver, and grey balls represent Si, Ge, Ag, and C atoms, respectively. The H atoms are omitted for clarity.

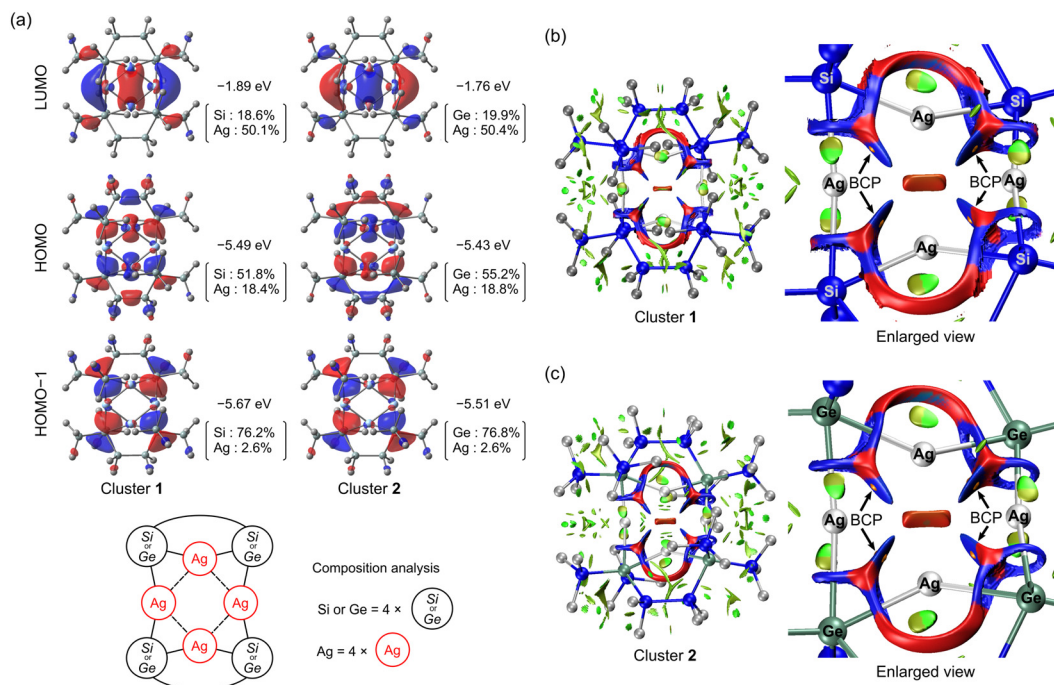


Fig. 2 (a) Molecular orbitals and their composition determined by natural atomic orbital analysis. RDG isosurface map plotted at an isosurface value of 0.3 with four BCPs (shown by orange spheres) observed along the Ag–Ag bond in clusters (b) **1** and (c) **2**.



energy of the HOMO–1 is close (within 0.2 eV) to that of the HOMO; therefore, the HOMO–1 is also shown. The HOMO and HOMO–1 are largely localised on the silyl and germlyl ligands. The HOMO is partially distributed on two opposing Ag atoms, and the LUMO is predominantly localised on the Ag atoms and their spatial centre. Natural atomic orbital (NAO) analysis revealed that the HOMO and HOMO–1 mainly originate from the four terminal Si or Ge atoms bonded to four Ag atoms and are composed of Si 3p and Ge 4p orbitals (Tables S4–S7, ESI†). The Ag orbitals contribute minimally (2.6%) to the HOMO–1 for both clusters but non-negligibly to the HOMO (approximately 18.4% for cluster 1 and 18.8% for cluster 2), indicating the important role of Ag. Specifically, the contributing Ag orbitals are 5s–4d hybridised orbitals. The contribution from the Ag atoms to the LUMO is 50.1% (cluster 1) and 50.4% (cluster 2) and mainly involves the 5s and 5p orbitals. A Hirshfeld analysis for the LUMO is provided in the ESI† that validates the NAO analysis.

The noncovalent d^{10} Ag intermetallic interactions in our clusters were characterised by performing a reduced density gradient (RDG) analysis¹⁰ and Bader's quantum theory of atoms in molecules (QTAIM) analysis¹¹ using Multiwfn software.¹² Electronic wavefunctions were generated based on the aforementioned structures optimised using DFT. The same level of theory was employed as in a recent study on metal–metal (M–M) interactions in d^{10} coinage-metal clusters, including Ag_4 clusters.¹³ In the referenced study, red RDG isosurface maps were reported between two metal centres, indicating repulsive interactions. In our case, as shown in Fig. 2 and Fig. S13 (ESI†), we observed blue RDG isosurface maps, indicating that the interactions between the Ag centres are as strong as hydrogen bonds, while red regions were observed at the centre of the Ag–Si (or Ge)–Ag triangles and the cluster centre. There are bond critical points (BCPs)^{11b} within the blue region, suggesting an attractive interaction. Table 1 shows representative values of the topological parameters of the electron density $\rho(r)$ at the BCPs. These values are consistent for the four BCPs in each cluster. The ratio of the Lagrangian kinetic energy (G_{CP}) to the absolute potential energy density ($|V_{CP}|$) is often used as a reliable indicator for characterising the type of interaction.¹⁴ Our clusters have $|V_{CP}|/G_{CP}$ values of ~ 1.25 , indicating that the Ag–Ag interaction is not purely closed-shell but possesses some electron-shared (covalent) character. We used a previously reported method¹⁵ to estimate the strength of argentophilic interactions (E_{int}) on the basis of the ρ values at a BCP as $-23.7 \text{ kcal mol}^{-1}$ (cluster 1) and $-22.9 \text{ kcal mol}^{-1}$ (cluster 2). The ρ values for our clusters are larger than the range reported in the referenced study, which may reduce the accuracy of the estimated E_{int} . However, the larger ρ and E_{int} values calculated for our clusters indicate strong

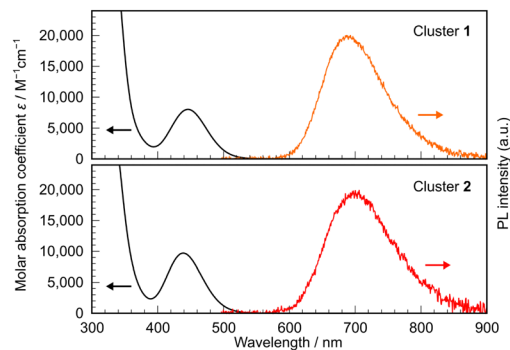


Fig. 3 UV-vis absorption and PL spectra of clusters 1 and 2.

argentophilic interactions in our systems. The underlying reasons for this enhanced interaction strength will be investigated in future studies.

To investigate the effect of the short d_{Ag-Ag} s of 1 and 2 on the photophysical performance of these clusters, a series of photophysical measurements was performed on solutions of 1 and 2 in methylcyclohexane. Fig. 3 shows the corresponding UV-vis absorption spectra. The spectra of clusters 1 and 2 contain broad absorption bands around 450 nm, with peak maxima (λ_{abs}) at 446 nm and 439 nm, respectively. We performed time-dependent (TD)-DFT calculations at the same level of theory as the structural optimisations. The results revealed that these absorptions are mainly attributable to the HOMO $\rightarrow$ LUMO-based $S_0 \rightarrow$ lowest singlet excited state (S_1) transition, with a smaller contribution from the HOMO–1 $\rightarrow$ LUMO-based $S_0 \rightarrow$ second lowest singlet excited state (S_2) transition. This result is obtained because the oscillator strength (f) for the $S_0 \rightarrow S_1$ transition (0.0698 for 1 and 0.0862 for 2) is considerably larger than that for the $S_0 \rightarrow S_2$ transition (0.0090 for 1 and 0.0119 for 2) (see the ESI† for details), even though the S_2 energies (2.99 eV for 1 and 2.97 eV for 2) are close to those of S_1 (2.78 eV for 1 and 2.85 eV for 2). Neither cluster exhibits noticeable emissions at room temperature but both clusters exhibit bright red emissions in glassy solutions at the temperature of liquid nitrogen. Fig. 3 shows the PL spectra of both solutions containing broad, single-peaked emissions with λ_{MAX} at approximately 690 nm (1) and 700 nm (2), which are among the longest wavelengths recorded for emissions from Ag_4 clusters to date. The PL quantum yields (Φ_{PL}) are 0.63 (1) and 0.50 (2), and the PL lifetimes (τ) are 102 μ s (1) and 58 μ s (2) (Fig. S13, ESI†). Given the large Stokes shift, even in frozen matrices, and the relatively long τ , these emissions can be assigned to phosphorescence. The phosphorescence rate constants (k_r^T s) are $6.1 \times 10^3 \text{ s}^{-1}$ (1) and $8.7 \times 10^3 \text{ s}^{-1}$ (2), whereas the non-radiative rate constants (k_{nr}^T s) are $3.6 \times 10^3 \text{ s}^{-1}$ (1) and $8.7 \times 10^3 \text{ s}^{-1}$ (2). Interestingly, both k_r^T and k_{nr}^T are larger for the Ge-containing cluster 2 than for the Si-based cluster 1. Considering that k_r^T and k_{nr}^T are rate constants for spin-forbidden transitions, the higher values of these constants for 2 than 1 can be attributed to the so-called heavy atom effect of Ge. A plausible reason for the relatively high Φ_{PL} s (resulting from the small k_{nr}^T) is the presence of bulky $-SiMe_3$ ligands densely packed around the clusters (Fig. S11

Table 1 Topological parameters at the BCPs appearing between the metal atoms in clusters 1 and 2

	Cluster 1	Cluster 2
Density of all electrons: ρ	0.041	0.040
Laplacian of electron density: $\nabla^2\rho$	0.090	0.091
Lagrangian kinetic energy: G_{CP}	0.030	0.030
Potential energy density: V_{CP}	−0.038	−0.038
Energy density: H_{CP}	−0.008	−0.008
$ V_{CP} /G_{CP}$	1.254	1.250



and S12, ESI†), which may contribute to restricting large-scale structural rearrangements.

Importantly, both complexes exhibit red emissions that are comparable to the longest-wavelength emissions recorded for Ag₄ clusters. To explore the reason for this result, we compared our results with those for previously reported emissive clusters. Among the 157 CSD entries, 32 Ag₄ clusters have been reported to exhibit emissions, as summarized in Table S1 (ESI†). Most of the emissions of these Ag₄ clusters originate from $\pi\pi^*$ transitions of the ligands, indicating a large ΔE_g for Ag centres (as mentioned in the Introduction), even for those forming tetra-nuclear clusters. For some clusters with an excited state involving a metal centre, such as the states involved in ligand-to-metal charge transfer¹⁶ and metal-metal-to-ligand charge-transfer,¹⁷ emissions with longer wavelengths than those associated with $\pi\pi^*$ transitions have been reported. However, clusters with red emissions approaching wavelengths of 700 nm are scarce. In the solid state, where molecules are aggregated, six reported complexes have been reported with λ_{MAX} beyond 600 nm—five at cryogenic temperatures, one at 298 K, and another at both 298 K and 77 K.^{17,18} Among these clusters, the longest λ_{MAX} of 707 nm (77 K) was reported for a heterometallic Pt₂Ag₄ cluster.^{18a} By contrast, in solution, where molecules are isolated, only three clusters have been reported to exhibit λ_{MAX} s beyond 600 nm (at 298 K or ambient temperature), with the longest-wavelength emissions reaching 628 nm at most.^{16,18c} The observed emission wavelengths of our clusters are only shorter than those of the Pt-containing cluster and correspond to the longest λ_{MAX} recorded in an isolated state. On the basis of our DFT calculations, this long-wavelength phosphorescence emission is attributed to a LUMO $\rightarrow$ HOMO-based transition with considerable involvement of Ag orbitals, particularly because of the cluster-centred σ -type LUMO. We confirmed that, in such cases, the LUMO energy decreases as $d_{\text{Ag-Ag}}$ decreases in a square-planar Ag₄ cluster (Fig. S14, ESI†). Although the long-wavelength emission results primarily from phosphorescence (rather than fluorescence), which induces a large Stokes shift as previously discussed, the low-energy LUMO of our clusters, realised by the short $d_{\text{Ag-Ag}}$, likely contributes to the further red-shift of the emission wavelength.

To broaden the scope of this study, we performed theoretical calculations on cluster 1 in which the Ag atoms were replaced with the congeners Cu and Au. The calculated Cu–Cu and Au–Au distances are short (2.49 Å and 2.75 Å, respectively), and the corresponding phosphorescence energies are 1.00 eV (for the Cu₄ cluster) and 1.06 eV (for the Au₄ cluster), reaching the near-infrared (NIR) region. These values are notably lower than that of the Ag₄ cluster 1 (1.68 eV). These theoretical results provide further evidence of the versatility of our molecular design concept, demonstrating that short M–M distances effectively induce red-shifted emission by enhancing M–M interactions.

In conclusion, we developed Ag₄ clusters with short Ag–Ag distances that exhibit red emission reaching an λ_{MAX} of 700 nm. Our experimental and theoretical investigations underscore the role of both Ag–Ag distances and interactions in influencing the

electronic structure and photophysical properties of Ag₄ clusters. Subsequent theoretical calculations suggest that employing congeners, such as Cu or Au, could yield even longer-wavelength emissions reaching the NIR region. Therefore, this study deepens our fundamental understanding of Ag clusters and broadens our insight into the molecular design of coinage-metal-based emitters.

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Data availability

Crystallographic data for clusters 1 and 2 have been deposited at the CCDC under No. 2400625 and 2400626, respectively. Data supporting the results presented in this article are presented in ESI.†

Conflicts of interest

There are no conflicts to declare.

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