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Linker driven site-specific catalysis in atomically precise silver cluster assemblies†

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Metal nanoclusters (NCs) exhibit potential as catalysts for electrochemical studies, providing atomic-level insights into mechanisms. However, it remains elusive to construct an integrated catalyst with a molecular-level understanding of its mechanism, especially in silver cluster assemblies. In this study, we have shown that atomically precise Ag_{12} cluster assemblies Ag_{12} -py, Ag_{12} -pyz, Ag_{12} -bpy, Ag_{12} -bpa, Ag_{12} -azopy, (where Ag_{12} = secondary building unit, Py = pyridine, Py = pyrazine, Py = pyrazine units: the cluster core and the linkers. The atomic resolution of such catalysts allows tracing the reaction process Py = pyrazine units: the cluster core and the linkers. The atomic resolution of such catalysts allows tracing the reaction process Py = pyrazine units: the cluster assembly and linker synergy can be accurately elucidated to dominate in the series. Taking advantage of the pyrazine linker due to its lower basicity and the isotropic nature of intercluster interactions in Py = pyrazine linker due to its lower basicity and selective hydrogen adsorption at the sulfur site, different from others in the series with nearly five times higher efficiency. This work on a series of silver cluster assemblies provides a substantial structural model to understand the catalyst's active site and activity, further driving advancements in functional cluster-based assemblies.

Introduction

Atomically precise noble metal nanoclusters (NCs), which bridge molecular organometallic complexes and plasmonic metal nanoparticles (NPs), 1,2 comprising only a few atoms, allow excellent control over the total structure, exhibiting unique electronic and optical properties, including molecular-like transitions, tunable photoluminescence, and high catalytic activity. 3-5 The manipulation of these properties is primarily achieved through control over the cluster core and protective ligands. The distinctive geometry and electronic structure make them particularly attractive in catalysis, due to their high

surface-to-volume ratio, atomic precision structure, rich surface chemistry, and low coordination number.⁵⁻⁸ These clusters can indeed be used as ideal model catalysts for correlating catalytic properties with their atom packing structures.⁸⁻¹² However, the major challenge is the insufficient stability leading to aggregation. Therefore, it requires strategies to stabilize NC structures while preserving cluster integrity.¹³⁻¹⁶

To address these challenges, recent progress has been made in utilizing NCs as nanoscale building blocks for superstructures^{17–19} through assembly processes by employing bridging ligands, known as cluster-assembled materials (CAMs).^{20–22} These CAMs not only enhance the stabilization of NCs but also merge the functionalities of NCs and linkers.²³ They exhibit superior properties and multiple tunable components, compared to their cluster building blocks.^{24–26} This approach provides a potential solution for stabilizing NCs while simultaneously enhancing catalytic performance by facilitating improved electron transport and chemical interactions between clusters.

The electrocatalytic hydrogen evolution reaction (HER) is a fascinating and sustainable strategy to produce molecular hydrogen, which is considered an excellent fuel for the future. Hydrogen has high energy density and produces water as the sole byproduct upon combustion, making it an ideal candidate to replace fossil fuels, which cause many detrimental

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environmental issues.^{27–29} Despite the inherent inertness or low HER activity of coinage metals such as gold, silver, and copper, the nanoscale regime introduces distinctive material properties, deviating from their bulk counterparts and offering potential avenues for enhanced catalytic activity.^{30–34}

Ligand-protected metal NCs, such as Ag, Au, and Pt, have been minimally explored as catalysts, 13,35-39 and their assembled materials remain largely unstudied.40-43 A more in-depth investigation is needed to understand the relationship between the structure of the electrocatalyst and catalytic performance at the atomic level.44-50 Clusters with identical surface ligands and core atoms, but different linkers, offer a unique opportunity to isolate and study the factors influencing overall performance. In this context, we focus on the synthesis and catalytic application of five silver cluster assemblies such as Ag_{12} -py $\{[Ag_{12}(C_6H_{11}S)_6(CF_3COO)_6(C_5H_5N)_6]\cdot$ $4H_2O_{n}^{25}$, Ag_{12} -pyz $[Ag_{12}(C_6H_{11}S)_6(CF_3COO)_6(C_4H_4N_2)_3]_n$, Ag_{12} - $[Ag_{12}(C_6H_{11}S)_6(CF_3COO)_6(C_{10}H_8N_2)_3]_n$ Ag₁₂-bpa $[Ag_{12}(C_6H_{11}S)_6(CF_3COO)_6(C_{12}H_{12}N_2)_3(DMF)_2]_n$, and Ag_{12} -azopy $[Ag_{12}(C_6H_{11}S)_6(CF_3COO)_6(C_{10}H_8N_4)_3(DMF)_2]_n$. By working with precise Ag₁₂ structured systems, we obviated the problem of accurately analyzing the origin of the difference in the catalytic activities, which plagues the study in structure-catalysis analysis. These assemblies exhibit varying activity levels, where the gradual lengthening of ligand systems from pyridine to azopyridine alters cluster arrangement. The Ag₁₂-pyz CAM displayed very good catalytic performance for the HER even at lower over-potentials. The low basicity of the pyrazine linker increases the electropositivity of the silver atom, shifting the catalytic active site from silver to sulfur. We propose that the linker basicity and assembly-induced cluster arrangements in these five assemblies define the active center and activity. This structure–property correlation, combined with experimental and theoretical insights, offers a promising model for cluster chemistry and electrocatalysis research.

Results and discussion

Ag₁₂-py, Ag₁₂-pyz, Ag₁₂-bpy, Ag₁₂-bpa, and Ag₁₂-azopy were prepared by a "one pot method". The $(Ag-SCy)_n$ (SCy-cyclohexanethiol) and AgCF₃COO precursors were dissolved in solvent, followed by a dropwise addition of different linkers to yield five distinct assembly patterns: 2 three-dimensional extended Ag₁₂pyz and Ag₁₂-bpa CAMs and 2 two-dimensional extended Ag₁₂-bpy and Ag₁₂-azopy CAMs, created by covalently linking the clusters with bidentate linkers (pyz = pyrazine, bpy = 4,4'-bipyridine, bpa = 1,2-bis(4-pyridyl)ethane, and azopy = 4,4'-azopyridine). The inter-cluster distance and dimensionality depend on the linker length and flexibility. Long rigid linkers, such as bpy and azopy, tend to facilitate two-dimensional assemblies (Fig. 1 and 2). In contrast, more flexible linkers like bpa and the shorter rigid pyz linker promote the formation of three-dimensional structures. Non-covalent interactions extended the clusters with monodentate pyridine in Ag_{12} -py (py = pyridine) (Fig. 1).

Single Crystal X-Ray Diffraction (SCXRD) data of Ag_{12} -pyz crystals (Table S1 and Fig. S1a†) show that they crystallize in the

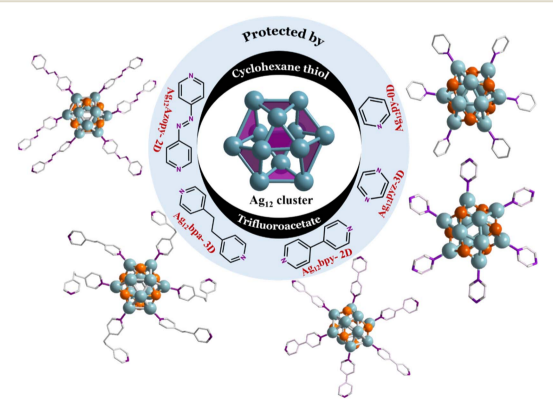


Fig. 1 (a) Schematic representation (cluster core with the linker) of the five silver nanocluster assemblies. The core constituting Ag_{12} (center) is protected by cyclohexane thiol and trifluoroacetate ligands. The assemblies (right to left) are Ag_{12} -py (0D NC), Ag_{12} -pyz (3D CAM), Ag_{12} -bpy (2D), Ag_{12} -bpa (3D), and Ag_{12} -azopy (2D). The color code: Ag, blue; S, orange; S, violet; S,

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space with the unit formula group $[Ag_{12}(C_6H_{11}S)_6(CF_3COO)_6(C_4H_4N_2)_3]_n$. The cluster core constitutes 12 Ag atoms held together by twelve argentophilic interactions with an ~Ag···Ag distance of 2.93 Å (Table S5†), which is protected by six primary cyclohexane thiol groups in a μ^4 fashion with an Ag−S distance of ~2.51 Å and six auxiliary CF₃COO[−] ligands via oxygen in $\mu^2 - \eta^2$ mode with an Ag-O distance of $\sim 2.40 - 2.51$ Å. This Ag₁₂(C₆H₁₁S)₆(CF₃COO)₆ node connects with 6 pyrazine linkers via Ag-N bonds (~2.30 Å) (Fig. S2†) to further connect 6 other clusters (Fig. S3†). A simple topology of this can be viewed in Fig. S4,† which repeats in three dimensions (Fig. 2b). The shortest Ag···Ag distance between two clusters in all three crystallographic directions is 7.40 Å (Fig. S5†).

Ag₁₂-bpy crystallized in the *Pnnm* (no. 58) space group (Table S2 and Fig. S1b†), with the cluster unit formula $[Ag_{12}(C_6H_{11}S)_6(CF_3COO)_6(C_{10}H_8N_2)_3]_n$. The 12 Ag atoms that constitute the core are held by 20 argentophilic interactions (~Ag···Ag distance of 3.15 Å) (Table S5†). The silver core is protected by 6 cyclohexane thiol (Ag-S distance ~2.47 Å) and six CF_3COO^- ligands (Ag-O distance $\sim 2.20-2.67$ Å) in a similar fashion and further connected by six 4,4'-dipyridyl linkers (Fig. S6†) (Ag-N bonds \sim 2.24-2.30 Å) to form a two-dimensionally extended solid (Fig. 2c and S7†), with the shortest Ag···Ag distances between two clusters in the bc plane being 11.57 and 11.38 Å (Fig. S8†).

 Ag_{12} -bpa $[Ag_{12}(C_6H_{11}S)_6(CF_3COO)_6(C_{12}H_{12}N_2)_3(DMF)_2]_n$ crystallized in the C2/c (no. 15) space group (Table S3 and Fig. S1c†). The core which is composed of 12 silver atoms is held together by 18 argentophilic interactions (~Ag···Ag distance of 3.10 Å) (Table S5†). The 6 cyclohexane thiol and 6 trifluoroacetate ligands protect the cluster in a similar fashion (Ag-S distance ~2.49 Å and Ag-O distance \sim 2.43-2.75 Å). The bpa linker *via* Ag-N bonds

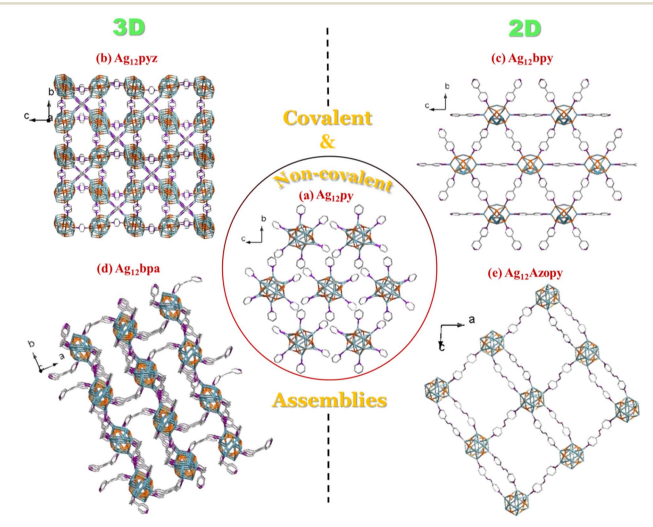


Fig. 2 Schematic representation of the four covalent assemblies (covalent bonding between the Ag of the cluster and the N of the different Nbased linkers) and a non-covalent assembly. (a) Ag₁₂-py NC extends via non-covalent interactions with the pyridine ligand, viewed along the a direction. (b) The Ag₁₂ clusters in Ag₁₂-pyz extend in three dimensions, and the figure is viewed along the a direction and slightly tilted to see the three-dimensional representation. (c) The bpy linker in Aq₁₂-bpy extends the cluster in two-dimensional hexagonal-like representation viewed along the a direction. (d) The flexibility of the bpa linker in Ag_{12} -bpa extends the cluster three-dimensionally, which is viewed along the c direction and slightly tilted to see the assembly extended in the c direction. (e) The azopy linker in Ag_{12} -azopy extends the cluster two-dimensionally, which is viewed along the b direction. The color code: Ag, blue; S, orange; N, violet; C, grey. All O, F, and H atoms are omitted for clarity.

(\sim 2.27–2.29 Å) connects each cluster to six others (Fig. S9 and S10†) to form a three-dimensionally extended solid (Fig. 2d), with the shortest Ag···Ag distances between the neighboring clusters in the bc plane being 9.89 and 12.02 Å (Fig. S11†).

Ag₁₂-azopy $[Ag_{12}(C_6H_{11}S)_6(CF_3COO)_6(C_{10}H_8N_4)_3(DMF)_2]_n$ crystallized in the C2/c (no. 15) space group (Table S4 and Fig. S1d†). The 12 Ag atoms constituting the cluster core are held by 20 argentophilic interactions (\sim Ag···Ag distance of 3.15 Å) (Table S5†). The cyclohexane thiol and trifluoroacetate ligands protect the cluster, and this is further covalently linked by 6 azopy linkers (Fig. S12†) to form a two-dimensionally extended solid (Fig. 2e and S13†). The shortest Ag···Ag distances with the neighbouring clusters in the ac plane are 13.36 and 13.46 Å (Fig. S14†).

The Ag_{12} -NC, $\{[Ag_{12}(C_6H_{11}S)_6(CF_3COO)_6(C_5H_5N)_6]\cdot 4H_2O\}$, a non-covalently extended solid that is protected by pyridine (Fig. 2a, S15 and S16†), was earlier synthesized by us. 25 This aids in correlating with the four new, two- and three-dimensionally extended solids (Fig. 1 and 2). Overall, five assemblies having a similar metal core and protecting ligands, differing in their assembly pattern, have been studied, with the shortest intercluster distance in Ag₁₂-pyz and the longest in Ag₁₂-azopy, based on the linker connecting them. More compact assemblies or higher dimensional structures tend to show shorter Ag···Ag inter- and intra-cluster distances. The shortest linker pyz connecting the clusters three-dimensionally (Ag₁₂-pyz) not only has reduced inter-cluster Ag···Ag distances but also has reduced intra-cluster Ag···Ag distances and the series can be found in Table S6.† Covalent assemblies offer a tunable approach to achieving desired structural compactness, which is less controllable in non-covalent assembly.

The Powder X-Ray Diffraction (PXRD) pattern verified the crystallinity and phase purity of all the bulk samples, and the resulting unaltered pattern indicates its structural stability (Fig. S17-S21†). The Scanning Electron Microscopy (SEM) images of Ag₁₂-pyz show an octahedral morphology (Fig. S22†), Ag₁₂-bpy and Ag12-azopy show a hexagonal morphology (Fig. S23 and S25†) and Ag₁₂-bpa has a block-like morphology (Fig. S24†), all of which are similar to their optical microscope images (Fig. S1†). The chemical composition of these samples was validated by SEM Energy-Dispersive X-Ray Spectroscopy (EDS) analysis, confirming the presence of all the desired elements (Fig. S22-S25†). X-Ray Photoelectron Spectroscopy (XPS) analysis further elucidated the chemical identity and oxidation state of the respective elements in all the samples (Fig. S26-S29†). The survey spectra confirmed the presence of all the desired elements, namely Ag, S, N, O, and F. The binding energies of Ag $3d_{5/2}$ and Ag $3d_{3/2}$ were 368.85 eV and 374.85 eV for Ag₁₂-pyz, 368.57 eV and 374.57 eV for Ag₁₂-bpy, 368.37 eV and 374.37 eV for Ag₁₂-bpa, and 368.59 eV and 374.59 eV for Ag₁₂-azopy, respectively, which confirms the presence of silver in the +1 oxidation state. Furthermore, the binding energy of S, approximately at 163 eV, indicates the presence of bridging sulfur. The four CAM structures (Ag₁₂-pyz, Ag₁₂-bpy, Ag₁₂-bpa and Ag₁₂azopy) exhibit enhanced thermal stability up to ~130 °C as determined by thermogravimetric analysis (TGA) (Fig. S30a†). However, they display varying degrees of thermal structural integrity at elevated temperatures, as indicated by PXRD analysis (Fig. S30b \dagger). Ag₁₂-pyz assembly exhibits enhanced structural stability at higher temperatures.

The electrochemical studies were carried out in $0.5~M~H_2SO_4$ with pH = 0.3 as the electrolyte; all five assemblies were loaded on carbon cloth for fabricating the working electrode. Linear sweep voltammetric (LSV) analysis of the samples carried out at a scan rate of $5~mV~s^{-1}$ revealed the overpotentials of the materials at a current density of $10~mA~cm^{-2}$ (Fig. 3a). It was observed that the Ag_{12} -pyz CAM delivers a much lower overpotential of 122~mV compared to Ag_{12} -azopy, Ag_{12} -py, Ag_{12} -bpa, and Ag_{12} -bpy, which show overpotentials of 205, 230, 401, and 575~mV, respectively, to drive the same current density of $10~mA~cm^{-2}$. The studies regarding the current density-dependent activity of the materials were carried out by calculating the overpotential delivered by the material at different current densities. The obtained values are shown in the bar diagram (Fig. 3b).

The charge transfer kinetics at the electrode–electrolyte interface were analyzed using the Tafel slopes of the materials obtained from the iR-corrected LSV plots, which were found to be 117, 163, 178, 235, and 242 mV dec⁻¹ for Ag₁₂-pyz, Ag₁₂-azopy, Ag₁₂-py, Ag₁₂-bpa, and Ag₁₂-bpy, respectively (Fig. 3c). The values of the Tafel slopes suggest that Ag₁₂-pyz shows the fastest charge-transfer kinetics among all the samples and follows the Volmer–Heyrovsky mechanism with the Heyrovsky step as the rate-determining step. ⁵¹ Electrochemical impedance spectroscopic analysis revealed the charge transfer resistance ($R_{\rm ct}$) at the electrode–electrolyte interface. Ag₁₂-pyz showed an $R_{\rm ct}$ as low as 0.68 Ω , whereas for Ag₁₂-azopy, Ag₁₂-py, Ag₁₂-bpa, and Ag₁₂-bpy, the $R_{\rm ct}$ values were found to be 3.1, 3.52, 5.1, and 11.1 Ω , respectively (Fig. 3d).

From the LSV, Tafel slope and R_{ct} analysis, it was observed that, among the five catalysts, Ag12-pyz exhibits the best HER activity. We also conducted some experiments to analyze the stability of the Ag₁₂-pyz catalyst. The dynamic stability of Ag₁₂pyz was studied using accelerated degradation (AD) studies for 1000 cycles at a 200 mV s⁻¹ scan rate. The material maintained its stability after the AD process (Fig. 3e). Chronoamperometric analysis carried out at a constant potential of -0.2 V also proved the high static stability of the material for 28 hours (Fig. 3f). The amount of silver in the electrolyte of chronoamperometric experiments was found to be 0.626 ppm using ICP-MS analysis, which further corroborates the stability of the catalyst. The stability of Ag₁₂-pyz in water is shown in Fig. S31.† The postcatalytic PXRD of Ag₁₂-pyz coated on carbon tape confirms that the cluster is intact after the catalysis cycle (Fig. S32†). The SEM-EDS analysis and XPS survey spectrum also confirm the presence of all the necessary elements: Ag, S, O, N, and F after the catalysis cycle (Fig. S33 and S34†). The binding energy values of silver at 368.9 and 374.9 eV are similar to that of the pre-catalytic Ag₁₂-pyz CAM, which further confirms the silver oxidation state of +1 and also the catalyst stability.

To gain further insight into the observed catalytic activity, computationally we measured the hydrogen adsorption free energy ($\Delta G_{\rm H}$) as a descriptor for understanding HER reactivity.⁵² Initially, we performed the adsorption of a hydrogen atom at

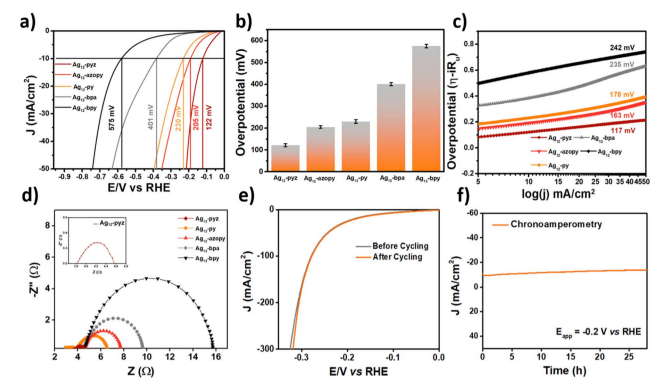


Fig. 3 HER performance of the catalysts. (a) LSV curves of Ag₁₂-pyz, Ag₁₂-azopy, Ag₁₂-py, Ag₁₂-bpa, and Ag₁₂-bpy, (b) bar diagram representing the current density at different overpotentials of Ag_{12} -pyz, Ag_{12} -azopy, Ag_{12} -py, Ag_{12} -bpa, and Ag_{12} -bpy, (c) Tafel slopes of all five assemblies, (d) Nyquist plots of all the structures (inset: Aq₁₂-pyz), (e) polarization curves of Aq₁₂-pyz before and after 2000 cycles, and (f) chronoamperometry measurements for stability.

various accessible sites among the Ag and S core atoms within the optimized clusters. The $\Delta G_{\rm H}$ energies at different conformers of the clusters are depicted in Fig. 4a and b. Based on the optimized structures after hydrogen adsorption, it is shown that except for the Ag₁₂-pyz, hydrogen adsorption at the Ag site is energetically favourable compared to that at the S site. In the Ag_{12} -azopy, Ag_{12} -py, Ag_{12} -bpa, and Ag_{12} -bpy clusters, hydrogen adsorption is more favourable by over 5.09, 6.8, 7.55, and 0.57 kcal mol⁻¹, respectively, at the bridge Ag-Ag sites compared to the S sites (Table 1). It is important to note that, according to the Sabatier principle, both highly exothermic and highly endothermic hydrogen adsorption are detrimental to the HER. In contrast, Ag₁₂-pyz shows hydrogen adsorption at the apical S sites, which is energetically favourable by 5.8 kcal mol⁻¹ compared to adsorption at the bridge Ag-Ag site. The calculated $\Delta G_{\rm H}$ values from the optimized structures (Fig. S35†) are shown in Fig. 4a and b, indicating that Ag₁₂-pyz could exhibit more effective HER activity compared to the others, aligning well with the experimentally measured overpotential values.

These energetic differences are further corroborated by the observation that the apical S sites in the Ag₁₂-pyz cluster exhibit a higher negative charge, making them more prone to hydrogen adsorption compared to those in other analogues. After hydrogen adsorption at the S sites in Ag₁₂-pyz, computed Bader charges indicate that it carries a negative charge (\sim -0.155e), suggesting a greater electronic interaction of H with the sulphur active site than the others. For Ag₁₂-py, Ag₁₂-bpy, Ag₁₂-bpa, and

Ag₁₂-azopy, the charges of hydrogen adsorbed on the S sites are -0.145e, -0.133e, -0.150e, and -0.133e, respectively. Changes in the linker within assemblies can influence basicity, which in turn affects catalytic activity. Specifically, linkers with higher basicity, such as py, bpy, bpa, and azopy, increase the electron density at Ag sites. This enhancement in electron density favors hydrogen binding at the Ag sites while reducing adsorption at the S sites. Overall, the trends in the binding energy at the S sites and the lower basicity of pyz result in superior performance in the HER compared to other linkers. It is important to note that the large negative charge density on the adsorbed S-H sites indicates the favourable charge transfer occurring from the catalyst surface to the adsorbed hydrogen. Hydrogen desorption would likely occur by the further adsorption of a proton from the electrolyte solution, followed by electron transfer, in a process known as the 'Heyrovsky' pathway. The adsorption of an additional hydrogen atom (Fig. S36†) in these clusters is energetically more favourable than the adsorption of a single hydrogen atom, further indicating that the HER follows the Volmer-Heyrovsky pathway as suggested by the Tafel slope analysis. Interestingly, the binding energy of the second hydrogen over the Ag₁₂-pyz cluster is less stable compared to the other studied clusters, resulting in superior HER activity as observed experimentally.

The highest occupied molecular orbitals (HOMO) in these clusters are predominantly localized on the Ag and S atoms, with a minor contribution from the ligand. In contrast, the

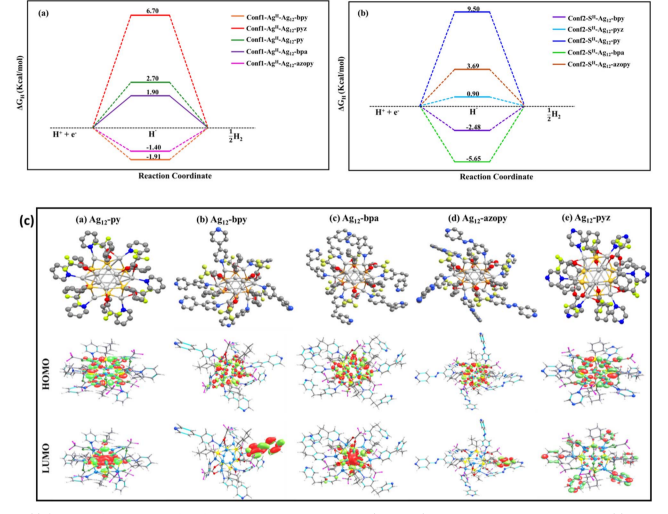


Fig. 4 (a) Calculated free energy profile of the HER at equilibrium potential ($U=U_{\rm RHE}$): hydrogen adsorption at the Ag site, (b) hydrogen adsorption at the S site. (c) Optimized structure and occupied and unoccupied molecular orbitals, (a) Ag₁₂-py, (b) Ag₁₂-bpy, (c) Ag₁₂-bpa, (d) Ag₁₂-azopy, and (e) Ag₁₂-pyz clusters. Color code: Ag, light grey; S, yellow; O, red; N, blue; C, grey; F, greenish yellow.

lowest unoccupied molecular orbitals (LUMO) exhibit a shift from the core of the cluster to the ligand. In Ag_{12} -py and Ag_{12} -bpa, the LUMO is primarily concentrated within the core, while in Ag_{12} -bpy, Ag_{12} -azopy, and Ag_{12} -pyz clusters, the LUMO is dominated by ligand-centric contributions. This inversion in the distribution of the unoccupied orbital is attributed to the structural variation introduced by the different ligands (Fig. 4c). Fig. S37† presents the calculated total density of states (TDOS) and partial density of states (PDOS) for these clusters. From the PDOS spectra, it is evident that the valence band in these clusters is primarily occupied by contributions from the core Ag

and S atoms, while the conduction band is predominantly influenced by the ligands. As we examine Ag_{12} -bpy, Ag_{12} -bpa, and Ag_{12} -azopy assemblies, the contribution from the ligands in the conduction band increases while the contribution from the Ag–S core decreases. Notably, in Ag_{12} -pyz, the pyrazine ligand makes a substantial contribution to the conduction band, whereas in Ag_{12} -py, both the pyridine ligand and the Ag–S core contribute to the conduction band.

Further, observing the theoretical band gap of the assemblies (Table S7 \dagger), Ag₁₂-pyz manifests a shorter band gap (2.01 eV). This shortest band gap correlates with the favourable

Table 1 Calculated ΔG_H values

$\Delta G_{ m H}$ (kcal mol ⁻¹)									
Ag ₁₂ -pyz		Ag ₁₂ -bpy		Ag ₁₂ -bpa		Ag ₁₂ -azopy		Ag ₁₂ -py	
conf1-Ag ^H 6.7	conf2-S ^H 0.9	conf1-Ag ^H -1.91	$conf2-S^H$ -2.48	conf1-Ag ^H 1.90	conf2-S ^H -5.65	conf1-Ag ^H -1.40	conf2-S ^H 3.69	conf1-Ag ^H 2.7	conf2-S ^H 9.5

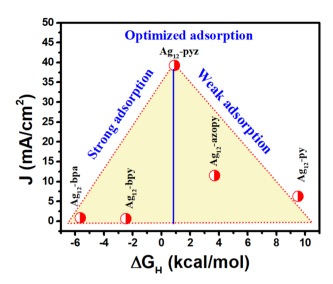


Fig. 5 Volcanic HER activity trend for the five Aq₁₂ cluster assemblies.

experimental conductivity of 4.1×10^{-6} S m⁻¹ (Fig. S38†). Therefore, it indicates that pyrazine plays a substantial role in the stabilization of the cluster that appears from the lowest LUMOs of the Ag₁₂-pyz CAM. These findings suggest that the electronic structure of the cluster can be effectively manipulated by changing the ligand structure.

From the experimental and theoretical analysis of the cluster structures, we made certain important observations. First, there is a systematic change in the bond distances of Ag-Ag and Ag-S upon variation of the ligand. The experimental crystal structure and theoretical bond length analyses show the Ag₁₂-pyz cluster has a shorter average Ag···Ag interaction of 2.93 Å (2.89 Å theoretically), compared to the others (Table S5 and Fig. S35†). This might have resulted in a slightly longer Ag-S bond in Ag₁₂pyz (Table S5†). These structural differences corroborate the observed site preference. The apical S-sites in Ag₁₂-pyz with a higher negative charge make it more prone to hydrogen adsorption compared to the others. It is also true that the basicity of the ligands in the series plays an important role in altering the active site based on the electron availability. The higher basicity of py, bpy, bpa, and azopy ligands increases the electron density on the Ag center, thereby favouring hydrogen adsorption at these sites. However, in the case of pyrazine, the reduced basicity alters the hydrogen adsorption site and maximizes HER activity through the S site. Second, the distribution of canonical orbitals changes, which varies with the ligand, affecting the structure-property relationship. Third, the intercluster interactions have a crucial effect on the conductivity of the materials. It has been observed that the material conductivity varies as a function of inter-cluster distance.53,54 The measured inter-cluster distances are as follows: Ag₁₂-pyz at 7.40 Å, Ag_{12} -bpa at 9.89 Å, Ag_{12} -py at 10.1 Å, Ag_{12} -bpy at 11.38 Å, and Ag_{12} -azopy at 13.36 Å (considering the shortest distance, see Fig. S5, S8, S11, S14 and S16†). Ag₁₂-pyz with the shortest Ag-Ag inter-cluster distance of 7.4 Å should possess the highest material conductivity, thereby reflecting faster charge transfer kinetics at the electrode-electrolyte interface. Moreover, the

isotropic nature (equal along three crystallographic axes) of the Ag-Ag inter-cluster distance in Ag₁₂-pyz allows the equivalent spatial distribution of the charge transfer process at the interface. In addition, the 3D structure of Ag₁₂-pyz allows effective mass transfer at the interface. However, for the other four sets of cluster assemblies, the excessive increase in the Ag-Ag intercluster distance (by ~3 Å) (Table S6†) does not significantly allow for altered electronic properties (such as conductivity) and therefore it can be considered that each Ag-cluster behaves independently. Moreover, the anisotropic behaviour of the Ag-Ag inter-cluster distance in different crystallographic directions might act as a barrier for improved electronic properties. Therefore, it can be said that for Ag₁₂-pyz, the inter-cluster distance and dimensionality play a crucial role in the HER kinetics, while for the other clusters, the HER kinetics is mainly governed by the H-adsorption energetics. Thus, tuning the ligand can significantly impact the cluster arrangements and the catalytic properties. Moreover, we have compared our cluster assemblies with other silver-based NC catalysts reported in the literature, in terms of the overpotential required for reaching a current density of 10 mA cm⁻² (Table S8†). As can be seen, Ag₁₂-pyz displays the best performance among the series.

The experimentally obtained HER activity trend was further rationalized by the current density value at -0.2 V vs. RHE against the calculated free energy of H adsorption (Fig. 5). It is well known that for the HER, any electrocatalyst with a highly negative ΔG_{H} value would bind the active H* very strongly and weakly with a positive ΔG_{H} value. Very strong and weak H* adsorption would slow down the desorption and adsorption process, thereby reducing the catalytic activity. Hence, H* adsorption that is neither too strong nor too weak leads to the best catalytic activity according to the 'Sabatier' principle. Here, various nitrogen linkers with their characteristics of basicity and electronic structure play a significant role in H adsorption over the S site and the Ag_{12} cluster with a pyrazine stabilizer possesses optimum adsorption energy, facilitating a facile adsorption and desorption process.

Conclusions

In this study, five assemblies were studied: Ag₁₂-py, Ag₁₂-pyz, Ag₁₂-bpy, Ag₁₂-bpa, and Ag₁₂-azopy. Ag₁₂-pyz showed superior catalytic activity for the HER compared to the others in the series. Theoretical analysis on hydrogen adsorption free energy showed more favourable hydrogen adsorption for Ag₁₂-pyz, enhancing its HER performance compared to the others. Further, the active center for catalytic hydrogen adsorption in Ag₁₂-pyz was observed to be at the apical S site, while it is at the Ag-Ag site for the other 4 assemblies. This alteration of the active site is attributed to intercluster assembly and the basicity of the linker. The isotropic nature of the Ag-Ag inter-cluster distance in Ag₁₂-pyz allows the equivalent spatial distribution of the charge transfer process. The relatively less basic pyrazine linker renders the silver more electropositive, a conclusion also supported by structural analysis, which showed the shortest Ag-Ag interactions in Ag₁₂-pyz, thus favoring its positioning at the apical S site. Variations in catalytic activity and active site **Edge Article Chemical Science**

preference among these assemblies were achieved by systematically altering the ligands from pyridine to azopyridine. This work, offering a molecular-level correlation between linkerinduced assembly, activity and active site switching in cluster assemblies, highlights a significant toolkit for selectively designing cluster assemblies as catalysts that hold broad implications for tailored catalytic activity in various applications.

Data availability

The data will be available from the authors on request for academic purposes.

Author contributions

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

Conflicts of interest

The authors declare no competing financial interest.

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