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Tuning of aromaticity and reactivity in gold-substituted cyclopropenyl cations†

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This study investigate the structural, electronic, and reactivity properties of the aurocarbon $C_3H_2Au^+$, $C_3HAu_2^+$, and $C_3Au_3^+$ clusters. A potential energy surface exploration indicates that the most stable isomers adopt cyclic structures, supporting the gold–hydrogen analogy in these cationic systems. These cyclic forms preserve the characteristic geometry of the cyclopropenyl cation, with covalent C–Au interactions confirmed by multiple bonding analyses. All three systems exhibit a delocalized $3c-2e$ π -bond across the carbon ring, consistent with an aromatic behavior. The substitution of hydrogen by gold alters local reactivity patterns.

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Introduction

Gold clusters and their compounds have attracted attention in nanoscience and nanotechnology due to their applications in catalysis, medicine, and electronics.^{1–5} Among these, aurocarbons, a class of gold clusters containing carbon atoms, have emerged as promising candidates for facilitating diverse chemical transformations. For instance, they play a key role in the environmentally relevant conversion of carbon monoxide to carbon dioxide.⁶ Density functional theory (DFT) studies by Tiwari revealed that small aurocarbons, such as C_4H_3Au and C_4Au_4 , exhibit enhanced catalytic activity for CO oxidation, particularly when CO pre-adsorbed on all gold sites in C_4Au_4 .⁷ These findings suggest that carbon-intercalated gold clusters may serve as efficient catalysts. Beyond CO oxidation, aurocarbons have also shown potential for

SO_2 reduction with CO and for acetylene hydracyanation.^{8,9} Their stability is often attributed to aurophilic interactions (strong mutual attractions between gold atoms). This phenomenon was explored in detail by Zaleski-Ejgierd and Pyykkö, who compared the bonding characteristics of aurocarbons with their carbon-only counterparts.¹⁰

Naumkin has also extensively investigated the formation, reactivity, and properties of carbon–gold nanoclusters. In 2006, he reported the C_5Au_{12} system and its charged derivatives ($C_5Au_{12}^-$ and $C_5Au_{12}^+$). These core–shell aurocarbons consist to a tetrahedral C_5 radical (carbon core) encapsulated by a 12-atom gold shell, mimicking the bonding environment of the C_5H_{12} molecule. Using DFT calculations, the study evaluated the structural stability, electronic properties, and bonding interactions of neutral and ionic forms. Two main classes of isomers were identified: an icosahedral-like (I_h) isomer and an octahedral-like (O_h) isomer, both resembling similar structures found in the Au_{13} cluster and C_5H_{12} .¹¹ Building on this approach, Naumkin later explored core-doping in the $C_{10}Au_{18}$ system. He reported a structural analogue of di-*tert*-butyl-acetylene $C_{10}H_{18}$ with near- C_3 symmetry. A key finding was that such clusters allow flexible tuning of catalytic properties: the C_{10} dopant can selectively alter the electron-donating capacity of the gold cluster while maintaining its electron-acceptor efficiency.¹²

Ramachandran and Naumkin further investigated the structure, stability, and electronic properties of C_4Au_n clusters, where $n = 4, 6$, and 8 . They classified the isomers into four categories: hydrocarbon analogues, carbon chains, carbon rings adsorbed on gold surface, and carbon cores encapsulated by gold atoms. The most stable structures typically featured linear or surface-bound carbon moieties rather than

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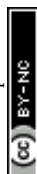
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encapsulated cores. For example, the lowest-energy isomers of C_4Au_4 resembles a gold analogue of 1,2,3-butatriene, while the cyclobutadiene-like structure (C_4Au_4) lies 36.9 kcal mol^{−1} higher in energy. Despite their lower thermodynamic stability, encapsulated-carbon isomers exhibit variations in ionization energies, electron affinities, and HOMO–LUMO gaps, properties that make them attractive for the rational design of tunable nanocatalyst.¹³

The structural characterization of aurocarbons clusters has been widely explored. In 2015, Meng *et al.* determined the structures of four AuC_3H^- isomers using emission spectroscopy and *ab initio* calculations.¹⁴ Ticknor reported the formation of AuC_n^+ species *via* laser vaporization and mass spectrometry¹⁵ techniques that also yielded AuC_n^+ ($n = 1–11, 18$), $Au_2C_n^+$ ($n = 1–16$), $Au_3C_n^+$ ($n = 1–10$), and AuC_3H^- clusters, which are single-gold analogues of the C_3H_2 carbene.^{14,16} Theoretical and experimental efforts have further examined gold carbide clusters and the synthesis of gold–acetylenes complexes ($Au-C \equiv C-Au$).^{8,17,18}

Beyond structural aspects, several studies have proposed an analogy between gold and hydrogen in these clusters. This so-called gold–hydrogen analogy suggests that hydrogen can mimic the behavior of a gold atom within a cluster.^{19–21} Zheng *et al.* investigated this concept by analyzing auro-acetylide species (AuC_2H , AuC_2Au , and Au_2C_2H) using photoelectron spectroscopy and DFT calculations, finding that AuC_2H and AuC_2Au share similar electronic and bonding properties.¹⁸ Wang *et al.* expanded this idea, exploring the formation of aurosilicon, auroboron, and aurocarbon clusters, all characterized by strong Au–E covalent bonds ($E = Si, B, C$).^{18,21,22} In addition, the combination of photoelectron imaging spectroscopy and DFT calculations has been used to elucidate the structures of four AuC_3H^- isomers.¹⁴ These techniques were subsequently applied to study AuC_4^- , AuC_6^- , and AuC_nH^- ($n = 2, 4$, and 6) clusters, leading to the discovery of auropolynes ($Au(C_2)_nAu^-$, $n = 1–3$) *via* high-resolution photoelectron imaging and *ab initio* calculations.²³

Further insight into bonding in gold–carbon systems comes from Armentrout's study of the gold carbene cation ($AuCH_2^+$), which used molecular orbital analysis to reveal covalent C–Au bonding.²⁴ Megha *et al.* revisited the gold–hydrogen analogy by studying CO and O₂ adsorption on pristine and hydrogen-doped gold clusters (Au_7H), confirming the analogy in the most stable planar isomer, although deviations were obtained in higher-energy structures.¹⁹ Tang *et al.* synthesized and characterized phenylgold (C_6H_5Au) and its anion ($C_6H_5Au^-$) using laser vaporization, photoelectron spectroscopy, and DFT calculations.²⁵ Tsipis and Gkarbounis investigated aromatic aurocarbons (C_6Au_6), the benzene analogue, predicting their stability and analyzing their magnetic properties. Their calculations indicated that coinage-metal-substituted benzenes exhibit reduced aromaticity compared to benzene.²⁶ More recently, Lu examined M_5C_6 systems ($M = Cu, Ag$ and Au), concluding that $Au_5C_6^-$ exhibits σ -aromaticity.²⁷

In this work, we investigate the sequential H-to-Au substitution in $C_3H_3^+$,²⁸ leading to the fully substituted species $C_3Au_3^+$.

Cyclic structures analogous to $C_3H_3^+$ emerge as the lowest-energy configurations, displaying marked electron delocalization and tunable reactivity. To characterize the bonding and electronic properties of $C_3H_2Au^+$, $C_3HAu_2^+$ and $C_3Au_3^+$, we employed Natural Population Analysis (NPA),²⁹ Wiberg bond Indices (WBI),³⁰ the Electron Localization Function (ELF),³¹ and Adaptive Natural Density Partitioning (AdNDP).³² Aromaticity was further assessed *via* magnetically induced current density (MICD) and induced magnetic field (B^{ind}) analysis,³³ both confirming the presence of diatropic ring currents. Finally, reactivity trends were examined using local hyper-softness (LHS),^{34–39} revealing that even a single H-to-Au substitution alters local reactivity patterns.

Computational details

The potential energy surfaces (PESs) of the clusters were explored using a modified genetic algorithm, as implemented in GLOMOS,^{40–42} which has been previously validated in related studies.^{43–48} Initial geometries were optimized at both the PBE0/def2-TZVP and PBE0-D3⁴⁹/def2-TZVP^{50,51} levels. The same isomers were located with both levels, with relative energy differences of less than 2 kcal mol^{−1} (see Fig. S1–S4, ESI†). These results indicate that, for the systems examined here, dispersion interactions do not alter the overall topology of the PES, although they do introduce slight shifts in the relative energies of the isomers. The structure were further refined at the MP2/def2-TZVP level,⁵⁰ incorporating effective core potentials for gold.⁵² Harmonic frequency analysis confirmed that all structures correspond to local minima. All calculations were carried out with Gaussian 16.⁵³ To analyze stability, bonding characteristics, and electron delocalization, we computed NPA charges,²⁹ WBI,³⁰ ELF,³¹ and AdNDP³² analysis using Multiwfn.⁵⁴

MICD⁵⁵ was computed *via* the response function⁵⁶ and the perturbing operator for the magnetic field using DIRAC 19.⁵⁷ The wavefunction was obtained with the four-component Dirac–Coulomb Hamiltonian using unrestricted kinetic balance,⁵⁷ employing the PBE0 functional⁵¹ and the cc-pVDZ⁵⁸ basis set for carbon and hydrogen atoms. Gold atoms were treated with the Dyall double- ζ basis set.⁵⁹ Shielding tensors required for the calculation of the induced magnetic field (B^{ind})³³ were computed with ORCA⁶⁰ using the PBE0 functional and the def2-QZVP^{52,61} basis set. Isolines maps and isosurfaces were visualized using VisIt.⁶²

Conceptual density functional theory (c-DFT)⁶³ was used to evaluate local reactivity using LHS,^{34–39} a descriptor that is more accurate and size-consistent than local softness, dual descriptors, or fukui functions.⁶⁴ LHS^{65,66} values were computed at the MP2/def2-QZVP level using Gaussian 16. To simplify the calculations, we adopted the frontier molecular orbital approximation (FMOA) rather than the finite difference approximation (FDA), ensuring a reliable reactivity description even in systems with orbital degeneracy.⁶⁷ The robustness of the dual descriptor, which underlies LHS, has been validated in systems as diiodine, where relativistic effects are significant,



confirming that FMOA-based dual descriptors provide a reliable qualitative description.⁶⁸

Results and discussion

A PES exploration identified global minima and low-energy isomers for the singlet and triplet states of the C_3HAu_2^+ , $\text{C}_3\text{H}_2\text{Au}^+$ and C_3Au_3^+ systems. In each case, the most stable isomer preserves the three-carbon cyclic geometry of the cyclopropenyl cation ($\text{c-C}_3\text{H}_3^+$), with energy differences of less than 33 kcal mol⁻¹ relative to linear isomers resembling the propargyl cation ($\text{l-C}_3\text{H}_3^+$) (Fig. 1, additional isomers in Fig. S1–S4, ESI†). In other words, these results support the gold–hydrogen analogy in cationic systems.

In the most stable isomers, the C_3 ring remains intact, maintaining constant C–H bond lengths, while the C–C bond lengths increase slightly from 1.37 to 1.39 Å upon H-to-Au substitution (Fig. 1). The resulting C–Au bonds (~ 1.88 Å) are consistent with reported Au–CO bond lengths (1.84–2.03 Å);⁶⁹ WBI confirm covalent C–Au interactions ($\text{WBI}_{\text{C–Au}} = 0.92$) and reveal a decrease in $\text{WBI}_{\text{C–C}}$ values from 1.43 in C_3H_3^+ to 1.34 in C_3Au_3^+ . Furthermore, replacing H with Au increases charge accumulation on the carbon skeleton, with NPA charges indicating a shift toward more positive on the Au atoms and more negative values on the C atoms.

ELF values close to 1 confirm covalent bonding across the systems (Fig. 2). Among these, C–Au bonds exhibit the lowest ELF values (0.80–0.90), deviating from ideal covalent values yet remaining within the covalent range. In Au-substituted systems, C–C bonds display slightly reduced ELF values, consistent with their elongation. These trend in ELF agree with the WBI data, collectively confirming the presence of strong C–H and C–Au covalent bonds and a weakening of C–C bond orders upon Au substitution.

AdNDP analysis of C_3Au_3^+ (Fig. 3) identifies three 2c–2e σ -bonds between carbon atoms, three C–Au σ -bonds, and a

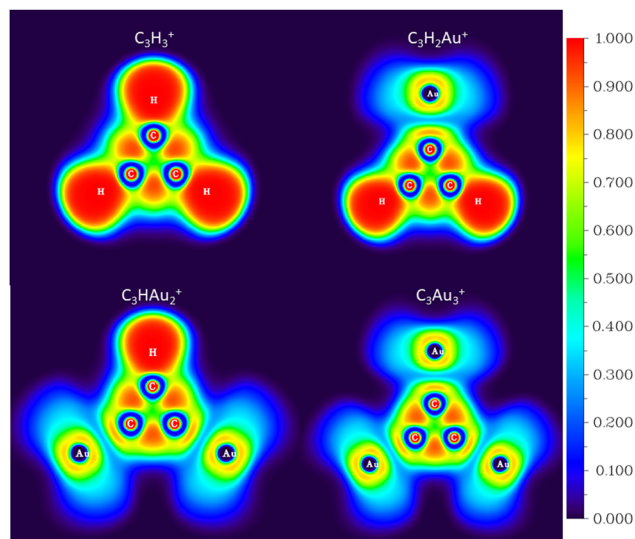


Fig. 2 Color-filled map ELF maps of C_3H_3^+ , $\text{C}_3\text{H}_2\text{Au}^+$, C_3HAu_2^+ , and C_3Au_3^+ .

delocalized 3c–2e π -bond across the carbon ring, satisfying Hückel's rule and confirming aromaticity. Each Au atom contributes five 5d orbitals corresponding to lone pairs, resulting in a total of fifteen. These orbitals exhibit slightly reduced occupation numbers (ONs) between 1.96 and 1.91 $|e|$, indicating partial charge transfer to the carbon atoms but no participation in π -delocalization. This bonding pattern, including the presence of a 3c–2e π -bond, is also present in the related complexes $\text{C}_3\text{H}_2\text{Au}^+$ and C_3HAu_2^+ (Fig. S5 and S6, ESI†).

These results are consistent with the NPA charge analysis, which shows that gold substitution increases the negative charge on the carbon atoms and the positive charge on the gold atoms. Accordingly, the AdNDP analysis confirms that the C_3 ring in C_3Au_3^+ retains an aromatic character similar to that of the prototypical C_3H_3^+ system (Fig. S7, ESI†), with a delocalized 3c–2e π -bond localized over the carbon atoms.

During the review process, one of the reviewers inquired about the effect of adding an electron to C_3H_3^+ and C_3Au_3^+ , a particular insightful question. The contrasting behavior of the resulting C_3H_3 and C_3Au_3 clusters arises from fundamental differences in electron delocalization and orbital symmetry, which govern their geometric and electronic structures. A key

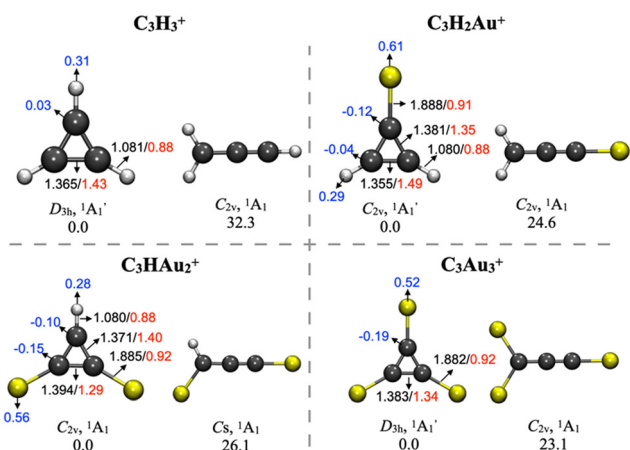


Fig. 1 Relative energies (kcal mol⁻¹) of the low-energy isomers of C_3H_3^+ , $\text{C}_3\text{H}_2\text{Au}^+$, C_3HAu_2^+ , and C_3Au_3^+ , calculated at the MP2/def2-TZVP level. Bond lengths (Å, black), natural population analysis charges ($|e|$, blue), and Wiberg bond indices (red) are indicated.

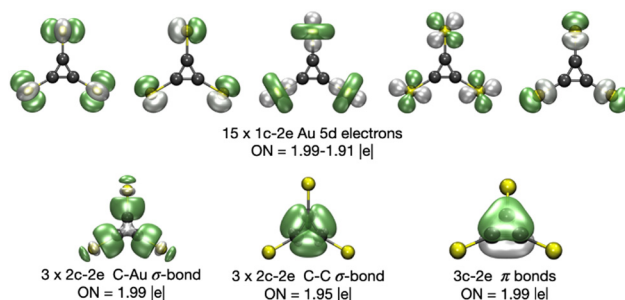


Fig. 3 Adaptive natural density partitioning plots of C_3Au_3^+ at the PBE0-D3/def2-TZVP level, using pseudopotentials for Au atom.



aspect lies in the nature of the molecular orbital populated upon reduction of the corresponding cation.

For C_3H_3^+ , the added electron occupies a degenerate π -antibonding orbital (Fig. S8, ESI[†]). This electron configuration violates Hückel's $4n + 2$ rule for aromaticity, and the presence of an unpaired electron in a π^* -orbital destabilizes the system. This electronic degeneracy triggers a Jahn–Teller distortion, lowering the symmetry from D_{3h} to C_{2v} , disrupting π -delocalization and eliminating aromatic stabilization. Geometry optimizations at the PBE0-D3/def2-TZVP level yields two imaginary frequencies at the D_{3h} geometry. Upon relaxation, the molecule adopts a lower-symmetry C_{2v} geometry, characterized by bond length alternation (1.38 Å vs. 1.42 Å), consistent with π -antibonding interactions and the loss of aromatic character. In contrast, the addition of an electron to C_3Au_3^+ populates a σ -bonding orbital (Fig. S8, ESI[†]). So, occupation of the σ -orbital does not induce the destabilization observed in the hydrocarbon analogue. Consequently, the neutral C_3Au_3 retains its D_{3h} symmetry and shows no imaginary frequencies, consistent with a true local minimum.

To further evaluate aromaticity across the aurocarbon series, MICD analysis was performed. All systems exhibit diatropic ring currents (Fig. 4), supporting their aromatic character. The strength of the current density was integrated in the xy plane (orange line in Fig. 4), which bisects a representative C–C bond in each system. These values are shown in Fig. 4 and further support the aromatic behavior of the molecules. Interestingly, the addition of Au results in a gradual decrease in current density strength. The diatropic response around the Au atoms

overlaps with that of the C_3 core, forming a net diatropic current. This effect is particularly evident in $\text{C}_3\text{H}_2\text{Au}^+$ and C_3HAu_2^+ (see Fig. S9a, ESI[†]), where current density plots ≥ 2 Bohr above the molecular plane reveal this combined delocalization pattern. Notably, C_3Au_3^+ shows a weaker diatropic current density (7.9 nA T^{-1}) compared to C_3H_3^+ (9.1 nA T^{-1}), consistent with the trend in Fig. S9a and b (ESI[†]). In contrast, both C_4H_4 and C_4Au_4 exhibit paratropic current densities, consistent with their antiaromatic character. For C_4H_4 , the integrated current density strength was calculated as -22.8 nA T^{-1} , in good agreement with the previously reported value of -19.9 nA T^{-1} .⁷⁰ For C_4Au_4 , a paratropic current density of approximately -14.7 nA T^{-1} was obtained. The corresponding current density plots for both species are shown in Fig. S9b (ESI[†]).

Isolines maps of the z -component of the induced magnetic field (B_z^{ind} , equivalent to NICS_{zz})⁷¹ also reveal diatropic ring currents in all systems (Fig. 5). Substitution of protons by Au atoms has minimal impact on the electron delocalization within the C_3 ring. (Fig. 5). However, due to Au's large atomic radius, localized diatropic regions appear around the heavy atoms, likely due to contributions from the 5d electrons. As more protons are replaced by gold, these diatropic regions become increasingly pronounced. Fig. S10 (ESI[†]) shows B_z^{ind} isosurfaces for C_3H_3^+ , $\text{C}_3\text{H}_2\text{Au}^+$, C_3HAu_2^+ , and C_3Au_3^+ , revealing a broader magnetic response in C_3Au_3^+ . Finally, Fig. S11 (ESI[†]) shows the B_z^{ind} isosurfaces for C_4H_4 and C_4Au_4 , confirming that their antiaromatic character, evidenced by a paratropic magnetic response, persists even after hydrogen-to-gold substitution.

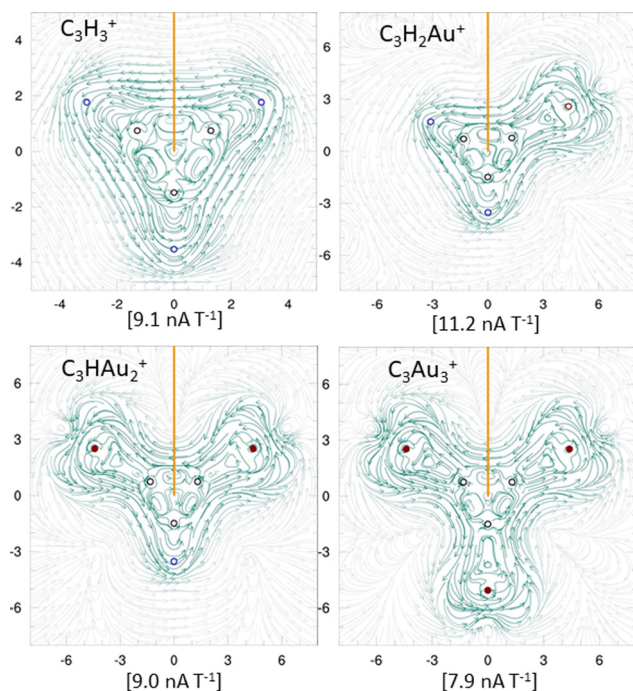


Fig. 4 Magnetically induced current density maps for: (A) C_3H_3^+ , (B) $\text{C}_3\text{H}_2\text{Au}^+$, (C) C_3HAu_2^+ , and (D) C_3Au_3^+ . Counterclockwise currents correspond to diatropic behavior. Black, blue, and red spheres represent C, H, and Au atoms, respectively. Numbers in brackets indicate current strength (nA T^{-1}), and the orange line denotes the integration plane.

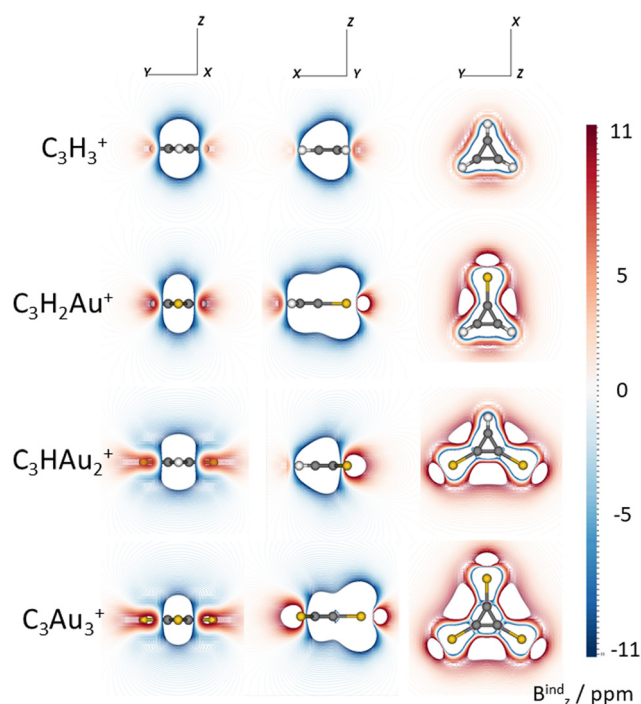


Fig. 5 Isolines maps of the z -component of the induced magnetic field (B_z^{ind}) for all systems. Units are ppm. Carbon, hydrogen, and gold atoms are represented as black, white, and yellow spheres, respectively.



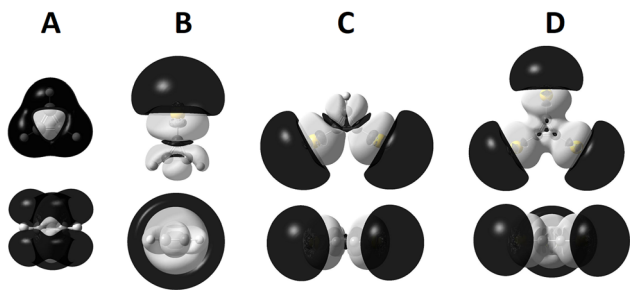


Fig. 6 Isosurfaces of local hyper-softness (LHS) at $\pm 0.00075 |e|^3 \text{ Hartree}^{-2} \text{ Bohr}^{-3}$: (A) C_3H_3^+ , (B) $\text{C}_3\text{H}_2\text{Au}^+$, (C) C_3HAu_2^+ , (D) C_3Au_3^+ . Upper panels show front views; lower panels show side views. Dark lobes correspond to positive LHS values ($+0.00075$), and white lobes to negative values (-0.00075). Additional isosurface cuts are provided in the ESI† (Fig. S12).

Do the geometric and electronic changes induced by H-to-Au substitution affect the reactivity of aurocarbons? To address this question, we performed LHS analysis to identify covalent nucleophilic and electrophilic sites. Fig. 6 shows the LHS isosurfaces, which aligns with the symmetric irreducible representations of their corresponding point groups: D_{3h} for C_3H_3^+ and C_3Au_3^+ , and C_{2v} for $\text{C}_3\text{H}_2\text{Au}^+$ and C_3HAu_2^+ . The cyclopropenyl cation (Fig. 6(A)) acts as an electrophile, accepting electron density above and below the molecular plane. Replacing H with Au localizes the electrophilicity character at the gold atoms while enhancing nucleophilic behavior in their vicinity (Fig. 6(B)–(D)). This dual reactivity pattern persists throughout the substitution series: The Au-decorated vertices retain pronounced electrophilic character, while nucleophilic regions emerge around the Au atom. Thus, H-to-Au substitution shifts the electrophilic behavior from a delocalized, planar nature to a more localized, vertex-centered pattern.

To further quantify this trend, we computed the electrophilicity index (ω),⁷² which estimates the energy stabilization upon electron saturation of an electrophile. As shown in Table S1 (ESI†), C_3H_3^+ exhibits the highest ω value, serving as the reference. The following trend is found:

$$\omega(\text{C}_3\text{H}_3^+) > \omega(\text{C}_3\text{Au}_3^+) > \omega(\text{C}_3\text{H}_2\text{Au}^+) > \omega(\text{C}_3\text{HAu}_2^+)$$

We also evaluated the electron-donating ($-\omega^-$) and electron-accepting (ω^+) powers.⁷³ $\text{C}_3\text{H}_2\text{Au}^+$ emerges as the most electron-donating species, whereas C_3H_3^+ , consistent with its electrophilic nature, shows minimal electron-donating capability. These trends are further summarized using the net electrophilicity index ($\Delta\omega^\pm$):⁷⁴

$$\Delta\omega^\pm(\text{C}_3\text{H}_3^+) > \Delta\omega^\pm(\text{C}_3\text{Au}_3^+) > \Delta\omega^\pm(\text{C}_3\text{H}_2\text{Au}^+) > \Delta\omega^\pm(\text{C}_3\text{HAu}_2^+)$$

Finally, the softness index (S)⁷⁵ shows the opposite trend:

$$S(\text{C}_3\text{HAu}_2^+) > S(\text{C}_3\text{Au}_3^+) > S(\text{C}_3\text{H}_2\text{Au}^+) > S(\text{C}_3\text{H}_3^+)$$

Altogether, these results confirm that while C_3H_3^+ is a prototypical electrophile, its Au-substituted analogues display

a richer dual character that becomes more pronounced with increasing Au content.

Conclusion

Sequential H-to-Au substitution in C_3H_3^+ preserves the cyclic geometry of the parent cyclopropenyl cation, as confirmed by potential energy surface analyses. The structural similarity between cyclic and linear isomers reinforces the gold–hydrogen analogy in these cationic systems. Substitution leads to elongated C–C bonds (1.37 – 1.39 Å) and the formation of covalent C–Au bonds (~ 1.88 Å), supported by Wiberg bond indices ($\text{WBI} > 0.9$) and charge transfer from Au to C. Electron Localization Function analysis further validates the covalent nature of the C–Au interactions, albeit with minor deviations attributed to the higher electronegativity of gold.

Aromaticity is preserved across all substituted clusters. This is evidenced by AdNDP, magnetically induced current density, and induced magnetic field analyses, all of which confirm the presence of diatropic ring currents. Gold's large atomic radius induces localized diatropic regions, particularly in C_3Au_3^+ , thereby altering the electron delocalization pattern while maintaining aromatic behavior.

So, H-to-Au substitution reshapes reactivity. The Au atoms act as electrophilic sites, while the adjacent carbon atoms exhibit enhanced nucleophilic character. This dual behavior becomes more pronounced with increasing substitution, gradually transforming the prototypical nucleophilic C_3H_3^+ into a system with vertex-centered, tunable reactivity. These adaptable electronic features make aurocarbons promising candidates for catalytic applications that demand site-specific reactivity.

Author contributions

Erick Cerpa: conceptualization, writing – review & editing, investigation; Alba Vargas-Caamal: writing – original draft, investigation; Luz Diego: writing – review & editing, investigation; David Arias-Olivares: writing – review & editing, investigation; Diego V. Moreno: software, editing, investigation; Jorge I. Martínez-Araya: writing – review & editing, investigation; Rafael Islas: writing – review & editing, investigation, formal analysis; Gabriel Merino: writing – review & editing, supervision.

Data availability

The data supporting this article have been included as part of the ESI.†

Conflicts of interest

There are no conflicts to declare.

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