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100% selective cyclotrimerization of acetylene to benzene on Ag(111)†

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Benzene, a high-volume chemical, is produced from larger molecules by inefficient and environmentally harmful processes. Recent changes in hydrocarbon feedstocks from oil to gas motivate research into small molecule upgrading. For example, the cyclotrimerization of acetylene reaction has been demonstrated on Pd, Pd alloy, and Cu surfaces and catalysts, but they are not 100% selective to benzene. We discovered that acetylene can be converted to benzene with 100% selectivity on the Ag(111) surface. Our temperature programmed desorption experiments reveal a threshold acetylene surface coverage of ~one monolayer, above which benzene is formed. Furthermore, additional layers of acetylene increase the amount of benzene produced while retaining 100% selectivity. Our scanning tunneling microscopy images show that acetylene prefers square packing on the Ag(111) surface at low coverages, which converts to hexagonal packing when acetylene multilayers are present. Within this denser layer, features consistent with the proposed C₄ intermediates of the cyclotrimerization process are observed. Density functional theory calculations demonstrate that the barrier for forming the crucial C4 intermediate generally decreases as acetylene multilayers are formed because the multilayer interacts more strongly with the surface in the transition state than in the initial state. Given that acetylene desorbs from Ag(111) at \sim 90 K, the C₄ intermediate on the pathway to benzene must be formed below this temperature, implying that if Ag-based heterogeneous catalysts can be run at sufficiently high pressure and low enough temperature, efficient and selective trimerization of acetylene to benzene may be possible.

Introduction

Benzene is a key compound in the chemical industry with a market value of \sim \$40 billion in 2022 which is projected to reach \sim \$80 billion by 2030.^{1,2} Benzene is widely used as a chemical intermediate for the manufacturing of various consumer goods such as cleaning products, clothing, packaging, building materials, and pharmaceuticals.³ Currently, benzene is mainly produced *via* three different methods which rely heavily on petroleum and coal, namely, catalytic reforming, hydrodealkylation of toluene, and steam cracking, each of which contribute \sim 30% to the market supply.⁴ These processes form benzene from >C₆ molecules and only the hydrodealkylation of toluene provides high conversion (\sim 90%) and selectivity (\sim 95%).⁵ The high cost and environmental impact of current benzene production methods – including severe operating conditions, high energy demand, and environmental

pollution – incentivize the search for alternative routes. 6,7 Furthermore, the recent surge in shale gas production in the United States has created a plentiful supply of $C_{1,2,3}$ species; therefore, identification of methods for converting these light hydrocarbons into value added C_{6+} products would be timely. Thus, acetylene trimerization to benzene may be attractive in its own right, and is also representative of a class of C–C coupling reactions that are of increasing technological interest.

Previous work has shown that it is possible to produce benzene from acetylene on a variety of metal surfaces under both ultra-high vacuum (UHV) and reaction conditions. 9-35 For example, Tysoe *et al.* showed acetylene forms benzene and ethylene on a Pd(111) surface and that increasing the surface acetylene coverage increased the yield of benzene. 36 In addition, the authors observed a threshold acetylene coverage of 0.3 ML, below which benzene formation does not occur. They postulated that the reactant molecules are too far apart and steric crowding is necessary to form benzene in the low coverage regime. This postulate was supported by experiments with coadsorbed NO, a spectator molecule, which helped compress the 2D acetylene monolayer and increased the yield of benzene for the same surface coverage of acetylene. Furthermore, the authors provided evidence for the existence of a C₄ intermediate

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along the pathway to benzene formation. Kyriakou et al. demonstrated that acetylene trimerizes to benzene on the Cu(111) surface but the reaction was not fully selective to benzene, as acetylene was also converted to ethylene, butadiene, and cyclooctatetraene.³⁷ Using isotopically labeled experiments Kyriakou et al. demonstrated that the formation of butadiene requires C-H bond scission whereas formation of benzene occurs through a pure C-C coupling mechanism. The authors hypothesized that the C4 metallocycle is a key intermediate which can be hydrogenated to produce butadiene or coupled with an additional acetylene molecule to form benzene. Moreover, Baddeley et al. showed that PdAu bimetallic surfaces are active for acetylene cyclotrimerization to benzene, among other products. They demonstrated that PdAu surfaces are more active than pure Pd(111) and observed a dependence of the benzene yield on Pd coverage, theorizing two main factors dictated the effectiveness of the alloy system: rehybridization of the adsorbed acetylene and weaker adsorption of benzene on PdAu, which facilitates reaction.35

More recently, Boudjahem et al. reported that nanoparticle Ni/SiO₂ catalysts perform acetylene cyclotrimerization to benzene in the presence of hydrogen.³⁸ The authors found that the partial pressure of hydrogen and acetylene, the reaction temperature, and Ni loading all determine the conversion of acetylene and the product distribution. By lowering the Ni loading and reaction temperature, they were able to increase the benzene selectivity to a maximum of \sim 52% in the presence of water vapor at conversions of \sim 3%. Together, these studies provide an understanding of the mechanism behind acetylene cyclotrimerization to benzene and the critical role of the C4 intermediate. However, on extended metal surfaces like Cu and Pd, 100% selectivity towards benzene remains elusive due to their ability to break C-H bonds leading to by-products such as ethylene, ethane, and butane in addition to the cyclotrimerization to benzene pathway.

Given the Au-based findings described above, we examined acetylene cyclotrimerization to benzene on Ag(111) because it is generally inert like Au,39 but considerably cheaper. Furthermore, low index Ag surfaces do not reconstruct to large surface unit cells like the $22 \times \sqrt{3}$ herringbone reconstruction exhibited by Au(111), which makes Ag much simpler to accurately model with theory. Using a combination of UHV surface science experiments and density functional theory (DFT) based calculations we discovered that acetylene cyclotrimerizes to benzene with 100% selectivity when densely packed on the Ag(111) surface. Our temperature programmed desorption (TPD) results demonstrate a clear threshold of acetylene surface coverage (>1 ML) for the formation on benzene and reveal that additional acetylene layers promote the conversion to benzene. STM results show the progression of surface acetylene structures and features consistent with the formation of C4 intermediates as the acetylene surface coverage is raised. Our DFT calculations indicate that formation of the C₄ intermediate is the crucial step in determining acetylene reaction vs. desorption. Furthermore, these DFT calculations reveal how multilayers facilitate C4 formation by strengthening interactions with the surface in the transition state more than the initial state, which lowers the C₄

formation barrier. Together these results demonstrate the possibility of 100% selective benzene formation on Ag based catalysts and provide mechanistic insights for potential catalyst development.

Methods

Temperature-programmed desorption (TPD)

TPD studies were conducted using a UHV chamber with a base pressure $< 1 \times 10^{-10}$ mbar equipped with a Hiden Hal RC 201 mass spectrometer which could be advanced to \sim 1 mm from the crystal face. The Ag(111) crystal could be cooled to cryogenic temperatures (\sim 85 K) using liquid nitrogen cooling and heated to 750 K *via* resistive heating of tungsten support wires. The Ag(111) crystal was cleaned using multiple Ar⁺ bombardment cycles (\sim 2 μ A drain current, 1.5 kV) and annealing to 750 K. Acetylene (atomic absorption grade, Airgas) was exposed to the crystal at 90 K. TPDs were performed with a linear heating rate of 1.5 K s⁻¹. Masses tracked: acetylene (m/z = 26), ethylene (m/z = 28), ethane (m/z = 30), 1,3-butadiene (m/z = 54), butene (m/z = 56), benzene (m/z = 78), and cyclooctatetraene (m/z = 104).

Scanning tunneling microscopy (STM)

STM experiments were performed on a low-temperature scanning tunneling microscope (Infinity by Scienta Omicron) in UHV with a base pressure $< 1 \times 10^{-11}$ mbar in the scanning chamber and $\sim 1 \times 10^{-10}$ mbar preparation chamber. The Ag(111) single crystal sample was cleaned with a series of Ar⁺ bombardment and 1000 K annealing cycles. Acetylene was exposed to the sample at ~ 25 K, while STM imaging was performed at ~ 13 K with a W tip.

Computational methods

The DFT calculations were performed with the Vienna *Ab initio* Simulation Package (VASP). 40,41 The Perdew–Burke–Ernzerhof generalized gradient approximation (GGA-PBE) 42 exchange–correlation functional and the Tkatchenko–Scheffler method 43 for van der Waals interactions were used in all the calculations. The cutoff energy was set to 400 eV. A $10 \times 7 \times 1$, $7 \times 7 \times 1$, or $3 \times 5 \times 1$ k-point mesh was used for $2 \times 3 \times 4$, $3 \times 3 \times 4$, and $7 \times 4 \times 4$ unit cells, respectively. The top two layers and adsorbates were fully relaxed during the optimization. The transition states were optimized by the dimer method 44 and were confirmed by frequency calculations. The structures were visualized by VMD. 45 For STM image simulations, the local density of states was sampled over an energy range corresponding to a bias voltage of -300 mV.

Results and discussion

We began by studying the effect of acetylene surface coverage on the selectivity and activity of the acetylene cyclotrimerization reaction on Ag(111) with UHV TPD. This technique enables determination of the reaction selectivity as well as the overall conversion of acetylene to benzene. We exposed the 90 K Ag(111) sample to various Langmuir (1 L = 1 \times 10⁻⁶ torr s)

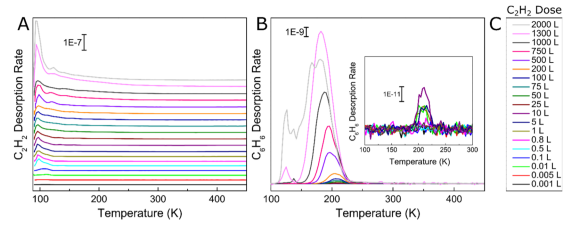


Fig. 1 TPD traces of (A) increasing Langmuir doses of C_2H_2 on Ag(111) and (B) corresponding C_6H_6 production. (C) Legend for C_2H_2 and C_6H_6 traces. All C_2H_2 doses were performed at 90 K and the inset highlights benzene production from acetylene doses of 10 L and below.

doses of acetylene, ramped the temperature at 1.5 K s⁻¹, and tracked the production of benzene with TPD as seen in Fig. 1. Fig. 1A shows acetylene desorption as a function of acetylene coverage and two distinct acetylene desorption features at ~90 K and \sim 130 K are seen. The lower temperature monolayer peak saturates at ~10 L acetylene while the higher temperature desorption peak (attributed to acetylene stabilized by C4 intermediate formation, discussed later) keeps growing as the acetylene coverage increases beyond 10 L. The broad multilayer peak centered at ~100 K continues to grow with increasing amounts of acetylene deposited, albeit at a slower rate. Importantly, very small amounts of benzene production occur below 1 L acetylene as seen in Fig. 1B, consistent with the surface acetylene coverage dependence observed in previous studies. 46,47 The growth of the reactively formed benzene peak is also consistent with the previous benzene adsorption studies on the Ag(111) surface.48 However, to the best of our knowledge this is the first report of acetylene cyclotrimerization to benzene on any Ag surface. A previous study on Ag(110) concluded that adsorbed acetylene desorbs without reaction between 100 and 160 K,49 and another study on Ag(111) showed the reversible desorption of acetylene without trimerization.50 However, our results are still consistent given that both of these previous studies examined much lower acetylene surface coverages than the range studied herein. Furthermore, Ag atoms supported on MgO thin films have also been shown to be inactive for the cyclotrimerization of acetylene whereas Pd and Rh atoms show 100% selective acetylene to benzene conversion.19

To quantify the reaction selectivity using TPD, we tracked acetylene and benzene at m/z 26 and 78 and also m/z of 28, 30, 54, 56 and 104 for ethylene, ethane, 1,3 butadiene, butene and cyclooctatetraene, respectively, which were common side-products in previously reported acetylene trimerization studies. TPD traces corresponding to these products were all flat, with the exception of ethylene (Fig. S2†). However, the observed ethylene appears to come from the chamber background as opposed to being reactively formed on the surface (Fig. S3†). Specifically, experiments in which acetylene and fully

deuterated benzene were deposited on Ag(111) and TPD was performed show that the ratio of C_6D_6 (84) to C_6D_5H (83) on the surface closely resembles the ratio observed during a background C_6D_6 dose revealing that no scrambling of hydrogen into the fully deuterated benzene occurs as seen in Fig. S3.† This result indicates that C–H bonds are not broken on the Ag(111) surface and therefore there is no supply of H needed to form ethylene on the Ag(111) surface. Further evidence for the observed ethylene originating from background adsorption comes from the fact that the Ag(111) surface is equally active for many reaction cycles without cleaning in between, indicating that there is no buildup of surface carbon, as would be expected if ethylene were a real product that required a supply of H from acetylene.

This demonstration that the acetylene trimerization to benzene is 100% selective on Ag(111) is supported by literature reports of the inability of the Ag(111) surface to break C-H bonds⁵¹⁻⁵⁴ which is a requirement to form products like ethylene and butadiene. ^{17,21,29,36} Furthermore, given that benzene is aromatic while cyclooctatetraene is not, benzene is the preferred product both enthalpically and entropically. ^{55,56} Specifically, enthalpically, due to its aromatic ring, benzene is more thermodynamically stable product than cyclooctatetraene. Entropically the formation of a smaller gas phase molecule (benzene) *versus* cyclooctatetraene makes the formation of benzene thermodynamically more favorable.

We further investigated the reactivity of Ag(111) towards acetylene cyclotrimerization by integrating the acetylene and benzene TPD peaks then applying correction factors to account for fragmentation pattern, ionization cross-section, and mass spectrometer quadrupole sensitivity.⁵⁷ We then calculated the experimental conversion of acetylene as a ratio between the acetylene converted to benzene and the total amount of acetylene on the surface, as depicted in Fig. 2. The monolayer coverages of acetylene and benzene were defined by the saturation point for the low-temperature (90 K) acetylene peak and a separate uptake for benzene on Ag(111) respectively (see Fig. S2†). Fig. 2A demonstrates that acetylene conversion to

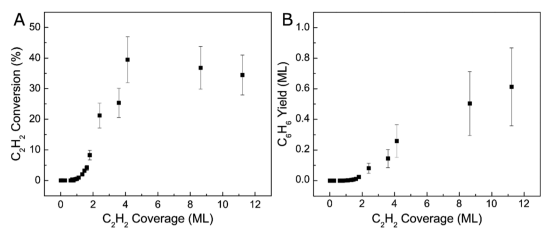


Fig. 2 Coverage-dependent acetylene cyclotrimerization to benzene on Ag(111). (A) Acetylene conversion with respect to acetylene surface coverage and (B) benzene yield in monolayers plotted against acetylene surface coverage.

benzene is very low when the acetylene surface coverage is low. However, the conversion increases abruptly when a saturated acetylene monolayer is formed and reaches a maximum of \sim 40% at \sim 4 ML. We hypothesize this phenomenon is due to the additional acetylene layers which facilitate tighter packing of acetylene molecules, leading to lower barriers for coupling, which will be explored further with DFT.31 Our TPD data indicate that additional acetylene adsorption beyond 4 ML still produces more benzene as shown in Fig. 2B, but does not improve the conversion. This may be due to the fact that acetylene does not grow strictly layer-by-layer on Ag(111), or due to effects beyond the second layer.⁵⁸ Moreover, the total benzene yield continues to increase with acetylene coverage and does not saturate up to 12 ML acetylene. These results highlight the previously unconsidered role of 3D molecular overlayers on the conversion of acetylene to benzene.

In order to probe the robustness of Ag, we examined if the acetylene trimerization reaction poisoned the Ag(111) surface since catalyst deactivation was common in previously reported

trimerization systems like Ni/SiO₂, SiO₂/Si(100) and fluorine on alumina.^{17,38,59} Specifically, we performed five consecutive cycles of TPD experiments without cleaning the Ag(111) in between as shown in Fig. 3. As is evident from the benzene desorption TPD results, the benzene yield in each TPD experiment is approximately the same, demonstrating that Ag(111) is not poisoned by the buildup of carbonaceous fragments like other active trimerization surfaces. This result is consistent with the inability of the Ag(111) surface to break C-H bonds which leads to carbon deposition.⁶⁰ Lack of poisoning is also consistent with 100% selectivity because side products such as ethylene would lead to carbon build up and result in deactivation.

We then studied acetylene cyclotrimerization to benzene at the molecular level by conducting STM experiments. Fig. 4 panels (A) and (E) show large scale and atomically resolved images of the clean Ag(111) surface before C_2H_2 exposure. At low coverages (B + F) acetylene forms 2D islands of square-packed acetylene molecules on Ag(111) at 25 K. At monolayer coverage (which corresponds to packing density of 1 acetylene

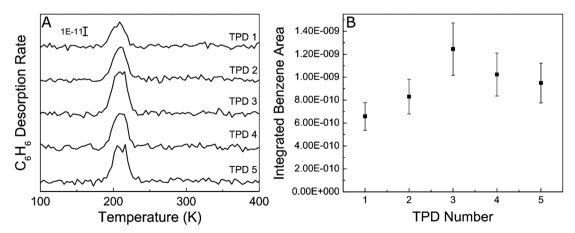


Fig. 3 (A) Consecutive TPD experiments showing C_6H_6 yield after 10 L C_2H_2 exposure on Ag(111) at 90 K. (B) Benzene desorption peak areas calculated by integrating the benzene desorption peaks in panel (A). The error bars were calculated repeating 10 L acetylene exposures 5 times and taking the % error of the benzene yields.

per 3.2 Ag surface atoms as calculated from STM experiments) (C) these islands merge into a complete monolayer composed of separate rotational domains of the square-packed acetylene structure due to their alignment with one of the three equivalent high-symmetry axes of the Ag(111) lattice. Four of these rotational domains can be seen in Fig. 4C and the inset in panel (G) shows the square packing structure of the acetylene monolayer. At acetylene coverages above one monolayer, growth of a second layer of acetylene is observed (D) which coincides with the appearance of dumbbell-shaped protrusions consistent with the proposed C4 intermediates within the first acetylene monolayer (H). Moreover, panel (H) reveals that the regions containing the intermediates have converted from a square to a hexagonal packing structure after the first monolayer is reached. These data indicate that the addition of >1 ML acetylene to Ag(111) causes a denser, hexagonal acetylene packing structure (1 acetylene per 2.7 Ag surface atoms) to form, which coincides with the appearance of dumbbell shaped protrusions which we speculate are the proposed C4 intermediates, and we investigate this point further with DFT.

In order to further examine the structure and stability of various configurations and coverages of acetylene, DFT was used to calculate energetics and generate simulated STM images. We first studied the acetylene adsorption energy in several structures and for different coverages, initially focusing on mimicking the square-packed structures seen in STM. The calculations generally predict stronger adsorption as the acetylene packing density increases, which is consistent with the experimental result that acetylene forms compact islands

surrounded by bare Ag(111) at <1 ML coverage (see Fig. 4B). For example, the adsorption energy of a single, isolated acetylene is -0.29 eV, while the presence of a co-adsorbed, parallel acetylene in a 2 \times 3 unit cell strengthens the adsorption energy to -0.35 eV (Fig. S5a†). This strengthening is likely due to van der Waals interactions between acetylene molecules and explains the islanding observed in experiments (Fig. 4B).

To mimic the structure seen in experimental STM at higher acetylene coverage containing the proposed C_4 intermediates, the simulated STM image of 3 C_2H_2 molecules and 2 C_4H_4 intermediates in a 7 \times 4 unit cell is shown in Fig. S5.† This simulated STM image is similar to the features seen in Fig. 4H, indicating that the large bright species in Fig. 4H are likely the C_4 intermediates surrounded by acetylene molecules which appear as smaller protrusions in the simulation.

To understand critical steps in the reaction, we studied the full acetylene trimerization pathway to benzene on a 3 \times 3 surface cell (Fig. 5). At 100 K and 1 atm, the transition state for forming the C_4 intermediate is very close to zero in free energy (-0.01 eV), where the zero is set to the acetylene molecules in the gas phase. Therefore, the competition between reaching this transition state and acetylene desorption will control the reaction efficiency. Even small changes in the energy of this transition state could move it above or below the energy zero and therefore have a large impact on the conversion to benzene ν s. desorption. Furthermore, Fig. 5 illustrates that once the C_4 intermediate is formed, the rest of the reaction pathway to benzene is significantly below 0 eV, generally downhill in energy, and features no large barriers, further indicating that

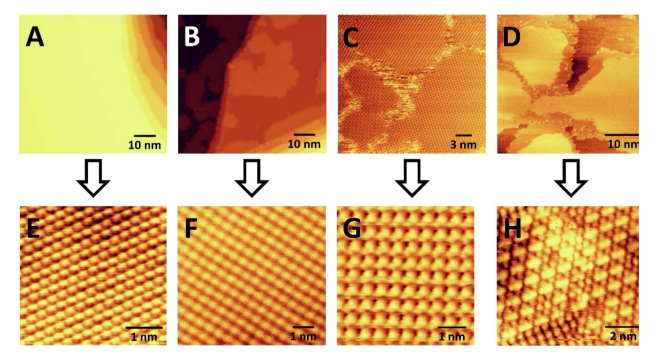


Fig. 4 STM images of (A) pure Ag(111), (B) 1 L C_2H_2 on Ag(111), (C) 3 L C_2H_2 on Ag(111), (D) 10 L C_2H_2 on Ag(111), and (E) atomically resolved STM image of pure Ag(111). STM images in panels (F) and (G) show the square packing of C_2H_2 at lower acetylene coverage. Panel (H) shows the hexagonal packing of the C_2H_2 at higher coverage and the appearance of dumbbell shaped intermediates. C_2H_2 was exposed to the surface at \sim 25 K and the images were taken at \sim 13 K. Imaging conditions: (A) and (E) 0.2 nA and 600 mV, (B) 0.02 nA and 900 mV, (F) 0.03 nA and 900 mV, (C) 0.1 nA and \sim 700 mV, (G) 0.1 nA and \sim 500 mV, (D) and (H) 0.1 nA and 800 mV.

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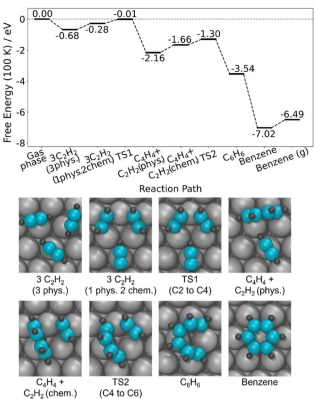


Fig. 5 Calculated reaction pathway of acetylene trimerization to benzene on Ag(111). The optimized structures along the pathway in a 3 imes 3 unit cell are shown. Free energies calculated at 100 K, 1 atm.

the formation of the C4 intermediate has the largest degree of rate control.⁶¹ These DFT calculations also indicate that in order to form the C₄ intermediate, two physiosorbed acetylene molecules must first become chemisorbed and bind to one Ag atom. For a single acetylene molecule, the chemisorbed state is 0.09 eV less stable than the physisorbed state, and this difference is expected to depend on the structure and packing density of the acetylene monolayer. Therefore, understanding the energetics of moving from the all-physisorbed state to a state with two chemisorbed acetylene molecules and then to the transition state for C₄H₄ formation as a function of acetylene coverage and structure should provide deeper insight into the reaction pathway and ultimate conversion to benzene.

We therefore studied the C4 formation step with DFT for several different acetylene packing structures in order to understand the influence of acetylene packing density and structure on benzene formation. Structures where all acetylene molecules are directly interacting with the surface were formed by placing two and three acetylene molecules in different unit cells and keeping the packing density ≤ 1 acetylene per 3 surface Ag atoms (for example, Fig. S5a†). Of these structures, three acetylene molecules in a 3×3 unit cell give the most stable configurations for both the all-physisorbed state and the state with two chemisorbed acetylene molecules. When increasing the packing density (see the structures in Fig. 6), our DFT calculations predict that acetylene molecules start forming the second layer when the packing density reaches 1 acetylene per 3 surface Ag atoms and there are three acetylene layers when the packing density is 1:1.

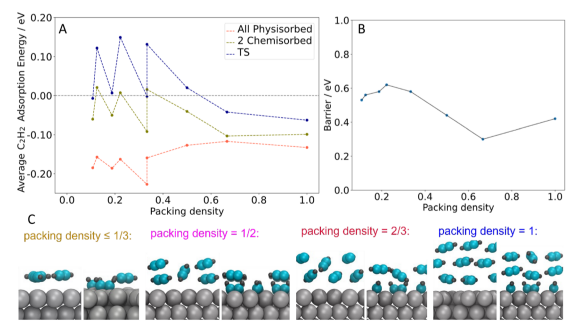


Fig. 6 The C_4 intermediate formation step with different packing densities. (A) Free energies for C_2H_2 in various states on the Ag(111) surface as a function of packing density. (B) C₄ formation barrier, calculated from the lowest energy state to the transition state (TS1), as a function of packing density. All packing densities given in this work are in terms of C_2 unit per surface Ag atom. (C) Side views of DFT optimized structures of C₂H₂ adsorption on the Ag(111) surface with different packing densities.

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To examine the mechanism through which physisorbed multilayers may promote the formation of benzene, we calculated the average acetylene adsorption energy for the all-physisorbed structure (initial state, lowest energy), the state with two chemisorbed acetylene molecules (precursor to the transition state for C–C coupling), and the transition state for C₄ formation (Fig. 6A). This allowed us to understand how the absolute and relative stabilities of these states change as a function of coverage. We also calculated the barrier for C_4 formation, defined as the difference in energy between the transition state and the most-stable state, which is always the all-physisorbed state (Fig. 6B).

For the coverages where acetylene forms a single layer (packing density ≤ 1 acetylene per 3 surface Ag atoms), the energies of all three states are highly correlated, showing very similar shifts (see Fig. 6A). These shifts are primarily due to the number of acetylene molecules in the unit cell: the structures in question all have two or three acetylene molecules present, and when there are three present the energy is shifted down. However, these shifts have only a small effect on the barrier (see Fig. 6B).

On the other hand, when >1 ML of acetylene is present (packing density ≥ 1 acetylene per 3 surface Ag atoms, all in 2×3 unit cells), the average stability in the physisorbed state shifts in the opposite direction of the chemisorbed state and transition state (see Fig. 6A). The stabilization of the chemisorbed state at higher coverages is likely due to the second layer of acetylene moving closer to the surface during the chemisorption process, strengthening its interaction with the surface (see Fig. 6C). Examining the stability of just the physisorbed molecules in each state supports this picture (see Fig. S7†). This trend applies to the transition state as well, resulting in a significant decrease in the barrier (Fig. 6B). The lowest barrier is 0.3 eV, from 4 acetylene molecules in a 2×3 unit cell (2 acetylene layers, packing density = 2 acetylene per 3 surface Ag atoms corresponding to \sim 2 ML in experiment) Together, these results indicate that when multilayers of acetylene are present, there is a driving force towards chemisorption: the increased energy for the chemisorbed or reacting molecules themselves is partially compensated by a stronger interaction between the second layer of acetylene and the Ag surface. This explains the effect of acetylene multilayers on the reaction, and also explains the DFT results which demonstrate how the reaction barriers decrease as the acetylene packing density increases beyond 1 acetylene per 3 surface Ag atoms, while the barriers are not sensitive to acetylene coverage at lower coverages than this. Additionally, the predicted trend is consistent with the experimental finding that benzene production begins around 1 ML acetylene, which corresponds to a packing density of roughly 1 acetylene per 3 surface Ag atoms (the square packing structure observed in STM contained 1 acetylene per 3.2 surface Ag atoms vs. the hexagonal packing structure which has 1 acetylene per 2.7 surface Ag atoms). Together, these results highlight the importance of the change in packing structure to a higher density, as occurs when the acetylene monolayer transitions from square to hexagonal, offering a lower reaction barrier that also benefits from multilayer acetylene adsorption.

Our results show a clear trend of a generally decreasing energetic barrier with increasing packing density, with quite significant variations in the barrier height. However, due to the complexity of the multilayer structures, we expect some quantitative error in the barrier heights due to the structures we use, in addition to the intrinsic error from DFT. Indeed, even our lowest barrier would suggest very slow formation of the C₄ intermediates at 25 K, and yet STM suggests that these intermediates are formed at low temperature. In experiment, C₄ intermediates may form at locations where the local molecular packing structure leads to particularly low reaction barriers.

To explain the high selectivity and lack of poisoning observed experimentally, we also calculated the possibility of C–H bond cleavage in acetylene on Ag(111). As shown in Fig. S8,† the transition state and product of C–H cleavage have much higher energies than those in the trimerization pathway. In particular, desorption (0.29 eV) is far more favorable than C–H bond cleavage (barrier of 1.43 eV). This implies that C–H bonds in acetylene are unlikely to be broken on Ag(111) and explains the experimental findings that acetylene can be transformed to benzene with 100% selectivity on Ag(111) versus more reactive surfaces like Pd(111) and PdAu alloys which are capable of C–H activation and are thus considerably less selective to benzene.

Conclusions

Using temperature programmed desorption, molecular scale imaging, and DFT, we discovered that acetylene cyclotrimerizes to benzene with 100% selectivity on the Ag(111) surface. Specifically, benzene formation requires a critical surface coverage of acetylene > 1 ML, after which the benzene formation dramatically increases. Importantly, the Ag(111) surface does not deactivate after multiple reaction cycles exhibiting resilience to poisoning unlike the Cu, Ni, Si and Pd based cyclotrimerization systems reported in the literature. STM imaging demonstrated that above the critical coverage of acetylene, the acetylene monolayer converts from a square to a denser hexagonal packing structure, and bright features consistent with the proposed C₄ intermediates are observed. DFT results indicate that reaching the transition state for forming the C4 intermediate is most important for determining the reaction efficiency and that this transition state, as well as the transformation of physisorbed acetylene to chemisorbed acetylene, can be promoted by the adsorption of additional acetylene layers. We attribute the lack of any side products such as ethylene and butadiene to the inability of the Ag(111) surface to perform dehydrogenation, which has a much higher calculated energy barrier. Together, these results provide fundamental understanding of the 100% selective acetylene cyclotrimerization to benzene on Ag(111) which we hope informs the design of Ag-based nanoparticle catalysts for this reaction which will be of greater importance as the move away from heavy hydrocarbon feedstocks continues.

Data availability

The datasets supporting this article have been uploaded as part of the ESI. \dagger

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Author contributions

Volkan Cinar carried out the TPD experiments, data analysis and writing of the original manuscript. Elizabeth E. Happel performed the STM experiments. Nipun T. S. K. Dewage conducted control experiments. E. Charles H. Sykes supervised the TPD and STM experiments, data curation and writing of the original draft. Shengjie Zhang carried out the DFT calculations under Matthew M. Montemore's supervision who also edited the manuscript.

Conflicts of interest

There are no conflicts of interest to declare.

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Notes and references

- 1 Benzene Market Global Industry Analysis, Size Share, Growth, Trends, Regional Outlook, and Forecast 2022 - 2030, https:// www.precedenceresearch.com/benzene-market.
- 2 A. Mandaokar, Benzene Market Research Report Forecast Till https://www.marketresearchfuture.com/reports/ benzene-market-928.
- 3 Q. Meng, J. Yan, R. Wu, H. Liu, Y. Sun, N. N. Wu, J. Xiang, L. Zheng, J. Zhang and B. Han, Sustainable Production of Benzene from Lignin, Nat. Commun., 2021, 12(1), 1-12, DOI: 10.1038/s41467-021-24780-8.
- 4 W. Fruscella, Kirk-Othmer Encyclopedia Chemical Technology, 2001, vol. 3, pp. 596-624.
- 5 National Library of Medicine, Benzene, https:// pubchem.ncbi.nlm.nih.gov/compound/Benzene.
- 6 E. Kintisch, Making Dirty Coal Plants Cleaner, Science, 2007, 317(5835), 184-186, DOI: 10.1126/science.317.5835.184.
- 7 J. Chow, R. J. Kopp and P. R. Portney, Energy Resources and Global Development, Science, 2003, 302(5650), 1528-1531, DOI: 10.1126/science.1091939.
- 8 A. Niziolek, O. Onel and C. Floudas, Production of Benzene, Toluene, and Xylenes from Natural Gasvia Methanol: Process Synthesis and Global Optimization, AIChE J., 2016, 62(5), 1401-1843, DOI: 10.1002/aic.15144.

- 9 J. C. Bertolini, J. Massardier and G. Dalmai-Imelik, Evolution of Adsorbed Species during C₂H₂ Adsorption on Ni(111) in Relation to Their Vibrational Spectra, J. Chem. Soc., Faraday Trans. 1, 1978, 74(111), 1720-1725, DOI: 10.1039/ F19787401720.
- 10 W. Sesselmann, B. Woratschek, G. Ertl, J. Küppers and H. Haberland, Low Temperature Formation of Benzene from Acetylene on a Pd(111) Surface, Surf. Sci., 1983, 130(2), 245-258, DOI: 10.1016/0039-6028(83)90359-X.
- 11 K. G. Pierce and M. A. Barteau, Cyclotrimerization of Alkynes on Reduced TiO₂ (001) Surfaces, J. Phys. Chem., 1994, (98), 3882-3892.
- 12 C. J. Baddeley, M. Tikhov, C. Hardacre, J. R. Loinas and R. M. Lambert, Ensemble Effects in the Coupling of Acetylene to Benzene on a Bimetallic Surface: A Study with Pd{lll}/Au, J. Phys. Chem., 1996, 100(6), 2189-2194, DOI: 10.1021/jp9517054.
- 13 A. F. Lee, C. J. Baddeley, C. Hardacre, G. D. Moggridge, R. M. Ormerod, R. M. Lambert, J. P. Candy and J. M. Basset, Structure-Reactivity Correlations in the Catalytic Coupling of Ethyne over Novel Bimetallic Pd/Sn Catalysts, J. Phys. Chem. B, 1997, 101(15), 2797-2805, DOI: 10.1021/jp9638346.
- 14 V. S. Lusvardi, K. G. Pierce and M. A. Barteau, Steady-State Catalytic C-C Bond Formation on Reduced TiO2 Surfaces, I. Vac. Sci. Technol., A, 1997, 15(3), 1586-1591, DOI: 10.1116/1.580637.
- 15 W. T. Tysoe, Palladium-Catalyzed Acetylene Cyclotrimerization: From Ultrahigh Vacuum to High-Pressure Catalysis, Isr. J. Chem., 1998, 38(4), 313-320, DOI: 10.1002/ijch.199800036.
- 16 J. Storm, R. M. Lambert, N. Memmel, J. Onsgaard and E. Taglauer, Catalysis with Monometallic and Bimetallic Films: Trimerization of Acetylene to Benzene on Pd/ Ru(001) and (Pd + Au)/Ru(001), Surf. Sci., 1999, 436(1), 259-268, DOI: 10.1016/S0039-6028(99)00672-X.
- 17 E. Sorek, G. Arbiv and M. Asscher, Medium-Pressure Reactivity of Acetylene on Pd-Cu Alloy Nanoparticles Supported on Thin Silica Films, Langmuir, 2020, 36(28), 8066-8074, DOI: 10.1021/acs.langmuir.0c00592.
- 18 W. Gan, L. Geng, B. Yin, H. Zhang, Z. Luo and K. Hansen, Cyclotrimerization of Acetylene on Clusters Co_n⁺/Fe_n⁺/ $Ni_n^+(n = 1-16)$, J. Phys. Chem. A, 2021, 125(48), 10392-10400, DOI: 10.1021/acs.jpca.1c09015.
- 19 K. Judai, A. S. Wörz, S. Abbet, J. M. Antonietti, U. Heiz, A. Del Vitto, L. Giordano and G. Pacchioni, Acetylene Trimerization on Ag, Pd and Rh Atoms Deposited on MgO Thin Films, Phys. Chem. Chem. Phys., 2005, 7(5), 955-962, DOI: 10.1039/ b414399j.
- 20 S. Abbet, A. Sanchez, U. Heiz, W. D. Schneider, A. M. Ferrari, G. Pacchioni and N. Rösch, Acetylene Cyclotrimerization on Supported Size-Selected Pd_n Clusters ($1 \le n \le 30$): One Atom Is Enough!, J. Am. Chem. Soc., 2000, 122(14), 3453–3457, DOI: 10.1021/ja9922476.
- 21 W. T. Tysoe, G. L. Nyberg and R. M. Lambert, Low Temperature Catalytic Chemistry of the Pd(111) Surface:

Benzene and Ethylene from Acetylene, *J. Chem. Soc. Chem. Commun.*, 1983, (11), 623–625, DOI: 10.1039/c39830000623.

Chemical Science

- 22 T. Okada, Y. Kim, M. Trenary and M. Kawai, Identification at the Single Molecule Level of C₂H_x Moieties Derived from Acetylene on the Pt(111) Surface, *J. Phys. Chem. C*, 2012, **116**(34), 18372–18381, DOI: **10.1021/jp307676f**.
- 23 T. V. W. Janssens, S. Völkening, T. Zambelli and J. Wintterlin, Direct Observation of Surface Reactions of Acetylene on Pd(111) with Scanning Tunneling Microscopy, *J. Phys. Chem. B*, 1998, **102**(34), 6521–6528, DOI: **10.1021**/**ip981383e**.
- 24 J. Dunphy, M. Rose, S. Behler, D. Ogletree, M. Salmeron and P. Sautet, Acetylene Structure and Dynamics on Pd(111), *Phys. Rev. B: Condens. Matter Mater. Phys.*, 1998, 57(20), R12705–R12708, DOI: 10.1103/PhysRevB.57.R12705.
- 25 E. F. Lutz, The Cyclic Trimerization of Acetylenes Over a Ziegler Catalyst, *J. Am. Chem. Soc.*, 1961, 83(11), 2551–2554, DOI: 10.1021/ja01472a029.
- 26 N. Zelinsky, Uber Die Kontakt-Kondensation Des Acetylens, *Eur. J. Inorg. Chem.*, 1924, 57, 264–276.
- 27 H. Öberg, Y. Nestsiarenka, A. Matsuda, J. Gladh, T. Hansson, L. G. M. Pettersson and H. Öström, Adsorption and Cyclotrimerization Kinetics of C₂H₂ at a Cu(110) Surface, *J. Phys. Chem. C*, 2012, 116(17), 9550–9560, DOI: 10.1021/jp300514f.
- 28 J. R. Lomas, C. J. Baddeley, M. S. Tikhov and R. M. Lambert, Ethyne Cyclization to Benzene over Cu(110), *Langmuir*, 1995, 11(8), 3048–3053, DOI: 10.1021/la00008a033.
- 29 T. M. Gentle and E. L. Muettertles, Acetylene, Ethylene, and Arene Chemistry of Palladium Surfaces, *J. Phys. Chem.*, 1983, 87(14), 2469–2472, DOI: 10.1021/j100237a007.
- 30 N. R. Avery, Adsorption and Reactivity of Acetylene on a Cu(110) Surface, J. Am. Chem. Soc., 1985, (6), 6711–6712.
- 31 R. M. Lambert and R. M. Ormerod, Heterogeneously Catalysed Cyclisation Reactions of Ethyne over Single Crystal Palladium and Palladium Catalysts, *Mater. Chem.*, 1991, 29, 105–115.
- 32 J. Szanyi and M. T. Paffett, Di- and Trimerization of Acetylene over a Model Sn/Pt Catalyst (V3xV3) R30° Surface Alloy, *J. Am. Chem. Soc.*, 1995, 27, 1034–1042.
- 33 C. Xu, J. W. Peck and B. E. Koel, A New Catalysis for Benzene Production from Acetylene under UHV Conditions: Sn/Pt(111) Surface Alloys, *J. Am. Chem. Soc.*, 1993, 115(2), 751–755, DOI: 10.1021/ja00055a054.
- 34 R. M. Ormerod, R. M. Ormerod, R. M. Lambert, R. M. Lambert, H. Hoffmann, H. Hoffmann, F. Zaera, F. Zaera, J. M. Yao, J. M. Yao, D. K. Saldin, D. K. Saldin, L. P. Wang, L. P. Wang, D. W. Bennett, D. W. Bennett, W. T. Tysoe and W. T. Tysoe, NEXAFS Identification C₄H₄ on Pd(111), *Surf. Sci.*, 1993, 5, 277–286.
- 35 C. J. Baddeley, R. M. Ormerod, A. W. Stephenson and R. M. Lambert, Surface Structure and Reactivity in the Cyclization of Acetylene to Benzene with Pd Overlayers and Pd/Au Surface Alloys on Au{111}, *J. Phys. Chem.*, 1995, 99(14), 5146–5151, DOI: 10.1021/j100014a039.
- 36 W. T. Tysoe, G. L. Nyberg and R. M. Lambert, Photoelectron Spectroscopy and Heterogeneous Catalysis: Benzene and

- Ethylene from Acetylene on Palladium (111), *Surf. Sci.*, 1983, 135(1-3), 128-146, DOI: 10.1016/0039-6028(83)90215-7.
- 37 G. Kyriakou, J. Kim, M. S. Tikhov, N. Macleod and R. M. Lambert, Acetylene Coupling on Cu(111): Formation of Butadiene, Benzene, and Cyclooctatetraene, *J. Phys. Chem. B*, 2005, **109**(21), 10952–10956, DOI: **10.1021**/jp044213c.
- 38 A. G. Boudjahem, S. Monteverdi, M. Mercy and M. M. Bettahar, Acetylene Cyclotrimerization over Ni/SiO₂ Catalysts in Hydrogen Atmosphere, *Appl. Catal.*, *A*, 2003, 250(1), 49–64, DOI: 10.1016/S0926-860X(03)00221-7.
- 39 G. O. Kayode and M. M. Montemore, Factors Controlling Oxophilicity and Carbophilicity of Transition Metals and Main Group Metals, *J. Mater. Chem. A*, 2021, 9(39), 22325–22333, DOI: 10.1039/d1ta06453c.
- 40 G. Kresse and J. Hafner, Ab Initio Molecular Dynamics for Liquid Metals, *Phys. Rev. B: Condens. Matter Mater. Phys.*, 1993, 47(1), 558–561, DOI: 10.1103/PhysRevB.47.558.
- 41 G. Kresse and J. Furthmüller, Efficiency of Ab-Initio Total Energy Calculations for Metals and Semiconductors Using a Plane-Wave Basis Set, *Comput. Mater. Sci.*, 1996, **6**(1), 15–50, DOI: **10.1016/0927-0256(96)00008-0**.
- 42 J. P. Perdew, K. Burke and M. Ernzerhof, Generalized Gradient Approximation Made Simple, *Phys. Rev. Lett.*, 1996, 77(18), 3865–3868, DOI: 10.1103/PhysRevLett.77.3865.
- 43 A. Tkatchenko and M. Scheffler, Accurate Molecular van Der Waals Interactions from Ground-State Electron Density and Free-Atom Reference Data, *Phys. Rev. Lett.*, 2009, **102**(7), 6–9, DOI: **10.1103/PhysRevLett.102.073005**.
- 44 A. Heyden, A. T. Bell and F. J. Keil, Efficient Methods for Finding Transition States in Chemical Reactions: Comparison of Improved Dimer Method and Partitioned Rational Function Optimization Method, *J. Chem. Phys.*, 2005, **123**(22), 224101, DOI: **10.1063**/1.2104507.
- 45 W. Humphrey, A. Dalke and K. Schulten, VMD: Visual Molecular Dynamics, *J. Mol. Graph.*, 1996, **14**, 33–38.
- 46 A. F. Lee, R. Mark Ormerod, K. Wilson, R. M. Lambert, A. Goldoni and A. Baraldi, A Fast XPS Investigation of NO-Promoted Acetylene Cyclotrimerisation on Pd{111}, Surf. Sci., 2002, 501(1-2), L165-L170, DOI: 10.1016/S0039-6028(01)01982-3.
- 47 J. Dvorak and J. Hrbek, Adsorbate Ordering Effects in the Trimerization Reaction of Acetylene on Cu(100), *J. Phys. Chem. B*, 1998, **102**(47), 9443–9450, DOI: **10.1021/jp981956n.**
- 48 T. J. Rockey, M. Yang and H. L. Dai, Adsorption Energies, Inter-Adsorbate Interactions, and the Two Binding Sites within Monolayer Benzene on Ag(111), *J. Phys. Chem. B*, 2006, **110**(40), 19973–19978, DOI: **10.1021/jp062225n**.
- 49 E. M. Stuve, R. J. Madix and B. A. Sexton, Characterization of the Adsorption and Reaction of Acetylene on Clean and Oxygen Covered Ag(110) by EELS, *Surf. Sci.*, 1982, 123(2–3), 491–504, DOI: 10.1016/0039-6028(82)90342-9.
- 50 D. L. Molina, M. Muir, M. K. Abdel-Rahman and M. Trenary, The Influence of Palladium on the Hydrogenation of Acetylene on Ag(111), *J. Chem. Phys.*, 2021, **154**(18), 184701, DOI: **10.1063**/5.0050587.

51 H. Y. Gao, J. H. Franke, H. Wagner, D. Zhong, P. A. Held, A. Studer and H. Fuchs, Effect of Metal Surfaces in On-Surface Glaser Coupling, *J. Phys. Chem. C*, 2013, **117**(36), 18595–18602, DOI: **10.1021/jp406858p**.

Edge Article

- 52 J. Björk, Y. Q. Zhang, F. Klappenberger, J. V. Barth and S. Stafström, Unraveling the Mechanism of the Covalent Coupling between Terminal Alkynes on a Noble Metal, *J. Phys. Chem. C*, 2014, **118**(6), 3181–3187, DOI: **10.1021/jp5002918**.
- 53 Y. Q. Zhang, N. Kepčija, M. Kleinschrodt, K. Diller, S. Fischer, A. C. Papageorgiou, F. Allegretti, J. Björk, S. Klyatskaya, F. Klappenberger, M. Ruben and J. V. Barth, Homo-Coupling of Terminal Alkynes on a Noble Metal Surface, *Nat. Commun.*, 2012, 3, 1–8, DOI: 10.1038/ ncomms2291.
- 54 J. Liu, Q. Chen, Q. He, Y. Zhang, X. Fu, Y. Wang, D. Zhao, W. Chen, G. Q. Xu and K. Wu, Bromine Adatom Promoted C-H Bond Activation in Terminal Alkynes at Room Temperature on Ag(111), *Phys. Chem. Chem. Phys.*, 2018, 20(16), 11081–11088, DOI: 10.1039/c7cp07972a.
- 55 D. H. Lo and M. A. Whitehead, Molecular Geometry and Bond Energy. III. Cyclooctatetraene and Related Compounds, *J. Am. Chem. Soc.*, 1969, **91**, 238–242.

- 56 G. E. Davico, V. M. Bierbaum, C. H. DePuy, G. B. Ellison and R. R. Squires, The C-H Bond Energy of Benzene, *J. Am. Chem. Soc.*, 1995, 117(9), 2590–2599, DOI: 10.1021/ja00114a023.
- 57 C. G. F. Siler, T. Cremer, J. C. F. Rodriguez-Reyes, C. M. Friend and R. J. Madix, Switching Selectivity in Oxidation Reactions on Gold: The Mechanism of C-C vs C-H Bond Activation in the Acetate Intermediate on Au(111), ACS Catal., 2014, 4(9), 3281–3288, DOI: 10.1021/cs500803n.
- 58 G. Gonella, H. L. Dai and T. J. Rockey, Tetracene Monolayer and Multilayer Thin Films on Ag(111): Substrate-Adsorbate Charge-Transfer Bonding and Inter-Adsorbate Interaction, *J. Phys. Chem. C*, 2008, 112(12), 4696–4703, DOI: 10.1021/jp709826q.
- 59 V. M. Allenger, J. R. Brown, D. Clugston, M. Ternan and D. D. McLean, Catalyst Deactivation in Acetylene Polymerization, *Appl. Catal.*, 1988, 39(C), 191–211, DOI: 10.1016/S0166-9834(00)80949-5.
- 60 W. H. Weinberg and T. E. Felter, An XPS and UPS Investigation of the Adsorption of Ethylene on the (111) Surface of Silver, Surf. Sci. Lett., 1980, 97, L313–L319.
- 61 C. T. Campbell, The Degree of Rate Control: A Powerful Tool for Catalysis Research, *ACS Catal.*, 2017, 7(4), 2770–2779, DOI: 10.1021/acscatal.7b00115.