Ethene to Graphene: Surface Catalyzed Chemical Pathways, Intermediates, and Assembly

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ABSTRACT: Diverse technologies from catalyst coking to graphene synthesis entail hydrocarbon dehydrogenation and condensation reactions on metals and assembly into carbon overlayers. Imperative to gaining control over these processes, through thermal steering of the formation of polyaryl intermediates and the controlled prevention of coking, is the exploration and elucidation of the detailed reaction scheme that starts with adsorbed hydrocarbons and culminates with the formation of extended graphene. Here we use scanning tunneling microscopy, high-resolution electron energy loss and thermal desorption spectroscopies, in combination with theoretical simulations to uncover the hierarchy of pathways and intermediates underlying the catalyzed evolution of ethene adsorbed on Rh(111) to form graphene. These investigations allow formulation of a reaction scheme whereby, upon heating, adsorbed ethene evolves via coupling reactions to form segmented one-dimensional polyaromatic hydrocarbons (1D-PAH). Further heating leads to dimensionality crossover (1D → 2D) and dynamical restructuring processes at the PAH chain ends with subsequent activated detachment of size-selective carbon clusters. Rate-limiting diffusional coalescence of these dynamically self-evolved precursors culminates (<1000 K) in condensation into graphene of high structural perfection.

INTRODUCTION

The formation of solid carbonaceous, carbidic, or graphitic material known as coke,1,2 derived through anaerobic processing of organic material on metal surfaces at elevated temperatures, has been known since antiquity and patented since the late 16th century for iron ore smelting and making of steel.3 Later, it has been found that coking processes, involving catalytic dehydrogenation, polymerization, and cyclization of hydrocarbons, are detrimental to the effective operation of heterogeneous catalytic reactors, necessitating regeneration or exchange of fouled catalytic material.1,2 On the other hand, as shown here, somewhat serendipitously, the thermally activated surface-catalyzed reaction processes that underlie undesirable coking may be steered toward the synthesis of a graphene overlayer of macroscopic lateral dimensions, highly valued for its unique physical and chemical properties.4,5

Following the isolation of graphene in 2004,1 top-down methods,6−10 including plasma etching lithography and carbon nanotube unzipping, as well as bottom-up techniques,11−14 based on chemical vapor deposition (CVD) or the application of surface-supported organic synthesis routes (e.g., coupling of polyphenylene precursors on metal substrates) have been employed in preparation of graphene structures with ever increasing perfection (see ref 15 for discussion of recent atomically precise bottom-up preparations). Early investigations yielded a wealth of information about the surface chemistry (e.g., catalyzed C−H bond activation and coupling reactions) of small alkane and alkene molecules on metal surfaces (see refs 16 and 17, and references therein). More recent studies (in particular on ethene deposited on transition metal surfaces) have utilized modern surface science techniques (low-energy electron microscopy, low-energy electron microscopy,18,19 X-ray photoelectron spectroscopy,20 and scanning tunneling microscopy (STM)21), extended observations into the graphene formation domain, and resulted in valuable insights into the growth kinetics, suggesting the role of carbon clusters (comprised of 5,18,19 21,22,23 or 2421 atoms) whose attachment, rather than that of monomers, leads to ripening and graphene growth; the carbon-cluster hypothesis has been incorporated in a phenomenological kinetic Monte Carlo simulation of the growth process.22
In the next section, we present our results and discussion. We begin with an overview of the reaction scheme starting from ethene and ending with graphene and then discuss in detail experimental and theoretical results pertaining to the different steps in the general reaction scheme. These include ethene deposition and early reaction stages; formation of segmented one-dimensional polyaromatic hydrocarbon (1D-PAH) narrow ribbons; kinking and one-dimensional to two-dimensional (1D → 2D) dimensionality crossover of the 1D-PAH chains; theoretically simulated pathways and microscopic formation mechanisms of the self-evolved 24-carbon-cluster precursors and graphene growth. After the concluding remarks, we describe the experimental methods, and details of the computational techniques used in our investigations.

RESULTS AND DISCUSSION

Reaction Scheme. The research reported herein focuses on gaining insights into the atomic-scale mechanisms of bottom-up chemical pathways that govern the molecular assembly processes resulting in formation of extended graphene structures. Whereas most bottom-up processes start from already rather well-developed polyaryl or oligophenylene precursors (via ex situ synthetic chemistry), we uncover the in situ self-formation of the precursors as integral part of the reaction scheme, starting from the decomposition of a simple hydrocarbon. A scheme derived from our joint experimental and theoretical investigations of the surface-catalyzed thermally activated reaction that transforms ethene adsorbed on Rh(111) into graphene is shown in Figure 1. This chemical transformation results from annealing the initial C2H4/Rh(111) system at successively higher temperatures; alternatively, it evolves in the course of in situ experiments conducted on a metal surface dosed with ethene at elevated temperatures that match the annealing temperatures used. The scheme shows the key reaction intermediates and their appearance temperatures for the entire ethene to graphene transformation with the corresponding experimental and theoretical findings discussed and illustrated in Figures 2, 3, and 4 below, as well as in the Supporting Information (SI). The reaction scheme presented in this paper has been validated for varying initial ethene exposures up to 1 monolayer. Herein, we present the results pertaining to an initial feedstock exposure of near saturation coverage.

For ease of navigation, we overview here the reaction steps and intermediates/precursors revealed in this study: (i) Adsorption of ethene and its subsequent partial dehydrogenation yielding ethylidyne (≡C=CH2; Figure 2, 300 K), resulting in the formation of a C4 metallocone coordinated to surface metal atoms (but-2-en-1,4-diylidene, M > C=C≡C−C < M; Figure 2, 370 K). (ii) Formation of benzene-like C6 intermediates (Figure 2, 370 K) and subsequent growth, through successive C4 additions via surface-assisted Diels–Alder condensation, of kinked linear 1D-PAH chains, consisting of ≤3 nm long 12-benzene-ring segments with adjacent segments joined by a 120° kink with respect to each other (Figure 1, rightmost configuration in the top row; Figure 2, 470 K). (iii) Chain-end restructuring with subsequent detachment and dehydrogenation of a coronene-like terminal unit (Figure 1, rightmost configuration in the bottom row; Figure 2, 570 K; see microscopic detachment mechanism in Figure 4) that results in the emergence of a unique size-selective carbon species (Figure 1, middle of bottom row; Figure 2, 670 K)), identified as the key self-evolved precursor, whose diffusional assembly culminates in the growth of an extended graphene adlayer (Figure 2, 770−970 K). To complement Figure 2b, a collection of STM images acquired at various magnifications for all annealing stages described above is reported in SI4 with the view to provide in-depth insights on the various temperature-dependent intermediates.

Activating and unveiling the above sequence of reaction steps was made possible through the implementation of a carefully programmed sequence of temperature windows, which enabled access to the individual reaction stages, one at a time, thus deconvoluting the complex reaction scheme. In contrast, when dosing the Rh(111) surface with ethene immediately at high temperatures (e.g., T ≥ 520 K), a number of dehydrogenation steps are activated simultaneously and as a result a thin amorphous carbon film forms; see a similar result for a Ru(0001) surface in ref 25.

Ethene Deposition and Early Reaction Stages. The widespread of activation energies associated with the catalytic dehydrogenation processes occurring at different stages of the ethene to graphene transformation is reflected in the broad temperature range of the hydrogen Thermal Desorption Spectroscopy (TDS), measured after saturating the surface with ethene at 330 K (Figure 2a). The first TDS regime,

![Figure 1. Schematic of the pathway describing the evolution of adsorbed ethene (top left) to graphene (bottom left). The sequence of intermediates identified in this study and their respective appearance temperatures are indicated. At the final stages, following heating to T > 570 K (see bottom row), coronene-like clusters form. The diffusional coalescence of these precursors results in graphene growth at T > 700 K.](image-url)
occurring for $T \leq 450$ K, consists mainly of a prominent peak at $T \approx 400$ K, whereas the second one, at $450 \leq T \leq 700$ K, is rather featureless. Early studies revealed that ethene adsorption at 300 K on Rh(111) proceeds initially via the formation of ethylidyne, accounting for 25% hydrogen loss (one out of every four H atoms of ethene). The TDS in Figure 2a hence monitors the loss of the three remaining H atoms in proportions of 50% and 25% for the low- and higher-temperature regimes, respectively.

Following deposition of ethene on Rh(111) at room temperature, the ethylidyne species are readily observable in our STM images as protrusions with an apparent height of 80 pm (Figure 2b). As expected, these form an ordered $(2 \times 2)$ structure of ethylidyne seen as 80 pm protrusions (see also HREELS data in agreement with earlier studies). At 370 K, disordered species, consisting of H atoms (apparent height, 40 pm) and covalently linked intermediates: $C_4$ (140 pm) and $C_6$ (100 pm, see SI5) (0.11 V; 0.11 nA). At 470 K, 1D-PAHs of various lengths, made of 12-benzene-ring linear segments (1.11 V; 0.11 nA). At 520 K, 1D-PAHs with increased number of kinks, leading to an apparent broadening and reduced height (1.07 V; 0.11 nA). At 570 K, 1D-PAHs shortened by detachment of dome-shaped coronene-type clusters (0.60 V; 0.12 nA). At 670 K, almost complete conversion of the PAH chains into size-selective carbon precursors that diffuse and coalesce into graphene (0.45 V; 0.12 nA). At 770 K, bimodal distribution of carbon precursors and small, defect-rich graphene flakes (1.00 V; 0.11 nA). At 970 K, extended graphene islands of high structural quality (1.00 V; 0.11 nA). STM images from 300 to 670 K are $22 \times 22$ nm$^2$ and are displayed with the same topographic contrast (identical $z$-scale). STM images for 770 and 970 K are $35 \times 35$ nm$^2$ and are displayed with the same topographic contrast (but different to previous ones). (c) The relaxed structure of a doubly kinked 19 benzene-ring long 1D-PAH, obtained from first-principle simulations, underlying the simulated STM micrograph shown in (d) for the 470 K panel in (d). (d) Simulated STM images of the relevant intermediates at each temperature based on the DFT-optimized geometries presented in SI7: $CH_3$ at 300 K, $C_4$ and $C_6$ species at 370 K, 1D-PAH at 470 K, and coronene-like precursors at 670 K.

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previously reported from which we infer that upon adsorption at 300 K the initial C—H bond cleavage induces a C—C bond order change to 1, and results in an upright adsorption geometry (see Figure 1, transformation at 300 K). This assignment is further confirmed by the excellent agreement between the corresponding experimental HREEL spectrum (300 K in Figure 3a) and the simulated one, computed from the DFT-optimized geometry of ethylidyne adsorbed on Rh(111) surface (Figure 3b, SI3, and SI7).

Thermal annealing of the ethylidyne/Rh(111) system to 370 K results in disordered arrangement of protrusions exhibiting various lateral dimensions and heights (Figure 2b). Close inspection reveals essentially the presence of three distinct species classified by their apparent height. The smaller and most abundant protrusions (40 pm) and the intermediate height protrusions (100 pm) are assigned to adsorbed hydrogen atoms and to C6 rings, respectively, on the basis of independent STM measurements, whereby molecular hydrogen and benzene are exposed to the Rh surface at room temperature (see SI5). The third species, which has the largest apparent height (140 pm) is identified as a C4 unit (see below).

The sequential transformation of C2 species (ethylidyne) into C6 intermediates points to a surface-assisted 2 + 2 + 2 cyclization mechanism (cyclotrimerization). Indeed, Rh (as well as other transition metals) is known to catalyze the formation of benzene. The C6 species is found to coexist with a C4 one (see Figure 2b (370 K) and refs 30 and 31); C4 is a metallocycle formed in 2 + 2 addition of ethylidyne, known as a reactive intermediate in organometallic catalysis, for example, cyclotrimerization of ethyne.32 We note here that the H2 TDS (Figure 2a) indicates a C/H ratio near 1:1 at 370 K. It is likely, however, that the intermediates are not perfectly stoichiometric because saturated benzene is known to be unreactive toward the formation of further surface-bound intermediates.

Experimental evidence for the involvement of C6 and C4 species in the temperature range of 370−420 K is confirmed by HREELS. The full set of observed vibrational modes in the HREEL spectrum for 420 K (Figure 3a) is only appropriately described in our DFT-based simulations (Figure 3b) when...
considering the coexistence of both C4 and C6 species adsorbed on the Rh(111) surface. Of particular significance is the extended vibrational structure observed upon annealing to 420 K between 250 and 600 cm⁻¹ (gray-shaded areas in Figures 3a and 3b) indicative of Rh–C stretch frequencies of small planar cyclic or inclined metallocyclic hydrocarbons that develop during the first growth steps. It has to be noted that this characteristic vibrational structure between 250 and 600 cm⁻¹ can only be reproduced by superposition of the corresponding C4 and C6 calculated bands. Hence, the judicious combination of microscopy, spectroscopy, and computational methods concur to support a cyclo-additive mechanism for the first stage of C2 coupling on Rh(111), yielding C4 and C6 species.

**Formation of Segmented 1D Polyaromatic Hydrocarbon Narrow Ribbons.** Following a further annealing of the ethene/Rh(111) system to 470 K, STM topographies show a transformation into chains of linear segments oriented along the three main substrate directions [110], [011], and [101] (Figure 2b) with an apparent height of 100 pm. We identify

**Figure 4.** Size-selective PAH fragmentation into coronene-like clusters. (a–e) Selected STM images following a 570 K annealing of the ethene/Rh(111) system. (a) 2D extension of a PAH chain-end, forming a coronene-type termination; (b) Chain-end dehydrogenation yielding a coronene-like carbon cluster termi

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these structures as one-dimensional linear polyaromatic hydrocarbons (1D-PAHs, see Figure 1, transformation at 470 K) based on several observations (details in SI5): (1) The chains exhibit a longitudinal corrugation periodicity of 2.5 Å, in agreement with edge-connected C₆ rings. (2) The lateral width of the chains matches that of benzene molecules and is smaller than that of coronene molecules. In addition, the 1D-PAHs, benzene, and coronene have all similar apparent electronic heights of 100 pm. (3) STM images of tetracene molecules (as a model for 1D-PAHs with a fixed length of four carbon rings) on Rh(111) show a remarkable resemblance to the chains. (4) Taking into account the hydrogen balance from TDS (Figure 2a) and the fact that no carbon is lost to the gas phase after prolonged growth at 450 K, the evolved structure is that of the 1D-PAH shown in Figure 2c (see SI7) with a 12 ≤ 4 Å²). The highest carbon atoms with respect to the underlying metal surface are found at the edges and in the middle of the linear segment, reflecting a maximal incommensurability (see SI7). The charge transfer from the metal surface to the carbon atoms and the nature of the electronic hybridization are also weakly modulated in a manner that is correlated with the height variations. The bonding is of sp² character where the chain is high above the surface, whereas it gains some sp³ character when the carbon atoms are closer to the metal surface, reflecting stronger binding with a consequent rehybridization (sp³ → sp²).

With the identification of ethyldyne at 300 K and 1D-PAH intermediates at 450 K, the first H₂ desorption regime (see Figure 2a) is associated with the dehydrogenation of ethene and condensation of aliphatic hydrocarbons into 1D-PAH. The second desorption regime at elevated temperatures (450−700 K) is governed by the thermal evolution of 1D-PAHs into extended graphene following a 1D to 2D structural transition. After prolonged growth at 450 K, the evolved structure is that of mutually avoiding 1D-PAHs (see SI6), indicating that further condensation into 2D-PAH intermediates is inhibited by steric repulsion, and requiring higher temperatures for continuation of the process. The existence of two different growth regimes and their respective interpretation is confirmed by our HREELS investigations. Records of the integrated signals for the various C−H vibrational frequencies from HREEL spectra, acquired after successive annealings, summarize the spectroscopic findings, as shown in Figure 3c. From this, we conclude that at 300 K the predominant frequencies relate solely to aliphatic carbons, as expected. Their corresponding intensities are seen to decrease with temperature, concomitant to the emergence of aromatic carbon fingerprints. Above 420 K, the aliphatic contributions vanish while the aromatic ones are maximized. With further annealing, the aromatic intensities decrease almost monotonically over a wide temperature range (up to 700 K), as a consequence of the slow, featureless dehydrogenation (C−H cleavage) observed in the second regime of the TDS data. There remains, however, the question as to how the 1D to 2D structural transition proceeds at high temperatures.

**Kinking and 1D → 2D Dimensionality Crossover.**

Further kinking of the 1D-PAHs and their consequent broader appearance occurs after annealing at 520 K (Figure 2b) with the resulting zigzags aligned with the three main symmetry directions of the substrate. While the chain density remains constant upon annealing from 470 to 520 K (ruling out coalescence), an apparent shortening of the 1D-PAHs by 25% without change in the apparent height is consistent with the introduction of kinks. This intramolecular reshaping is a consequence of simple thermodynamic arguments (Clar’s rule) and can occur by rearrangement of a single, weak, edge-connecting C−C bond while keeping the molecular backbone intact (see SI8). Annealing to 570 K drives a further shortening of the 1D-PAHs (Figure 2b). In parallel, several chain ends become clearly broader, indicative of the onset of structural reshaping into the second dimension. The increased propensity of kinking can bring the chain ends to curl up (see Figure 1, transformations at 520 and 570 K), leading to close proximity of adjoining C−H bonds and allowing for C−C bond formation at these temperatures; recall the activated formation of graphene nanoribbon on Au substrates at around 670 K. Inspection of Figure 2b (at 520 K, and particularly 570 K) reveals that the broader PAH chain ends take two distinct...
appearances: one in which the apparent height remains unchanged with respect to the chain segments (100 pm) and the other exhibiting an apparent height increase to 150 pm. In addition, these protruding chain ends have a lateral extension of 1 nm, a width reminiscent of that of the size-selective carbon precursors first identified on Rh(111) \(^{21}\) and later on Ru(0001).\(^{36}\) Examination of the STM records at 570 K (Figure 2b) confirms that the density of PAH chains decreases at the expense of smaller and detached particles of quasi-circular shape, exhibiting a diameter of 1 nm and a height of up to 150 pm, a carbon precursor species with a varying hydrogen content (see below). We note here the ambiguous structural assignment of the carbon precursors: following the work of ref \(^{21}\), several reports have confounded the identity of the size-selective carbon precursors on the Rh(111) on grounds of DFT calculations,\(^{22,23}\) and to-date no experimental validation has yet emerged. Our present microscopy data precludes further identification. Nevertheless, a lateral extension of 1 nm suggests a polyaromatic precursor of coronene-type dimensions.

On the basis of these microscopic observations, the following mechanism for the formation of graphene emerges: under appropriate thermal conditions, the 1D-PAH chains-ends undergo 2D-restructuring by a combination of kink formation and curling-up. Coronene-type terminations become apparent in the STM topographs (Figure 4a). Continued 2D-PAH growth beyond this critical size entails detachment of coronene-type species with varying hydrogen content. To substantiate the detachment process, we performed DFT calculations assuming a coronene structure for the carbon precursor (mindful of its contentious structural identification). The results of our DFT calculations reveal that the detachment process may be induced by hydrogen released during the 2D-restructuring. Concomitantly, peripheral dehydrogenation of the detaching species leads to carbon–metal hybridization with the underlying Rh substrate, forcing the detaching units to adopt a dome-shaped configuration (hence the increased apparent height, Figure 4b).\(^{41,45}\) Dehydrogenation may be completed before or after the detachment of the coronene-like unit, resulting in the STM images in Figure 4 c.d. These detached, size-selective entities on the Rh(111) surface that act as precursors to the further growth of graphene\(^{21}\) can be identified by constant-height STM that seem to possibly resolve \(C_6\) rings (Figure 4e). The dehydrogenation processes associated with the above restructuring and disjoining processes of the 1D-PAHs lead to gentle release of \(H_2\) to the gas phase, correlating with the TDS findings in the range of 450 to 700 K (Figure 2a).

The slow dehydrogenation process and the concomitant increase in lateral extension of the aromatic backbone are clearly evidenced in the HREELS spectra (Figure 3a) for temperatures above 450 K. Indeed, the intensity of the aromatic C–H stretch peak at 3000 cm\(^{-1}\) slowly decreases with temperature and vanishes at 700 K (as expected from our hydrogen TDS data, Figure 2a). In a concerted manner, one also observes the decrease in intensity of the 750 cm\(^{-1}\) peak, which arises from C–H out-of-plane wagging and C–C-breathing modes (aromatic). While we can rationally infer that the C–H wag component must also vanish by 700 K with the loss of hydrogen, the C–C breathing component must remain present at even higher temperatures, when graphene flakes emerge. In addition, the C–C breathing modes have mainly dipolar contributions parallel to the surface, and HREELS (operated in specular geometry) is sensitive only to perpendicular dipolar contributions. Hence, the growth of graphene into ever larger flakes is accompanied by a decrease in HREELS intensity for the C–C breathing modes and a shift to lower frequency. Interestingly, the C–C breathing modes are also seen to broaden with temperature. This is to be expected due to it averaging over the many Rh–C bonding configurations imparted by the lattice mismatch between graphene and the Rh substrate, as discussed previously.

**Microscopic Formation Mechanism of the Self-Evolved Precursors and Graphene Growth.** The pathway of the coronene-like detachment process, obtained from first-principles simulations, is displayed in Figure 4f (see SI9). The starting configuration (0) corresponds to a curved-up (surface-lifted) PAH chain-end configuration. The detachment of this terminal unit requires activation (weakening) of the C–C bonds bridging the PAH and the attached coronene through their partial hydrogenation that thus acquire sp\(^3\) character. This process is akin to a hydrogenolysis-type mechanism defined as a chemical reaction whereby a carbon–carbon undergoes “lysis” by hydrogen.\(^{38}\) The detachment process of the carbon cluster from the thickened PAH chain-ends was found to occur at about 570 K (see Figure 2b (570 K)). At this temperature, hydrogen evolution on the surface is still observed, as evidenced by the measured hydrogen desorption spectrum displayed in Figure 2a; in fact, the hydrogen desorption signal vanishes only at temperatures well above that of the detachment process, that is, above 700 K (see Figure 2a). Therefore, it is most reasonable to expect that hydrogen atoms in sufficient abundance are available for capture and local hydrogenation/activation of the bonds linking the carbon cluster to the polyaromatic hydrocarbon chain as conjectured by the microscopic detachment mechanism simulated and described in Figure 4f.

Thereafter, the detachment mechanism depicted in Figure 4f involves successive breaking of these C–C bonds. The evolution of the detachment process entails pivotal in-plane rotations of the detachment 24-carbon-atom cluster (see Figure 4f, and Figures S26 and S27 where alternative detachment pathways are shown with overall characteristics similar to the pathway displayed in Figure 4f). Such pivotal cluster rotations have been discovered earlier\(^{39}\) to underlie cluster diffusional (and superdiffusional, that is, Lévy flights) motion\(^{40–42}\) on incommensurate solid surfaces (as is the case for the interface between the size-selective cluster and the Rh(111) surface mesh). After detachment, the PAH chain residues are likely to diffuse, merge, and perpetuate the production of size-selective clusters. Increased annealing leads to thermal diffusion (involving the aforementioned in-plane cluster rotations) of the precursors,\(^{21}\) which coalesce into extended graphene.

The microscopic pathways described here operate over a wide temperature range and allow for a slow release of the remaining 25% hydrogen to the gas phase, compatible with our TDS data (Figure 2a). The surface-catalyzed thermally activated chemical transformation of adsorbed ethene into graphene (Figure 1), which we found to proceed via coupling reactions that lead to formation of segmented 1D-PAH chains with a subsequent dimensionality crossover (1D → 2D) and restructuring, resulting in eventual detachment of self-evolved precursors (Figure 4), culminates at 770 to 990 K with diffusional self-assembly and condensation of the precursors to form extended graphene monolayer patches with a high degree of structural perfection, as displayed in Figure 2b. We note here that the maximum graphene coverage attainable under our experimental conditions is limited by the number of ethylene
molecules deposited at 300 K to 21% of the total Rh surface, because no carbon atoms are lost to the gas phase nor to the bulk of the Rh support during our annealing experiments.

**CONCLUSIONS**

When employing an appropriate programmed sequence of temperature windows, thereby regulating and controlling the subtle balance between kinetics and thermodynamics, the surface-catalyzed chemical transformation of adsorbed ethene into graphene (Figure 1), studied here on single crystal Rh(111) under ultrahigh vacuum conditions, can be directed toward the controlled formation of polyaryl intermediates (and products) with precise morphologies: 1D-PAH chains, size-selective 2D-PAH clusters, and extended single-layer graphene. The observed dimensionality crossover (1D → 2D), resulting from polyaryl dynamical restructuring, ushers the emergence of a self-evolved precursor resulting from detachment of a coronene-type end-chain-unit (Figure 4) and culminates with rate-limiting diffusional self-assembly and condensation into extended graphene monolayer patches with a high degree of structural perfection (Figure 2b). These results suggest future explorations of the paradigm of dynamically self-evolved precursors in surface-catalyzed reactions through judicial selection of substrate and adsorbate materials, surface processing, and systematic programmed variation of the ambient reaction conditions.

**EXPERIMENTAL METHODS**

Methodology for the Low-Temperature STM Measurements. Experiments were performed in an ultrahigh vacuum (UHV) surface analysis system with a base pressure below 1 × 10⁻¹⁰ mbar, consisting of a preparation chamber allowing for standard sample preparation and characterization by Auger electron spectroscopy (AES), and a microscope chamber housing a CreaTec low-temperature STM. The Rh(111) crystal was cleaned by repeated cycles of Ar⁺ sputtering and annealing in oxygen (3 × 10⁻⁷ mbar) at 1000 K and finalized by flash-annealing in vacuum at 1000 K. STM acquisition was performed at liquid nitrogen (LN₂) temperature using the constant-current mode and homemade W tips. The cleanliness of the surface was monitored by STM and AES. Research-grade ethene, C₂H₄ (99.95% purity), was exposed to the Rh surface up to saturation levels (approximately 5 mbar × 6 mbar/s) at room temperature, the heating current was increased slowly, by 0.1 mA (∼0.6 mW) at fixed time steps of 10 s, leading to temperature increases of ∼0.05 K. The induced drift was corrected via lateral feedback. The cleanliness of the surface was monitored by STM, AES, and QMS. Ethene, C₂H₄ (99.5% purity), was exposed to the Rh surface. First, when dosing at room temperature, exposures of 27 L were applied, after which the substrate was annealed to higher temperatures. Second, when dosing at elevated temperatures, ethene was exposed continuously at approximately 5 × 10⁻⁹ mbar during the STM acquisition.

Methodology for the HREELS Measurements. Within a dedicated experimental system, composed of a preparation chamber with base pressure below 5 × 10⁻¹⁰ mbar and an analysis chamber with base pressure below 1 × 10⁻¹⁰ mbar, HREELS measurements were carried out in the specular direction (θi = θf = 45°) with a primary beam energy of 4 eV and a typical elastic peak resolution of about 50 cm⁻¹ (6.2 meV fwhm). A maximum likelihood-based resolution enhancement method was used to recover the spectra from the instrumental broadening, leading to an improved resolution of about 40 cm⁻¹. Spectra were normalized to the intensity of the elastic peak. Ethene (C₂H₄, 99.995% purity) was dosed at room temperature (300 K) for an exposure of 2, 5, and 10 L, enough to ensure saturation coverage. The substrate was then flash-annealed to progressively higher temperatures and HREEL spectra collected after allowing the sample to cool to room temperature. Similar behavior is observed following each of the exposures; the 5 L experiment is discussed as a typical example of the spectral evolution seen upon sequential flash annealing.

**COMPUTATIONAL DETAILS**

Density Functional Calculations. Calculations were performed using the VASP-DFT package with a plane-wave basis (kinetic-energy cutoff of 400 eV), PAW pseudopotentials, and the PBE generalized gradient approximation (GGA) for the exchange-correlation potential. Because the unit cell used here is rather large, we have used in most of our calculations a single k-point sampling (Γ-point) of the surface Brillouin zone (SBZ); we have checked that the results remain very similar by employing a (3 × 3 × 1) SBZ sampling. In optimization of the various structures, convergence was achieved when the calculated forces were smaller than 0.005 eV/Å. In the simulations, the Rh(111) substrate surface consisted of three layers with the optimized Rh lattice parameter a = 2.718 Å, agreeing with the experimental values (a = 2.69 Å). In structural relaxations, the bottom layer of the substrate slab was held fixed. In simulations of the configuration of long linear 1D-PAH chain with two kinks (see Figure 2b), the supercell had a (18 × 10) lateral periodicity of the three-layer Rh(111) slab, and a vacuum region that was taken large enough to ensure no interaction between periodic replicas; for the bare Rh(111) slab, the vacuum region was taken as 20.7 Å. In simulations of configuration, the 24 carbon atom cluster (or coronene) configuration with a 1D-PAH tail (see Figure 4), the supercell had an (10 × 8) lateral periodicity of the three-layer Rh(111) slab, and in simulation of the C₆H₆ and C₄H₄ intermediates (see Figure 2d) the supercell had an (12 × 6) lateral periodicity.

Simulations of Reaction Pathways. In simulations of the reaction pathways (in particular the detachment of the 24-carbon atom cluster from the PAH tail (see Figure 4f)), we...
have used steered first-principles simulations, where a certain reaction coordinate (or reaction coordinates, which may consist of certain interatomic distances, angles, and/or combinations thereof) is varied in a controlled way, whereas all other nuclear degrees of freedom are allowed to fully relax after each increment of the reaction coordinate. The relaxation process is performed by using an algorithm based on a conjugated gradient method with the total energy evaluated at each step of the relaxation process quantum mechanically using the density functional theory (DFT) described above. In this way, the activation energy barriers are determined; once the top of an activation barrier is reached, the steering of the reaction is stopped and the system’s nuclear coordinates are allowed to evolve freely on the concomitantly first-principles-calculated potential energy surface until the nearest potential energy minimum is reached. If the minimum reached corresponds to the desired reaction products, the simulation is stopped. On the other hand, if the potential energy minimum reached is a local one, steering of the reaction resumes along a judiciously chosen reaction coordinate, and the next activation barrier is determined. Often several alternative reaction coordinates are employed and the one yielding the lowest activation energy barrier is selected. These calculations yield results that are the same as, or close to, those obtained by other methods (e.g., the nudged elastic band and variants thereof; see discussion in ref 47).

In the present study, we restricted ourselves to consideration of the enthalpy contribution to the reaction profile. Free-energy reaction profiles (that is, including entropic effects) can be obtained through a sequence of canonical (constant-temperature) first-principle (Born–Oppenheimer) molecular dynamics (FPBOMD) simulations while constraining the reaction coordinate to prescribed values (see Results and Discussion above), and calculation of the reversible work done in taking the system along the reaction path between the reactant and product states. However, the exceedingly large computational demands of such first-principle calculations limits the methods to systems much smaller than the one investigated here. Furthermore, because the reaction that we discuss here involves breaking of a rather strong bond (the C–C bond) it is reasonable to assume that potential energy (enthalpy) effects will dominate the reaction profile and that the inclusion of entropic ones (mainly vibrational entropy of the adsorbed complex) will not modify our results in an essential way.

van der Waals (vdW) Interactions. Inclusion of vdW interactions 48 resulted in changes in the total energies of the systems being investigated. However, structural relaxations with and without the inclusion of vdW interactions yielded essentially the same optimal structures. Moreover, the energy barriers determined from our simulations show very small dependence on the vdW interactions. See further details in the Supporting Information.

Calculations of STM Images. STM images were simulated by using the Tersoff–Hamann approximation 49 in which the tunneling current is considered to be proportional to the integrated local density of states within a given energy window determined by the applied bias on the sample; we have used an energy window of 1 eV below the Fermi energy, that is, $E_F - 1$ eV $< E < E_F$ corresponding to a bias voltage of $-1$ V. These STM images are topographic images (that is, the color variations correspond to different heights) because the images correspond to those recorded in constant current scans.

Simulations of HREEL Spectra. C$_2$H$_3$ (ethyldiene), C$_6$H$_4$ and C$_{10}$H$_{16}$ isolated species (all on 4 × 4 cell sizes) were optimized using a higher convergence criterion to obtain the simulated spectra. In particular, the convergence criterion for the electronic self-consistent cycle was fixed at $10^{-6}$ eV/Å and the forces on all ions were required to be smaller than 0.015 eV/Å. The Born effective charges (BEC) matrix for each system were obtained employing density functional perturbation theory inherent in VASP. Intensities were calculated using a similar procedure as found in reference 50 where the BEC tensor was projected to the perpendicular direction with respect to the surface. The intensity-weighted harmonic eigenfrequencies were broadened by applying a Gaussian function with a fwhm of 40 cm$^{-1}$, corresponding to the resolution of the experimental HREELS setup. In addition, the spectra were corrected for the spectrometer transfer function, as described in ref 51.

ASSOCIATED CONTENT

Supporting Information

The Supporting Information is available free of charge on the ACS Publications website at DOI: 10.1021/acs.jpcc.7b01999.

Methodology of the experiments; methodology of the DFT calculations; HREEL spectroscopy studies; high-resolution STM images of the ethene/Rh(111) system after deposition and after annealing to successively higher temperatures; the initial stages of ethene condensation on Rh(111) as observed by STM; time-resolved STM data on the 1D-PAH chain growth at 455 K; the intermediates identified during ethene condensation on Rh(111) as modeled by DFT calculations; 1D-PAH intramolecular reshaping; DFT calculation results on the formation of coronene-like clusters—mechanistic details and alternative pathways (PDF)

STM time series of the ethene decomposition at 455 K, dynamics of 1D-PAH growth (AVI)

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B.W. and M.K. contributed equally. R.S., F.E., B.W., U.H., and U.L. conceived and designed the study. B.W., M.K., C.J.B., M.J.T., and M.C. prepared specimens, took STM images, and performed image processing. F.G. and C.J.B. prepared spectra. F.G. and C.J.B. prepared simulated spectra. M.J.T., and M.C. prepared specimens, took STM images, and conducted image processing. F.G. and C.J.B. prepared specimens, took HREELS spectra and conducted spectroscopic analysis. B.Y. and U.L. conducted and analyzed DFT calculations. J.A.G.T. and H.F. computed the HREEL spectra from the DFT calculations. All of the authors discussed the

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experimental results. All of the authors contributed to the interpretation of the data. B.W. and M.K. contributed to the initial elaboration of the manuscript; R.S., U.L., and F.E. wrote the manuscript.

Notes

The authors declare no competing financial interest.

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