Structural, Electronic, and Vibrational Properties of a Two-Dimensional Graphdiyne-like Carbon Nanonetwork Synthesized on Au(111): Implications for the Engineering of sp-sp² Carbon Nanostructures

Andi Rabia, Francesco Tumino, Alberto Milani, Valeria Russo, Andrea Li Bassi, Nicolò Bassi, Andrea Lucotti, Simona Achilli, Guido Fratesi, Nicola Manini, Giovanni Onida, Qiang Sun, Wei Xu, and Carlo S. Casari*

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ABSTRACT: Graphdiyne, atomically thin two-dimensional (2D) carbon nanostructure based on sp-sp² hybridization is an appealing system potentially showing outstanding mechanical and optoelectronic properties. Surface-catalyzed coupling of halogenated sp-carbon-based molecular precursors represents a promising bottom-up strategy to fabricate extended 2D carbon systems with engineered structure on metallic substrates. Here, we investigate the atomic-scale structure and electronic and vibrational properties of an extended graphdiyne-like sp-sp² carbon nanonetwork grown on Au(111) by means of the on-surface synthesis. The formation of such a 2D nanonetwork at its different stages as a function of the annealing temperature after the deposition is monitored by scanning tunneling microscopy (STM), Raman spectroscopy, and combined with density functional theory (DFT) calculations. High-resolution STM imaging and the high sensitivity of Raman spectroscopy to the bond nature provide a unique strategy to unravel the atomic-scale properties of sp-sp² carbon nanostructures. We show that hybridization between the 2D carbon nanonetwork and the underlying substrate states strongly affects its electronic and vibrational properties, modifying substantially the density of states and the Raman spectrum compared to the free standing system. This opens the way to the modulation of the electronic properties with significant prospects in future applications as active nanomaterials for catalysis, photoconversion, and carbon-based nanoelectronics.

KEYWORDS: graphdiyne, 2D carbon nanonetwork, sp-sp² carbon, 2D materials, scanning tunneling microscopy, Raman spectroscopy, density functional theory

INTRODUCTION

Two-dimensional (2D) carbon systems with mixed sp-sp² hybridization, i.e., graphene and graphdiyne, aroused great interest in the scientific community over the past 30 years as novel 2D carbon structures,¹⁻³ paving the way for the ultimate goal of fabricating sp-hybridized carbon fragments, whose structural, optical, and transport properties were deeply explored.⁴⁻⁶ These systems can form a variety of 2D crystals with a different structure, sp/sp² ratio, density, and porosity and have been predicted to possess peculiar electronic properties, such as multiple Dirac cones in graphene.⁷

Synthetic chemistry attempts to develop sp-sp² carbon networks have been carried out by developing efficient methodologies based on a monomer-to-polymer approach. In this respect, one of the most used protocols has been the oxidative acetylenic coupling of macrocyclic carbon-rich precursors.⁸ However, direct coupling of the monomers reported the cross-linking of the chains,⁹ which precludes the development of well-defined extended structures. To tackle this issue, Haley¹⁰ proposed the isolation of the reactive acetylene moieties and the assembly of the structure via an intramolecular cyclization approach, obtained through sequential Sonogashira cross-coupling reactions.¹¹ A graphdiyne film has been for the first time produced by Li et al.¹² on a copper substrate through cross-coupling reaction, then several works reported the fabrication in the form of flakes or powder and tested for potential applications in the field of photocatalysis and nanoelectronics.¹³⁻¹⁷

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The investigation of the structure of these systems is of fundamental importance since for a long time graphdiyne has been considered elusive and the poor stability of the sp carbon phase is still an important issue opposing the realization of extended and ordered 2D structures. In this context, Raman spectroscopy stands as the election technique to investigate the structure of carbon-based materials, proving to get access to the presence of sp carbon and to the structural properties and local bond order. The characteristic Raman signal of sp carbon and its structure-related behavior allows a detailed investigation of sp and sp-sp² carbon systems.²⁸,¹⁹

Despite the significant advancement in the synthesis and the investigation of these materials, the fabrication of extended graphdiyne monolayers and the imaging of their atomic-scale structure is still a challenge. Scanning tunneling microscopy and spectroscopy (STM/STS) in combination with theoretical modeling has the potential to unravel the atomic-scale structure and to provide a deep insight into the surface electronic properties. The atomic-scale imaging by means of scanning probe techniques usually requires material growth in ultra-high vacuum (UHV) conditions on atomically flat surfaces and in situ characterization. In this framework, the rapidly growing on-surface synthesis technique has demonstrated tremendous potential in the high-precision bottom-up construction of low-dimensional carbon nanostructures and in the atomic-scale imaging and characterization by surface science techniques.²⁰−⁲⁵ Its great advantage relies on fostering selective chemical reactions between molecular precursors on metal surfaces under ultra-high vacuum (UHV) conditions.²⁶,²⁷ To this aim, molecular precursors are designed to metal surfaces under ultra-high vacuum (UHV) conditions, through the exposition to an organic molecular evaporator (OME) and its structure-related behavior allows a detailed investigation of sp and sp-sp² carbon systems.²⁸,¹⁹

![Figure 1](https://dx.doi.org/10.1021/acsnano.0c02665)

**Figure 1.** (a) Scheme showing the deposition of 1,3,5-tris(bromoethyl)benzene (tBEP) by an organic molecular evaporator (OME) and reporting the different steps of the on-surface synthesis process on Au(111); adsorption (1°), dehalogenation process (2°), and coupling through Au adatom (3°). (b) Large-scale STM image showing the sp-sp² carbon nanonetwork formed after the deposition of the tBEP molecular precursor on Au(111) (1.3 V, 0.5 nA).

The fabrication of sp² carbon systems is achieved through on-surface Ullman coupling reaction of aryl halides, an efficient strategy for the formation of sp carbon nanostructures is represented by on-surface dehalogenative/dehydrogenative homocoupling reaction of precursors functionalized with alkynyl halides.²⁷−⁴³ However, accessing the nature of molecular bonds and hybridization state represents a key factor to comprehensively understand the properties of the novel sp-sp² carbon structures. In a previous work, we have adopted Raman spectroscopy for the identification of sp hybridization and investigation of on-surface formation mechanism of sp-sp² carbon atomic wires on Au(111) substrate.⁴⁴

Herein, we report on the nanoscale structure and electronic and vibrational properties of a carbon monolayer nanonetwork whose structure resembles γ- or α-graphdiyne, while differing by the presence of aromatic rings with threefold hydrogen terminated bonds. This 2D carbon nanonetwork, characterized by sp-sp² hybridization, was grown on Au(111) under UHV conditions, through the exposition to an organic molecular precursor with three alkynyl bromide groups.⁴⁵ By a combination of high-resolution STM imaging, Raman spectroscopy, and DFT simulations, we unveil the structure at different stages of the formation, i.e., from metal organic nanostructure comprising Au adatoms to the pure sp-sp⁴ carbon nanonetwork obtained after the release of Au atoms by thermal annealing. Insight on the 2D surface electronic structure is provided by first principle calculations. Raman spectroscopy and DFT calculations reveal the C–C vibrational modes of this one-atom thick 2D sp-sp² carbon nanonetwork on Au(111). The present work provides an example of the great potentiality of Raman spectroscopy in studying the atomic-scale structure of novel carbon nanostructures based on sp-sp² hybridization, e.g., graphyne and graphdiyne, complementing the high-resolution STM imaging. The interaction between the 2D carbon nanonetwork and the underlying Au(111) surface involves a charge transfer and has a fundamental role in modifying both the electronic and vibrational properties of this system with respect to what expected for the freestanding counterpart. The availability of a 2D carbon-based semiconductor on a metal may open new opportunities in the field of catalysis with novel non precious
The graphdiyne-like 2D sp-sp\textsuperscript{2} carbon monolayer is obtained on Au(111) after UHV deposition at room temperature of the precursor molecules shown in Figure 1a. The on-surface synthesis mechanism can be easily sketched in three steps\textsuperscript{40} (see Figure 1a), which take place simultaneously: (i) adsorption of the molecules on the Au(111) surface, (ii) cleavage of C−Br bonds followed by the diffusion of radicals on the 2D template, and (iii) coupling of the sp carbon chains through gold adatoms. Following this scheme, after deposition, the molecules self-assemble and form an organometallic 2D nanonetwork (i.e., metalated system) based on sp-sp\textsuperscript{2} carbon catalyzed by the Au(111) surface. The substrate is subsequently annealed to remove the Au atoms and promote the C−C homocoupling reaction\textsuperscript{5} to form an all-carbon nanonetwork. The obtained nanostructure has three sp carbon diaacetylenic chains connecting sp\textsuperscript{2} six-membered aromatic rings in a threefold organization instead of the sixfold structure of \(\gamma\)-graphdiyne (the remaining threefold dangling bonds are saturated by H atoms). The resulting structure shows a large-scale hexagonal organization very similar to the \(\alpha\)-graphdiyne, except for the presence of sp\textsuperscript{2} aromatic rings in place of single C atoms (see the Supporting Information).

In the following sections, we present the results of STM and Raman investigation of this system. DFT calculations of the electronic and vibrational properties allow us to discuss STM images and Raman spectra and to unveil the fundamental role of the Au(111) surface, which affects in a relevant way the density of states and the Raman spectrum.

**Structural and Electronic Properties.** In the STM image in Figure 1b, we report the formation of sp-sp\textsuperscript{2} carbon nanonetwork, which consists in a honeycomb structure extending on the gold surface. Investigating the coverage obtained on the sample, by acquiring large-scale STM images at different regions of the substrate, we observe an ordered sp-sp\textsuperscript{2} carbon nanonetwork over most of the surface. However, there exist fractions of network where the regularity is disrupted by unreacted or possibly degraded molecules deposited as a second layer (Figure 1b). Boundary regions between the sp-sp\textsuperscript{2} nanonetwork and uncovered Au(111) surface show a disordered morphology (Figure 1b), probably because of the random endcapping of highly reactive sp carbon chains with substrate atoms.

We studied the effect of thermal annealing on the morphology and structure by acquiring in situ STM images at room temperature (RT). To this end, we have annealed the sample to temperatures up to 580 K (Figure 2) in UHV. The STM image of the sample annealed at 370 K (Figure 2a) reports a lower coverage than the as-deposited sample (Figure 1b), and this is an indication of the detachment of the less-strong interacting molecules with the substrate. Then, after increasing the annealing temperature in the range of 480−580 K (Figure 2b−c), we observe the formation of an amorphous phase coexisting with the regular 2D nanonetwork. Such a patchy structure might be established by the diffusion and thermal activation of the reactive sp carbon atomic chains originating the onset of the disordered phase, as also revealed by the Raman spectra discussed in the next section. The presence of the amorphous phase further increases when the sample is annealed at 580 K, as shown in Figure 2d, making it difficult to image the surface.

In addition to the morphology, we studied the molecular structure of the sp-sp\textsuperscript{2} carbon nanonetwork by high-resolution STM images and DFT calculations. The sp-sp\textsuperscript{2} carbon nanonetwork has been obtained through on-surface coupling of \(\gamma\)BEP molecules where the substrate catalyzes the reaction by detaching the bromine atoms and substituting them with surface gold adatoms. The crucial role of the surface atoms is pointed out in the STM image acquired on the sample annealed at 370 K (Figure 3a), where, notably, the color contrast enhances the round and bright protrusions associated to the gold adatoms. In Figure 3a, gold adatoms are circled with green color and they form a Kagome-like structure.

A further support to the structural analysis of metalated sp-sp\textsuperscript{2} carbon nanonetwork has been provided by the fast Fourier transform (FFT) (Figure 3b), which reports a hexagonal symmetry and a periodicity of \(\sim 21\) Å. As the annealing temperature increases to 480 K, the hexagons in Figure 3c show protrusions associated to phenyl rings and intermediate gold atoms, while it is difficult to resolve the atomic structure of short sp carbon atomic chains due to the limited resolution of the STM signal. Nevertheless, at 480 K, we observe the disappearance of Au atoms in some sp carbon chains, which are highlighted by white arrows in Figure 3c. Indeed, by extracting the topographic profile (Figure 3d,e) of the metalated and C−C coupled atomic chains (red and blue...
constant corresponding to 20.85 Å (where the theoretical lattice and C−C bond lengths are 136 and 125 pm, respectively.

The theoretical simulation of the STM images, reported in Figure 4, are in agreement with the experiments, showing bright protrusion in correspondence of Au atoms in the chain for the metalated nanonetwork (Figure 4c). The theory predicts also a slightly enhanced STM contrast on the phenyl groups with respect to the attached “arms”.

For the all-carbon case (Figure 4d), the model shows a bright contrast of the atomic carbon chains along the rotated [112] direction and a less intense signal in proximity to one end of the four oblique chains. This effect can be related to the different alignment of the atomic carbon chains with the underlying substrate in the adopted structural model. The resulting modulation of the heights of carbon atoms gives rise to a geometrical effect in the simulation.

In order to investigate the variation of the electronic properties induced by the adsorption of precursors, by the formation of the 2D nanonetwork on the surface, and by the removal of the gold atom upon annealing, we have analyzed the projected density of states (PDOS) integrated in the whole Brillouin zone for both the 2D nanonetworks supported on the Au(111) surface, in comparison with the freestanding system. Figure 5 reports the integrated PDOS of the phenyl groups of the adsorbed polymer are 125 and 199 pm for the C−C triple bond and C−Au bond, respectively. Moreover, the interaction between the Au adatom and the substrate induces a corrugation of the molecular nanonetwork leading to a smaller distance between the Au atom in the chain and the substrate (2.46 Å) compared to the average phenyl−substrate distance (2.70 Å). These data are in agreement with previous theoretical findings for the analogous 1D system.

The C−C coupled system is decoupled from the substrate and displays a negligible out-of-plane and in-plane distortion and a larger distance from the substrate that amount to 2.78 Å. The C−C single and triple bond lengths are 136 and 125 pm, respectively.
the molecule and of the sp chain aligned with the [112̅] direction. The latter is also resolved with respect to the magnetic quantum number (m) of the p orbitals.

In the metalated 2D nanonetwork, the states of the gold atom in the sp chain (green line in Figure 5a) display a relevant superposition with the d band of the Au(111) surface (gray shaded area). The chain–substrate interaction is also the driving force in the charge transfer process that determines an increase of electronic charge (+0.68 e) on the molecule with respect to the freestanding system. The energy separation related to the HOMO–LUMO gap is still visible, even though the hybridization with the states of the substrate induces a metal behavior in the 2D overlayer.

The analysis of the m components of the PDOS allows us to identify the contribution of p states of the sp chain along the different directions. In making the projection, we consider the average of the C atoms of a specific sp chain and align the γ axis along the chain direction. Indeed, considering all the sp chains would result in averaging in-plane. Hence, p_x, p_y, and p_z orbitals are aligned along [110], [112], and [111] directions, respectively.

In particular, the states closer to the Fermi level have p_x symmetry and form out-of-plane π-states. Other π-states (orthogonal to the sp chain axis) now lying parallel to the surface are formed by the p_y orbital; those are more separated in energy, being located below −2 eV and above 2 eV. Noticeably, p_y states are sharper in the PDOS than p_x states due to a smaller hybridization with the Au(111) orbitals. Similar observations could be drawn for a linear sp-sp² system. Conversely, p_x states sit farther from the Fermi level both in the occupied (valence band) and unoccupied (conduction band) part, as expected from their σ character and direction along the chain axis.

Upon removal of the Au adatom, the 2D nanonetwork is decoupled from the substrate and maintains the energy gap around the Fermi level, although reduced with respect to the freestanding nanonetwork (∼1.6 eV versus ∼2.1 eV). The lack of the metallic character related to the hybridization with the substrate is associated to a lower electronic charge on the carbon nanonetwork with respect to the metalated case and comparable to that of the freestanding layer (∼0.012 e on the supported 2D carbon nanonetwork). The ordering of the different p components is the same as observed for the metallic system. In particular, the occupied p states of the sp carbon chain lie in correspondence of the d band of Au(111).

The theoretical calculation can also provide a simulation of STS images as the DOS at the Γ point, which is representative of the states with the maximum interaction with the tip due to their slow decay in vacuum (see the Supporting Information).

DFT calculations reveal that the interaction between the 2D carbon nanonetwork and the underlying Au(111) surface consistently affects the electronic properties in comparison with the freestanding nanostructure. As detailed above and in the Supporting Information, the metalated nanostructure acquires a metallic character due to a consistent charge transfer while the graphdiyne-like nanonetwork displays a modified gap with respect to the freestanding system.

In these sp carbon-based systems, peculiar structure-dependent conjugation effects and relevant electron–phonon coupling lead to a strong relationship between electronic and vibrational properties. The effect of the interacting Au(111) surface on the vibrational properties of the 2D carbon system is addressed by comparing the experimental Raman spectrum with DFT calculations, as discussed in the next section.

**Raman Spectroscopy and Vibrational Properties.** The Raman spectra of the precursor and the graphdiyne-like 2D carbon nanonetwork are here discussed and interpreted in view of DFT calculations of the Raman response. To provide a reliable interpretation of the experimental trends, we performed DFT calculations on the precursor molecule and on a molecular model describing the final 2D all-carbon nanostructure grown on the substrate.

We start by discussing the spectrum of the molecular precursor (see Figure 6) measured in powder form prior to sublimation. In particular, FT-Raman spectroscopy has been employed to acquire the Raman spectrum of the organic molecule and to avoid the strong luminescence background. By comparing the FT-Raman spectrum of the initial molecule (black) in Figure 6 with the simulated spectrum of the
precursor (red), a very good agreement is found, demonstrating the accuracy of the adopted level of theory in providing an accurate interpretation of the Raman response. The experimental spectrum shows three main spectral features: (i) the sp carbon stretching mode at 2203 cm$^{-1}$, (ii) the peak associated to the sp$^2$ carbon in the phenyl rings at 1581 cm$^{-1}$, and (iii) few weak peaks in the low-wavenumber region (950–1350 cm$^{-1}$), which show a very good correspondence with the peaks predicted in the theoretical spectrum. On the other hand, the two broad bands observed at about 1500 and 1150 cm$^{-1}$ can be attributed to impurities, also possibly due to degradation processes. DFT calculations allow us to assign the peaks in detail, starting from the peak at 2236 cm$^{-1}$ (computed at an unscaled wavenumber of 2319 cm$^{-1}$), which is due to the C–C stretching mode of the triple bond. The peak at 1581 cm$^{-1}$ (computed at an unscaled 1641 cm$^{-1}$) is attributed to the C–C stretching mode localized on the phenyl groups, while the bands at lower wavenumber values (1370, 1201, and 1014 cm$^{-1}$ unscaled values) are associated respectively to C–C stretching of the phenyl coupled to C–C stretching on the three branches and different combination of C–C and C–H bending modes on the phenyl also coupled to the contribution of C–Br stretching.

Ex situ Raman spectra of the 2D nanonetwork on Au(111) annealed at 480 and 580 K have been acquired to follow the evolution as a function of the temperature (see Figure 7). The experimental spectra of the 2D nanonetwork on Au(111) (Figure 7) appear to be significantly modified compared to the spectrum of the precursor molecule (Figure 6). Indeed, the weak signals in relatively low frequency range (950–1350 cm$^{-1}$) disappear while the peaks associated to sp carbon and sp$^2$ carbon in the phenyl ring become broader. By fitting the sp carbon band with two Gaussians (Figure 7c) (one centered at 2200 cm$^{-1}$ and the other one at 2165 cm$^{-1}$), we observe that the contribution of the lower frequency peak (2165 cm$^{-1}$) increases after annealing the sample at 580 K. Concerning the spectral region associated to the sp$^2$ carbon in the phenyl ring, we observe a broad band shifted to a lower frequency (from 1581 to 1565 cm$^{-1}$) compared to the peak of the precursor molecule. The broadening of this band is further enhanced at 580 K probably due to the increased disorder observed in STM images and discussed below.

We computed the Raman spectrum of the all-carbon 2D nanonetwork on Au(111) by adopting the simplified model in which small Au$_4$ clusters are interacting with the sp domains of a fragment of the 2D nanonetwork, as shown in Figure 7b (see Supporting Information, Figure S3 for an extended discussion). DFT calculations predict a double peak associated to the sp carbon chains in a high frequency range, namely, at 2145 and 2178 cm$^{-1}$. This doublet falls at a lower wavenumber with respect to the sp carbon peak predicted for the precursor at 2236 cm$^{-1}$ and it is associated now to collective stretching modes involving the different sp carbon domains, as described for the sp carbon chain by the effective conjugation coordinate (ECC) model.$^{3,18}$ The modulation of their frequency is due to the interactions with the gold clusters, which can affect the stretching force constants of the C–C bond most involved in the interaction with gold, as also found in similar systems.$^{18}$ As a result, the whole collective C–C stretching mode of the sp carbon chains is perturbed by the non-bonding interactions, and the peaks are shifted to a lower wavenumber with respect to those predicted for the unperturbed precursor molecule. This result is in agreement with what we observe in the experimental spectra, where the sp carbon band can be fitted by two peaks at a slightly lower wavenumber with respect to...
the precursor molecule. Therefore, DFT calculations allow us to give an interpretation to the Raman spectra indicating the formation of the 2D nanostructure on Au(111) as also observed by STM imaging. Moreover, DFT calculations predict other two weak peaks at 1939 and 1969 cm\(^{-1}\). The normal mode associated to these peaks can be described as C–C stretching vibrations mainly localized on the C–C bonds which are more strongly interacting with the gold cluster. Due to their weak intensity, these peaks are difficult to observe in the experimental spectra, and they could possibly correspond to the weak bump at 2025 cm\(^{-1}\) barely observable in both spectra.

The comparison with DFT calculations allows us to identify the Raman spectrum of the 2D nanostructure on Au(111). The two main peaks at slightly lower wavenumbers (2165 and 2200 cm\(^{-1}\)) than the precursor molecule provide a clear signature of the occurrence of homocoupling of the molecules on the substrate. Although STM images provide a clear evidence of the metalated 2D nanostructure formed right after the deposition of the molecules on Au(111), we were not able to record an ex situ Raman spectrum due to a low signal-to-noise ratio. Such a behavior was observed also in the case of 1D nanostructures based on sp-sp\(^2\) carbon studied in a previous work,\(^{46}\) and we think that annealing the system in UHV can improve the signal-to-noise ratio, possibly due to higher stability of the whole nanonetwork after exposure to the atmosphere. However, from STM images, we notice a wide temperature window for the homocoupling reaction and the metalated domains are still present after annealing at 480 K and much less at 580 K. The progressive transformation of the structure with the thermal annealing is reflected in the Raman spectra where the peaks at 2165 and 2200 cm\(^{-1}\) exhibit a change in intensity and width with the temperature (Figure 7c). Lastly, the peak at 1565 cm\(^{-1}\), which shows broadening at 580 K with the onset of a pronounced shoulder, can be interpreted as a superposition of the signal coming from the aromatic rings and a combination of D and G bands typical of the sp\(^2\)-carbon amorphous phase, in agreement with the disordered phase observed in the STM image in Figure 2d.

The interaction with Au turns out to modify significantly the vibrational properties and the Raman spectrum. By comparing DFT calculations of the sp-sp\(^2\) system on Au(111) with the freestanding case, three main effects can be seen: (i) the ECC peak of sp carbon stretching mode redshifts, (ii) the ECC peak splits in a doublet, and (iii) a new band at about 2000 cm\(^{-1}\) is present, which is split in two peaks as well. No substantial differences are seen in the peaks at about 1600 cm\(^{-1}\) related to the sp\(^2\)-carbon stretching in the phenyl rings.

The previous work carried out by us\(^{44}\) reported the redshift and the splitting of the sp carbon fingerprint (ECC modes) and the activation of new Raman frequencies not originally present in freestanding nanostructure. The overall redshift of the sp carbon modes and the new Raman peak observed at significantly lower frequencies is attributed to bond softening promoted by the interaction with gold atoms.

CONCLUSIONS

We characterized the atomic-scale structure and the vibrational properties of a carbon sp-sp\(^2\) 2D nanonetwork on Au(111) by means of STM and Raman spectroscopy. In particular, high-resolution STM imaging combined with Raman spectroscopy allowed us to identify the metalated nanostructure formed right after the deposition and the subsequent formation of sp-sp\(^2\) 2D nanonetwork upon annealing in UHV. Raman spectra reported the redshift and the splitting of the sp carbon fingerprint (ECC modes) and the activation of new Raman frequencies not originally present in freestanding nanostructure. The overall redshift of the sp carbon modes and the newly formed peak observed at significantly lower frequencies is attributed to bond softening promoted by the interaction with gold atoms.

The DFT calculations of the electronic properties of both metalated and C–C coupled nanonetwork on Au(111) show a strong contribution of the substrate states and shifting of molecular states. Indeed, the metalated nanonetwork exhibits states at the Fermi level related to the charge transfer from the substrate to the 2D layer, while for the C–C coupled system, the interaction with the substrate is less important but still modifies the gap and the vibrational properties. Although we obtained sp-sp\(^2\) 2D nanonetworks with different domain sizes, we did not observe any variation of the properties with the size since the sp conjugation is limited in the short sp carbon chain and the phenyl ring. A larger extension of the \(\pi\)-electron conjugation could be achieved through the control of the oligomeric unit by a proper design of the molecular precursor, finely tuning the properties of the sp-sp\(^2\) 2D nanonetwork.

Although different research groups have proposed novel graphdiyne-like nanostructures obtained by the on-surface synthesis,\(^{39,40,49,50}\) the characterization of their vibrational properties by Raman spectroscopy is lacking. On the line opened by the present work, UHV-STM and Raman spectroscopy represent a powerful and non-destructive combination of techniques for the characterization of novel
sp carbon systems, such as graphdiyne produced by means of the on-surface synthesis on metal surfaces. These systems can display unique properties resulting from the interaction with the metal surface whose presence is a key factor to catalyze the formation of the 2D layer itself. The control of such interaction through an accurate selection of the systems could open to the design of novel 2D carbon materials beyond graphene. The choice of the precursor molecule plays an important role in determining the final properties of the nanostructure.\(^\text{25}\) For instance, the content and architecture of the linear sp carbon structures defines the sp/sp\(^2\) ratio and pore size. The chemical functionalization of the initial molecule, such as with halogens, tailors the interaction with the substrate, which catalyzes the on-surface synthesis and modifies the electronic and vibrational properties of the resulting system.

Compared with the γ-graphdiyne structure, our carbon nanonetwork shows a threefold connection of sp\(^6\) six-membered aromatic rings through diacetylenic linkages and the remaining threefold dangling bonds are saturated by H atoms. Accordingly, it possesses a uniformly larger pore size than the remaining threefold dangling bonds are saturated by H atoms. The latter suggests a higher band-gap energy compared to γ-graphdiyne, as confirmed by the prediction of about 1.2 eV for γ-graphdiyne\(^\text{19}\) with respect to 2.1 eV for the structure reported here. In addition, the electronic properties of this 2D graphdiyne-like carbon nanonetwork are strongly affected by the interaction with the metallic substrate resulting in a decrease of the band gap to about 1.6 eV. The possibility to modulate the pore size and electronic properties of a 2D graphdiyne-like nanomaterial represents a great potential for future applications in catalysis\(^\text{16,52}\) and nanoelectronics.\(^\text{55}\)

### EXPERIMENTAL AND COMPUTATIONAL DETAILS

The experiments were carried out in two interconnected ultra-high vacuum chambers with a base pressure of less than \(5 \times 10^{-11}\) mbar. The first experimental setup was used for the STM measurements, while the second setup was devoted to sample preparation.

The Au(111) surface was prepared by repeated cycles of Ar\(^+\) ion sputtering followed by annealing at 720 K. The herringbone reconstruction characterizes the clean Au(111) surface with atomically flat terraces as confirmed by STM measurements. The molecular precursor, i.e., 1,3,5-tris(bromothiophenyl)benzene (tBEP), was thoroughly degassed prior to the deposition onto the cleaned Au(111) surface kept at room temperature (RT). The tBEP precursor was thermally sublimated onto the Au(111) surface by means of an Omicron variable-temperature scanning tunneling microscope. STM images were taken in constant-current mode at room temperature, with a chemically etched tungsten tip. The bias for tunneling current is applied to the sample and typical voltages are from \(-1.5\) to \(+1.5\) V and tunneling current in the range of \(0.2\)–\(1.0\) nA.

Density functional theory calculations were performed by adopting standard norm-conserving pseudopotentials and an atomic-oriﬁcal basis set, which includes double-zeta and polarization orbitals, as implemented in the SIESTA package.\(^\text{53}\)

The exchange and correlation potential was treated with the generalized gradient approximation (GGA-PBE),\(^\text{44}\) and the molecular-surface van der Waals interaction was introduced via a DFT-D2 Grimme potential.\(^\text{55}\) The mesh cutoff for the real space grid was set to 450 Ry, and a \(3 \times 3\) and \(4 \times 4\) sampling of the Brillouin zone was adopted for the metalated and C–C coupled system, respectively, corresponding to approximately a \(21 \times 21\) grid in the primitive Brillouin zone of Au(111).

A five times denser grid was used for the calculation of the density of states (DOS). The molecular layers were fully relaxed until the forces reached the tolerance value of 0.04 eV Å\(^{-1}\). The substrate atoms were kept fixed to the coordinates of the relaxed ideal clean Au(111) surface, neglecting the \(22 \times \sqrt{3}\) reconstruction. Along the \(z\) direction, we consider six gold layers, with an interspersed vacuum of 38 Å. The STM simulations were performed in a Tersoff–Hamann approach, assuming a constant density of states for the tip. We integrated the electronic density of the empty states in an energy interval of 0.5 eV just above the Fermi level. STM images were simulated at a constant distance of 3 Å from the surface and applying a 2 Å-wide Gaussian spatial broadening to the electronic density to mimic finite experimental resolution.

MicroRaman measurements were conducted, ex situ, using a Renishaw InVia spectrometer coupled with an Argon laser (514.5 nm). With the power set at 5 mW, we have acquired 100 spectra for each sample aiming at an adequate signal-to-noise ratio. At this excitation wavelength, we have a fluorescent background coming from the Au(111) surface and concurrent with the Raman signal of the sp\(^2\) carbon atomic sheet. The background has been removed by subtracting the signal acquired on the pristine clean Au(111) surface under the same experimental conditions. The Raman spectrum of the molecular precursor in powder form was obtained by FT-Raman (NICOLET NEXUS NXR 9650) with 1064 nm excitation wavelength.

The simulations of the Raman spectrum of the precursor and the carbon nanonetwork have been carried out by applying the approach successfully adopted in a previous paper\(^\text{46}\) on a similar system, using finite-dimension molecular models that properly describe the real system. Modeling the present system is indeed far from being trivial. An extended computational investigation has been carried out to this aim as reported in detail in the Supporting Information. After the calculation of the molecular precursor, DFT calculations have been performed for an isolated fragment of the 2D crystal interacting with three Au clusters in order to model the interaction expected between the Au surface and the sp domains.\(^\text{44}\) This model is reported in the section dedicated to Raman Spectroscopy and Vibrational Properties. The Gaussian09 package\(^\text{56}\) has been used to carry out these calculations, adopting the PBE0 functional and a cc-pVTZ basis set for C and H and ECP60MDF pseudopotential together with VTZ basis set for Au.\(^\text{57}\) The computed vibrational frequencies have been rescaled by a factor of 0.963, determined by adjusting the phenyl stretching mode predicted by DFT at 1641 cm\(^{-1}\) to the value 1581 cm\(^{-1}\) observed experimentally for the precursor.

### ASSOCIATED CONTENT

#### Supporting Information

The Supporting Information is available free of charge at https://pubs.acs.org/doi/10.1021/acsanm.0c02665.

Theoretical simulation of STS spectra and a detailed discussion on methods and models adopted for the computation of Raman spectra of both the molecular precursor and 2D system on Au(111)\(^\text{(PDF)}\).

### AUTHOR INFORMATION

#### Corresponding Author

Carlo S. Casari — Department of Energy, Politecnico di Milano, Milano I-20133, Italy; orcid.org/0000-0001-9144-6822; Phone: +39 022396331; Email: carlo.casari@polimi.it

#### Authors

Andi Rabia — Department of Energy, Politecnico di Milano, Milano I-20133, Italy; orcid.org/0000-0002-7942-7626

Francesco Tumino — Department of Energy, Politecnico di Milano, Milano I-20133, Italy; orcid.org/0000-0002-4846-2701

Alberto Milani — Department of Energy, Politecnico di Milano, Milano I-20133, Italy; orcid.org/0000-0001-6026-5455

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Valeria Russo — Department of Energy, Politecnico di Milano, Milano I-20133, Italy
Andrea Li Bassi — Department of Energy, Politecnico di Milano, Milano I-20133, Italy
Nicolò Bassi — Department of Energy, Politecnico di Milano, Milano I-20133, Italy
Andrea Lucotti — Dipartimento di Chimica, Materiali e Ingegneria Chimica “G. Natta”, Politecnico di Milano, Milano I-20133, Italy
Simona Achilli — ETSF and Dipartimento di Fisica “Aldo Pontremoli”, Università degli Studi di Milano, Milano I-20133, Italy
Guido Fratesi — ETSF and Dipartimento di Fisica “Aldo Pontremoli”, Università degli Studi di Milano, Milano I-20133, Italy
Nicolò Manini — ETSF and Dipartimento di Fisica “Aldo Pontremoli”, Università degli Studi di Milano, Milano I-20133, Italy
Giovanni Onida — ETSF and Dipartimento di Fisica “Aldo Pontremoli”, Università degli Studi di Milano, Milano I-20133, Italy
Qiang Sun — Interdisciplinary Materials Research Center, College of Materials Science and Engineering, Tongji University, Shanghai 201804, P. R. China; orcid.org/0000-0003-4903-4570
Wei Xu — Interdisciplinary Materials Research Center, College of Materials Science and Engineering, Tongji University, Shanghai 201804, P. R. China

Complete contact information is available at:
https://pubs.acs.org/10.1021/acsanm.0c02665

Notes
The authors declare no competing financial interest.

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