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Tailoring Large Pores of Porphyrin Networks on Ag(111) by Metal-Organic Coordination

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Abstract: The engineering of nanoarchitectures to achieve tailored properties relevant for macroscopic devices is a key motivation of organometallic surface science. To this end, understanding the role of molecular functionalities in structure formation and adatom coordination is of great importance. In this study, the differences in formation of Cu-mediated metal-organic coordination networks based on two pyridyl- and cyano-bearing free-base porphyrins on Ag(111) are elucidated by use of low-temperature scanning tunneling microscopy (STM). Distinct coordination networks evolve via different pathways upon codeposition of Cu adatoms. The cyano-terminated module directly forms 2D porous networks featuring fourfold-coordinated Cu nodes. By contrast, the pyridyl species engage in twofold coordination with Cu

and a fully reticulated 2D network featuring a pore size exceeding 3 nm² only evolves via an intermediate structure based on 1D coordination chains. The STM data and complementary Monte Carlo simulations reveal that these distinct network architectures originate from spatial constraints at the coordination centers. Cu adatoms are also shown to form two- and fourfold monoatomic coordination nodes with monotopic nitrogen-terminated linkers on the very same metal substrate—a versatility that is not achieved by other 3d transition metal centers but consistent with 3D coordination chemistry. This study discloses how specific molecular functionalities can be applied to tailor coordination architectures and highlights the potential of Cu as coordination center in such low-dimensional structures on surfaces.

Introduction

Supramolecular coordination chemistry is a vivid field of research as the combination of organic or metal–organic ligands and metal centers yields structures and properties that are not achievable by the individual building blocks. Recent examples highlighting the potential of metal–organic compounds include reports alluding to information storage^[1] and processing,^[2] energy storage,^[3] catalysis,^[4] and molecular electronics.^[5] Considerable efforts have been dedicated to adapting design principles from such three-dimensional (3D) supramolecular structures to a two-dimensional (2D) environment represented

by surfaces in a controlled ultrahigh-vacuum (UHV) setting. [6] Also in this 2D scenario, the functionality of metal-organic coordination networks is represented by the combination of metal nodes, offering, for example, active sites for energy conversion chemistry, [7] and the ligands, dictating the pore sizes, the confinement of adsorbates, [8] and the (magnetic) coupling between the metals. [9] Specifically, the coordination number and symmetry at the nodes are decisive for the topology of the resulting metal-directed architectures. [10] Despite the many reports on surface-anchored metal-organic architectures, [1-10]

■ok?■■ strategies to engineer and deliberately tailor assemblies still need to be further developed and refined. For example, extended 2D networks featuring a grid-like structure with square pores and mononuclear nodes are rarely reported.^[11] In this respect, molecules offering fourfold symmetry might yield advantages compared to the frequently applied ditopic linear linkers. Hereby, tetrapyrroles as porphyrins are ideal candidates, which proved to be stable and versatile building blocks for self-assembled molecular structures on surfaces.^[12] The central macrocycles, which host two hydrogens or a metal center, add functionality to the system, as they can be used as molecular switches, [13] can undergo metalation reactions directly on a surface, [14] and have potential for heterogeneous catalysis [15] and spintronics. [16] Most importantly, the tetrapyrrole macrocycle can be substituted by a wide variety of terminal moieties, offering vast possibilities to steer intermolecular and metal-organic interactions.[11a, 12a-c, 17]

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Herein we present a low-temperature scanning tunneling microscopy (STM) study that compares the Cu-directed assemblies of two de novo-synthesized porphyrins, functionalized with biphenylene-cyano (2H-TPCN) and phenylene-pyridyl substituents (2H-TPyPP), respectively, on Ag(111). Although both molecules feature nitrogen-terminated ligands and assemble into similar organic arrays, they respond markedly differently to the copper atoms. TPCN directly forms 2D metal-organic networks with small pores and fourfold coordination nodes, whereas TPyPP follows a hierarchic pathway from 1D metal coordination chains to an open porous 2D metal-organic network with linear twofold-coordinated metal centers. Monte Carlo simulation and comparison of Co- and Cu-directed networks, indicate that the origin of the two- and fourfold coordination motif results from an interplay between the steric hindrance at the coordination center and the inherent coordination properties of Cu.

Results

2H-TPCN and 2H-TPyPP modules on Ag(111)

The porphyrins investigated in this study are tetrapyrrolic macrocycles substituted at all four *meso* positions either with biphenylene–cyano or with phenylene–pyridyl moieties. Structural models of these two porphyrin derivatives, namely tetra[(4-cyanophenyl)phen-4-yl]porphyrin (2H–TPCN) and tetra[(4-pyridylphenyl)phen-4-yl]porphyrin (2H–TPyPP) are depicted in Figure 1 a and e sok?

characterization of 2H-TPCN on Ag(111) and BN/Cu(111),[11c,18] whereas 2H-TPyPP is addressed in this study for the very first time. Compared to commercially available tetraphenyl- (TPP) or tetrapyridylporphyrins (TPyP) featuring only one phenyl or pyridyl unit in each meso substituent, these novel modules introduce an additional degree of conformational freedom as the terminal ring R² can rotate around the C-C single bond between the two phenyl rings R¹ (cf. Figure 1a and e). Upon surface confinement, this enhanced molecular flexibility strongly influences the molecular self-assembly and coordination characteristics (see below). The adsorption of TPP and TPyP species on Ag(111) induces a saddle-shaped macrocycle deformation where the terminal rings are rotated typically 50-60° out of the surface plane. [13,19] Saddle-shape deformations—induced by steric hindrance between the macrocycle and rotated moieties R¹—are also expected for both 2H-TPCN and 2H-TPyPP. However, as R¹ acts as a spacer, geometrically decoupling the terminal rings R² from the macrocycle, a rather parallel alignment of R² with the Ag(111) surface is anticipated as both, individual benzene and pyridyl rings adsorb on Ag(111) in a planar fashion.[20]

Figure 1 b, c, f and g show high-resolution STM images of 2H–TPCN and 2H–TPyPP/Ag(111) representing occupied states. Both species present four peripheral lobes associated to the *meso* substituents and a donut shape that is assigned to the macrocycle. The latter shows twofold symmetry and appears with a depression in the center, as observed previously for free-base TPP on Ag substrates.^[13] The elongated *meso* substituents of 2H–TPCN are reflected in the larger apparent size of the molecule and an increased intermolecular distance com-

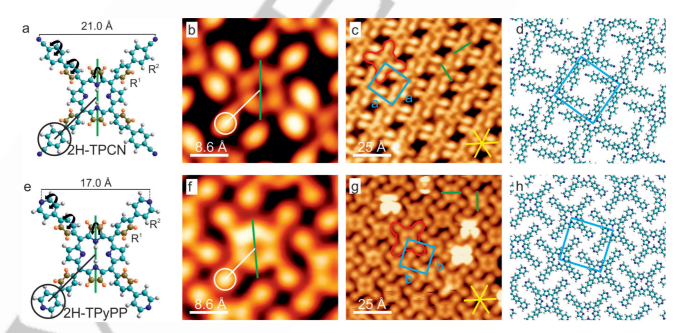


Figure 1. Structural models of the porphyrin derivatives (a, e). The higher parts of rotated molecular moieties are highlighted in orange for better comparison to the magnification on single molecules (b, f) within self-assembled, dense-packed islands on Ag(111) (c, g). d, h) Sketch of the assembly structure. The green lines highlight the molecular axis through the upward rotated pyrroles. A single molecule is outlined in red in (c) and (g) as a guide to the eye and the blue squares indicate the unit cells that include one molecule for both compounds. The substrate dense-packed directions are represented by yellow lines. Scan parameters: b) U = -1 V, I = 30 pA; c) U = -1 V, I = 70 pA; f, g) U = -1 V, I = 0.1 nA. The models were created with HyperChem and the molecular dimensions were extracted after geometry optimization of the free molecule within the semi-empirical AM1 framework.

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pared to 2H-TPyPP (see below). Nevertheless, the overall appearance of both species is rather similar, as the cyano group does not contribute considerably to the STM contrast. [21] Based on a comparison of submolecular features, presented in Figure 1b and f with structural models, a tentative conformation of the meso substituents' orientation can be inferred. Whereas R² looks disk-like, indicating a planar adsorption, R¹ appears as a narrow bridge connecting the macrocycle and R². The asymmetric appearance of R¹ with respect to the axis through the meso position of the macrocycle (white lines in Figure 1b and f) provides an indication for the rotation of R¹. The upper part of the phenyl group dominates the image contrast, in full agreement with previously reported high-resolution STM data (see the Supporting Information of ref. [22]). Steric hindrance between the rotated R¹ and the pyrroles of the macrocycle leads to its saddle-shape deformation. R² appears as a broad protrusion symmetric with respect to the axis connecting opposing legs, in line with the contrast reported for terminal pyridyl groups adsorbed parallel to the Ag(111) surface. [22] We thus conclude that the R² rings are aligned approximately parallel to the surface. Note that the larger apparent height of the legs compared to the macrocycle prevails only at small bias voltages and thus is assigned to an electronic effect (cf. Figure S1 in the Supporting Information). As discussed below, the adsorption geometry of the terminal pyridyl rings in TPyPP is decisive for the distinct coordination behavior from TPyP, in which the pyridyl group is rotated considerably out of the surface plane.

Self-assembly of 2H-TPCN and 2H-TPyPP on Ag(111)

After room-temperature deposition on Ag(111), both modules self-assemble into highly ordered, extended two-dimensional islands (Figure 1 c and g; corresponding structural models are shown in Figure 1 d and h). Both assemblies feature a square unit cell with internal angles of $(90 \pm 1)^{\circ}$ (marked in blue in Figure 1 c and g) with side lengths $a = (20.4 \pm 0.5) \text{ Å}$ for 2 H-TPCN and $b = (18.2 \pm 0.5)$ Å for 2H-TPyPP. In addition, a distinct metastable structure characterized by a rhombic unit cell can be achieved when depositing 2H-TPCN at high flux (see the Supporting Information, Figures S2 and S3). For both porphyrin modules, the molecular axis through the two upwardbent pyrroles of the macrocycle (green lines in Figure 1a and e) is either aligned with the $\langle \bar{1}\bar{1}2\rangle$ or the $\langle \bar{1}10\rangle$ highsymmetry directions of the Ag(111) lattice (green lines in Figure 1 b and f). Whereas 2H-TPCN mostly aligns along $<\bar{1}\bar{1}2>$, as reported for Co-TPP/Ag(111), [19b] no preference is discernible for 2H-TPyPP. Despite these distinct azimuthal orientations induced by the Ag(111) surface, site-specific interactions do not prevail over lateral intermolecular interactions and the 2H-TPCN and 2H-TPyPP arrays are not commensurate with the underlying Ag(111) lattice, as revealed by bias-dependent imaging and dI/dV spectroscopy (see the Supporting Information, Figure S1). Both assemblies are stabilized by lateral noncovalent interactions between neighboring nitrogen-phenylene groups.

Formation of metal-organic coordination networks

To investigate the response of the porphyrin species to metal adatoms, Cu was deposited with submonolayer, monomolecular coverages at room temperature. Figure 2 shows the fully reticulated metal-organic coordination networks and the corresponding structural models. For TPCN, a highly regular porous network with a rectangular unit cell of size $c = (21.9 \pm 0.5) \text{ Å}$ and $d = (24.0 \pm 0.5)$ Å evolves, featuring domains that extend over hundreds of square nanometers with a low defect density (Figure 2a). The long-range order and the symmetry of the network are reflected in the autocorrelation plot and a sharp fast Fourier transform (FFT) pattern (Figure 2b). High-resolution images (Figure 2c) reveal details of the network structure, with a pore size of approximately 86 Å² exposing bare Ag. Clearly, every node—assigned to a Cu adatom—links four TPCN units through their carbonitrile termini, resulting in a 1:1 stoichiometric ratio between TPCN and Cu. The projected N-Cu distance is (1.6 \pm 0.5) Å, in agreement with previously reported interfacial cyano-Cu coordination.[23] Within the network, the molecules appear slightly compressed compared to the organic phase; their aspect ratio deviates from unity, resulting in an "X"-like shape.

This is also reflected in the rectangular unit cell, which differs from the square unit cell reported for Gd-coordinated TPCN networks. [18] The reduced symmetry might be induced by the flexibility of the meso substituents combined with the favorable hollow site absorption of Cu adatoms on Ag(111).[24] Indeed, a simple model overlay of the coordination network onto a lattice representing the Ag(111) substrate demonstrates that a highly regular, commensurate (8×80) structure can evolve with Cu adatoms exclusively at hollow sites (see the Supporting Information, Figure S5 for the details). Apparently, the energy gained by formation of the commensurate network—enabled by the specific dimensions of 2H-TPCN—exceeds the energy costs for deforming the molecule. As usual for 3d transition metals, the coordination center is not visualized in STM. [25] However, an indirect fingerprint of the metal coordination is observed, as the coordinated terminal groups of TPCN appear higher than those that are noncoordinated (see the Supporting Information, Figure S4). For TPCN, fully reticulated metal-organic coordination networks were obtained, coexisting with dense-packed organic islands and large Cu clusters on the Ag(111) support under all employed preparation conditions. Thus, the yield for the metal-organic network formation is not optimal at the given preparation temperature and copper flux. However, additional architectures based on a simultaneous expression of metal-organic and organic bonding motifs were never observed for TPCN and Cu. [26] This was also confirmed by the Monte Carlo simulations (see below).

For TPyPP, an extended metal–organic coordination network also evolves upon exposure to Cu (Figure 2e and f), characterized by a molecule/Cu adatom ratio of 1:2 (Figure 2e and g). All four pyridyl termini of 2H–TPyPP are connected with the adjacent molecules by pyridyl–Cu–pyridyl coordination motifs, forming a square unit cell with a side length of e= (28.2 ± 0.5) Å. The projected N–Cu distance amounts to

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Figure 2. Formation of metal-organic networks upon deposition of Cu atoms. The blue squares indicate the unit cells and one molecular unit is outlined in red as a guide to the eye in (c) and (g). For TPCN (a-d) every coordination node is surrounded by four molecules and the unit cell consists of one molecule and one Cu atom. In contrast, the unit cell of TPyPP/Cu (e-h) consists of one molecule and two Cu atoms. The regularity of the metal-organic networks is reflected in autocorrelation plots (upper images in b and f) and sharp spots in FFT images (lower images in b and f). d, h) Model sketches of the networks. Differences in the molecular appearance are assigned to the interaction of the macrocycle with Cu adatoms. The yellow stars represent the substrate's dense-packed directions. Scan parameters: a) U = 0.7 V, I = 50 pA; c) U = 0.2 V, I = 80 pA; g) U = -0.2 V, I = 80 pA.

(1.9 $\pm\,$ 0.5) Å, in agreement with the previous reports. $^{[25a,27]}$ Similar head-on, twofold Cu-mediated coupling motifs of pyridyl moieties are well-known in surface-confined coordination chemistry. [11a,22b,25a,27,28] Compared to the dense-packed organic arrays, TPyPP modules within the metal-organic network are rotated by 45°. Apparently, the energy arising from metal coordination exceeds the energy penalty from deviation from the original adsorption orientation, underlining the weak site-specific molecule-substrate interaction. The network domains extend over hundreds of square nanometers and exhibit longrange regularity (see autocorrelation plot and FFT pattern in Figure 2 f). However, high-resolution STM data reveal that the pores vary in size and shape (Figure 2 g). This local disorder is attributed to the flexibility of both the pyridyl-Cu-pyridyl motif, which features bond angles deviating from 180 $^{\circ}$, and the meso substituents. [22b,27] Thus, a variety of pore shapes that deviate from a perfect square can coexist, which classifies this architecture as a 2D short-range disordered crystalline network. [29] The average pore size amounts to 340 Å². To our knowledge, such a large area is unprecedented for homomolecular surface-supported porphyrin-based architectures. Consequently, the Cu-directed TPyPP network might serve as a template to trap and order large adsorbates or even molecular aggregates. [30] To this end, the intrinsic flexibility opens perspectives for hosting and sorting specific molecular guest species, enabling an adaptive behavior of the pores, thus representing a two-dimensional analogue of a "soft porous crystal".[31] In contrast to the fully reticulated TPCN coordination architecture that evolves directly from the organic islands, the

TPyPP coordination follows a hierarchic pathway upon increasing the (local) density of Cu adatoms. After depositing small amounts of Cu adatoms onto a submonolayer of 2H-TPyPP/ Ag(111), a porous array appears that is characterized by chainlike sub-structures (Figure 3a). A close inspection reveals that it expresses simultaneously metal-organic and organic bonding motifs like those described in ref. [45]. As visualized in the corresponding structural model (Figure 3d), TPyPP tectons are dense-packed in one direction (organic bonding, marked with f) and form a head-on configuration along the other direction (marked with e), which is assigned to a pyridyl-Cu-pyridyl coordination bond, in analogy to the fully reticulated network (cf. Figure 2). The network is thus formed by 1D metal-organic chains that mutually interact through lateral noncovalent interactions between neighboring nitrogen-phenylene groups and follow the dense-packed substrate directions. The structure features a rhomboid unit cell of size $e = (28.0 \pm 0.5) \text{ Å}$ and f =(18.2 \pm 0.5) Å including an angle of (60 \pm 1) $^{\circ}$ and a TPyPP/Cu adatom ratio of 1:1.

When the Cu dosage is increased and (locally) exceeds a 1:1 ratio of Cu adatoms to molecules, the organic bonds are transformed into pyridyl–Cu–pyridyl coordination bonds and the structure evolves into a fully reticulated 2D metal–organic network. Figure 3 b and the corresponding structural model in Figure 3 d show the transition from 1D coordination chains to a 2D coordination network. Both architectures coexist locally and TPyPP can form coordination bondsfrom Cu to two, three, or four adjacent molecules. Clearly, the *meso* substituents engaged in coordination bonds appear brighter than their non-

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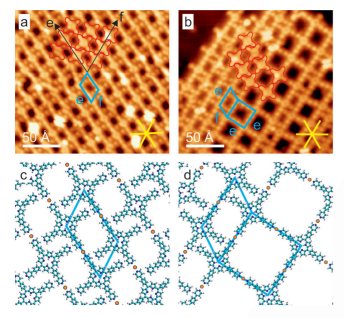


Figure 3. For TPyPP, depending on the (local) Cu density, metal coordination evolves in one or two directions. a) 1D coordination along one specific direction, indicated here by the black arrow labeled with "e", coexists with organic interactions along "f". Coordination will firstly be completed along one direction before starting in another direction (as shown in b). c, d) Models. As a guide to the eye, some molecules are outline in red. The yellow stars represent the substrate dense packed directions. Scan parameters: a–c) U=-0.7 V, I=0.1 nA.

coordinated counterparts (cf. Figure 3 a, c and Figure S4 in the Supporting Information), in line with the results discussed for TPCN.

Monte Carlo modeling

The experimental results reveal striking differences in the formation of metal-organic coordination networks and the corresponding coordination motifs, despite the similarities of the systems, that is, mononuclear Cu nodes coordinated to nitrogen of quasi-fourfold symmetric porphyrins on Ag(111). To rationalize the experimental findings, Monte Carlo modeling was performed, as structure formation is known to be correctly reproduced in such simple simulations for a variety of functional tectons, including porphyrins^[17,32] and phthalocyanines.^[33] For these simulations, both TPyPP and TPCN are represented by a fourfold symmetric cross and metal adatoms are depicted as circles (e.g., inset in Figure 4a), following a representation previously introduced for similar systems.[34] Within the description of this simplified model, the two porphyrin species are identical. To distinguish the two modules, TPyPP is restricted to form one- and twofold coordination nodes only, as observed in the experiment, and TPCN is allowed to engage in one-, two-, three-, or fourfold coordination. Snapshots of the Monte Carlo simulations are depicted in Figure 4a-d (see Methods section for details). Naturally, the metal-free dense-packed islands are identical for the two species (Figure 4a). After including a small amount of adatoms, TPyPP shows forms metal-organic chains (Figure 4b) that evolve into an open-porous 2D coordination network (Figure 4c) when more metal is added. In contrast, for TPCN the addition of small amounts of adatoms already leads to the formation of a 2D coordination network (Figure 4d). Importantly, the simulations reflect all assemblies observed experimentally and correctly reproduce a key difference between TPyPP and TPCN, namely the fact that 1D metal–organic chains emerge exclusively from TPyPP, even though they would be allowed for TPCN.

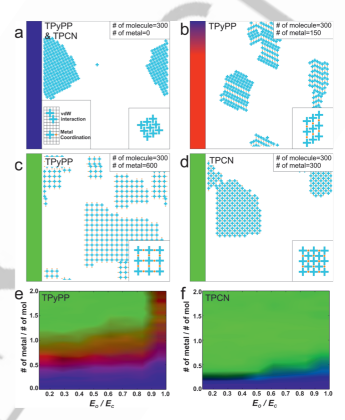


Figure 4. a–d) Snapshots of the structure formation of porphyrin linkers with metal atoms modeled in Monte Carlo simulations. a) 300 molecules and no metal atoms. Without metal atoms both species yield the same assembly. b) 300 TPyPP and 150 metal atoms. c) 300 TPyPP and 600 metal atoms. d) 300 TPCN and 300 metal atoms. The lower-left inset in (a) highlights the two interactions included in the simulations. The bottom-right insets in (a–d) provide a magnification of the dominating pattern. e, f) Phase transition diagrams for TPyPP and TPCN respectively, as a function of the relative number of units (no. of metal atoms)/(no. of molecules) and the ratio of interaction energies E_0/E_C . Due to the finite island sizes, the transition to the coordinated phase (green) proceeds for TPCN already at values of (no. of metal atoms)/(no. of molecules) < 1 (cf. Figure S7 in the Supporting Information). ■ ■ ok? ■

To further characterize the system, additional Monte Carlo simulations were carried out for different interaction energy ratios $E_{\rm O}/E_{\rm C}$ (van der Waals interaction vs. metal-coordination). The results are summarized in phase transition diagrams for TPyPP and TPCN, shown in Figure 4e and 4f, respectively. To generate these plots, the resulting structures are color-coded; blue for the dense-packed organic network, red for the 1D coordination chains, and green for the 2D metal-organic network. Note that the 2D fully reticulated coordination networks are different for the two species (TPyPP in a twofold fashion





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and TPCN in a fourfold fashion), despite being both represented in green (see insets in Figure 3 c and d, respectively). The most prominent difference between the two diagrams is the lack of 1D coordination for TPCN—although it is allowed—regardless of the ratio of E_0/E_0 evidenced by the lack of red in Figure 4 f. Apparently, a separation of organic islands and fully reticulated fourfold coordinated arrays is energetically favored over mixed organic/twofold coordination assemblies. Furthermore, the simulations yield no 1D metal-organic chains for TPCN at any probed temperature (see Methods section and Figures S7 and S8 in the Supporting Information), thus ruling out any influences of the experimental preparation conditions. The TPyPP structure formation proceeds via 1D chains (red, Figure 4 d) for all calculated E_0/E_c ratios, despite the possibility to directly form 2D coordination networks. Therefore, in the case of terminal groups expressing only in a twofold coordination motif, the observed chaining seems energetically clearly favored even for large relative values of E_c . In the modeling, the only difference between the two species is the permitted coordination geometry at the node; no effects of the electronic structure are considered. As the simulations qualitatively reproduce the experimentally observed networks, it suggests that the different formation pathways and the resulting network topologies are mainly determined by distinct spatial constraints at the coordination center.

Discussion

Based on the above findings, the following conclusions can be drawn. Without steric hindrance, Cu adatoms favor a fourfold coordination to the nitrogen termini of porphyrins on Ag(111). In the experiments, this situation is realized for the slender cyano moieties of TPCN. Restricting the nodal symmetry to twofold by introducing spatial constraints due to planar or near-planar pyridyl groups in TPyPP, assemblies featuring 1D coordination chains can be achieved for molecule to adatom ratios below 1:1. Increasing the adatom concentration induces formation of additional pyridyl links, thus yielding a fully reticulated porous coordination network. Regarding ligands, the observed linear pyridyl-Cu-pyridyl motif was tentatively assigned to steric hindrance by several studies.[11a,22b,25a,27] Only when relaxing these constraints by rotating terminal pyridyl rings out of the surface plane, for example, by using the TPyP modules, a fourfold coordination to mononuclear centers can be achieved^[35] (see the Supporting Information, Figure S6). Such square-planar motifs are well known for pyridyl complexes in 3D coordination chemistry, [36] but uncommon in a 2D environment.[37] Their rare occurrence on metallic supports might be attributed to several aspects: A large adatom-nitrogen distance with respect to the surface induced by the rotation of the pyridyl ring out of the surface plane, weakening the pyridyl-adatom interaction, the nature of the coordinating metal center (see below) or simply the limited number of studies addressing molecular modules featuring rotated terminal pyridyl moieties. To achieve fourfold coordination with coplanar adsorbing moieties, a terminal group inducing minor steric constraints (e.g., cyano) is required. In this sense, the pyridylphenylene substitution of the de novo-synthesized 2H–TPyPP providing rotational flexibility to the termini is crucial for the formation of linear pyridyl–Cu–pyridyl binding motifs and for the construction of large-pore Cu-mediated coordination networks. Spatial constraints at the coordination center, which can be tuned by the geometric footprint of the terminal moieties, can be deliberately exploited to control the coordination number and thus the topology of the network architectures. Additionally, the Monte Carlo simulations show that the spatial constraints of the pyridyl groups not only influence the final architectures, but also induce an energetic preference for the formation of 1D coordination chains and therefore are the origin of the hierarchic assembly protocol in the case of TPyPP.

Regarding the role of the metal center, our experiments reveal that single Cu adatoms - somewhat neglected in on-surface coordination chemistry to date - can link four ligands in a quasi square-planar arrangement. Fourfold mononuclear 3dmetal nodes on metal substrates reminiscent of the squareplanar coordination motif have been observed for Mn,[38] Fe,[35a] Co^[21c,35b] and Ni.^[9] Recently, on-surface fourfold coordination was achieved in porphyrin-based metal-organic networks by either applying lanthanide centers[18] that support high coordination numbers^[39] or by introducing a boron nitride spacer layer.[11c] To our knowledge, a coordination number of four on surfaces was only reported for Cu dimers^[40] in metal-organic networks and for distorted Cu-carboxylate complexes.[41] This study introduces the first coordination network based on a square-planar motif based on monoatomic Cu centers. Accordingly, on Ag(111), Cu adatoms can form coordination bonds to two, [22b, 25a, 27, 42] three, [43] or four nitrogen atoms. This diversity in on-surface coordination numbers discriminates Cu from other 3d metals such as Co, for which threefold coordination reminiscent of the trigonal planar motif known from 3D coordination chemistry prevails. Even for cross-like TPCN molecules on Ag(111), Co-coordination results in a random metalorganic network in which three- and fourfold nodes coexist, [18] thus ruling out a dominating role of the molecular symmetry on the resulting metal-organic architecture. Indeed, a quantitative analysis of coordination geometries of d-block metals in 3D supramolecular complexes and solid-state structures shows a frequent occurrence of square planar and square pyramidal motifs for Cu, which only play a negligible role for Co. [44] Of course, one should be well aware that the metallic surface can drastically influence the coordination behavior, allowing for non-integer oxidation states, coordination spheres unachievable in solution, and coordinatively unsaturated centers exposing apical sites to vacuum. This is exemplified by the coordination of cyano to Co with dicarbonitrile–polyphenyl linkers^[8a,45] or to Cu with 9,10-anthracenedicarbonitrile molecules. [46] Furthermore, the metallic surface might mimic an additional ligand^[47] and thus reduce the coordination number in the 2D adsorbate systems. [8a] From this point of view, the twofold pyridyl-Cu-pyridyl link translates to a T-shaped coordination sphere of Cu with three ligands (two pyridyl groups and Ag). Indeed such T-shaped motifs are observed for Cu centers in 3D complexes, in contrast to Co preferring tetrahedral or pyramidal geometries. [44,48] Additionally, TPyP molecules form a four-





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fold coordination motif with Fe^[35a] but a twofold coordination with Cu^[11a] on Au(111), which fits observations from 3D chemistry where Fe, similar to Co, strongly prefers coordination to four or more partners. Thus, in addition to the important steric constraints induced by the pyridyl rings adsorbed approximately parallel to the surface, as discussed above, the naturally preferred coordination geometries of Cu itself might contribute to the stabilization of the linear pyridyl–Cu–pyridyl motif prevalent on coinage metal surfaces and at the same time support fourfold coordination in the case of TPCN.

Conclusion

In summary, by exploiting the preferred coordination geometries of copper in combination with a deliberate porphyrin functionalization, we were able to design extended, 2D, grid-like metal–organic coordination networks on Ag(111). Both TPCN and TPyPP thus offer a basis for the fabrication of bimetallic^[18] and mixed-valence^[11a] open porous networks through orthogonal insertion of metals.^[18] Additionally, for TPyPP, the large pore size and the flexibility of the pyridyl–Cu–pyridyl links result in 2D structure reminiscent of a soft porous crystal, providing opportunities to act as a template for the selective adsorption of molecular guests or nanostructures.^[12g]

To rationalize the formation of distinct Cu-mediated structures from TPCN and TPyPP, we performed Monte Carlo simulations and related the experimental findings to reports on Comediated coordination networks and metal-organic complexes in solution chemistry. This comparison reveals that the choice of coordinating metal is decisive for the emerging coordination motif; for example, replacing Co by Cu in TPCN coordination assemblies results in a highly regular network rather than a random structure. Here, coordination geometries in 3D metal-organic complexes can provide some clues for an appropriate selection of suitable metal for a targeted motif. Co preferentially binds in a tetrahedral fashion and therefore is not a promising candidate for the formation of linear, twofold coordination motifs on surfaces. In contrast, Cu was identified by our study as a versatile center supporting different coordination numbers and geometries. By using TPCN, we achieved the first surface-based coordination network based on a fourfold motif and a mononuclear Cu center. Additionally, our study indicates that the ligand properties (e.g., rotated pyridyl vs. planar pyridyl vs. cyano moieties) must fit the targeted nodal geometry and thus can be used to tailor the resulting network structure and their formation pathway through spatial constraints. Furthermore, we demonstrated the benefits of basic Monte Carlo simulations in selecting suitable molecular modules for metal-organic architectures prior to the actual experiment. Consequently, our study introduces prospects for the programmed design and selection of molecular and monoatomic building blocks for surface-confined supramolecular networks and thus contributes to controlled engineering of metal-organic/organic architectures.

Experimental Section

Experimental procedures

All experiments were performed in a custom-designed ultrahigh vacuum chamber housing a commercial STM from Createc operated at 5 K. The base pressure during experiments was below 3× 10⁻¹⁰ mbar. Repeated cycles of Ar⁺ sputtering and annealing to 725 K were used to prepare the Ag(111) single crystal. 2H–TPyPP and 2H–TPCN molecules were dosed from a thoroughly degassed quartz crucible held at 760 K. During deposition, the sample was kept at room temperature. Cu was evaporated from a home-built, water-cooled cell by resistively heating a W filament supporting a Cu wire of high purity (99.9999%). All STM images were recorded in constant current mode by using an electrochemically etched tungsten tip prepared by sputtering and controlled dipping into the Ag(111) substrate. In the figure captions, voltage *U* refers to

the bias voltage applied to the sample. The WsXM program

(www.nanotec.es) was used to process the STM raw data.

Monte Carlo simulation

A square lattice with 100×100 points is used as substrate due to the shape of the molecules. Both TPyPP and TPCN are represented by a cross that occupies five lattice sites, whereas a metal atom fills one lattice site (e.g., inset in Figure 4a). Only two intermolecular interactions are considered and are limited to be short-ranged and directional, that is, they reach one lattice site along the direction of the molecular substituents. The first is the van der Waals interaction with energy E_0 , which occurs when two molecules align in a dense-packed fashion (see top right inset in Figure 4a) and the second describes metal coordination with interaction energy E_{c} between molecules and metal adatoms. It can only be formed when the metal atom is placed on a lattice site right at the end of a molecular substituent. For TPyPP, the metal atom is restricted to form one- or twofold coordination to mimic steric hindrance. Thus, a metal atom can express coordination bonds with a maximum of two TPyPP units, which have to be on opposite sides of the metal atom. In contrast, a metal atom can coordinate up to four TPCN units. The molecule-substrate interaction is neglected, which is a reasonable approximation, as the experimental observations reveal no relevant site-specific molecule-substrate bonding. In the simulations, E_C is kept constant at a value of 40 (expressed in kT units of energy), while E_0 is varied from 5 to 40. These values are selected by considering both theoretical values of the bonding energies and experimental results. According to previous reports, the bonding energy of a Cu-N coordination bond varies from 0.5 to 2 eV for 3D systems^[49] and the energies of noncovalent bonds, here T-type^[50] or PARI interactions,^[51] range from 40 to 80 meV. As described above, molecules are deposited at room temperature, which means the formation temperature of the self-assembly should be less than or equal to this value. However, stable densepacked islands of TPP have been observed at room temperature, [52] indicating that at this temperature the bonding energies already exceed the thermal energy. Therefore it is reasonable to choose a temperature range of 200-250 K in the simulation. By using these values to express the bonding energies in kT units, we get $E_{\rm c} = 40$ and $E_{\rm o} = 5$ (see above). The number of molecules is kept constant at 300. The simulation procedure follows the protocol described in ref. [53].

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Porphyrins

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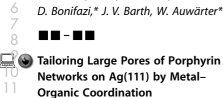
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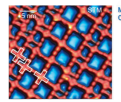
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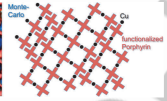
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F. Bischoff, Y. He, K. Seufert, D. Stassen, D. Bonifazi,* J. V. Barth, W. Auwärter*









The grids are alright: A scanning tunneling microsopy study combined with Monte Carlo modeling of two similar, but distinctly functionalized, porphyrin tectons reveals how steric hindrance at Cu coordination nodes guides the assembly of porous grid-like networks with unprecedented morphology and pore size.

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