Adatom mediated adsorption of N-heterocyclic carbenes on Cu(111) and Au(111)

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Abstract
The adsorption of N-heterocyclic carbenes (NHCs) on Cu(111) and Au(111) surfaces is studied with density-functional theory. The role of the molecular side groups as well as the surface morphology in determining the adsorption geometry are explored in detail. Flat-laying NHCs, as observed experimentally for NHC with relatively small side groups, result from the adsorption at adatoms and give rise to the so-called ballbot configurations, which are more stable than adsorption on flat surfaces and provide an efficient precursor for the formation of bis(NHC) dimers. On Au(111), the resulting (NHC)2Au complexes are purely physisorbed and thus mobile. On the more reactive Cu(111), in contrast, the central Cu atom in the (NHC)2Cu dimer is still covalently bound to the surface, resulting in a mobility, which has to be thermally activated.

KEYWORDS
adatoms, DFT, M(111) surfaces, N-heterocyclic carbenes, side groups

1 | INTRODUCTION

N-heterocyclic carbenes (NHCs)\(^1,2\) have been recently attracted much attention as suitable ligands for stabilization and modification of metal surfaces and nanoparticles.\(^3-6\) NHCs are organic molecules, which include in their central ring one or more N atoms and the so-called carbene C atom, that is, a neutral carbon atom with a valence of two and two unshared valence electrons (see Figure 1), giving rise to a high reactivity. They find applications in various chemistry fields\(^2\) as well as in the electronic device technology.\(^7\) In many instances, NHCs form highly ordered self-assembled monolayers on planar metal\(^7-19\) and silicon surfaces\(^20\) with considerable thermal and chemical stability\(^7,8,20-22\) and a remarkable influence on the surface properties.\(^2,15,19,20,22-25\) In particular, such systems appear to be promising in connection with doping of (organic) semiconductors, as similar molecules have been shown to allow immobilization of charge carriers via covalent anchoring.\(^26-28\)

In the last decade, NHCs have been widely studied on flat Au surfaces.\(^7,9,13,14,22,23,25,29,30\) However, the interaction with more reactive surfaces has received less attention. Hitherto, there is merely a handful of investigations on Cu and Ag surfaces.\(^10,12,16,31,32\) Recently, particular attention has been paid to the different binding modes of the NHCs on metallic surfaces and to the role of their side groups in determining these modes. Jiang et al.\(^10\) and Wang et al.\(^9\) reported flat-laying adsorption configurations for NHC with relatively small N-bound organic substituents on coinage metals\(^10\) and Au(111),\(^9\) respectively. It has been furthermore shown that NHCs can easily bind to under coordinated metal adatoms on the respective metal surfaces.\(^7,12,32\) NHCs with bulkier N-substituents prefer upright adsorption geometries, displaying high surface mobility.\(^9\) In line with these studies, but for another class of NHC molecules, Larrea et al.\(^12\) reported the self-assembled monolayers of three NHCs, namely benzimidazolium bicarbonates bearing methyl, ethyl, and isopropyl wing tip groups on Au(111) and Cu(111) surfaces (denoted NHC1, NHC2, and NHC3 in Reference [12] and in the following, see Figure 1). They found again that bulky isopropyl substituents (in case of NHC3) are responsible for allowing the molecules adopt upright geometries on both surfaces, while the less bulky NHCs (NHC1 and NHC2) lie flat on planar surfaces.
Very recently, the work of Deng et al.\textsuperscript{32} was published, which reports on DFT calculations on NHC\textsubscript{1} adsorption (labeled NHC\textsuperscript{Me} in Reference [32] at different solid surfaces, including Au(111) and Cu(111), suggesting that adsorption at adatoms plays a decisive role for the formation of (NHC\textsubscript{1})\textsubscript{2} dimer complexes.

In the present work, we extend this DFT study to NHC\textsubscript{2} and NHC\textsubscript{3} in order to investigate the influence of the steric bulk effect of the side groups, which has been frequently discussed in experimental studies.\textsuperscript{7–10,14,16} Surface adatoms are confirmed to play a decisive role in determining the adsorption configuration independent on the size of the molecular side groups. Furthermore adatom attached NHC are characterized as \textit{bistable} complexes with both \textit{upright} and \textit{flat-laying} configurations, providing an efficient precursor for the observed formation of bis(NHC) dimers.

2 | METHODOLOGY

Total-energy density functional calculations are performed using the Quantum-ESPRESSO package.\textsuperscript{33,34} The electron–electron exchange-correlation energy is modeled using the Perdew-Burke-Ernzerhof (PBE) exchange-correlation (XC) functional\textsuperscript{35} complemented with the non-empirical Tkatchenko-Scheffler dispersion correction.\textsuperscript{36} The Kohn–Sham orbitals are expanded in a plane-wave basis functions with 60 Ry energy cut-off. The Cu(111) and Au(111) surfaces are modeled using periodically repeated slabs with a thickness of four atomic layers separated by 20 Å-thick vacuum. Thereby, the bottom two layers are fixed at their bulk positions during structure optimization. If not stated otherwise, the substrate two top layers and the adsorbate are allowed to relax freely. Supercells up to 9/\textsubscript{C2}×9/\textsubscript{C2} surface unit cells were used. Thanks to their large lateral dimensions (23.16 Å for Cu(111) and 26.40 Å for Au(111)), Γ-only k-point meshes came out be sufficient for a proper description of the metallic surfaces with NHC molecules on top. In fact, we did not find considerable change in adsorption energy trends and bond lengths for these molecules if compared with larger Monkhorst-Pack (MP) k-point grids (see Tables SI and SII).

Atomic structure optimization is performed with convergence criteria of 3 meV/Å for the atomic forces and 10\textsuperscript{-7} eV for total energy difference. The adsorption energies ($E_{\text{ads}}$) are given by

$$E_{\text{ads}} = E_{\text{sys}} - E_{\text{sur}} - E_{\text{mol}},$$

where $E_{\text{sys}}$, $E_{\text{sur}}$, and $E_{\text{mol}}$ are the energies of the adsystem, the surface (with and without adatoms) and the gas-phase molecule, respectively.

For a few selected structures, the migration barriers on the surface have been determined via the corresponding saddle-point structures, which are determined under the constraint that the carbene carbon atom is laterally fixed on its position along the \langle 1 1 0 \rangle direction of the surface (see also Appendix).

3 | RESULTS AND DISCUSSIONS

3.1 | Single N-heterocyclic carbene molecules on planar surfaces

We start our DFT calculations by carefully determining the most stable structures of the three NHCs on pristine Au(111) and Cu(111) surfaces. Since the registries of the molecules depend on the flexible orientation of the side groups and the orientation of the molecule itself and provide a flat energy landscape, a large variety of plausible starting configurations have to be used and fully relaxed to identify the ground state geometries and possible intermediate states. This includes different spatial geometries (flat-laying, perpendicular and tilted), different rotational degrees of freedom and different surface-
related registries (top, bridge, fcc and hcp hollows) of the adsorbed NHC molecule (see Figure S1 and Tables SIII and SIV for details).

In agreement with previous works,9,16,30 the most stable structures of the three NHCs on ideal surfaces are characterized by a formation of a single covalent bond between the NHC’s carbene carbon atom (see Figure 1) and one atom from the metal surfaces (see Figure 2, Figure S3, and the charge density difference plots in Figure S6). Such geometries are denoted as surface-bound species in the following. Table 1 compares the characteristic structural parameters, the adsorption energies and the contributions of the dispersion interactions to the adsorption energies for the three surface-bound NHCs in their most stable states.

The calculated $d_1$ values for the adsorption heights are in very good agreement with those reported for other NHC on Au(111),8,14,15,25,29,30,32 and slightly longer than that reported in Reference [31] (1.97 Å) for an NHC on Cu(111). However, this value is apparently shorter than that reported in the recent work of Deng et al.32 (2.21 Å) and might be related to the laterally smaller supercells used in Reference [31]. In any case, for a given substrate the contribution of the covalent interactions ($\approx 0.85$ eV on Cu(111) [−1.05 eV on Au(111)]), and by this the bond length ($d_1 = 2.05$ Å) (−2.15 Å) are almost the same for the three NHCs. The most striking effect onto structure is thus given by the influence of the substrate, as the bond lengths on the more reactive Cu(111) surface are shorter by about 5% compared to adsorption on Au(111) surfaces. Consequently, the adsorption energies for the three NHCs are slightly higher on Cu(111) compared to Au(111). A similar trend was observed for NHCl3 in Reference [12]. However, an opposite tendency was reported by Deng et al.32 for NHCl1, where surprisingly high adsorption energies on Au(111) were obtained.

Our calculations clearly show, however, that the adsorption energies are increasing with the size of the side groups. Apparently, this is related to the increased contribution of the dispersion interactions of 53% (45%), 58% (52%) and 63% (58%) for NHC1, NHC2, and NHC3 on Cu(111) (Au(111)), respectively. The vertical shift of the metal atom on which the molecule anchors ($d_2$) shows the same tendency as well, further highlighting the crucial role of the dispersion interactions to determine adequate molecular geometries and reliable adsorption energies, also in case of covalently bound species.

It is important to note that our calculations found no indications for any flat-laying geometries. Irrespective of the start structure, the geometries of all studied molecules have been finally converged to upright configurations (with $\theta < 10^\circ$; see Table 1). Vertical geometries ($\theta = 0^\circ$) can be achieved within a few meV. In contrast, forcing the molecules to adopt flat-laying geometries would require energies in the order of 1 eV (see Figure S2 for details). This is true not only on Au(111),14,30 but also on Cu(111). In both cases, flat-laying adsorption geometries can be safely ruled out on pristine metallic substrates and thus cannot account for the different binding modes reported by Larrea et al.12 for these NHCs.

3.2 Adsorption at Cu/Au adatoms

Adsorption at step edges or at omnipresent adatoms might facilitate a flat adsorption scheme (see Figure 3). Such scenarios were previously reported for isonitriles,37 thiols38 and NHCs on Au(111).9 According to Wang et al.,9 the energy cost for the extraction of an Au atom (to form an adatom), is reduced by one order of magnitude (to the 100 meV regime) if small-sized NHC are attached. This premise has been verified by STM height measurements,9 which confirmed the adsorption of NHCs at adatoms on the Au(111) surface. On Cu (111), extracting Cu atoms from steps and their incorporation in the molecular assembly is reported to be a thermally activated process.39

Adatoms are also reported to play a key role for the formation of NHC self-assembled monolayers.40 The presence of adatoms explains the high mobility of NHCs on Au(111) surface despite strong NHC–Au bond formation. As suggested by Wang et al. for 1,3-dimethylimidazol-2-ylidene (IMe) molecule and by Larrea et al.12 for NHC1, NHC2, and NHC3, the formation of NHC–adatom complexes enables the so-called ‘ballbot’-type motion of these complexes. It offers low barriers to diffuse, much lower than surface-bound molecules and even lower than the adatom alone, thereby promoting the molecular self-assembly. As a
simple test for this premise, we calculate the migration barriers of NHC3 along the <110> direction of Au(111). Barriers of about 0.9 eV for single surface-bound NHC3 are lowered to about 0.2 eV for the NHC3-Au ‘ballbot’ complex (see Figure S4).

3.2.1 | N-heterocyclic carbene ballbot species on Cu(111) and Au(111) surfaces

In the following, we show that this NHC ballbot species, that is, adsorption on surface (M) adatom on M(111) surface, M@M(111), is actually able to explain the different adsorption modes experimentally observed. In agreement with Wang et al.9 single Au [Cu] adatoms are found to be most stable when they occupy fcc hollow sites of the surface at a distance of \( d_0 = 2.00 \text{ Å} \) (1.86 Å). In order to decorate these adatoms with NHCs, we tested a wide variety of plausible start geometries of NHC1, NHC2, and NHC3 in various ballbot configurations. Notably, in contrast to the surface-bound species, we found for each species stable upright (standard ballbot) and flat-laying structures on both metal surfaces (see Figure 3).

Table 2 presents the structural geometries and the adsorption energies of the most stable structures of the three NHCs on Cu(111) and Au(111) in the upright and flat-laying binding modes (see also Figure S5). Compared to the surface-bound species, NHC ballbot species in the

<table>
<thead>
<tr>
<th>Configuration</th>
<th>( E_{\text{ads}} ) (eV)</th>
<th>( E_{\text{vdW}} ) (eV) (% ( E_{\text{ads}} ))</th>
<th>( d_1 ) (Å)</th>
<th>( d_2 ) (Å)</th>
<th>( \theta ) (degree)</th>
<th>( \phi ) (degree)</th>
</tr>
</thead>
<tbody>
<tr>
<td>NHC1</td>
<td>−1.96</td>
<td>−1.89</td>
<td>−1.03 (53%)</td>
<td>2.05</td>
<td>0.43</td>
<td>3</td>
</tr>
<tr>
<td>NHC2</td>
<td>−2.28</td>
<td>−2.16</td>
<td>−1.32 (58%)</td>
<td>2.05</td>
<td>0.52</td>
<td>7</td>
</tr>
<tr>
<td>NHC3</td>
<td>−2.58</td>
<td>−2.43</td>
<td>−1.62 (63%)</td>
<td>2.05</td>
<td>0.60</td>
<td>0</td>
</tr>
</tbody>
</table>

FIGURE 3  (A) Side-view of, for example, an Au adatom residing on the fcc hollow site of planar surface. Its height assuming no molecular adsorption is considered as a reference (\( d_3 \)). (B,C) side-views of the most stable structures of a N-heterocyclic carbene (NHC) (e.g., NHC2) adsorbed on an adatom (ballbot species) in an upright/a flat-laying geometry. The vertical upwards shift of the adatom after the NHC adsorption is denoted as \( d_3 \). (D) Is like (C) but in a top-view. The azimuthal orientation of the molecular symmetry axis (SA) with respect to the <110> direction of the surfaces is denoted as \( \phi \). The adsorption parameters of NHC1, NHC2, and NHC3 are given in Table 2 and 3 for Cu(111) and Au (111), respectively.
upright geometries are further stabilized by about 0.85, 0.80, and 0.67 eV for NHC1, NHC2, and NHC3, respectively (Figure 5A). This can be related to the fact that the metal s- and d-orbitals are more accessible in the ballbot species, enabling a stronger donor-acceptor bond.14 Thereby, the length of the formed C—Cu bond (about 1.93 Å) is apparently shorter than that for the surface-bound species (2.05 Å). This tendency is also reflected in the far smaller upwards shift of the adatom (d3) compared to that (d2) shown in Table 1 for the same NHCs in surface-bound species.

In parallel, the flat-laying ballbot geometries become possible and even most stable for NHC1 and NHC2 (see also Figure 5A). In fact, further 0.32 and 0.20 eV, respectively, are gained in comparison with upright adsorption on Cu@Cu(111). However, in case of NHC3, the upright adsorption remains favored by 0.17 eV. These results are fully in line with the measurements of Larrea et al.12 and can be understood in the light of a subtle balance between the covalent C—Cu bond on the one hand and the van der Waals (vdW) energy interactions between the molecule and planar surface on the other hand. Going from upright to flat-laying geometries yields a twist in the Cu—C bonds accompanied with energy-loss of 0.54, 0.53 and 1.05 eV, for NHC1, NHC2, and NHC3, respectively. This is reflected in the increased length of the C—Cu bond (d1) and upwards shifts Cu adatom (d3) in the latter. In parallel, it gives rise to higher vdW interactions for the respective species by 0.86, 0.73, and 0.88 eV. Without adatoms (i.e. for surface-bound species), the molecule (or at least parts of it) would be simply too close to the surface atoms, so that flat-laying geometries are not possible from sterical reasons. It is thus intuitively clear that the upright configuration provides the mobile version of any NHC ballbot; it will become dominating at elevated temperatures, where thermal fluctuations of the side groups decrease the influence of vdW interactions.

The results drawn for Cu(111) can be more or less transferred to the Au(111) surface, see Table 3. Compared to the surface-bound species, NHC ballbot species in the upright geometries are more stable by about 1 eV, in agreement with previous studies.14,16,29,30 The length of the formed C—Au bond (2.03 Å) is again reduced (2.15 Å in the surface-bound species). The calculated length of d1 is in a very good agreement with values presented in References [13, 14, 16, 23, 29]. A comparison between upright and flat-laying geometries indicates that the length of d1 is longer in the latter by about 2%. The tilting angles of the three NHCs on Au(111) show, however, a different tendency compared to Cu(111): The flat-laying molecules are completely parallel to the surface θ~90°, while the upright configurations exhibit different tilting angles of 6°, 18°, and 35° for NHC1, NHC2, and NHC3, respectively. Such unexpected geometries indicate that the size of the side groups plays a more decisive role. The molecular geometries are the result of complicated molecular-surface interactions and their detailed balance with respect to the size of the molecular side groups.

In addition, in this case the calculations are in agreement with the experimental observations.12 The adsorption energies of NHC1 and NHC2 (NHC3) in the flat-laying geometries are more stable (less stable) than the upright ones by 0.41 and 0.20 (0.09) eV (see Figure 5B). Notably, a weak dependence of the adsorption energies for NHC3 on the tilting angle was previously reported.14 In agreement with their near-edge X-ray absorption fine-structure spectroscopy (NEXAFS) measurements for NHC3 on Au(111), the calculations show that NHC3 adopt an intermediate tilting angle of θ = 35° nicely reflecting the experimentally derived value of about 40°.

3.2.2 | Bis(N-heterocyclic carbene) complexes on Cu(111) and Au(111)

Our calculations in the previous section demonstrate that single NHC ballbot species are energetically more favorable than surface-bound species. Another conceivable adsorption scenario is the formation of flat-laying bis(NHC) species, where both NHC molecules are bound to

![Figure 5](image-url)
the same surface atom (NHC-adatom-NHC, see Figure 4). For different small-sized NHCs (lMe and NHC1), such species have been previously reported experimentally on Au(111) as well as on Cu(111) surfaces, see for example, References [9, 10, 12, 13]. There are some indications that the formation of such complexes is promoted by thermal activation or by intermolecular interactions.12,14

Table 4 summarizes the calculated adsorption geometries and energies for the most stable bis(NHC) structures for NH1, NH2, and NH3 on both substrates. In all cases, the adsorption energies per molecules are comparable to those obtained for the ballbot species (cf. Tables 2 and 4). They are, thus, again much larger than those obtained for surface-bound NHC (see also Figure 5 for an overview): bis(NHC) dimer formation is energetically more favorable by 0.8 eV (NH3) to 1.3 eV (NH1, NH2) on Au(111) and 0.5–1 eV on Cu(111), respectively. This indicates that the formation of ballbot species, in particular in flat-laying geometry, provides a highly efficient precursor for the formation of bis(NHC) species, independent on the size of the molecular side group. On Au(111), the adsorption energy per molecule for the (NH3)Au dimers are even slightly higher (by 10–80 meV) than the adsorption energies of the single NHC flat-lying ballbot species. In other words, the (NH3)Au dimers are highly probable and expected to be formed even prior to a separate ballbot configuration at already present, but still ‘unoccupied’ adatoms.

On Cu (111), the situation is more complex. The adsorption energies per molecule for (NH3)Cu are slightly smaller than for the respective most stable ballbot species (cf. Figure 5A), suggesting that on Cu(111) the formation of dimers requires high NHC coverage. At least at low temperatures, all available Cu adatoms should be first decorated with a single NHC, before the formation of bis(NHC)Cu complexes sets in. We have to note, however, that further alternative formation scenarios exist and that dimer formation could be promoted kinetically at higher temperatures when the bridging Cu adatom is lifted and the potential (NH3)2Cu dimers become mobile. An alternative formation scenario for bis(NHC) dimers has been indeed suggested in Reference [13] and is confirmed by our present calculations: Starting from NHCs binding to the same surface atom, structural relaxation always

<table>
<thead>
<tr>
<th>$E_{\text{ads}}$ (eV)</th>
<th>$E_{\text{vdW}}$ (eV) (% of $E_{\text{ads}}$)</th>
<th>$d_1$ (Å)</th>
<th>$d_2$ (Å)</th>
<th>$\theta$ (degree)</th>
<th>$\phi$ (degree)</th>
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<tbody>
<tr>
<td><strong>Upright</strong></td>
<td><strong>Flat</strong></td>
<td><strong>Upright</strong></td>
<td><strong>Flat</strong></td>
<td><strong>Upright</strong></td>
<td><strong>Flat</strong></td>
</tr>
<tr>
<td>NHC1  &amp; -2.83 &amp; -3.24 &amp; -0.41 (15%) &amp; -1.28 (40%) &amp; 2.03 &amp; 2.07 &amp; 0.13 &amp; 0.10 &amp; 6 &amp; 89 &amp; - &amp; 30</td>
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<tr>
<td>NHC2  &amp; -3.15 &amp; -3.35 &amp; -0.70 (22%) &amp; -1.42 (43%) &amp; 2.03 &amp; 2.07 &amp; 0.11 &amp; 0.14 &amp; 18 &amp; 89 &amp; - &amp; 30</td>
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<tr>
<td>NHC3  &amp; -3.32 &amp; -3.23 &amp; -0.95 (29%) &amp; -1.62 (50%) &amp; 2.04 &amp; 2.07 &amp; 0.10 &amp; 0.15 &amp; 35 &amp; 89 &amp; - &amp; 30</td>
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**TABLE 3** Adsorption parameters of most stable structures of ballbot species of NHC1, NHC2, and NHC3 on Au(111). The definition of $d_1$, $d_3$, the angles $\theta$ and $\phi$, were presented in Table 2

<table>
<thead>
<tr>
<th>$E_{\text{ads}}$ (eV)</th>
<th>$E_{\text{vdW}}$ (eV) (% of $E_{\text{ads}}$)</th>
<th>$d_1$ (Å)</th>
<th>$d_2$ (Å)</th>
<th>$\theta$ (degree)</th>
<th>$\phi$ (degree)</th>
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<tbody>
<tr>
<td><strong>Cu(111)</strong></td>
<td><strong>Au(111)</strong></td>
<td><strong>Cu(111)</strong></td>
<td><strong>Au(111)</strong></td>
<td><strong>Cu(111)</strong></td>
<td><strong>Au(111)</strong></td>
</tr>
<tr>
<td>NHC1  &amp; -2.96 &amp; -3.25 &amp; -1.59 (54%) &amp; -1.32 (41%) &amp; 1.97 &amp; 2.05 &amp; 0.49 &amp; 1.37 &amp; 85 &amp; 90</td>
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<tr>
<td>NHC2  &amp; -3.22 &amp; -3.43 &amp; -1.80 (56%) &amp; -1.50 (44%) &amp; 1.96 &amp; 2.04 &amp; 0.45 &amp; 1.22 &amp; 85 &amp; 90</td>
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<tr>
<td>NHC3  &amp; -3.06 &amp; -3.25 &amp; -2.15 (70%) &amp; -1.70 (53%) &amp; 1.99 &amp; 2.06 &amp; 0.56 &amp; 1.39 &amp; 85 &amp; 90</td>
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</table>

**TABLE 4** Formation of (NH3)Cu [(NH3)Au] complexes from NHC1, NHC2, and NHC3 on Cu(111) (Au(111)) surfaces. The contributions of the vdW interactions ($E_{\text{vdW}}$) to the adsorption energy per molecule ($E_{\text{ads}}$) are also given. The adsorption geometry parameters ($d_1$, $d_3$, and $\theta$) have been defined in Figure 3 and Figure 4

**FIGURE 4** Side views of the most stable adsorption configuration of bis(N-heterocyclic carbenes (NHC)) complexes on Cu(111) and Au(111) surfaces: (A) (NHC1)2Cu and (B) (NHC3)2Au. The height of the adatom without molecular adsorption is considered as a reference ($d_0$). The definitions of $d_1$, $d_3$, and $\theta$ were presented in Figure 3
converges to geometries, where the metallic atom is pulled out of the surface, leaving a vacancy behind (see Supplementary movie). In case of Au(111), the surface atom is completely detached from the surface layer, resulting barrier-free in an almost planar, physisorbed NHC–Au–NHC dimer configuration, see also Figure 4B, where the central Au atom lay in plane with the two NHC molecules, in agreement with NEXAFS measurements reported in Reference [14].

On Cu(111) our calculations find that a similar planar configuration is less stable, by about 0.2 eV higher in energy than the most stable structure, where the Cu atom within the (NHC)2Cu dimer is still covalently bound to two surface atoms (bridging position, see also Figure 4A). The central Cu adatom resides below the two NHCs, hindering them to adsorb completely parallel to the surface ($\theta = 85^\circ$). Such result is in line with STM measurements reported in Reference [10], where the central Cu atom is observed to appear less bright than the two NHC molecules. This again contributes in rationalizing the different, strongly surface depending binding modes observed by Larrea et al.12

4 | CONCLUSIONS

In summary, we performed vdW-corrected DFT calculations on the adsorption of N-heterocyclic carbenes with different side groups on Au(111) and Cu(111) surfaces. Special emphasis was given to factors that determine flat-lying or upright bonding configurations. For this purpose, we performed a comprehensive analysis for three possible adsorption scenarios, namely the surface-bound, ballbot and bis(NHC) dimers. Based on the detailed understanding of the trends in the geometries, adsorption energies, and the vdW contributions, we provided an explanation for a recent experimental observation reporting different bonding modes of NHCs on Au(111) and Cu(111) surfaces.

In all studied structures, the calculations show that the adsorption energies increase by binding to the more reactive Cu(111) surface and by increasing the size of the side groups. In the surface-bound model, that is, NHCs directly bound to the ideal (111) surface, the molecules are calculated to be covalently anchored to both Au and Cu surfaces in an upright configurations exclusively. Flat-lying NHC, as observed experimentally for NHC with relatively small side groups (NHC1 and NHC2), require adsorption at adatoms, resulting in a first step in the so-called ballbot configuration. These ballbot species bind significantly stronger to both surfaces and provide a highly efficient precursor for the formation of bis(NHC) dimers. Here, the central metal atom is lifted; on Au(111) it is even fully removed from the surface, so that the (NHC)2Au complex forms a planar 2D molecular island which is purely physisorbed to the surface in a very efficient way. Consequently, the bis(NHC) complexes are found to be thermodynamically favorable on Au(111). On Cu(111), in contrast, they require thermal activation or high NHC coverages. Otherwise, single ballbot configuration are preferred.

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CONFlict OF INTEREST

The authors declare no conflict of interest.

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