Binding Behavior of Carbonmonoxide to Gold Atoms on Ag(001)

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Abstract
The adsorption behavior of single CO molecules at 4 K bound to Au adatoms on a Ag(001) metal surface is studied with scanning tunneling microscopy (STM) and inelastic electron tunneling spectroscopy (IETS). In contrast to earlier observations two different binding configurations are observed—one on top of a Au adatom and the other one adsorbed laterally to Au on Ag(001). Moreover, IETS reveals different low-energy vibrational energies for the two binding sites as compared to the one for a single CO molecule bound to Ag(001). Density functional theory (DFT) calculations of the adsorption energies, the diffusion barriers, and the vibrational frequencies of the CO molecule on the different binding sites rationalize the experimental findings.

Keywords CO · Au · Ag(001) · Adsorption · STM · STS · IETS · DFT

1 Introduction
One of the initial steps in a heterogeneous catalytic reaction is the binding of molecules to surfaces. Characterizing the chemical nature of the molecules as they bind to metal surfaces or model systems involving oxide supported metal nanoparticles is a key issue. Infrared spectroscopy among other techniques is the tool often used [1–7]. However, if characterization via ensemble averaging is not sufficient, and characterization at the individual molecule level is necessary, one has to resort to other techniques. A technique that has proven very useful in this respect is inelastic electron tunneling spectroscopy, IETS [8–11]. Here we study the adsorption of CO on or near Au atoms supported on a Ag(001) surface [12–16].

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The binding behaviour of CO, representing a prototype molecule, to various metal substrates has been intensively studied in the past. Nevertheless, it is still of interest to many applied experiments like the AgAu alloy catalysts for selective oxidation reactions [17–19].

Of considerable significance for the present investigation of the CO binding behaviour to individual Au atoms on Ag(001) are the reports concerning (I) the adsorption of CO on Ag(001) [16] and (II) the observation of metal carbonyl formation on individual Cu and Fe adatoms on Ag(110) [10, 14]. CO on Ag(001) is found to be very weakly bound and is still mobile at temperatures of 6 K [16].

In contrast to the CO adsorption behaviour observed so far for single adatoms and metal islands, where CO molecules are predominantly bound on top of the Au adatoms and to the rim of Au islands [20–22], we find two different configurations for a single Au adatom—one on top and the other one at the side of individual Au atoms on Ag(001). Moreover, IETS reveals different low-energy vibrational energies observed between tunneling voltages of 10 and 30 meV for the two adsorption sites of CO to Au as compared to the one at 16 meV for the single CO molecule bound to Ag(001). The latter value corresponds closely to the one reported previously [16], identified as the frustrated rotational mode of CO. DFT calculations of the adsorption energies, the diffusion barriers, and the vibrational frequencies of the CO molecule on the different adsorption sites contribute to a
quantitative insight into the CO adsorption behaviour on single Au adatoms.

2 Experimental and Computational Details

The experiments have been performed in a home-built ultra-high-vacuum (UHV) system equipped with a low-temperature scanning tunneling microscope (LT-STM) which operates at liquid He temperature (4 K). Moreover, low-energy electron diffraction (LEED), sample cleaning and gas dosing facilities are incorporated in the UHV chamber. The Ag(001) single crystal samples have been cleaned prior to Au deposition by several cycles of Ar sputtering at 800 eV and annealing at 673 K in UHV. The sample cleanliness has been checked with STM. The Au atoms are deposited in situ at 5 K by ohmic heating of a gold coated filament located close to the STM head. CO molecules are dosed in situ at 12 K through a gas line which is brought near the sample via a movable manipulator. STM and IETS measurements are performed at 4.2 K utilizing Pt–Ir tips. The differential conductance \( \frac{dI}{dV} \) was measured using lock-in detection with a 2 mVpp modulation at 6.56 kHz. The second derivative of the tunneling current has been obtained by numerical differentiation.

We have performed spin-polarized density functional theory (DFT) calculations as implemented in the GPAW package using a grid based basis set [23]. A grid spacing of 0.15 Å was used in all calculations. The electron–electron interactions were treated using the PBE exchange-correlation functionals [24], while the electron occupations were smeared using the Fermi–Dirac expression with a smearing width of 0.1 eV to aid convergence. To study the Ag(001) surface, we considered a 5 \( \times \) 5 surface unit cell with four atomic layers and at least 8 Å vacuum in the surface normal direction. For all calculations, the bottom two layers were fixed to bulk spacing while the top two layers and any adsorbate were allowed to optimize until forces on the atoms were less than 0.05 eV/Å. Brillouin zone sampling was done using a 3 \( \times \) 3 \( \times \) 1 k-point mesh for all calculations. The energetic barriers for the diffusion of a CO molecule in the presence of an adsorbed Au atom on Ag(001) surface were calculated using the climbing-image nudged elastic band method [25].

3 Results

3.1 Experimental Findings

Figure 1a shows a typical STM topographic image, obtained with a CO terminated tip (see Fig. S1 in the Supplementary Material and [12]), after deposition of 0.005 monolayer (ML) of Au and CO onto a clean Ag(001) surface at 5 K. CO
molecules appear as dark round depressions at a bias voltage below 100 mV [12, 26, 27]. The bright protrusions indicate the presence of the gold adatoms [20] with an apparent height of around 1.5 Å at a bias voltage of 40 mV. They are distributed randomly over the surface. Moreover, alongside of three Au adatoms, dark depressions appear which are assumed to reflect the adsorbed CO molecules to the Au adatoms. In addition, two Au adatoms exhibit a halo which reflects the adsorption of a CO molecule on top of the Au adatom. Figure 1b illustrates these observations in a simplified graphic with CO molecules (black) and Au adatoms (yellow) arranged according to their respective experimentally observed positions and contrasts shown in Fig. 1a. The crosses indicate the location of the STM tip during the IET spectroscopic measurements the results of which are displayed in Fig. 1c.

In addition to Au atoms with single co-adsorbed CO molecules, Au adatoms with two, three, and four laterally adsorbed CO molecules are observed as depicted in Fig. 2. Figure 2a–c show the corresponding topographic images and Fig. 2d–f highlight the positions of the adsorbed Au adatoms and of the CO molecules with yellow and red circles, respectively. The distance between the centers of the red and yellow circles are found to be 6.0 ± 0.3 Å. The angle between the line connecting these centers with respect to the nearest high symmetry orientation vectors of the Ag(001) substrate are found to be only either plus or minus 14° ± 1°, unless the number of co-adsorbed CO molecules is higher than four, which may happen with increasing CO coverage (see Supplementary Material Fig. S2).

Figure 1c summarizes the results of the IET spectroscopic measurements for the different measuring positions indicated in Fig. 1a, b. Figure 1c shows from bottom to top the IET spectra obtained on the clean Ag(001) surface (black, position indicated by the cross in the upper left of Fig. 1b), on a single Au adatom (yellow), on a Au adatom with a co-adsorbed CO molecule (red), on a Au adatom with a CO molecule on top of it (magenta), and on a single CO molecule on Ag(100) (blue). The spectra have been vertically displaced to provide better mutual distinction. They represent the second derivative of the average tunneling conductance dI/dV, obtained by summing up the signals of a voltage scan from −40 to +40 mV and vice versa, and subsequently applying a 5 point binomial smoothing.

The IET spectrum for the clean Ag(001) surface (bottom spectrum, black curve) is featureless, pointing to a metallic tip without a CO functionalization. With a CO functionalized tip the low-lying vibrational excitation observed for CO on Ag(001) (top spectrum, blue line) is present [16] (spectrum not shown). As expected, the IET spectrum obtained on top of a Au adatom (yellow) is also featureless within the chosen bias voltage range. However, the IET spectrum for a CO molecule adsorbed laterally to a Au adatom (red) exhibits a vibrational excitation around 30 meV, whereas the spectrum recorded on a Au adatom with a CO molecule adsorbed on top of it (magenta) shows a prominent vibrational excitation around 10 meV. Finally, for CO adsorption on the clean Ag(001) surface the IET spectrum (blue) displays a vibrational excitation around 16 meV. A comparison of these observed vibrational energies with the values and their interpretation in terms of the vibrational modes [internal stretching (IS), metal–C stretching (MC), frustrated rotation (FR), frustrated translation (FT)], reported in the existing literature, is presented in Table 1.

The value of 16 meV for the FR mode of CO on Ag(001) is fully in line with the value reported in the literature [16]. The value for the FR mode of CO atop of a single Au on Ag(001) found in our experiment (around 10 meV) is nearest to the case of CO on single Fe atoms on Ag(110) where the CO molecule on top of the Fe atom is believed to have a slightly tilted on top configuration [14]. A qualitatively similar redshift to lower energy with respect to the case of CO on Ag(001) can be observed for the IS mode for CO adsorbed to a single Au atom on a

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**Fig. 2** STM measurements showing adsorption sites of CO at the side of Au atoms on Ag(001). STM tunneling parameters: a, d, I_T = 100 pA, V_S = 30 mV, and b, c, e, f: I_T = 200 pA, V_S = 40 mV, image size all 2.0 nm × 2.0 nm. Yellow and red circles highlight the positions of the adsorbed Au adatoms and of the CO molecules, respectively.
In all other cases presented here from literature the FR mode values are shifted to higher values—these values resemble the one found in our experiment for the case of CO adsorbing to the side of a Au atom on Ag(001).

3.2 Computational Analysis

Density functional theory (DFT) calculations have been performed to rationalize the experimental findings concerning the energetically preferred adsorption site of Au on a Ag(001) surface, the adsorption site of CO on Ag(001),

Table 1 Vibrational modes of CO (meV), depending on local environment: internal stretching (IS), metal–C stretching (MC), frustrated rotation (FR), frustrated translation (FT), as reported in the existing literature

| Vibrational modes of CO (meV) | IS    | MC | FR | FT | References |
|-------------------------------|-------|----|----|----|------------|
| Experimental                  |       |    |    |    |            |
| Gas phase (1)                 | 266   |    |    |    | [5]        |
| On Au(110)                    |       |    |    |    | [28]       |
| On Ag(100)                    | 263   | 33 | 16 | 5  | [16]       |
| On Ag(100) (2)                | 256   | 51 |    |    | [29]       |
| On Ag(110)                    | 31    |    | 8  |    | [10]       |
| On Pd(110) (0.1ML CO) (2)     | 263   | 46 |    |    | [30]       |
| On Pd(110) (1.0ML CO) (2)     | 248   | 42 | 9  |    | [31]       |
| On Ni(110) (2)                | 48    | 8  |    |    | [32]       |
| On Cu(100)                    | 256   | 43 | 36 | 4  | [13, 33, 34]|
| On Cu(100)                    | 35    | 5  |    |    | [35]       |
| On Cu(110)                    | 259   |    |    |    | [36]       |
| On Cu(110)                    | 236   | 3–4 | 3  |    | [37, 38]   |
| On Cu(111)                    | 241   | 35 | 5  |    | [39]       |
| On Fe(110) (2)                | 234   | 57 |    |    | [40]       |
| On single Au on NiAl(110)     |       |    |    |    | [41]       |
| On single Pd on NiAl(110)     |       |    |    |    | [42]       |
| On single Au on MgO/Ag(100) (1) | 230  |    |    |    | [5]        |
| On Au clusters on MgO/Ag(100) (1) | 263  |    |    |    | [5]        |
| On single Fe on Ag(110)       | 236   | 10 | 4  |    | [14]       |
| On single Cu on Ag(110)       | 262   | 36(32) | 5   |    | [14]       |
| On single Au on Ag(100)       | 11    | 2  |    |    | This publication |
| At the side of single Au on Ag(100) | 30  | 5  |    |    | This publication |
| Theoretical                   |       |    |    |    |            |
| On Cu(100)                    | 249   | 47 | 2.8|    | [43]       |
| On Cu(100)                    | 245   | 30(29) |    |    | [35]       |
| On Cu(110)                    | 261   | 50 | 38(36) | 8(7) | [44]       |
| On Cu(111)                    | 246   | 47 | 33 | 2–6 | [45]       |
| On Ag(100)                    | 254   | 27 | 17 | 4  | [16]       |
| On Ag(100)                    | 251   | 30 | 16–19 | 3  | [43]       |
| On Ag(110)                    | 262   | 33 | 29(26) | 9(8) | [44]       |
| Gas phase                     | 266   |    |    |    | This publication |
| On Ag(100)                    | 253   | 28 | 19, 23 |    | This publication |
| On single Au on Ag(100)       | 248   | 36 | 11, 19 |    | This publication |
| On Ag(100) 2NN atop at the side of single Au | 252  | 30 | 20, 29 |    | This publication |
| On Ag(100) 3NN bridge at the side of single Au | 240  | 27 | 20, 25 |    | This publication |

(1) Estimated from IR measurements. Units translated from cm\(^{-1}\) to meV (1 meV = 8.065 cm\(^{-1}\)). (2) Estimated from HREELS measurements. (3) Estimated from He atom scattering measurements, vibrational mode along the [001][1\(\bar{1}0\)] direction. (4) Estimated from the drawn curve in the figure of the cited paper. The value is not mentioned within the text of the cited paper. (5) Value in parentheses from the negative bias range of IETS measurements. (6) Vibrational mode along the [001][1\(\bar{1}0\)] direction.
the co-adsorption of a Au adatom and a CO molecule on Ag(001), the vibrational frequencies of different adsorption sites, and the diffusion of CO in the presence of a Au adatom on Ag(001).

The lattice constant for bulk Ag was calculated to be 4.17 Å in reasonably good agreement with the experimental value of 4.09 Å [46]. A Au adatom prefers to bind to the hollow site on the Ag(001) surface with a binding energy of −2.93 eV (calculated with respect to the bare Ag surface and an isolated Au atom in gas phase). The adsorption at bridge and atop sites were found to be less stable with binding energies of −2.28 eV and −1.92 eV, respectively.

A CO molecule, on the other hand, prefers to bind to the atop site on an Ag surface with a binding energy of −0.26 eV (calculated with respect to the bare Ag surface and an isolated CO molecule in gas phase). The adsorption at the bridge site is almost degenerate in energy with that at the atop site with binding energy of −0.25 eV, while the adsorption at the hollow site is least preferred with a binding energy of −0.19 eV. For the CO molecule adsorbed at the atop site on the Ag surface, we calculate vibrational frequencies corresponding to the IS mode (253 meV), MC stretching mode (28 meV) and two possible FR modes (19 meV and 23 meV). In agreement with the experiments, a red shift to lower energy is observed in the IS mode of CO, on going from gas phase (266 meV) to the adsorbed state (253 meV).

Figure 3 shows five adsorption sites for a CO molecule and their respective binding energies on the Ag(001) surface in the presence of an adsorbed Au adatom, and the energy profile for the diffusion of the CO molecule between these sites. The binding energies (in black) are calculated with respect to an isolated CO molecule in gas phase and a Au adatom adsorbed (at the hollow site) on Ag(001). The numbers in black are according to the ordinate scale, while the numbers in red are the energy barriers for the forward process for every individual diffusion step.

The binding energies suggest that sites around Au, where CO is in fourth nearest neighbor (4NN), third nearest neighbor (3NN), or second nearest neighbor (2NN) positions, are degenerate in energy and are comparable to the binding energy of CO on a bare Ag(001) surface (−0.26 eV), i.e., the CO molecules at these sites do not feel the presence of the Au adatom. The adsorption of CO at the first nearest neighbor (1NN) atop Ag position is less stable with a binding energy of −0.18 eV. The experimental STM measurements do not show the presence of CO molecules on the Ag surface closer to Au than the 4NN bridge site position.

However, the different experimental vibrational spectra as well as the corresponding DFT calculations [for CO on Ag(001): calculated values (meV) = 19, 23 and for CO on Ag(001) near a Au adatom: calculated values 20, 25 and 29 meV] underline that the dynamical properties, i.e., the vibrational properties of the CO molecule are slightly different for the CO on Ag(001) and the CO near a Au adatom on Ag(001). This fact indicates that the Au adatom with its contribution to the local density of states is somehow in contact with the CO molecule. In a similar way, the interaction of CO molecules with surface state electrons on Ag(111) has revealed a preferred adsorption at 5 K near the minima of the standing wave pattern [47].

![Fig. 3](image_url) Energy profile for the diffusion (green/blue arrows) of CO on the Ag(001) surface in the presence of an adsorbed Au adatom. The energy scale is referenced to the sum of the total energies of a CO molecule in gas phase and Au/Ag(001) surface, i.e. \( E_{CO} + E_{Au/Ag(001)} \). Numbers in red represent the energy barriers for individual diffusion steps in the forward direction. The atoms C, O and Au are shown in purple, red and yellow colors, respectively. The Ag atoms on the top and second layers of the surface are shown in light and dark gray colors, respectively. We show the diffusion of CO from the fourth nearest neighbor (4NN) bridge site to the third nearest neighbor (3NN) bridge site to the second nearest neighbor (2NN) atop site. From the 2NN atop Ag site it can diffuse to the atop Au position, directly, or via a first nearest neighbor (1NN) atop Ag site. All neighboring positions are counted with respect to the position of the adsorbed Au atom.
Additional electronic multimedia files (mpg) showing the frustrated rotation vibrational modes and frequencies for the CO molecule adsorbed (i) on the bare Ag surface, (ii) on top of an adsorbed Au atom on the Ag surface and (iii) on the side of an adsorbed Au atom on the Ag surface, are given in the Supplementary Material. The energy barriers for the diffusion of CO over the Ag surface from 4NN bridge site to 3NN bridge site to 2NN atop site are very low (0.03 to 0.04 eV). The CO at the 2NN atop site can diffuse to the most energetically preferred adsorption site, atop Au position via two pathways, viz., (i) 2NN atop Ag → atop Au with a barrier of 0.25 eV or (ii) 2NN atop Ag → 1NN atop Ag → atop Au with barriers 0.20 eV and 0.06 eV, respectively. This suggests that the CO molecules have to overcome a high barrier in order to reach the atop Au position. This may not be possible at the low temperatures of the experiment. We suggest that the direct adsorption by gas-phase deposition or tip assisted diffusion of CO [48] are two possible explanations for the existence of the CO molecules on top of Au/Ag(001).

4 Discussion

The very low-diffusion barriers for CO on Ag(001) together with tip-assisted diffusion are most probably the reason for the experimentally observed agglomeration of CO molecules around Au adatoms, as shown in Fig. 2. The calculated vibrational frequencies for CO adsorbed on bare Ag and that adsorbed to the side of Au are almost the same, i.e., DFT does not find a difference between these two sites, which is also evidenced by the results of the binding energies for the two sites. The vibrational frequencies of CO on top of Au are different. Our calculations predict for the three cases [CO on Ag, CO on Au(top)/Ag, and CO on Au(side)/Ag] 8 distinct IS, MC, and FR vibrational modes. In the experiment, however, only one mode (MC or FR, see Table 1) is observed for each adsorbate configuration.

The IETS selection rules depend strongly on the local density of states induced by the adsorbates at the Fermi level and on the local symmetry [49]. We speculate that in the CO/Ag case only the FR mode at 19 meV is observed, in the case of CO adsorbed on top of a Au atom only the 11 meV FR mode is seen, and on CO adsorbed at the side of Au the 30 meV (MC) and/or 29 meV (FR) modes are present. With these assumptions a nearly quantitative correspondence between measured and calculated vibrational frequencies is obtained.

5 Conclusions

To summarize, the binding behaviour of a CO molecule to a Au adatom on a Ag(001) surface has been investigated by low-temperature STM and IETS. In addition to the binding site of the CO molecule on bare Ag(001), two different binding configurations of the CO molecule have been identified, one atop of a Au adatom and another one laterally adsorbed to a Au adatom. Both configurations exhibit different frustrated rotational vibrational frequencies detected with local IET spectroscopy. The measured vibrational frequency for CO/Ag(001) confirms earlier results, reported in Table 1. Theoretical modeling with DFT concerning the adsorption energies, the diffusion barriers, and the vibrational frequencies of the CO molecule on the different binding sites rationalizes the experimental findings. The chemical verification and identification of atomic species by IETS is a very powerful tool and although first measurements have already been shown by Stipe et al. [8] more than 20 years ago, this method is still not fully established and should be applied to many more examples. The present study enlarges our knowledge about the characteristics of molecular binding sites at the atomic level.

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Conflict of Ethical Standards

Conflict of interest The authors declare that they have no conflict of interest.

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References

1. Bäumer M, Freund HJ (1999) Prog Surf Sci 61(7):127
2. Haruta M (1997) Catal Today 36(1):153
3. Valden M, Lai X, Goodman DW (1998) Science 281(5383):1647
4. Sanchez A, Abbet S, Heiz U, Schneider WD, Häkkinen H, Barnett RN, Landman U (1999) J Phys Chem A 103(48):9573
5. Sterrer M, Yulikov M, Risse T, Freund HJ, Carrasco I, Illas F, Di Valentini C, Giordano L, Pacchioni G (2006) Angew Chem Int Ed 45(16):2633
6. Risse T, Shaikhutdinov S, Nilius N, Sterrer M, Freund HJ (2008) Acc Chem Res 41(8):949
7. Schneider WD, Heyde M, Freund HJ (2018) Chem Eur J 24(10):2317
8. Stipe BC, Rezaei MA, Ho W (1998) Science 280(5370):1732
9. Stipe BC, Rezaei MA, Ho W (1999) Phys Rev Lett 82(8):1724
10. Ho W (2002) J Chem Phys 117(24):11033
11. Reed MA (2008) Mater Today 11(11):46
12. Bartels L, Meyer G, Rieder KH (1997) Appl Phys Lett 71(2):213
13. Lauhon LJ, Ho W (1999) Phys Rev B 60:R8525
14. Lee H, Ho W (2000) Phys Rev B 61(24):R16347
15. Heinrich AJ, Lutz CP, Gupta JA, Eigler DM (2002) Science 298(5597):1381
16. Arafune R, Shin J, Jung J, Minamitani E, Takagi N, Kim Y, Kawai M (2012) Langmuir 28(37):13249
17. Shirman T, Lattimer J, Luneau M, Shirman E, Reece C, Aizenberg M, Madix RJ, Aizenberg J, Friend CM (2018) Chem Eur J 24(8):1833
18. Pia G, Sogne E, Falqui A, Delogu F (2018) Sci Rep 8(1):15208
19. Zugic B, van Spronser MA, Heine C, Montemore MM, Li Y, Zakharov DN, Karakalos S, Lechner BA, Crumlin E, Biener MM, Frenkel AI, Biener J, Stach EA, Salmeron MB, Kaxiras E, Madix RJ, Friend CM (2019) J Catal 380:40
20. Yang B, Lin X, Gao HJ, Nilius N, Freund HJ (2010) J Phys Chem C 114(19):8997
21. Wallis TM, Nilius N, Ho W (2016) J Phys Chem Lett 7(22):4683
22. Nilius N, Risse T, Sterrer M, Shaikhutdinov S, Freund HJ (2014) Structure and bonding, vol 162. Springer, Cham, p 91
23. Enkovaara J, Rostgaard C, Mortensen JJ, Chen J, Dulak M, Ferrighi L, Gavnholt J, Glansad C, Haikola V, Hansen HA, Kristoffersen HH, Kaisma M, Larsen AH, Lehtoavaara L, Ljungberg M, Lopez-Acevedo O, Moses PG, Ojanen J, Olsen T, Petzold V, Romero NA, Stausholm-Moller J, Strange M, Tritsaris GA, Vanin M, Walter M, Hammer B, Häkkinen H, Madsen GKH, Nieminen RM, Norskov JK, Puska M, Rantala TT, Schiotz J, Thygensen KS, Jacobsen KW (2010) J Phys Condens Matter 22(25):253202
24. Perdew JP, Burke K, Ernzerhof M (1996) Phys Rev Lett 77:3865
25. Henkelman G, Uberuaga BP, Jónsson H (2000) J Chem Phys 113:9901
26. Niemi E, Nieminen J (2004) Chem Phys Lett 379(1):200
27. Nieminen JA, Niemi E, Rieder KH (2004) Surf Sci 552(1):L47
28. Xu C, Chiang CL, Han Z, Ho W (2016) Phys Rev Lett 116:166101
29. Burghaus U, Vattuone L, Gambardella P, Rocca M (1997) Surf Sci 374(1–3):1
30. Oh J, Lim H, Arafune R, Jung J, Kawai M, Kim Y (2016) Phys Rev Lett 116:056101
31. Kato H, Kawai M, Yoshinobu J (1999) Phys Rev Lett 82:1899
32. Voigtländer B, Bruchmann D, Lehwald S, Ibach H (1990) Surf Sci 225(1):151
33. Andersson S (1979) Surf. Sci. 89(1):477
34. Arafune R, Hayashi K, Ueda S, Uehara Y, Ushioda S (2006) Surf Sci 600(18):3536
35. Pascual JJ (2005) Eur Phys J D 35(2):327
36. Woodruff D, Hayden B, Prince K, Bradshaw A (1982) Surf Sci 123(2):397
37. Lee HJ, Ho W (1999) Science 286(5445):1719
38. Braun J, Weckesser J, Ahner J, Mocuta D, Yates JT, Wöll C (1998) J Chem Phys 108(13):5161
39. Vitali L, Ohmann R, Kern K, Garcia-Lekue A, Frederiksen T, Sanchez-Portal D, Arnau A (2010) Nano Lett 10(2):657
40. Erley W (1981) J Vac Sci Technol 18(2):472
41. Wallis TM, Nilius N, Ho W (2003) J Chem Phys 119(4):2296
42. Nilius N, Wallis TM, Ho W (2002) J Chem Phys 117(24):10947
43. Ortigoza MA, Rahman TS, Heid R, Bohnen KP (2010) J Phys Condens Matter 22:39
44. Lorente N, Ueba H (2005) Eur Phys J D 35(2):341
45. Rossen ETR, Flipse CFJ, Cerdá J (2013) Phys Rev B 87:235412
46. Kittel C (1953) Introduction to solid state physics. Wiley, Hoboken
47. Kulawik M, Rust HP, Heyde M, Nilius N, Mantooth B, Weiss P, Freund HJ (2005) Surf. Sci. 590(2):253
48. Li J, Berndt R, Schneider WD (1996) Phys Rev Lett 76(11):1888
49. Lorente N, Persson M, Lauhon LJ, Ho W (2001) Phys Rev Lett 86(12):2593

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