Bridge-bonded methylthiolate on Au(111) observed with the scanning tunneling microscope

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We report the discovery of bridge-bonded methylthiolate, SCH₃, along the step edges of the Au(111) surface. Real-space imaging with a scanning tunneling microscope reveals the presence of bridge-bonded SCH₃ along both the [110] and the [112] oriented step edges. The nearest neighbour distances of SCH₃ along these steps are 2a and √3a, respectively. The Au(111) terrace is covered with the usual CH₃SAuSCH₃ staples. The bridge-bonded alkanethiolate is expected to play a rather significant role in the formation of thiol-passivated Au nanoclusters because of the high fraction of atoms in similar low-coordination sites.

Introduction

Alkanethiol passivated gold surfaces and nanoclusters lead to a rich variety of surface reconstructions and, in the latter case, size-dependent electronic and optical properties.¹⁻⁴ Due to their very small sizes, the determination of the structure of these nanoclusters has been a great challenge.⁵⁻¹² Recent studies demonstrate that even for the magic number Auₙ(SR)ₙ clusters, there is structural polymorphism.⁷,¹² In contrast to naked Au nanoclusters, thiolate passivation does not just protect the clusters from interacting with other clusters or the surroundings, but also directly influences the structure of the clusters via the S–Au bonding at the interface.¹³ The current view on the S–Au bonding at the thiolate–Au interface is based on the RS–Au–SR staple motif which has been found on both the surface of Au nanoclusters⁶ using X-ray diffraction and extended Au(111) surfaces by direct imaging with a scanning tunneling microscope (STM).¹⁴⁻¹⁸ Variations of the RS–Au–SR staple motif, such as the dimeric¹³ and trimeric¹⁹ staple motifs, have also been introduced to explain the structure of passivated Au nanoclusters. An interesting, but less conventional, structural motif containing SR bridge-bonded to surface Au atoms was reported recently.¹¹ This bridge-bonded SR, if existing on the surface of Au nanoclusters, is expected to have a great impact on our understanding of the thiolate–Au interface. Herein we report the discovery of bridge-bonded SR on the extended Au(111) surface by imaging with the STM. The capability of STM to perform real space imaging with atomic resolution makes it a useful complementary technique to diffraction analysis, providing important local structural information at the atomic level.

Methods

A (111)-oriented Au single crystal is used for the experiment. The sample is cleaned by standard ion sputtering and thermal annealing cycles. The clean Au(111) is then exposed to dimethyl-disulfide (DMDS) at room temperature to reach full surface coverage. This is followed by thermal annealing at 350 K to desorb some molecules so that the remaining molecules form the striped phase.¹⁸ After cooling to 77 K, the sample is imaged with an Omicron low temperature STM (LT-STM) with an electrochemically etched tungsten tip.

Results and discussion

Fig. 1 shows an STM image of the Au(111) sample with ~0.2 monolayers (ML) of methylthiolate. On the flat (111) terraces, we find the usual rows of CH₃S–Au–SCH₃ staples aligned in one of the (112) directions.¹⁸ For each CH₃S–Au–SCH₃ staple, high-resolution STM images show three protrusions: two from the CH₃ groups and one from the Au adatom¹⁵,¹¹ with S not visible under normal imaging conditions. The inset in Fig. 1, for example, shows a row of four CH₃S–Au–SCH₃ staples. The Au adatom becomes less visible for thiols with longer alkane chains.¹⁹,²⁰ The distance between the two staples along the row in the [112] direction is √3a. The distance between the two methyl groups
within the same staple, measured to be 0.66 nm, is a pure property of the staple, and has no specific relation to the lattice constant of gold. The occurrence of paired-spots, representing the two arms of RS–Au–SR, is a distinctive feature for the presence of RS–Au–SR staples on Au(111). In addition to the standard CH$_3$S–Au–SCH$_3$ staples found on the terraces, we observe regularly-spaced spots along the step edges, organized in a different manner from those associated with the staples. In Fig. 1, two sections of the [110] oriented step edges are highlighted with blue rectangular boxes. These steps consist of close-packed Au atoms. Fig. 2a shows a faceted step edge consisting of segments of [110] steps. Fig. 2b presents a height profile taken along the double-headed arrow, A–B. This allows an accurate measurement of the distance between the nearest neighboring spots along the step. The measured distance is 0.58 ± 0.01 nm which is significantly smaller than the distance (0.66 nm) between the two methyl groups within one staple. Therefore, the spots along the [110] step are not paired. They are individual spots equally spaced along the step. 0.58 ± 0.01 nm is consistent with 2$\alpha$ where $\alpha$ is the distance between the nearest neighboring Au atoms. Thus, we conclude that along the [110] step there is one spot for every two Au atoms.

There are not many detailed studies on the organization of alkanethiolate along atomic steps on Au(111). Step decoration by methylthiolate on Au(111) has been reported on two previous occasions$^{17,21}$ with no structural model proposed. In an STM study of ethylthiolate on Au(111), Clair et al. reported the decoration of the [110] step edges by single thiolate rows.$^{22}$ The single row of ethylthiolate reported by Clair et al. was assumed to have a similar bonding configuration to that occurring in a staple. However, the remarkable similarity between the ethylthiolate rows observed by Clair et al. and the methylthiolate rows shown in Fig. 1 and 2 suggests that they are the same type of species. The nearest neighbor distance between the ethylthiolates along the [110] step is 0.55 nm$^{22}$ which is also very close to the distance (0.58 nm) measured in this study for the methylthiolates allowing for measurement uncertainties. The distance measured in our study is calibrated against the distance (0.5 nm) between staples along a staple row and it has an uncertainty of ±0.01 nm.

In Fig. 2c, we show a [112] oriented step. There is a CH$_3$S–Au–SCH$_3$ staple row sitting on the upper terrace at this step edge, along the direction P–Q. This staple row has the same appearance as all other staple rows on the terrace. Interestingly, one can see a single row of spots, M–N, right next to the staple row, P–Q. A few circles are drawn onto the image to assist the eye to locate this single row of spots. Again, here we have a row of spots showing no signs of pairing. These spots along M–N are regularly spaced at a distance of 0.50 ± 0.01 nm, which corresponds to $\sqrt{3}$a. The spots along M–N appear 0.07 ± 0.01 nm lower than the methyl groups along the P–Q staple row, and are 0.16 ± 0.01 nm higher than the methyl groups on the lower terrace. This gives further evidence that the spots along the step edges are different from those found either on the upper or the lower terraces. For the spots along both the [110] and [112] step edges, the distance between the neighboring spots is determined by the geometry of the gold substrate: 2a along the [110] direction and $\sqrt{3}$a along the [110] direction. Based on these observations and the above discussion, we come to the conclusion that each spot along the step edge corresponds to a single SCH$_3$ species bonded to Au atoms along the step, and there is...
no association of two adjacent SCH₃ groups such as those found within a staple.

In Fig. 3, we present a structural model showing the proposed bonding of the SCH₃ group to Au atoms at steps. Considering the nearest neighbor distances and the likelihood that the SCH₃ group would occupy a high symmetry site, there are only two possible bonding sites for SCH₃ along the [110] step: monodentate bonding with S bound to a single Au atom or bidentate bonding with S bridging between two Au atoms. Adsorption energetics suggests that the bidentate form is much more favored.²³ Along the [110] step, the shortest distance possible between two bridge-bonded SCH₃ groups is thus 2a, as 1 a is too short a distance to accommodate two neighboring methyl groups. This bonding scheme is similar to the bonding of SCH₃ within the staple, where S interacts directly with two gold atoms: one is the Au adatom in the staple and the other the Au atom within the top layer of Au(111). Along the [110] step, the S atom is bonded to two Au atoms in identical positions and the S-C bond is expected to be perpendicular to the step edge. This makes the CH₃ group along the step appear at a height between the methyl groups on the upper terrace and those on the lower terrace.

On the (111) plane of fcc metals, there are two types of [110] oriented steps: (i) a step presenting a {111} microfacet and (ii) a step presenting a {100} microfacet.²⁴ The two types of steps are energetically different from the {111}-faceted step having lower energy. This energy difference is expected to influence molecular adsorption. We do not know which type of step is the one observed in Fig. 1, although the two [110] steps in the figure belong to the same type. It would be of interest to find out if thiolate interacts differently with the two types of steps. In a previous study using a zero-gradient stepped surface, we have shown that the adsorption of C₆₆ molecules on Au(111) exhibits clear step type dependence.²⁵

The [112] step has a zig-zag structure exposing micro-faceted [110] steps. For the reason as discussed above, we have alternating {111}- and {100}-faceted steps along the [112] direction. Each {111}- or {100}-faceted step is only two-atoms long, The SCH₃ group hence has a choice between the two types of steps. In Fig. 3, we have arbitrarily put SCH₃ onto one of the bridging sites. As can be seen in Fig. 3, the horizontal step at the bottom of the diagram is a [112] oriented step. Along this step, SCH₃ can find identical bridge-bonding sites to those presented by a [110] oriented step. Applying the same bridge-bonding scheme for the [110] step, we can generate a row of SCH₃ parallel to the [112] direction with the nearest neighboring SCH₃ groups separated by √3a. The model in Fig. 3 is in good agreement with what is observed in STM images. For example, the position of SCH₃ attached to the [112] step relative to the position of the first staple row on the upper terrace is correctly given by the model.

From the STM images, we find that the step edges are almost fully covered with thiolate while the coverage on the terrace is lower. This suggests preferential binding of thiolate at step edges in competition with the formation of the RS–Au–SR staple on terraces. Indeed, the bridge-bonded species have a higher thermal stability than the staples. We find that at 210 K when the staple rows on the terraces are fully disintegrated to form a liquid-like layer, the bridge-bonded species along the steps remain static. This finding is important when one attempts to explain the formation of staples that require the supply of Au adatoms. Since adatoms on surfaces usually come from steps, the passivation of steps by bridge-bonded thiolate may play a key role in the formation of SAMs on Au(111). At RT, no ordered CH₃S–AuSCH₃ staple rows are observed on Au(111). Instead, the STM image shows a disordered layer indicating rapid movement of adsorbed species. CH₃S–AuSCH₃ can break apart into CH₃S–Au and SCH₃ or 2SCH₃ and Au. There is a possibility that when SCH₃ moves close to a step already occupied by bridge-bonded thiolate, it may combine with a thiolate at the step to form CH₃S–AuSCH₃ which moves away from the step into the terrace. If the temperature is not sufficiently high, the above proposed reaction step does not occur and the step is protected from being etched. For alkanethiol molecules with long alkane chains, the diffusion of the rather bulky molecules at RT is slow on Au(111). Thus, Au atoms might get extracted locally from the flat atomic terraces and hence etch pits are formed. For short chain alkanethiols deposited onto Au(111) using their vapors under vacuum, etch pits in general do not appear. However, etch pits can form if the Au(111) is dipped into a liquid solution of short chain thiols. This is because the extraction of Au atoms from steps is too slow in comparison to the uptake of the thiol molecules in such an environment.

Bridge-bonded thiolate is likely to be a more common species considering that there is now evidence for the presence of both methythiolate and ethylthiolate on Au(111), as well as on the surface of gold nano-particles.¹¹ In the following, we provide some additional information on the assembly of thiolate in the vicinity of steps on Au(111). Data presented below are from the Au(111) surface fully covered with a layer of mixed methythiolate and propylthiolate. The sample was prepared by exposing a (111) oriented Au film to methyl-propyl-disulfide, CH₃S–S(CH₂)₄CH₃, at RT with STM imaging also conducted at RT. No thermal annealing was conducted between deposition
and imaging. The initial motivation for studying the mixed molecular layer came from our earlier work with methyl- and ethyl-thioliate on Au(111),\textsuperscript{17} where we demonstrated that both methyl- and ethyl-thioliate form a $3 \times 4$ phase at saturation coverage, instead of the well-documented ($\sqrt{3} \times \sqrt{3}$) – R30° phase. By depositing CH$_3$S–S(CH$_2$)$_2$CH$_3$ onto Au(111), we were hoping to observe domain-separated $3 \times 4$ methylthiolate and ($\sqrt{3} \times \sqrt{3}$) – R30° propylthiolate on the same sample. But it was only discovered later that propylthiolate also forms the $3 \times 4$ phase, albeit with a lower degree of crystallinity.\textsuperscript{20}

Methylthiolate and propylthiolate tend to phase separate on Au(111), but the phase separation is not extensive and there are regions of the sample having a mixture of both.\textsuperscript{26} Whilst both ethyl- and propyl-thioliate monolayers are stable at RT,\textsuperscript{19,20} the mixed methyl- and propyl-thioliate layer is rather fluidic with constant molecular movement. This is mainly due to the contribution of the methylthiolate. Moving from methylthiolate to ethyl and propylthiolate, the increase of the alkane chain length offers some extra stability to the adsorbate due to the contribution of the van der Waals interaction. In addition to the van der Waals interaction, there is also a possibility that the methylthiolate staple is intrinsically less stable. The reason for nonobservance of a stable structure at RT for a full layer of methylthiolate on Au(111) is likely that the CH$_3$S–S(CH$_2$)$_2$CH$_3$ staple breaks down to (CH$_3$)$_2$S–Au–S(CH$_3$) (a negative-ion-like species) and Au. We expect an abrupt change in the stability of the negative-ion-like species when the number of carbon in the thiolate increases from one to two or more. A relatively stable adsorbate structure is found along step edges as shown in Fig. 4 which is an STM image acquired at RT from an area enriched with propylthiolate. Due to the mixing of methylthiolate and propylthiolate on the surface, it is not easy to identify which feature is due to methylthiolate and which is due to propylthiolate. However, a methylthiolate-rich region is expected to be completely disordered at RT. Since relatively stable staple rows are observed in Fig. 4, we believe that this region consists of mainly propylthiolate. The image consists of an upper terrace and a lower terrace with steps in between. There is a segment of the [110] oriented step and a segment of the [112] oriented step. It can be seen that the [110] step is decorated by bridge-bonded thioliates with a regular spacing of 0.58 nm. There is a staple row of RS–Au–SR along the [112] step on the upper terrace. In this image, the resolution is not high enough to resolve the bridge-bonded species along the [112] step. From our observation, the first row of staples on the upper terrace is formed at the step edge. On some narrow terraces, no ordered structure exists apart from one single row of staples pinned to the [112] step edge. More rows are formed subsequently during scanning and eventually a whole terrace can be covered with the staple rows. There is also a region of the disordered phase on the lower terrace. As the area is scanned continuously, the striped phase is observed to grow by consuming the disordered phase. Both the [110] and [112] oriented steps are stable. However, not all steps are stable. Fig. 5 shows an STM image covering a larger area than that of Fig. 4. In Fig. 5, we can see that near the upper right hand corner, there is an area with a high density of [112] oriented steps. All the [112] oriented steps have a common feature that there is a staple row running along the step edge on the upper terrace. Areas marked by white rectangles consist of steps not along any major crystallographic directions. The hairy/spiky appearance of these steps indicates that there are atoms/molecules constantly entering as well as leaving these steps. The terraces next to these irregular and changing steps...
do not have an ordered structure. It is likely that these irregular steps are the providers of gold atoms for the formation of the staples. The [112] oriented steps are not the stable steps on the clean Au(111) surface. However, they seem to be abundant on the thiolate covered surface. These [112] oriented steps are relatively long and very straight suggesting a significant stabilizing effect from the step-bound thiolates. From the structural model of Fig. 3, it can be seen that the [113] step is more open than the [110] step. Au atoms along the [112] step have a lower coordination number. This may enhance the bonding to the thiolates as a result of a higher charge density offered by the [112] step. It is also possible that many of the [112] oriented steps are transformed from irregular steps mediated by the diffusing SR.

Since the discovery of the RS–Au–SR staple motif on both the surfaces of Au nano-clusters and the Au(111) crystal surface, numerous studies have demonstrated that such a staple motif applies almost universally to all small Au clusters. However, recent work reports the finding of bridge-bonded thiolate on the (001) surface of a tetragonal Au nanocrystal. The direct observation of bridge-bonded thiolate along step edges on Au(111) is significant for the understanding of the structures of larger Au clusters with a fcc lattice. Such clusters have exposed (111) and (100) facets with edges similar to those found on planar Au(111) surfaces. Even for small clusters which are not fcc, there is still a possibility of forming bridge-bonded SR along the edges of the cluster surfaces.

It is interesting to speculate whether allylthiolates with longer chains such as butylthiolate would form the same kind of bridge-bonded species along step edges on Au(111). It is known that the van der Waals interaction between the allyl chains plays an important role in the self-assembly of allylthiolate monolayers on Au(111). Most strikingly, at maximum coverage, thiolates with three or fewer carbon atoms form the $3 \times 4$ phase, while thiolates with more than three carbon atoms in the chain take a different $2\sqrt{3}\times 3$-rect. structure. We expect that the formation of the bridge-bonded species along step edges may also show some chain length dependence.

Conclusions

To summarize, we have found clear evidence of bridge-bonded methylthiolate along the [110] and the [112] step edges on Au(111). There is also some indication of bridge-bonded propylthiolate from the experiment involving mixed methyl- and propylthiolate, but further confirmation of such species is needed in future studies. This bridge-bonded SR is expected to play an important role in the assembly of SAMs on both the planar (111) surface and on the surfaces of gold nano-particles. More work is needed in this area to search for bridge-bonded SR at saturation coverage and also for thiolates with longer chains. Because of the importance of surface steps in providing free-diffusing Au atoms, understanding the nature of step decoration by bridge-bonded thiolate would be an important achievement in terms of a better explanation of alkanethiol self-assembly.

Conflicts of interest

There are no conflicts to declare.

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References

1. M. Zhu, C. M. Aikens, F. J. Hollander, G. C. Schatz and R. Jin, Correlating the crystal structure of a thiol-protected Au-25 cluster and optical properties, J. Am. Chem. Soc., 2008, 130, 5883.
2. H. C. Weissker, et al., Information on quantum states pervades the visible spectrum of the ubiquitous Au$_{144}$SR$_{60}$ gold nanocluster, Nat. Commun., 2014, 5, 3785.
3. R. Philip, P. Chantharasupawong, H. F. Qian, R. C. Jin and J. Thomas, Evolution of nonlinear optical properties: From gold atomic clusters to plasmonic nanocrystals, Nano Lett., 2012, 12, 4661.
4. Y. Negishi, et al., A critical size for emergence of nonbulk electronic and geometric structures in dodecanethiolate-protected Au clusters, J. Am. Chem. Soc., 2014, 137, 1206–1212.
5. A. Dass, S. Theivendran, P. R. Nimmala, C. Kumara, V. R. Jupally, A. Fortunelli, G. Barcaro, X. Zuo and B. C. Noll, Au$_{133}$[SPh-tBy]$_{52}$ nanomolecules: X-ray crystallography, optical, electrochemical, and theoretical analysis, J. Am. Chem. Soc., 2015, 137, 4610.
6. P. D. Jadzinsky, G. Careo, C. J. Ackerson, D. A. Bushnell and R. D. Kornberg, Structure of a thiol monolayer-protected gold nanoparticle at 1.1 angstrom resolution, Science, 2007, 316, 3754.
7. K. M. O. Jensen, P. Juhas, M. A. Tofanelli, C. L. Heinecke, G. Vauhgan, C. J. Ackerson and S. J. L. Billinge, Polymorphism in magic-sized Au$_{144}$SR$_{60}$ clusters, Nat. Commun., 2016, 7, 11859.
8. C. Zeng, Y. Chen, K. Iida, K. Nobusada, K. Kirschbaum, K. J. Lambright and R. Jin, Gold quantum boxes: on the periodicities and the quantum confinement in the Au$_{238}$, Au$_{356}$, Au$_{44}$ and Au$_{52}$ magic series, J. Am. Chem. Soc., 2016, 138, 3950.
9. C. Zeng, C. Liu, Y. Chen, N. Rosi and R. Jin, Atomic structure of self-assembled monolayer of thiolate on a tetragonal Au$_{25}$ nanocrystal, J. Am. Chem. Soc., 2016, 138, 8710.
10. H. W. Dong, L. W. Liao and Z. K. Wu, Two-way transformation between fcc- and non-fcc-structured gold nanoclusters, J. Phys. Chem. Lett., 2017, 8, 5338.
11. C. Zeng, H. Qian, T. Li, G. Li, N. L. Rosi, B. Yoon, R. N. Barnett, R. L. Whetten, U. Landman and R. Jin, Total structure and electronic properties of the gold nanocrystal Au$_{49}$SR$_{34}$, Angew. Chem., Int. Ed., 2012, 51, 13114.
12. S. Tian, et al., Structural isomerism in gold nanoparticles revealed by X-ray crystallography, Nat. Commun., 2015, 6, 8667.
13. H. Hakkinen, The gold–sulfur interface at the nanoscale, *Nat. Chem.*, 2012, 4, 443.

14. O. Voznyy, J. J. Dubowski, J. T. Yates and P. Maksymovych, The role of gold adatoms and stereochemistry in self-assembled methylthiolate on Au(111), *J. Am. Chem. Soc.*, 2009, 131, 12989.

15. P. Maksymovych, D. C. Sorescu and J. T. Yates, Gold-adatom-mediated bonding in self-assembled short-chain alkanethiolate species on the Au(111) surface, *Phys. Rev. Lett.*, 2006, 97, 4.

16. Q. Guo and F. Li, Self-assembled alkanethiol monolayers on gold surfaces: resolving the complex structure at the interface by STM, *Phys. Chem. Chem. Phys.*, 2014, 16, 19074.

17. L. Tang, F. Li, W.-C. Zhou and Q. Guo, The structure of the methylthiolate and ethylthiolate monolayers on Au(111): absence of the (root 3 × root 3) R30 degrees phase, *Surf. Sci.*, 2012, 606, L31.

18. L. Tang, F. Li and Q. Guo, Complete structural phases from self-assembled methylthiolate monolayers on Au(111), *J. Phys. Chem. C*, 2013, 117, 21234.

19. F. Li, L. Tang, O. Voznyy, J. Gao and Q. Guo, The striped phases of ethylthiolate monolayers on the Au(111) surfaces: a scanning tunneling microscopy study, *J. Chem. Phys.*, 2013, 138, 194707.

20. J. Gao, F. S. Li and Q. Guo, Balance of forces in self-assembled monolayers, *J. Phys. Chem. C*, 2013, 117, 24985.

21. P. Maksymovych, O. Voznyy, D. B. Dougherty, D. C. Sorescu and J. T. Yates Jr, Gold adatoms as a key structural component in self-assembled monolayers of organosulfur molecules on Au(111), *Prog. Surf. Sci.*, 2010, 85, 206–240.

22. S. Clair, Y. Kim and M. Kawai, Coverage-dependent formation of chiral ethylthiolate-Au complexes on Au(111), *Langmuir*, 2010, 27, 627–629.

23. Y. Yourdshahyuan and A. M. Rappe, Structure and energetics of alkanethiol adsorption on the Au(111) surface, *J. Chem. Phys.*, 2002, 117, 825.

24. S. Rousset, V. Rapain, G. Baudot, Y. Garreau and J. Lecoeur, Self-ordering of Au(111) vicinal surfaces and application to nanostructure organized growth, *J. Phys.: Condens. Matter*, 2003, 15, S3363–S3392.

25. L. Tang, X. Zhang and Q. Guo, Site-specific chemistry directed by a bi-functional nanostructured surface, *Langmuir*, 2010, 26, 4860.

26. J. Gao, F. S. Li and Q. Guo, Mixed methyl- and propylthiolate monolayers on the Au(111) surface, *Langmuir*, 2013, 29, 11082.

27. G. E. Poirier, Characterization of organosulfur molecular monolayers on Au(111) using scanning tunneling microscopy, *Chem. Rev.*, 1997, 97, 1117.

28. R. G. Nuzzo, L. H. Dubois and D. L. Allara, Fundamental studies of microscopic wetting on organic-surfaces. 1. Formation and structural characterization of a self-consistent series of polyfunctional organic monolayers, *J. Am. Chem. Soc.*, 1990, 112, 558.