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### Probing Phase Evolutions of Au-Methyl-Propyl-Thiolate Self-Assembled Monolayers on Au(111) at Molecular Level

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#### **ABSTRACT**

A self-assembled monolayer (SAM) consisting of a mixture of CH<sub>3</sub>S-Au-SCH<sub>3</sub>, CH<sub>3</sub>S-Au-S(CH<sub>2</sub>)<sub>2</sub>CH<sub>3</sub> and CH<sub>3</sub>(CH<sub>2</sub>)<sub>2</sub>S-Au-S(CH<sub>2</sub>)<sub>2</sub>CH<sub>3</sub> was studied systematically using scanning tunneling microscopy and density functional calculations. We find that the SAM is subject to frequent changes at the molecular level on the time scale of ~minutes. The presence of CH<sub>3</sub>S or CH<sub>3</sub>S-Au as a dissociation product of CH<sub>3</sub>S-Au-SCH<sub>3</sub> plays the key role in the dynamical behavior of the mixed SAM. Slow phase separation takes place at room temperature over hours to days leading to the formation of methyl-thiolate-rich and propyl-thiolate-rich phases. Our results provide new insights into the chemistry of the thiolate-Au interface, especially for ligand exchange reaction in the RS-Au-SR staple motif.

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#### INTRODUCTION

Alkanethiol passivated gold nanoclusters represent an important class of materials with size-dependent electronic and optical properties.<sup>1-4</sup> Ever since the discovery of thiol-passivated Au nanoclusters,<sup>5</sup> there has been a sustained high level of interest both in producing Au nanoclusters with precise size control and accurate structural determination of such clusters.<sup>6-15</sup> Despite the well-documented stability of thiol-passivated Au nanoclusters, these clusters are found to be able to transform by ligand exchange <sup>16,17</sup> and even intercluster reactions.<sup>15</sup> Many different processes have been proposed to explain ligand exchange and cluster transformation, <sup>18,19</sup> but there is still a lack of consensus over the surface chemistry of the thiolate-Au interface.<sup>20</sup> In particular, when the protecting layer consists of more than one type of thiolates, it is not yet clear how the two different types of thiolates interact with each other and how this interaction affects the stability of the protected Au cluster.

There is a long history of research into self-assembled monolayers (SAM) of alkanethiols on the extended Au(111) surface. The fundamental structural motif, the RS-Au-SR staple motif, is found on both Au(111) <sup>23,26</sup> and the surface of Au nanoclusters. Furthermore, the application of scanning tunneling microscopy (STM) to SAMs on Au(111) has a great advantage that it images the surface in real space with atomic scale resolution and is hence able to provide important structural

information of the S-Au interface, which is otherwise difficult to obtain from passivated Au nanoclusters. Here, we present the findings from our recent study of mixed methyl- and propyl-thiolate monolayers on Au(111) using STM. We focus on investigating the mixing and phase-separation of the two alkyl-thiolate species. Our mixed SAM has three different building blocks: CH<sub>3</sub>(CH<sub>2</sub>)<sub>2</sub>S-Au-S-(CH<sub>2</sub>)<sub>2</sub>CH<sub>3</sub>, CH<sub>3</sub>S-Au-S-CH<sub>3</sub> and CH<sub>3</sub>S-Au-S-(CH<sub>2</sub>)<sub>2</sub>CH<sub>3</sub>. Because of different chain lengths between CH<sub>3</sub>S and CH<sub>3</sub>(CH<sub>2</sub>)<sub>2</sub>S, ligand exchange reaction can be followed more easily as a result of clear contrast change in STM images. Therefore, we are able to monitor, on the molecular level, phase evolutions within the SAM in real time.

#### EXPERIMENTAL AND COMPUTATIONAL METHODS

The experimental details have been described in an earlier publication. <sup>47</sup> The mixed SAM is prepared by exposing the gold sample to a background of methyl-propyl disulfide (MPDS) vapor at room temperature (RT) at a pressure of 1×10<sup>-5</sup> mbar. Saturation coverage is reached after ~ 15 minutes' exposure. The exact exposure is subject to uncertainties in the measurement of the MPDS pressure using an ion gauge. The gold sample is a (111)-oriented Au film deposited on a highly-oriented pyrolitic graphite substrate. The Au film is cleaned using cycles of Ar<sup>+</sup> ion sputtering and thermal annealing. STM imaging was performed at both RT and 110 K using

electrochemically etched tungsten tips. All STM images reported here, were obtained by using about -0.6 V sample bias and 0.05 nA tunneling current. STM tip is far away from the surface molecules in order to avoid the tip effect. The molecular coverage is obtained by finding the density of the thiolate which is then normalized by the density of surface gold atoms.

The DFT calculations were performed with the Vienna *ab initio* Simulation Package (VASP).<sup>59,60</sup> The electron-ion interactions were described using the projected augmented wave (PAW) method.<sup>61</sup> The exchange-correlation energy was calculated with the general gradient approximation (GGA) functionals of Perdew-Burke-Ernzerhof (PBE).<sup>62</sup> An energy cutoff of 400 eV was selected for the plane-wave expansion. The dispersion corrections of the molecules and Au surface interactions were included by the van der Waals density functional (vdw-DF) proposed by Dion.<sup>63-65</sup> The Au surfaces were modeled with periodic slabs consisting of five atomic layers. A vacuum of 15 Å was adopted to avoid the periodic image interactions normal to the surface. A Monkhorst-Pack grid of  $2 \times 4 \times 1$  was employed to sample the surface Brilloun zone. In all cases, the top three layers of atoms were allowed to relax in three dimensions. The first principles STM simulations were conducted with the Tersoff-Hamann method and the bSKAN code. <sup>66,67</sup>

#### **RESULTS AND DISCUSSION**

#### Full coverage $3 \times 4$ phase

We begin by examining the structure of the mixed SAM at the coverage corresponding to 1/3 monolayer (ML) of thiolate (treating both methyl-thiolate and propyl-thiolate equally). Here 1 ML is defined as one thiolate per Au atom. 1/3 ML is thus the maximum coverage that can be achieved on Au(111). Figure 1a and b shows two STM images of the mixed monolayer at RT. It can be seen that complete phase separation into separate methyl-thiolate and propyl-thiolate domains does not happen. The nearly random mixture of methyl-thiolate and propyl-thiolate results in a rather disordered appearance of the STM images. However, as shown previously using Fourier transform of the images,<sup>47</sup> the bright spots in the images have excellent positional correlation and all spots can be matched onto a 3 × 4 lattice which is common for monolayers of methylthiolate, 46,48 ethylthiolate 42,46 and propylthiolate. 55 Previous studies of the  $3 \times 4$  phase of propyl-thiolate SAM <sup>55</sup> and methyl-thiolate SAM, 46 show that the propyl/methyl chains can be divided into two groups as illustrated in the ball model of Fig. 1c. The  $\beta$  chain tilts more towards the surface normal and appears taller than the  $\alpha$  chain. Therefore, for the mixed SAM, we expect to see spots in STM images having four different levels of contrast from i) the \beta propyl chain; ii) the  $\alpha$  propyl chain; iii) the  $\beta$  methyl chain and iv) the  $\alpha$  methyl chain. Under our imaging conditions, the "visible" spots in the images from the 3 × 4 phase of the mixed SAM are from either the  $\alpha$  or the  $\beta$  propyl chains with the  $\beta$  propyl chain appearing the tallest. The methyl-thiolate species, being significantly shorter than propyl-thiolate, appears dark and "invisible". The 3 × 4 lattice is occupied by a rather random mixture of CH<sub>3</sub>(CH<sub>2</sub>)<sub>2</sub>S-Au-S-(CH<sub>2</sub>)<sub>2</sub>CH<sub>3</sub>, CH<sub>3</sub>S-Au-S-CH<sub>3</sub> and CH<sub>3</sub>S-Au-S-(CH<sub>2</sub>)<sub>2</sub>CH<sub>3</sub>. This results in the apparent lack of ordering in the two STM images. The images also show the presence of a number of voids within the SAM. These voids are not patches of bare Au(111), otherwise, thiolate from the surrounding area with a high adsorbate density would diffuse into the voids. The lack of ordered structure inside the voids indicates that the voids are filled by SCH<sub>3</sub> or Au-SCH<sub>3</sub>, both are dissociation products of CH<sub>3</sub>S-Au-S-CH<sub>3</sub>. CH<sub>3</sub>S-Au-SCH<sub>3</sub> itself is not stable as a complete unit at RT.<sup>68</sup> SCH<sub>3</sub> or Au-SCH<sub>3</sub> inside the voids are mobile at RT and they provide the required surface pressure to prevent the voids from collapsing.

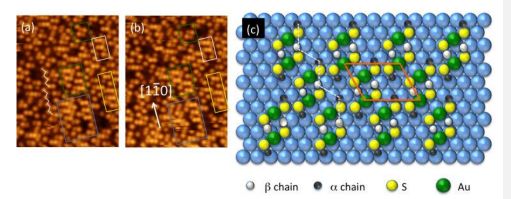


Figure 1. (a) and (b) STM images,  $5.5 \text{ nm} \times 7.5 \text{ nm}$ , acquired from the same area of the sample at RT. Image in (b) was collected three minutes after the image in (a). Rectangular boxes are drawn onto the images to highlight areas where only minor changes have taken place. Outside the boxes, changes are so extensive that direct comparison between the two images becomes difficult. In both images, the spots are from propyl chains. The brightest spots are the  $\beta$  chains. The methyl chains are in the dark voids and not resolved under the tunnel condition applied. S and Au are not visible in these images. (c) Ball model of the  $3 \times 4$  phase of alkanethiolate monolayers at 1/3 ML coverage. The zigzag line formed by connecting the  $\alpha$  and  $\beta$  chains is directly comparable with experimental findings shown in (a).

According to the staple motif, each spot in the STM image comes from one branch of the staple. The  $CH_3(CH_2)_2S$ -Au- $S(CH_2)_2CH_3$  staple can be identified easily by finding a pair of related spots. For the  $CH_3S$ -Au- $S(CH_2)_2CH_3$  staple, we can also find the location of the  $SCH_3$  branch by knowing where its partner, S- $(CH_2)_2CH_3$ , is located. There are local areas, inside the blue rectangle in Fig. 1a for example, where the composition is significantly rich in propyl-thiolate and the  $\beta$  chains are exclusively propyl chains. Therefore, some minor segregation of propyl-thiolate from the mixed

layer has already taken place.

To summarize the features of the  $3 \times 4$  mixed SAM, most part of the SAM consists of a rather random mixture of  $CH_3(CH_2)_2S$ -Au-S( $CH_2$ )<sub>2</sub> $CH_3$  and  $CH_3S$ -Au-S( $CH_2$ )<sub>2</sub> $CH_3$ . Although there are not complete separations of the methyl- and propyl-thiolate, there are local areas enriched either in methyl-thiolate or propyl-thiolate. The areas heavily enriched with methyl-thiolate are covered by a liquid-like  $SCH_3$  or  $AuSCH_3$  and appear as voids within the  $3 \times 4$  SAM.

When the mixed SAM with the  $3 \times 4$  structure is continuously monitored with the STM, we find that the local areas highly-enriched with propyl-thiolate are more stable and do not change much with time. The two images shown in Fig. 1 are collected from the same area with image in (b) acquired three minutes after that in (a). Significant changes are observed within this three minutes' period. In fact, only a small fraction of the scanned area, mostly regions highly enriched with propyl-thiolate, maintains the original structure. Rectangular boxes are added to the images to assist the eye to identify areas subjected to minor or no changes in between the two scans. The changes can sometimes be captured in real time by the STM as shown in Fig. 2.

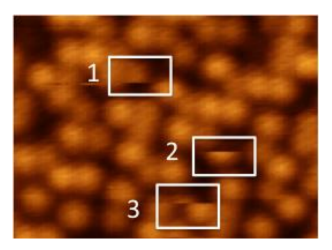


Figure 2. STM image showing real time changes captured by the STM. The fast scan direction is the horizontal direction. After each scan line, the STM tip moves upwards by a small step. Inside the rectangular boxes we highlight instantaneous changes that are recorded by the STM.

Inside rectangular box 1, we see the sudden appearance of a thiolate. We see the top <sup>3</sup>/<sub>4</sub> of the thiolate with the bottom <sup>1</sup>/<sub>4</sub> of it not showing up because the thiolate was not in that location when the tip passed by. This is a typical phenomenon when an atom/molecule moves from somewhere else to a position such that part of the atom/molecule falls in front of the advancing STM tip. Inside box 2, we see a thiolate disappearing after the bottom half of it has been imaged by the STM. In box 3, we can identify a thiolate jumping from its initial location to a new location to the upper left. In this case, the bottom half of the thiolate was seen by the STM in its original location and the top half of the same thiolate seen in the new location.

The events of mobile thiolates shown in Fig. 2 are the ones captured by the STM while in action and are only a small fraction of the total events. When comparing two STM images taken just a few minutes apart, the majority of the thiolates are observed to have changed positions. This high activity of thiolate displacement, however, is not a common feature for the short chain alkanethiol monolayers. When the 3 × 4 phase of a pure propyl-thiolate monolayer <sup>55</sup> or of an ethyl-thiolate monolayer <sup>42,-46</sup> is imaged with the STM at RT, it appears stable with little changes occurring. The facile movement of the propyl and methyl chains inside the mixed SAM is hence a result of the presence of methyl-thiolate. The mobile AuSCH3 and SCH3 are able to interact with CH<sub>3</sub>(CH<sub>2</sub>)<sub>2</sub>S-AuS(CH<sub>2</sub>)<sub>2</sub>CH<sub>3</sub> and CH<sub>3</sub>(CH<sub>2</sub>)<sub>2</sub>S-AuSCH<sub>3</sub> via ligand exchange reactions. When molecules are observed to change their location or conformation under the STM, there are several potential contributions to these changes. The STM tip, for example, has the capability to initiate changes due to either electronic excitations by the tunneling electrons or the localized electric static force. Under many situations, one can identify which is the dominating contribution to changes by altering the tunneling current of the strength of the electric field beneath the tip. We have not investigated how the observed changes respond to tunnel current or electric field. However, we have clear evidence that the changes are due to the presence of AuSCH<sub>3</sub> or SCH<sub>3</sub> which are mobile on Au(111) at RT. When the sample temperature

is decreased and the mobility of AuSCH<sub>3</sub> or SCH<sub>3</sub> is frozen, the molecular layer stops changing.

#### Nearly stoichiometric striped phase

When the mixed layer with the  $3 \times 4$  phase is left at RT in vacuum for two days, there appears to be a gradual decrease in coverage, presumably due to expansion of the monolayer and loss of the thiolate to the surrounding sample holder. There is also a small probability of desorption via the formation of dimethyldisulphide. The  $3 \times 4$ phase eventually becomes disordered before the appearance of a striped phase. Figure 3a shows an STM image acquired at RT showing alternating bright and dim rows of the striped phase. Inset shows the height profile across line A-B. The regular brightdim row feature can be explained as due to an ordered arrangement of CH<sub>3</sub>S-Au-S(CH<sub>2</sub>)<sub>2</sub>CH<sub>3</sub>. Figure 3c is a structural model derived from DFT calculations and Fig. 3d shows a simulated STM image according to the structural model. The striped phase according to the model consists of 100% CH<sub>3</sub>S-Au-S(CH<sub>2</sub>)<sub>2</sub>CH<sub>3</sub> staple rows. The staples are organized in such a way that there are alternating SCH<sub>3</sub> and S(CH<sub>2</sub>)<sub>2</sub>CH<sub>3</sub> rows, all aligned in the  $[11\overline{2}]$  direction. The STM image in Fig. 3a shows that although the overall structure is consistent with the model of Fig. 3c, the real structure at RT is not as rigid. Along the bright S(CH<sub>2</sub>)<sub>2</sub>CH<sub>3</sub> rows, one can see gaps as a result of substitution of S(CH<sub>2</sub>)<sub>2</sub>CH<sub>3</sub> by SCH<sub>3</sub>. Similarly, along the dim SCH<sub>3</sub> rows, we can find SCH<sub>3</sub> being substituted by S(CH<sub>2</sub>)<sub>2</sub>CH<sub>3</sub>. Overall, the chemical composition of the striped phase is close to 1:1 for the SCH<sub>3</sub>/S(CH<sub>2</sub>)<sub>2</sub>CH<sub>3</sub> ratio. The striped phase shown in Fig. 3 is under dynamical equilibrium at RT. Frequent displacement of thiolate is observed and as a result two subsequent images taken from the same area never look the same. The changes are however limited to the extent that the bright rows remain bright and dim ones remain dim.

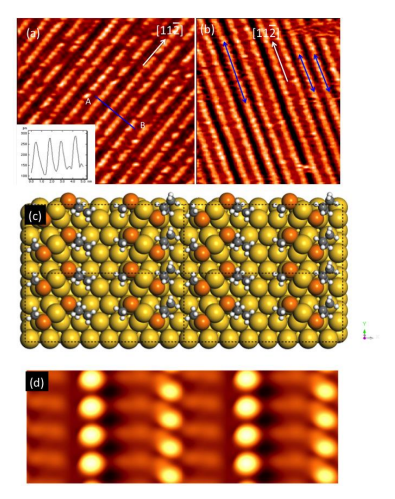


Figure 3. (a) STM image,  $10 \text{ nm} \times 10 \text{ nm}$ , showing the nearly stoichiometric striped phase with alternating bright and dim rows. There is also evidence of row-pairing. (b) Some rows are not paired

with their neighbors. (c) Structural model derived from DFT. (d) Simulated STM image according to the structural model in (c).

#### Striped phases enriched with either $SCH_3$ or $S(CH_2)_2CH_3$

As described earlier, the nearly stoichiometric striped phase is under dynamic equilibrium. Given enough time, this striped phase changes further leading to clear phase separation. Phase separation gives rise to domains dominated by methylthiolate and separate domains dominated by propyl-thiolate. The domains heavily enriched with methyl-thiolate have no ordered structure at RT. However, under repeated scanning by the STM tip, an ordered striped phase can be formed locally as demonstrated in Fig. 4. The formation of the stripes shown in Fig. 4 is tip-induced because once scanning is terminated, the stripes get dissolved quickly into the disordered background. Stripes similar to those shown in Fig. 4 are observed to be stable at lower temperatures. Figure 5a shows an STM image from a striped phase of a methyl-thiolate-rich area at 110 K. Thus, tip-induced striped phase is very similar to the self-formed striped phase under low enough temperature. The reason that the striped phase of methyl-thiolate is not stable at RT is probably due to the entropic effect. When scanning at RT, there is expected to be an attractive tip-molecule interaction. This interaction is electrostatic in nature due to the electric field within the tunnel junction. This attractive interaction tends to pull molecules towards the area below the STM tip. This is equivalent to a compressive pressure acted on the molecules. This "compressive pressure", we believe, forces the molecules to assemble into the striped phase.

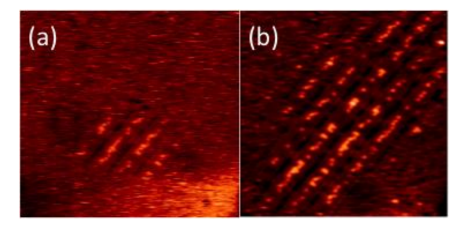


Figure 4. STM-induced formation of a striped phase from a disordered, methyl-thiolate-rich, area at RT. The stripes are parallel to the  $[11\overline{2}]$  direction. (a) The early stage where four short segments of thiolate rows are formed. (b) Later on, the segments grow longer with new rows also formed. Image size: 20 nm  $\times$  20 nm.

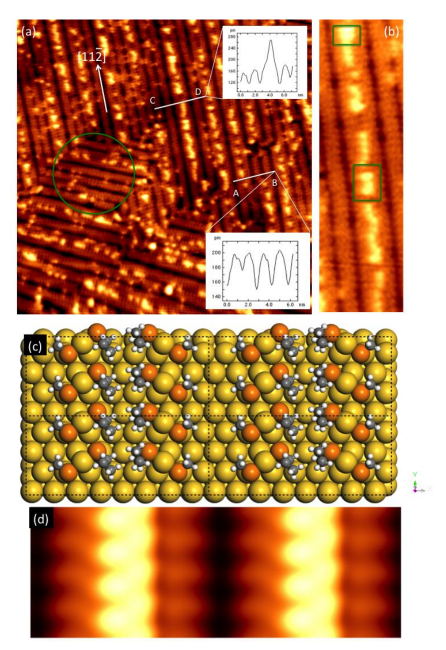


Figure 5. (a) STM image of a striped phase of the mixed SAM. The SAM was prepared at RT and it was cooled down to 110 K. Image was collected at 110 K. The striped phase was formed when temperature is below 190 K. This area is heavily enriched with methyl-thiolate. Insets show height profiles across the staple rows. Area inside the green circle highlights a nearly pure methyl-thiolate phase. (b)  $\beta$  chains of propyl-thiolate appear much taller than the methyl-chains. (c) Structural model derived from DFT calculations. (d) Simulated STM images according to the model in (c).

In Fig. 5a, we first concentrate on a particular area inside the green circle. The height corrugation of the rows, 40 pm, is the same as that found for the CH<sub>3</sub>S-Au-SCH<sub>3</sub> rows. <sup>48</sup> We therefore identify these rows as CH<sub>3</sub>S-Au-SCH<sub>3</sub> rows. The tall features, 120 pm taller than the CH<sub>3</sub>S-Au-SCH<sub>3</sub> rows, are from S(CH<sub>2</sub>)<sub>2</sub>CH<sub>3</sub>. The tall features are mostly found in between two rows as shown in Fig. 5b. Figure 5c gives a model based on DFT calculations. According to the model, two adjacent rows are paired with an inter-row distance of 3.5a. Each row consists of CH<sub>3</sub>S-Au-S(CH<sub>2</sub>)<sub>2</sub>CH<sub>3</sub> staples. The S(CH<sub>2</sub>)<sub>2</sub>CH<sub>3</sub> branches from the two rows in the pair have a head-to-head arrangement. Figure 5d shows a simulated STM image according to the model in (c). The simulated image is in good agreement with the experimental findings. For two staple rows in a pair, there are thus two rows of S(CH<sub>2</sub>)<sub>2</sub>CH<sub>3</sub>. The experimentally observed images appear more complicated, because the stoichiometry of the staples in each of the paired rows is not necessarily the same. At the top of the image in Fig. 5b, the area inside the green box shows features almost identical to that in Fig. 5c. However, rather than an exact head-to-head CH<sub>3</sub>S-Au-S(CH<sub>2</sub>)<sub>2</sub>CH<sub>3</sub>:CH<sub>3</sub>(CH<sub>2</sub>)<sub>2</sub>CH<sub>3</sub>S-Au-SCH<sub>3</sub> configuration, we find many instances of CH<sub>3</sub>S-Au-SCH<sub>3</sub>:CH<sub>3</sub>(CH<sub>2</sub>)<sub>2</sub>S-Au-SCH<sub>3</sub> and CH<sub>3</sub>S-Au-S(CH<sub>2</sub>)<sub>2</sub>CH<sub>3</sub>:CH<sub>3</sub>S-Au-SCH<sub>3</sub> configurations. Therefore, instead of seeing two regular rows of S(CH<sub>2</sub>)<sub>2</sub>CH<sub>3</sub>, we see the frequent interruptions of the S(CH<sub>2</sub>)<sub>2</sub>CH<sub>3</sub> row by SCH<sub>3</sub>. Figure 6 shows a simulated STM image based on a structural model

allowing the substitution of S(CH<sub>2</sub>)<sub>2</sub>CH<sub>3</sub> by SCH<sub>3</sub>. The simulated STM image in Fig. 6 is in good agreement with the observed image in Fig. 5a.

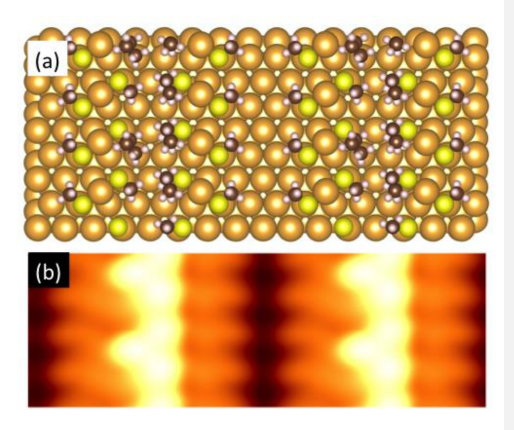


Figure 6. (a) The same structural model as that in Fig. 5c but with the substitution of some  $S(CH_2)_2CH_3$  by  $SCH_3$ . (b) Simulated STM image based on the model in (a).

In areas where propyl-thiolate is the majority of the population, we find striped phases similar to that shown in Fig. 5c even at RT. Figure 7 shows an STM image from an area enriched with propyl-thiolate. The staple rows have similar characteristics as that given by the simulation in Fig. 5c. If we substitute some  $\alpha$  SCH<sub>3</sub> by S(CH<sub>2</sub>)<sub>2</sub>CH<sub>3</sub> and some  $\beta$  chains by SCH<sub>3</sub> in Fig. 5c, the simulated image would be rather similar to that

shown in Fig. 7. It is evident that there is plenty of movement inside the striped SAM of Fig. 7. The SCH<sub>3</sub>/S(CH<sub>2</sub>)<sub>2</sub>CH<sub>3</sub> ratio evaluated from Fig. 7 is 4:1. The presence of around 25% of methyl-thiolate is once again responsible for the frequent changes of the propyl-rich SAM observed at RT. We found previously that striped phases of pure propyl-thiolate are stable, not showing displacement of individual thiolate at RT.<sup>47</sup>

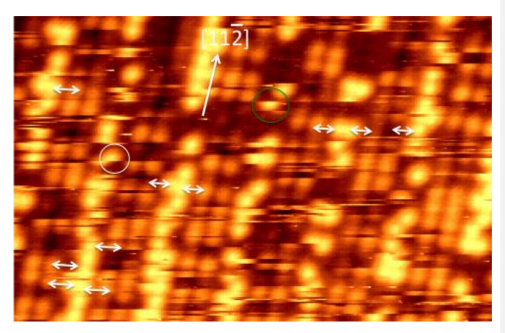


Figure 7. STM image,  $12 \times 7.5$  nm, of a striped phase of staples enriched with propyl-thiolate imaged at RT. The overall striped structure remains more or less unchanged with time, but frequent changes at the individual thiolate level are observed. Arrows indicate two thiolate branches belonging to the same staple. Circles highlight displacement of thiolate as captured by STM in real time.

Our study of the mixed methyl- and propyl-thiolate SAM clearly demonstrates that the SAM is an active layer at room temperature undergoing frequent changes. It has

been known for some time that the gold-thiolate interface is not rigid.<sup>69</sup> When phase separated domains of 1-undecanethiol and 11-mercaptoundecanoic acid are prepared on Au(111), the two thiolate species inter-diffuse and a mixed phase is obtained after several hundred hours at 353 K. 70 Lateral diffusion of thiolates on the surface of gold nano-clusters has also been studied. <sup>71</sup> However, several important questions remain to be answered. What is moving inside a SAM? Does the staple move as a single unit or is the dissociation of the staple a prerequisite for diffusion? If so, is RS-Au or just RS the mobile unit? Our study does not answer all of the above questions. Nevertheless, findings from the present study demonstrate the unequal role of the thiolates. Ligand exchange within the methyl- and propyl-thiolate layer is dominated by the properties of the methyl-thiolate. The methyl-thiolate staple is not stable at RT.<sup>68</sup> The dissociation of the staple should produce RS-Au, RS and even free Au adatoms. These dissociation products can then attack the CH3(CH2)2S-Au-S(CH<sub>2</sub>)<sub>2</sub>CH<sub>3</sub> staple leading to ligand exchange. Of course, at high enough temperatures, the CH<sub>3</sub>(CH<sub>2</sub>)<sub>2</sub>S-Au-S(CH<sub>2</sub>)<sub>2</sub>CH<sub>3</sub> staple is expected to break as well. Loss of methyl-thiolate from the surface through thermal desorption also takes place at high enough temperatures leading to the formation a pure propyl-thiolate SAM.<sup>55</sup> Although the S-Au bond within the staple can be broken via thermal activation, the energy required to break this bond increases with the chain length of the alkanthiol. For longer chain lengths, the van der Waals interaction provides a restoring force to repair the broken S-Au bond before the complete separation of RS chain from the staple. This restoring force is most effective for SAMs on the flat Au(111) surface, but less effective on the surface of nanoclusters. This is because on the surface of nanoclusters, the curved surface of the cluster is unable to keep a uniform distance between adjacent alkyl-thiolate chains, hence a weakened Van der Waals force.

The STM as an imaging technique has its drawbacks that each image requires a rather lengthy time to collect. This makes the STM unsuitable to follow changes that occur too rapidly. There are, however, ways to overcome this shortcoming. For example, by recording tunnel current as a function of time with the feedback loop disabled, temporal resolution of the order of 20 µs has been achieved 72 when studying the system of decanethiol monolayers on Au(111). 73

#### **CONCLUSIONS**

The interaction of short chain alkanethiols including  $CH_3SH$ ,  $CH_3CH_2SH$  and  $CH_3(CH_2)_2SH$  with the Au(111) surface produces similar structures based on the RS-Au-SR staple motif. They all have an ultimate  $3 \times 4$  phase at 1/3 ML of coverage. At coverage below 1/3 ML, the SAMs organize into similar striped phases. Among the three Au-thiolate staples we have investigated so far,  $CH_3(CH_2)_2S$ -Au- $S(CH_2)_2CH_3$  is the most stable. The methyl-thiolate SAM has the lowest stability. By adding methyl-

thiolate into propyl-thiolate, a mixed  $3 \times 4$  phase containing nearly equal amount of SCH<sub>3</sub> and SCH<sub>3</sub>(CH<sub>2</sub>)<sub>2</sub> is formed at 1/3 ML at RT. CH<sub>3</sub>S-Au-SCH<sub>3</sub> readily breaks apart at RT giving rise to SCH<sub>3</sub> and AuSCH<sub>3</sub>. It is not clear if we have both SCH<sub>3</sub> and AuSCH<sub>3</sub> on the surface or just SCH<sub>3</sub>. What is clear is that SCH<sub>3</sub> or AuSCH<sub>3</sub> is mobile on the surface and can react with CH<sub>3</sub>(CH<sub>2</sub>)<sub>2</sub>S-Au-S(CH<sub>2</sub>)<sub>2</sub>CH<sub>3</sub> by the ligand exchange process. This ligand exchange makes the SAM in dynamical equilibrium at RT. The mixed SAM when left at RT in vacuum slowly phase separates into methylthiolate-rich and propyl-thiolate-rich domains. The phase separation however, does not produce either a pure methyl-thiolate phase or a pure propyl-thiolate phase.

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## TOC Graphic

