Nanoscale chemical imaging of reversible photoisomerization of an azobenzene-thiol self-assembled monolayer by tip-enhanced Raman spectroscopy

Journal Article

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Publication date: 2018-01-22

Permanent link: https://doi.org/10.3929/ethz-b-000220746

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Originally published in: Angewandte Chemie. International Edition 57(4), <u>https://doi.org/10.1002/anie.201710443</u>

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Nanoscale chemical imaging of reversible photoisomerization of an azobenzene-thiol self-assembled monolayer by tip-enhanced Raman spectroscopy

Li-Qing Zheng, ^[1] Xing Wang, ^[1] Feng Shao, ^[1] Martin Hegner, ^[2] and Renato Zenobi*^[1]

Abstract: An understanding of the photoisomerization mechanism at the molecular scale of molecules bound to a metal surface is required for designing photo switches at surfaces. It has remained a challenge to correlate the surface structure and isomerization of photo switches at ambient conditions. Here, the photoisomerization of a self-assembled monolayer of azobenzene-thiol molecules on an Au surface was investigated using scanning tunneling microscopy and tip-enhanced Raman spectroscopy. The unique signature of the cis isomer at 1525 cm⁻¹ observed in tip-enhanced Raman spectra was clearly distinct from the trans isomer. Furthermore, tip-enhanced Raman images of azobenzene-thiols after ultraviolet and blue light irradiation are shown with nanoscale spatial resolution, demonstrating a reversible conformational change. Interestingly, the cis isomers of azobenzene-thiol molecules were preferentially observed at Au grain edges, which is confirmed by density functional theory calculation.

Molecular switches, which undergo molecular-scale motions upon external stimuli, are very promising for molecular electronics and high-density data storage. However, a deeper understanding of their mechanical and electronic properties at the molecular level is required for developing applications in the areas of information processing, energy and biology. ^[1-4] Of the various molecular switches studied, azobenzene and its derivatives adsorbed on surfaces, one of the most thoroughly investigated classes of molecular switches, [5-7] have attracted considerable attention in the last few years thanks to their practical applications as nanomotors ^[8], adaptive imaging ^[9] and energy storage devices ^[10] when binding to adaptive surfaces. The photoisomerization of azobenzene can be triggered by illumination with light of the appropriate wavelength. Azobenzene undergoes trans-to-cis isomerization when irradiated with ultraviolet (UV) light (365 nm). The conformational change can be reversed by blue light (450 nm), but it can also be caused thermally in the dark owing to the greater thermodynamic stability of the trans isomer. [11-14] Moreover, the isomerization can be induced mechanically ^[15], thermally ^[16] and

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electrostatically [17].

Photoisomerization of different molecular switches, for example 3,3',5,5'-tetra-tert-butyl-azobenzene (TBA) ^[18], 3,5-ditert-butyl-N-(3,5-di-tertbutylbenzylidene) aniline (TBI) [19] or azobenzene thiol (ABT) [20], adsorbed on an Au (111) surface has already been demonstrated. The mechanism governing the isomerization behavior of azobenzenes in solution, which is based on n- π^* ^[21,22] or π - π^* ^[23] electronic transition, is well understood from ultrafast time-resolved studies such as coherent resonant Raman scattering [24-26]. However, the mechanism of photoisomerization of azobenzenes is quite different when they are in direct contact with a metal surface, in contrast to the photoswitching in solution. The observation that azobenzene fails to isomerize on Au (100) or Cu (111) shows a strong substrate dependence. [27] To date, questions about the switching mechanism of surface-bound molecules under ambient conditions remain.

For a complete understanding of the isomerization processes of ABT on an Au surface, it is important to provide information on both the surface structure and the molecular fingerprint with nanometer spatial resolution, so that the relationship between structure and chemical behavior can be established. [28-30] Tip-enhanced Raman spectroscopy (TERS) is a near-field optical technique that combines the advantages of scanning probe microscopy (SPM) and plasmon-enhanced ^[31-35] In this methodology, an Raman spectroscopy. electrochemically etched Ag or Au tip amplifying the electromagnetic field of incident laser light enhances the Raman signal of sample molecules located close to the tip. TERS possesses high spatial resolution and high sensitivity, down to the single molecule level. [36, 37] TERS is ideally suitable to study defects and step edges, which may play a significant role in surface reactions. [38] Only very few studies have so far addressed the photoswitching properties of metal-surface-bound ABT at ambient conditions, where molecular switches are used in practical applications. [2,39] In these two works, the photoisomerization of ABT and an ABT derivative was probed by surface-enhanced Raman spectroscopy [13] and scanning tunneling microscopy (STM) [39] at ambient conditions, respectively. The reversibility ^[13] and a cooperative mechanism ^[39] of the photoisomerization were shown.

Here, we report the investigation of the isomerization mechanism of an ABT self-assembled monolayer (SAM) on a template stripped (TS) Au surface ^[40] (Figure S1) using ambient STM-TERS, taking advantange of the strong scattering properties of ABT. The reversible photoisomerization behavior of an ABT SAM on an Au surface was demonstrated, based on the presence and absence of the characteristic vibrational mode at 1525 cm⁻¹ of cis isomer after UV irradiation (*ex situ*). Furthermore, TERS maps were used to follow the isomerization

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of ABT, and to obtain the local information of the reactivity. Interestingly, the cis isomers of ABT molecules were preferentially observed at the grain edges of Au. This is ascribed to the lower reaction energy of trans-to-cis isomerization at Au steps due to the shift of d bands and different molecular orientation. These results underline that grain edges play a significant role in the photoswitching behavior of surface-bound molecules.



Figure 1. A. TER spectra of an ABT SAM before and after *ex-situ* or irradiation for 5 mins, as well as a spectrum with the tip retracted. The blue bars indicate the Raman peak at 1525 cm⁻¹ and 1107 cm⁻¹ which are characteristic for the cis isomer. B. Simulated Raman spectra of the trans and cis isomers of ABT with different mole fractions of the two isomers.

ore and after UV irradi 1A. h/ 07 cm ht. The p sample was irradiated with N) mod omer, respectively. in ER spectra and their correlation with the transazobenzene thiol molecules, B3LYP/6ioñ spectrum on the mole i ctic Figure 1B: The calculated free ct for the anharmonicity. [41] The cis scaling factor of 0.97 to corre

er and the consequent ity. ^[13] To compensate the loss of conjugation of the cis ison tensity. for this intensity difference, the intensistence of the sectrum was magnified 10 times. To f ity of the iay etter view of the rmalized to the most modes, all the Raman pe<mark>aks</mark>. n ectrum culate m⁻¹ 2 45 cm⁻¹ omponents. The band at anc ing e ri Inc etc fro ¹ is the H-C-H bending 65 cm⁻¹ is predominately m aky at 1 hing le of the -C-H bending mode of the alkyl group. he The band at 1600 is herization of ABT, the v_{sym} (N=N) frequency cm⁻¹ to 1525 cm⁻¹, while, the C-N stretching/ring shifts from 1145 cm⁻¹ to 1107 cm⁻¹. Since the shifts from 1465 cm⁻¹ to 15 R SAM de isomerization of ABT. Based on the simulated spectra for nola The low intensity of the peak at 1107 cm⁻¹ could be due to RS

The photoisomerization behavior of an ABT SAM on a semitransparent Au surface was first verified using UV-Vis spectroscopy. The absorbance difference at 340 nm of the ABT SAM before and after UV irradiation is shown in Table. S2. The absorbance at 340 nm decreases upon irradiation by UV light, confirming that trans-to-cis photoisomerization on the Au surface takes place. Moreover, when the irradiation time is prolonged from 5 mins to 15 mins, the value remains the same. This indicates that the steady-state trans-to-cis isomerization yield of the ABT SAM is already reached after a UV irradiation time of 5 mins. Upon irradiation with blue light for 15 mins, the absorbance at 340 nm recovers, indicating cis-to-trans isomerization.

The lifetime of the cis isomers on an Au surface is of great importance. The absorbance change at 340 nm of an ABT SAM on an Au surface after UV irradiation as a function of time is shown in Figure S4. The plot was fitted to exponential function. The thermal back reaction on the surface is dominated by first-order kinetics with a reaction rate constant of $5.67 \times 10^{-5} \text{ s}^{-1}$. The lifetime of the cis isomer on the Au surface is thus around 5 h, which is sufficient for acquisition of a TERS map. Isomerization of ABT in ethanol solution is shown in Figure S5.

TERS maps of an ABT SAM on a TS Au surface before and after irradiation of UV light were collected. Since the characteristic peak of the cis isomer is at 1525 cm⁻¹, which is in the region where spurious peaks from carbonaceous

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decomposition products usually appear, a very low laser power and a short exposure time were used to collect the TERS maps. Figure 2 shows the TERS peak intensity ratio (1525 cm⁻¹ peak / 1143 cm⁻¹ peak) maps of the ABT SAM on the Au surface before and after UV irradiation for different times. A dark blue color corresponding to 0 in the scale bar indicates that there is no Raman peak at 1525 cm⁻¹ in that particular region. In Figure 2A, we can barely see the cis peak. After the ABT SAM was irradiated with UV light for 5 mins, the Raman peak at 1525 cm⁻¹ appeared much more frequently than before, indicating that part of the ABT molecules underwent a trans-to-cis isomerization. (Figure 2B) This peak is irregularly distributed, and present in 29 % of all the pixels. To assess the variability of the peak at 1525 cm⁻¹ associated with the isomerization of ABT, principal component analysis (PCA) on Figure 2B was performed. (Section S3) PC1 corresponds to the TER spectrum of trans isomer, while PC2 may correspond to that of trans and cis isomers. Only a very small contribution from PC3, which stems from carbon contamination, is present in the spectra. This suggests that sample decomposition is minor.



Figure 2. TERS peak intensity ratio (1525 cm⁻¹ peak / 1143 cm⁻¹ peak) maps of ABT SAM before (A) and after UV irradiation for 5 mins (B) and 15 mins (C). The sample was prepared in the following way: ABT molecules were first self-assembled on a TS Au substrate, followed by irradiation with UV light for 5 mins and 15 mins. TERS peak intensity ratio map of the ABT SAM after UV irradiation (D). The sample was prepared in the following way: ABT in ethanol was first irradiated with UV light for 5 mins and then deposited onto a TS Au substrate. The size of the maps is 200 x 100 mm² with a 4.8 nm pixel size. The acquisition time for each spectrum is 1 s.

When the exposure time was increased to 15 mins, the presence of this peak in the whole map was slightly decreased to 26 %, which is a bit surprising. We interpret this to be due to the thermal back reaction during the TERS mapping (Figure 2C). From the data in Figure 2C, the spatial resolution was estimated to be around 10 nm based on a full-width at half-maximum (FWHM) analysis (Figure S8).

Another sample preparation method was tested to check the different isomerization rate of ABT in ethanol solution and in the form of a SAM on the Au surface. ABT in ethanol solution was first inediated with UV light for 5 mins, and then deposited on an Au surface, which takes ca. 120 mins. Finally, the TERS map of the ABT SAM sample was collected. (Figure 2D) The proportion of the cis peak in the whole map is 23 %, which is smaller than that in Figure 2B. This indicates that part of the cis isomers of ABT molecules must have reacted back to the trans isomers, while assembly on the Au surface was carried out. When ABT molecules are bound to a metal surface, the lifetime of the cis isomers decreases, as demonstrated above.

The ABT SAM on the Au su irradiated with UV w liated with TM image are shown in the cis peak in the that in Figure 2B and ng ST on of niv m than dic f th ck m., cis 72 all cor (he) paring the h th late grain edge. Me 00 cm⁻¹) map peak intensity ratio (1525 cm⁻¹ / 1600 cm⁻¹) map. (Figure S9) of 41 there is S11B and S11C), which indicates that the complementary nsity ratio maps shown in Figure S9 is not induced beak inte by different IS S



Figure 3. A. TERS peak intensity ratio (1525 cm⁻¹ peak / 1143 cm⁻¹ peak) map of an ABT SAM after UV irradiation for 5 mins followed by blue light irradiation for 5 mins; B. STM image of ABT SAM on the Au surface acquired together with the TERS map. The size of the map is 200 x 200 nm² with a 4.8 nm pixel size. The acquisition time for each spectrum is 1 s.

To confirm that the isomerization behavior at grain edges of Au is different from that on terraces, several high-resolution TERS measurements going across a terrace and a grain edge on Au were performed, and the results are shown in Figure 4, Figure S12 and Figure S13. We observed that the peak at 1525 cm⁻¹ has a strong preference to be observed at Au grain edges (Figure 4B, Figure S12B and Figure S13B). Moreover, all of the peak intensity ratios between trans peaks and the 1600 cm⁻¹ peak in the TER spectra of ABT adsorbed at the Au grain edges after UV irradiation (shown in Figure 4B) are smaller than that of ABT adsorbed on Au terraces (Figure S14), which also indicates that trans-to-cis isomerization of ABT took place predominantly at the grain edge of Au.

To understand the excitation mechanism in the photoisomerization, the electronic structure of ABT SAM on the Au surface and the density of states of the Au substrate should be taken into account. Tegeder and coworkers proposed a substrate-mediated charge transfer process for the photoisomerization of TBA on an Au surface ^[7]. UV light first generates hot holes in the Au d-band, which relax to the top of the d-band via an Auger-decay subsequently. The hot holes undergo a charge transfer process to the HOMO of TBA, leading to the formation of transient positive ions, which may subsequently result in the isomerization of TBA. According to density functional theory (DFT) calculations (Figure S15), the energy levels of the step edge Au atoms shift to higher values compared to those of Au atoms on the terrace due to unsaturation and a decrease in inter-atom coupling ^[43]. The higher the d-band center, the higher the reactivity. ^[44] Moreover, we calculated the total energy of trans-ABT isomers and cis-ABT isomers adsorbed on an Au terrace and a 2-atom step edge, respectively. (see Figure S16) The reaction enthalpy of trans-to-cis isomerization at the 2-atom step edge is 0.61 eV, which is 13% lower than that at the terrace (0.72 eV). Therefore, the cis isomers of ABT on step edges are thermodynamically more stable compared to those on terrace sites. It is fairly well established, for example, that the cis isomers of TBI are preferentially found on the herringbone reconstruction of Au, as observed from STM images due to the slight corrugation at these sites. ^[20]



Figure 4. A. STM image of the Au sunace covered with an ABT SAM after UV irradiation for 5 mins, obtained using an Ag tip; B. line-trace TER spectra along the crosses indicated in A. One spectrum was acquired every 2 nm of surface distance. The exposure time for per spectrum is 1 s.

Also, because of the completely different geometry at grain edges, the molecular orientation of ABT molecules adsorbed at these sites is different from that of ABT molecules adsorbed at

the terraces of Au, which could favor the trans-to-cis photoisomerization of ABT (Figure S17).

bstrate the in photoisomerization of an ABT SAI TM-TERS. TER ้นร spectra of ABT SAM before and U) ce of the ene-th the mer. ino ecules after UV light m imaging. Furthermore, we Ы hn i e p it . is h the cis-to-trans b eaction is sl e shift of d bands to higher energy and ibed en ov D substrate-mediated han photoisomeri of ABT on an Au surface. The ability to spatially distinguish the molecular vibrational features of elecules adsorbed at different surface sites will contribute to a sta switche hò

Acknowledgements

L.-Q. Z. tharlks the Chinese Scholarship Council for a Ph.D. student fellowship. We thank the High Performance Computing Team at ETH Zurich for help with the DFT calculations, Dr. Ewellina Lipiec and Jacek Szczerbinski for helpful discussions, and Dr. Alessandro Lauria for help with recording UV-Vis

Keywords: photoisomerization • azobenzene thiol • tipenhanced Raman spectroscopy • grain edges • substrate

mediated charge transfer

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Entry for the Table of Contents

Layout 1:

COMMUNICATION

The reversible photoisomerization of an azobenzenes-thiol (ABT) self-assembled monolayer was demonstrated by tipenhanced Raman spectroscopy. The cis isomers of ABT were preferentially observed at Au grain edges.



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