

Effect of the pH in the Growth of Benzotriazole Model Layers at Realistic Environmental Conditions.

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Abstract

The growth of benzotriazole (BTAH) via solution deposition onto copper monolayers prepared via underpotential deposition (UPD) on Au(111)/mica substrates has been investigated using X-ray photoelectron spectroscopy (XPS) and ambient scanning tunnelling microscopy (STM) as a function of solution pH and immersion time. Adsorbed species have been found sensitive to the solution pH with a higher benzotriazole surface concentration following deposition from an acidic environment. Although the layers prepared at different pH are chemically different, as highlighted by XPS, similar morphologies are recorded via STM. These results are critically discussed in the light of some of the adsorption models previously reported.

Keywords: copper; oxidation; passive films; XPS; STM

1. Introduction

Copper and its alloys are widely used in a variety of applications, from electrical wiring and plumbing, to electronic and microelectronics, to ornaments, due to desirable properties such as high thermal and electrical conductivity, ductility, malleability, and corrosion resistance [1]. Copper behaves like a noble metal because its surface is passivated by a naturally protective film [2], mainly consisting of cuprous oxide (Cu_2O), which can protect the copper surface from corrosion. However, when copper is exposed to environments exhibiting harsh conditions, it may be susceptible to corrosive phenomena. In aqueous environment, copper corrosion has often been associated with the presence of chloride ions alone, or in combination with other reactive molecular species, to the extent that it cannot be completely prevented. One useful approach to minimize corrosion is to use corrosion inhibitors, additives which, in low concentration, decrease the deterioration of

the surface layers. Benzotriazole (BTAH, figure 1) coatings have long been established as an efficient method of passivating copper surfaces [3-6].

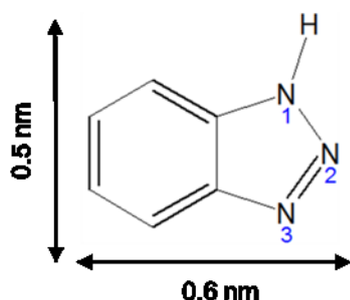


Figure 1. Benzotriazole (BTAH) molecular structure with approximate molecular dimensions.

Since the early work of Cotton *et al.* [7, 8] there has been ever growing interest in understanding the mechanism of corrosion inhibition and therefore in the interaction of BTAH with both oxide-free [7-35] and oxide-covered [7-10, 13-18, 28, 32], copper surfaces. However, the origin of the particular efficacy of BTAH, in term of its detailed bonding, is not really explained and controversial data and interpretation are reported [4-6]. BTAH has been proposed to interact with the copper substrate resulting in the formation of a copper metal complex through the nitrogen atoms [7-8, 16-18, 32] capable of generating a polymer chain [9, 19, 22, 26-28, 31-32]. BTAH was also observed to adopt different orientation on copper surfaces, i.e. adopting an upright/tilted [10, 13, 23-25, 32-36] or flat orientation [14-15, 32-34, 36, 37 on Au(111)] with respect to the surface or a mix of these orientations [14-15, 33-34]. The majority of previous studies focus on ideal model systems such as clean copper surfaces, or theoretical calculations. In attempts to simulate real application conditions, several studies were undertaken in a liquid acidic environment on copper single crystals [15-17]. Studies conducted at real conditions usually focus on bulk copper samples, the surface of which is considered, evaluating the influence of the pH in aqueous solution [9, 11, 26-27, 30, 38], synergistic effects due to the presence of other additives [30, 39], static and flow conditions [29]. Nevertheless, the investigation of the interaction of benzotriazole with real or realistic copper surfaces is made more complex both because of the multiplicity of the possible interactions occurring within the system studied, and by the difficulty of performing surface science measurements under realistic conditions.

The aim of this research is to study the interaction between benzotriazole and copper containing substrates exposed to realistic operating conditions, at room temperature. X-ray Photoelectron Spectroscopy (XPS) and ambient Scanning Tunnelling Microscopy (STM) measurements were carried out upon immersion of the samples in benzotriazole containing solutions to gain a chemical and morphological characterisation of the system. The samples consisted of copper monolayers prepared via underpotential deposition on Au(111)/mica substrates. The use of well-defined copper monolayers allows one to obtain information on the benzotriazole-copper interface, as, by limiting the interaction of BTAH with a single layer of copper atoms,

3D film growth is minimised, and BTAH interaction with the sample surface layer is rendered more evident. As a result, the gap between model studies, undertaken under ideal conditions, and realistic studies, performed at realistic conditions, can be bridged. The Cu – BTAH interaction has been considered as a function of exposure parameters such as solution pH and immersion time.

2. Materials and Methods

Samples consisted of copper monolayers prepared on Au(111)/mica substrates (300 nm epitaxial, Georg Albert PVD, Silz, Germany). Substrates annealed using natural gas flame prior to use and copper overlayers prepared by underpotential deposition (UPD), to yield a (5×5) chlorine terminated copper monolayer (set potential of 0.2 V) [40, 41, 42]. UPD layers were produced in a hydrochloric acid solution (10%) containing 50 mM CuSO₄ (Sigma-Aldrich, 99.999%) and 10 mM KCl (Fluka, ≥ 99.5%). A cyclic voltammogram was recorded to ascertain the quality of the Au/mica substrate prior to perform the metal deposition (a typical cyclic voltammogram is shown in S11). Inhibitor stock solutions were prepared by dissolving 10 mmol (1.192 g) of BTAH (Sigma-Aldrich, 99%) in 1000 ml of NaCl solution 0.5 M, at room temperature, under magnetic stirring; HCl (1%) and NaOH (1%) were used to obtain the desired pH.

XPS measurements were performed on a Scienta ESCA 300 spectrometer, using a conventional Al anode (K α 1486.6 eV, 10 kV, 20 mA). The detection system consisted of a large hemispherical analyser coupled to a multichannel plate/video camera. The instrument was calibrated on a daily basis to the Au 4f and Ag 3d photoelectron lines. The binding energy scale was referenced to the Au 4f_{7/2} core level peak at 84.0 eV. All spectra were analysed using the CasaXPS [43] software package.

STM topographs were collected on a nanoscope Pico STM, in constant current mode, at ambient conditions, using mechanically cut Pt/Ir tips (80:20 wire, Advent Research Materials Ltd., 0.25 mm diameter). Images were processed using the WSxM software package [44].

Substrates were immersed in the BTAH containing solutions for the necessary time, in static conditions and at room temperature; after removal from the solutions and prior to XPS and STM analysis, substrates were rinsed in deionised water and dried in a nitrogen flow.

3. Results

3.1 XPS measurements

Figure 2a shows the Cu 2p_{3/2} region before and after immersion of Cu(UPD)/Au(111)/mica layers into BTAH solutions at pH 6 for different lengths of time. After the copper layer is prepared and before immersion, the Cu 2p_{3/2} region can be fitted with a peak having maximum at 932.3 eV, with a small shoulder at 935.5 eV; both peaks have a FWHM of 1.87 eV (see the peak fitting for the as prepared layers in table S1 and S12). The Cu 2p_{3/2} region changes after immersion in the BTAH solution at pH 6 for 30 minutes. The total area

decreases, the peak maximum shifts to 932.7 eV, a new peak appears at 935.6 eV, the FWHM increases to 1.92 eV and a raised background, due to satellite features, is seen between 940 eV and 948 eV. After longer immersion times, the maximum of the Cu $2p_{3/2}$ shifts to slightly lower binding energy values, the overall peak areas, including the satellite regions, decrease in intensity. Figure 2b shows the Cu $L_3M_{45}M_{45}$ (referred to as LMM for simplicity) Auger transition recorded after preparing the copper UPD layer and after immersion into the BTAH solution at pH 6 for different times. It has been reported that the copper LMM Auger transition is comprised of four peaks, the most intense one at a kinetic energy of ca. 918.4 eV; with other transitions at 921.1 eV, 916.0 eV and 914.0 eV [45]; the Auger spectra from a sputtered copper single crystal is reported in SI3 for comparison. Before exposure to BTAH, the Cu LMM main peak is recorded at 918.1 eV, with shoulders at 920.5 eV and a raised background at lower kinetic energy (see table S1.) The shape of the transition is very similar to that recorded for CuI (SI3), where copper is in the +1 oxidation state. After exposure to the BTAH solution, the overall intensity of the transition decreases, the maximum of intensity shifts to a kinetic energy of 917.9 eV and a new feature appears at ca. 914.8 eV. Upon further exposure, the two observed peaks keep decreasing in intensity with the peak at 917.9 eV initially decreasing faster than that at 914.8 eV.

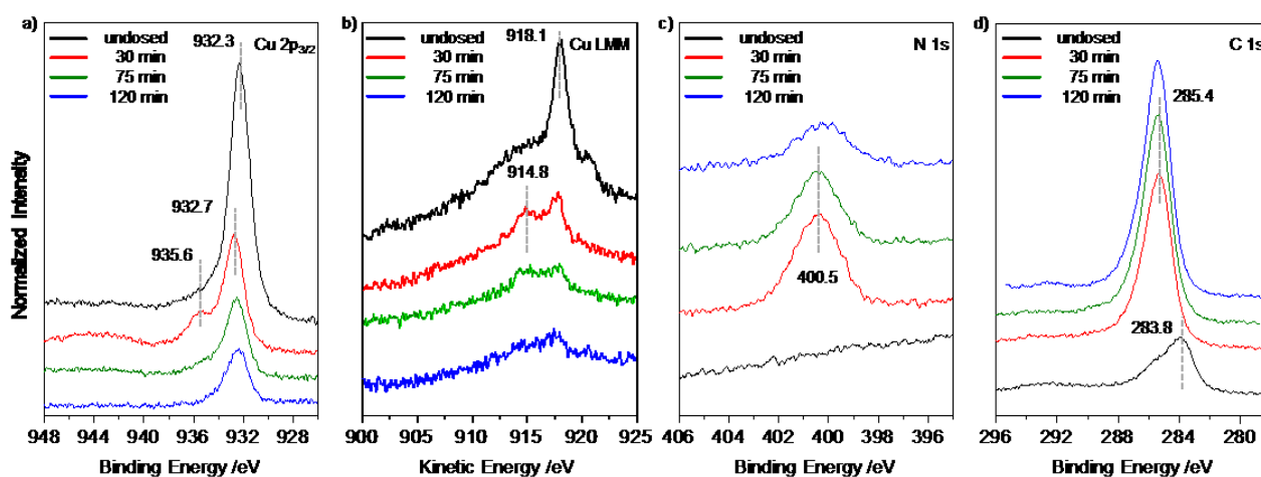


Figure 2. a) Cu $2p_{3/2}$ XP spectra before and after immersion in BTAH solution at pH 6 for increasing times; b) corresponding Cu LMM Auger transitions, c) N 1s and d) C 1s spectra.

Figure 2c and 2d show the N 1s and C 1s regions (see SI 2b and c, where the peak fitting is reported). Before exposing to BTAH, the N 1s region shows a flat line; after immersing into the solution, a single N 1s peak is recorded at 400.5 eV, with a FWHM of 2.22 eV. The peak becomes a little broader and less intense with increasing immersion times. Benzotriazole is the only source of nitrogen in the system. The C 1s region shows a peak with maximum at 283.8 eV (1.77 eV FWHM) and some other additional components before immersion. After immersion the maximum of the C 1s shifts to 285.4 eV, the FWHM increases to 2.10 eV. The peak increases in intensity and a raised background with maximum at 292.4 eV develops with increasing immersion times. Only a small amount of oxygen is present before immersion (S2 2d peak maximum at

532.1 eV, 1.85 eV FWHM); after immersion the O 1s peak increases with the immersion time, maximum at 533.5 eV, with a shoulder at 536.7 eV, which progressively decreases in intensity; ca. 2.5 eV FWHM. Before exposure to the BTAH solutions, a Cl 2p at 197.5 eV was seen (fittings in SI 2e). This is due to the chloride termination over the copper layer produced during the under potential deposition process [40, 41, 42]. The peak can be resolved into the 3/2 and 1/2 components and has a narrow FWHM (1.89 eV). A shoulder to higher binding energy (with a possible $2p_{3/2}$ peak at ca. 201 eV), might indicate a small amount of an oxychloride species [46]. Upon immersion in the BTAH solution, after the first 30 minutes the total signal seems to increase a little, the FWHM increases (2.43 eV); both the total signal and FWHM decrease following longer immersion. In particular, the tail at higher binding energy decreases at a faster rate.

Traces of Na were seen after immersion, as those ions were present in the BTAH stock solution.

As shown in figure 3a, the Cu $2p_{3/2}$ XP spectra after immersion in the BTAH solution at pH 5 show very similar features to those observed after immersions in the pH 6 solutions (peak fitting is reported in SI 4a). The peak maximum is recorded at 932.7 eV (2.32 eV FWHM), with a shoulder at 935.7 eV and a raised background between 940 eV and 948 eV due to satellite features. With increased exposure, the intensity of all peaks decreases. However, the Cu LMM region shows some differences (figure 3b). After immersion, two peaks are recorded at kinetic energies of 917.8 eV and 915.2 eV. The peak at 915.2 eV has a larger area than that at 917.8 eV after the initial exposure. With increasing exposures, the Auger transition is attenuated with the peak at 915.2 eV decreasing faster, similarly to what observed after immersion at pH 6. Likewise, after immersion, an N 1s peak with maximum at 400.5 eV (SI 4b), a C 1s with maximum at 285.4 eV (SI 4c) and an O 1s peak with maximum at 533.5 eV (SI 4d) were observed. Both the N 1s and C 1s peak decreased in intensity with longer immersion times, whilst the O 1s increased. Similarly to the observed behaviour after immersion at pH 6, the Cl 2p increases after 30 min and then decreases.

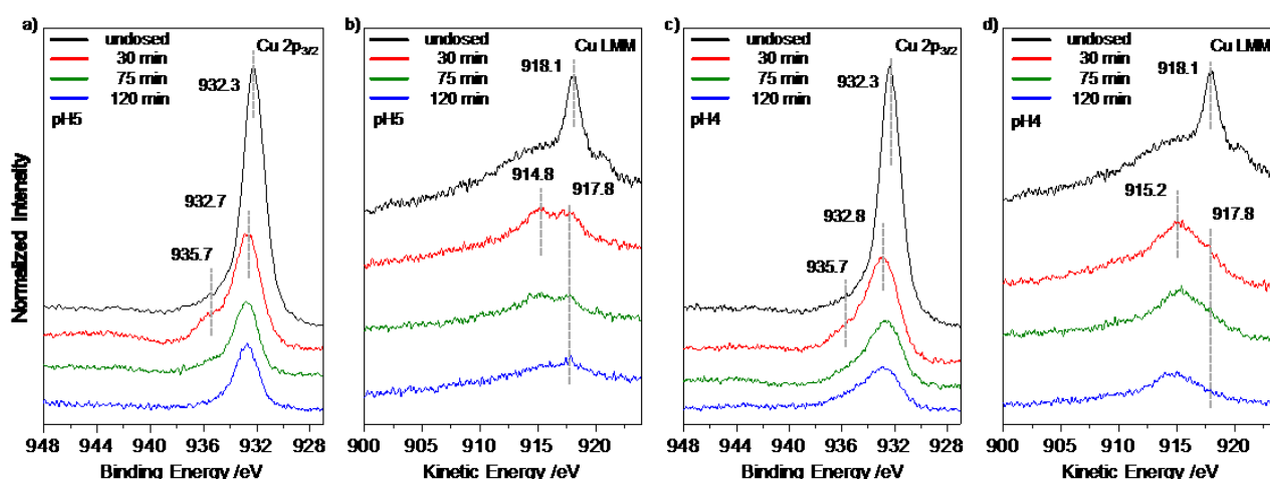


Figure 3. a) Cu $2p_{3/2}$ XP spectra before and after immersion in BTAH solution at pH 5 for increasing times; b) corresponding Cu LMM Auger transitions spectra; c) Cu $2p_{3/2}$ XP spectra before and after immersion in BTAH solution at pH 4 for increasing times; d) corresponding Cu LMM Auger transitions spectra.

Figures 3c and 3d show the Cu 2p_{3/2} XP and Cu LMM Auger transition spectra recorded after immersing the Cu(UPD)/Au(111)/mica substrates into the BTAH solution at pH 4 for increasing times (peak fittings are reported in SI 5a). After the initial 30 minutes exposure, the Cu 2p_{3/2} shows a maximum at 932.8 eV, with a shoulder at 935.7 eV, and the peak's FWHM increases to 2.63 eV. A weak satellite structure is still present. With increasing exposure to BTAH, both peaks are attenuated and their FWHM increase. The Cu LMM region shows a different behaviour yet again. A peak with a large FWHM is recorded at a kinetic energy of 915.2 eV with a raised background at the lower kinetic energy side. The peak recorded at 917.8 eV is now very weak. With increasing exposures, the Auger transition is attenuated with the peak at 917.8 eV decreasing faster, and the shoulder at 914 eV becoming more evident. After immersion, the N 1s peak shows a maximum at 400.6 eV, with a shoulder at 403.1 eV (SI 5b) and a C 1s with maximum at 285.3 eV (SI 5c). The O 1s peak shows a maximum at 532 eV (SI 5d). The N 1s decreased in intensity with longer immersion times, whilst the C 1s stayed approximately constant and the O 1s increased in intensity. The Cl 2p decreases in intensity, this time however, the tail at high binding energy is more persistent, but the signal decreases faster than for the other solutions.

3.2 STM measurements

Figure 4a recorded after immersing a sample in the BTAH solution at pH 6 for 75 minutes is a typical example of surface topography. BTAH molecular features appear as elongated protrusions and organise in chain-like structures which follow essentially three propagation directions related by a 120° rotation. The peak to peak distance between the elements composing a chain-like feature is 0.35 ± 0.02 nm. This value implies that adsorbed molecules are to some extent in registry with the underlying (5×5) Cl terminated Cu layer which is reported to have an average copper adatom spacing of 0.36 nm [40]. When compared with the dimensions of gas phase BTAH (figure 1), this suggests that the molecules are oriented upright or perhaps approximately perpendicular with respect to the surface. The distance between two adjacent BTA units may indicate the possibility of π - π interactions between the aromatic systems [47]. Although the overall propagation occurs along three rotationally equivalent surface directions, chain-like features appear to deviate from a straight line, and exhibit irregular widths.

Statistical analysis undertaken at the different immersion times, normalised to a 10×10 nm² area, shows a tendency for BTAH features to agglomerate in short ensembles, composed of 1 to 3 units (figure 4b). However, the increase in exposure times seems to favour the presence of features comprised of a higher number of elements; in fact, chains with fewer units are always more numerous but, with increasing immersion time, the number of chains exhibiting longer dimensions increase. As expected, the surface coverage increases with increasing the immersion time, going from ca. 38% at 30 minutes to ca. 66% at 75 minutes, to ca. 89% at 120 minutes.

STM images collected after immersing the samples in BTAH solutions at pH 5 and pH 4 show essentially the same topographic features and behaviour with increasing exposure times. Additional images are reported in SI6.

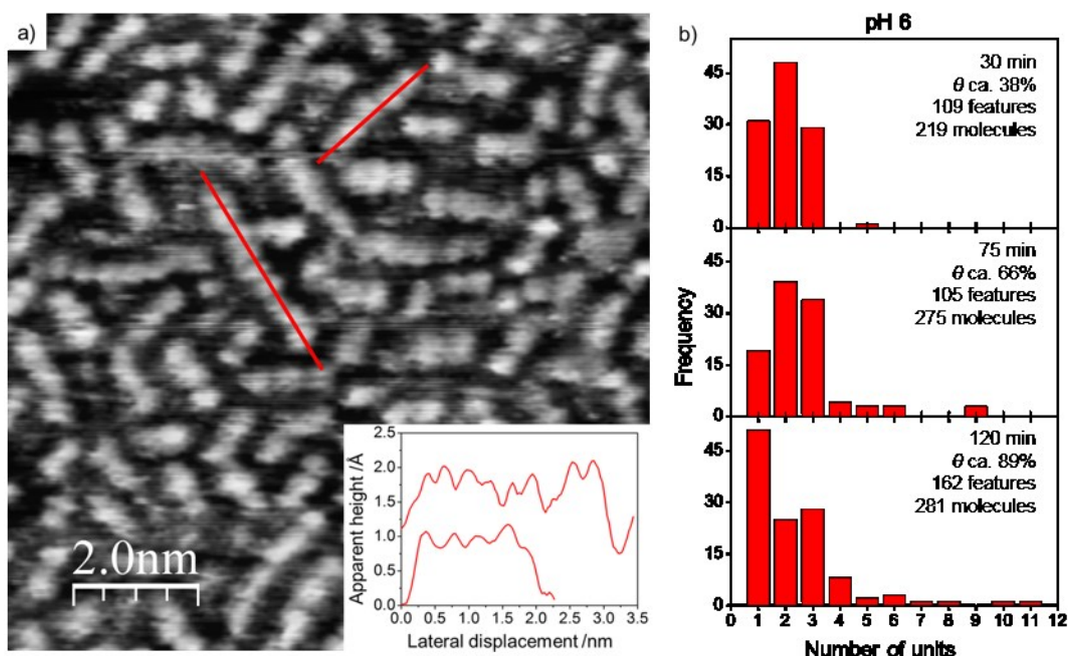


Figure 4. a) STM topography of a (5×5)Cl-terminated (1×1)Cu(UPD)/Au(111)/mica sample immersed into a BTAH solution at pH 6 for 75 minutes. $10 \times 10 \text{ nm}^2$, 0.2 V, 0.075 nA. In the inset, line profiles as shown in picture. b) Statistics on length of chains at pH 6, normalized to a 100 nm^2 area.

4. Discussion

The choice of preparing copper monolayers by underpotential deposition (UPD) is motivated by the necessity to control the amount of copper and evaluate its oxidation state reliably. In fact, through UPD, copper monolayers can be prepared and have been demonstrated to have a well-defined geometry with no intermixing between the two metals [40, 41, 42]. Another advantage of these preparations is the absence in the Cu 2p XP and LMM regions of a contribution due to the bulk, which are indeed present when using single crystals, or bulk specimens. The presence of copper is confirmed by a Cu $2p_{3/2}$ XP peak at 932.3 eV and fine structure of the Auger $L_3M_{45}M_{45}$ transition. For similar preparations binding energies of ca. 932.5 eV [48] and 931.1 eV [49] are reported. Following such preparation method, the copper layer is capped by a layer of chloride atoms, therefore copper is expected to be oxidised. As a reference, Cu $2p_{3/2}$ binding energies and Cu LMM kinetic energies for copper and relevant copper chlorides and oxides are reported in Table 1. The most intense component of the Auger LMM transition is recorded at a kinetic energy of 918.1 eV, a value much closer to that of Cu(0), than of Cu(I). However, the line-shape generated by bulk CuCl is very

similar to that of metallic copper, albeit much less intense and broadened [50, 51]. A Cl 2p_{3/2} peak at 197.3 eV confirms the presence of chlorine negatively charged. This value is much lower than those previously reported for chlorine in bulk CuCl and CuCl₂, as evidenced by the comparison with the binding energy values reported in Table 1, indicating that chlorine is a surface species, other than negatively charged. After preparation copper atoms are essentially in the +1 oxidation state and are capped by a layer of chloride atoms. A Cu:Cl ratio of ca. 1:2.2 is estimated, corresponding to a CuCl·Cl surface species. The attenuation of the copper signal may be explained by the fact that copper atoms are covered by chlorine species.

The fine structure of the Cu LMM Auger transition is different in shape, less resolved and broader than that for exhibited by bulk copper. The geometry of the layers, which is not directly accessible through XPS measurements alone, is discussed elsewhere [42]; however, the superposition of these layers is enough to create several slightly different copper chemical environments, which result in a broader distribution of kinetic energies for the Auger electrons, reflecting in an increased FWHM for the whole LMM transition.

Upon immersion of the samples in the BTAH containing solutions, the Cu 2p_{3/2} shows a shift towards higher binding energy, 932.7 eV; this shift alone may be not enough to warrant a change in the copper oxidation state from +1 to +2, albeit it may indicate a change in the copper chemical environment. A Cu 2p_{3/2} component developed at higher BE than for the original copper substrate, 938.4 eV, was previously attributed to the Cu(I)-BTA polymeric complex [17, 59, 60]. However, in this work, the development of a shoulder at ca. 935.7 eV and of a satellite structure at ca. 9 eV higher binding energy is indicative of the presence of some copper in the +2 oxidation state [61, 62].

	Cu 2p _{3/2} BE /eV	LMM KE /eV	α' /eV	O 1s BE /eV	Cl 2p _{3/2} BE /eV	Ref.
Cu UPD	~ 932.5					48
	931.1					49
CuCl	931.8				199.5	50
	932.2	915.6	1847.8			52
	932.4	915.5	1847.9		199.7	54
	934.6				199.8	51
CuCl ₂	933.8	915.5	1850.1			52
	933.8	916.6	1850.4		198.0	53
	933.9				199.2	53
	932.4	919.0	1851.4			52
Cu	932.9	918.3	1851.2			54
	932.6	918.0	1850.6			55
	932.6	918.4	1851.0			56
	932.6	918.5				57
Cu ₂ O	932.0	917.1	1849.1			52
	932.5	915.8	1848.3	530.8		55
	932.4	916.5	1848.9			56
	933.3	916.3	1849.6			57
	932.3	916.8	1849.1	530.2		58
CuO	933.6					52
	933.6	917.9	1851.5	529.4		53
	932.7	917.1	1850.7	529.8		53
	934.0	918.0	1850.7	529.9		55
	933.6	917.1	1851.1			56
	933.6	917.8	1851.4	530.3		57
	933.9			529.7		58

Table 1. Binding energy, kinetic energy and Auger parameters values for copper and selected compounds.

Therefore a mix of Cu(I) and Cu(II) is now present, with the majority still being in favour of Cu(I). This is observed consistently for the initial exposure at every pH; with increasing immersion times both the shoulder at higher binding energy and the satellite structure tend to be less pronounced. The LMM Auger transition shows the peak related to copper in a chloridic environment decreasing in intensity and a new feature appearing at 914.8 eV. This second feature was previously attributed to a Cu(I)BTA compound [63, 64, 65]. From the data reported in Table 1, for bulk compounds, the Cu LMM for Cu(I) is expected at an average kinetic energy of 916.5 eV for Cu₂O and 915.5 eV for CuCl, whereas at a kinetic energy of ca. 917.6 eV for CuO and 916.0 eV for CuCl₂ for Cu(II). Thus, other than to the oxidation state, the LMM seem to be dependent upon the chemical environment of the copper atoms. A clear peak indicating the presence of Cu(II) could not be observed in the LMM region. Thus, considering both the Cu 2p_{3/2} and Cu LMM regions, after the initial immersions in the BTAH solution, the vast majority of copper atoms are essentially still in the +1 oxidation state, even though some Cu(II) is also present.

As immersion time increases, especially for pH 6 and pH 5, the amount of Cu(II) decreases until the only detectable copper oxidation state remaining on the surface is +1, both in form of a Cu(I)BTA and copper in a chloridic environment. For pH 4, the signal related to Cu(II) seems more persistent. Changes in oxidation states for the Cu-BTA compound were previously observed for Cu₂O surfaces initially covered in a Cu⁺-BTA surface compound which was gradually oxidized to a Cu²⁺-BTA surface compound at increased immersion times in BTA containing solutions [16]. A similar behaviour was recorded for copper-nickel alloys exposed to BTA, where the Cu(I)BTA film was found to rapidly oxidise to Cu(II)BTA upon removal from the solution [17]. However, in these cases, a change in oxidation state from +1 to +2 was observed. A further explanation for the appearance of some Cu(II) will be discussed later. In the present case other than a change in oxidation state, a decrease in the total copper signal and in particular of the component related to Cu(II) was detected. Moreover, the N 1s peak area is seen to decrease following a similar trend. This is probably due to the presence of a capping layer forming after the successive exposures to the BTAH solutions, rather than due to a change in oxidation state. In fact, the decrease in area of the Cu 2p_{3/2} and N 1s peaks correlates with a relative increase of the O 1s and C 1s peak areas. This may indicate the possible growth of an adventitious contamination layer due to the adsorption of molecules from the atmosphere occurring while the samples are removed from the solutions, dried under a N₂ flow and brought into the vacuum chamber for XPS analysis. A schematic illustration of the adsorption mechanism is shown in figure 5.

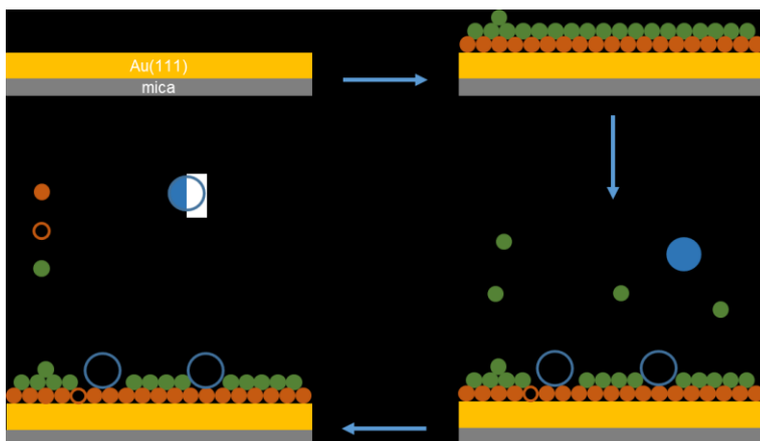


Figure 5. Schematic illustration of BTAH adsorption mechanism on a (5×5)Cl-terminated (1×1)Cu(UPD)/Au(111)/mica sample.

The most important difference, noted upon exposing the substrates to BTAH solutions at different pH after the initial immersion, is the variation in the relative intensities of the peaks comprising the copper LMM Auger transition, due to Cu-BTA (915.2 eV) and copper in a chloridic environment (917.8 eV), as reported in figure 6a for ease of comparison.

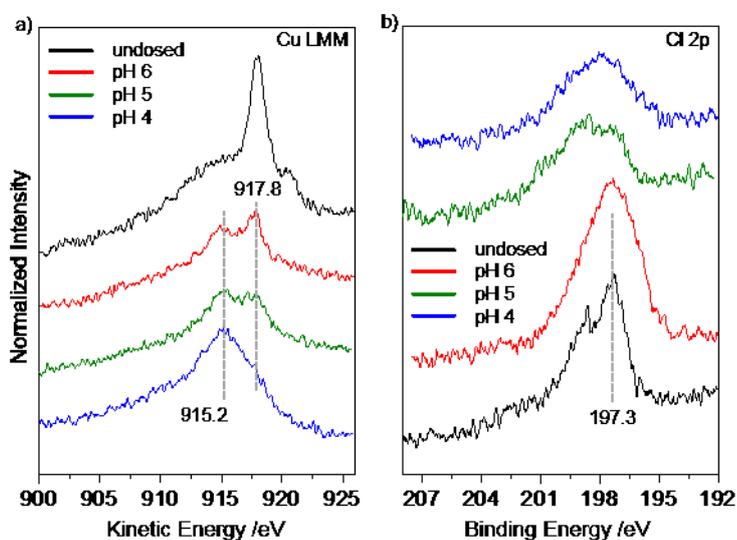


Figure 6. a) Cu LMM Auger transitions and b) Cl 2p regions before and after 30 min exposure to BTAH solutions at different pH.

A peak fitting done according to the method proposed by Finšgar and co-workers [64], shows that the peak related to the Cu-BTA species integrates roughly the same amount at pH 6 and 5, whereas increases significantly at pH 4, indicating that the Cu(I)BTA surface complex forms more favourably in the more acidic environment (the peak fitting is reported in SI7). Incidentally, Cohen and co-workers reported the formation of thicker Cu-BTA overlayers on samples treated at more acidic pH [63]. The changes in the

copper LMM regions are accompanied by a broadening of the Cl 2p_{3/2} region (figure 6b), which indicates the presence of additional environments for the chlorine atoms, albeit negatively charged.

The peak related to copper in a chloridic environment decreases with the increased acidity of the solution. This observation prompts a consideration on how the relative stabilities of copper and BTAH vary with pH. Although copper does not react with hydrochloric acid, because the reduction potential of copper is higher than that of hydrogen, copper specimens immersed in 0.5 M HCl solution [66, 67, 68] exhibit some weight loss, which is ascribed to acidic corrosion. Bulk copper is reported to go in solution as the soluble chloride complex CuCl₂⁻, through a series of surface reactions involving CuCl, which has a poor adhesion to metallic copper and hence is easily transferred to the solution [66, 67, 68]. Potential-pH diagrams report that in aqueous solution the stability of CuBTA and CuCl₂⁻ show an inverse relationship for 4 < pH < 6; at 0 volts, for a 10⁻² total activity of dissolved BTAH species in 1 M NaCl, CuBTA is the more stable species at higher pH values, whereas the water soluble CuCl₂⁻ species is more stable at pH < 4 [69]. Moreover, also the solution chemistry of CuCl is pH sensitive [70]: copper(I) chloride goes in solution as a H_xCuCl_{1+x} (x=1, 2) complex for pH < 2.5. At 2.5 < pH < 5, CuCl disproportionates in CuCl₂ + Cu. At pH > 5.0 copper(I) chloride undergoes alkaline hydrolysis.

In our experimental framework the acidity of the solution was kept at 4 < pH < 6; therefore, assuming that the reactivity of the (5×5) Cl capped Cu (1×1)/Au(111)/mica system has some similarities with the solution chemistry of bulk CuCl, at the operation conditions, there would be a potential for the Cl-capped Cu layer to disproportionate at lower pH or to dissolve into solution at higher pH. However, the UPD layer is reported as stable from neutrality to pH 1 [42]. Nevertheless, after 30 minutes of immersion in the BTAH solutions, with decreasing pH a decreasing amount of copper in a chloridic environment is revealed by measurements in the Cu LMM Auger region.

BTAH has a pK_a of 8.4 at 25 °C in water [5, 27, 69, 71, 72], which means that at pH values lower than pK_a the prevalent species is BTAH in its neutral form; at pH < 1 is in the protonated form, BTAH₂⁺; above 8.4 is in the anionic form, BTA⁻, instead. Therefore at our experimental conditions BTAH is prevalent as a solvated neutral species. UHV experiments demonstrated that BTAH can adsorb rather strongly on clean [14, 15, 23, 24, 25] and oxidised copper surfaces [14, 15], as an anionic species. Kokalji and co-workers calculated that at the metal/vacuum interface (representative of experiments undertaken in UHV conditions), chlorine adsorbs stronger than BTAH on Cu(111), whereas the opposite should happen for the metal/water interface, because the chloride ion is much smaller and solvates much stronger in water than BTA⁻ [73]. Because of the pK_a value, deprotonation in a solution with 4 < pH < 6 is likely to be a copper assisted phenomenon. Hence adsorption of BTAH on copper, in the form of BTA⁻, has to occur through displacement of chlorine atoms, and likely surface defects act as sites for the initiation of the reaction, as a defect-free Cl-Cu-Au/mica system is otherwise stable [42]. In our specific reaction environment, chlorine atoms may have some inherent degree of mobility, as it has been reported that their migration may be somehow linked with the formation of the BTA protective layer, which acts as a surface passivator for copper [27, 74, 75]. Moreover, two different factors have to be considered upon BTAH adsorption: on the one hand a chemical displacement of the

chloride atoms, which go in solution, due to the adsorption of benzotriazole on the copper; on the other hand, after adsorption, the steric hindrance of adsorbed BTAH, BTA, may induce a physical shift of the chloride atoms, creating local accumulation of negative charge, to which the surface copper atoms will respond with local accumulation of positive charge. This may result in areas on the surface with a stoichiometry described as CuCl_x , with $x > 1$, to which a more oxidised copper signal is attributed. This generates on the XP signal a $\text{Cu } 2p_{3/2}$ component at higher binding energy than for Cu(I) and a broadening of the $\text{Cl } 2p$ peak.

The Auger features (figure 6a) seem to indicate that BTA-Cu is less sensitive than copper in a chloridic environment to the variation of pH: after 30 minutes of immersion, the amount of BTA-Cu is approx. the same, whereas the amount of copper in a chloridic environment species decreases with the pH. The FWHM of the Cu LMM Auger feature slightly increases after immersion in the solution at pH 4, indicating the possibility of an increased amount of different chemical environments for the BTA-Cu species. As in NaCl 1 molar solution CuCl_2^- is favoured as the pH decreases [69], it is conceivable to think that some of the initial Cl-capped-Cu may go in solution more easily at lower pH values. Some changes in the $\text{Cl } 2p$ peak point towards the chlorine related to copper in a chloridic environment peak decreasing faster than the tail, which is related to the oxychloride species. Bulk copper(II) oxychloride, $\text{Cu}_2\text{Cl(OH)}_3$, is rather insoluble in water [76], so one may think that if a surface ensemble/compound the like forms, than it probably remains adsorbed more persistently than the chloride ions, related to copper in a chloridic environment, which will be displaced preferentially. The presence of some oxychloride-like species may be a further explanation of the presence of some Cu(II) .

In general, for subsequent immersions in the same solution, the XPS signals vary in a similar way with increasing immersion times. The $\text{Cu } 2p_{3/2}$ signal changes shape, as previously described, and decreases, whilst the $\text{N } 1s$ increases after the first measurement, as BTAH adsorbs, and then decreases. The $\text{Cl } 2p$ decreases, the $\text{C } 1s$ increases dramatically after the first immersion, then stays approx. constant, the $\text{O } 1s$ generally increases. The peak shapes do not change drastically after the first immersion, indicating that the chemistry of the system is pH dependent rather than exposure dependent. In particular, formation of a full BTA-Cu layer does not seem to occur, as will be elucidated later.

A further insight into the copper oxidation state is offered by evaluating the modified Auger parameter, α' [46], defined as the sum of the binding energy of the most intense $\text{Cu } 2p_{3/2}$ photoelectron line and the kinetic energy of the most intense Cu LMM Auger line, for each of the preparations. The respective peak positions were read on the raw spectra and the results are summarised in Table 2. After preparation of the Cl-capped Cu layer, an α' parameter of 1849.7 eV is calculated. When comparing with the values reported in Table 1, this is closer to α' of bulk CuCl_2 , however the shape of both $\text{Cu } 2p_{3/2}$ and LMM clearly indicate that the most prevalent copper oxidation state is +1. Although α' parameters evaluated for the samples immersed in the BTAH solutions at pH 6 and pH 5 are all consistent with copper being in +2 oxidation state, the shape of the $\text{Cu } 2p_{3/2}$ shows that only a small amount of Cu(II) is present, with the majority still being in favour of Cu(I) . The discrepancy is likely due to the fact that the values calculated in the present work, regarded as thin film surface species, are compared with values obtained for bulk compounds. In fact the morphology of the

adsorbate has been shown to give unexpected shifts in the Auger lines [77]. Therefore, not only the energies of the photoelectron and Auger lines, but also their shapes have to be taken into account. The α' parameters evaluated for pH 4 are more indicative of the presence of Cu(I) instead, highlighting the different behaviour in the more acidic solution.

preparation		Cu 2p _{3/2} BE /eV	Cu LMM KE /eV	α' /eV
UPD		932.3	917.4	1849.7
pH 6	30 min	932.3	917.3	1849.6
	75 min	931.9	917.6	1849.5
	120 min	931.8	918.0	1849.8
pH 5	30 min	932.7	917.8	1850.5
	75 min	932.8	917.8	1850.6
	120 min	932.7	917.8	1850.5
pH 4	30 min	932.9	915.2	1848.1
	75 min	932.7	915.7	1848.4
	120 min	932.8	914.7	1847.5

Table 2. Cu 2p_{3/2} binding energy, Cu LMM kinetic energy and modified Auger parameter values evaluated from the raw spectra.

STM images show that the layers prepared by immersion at different pH exhibit very similar topographical features. Benzotriazole adsorbed species are imaged as elliptical shapes with an average separation of 0.35 ± 0.02 nm, a value which well-agrees with the average spacing of a chloride terminated overlayer with a (5×5) geometry, over a Cu (1×1) layer, commensurate with the underlying Au(111) [42]. Adsorption has to occur by displacement of the chlorine atoms and may be favoured in solution because chloride is solvated by water more effectively than BTAH [73]. Molecular features are seen to propagate in three directions rotated by 120° with respect to each other. Molecular ensembles, however, do not have a regular width and don't propagate in a straight line. Adsorption of an upright Cu(I)-BTA compound in bidentate configuration, whereby each deprotonated BTAH molecule coordinates with two copper atoms through either the N₁ and N₂ nitrogen atoms, or the equivalent N₂ and N₃, can explain the variations seen. In fact, even though chemically indistinguishable, adsorption through one of the two couples will result in a molecule tilting in the opposite direction as if adsorption were to occur through the other. Considering two molecules adsorbed upright and with the planes containing the aromatic systems parallel to each other, successive adsorption through N₁ and N₂ (or the equivalent N₂ and N₃) will result in a geometry which may favour the superposition of the π systems leading to a π - π interaction. However, the alternation of a molecule adsorbed through N₁ and N₂ with one adsorbed through N₂ and N₃ will result in a geometry not compatible with a π - π superposition. It has been reported that, in general, in a π - π interaction, aromatic rings are parallel displaced in an offset or slipped stacking configuration more often than not [47]. Such parallel-displaced structures are also thought to have a contribution from a σ - π interaction, the more favourable with increasing offset. If this were the case, a chain

with regular width would be expected. The balance between $\sigma - \pi$ and $\pi - \pi$ [78, 79] and surface – molecule interactions has been reported to have an effect at much larger range than that observed in the present study. For example, they have been shown to produce unexpected 2D structures, like supramolecular corrals, on which the closest separation between the observed features is calculated as 3.2 Å for dimer pairs [80], and 3D Kagome-like lattices, where π - π stacked molecular dimers adsorb perpendicularly over a layer of flat-lying molecules [81].

Perhaps a further factor which can have an effect in the molecular orientation is due to the varying spacing of the (5×5) chloride layer, i.e. when molecules would be too close to each other an offset configuration would be preferred. The featureless areas on the STM images are occupied by the chlorine atoms, which are not resolved at these specific scanning conditions.

With increasing exposure, STM images shows a small increase in surface coverage, however XPS seems to show a decrease of the copper, chlorine and nitrogen signals, accompanied by an increase of the carbon and oxygen ones. Because adsorption phenomena normally occur on a much quicker timescale than our first measurement (i.e. 30 minutes of immersion time), our results are more representative of an equilibrium state between molecules adsorbing on the surface and going back into solution. The increase in coverage evaluated from both XPS measurements and STM images is not linear, suggesting that when some adsorption sites are occupied by BTAH, some adjacent sites may be destabilised, therefore further adsorption is rendered more difficult. Such destabilisation effects could be electronic, because the remaining chloride ions may be more difficult to displace from potential adsorption sites, because they more strongly bound, or physical, because the presence of BTAH molecules hinders further BTAH from adsorbing. The competitiveness of adsorption between BTAH and chlorine may account for the observation of short molecular ensembles, which grow very slowly with prolonged immersion times. The increased number of chemical species present in the system, copper in a chloridic environment, Cu-BTA, adventitious contaminants, renders extremely difficult to establish a precise Cu:BTA stoichiometry via XPS. However, on the basis of the STM images, as molecules are through to adsorb upright, a Cu:BTA \approx 2:1 stoichiometry would correspond to a bidentate adsorption (through N₁ and N₂, or equivalent N₂ and N₃). A Cu:BTA \approx 1:1 stoichiometry would be expected for monodentate adsorption (through the N₁ (N₃) or N₂ atom) instead. Computational studies report that for neutral BTAH, “bridge-N₂+N₃” adsorption is slightly more favourable than “top-N₃” and “top-N₂”, whereas bidentate adsorption is strongly favoured for dehydrogenated BTA[⊙] [33, 34, 73, 82].

Previous STM studies have proposed that on Cu(100) in a H₂SO₄ electrochemical environment, depending on the potential employed, benzotriazole can adsorb in different configurations: an ordered phase of flat lying molecules and disordered chains with a spacing of 0.42 nm, compatible with the 4.13 Å lattice spacing along the *c*-axis of crystalline BTA, along which the molecules are oriented almost parallel to each other [19]. This suggests a parallel upright stacking of BTA molecules for the disordered chains, adsorbed through the nitrogen lone pairs [19]. On Cu(111), a poorly ordered adlayer phase containing individual BTAH chains with a interspacing of approximately 4 nm along the chain, orientated upright with respect to the surface was

observed [20]. In a H_2SO_4 electrochemical environment, the observation that BTAH formed more ordered adlayers, for a wider range of potentials, on Cu(100) than on Cu(111) was linked to the increased resistance of the (100) BTA covered surface towards the adsorption of chlorided ions [83]. In a hydrochloric acid environment, BTAH was seen to adsorb with its molecular plane upright or slightly tilted in a double row structure [21]. Also in a HClO_4 environment, BTAH molecules were found to adopt an upright adsorption geometry and being stacked into straight molecular rows on Cu(111), whereas on Cu(110) and Cu(100) a flat-lying orientation was detected [84].

Under UHV conditions on Cu(111), BTAH was found to adsorb predominantly as a $\text{Cu}(\text{BTA})_2$ dimer, whereas some monomers were also seen [28, 29]. In the low coverage regime features widths of ca. 0.5 nm and ca. 1 nm, for Cu-BTA and BTA-Cu-BTA units respectively, were reported. On oxygen free and oxygen covered Cu(110), BTAH was seen to adsorb forming ordered $c(4 \times 2)$ domains and disordered phases respectively, with preference for adsorption on the oxygen covered areas [14, 15].

5. Conclusions

The behaviour of Cl-capped Cu(111) monolayers prepared via UPD methods on Au(111)/mica immersed in BTAH solutions at different pH was investigated via X-ray photoelectron spectroscopy (XPS) and scanning tunnelling microscopy (STM).

XPS revealed that a Cu(I)BTA surface compound is formed upon immersion in the BTAH solutions. While a small amount of copper in a +2 oxidation state is seen initially, Cu(I) is the species always prevalent. The layers appear to be chemically very similar; however, the interaction with BTAH seems more favourable at lower pH, when BTAH is more effective in displacing the chlorine atoms from the copper layer. As the amount of BTAH grows very slowly with increasing exposure, this may represent an equilibrium situation in which BTAH molecules and chloride ions are coadsorbed on the copper layer and competitive.

STM measurements show that BTAH adsorbs as upright species, parallel stacked, with an intermolecular distance compatible with a π - π interaction. Although a BTA-Cu metalorganic compound forms upon adsorption, this is in the form of discrete units and an extended metal-organic polymer was not identified.

The observed features appeared having similar morphology, independently of pH and immersion time.

This study contributes to a further understanding of the interaction between benzotriazole and copper in solution, with focus on the species adsorbed on the copper, by highlighting its complexity and pH dependency.

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Supplementary material: cyclic voltammetry, fitted XP and Auger spectra, reference Auger spectra, additional STM images.

Author's declaration

All authors have approved the final version of the manuscript.

The authors declare no conflict of interest.

Data availability

The raw data required to reproduce these findings are available to download from

<http://dx.doi.org/10.17630/3175f32b-2508-441f-9f3e-e86c5604f076>

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