

Tribological characteristics of self-assembled nanometer film deposited on phosphorylated 3-aminopropyltriethoxysilane

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Abstract. Thin films deposited on the phosphonate 3-aminopropyltriethoxysilane (APTES) self-assembled monolayer (SAM) were prepared on the hydroxylated silicon substrate by a self-assembling process from specially formulated solution. Chemical compositions of the films and chemical state of the elements were detected by X-ray photoelectron spectrometry (XPS). The thickness of the films was determined with an ellipsometer, while the morphologies and nanotribological properties of the samples were analyzed by means of atomic force microscopy (AFM). As the results, the target film was obtained and reaction might have taken place between the thin films and the silicon substrate. It was also found that the thin films showed the lowest friction and adhesion followed by APTES-SAM and phosphorylated APTES-SAM, whereas silicon substrate showed high friction and adhesion. Microscale scratch/wear studies clearly showed that thin films were much more scratch/wear-resistant than the other samples. The superior friction reduction and scratch/wear resistance of thin films may be attributed to low work of adhesion of nonpolar terminal groups and the strong bonding strength between the films and the substrate.

Keywords. Thin films; friction; adhesion; scratch/wear.

1. Introduction

Microelectromechanical systems (MEMS) and emerging nanoelectromechanical systems (NEMS) are expected to have a major impact on our lives, much like the way that the integrated circuit has affected information technology (Anonymous 1977; Roukes 2001). However, due to the large surface area to volume to volume ratio in MEMS/NEMS devices as the size scale shrinks, currently many potential applications for MEMS/NEMS are not really practical. Many studies have revealed the profound negative influence of stiction, friction and wear on the efficiency, power output, and steady-state speed of micro/nanodynamic devices (Bhushan 1998; Bhushan 1999; Kayali *et al* 1999; Arney 2001). The self-assembled monolayer (SAM) has gained growing interest over the past years because it has advantageous characteristics of well-defined structure, strong head group-substrate binding and dense packing of hydrocarbon chains. Indeed SAM considerably reduces friction and adhesion and is found useful in various microelectromechanical systems (MEMS) devices (Depalma and Tillman 1989; Bhushan *et al* 1995; Tsukruk *et al* 1996; Boshui *et al* 1998; Dugger *et al* 2000).

A number of studies have been done on the nanotribological properties of different SAMs (Cha and Kim

2001; Bhushan 2005), but the study of the rare earth films on the nanotribological behaviour is still lacking. In this study, thin films deposited on the phosphorylated 3-aminopropyltriethoxysilane (APTES) self-assembled monolayer (SAM) were prepared on the silicon substrates. Different means, such as X-ray photoelectron spectroscopy (XPS), atomic force microscopy (AFM), ellipsometer and contact angle measurements, were applied to investigate the structure and nanotribological properties of the prepared films.

2. Experimental

2.1 Preparation of thin films on the phosphorylated APTES-SAM

APTES was purchased from Aldrich Chemical Company, Inc. A single-crystal silicon wafer polished on one side was used as substrate for the SAM transfer. Other reagents were of analytical grade. Deionized water was used throughout the experiment. For the preparation of the target solution, APTES, toluene, ethanol, acetonitrile, phosphorus oxychloride, and collidine were commercially obtained and used without further purification. The silicon substrates were cleaned with ‘piranha’ solution ($H_2SO_4:H_2O_2 = 7:3$ (v:v)) (caution: this solution reacts violently with organics), then exposed to a solution of APTES in toluene (2%V:V, 24 h,

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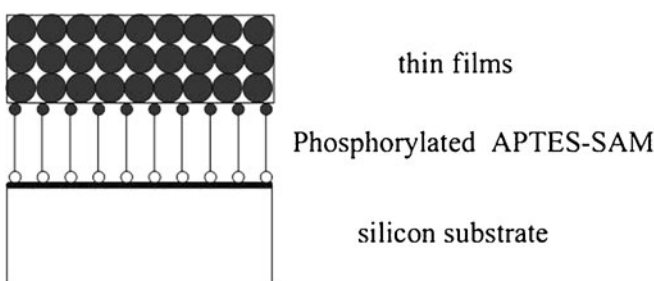


Figure 1. Schematic of growth of thin films.

room temperature), followed by POCl_3 in acetonitrile (0.2 M POCl_3 , 0.2 M collidine, 20 min, room temperature), according to the described procedure (Katz *et al* 1991; Buscher *et al* 1996). This treatment resulted in surface richness in phosphonate groups ($-\text{PO}(\text{OH})_2$), which adsorbed a layer of thin films when immersed in pre-prepared solution, then naturally cooled in a desiccator. Schematic of growth of thin films on phosphorylated APTES-SAM is shown in figure 1.

2.2 Description of apparatus and test procedures

Chemical compositions of the films and chemical state of the elements were analyzed by a PHI-5702 XPS system, using $\text{Mg-K}\alpha$ radiation operating at 250 W and pass energy of 29.35 eV. The binding energy of C 1 s (284.6 eV) was used as the reference. The resolution for the measurement of the binding energy is about ± 0.3 eV. The static contact angles were measured in ambient air (relative humidity 40%) using an OCA-20 contact angle measurement device (DataPhysics Instruments GmbH). Distilled water was used as the spreading reagent. Measurements were done at least on three samples, and were made at a minimum of three different spots on each sample. The contact angles were typically reproducible to within $\pm 2^\circ$. The thickness of the films was measured on an ellipsometer (V-VASE with AutoRetarder from J.A. Woollam Co., polarizer-retarder-sample-rotating analyzer configuration) which was equipped with a He-Ne laser (632.8 nm) set at an incident angle of 70° . The index of refraction for the refraction was taken to be 1.45. Thickness data was obtained by averaging five measurements at different spots of each sample surface. The thickness was recorded to an accuracy of ± 0.3 nm.

The surface morphology and the nanotribological properties of the prepared films were investigated using an SPM-9500 atomic force microscope (NanoScope IIIa) produced by Shimadzu Corporation (Kyoto, Japan). Square pyramidal Si_3N_4 tips with a nominal 50 nm radius mounted on gold-coated triangular Si_3N_4 cantilevers with spring constants of 0.6 N/m were used. The adhesion and friction data were measured ten times at each interesting location and average data values were obtained. Adhesive forces were measured using the so called ‘force calibration plot’. Friction forces

were measured according to the procedures given by Binnig *et al* (1986). By following the friction force calibration procedures developed by Glosli and Gary McClelland (1993), voltages corresponding to friction forces can be converted to force units. Coefficient of friction is obtained from the slope of friction force data measured as a function of normal loads.

For the scratch and wear tests, specially fabricated microtips were used. These microtips, consisted of single-crystal natural diamond, ground to the shape of a three-sided pyramid, with an apex angle of 80° and tip radius of about 50 nm, were mounted on a platinum-coated stainless steel cantilever beam whose stiffness was 50 N/m. Samples were scanned orthogonal to the long axis of the cantilever with loads ranging from 20 to 100 μN to generate scratch/wear marks. Observations of the sample surface before and after the wear tests were done by scanning parallel to the long axis of the cantilever with loads ranging from 0.5 to 1 μN . The parallel scans enabled near-zero wear of the sample surface and also eliminated post-data analysis errors in surface feature height. Scratch/wear tests were performed over a scan area of $2 \times 2 \mu\text{m}$ at a scan rate of 10 Hz. The reported scratch/wear depths are an average of six runs at separate instances. In this study, all the measurements were carried out in ambient conditions (22°C, RH 40–44%).

3. Results and discussion

3.1 Characterization of the prepared films

Figure 2 shows a series of AFM images taken over regions $1.0 \times 1.0 \mu\text{m}$ of specimens at various stages of the film deposition process, where panel (a) refers to the bare cleaned silicon substrate, (b) APTES-SAM on the silicon substrates, (c) the phosphorylated APTES-SAM, and (d) to the as-deposited thin films on the phosphorylated APTES-SAM. It can be seen that the surface of the silicon substrates (figure 2(a)) is clean and smooth with surface root-mean-square (rms) roughness in the range of 0.2–0.3 nm. Figure 2(b) is uniform and homogenous with surface rms roughness about 0.522 nm. The phosphorylated APTES-SAM (figure 2(c)) becomes more and more uniform and homogeneous with rms roughness about 0.393 nm. The possible reason is that the size of $-\text{PO}(\text{OH})_2$ groups is bigger than the terminal groups $-\text{NH}_2$ which leads to the $-\text{PO}(\text{OH})_2$ terminal molecules provided a more densely packed arrangement than the $-\text{NH}_2$ terminal molecules. After the deposition of the thin films (figure 2(d)) on the phosphorylated APTES-SAM, many differences are visible in the corresponding AFM images, viz. the surface of the as-deposited films is rough and quite densely round-looking particles with the rms roughness to be about 0.863 nm, which shows wide potential application in lubrication and wear protection.

The XPS spectra applied to detect the chemical states of some typical elements for the prepared films are shown in

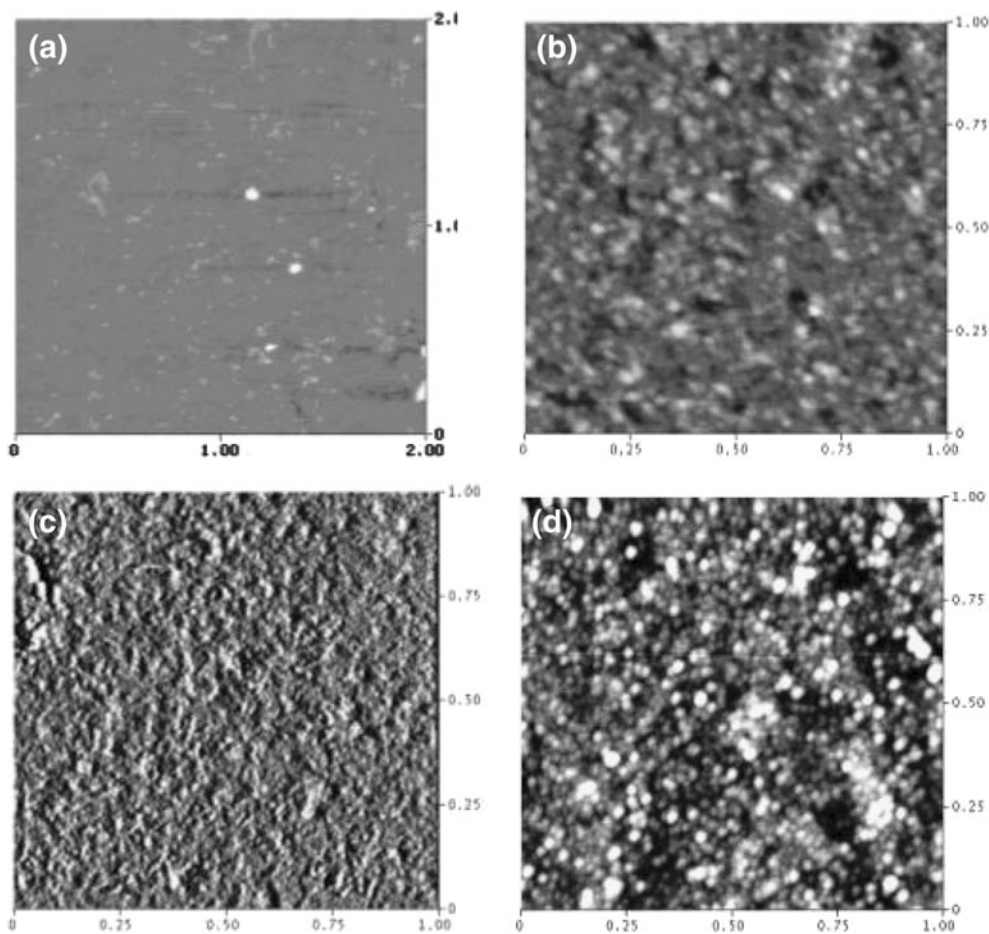


Figure 2. AFM images of (a) bare silicon substrate, (b) APTES-SAM, (c) phosphorylated APTES-SAM and (d) thin films.

figures 3–6. XPS survey and single scan spectra of APTES-SAM (figure 3) shows contributions from the substrate and the film are: silicon (15.5%), carbon (56.3%), oxygen (20.2%), and nitrogen (8.0%). Nitrogen is detected indicating successful APTES-SAM deposition, since this element is contained only in this film material. Figure 4 shows single-scan XPS spectra of N 1s peak decomposed into two different nitrogen species occurring in different binding states. The peak at 400.8 eV is assigned to the protonated aliphatic amino groups, while that at 399.5 eV may be ascribed to the aliphatic amino groups. This interpretation is consistent with the one given by Bierbaum *et al* (1995).

After *in situ* phosphorylation of the APTES-SAM, the P2_p at 134.5 eV is observed (figure 5), which is assigned to the P-atoms in –PO(OH)₂ group and N signal is absent. It indicates that the terminal –NH₂ group in the APTES-SAM has been phosphorylated and transformed to –PO(OH)₂ group successfully and completely.

The contact angles of distilled water on silicon substrate and the prepared films were measured using a contact angle measurement. The APTES-SAM has a contact angle of $50 \pm 2^\circ$, which is consistent with a moderately polar surface where

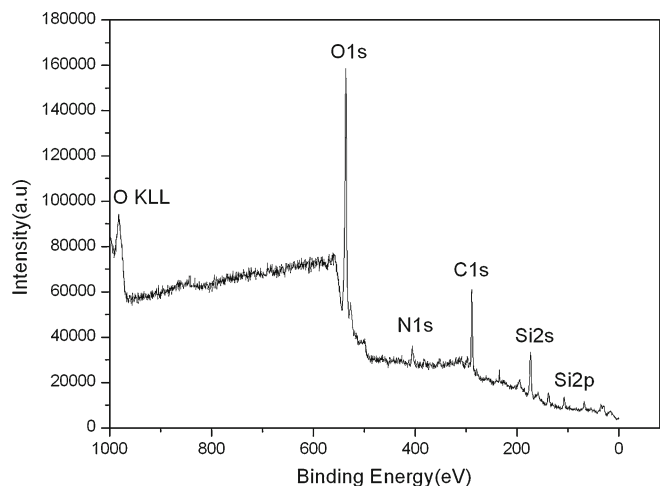


Figure 3. XPS survey and single scan spectra of APTES film.

the amino group is oriented upward. After phosphorylated *in situ* for 20 min, a contact angle of $24 \pm 2^\circ$ is recorded for the phosphorylated surface. The contact angle of thin films

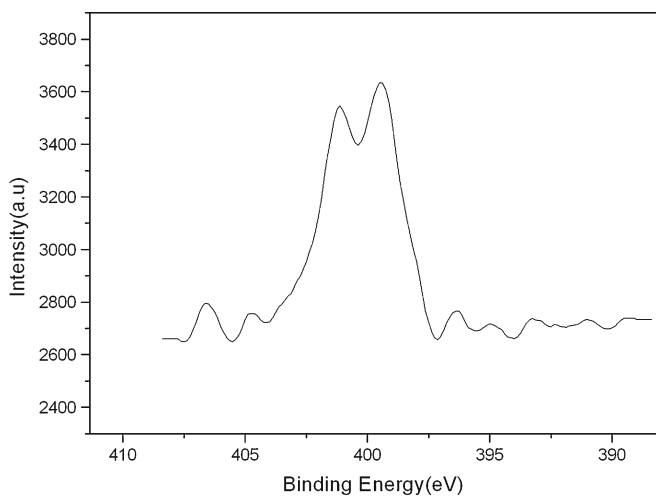


Figure 4. XPS spectrum of the N 1 s region of an APTMS film.

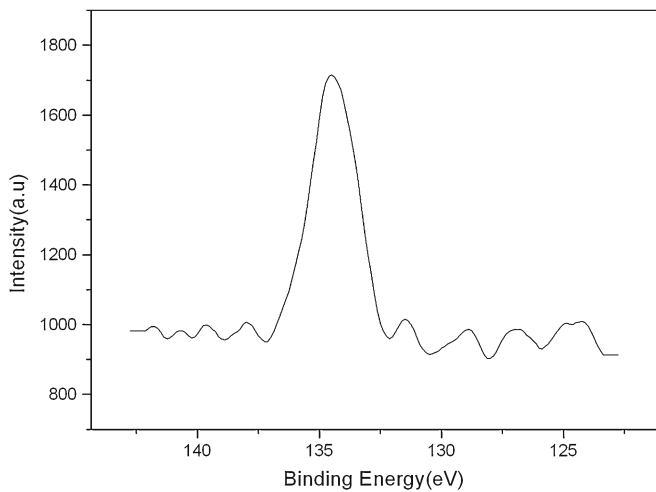


Figure 5. XPS spectrum of the P_{2p} region in APTMS film.

deposited on the phosphorylated APTES-SAM increases to $66 \pm 2^\circ$.

In our work, the thickness of the prepared films on glass substrates is determined with ellipsometer. The averaged thickness of the APTES-SAM is about 7.5 nm, which matches with the projection of a normally extended molecular chain on the surface. The thickness is hardly changed after the $-NH_2$ is phosphorylated to $-PO(OH)_2$ group, which indicates that a monolayer of phosphorylated APTES has been prepared on glass substrates. The thickness increases by about 15 nm after the phosphorylated APTES-SAM is immersed in solution, which also shows that thin films have been successfully obtained.

3.2 Nanotribological properties of prepared films

3.2a Adhesion, friction and work of adhesion: Figure 6 shows the average values of the adhesion force and

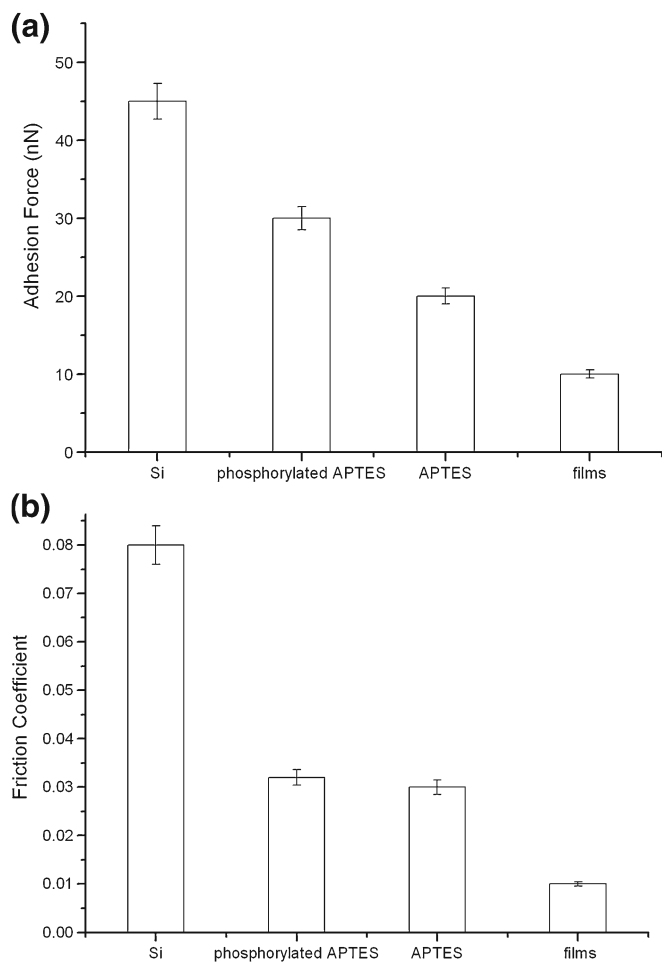


Figure 6. (a) Adhesive forces of silicon and prepared film and (b) coefficients of friction of silicon and films.

coefficients of friction of four kinds of flat surfaces measured by contact mode AFM under an applied normal load of 20 nN and a scan rate of 10 Hz. It shows that SAMs can reduce the adhesive and frictional forces of silicon substrate. In particular, thin films exhibit the lowest adhesive force and coefficient of friction. It means that the prepared films can be used as effective molecular lubricants for micro/nanodevices fabricated from silicon. Based on the data, the ranking of adhesive forces F_a is in the following order: $F_{a-Si} > F_{a-phosphorylated\ APTES} > F_{a-APTE} > F_{a-films}$. And the ranking of the coefficients of friction is in the following order: $\mu_{Si} > \mu_{phosphorylated\ APTES} \approx \mu_{APTES} > \mu_{films}$.

Based on Young–Dupre equation, work of adhesion W_a (the effort required to pull apart the unit area of the solid interface (Israelachvili 1985)) was summarized:

$$W_a = \gamma_{1a} (1 + \cos \theta_1), \quad (1)$$

where γ_{1a} is the surface tension of liquid–air interface and θ_1 is the water contact angle of liquid and flat. From figures 6 and 7, it is clear that the adhesive force and friction coefficient decreases as the work of adhesion decreases. It implies

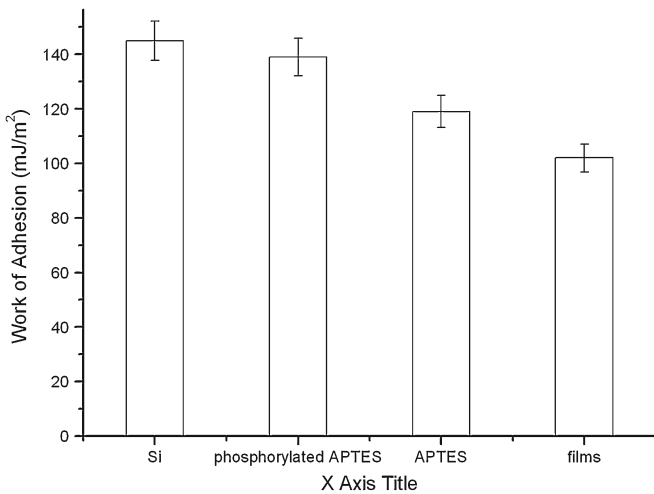


Figure 7. Work of adhesion of friction of silicon and films.

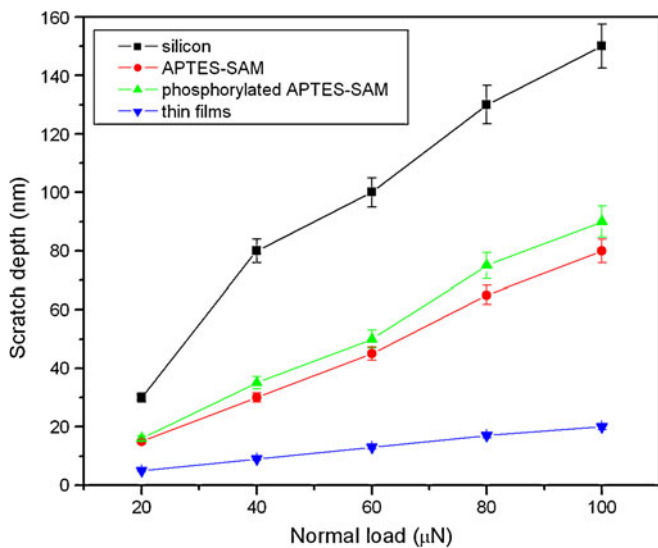


Figure 8. Scratch depths for 10 cycles as a function of load.

that the capillary force acting between the tip and the flat surfaces affected seriously on nano-adhesion and nano-friction. From figure 6, it can be also found that APTES-SAM and phosphorylated APTES-SAM have polar surface groups ($-NH_2$ and $-PO(OH)_2$ groups), thus lead to larger W_a and eventually larger adhesive forces. The films do not have polar surface groups, thus have a smaller W_a and adhesive force than APTES-SAM and phosphorylated APTES-SAM.

3.2b Scratch/wear tests: As explained earlier, the scratch tests of silicon and the prepared films were studied by making scratches for 10 cycles with varying loads. Figure 8 shows a plot of scratch depth vs. normal load for various samples. Scratch depth increases with increasing normal load. APTES-SAM and phosphorylated APTES-SAM show similar scratch resistance. From the data, it is clear that the thin

films show the best scratch resistance compared with the silicon substrate, APTES-SAM and phosphorylated APTES-SAM. The increase in scratch depth with normal load is very small and all depths are less than 20 nm, while the silicon substrate and APTES-SAM finally reach depths of about 80 nm and 150 nm, respectively.

Wear tests were conducted on the samples by wearing the same region for 30 cycles at a normal load of 20 μ N, while observing wear depths at different intervals (1, 5, 15, 20 and 30 cycles). This would give information as to the progression of wear of the films. The wear depths observed are plotted against the number of cycles in figure 9. For all the samples, the wear depth increases almost linearly with increasing number of cycles. This suggests that material is removed layer by layer in all the materials. Here also, thin films exhibit lesser increase in wear depth (slope) than the other samples. APTES-SAM wears less than the silicon substrate, which shows similar wear characteristics to the phosphorylated APTES-SAM. Combined with the scratch/wear data and the thickness of the films, it can be found that the scratch/wear depth of APTES-SAM and phosphorylated APTES-SAM are much more than their thickness (7.5 nm). On the contrary, the scratch/wear depth of thin films does not exceed its thickness (15 nm) all the time. It shows that APTES-SAM and phosphorylated APTES-SAM are seriously destroyed with the increase of normal load and number of cycles, while thin films are still not completely worn at last. The scratch/wear results all indicate that thin films have better surface mechanical properties than silicon substrate, APTES-SAM or phosphorylated APTES-SAM.

The $-PO(OH)_2$ group on the APTES-SAM provides net negative charge, thus promoting the process of self-assembly. Then $-PO(OH)_2$ can provide a P=O bond as complexing group to react after adsorption, thus improving the bonding strength between the films and the silicon substrates. In

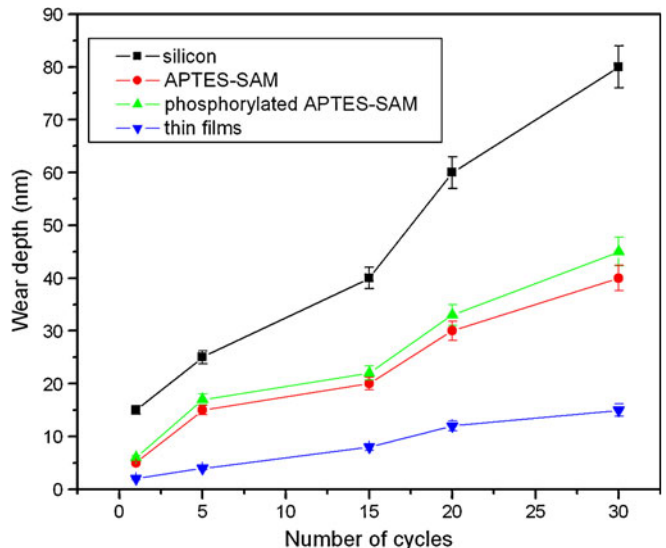


Figure 9. Wear depths as a function of number of cycles.

addition, Hertzian cone cracks can occur when the normal stress exceeds a critical value as the AFM tip slides over the surface. Friction forces during sliding reduce this critical value. In short, strong bonding strength between the films and the silicon substrate and low coefficient of friction are responsible for the superior scratch/wear resistance of thin films.

4. Conclusions

In this work, the nanotribological properties of silicon substrate, APTES-SAM, phosphorylated APTES-SAM, and the thin films were characterized by an AFM. The thin films showed the lowest friction and adhesion followed by APTES-SAM and phosphorylated APTES-SAM, while silicon substrate showed high friction and adhesion. Microscale scratch/wear studies clearly showed that thin films were much more scratch/wear resistant than the other samples. The superior friction reduction and scratch/wear resistance of thin films may be attributed to low work of adhesion of non-polar terminal groups and the strong bonding strength between the films and the substrate.

It is thus concluded that the prepared thin films could be used for protection from scratching as well as reducing friction.

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