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Self-Healing Fluoropolymer Brushes as Highly Polymer-Repellent Coatings

Zhanhua Wang, ab and Han Zuilhofbc, *

The fouling of surfaces by organic polymers can be strongly reduced by applying a 75 nm covalently bound fluorinated polymer brush onto the surface. This strong reduction can be repaired, even more than 10 times, after the polymer brush has been damaged (e.g. by strong base) using the self-repairing character of these brushes via molecular reorganization at the surface-air interface at slightly elevated temperatures.

The fouling of surfaces by biological entities (from isolated proteins and other biopolymers to entire organisms) has been studied in great detail. This has led to a deep understanding of at least the initial stages of biofouling and the design of novel materials that successfully repel such fouling. Probably the best known examples are the first-generation antifouling polyethylene oxide polymers and the currently state-of-the-art zwitterionic polymer brushes. Such progress has been nearly absent in the field of fouling by polymers in non-aqueous, organic media. Such fouling is of significant industrial importance, e.g. in food processing, paper manufacturing, and high-resolution 3D printing. Yet, there is only very little literature on this topic that probes the mechanisms behind it down to a molecular level.

Recently, we reported the first systematic study of fouling by a wide range of polymers with variable molecular weight onto high-quality fluorinated monolayers onto ultraflat Si surfaces. This choice was based on the high degree of control that is available in the construction of such monolayers, whereas the ultraflat substrate rigorously decouples the contributions of molecular structure and surface roughness. In addition, the choice of Si was practically driven by the fact that small-scale orifices used in industry are increasingly lithographically prepared from Si. Fouling over time by polymers therefore limits their long-term use. In this first study we found that partially fluorinated monolayers, especially mono-fluoro alkyne-derived Si-CH=CH-C\textsubscript{13}H\textsubscript{28}CH\textsubscript{2}F monolayer F1, showed excellent antifouling behavior against a range of polymers with different molecular weights even on atomically flat Si surfaces. However, while extremely successful for most of the tested polymers, in two regimes limitations were observed. First, for a variety of polymers fouling was still observed in solvents of low polarity. Second, for a special polymer P2VP, heavy fouling was also found on the F1 monolayer due to the strong interaction between the single C-F bond and protonated N atom in P2VP. Finally, a generic limitation of organic monolayers is their susceptibility to mechanical damage, and the absence of self-repair. Self-repair is a highly desirable property, and significant successes in the field of self-repairing and antifouling surface have been achieved by e.g. the groups of Aizenberg (based on liquid captured within nanostructures), Minko (using flexible polymers), Sun (using layer-by-layer assembled films), and Zeng (using catechol-mediated hydrogen bonding interactions and aromatic interactions). Challenges in this field are still the evaporation of the infused liquid, the necessity of complicated fabrication of the required micro/nano structures or the limitation to use in an aqueous environment, use of preferably ultrathin surface coatings, and overall: the full repair of surface properties. Therefore, long-term anti-fouling coatings with a self-healing function pre-incorporated inside the materials by facile methods that can be widely used still deserve significant attention.

In the current communication we aim to address all these issues by the development of a covalently grafted fluoropolymer brush (75 nm thickness) onto atomically flat Si(111) surfaces using atom transfer radical polymerization (ATRP; Fig. 1a). The surface functionalization and initiator immobilization were achieved by standard methods onto atomically flat, oxide-free Si(111) surfaces, and the resulting surfaces – like all surfaces under study – were characterized in detail by static water contact angle (CA) and X-ray photoelectron spectroscopy (XPS) (Table S1 and Fig. S1). Next,
the poly(2-perfluorooctylethyl methacrylate) (PMAF17) brush was
grafted to a carefully controlled thickness (75 nm) onto
the initiator-immobilized Si surface via surface-initiated
ATRP. This polymer brush was selected because of its high
hydrophobicity (CA = 121°) and ultra-low surface energy. In
addition, rearrangement of the polymer segments above the
glass transition temperature (T_g) of the PMAF17 brush (T_g of
bulk PMAF17 = 40 °C) was hypothesized to induce the
fluorinated tail to come to the top of the surface to decrease
the surface interaction between the brush and air. This
property will endow the PMAF17 brush with self-healing
character, due to the reregulating chemical composition at the
surface via the movement of the polymer segments (Fig. 1b).
Furthermore, such covalent grafting endows fluoropolymer
brushes with an improved stability and durability.

The brush-coated surface is rather flat, as displayed by a
low surface roughness (see below: RMS = 0.87 nm, Fig. 4 C1),
compared to its thickness (75 nm). This thickness was chosen
as an optimum between two opposing trends: on the one
hand ultrathin coatings will be most conformal to surface
structures, and be useful on nanopatterned surfaces without
effectively removing that structuring. On the other hand we
reasoned that a minimal thickness is required, as only for
polymer coatings with a thickness > ca. 50 nm we expected the
fluoropolymer brush to possess roughly the same T_g as the
Corresponding bulk polymer. This means that above this
threshold thickness the thermo-induced movement of the
polymer brush is independent on the substrate, which is
favorited for the self-healing process. Furthermore, the thicker
the polymer brush, the more times the brush can be repaired,
eventually leading to longer service lifetimes. Making thicker
PMAF17 brushes is trivial by using e.g. longer reaction times in
the polymerization step, so the challenge is in getting polymer
layers as thin as possible while still displaying this character.
Finally, we chose the two most rigorously fouling conditions
found in our previous study 13 PS in highly apolar toluene and
strongly interacting P2VP in DMF as two model systems to
study the antifouling property of the PMAF17 brush.
Unmodified silicon surface, and hexadecenyl monolayers with
one (F1) or 17 fluorine atoms (Si-CH=CH-C6H12-C6F17; F17) (see
Scheme S1) were used as references.

To study the polymer antifouling properties modified
surfaces were dipped overnight in a solution containing the
polymer, taken out, and cleaned with the same solvent (Figure
1). The polymer fouling was studied by XPS, CA and a
previously developed bimodal atomic force microscopy (AFM)
based approach. 13 For each surface, the fouling experiments
were conducted on three different samples, and on each
sample the ellipsometry, XPS and AFM measurements were
carried out at at least three different places. Fig. 2 shows the
increased thickness of the four different surfaces after our
fouling protocol determined from the attenuation of XPS
signals of substrates in the presence of an overlayer (here: the
fouling polymer). The increased thickness on the unmodified
silicon and the F1 and F17 monolayers is determined by the
increased atomic C/Si ratio, 27 while for the PMAF17 brush the
increased thickness can be calculated by the increased C/F
ratio, since the fouling polymers do not contain any fluorine
atom in the polymer chain. As a generic finding, the fouling
deposit on the PMAF17 brush is < 0.1 nm for both PS
in toluene and P2VP in DMF. This result is evidently better
than for those of unmodified Si, and the F1 and F17
monolayers, indicating the as-prepared fluoropolymer brush
possesses superior antifouling towards the studied polymers.
The best antifouling properties were achieved by the
fluoropolymer brush due to its stronger hydrophobic
interaction with the solvent compared with the references.
Surface morphology surveys of these surfaces by tapping mode AFM measurements further confirmed the polymer adsorption behavior. As shown in Fig. 3, after the fouling study, no polymer particles were found for the PMAF17 brush, which further confirmed its superior antifouling properties towards polymers. In contrast, polymer particles are clearly observed on the F1 and F17 monolayers, resulting in higher roughness and confirming significant fouling of these monolayers under these conditions.

To determine the degree of fouling, we calculated the fraction of the surface that has a height (as determined by AFM) above a certain threshold value, and assume that exceedance of this threshold value is the result of polymer fouling as the substrate is atomically flat and the polymer brush smooth. The determined degree of fouling for the surfaces using a threshold value of 3 nm is shown in Fig 3. Dark blue regions correspond to areas above the threshold, and are labelled as fouled by polymer, while the yellow areas are classified as non-fouled areas. It is clear to see that for these two model systems, the fouled surface fraction of the PMAF17 brush is by far the lowest (less than 0.3%) compared with F1 and F17 monolayers, further confirming its superior anti-fouling properties, as in fact the F1 and F17 monolayers scored best as non-fouling coatings within a range of monolayers. The fouled surface fraction should, of course, decrease when increasing the threshold, but the variation from surface to surface does not vary substantially. The surface coverage on the PMAF17 brush is always lowest irrespective of the precise threshold chosen (Fig. S2). In addition, we also tested the anti-fouling property of the PMAF17 brush against some other polymers in different solvents (poly(acrylic acid) in methanol; poly(hydroxyl propyl methacrylate) in acetone; poly(4-chloro styrene) in ethyl acetate; polystyrene in dichloromethane; poly(adipic anhydride) in dimethylformamide; poly(N-isopropyl acrylamide) in trifluoroethanol). The AFM results in Fig. S3 illustrates that PMAF17 brushes display excellent antifouling towards all these polymers studied.

Compared with monolayers, another advantage of the polymer brush is the long-term stability. In order to investigate the stability of our fluoropolymer brush under stringent conditions, we immersed the PMAF17 brush into a swirling pH 11 solution for 24 h under standardized stability testing. Subsequently, the surface was washed by water, ethanol and dichloromethane, dried by argon and then characterized by CA, AFM and XPS. The water and toluene static CA of this damaged brush were still 118° and 71° (Fig. 4 A2 and B2), respectively, which illustrates that the fluoropolymer brush is highly stable in basic environment, and that neither the polymer brush nor the attachment to the surface are significantly damaged. However, the 3° and 4° decrease in water and toluene static CA does indicate some polymer breakdown, which was confirmed by a slight increase of the surface roughness (Fig. 4 C2). This increased surface roughness will likely also contribute to the relatively have wetting angle of the chemically degraded surface.

We hypothesized that above the T_g of the brush polymer (here: 40 °C) the polymer chain might display sufficient mobility to reorient itself and reform an optimal surface. The driving force for this process should then be the low surface

![Fig. 3. Top: Tapping mode AFM images (5 × 5 µm²) in air of the PMAF17 brush, F1 and F17 monolayers after 12 h immersion in polymer solution. Below: Corresponding surface coverage of the modified surfaces assuming a threshold of 3 nm.](image-url)

![Fig. 4. Static contact angle (water: A1-A3 and toluene B1-B3), tapping-mode AFM image (C1-C3) and XPS carbon narrow scan (D1-D3) of the original, damaged and repaired PMAF17 brush.](image-url)
energy of the fluorinated materials, which should cause undamaged fluorinated tails to come to the top of the surface during heating and repair the hydrophobicity and antifouling character of the surface. Indeed, and actually better than expected, the water and toluene static CA of the damaged polymer brush recovers to 121° and 76° after being heated at 120 °C for 2 h (Fig. 4 A3 and B3). 120 °C was selected here because this temperature is far above the Tg of the fluorinated methacrylate, and thus allows facile chain reorientation. We noticed that, apart from a return of the hydrophobicity, also the original low RMS roughness recovers (Fig. 4 C3), together with regaining the original XPS spectra, specifically the Cls XPS spectrum (see below). We thus hypothesize that, apart from reorientation of individual F17 side chains, also the polymer backbone rearranges. From studies of Takahara and co-workers it follows that C8F17 chains in such a brush preferentially form a tightly packed structure due to the low surface free energy, in which these side chains are oriented almost perpendicular to the substrate to afford a lamellar structure (Fig. 5A). Based on the changes in contact angle, increased polymer fouling, increased roughness and loss of F in the survey XPS spectrum, we reason that in addition to the loss of side chains during the damage process also this molecular orientation is partially lost, leading to e.g. a higher surface roughness (Fig. 5B). However, rearrangement of the polymer segments during heating will repair the top-layer molecular orientation (Fig. 5C), and the concomitant surface properties. The damaged brush hidden below may still influence the layer structure a bit, but will initially not influence the surface characteristics. Only upon further increased and repeated damage do these polymer reorientation processes not suffice anymore, and more permanent damage results.

![Schematic illustration of the polymer segment orientation of the PMAF17 brush during the damage and repair process.](image)

To confirm that these molecular reorientation process are indeed responsible for the damage-repair process detailed XPS studies were performed. Fig. 4 D1-D3 shows the C 1s narrow scan of the original, damaged and repaired fluoropolymer brush. The initial fraction of C atoms connected to fluorine is 57.3%, which is close to the theoretical value (57.1%; 8 out of the 14 C atoms in the monomer). The Cls XPS data clearly show that this value decreases to 49.2% upon alkaline damage, suggesting partial loss of fluorinated tails by basic hydrolysis of the ester group in the polymer backbone. This ratio recovers to 55.7% after heating, i.e. close to the initial value. We interpret this as that the carboxylic acid groups that are formed during the hydrolysis process are buried again by the regenerated top-layer fluorinated tail induced by the rearrangement of the polymer segments during heating. This damage-repair cycle of hydrophobicity can be repeated many times (Fig. 6D). The overall thickness of the PMAF17 brush decreased significantly after 12 damage-repair cycles, from 75 nm to 43 nm (Table S2). In line with the continuing accumulation of carboxylic acid moieties, the SCA decreases more dramatically after several alkaline treatments, but still the surface hydrophobicity and smoothness can be completely repaired. Apart from stability in a basic environment, the PMAF17 brush also possesses good stability in acidic media and under UV exposure. The water static CA only decreases to 116° and 119° at pH 3 for 24 h, and under UV (254 nm, 3 cm above the sample) exposure for 1 h, respectively. The CA of these damaged brushes in these two cases also recovers to the same values as the freshly prepared ones after heating at 120 °C for 2 h (Table S3 and S4). It is worth mentioning here that our surfaces can also be repaired several times at acidic environment or UV exposure. Here we only took basic environment as an typical damage condition to investigate the molecular event responsible for the damage-repair process and antifouling property of the brushes after self-healing, as the acid conditions or UV irradiation are typically less damaging for these surfaces, and base-induced damage thus displays the regeneration potential of the polymers under study most clearly.

Next, the polymer fouling behavior on the original, damaged and repaired fluoropolymer brushes was investigated by AFM and XPS. Here we used PS in toluene as a model system. As described above, the originally prepared brush displays excellent anti-fouling behavior (Fig. 6 A1 and B1), while heavy fouling was observed on the surface damaged at pH 11 for 24 h (Fig. 6 A2 and B2). Such heavy fouling was confirmed by polymer particles that could be found on the surface. And also the RMS surface roughness and the fouled surface fraction of the damaged brush increases dramatically compared with the freshly prepared brush [original RMS and surface coverage: 1.17 nm and 0.76%; RMS and surface coverage after damage: 2.70 nm and 6.18%]. In contrast, upon heating of a damaged surface at 120 °C for 2 h and cooling down, immersion of the resulting repaired surface into a fouling polymer solution did not yield any remaining fouling polymer particles after rinsing. Furthermore, the repaired surface also obtains the same surface roughness and fraction of surface fouling as the freshly prepared one (Fig. 6 A3 and B3). In addition, the XPS results determined by the atomic C/F ratio after fouling study exhibit the same trend as we got from the AFM measurements (Fig. 6C). These observations indicate that basically no fouling happens on the repaired fluoropolymer brush, which illustrates that the repaired PMAF17 brush displays the same highly favorable anti-fouling behavior as the undamaged one. We attribute the intermediate fouling upon damage of the brush to the reduced surface oleophobicity, which results from the hydrolysis of some ester bonds to carboxylates as also confirmed by XPS (see Fig 4 D2 versus D1 and D3).
Fig. 6. A1- A3: Tapping-mode AFM images of the original, damaged and repaired PMAF17 brushes after immersion into PS-toluene solution for 12 h, the inset value is AFM determined RMS. B1 – B3: Surface coverage of the original, damaged and repaired PMAF17 brushes after immersion into PS-toluene solution for 12 h, the inset value is AFM

determined RMS. C – The increased thickness by XPS of the original, damaged and repaired PMAF17 brushes after fouling study in PS-toluene solution. D – The static contact angle changes of the PMAF17 brush between pH 11 h and 120 °C 2 h.

In summary, a highly polymer-repellent fluoropolymer brush with excellent stability in basic/acidic environment or UV exposure was successfully prepared by surface-initiated ATRP. Upon damage at basic/acidic media or UV exposure, the hydrophobicity and antifouling character of the brush can simply be repaired many times by heating. It is likely that this type of chemistry can be combined with e.g. micro- and nanostructuring and other advanced materials properties to yield even better long-term antifouling behavior under harsh environments.33-36 Studies on the influence of the molecular architecture and degree of fluorination on the self-healing and antifouling properties are currently ongoing in our lab.

Notes and references
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