**Effect of H$_2$S Plasma Treatment on the Surface Modification of a Polyethylene Terephthalate Surface**

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**Abstract:** H$_2$S plasma created by an electrode-less radio-frequency discharge was used to modify the surface properties of the polymer polyethylene terephthalate. X-ray photoelectron spectroscopy, secondary ion mass spectrometry and atomic force microscopy were used to determine the evolution of the surface functionalities and morphology. A very thin film of chemically bonded sulfur formed on the surface within the first 10 s of treatment, whereas treatment for more than 20 s caused deposition of higher quantities of unbonded sulfur. The sulfur concentration reached a maximum of between 40 and 80 s of plasma treatment; at longer treatment times, the unbonded sulfur vanished, indicating instability of the deposited sulfur layer. Large differences in the surface morphology were observed.

**Keywords:** hydrogen sulfide (H$_2$S); plasma treatment; polymer; polyethylene terephthalate (PET); surface modification

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1. **Introduction**

Plasma created in various gases are often used for altering the surface properties of polymer materials. Oxygen and nitrogen-containing plasmas (O$_2$, CO$_2$, N$_2$, NH$_3$) or their mixtures with noble gases, such as He and Ar, are commonly used for surface hydrophilization [1–5]. In contrast, fluorine-containing plasmas (CF$_4$, SF$_6$) are used for surface hydrophobization [6,7]. Recently, a few papers have reported on tailoring the surface properties by SO$_2$ plasma [8–10]. SO$_2$ plasma could be useful for polymer modification in biomedical applications, e.g., preparation of antithrombogenic surfaces or for altering cell adhesion. However, there are almost no reports in the scientific literature regarding the use of other sulfur-containing plasmas, such as H$_2$S, for polymer modification.

An interesting application of gaseous plasma created in H$_2$S is the decomposition of this hazardous gas into hydrogen and sulfur. Such decomposition has attracted serious attention recently because it enables the production of hydrogen from a hazardous H$_2$S waste gas, which is a byproduct of oil refinement [11–17]. Currently, the conventional treatment method for H$_2$S destruction is the Claus process, which produces sulfur, whereas hydrogen is converted to water and therefore lost [12,18]. This is the reason why plasma has been investigated as a promising alternative technique for H$_2$S decomposition into hydrogen and sulfur and thus for hydrogen production, which is of strong commercial interest [19–21].

Another possible application of H$_2$S plasma is the synthesis of thiol (-SH) groups on the polymer surfaces upon treatment in H$_2$S plasma. Such -SH functionalities can serve as anchoring sites for the immobilization of macromolecules in biosensors and disease diagnostics, or they can act as a support for gold nanoparticles [22–25]. Classical methods for the synthesis of thiol groups are based on wet chemistry procedures that use different solvents, require long reaction times and depend on substrate surface properties [22]. Therefore, dry, solvent-free processes, like gaseous plasmas, are a good alternative to classical methods. Thiry et al. synthesized a plasma polymer film containing -SH
When comparing the carbon spectra of various plasma-treated samples, we did not find a significant difference in the shape of the spectra. The only difference was observed when comparing the spectra of plasma-treated samples with the untreated sample. The local minimum at 286 eV between the C-C and C-H groups [22–25]. He did not use H2S plasma but performed plasma polymerization using propanethiol (CH3CH2CH2SH) as a precursor. Furthermore, other authors reported the formation of sulfur-rich surfaces by plasma polymerization using thiophene as a precursor [26–30].

The literature survey indicates interesting applications of H2S plasma in various fields, from oil chemistry to biopolymers; therefore, it is worth investigating the mechanisms of the interaction with solid materials. In this paper, we report observations on the surface modification of polyethylene terephthalate (PET) polymer treated in H2S plasma.

2. Results and Discussion

2.1. Plasma Characterization

Figure 1a shows an optical emission spectrum (OES) of radiofrequency (RF) H2S plasma created at a pressure of 30 Pa and RF power of 150 W. The spectrum is rich in S2 molecular transitions. Only a few atomic lines are observed, and they correspond to H α, H β, and S. The transitions at approximately 600 nm and 750 nm correspond to the H2 Fulcher band [31], whereas the transitions between 280 and 620 nm correspond to S2 molecular transitions [32]. The S2 emission band B-X from 283 to 306 nm is particularly strong. Figure 1b shows the OES spectrum of plasma during sample treatment. The samples were placed in the middle of the RF coil. The spectrum is similar to the spectrum obtained for the empty tube.

![Image of OES spectra](image)

**Figure 1.** OES spectra of H2S plasma (a) without; and with (b) PET samples.

2.2. Surface Chemistry

Figure 2 shows the surface composition of PET samples treated in H2S plasma for various periods as deduced from the XPS (X-ray photoelectron spectroscopy) survey spectra. The samples were not rinsed before analysis to remove loose material but were rather quickly transferred to an XPS chamber to minimize aging effects. Plasma treatment resulted in the appearance of a significant sulfur content at the surface. The sulfur content was not constant, and it depended on the treatment time. At lower treatment times, the sulfur content was approximately 14 atomic %. At 40 s of treatment, the sulfur content increased significantly, and it reached a maximum of 40 atomic % at 80 s of treatment. At longer treatment times, the sulfur concentration slowly decreased, and at 640 s of treatment, the content was only 9 atomic %. The opposite variation of the concentration was observed for carbon and oxygen, i.e., the minimum concentration of carbon and oxygen was observed at the maximum sulfur concentration.

To explain the unusual behavior of the sulfur concentration, we recorded high-resolution XPS spectra to obtain additional information regarding the surface chemical composition. Figure 3 shows the high-resolution XPS spectra of the carbon and sulfur peaks for the selected treatment periods. When comparing the carbon spectra of various plasma-treated samples, we did not find a significant difference in the shape of the spectra. The only difference was observed when comparing the spectra of plasma-treated samples with the untreated sample. The local minimum at 286 eV between the C-C and
We observe the following major characteristic PET fragments: m/z 76 (C7H5O2+), 104 (C7H4O+), 149 (C10H9O4+), and 193 (C10H8O4+), which correspond to characteristic molecular fragments of PET polymers [34]. In the negative SIMS spectrum of the untreated sample (Figure 5a), we observe the following major characteristic PET fragments: m/z 76 (C6H4−) and 121 (C7H5O2−). For the plasma-treated sample (the positive SIMS spectrum is shown in Figure 4b), we

Figure 2. XPS surface composition of PET samples treated in H2S plasma for various periods.

Figure 3. Comparison of the high-resolution XPS spectra of the (a) carbon C1s peak; and (b) the sulfur S2p peak at various treatment periods.

Further information was obtained from Figure 3b, which shows the comparison of the high-resolution sulfur peaks. Sulfur peaks for plasma-treated samples were positioned at approximately 163.3 eV, indicating that sulfur was bonded either to another sulfur atom or to carbon. The absence of peaks at higher binding energies close to 170 eV indicated that sulfur was not bonded to oxygen atoms in PET. According to the results shown in Figure 3, sulfur was bonded to carbon atoms in -C-SH or similar groups (C-S-S, C=Si, C-S-S). Unfortunately, the chemical shifts corresponding to different sulfur functional groups are too small to allow for reliable conclusions about the surface chemistry involved. Nevertheless, the maximum sulfur concentration on the surface was so high that it could not be explained solely by surface functionalization with sulfur-containing functional groups. The XPS technique gives the concentration of elements averaged over the investigation depth, which is estimated to several nm.

To obtain more information regarding the chemical changes at the plasma-treated surface, we performed SIMS (secondary ion mass spectrometry) measurements. Positive and negative SIMS spectra for an untreated sample and a plasma-treated sample, where the maximum sulfur content was observed by XPS, are shown in Figures 4 and 5 respectively.

C-O peaks was less pronounced, which was due to the appearance of a new sub-peak corresponding to the C-S bonds at approximately 285.5 eV [33].
can see that all characteristic PET peaks remain in the spectrum. Furthermore, a new peak at \( m/z = 45 \), assigned to CHS\(^{+}\), appeared as a consequence of plasma treatment. Additional information for the plasma-treated sample can be found in the negative SIMS spectrum shown in Figure 5b, where we can see characteristic peaks arising from S\(_4\)\(^−\) (\( m/z \) 128), S\(_3\)\(^−\) (\( m/z \) 96), S\(_2\)\(^−\) (\( m/z \) 64) and S\(^−\) (\( m/z \) 32) molecules, which indicates that the sulfur accumulates on the polymer surface. Other peaks can be found as well: HS\(^−\) (\( m/z \) 33), C\(_2\)HS\(^−\) (\( m/z \) 57) and S\(_2\)C\(_2\)\(^−\) (\( m/z \) 88).

![Figure 4. Positive SIMS spectra for (a) untreated PET; and (b) PET treated for 80 s.](image)

![Figure 5. Negative SIMS spectra for (a) untreated PET; and (b) PET treated for 80 s.](image)

Figures 6 and 7 show the variation of the relative intensities of the main positive and negative SIMS signals with treatment time. The diagrams in Figure 6 show the intensities of the fragments correlated with the PET polymer, whereas Figure 7 shows the fragments containing sulfur atoms. The intensity of the characteristic PET positive signals (Figure 6a) at first decrease with increasing treatment time, reaching a minimum, whereas, at longer treatment times, the original intensities are almost restored. A similar conclusion can be drawn for the variation of the negative signals of the characteristic PET fragments with treatment time (Figure 6b). However, for fragments containing sulfur, we observe the opposite variation with treatment time (Figure 7)—there is a distinctive maximum, similar to the one observed in the case of XPS measurements.

By comparison of the results obtained by XPS and SIMS, we can explain such a high concentration of sulfur on the polymer surface by its accumulation rather than chemical bonding to carbon atoms from the polymer. The surface-reaction mechanism is thus as follows: the initial step in the interaction of plasma with the polymer is chemical binding of sulfur to carbon atoms. This conclusion can be drawn from the behavior of C\(_2\)HS\(^−\), CHS\(^+\) and C\(_2\)S\(_2\)\(^−\) in Figure 7. The surface quickly saturates with sulfur, forming a thin film of chemically bonded sulfur. The necessary treatment time for saturation of the surface with sulfur atoms chemically bonded to carbon can be estimated from Figure 7 as almost 20 s. Once the concentration of chemically bonded sulfur reaches an almost constant value (treatment
time over 20 s), simultaneously with chemical bonding, the sulfur atoms also accumulate on the surface of the treated polymer. The deposition of sulfur is revealed from the appearance of the \( S_2^- \) and \( S_3^- \) peaks in the negative SIMS spectra. The intensity of these spectral features increases strongly for the first 40 s of plasma treatment and then decreases slowly, as revealed from Figure 7. This layer of sulfur, which is not chemically bonded to carbon of the polymer substrate, may be in the form of polysulfides \(-S_x^-\) because sulfur tends to catenate (bind to itself by the formation of chains), but the formation of any type of polysulfide is almost impossible to confirm by our experimental techniques. The appearance of unbonded sulfur has been described in several reports, in which plasma-assisted decomposition of \( \text{H}_2\text{S} \) was investigated to develop a method for destroying this environmentally problematic gas, which is produced in the oil refinement industry [13,17–20]. These studies found direct decomposition into \( \text{H}_2 \) and S, which was deposited on the reactor wall [19]. A similar deposition on the walls of the plasma reactor was observed in our study, as revealed from Figure 7.

![Figure 6](image1)

**Figure 6.** Variation of the relative SIMS intensities of PET fragmented ions: (a) positive; and (b) negative.

![Figure 7](image2)

**Figure 7.** Variation of the relative SIMS intensities of the positive and negative fragments, which are linked with sulfur.

It was unexpected that, at longer treatment time, the quantity of sulfur at the surface was significantly reduced. The deposited sulfur layer must be unstable in our experimental conditions because both the XPS and SIMS results clearly demonstrated slow but continuous decreasing of the S concentration for prolonged treatment periods. The layer formed upon the first minute of plasma treatment must have been removed from the surface upon prolonged treatment. The reason for this
may be degradation as a result of the higher surface temperature because polysulfides (R-Sx), as well as
other sulfur compounds, such as polysulfanes (H2Sx), are not thermally stable [35,36]. Furthermore, the
process of degradation may be accelerated by UV radiation from plasma [35]. To estimate the possible
thermal effect, we measured the sample temperature during plasma treatment. The result is shown
in Figure 8. This figure shows that the bulk temperature of the sample reached more than 90 °C
at 200 s of treatment, which is enough to cause degradation of thermally unstable compounds and
desorption to the gas phase. Our results presented in Figures 2, 6 and 7 indicate that even a somewhat
lower temperature is high enough to facilitate desorption of sulfur compounds under low-pressure
conditions. Thiry et al. observed a similar effect: at treatment conditions where the surface temperature
was low (30 °C–35 °C), the sulfur concentration was much higher than at conditions where the surface
temperature was moderate (60 °C–90 °C) [25]. He explained this effect as being caused by trapped
H2S molecules in the surface film, which were released at high temperatures.

![Figure 8. Sample temperature during plasma treatment.](image)

2.3. Surface Morphology, Etching and Aging

Any deposition of a foreign material on the substrate may or may not be reflected in modified
surface morphology, depending on the growth mechanisms. To investigate the morphological changes,
we performed AFM (atomic force microscopy) analyses. Figure 9 shows an AFM image of an
untreated PET sample, whereas Figures 10 and 11 show AFM images of selected samples recorded
on 5 × 5 μm² and 2 × 2 μm² areas, respectively. The corresponding surface roughness is shown in
Table 1. The roughness evolution deduced from the AFM images roughly coincides with the behavior
of the sulfur concentration determined by XPS and SIMS. Initially (up to the treatment time of 10 s),
the surface is quite smooth without any special features (Figures 10a and 11a). At 20 s of treatment, the
formation of the first particles at the surface is initiated (Figures 10b and 11b). At 40 s of treatment, the
surface is fully covered with particles, and their lateral size and height have increased (Figures 10c
and 11c). These particles can be attributed to large sulfur clusters on the surface, but as shown later in
the text, this conclusion does not agree with other observations. After 80 s of treatment, the surface
roughness has decreased; however, the surface is still fully covered with circular features (Figures 10d
and 11d). At longer treatment periods, when the sulfur concentration measured by XPS and SIMS has
decreased, the lateral size of the particles has also decreased (Figures 10e,f and 11e,f).
Figure 9. AFM images of untreated PET sample: (a) 5 × 5 μm²; and (b) 2 × 2 μm².

Figure 10. AFM images (5 × 5 μm²) of samples treated for various periods.
Figure 10. AFM images (5 × 5 μm²) of samples treated for various periods.

Figure 11. AFM images (2 × 2 μm²) of samples treated for various periods.

Table 1. Surface roughness of the samples treated for different periods as measured by AFM over an area of 2 × 2 μm² and 5 × 5 μm².

| Treatment Time (s) | Roughness (nm) | Measured on 5 × 5 μm² Area | Measured on 2 × 2 μm² Area |
|-------------------|----------------|----------------------------|---------------------------|
| 0                 | 1.2            | 1.2                        |                           |
| 10                | 1.2            | 1.2                        |                           |
| 20                | 2.4            | 1.6                        |                           |
| 40                | 1.7            | 1.4                        |                           |
| 80                | 2.3            | 2.1                        |                           |
| 160               | 3.3            | 3.1                        |                           |
| 320               | 3.8            | 3.2                        |                           |

Even though the AFM results show a good correlation with the XPS, and especially the SIMS, the clusters observed in the AFM images are not associated with the sulfur deposit. The sample with the maximum sulfur concentration was aged in air for six months and then analyzed again (Table 2). The XPS results of the aged sample showed a significant decrease in the sulfur concentration. After half a year of aging, the concentration was only approximately 6 atomic % (initially, it was 40 atomic %). The AFM images showed that the surface morphology remained unchanged, and clusters are still observed (Figure 12).
was approximately 1.5 nm/s. Although the etching rate was small, it caused changes to the surface.

Therefore, we can only conclude that the moderate hydrophobic nature of this polymer is preserved upon treatment with H$_2$S plasma.

Because the surface roughness and morphology in combination with the surface functionalization may have a significant effect on the surface wettability, we also measured the water contact angles on the plasma-treated samples. The contact angles are shown in Figure 13. For untreated PET, the angle was approximately 73°, whereas for plasma-treated samples, the angle was between 57° and 72°. Comparing the contact angles with the XPS, SIMS and AFM results, it is difficult to find any correlation. Therefore, we can only conclude that the moderate hydrophobic nature of this polymer is preserved upon treatment with H$_2$S plasma.

Therefore, the observed surface morphology cannot be explained by sulfur deposition or by intensive etching of the polymer because no oxygen lines were observed in the OES spectra (Figure 1b). Furthermore, the etching rates of polymer exposed to hydrogen radicals and ultraviolet (UV) or vacuum ultraviolet (VUV) radiation are small [37], but they are unknown for SH radicals. Therefore, we performed weight loss measurements to estimate the etching rate. Selected samples were first weighed, treated in plasma and rinsed with toluene to remove sulfur deposits, and then weighed again. Knowing the surface area, sample density and treatment time, we calculated the etching rate, which is the mass of the sample divided by the surface area, sample density and treatment time.

\[
\text{Etching Rate} = \frac{\Delta m}{A \cdot \rho \cdot t}
\]

The etching rate is given in units of nm/s. Knowing the surface area, sample density and treatment time, we calculated the etching rate, which is the mass of the sample divided by the surface area and treatment time.

| Aging Period | Carbon (atomic %) | Oxygen (atomic %) | Sulfur (atomic %) |
|--------------|------------------|------------------|------------------|
| 0 days       | 46.9             | 13.1             | 40.0             |
| 1 day        | 65.0             | 18.5             | 16.4             |
| 2 days       | 67.2             | 19.1             | 13.7             |
| 6 months     | 73.8             | 19.8             | 6.4              |

The etching rate of polymer exposed to hydrogen radicals and ultraviolet (UV) or vacuum ultraviolet (VUV) radiation are small [37], but they are unknown for SH radicals. Therefore, we performed weight loss measurements to estimate the etching rate. Selected samples were first weighed, treated in plasma and rinsed with toluene to remove sulfur deposits, and then weighed again. Knowing the surface area, sample density and treatment time, we calculated the etching rate, which was approximately 1.5 nm/s. Although the etching rate was small, it caused changes to the surface morphology. The etching was highly non-uniform due to sulfur deposition. Deposition of sulfur caused certain areas of the surface to be covered with sulfur and were therefore protected, whereas the other parts of the surface were exposed to etching. At longer treatment periods, when sulfur was removed, the surface was etched more uniformly, and the surface morphology was changed again.

![AFM images](image_url)

**Figure 12.** AFM images (a) 2 × 2 μm$^2$; and (b) 5 × 5 μm$^2$ of the PET sample treated for 80 s after six months of aging.

**Table 2.** Surface composition of the sample with the maximum sulfur concentration (treated for 80 s) after aging for different periods (in atomic %).

2.4. Surface Wettability

2.5. Surface Roughness and Morphology

2.6. Surface Functionalization

2.7. Surface Aging
were treated in the discharge tube presented schematically in Figure 14. The tube was made from Pyrex glass and was 80 cm long and 4 cm in diameter. The discharge tube was pumped with a rotary pump operating at a nominal pumping speed of 80 m³·h⁻¹. Hydrogen sulfide (H₂S) gas was leaked into the experimental system on the opposite side, as shown in Figure 14. The pressure was set to 30 Pa, so the gas flow rate was 400 sccm. A coil of 6 turns was mounted in the center of the Pyrex tube, as shown in Figure 14. Plasma was created by an RF generator coupled to the coil via a matching network. The matching network consisted of two vacuum capacitors: one connected in series and another in parallel. The generator operated at the standard frequency of 13.56 MHz, and its nominal power was set to 150 W. At these discharge conditions, the plasma was ignited in the low-density mode (i.e., E-mode) [38]. Samples were treated in H₂S plasma for various periods of 10, 20, 40, 80, 160, 320 and 640 s. After treatment, the samples were characterized using atomic force microscopy (AFM), X-ray photoelectron spectroscopy (XPS), time-of-flight-secondary ion mass spectrometry (ToF-SIMS), and contact angle measurements. Some samples were stored and re-characterized after prolonged time to monitor ageing effects.

3. Materials and Methods

3.1. Plasma Treatment

Biaxially oriented polymer PET from Goodfellow Ltd. (Huntingdon, UK) was used. The polymer film with a thickness of 0.125 mm was cut to small square pieces with a size of 1 × 1 cm². The samples were treated in the discharge tube presented schematically in Figure 14. The pressure was set to 30 Pa, so the gas flow rate was 400 sccm. A coil of 6 turns was mounted in the center of the Pyrex tube, as shown in Figure 14. Plasma was created by an RF generator coupled to the coil via a matching network. The matching network consisted of two vacuum capacitors: one connected in series and another in parallel. The generator operated at the standard frequency of 13.56 MHz, and its nominal power was set to 150 W. At these discharge conditions, the plasma was ignited in the low-density mode (i.e., E-mode) [38]. Samples were treated in H₂S plasma for various periods of 10, 20, 40, 80, 160, 320 and 640 s. After treatment, the samples were characterized using atomic force microscopy (AFM), X-ray photoelectron spectroscopy (XPS), time-of-flight-secondary ion mass spectrometry (ToF-SIMS), and contact angle measurements. Some samples were stored and re-characterized after prolonged time to monitor ageing effects.

Figure 13. Water contact angle variation vs. plasma treatment period.

Figure 14. Schematic diagram of the experimental system: 1—H₂S gas source; 2—leak valve; 3—discharge tube; 4—coil; 5—sample; 6—plasma; 7—matching network; 8—RF generator; 9—vacuum gauge; 10—catalyzer; 11—vacuum pump; 12—flange; 13—OES spectrometer.
3.2. Plasma Characterization

Plasma was characterized using optical emission spectroscopy (OES). OES measurements were performed in a quartz tube with a 16-bit Avantes AvaSpec 3648 fiber optic spectrometer (Avantes Ltd., Leatherhead, Surrey, UK). The nominal spectral resolution was 0.8 nm, and spectra were recorded in the range from 200 to 1100 nm. A combined deuterium tungsten reference light source was used to determine the spectral response of the spectrometer. The measured OES spectra were calibrated with this spectral response. Because the light emitted by H\textsubscript{2}S plasma at our discharge parameters was weak, the integration time used to record the OES spectra was 10 s.

3.3. Temperature Measurements

Temperature of the sample during plasma treatment was measured by a chromel-alumel thermocouple. The thermocouple tip was placed between two pieces of PET film, which were pressed together to make good contact with the thermocouple.

3.4. Weight-Loss Measurements

The etching of the polymer during plasma treatment was monitored by measuring the weight loss of the selected polymer samples. The samples were weighed just before mounting into the plasma reactor and again just after plasma treatment. A Radwag XA 110 (Radwag, Radom, Poland) professional microbalance was used. The accuracy of the measurements was, according to the producer, 0.01 mg. Samples were washed in toluene and dried before weighing to remove any impurities, deposits or degradation products from the surface.

3.5. AFM Measurements

An AFM (Solver PRO, NT-MDT, Moscow, Russia) was used to characterize the topology of the samples. All measurements were performed in tapping mode using ATEC-NC-20 tips (Nano And More GmbH, Wetzlar, Germany) with a resonance frequency of 210–490 kHz and force constant of 12–110 Nm\textsuperscript{–1}. The surface roughness was calculated from the AFM images taken over an area of 2 × 2 μm\textsuperscript{2} and 5 × 5 μm\textsuperscript{2} using the program Spip 5.1.3 (Image Metrology A/S, Hørsholm, Denmark). The surface roughness was expressed in terms of the average roughness (Ra).

3.6. XPS Measurements

XPS characterization of the polymer samples was performed to determine their chemical composition after plasma treatment using an XPS (TFA XPS Physical Electronics, Münich, Germany). The samples were excited with monochromatic Al K\textsubscript{α1,2} radiation at 1486.6 eV over an area with a diameter of 400 μm. Photoelectrons were detected with a hemispherical analyzer positioned at an angle of 45° with respect to the normal of the sample surface. XPS survey spectra were measured at a pass-energy of 187 eV using an energy step of 0.4 eV, and high-resolution spectra were measured at a pass-energy of 23.5 eV using an energy step of 0.1 eV. An additional electron gun was used for surface neutralization during the XPS measurements. All spectra were referenced to the main C1s peak of the carbon atoms, which was assigned a value of 284.8 eV. The measured spectra were analyzed using MultiPak v8.1c software (Ulvac-Phi Inc., Kanagawa, Japan, 2006) from Physical Electronics, which was supplied with the spectrometer.

3.7. ToF-SIMS Measurements

ToF-SIMS analyses were performed using a ToF-SIMS 5 instrument (ION-TOF, Münster, Germany) equipped with a bismuth liquid metal ion gun with a kinetic energy of 30 keV. The analyses were performed in an ultra-high vacuum of approximately 10\textsuperscript{–7} Pa. The SIMS spectra were measured by scanning a Bi\textsubscript{3+} cluster ion beam with a diameter of 1 μm over a 100 × 100 μm\textsuperscript{2} analysis area. The positive secondary ion mass spectra were calibrated using CH\textsubscript{2}+, CH\textsubscript{3}+, and C\textsubscript{2}H\textsubscript{3}+, and the
negative secondary ion mass spectra were calibrated using C−, C2−, and C3−. An electron gun was used for charge compensation on the sample surfaces during the analysis.

3.8. Contact Angle Measurements

The surface wettability was measured immediately after plasma treatment by determining the water contact angle (WCA) with a demineralized 3 µL water droplet. Contact angles were measured by See System (Advex Instruments, Brno, Czech Republic). For each sample, five measurements were taken to minimize the statistical error. The contact angles were determined by the software supplied by the producer.

4. Conclusions

PET polymer was modified with radicals created in the gaseous plasma of H2S. Plasma was sustained by an electrode-less RF discharge created in the E-mode, which enabled ionization and dissociation of H2S molecules to form H and HS radicals. Combination of different surface-sensitive techniques allowed for insight into the surface chemistry as well as morphological changes upon plasma treatment. A couple of competitive processes were identified: (i) chemical bonding of sulfur atoms to carbon on the substrate surface; and (ii) deposition of a thin sulfur film. The first process was irreversible because the concentration of the sulfur bonded to carbon atoms remained unchanged even after prolonged aging. The thin sulfur film, however, was unstable. The concentration of the chemically unbonded sulfur decreased with prolonged plasma treatment time and almost vanished after storing for several months.

The concentration of total sulfur on the polymer surface increased with treatment time for the first minute, which was explained by the accumulation of sulfur on the surface. This layer is thermally unstable and degrades spontaneously, even at room temperature. Slow decrease of the sulfur concentration for prolonged plasma treatment times (over a minute) was observed and was attributed to the increasing polymer temperature. After several minutes of plasma treatment, the sulfur concentration significantly decreased and reached a few atomic % after approximately 10 min. The results indicate the existence of an optimal treatment time to obtain a large concentration of sulfur. At our experimental conditions, the optimal time is approximately one minute, but this value is different under other conditions due to the different heating of the polymer samples. Large changes of the surface morphology (formation of spherical features) were observed, but were not directly related to sulfur deposition.

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References

1. Chiper, A.; Borcia, G. Argon versus helium dielectric barrier discharge for surface modification of polypropylene and poly(methyl methacrylate) films. Plasma Chem. Plasma Process. 2013, 33, 553–568. [CrossRef]

2. Borcia, C.; Punga, I.L.; Borcia, G. Surface properties and hydrophobic recovery of polymers treated by atmospheric-pressure plasma. Appl. Surf. Sci. 2014, 317, 103–110. [CrossRef]

3. Jacobs, T.; Carbone, E.; Morent, R.; de Geyter, N.; Reniers, F.; Leys, C. Surface modification of polymer films with a remote atmospheric pressure d.c. glow discharge: Influence of substrate location. Surf. Interface Anal. 2010, 42, 1316–1320. [CrossRef]

4. Mafra, M.; Belmonte, T.; Foncin-Epaillard, F.; Maliska, A.; Cvelbar, U. Treatment of hexatriacontane by Ar-O2 remote plasma: Formation of the active species. Plasma Process. Polym. 2009, 6, S198–S203. [CrossRef]
5. Belmonte, T.; Pintassilgo, C.D.; Czerwiec, T.; Henrion, G.; Hody, V.; Thiebaut, J.M.; Loureiro, J. Oxygen plasma surface interaction in treatments of polyolefins. Surf. Coat. Technol. 2005, 200, 26–30. [CrossRef]

6. Barni, R.; Zanini, S.; Beretta, D.; Riccardi, C. Experimental study of hydrophobic/hydrophilic transition in SF6 plasma interaction with polymer surfaces. Eur. Phys. J. Appl. Phys. 2007, 38, 263–268. [CrossRef]

7. Bastos, D.C.; Santos, A.E.F.; da Fonseca, M.D.; Simao, R.A. Inducing surface hydrophobization on cornstarch film by SF6 and HMDSO plasma treatment. Carbohydr. Polym. 2013, 91, 675–681. [CrossRef] [PubMed]

8. Fatyeyeva, K.; Poncin-Epaillard, F. Sulfur dioxide plasma treatment of the clay (laponite) particles. Plasma Chem. Plasma Process. 2011, 31, 449–464. [CrossRef]

9. Holländer, A.; Kröpke, S. Polymer surface treatment with SO2-containing plasmas. Plasma Process. Polym. 2010, 7, 390–402. [CrossRef]

10. Siow, K.S.; Britcher, L.; Kumar, S.; Griesser, H.J. Sulfonated surfaces by sulfur dioxide plasma discharge treatment of polymer films. Plasma Process. Polym. 2009, 6, 583–592. [CrossRef]

11. Zhao, L.; Wang, Y.; Jin, L.; Qin, M.; Li, X.; Wang, A.; Song, C.; Hu, Y. Decomposition of hydrogen sulfide in non-thermal plasma aided by supported CdS and ZnS semiconductors. Green Chem. 2013, 15, 1509–1513. [CrossRef]

12. Nunnally, T.; Gutsol, K.; Rabinovich, A.; Fridman, A.; Starikovsky, A.; Gutsol, A.; Potter, R.W. Dissociation of H2S in non-equilibrium gliding arc “tornado” discharge. Int. J. Hydrogen Energ. 2009, 34, 7618–7625. [CrossRef]

13. Reddy, E.L.; Biju, V.M.; Subrahmanyam, C. Production of hydrogen from hydrogen sulfide assisted by dielectric barrier discharge. Int. J. Hydrogen Energ. 2012, 37, 2204–2209. [CrossRef]

14. Reddy, E.L.; Karuppiiah, J.; Subrahmanyam, C. Kinetics of hydrogen sulfide decomposition in a DBD plasma reactor operated at high temperature. J. Energ. Chem. 2013, 22, 382–386. [CrossRef]

15. John, S.; Hamann, J.C.; Muknahallipatna, S.S.; Legowski, S.; Ackerman, J.F.; Argyle, M.D. Energy efficiency of hydrogen sulfide decomposition in a pulsed corona discharge reactor. Chem. Eng. Sci. 2009, 64, 4826–4834. [CrossRef]

16. Sassi, M.; Amira, N. Chemical reactor network modeling of a microwave plasma thermal decomposition of H2S into hydrogen and sulfur. Int. J. Hydrogen Energ. 2012, 37, 10010–10019. [CrossRef]

17. Huang, L.; Xia, L.; Dong, W.; Hou, H. Energy efficiency in hydrogen sulfide removal by non-thermal plasma photolysis technique at atmospheric pressure. Chem. Eng. J. 2013, 228, 1066–1073. [CrossRef]

18. Reddy, E.L.; Biju, V.M.; Subrahmanyam, C. Production of hydrogen and sulfur from hydrogen sulfide assisted by nonthermal plasma. Appl. Energ. 2012, 95, 87–92. [CrossRef]

19. Gutsol, K.; Nunnally, T.; Rabinovich, A.; Fridman, A.; Starikovsky, A.; Gutsol, A.; Kemoun, A. Plasma assisted dissociation of hydrogen sulfide. Int. J. Hydrogen Energ. 2012, 37, 1335–1347. [CrossRef]

20. Zhao, G.-B.; John, S.; Zhang, J.-J.; Hamann, J.C.; Muknahallipatna, S.S.; Legowski, S.; Ackerman, J.F.; Argyle, M.D. Production of hydrogen and sulfur from hydrogen sulfide in a nonthermal-plasma pulsed corona discharge reactor. Chem. Eng. Sci. 2007, 62, 2216–2227. [CrossRef]

21. Reddy, E.L.; Karuppiiah, J.; Biju, V.M.; Subrahmanyam, C. Catalytic packed bed non-thermal plasma reactor for the extraction of hydrogen from hydrogen sulfide. Int. J. Catalytic. Res. 2013, 37, 1280–1286. [CrossRef]

22. Thiry, D.; Francq, R.; Cossement, D.; Guillaume, M.; Cornil, J.; Snyders, R. A detailed description of the chemistry of thiol supporting plasma polymer films. Plasma Process. Polym. 2014, 11, 606–615. [CrossRef]

23. Thiry, D.; Britton, N.; Konstantinidis, S.; Dauchot, J.-P.; Guillaume, M.; Cornil, J.; Snyders, R. Experimental and theoretical study of the effect of the inductive-to-capacitive transition in propanethiol plasma polymer chemistry. J. Phys. Chem. C 2013, 117, 9843–9851. [CrossRef]

24. Thiry, D.; Britton, N.; Konstantinidis, S.; Dauchot, J.-P.; Denis, L.; Snyders, R. Altering the sulfur content in the propanethiol plasma polymers using the capacitive-to-inductive mode transition in inductively coupled plasma discharge. Appl. Phys. Lett. 2012, 100. [CrossRef]

25. Thiry, D.; Aparicio, F.J.; Britton, N.; Snyders, R. Concomitant effects of the substrate temperature and the plasma chemistry on the chemical properties of propanethiol plasma polymer prepared by ICP discharges. Surf. Coat. Technol. 2014, 241, 2–7. [CrossRef]

26. Akhavan, B.; Jarvis, K.; Majewski, P. Plasma polymerization of sulfur-rich and water-stable coatings on silica particles. Surf. Coat. Technol. 2015, 264, 72–79. [CrossRef]

27. Akhavan, B.; Jarvis, K.; Majewski, P. Development of oxidized sulfur polymer films through a combination of plasma polymerization and oxidative plasma treatment. Langmuir 2014, 30, 1444–1454. [CrossRef] [PubMed]
28. Kiesow, A.; Heilmann, A. Deposition and properties of plasma polymer films made from thiophenes. *Thin Solid Films* **1999**, *343–344*, 338–341. [CrossRef]
29. Bhat, N.V.; Wavhal, D.S. Preparation and characterization of plasma-polymerized thiophene films. *J. Appl. Polym. Sci.* **1998**, *70*, 203–209. [CrossRef]
30. Yu, Y.-J.; Kim, J.-G.; Boo, J.-H. Plasma-polymerized thiophene films for corrosion protection in microelectronic devices. *J. Mater. Sci. Lett.* **2002**, *21*, 951–953. [CrossRef]
31. Kregar, Z.; Biscan, M.; Milosevic, S.; Mozetic, M.; Vesel, A. Interaction of argon, hydrogen and oxygen plasma early afterglow with polyvinyl chloride (PVC) materials. *Plasma Process. Polym.* **2012**, *9*, 1020–1027. [CrossRef]
32. Pearse, R.W.B.; Gaydon, A.G. *The Identification of Molecular Spectra*; Chapman and Hall: London, UK, 1984.
33. Thiry, D.; Francq, R.; Cossement, D.; Guerin, D.; Vuillaume, D.; Snyders, R. Establishment of a derivatization method to quantify thiol function in sulfur-containing plasma polymer films. *Langmuir* **2013**, *29*, 13183–13189. [CrossRef] [PubMed]
34. Lang, F.R.; Pitton, Y.; Mathieu, H.J.; Landolt, D.; Moser, E.M. Surface analysis of polyethyleneterephthalate by ESCA and TOF-SIMS. *Fresen. J. Anal. Chem.* **1997**, *358*, 251–254. [CrossRef]
35. Lee, T.C.P. *Properties and Applications of Elastomeric Polysulfides*; Rapra Technology: Shrewsbury, UK, 1999.
36. Kozhevnikov, V.F.; Payne, W.B.; Olson, J.K.; McDonald, C.L.; Inglefield, C.E. Physical properties of sulfur near the polymerization transition. *J. Chem. Phys.* **2004**, *121*, 7379–7386. [CrossRef] [PubMed]
37. Cho, K.; Setsuhara, Y.; Takenaka, K.; Shiratani, M.; Sekine, M.; Hori, M. Effects of photoirradiation in UV and VUV regions during plasma exposure to polymers. *Thin Solid Films* **2011**, *519*, 6810–6814. [CrossRef]
38. Zaplotnik, R.; Vesel, A.; Mozetic, M. Transition from E to H mode in inductively coupled oxygen plasma: Hysteresis and the behaviour of oxygen atom density. *Eur. Phys. Lett.* **2011**, *95*. [CrossRef]

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