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# Kinetics, Chemistry and Morphology of Syngas Photo-Initiated Chemical Vapor Deposition

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**KEYWORDS:** photo-initiated chemical vapor deposition (PICVD), syngas, photo-polymerization, deposition rate, coating.

**ABSTRACT:** Syngas is the product of gasification processes and is used for the production of petrochemicals. Little attention has been given to its use for the production of oligomeric thin films at ambient conditions. Herein, the nature of photo-initiated chemical vapor deposition of films made from syngas using high wavelength ultraviolet light is discussed, including an exploration into the oligomeric films' structure, synthesis mechanism and growth kinetics. Specifically, X-ray photoelectron spectroscopy and time-of-flight secondary ion mass spectrometry analyses provide insight into the chemical structure, illustrating the effect of photo-generated radicals in the formation of aliphatic, anhydride and cyclic structures. The films are covalently bonded to the substrate and chemically uniform. Electron and atomic force microscopy identify an island-like morphology for the deposit. These insights into to the mechanism and structure are linked to processing parameters through a study on the effect of residence time and treatment duration on deposition rate, determined through profilometry.

## 1. INTRODUCTION

Surface modification techniques have gained a lot of interest due to their ability to tailor surface properties without altering the bulk of the material. Hybrid organic-inorganic coatings are of interest in various fields such as optics, environmental sciences, energy, mechanics, catalysis and sensors, because they can tailor surface wettability as well as the chemical specifications of materials.<sup>1-5</sup> Several techniques such as chemical vapor deposition (including plasma enhanced and thermally activated), atomic layer deposition and sol-gel techniques have been used for thin film synthesis.<sup>2,6</sup> Among these techniques, photo-initiated chemical vapor deposition (PICVD) has considerable superiority given its simple reactor design, ease of use and control, scalability, versatility, affordability and low environmental footprint. The solvent-free deposition nature, as well as the ability to work without any high vacuum or high temperatures make this process an attractive alternative to other gas phase surface modification approaches.<sup>7-10</sup>

Most studies in this field use vacuum ultraviolet (VUV) lamps emitting at wavelengths shorter than 200 nm as initiation sources.<sup>11-19</sup> However, despite the ability of light at these wavelengths to cleave most chemical bonds due to their high energy photons (5-15 eV),<sup>20</sup> light transmission becomes prohibitive (i.e. specialized reactor windows, such as MgF<sub>2</sub> and LiF, or designs are compulsory.<sup>2,21</sup>) Given these limitations, UVC light (200-280 nm),<sup>22</sup> which can be transmitted through affordable quartz windows and generated via commercially ubiquitous germicidal lamps, merits investigation. Its application to PICVD requires the selection of appropriate precursors containing chemical bonds reactive at the wavelength of the UV source.

In general, precursor selection depends on (1) the modification technique, (2) the reactor type and (3) the required chemical and physical bulk properties corresponding to the application. In PICVD, precursors should either absorb UV or, in the case of precursor mixtures, react with the radical/excited species produced upon UV absorption of other monomers. Thus, absorption cross section, absorption

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2 coefficient and ionization potential drive precursor selection.<sup>2,23</sup> Window composition is one of the  
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4 main features of reactor design. In PICVD processing with UVC lamps, there is no need for special  
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6 materials and quartz windows are applicable. Many chemical compounds have been investigated for  
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8 PICVD process such as methyl methacrylate, polyethyleneglycol diacrylate, trimethylolpropane  
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10 tris(mercaptopropionate), ethylene, styrene, butadiene, etc.<sup>1,24-29</sup> However, these need to be excited  
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12 either in the presence of photoinitiators or photosensitizers, or with high energy UV lamps; this makes  
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14 the process very expensive.  
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17  
18 To overcome these issues, we looked to Fischer-Tropsh synthesis. In this process, catalysts are used  
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20 to combine the binary mixture of CO and H<sub>2</sub>, known as synthesis gas or syngas, into hydrocarbons.  
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22 Catalysts deactivate during this process, due to the formation of lightly polymerized carbon species.<sup>30-34</sup>  
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24 This surface fouling can be reimagined as surface engineering - this triggered us to consider syngas as  
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26 an alternative reactive precursor in PICVD. Very few researchers have attempted to study the photo-  
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28 polymerization of syngas components,<sup>9,35</sup> and most have focused on carbon monoxide photochemical  
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30 reactions in the early Earth's atmosphere or with low wavelength VUV lamps.<sup>35-37</sup> Previously, our  
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32 group demonstrated that syngas PICVD could be used to deposit thin films with various functionalities  
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34 on copper substrates<sup>9</sup> and nanomaterials,<sup>38-41</sup> but the deposition mechanism and kinetics remain elusive.  
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36 Herein, we report on the photochemical reaction of CO and H<sub>2</sub> molecules to bring new insights into the  
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38 mechanism and structure of the deposited film, and relate these features to key processing parameters.  
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40 We present in terms of (i) chemical groups, (ii) density of generated products, (iii) morphology and  
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42 topography of the deposited thin film, as well as (iv) film growth with respect to the irradiation time  
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44 (and conversely, reagent residence time). These findings allow us to derive a reaction mechanism for  
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46 the deposition process and a scheme for the chemical structure of the film, as well as kinetics to explain  
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48 the deposition rate of the oligomeric film.  
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## 2. EXPERIMENTAL

### 2.1. Sample preparation:

Substrates were type N, orientation 110 silicon wafers with silicon oxide on the top surface (University Wafer), cut into 1 cm × 1 cm coupons. Before experiments, the coupons were cleaned by soaking in isopropanol (Fisher, 99.9% purity) in an ultrasonic bath for 10 minutes, and dried with compressed air.

### 2.2. Experimental methodology:

Before mounting onto a sample holder, we masked one side of the silicon coupon with Kapton tape to serve as a reference (Kapton® Polyimide for soldering and splicing; 7648A73 McMaster-Carr). The sample was inserted into the PICVD reactor (**Figure 1S**), which consists of a 225 cm<sup>3</sup> plug flow quartz tube reactor with standard 24/40 taper joints (Montreal Glassblowing Inc.), surrounded by 28 UVC mercury (germicidal) lamps mounted in a UV-cabinet (custom made by Daavlin Co.). The UVC lamps have a main emission peak at a wavelength of 253.7 nm (measured via Ocean Optics Spectrometer/USB4000-XR1-ES), with a light intensity of 0.012 W/cm<sup>2</sup> at the reactor distance of 33 cm (measured via ILT1700 Research Radiometer coupled with a SED240/QNDS2/ W254 nm sensor, International Light Technologies).

Before experiments, the reactor was purged for 5 minutes with 0.4 L/min of argon (Air Liquide, 99.99%). The prescribed amount of syngas (CO and H<sub>2</sub>, Air Liquide, 99.99%) was then fed to the reactor at an inlet pressure of 40 psi. The reactor was maintained at atmospheric pressure. The UV cabinet was then turned on and treatment duration time was set according to the experimental plan (**Table 1S**). Experiments were initiated at room temperature (~22 °C). However, over the course of treatment (2h), the temperature increased to 48 °C because of the heat given off by the UV lamps. Each experiment was repeated 3 times to ensure reproducibility.

### 2.3. Characterization techniques:

Elemental analysis of deposited films was done via a VG ESCALAB 3 MKII X-ray photoelectron spectroscope (XPS) equipped with a non-monochromatic Mg-K $\alpha$  radiation source, operated at 300 W (15 kV, 20 mA). Survey scans were conducted at an X-ray incident angle of 15° with a penetration depths of ~10 nm. The pass energy was 100 eV for survey scans and 20 eV for high-resolution scans in 1.00 eV and 0.05 eV increments, respectively. Pressure during analysis was kept under  $5 \times 10^{-9}$  Torr and the analyzed surface was 2 mm  $\times$  3 mm. Samples were stored under vacuum overnight prior to analysis. The XPS spectra were acquired normal to the sample surface and analyzed via the Avantage XPS software package. The elemental distribution of the samples was determined based on the peak area comparison (C1s, O1s, etc.), normalized to their corresponding sensitivity factors, after removal of the scattered electrons background. In the case of high-resolution spectra, binding energies were referenced to the C1s peak at 285.0 eV to adjust for possible charging effects, and the Shirley method was applied for background noise subtraction. According to the data trend for each distribution of binding energy, the baseline was manually placed. Each curve is represented by its maximum BE in this discussion. Species elemental distribution is obtained via Gaussian/Lorentzian curve fitting on the original curve. The number of sub-curves and their corresponding species were obtained with full width at half maximum (FWHM) = 1.6, 1.8, 2.2 and 2.7 eV for C, O, Si and Fe respectively. These are the most frequent FWHM that we observed with this machine for each elements and normally have been used for all samples.

Static time-of-flight secondary ion mass spectrometry (TOF-SIMS) analysis was carried out with a ION-TOF SIMS IV (from ION-TOF GmbH, Germany) equipped with a 25 keV Bi<sub>3</sub> liquid metal ion source as the primary ion beam source, in high current bunched mode. A bunching system gives pulse durations of 21.9 ns with a mass resolution  $M/\Delta M$  better than 8,000 around  $m/z$  100 in positive and negative SIMS modes. Depth profiling (10 Å maximum) was performed in non-interlaced mode, where

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2 the analysis and sputtering occurred with a  $50\ \mu\text{m} \times 50\ \mu\text{m}$  analysis area confined within a  $500\ \mu\text{m} \times$   
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4  $500\ \mu\text{m}$  sputter area, keeping the total dose below  $5 \times 10^{12}$  ions  $\text{cm}^{-2}$  (so called static conditions).  
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6 Imaging of the samples was also performed in high mass resolution mode over an area of  $500\ \mu\text{m} \times$   
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8  $500\ \mu\text{m}$  with an imaging pixel density of  $128\ \text{pixels} \times 128\ \text{pixels}$ . Three measurements at adjacent spots  
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10 were performed on the samples and positive and negative ion spectra were compared to confirm  
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12 uniformity. Ion spectra were internally calibrated using  $\text{H}^+$ ,  $\text{H}_2^+$ ,  $\text{CH}_3^+$ ,  $\text{C}_2\text{H}_2^+$ ,  $\text{C}_3\text{H}_5^+$  and  $\text{H}^-$ ,  $\text{C}^-$ ,  $\text{CH}^-$ ,  
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14  $\text{C}_2\text{H}^-$ , and  $\text{C}_4\text{H}^-$  peaks, respectively.  
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18 Surface morphology of the deposited film was examined with a Hitachi ultrahigh-resolution cold  
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20 field emission scanning electron microscope (Hitachi SU8230), equipped with an X-ray energy-  
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22 dispersive spectroscope (EDS) (Bruker® Quad detector-Bruker® 4SD Argus system) operated at an  
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24 accelerating voltage of 1-2.5 kV. Samples were mounted on metal stubs using conductive double-sided  
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26 carbon tape.  
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30 Topography was investigated via a Multi-Mode 3 Atomic Force Microscope (Bruker AS-130)  
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32 (AFM) operated in tapping mode. Microfabricated V-shaped silicon cantilevers (Applied Nano Inc.)  
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34 with a theoretical spring constant of 37 N/m, a resonance frequency of  $\sim 300$  kHz and a silicon conical  
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36 tip (radius = 6 nm) were used. Squares of  $5\ \mu\text{m}$  in size were scanned in air under constant applied force  
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38 conditions at a 0.8-1 Hz scan rate. Nanoscope analysis software was used for image processing and  
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40 data analysis; images with  $512 \times 512$  pixel resolution were obtained.  
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44 The film thicknesses were measured using stylus profilometry with a Veeco Dektak 150 surface  
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46 profiler. In order to confirm the profilometry measurements, a few samples were measured by an ex  
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48 situ M44 variable angle spectroscopic ellipsometer at angles of  $65^\circ$ ,  $70^\circ$ ,  $75^\circ$  in the wavelength range  
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50 of 200-1000 nm. The applied optical model consisted of three components: the silicon substrate, the  
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52 native  $\text{SiO}_2$  layer and the film bulk layer. The Cauchy function with Urbach tail was incorporated for  
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bulk components modulation. The model also took into account possible thickness inhomogeneity within the sampled area.

### 3. RESULTS AND DISCUSSION

#### 3.1. Chemical characterization of deposited films

High resolution X-ray photoelectron spectroscopy (XPS) and time-of-flight secondary ion mass spectrometry (TOF-SIMS) analyses provided insight into the chemical composition of the deposited films. In order to gain information on the chemical bonding states and structure of the film, the XPS peaks for bare and coated silicon samples were deconvoluted (**Figure 1**). A summary of the high-resolution spectral deconvolutions along with their possible structure assignment and At % is presented in **Table 1**.

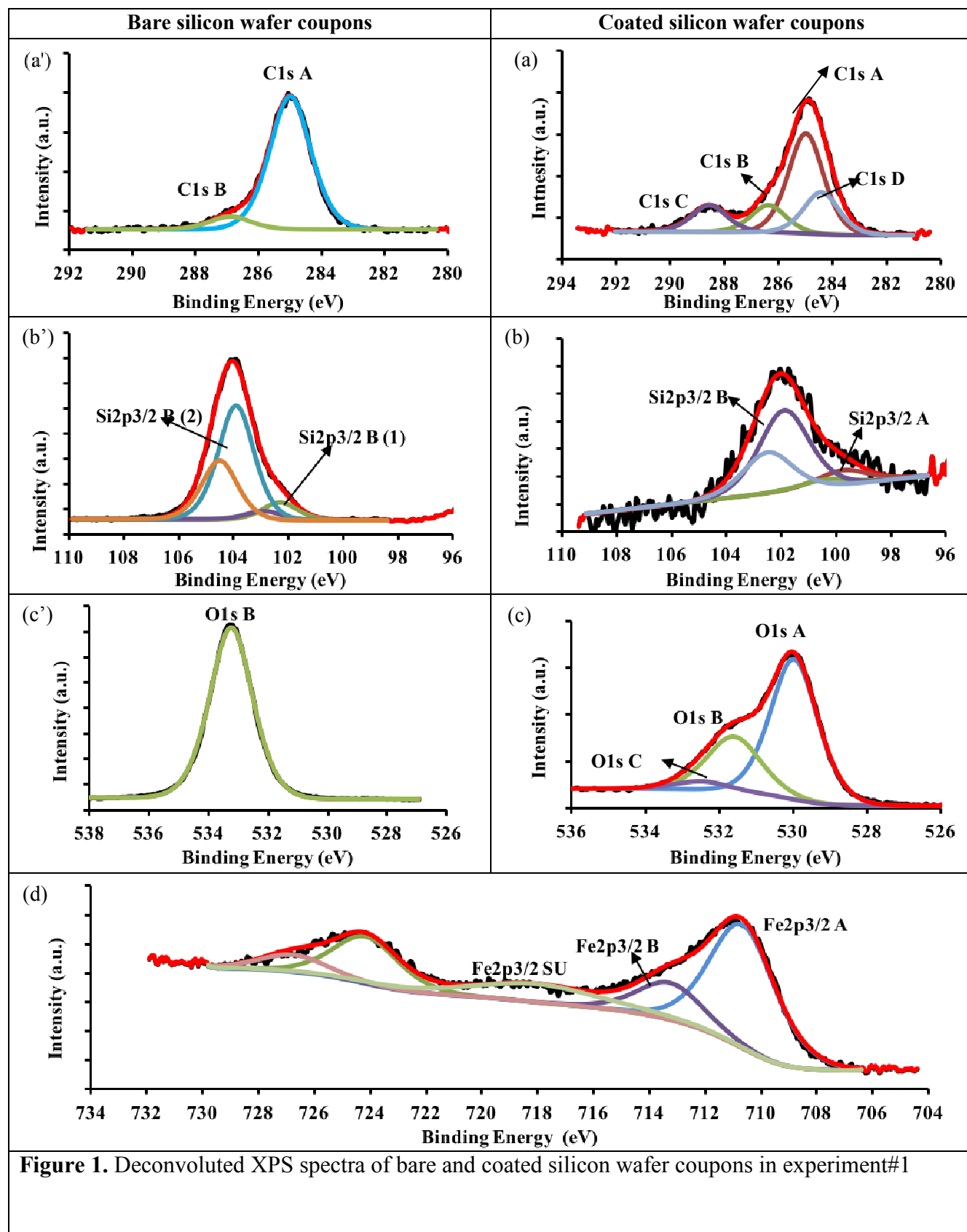
**Table 1.** XPS high-resolution spectral deconvolution peaks and their origins.

Peak Name	Centered Binding Energy [eV]	Possible Structure <sup>(1)</sup>	Elemental atomic %	
			Bare sample	Coated sample <sup>(2)</sup>
C1s A	285.0	*C-*C, *C-H, *C=*C, etc.	6.4	15.0
C1s B	286.4	*C-O-*C, *C-O-Fe,	1.2	2.4
C1s C	288.6	O-*C-O, O=*C-O, O=*C-O-Fe	-	3.0
C1s D	284.4	Si-*C	-	1.9
O1s A	530.0	Fe-*O	-	36.7
O1s B	531.6	Fe-*O-H, C=*O, Si-*O-H	64.1	8.6
O1s C	532.5	C-*O-Fe, *O=C-*O-Fe	-	6.4
Si2p3/2 A	99.8	*Si-*Si, *Si-Fe	-	1.1
Si2p3/2 B	101.9	H <sub>3</sub> C-*Si(O)-CH <sub>3</sub>	-	2.0
Si2p3/2 B (1)	102.4	Si <sup>+2</sup> Polycrystalline Si in Si-O <sub>x</sub>	4.5	-
Si2p3/2 B (2)	103.9	Si <sup>+4</sup> Polycrystalline Si in Si-O <sub>x</sub>	23.8	-
Fe2p3/2 A	710.7	*Fe-O, O=C-O-*Fe	-	13.5
Fe2p3/2 B	713.3	C-O-*Fe	-	4.8
Fe2p3/2 SU	718.0	C-O-*Fe, O=C-O-*Fe	-	4.6

<sup>(1)</sup> \*shows the elements correspond to that binding energy.

<sup>(2)</sup> Elemental atomic % correspond to the experiment #1 (**Table 1S**).





(Table 1S)

Before treatment, the silicon wafer surface was mainly composed of silicon, oxygen and contained a small amount of carbon, most likely due to contaminants<sup>42,43</sup> (~7.6%, **Table 1**). These contaminants (**Figure 1a'**) exhibited a high-resolution C1s peak at binding energy BE  $\cong$  285.0 eV and 286.9, which were assigned to C–C and C–H bonds (such as –CH, –CH<sub>2</sub>, –CH<sub>3</sub> and C<sub>2</sub>H<sub>2</sub>, C<sub>2</sub>H<sub>5</sub>,...) and C–O–C bonds, respectively. Silicon sub-oxides such as SiO (Si<sup>2+</sup>), Si<sub>2</sub>O<sub>3</sub> (Si<sup>3+</sup>) and SiO<sub>2</sub> (Si<sup>4+</sup>) were assigned to BE of 101.4–102.0, 102.1–102.7 and 103.5–104.1 eV within the Si2p peak, respectively.<sup>45,46</sup> Thus, the broad distribution of Si2p spectra at 102.2–104.2 eV was attributed to the various Si–O<sub>x</sub> species (**Figure 1b'**). The oxygen peak at BE  $\cong$  533.3 eV represented oxygen in the oxide form (**Figure 1c'**). High-resolution spectral deconvolution of Si2p and O shows that the atomic percentage of oxygen, and silicon did not match. Indeed, in terms of stoichiometry, the Si/O ratio should be 1/2, but the amount of oxygen was determined to be higher (for 28.3 At % Si, the corresponding atomic percentage for oxygen should be 56.6% (**Table 1**). The observed higher oxygen amount (+7.5%, **Table 1**) was attributed to surface hydroxyl groups (Si–OH).<sup>45,47</sup>

After PICVD treatment, the C1s peak increased significantly, and the Si2p peak was attenuated, which was in agreement with survey results (**Figures 2S** and **3S**). The C1s spectrum (**Figure 1a**) showed a group of overlapping peaks with BE from 282 to 292 eV, attributed to four significantly different surface carbon phases (identified as A, B, C and D). The dominant Phase A peak (C1s A, BE  $\cong$  285.0 eV) was attributed to aliphatic carbon (e.g. –CH, –CH<sub>2</sub>, –CH<sub>3</sub>), as well as polymerized C<sub>n</sub>H<sub>m</sub> chains (e.g. C $\equiv$ C, C=C, C–C). Phase B (C1s B, BE  $\cong$  286.4 eV) contained C=CO–C and C–O–C, though it also could be assigned to C–O–Fe. Phase C (C1s C, BE  $\cong$  288.6 eV) could correspond to O=C–C, O–C–C, and also O=C–O–Fe.<sup>6</sup> This range of BE (286.0–289.0 eV) is generally attributed to carbonates. Phase D (C1s D, BE  $\cong$  284.4 eV) was designated as a carbidic carbon layer, in the form of Si–C bonds. This means that the organic films we deposited had bonded with the silicon substrate

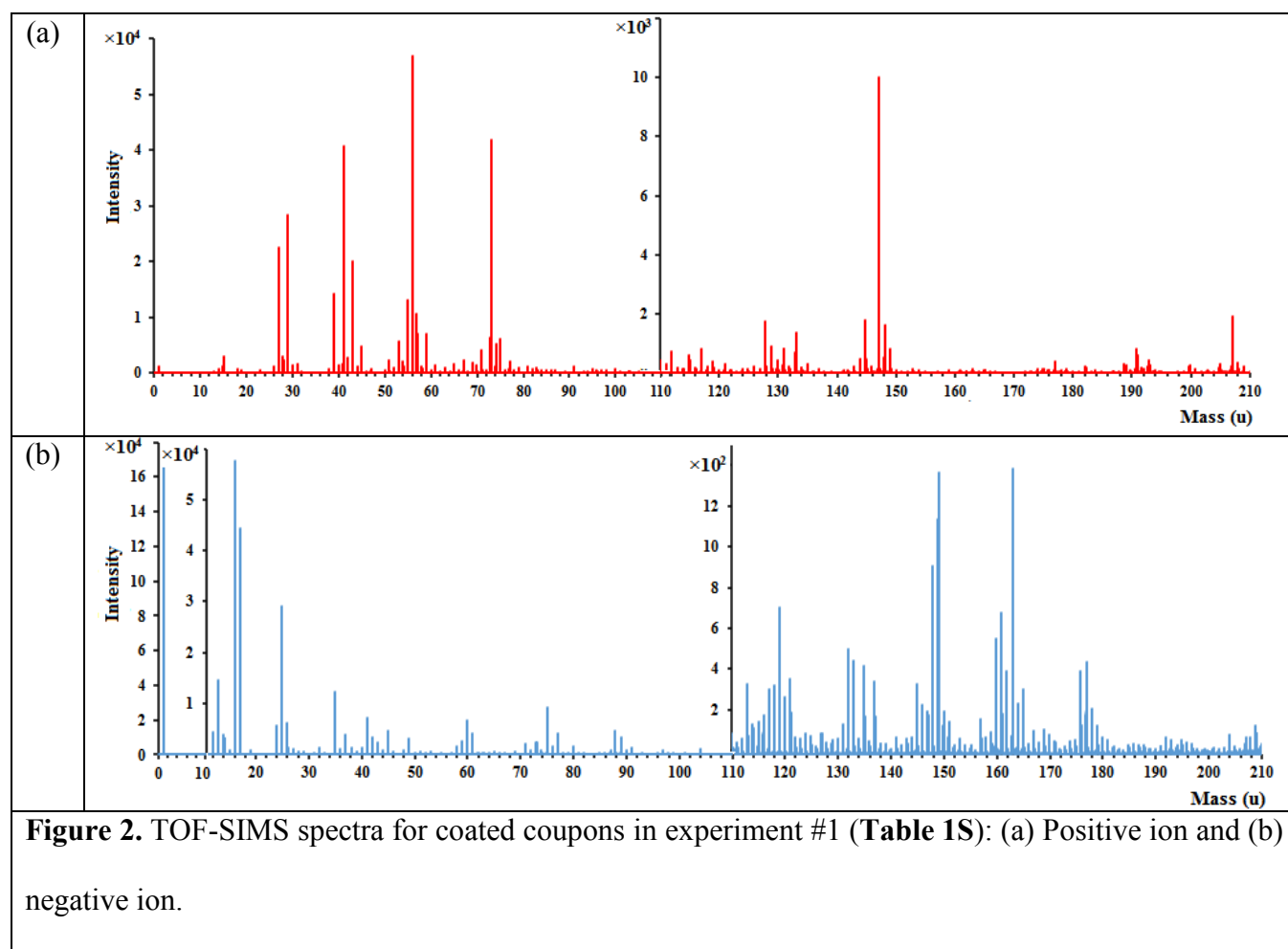
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2 covalently. Since there is no C1s peak on the untreated sample at  $BE \leq 284.4$  (**Figure 1a'**), this  
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4 covalent bond was formed during the PICVD process, which was confirmed by the Si2p spectrum: in  
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6 **Figure 1c**, the attenuation of the doublet Si2p<sub>3/2</sub> and Si2p<sub>1/2</sub> in Si-O<sub>x</sub> (102-104 eV) was observed  
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8 after the organic grafting. In grafted silicon samples, the Si2p spectrum consisted of two main peaks  
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10 and their doublet (A, B) located at  $BE \cong 99.7$  eV, and  $BE \cong 101.8$  eV. The first peak and its  
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12 corresponding doublet exhibited the characteristic of Si2p<sub>3/2</sub> with  $BE \cong 99.7$ – $99.9$  eV and Si2p<sub>1/2</sub>  
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14 with  $BE \cong 100.2$ – $100.7$  eV. This corresponded to elemental silicon in the form of Si–Si or even Fe–Si  
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16 bond in the structure of the deposited film.<sup>46,48</sup> The second peak was more alike to the binding energy  
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18 in organic silicon  $\sim 102$  eV (Si2p<sub>3/2</sub> and Si2p<sub>1/2</sub> of 101.7 eV and 102.4 eV respectively).<sup>46</sup> Based on  
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20 the elemental atomic percentage (**Table 1**), we propose the H<sub>3</sub>C–Si(–O)–CH<sub>3</sub> structure for this peak.  
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22 Other researchers have shown that such covalent bonding is possible through plasma polymerization  
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24 under low pressure conditions.<sup>49</sup> Here, we demonstrate the formation of a strong covalent bond under  
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26 gentler, atmospheric PICVD processing conditions.  
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33 Using Gaussian fitting and literature reports,<sup>46,50</sup> the O1s spectrum (**Figure 1b**) can be divided into  
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35 three peak areas: A, B and C. A ( $BE \cong 530$  eV) is attributed to oxide species. B ( $BE \cong 531.6$  eV) is  
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37 assigned to hydroxyl groups such as H–O–Fe, H–O–Si, –O=C, and C–O–O agents in ether and alcohol.  
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39 C ( $BE \cong 532.5$  eV) corresponds to C–O–Fe, O=C–O–Fe species.  
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43 Peaks in the BE range of 705 to 740 eV have been attributed to iron.<sup>46</sup> In syngas PICVD, Fe can  
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45 come from carbon monoxide cylinders, in the form of iron pentacarbonyl (Fe(CO)<sub>5</sub>). This compound is  
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47 generated in small concentrations over extended periods in CO cylinders<sup>51,52</sup> and happens to be photo-  
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49 active in the UVC range.<sup>53</sup> The peak deconvolution of the Fe2p spectrum (**Figure 1d**) showed two  
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51 main phases. Phase A was composed of Fe2p<sub>3/2</sub> ( $BE \cong 710.2$  eV) and Fe2p<sub>1/2</sub> ( $BE \cong 724.5$  eV);  
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53 these were assigned to iron oxide, either in the free state or in chemical bonds with the organic film  
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55 (e.g. O–Fe, and O=C–O–Fe), with satellite peaks at  $BE \cong 718.3$  eV (Fe2p<sub>3/2</sub> satellite) and  $BE \cong 732.5$   
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eV (Fe2p1/2 satellite). Based on the literature,<sup>46,54-57</sup> for iron sub-oxides such as FeO (Fe<sup>2+</sup>), Fe<sub>2</sub>O<sub>3</sub> (Fe<sup>3+</sup>) and Fe<sub>3</sub>O<sub>4</sub> (Fe<sup>8/3+</sup>), binding energy of Fe2p3/2 spectra are 709.1-710.7, 710.6-711.5 and 707.90-710.7 eV, respectively. Thus, the binding energies in phase A corresponded to either Fe<sup>3+</sup> in Fe<sub>2</sub>O<sub>3</sub> or Fe<sup>8/3+</sup> in Fe<sub>3</sub>O<sub>4</sub>. Phase B corresponded to the Fe2p3/2 and Fe2p1/2 peaks located at BE  $\cong$  713.4 and 726.3 eV, which can be assigned to Fe–O–C and Fe–(OH)<sub>3</sub>. This indicates that Fe(CO)<sub>5</sub> is fully decomposed and converted to iron oxide which had been incorporated in the chemical structure of deposited film.

A complementary characterization of chemical bonds between the silicon wafer and the carbon film was then performed by TOF-SIMS analysis. The latter provided more chemical information on the deposited film and further supported our XPS results (**Figure 2**).



1  
2 TOF-SIMS results can be divided into three main categories:  
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4 (1) Hydrocarbons: Both in the acquired positive and negative spectra, the expected  $C_nH_m$  fragments  
5 of light polymer-like species were detected. Positive ion spectra suggested that a great number of  
6  $C_2H_2^+$ ,  $C_2H_3^+$ ,  $C_2H_5^+$ ,  $C_3H_3^+$ ,  $C_3H_7^+$ ,  $C_4H_9^+$ ,  $C_5H_9^+$ ,  $C_6H_5^+$ ,  $C_6H_6^+$ ,  $C_7H_7^+$ ,  $C_8H_7^+$ ,  $C_8H_{13}^+$ ,  $C_9H_7^+$ ,  $C_9H_{15}^+$ ,  
7  $C_{10}H_8^+$ ,  $C_{11}H_9^+$ , up to  $C_{12}H_8^+$  ions at  $m/z = 26, 27, 29, 39, 43, 57, 69, 77, 78, 91, 103, 109, 115, 123,$   
8  $128, 141,$  and  $152,$  respectively, were generated by the PICVD treatment. These fragments could be  
9 either linear or cyclical. Similar observations have been made in the negative iron spectrum at  $m/z =$   
10  $24, 37, 48, 49, 51, 60, 61, 64, 65, 72, 78, 83,$  and  $96$  corresponding to  $C_2^-$ ,  $C_3H^-$ ,  $C_4^-$ ,  $C_4H^-$ ,  $C_4H_3^-$ ,  $C_5^-$ ,  
11  $C_5H^-$ ,  $C_5H_4^-$ ,  $C_5H_5^-$ ,  $C_6^-$ ,  $C_6H_6^-$ ,  $C_7^-$ , and  $C_8^-$  ions, respectively. Moreover, some fragments with O in  
12  $C_nH_mO_y$  formula like  $COOH^-$ ,  $C_3H_3O_2^-$ ,  $C_5H_5O^-$ ,  $C_6H_5O^-$ , and  $C_7H_6O^-$  were detected at  $m/z = 44, 71, 81,$   
13  $93,$  and  $106,$  respectively. These species were related to the carboxylic acid, anhydride and ketone  
14 **(Figure 4S).**  
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30 (2) Iron: The spectrum obtained from the positive ions was dominated by peaks representing iron  
31 ions. The strongest peaks at  $m/z = 56$  corresponded to  $Fe^+$ ; whereas, the other peaks at  $m/z = 57, 68,$   
32  $69, 70, 71, 72, 73, 80, 84, 85, 86, 89, 96, 101, 112, 124, 128, 132,$  and  $136$  in the positive ion spectrum  
33 corresponded to different species in the form  $Fe^+$ ,  $FeH^+$ , and  $Fe_xO_yH_mC_n^+$ , e.g.,  $FeH^+$ ,  $CFe^+$ ,  $FeCH^+$ ,  
34  $CH_2Fe^+$ ,  $FeCH_3^+$ ,  $FeO^+$ ,  $FeOH^+$ ,  $C_2Fe^+$ ,  $COFe^+$ ,  $FeCOH^+$ ,  $CH_2OFe^+$ ,  $FeCOH_3^+$ ,  $C_2OFe^+$ ,  $FeCOOH^+$ ,  
35  $Fe_2^+$ ,  $CFe_2^+$ ,  $Fe_2O^+$ ,  $C_5OFe^+$ , and  $C_2Fe_2^+$ , respectively. Despite these peaks, there was no peak related to  
36 the  $Fe(CO)_5$  in the negative/positive ion spectra of TOF-SIMS confirming our XPS results indicating  
37 this compound was fully decomposed. Several combinations of iron oxide species (e.g.  $Fe_2O_2^+$ ,  $Fe_3O^+$ ,  
38  $Fe_3O_2^+$ ,  $Fe_4O_2^+$ ) were also found at  $m/z$  ranging from 148 to 503. These fragments were also observed  
39 in the negative ion mode, as well as fragments in the form of  $FeOH^-$ ,  $Fe_xO_yH_mC_n^-$  and  $OH^-$  species.  
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54 (3) Silicon: The relevant negative and positive ions spectra for silicon showed the presence of  $Si^+$ ,  
55  $SiC^+$ ,  $SiHO^+$ ,  $SiOCH_3^+$ ,  $SiOC_2H_6^+$  ( $m/z = 28, 44, 45, 59, 74,$  respectively) and their relevant negative  
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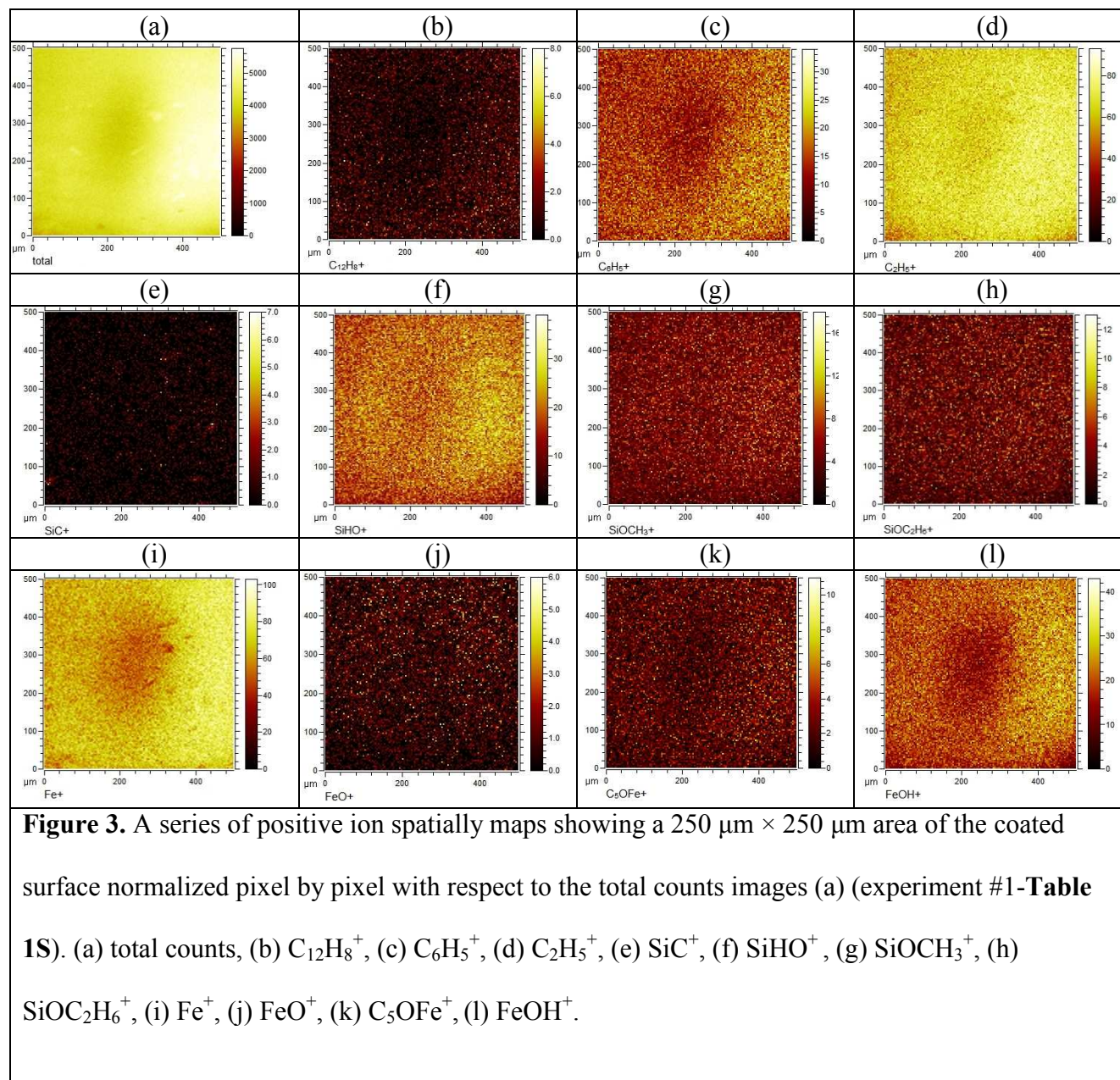
ions such as  $\text{SiCO}^-$ ,  $\text{SiC}_2\text{O}^-$ ,  $\text{SiC}_2\text{OH}^-$  (at  $m/z = 56, 68, 69$ , respectively). These results corroborated the suspected structure obtained from XPS.

In addition, to confirm the chemical structure of the thin film, TOF-SIMS helped us to investigate the abundance and homogeneity of deposited chemical moieties. Thus, ion imaging was carried out on a deposited film (experiment #1, **Table 1S**) to identify the ion fragments in the film and their micro-scale spatial distribution (**Figure 3**). We have applied a pixel-by-pixel normalization with respect to the total ion counts. Integrated mass spectrum at each pixel (**Figure 3a**) for all images has been performed. This removed any effect of preferential sputtering and secondary ion yield effects due to topography.

**Figure 3b-d** show the spatial distribution of some light hydrocarbons.  $\text{C}_2\text{H}_2^+$ ,  $\text{C}_2\text{H}_5^+$ ,  $\text{C}_3\text{H}_3^+$ , ...  $\text{C}_{12}\text{H}_8^+$  were observed with dominant distribution of  $\text{C}_2 > \text{C}_3 > \text{C}_4$  aliphatic branches. The maximum number of carbon in the polymer reached  $\text{C}_{12}$ . Also, aromatic structures were detected in the deposited film, most likely phenol, which is in agreements with previous results.<sup>9</sup> Results from these images and TOF-SIMS ion spectra showed formation of acid anhydride, aldehyde, ketones, carboxylic, and hydroxyl groups as well.

**Figure 3e-h** are representative of the spatial distribution of Si ions at the interface between the deposited film and substrate. The SiC distribution images corroborated the formation of covalent bonds between the oligomeric film and the substrate, previously identified by XPS. Moreover, the structures of these covalent bonds are mainly in the form of  $\text{SiOCH}_3$  and  $\text{SiO}(\text{CH}_3)_2$  at the silicon wafer/deposited film interface. Considering the presence of  $\text{SiHO}^+$  but not  $\text{SiO}_2$  in the sample, the interface consisted mainly of SiOH groups.

**Figure 3i-l** represent some of the iron fragments in the deposited film. Most of the iron in the film was in the form of iron oxide (especially  $\text{Fe}^+$ ) and iron hydroxyl ( $\text{FeOH}$ ). Moreover, some iron was in the form of  $\text{COFe}^+$ ,  $\text{C}_3\text{OFe}^+$ ,  $\text{C}_5\text{OFe}^+$ , etc. while the ion spectra detected Fe bonding to carbon ( $\text{FeC}^+$  and  $\text{CH}_2\text{Fe}^+$ , etc.).



### 3.2. Proposed mechanism

Based on the in-depth characterization of the chemical structure of the deposited films, we can shed a light on their growth mechanism. Based on the chemical characterization, C, O, H, Fe, Si are the elements existing in the samples. Si and partial O are from the substrate, while the other elements C, H,

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2 Fe and partial O are mainly coming from the reactive species. **Figure 4a** shows the chemical structure  
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4 of the outmost layer of the silicon wafer substrate, which mainly consists of SiO<sub>2</sub> and SiOH, and  
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6 chemisorbed water. Since the reaction depends on the UV lamps, the absorption cross sections of the  
7  
8 participating reactive species are investigated.  
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11 Hydrogen (H<sub>2</sub>) has an ionization potential of 15.42 eV and a dissociation energy of 4.52 eV.<sup>37,58</sup>  
12  
13 Thus, photons at wavelengths below 80.4 nm and 274.3 nm are capable of ionizing and dissociating H<sub>2</sub>,  
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15 provided they can be absorbed. However, H<sub>2</sub> does not absorb light above 110.8 nm.<sup>37</sup> Given that the  
16  
17 UVC lamps used in this study emit at 253.7 nm (secondary peak at 185 nm), there will be no energy  
18  
19 absorption. In other words, hydrogen cannot participate in the reaction, unless excited by other  
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21 molecules and radicals.  
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25 Carbon monoxide has an ionization potential of 14.01 eV<sup>59</sup> and a dissociation energy of 11.14 eV.<sup>58</sup>  
26  
27 This corresponds to a potential for ionization and dissociation at wavelengths below 88.5 nm and 112.3  
28  
29 nm, respectively. CO can absorb light at wavelengths shorter than 200 nm,<sup>37</sup> which means that the  
30  
31 secondary emission peak at 185 nm can transmit 6.7 eV of energy to the molecule. While this is  
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33 insufficient for ionization or full dissociation, it is enough to dissociate a constituent bond (3.7 eV) in  
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35 CO or cause CO to activate through vibrational excitation.<sup>35</sup>  
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39 Iron pentacarbonyl (Fe(CO)<sub>5</sub>) fed concurrently with the CO gas,<sup>51,52</sup> has an ionization potential of  
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41 8.35 eV and dissociation energy of 0.37 eV,<sup>60,61</sup> corresponding to wavelengths of 148.5 nm and 3350.5  
42  
43 nm, respectively. Fe(CO)<sub>5</sub> has strong absorption at 253.7 nm wavelength. It is therefore able to  
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45 dissociate, resulting in the formation of extra radicals.<sup>53,62</sup>  
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49 Thus, the main reactive components following UV irradiation are vibrationally excited CO (CO\*), or  
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51 CO radicals (CO•) with dissociated bonds and CO• stemming from Fe(CO)<sub>5</sub> dissociation. Based on  
52  
53 these postulations, two colliding CO\* molecules will form a highly unstable unit. Once excess energy  
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55 has been dissipated via collisions of the nascent cluster with H<sub>2</sub> molecules, lightly polymerized carbon  
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species (mostly consisting of aliphatic chains with  $C_2 > C_3 > C_4$ ) will be formed.<sup>35,36,63</sup> Iron pentacarbonyl can also play a catalytic role for isomerization of olefins and can increase the formation of alcohols on the film instead of carboxylic groups.<sup>64</sup>

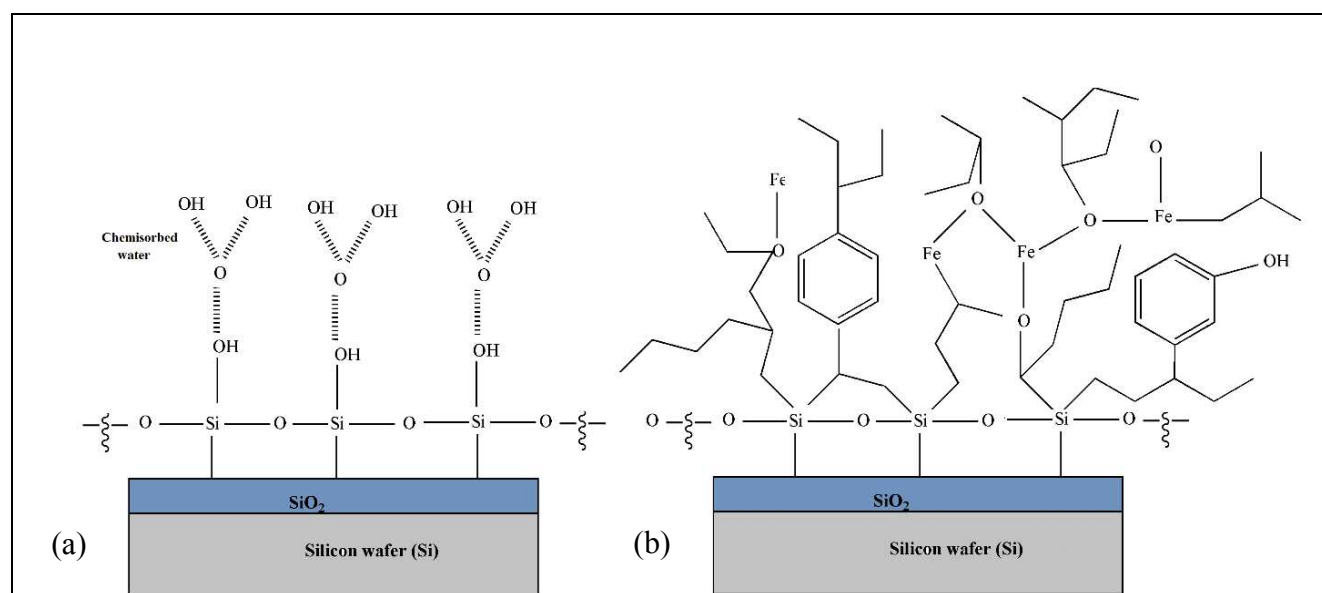
Based on the previously identified chemical bonds, as well as the existing Fisher-Tropsch literature,<sup>35,36,53,64</sup> we present a series of possible chemical reactions. Initiation mechanisms are based on the absorption cross section, while propagation and termination are based on the structure we obtained and known syngas chemical reactions:

<b>Initiation</b>	$CO(\pi\text{-bond dissociation}) + H_2 \leftrightarrow HCO^\bullet + H^\bullet$		(1)
	$CO(\pi\text{-bond dissociation}) + H_2 \leftrightarrow H_2CO^\bullet$		(2)
	$2CO^\bullet + H_2 \leftrightarrow CH_2^\bullet + CO_2$		(3)
	$H^\bullet + CO^\bullet \leftrightarrow HCO^\bullet$		(4)
	$Fe(CO)_5 + hv \leftrightarrow Fe + 5CO^\bullet$		(5)
	$Fe(CO)_5 + hv \leftrightarrow Fe(CO)_4^\bullet + CO^\bullet$		(5)
	$Fe(CO)_4^\bullet + hv \leftrightarrow Fe(CO)_3^\bullet + CO^\bullet$		(6)
<b>Propagation</b>	$HCO^\bullet + H_2 \rightarrow H_2CO^\bullet + H^\bullet$		(7)
	$CH_2^\bullet + H_2 \leftrightarrow CH_3^\bullet + H^\bullet$		(8)
	$CH_3^\bullet + 2CO \leftrightarrow CH_3O^\bullet + CO^\bullet$		(9)
	$CH_3O^\bullet + HCO^\bullet \rightarrow CH_3OH + CO^\bullet$		(10)
	$CH_3O^\bullet + H_2CO^\bullet \rightarrow CH_3OH + HCO^\bullet$		(11)
	$2CH_2^\bullet + H_2 \rightarrow C_2H_2 + 2H^\bullet$		(12)
	$H_2CO^\bullet + CO \leftrightarrow CO^\bullet + H_2CO^\bullet$		(13)
<b>Termination</b>	$2CH_3O^\bullet + 2H_2 \rightarrow C_2H_5 + 2H_2O + HO^\bullet$		(14)
	$CO^\bullet + HCO^\bullet + CH_3O^\bullet \rightarrow CH_3OOH$		(15)
	$Fe(CO)_2^\bullet + HO^\bullet \rightarrow FeOH + 2CO^\bullet$		(16)
	$CO^\bullet + 2HO^\bullet \rightarrow CO_2 + H_2$		(17)
	$nCO + (2n + 1)H_2 \rightarrow C_nH_{2n+1} + nH_2O$		Paraffins (18)
	$nCO + (2n)H_2 \rightarrow C_nH_{2n} + nH_2O$		Olefins (19)
	$CO + H_2O \rightarrow H_2 + CO_2$		Water-gas shift (20)

Reaction (7) is endothermic ( $16.4 \text{ kcal mol}^{-1}$ ), which makes this reaction most likely irreversible. It needs  $20 \text{ kcal mol}^{-1}$  activation energy, provided mostly through exothermic reactions (such as (4), which gives off  $17 \text{ kcal mol}^{-1}$ ).<sup>35</sup> On the other hand, for most free-radical polymeric chain reactions,

vibrational excitation is key to provide the driving force for propagation step. As such, reaction (13) must maintain this driving force by transferring vibrational energy to CO, otherwise the reaction would move towards termination.<sup>35</sup> These pathways are based on observations by Roussel and Back, who had conducted research on the photochemistry of CO/H<sub>2</sub> mixtures irradiated with light at 193.3 nm (similar to the secondary emission peak of our Hg lamps).

This proposed series of reactions leads to the formation of light paraffins, olefins and iron oxide on the surface of the Si substrate (with water as an expelled by-product) (**Figure 4b**). Most reactive species are bonded to iron and also can react with either hydroxyl groups on the Si surface or directly with Si to bond covalently. XPS and TOF-SIMS results indicate that the film is composed of aliphatic branches and aromatic structure (most likely phenol, given the presence of C<sub>6</sub>H<sub>5</sub>O<sup>-</sup> at m/z = 93, supported by previous analyses<sup>9</sup>), as well as acid anhydride, ketones, carboxylic, and hydroxyl moieties.



**Figure 4.** Schematic of (a) bare silicon wafer (b) deposited oligomeric film on silicon wafer.

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2 Chemical characterization results showed uniform distribution of chemical bonds in the deposited  
3 film. These observations triggered us to explore if we have same uniformity in the morphology and  
4 texture of the deposited film.  
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### 9 3.3. Surface morphology and topography

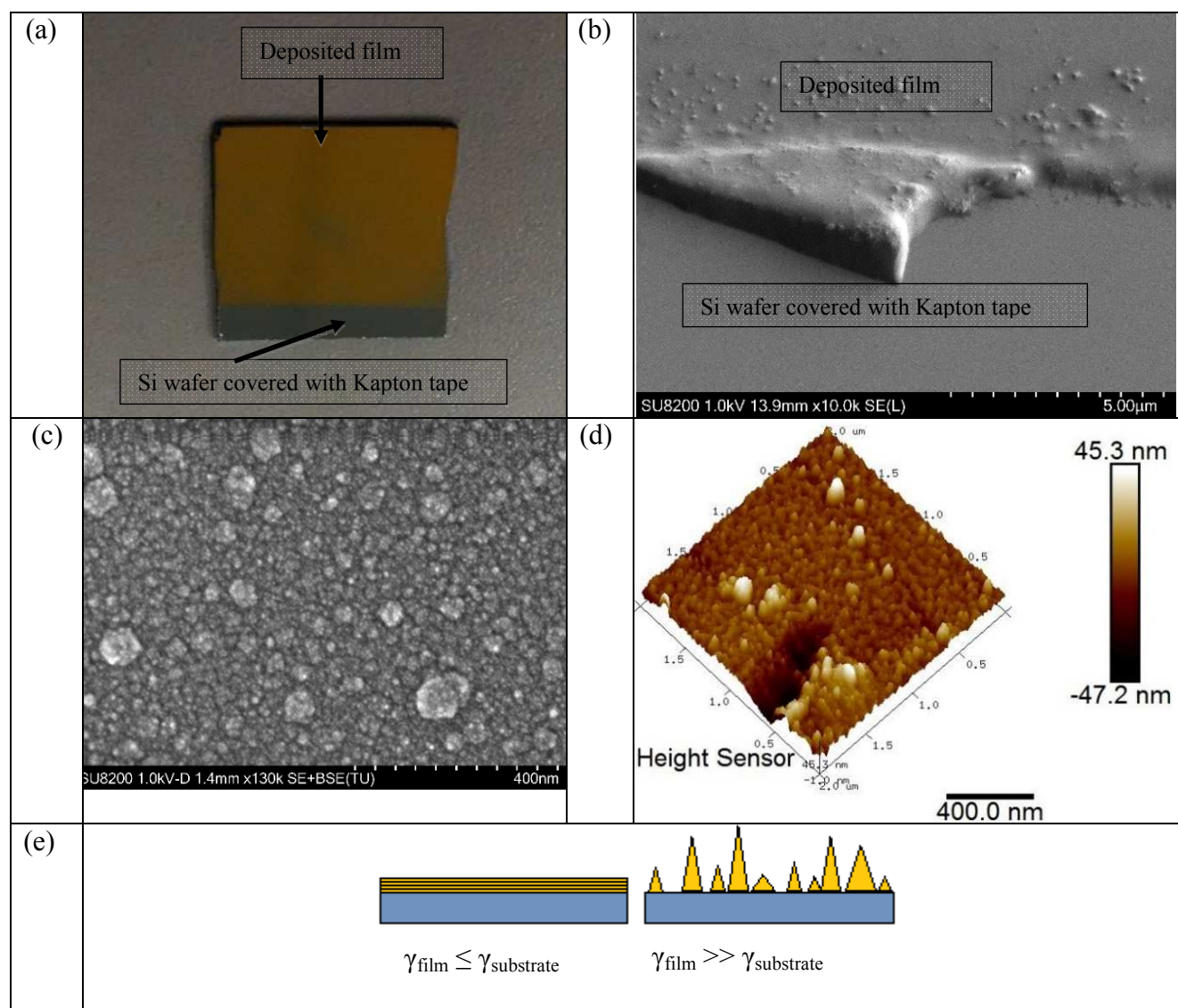
10 The electron micrographs of a coated silicon substrate (**Figure 5b, 5c**) reveal the formation of a  
11 textured, particle-like morphology with many micro patterns and trenches along the deposited film.  
12 This surface morphology is corroborated by AFM (**Figure 5d**), showing that the features have a trench  
13 size on the order of 40 nm. Such island-like deposition is consistent with previous investigations on  
14 PICVD.<sup>9</sup> This textured morphology is observed even for very low film thickness (treatment duration  $t$   
15 = 30min). The trenches are filled in as treatment duration increases, leading to a more uniform film (see  
16 Supplementary **Figure 5S** profilometry results).  
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28 This kind of trench morphology of film formation is common especially in the case of transient  
29 metals,<sup>65</sup> and can be explained in our case by the involvement of iron and iron oxide in the deposition  
30 reaction. Film deposition in this case follows the Volmer–Weber (VW) growth mode (3D morphology,  
31 island growth) and allows us to gain information on the surface tension of the film through Bauer's  
32 criterion ( $\gamma_{\text{film}} \gg \gamma_{\text{substrate}}$ ,  $\gamma_{\text{substrate}} = 44.16 \text{ mJ/m}^2$ , **Figure 5e**).<sup>66</sup> Control of the film morphology (and  
33 therefore macroscopic properties such as wettability) is the focus of on-going work, namely through the  
34 action of treatment pressure.  
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45 Through energy-dispersive X-ray spectroscopy (EDS), SEM measurements allowed us to further  
46 confirm the uniform elemental distribution and composition of the films (**Figure 6S**). Bare silicon  
47 substrates show mainly Si and O, with some C (**Figure 6Sa**) – C attributable to contamination,<sup>42,43</sup>  
48 while the coated sample (**Figure 6Sb**) exhibits an increase in the C peak and the presence of an Fe peak  
49 in the deposited film. The mapping analyses of bare and coated samples in **Figures 6Sc-f** confirm the  
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formation of a film (clear contrast with the reference area covered with Kapton tape during treatment).  
As we move further to the right-hand side, the amounts of Fe and C increase.

The film also exhibits stability to solvent attacks, resisting immersion for 1 hour in chloroform (Table 2S), in agreement with previous work showing stability with respect to solvents.<sup>9</sup>



**Figure 5.** (a) Coated silicon coupon, (b) and (c) FESEM images of deposited film, (d) 3D AFM image of deposited thin film in experiment #1, (e) Bauer's thermodynamic film growth study with respect to surface tensions.

### 3.4. Processing parameters

In an effort to control the thickness of the films deposited through syngas PICVD, we conducted a series of experiments focusing on two main parameters (i) residence time and (ii) treatment duration of experiments.

The time during which the interactions between the UV light and the gas precursors occur, resulting in the production of reactive species, depends on the gas residence time ( $\tau$ ). It can be calculated from Equation 1, in which  $V$  stands for the reactor volume and  $Q$  for the gas precursors volumetric flow rate. For a constant reactor volume, decreasing the flow rate leads to longer precursor residence time and saves consumption of gas precursors, which can be important from an operational point of view.

$$\tau = \frac{V}{Q}$$

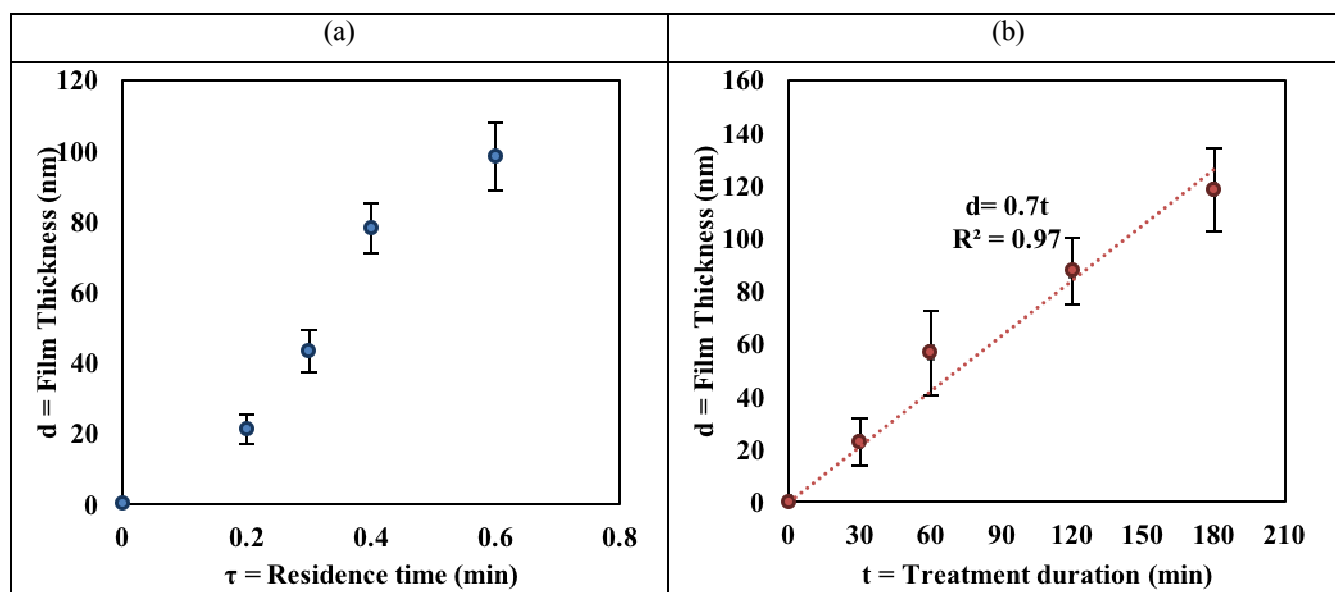
Equation 1

Using a constant gas feed pressure (40 psi), a volumetric precursor ratio (CO/H<sub>2</sub>) of 1 and a total gas flow rate varying from 0.4 to 1 L/min for a 120 min treatment duration, we varied the reactor residence times from 0.6 to 0.2 min (**Table 1S**). Film thickness (assessed through profilometry) and elemental distribution (determined from XPS measurements) are given as a function of residence time in **Figures 6a and 2S**.

As deduced from **Figure 6a**, the film thickness increased with residence time. Linear regression ( $R^2 = 0.95$ ) indicates accumulation at a rate of approximately 165 nm of film per minute of residence time (note that  $\tau = 0$  corresponds to an untreated, bare substrate), though the shape of the curve more closely resembles a sigmoid (indicating diminishing returns at longer residence times). The bare substrate was composed primarily of Si and O, along with some C attributable to contaminants<sup>42,43</sup> (**Figure 2S**). For coated substrates ( $\tau = 0.2 - 0.6$  min), the C atomic percentage increased to a uniform value of approximately 30%, while Si decreases significantly, indicating that the substrate was masked. The

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2 elemental distribution did not vary as a function of  $\tau$ , indicating that film composition and thickness  
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4 were independent. Also, despite XPS' shallow probing depth ( $\sim 10$  nm), Si remained visible on the  
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6 thicker films which is explained by the morphology of the film (see section 3.3).  
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9 In order to increase the thickness of the deposited films while decreasing precursor consumption (i.e.  
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11 at longer residence time), we increased the treatment duration. **Figure 6b** shows the effect of treatment  
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13 duration (30 to 180 min treatment duration, at a fixed residence time of  $\tau = 0.6$  min) on film thickness.  
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15 Full thickness profiles are available in Supplementary **Figure 5S**. As expected, film thickness increased  
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17 linearly with treatment duration,<sup>18,44</sup> at a rate of 0.7 nm/min for  $\tau = 0.6$  min.  
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**Figure 6.** (a) Profilmometry film growth results vs. residence time (experiments # 1 to # 4, **Table 1S**; error bars indicate the 95% confidence interval (95%CI)); (b) Variation of film thickness as a function of treatment duration (Data correspond to experiments # 5 to # 8, **Table 1S**; error bars indicate the 95%CI).

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As already noted for residence time experiments, the amount of Si significantly decreased in the treated samples compared to the bare one. Furthermore, the Si At % decreased with increasing deposition time until 120 min, where it reached a plateau (**Figure 3S**). This would warrant that by

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2 increasing the treatment duration, trenches are masked in the deposited film and a more uniform film is  
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4 achieved, which is confirmed by profilometry analyses (Supplementary results, **Figure 5S**).

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6 These findings are important in two respects: first, both the residence time and treatment duration  
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8 affect the film growth and thickness. Second, lower treatment duration or residence times lead to a less  
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10 homogeneous film texture (Si visible at low  $t$  and low  $\tau$ ). Finally, combining longer residence time and  
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12 varying treatment duration allows for control over the thickness of the film while limiting gas precursor  
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14 consumption. After 60 min of treatment, there is no statistically significant evolution in the elemental  
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16 composition of the film (**Figure 3S**). We suspect that, before 60 min, the observed difference is  
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18 attributed to the fact that the substrate is not completely covered (trenches in the film).  
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22 These treatment duration experiments also provide insight into the reaction mechanism. As described  
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24 in the experimental section, temperature increases as a function of time during treatment, because of  
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26 heat given off by the UVC lamps. The fact that the deposition rate does not decrease as a function of  
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28 time (and therefore increasing temperature, and correspondingly decreased reagent adsorption) may  
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30 indicate that the overall reaction mechanism is kinetically limited. However, there is also evidence to  
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32 suggest that mass transport may be limiting the overall mechanism, given that one key reagent (iron  
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34 pentacarbonyl) is present at very low concentrations. On-going work is focusing on decoupling time  
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36 and temperature to better assess this phenomenon.  
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42 We have shown that film composition is independent of residence time and treatment duration, at a  
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44 fixed precursor ratio. Given the insights gained from the reaction mechanism, there is reason to believe  
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46 that varying the precursor ratio may alter the composition and/or growth of the organic film. To  
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48 investigate this, the gas ratio ( $\text{CO}/\text{H}_2$ ) was varied from 0.3 to 3, while the residence time and treatment  
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50 duration were kept constant at 0.3 min and 2h respectively (**Table 1S**). **Figures 7S** and **8S** show that  
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52 the variation of gas ratio does not a significant effect both in terms of film growth and chemical  
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54 composition. The absence of a measurable chemical change after changing the precursor ratio is  
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2 intriguing, as our previous work had indicated changes in macro-level properties (wettability) as a  
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4 function of ratio.<sup>9</sup> To confirm, we tested three samples with CO/H<sub>2</sub> ratio = 0.3, 1 and 3 (minimum,  
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6 middle, and maximum) with TOF-SIMS. Three spots were analyzed on each sample and their spectra  
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8 were compared to ensure they were representative the whole sample (**Figures 9S and 10S**); these  
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10 corroborate the high resolution XPS results. Based on these findings, we can conclude that any  
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12 wettability change previously observed is not likely to be a result of a chemical change, but rather a  
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14 morphology change on the surface.  
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#### 19 **4. CONCLUSIONS**

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21 Through this work, we have gained insight into the growth mechanism of a novel deposition  
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23 approach (syngas PICVD) and proposed a reaction scheme. Both XPS and TOF-SIMS reveal the  
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25 formation of covalently bonded oligomeric film with aliphatic branches and aromatic structure (most  
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27 likely phenol), as well as acid anhydride, ketones, carboxylic, and hydroxyl moieties. Moreover, we  
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29 have shown that film thickness can be controlled through treatment duration (as expected), but also  
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31 through residence time (a feature specific to atmospheric pressure deposition processes). This gives us  
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33 an ability to control thickness while minimizing precursor consumption. Interestingly, altering the  
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35 precursor gas ratio does not affect the final composition of the film. However, equal parts seem best  
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37 suited to provide a slightly thicker film.  
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42 Despite these significant findings, a number of potential challenges still exist and are the focus of on-  
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44 going work, including: (i) the control over Fe contaminants and their role in dictating the reaction  
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46 regime, (ii) understanding the effect of operating pressure, (iii) relating macroscopic surface properties  
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48 (wettability) to the observed microstructure, and (v) enhancing functional moieties through additives  
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50 (H<sub>2</sub>O<sub>2</sub>, NH<sub>3</sub>, etc.). Overall, the development of syngas PICVD shows promise for application in the  
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52 coatings industry.  
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## ■ ASSOCIATED CONTENT

### Supporting Information

The Supporting Information is available free of charge on the ACS Publications website at DOI:

The profilometry thickness profiles, XPS survey results vs. residence time and treatment duration, as well as TOF-SIMS results due to the ratio variation and EDS micrographs of treated sample.

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### Author Contributions

The manuscript was written through contributions of all authors: Donya Farhanian. Author-One, Prof. Gregory De Crescenzo. Author-Two, Prof. Jason R. Tavares. Corresponding Author\*. All authors have given approval to the final version of the manuscript.

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## Notes

The authors declare no competing financial interest.

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