Fabrication of Nanoscale Oxide Textured Surfaces on Polymers

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Abstract: Nanoscale textured surfaces play an important role in creating antibacterial surfaces, broadband anti-reflective properties, and super-hydrophobicity in many technological systems. Creating nanoscale oxide textures on polymer substrates for applications such as ophthalmic lenses and flexible electronics imposes additional challenges over conventional nanofabrication processes since polymer substrates are typically temperature-sensitive and chemically reactive. In this study, we investigated and developed nanofabrication methodologies to create highly ordered oxide nanostructures on top of polymer substrates without any lithography process. We developed suitable block copolymer self-assembly, sequential infiltration synthesis (SIS), and reactive ion etching (RIE) for processes on polymer substrates. Importantly, to prevent damage to the temperature-sensitive polymer and polymer/oxide interface, we developed the process to be entirely performed at low temperatures, that is, below 80 °C, using a combination of UV crosslinking, solvent annealing, and modified SIS and RIE processes. In addition, we developed a substrate passivation process to overcome reactivity between the polymer substrate and the SIS precursors as well as a high precision RIE process to enable deep etching into the thermally insulated substrate. These methodologies widen the possibilities of nanofabrication on polymers.

Keywords: block copolymer; solvent vapor annealing; sequential infiltration synthesis; nanotexture; surface passivation

1. Introduction

Oxide nanoscale structures play a central role in optical, electrical, and biomedical nanotechnological devices and sensors due to their tunable optoelectronic properties, high surface-to-volume ratio, and good stability [1,2]. In particular, surface texturing with high-aspect-ratio oxide nanostructures significantly enhances surface-based properties in sensors [3,4], antibacterial surfaces [5,6], hydrophobic surfaces [7], and optical lenses [8,9]. Oxide surface texturing has been demonstrated by several nanostructure formations and patterning techniques, including photolithography [10], nanoimprint lithography [11,12], colloid assembly [13,14], and mesoporous silica layers [15]. A high-aspect-ratio oxide nanostructure can enhance properties such as light transmission and surface hydrophobicity as well as act as an antireflective layer due to the low effective refractive index of the metal oxide/air sub-wavelength nanostructure [11]. If the nanostructure is designed to create an effective refractive index gradient, it can exhibit wide angular broadband anti-reflective behavior, such as moth eyes exhibit in nature and as has been mimicked by several man-made approaches [12,16–23].

Recent work has demonstrated oxide nanoscale structures on silicon substrates [23]. However, surface texturing of polymer substrates imposes new challenges compared to inorganic substrates due to additional constraints on the processing temperatures as well as polymer reactivity. With the growing demand for lightweight and cost-effective devices, there is a need for new nanofabrication processes for polymeric substrates [24].
In the past few decades, block copolymer (BCP) self-assembly has emerged as an excellent nanofabrication technique for simple and scalable nanoscale patterning [25–29]. BCP self-assembly yields highly ordered and uniform nanostructures with periodicity ranging between 10 and 100 nm [29]. BCP patterning is typically performed by casting a thin BCP layer on the substrate and annealing it to induce phase separation [28]. The BCP pattern can then be transferred to the underlying inorganic layer, resulting in a textured surface that can enhance hydrophobicity or anti-reflective properties [30,31]. However, pattern transfer of the BCP layer into high aspect ratio structures can impose a challenge due to the low etch contrast between typical BCP organic-organic domains.

Recently, sequential infiltration synthesis (SIS) has emerged as a novel technique for selectively growing metal oxides within the polar domain of block copolymers, enabling a high etch contrast between the blocks and efficient pattern transfer to form high-aspect-ratio structures [23,32–36]. SIS is based on atomic layer deposition (ALD) chemistry, where high precursor partial pressures and long exposure times result in precursors’ sorption and diffusion within the polymer [37]. The favorable interaction between the organometallic precursors and the polar domains of BCP can lead to the selective growth of metal oxides such as AlOₓ [35], ZnO [38], TiOₓ [39], and SnOₓ [40] in only one domain of the block copolymers [25,32,33,36]. Following the SIS process, the BCP film can be removed, resulting in a metal oxide nanostructure templated by the BCP morphology with high fidelity [25]. SIS has been successfully implemented into nanofabrication processes, such as semiconductor patterning [25,38], porous membrane fabrication [32,41,42], and anti-reflective layers [23,43].

Until now, SIS-based metal oxide nanostructures were fabricated onto silicon wafers or glass substrates, which are chemically inert and can withstand high-temperature processes. On the other hand, polymer substrates, such as diglycol, thiosulfonate, polycarbonate, and others, are not only sensitive to temperature but also can interact with the organometallic precursors used in the SIS process. Combining BCP self-assembly and SIS could offer a simple, scalable, and efficient way to fabricate high-aspect-ratio nanoscale oxide textures over a polymeric substrate.

In this study, we aim to expand SIS processes to polymer substrates and to enable nanoscale texturing of oxides onto polymers. Working with polymeric substrates poses a two-front challenge. First, there are competing temperature needs for optimal BCP assembly, SIS, and etching processing vs. the polymer substrate Tg and the polymer/oxide layer thermal expansion properties. Second, unlike silicon or glass inert substrates, polymers are potentially chemically reactive through outgassing or terminal edge moieties. The approach and methodologies developed in this work mitigate these challenges by using a combination of polymer substrate passivation processes to prevent undesired interactions between SIS precursors and the polymer substrates, room-temperature BCP assembly processes, and composite SIS patterning that overcomes pattern transfer challenges.

2. Experimental Details

2.1. Materials

BCP: polystyrene-block-poly(methyl methacrylate) (PS-b-PMMA) (Mn: 140k-b-65k, PDI = 1.16), (Mn:46.1k-b-21k, PDI = 1.09) (Mn: 68k-b-33.5k, PDI = 1.08), metalorganic precursors (trimethyl aluminum (TMA), diethyl zinc (DEZ)) and solvents (toluene, tetrahydrofuran, chloroform, etc.) were purchased from Polymer Source, STREM chemicals, and Fisher Scientific, respectively, and used as received.

2.2. Substrate Preparation

Discs of diglycol (MR-8™), thiosulfonate (CR-39™), and polycarbonate with a diameter and thickness of 20 mm and 2 mm, respectively, low resistivity (<0.005 Ω.cm) 4″ Si (100) substrates, and glass slides were coated with 350 nm of SiO₂ using chemical vapor deposition (MC380X box coater, Satisloh, Germantown, WI, USA) at Shamir Optics, Israel.
SiO$_2$/Si and SiO$_2$/glass substrates were used as the reference standards. The surfaces of all the SiO$_2$-coated substrates were cleaned in O$_2$ plasma for 5 min prior to further processing.

2.3. BCP Templates

A thin layer of poly(styrene-$r$-methyl methacrylate-$r$-glycidyl methacrylate) (P(S-$r$-MMA-$r$-GMA)) containing ~4 mol% of glycidyl methacrylate with styrene mole fractions of 77% (PG-4 76%PS) was used for tuning the surface interactions of PS-$b$-PMMA with SiO$_2$ substrate by spin-coating 0.3 wt. % of it in toluene. P(S-$r$-MMA-$r$-GMA) was synthesized by reversible addition fragmentation chain transfer polymerization with Azobisisobutyronitrile (AIBN), which is a thermal and photoinitiator. UV crosslinking was performed using Spectrolinker™ UV Crosslinkers (XL-1000 model) at a wavelength of 254 nm in N$_2$ atmosphere. A layer of PMMA cylinder-forming PS-$b$-PMMA was fabricated by spin-coating 2 wt. % solution of PS-$b$-PMMA in toluene. The BCP layer was solvent vapor annealed (SVA) by exposing it to saturated vapors of tetrahydrofuran in a closed chamber containing ~100 mL of solvent at room temperature for durations ranging from 5 to 30 min. After exposure, the SVA was immediately quenched by taking it out of the annealing chamber, and the film was kept inside a fume hood for ~4 h to complete the evaporation of trapped THF vapors.

2.4. Sequential Infiltration Synthesis

AlO$_x$ and ZnO SIS were performed in a Savannah S100 ALD system (Veeco, New York, NY, USA). BCP films were loaded into the ALD reactor at 80 °C. To achieve thermal equilibrium and remove excess moisture, the samples were subjected to 20 sccm of N$_2$ flow at 0.3 Torr for at least 30 min. The SIS process was carried out as follows: upon TMA pulse, the chamber was closed in a static mode for 300 s, enabling precursor diffusion into the polymer film (exposure). The exposure step was followed by an N$_2$ purge step for 350 s to remove excess reactants. A similar exposure/purge process was used for water. The full AlO$_x$ SIS cycle, TMA/purge/H$_2$O/purge = 300s/350s/300s/350s, was repeated several times. In a similar approach, AlO$_x$/ZnO composite cylinders were fabricated by 2 cycles of AlO$_x$ SIS, followed by 6 cycles of ZnO SIS (DEZ/purge/H$_2$O/purge) using the same sequence as AlO$_x$ SIS. After completion of the SIS process, the BCP template was removed by O$_2$ plasma etching (50 W, 0.4 mbar for 10–20 min).

2.5. Passivation Process

The SiO$_2$/polymer substrate was positioned over three adjacently placed Teflon discs to avoid the direct thermal contact of the polymer substrate with the ALD chamber. The passivation was performed by 100 cycles of Al$_2$O$_3$ ALD cycles using the following sequence: TMA exposure/purge/H$_2$O exposure/purge for 10s/10s/10s/10s, respectively, at 80 °C.

2.6. Reactive Ion Etching

Vertically aligned SiO$_2$ nanorods were fabricated by inductively coupled plasma-reactive ion etching using a Plasma-Therm (Oxford Instruments, Model: 790) with a CHF$_3$ (5 sccm, 12 mTorr, 97 W rf power) and CF$_4$ + O$_2$ (36 sccm of CF$_4$, 2 sccm of O$_2$, 40 mTorr, 175 W rf power) mixture for different durations at room temperature.

2.7. Characterizations

The morphology of the sample was studied by a high-resolution scanning electron microscope (Ultra-Plus HRSEM, Zeiss, Oberkochen, Germany) at 1 kV. SEM cross-sectional samples on a SiO$_2$/Si substrate were prepared by cleaving the substrate along a crystallographic direction. SiO$_2$/polymer lenses were coated with 3 nm of Pt using BAF (Leica BAF 060) and cleaved after dipping them inside liquid nitrogen. Cross-section images were recorded by tilting the sample by 70°.
3. Results and Discussion

3.1. Process Overview

The fabrication methodology and processes for the nanoscale texturing of oxide layers on polymer substrates are schematically illustrated in Figure 1. The process was designed to enable the desired interaction between the BCP layer and SIS precursors while hindering interactions between these precursors and the polymer substrate. MR-8™ was chosen as the polymer substrate model system due to its wide use in ophthalmic lenses. Other polymer substrates that were examined, CR-39™ and polycarbonate, gave similar results, but for simplicity, we used MR-8™ as a model system. (1) First, we performed an ALD passivation process on the SiO₂/MR-8™ polymer substrate to passivate the polymer substrate and prevent undesired interactions in the SIS stage. (2) We self-assembled polystyrene-block-poly(methyl methacrylate) (PS-b-PMMA) thin films on the passivated SiO₂/MR-8™ polymer substrates using a room-temperature process, which includes random copolymer mat UV crosslinking to neutralize the interaction between the SiO₂ surface and the BCP layer [44], followed by BCP solvent vapor annealing (SVA) to promote BCP self-assembly. (3) We selectively grew AlOₓ and ZnO in the cylindrical PMMA domains with the AlOₓ/ZnO SIS process. Following SIS, we removed the polymer template with oxygen plasma to obtain AlOₓ/ZnO composite nanorods that can be used as a hard mask for deep reactive ion etching (RIE). (4) We performed an RIE process to break through the Al₂O₃ passivation layer and transfer the AlOₓ/ZnO nanorod hard masks into the underlying SiO₂ layer. The main steps of this process on non-sensitive substrates, SiO₂/Si and SiO₂/glass, are presented in Figures S1 and S2.

![Figure 1](image_url)

Figure 1. Schematic illustration of the developed processes. The process starts with SiO₂/MR-8™ polymeric substrate passivation using Al₂O₃ ALD, followed by room-temperature BCP assembly—random copolymer mat layer crosslinking using UV exposure and SVA of cylinder-forming PS-b-PMMA to promote self-assembly. Growth of AlOₓ and ZnO within the BCP layer is performed using TMA/H₂O and DEZ/H₂O SIS processes, followed by polymer template removal. Finally, the pattern is transferred to the SiO₂ layer with RIE.

3.2. Room-Temperature BCP Assembly

3.2.1. Random Copolymer Mat Crosslinking Using UV Treatment

Oxide surfaces, such as SiO₂ and Al₂O₃, are favorably wetted by the polar domains of BCP [45]. To induce vertical-oriented assembly, random copolymer mats are commonly used to create a surface that is wetted by both blocks [46]. The required polymer mat
layer crosslinking is typically performed with thermal annealing at elevated temperatures (190 °C–250 °C). For example, Figure S3 shows the highly ordered perpendicular orientated self-assembly of PS-b-PMMA 46.1k-b-21k on a thermally (230 °C) crosslinked PG-4 76% mat. To create a room-temperature BCP assembly process, we first probed room-temperature polymer mat crosslinking using UV exposure (λ = 254 nm) in an inert environment. PG-4 mats are known to be UV crosslinkable due to the presence of epoxy side groups [47].

The mat layer quality of SiO₂/Si using the self-assembly of an ~80 nm thick PS-b-PMMA 140k-b-65k film treated with 15 min of tetrahydrofuran (THF) SVA was examined. Figure 2 presents the resulting PS-b-PMMA morphology as a function of UV treatment. The dark domains correspond to PMMA, while the bright domains correspond to PS. When the random copolymer mat, PG-4 76%, was not treated with UV, the resulting PS-b-PMMA assembly lacked long-range order (Figure 2a). Applying a short, 20 sec UV treatment of a 120 mJ/cm² dose led to the BCP self-assembly of perpendicular-orientated PMMA cylinders with 42 ± 4 nm diameters and an average grain size of 202 ± 6 nm (Figure 2b). Extensive UV treatment (1 h of exposure with a dose of 27,100 mJ/cm²; Figure 2c), resulted in perpendicular assembly but with a lower degree of long-range order.

![Figure 2](image-url)

**Figure 2.** Random copolymer mat crosslinking on SiO₂/MR-8 polymer substrates. SEM images of PS-b-PMMA self-assembled film (15 min THF SVA) on a PG-4 76% mat layer: (a) without UV treatment, (b) with 20 secs of UV exposure of a 120 mJ/Cm² dose, and (c) with 1 h of UV exposure of a 27,108 mJ/Cm² dose.

### 3.2.2. BCP Solvent Annealing

In order to establish a room-temperature BCP assembly process, we optimized the SVA process for PS-b-PMMA 140k-b-65k. Vapor infiltration into the polymer film leads to higher chain mobility due to polymer plasticization. It lowers the glass transition temperatures ($T_g$) and the effective Flory–Huggins parameter of the blocks ($\chi_{eff} < \chi$), as well as creating interface interaction screening, resulting in room-temperature BCP assembly [46,48–51]. Figure 3 shows the surface morphology of ~80 nm thick PS-b-PMMA 140k-b-65k after 15 min of SVA using chloroform, acetone, and THF. While chloroform and acetone SVA resulted in disordered assembly and parallel cylinder assembly, respectively (Figure 3a,b), THF SVA resulted in perpendicular PMMA cylinder assembly with 71 ± 2 nm periodicity and a high degree of order. THF is considered a good solvent for both blocks [49,52], swelling both blocks and creating similar effective interfacial energies that lead to perpendicular assembly. Interestingly, when the THF SVA is performed for longer or shorter times than 15 min, the assembly becomes disordered (Figure S4). This “sweet spot” time interval indicates that short SVA might be insufficient to swell both blocks, while long SVA could lead to non-uniform surface interactions due to the differences in polymer block/solvent interactions ($\chi_{PS-THF} = 0.32$ and $\chi_{PMMA-THF} = 0.8$) [53].
3.2.3. SIS on BCP/SiO$_2$/MR-8 Polymer

In order to develop oxide nano-textured surfaces on polymeric substrates, we examined the SIS process on BCP/SiO$_2$/MR-8$^{\text{TM}}$ polymer stacks. A 500 nm SiO$_2$ layer was deposited on an MR-8$^{\text{TM}}$ substrate with physical vapor deposition. The prepared substrates (SiO$_2$/MR-8$^{\text{TM}}$) were then coated with PG4 76% mat and PS-\textit{b}-PMMA 140k-\textit{b}-65k layers, as described in Section 2. BCP assembly on the top of SiO$_2$/MR-8$^{\text{TM}}$ substrates displayed hexagonally ordered perpendicular PMMA cylinders in a PS matrix (Figure 4a). The thermal stability of the BCP layer was confirmed by placing BCP/SiO$_2$/MR-8 samples at 80 °C for 3 h in the ALD chamber without any reaction (Figure 4b). The retention of the BCP self-assembled structure and the absence of SiO$_2$/MR-8 polymer degradation during thermal treatment make the PS-\textit{b}-PMMA/SiO$_2$/MR-8 stack compatible for the SIS process. However, when we performed between one and eight cycles of AlO$_x$ SIS on the PS-\textit{b}-PMMA/SiO$_2$/MR-8$^{\text{TM}}$ stack, no traces of the self-assembled BCP layer nor a hybrid AlO$_x$-BCP layer (Figure 4c) could be identified on the surface.

![Figure 3](image1.png)

**Figure 3.** SVA of PS-\textit{b}-PMMA 140k-\textit{b}-65k on PG4 76%/SiO$_2$/Si substrates. SEM images after 15 min of BCP SVA using: (a) chloroform, (b) acetone, and (c) THF.

![Figure 4](image2.png)

**Figure 4.** SEM images of PS-\textit{b}-PMMA over SiO$_2$/MR-8 substrate: (a) self-assembled, (b) after 3 h of heat treatment at 80 °C, and (c) after eight cycles of AlO$_x$ SIS. (d) SEM image of AlO$_x$ nanocylinders on SiO$_2$/Si substrate fabricated using the same process as (c).
In comparison, when the same SIS process on PS-b-PMMA/SiO$_2$/Si and PS-b-PMMA/SiO$_2$/glass stacks was performed without the presence of the MR-8$^\text{TM}$ polymer substrate, selective growth of AlO$_x$ within the PMMA cylinders could easily be observed. Subsequent etching of the BCP film on SiO$_2$/Si substrate with O$_2$ plasma resulted in an ordered AlO$_x$ nanocylinders array, templated by the PMMA domains, with 42 ± 4 nm diameter AlO$_x$ cylinders (Figure 4d and Figure S1). Moreover, when these nanocylinder arrays were used as hard masks for RIE of the underlying SiO$_2$ layer on Si or glass substrates, textured SiO$_2$ surfaces with tunable-aspect-ratio SiO$_2$ nanorods were obtained (Figures S2 and S5). The vertical AlO$_x$ nanocylinders and the resulting SiO$_2$ nanorods indicate that the PMMA is vertically assembled in the BCP layer and that the SIS process efficiently grew AlO$_x$ in the PMMA domains.

While SIS on BCP/SiO$_2$/Si and on BCP/SiO$_2$/glass substrates resulted in selective growth of AlO$_x$ within the PMMA domains, SIS processes on the BCP/SiO$_2$/MR-8 polymer substrate did not yield any ordered structure on the top surface. Other polymer substrates that were examined, CR-39$^\text{TM}$, and polycarbonate, also exhibited the same behavior. We attribute this phenomenon to interactions of the organometallic precursor, TMA, with reactive moieties in the polymer substrate. For example, MR-8$^\text{TM}$ polymer has an abundance of reactive groups, such as hydroxyl and cyano groups, which can react with TMA and release byproduct molecules. These molecules can partly be deposited on the top BCP layer, obscuring the self-assembled layer and hindering SIS growth within the PMMA domains. SEM characterization of the post-SIS surface and scratching tests (data not shown) indicated the presence of additional organic materials at the top surface.

To overcome this limitation and enable SIS-based patterning on polymer substrates, we performed a surface passivation process. The SiO$_2$/MR-8$^\text{TM}$ substrate, prior to random copolymer mat and BCP deposition, was exposed to 100 cycles of Al$_2$O$_3$ ALD (80 °C, TMA/purge/H$_2$O/purge = 10s/10s/10s/10s), resulting in Al$_2$O$_3$ deposition on the SiO$_2$/MR-8$^\text{TM}$ surface (see illustration in Figure 1). Figure 5 displays SEM imaging of PS-b-PMMA 140k-b-65k self-assembly and SIS processes on passivated SiO$_2$/MR-8$^\text{TM}$ substrates. Following passivation, we self-assembled PS-b-PMMA 140k-b-65k on the crosslinked random copolymer mat, as described in Section 2, resulting in highly ordered perpendicular cylinder morphology (Figure 5a). Eight cycles of AlO$_x$ SIS on the self-assembled layer resulted in selective growth of AlO$_x$ in the PMMA domains, as can be seen from the reverse in contrast to the BCP domains (Figure 5b). We also examined the AlO$_x$/ZnO SIS process, where two cycles of AlO$_x$ SIS were performed to promote ZnO growth in the PMMA domains using six cycles of ZnO SIS. When we removed the BCP layer using oxygen plasma, we obtained highly ordered AlO$_x$ and AlO$_x$/ZnO nanorod arrays on the passivated SiO$_2$/MR-8$^\text{TM}$ substrates (Figures 5c and 5d, respectively). A cross-sectional SEM image of AlO$_x$/ZnO nanorod arrays is presented as an inset of Figure 5d. The nanorods are estimated to be 25-35 nm in height and 35-40 nm in width. These BCP-templated metal oxide nanorods can now be used as a hard mask for pattern transfer to the underlying SiO$_2$ layer.

3.2.4. Pattern Transfer into SiO$_2$ Underlying Layer

In order to transfer the BCP-templated metal oxide structure into the SiO$_2$ layer, we performed CHF$_3$ RIE, which is known for its anisotropic etch of SiO$_2$ and good Al$_2$O$_3$/SiO$_2$ selectivity [54,55]. However, due to the SiO$_2$/MR-8$^\text{TM}$ passivation, there was a nanometric Al$_2$O$_3$ layer between the metal oxide nanorods that needed to be etched to reach the SiO$_2$ layer. When we performed CHF$_3$ RIE on AlO$_x$ nanorods/Al$_2$O$_3$/SiO$_2$/MR-8$^\text{TM}$, the etch budget in the AlO$_x$ nanorods was insufficient to enable Al$_2$O$_3$ passivation breakthrough and maintain an efficient hard mask for SiO$_2$ etching. On the other hand, when we performed CHF$_3$ RIE on AlO$_x$/ZnO composite nanorods/Al$_2$O$_3$/SiO$_2$/MR-8$^\text{TM}$, the higher etch resistance of the AlO$_x$/ZnO composite nanorods enabled the nanorod to maintain its structure through the Al$_2$O$_3$ passivation breakthrough etch, leading to efficient pattern transfer.
3.2.4. Pattern Transfer into SiO₂ Underlying Layer

In order to transfer the BCP-templated metal oxide structure into the SiO₂ layer, we performed CHF₃ RIE, which is known for its anisotropic etch of SiO₂ and good Al₂O₃/SiO₂ selectivity [54,55]. However, due to the SiO₂/MR-8TM passivation, there was a nanometric Al₂O₃ layer between the metal oxide nanorods that needed to be etched to reach the SiO₂ layer. When we performed CHF₃ RIE on AlOₓ nanorods/Al₂O₃/SiO₂/MR-8TM, the etch budget in the AlOₓ nanorods was insufficient to enable Al₂O₃ passivation breakthrough and maintain an efficient hard mask for SiO₂ etching. On the other hand, when we performed CHF₃ RIE on AlOₓ/ZnO composite nanorods/Al₂O₃/SiO₂/MR-8TM, the higher etch resistance of the AlOₓ/ZnO composite nanorods enabled the nanorod to maintain its structure through the Al₂O₃ passivation breakthrough etch, leading to efficient pattern transfer.

We further developed the RIE process to account for the low thermal conductivity of the polymer substrate. Figure 6a,b present SEM images of the SiO₂ surface formed after exposing the BCP-templated AlOₓ/ZnO nanorods to two modes of CHF₃ RIE: continuous RIE for 3 min (Figure 6a), and loop RIE for 14 min, which is built from loops of 30 s of RIE and a 1 min purge (Figure 6b). The continuous 3 min etching resulted in shallow SiO₂ texturing, as implied by the low contrast of the structure. The low thermal conductivity of the polymer substrate surface leads to substrate heating, enhancing isotropic etching, which results in nanocylinder mask etching. The loop RIE process, on the other hand, provides sufficient time between the plasma periods to dissipate the heat generated on the surface. Figure 6b clearly shows the intactness of the hexagonally ordered cylindrical array formed after the loop RIE. The corresponding cross-sectional image (Figure 6c) shows ~80 nm high and ~40 nm wide (~2:1 aspect ratio) oval-shaped nanorods, fabricated in the SiO₂ layer by room-temperature BCP self-assembly, AlOₓ/ZnO SIS, and loop RIE. Further optimization of RIE chemistry for deep and/or conical-shaped high-aspect-ratio nanostructures is viable using the methodology presented in Figure S5.
Further optimization of RIE chemistry for deep and/or conical-shaped high-aspect-ratio nanostructures is viable using the methodology presented in Figure S5.

Figure 6. SEM images of AlO$_x$/ZnO nanocylinders over an SiO$_2$/MR-8 polymer substrate after (a) continuous 3 min of CHF$_3$ RIE, (b) 14 min of loop CHF$_3$ RIE. (c) Cross-sectional view of a textured SiO$_2$ surface generated with the process shown in (b). Scale bar is 100 nm in all images.

4. Conclusions

In this research, a new methodology for creating highly ordered and periodic, nanoscale oxide texture on top of polymeric substrates was developed. These nanoscale textures can be used to enhance various properties, such as anti-reflectiveness and superhydrophobicity. The texturing is based on transferring the pattern of BCP onto metal oxide nanorods via SIS and onto the oxide layer via RIE. Our approach provided solutions to the challenges that nanoscale oxide texturing of polymer substrates poses. We have developed a series of suitable low-temperature (<80 °C) processes geared towards temperature-sensitive substrates. The polymer substrates were 3D-passivated with an ALD-processed Al$_2$O$_3$ barrier layer. On top of preventing undesired interactions between the SIS organometallic precursors and the polymer substrates, the passivation layer can be used as a general basis layer, reducing the need to tune the process for each polymer substrate chemistry. In addition, in order to enable a differential etch rate in the RIE step, an AlO$_x$/ZnO composite was generated in the PMMA cylinder phase. This combined barrier-differential etch approach expands the standard SIS toolbox to enable new pathways for nanofabrication on a wide array of lighter-weight polymers over traditional SIS-processed glass and silicon substrates.

Supplementary Materials: The following are available online at https://www.mdpi.com/article/10.3390/polym13132209/s1. Figure S1: SEM images of AlO$_x$ cylinders pattern formed on (a) SiO$_2$/Si substrate and (b) SiO$_2$/glass substrate; Figure S2: Top (a,c) and cross-sectional (b,d) SEM images of SiO$_2$ nanostructures formed by etching the AlO$_x$ nanocylinder pattern on SiO$_2$/Si substrate.
(a,b) and SiO2/glass substrate (c,d) using 10 min CHF3 RIE; Figure S3: Top view SEM image of (a) PS-b-PMMA (46.1k-b-21k) block/copolymer after thermal annealing at 230 °C; Figure S4: SEM image of PS-b-PMMA (140k-b-65k) over SiO2/Si substrate vapor annealed for 10 and 20 min in tetrahydrofuran; Figure S5: SEM image of SiO2 nanorods array fabricated by different RIE chemistry: anisotropic etching of SiO2 by CHF3 for (c) 5 min, (b) 10 min; isotropic etching of SiO2 by CF4 + O2 (c) 3 min, (d) 4 min, and combination of CHF3 and CF4 + O2 (e) 5 min and 1 min, (d) 10 min and 0.5 min, respectively.

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