Directed Self-Assembly of Block Copolymers for High Breakdown Strength Polymer Film Capacitors

Saumil P. Samant, † Christopher A. Grabowski, ‡ Kim Kisslinger, § Kevin G. Yager, # Guangcui Yuan, § Sushil K. Satija, ‡ Michael F. Durstock, ‡ Dharmaraj Raghavan, †∥ and Alamgir Karim*†∥

†Department of Polymer Engineering, University of Akron, Akron, Ohio 44325, United States
‡Air Force Research Laboratory, Wright Patterson Air Force Base, Dayton, Ohio 45433, United States
§Center for Functional Nanomaterials, Brookhaven National Laboratory, Upton, New York 11973, United States
∥Center for Neutron Research, National Institute of Standards and Technology (NIST), Gaithersburg, Maryland 20899, United States
#Department of Chemistry, Howard University, Washington, D.C. 20059, United States

ABSTRACT: Emerging needs for fast charge/discharge yet high-power, lightweight, and flexible electronics requires the use of polymer-film-based solid-state capacitors with high energy densities. Fast charge/discharge rates of film capacitors on the order of microseconds are not achievable with slower charging conventional batteries, supercapacitors and related hybrid technologies. However, the current energy densities of polymer film capacitors fall short of rising demand, and could be significantly enhanced by increasing the breakdown strength ($E_{BD}$) and dielectric permittivity ($\varepsilon_r$) of the polymer films. Co-extruded two-homopolymer component multilayered films have demonstrated much promise in this regard showing higher $E_{BD}$ over that of component polymers. Multilayered films can also help incorporate functional features besides energy storage, such as enhanced optical, mechanical, thermal and barrier properties. In this work, we report accomplishing multilayer, multicomponent block copolymer dielectric films (BCDF) with soft-shear driven highly oriented self-assembled lamellar diblock copolymers (BCP) as a novel application of this important class of self-assembling materials. Results of a model PS-b-PMMA system show ~50% enhancement in $E_{BD}$ of self-assembled multilayer lamellar BCP films compared to unordered as-cast films, indicating that the breakdown is highly sensitive to the nanostructure of the BCP. The enhancement in $E_{BD}$ is attributed to the “barrier effect”, where the multiple interfaces between the lamellae block components act as barriers to the dielectric breakdown through the film. The increase in $E_{BD}$ corresponds to more than doubling the energy storage capacity using a straightforward directed self-assembly strategy. This approach opens a new nanomaterial paradigm for designing high energy density dielectric materials.

KEYWORDS: block copolymer, directed self-assembly, cold zone annealing—soft shear, lamellae, breakdown strength, barrier effect, dielectric, capacitor

INTRODUCTION

Energy storage is a fundamental issue driving the development of new materials and their associated processing and fabrication into devices remains a major challenge in energy research today. Among the various energy storage technologies are batteries, fuel cells, supercapacitors and solid-state film capacitors. While batteries and fuel cells have high energy-storage capacities, they are bulky and not suitable for high-power applications because of very slow charge–discharge characteristics. Supercapacitors have moderate energy densities but low operating voltages and high leakage currents limit their application in pulsed power systems. In contrast, electrostatic or solid-state film capacitors possess the advantage of high power density due to relatively fast (ms–μs) energy storage and discharge rates. However, the energy storage density of these capacitors is limited by the dielectric properties of the materials that separate the oppositely charged electrodes, falling significantly short of rising demand in advanced applications.  

There is a critical need to not only develop new dielectric materials, but also to control their morphologies at the nanoscale in ways that can significantly increase their energy storage capacity thereby reducing the size, weight, and cost of electronic devices and power systems.  

The maximum energy density ($U$) that can be stored within a capacitor is governed by the product of dielectric permittivity ($\varepsilon_r$) and breakdown strength ($E_{BD}$) of the dielectric material.
where $\varepsilon_0$ (8.85 × 10⁻¹² F/m) is the permittivity of free space.

Polymer films are widely used as capacitor dielectric materials because of their desirable properties, such as high $E_{BD}$, low loss, flexibility, solvent processability, and graceful failure as opposed to the catastrophic failure of ceramic insulators.³⁻⁵ Commercially available capacitors most frequently employ semicrystalline biaxially oriented polypropylene (BOPP) films, which features a theoretical maximum energy storage of ~4 J/cm³, while fully packaged BOPP capacitors exhibit storage in the 1–2 J/cm³ range. This energy density falls short of current demands, thus requiring enhancement in $\varepsilon_r$ and $E_{BD}$. Although significant efforts have been made to increase the $\varepsilon_r$ of polymer film capacitors using nanocomposites or nanodielectrics,¹⁶⁻¹⁰ enhancement of $E_{BD}$ without the use of fillers has not received similar attention. As seen from eq 1, the energy density has a quadratic dependence on $E_{BD}$ and thus enhancement of $E_{BD}$ holds great potential for improving the current energy densities of polymer film capacitors.

In polymers, dielectric breakdown is traditionally assumed to occur by an electron avalanche mechanism via formation and propagation of electrical trees that resemble fractal conducting channels within the material.¹¹,¹² These electrical trees are initiated by structural defects in the material and propagate between the electrodes leading to breakdown. If the propagation of these electrical trees is impeded by a barrier, the breakdown process can be forestalled, thus enhancing the overall dielectric strength. This “barrier effect” is a well-known concept applied in high-voltage engineering to increase the breakdown voltage.¹³⁻¹⁵ It is hypothesized that the presence of a barrier, that is, a material of different permittivity or conductivity than the matrix, leads to formation of space charge near the interface on application of an external field, which causes redistribution of the field along the barrier-matrix interface. Vogelsang and co-workers have predicted using a deterministic model that the presence of a barrier (mica) significantly impedes the breakdown process in polymers (epoxy) as the electrical trees are forced along an extended tortuous pathway along the barrier-matrix interfaces.¹⁶,¹⁷ Pioneering work by Wolak and co-workers demonstrated the barrier effect in multilayered polymer films obtained using forced assembly multilayer coextrusion.²¹ By fabricating multilayered PC-coP 2VDF composite films, a marked increase in the $E_{BD}$ of multilayered films was observed over the expected linear weighted-average of individual polymer control films. Zhu and co-workers have further explored the multilayered coextrusion, demonstrating that layered films have increased dielectric lifetimes as compared to blend films due to the barrier structure.¹⁹ Layered/interwoven architectures can also be generated through the dispersion and alignment of high aspect ratio, nanometer thick sheets of montmorillonite within a polymer matrix. The breakdown strength of such nanolaminates was found to increase by 30–50% relative to the unfilled polymer film.²⁰,²¹

An undetermined issue with the coextruded multilayers is that of layer homogeneity due to spatial thickness variations, large micron-sized defects and layer delamination because of poor adhesion. In contrast to macroscopic processing, self-assemblying polymer systems generate much more uniform and well-controlled nanostructured layers. In principle, self-assembled multilayered structures can be obtained using lamellae-forming symmetric block copolymer (BCP) films with the lamellar microdomains of the BCP aligned parallel to the substrate over large areas. To this end, we devised a novel strategy to create a polymer capacitor film having a highly ordered multilayered structure, with relatively defect-free, sharp nanoscale interfaces using directed self-assembly (DSA) methods applied to lamellae-forming BCP thick (>1 μm) films. BCP films offer extraordinary tunability of dielectric contrast, orientation, layer thickness and interfacial width with self-assembly processes, from the intrinsic nanoscale molecular level to macroscopic thickness for device-level compatibility. While BCP self-assembly and resulting BCP-templated films have been extensively studied for many diverse electronic applications,²² the dielectric performance of self-assembled BCPs has not been explored and their use as the dielectric media in solid-state film capacitors has not been reported to date. In this paper, we report the very first application of self-assembled BCPs as potential solid-state capacitor materials and demonstrate that the dielectric strength ($E_{BD}$) is highly sensitive to the BCP nanostructure. These findings have significant ramifications in optimizing the breakdown strength of block copolymer structures and can guide future capacitor device development.

### RESULTS AND DISCUSSION

In thin (<500 nm) films, the parallel (to substrate) lamellar morphology can be readily achieved with simple uniform oven annealing in select low molecular weight BCP systems on preferential substrate surfaces.²³⁻²⁵ However, we have determined that highly ordered parallel lamellae could not be obtained with oven annealing for micron thick, high molecular weight BCP films on nonmodified conductive substrates desired for capacitor studies (shown later). Instead, this can be potentially achieved using directed self-assembly (DSA) techniques with high degree of directive forces applied to the BCP films. Most DSA methods employing solvent vapor,²⁶ shear,²⁷,²⁸ magnetic²⁹ or electric fields³⁰,³¹ are batch processes while others require topographical patterning³²,³³ or solvent immersion, such as the recently developed direct immersion annealing (DIA) process.³⁴ For large-area fabrication of capacitor films, it is useful to develop a DSA method that does not require substrate modification and is also compatible with roll-to-roll (R2R) processing. To this end, we employ a unique cold zone annealing-soft shear (CZA-SS) method that was previously developed for unidirectional alignment of cylindrical block copolymer films.³⁵,³⁶ Herein, we demonstrate its novel application to fabricate highly ordered micron thick lamellar BCP dielectric films (BCDFs) aligned parallel to conductive Aluminum substrate for solid-state capacitor applications.

**BCP Film Processing.** The effect of a self-assembled multilayered lamellar structure on the breakdown properties was studied using the model BCP system polystyrene-b-poly(methyl methacrylate) (PS-b-PMMa), where the individual blocks have disparate dielectric breakdown values (PS ≈ 250 V/μm, PMMA ≈ 500 V/μm), but relatively close dielectric permittivity at 1 kHz [PS ($\varepsilon_r$ ≈ 2.6), PMMA ($\varepsilon_r$ ≈ 3.2)]. To elucidate the generality of the concept, we also studied another BCP system, polystyrene-b-poly(2-vinylpyridine) (PS-b-P2VP), wherein the blocks have closer dielectric strength (PS ≈ 250 V/μm, P2VP ≈ 380 V/μm), but much higher dielectric contrast [P2VP ($\varepsilon_r$ ≈ 5)]. Approximately 1 μm thick BCP films of lamellae-forming PS-b-PMMa (19.5-b-38.1 kg mol⁻¹ and 33-b-
33 kg mol$^{-1}$) and PS-b-P2VP (25-25 kg mol$^{-1}$) were cast onto smooth aluminum-coated quartz substrates, which act as the bottom electrode, using a flowcoating technique (Figure S1). The films were then subjected to the CZA-SS process of directed self-assembly as detailed in a previous publication$^{35}$ and Supporting Information. Briefly, a cross-linked polydimethylsiloxane (PDMS) pad is placed over the BCP films that are linearly translated over the cold block-hot wire-cold block CZA setup as shown in Figure 1.a. The temperature of the hot rod ($T_{\text{max}}$) was set to 205 °C (well above the $T_g$ of all polymers under investigation) whereas the cold blocks were kept at 26 °C giving an in-plane temperature gradient of $\sim 45$ °C/mm as shown in Figure 1.b. The films were translated with a velocity of $V = 5 \mu$m/s, which corresponds to a $T_g$-to-$T_g$ annealing time of 21 min on the temperature gradient curve. Since, the coefficient of thermal expansion of PDMS is about 4 times that of the BCP film (Figure 1.b), a thermal gradient-induced differential expansion and contraction of PDMS over the BCP results in a single oscillatory shear cycle, generating a shear stress of about 120 kPa at the PDMS-BCP film interface.$^{35}$ This shear stress directs the alignment of the BCP lamellae parallel to the substrate.

**Morphology of BCP Films.** By processing the film using CZA-SS, we obtain a completely parallel (to the substrate) lamellar morphology within the entire $\sim 1 \mu$m thick PS-b-PMMA film as shown in the cross-sectional TEM image in Figure 1.e. The dark phase in the TEM images corresponds to PS and light phase is the PMMA. For comparison, an "as-cast" film (no annealing) resembles a homogeneous single-phase morphology as illustrated in the cross-sectional TEM image in Figure 1c.

To underscore the importance of CZA-SS on BCP orientation and on the resulting breakdown properties, control samples were subjected to comparable thermal annealing at 205 °C ($\gg T_g$) in a vacuum oven and the morphologies were compared to CZA-SS samples. At relatively short times (6 h), the oven-annealed films exhibited predominantly perpendicular (to substrate) lamellar domains with some fraction of parallel lamellae near the substrate and air interfaces; at longer annealing times (96 h), the fraction of perpendicular lamellae decreased and parallel lamellar orientation dominated. This can be ascribed to the preferential wetting of PMMA on the substrate surface due to the polar interaction of PMMA with the native oxide surface on the Aluminum via the carbonyl bond (\text{\text{\text{--C=O}}}), similar to known interaction with SiO$_2$ surface on silicon substrates, while the lower surface tension PS covers the film-air interface.$^{24}$ This is followed by propagation of the surface interaction induced parallel lamellar ordering from both the substrate and air interface via thermal annealing. However, the strength of the surface interactions

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**Figure 1.** Thermal processing of BCP films and cross-sectional TEM imaging of PS-b-PMMA films. (a) Schematic for directed self-assembly of BCP film using cold zone annealing-soft shear (CZA-SS). Reprinted with permission from 35. Copyright 2012 American Chemical Society. (b) Sharp thermal gradient profile induced by the CZA setup and linear thermal expansion coefficients of materials. Cross-sectional TEM images (no stain) of PS-b-PMMA (19.5-b-18.1 kg mol$^{-1}$) film on Al coated quartz substrate: (c) as-cast, (d) oven annealed for 96 h at 205 °C under vacuum, and (e) CZA-SS at $T_{\text{max}} = 205$ °C, $V = 5 \mu$m/s. (Scale bar: 0.2 μm).
diminish with increasing distance and lead to mixed orientations away from the surfaces, especially in thick films. Hence, even after 96 h annealing, the lamellae did not orient completely parallel throughout the film, exhibiting some fraction of perpendicular lamellar component as seen in Figure 1.d.

The internal ordering and average interfacial structure within the CZA-SS processed films was further examined through neutron reflectivity (NR) studies on a deuterated BCP, dPS-b-PMMA (19.5-b-18.1 kg mol\(^{-1}\)). The PS block is deuterated for neutron contrast; the self-assembly behavior of this BCP is essentially identical to its nondeuterated analogue. A NR curve...
of a CZA-SS processed film, along with a best fit, is shown in Figure 2.a.

Five orders of well-defined Bragg reflection peaks are seen, indicative of highly ordered parallel lamellar morphology; that is, domains of dPS and PMMA alternating in the film normal direction. On fitting the data to a neutron scattering length density (SLD) profile, we obtain a composition profile normal to the substrate as seen in the inset of Figure 2.a and enlarged in Figure 2.b. The film consists of approximately 44 repeating units having an average width of 27 nm, which is comprised of a dPS (~13 nm, SLD ≈ 5.7 × 10^{-6} Å^{-2}) and PMMA (~14 nm, SLD ≈ 0.9 × 10^{-6} Å^{-2}) block. The PMMA preferentially wets the native oxide-covered Aluminum substrate, while the lower surface tension of PS causes this material to cover the PMMA layers (Figure 2b). To study the effect of molecular weight on the BCP ordering and subsequently on the breakdown characteristics, we also used a higher molecular weight PS-b-PMMA (33-b-33 kg mol^{-1}) BCP and subjected it to the same processing conditions as the lower molecular weight PS-b-PMMA.

Grazing-incidence small-angle X-ray scattering (GISAXS) experiments were performed to investigate the orientation and long-range ordering within the films under different processing conditions for all BCP systems under this study. The grazing-incidence angle (0.18°) was kept above the film critical angle (0.16°) thereby ensuring the measurement probes the entire film thickness. Two-dimensional GISAXS images and their corresponding intensity profiles for the higher molecular weight PS-b-PMMA (33-b-33 kg mol^{-1}) system are shown in Figure 3. Horizontal line cuts along q_x (in the X–Y plane of the scattering geometry) and vertical line cuts along q_z as indicated by the boxed areas in Figure 3a allow quantification for in-plane (perpendicular lamellae) and out-of-plane (parallel lamellae) orientation, respectively, as shown in Figure 3e–f.

The as-cast film exhibits a very broad first order peak in q_z and no distinct peaks in q_x which is typical for poorly ordered materials lacking well-defined domains (Figure 3a), whereas oven annealing results in the appearance of peaks along both q_z and q_x indicating the coexistence of perpendicular and parallel lamellar orientation (Figure 3b, c). For the 6 h oven-annealed sample, up to 4 orders of in-plane Bragg peaks with q_z values in the ratio 1:2:3:4 are observed, while none out-of-plane (q_x), indicative of predominantly perpendicular lamellae throughout the film. At longer annealing times (96 h), there is an appearance of higher-order out-of-plane (q_x) peaks along with the in-plane (q_z) peaks indicating a mixed lamellar morphology. The diffuse isotropic rings connecting the in-plane and out-of-plane Bragg peaks is a signature of lamellae with mixed orientations. Thus, with increasing annealing time, parallel oriented BCP domains are favored over vertical orientation, as discussed previously. Nevertheless, there exists a point where the morphology does not evolve further with additional annealing. In contrast to oven annealing, CZA-SS processed films exhibit no distinct peaks along q_z indicating there is no perpendicular lamellae component, while higher-order peaks are observed along q_x (marked arrows) indicating a morphology with well-defined lamellar domains oriented parallel to the substrate.

Figure 3. Probing the ordering and orientation of BCP films using GISAXS. 2-Dimensional GISAXS images of (a) as-cast, (b) short time oven-annealed (6 h), (c) long time oven annealed (96 h), and (d) CZA-SS processed films with corresponding (e) in-plane and (f) out-of-plane integrated intensity profiles for PS-b-PMMA (33-b-33 kg mol^{-1}).
substrate. The presence of a faint bump in the CZA-SS intensity curve (green), around the position of first maxima of the oven-annealed curves in Figure 3e, can be attributed to a slight in-plane modulation in the parallel lamellar structure; however, considering the intensity and the breadth of the bump in comparison with the Bragg peaks of oven annealed curves, it can be concluded that there is no significant vertical component and that the parallel ordering is near perfect. Similar GISAXS trends are observed for the low molecular weight PS-b-PMMA films (19.5-b-18.1 kg mol⁻¹) films (see Figure S2); in fact the qx integrated intensity profile for the CZA-SS sample shows no sign of in-plane modulations indicating a perfect parallel lamellar structure.

The processing-morphology trend for the PS-b-P2VP films is very similar to that for PS-b-PMMA as evidenced by the cross-sectional TEM images and complementary GISAXS data shown in Figure 4. The TEM of as-cast films resembles a homogeneous single-phase system, while GISAXS shows very broad spots in qx indicating domains lacking any long-range order. Short time oven annealing results in morphology with mixed lamellar orientations having a predominant perpendicular component. The CZA-SS processed films show extremely well ordered and highly oriented parallel lamellae (domain size ~37 nm) that resemble a multilayered structure of alternating PS and P2VP layers. These results clearly underscore the importance of the CZA-SS process and its general applicability.
to continuously fabricate highly oriented, well-ordered lamellar structures in micron thick BCP films on nonmodified, conductive substrates at much faster rates than conventional annealing and potentially over large areas. For BCPs, the domain size $L_0$ or the lamellar thickness is a function of the molecular weight ($N = \text{degree of polymerization}$) and the interaction parameter between the two blocks ($\chi$); $L_0 \propto \chi^{1/6} N^{2/3}$. In this work, we used two different molecular weights of PS-$b$-PMMA, thereby increasing the domain size from 27 to 37.5 nm and decreasing corresponding number of lamellar layers from ~88 to ~63 for PS-$b$-PMMA (19.5-b-18.1 kg mol$^{-1}$) and PS-$b$-PMMA (33-b-33 kg mol$^{-1}$) respectively, while keeping film thickness constant around 1 $\mu$m. Recent trends in BCP lithography have pushed the lower limits of lamellar thicknesses achievable down to ~5 nm. Thus, by changing the block chemistry ($\chi$), molecular weight or even film thickness, the lamellar thicknesses and corresponding number of layers can be readily tuned. Since, CZA-SS operates via a shear mechanism, it has broad applicability over a range of chemistries, molecular weights and film thicknesses to achieve highly ordered self- assembled BCDFs, as long as the shear stress is sufficient to direct the self-assembly.

**Dielectric Performance of BCP Films.** BCP films with representative unorderd, perpendicular and parallel lamellar ordering were subjected to dielectric breakdown experiments to characterize dielectric strength. These were obtained by testing the film as-cast, annealing the film for 6 h under vacuum, and applying the CZA-SS procedure, respectively. Films annealed for 96 h that exhibit a mixture of lamellar orientations were also examined. Figure 5.a-d summarizes the dielectric failure data for PS (100 kg mol$^{-1}$), P2VP (77 kg mol$^{-1}$), PMMA (88 kg mol$^{-1}$) homopolymer control films, 33-b-33 kg mol$^{-1}$ PS-$b$-PMMA, 19.5-b-18.1 kg mol$^{-1}$ PS-$b$-PMMA, and 25-b-25 kg mol$^{-1}$ PS-$b$-P2VP films, respectively.

A two-parameter Weibull cumulative probability function was employed to fit the data and is given by $P(E) = 1 - \exp\left(-\left(E/E_{BD}\right)^\beta\right)$, where $P(E)$ is the cumulative probability for failure, $E$ is experimental breakdown strength, $E_{BD}$ reflects the electric field where there is 63.2% probability for failure, $\beta$ is the shape parameter associated with the least-squares fit of the distribution. Data from these fits are summarized in Table 1. The PS and PMMA homopolymer films yielded $E_{BD}$ values of 251 V/$\mu$m and 494 V/$\mu$m, respectively (Figure 5.a). Thus, a film comprised of 1:1 PS: PMMA would be expected to have characteristic breakdown strength of ~375 V/$\mu$m by simple rule-of-mixtures. As-cast 33-b-33 kg mol$^{-1}$ PS-$b$-PMMA films yielded a breakdown strength value of 403 V/$\mu$m, which is comparable to this rule-of-mixture estimate (Figure 5b). Films annealed for 6 h exhibited a significant drop off in breakdown strength (316 V/$\mu$m), while films annealed for 96 h showed an enhancement (529 V/$\mu$m) as compared to the as-cast film. CZA-SS processed films exhibited the best performance with an $E_{BD}$ value of 599 V/$\mu$m, which is a 60% improvement above the rule-of-mixtures estimation. Similar trends were observed for the lower molecular weight PS-$b$-PMMA (19.5-b-18.1 kg mol$^{-1}$), with $E_{BD}$ values approximately 10% lower when comparing equivalent sample morphologies. Here, the as-cast film yielded an $E_{BD}$ value of 346 V/$\mu$m, while the 6h and 96h films had breakdown strengths of 286 V/$\mu$m and 476 V/$\mu$m, respectively. The CZA-SS sample again exhibited the highest breakdown strength with a value of 520 V/$\mu$m. $\beta$ values were observed to be unaffected by molecular weight or morphology for PS-$b$-PMMA films, falling in the range of 4–7 for all films under investigation. Results for breakdown tests performed on 25-b-25 kg mol$^{-1}$ PS-$b$-P2VP films (as-cast, annealed for 6 h, and CZA-SS processed) are shown in Figure 5.d. Here, the breakdown strength trend matches the one observed for the PS-$b$-PMMA, which proves the general dependence of the breakdown on the nanostructure of the BCP films irrespective of the disparity in dielectric properties of individual blocks; however, the difference between CZA-SS (390 V/$\mu$m) and as-cast (344 V/$\mu$m) films was smaller (~14% enhancement) as compared to the PS-$b$-PMMA samples. It is interesting to note that although the morphology and layer thickness for both CZA-SS processed 25-b-25 kg mol$^{-1}$ PS-$b$-P2VP and 33-b-33 kg mol$^{-1}$ PS-$b$-PMMA films is similar (~37 nm), there is disparity in the breakdown strength enhancement. This can be attributed to the breakdown strength of P2VP (382 V/$\mu$m) component, which is much lower than PMMA and thus limits the breakdown strength enhancement available to PS-$b$-P2VP as compared with PS-$b$-PMMA.

Focusing on the high-breakdown strength PS-$b$-PMMA system, we discuss the results in more detail, outlining the likely mechanism of $E_{BD}$ enhancement. The results of Figure 5 clearly demonstrate how BCP morphology impacts breakdown strength, as we observe significant differences in $E_{BD}$ for 6 h oven annealed PS-$b$-PMMA samples when compared to CZA-SS processed films. The compositions of the films are identical—only their sample morphologies are changed (perpendicular lamellae vs parallel lamellae). The dielectric strength for both molecular weight as-cast PS-$b$-PMMA films falls within 10% of the rule-of-mixtures breakdown estimate. This supports the idea that as-cast films behave similarly to a single-phase polymer system whose dielectric behavior is an average of its constituents. Forcing lamellae to align predominantly perpendicular within the film—as in the 6 h annealed films—yielded breakdown values closer to neat PS (250 V/$\mu$m). Since failure is more likely to occur in the weaker breakdown component (PS) and the perpendicular lamellae can span much of the entire sample thickness, it is likely both breakdown initiation and propagation events take place within PS blocks only. In contrast, the films with parallel lamellae obtained via CZA-SS gave $E_{BD}$ values higher than their as-

### Table 1. Summary of Characteristic Dielectric Breakdown Strength ($E_{BD}$) and Shape Parameter $\beta$ Measured for PS, PMMA, PS-$b$-PMMA, and PS-$b$-P2VP Films

<table>
<thead>
<tr>
<th>Sample</th>
<th>$E_{BD}$ (V/$\mu$m)</th>
<th>$\beta$</th>
</tr>
</thead>
<tbody>
<tr>
<td>100k PS</td>
<td>251</td>
<td>7.1</td>
</tr>
<tr>
<td>77k P2VP</td>
<td>382</td>
<td>8.5</td>
</tr>
<tr>
<td>88k PMMA</td>
<td>494</td>
<td>10.2</td>
</tr>
<tr>
<td>19.5k-b-18.1k PS-$b$-PMMA</td>
<td>289</td>
<td>4.1</td>
</tr>
<tr>
<td>6 h oven</td>
<td>346</td>
<td>5.4</td>
</tr>
<tr>
<td>as-cast</td>
<td>467</td>
<td>6.4</td>
</tr>
<tr>
<td>96 h oven</td>
<td>520</td>
<td>4.6</td>
</tr>
<tr>
<td>CZA-SS</td>
<td></td>
<td></td>
</tr>
<tr>
<td>33k-b-33k PS-$b$-PMMA</td>
<td>316</td>
<td>6.4</td>
</tr>
<tr>
<td>6 h oven</td>
<td>403</td>
<td>5.8</td>
</tr>
<tr>
<td>as-cast</td>
<td>529</td>
<td>6.2</td>
</tr>
<tr>
<td>CZA-SS</td>
<td>599</td>
<td>6.1</td>
</tr>
<tr>
<td>25k-b-25k PS-$b$-P2VP</td>
<td>322</td>
<td>4.9</td>
</tr>
<tr>
<td>6 h oven</td>
<td>344</td>
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</tr>
<tr>
<td>as-cast</td>
<td>390</td>
<td>10.4</td>
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We hypothesize that for the electrical tree to penetrate across the multilayered BCP film, it must overcome the block interfacial barriers and in order to do so, it must break bonds at the interface. In previous studies of the barrier effect, it has been shown that the tree propagation is highly sensitive to the interfacial strength, that is, the type of bonding between the barrier and the matrix. In the case of the highly ordered and aligned BCP, the barrier interfaces are sharp and constituted by strong covalent bonds between the two blocks. Hence, we believe the \( E_{BD} \) values are highest for CZA-SS processed parallel lamellar films where the multiple sharp interfaces of the lamellae with strong covalent bonds are highly oriented throughout the film, normal to direction of tree propagation, in contrast to 6 h oven annealed films where the interfaces are parallel to the direction of tree propagation. The high lateral coherence of the parallel lamellae must force the tree along a tortuous pathway along the interfaces until, at a higher field strength (enhanced \( E_{BD} \)) it overcomes the barrier to propagate toward the other electrode, causing breakdown. Higher molecular weight PS-b-PMMA films (33-33 kg mol\(^{-1}\)) exhibited consistently higher breakdown strengths for all sample morphologies, which is consistent with previous experiments performed on semicrystalline and amorphous homopolymers. For instance, the dielectric strength of polystyrene has been shown to monotonically increase with increasing molecular weight, where the largest changes occur around 20 kg mol\(^{-1}\)—close to the entanglement molecular weight (16 kg mol\(^{-1}\)). A 10% difference in \( E_{BD} \) was reported when comparing polystyrene films of molecular weight 19 and 33 kg/mol. Combined with similar differences expected for PMMA breakdown strength, we can explain the discrepancies between the lower and higher molecular weight BCP films. This behavior may be caused by differences in regularity of...
molecular conformation or density of chain ends within the film.

The improvement in breakdown strength facilitated by the layered film architecture comes at no cost to the relative permittivity and loss of the films. The permittivities of BCP films essentially follow a mean-field approximation and the values lie between those of component homopolymers. Dielectric spectroscopy experiments demonstrate that all PS-b-PMA and PS-b-P2VP samples, regardless of morphology and molecular weight, have comparable permittivity spectra as shown in Figure 6a, b and Figure S4. This is expected as the relative composition of the PS and PMMA/P2VP block fraction remains constant for all samples; only the morphology is variant. Therefore, we have successfully improved breakdown strength with no deleterious effect to the relative permittivity or loss ($\tan\delta$), which is of critical importance when considering the energy storage performance of the films.

The maximum theoretical energy storage densities of the BCDFs with different morphologies and respective homopolymer controls are calculated based on eq 1 using the experimentally obtained $E_{BD}$ and $\varepsilon_r$ values. The energy density trends are plotted in Figure 7. The CZA-SS processed films with parallel lamellae always exhibit higher energy densities than corresponding as-cast or oven annealed films for both PS-b-PMA and PS-b-P2VP. In fact, the PS-b-PMA films with layered architecture exhibit more than double (~2.25 times) the energy storage density (4.6 J/cm$^3$) as compared to the as-cast films (2 J/cm$^3$) and ~3.5 times that of oven annealed films (1.3 J/cm$^3$) with predominantly perpendicular lamellae. They also exhibit higher energy storage performance than the individual PS and PMMA homopolymer components. It is noteworthy that although the BCP systems used here comprise of amorphous polymer components rather than semicrystalline polymer capacitor materials, yet the maximum energy densities obtained using the self-assembly approach are comparable or even higher (4.6 J/cm$^3$) than the industrial-standard semicrystalline BOPP films (4 J/cm$^3$). With a judicious choice of BCP comprising a high dielectric strength ($E_{BD}$) block, say semicrystalline PP, and another block with high permittivity ($\varepsilon_r$), say PVDF, the resulting energy storage density (a product of the two quantities) could be significantly enhanced through this self-assembly strategy in future applications of this technology. Thus, the BCP strategy allows for the design of block specific functionality and is not only significant from the energy storage point of view, but the weaker dielectric component may be used to incorporate other useful properties such as high mechanical strength, UV resistance, high refractive index, low oxygen permeability, which permit use of these films in other multifunctional technologies. For instance, in the present system, one may consider that in the case of higher molecular weight PS-b-PMA, we enhanced the breakdown strength of the stronger component (here PMMA) by ~21% which is notable, while the PS layers can bring auxiliary function such as better UV resistance for outdoor applications.

**CONCLUSION**

We have successfully demonstrated a new material design and processing method to fabricate block copolymer dielectric film (BCDF) capacitors with multilayered (lamellar) nanostructured dielectric media via directed self-assembly. CZA-SS is a powerful technique to fabricate highly ordered BCP on nonmodified conductive (rigid or flexible) substrates and is also viable with high-throughput roll-to-roll processing as compared to other directed self-assembly methods. The highly ordered lamellar structures obtained by CZA-SS yield up to ~50% increase in $E_{BD}$ of the nanostructured BCP film when compared to an as-cast BCP film, due to the barrier effect. This translates to a ~ 2.25 times increase in the maximum energy storage density since it has a quadratic dependence on the $E_{BD}$ ($U = 1/2\varepsilon_r E_{BD}^2$). With the potential to synthesize new BCPs having optimum dielectric properties, it is possible to obtain a catalog of tailored BCP materials. Using the simple but robust self-assembly strategy, these BCP materials can be used to fabricate multilayered high-energy density, pulsed power, flexible capacitor films for use in directed energy, electric vehicles and future high-power flexible electronics.

**METHODS**

**Film Preparation.** Diblock copolymers PS-b-PMA ($M_w = 66$ kg mol$^{-1}$), deuterated PS-b-PMA ($M_w = 37.6$ kg mol$^{-1}$) with the PS block deuterated, PS-b-P2VP ($M_w = 50$ kg mol$^{-1}$), and homopolymers PS (100 kg mol$^{-1}$), PMMA (88 kg mol$^{-1}$), and P2VP (77 kg mol$^{-1}$) were purchased from PolymerSource Inc. and used as received. Electron beam physical vapor deposition was utilized to coat 100 nm thickness aluminum films (roughness ~2 nm) onto quartz slides (GM Associates Inc.) to form the bottom electrode for breakdown testing. Polymer samples were dissolved in toluene and 10% (w/w) solutions were flow-coated (Figure 1) onto UV-Ozone (Novoscan Tech., Inc.) cleaned Au-coated (1 in. x 3 in.) quartz slides. UV-ozone treatment removes organic impurities from the substrate surface, forms an oxide layer and thereby increases wettability of polymer film. Film thicknesses were measured using a Bruker DektakXT profilometer. Prior to processing, the films were dried in a vacuum oven at 65 °C for 12 h to remove residual solvent.

**Preparation and Installation of PDMS Pad.** Syrgan-182 (Dow Corning Corp., USA) was used to make the polydimethylsiloxane (PDMS) films. The PDMS elastomer/curing agent (20:1 weight ratio) were mixed and degassed in vacuum followed by pouring onto a glass slide and thereby increases wettability of polymer film. After polymerization (30 min at 80 °C), the PDMS was then peeled off the glass to obtain ~0.5 mm thick films. PDMS pad of dimensions similar to the film (~2.5 cm lateral size) was cut out and manually placed onto the BCP film by light pressing. The PDMS is prepared on a smooth glass surface, hence the

![Figure 7. Energy density trends of homopolymer films and BCP dielectric films (BCDFs). Maximum theoretical energy storage performance of component homopolymer and BCP dielectric films under different processing conditions, calculated from eq 1 using experimentally obtained values of breakdown strength ($E_{BD}$) and relative permittivity ($\varepsilon_r$).](image-url)
smooth PDMS surface allows for conformal contact with the film during the CZA-SS process.

**Film Processing.** The BCP sample with PDMS pad was then translated across the CZA-SS gradient assembly (Supporting Information) at a speed of 5 μm/s using a syringe pump. The CZA-SS procedure was performed under ambient conditions. After CZA-SS, the PDMS was manually peeled off the film after the film is cooled below Tg of both blocks. Below Tg, the film is in the glassy state and adhered to the substrate, hence stable during the peeling process. Conventional oven annealing was carried out in an oven at 205 °C under vacuum.

**Electron Microscopy.** TEM samples of the film cross sections were made via the in situ lift-out method using a FEI Helios 600 dual-beam FIB and imaged with a JEOL 1400 LAB6 TEM at 120 kV. All images are brightfield images.

**Neutron Reflectivity.** The Neutron reflectivity (NR) experiments were carried out on the NG7 horizontal reflectometer at NIST Center for Neutron Research. A 4.76 Å collimated neutron beam with a wavelength divergence of 0.18 Å was used. Scans were made over a wave vector magnitude (qz) range from 0.005 to 0.16 Å−1. A pure quartz substrate was used in place of Al-coated quartz for the NR measurements for ease of data fitting. Data fitting and conversion of the NR spectra to the scattering length density (SLD) and concentration profile was done using Reflex package software (NIST CNR).

**Grazing-Incidence Small-Angle X-ray Scattering.** GISAXS experiments were performed at beamline X9 of the National Synchrotron Light Source, Brookhaven National Laboratory. An X-ray beam of 7.35 (8-1D) or 14.1 keV (X7) impinged on the films (in vacuum) at an angle well above the critical angle of the films, to probe bulk of the film. Data was collected on a Dectris Pilatus 1 M pixel-array detector. Data analysis and conversion to q-space was done using GIXSGUI software in MatLab, developed at beamline 8-1D.

**Dielectric Testing.** All samples were annealed at 80 °C for 12 h prior to breakdown testing to remove residual solvent and adsorbed moisture that may be present within the film. Dielectric breakdown trials were performed using a 10 kV Spellman SL300 high voltage supply controlled by a ramping circuit (Figure S3). The ramp was set such that a breakdown event occurs at ~20 s in accordance to ASTM standard protocol for short-term dielectric stress tests (which corresponds to 50–300 V/s, depending on the sample).13 Once >1 mA current passes through the device, a silicon rectifier switch activates and breaks the circuit. The breakdown voltages were read from a Fluke 289 multimeter set in peak capture mode. Our experimental geometry features a copper rod with a hemispherical end (radius of curvature = 2.5 mm) that makes direct contact with the BCP film. Breakdown is confined to a small sample region (area ~0.1 cm²) in an attempt to remove the influence of film heterogeneity by spatially localizing the electric field. At least 15 breakdown trials were performed for each film to conduct Weibull failure analysis. The copper contact rod was polished after every 15 breakdowns using diamond paste to remove pitting. All experiments were conducted at room temperature in an N₂ purged environment where relative humidity was observed in the range of 10–20%. Breakdown voltages were converted to breakdown strength by measuring film thickness near each test site via profilometry.

Permittivity measurements were conducted using a Novocontrol Impedance Spectroscopy Analyzer and facilitated by coating the BCP films with a 3 mm diameter circular aluminum top electrode.

**ASSOCIATED CONTENT**

Supporting Information

The Supporting Information is available free of charge on the ACS Publications website at DOI: 10.1021/acsami.5b11851

Schematic of the flow coating method, CZA-SS experimental procedure, GISAXS data of PS-b-PDMA (19.5-8-18.1 kg mol⁻¹) films, schematic of dielectric breakdown test setup and dielectric spectroscopy of PS-b-PMMA (33-b-33 kg mol⁻¹) films (PDF)

**AUTHOR INFORMATION**

**Corresponding Author**

E-mail: alamgir@uakron.edu.

**Author Contributions**

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**Notes**

Certain commercial equipment, instruments, or materials (or suppliers, or software, etc.) are identified in this paper to foster understanding. Such identification does not imply recommendation or endorsement by the National Institute of Standards and Technology nor does it imply that the materials or equipment identified are necessarily the best available for the purpose. The authors declare no competing financial interest.

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