Dynamics of Photogenerated Surface Charge on BiFeO$_3$ Films

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ABSTRACT We report on the spatial and temporal evolution of photoinduced charge generation and carrier separation in heteroepitaxial BiFeO$_3$ thin films deposited on Nb:SrTiO$_3$ as measured in ambient at room temperature with Kelvin probe and piezoresponse force microscopy. Contributions from the self-poled and ferroelectric polarization charge are identified from the time evolution of the correlated surface potential and ferroelectric polarization in films as grown and following poling, and at different stages and intensities of optical illumination. Variations in the surface potential with bias voltage, switching history, and illumination intensity indicate how both bulk ferroelectric photovoltaic and the domain wall offset potential mechanisms contribute to the photogenerated charge.

KEYWORDS: ferroelectric photovoltaics - BiFeO$_3$ - Kelvin probe microscopy

Ferroelectric semiconductors are attractive candidate photovoltaics for replacement of conventional silicon-based bipolar junctions owing to efficient ferroelectric polarization-driven carrier separation.1−3 Much attention has been focused on the bulk photovoltaic effect in traditional ferroelectric materials4 such as BaTiO$_3$ and Pb(Zr,Ti)O$_3$, which possess band gaps in the ultraviolet,5 and on the local photoinduced charge dynamics on the surfaces of BaTiO$_3$.6 Among ferroelectric oxide perovskites, BiFeO$_3$ (BFO) is distinguished by having a band gap in the visible (ca. 2.6−2.8 eV6,7) and possessing room-temperature intrinsic multiferroic properties permitting several modes of investigating and controlling the photovoltaic effect. The photovoltaic effect in BFO has both bulk and domain wall contributions; the former is due to ferroelectric polarization5 and the latter also involves ferroelastic domains.2 Potential offsets owing to the presence of domain walls enable a more efficient separation of carriers, and a net potential difference evolves across the entire sample.2 Alexe et al.,3 using a scanned local probe to collect the enhanced photogenerated carriers, reported that the anomalous photoexcited carriers in a BFO single crystal exist over the entire crystal and do not recombine within the ferroelastic domains.

The understanding of issues relating to the dynamic processes of charge generation and transport in BFO remains limited, however. Collection and analysis of the evolution of the surface potential (SP) owing to the photoinduced charge permit study of the effects of polarization on exciton transport and quantitative analysis of the effect of domain wall number. Kelvin probe force microscopy (KPFM)8−13 remains a powerful noncontact local probe of SP, including SP arising from photoexcited charge.8 Here we report on the use of KPFM, in conjunction with piezoresponse force microscopy (PFM), to probe the local spatiotemporal evolution of the SP in BFO at different stages of illumination and switching arising from the separation of carriers. We describe quantitatively both the influences of the orientation of ferroelectric polarization and of the domain wall on the measured SP of the BFO film associated with piezoresponse force microscopy (PFM). A direct visualization of charge generation and recombination enables further insight into the roles of ferroelectric polarization and domain walls on charge transfer and diffusion in BFO.

RESULTS AND DISCUSSION

Following the writing of a small square pattern with $V_{tip} = -9.8 \text{ V}$ within a larger square pattern ($V_{tip} = +9.8 \text{ V}$, contrast in the SP (Figure 1a), the plane-normal component of piezoelectric displacement amplitude (Figure 1b) obtained sequentially in the dark can be discerned. The as deposited, unpoled area of the film exhibits a preferential outward film-normal component of polarization (as seen in the perimeter of the scan area that appears orange on the color
map) due to self-poling caused by the electric field that arises due to the interfacial potential barrier between the BFO and NSTO. The large SP differences between (i) the outer and inner regions of poling as seen by the yellow and black areas in Figure 1a and (ii) each in relation to the surrounding unpoled area are due to the induced ferroelectric polarization charges and screened charges under given biases, respectively. The SP contrast (Figure 1a) is much stronger than that in the PFM image (Figure 1b), and suggests complementarity of KPFM in analyzing photogenerated charges. Macroscopically, these films exhibit a significant photovoltaic response and nonzero short-circuit current density ($J_{sc}$ = $-4.5 \times 10^{-6}$ mA/cm$^2$ and an open circuit voltage $V_{oc}$ of 0.26 V under an illumination intensity of $\sim 10 \text{ mW/cm}^2$) owing to the self-polarization of BFO$^9$ (Figure 1c). The photovoltaic efficiency amounts to $\sim 1.2 \times 10^{-3}\%$, much smaller than other reported values for BFO$^{14}$ because of the low transmissivity of the Pt top electrode. The data in Figure 1d are plotted on a semilog scale where the change in conductivity upon illumination is the result of excitation across a BFO/NSTO p–n junction,$^{15}$ in addition to the strong rectifying character.

Since KPFM can be taken as a measure of the electrostatic interactions between the conductive tip and sample, contributions to the obtained SP from the BFO and NSTO can be analyzed as parallel capacitances.$^{11,16,17}$ Schematic interfacial band diagrams are sketched (Figure 2), where the BFO film and NSTO substrate are assumed to be nondegenerate semiconductors$^{15}$ and the Fermi level of NSTO is in the vicinity of the bottom of the conduction band. A Schottky-like barrier is at the tip–sample interface which is modulated by the local ferroelectric polarization$^{18,19}$ and the polarization causes the conduction and valence bands to bend at the interface and produce surface charge with opposite sign on the different domain at low bias$^{19}$ (Figure 2a). When tip is brought in contact with the sample (in the absence of Fermi level pinning) Fermi level alignment takes place (Figure 2b), inducing an electrostatic force that depends on the SP difference (SP$_1$).$^{16}$ In the KPFM measurement, an external voltage $V_{dc}$ is applied to the probe in order to nullify SP$_1$ (flatband case, Figure 2c).

To relate the KPFM measurement to the photovoltaic effect of BFO, we sketch a band diagram under illumination for the as-deposited film (Figure 2d), where a photon is incident upon the top surface of BFO and an exciton is separated, producing a gradient in the quasi-Fermi levels.$^{20}$ Here, a surface photovoltage ($V_{ph}$) produces a surface photopotential (SP$_{ph}$).
When a negative bias is applied under illumination, the SP decreases by about 0.16 V, with an exponential-like onset of larger SP. These positive shifts from dark to light arise from charge separation of the photogenerated electron–hole pairs with s drifting of holes to the surface and electrons to the NSTO substrate under the self-poled polarization-induced internal electric field, owing to the bulk photovoltaic effect.

To quantify the observed dynamic charging, measurements of the time-dependent SP evolution were performed. The sample was illuminated at the saturation power (158 mW/cm²) and the time evolution of SP was measured at different locations on the sample surface. The average SP is seen to vary as the light is switched on and off (Figure 3d). During an on/off cycle, the SP saturates more quickly in the off/on transition as compared to the characteristic decay time scale in the on/off transition, corresponding to the charging and discharging processes. Both measured SP changes trend according to a single exponential, that is, \( \text{SP}(t) = \text{SP}_0 e^{-t/\tau_d} + \text{SP}_1 e^{-t/\tau_l} \) where \( \text{SP}_0 \) and \( \text{SP}_1 \) are the initial values, and \( \tau_d \) and \( \tau_l \) are mean onset and decay lifetimes for dark-to-light and light-to-dark. Estimated lifetimes for charging and discharging (7.6 and ~31.5 s, respectively) indicate that while the surface retains charge on the time scale of seconds, the discharging time appears to be longer than for charging and may be explained by the release of electrons from shallow traps.
traps\(^9\) (i.e., oxygen vacancies\(^21\)), or electrostatic potential offset across the domain walls of BFO film.\(^2,4\) (This asymptotic charging and discharging behavior is obviously orders of magnitude slower than the photoexcited charge generation and recombination.) The full width at half-maximum (fwhm) is seen to decrease with increasing illumination intensity and is accompanied by a saturation of the fwhm at higher intensity. We propose that the narrowing of the line width can be accounted for by polar asymmetry in the surface charge dynamics: it is straightforward to show that the observed reduction in standard deviation of the SP distribution for increasing illumination intensity is consistent with a recombination of photoinduced

Figure 3. (a) Distributions of surface potential values as collected from a single 5 × 5 \(\mu m^2\) area of the BFO/NSTO heterojunction in the dark and under illumination (48, 158, and 166 mW/cm\(^2\)). (b) KPFM image in the dark (bottom portion of scan) and under illumination (top), 158 mW/cm\(^2\). (c) Corresponding SP profile of panel b; the arrows in panels b and c denote the slow scan axis, and scan progression, respectively. (d) Time evolution of the surface potential of BFO turning on and off the illumination (158 mW/cm\(^2\)), indicative of exponential charging and discharging processes.

Figure 4. (a,c) Demonstration of the effect of direction of polarization on SP profile under \(V_{\text{tip}} = \pm 9.8\) V. The inset images are KPFM images under different illumination intensities. (b and d), corresponding poled and unpoled area surface potential as a function of light intensity from panels a and c, respectively. The insets of panels b and d illustrate the photogeneration processes under different polarization orientation.
negative charge on a faster time scale than positive charge.

It is well-known that the photovoltaic response in a ferroelectric can be controlled by electric field as a result of the reversed domain switching.\textsuperscript{1,2,5} To understand the effect of direction of polarization on the observed photoinduced charges, the SP was measured prior to and following reorientation of polarization. Shown in Figure 4a,c are the spatial profiles of SP for the poled area for downward and upward polarization under different selected values of illumination intensity, respectively. As expected, the poled area exhibits a different SP behavior compared with the as-deposited (unpoled) portions. Meanwhile, the diode-forward diode-rectifier effect. With increasing illumination intensity, the observed SP arises from (i) charges injected from the conductive tip, from (ii) screened polarization charges of the oriented dipoles, and from (iii) the photoinduced charges. In addition, the SP is observed to increase with an increase in the illumination intensity, as shown in Figure 4 panels b and d for the as-grown (unpoled), positively and negatively poled films, respectively, in accordance with the photovoltaic effect. With increasing illumination intensity, the SP of the negatively poled area saturates with lower illumination intensity, while the positively poled area requires higher illumination intensity to reach saturation. These observations indicate that the photoinduced electron—hole pairs are separated and recombine with the injected charges from the tip. Under positive bias (here, polarization is downward, as shown in the inset of Figure 4b), the injected and photoinduced holes are compensated by the produced negative polarization charges, thus the SP increases slowly as a function of intensity. Under negative bias (here, polarization is upward, as shown in the inset of Figure 4d), the photoinduced holes and produced positive polarization charges recombinced with the injected negative charges (i.e., electrons) at the surface and led to the sharp increase and saturation of SP (Figure 4d).

To quantify the effects of polarization direction on the evolution of surface charge density, time dependencies of the SP of the poled area in the dark and under illumination were also measured (Figure 5). For the positively poled area (Figure 5a) the SP increased sharply in the poled area when the illumination is turned on. However, removing the illumination caused the SP to relax more slowly, with average lifetimes for charging and discharging of 19.5 and 144.4 s, respectively. Our observation of a longer lifetime for the poled area as compared with that for the unpoled area (Figure 5a) suggests that a further increase of surface potential could be limited because of the suppression or delay of charge migration into the film by the positively poled field. One possible mechanism is that the contribution from the photoexcited holes is enhanced by injected holes under illumination and the negative polarization charge becomes a barrier during the discharging process, resulting in an increased macroscopic current density. For the negatively poled area (Figure 5b), the SP increased significantly following illumination with average charging lifetimes of \( \sim 93.7 \) s for the initial illumination stage and \( \sim 75.8 \) s for the second illumination stage. The sharp increase in SP could be attributed to the recombination of photoinduced holes with the injected negative charges on the surface of BFO, and further compensation of the SP by the positive polarization charges. When the light is switched off, the SP decays exponentially with average lifetimes of \( \sim 236.2 \) and \( \sim 104.3 \) s for the first and second dark stages, respectively. The longer observed lifetime for the discharging that accompanies the light-to-dark transition can be ascribed to the photoinduced holes depleted with the injected electrons from the tip where positive polarization charge plays the same role—that of a barrier for the relaxation of the SP. Under the negatively poled condition the photoinduced electrons move to the bottom electrode and induce a transit current. We remark on the additional domain wall contribution to the ferroelectric photovoltaic effect that is due to nanoscale steps of the electrostatic potential occurring at the ferroelectric domain walls in BFO.\textsuperscript{2,22} Such electrostatic potential offset causes a band bending across the domain wall,\textsuperscript{6} similar to the band bending in the depletion layer of a \( p-n \) junction. Here the variations in the local SP were mapped as functions of illumination intensity and applied voltage, including switching. We applied a series of dc biases ranging from \(-10\) to \(+10\) V with a step of \( 2 \) V using line poling,\textsuperscript{23} and subsequent collection of PFM and KPFM images over an area of \( 10 \times 10 \) \( \mu \)m\(^2\) (Figure 6a and 6b, respectively). A poled area with a step of \( 8 \times 0.72 \) \( \mu \)m\(^2\) was observed through the out-of-plane phase image (Figure 6a), indicating that the domain was switched with sufficiently large dc bias. Since the number of domain walls changes under different values of dc bias, this confirms that through measurement of photogenerated current and corresponding SP that the open-circuit voltage increases linearly with the number of domain walls.\textsuperscript{2,24}
We note that the SP behavior in the dark is consistent with reported observations in heteroepitaxial PbTiO$_3$ films,$^{11}$ and that the origin of the SP under dark conditions is from both polarization and screened charges. Once the illumination is turned on and its intensity increased, the SP profile evolutions reflect the photovoltaic effect based on the variation of domain wall number (Figure 6c). The surface charge density and excess screening charge can be estimated by treating these contributions as parallel capacitances,$^{24}$ and using the measured remnant polarization of 52 $\mu$C/cm$^2$, and relative dielectric constant of 220 for these films (Figure 6d). With the onset of illumination, the negative contribution increases gradually, indicating that the injected electrons were recombined with the photoinduced holes. With increased negative dc bias voltage, the domain wall number is reduced and the corresponding domain wall contribution to the photocurrent should also decrease. As for the positively poled area, we observe that the SP changes are different below and above the coercive voltage ($\sim$5 V). Below the coercive voltage, the SP increases linearly for $+2$ and $+4$ V and is seen to saturate when the illumination intensity is $\sim$158 mW/cm$^2$, indicating that photoinduced holes combine with the injected holes. Above the coercive voltage, the domain walls begin to move and the domain wall number gradually decreases in the poled area with higher bias voltage. As a result, the dissociation of photoinduced electron–hole pairs across the domain wall is reduced, causing the SP to begin to decrease for the area poled using $+6$ and $+8$ V. Interestingly, as the bias voltage increased from $+8$ to $+10$ V, the SP is seen to decrease at low illumination intensity ($<20.5$ mW/cm$^2$), but is seen to increase for high illumination intensity ($>20.5$ mW/cm$^2$) (This is not observable in Figure 4b where only one fixed voltage was applied.) Under low illumination intensity, the SP decreased because of reduced overscreening charge density.$^{11}$ However, when the illumination intensity is increased ($>20.5$ mW/cm$^2$), the photoinduced holes compensate the overscreened charges and dominate the SP, as seen by a modest increase for the area poled at $+10$ V.

Finally we remark that molecular adsorbates are known to contribute to the surface potential and specifically to the ferroelectric stability in ultrathin films and nanostructures,$^{25}$ in fact, the slow loss of electrostatic force microscopy contrast following poling for bulk-like BaTiO$_3$ takes place on the time scale of many tens of hours,$^{26}$ and not minutes as we observe presently. Nevertheless rigorous experimental and theoretical studies are needed to understand the contribution from adsorbates in the presence of a local field in ambient conditions to the time evolution of the SP contrast.

**CONCLUSIONS**

In summary, we demonstrate the value of combining KPFM and PFM in mapping the evolution of SP arising from the dynamics of a photoinduced charge within a semiconducting ferroelectric film. The time-dependence to the SP is obtained for the as-grown film and is attributed to a relaxation of both free and screened charges, whereas for the poled area the time
dependence is impacted by the direction of polarization, with contributions both from the bulk ferroelectric photovoltaic effect and from number-dependent domain walls.

**METHODS**

**Sample Preparation.** High quality epitaxial BFO films (~140 nm thick) were grown on 0.5 wt % Nb-doped SrTiO3 (001)-oriented (NSTO) single crystal substrates at 550 °C via pulsed laser deposition using a KrF excimer laser (~248 nm). The partial oxygen pressure was about 50 mTorr and the energy fluence of the KrF excimer laser at the target was ~1 J/cm². Thickness was determined via measurement of a step height and validated using cross-section SEM.

**Macroscopic Photovoltaic Measurements.** Semitransparent Pt top electrodes (ca. 10–20 nm in thickness and ~100 µm in diameter) were sputter-deposited on the BFO films for macroscopic photovoltaic and ferroelectric testing. The macroscopic photovoltage and photocurrent measurements were collected using an electrometer (Keithley 6487) under illumination through the top electrodes.

**Kelvin Probe and Piezoresponse Force Microscopy.** KPFM and PFM images were recorded (Asylum Research MFP-3D) using a Pt-coated cantilever (Olympus AC240, nominal spring constant ∼70 kHz) with the NSTO substrate held electrically to ground. All of the data shown were obtained using the identical AFM cantilever. To produce the photogenerated charge, samples were illuminated using a 150 W tungsten halogen lamp. To determine the local ferroelectric contribution to the photovoltaic effect, a voltage was applied to the tip during the first scan using a contact mode to switch the polarization of BFO film. The SP signal was recorded in successive scans, first in the dark and again immediately under steady-state illumination. To eliminate the pyroelectric contribution to the measured potential, data were collected using relatively slow scanning rates (0.3–0.5 Hz) and analyzed from within a region of uniform illumination intensity by avoiding a strong focus. For the KPFM and PFM data shown in Figure 1, the scan rate for poling was 1 Hz; however, all other data shown in the manuscript was collected following the use of the slower scanning rate for poling (0.3 Hz).

**Conflict of Interest:** The authors declare no competing financial interest.

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**Supporting Information Available:** X-ray diffraction characterization of the film; macroscopic ferroelectric hysteresis loops, leakage currents and the local ferroelectric properties; PFM response collected using a slower scan rate for poling as described in the Methods section; additional KPFM experimental results. This material is available free of charge via the Internet at https://pubs.acs.org.

**REFERENCES AND NOTES**


