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Engineered semiconductor-dielectric interfaces in polymer ferroelectric transistors†

Arash Ghobadi,^a Thomas B. Kallaos,^a Indeewari M. Karunarathne,^a Dilan M. Gamachchi,^a Andrew C. Meng,^{ab} Joseph C. Mathai,^c Shubhra Gangopadhyay^c and Suchismita Guha^{*,ab}

Polymer ferroelectrics are witnessing a renewed interest in organic transistors due to their multi-conductance states. Although their high dielectric constant allows low operating voltages, the polarization fluctuation due to the energetic disorder at the interface reduces the carrier mobility in organic transistors. Here, two copolymers of poly(vinylidene fluoride) (PVDF) with trifluoroethylene (TrFE) and hexafluoropropylene (HFP) as the dielectric layer, and a donor–acceptor copolymer as the active semiconductor layer are used in bottom-gate top-contact transistor architectures. We investigate the impact of the dielectric thickness, external poling, and an added interfacial ultrathin Al₂O₃ layer at the semiconductor–dielectric surface on the performance of organic field-effect transistors (FETs). Although poling the dielectric layer significantly enhances the carrier mobility in PVDF–TrFE-based FETs, it has a minimal effect on PVDF–HFP-based devices. Poled PVDF–TrFE devices with a thickness of 45 nm show the highest saturation carrier mobility, exceeding 1 cm² V^{−1} s^{−1}. The subthreshold swing (SS), which is primarily governed by the trap states at the semiconductor–dielectric interface, is seen to significantly improve when an atomic layer deposited Al₂O₃ film with varying thickness between 2 nm and 12 nm is deposited on PVDF–HFP. In the linear region of operation, PVDF–HFP based FETs with Al₂O₃ yield SS values below 80 mV dec^{−1}. The trap density of states at the semiconductor–dielectric interface was evaluated, providing deeper insight into charge trapping and transport mechanisms.

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Introduction

The high dielectric constant (κ) of oxide and ferroelectric dielectrics provides a path towards low-operating voltage organic field-effect transistors (FETs),¹ especially since they work in the accumulation region. At the semiconductor–dielectric interface, the dielectric layer not only influences the morphology of the semiconducting layer, but it also changes the density of states (DOS) due to local polarization effects.^{2–4} Polarizable gate dielectrics, for example, broaden the DOS at the semiconductor–dielectric interface due to charge–dipole coupling and increase the

localization of carriers at the interface.^{5,6} These effects reduce the carrier mobility in FETs at the cost of achieving low operating voltages with high κ dielectrics. Another route for low operating voltage organic FETs is to reduce the thickness of the dielectric layer. Several strategies involving stacked self-assembled monolayers and AlO_x as ultrathin layers, with thicknesses typically less than 20 nm, have demonstrated operating voltages less than 5 V in organic FETs.^{7–9} However, most of these examples use small molecule organic semiconductors. Moreover, such stacked dielectrics require oxidation or anodization processes to form the AlO_x layer. In addition to low voltage, practical applications of organic FETs demand low subthreshold swing (SS) and high carrier mobilities. Although the carrier mobility in polymeric semiconductors is limited due to the hopping nature of transport between disordered-localized states,^{10,11} other factors such as a polarizable dielectric layer^{12,13} and the source/drain contacts^{14,15} strongly influence charge transport properties in FET architectures. Hence, the same semiconductor may display a wide range of carrier mobilities as κ is tuned from low to high. Several strategies for contact modification including adding self-assembled monolayers have resulted in a remarkable reduction of contact resistance in the tens of Ω cm.^{16–18} Approaches for reducing the Schottky barrier by de-pinning the Fermi level in organic FETs have

^a Department of Physics and Astronomy, University of Missouri, Columbia, MO, 65211, USA. E-mail: guhas@missouri.edu

^b MU Materials Science and Engineering Institute, University of Missouri, Columbia, MO, 65211, USA

^c Department of Electrical Engineering and Computer Science, University of Missouri, Columbia, MO, 65211, USA

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involved introduction of oxygen at the metal-semiconductor surface in p-type transport¹⁹ and adding ultrathin Al₂O₃ on the metal contacts for both p- and n-type transport in organic FETs.²⁰

Ideally, ferroelectric dielectrics could be of great benefit for low operating voltage organic FETs, especially if the polarization fluctuation can be reduced, they are thin, they are solvent compatible with π -conjugated small molecules and polymers, and the trap-DOS at the semiconductor-ferroelectric interface can be lowered. These conditions would permit enhanced switching efficiencies by lowering SS. Further, the choice of a solution processable dielectric and a semiconducting layer allows low-cost fabrication methods. This work focusses on solution processable ferroelectrics and a donor-acceptor copolymer, diketopyrrolopyrrole (dithienylthieno[3,2-*b*]thiophene (DPP-DTT)), for improving the semiconductor-dielectric interface in organic FETs. Since the devices are bottom-gate with top-contact, no additional contact treatment was performed.

Among polymer ferroelectrics, the copolymers of polyvinylidene fluoride (PVDF) have been extremely popular as dielectric layers in organic FETs for applications in memory devices,²¹⁻²⁴ sensors,²⁵⁻²⁷ and more recently in neuromorphic devices.^{28,29} Copolymers of PVDF with trifluoroethylene (PVDF-TrFE) and hexafluoropropylene (PVDF-HFP) have differences in their ferroelectric properties, although their κ values are similar at room temperature. The ferroelectricity in PVDF and its copolymers originate from the β phase, where the carbon atoms are in an all-*trans*-configuration, and the paraelectric α phase arises from a *trans-gauche* conformation.³⁰ PVDF-TrFE upon spin-casting from a solution is already in the β phase and, therefore, metal-insulator-metal capacitors made with PVDF-TrFE show typical hysteresis loops in the polarization *versus* voltage curves, expected from a ferroelectric film.^{31,32} PVDF-HFP, on the other hand, is an elastomer with a higher component of the α phase compared with the β phase.^{33,34} Typical saturation polarization values are $6 \mu\text{C cm}^{-2}$ and $3 \mu\text{C cm}^{-2}$ for PVDF-TrFE and PVDF-HFP, respectively (see ESI†).³⁵ With similar values of κ but differences in their ferroelectric properties, PVDF-TrFE and PVDF-HFP allow an excellent comparison for monitoring the semiconductor-dielectric interfaces in organic FETs.

Our prior work demonstrated that by externally poling the PVDF-TrFE film in the vertical direction, prior to the fabrication of organic FETs, and using DPP-DTT as the active semiconductor layer, improves both carrier mobility and SS if the poling field is $< 100 \text{ mV m}^{-1}$. Differential phase contrast (DPC) images in the scanning transmission electron microscope (STEM) indicated that the vertically poled PVDF-TrFE films have uniform ferroelectric domains throughout the bulk of the film.³⁶ Polarization domains are also seen in the DPC images from the unpoled film, except they are significantly more disordered. The higher mobility in the poled devices could be rationalized by a lower trap DOS, as determined by the Grünwald's method,^{37,38} compared with the unpoled devices. The operating voltage in this study was 20 V. Further reduction of the operating voltage demands lowering the thickness of the polymer ferroelectric layer.

In this work we use two approaches for improving the ferroelectric-DPP-DTT interface. The first is to reduce the

dielectric thickness to below 50 nm. The second approach is to add a thin (2–12 nm) atomic layer deposited Al₂O₃ layer on top of the ferroelectric layer. The operating voltages are below 8 V for both PVDF-TrFE and PVDF-HFP based FETs, which operate as p-type devices. Vertically poling the PVDF-TrFE layer (with thickness $\sim 45 \text{ nm}$) improves carrier mobilities to $> 1 \text{ cm}^2 \text{ V}^{-1} \text{ s}^{-1}$. However, there is a marginal difference in the carrier mobilities between the vertically poled and unpoled PVDF-HFP films. In an effort to further lower the SS, we deposited a thin layer of Al₂O₃ by atomic layer deposition on top of the ferroelectric layer. No significant improvement in SS was observed with PVDF-TrFE when 2 nm, 3 nm, and 5 nm of Al₂O₃ were deposited. Additionally, the carrier mobilities are also seen to be lower than that of the unpoled devices. There is a significant improvement in SS for the PVDF-HFP devices with the layer of Al₂O₃, highlighting the role of the oxide layer in passivating defect sites. Trap DOS calculations show lower trap densities for PVDF-TrFE compared with PVDF-HFP FETs. A similar distribution of shallow traps in as-is PVDF-HFP and with the thin Al₂O₃ layer are observed, highlighting the influence of the oxide layer on the deep-level traps. Cross-sectional STEM images show the same morphology in the bulk of the semiconductor with and without the Al₂O₃ layer.

Experimental methods and analysis

Materials

Poly(vinylidene fluoride-trifluoroethylene) (PVDF-TrFE) (75 : 25) and poly(vinylidene fluoride-*co*-hexafluoropropylene) (PVDF-HFP) ($M_w = 455\,000$) were purchased from PolyK Technologies Inc. and Sigma-Aldrich, respectively. The donor-acceptor copolymer DPP-DTT was obtained from 1-Material Inc. (Dorval, Quebec, Canada). *N,N*-Dimethylformamide (DMF) and 1,2-dichlorobenzene (DCB) (anhydrous, 98%) were sourced from Sigma-Aldrich (St. Louis, MO, USA).

Device fabrication

The device structure is bottom-gate, top-contact with a patterned Al layer as the gate electrode, the PVDF copolymers as the dielectric layer, followed by DPP-DTT as the semiconductor. Top Au contacts served as the source and drain. A 50 nm Al gate electrode was thermally evaporated onto $1'' \times 1''$ glass substrates using a patterned mask following an organic cleaning process. The dielectric solutions were prepared by dissolving PVDF-TrFE and PVDF-HFP in DMF at a concentration of 100 mg mL^{-1} , heated to $80 \text{ }^\circ\text{C}$ for 3 hours, and subsequently stirred at room temperature overnight. The solutions were diluted to 50 mg mL^{-1} for the thinner dielectric films. Under a nitrogen atmosphere, the diluted solutions were statically dispensed onto the pre-deposited Al gate and spin-coated at 1600 rpm for 60 seconds to ensure uniformity. The coated substrates were then annealed at $70 \text{ }^\circ\text{C}$ for 10 minutes in a nitrogen environment to facilitate solvent evaporation, resulting in a dielectric film thickness of approximately 45 nm.



Each substrate was placed on a hotplate under ambient conditions for the poling process. A thin Al strip was positioned on top of the PVDF-TrFE or PVDF-HFP film for the application of an electric field (40 MV m^{-1} for the thinner films) between the underlying Al gate and the external Al contact. The poling was conducted at $135 \text{ }^\circ\text{C}$, close to the Curie temperature of PVDF-TrFE and PVDF-HFP, to enhance dipole reorientation. The applied field was maintained as the substrates cooled to room temperature to preserve the dipole alignment.

DPP-DTT solution was prepared by dissolving DPP-DTT in DCB at a concentration of 5 mg mL^{-1} . The solution was subjected to sequential heating at $100 \text{ }^\circ\text{C}$ for 1 hour, $130 \text{ }^\circ\text{C}$ for 1 hour, and $145 \text{ }^\circ\text{C}$ for over 12 hours while stirring at 200 RPM. After overnight stirring at room temperature, the solution was filtered through a $0.45 \text{ }\mu\text{m}$ PTFE filter and reheated for 30 minutes before spincoating. Under a nitrogen atmosphere, $75 \text{ }\mu\text{L}$ of the DPP-DTT solution was dynamically spin-coated onto the ferroelectric layer at 900 rpm for 60 seconds. A Teflon tape was employed to confine the semiconducting film to a defined channel region during spin-coating and was removed before annealing the film at $120 \text{ }^\circ\text{C}$ for 1 hour under nitrogen. Finally, a 50 nm Au layer was thermally evaporated through a patterned mask to define the source and drain electrodes. Each substrate contained four devices with a fixed channel width of $1000 \text{ }\mu\text{m}$ and channel lengths of $50 \text{ }\mu\text{m}$, $75 \text{ }\mu\text{m}$, $100 \text{ }\mu\text{m}$, and $125 \text{ }\mu\text{m}$.

Al_2O_3 deposition

Al_2O_3 layers of varying thicknesses were deposited on top of PVDF-TrFE and PVDF-HFP using a Cambridge NanoTech S200 atomic layer deposition (ALD) system. The process employed trimethylaluminum (TMA) and deionized water (DI-water), both obtained from Sigma-Aldrich. The chamber was maintained at $150 \text{ }^\circ\text{C}$, which was the optimum temperature that preserves the integrity of the underlying dielectric layer while ensuring sufficient Al_2O_3 growth. Following sample loading, a continuous nitrogen (N_2) flow of 5 sccm was introduced into the chamber and subsequently increased to 20 sccm . The deposition was performed over 22, 33, 55, 88, and 132 ALD cycles to achieve film thicknesses of 2, 3, 5, 8, and 12 nm, respectively. Each cycle consisted of a 0.01 s pulse of DI-water, a 20 s N_2 purge, a 0.015 s pulse of TMA vapor, followed by another 20 s N_2 purge. Upon completion, the N_2 flow rate was reduced to 5 sccm .

Characterization

Current-voltage measurements were performed at room temperature using a Keithley 4200A-SCS parameter analyzer. Scanning/transmission electron microscope (STEM) imaging was performed using a probe aberration corrected ThermoFisher Spectra 300 scanning/transmission electron microscope. TEM cross-section lift out and thinning were performed in a ThermoFisher Helios Hydra UX DualBeam plasma focused ion beam (PFIB) using Xe^+ as the ion source at 30 kV accelerating voltage. Sample thinning was performed using Ar^+ as the ion source at 30 kV , with final thinning performed at 8 kV . The capacitance and conductance measurements were carried out using an HP

4284 LCR meter. The AFM images were acquired by Bruker (model: Innova).

Transistor parameters and trap density of states analysis

The carrier mobility was extracted from the current-voltage transfer characteristics in the saturation and linear region using the standard MOSFET equations. The drain current (I_D) in the saturation region is given by: $I_D(\text{sat}) = (W/2L)\mu C_i(V_{GS} - V_{th})^2$, where W and L are the channel width and length, respectively. C_i is the capacitance/area of the dielectric, V_{GS} is the gate-source voltage and V_{th} is the threshold voltage. Hence, the saturation mobility is $\mu_{\text{sat}} = (2L/WC_i)(\partial\sqrt{I_D}/\partial V_{GS})^2$. In the linear region, $I_D(\text{lin}) = (W/L)\mu C_i[(V_{GS} - V_{th})^2 - V_{DS}^2/2]$, and thus, the carrier mobility in this region is $\mu_{\text{lin}} = (L/C_i W V_{DS})(\partial I_D/\partial V_{GS})$. The subthreshold swing (SS) is given by $\text{SS} = dV_{GS}/(d \log I_{DS})$, which gives an upper limit for the interface trap states (N_{int}) such that $\text{SS} = kT \log 10/e[1 + (e^2/C_i)N_{\text{int}}]$, where k is the Boltzmann constant and T is the temperature. For state-of-the-art MOSFETs, SS is typically $70 \text{ mV decade}^{-1}$ (dec).

The contact resistance was determined using the transmission line method (TLM). The contact resistance (R_C) normalized with W is related to the total resistance (R) in the channel and is given by $RW = R_CW + L/\mu_{\text{eff}}C_i(V_{GS} - V_{th})$, where μ_{eff} is the channel mobility free from the contact resistance.

The trap DOS was analyzed using the Grünwaldt's method.^{37,38} The gate-dependent dielectric-semiconductor interface potential $V_0 = V_0(U_g)$ is first determined from:

$$\exp\left(\frac{eV_0}{kT}\right) - \frac{eV_0}{kT} - 1 = \frac{e}{kT} \frac{\epsilon_i d}{\epsilon_s l \sigma_0} \left[U_g \sigma(U_g) - \int_0^{U_g} \sigma(\tilde{U}_g) d\tilde{U}_g \right]. \quad (1)$$

In eqn (1) e , k , and T are the elementary charge, the Boltzmann constant, and the absolute temperature, respectively; $U_g = |V_{GS} - V_{FB}|$ and V_{FB} is the flat-band voltage, which is assumed to be the turn-on voltage of the transistor. ϵ_i and ϵ_s are the dielectric constant of the ferroelectric dielectric and the semiconducting layer, respectively. d is the thickness of the semiconducting layer and l is the thickness of the dielectric layer. $\sigma(U_g)$, the field effect conductivity is defined as $\sigma(U_g) = L/WI_D(U_g)/V_{DS}$. σ_0 is the conductivity at $U_g = 0$. Eqn (1) is numerically evaluated using the measured field-effect conductivity to determine the interface potential. The solutions of V_0 are used to solve for the carrier density, $p(V_0)$, using a derivative method:

$$p(V_0) = \frac{\epsilon_0 \epsilon_i^2}{\epsilon_s l^2 e} U_{GS} \left(\frac{dV_0}{dU_{GS}} \right)^{-1}, \quad (2)$$

and the trap DOS ($N(E)$) is then obtain from

$$N(E) \approx \frac{1}{e} \frac{dp(V_0)}{dV_0}. \quad (3)$$

Eqn (1)–(3) were solved in open-source Python.



Results and discussions

Dielectric thickness and poling

The semiconductor was DPP-DTT and all FETs were fabricated using a bottom-gate, top-contact geometry. Fig. 1(a) shows a schematic of the FET where both PVDF-TrFE and PVDF-HFP were used as the dielectric layer. Vertical poling was achieved by applying an electric field of either 100 MV m^{-1} for the thicker dielectric layer ($\sim 125 \text{ nm}$) and 40 MV m^{-1} for the thinner dielectric layer ($\sim 45 \text{ nm}$) at the Curie temperature, as shown in Fig. 1(b). The field is applied in a direction such that the dipole moments of the ferroelectric domains point vertically downwards. This facilitates the accumulation of hole carriers at the semiconductor–dielectric interface, which is advantageous for p-type transport in DPP-DTT. Fig. 1(c)–(e) compares the transfer and output characteristics of a vertically poled and unpoled PVDF-TrFE FET. The dotted line in the transfer curve is a linear fit where the intercept yields V_{th} . The dielectric thickness was $\sim 125 \text{ nm}$. The vertically poled PVDF-TrFE FET demonstrates three times higher saturation carrier mobility compared with the unpoled device. Earlier work highlighted that vertically poled PVDF-TrFE based organic FETs show a significant improvement in carrier mobilities and other transistor properties.³⁹ An additional lateral poling of the bottom half of the PVDF-TrFE layer to an already vertically poled film minimizes the gate leakage current in TIPS-pentacene FETs, further enhancing the carrier mobility.⁴⁰ DPC images have revealed uniform polarization domains in vertically poled PVDF-TrFE films with almost no localization of the polarization domains at the interface.³⁶ This reduces the overall polarization fluctuation in transport, enhancing carrier mobility in vertically poled FETs.

Next, the thickness of the dielectric layer was reduced to $\sim 45 \text{ nm}$ for lowering the operating voltage of the FETs. Fig. 2

shows the transfer characteristics from four such devices with PVDF-TrFE and PVDF-HFP, both unpoled and poled. In all cases, the saturation characteristics are observed by -7 V . The output characteristics of these four devices are shown in ESI† (Fig. S1). The poled PVDF-TrFE FET shows a saturation carrier mobility (μ_{sat}) of $1.24 \text{ cm}^2 \text{ V}^{-1} \text{ s}^{-1}$ compared with $0.3 \text{ cm}^2 \text{ V}^{-1} \text{ s}^{-1}$ for the unpoled device. The histogram of μ_{sat} from several poled PVDF-TrFE devices shows a mean value $> 1 \text{ cm}^2 \text{ V}^{-1} \text{ s}^{-1}$ (Fig. S2, ESI†). The SS value is also lowered for the poled PVDF-TrFE FET. In contrast, poling does not seem to enhance the PVDF-HFP FET performance. There is almost no difference in μ_{sat} ($\sim 0.28 \text{ cm}^2 \text{ V}^{-1} \text{ s}^{-1}$) between poled and unpoled PVDF-HFP. Since the overall ferroelectricity in PVDF-HFP is lower than in PVDF-TrFE, external electrical field poling seems to have almost no impact in orienting the polarization domains.

Although the value of SS is an indicator of the trap density, it mainly probes the deeper bandgap states as the quasi-Fermi level is far from the band-edge in the subthreshold region. The trap DOS estimated from the Grünwald's method allows an estimation of the shallow trap states. Polymeric semiconductors are usually described by a Gaussian distribution of traps as sketched in Fig. 3(a). Since disorder in these systems results in electron localization, it has a Gaussian distribution. The mobility edge in a polymeric semiconductor acts as the band edge in a crystalline material. In addition to the shallow trap states, localized deep states are present. If the Fermi level is located in the region of the localized states (near the green peaks in Fig. 3(a)), conductivity at high temperatures can arise from thermal excitation of carriers across the mobility gap in addition to hopping of carriers due to the shallow trap states. We estimate the trap DOS using the Grünwald's method from

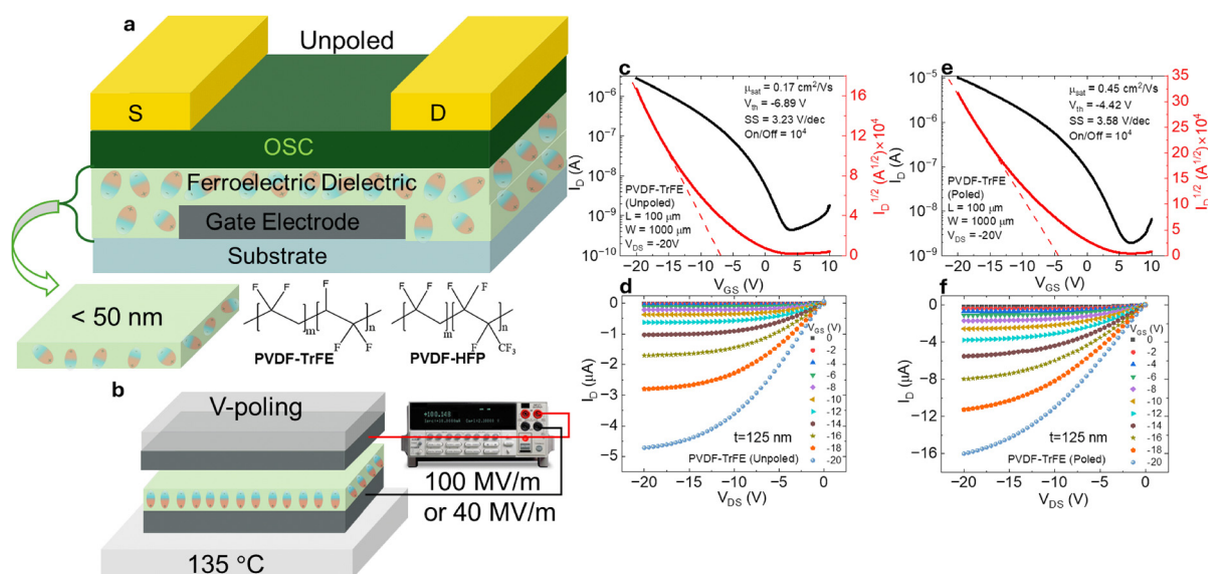


Fig. 1 (a) Device schematic where the dielectric thickness was reduced from 125 nm to below 50 nm. Chemical structure of PVDF-TrFE and PVDF-HFP. (b) Vertical poling of the dielectric layer was carried out by applying an electric field at a temperature of $\sim 135 \text{ }^\circ\text{C}$. (c) and (d) Transfer and output characteristics of an unpoled PVDF-TrFE/DPP-DTT FET, respectively. (e) and (f) Transfer and output characteristics of a vertically poled PVDF-TrFE/DPP-DTT FET, respectively. The dielectric thickness was $\sim 125 \text{ nm}$ for both poled and unpoled devices.



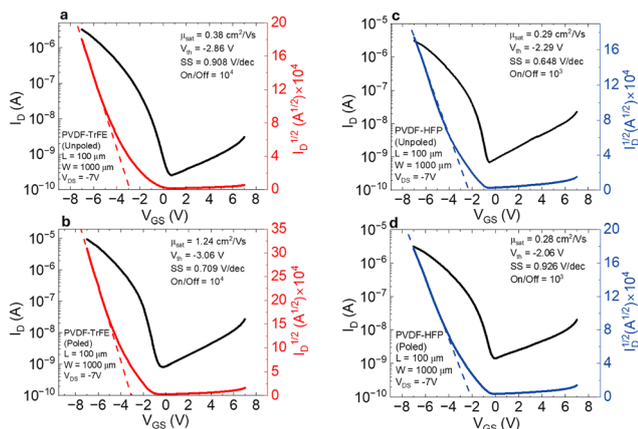


Fig. 2 Transfer characteristics of DPP-DTT FETs with a dielectric thickness of 45 nm. (a) Unpoled PVDF-TrFE, (b) poled PVDF-TrFE, (c) unpoled PVDF-HFP, and (d) poled PVDF-HFP.

the valence band edge (E_V), which is the HOMO level, for PVDF-TrFE and PVDF-HFP FETs, as illustrated in Fig. 3(b) and (c). The trap DOS is almost an order of magnitude higher near the band edge for the thicker unpoled (125 nm) PVDF-TrFE FET compared with the thinner (45 nm) unpoled one. Additionally, the trap DOS for the vertically poled PVDF-TrFE FET is lower compared with the unpoled one. On the other hand, unpoled and poled PVDF-HFP (of thickness 45 nm) FETs show similar trap DOS. It should be noted that vertically poled PVDF-TrFE displays a very different morphology with distinct domains compared with its unpoled counterpart.³⁹ Hence, it is not surprising that their trap DOS differ. We further estimate the interface trap density (D_{it}) from capacitance and conductance *versus* voltage measurements from PVDF-TrFE based metal-insulator-semiconductor (MIS) diodes for a comparison of the unpoled and poled layers. The D_{it} values are determined from the loss as the occupancy of the interface trap states change with the gate bias.⁴¹ Using a continuum of state model,⁴² the D_{it} values obtained at the flat-band voltage are $2.3 \times 10^{12} \text{ eV}^{-1} \text{ cm}^{-2}$ and $3.5 \times 10^{12} \text{ eV}^{-1} \text{ cm}^{-2}$ for the poled and unpoled PVDF-TrFE MIS diodes, respectively. See ESI† for details (Fig. S8 and S9).

The above results demonstrate that vertical poling has a dramatic effect on PVDF-TrFE FETs but not on PVDF-HFP FETs. The improved carrier mobility in poled PVDF-TrFE FET may be rationalized in terms of the lower trap DOS. Our results further suggest that poling does not change the inherent fraction of the crystalline β phase to the α phase; it merely orients the ferroelectric domains. Since the extent of the β phase is much lower in PVDF-HFP compared with PVDF-TrFE, the external poling field has a minimal effect. We next look at the effect of adding ALD grown ultrathin Al_2O_3 layer to the interface of both PVDF-TrFE and PVDF-HFP devices.

Addition of an ultrathin Al_2O_3 layer

Al_2O_3 as a dielectric material has been broadly used in both organic and inorganic transistors. Depending on the growth method of Al_2O_3 , κ can vary between 7–10.¹ ALD yields extremely smooth and high-quality films in short cycle times.⁴³ Since the κ values of Al_2O_3 and PVDF copolymers are almost identical, the addition of Al_2O_3 serves as an excellent platform to modulate the trap states at the semiconductor–dielectric interface and tune the polarization fluctuation of the polymer ferroelectric without affecting the overall κ of the dielectric stack. With ALD growth temperatures being beyond the Curie temperature of the PVDF copolymers, all films were unpoled. Al_2O_3 thicknesses between 2 nm and 12 nm were deposited on 45 nm thick PVDF-TrFE or PVDF-HFP prior to the deposition of DPP-DTT. Metal-insulator-metal capacitors were also fabricated, where the overall capacitance/area varied between 94–116 nF cm^{-2} for 2–12 nm Al_2O_3 on the PVDF copolymers, respectively.

The current–voltage characteristics of PVDF-TrFE and PVDF-HFP based FETs with 5 nm of Al_2O_3 are compared in Fig. 4. Overall, the PVDF-TrFE FET shows lower carrier mobility and similar SS compared with the unpoled (without Al_2O_3) devices. Other thicknesses of the Al_2O_3 layer with PVDF-TrFE also do not improve the FET performance (see ESI†). In contrast, significant improvements in SS and on/off ratio are observed for the PVDF-HFP FET with the 5 nm Al_2O_3 (Fig. 4(d)). To understand the effect of Al_2O_3 on PVDF-HFP, we tested several other thicknesses of Al_2O_3 layer between 2 nm and 12 nm. Current–voltage characteristics of other PVDF-HFP devices with varying

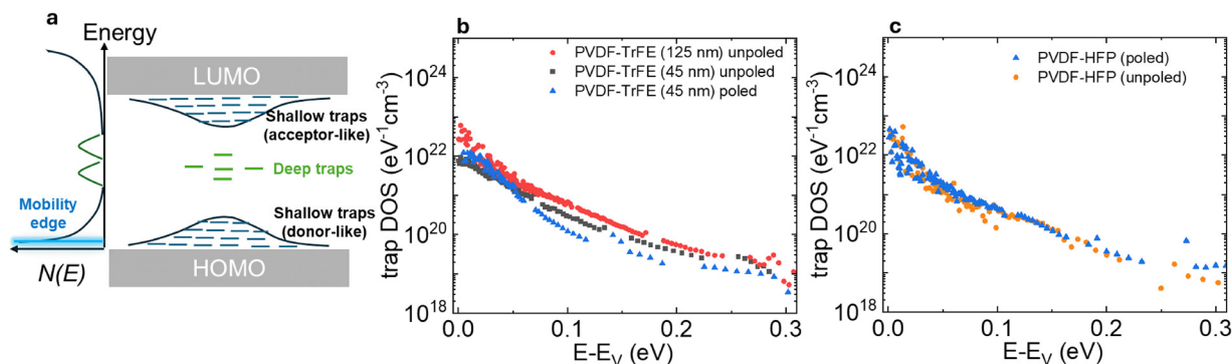


Fig. 3 Trap DOS in PVDF-TrFE and PVDF-HFP based DPP-DTT FETs. (a) Schematic DOS ($N(E)$) and trap states in polymeric semiconductors. Both deep and shallow traps are present. (b) Trap DOS in DPP-DTT FETs for two thicknesses of the PVDF-TrFE layer including a poled layer. (c) Trap DOS in DTT FETs for poled and unpoled PVDF-HFP layers.



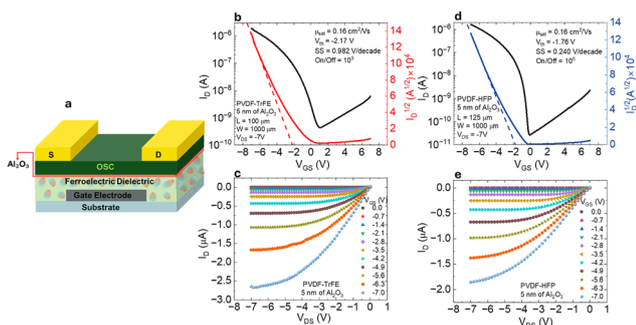


Fig. 4 FET characteristics with 5 nm Al_2O_3 . (a) Device geometry with the addition of Al_2O_3 layer. (b) and (c) Transfer and output characteristics of a DDP-DTT/PVDF-TrFE FET, respectively. (d) and (e) Transfer and output characteristics of a DDP-DTT/PVDF-HFP FET, respectively.

Al_2O_3 are shown in ESI†. In each case we find the SS value to be lower than 300 mV dec^{-1} .

Adding between 2 nm and 12 nm of Al_2O_3 on PVDF-HFP has a similar effect on the FET performance, although the carrier mobility decreases slightly for the 12 nm oxide layer. We discuss this reduction in mobility in terms of the contact resistance. The transfer characteristics of four PVDF-HFP FETs with varying Al_2O_3 thickness between 2 nm to 12 nm are plotted in Fig. 5(a). A histogram of the carrier mobilities and SS for the Al_2O_3 coated devices are shown in Fig. 5(b) and (c), respectively, for 30–40 devices. A clear reduction in the SS values is observed compared with the non- Al_2O_3 coated PVDF-HFP FETs. The average value of $\text{SS} = 400 \text{ mV dec}^{-1}$ for Al_2O_3 on PVDF-HFP compared with $\sim 800 \text{ mV dec}^{-1}$ for the non- Al_2O_3 coated devices (see Fig. S5, ESI†). It should also be pointed out that the average values are from all different thickness of Al_2O_3 , where the 12 nm thickness overall shows a slightly higher value of SS. These data further suggest that the optimum thickness of the Al_2O_3 layer is between 2 nm and 8 nm for a significant reduction in SS. The transfer curves for the 8 nm Al_2O_3 device are shown for different values of V_{DS} in Fig. 5(d), and the linear fits for extracting the SS values are shown in Fig. 5(e). The high off current with increasing V_{DS} is expected due to the high shallow trap DOS. At the lowest V_{DS} of -0.7 V , $\text{SS} = 75 \text{ mV dec}^{-1}$. A similar trend is also seen for the 2 nm Al_2O_3 on PVDF-HFP FET (see ESI†). This is a clear improvement over the non- Al_2O_3 coated PVDF-HFP devices which show $\text{SS} > 300 \text{ mV}$ in the linear region (Fig. S6, ESI†). With the addition of a thin Al_2O_3 layer, we observe some of the lowest values of SS reported for FETs using a polymer semiconductor and a solution processable polymer dielectric.

As a quick summary, ALD grown Al_2O_3 on PVDF-HFP significantly improves SS but slightly reduces the carrier mobility in DPP-DTT FETs. PVDF-TrFE based DPP-DTT FETs, however, show no improvement in SS and there is a further degradation in carrier mobility. Before discussing the differences in SS between the two PVDF systems, we first look at the parameters that govern carrier mobility. The contact resistance of these devices is significant. A comparison of the carrier mobility in the saturation and linear region of a PVDF-HFP FET with 12 nm of Al_2O_3 as a function of the gate voltage in Fig. 6(a) clearly

shows a large difference at higher gate voltages. Ideally, the two mobility curves should overlap. This discrepancy mainly arises due to the contact resistance. The bulk resistance (R_{Bulk}) contributes to the contact resistance (R_{C}) due to charge transport within the organic semiconductor, extending from the contact to the channel, as illustrated in Fig. 6(b). Charge carriers move from the source contact to the accumulation layer at the semiconductor–dielectric interface before reaching the drain contact for extraction. Consequently, both metal–semiconductor interface resistance, arising from the resistance associated with the charge-injection barrier, R_{int} , and R_{Bulk} play significant roles in determining R_{C} . Therefore, minimizing R_{C} requires reducing the contributions of both R_{int} and R_{Bulk} .^{15,44}

Using the TLM method as discussed in section: Experimental methods and analysis, we determine R_{C} for a PVDF-HFP FET without Al_2O_3 and with 12 nm of Al_2O_3 (Fig. 6(c) and (d)). The contact resistance increases by almost a factor of 14 when 12 nm of Al_2O_3 is present. Without Al_2O_3 , the contact resistance of the PVDF-HFP FET is approximately $7 \text{ k}\Omega \text{ cm}$. A similar effect is seen for PVDF-TrFE. Adding Al_2O_3 on PVDF-TrFE also increases R_{C} compared with as-is unpoled PVDF-TrFE FET, which is the main reason why we see a reduction in μ_{sat} for Al_2O_3 on PVDF-TrFE devices. Being top contact devices, adding a barrier layer or self-assembled monolayer for reducing the Schottky barrier, which reduces R_{int} , is not straightforward. Future direction involving organic ices as negative tone resists^{45,46} that leave behind thin insulating residues in a modified electron-beam lithography is being developed for deposition under the contacts for reducing R_{int} in top contact organic FETs.

Next, we discuss the impact of the Al_2O_3 layer on SS. There is a clear benefit to PVDF-HFP FETs, as seen in Fig. 5, but no significant improvement is observed in PVDF-TrFE FETs. Since the SS values are dictated by the trap states, we look at the trends for both trap DOS as well as the N_{int} values. Table 1 lists the SS, the normalized SS values, which is the product of SS and C_{i} , and N_{int} for DPP-DTT FETs with PVDF-TrFE and PVDF-HFP as the dielectric layer. All FETs have the same L and W values. The SS values noted here are from the transfer characteristics in the saturation region ($V_{\text{DS}} = -7 \text{ V}$). Lowest normalized SS is seen for the PVDF-HFP FETs with Al_2O_3 , which further indicates low N_{it} values. Table S1 in ESI† highlights benchmark SS values from other organic FETs in the literature.

We look more closely into the ALD growth process of Al_2O_3 on PVDF-TrFE and PVDF-HFP to see how that may affect the two copolymers differently and compare the trap DOS near the HOMO level of the semiconductor with and without Al_2O_3 . A schematic of the first ALD process for the growth of Al_2O_3 is shown in ESI† (Fig. S10), where the water cycle is followed by TMA, which is then evacuated, followed by re-introduction of water. The morphology of PVDF-TrFE and PVDF-HFP is quite different. The higher fraction of the β phase in PVDF-TrFE enhances its crystallinity compared with PVDF-HFP, which is more amorphous. Moreover, due to the HFP group, PVDF-HFP is more hydrophobic compared with PVDF-TrFE.³⁴ The ALD process typically relies on surface hydroxyl groups to facilitate



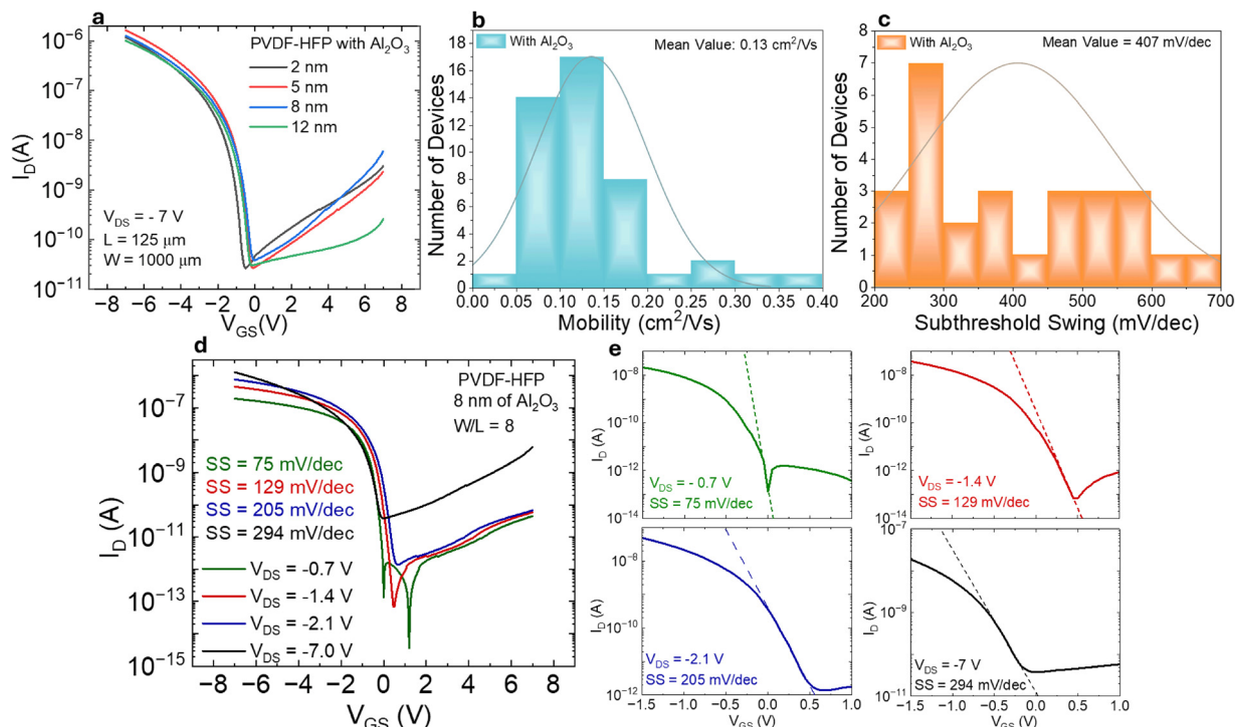


Fig. 5 Subthreshold characteristics with Al_2O_3 and PVDF-HFP. (a) Transfer current–voltage characteristics of PVDF-HFP devices with varying thicknesses of Al_2O_3 . (b) Histogram of μ_{sat} for PVDF-HFP FETs with Al_2O_3 . (c) Histogram of SS for PVDF-HFP FETs with Al_2O_3 . (d) Transfer current–voltage characteristics for varying values of V_{DS} for the PVDF-HFP FET with 8 nm of Al_2O_3 . (e) Linear fits for obtaining SS for the transfer characteristics shown in (d).

the reaction. Since both PVDF copolymers lack surface O–H groups, we hypothesize that the reaction of the water pulse in the ALD cycle might induce partial surface modification,

generating potential reaction sites for TMA. Further, the first pulse of water may hydroxylate the surface of amorphous PVDF-HFP more than crystalline PVDF-TrFE, allowing oxygen to diffuse. Once TMA reacts on the PVDF-HFP surface and with an additional water cycle, further oxidation may be possible, allowing more oxygen to diffuse. The last stage of the first ALD cycle for the growth of Al_2O_3 on PVDF-HFP and PVDF-TrFE is shown in Fig. 7(a) and (b), highlighting the diffusion of oxygen in PVDF-HFP.

HAADF-STEM (high-angle annular dark-field-STEM) images and STEM-EDS (energy dispersive spectroscopy) maps confirm

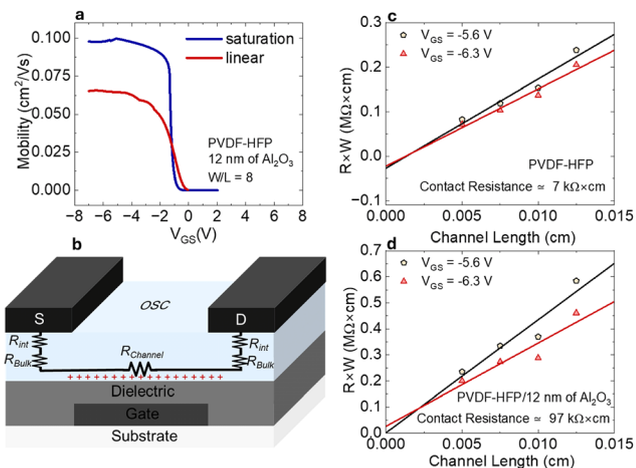


Fig. 6 Contact resistance in PVDF-HFP FETs. (a) Comparison of the linear and saturation carrier mobilities as a function of the gate voltage in a 12 nm Al_2O_3 on PVDF-HFP DPP-DTT FET. (b) Schematic of an organic FET showing the origin of contact resistance (R_{C}). Both R_{int} and R_{bulk} contribute to R_{C} . R_{C} estimation using the transmission line model for (c) PVDF-HFP/DPP-DTT FET and (d) PVDF-HFP/12 nm Al_2O_3 /DPP-DTT FET. The product of the total resistance and width versus channel length for two different gate-source voltages are shown. The intersection of the two straight lines representing the gate biases on the y-axis yields R_{C} .

Table 1 The subthreshold swing (SS), normalized SS, and the interface trap density (N_{it}) for DPP-DTT FETs with PVDF-TrFE and PVDF-HFP as the dielectric layer. The second column shows the nature of the ferroelectric layer, which is unpoled, poled, or with an Al_2O_3 layer, where the thickness is quoted in the bracket

Dielectric	Al_2O_3 Thickness (nm)	SS (V dec ⁻¹)	Normalized SS (nFV dec ⁻¹ cm ⁻²)	N_{it} (cm ⁻² eV ⁻¹)
PVDF-TrFE	Unpoled (0)	1.12 ± 0.05	110.8	1.11×10^{13}
	Poled (0)	0.70 ± 0.02	69.3	6.81×10^{12}
	Al_2O_3 (2)	1.31 ± 0.07	129.7	1.31×10^{13}
	Al_2O_3 (3)	1.45 ± 0.05	143.5	1.46×10^{13}
	Al_2O_3 (5)	0.98 ± 0.04	97	9.66×10^{12}
PVDF-HFP	Unpoled (0)	0.59 ± 0.02	55.4	5.27×10^{12}
	Poled (0)	0.67 ± 0.02	62.9	6.08×10^{12}
	Al_2O_3 (2)	0.26 ± 0.01	24.4	1.99×10^{12}
	Al_2O_3 (5)	0.24 ± 0.01	22.5	1.79×10^{12}
	Al_2O_3 (8)	0.29 ± 0.01	27.2	2.29×10^{12}
	Al_2O_3 (12)	0.27 ± 0.01	25.3	2.09×10^{12}



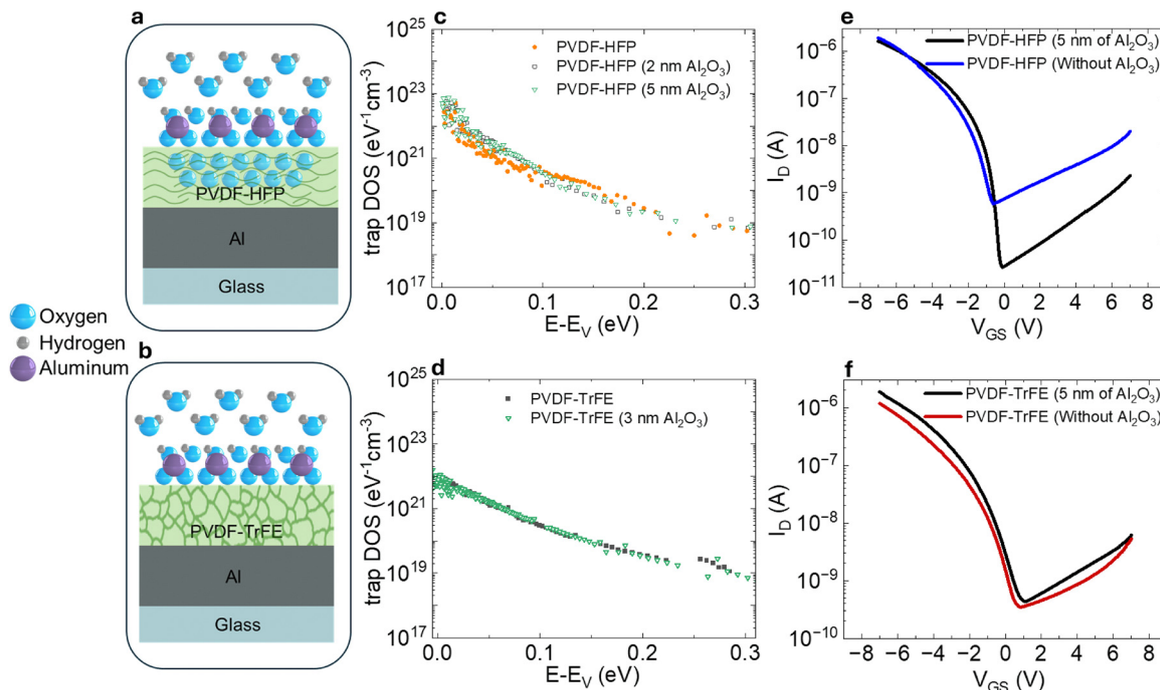


Fig. 7 ALD growth of Al_2O_3 on PVDF-HFP and PVDF-TrFE. (a) and (b) The final stage of the first cycle of the ALD process for the deposition of Al_2O_3 on PVDF-HFP and PVDF-TrFE, respectively. The green domains in PVDF-TrFE represent higher crystallinity. (c) and (d) Trap DOS for PVDF-HFP and PVDF-TrFE FETs, respectively, with and without Al_2O_3 . (e) and (f) Transfer characteristics of representative PVDF-HFP and PVDF-TrFE based FETs, respectively, with and without Al_2O_3 .

the presence of Al_2O_3 on PVDF-HFP samples (see Fig. S11, ESI[†]). While the bulk of the polymer film is similar with and without the Al_2O_3 layer, details of the interfacial structure are obscured due to holes at either the polymer–oxide (Fig. S11, ESI[†]) or polymer–electrode (Fig. S12, ESI[†]) interfaces, depending on sample thickness. These holes result from ion beam damage during the thinning process (Fig. S11, ESI[†]).

AFM images (included in ESI[†]) from PVDF-TrFE and PVDF-HFP films coated with 12 nm Al_2O_3 show some changes in morphology compared with their pristine counterpart. The changes are more significant in PVDF-TrFE where the large ferroelectric domains are somewhat disrupted in the presence of Al_2O_3 .

A surprising result is that the trap DOS, as shown in Fig. 7(c) and (d), with and without Al_2O_3 , are almost identical. This suggests that the shallow trap distribution hardly changes with the deposition of Al_2O_3 . Hence, the improvement in the FET characteristics with Al_2O_3 on PVDF-HFP arises from a modification of the deep level traps. The transfer curves of representative FETs with and without Al_2O_3 on PVDF-HFP and PVDF-TrFE are shown in Fig. 7(e) and (f). As discussed earlier, there is hardly any change in SS for PVDF-TrFE FETs with Al_2O_3 . The improved SS with Al_2O_3 on PVDF-HFP most likely then arises from filling of the interface states with oxygen, which influences the deep level discrete traps (Fig. 3(a)). Such a dependence of the interface states and the deep level states have been observed in other semiconductor–insulator interfaces such as ALD grown Al_2O_3 and GaN.⁴⁷ Our results, therefore, warrant other techniques such as deep level transient spectroscopy for a

complete picture of the spatial and energy profile of the trap states.⁴⁸ Such characterization strategies in the future will allow further improvement of organic FETs when modulating the semiconductor–dielectric interface.

It is conceivable that the oxide layer could be replaced with other thin insulating layers as recently demonstrated with negative-tone ethanol-based ice resist⁴⁹ or a self-assembled monolayer. Such strategies in the future will ensure cost-effective means for fabricating the entire device along with improved performance.

Conclusions

This work demonstrates that external electric field poling remains the most effective approach for reducing polarization fluctuations and enhancing the performance of polymer ferroelectric transistors based on PVDF-TrFE. Further, by reducing the thickness of the PVDF-TrFE film from 125 nm to 45 nm, the operating voltage of the FETs decreases below 8 V. Under optimized conditions, the saturation carrier mobility exceeds $1 \text{ cm}^2 \text{ V}^{-1} \text{ s}^{-1}$. In contrast, PVDF-HFP, with lower ferroelectric properties compared with PVDF-TrFE, does not exhibit improved FET properties with external poling.

The incorporation of an ultrathin ALD grown Al_2O_3 interfacial layer plays a critical role in modulating the semiconductor–dielectric interface with PVDF-HFP. SS as low as 75 mV dec^{-1} in the linear region is observed for PVDF-HFP based FETs with an interfacial Al_2O_3 layer. No significant improvement in SS or other FET properties are observed with



PVDF-TrFE. The differences between the two copolymers arise mainly from their specific morphology. The more amorphous nature of PVDF-HFP compared with PVDF-TrFE allows the penetration of oxygen within the film during the growth of Al₂O₃, resulting in passivating the interface traps and further influencing the deep level discrete traps in the polymer semiconductor. Our results suggest that Al₂O₃ treatment could be beneficial for other high κ organic dielectrics where passivation of interface trap states may be required.

Author contributions

A. Ghobadi and S. Guha conceived the work. A. Ghobadi was involved with the fabrication of devices, conducting electrical measurements, and analysing the data. S. Guha was involved with measurements and the analysis of the data. S. Gangopadhyay helped with the analysis and process development for device fabrication. J. Mathai performed ALD growth and helped in the setting up of electrical and AFM measurements. T. Kallaos performed the trap DOS calculations. I. M. Karunaratne, D. M. Gamachchi, and A. C. Meng carried out the lift-off processes and TEM measurements. The manuscript was written by S. Guha and A. Ghobadi with contributions from all authors. All authors have given approval to the final version of the manuscript.

Data availability

The data supporting the findings of this study are available within the article and/or have been included as part of its ESI.†

Conflicts of interest

There are no conflicts to declare.

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Notes and references

- B. Wang, W. Huang, L. Chi, M. Al-Hashimi, T. J. Marks and A. Facchetti, *Chem. Rev.*, 2018, **118**, 5690–5754.
- J. Veres, S. D. Ogier, S. W. Leeming, D. C. Cupertino and S. Mohialdin Khaffaf, *Adv. Funct. Mater.*, 2003, **13**, 199–204.
- T. Richards, M. Bird and H. Sirringhaus, *J. Chem. Phys.*, 2008, **128**, 234905.
- H. Chen, W. Zhang, M. Li, G. He and X. Guo, *Chem. Rev.*, 2020, **120**, 2879–2949.
- S. P. Senanayak, S. Guha and K. S. Narayan, *Phys. Rev. B: Condens. Matter Mater. Phys.*, 2012, **85**, 115311.
- A. Laudari and S. Guha, *J. Appl. Phys.*, 2015, **117**, 105501.
- F. Yang, L. Sun, J. Han, B. Li, X. Yu, X. Zhang, X. Ren and W. Hu, *ACS Appl. Mater. Interfaces*, 2018, **10**, 25871–25877.
- T. Wollandt, F. Letzkus, J. N. Burghartz and H. Klauk, *Adv. Electron. Mater.*, 2024, **10**, 2300841.
- H. Klauk, U. Zschieschang, J. Pflaum and M. Halik, *Nature*, 2007, **445**, 745–748.
- L. Torsi, A. Dodabalapur, L. J. Rothberg, A. W. P. Fung and H. E. Katz, *Science*, 1996, **272**, 1462–1464.
- N. F. Mott and E. A. Davis, *Electronic processes in non-crystalline materials*, Clarendon Press, Oxford University Press, Oxford, New York, 1979.
- I. N. Hulea, S. Fratini, H. Xie, C. L. Mulder, N. N. Iossad, G. Rastelli, S. Ciuchi and A. F. Morpurgo, *Nat. Mater.*, 2006, **5**, 982–986.
- A. Laudari and S. Guha, *J. Appl. Phys.*, 2019, **125**, 035501.
- C. Liu, Y. Xu and Y.-Y. Noh, *Mater. Today*, 2015, **18**, 79–96.
- M. Waldrip, O. D. Jurchescu, D. J. Gundlach and E. G. Bittle, *Adv. Funct. Mater.*, 2020, **30**, 1904576.
- J. W. Borchert, B. Peng, F. Letzkus, J. N. Burghartz, P. K. L. Chan, K. Zojer, S. Ludwigs and H. Klauk, *Nat. Commun.*, 2019, **10**, 1119.
- J. Zeng, D. He, J. Qiao, Y. Li, L. Sun, W. Li, J. Xie, S. Gao, L. Pan, P. Wang, Y. Xu, Y. Li, H. Qiu, Y. Shi, J.-B. Xu, W. Ji and X. Wang, *Nat. Commun.*, 2023, **14**, 324.
- A. Bilgaiyan, M. Abiko, K. Watanabe and M. Mizukami, *ACS Appl. Electron. Mater.*, 2025, **7**, 2583–2592.
- Y. Fu, J. Zhu, Y. Sun, S. Sun, K. Tie, J. Qi, Y. Wang, Z. Wang, Y. Hu, S. Ding, R. Huang, Z. Gong, Y. Huang, X. Chen, L. Li and W. Hu, *ACS Nano*, 2023, **17**, 15044–15052.
- A. Ghobadi, C. J. Mathai, J. Cook, G. Bian, S. Attar, M. Al-Hashimi, S. Gangopadhyay and S. Guha, *Adv. Electron. Mater.*, 2005, **11**, 2400503.
- S. J. Kang, Y. J. Park, I. Bae, K. J. Kim, H.-C. Kim, S. Bauer, E. L. Thomas and C. Park, *Adv. Funct. Mater.*, 2009, **19**, 2812–2818.
- S. Das and J. Appenzeller, *Nano Lett.*, 2011, **11**, 4003–4007.
- S. K. Hwang, I. Bae, S. M. Cho, R. H. Kim, H. J. Jung and C. Park, *Adv. Funct. Mater.*, 2013, **23**, 5484–5493.
- C.-F. Lin, Y.-S. Wu, W.-C. Chen, B.-H. Lin, Y.-C. Lin, C.-C. Kuo and W.-C. Chen, *ACS Appl. Electron. Mater.*, 2025, **7**, 1068–1079.
- R. Peng, B. Zhang, G. Dong, Y. Wang, G. Yang, J. Zhang, B. Peng, Y. Zhao and M. Liu, *Adv. Funct. Mater.*, 2024, **34**, 2316519.
- L. Lu, W. Ding, J. Liu and B. Yang, *Nano Energy*, 2020, **78**, 105251.
- C. Ribeiro, C. M. Costa, D. M. Correia, J. Nunes-Pereira, J. Oliveira, P. Martins, R. Gonçalves, V. F. Cardoso and S. Lanceros-Méndez, *Nat. Protoc.*, 2018, **13**, 681–704.
- X. Wang, W. Lu, P. Wei, Z. Qin, N. Qiao, X. Qin, M. Zhang, Y. Zhu, L. Bu and G. Lu, *ACS Appl. Mater. Interfaces*, 2022, **14**, 48948–48959.
- X. Wang, S. Yang, Z. Qin, B. Hu, L. Bu and G. Lu, *Adv. Mater.*, 2023, **35**, 2303699.
- A. J. Lovinger, D. D. Davis, R. E. Cais and J. M. Kometani, *Macromolecules*, 1986, **19**, 1491–1494.



- 31 G. Knotts, A. Bhaumik, K. Ghosh and S. Guha, *Appl. Phys. Lett.*, 2014, **104**, 233301.
- 32 R. C. G. Naber, C. Tanase, P. W. M. Blom, G. H. Gelinck, A. W. Marsman, F. J. Touwslager, S. Setayesh and D. M. de Leeuw, *Nat. Mater.*, 2005, **4**, 243–248.
- 33 J. Roy, B. Joseph, A. Ashraf K and R. B. Gangineni, *Thin Solid Films*, 2023, **781**, 139981.
- 34 D. M. Correia, J. Nunes-Pereira, D. Alikin, A. L. Kholkin, S. A. C. Carabineiro, L. Rebouta, M. S. Rodrigues, F. Vaz, C. M. Costa and S. Lanceros-Méndez, *Polymer*, 2019, **169**, 138–147.
- 35 E. Restuccia, A. Ghobadi and S. Guha, *Adv. Mater. Interfaces*, 2025, 2401035.
- 36 A. Ghobadi, S. S. Niavol, E. Restuccia, A. C. Meng and S. Guha, *Appl. Phys. Lett.*, 2024, **124**, 153301.
- 37 M. Grünewald, P. Thomas and D. Würtz, *Phys. Status Solidi B*, 1980, **100**, K139–K143.
- 38 W. L. Kalb and B. Batlogg, *Phys. Rev. B: Condens. Matter Mater. Phys.*, 2010, **81**, 035327.
- 39 A. Laudari, A. R. Mazza, A. Daykin, S. Khanra, K. Ghosh, F. Cummings, T. Muller, P. F. Miceli and S. Guha, *Phys. Rev. Appl.*, 2018, **10**, 014011.
- 40 A. Laudari, A. Pickett, F. Shahedipour-Sandvik, K. Hogan, J. E. Anthony, X. He and S. Guha, *Adv. Mater. Interfaces*, 2019, **6**, 1801787.
- 41 E. H. Nicollian and J. R. Brews, *MOS (Metal Oxide Semiconductor) Physics and Technology*, Wiley, New York, 2003.
- 42 M. Yun, S. Gangopadhyay, M. Bai, H. Taub, M. Arif and S. Guha, *Org. Electron.*, 2007, **8**, 591–600.
- 43 Y.-S. Min, Y. J. Cho and C. S. Hwang, *Chem. Mater.*, 2005, **17**, 626–631.
- 44 P. Darmawan, T. Minari, Y. Xu, S.-L. Li, H. Song, M. Chan and K. Tsukagoshi, *Adv. Funct. Mater.*, 2012, **22**, 4577–4583.
- 45 A. Elsukova, A. Han, D. Zhao and M. Beleggia, *Nano Lett.*, 2018, **18**, 7576–7582.
- 46 G. Yao, D. Zhao, Y. Hong, S. Wu, D. Liu and M. Qiu, *Nanoscale*, 2020, **12**, 22473–22477.
- 47 C. M. Jackson, A. R. Arehart, E. Cinkilic, B. McSkimming, J. S. Speck and S. A. Ringel, *J. Appl. Phys.*, 2013, **113**, 204505.
- 48 N. M. Johnson, D. J. Bartelink, R. B. Gold and J. F. Gibbons, *J. Appl. Phys.*, 1979, **50**, 4828–4833.
- 49 D. A. Chiaro, T. J. Hager, K. T. Renshaw, B. M. Moore, A. Ghobadi, R. I. Haque, A. Han, B. M. Broderick, S. Guha and G. M. King, *Nano Lett.*, 2025, **25**, 7107–7114.

