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3 Supporting Information

ABSTRACT: Monolayers of six alkylphosphonic acids ranging from C₆ to C₁₈ were prepared by vacuum evaporation and incorporated into low-voltage organic field-effect transistors based on dinaphtho[2,3-b:2′,3′-f]thieno[3,2-b]thiophene (DNTT). Similar to solution-assembled monolayers, the molecular order for vacuum-deposited monolayers improved with increasing length of the aliphatic tail. At the same time, Fourier transform infrared (FTIR) measurements suggested lower molecular coverage for longer phosphonic acids. The comparison of FTIR and vibration frequencies calculated by density functional theory indicated that monodentate bonding does not occur for any phosphonic acid. All monolayers exhibited low surface energy of ~17.5 mJ/m² with a dominating Lifshitz–van der Waals component. Their surface roughness was comparable, while the nanomechanical properties were varied but not correlated to the length of the molecule. However, large improvement in transistor performance was observed with increasing length of the aliphatic tail. Upon going from C₆ to C₁₈, the mean threshold voltage decreased from ~−1.37 to ~−1.24 V, the field-effect mobility increased from 0.03 to 0.33 cm²/(V·s), the off-current decreased from ~8 × 10⁻¹¹ to ~3 × 10⁻¹³ A, and for transistors with L = 30 μm the on-current increased from ~3 × 10⁻⁶ to ~2 × 10⁻⁶ A, and the on/off-current ratio increased from ~3 × 10⁸ to ~4 × 10⁹. Similarly, transistors with longer phosphonic acids exhibited much better air and bias-stress stability. The achieved transistor performance opens up a completely “dry” fabrication route for ultrathin dielectrics and low-voltage organic transistors.

KEYWORDS: organic field-effect transistors, alkylphosphonic acids, monolayers, DNTT, bias stress

1. INTRODUCTION

The focused improvement of organic field-effect transistors (OFETs) has allowed a whole host of novel demonstrations including radio frequency identification tags,¹,² analog and digital circuits,³,⁴ active matrix displays,⁵ and various sensor systems.⁶,⁷,⁸ However, when the application of OFETs in forthcoming areas such as wearable or disposable electronics is considered, low power consumption and low-voltage operation are necessary features, especially for applications powered by batteries or energy-harvesting devices.

A common approach to achieving low-voltage transistor operation is to increase the gate dielectric capacitance by choosing thin layers and/or materials with high relative permittivity (high-κ). To date, approaches have included single layers of inorganic or organic high-κ materials, inorganic/organic bilayers, and organic/inorganic composites.⁹,¹⁰ Reduction in the gate dielectric thickness to 10 nm or less typically involves a bilayer, where a medium-κ to high-κ inorganic layer is covered with an organic monolayer whose function is to suppress the leakage current, inhibit the surface —OH groups, and reduce the energy of the dielectric surface.¹¹–¹⁷ In such a case, the transistor operating voltage can be as low as 1.5 V, while the transistor is in the off state at 0 V.¹⁸–²⁰

As the thickness of the dielectric is reduced, small variations in its thickness lead to more pronounced variations in its capacitance. Consequently, procedures that inherently lend
themselves to good layer uniformity (atomic layer deposition) or self-limit the layer thickness (oxidation or use of monolayers) are advantageous. Aluminum oxide (AlOₓ) functionalized with aliphaphosphonic acids (CₙPA) is an established bilayer dielectric for low-voltage organic transistors. Such transistors have a bottom-gate structure where the aluminum oxide is commonly prepared by oxidation of the aluminum gate electrode and the assembly of the organic monolayer is performed in solvent-based solutions. A dry route to monolayer assembly has also been demonstrated for n-octylphosphonic acids (C₉PA). In such a case, several monolayers of C₉PA were thermally evaporated in vacuum, followed by thermal desorption of all molecules that were not chemically bonded to AlOₓ. In such a technique, nonuniformity in the as-deposited C₉PA thickness is eliminated during the second step of thermal desorption, leaving monolayer formation across the substrate. Thermal desorption provides an additional benefit of annealing that improves both the monolayer structure and transistor performance.

In this paper we report bottom-gate OFETs based on various AlOₓ/C₉PA dielectrics and an air-stable,25 organic semiconductor, dinaphtho[2,3-b:2′,3′-f]thieno[3,2-b]thiophene (DNTT). This was prompted by previous research on the corresponding transistor behavior. The achieved transistor performance opens up a completely new fabrication route for OFETs and metal (MIM) structures followed the same fabrication procedure. The captured force curve is used to calculate the DMT modulus, deformation, energy dissipation, and force of adhesion. To obtain the Young’s modulus, the retract curve is fitted by use of the Derjaguin–Muller–Toporov (DMT) model, for that reason called DMT modulus.

To probe liquid–surface interactions with maximum resolution, contact angles (at 22 °C) of small drops (four on each substrate) of diiodomethane (DIM, >99%, surface tension γ₆ ≈ 48.7 mN/m at 18.8 °C, ~1 μL), 1,2-ethanediol or ethylene glycol (EG, >99%, γ₇ = 47.7 mN/m at 18.8 °C, ~1 μL), and filtered water (FW, γ₈ = 73.4 mN/m at 18.8 °C, ~2 μL) placed on horizontal surfaces were measured by use of a contact angle goniometer (Kruus DSA30, Germany). Advancing angles (θₐ) were obtained for both “left” and “right” contact angles about 20–30 s after placement of the drop. 1 Surface energies (γ) of the probed solid surfaces were calculated from contact angles and interfacial energies (γ₉) of the three probe liquids by use of eqs 1–3 and an in-house Visual Basic program. Total surface energy is the sum of Lifshitz–van der Waals (also called dispersion or nonpolar) and acid–base (also called polar) contributions. The polar portion can be further subdivided into Lewis acid and Lewis base components.

\[ \gamma_i = \gamma_i^{LW} + \gamma_i^{AB} = \gamma_i^{LW} + 2(\gamma_i^{LW})^{0.5} \]  
\[ \gamma_i = \gamma_i^{LW} + \gamma_i^{L} = \gamma_i^{LW} + 2(\gamma_i^{LW})^{0.5} \]  
\[ (1 + \cos \theta) = 2\left[(\gamma_i^{LW})^{0.5} + (\gamma_i^{LW})^{0.5} + (\gamma_i^{LW})^{1.5}\right] \]  
In eqs 1–3, superscripts denote components of the surface energy: LW, Lifshitz–van der Waals; AB, acid–base; L, Lewis acid; and \( \gamma_i \), Lewis base. In FW, \( \gamma_i^{LW} = 21.8 \text{ mJ/m}^2 \) and \( \gamma_i^{AB} = 25.5 \text{ mJ/m}^2 \); in DIM, \( \gamma_i^{LW} = 50.8 \text{ mJ/m}^2 \) and \( \gamma_i^{AB} = 0 \text{ mJ/m}^2 \); and in EG, \( \gamma_i^{LW} = 29 \text{ mJ/m}^2 \) and \( \gamma_i^{AB} = 1.92 \text{ mJ/m}^2 \), and \( \gamma_i^{AB} = 47 \text{ mJ/m}^2 \).

Six samples featuring both bottom-gate organic field-effect transistors (OFETs) and metal–insulator–metal (MIM) structures were fabricated side-by-side with the exception of thermal evaporation and desorption of aliphaphosphonic acid (C₉PA). One C₉PA was used in each sample.

MIM and OFET structures followed the same fabrication procedure up to and including the phosphonic acid monolayer. All thermal evaporations (metals and organics) were performed in Mini-Spectrocs. First, 30 nm thick aluminum was evaporated on a glass substrate through a shadow mask. Part of each electrode was coated with 40 nm thick Au to prevent its oxidation. Next, all samples were exposed to UV/ozone to produce about 10 nm thick AlOₓ. AlOₓ was functionalized with C₉PA by the process described earlier. Some AlOₓ was left uncoated to provide reference AIOₓ MIM structures. Next, a 20 nm thick DNTT layer was thermally evaporated at a rate of 0.5 Å/s at room temperature through a shadow mask. All MIM structures were masked during this step. Finally, a 50 nm thick Au layer was evaporated through a shadow mask at a rate of ~3 Å/s to themselves to good layer uniformity (atomic layer deposition) or self-limit the layer thickness (oxidation or use of monolayers) are advantageous. Aluminum oxide (AlOₓ) functionalized with aliphaphosphonic acids (CₙPA) is an established bilayer dielectric for low-voltage organic transistors. Such transistors have a bottom-gate structure where the aluminum oxide is commonly prepared by oxidation of the aluminum gate electrode and the assembly of the organic monolayer is performed in solvent-based solutions. A dry route to monolayer assembly has also been demonstrated for n-octylphosphonic acids with varying length of the aliphatic tail (CₙPA). To provide insight into the structure of CₙPA monolayers and to monolayer assembly has also been demonstrated for n-octylphosphonic acids (C₉PA). In such a case, several monolayers of C₉PA were thermally evaporated in vacuum, followed by thermal desorption of all molecules that were not chemically bonded to AlOₓ. In such a technique, nonuniformity in the as-deposited C₉PA thickness is eliminated during the second step of thermal desorption, leaving monolayer formation across the substrate. Thermal desorption provides an additional benefit of annealing that improves both the monolayer structure and transistor performance.

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complete the transistors and MIM structures. The transistors have nominal channel lengths of 30, 50, 70, and 90 μm and a channel width of 1000 μm.

Transistor and MIM measurements were performed with an Agilent B1500A semiconductor device analyzer under dark ambient conditions. All fabricated devices were kept in oxygen- and moisture-free environment until their measurement, and they shared the same history. The gate dielectric capacitance of MIM structures was measured between 1 kHz and 1 MHz and extracted at 100 kHz. The MIM current density was measured as a function of applied voltage between −3 and 3 V. The transfer and output characteristics of the OFETs were measured in a sweep mode. The threshold voltage and field-effect mobility were extracted from the transfer characteristics measured in saturation by use of metal–oxide–semiconductor field-effect transistor (MOSFET) equations. Subthreshold slope, on-current, off-current, and on/off-current ratio were also extracted from the saturation curve. Mean values and standard deviations were calculated for all relevant transistor parameters. Bias stress was performed at VGS = −2 V for 1000 s with source and drain electrodes grounded. At certain intervals, the bias stress was briefly interrupted and transistor transfer characteristics in saturation were measured to allow transistor parameter extraction.

Calculations were performed with density functional theory (DFT) as implemented in the Quantum Espresso (QE) package, version 5.1.  

The core electrons were represented by use of projector-augmented wave potentials, and valence electrons were represented with a planewave basis with cutoffs of 50 and 400 Ry for the wave functions and charge densities, respectively. The PBE version of the generalized gradient approximation was used for the exchange and correlation functional. Self-consistent calculations used an electronic convergence threshold of 1 × 10−5 Ry. Isolated molecules were generated with the aid of the chemical drawing tool Avogadro 1.1.1.  

The α-alumina (0001) slabs were six AlO₆Al layers thick, with a hexagonal lattice parameter of 4.80 Å. The top surface was terminated with an extra Al(OH)₃ layer. Alkylphosphonic acid molecules with alkyl chain length from 2 to 18 carbon atoms were added on top of the relaxed Al₂O₃ (0001) hydroxylated surface with either monodentate or bidentate binding, by removing H atoms from the surface. The molecule, top AlO₆Al layer, and hydroxylated Al(OH)₃ layer were allowed to relax until a force convergence threshold of 1 × 10−3 au was reached. A single k-point was used for the relaxation of isolated molecules, while a 3 × 3 × 1 hexagonal k-point mesh was used when the surface was present. Periodic boundary conditions were implemented, and a vacuum layer of at least 7 Å was added on top of the molecules, to avoid self-interaction.

Vibration frequencies were calculated by use of the Phonon package in QE, and the self-consistency threshold was 1 × 10−14 Ry. For a few cases (free acids, C₈PA monodentate, and C₁₈PA monodentate), the self-consistency threshold was 1 × 10−8 Ry. The vibrational mode characters were assigned by visualizing individual vibrational modes with the Molden 5.3 package.  

It is important to note that for some frequencies it was difficult to identify independent vibrational modes due to the interference of a wide range of vibrational modes occurring at similar frequencies. The vibration spectra were created by a superposition of Gaussians. Each Gaussian is centered at an identified vibration frequency and has a standard deviation of 10 cm⁻¹, and the area under the curve is proportional to the intensity of the mode.

3. RESULTS AND DISCUSSION

3.1. Gate Dielectric: Electrical and Structural Measurements. Figure 1a shows the capacitance of AlO₃/C₉PA bilayers and the thickness calculated for each alkylphosphonic acid. Total dielectric capacitance (C_dielectric) consists of capacitance of the AlO₃ layer (C_ALO₃) and capacitance of the phosphonic acid (C_PA). The two capacitances are in series, therefore 1/C_dielectric = 1/C_ALO₃ + 1/C_PA. This relationship can be rearranged to find C_PA, which is then used to calculate C_PA thickness by using a relative permittivity value of 2.1. The capacitance of bare AlO₃ has a mean value of 0.60 μF/cm². AlO₃/C₉PA capacitance is lower, and a decrease in capacitance per unit area is seen as C_PA length increases. The increase in C_PA thickness from 0.83 nm for C₈PA to 2.49 nm for C₉₈PA is consistent with the increased linear length of C₉PA molecules when n increases from 8 to 18.

AlO₃ and AlO₃/C₉PA leakage current densities for various phosphonic acids are shown in Figure 1b. Bare AlO₃ dielectric displays a leakage current density of ~2 × 10⁻⁷ A/cm² at −3 V. At the same voltage, AlO₃ functionalized with phosphonic acid shows reduced leakage current density with values between ~6 × 10⁻⁸ and ~3 × 10⁻⁸ A/cm². Leakage current decreases as C_PA length increases. In summary, both capacitance and leakage current measurements confirmed that the thickness of C_PA increased with increasing n, while capacitance measurement determined that in all cases the C_PA thickness corresponds to about a monolayer.

Water contact angles of AlO₃/C₉PA as a function of C₉PA length are shown in Table 1a. After annealing to remove physisorbed molecules, all AlO₃/C₉PA surfaces are hydrophobic, with water contact angles greater than 110° regardless of the alkyl chain length. Although a maximum value of 112.0° ± 1.1° and a minimum value of 110.8° ± 1.3° were obtained for C₈PA and C₇₈PA respectively, all contact angles are the same within the error of measurement. These contact angles were compared to a C₇₈PA layer prior to annealing, whose thickness of ~20 nm was confirmed by atomic force microscopy (AFM). This layer exhibits significantly different contact angles for all three liquids.
monolayers exhibit slightly lower contact angles for FW and DIM and higher surface energies ranging between 25 and 30 mJ/m². However, this difference may result from the fact that ref 34 reports results for alkylphosphonic acids on sol-gel-derived hafnium oxide, while our results are for C₈PA on aluminum oxide.

AlOₓ/C₈PA surfaces were also investigated by AFM (see Table 2). In addition to determining the root-mean-square (RMS) surface roughness, the use of a new PeakForce QNM scanning mode enabled direct extraction of quantitative nanomechanical information such as force of adhesion (Fₐd), elastic modulus (E), deformation, and dissipation. The surface roughness (Rₙ) of ∼10 nm thick AlOₓ is 1.27 nm. The 3-h anneal at 160 °C does not affect its surface roughness, confirming that C₈PA desorption/annealing step would not lead to increased surface roughness of the AlOₓ/C₈PA dielectric due to increased roughness of AlOₓ. The surface roughness of AlOₓ functionalized with C₈PA is comparable or slightly higher but below 2 nm in most cases. The C₈PA layer prior to annealing has a roughness of 2.25 nm. There is no correlation between AlOₓ/C₈PA surface roughness and length of the phosphonic acid. In addition, no correlation is observed between C₈PA length and nanomechanical properties. Force values lie between ∼3 and ∼5 nN, moduli are between 22 and 63 GPa, deformation falls between ∼3 and ∼8 nm, and dissipation is between ∼200 and 1000 eV. Previously, the surface roughness of alkylphosphonic acids, solution-assembled on top of Si/AlOₓ, was affected by the length of the molecule, and the lowest surface roughness was achieved for C₈PA.¹⁴

In summary, AlOₓ/C₈PA surfaces after annealing exhibit comparable RMS surface roughness. While their mechanical properties on the nanometer scale (AFM tip size is 2 nm) vary by a factor of 2–5 (see Table 2), their macroscopic surface properties (see Table 1) are similar.

Fourier transform infrared (FTIR) spectroscopy provides structural information on a macroscopic scale, that is, the scale of the transistor channel. FTIR was performed on AlOₓ/C₈PA surfaces after annealing and on the reference AlOₓ surface (see Figure 2). A strong broad absorbance near 900 cm⁻¹ is dominated by Al–O vibrations in all samples. Weaker vibrations are observed in the region from 1000 to 1250 cm⁻¹ near 1450 cm⁻¹, and between 2800 and 3000 cm⁻¹. In general, the integral intensities of various peaks is higher for shorter-chain C₆PA and lower for longer chains. As the length of the phosphonic acid is increased, the position of C–H stretching is shifted to lower wavenumbers and their full width at half maximum (fwhm) is reduced. For C₈PA the peaks are centered at 2853 cm⁻¹ (fwhm = 20.7 cm⁻¹) and 2924 cm⁻¹.

After annealing to remove physisorbed molecules, all AlOₓ/C₈PA surfaces have low surface energy of ∼17.5 mJ/m² with dominating Lifshitz–van der Waals (dispersive) component. No correlation between surface properties and length of the aliphatic chain is observed. The small Lewis acid component and negligible Lewis base component indicate that the surface has some electron-pair-accepting ability. On the contrary, surface energy of the ∼20 nm thick C₆PA layer prior to annealing is much higher at ∼31 mJ/m². The larger surface energy suggests that some C₆PA head groups are found on the surface. In this case the Lewis base component dominates (the surface has electron-pair-donating ability), followed by the sizable Lifshitz–van der Waals energy, which is about 50% higher than that of C₈PA monolayers after annealing. While C₆PA molecules are likely to be “standing up” in the monolayers as a result of their bonding to AlOₓ, C₈PA molecules that form the top surface of a 20 nm thick layer are expected to assume random orientations, resulting in exposed —OH groups and different surface properties.

Comparing these results to those obtained for solution-assembled C₆PA, one would notice that solution-assembled

<table>
<thead>
<tr>
<th>(b) Surface Energies (mJ/m²)</th>
<th>γ⁺</th>
<th>γ⁻</th>
<th>γₛ</th>
<th>γₚ</th>
</tr>
</thead>
<tbody>
<tr>
<td>C₈PA</td>
<td>3.2</td>
<td>0.56</td>
<td>14.9</td>
<td>17.61</td>
</tr>
<tr>
<td>C₆PA</td>
<td>3.18</td>
<td>0.53</td>
<td>14.39</td>
<td>16.98</td>
</tr>
<tr>
<td>C₆PA</td>
<td>3.15</td>
<td>0.58</td>
<td>14.02</td>
<td>17.21</td>
</tr>
<tr>
<td>C₆PA</td>
<td>2.73</td>
<td>0.44</td>
<td>15.69</td>
<td>17.89</td>
</tr>
<tr>
<td>C₆PA</td>
<td>3.64</td>
<td>0.82</td>
<td>15.06</td>
<td>18.51</td>
</tr>
<tr>
<td>C₆PA</td>
<td>3.30</td>
<td>0.55</td>
<td>14.44</td>
<td>17.13</td>
</tr>
<tr>
<td>C₆PA</td>
<td>0.67</td>
<td>27.23</td>
<td>22.48</td>
<td>31.03</td>
</tr>
</tbody>
</table>

“FW, filtered water; EG, ethylene glycol; DIM, diiodomethane. γ⁺, γ⁻, γₛ, γₚ, Lifshitz–van der Waals; Lewis acid; Lewis base. *Prior to annealing (∼20 nm thick).”

### Table 2. Surface Roughness and Nanomechanical Properties Determined by AFM

<table>
<thead>
<tr>
<th>surface</th>
<th>roughness (nm)</th>
<th>force (nN)</th>
<th>modulus (GPa)</th>
<th>deformation (nm)</th>
<th>dissipation (eV)</th>
</tr>
</thead>
<tbody>
<tr>
<td>C₈PA</td>
<td>1.25</td>
<td>3.1 ± 0.7</td>
<td>63 ± 9</td>
<td>2.8 ± 0.6</td>
<td>353</td>
</tr>
<tr>
<td>C₆PA</td>
<td>1.53</td>
<td>3.6 ± 1.1</td>
<td>30 ± 9</td>
<td>3.2 ± 0.5</td>
<td>226</td>
</tr>
<tr>
<td>C₆PA</td>
<td>2.34</td>
<td>5.3 ± 1.6</td>
<td>28 ± 10</td>
<td>3.6 ± 0.7</td>
<td>1016</td>
</tr>
<tr>
<td>C₆PA</td>
<td>1.13</td>
<td>5.4 ± 1.5</td>
<td>29 ± 8</td>
<td>8.6 ± 2.5</td>
<td>277</td>
</tr>
<tr>
<td>C₆PA</td>
<td>1.82</td>
<td>2.8 ± 0.6</td>
<td>22 ± 5</td>
<td>4.0 ± 0.6</td>
<td>219</td>
</tr>
<tr>
<td>C₆PA</td>
<td>1.47</td>
<td>3.9 ± 1.0</td>
<td>37 ± 11</td>
<td>3.1 ± 0.1</td>
<td>945</td>
</tr>
<tr>
<td>C₆PA</td>
<td>2.25</td>
<td>3.3 ± 1.2</td>
<td>65 ± 31</td>
<td>4.2 ± 1.1</td>
<td>566</td>
</tr>
<tr>
<td>AlOₓ</td>
<td>1.27</td>
<td>4.9 ± 1.3</td>
<td>43 ± 10</td>
<td>4.4 ± 0.1</td>
<td>961</td>
</tr>
<tr>
<td>AlOₓ</td>
<td>1.22</td>
<td>4.9 ± 1.1</td>
<td>65 ± 50</td>
<td>4.5 ± 0.1</td>
<td>509</td>
</tr>
</tbody>
</table>

“Prior to annealing (∼20 nm thick).”
(FWHM = 31.9 cm⁻¹), while for C₃₄PA the peaks are found at 2850 cm⁻¹ (FWHM = 17.3 cm⁻¹) and 2920 cm⁻¹ (FWHM = 28.4 cm⁻¹). These peak positions are very similar to those reported for C₃₄PA monolayers prepared in ethanol on hafnium oxide. The shift in the peak location was previously interpreted as an improved molecular order within monolayers; that is, the increase in C₃₄PA length leads to stronger van der Waals interaction between aliphatic chains and results in a more ordered self-assembly with denser molecular packing. However, one would predict that the density of vacuum-deposited monolayers would be lower than that of solution-deposited monolayers as a result of the laws that govern physical vapor deposition. This is supported by the fact that the integral intensity of CH stretching vibrations confirms that a degree of order exists even for vacuum-deposited monolayers, and the molecular order improves with increasing length of C₃₄PA. The spread in nanomechanical properties suggests heterogeneous monolayer structure, such as presence of domains or nanopores. Previously, molecular dynamic simulations performed on solution-assembled C₃₄PA monolayers on aluminum oxide confirmed a change in the morphology from amorphous to quasi-crystalline with increasing length of C₃₄PA. In such a case, highly ordered domains with gaps between them exist for long C₃₄PA molecules.

In summary, CH₂ stretching vibrations confirm that a degree of order exists even for vacuum-deposited monolayers, and the molecular order improves with increasing length of C₃₄PA. The spread in nanomechanical properties suggests heterogeneous monolayer structure, such as presence of domains or nanopores. Previously, molecular dynamic simulations performed on solution-assembled C₃₄PA monolayers on aluminum oxide confirmed a change in the morphology from amorphous to quasi-crystalline with increasing length of C₃₄PA. In such a case, highly ordered domains with gaps between them exist for long C₃₄PA molecules.

3.2. Organic Field-Effect Transistors: As Fabricated and Under Bias Stress. The cross-section of the bottom gate Al/AIOₓ/C₃₄PA/DNTT/Au transistor is shown in Figure 3a. Figure 3b shows transistor transfer characteristics for as-fabricated transistors with various phosphonic acid monolayers. As C₃₄PA length increases, threshold voltage (Vₜ) and transistor off-current (Iₒff) decrease. Figure 3c depicts output characteristics of an OFET with C₃₄PA and nominal channel length of 30 μm. The behavior of all transistor parameters for as-fabricated transistors is shown in Figure 3d-i. Upon going from C₃₄PA to C₃₄PA, mean threshold voltage (Vₜ) decreases from -1.37 to -1.24 V, field-effect mobility (μ) increases from 0.03 to 0.33 cm²/(V·s), subthreshold slope (S) remains the same within the error of measurement (variation between 86 and 94 mV/decade), off-current (Iₒff) decreases from ~8 × 10⁻¹³ to ~3 × 10⁻¹² A, and for OFETs with L = 30 μm, on-current (Iₚₜ) increases from ~3 × 10⁻⁸ to ~2 × 10⁻⁶ A and on/off-current ratio (Iₒff/Iₚₜ) increases from ~3 × 10⁸ to ~4 × 10⁹.

Overall, increasing length of the phosphonic acid leads to a significant improvement of transistor parameters. This behavior is different from the behavior of transistors that incorporated solution-assembled C₃₄PA. For Si/HfO₂/C₃₄PA/pentacene/Au OFETs, threshold voltage became more negative with increasing C₃₄PA length, while mobility of C₃₄PA exceeded that of C₁₈₆PA. For Si/AIOₓ/C₃₄PA/pentacene/Au OFETs, voltage did not change significantly and mobility peaked for C₃₄PA. For Si/SiO₂/C₃₄PA/pentacene/Au OFETs, the lowest threshold voltage occurred for C₃₄PA, mobility decreased from C₃₄PA to C₃₄PA, and subthreshold slope was unaffected. For Al/AIOₓ/C₃₄PA/pentacene/Au OFETs, mobility peaked for C₃₄PA for oxygen-plasma AIOₓ and increased with increasing C₃₄PA length for mild-air-plasma AIOₓ. For Si/AIOₓ/C₃₄PA/pentacene/Au OFETs, both mobility and threshold voltage increased with increasing C₃₄PA length.

Comparison of our C₃₄PA transistors to recently reported DNTT transistors that use other gate dielectrics is as follows. Cross-linked poly(ethylene-alt-maleic anhydride) (PEMA) led to a mobility of 0.11 cm²/(V·s), while PEMA modified with
poly(maleic anhydride-alt-1-octadecane) resulted in mobility of 0.24 cm²/V·s.\textsuperscript{10} Octylamine-treated PEMA gate dielectric led to mobility of 0.17 cm²/V·s.\textsuperscript{40} Vapor-jet-deposited DNTT on polystyrene-buffered poly(tripropylene glycol diacrylate) dielectric achieved a mobility of 0.43 cm²/V·s.\textsuperscript{41} Field-effect mobilities in excess of 1 cm²/V·s were achieved when gate dielectrics incorporating organic monolayers were used.\textsuperscript{42−44} Threshold voltage and subthreshold slope depend on the thickness of the gate dielectric, and our values are consistent with the values achieved for other thin dielectrics. Similarly, the comparison of the on/off ratio is difficult because it depends on transistor dimensions.

A short bias stress lasting for 1000 s was also performed. During the bias stress, a voltage of −2 V was applied to the gate while source and drain electrodes were grounded. All samples were kept in dark ambient air for 3 days before the bias stress was performed. Figure 4a shows the evolution of threshold voltage with increasing bias stress time. While transistors incorporating C\textsubscript{16}PA and C\textsubscript{14}PA monolayers exhibit initial threshold voltage similar to that shown in Figure 3, transistors with shorter phosphonic acids exhibit lower initial V\textsubscript{T\textsubscript{th}}, indicating poorer ambient stability. Initial values of mobility and off-current are similar to those shown in Figure 3.

Bias stress leads to a more negative threshold voltage for all transistors; however, the rate of degradation decreases as length of the phosphonic acid increases, making transistors with C\textsubscript{14}PA almost a factor of 2 more stable than those with C\textsubscript{16}PA. Degradation in field-effect mobility (see Figure 4b) exhibits different behavior for various C\textsubscript{3}PA. For C\textsubscript{16}PA and C\textsubscript{14}PA, mobility exhibits only a minor change, reaching ∼93% of the initial value at 1000 s. For C\textsubscript{14}PA and C\textsubscript{12}PA, mobility first increases by about 20% and then decreases to about 80% of the initial value. For the shortest C\textsubscript{10}PA, mobility first increases by 30−35% and then decreases to about 60% of the initial value. Off-current decreases for all C\textsubscript{3}PA (see Figure 4c). While the decrease is almost negligible for longer C\textsubscript{3}PA, the drop in I\textsubscript{off} is much more pronounced for C\textsubscript{10}PA and C\textsubscript{12}PA.

It has been shown that bias-stress degradation of DNTT transistors with polystyrene-buffered poly(tripropylene glycol diacrylate) dielectric depends on environmental conditions.\textsuperscript{25} Since all our transistors were bias-stressed in the same laboratory environment (air, ∼40% relative humidity), differences in their degradation are ascribed to transistor structure.

The choice of C\textsubscript{3}PA affects gate dielectric capacitance (see Figure 1) and threshold voltage of the transistors (see Figure 3). Therefore, one should consider the induced capacitive charge at the beginning of the transistor bias-stress degradation. Since the gate-to-channel voltage is not known, the gate-to-source voltage V\textsubscript{GS} is used to approximate the accumulated charge; that is, Q = C |V\textsubscript{GS} − V\textsubscript{th}|. This charge is 0.41 μC/cm² for C\textsubscript{16}PA, 0.24 μC/cm² for C\textsubscript{14}PA, and 0.31−0.33 μC/cm² for the remaining C\textsubscript{3}PA. If transistor bias degradation were solely controlled by induced charge density, then transistors with C\textsubscript{16}PA, C\textsubscript{14}PA, and C\textsubscript{12}PA should exhibit similar degradation behavior. However, the results of Figure 4 show degradation that is clearly linked to length of the phosphonic acid instead of induced charge density.

The experimental results confirm strong correlation between initial and bias-induced transistor behavior and length of the C\textsubscript{3}PA monolayer. Overall, the transistors exhibit improved initial parameters and ambient bias-stress stability when C\textsubscript{3}PA length increases. FTIR vibrations between 2800 and 3000 cm\textsuperscript{-1} indicate a degree of order within the monolayer that improves with increasing C\textsubscript{3}PA length. Our discussion will now focus on DFT results and parts of the FTIR spectra that “probe” the bonding of C\textsubscript{3}PA to AlO\textsubscript{x}.

3.3. Density Functional Theory. As shown above, water contact angles and surface energy are the same for all AlO\textsubscript{x}/C\textsubscript{3}PA surfaces after annealing. Minor differences in RMS surface roughness are not correlated to the length of C\textsubscript{3}PA. However, FTIR spectra show differences and so does the transistor behavior. DFT was therefore used to calculate vibration frequencies of different C\textsubscript{3}PA molecules in their free state as well as bonded to stoichiometric AlO\textsubscript{3}. Here, monodentate and bidentate bonding was considered.

Figure 5 shows sections of the vibration spectra calculated from DFT. Vibrations exist below 700 cm\textsuperscript{-1}; however, this section is not shown as there are no experimental data to match it. For free acids, P−OH vibrations are located between ∼800 and ∼850 cm\textsuperscript{-1}, P=O is found near 1250 cm\textsuperscript{-1}, and CH\textsubscript{2}/CH\textsubscript{3} stretches occur between ∼2930 and ∼3050 cm\textsuperscript{-1}. These frequencies can be compared with calculations from self-consistent charge density-functional tight binding (SCC-DFTB),\textsuperscript{26} which is based on DFT but is a more approximate method. Such calculations gave P−OH vibrations in the range 625−683 cm\textsuperscript{-1}, P=O vibration of 1324 cm\textsuperscript{-1}, and CH\textsubscript{2}/CH\textsubscript{3} vibrations in the region between 2750 and 3000 cm\textsuperscript{-1}. For monodentate attachment of C\textsubscript{3}PA (see Figure 5), the region between 700 and 1000 cm\textsuperscript{-1} contains AlO, P−OAl, P−OH, and P−C vibrations. AlO−H bending and P=O stretching are both found near 1025 and 1150 cm\textsuperscript{-1}; however, AlO−H bending contributes more strongly to the vibration at 1025 cm\textsuperscript{-1}, while P=O stretch dominates the vibration near 1150 cm\textsuperscript{-1}. C\textsubscript{3}PA length does not affect the position of these peaks.
This ultimately leads to tridentate attachment and to an increase in the degree of interaction between the molecule and the surface. The two main CH bonds are pushed further apart due to an additional shift of CH:H vibrations. For HPO(OH)2 on alumina, SCC-DFTB gave the P=O frequency as 1325 cm⁻¹, which is very similar to their monodentate value.

The two main CH/CH₂ stretches are pushed further apart due to CH/H asymmetric stretching being shifted to slightly higher wavenumbers. The P=O frequency of 1150 cm⁻¹ can be compared to the SCC-DFTB value of 1320 cm⁻¹ for phosphonic acid HPO(OH)₂ on alumina and to an experimental value of 1278 cm⁻¹ for CpPA acid on silicon. DFT calculations of alklyphosphonic acids on silicon gave CH/CH₂ frequencies in the range 2925–3075 cm⁻¹. Finally, for bidentate attachment of CpPA (see Figure 5), the region between 700 and 1000 cm⁻¹ contains AlO, P=OAl, and P=C vibrational bands. AlO-H bending and P=O stretch are both found near 1100 cm⁻¹, and CpPA length does not affect the position or intensity of this peak. The two main CH/CH₂ stretches are pushed even further apart due to an additional shift of CH/H asymmetric stretching to higher wavenumbers. The second CH/CH₂ asymmetric stretch is also shifted to higher wavenumbers. For HPO(OH)₂ on alumina, SCC-DFTB gave the P=O frequency as 1325 cm⁻¹, which is very similar to their monodentate value.

Compared to the measured data of Figure 2, calculated positions of CH/CH₂ stretches are shifted to higher wavenumbers for all bonded phosphonic acids. This could be due to two reasons: (a) DFT approximation of exchange and correlation effects and (b) neglect of dynamic (finite temperature) effects. Inclusion of van der Waals interactions that partially account for electron correlation effects may improve the DFT results, as has been shown for the case of water. Overall, the most significant differences in vibration spectra between monodentate and bidentate attachments are in the regions 750–800 cm⁻¹ and 1000–1200 cm⁻¹. While the former is buried in the strong AlO peak of measured FTIR spectra, the second region can be used to analyze the experimental data. DFT also showed that the tilt of CpPA with respect to the surface normal decreases with increasing length of the molecule, and for any CpPA the bidentate attachment results in a smaller tilt when compared to the monodentate attachment (see Supporting Information). Tilt angles are smaller than those predicted by the SCC-DFTB method, but this is to be expected as our coverage is higher, forcing the molecules to be more upright.

Previous research has shown that phosphonic acids self-assemble from solutions strongly attach to aluminum oxide. Attachment is facilitated by the headgroup that reacts with surface hydroxyl groups of aluminum oxide. When the metal oxide surface possesses Lewis acidic sites, binding originates from coordination of P=O to such a site, followed by the condensation reaction between P=O and Al–OH moieties to produce P–O–Al bonds. This ultimately leads to tridentate attachment. On metal oxides lacking Lewis acidity, reaction between P=O and Al–OH moieties results in bidentate attachment. Tridentate attachment is also possible when a hydrogen bond between surface —OH and P=O moieties is formed. Another proposed mechanism involves protonation of surface —OH groups followed by formation of the P=O–Al bond. In such a case, the phosphorus atom is left with a positive charge. In addition, DFT calculations have shown that the thermodynamically preferred binding mode depends on surface structure of the aluminum oxide and the amount of residual water.

DFT calculations show that monodentate binding of CpPA to Al₂O₃ results in a strong PO–H vibration found below the region of CH₂/CH₃ stretches. Since measured FTIR spectra of all CpPA monolayers lack such vibrations, monodentate attachment is unlikely for vacuum-deposited monolayers. This is further supported by the presence of a broad vibration band near 1100 cm⁻¹, consistent with bidentate bonding. DFT assigns this vibration band to AlO–H bending and P=O stretching. The measured integral intensity of this band decreases with increasing CpPA length, and the contributing vibration frequencies move slightly apart. This is interpreted as reduced CpPA coverage because similar reduction in the integral intensity of CH₂ stretches is observed for longer CpPA. This also agrees with the expected random orientation of CpPA molecules during physical vapor deposition and the area each molecule is likely to occupy. Another possibility is that some P=O bonds disappear as a result of tridentate attachment. However, if tridentate attachment occurred for longer CpPA via protonation of surface —OH groups, the resulting immobile positive charge should lead to a more negative threshold voltage, which is not observed. Formation of a hydrogen bond between surface —OH and P=O moieties is more plausible, because it would lead to a shift in AlO–H and P=O vibration.
frequencies and no immobile charge. Consequently, tridentate attachment via hydrogen bond cannot be excluded.

3.4. Final Remarks. We would like to make a few remarks about the observed transistor behavior. Negatively charged immobile species in the presence of hydroxyl groups were demonstrated in the case of Si–OH when silicon dioxide was used as gate dielectric61 or as a result of electrochemical reactions that involve adsorbed water.57 The presence of Al–OH groups on the surface of AlOx could also lead to fixed negative charge located at the AlOxC/PAl interface. Since more Al–OH sites would be expected for longer C8PA when fewer molecules are attached to AlOx, lower threshold voltage is expected for longer C8PA. This is indeed observed for as-fabricated transistors. However, the presence of Al–OH sites does not explain the different ambient and bias-stress degradations.

Phosphonic acids possess a dipole moment that has been shown to control the threshold voltage of the transistors.56 The component normal to the AlOx surface is quite small if the molecule stands perpendicular to the surface.56 This component would change with length and tilt of the C8PA molecule. Bending of the longer molecules observed by DFT would complicate matters even more. The existence of dipole moments can potentially explain the observed differences in IVf for as-fabricated transistors. However, it cannot explain the different ambient and bias-stress degradations.

Another possibility involves the reduced density of C8PA monolayers for longer molecules. Lower molecular coverage can be intentionally induced by substrate heating during C8PA growth.22 Our previous experiments with C8PA and pentacene showed an increase in IVf, and a decrease in field-effect mobility with increasing C8PA growth temperature.22 This behavior is opposite to that observed in Figure 3, and therefore, varied molecular density is an unlikely cause of the change in threshold voltage.

Finally, changing morphology of the monolayers with increasing C8PA length could have a profound effect on the growth of DNTT. The large increase in field-effect mobility and the dissimilar degradation behavior must be controlled by the C8PA/DNTT interface and/or DNTT itself. Additional research is needed to understand how these vacuum-evaporated alkylphosphonic acids control the growth of DNTT.

4. CONCLUSION

Growth that self-limits the thickness of materials is desirable for ultrathin dielectrics for low-voltage transistors. In this paper, we showed that monolayers of alkylphosphonic acids (Cn−C18) can be prepared by vacuum evaporation and incorporated into organic field-effect transistors based on DNTT.

AlOxC/PAl bilayers (∼11–12 nm thick) exhibit low leakage current densities ranging between ∼6 × 10−8 and ∼3 × 10−8 A/cm2 at ∼3 V. The decrease in capacitance with increasing length of C8PA confirms monolayer formation for all phosphonic acids. Total surface energy of AlOxC/PAl surfaces after annealing is ∼17.5 mJ/m2 and independent of C8PA length. All AlOxC/PAl surfaces exhibit comparable RMS surface roughness. While their macroscopic surface properties are similar, the mechanical properties on the nanometer scale vary by a factor of 2–5. Similarly to solution-assembled monolayers, the CH2 stretching peaks narrow and shift to lower wavenumbers, confirming that molecular order improves with increasing length of C8PA. At the same time, reduced molecular coverage appears for longer CnPAs. The spread in nano-mechanical properties suggests a heterogeneous monolayer structure, such as the presence of domains or nanopores.

Performance of as-fabricated transistors is affected considerably by the chosen C8PA. Upon going from C8PA to C18PA, threshold voltage moves closer to zero by ∼10%, field-effect mobility increases by an order of magnitude, off-current decreases by ∼50%, and subthreshold slope does not visibly change. As a result, on-current and off/on-current ratio increase by 2 orders of magnitude for OFETs with L = 30 μm. Increasing C8PA length leads to a significant improvement of transistor parameters. Results of bias stress also confirm that degradation behavior is linked to length of the phosphonic acid instead of induced charge density. As C8PA length increases, transistors are less prone to bias stress. In addition, transistors with longer C8PA exhibit better air stability.

DFT calculations show that bonding of C8PA molecules (monodentate versus bidentate) results in different vibration frequencies between 1000 and 1200 cm−1. Strong PO–H vibration is also present for monodentate bonding. Comparison of DFT results and FTIR measurements leads to the conclusion that monodentate bonding does not occur for any C8PA.

■ ASSOCIATED CONTENT

5 Supporting Information

The Supporting Information is available free of charge on the ACS Publications website at DOI: 10.1021/acsami.6b08426.

Four figures showing images of relaxed phosphonic acids (C10H, C14H, and C18H) on alumina in monodentate and bidentate coordinations and comparison of DFT vibration spectra of phosphonic acids (C2−C18) free and bonded to alumina (PDF)

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■ ACKNOWLEDGMENTS

S.H. is a recipient of a doctoral training grant funded by the Engineering and Physical Sciences Research Council (EPSRC), Grant EP/L505080/1. J.C. was funded by a WISE Bridging the Gap project, University of Strathclyde. DFT results were obtained by use of the EPSRC-funded ARCHIE-WeSt High Performance Computer (www.archie-west.ac.uk); EPSRC Grant EP/K000586/1. FTIR analysis was developed within CENTEM project CZ.1.05/2.1.00/03.0088, cofunded by the ERDF and follow-up sustainability stage CENTEM+ (LO1402) under the National Sustainability Programme I.

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