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Surface engineering of high-k polymeric dielectric layers with a fluorinated organic crosslinker for use in flexible-platform electronics<sup>†</sup>

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High-*k* polymeric layers were prepared by combining various functional groups and were applied as gate dielectrics for practical organic field-effect transistors (OFETs). Crosslinking of the polymeric layers through UV-assisted organic azide fluorine-based crosslinkers induced dramatic improvements in the electrical performance of the OFET, such as field-effect mobility and bias-stress stability. Our synthesis and manufacturing method can be a useful technique for ensuring device operation stability and electrical property enhancement. With this analysis, we further applied our polymer-dielectric OFETs to flexible-platform-based electronic components, including unit OFETs and simple logic devices (NOT, NAND, and NOR gates). The outcomes of this research and development suggest a suitable method for the low-cost mass production of large-area flexible and printable devices, using a printing-based approach to replace current processes.

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# 1. Introduction

There has been growing interest in the development of nextgeneration electronics for applications in various fields, including wearable and biocompatible electronics.<sup>1,2</sup> Organic fieldeffect transistors (OFETs) are considered vital building blocks for next-generation electronics.<sup>3,4</sup> OFETs have been extensively studied for several decades and have been used in high-performance devices with various functionalities owing to their advantages of mechanical flexibility, low-cost processing, and large-area scalability.<sup>5</sup> However, implementing these high-performance OFETs in next-generation electronics still remains a challenge.

One of the most significant criteria for the practical application of OFETs in electronic components is improving OFET performance while maintaining operational stability. The

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operational stability of OFETs can be classified in various ways,<sup>6,7</sup> most of which are strongly determined by the gate dielectrics. The gate dielectrics in OFETs play a role in lowering the working voltage of the device, limiting the leakage current during operation, and affecting the trap density at the interface between organic semiconductors and insulators.<sup>8-11</sup> In particular, leakage current and trap density have a significant impact on the operational stability of the device. As the leakage current characteristic increases, the OFF current level increases, which greatly affects the ratio between ON and OFF currents.<sup>12</sup> Additionally, interface traps can create residual charges and shift operational behavior, causing operational instability.<sup>7</sup>

To date, the most common method for improving the operational stability of OFETs is the treatment of the surface with an inorganic oxide insulating material.<sup>13</sup> Inorganic oxide-based dielectrics exhibit excellent insulating properties owing to their high packing densities. When the surface is treated with hydrophobic organic materials, such as various self-assembled monolayer materials, the compatibility with organic semiconductors is enhanced and yields OFETs with good operational stability. However, these high-packing inorganic materials are usually manufactured using expensive vacuum deposition and hightemperature heat treatments.<sup>14,15</sup> Additional surface treatment methods often further complicate the manufacturing process. In addition, it is difficult to apply these types of insulating materials to plastic substrates because of their brittleness.<sup>16-18</sup>

Concerning the above, organic polymeric dielectrics have several advantages over dielectrics made from inorganic materials. Polymeric materials are naturally compatible with organic materials, thereby minimizing the interfacial trap number and



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improving the electrical performance of OFETs.<sup>8</sup> In particular, the desired properties (*e.g.*, mechanical flexibility, hydrophobicity, crosslinking, and solution-processability) can be easily realized through molecular design/synthesis.<sup>19–21</sup> Therefore, the common disadvantages of polymers, such as their low leakage properties, mechanical thermal stability, and low dielectric constant (k), can be readily resolved.<sup>22,23</sup>

Herein, we synthesized a high-k (k > 7) alkyl-based hydrophobic polymeric dielectric material, MBHCa-Ha, for use in the gate dielectrics of OFETs. The prepared material was further modified with an organic azide fluorine-based crosslinker (FPA-3F) to enhance the electrical performance of the OFET, namely the operational stability and interfacial properties. We investigated the effects of crosslinking in polymeric dielectric layers on the morphological and electrical properties and demonstrated a dramatic improvement in device operation. In addition, we manufactured flexible unit OFETs, NOT, NAND, and NOR gates, all of which suggested that the crosslinked layers contributed to robust and stable transistor operation.

# 2. Experimental section

#### 2.1 Materials

All chemical solvents and reagents were purchased from commercial suppliers, and all reactions were performed in oven-dried glassware. Methyl methacrylate (MMA), butyl acrylate (BA), mono(2-acryloyloxyethyl)succinate (HEA-suc), cyclohexyl methacrylate (Ca), hexyl acrylate (Ha), and ethyl acetate (EAc) were purchased from Sigma-Aldrich. 2,2'-Azobis(2methylpropionitrile) (AIBN) was purchased from TCI Chemical. From the first alphabet of the monomer, the synthesized polymer was named MBHCa-Ha.

### 2.2 Synthesis of the MBHCa-Ha polymer

Methyl methacrylate (12 g, 0.12 mol), butyl acrylate (8 g, 0.06 mol), HEA-suc (14 g, 0.06 mol), cyclohexyl methacrylate (10 g, 0.06 mol), hexyl acrylate (9.37 g, 0.06 mol), ethyl acetate (100 mL), and AIBN (0.1 g) were weighed and deposited into a 250 mL 4-neck round bottom flask connected to a reflux condenser, mechanical stirrer, and thermometer. After dissolving the monomers and initiator, purging with nitrogen gas was performed for at least 30 minutes. After internal displacement was complete, the solution flask was heated to 85 °C. Then the temperature was maintained for 9 h. The reaction was terminated by cooling the solution to room temperature ( $\sim 25^{\circ}$ C) and passing air over it. A viscous solution was obtained and purified by precipitation with hexane. Finally, after vacuum drying, a solid white MBHCa-Ha polymer was obtained. Yield: 32 g, 48%, GPC:  $M_{\rm n} = 509$  kDa,  $M_{\rm w} = 1095$  kDa, PDI = 2.15. <sup>1</sup>H NMR (300 MHz, CDCl<sub>3</sub>) δ 4.66-4.64 (br, 1H), 4.31-4.28 (br, 3H), 4.00-3.97 (br, 4H), 3.64-3.59 (m, 8H), 2.69-2.67 (br, 4H), 1.82-1.59 (br, 6H), 1.30-1.25 (m, 14H), 0.96-0.86 (m, 12H).

#### 2.3 Device fabrication

n-Doped silicon wafer with SiO<sub>2</sub> (thermally grown, 300 nm) wafers (Namkang Hi-tech) and polyethylene terephthalate (PET)

films (thickness 188 µm, KOLON Ind., Korea) were used as substrates for device manufacturing. The substrates were cleaned as follows. Si wafers were cleaned with boiled acetone for 15 min, then underwent ultrasonic cleaning with acetone and isopropyl sequentially for 20 min each, and finally UVozone treatment for 10 min. The PET was subjected to ultrasonic cleaning for 1 min, followed by UV-ozone treatment for 10 min. Dielectric solutions were then prepared by diluting 5 wt% MBHCa-Ha or MBHCa-Ha/FPA-3F in propylene glycol methyl ether acetate (PGMEA, Sigma-Aldrich). The weight ratio of MBHCa-Ha to FPA-3F was 95:5.

After the cleaning and solution preparation processes, Al electrodes were deposited by using a thermal evaporator (2-3 Å s<sup>-1</sup>, vacuum pressure of 10<sup>-6</sup> Torr, and substrate temperature of 25 °C) to a thickness of 30 nm on the cleaned substrates. The polymeric layers were coated by spin coating (2000 rpm for 30 s) or by a printing process using an electrostatic force-assisted dispensing method.24 A thermal annealing process at 120 °C was then conducted for 30 min to eliminate residual solvent. For the crosslinked sample, the crosslinking reaction was conducted under deep UV irradiation (~254 nm) in an N2-purged glove box.25 Synthesized organic semiconducting layers composed of 2,9-didecyldinaphtho[2,3-b:2',3'-f]thieno[3,2-b] thiophene (C10-DNTT) were deposited by organic molecular beam deposition as described in previous work.26 The device fabrication process was finished by the deposition of Au source/ drain (S/D) electrodes (50 nm) on the sample using a thermal evaporator (2 Å s<sup>-1</sup>, vacuum pressure of  $10^{-6}$  Torr, and substrate temperature of 25 °C).

#### 2.4 Characterization

The synthesized polymers were analyzed using various instrumental techniques. The <sup>1</sup>H NMR (nuclear magnetic resonance) spectrum was acquired using a Bruker Avance III 300 spectrometer, and the gel permeation chromatography (GPC) molecular weight was estimated using an Agilent Technologies 1260. Differential scanning calorimetry (DSC) and thermogravimetric analysis (TGA) were performed using TA Instruments DSC Q2000 and TGA Q500, respectively. Micro-images were captured using an optical microscope (Nikon ECLIPSE LV100ND). The surface topography of the polymeric layer was measured using atomic force microscopy (AFM, Multimode 8, Bruker). The contact angles and surface energies of the thin polymeric films were measured using a PerkinElmer LAMBDA-900 and extracted from the calculation of surface tension using two reference liquids, deionized (DI) water and diiodomethane (DII):

$$1 + \cos \theta = \frac{2(\gamma_{\rm s}^{\rm d})^{\frac{1}{2}}(\gamma_{\rm lv}^{\rm d})^{\frac{1}{2}}}{\gamma_{\rm lv}} + \frac{2(\gamma_{\rm s}^{\rm p})^{\frac{1}{2}}(\gamma_{\rm lv}^{\rm p})^{\frac{1}{2}}}{\gamma_{\rm lv}},\tag{1}$$

where  $\theta$  is the contact angle, and  $\gamma_{\rm lv}$ ,  $\gamma$ sd, and  $\gamma$ sp are the surface tensions of the reference liquid, corresponding dispersion, and polar components, respectively.<sup>27</sup> X-ray photoelectron spectroscopy (XPS) was conducted at the Yeungnam University Center for Research Facilities. The dielectric characteristics (leakage current density and *k*-value) were determined using an

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# 3. Results and discussion

Fig. 1a shows the molecular structures of MBHCa-Ha and FPA-3F. The synthesized polymer was purified by reprecipitation using n-Hex, and its chemical structure was determined from its <sup>1</sup>H NMR spectrum. The number average molecular weight  $(M_n)$ and polydispersity index (PDI) determined by GPC were 509 kDa and <2.15, respectively. The TGA and DSC curves shown in Fig. S1a and b<sup>†</sup> demonstrate the thermal decomposition characteristics of the polymer; the polymer exhibited a glass transition temperature  $(T_{\alpha})$  value of 27 °C and high thermal stability, with a weight loss of 5% at 263 °C. Generally, the polar groups cause polymer chains to be mutually attractive and so increase  $T_{\rm g}$ , whilst large nonpolar side chains tend to hold them apart and so lower  $T_{g}$ .<sup>28</sup> Thus, the synthesized polymer has a relatively low  $T_{\rm g}$  value. However, only polar side functional groups introduced in polymeric insulators can cause the strong interaction between side polar groups. These materials can cause high dielectric constant characteristics even if they are strongly crystalline or amorphous like PVDF-based polymers.29 These properties can cause strong polarization and yield operation instability issues in the device. Our synthesized polymer possessed several large non-polar side chains with one polar side group. This creates a lot of free volume and provides space for the polar side chain to move, resulting in a high k value.<sup>22</sup>

With prepared materials, polymeric thin films were deposited using spin-casting and fabricated through the photoinitiated crosslinking process (254 nm UV irradiation).<sup>20</sup> Fig. S2† shows that crosslinked patterned FMBHCa-Ha layers were successfully fabricated through line-patterned photomask UV irradiation, indicating the successful crosslinking reaction between MBHCa-Ha and FPA-3F. FPA-3F containing azide moieties functions as a crosslinker with MBHCa-Ha polymers according to the following mechanism described in Fig. S2.† 254 nm UV light irradiation activates the azide moieties ( $-N^3$ ) as a reactive singlet nitrene ( $-N^1$ ) and produces inert nitrogen gas.<sup>25</sup> The singlet nitrene group can make connections with alkyl groups (C-H bonds) of the polymer. Through this process, these crosslinked and non-crosslinked (single MBHCa-Ha layer) thin layers were used to assess the polymeric surface properties.

Fig. 1b displays the AFM images of MBHCa-Ha and FMBHCa-Ha. Both had similar topographies and exhibited extremely smooth surfaces (root-mean-square (RMS) roughness < 0.5 nm), regardless of the presence of the crosslinking agent, FPA-3F. The smooth surface provides favorable conditions for the growth of organic semiconducting crystals on the top and non-trapping dielectric surfaces during OFET operation.<sup>8</sup> The phase separation phenomenon caused by the aggregation of FPA-3F was not observed in the AFM image of FMBHCa-Ha, owing to uniform mixing between the fluorine-based FPA-3F and MBHCa-Ha; MBHCa-Ha comprising a hydrophilic functional group such as an acrylic-based side chain well mixed with FPA-3F.<sup>20</sup> The surface energies of the samples were determined

through contact angle measurements using two test liquids (DI water and DII). As shown in Fig. 1c, the pristine MBHCa-Ha thin film and the crosslinked FMBHCa-Ha film exhibited surface energy values of 41.10 and 40.61 mJ m<sup>-2</sup>, respectively. Similar to the AFM data, the surface energy and contact angle values of the MBHCa-Ha and FMBHCa-Ha were not significantly different.

To verify the insulating properties of the polymeric layers, metal-insulator-metal (MIM) capacitors were prepared with 30 nm of Al and the spin-cast polymeric layers (MBHCa-Ha and FMBHCa-Ha), as shown in the inset of Fig. 1d. The change in the leakage current density value as a function of the applied electric field is shown in Fig. 1d. Both cases displayed highly durable insulation properties until 4 MV cm<sup>-1</sup>, exhibiting leakage current density values of less than  $10^{-8}$  A cm<sup>-2</sup> at 4 MV cm<sup>-1</sup>. As the allowable leakage current density limit in the industry is  $10^{-6}$  A cm<sup>2</sup> at 2 MV cm<sup>-1</sup>, these results imply that the synthesized polymeric layers have relatively good insulation qualities and are suitable for use as gate insulation films for OFETs.

The *k*-values of the crosslinked and non-crosslinked polymers were extracted from the capacitance measurements in the frequency range of 0.1 to 1000 kHz. The *k*-values were calculated using the following equation:

$$k = \frac{C_{\rm i}d}{\varepsilon_0} \tag{2}$$

where  $C_i$  is the capacitance per unit area, d is the capacitor thickness, and  $\varepsilon_0$  is the vacuum permittivity. The calculated frequency-dependent k-values are shown in Fig. 1e. The k-values of the MBHCa-Ha and FMBHCa-Ha at 1 kHz were determined to be 7.9 and 7.2, respectively. These values are greater than those of typical polymer insulators ( $k \sim 2-4$ ) used in electronic devices.<sup>30</sup> The high-k properties of the polymers were induced by the dipolar segments (acrylate), which could be aligned under an electrical field.9 High-k and durable insulation properties are highly conducive to the manufacture of low power consumption devices and guarantee good insulation qualities in the gate insulator of OFETs, inducing good on/off current ratio and stability characteristics. Additionally, the dielectric constant values did not fall significantly, even at higher frequencies (the films exhibited almost constant k-values over a wide frequency range). The results indicate that these polymer layers can be applied to practical OFET gate dielectrics without causing operational instability phenomena such as hysteresis by dipolar disorientation.31

To test the availability of the synthesized polymeric layers, a bottom-gate top-contact OFET was fabricated. As shown in Fig. 2a, the Al gate electrodes were first coated on the substrates, and the fabrication of the OFET was completed by the sequential deposition of MBHCa-Ha/FMBHCa-Ha, C10-DNTT, and finally Au S/D electrodes. The channel length (*L*) and width (*W*) were 100  $\mu$ m and 1500  $\mu$ m, respectively. The *C*<sub>i</sub> values of the polymeric layers were measured using an LCR meter which yielded values of 43 and 47 nF cm<sup>-2</sup> for MBHCa-Ha and FMBHCa-Ha, respectively. The electrical properties of the fabricated devices were measured in terms of their transmission and output characteristics. The drain current in the saturation



Fig. 1 (a) Molecular structure of MBHCa-Ha and FPA-3F, polymeric thin film (MBHCa-Ha and FMBHCa-Ha) characteristics: (b) AFM image, (c) surface energy and contact angle values of DI water and DII, (d) leakage current density *vs.* electrical field (inset shows the capacitor device OM image), and (e) *k* values depending on frequency.

zone was monitored in response to a change in the gate voltage  $(V_{\rm G})$  from 20 V to -20 V, while a drain voltage  $(V_{\rm D})$  of -20 V was applied continuously to measure the transfer characteristics. The main electrical parameters of the OFETs, such as the field-effect mobility ( $\mu_{\rm FET}$ ), threshold voltage ( $V_{\rm th}$ ), and ON-current

 $(I_{on})$ /OFF-current  $(I_{off})$  ratio, were determined using the transfer characteristic graph (Fig. 2b) and the following equation:

$$I_{\rm D} = \frac{W}{2L} \mu_{\rm FET} C_{\rm i} (V_{\rm GS} - V_{\rm th})^2 \tag{3}$$



Fig. 2 (a) Schematic image of OFETs with polymeric dielectrics and electrical properties of devices: (b) transfer and (c) output curve (red: MBHCa-Ha, blue: FMBHCa-Ha).

Table 1	Electrical parameters of	of OFETs with	MBHCa-Ha/FMBH	ICa-Ha polymeric	gate dielectrics
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	$C_{\rm i} \left( \rm nF \ cm^{-2} \right)$	$\mu_{\rm FET} \left( { m cm}^2 \ { m V}^{-1} \ { m s}^{-1}  ight)$	$I_{\rm on}/I_{\rm off}$	$V_{\rm th}$ (V)	SS (mV dec <sup><math>-1</math></sup> )
МВНСа-На	43	$0.15\pm0.01$	${\sim}10^8$	$4.38\pm0.31$	412
<b>FMBHCa-Ha</b>	47	$1.33\pm0.05$	${\sim}10^8$	$-1.34\pm0.17$	612

where  $I_{\rm D}$  denotes the drain current. Table 1 summarizes the electrical performance parameters of the OFETs assembled in this study. For the output characteristics, the  $I_{\rm D}$  was measured as the  $V_{\rm D}$  changed under a stepped  $V_{\rm G}$  (0 V to -20 V at -5 V intervals), as shown in Fig. 2c.

The observed electrical properties (transfer and output curves) of the crosslinked devices were clearly superior to those of the non-crosslinked devices. The MBHCa-Ha devices exhibited an  $I_{\rm on}$  of  $\sim 1.5 \times 10^{-5}$  A, while the FMBHCa-HA devices exhibited an  $I_{\rm on}$  of  $\sim 1 \times 10^{-4}$  A, even with similar  $C_{\rm i}$  values. This order of magnitude difference between the on-current values caused a large difference in the  $\mu_{\rm FET}$  value (0.15 cm<sup>2</sup> V<sup>-1</sup> s<sup>-1</sup> for MBHCa-Ha and 1.33 cm<sup>2</sup> V<sup>-1</sup> s<sup>-1</sup> for FMBHCa-Ha) and the output curve. The only difference in device configuration was the addition of the crosslinking agent (FPA-3F) and the subsequent crosslinking reaction; the resulting performance improvements were attributed to the degree of crosslinking.

As shown in Fig. 3a, the addition of FPA-3F led to slight vertical phase separation by surface energy minimization.<sup>22</sup> The wide-range XPS data shown in Fig. 3b show the contents of the MBHCa-Ha and FMBHCa-Ha layers. As the pristine polymer MBHCa-Ha is only composed of oxygen and carbon (as per Fig. 1a), the observed atomic contents were primarily carbon and oxygen, while additional fluorine was observed in the crosslinked FMBHCa-Ha. Atomic weight distributions of the surface, extracted from the XPS data, showed that the percentages of carbon, oxygen, and fluorine were 72.47, 27.17, and 0.36%, respectively, for MBHCa-Ha. It can be confirmed that there was a decrease in oxygen-attached functional groups on the polymer surface owing to the addition of the crosslinking agent.

The crosslinking reaction and subsequent change in atomic weight distribution caused a significant change in the morphology of the organic semiconductor, C10-DNTT. Fig. 3c



Fig. 3 (a) Schematic illustration of FMBHCa-Ha in OFETs, (b) wide range XPS data of the polymeric layer, (c) topographic image of C10-DNTT deposited on MBHCa-Ha and FMBHCa-Ha, device operation stability: transfer curves with (d) sweep rate and (e) bias stress.



Fig. 4 (a) Schematic and real image of the flexible FMBHCa-Ha based OFET, (b) leakage current stability of FMBHCa-Ha under bending conditions (inset shows the OM image before/after bending 1000 cycles), (c) electrical characteristics of prepared devices: tranfer (left), output (middle), and bias-stress stability (right), and (d) distribution of field-effect mobility of flexible electronics.

shows a topographic image of the 50 nm C10-DNTT layers deposited on MBHCa-Ha and FMBHCa-Ha. Both cases exhibited mixed crystals composed of smooth pancake-shaped grains and sharp needle-like lamellar grains protruding on the surface. The direction of the lamellar grains indicates the orientation of C10-DNTT. As these individual grains were generated from different nuclei, the orientations of the grains were random.<sup>32</sup> However, the grains of C10-DNTT on FMBHCa-Ha were found to be much larger than those of C10-DNTT on MBHCa-HA. Although the surface energy values were similar between the two cases (as shown in Fig. 1c), the addition of the crosslinking agent caused an increase in the hydrophobic functional groups present on the surface, as indicated in the XPS data. Therefore, larger-grain crystals were formed. This result is similar to that in a previous report whereby an organic-inorganic hybrid material was synthesized with a similar surface energy of hydrophobic organic polymeric materials (e.g. cyclo olefin polymers), but with smaller crystals, than were formed in our hydrophobic organic polymer material.24,25,33 The resulting large-grain morphology of C10-DNTT, caused by deposition on our polymer dielectric, improves the charge transfer characteristics. This can be attributed to the presence of fewer charge-trapping sites, such as grain boundaries, between crystal grains.<sup>34</sup>

In addition to the electrical transport characteristics, the operation stability is also an important part. As shown in the transfer curve in Fig. 2b, regardless of whether a cross-linking agent was applied, both cases showed low hysteresis behavior.

Also, electrical operation stability tests were performed using two methods. Fig. 3d shows the sweep-rate-dependence transfer curve data for the MBHCa-Ha- and FMBHCa-Ha-based devices. The transfer curves were measured at different sweep rates of  $V_{\rm G}$ : fast sweep conditions were measured with a hold time of 0 s, a sweep delay of 0 s, and 101 data points; slow sweep conditions were measured with a hold time of 0 s, a sweep delay of 1 s, and 101 data points. There was no discernible variation in the degree of response of either device as a function of the gate bias sweep rate, implying that there was no dipole imbalance in the dielectric layers. In addition, the bias-stress stability was identified at ambient temperature and atmospheric pressure with a gate bias-stress (-20 V) at a specific time (0 min to 72 min). As shown in Fig. 3e, the shift in the transfer curve due to the gate bias stress was almost small in both cases (<1 V). The pristine case without the crosslinker induced the low trapping characteristics because of the presence of many non-polar alkyl groups in polymer's molecular structure. Besides, the addition of a fluorinated crosslinker case with surface modification still remained with excellent bias-stress stability properties. This was attributed to deep ionization energy due to the fluorinebased functional groups present on the surface of the crosslinked polymer (fluorine withdraws the electrons very well), which greatly increased the energy barrier between the ionization energy of dielectrics and the highest occupied energy level of C10-DNTT to be overcome for driving the p-type OFETs.35,36

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Additionally, the interfacial trap density  $(N_{\text{trap}})$  values of device through transfer curve data, including bulk and shallow traps, were calculated with the following equation:

$$\mathbf{V}_{\text{trap}} = \left(\frac{q\text{SS}\log(e)}{k_{\text{B}}T} - 1\right) \frac{C_{\text{i}}}{q},\tag{4}$$

where SS, q, and  $k_{\rm B}$  are the subthreshold swing, electronic charge and Boltzmann constant, respectively.<sup>35</sup> The calculated values were  $1.59 \times 10^{12} \, {\rm eV^{-1}} \, {\rm cm^{-2}}$  and  $2.73 \times 10^{12} \, {\rm eV^{-1}} \, {\rm cm^{-2}}$  for MBHCa-Ha OTFTs and FMBHCa-Ha OTFTs. These results from the electrical operation stability test indicate that our fabricated polymer layer with both non-crosslinked and cross-linked cases displayed stable operation under current operation conditions.

Based on the above analyses (performance and operation stability), we used the crosslinked polymer layers to manufacture practical flexible printed electronics. FMBHCa-Ha layers were printed on PET films to create a large-area OFET array, and its features were examined to confirm reproducibility (Fig. 4a). Before verifying the OFET properties, the leakage current density of the flexible MIM platform on the PET substrate was observed. Fig. 4b shows the leakage current density according to the bending cycles and the images captured while measuring the leakage current in a bending glass tube (radius 4 mm). Because the crosslinked polymeric layers were flexible, good insulation properties were maintained even under bending (leakage current density <  $10^{-8}$  A cm<sup>-2</sup> at 4 MV cm<sup>-1</sup> after 1000 bending cycles).



Fig. 5 (a) Flexible integrated logic device photo, OM, and circuitry images; NOT (left), NAND (middle), and NOR (right) gates, (b) VTC and voltage gain curve of NOT gates, and (c) input and output curves of NAND and NOR gates.

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A flexible OFET array was prepared, and its conventional electrical properties (transfer curve, output curve, and biasstress stability) were measured, as shown in Fig. 4c. Similar to rigid-platform OFETs, our p-type devices on flexible substrates exhibited stable operation and high uniformity in device performance. In addition, the transfer curves of the flexible OFET displayed robustness under bias-stress conditions, indicating excellent electrical stability. In particular, the OFETs on the flexible platforms exhibited average  $\mu_{\text{FET}}$  values of 0.94 cm<sup>2</sup> V<sup>-1</sup> s<sup>-1</sup>, with a standard deviation of 0.013 across 64 devices. The ability to fabricate a device with such excellent electrical stability with high reproducibility is promising for the potential large-scale integration of such a unit device.

To confirm this possibility, simple logic devices were prepared on the PET substrates. The manufactured circuit schematics and top-view OM images of the logic gates are shown in Fig. 5a. Diode-connected OFETs as the load transistors and standard ptype OFETs as the driving transistors were coupled in series to create NOT gates. The W/L ratios for the load and drive transistors were 2 (W: 100 µm; L: 200 µm) and 20 (W: 100 µm; L: 2000 µm), respectively. Fig. 5b depicts the operation behavior of our NOT gates, showing voltage transfer characteristics (VTC) and voltage gain curves with supply biases  $(V_{DD})$  of 10, 20, and 30 V. According to the VTC, the output voltage  $(V_{OUT})$  assumed values close to  $V_{\rm DD}$  under low input voltage  $(V_{\rm IN})$  conditions and values close to zero under high V<sub>IN</sub> conditions. Because the gate-source voltage of the driving transistor is small in a high  $V_{\rm IN}$  bias state, the load transistor is aggressively switched on, resulting in low current characteristics. Consequently, VOUT was reduced to zero. However, a low  $V_{\rm IN}$  bias state results in comparable gate-source voltages for both the drive and load transistors. In our device, the W/L ratio of the driving transistor was much larger than that of the load transistor, and  $V_{\rm OUT}$  was pushed up against  $V_{\rm DD}$ . Therefore, the "0" and "1" signals of the bias can be referred to as low and high voltage input or output conditions, respectively. The output of the NOT gates is the inverse of the input circumstance; therefore an input voltage signal of "0" results in an output voltage signal of "1", and vice versa. The VTC curve was used to extract the gain values of these NOT gates  $(dV_{OUT}/dV_{IN})$ , which showed maximum values of 9.89, 13.18, and 22.9 for 10, 20, and 30 V, respectively.

Furthermore, one load OFET and two drive OFETs were utilized to create integrated circuit NAND and NOR gates, as shown in Fig. 5a. Two p-type driving OFETs were connected in parallel with each other in the case of the NAND gates. We noticed that the output signal for the produced NAND gates was driven by various logic input signals (see the NAND output signal in Fig. 5c). Both OFETs were turned on when both inputs were in logic-low states, that is, when  $V_{\rm A}$  and  $V_{\rm B}$  were equal to 0 V, resulting in a straight link between  $V_{DD}$  and  $V_{OUT}$ . As a result, the output signal was logic-high. Similarly, when one of the inputs was logic-high and the other was logic-low, either of the parallel transistors was operated and  $V_{DD}$  and  $V_{OUT}$  were connected directly. Therefore, the output signal was logic-high. However, when both inputs were in logic-high states (10 V of input bias), none of the parallel transistors worked, causing the ground (GND) to connect directly to  $V_{OUT}$ . As a result, the output

signal of the NAND gates was in a logic-low state. Two p-type drive OFETs were connected in series to provide the NOR gates. The series-connected OFETs were turned on when both inputs were in a logic-low state of 0 V input bias at  $V_A$  and  $V_B$ , resulting in the establishment of a direct channel between  $V_{DD}$  and  $V_{OUT}$  (see the NOR output signal in Fig. 5c). In this case, the output signal was in a logic-high state. When any of the inputs was logic-high (greater than 30 V), one or more drive transistors were turned off, causing  $V_{DD}$  and  $V_{OUT}$  to be disconnected. Consequently, the signal output was logic-low. This operation coincided with that of conventional logic gate drives, indicating that our crosslinked dielectric OFETs were successfully applied to practical integrated circuits.

# 4. Conclusion

In this study, we synthesized high-k polymeric materials with various functional segments and applied a fluorine-based azide organic crosslinker to yield crosslinked dielectric layers. Using these prepared thin films, we could successfully manufacture OFETs with dramatically enhanced electrical properties. The crosslinking of the dielectric polymer induced a hydrophobic surface, decreasing the oxygen-based functional groups at the surface and yielding a favorable interfacial environment for the growth of organic crystals. In addition, this fluorine-based interface still remained the energy barrier between the dielectric and p-type organic semiconductors, thereby inducing excellent bias-stress stability. Moreover, we confirmed the flexibility of the crosslinked layers and applied them to the manufacture of flexible and printed electronic circuits for unit and integrated devices. Therefore, our research effectively demonstrates a method for the synthesis and preparation of polymeric dielectric layers for practical, flexible, and printed organic electronics.

# Conflicts of interest

There are no conflicts to declare.

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