Lyotropic Liquid-Crystalline Mesophase of Lithium Triflate–Nonionic Surfactant as Gel Electrolyte for Graphene Optical Modulator

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Supporting Information

ABSTRACT: Lithium salt (noncoordinating anions, such as lithium triflate (Ltf)) gel electrolytes may be key for the practical use of electrochemical devices. We introduce a new lyotropic liquid-crystalline (LLC) mesophase using Ltf, a small amount of water (as low as 1.3 water per Ltf), and nonionic surfactant ($C_{18}H_{37}(OCH_2CH_2)_{10}OH$, $C_{18}E_{10}$). The LLC phase forms over a broad range of Ltf/ $C_{18}E_{10}$ mole ratios, 2–18. The clear ethanol solution of the ingredients can be either directly spin-coated over a glass substrate to form a gel phase or it can be prepared as a gel by mixing Ltf, water, and $C_{18}E_{10}$. The mesophase leaches out surfactant molecules at low salt concentrations, but at a salt/surfactant mole ratio of above 8, the phase is homogeneous with a cubic mesostructure, fully transparent in the visible optical region, mechanically flexible, and an effective gel electrolyte. We have observed a large electrostatic doping on graphene with the Fermi energy level of ~1.0 eV using Ltf- $C_{18}E_{10}$



gel electrolytes. The Ltf-based gels demonstrate better properties than commonly used ionic liquid electrolyte in graphene optical modulators. The stability of the new gel electrolytes and their superior performance make them suitable electrolytes for use in graphene-based optical modulators.

INTRODUCTION

Lyotropic liquid-crystalline (LLC) mesophases consist of at least two components, such as a solvent and a surfactant. The solvent may be water,^{1,2} water–oil,³ aqueous solutions,⁴ ionic liquids,^{5–7} acids,^{8,9} and salts.^{10–12} The salts can be alkali metals (LiX, X is NO₃⁻, Cl⁻, Br⁻, I⁻),^{11,12} alkaline earth metals (CaX₂, X⁻ is Cl⁻ and NO₃⁻),¹² transition metals ($[M(H_2O)_n](NO_3)_2$) where M represents Mn(II), Co(II), Ni(II), Zn(II), and Cd(II)),¹⁰ or lanthanides (Eu(II) and La(III)).¹³ The common properties of these salts are their low melting point or high solubility.^{11,12} It is also important to note that the melting points of the salts drop¹⁰ and their solubility is enhanced^{11,12} in confined spaces, such as in the hydrophilic domains of the mesophases, known as soft confinement effect.¹⁴ Even though most lithium salts have a high melting point, because of their high solubility, they form LLC phases with nonionic surfactants.^{11,12}

The salt–surfactant LLC mesophases are important in the development of gel-electrolytes for practical electrochemical devices, ¹⁵ such as solar cells, ¹⁶ batteries, ¹⁷ supercapacitors, ⁸ and so on. Therefore, it is important to investigate the salt–surfactant mesophases, especially the lithium salt–surfactant mesophases, toward the development of gel-electrolytes for electrochemical systems. Moreover, the lithium salts of noncoordinating anions (such as BF₄⁻, PF₆⁻, and CF₃SO₃⁻) are very important due to their high ionic conductivity.^{18–20} However, the supersaturated solutions of lithium salts of these

ions have not been investigated as a solvent in the salt–surfactant LLC mesophases. For instance, $LiCF_3SO_3$ is a very soluble salt in water and can be a good candidate for a stable LLC mesophase, which may be used as the gel-electrolyte in various super capacitors and battery systems.

For example, a phosphoric acid-surfactant mesophase, used as a gel-electrolyte, has been recently tested in a graphene optical modulator, a supercapacitor consisting of two graphene electrodes and a gel-electrolyte, with excellent results.⁸ Polarizing the electrolyte by an applied electric field between the two graphene electrodes effectively dopes the graphene and significantly shifts its Fermi level to effectively modulate the optical transmittance of graphene.^{8,21} Major challenges in the device configuration include finding an electrolyte that (i) has a large electrochemical window, (ii) shifts the Fermi level of graphene greater than 1 eV for visible optoelectronic applications, (iii) reaches very large charge densities on graphene, (iv) is transparent in the visible region, and (v) is mechanically flexible for solid state device applications. For these reasons, ionic liquids have been considered target materials.²¹ However, gel-electrolytes that perform as well as ionic liquids may be more beneficial for use in practical applications.

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In this contribution, we have developed a new LLC mesophase of $\text{LiCF}_3\text{SO}_3-\text{C}_{18}\text{H}_{37}(\text{OCH}_2\text{CH}_{2})_{10}\text{OH}$ (denoted as Ltf-C₁₈E₁₀) that can be used as a gel electrolyte in graphene optical modulators. The mesophase is very stable over a broad range of Ltf/C₁₈E₁₀ mole ratios, characterized using various techniques (such as XRD, POM, FTIR, and conductivity measurements). Its performance, as LLC gel electrolytes, has been demonstrated in an optical modulation of graphene in the visible region. The new electrolyte developed here, applicable in graphene optical modulators, has many advantages such as ease of device fabrication, minimal electrolyte leakage, excellent electrochemical properties, high charge density on graphene, mechanical flexibility, produced under ambient conditions, transparent in the visible region, stable over long periods (more than 3 months as measured in this study), and so on.

EXPERIMENTAL PART

Preparation of Ltf-C₁₈E₁₀ **Solutions.** In a typical preparation, the required amount of Ltf was first dissolved in ethanol, and following that the required amount of $C_{18}E_{10}$ was added and stirred overnight at RT to obtain a clear solution. For example, the sample with $2Ltf/C_{18}E_{10}$ was prepared as follows. First, 2 mmol (312 mg) of Ltf was dissolved in 5 g EtOH. Subsequently, 1 mmol (711 mg) of $C_{18}E_{10}$ was added to the above solution. The clear solutions were then coated on glass slides via spin coating at 1000 rpm for 15 s to obtain LLC mesophases.

Preparation of Ltf-C₁₈E₁₀ **LLC Mesophases.** The required amounts of Ltf, water, and C₁₈E₁₀ were placed in a vial and sealed using Teflon tape. As an example, the sample with $4Ltf/C_{18}E_{10}$ and $3H_2O/Ltf$ was prepared by mixing 4 mmol (624 mg) of Ltf, 1 mmol (711 mg) of C₁₈E₁₀, and 12 mmol (216 mg) of H₂O in a vial. The vial was kept at 80 °C in a water bath for 24 h to obtain homogenized gels.

Synthesis of Graphene. Graphene layers on ultrasmooth copper foil substrates (Mitsui Mining and Smelting Company, B1-SBS) were synthesized using chemical vapor deposition (CVD), as described in our previous publications.^{8,21} We are able to synthesize single and multilayer graphene in a large area (several cm²). During the annealing step, the copper oxide on the copper substrate was removed by sending H₂ gas with a flow rate of 100 sccm. The graphene layer on the copper substrate was synthesized at 1035 °C and 5 Torr under a CH₄ flow rate of 10 sccm. After 30 s of growth, the growth was stopped and cooled to room temperature.

AC Conductivity Measurements. AC conductivity measurements were taken using a Gamry G750 potentiostat/ galvanostat operating at an AC voltage of 10 mV and 100 kHz. The FTO glass electrodes were prepared as described in our previous publications.¹⁶ The cell constant was determined using standard KCl solutions with a known conductivity, each time before the measurement of the gels. The solutions were dropped over the prestretched line over FTO glass. Upon evaporation of the solvent (~24 h), the counter and reference electrodes were attached to one side and working electrode was attached to the other side for the conductivity measurements.

XRD Measurements. Small-angle XRD patterns were collected using Miniflex diffractometer, equipped with high-power Cu K α source operating at 30 kV/15 mA and wavelength of 1.5405 Å. Film samples that were spin-coated over glass slides were used for the XRD measurements.

ATR-FTIR Measurements. The FTIR spectra were recorded using a Bruker Alpha Platinum ATR in the 400–4000 cm⁻¹ with a 4 cm⁻¹ resolution and conducting 32 scans. A few drops of solution were placed on the diamond ATR crystal, and the spectra were followed for 2 days until all ethanol was evaporated and no spectral changes were identified.

Determination of Water Amounts in the Gels. Around 1 g of the clear solution was placed onto a plastic tray and the mass of the solution was monitored for 2 days until there was no significant change in the mass of the sample.

Optical and Electrical Measurements. Graphene layers were transferred to glass substrates; then, graphene-based optical supercapacitors²¹ were fabricated. The graphene layers synthesized on a copper substrate were covered with Shipley 1813 photoresist liquid and then annealed at 60 °C overnight. To transfer the graphene from photoresist layer to a glass surface, the copper film attached to the graphene was chemically etched in 1 M FeCl₂ aqueous solution and then washed with excessive amount of water. Subsequently, the graphene layers attached to the photoresist were placed on a glass surface and heated to 80 $^\circ\bar{C}$ for a few minutes on a hot plate. Following this the photoresist layer was removed with acetone and washed with an ample amount of isopropanol. Another glass substrate with a ~100 nm gold film and graphene-covered glass substrate were then joined. The counter electrodes were separated by a 100 μ m thick plastic spacer, a double-sided adhesive tape stripe. The space between the graphene and gold electrodes was filled with an electrolyte. The ionic liquid electrolyte used in this study is diethylmethyl(2methoxyethyl) ammonium bis(trifluoromethylsulfonyl)imide. Under the applied bias voltage, the electrolyte generates electrical double layer (EDL) at the interface between graphene and electrolyte. The optical properties (transmission) of singlelayer graphene were measured in the visible and near-infrared wavelength as a function of the applied bias voltage using a Fourier transform infrared spectrometer (Bruker VERTEX 70v). Lithium-salt-based LLC gel electrolytes were inserted into the graphene supercapacitor by heating the device and electrolyte to 70 °C. ~20 μ L of the synthesized electrolyte was placed at the edge of the device; then, the electrolyte flows in the narrow space between the electrodes due to the capillary forces; see Figure S1 for the details. Capacitance and total resistance of the graphene capacitor were measured by using LCR Meter (Hewlett-Packard, 4284A). Bias voltages to graphene and gold electrodes were applied by using a Source Meter (Keithley, 2400).

RESULTS AND DISCUSSION

The LLC mesophases of lithium triflate (LiCF₃SO₃, Ltf) and brij 76 ($C_{18}H_{37}(OCH_2CH_2)_{10}OH$, $C_{18}E_{10}$) form over a broad range of salt/surfactant mole ratios (2–18, denoted as #-Ltf- $C_{18}E_{10}$, where # represents Ltf/ $C_{18}E_{10}$ mole ratio), corresponding to a ratio as high as 82% (w/w). This indicates that a major component of the mesophase could be the salt species. The mesophases can be prepared as a solution in excess ethanol, which can be completely evaporated upon spin coating over any substrate to obtain the gel LLC phase or prepared as a gel starting from salt, surfactant, and small amount of water. In general, when using the second method, it is more difficult to produce a homogeneous mixture of the LLC phase; however, it is more practical to use as a gel electrolyte in electrochemical applications.



Figure 1. XRD pattern of (a) 2-Ltf- $C_{18}E_{10}$ with aging (I) fresh, (II) 1 day, (III) 1 week, and (IV) 2 weeks. (b) 8-Ltf- $C_{18}E_{10}$ mole ratio with aging (I) fresh, (II) 1 day, (III) 1 week, and (IV) 5 weeks. (c) #-Ltf- $C_{18}E_{10}$ with increasing # from 2 (top) to 18 (bottom) with an increment of 2.



Figure 2. Normalized ATR-FTIR spectra of (a,b) #-Ltf- $C_{18}E_{10}$ with increasing Ltf (Ltf/ $C_{18}E_{10}$ mole ratio increases from 2 to 18 with four increments, bottom is the spectrum of pure $C_{18}E_{10}$). ATR-FTIR spectra of xLtf/ $C_{18}E_{10}$ during aging: x is (c) 2, (d) 10, and (e) 18 (fresh and 4 h). (f) Normalized spectra after 24 h of aging of the same samples.

Both mesophases, prepared as gels and solutions, were characterized using POM, XRD, and FTIR techniques to elucidate the structural details as well as AC conductivity measurements for the electrical characterization. Figure 1 shows a series of small-angle XRD patterns of samples with different concentration of Ltf. Notice that both films, prepared by spin coating or spreading over a substrate, diffract at small angles with no diffraction at high angles. In fact, both preparation methods provide the same LLC phase upon aging.

The samples, prepared using ethanol solution, have been monitored by recording their XRD patterns over 5 weeks to ensure homogeneous LLC phase through complete ethanol evaporation. Figure 1a,b shows the aging process of two samples, namely, 2 and 8 Ltf and 1 $C_{18}E_{10}$. The diffraction lines observed immediately after spin coating undergo some changes, while keeping the line position relatively constant. At low Ltf concentrations, there are two sets of diffraction lines at small angles, which eventually become a single set at higher concentrations; see Figure 1. The small-angle XRD pattern of 2-Lt- $C_{18}E_{10}$ displays lines at 1.48, 1.77, 2.99, and 4.54°, 2 θ values. The diffraction line at 1.77° likely corresponds to a free surfactant, which gradually disappears at higher Ltf concentrations. The free surfactant line appears upon coating a clear solution of the salt-surfactant, gradually increases after 1 h, and then gradually decreases over time with further aging the gel films. It is likely that the mesophase leaches out crystalline

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surfactants, which gradually become disordered or molten over time due to the thermal energy (melting point of free surfactant is ~38 °C). Note also that the XRD pattern of the free surfactant, spin-coated from its clear ethanol solution, diffracts at ~1.77°, which also gradually disappears over time, similar to our samples. Besides, the surfactant leach from the LLC media has also been observed in other lithium salt–pluronic LLC phases.²²

The 2-Ltf-C₁₈E₁₀ sample has also been monitored using POM, and it displays birefringent and dark domains, likely due to the free surfactant and LLC domains, respectively. It is worth noting that the freshly prepared films, before surfactant separation, display focal conic fan texture under POM. It means that the mesophase is 2D hexagonal at low salt concentrations but is metastable and undergoes phase separation into salt-rich LLC and salt-free surfactant domains over time. Therefore, the salt concentration in the LLC domains is much higher than $2Ltf/C_{18}E_{10}$. The free surfactant line gradually disappears with an increasing Ltf/C₁₈E₁₀ mole ratio in the samples, and at a mole ratio above $8Ltf/C_{18}E_{10}$, all diffraction lines can be indexed to a cubic mesophase. Also, this behavior correlates with the POM images that appear dark between the two crossed polarizers. Note that the diffraction lines gradually shift to lower angles with an increasing Ltf/ $C_{18}E_{10}$ mole ratio. If we consider the line at ~1.34°, 2 θ , originating from the (200) plane of the cubic phase, the unit -cell parameter increases from 120 to 142 Å by increasing Ltf/ $C_{18}E_{10}$ mole ratio from 2 to 18, respectively; see Figure 1c. However, the changes of the unit-cell parameters, at low Ltf/ C18E10 ratio, are negligibly small and gradual at higher ratios. It is likely that the Ltf/C₁₈E₁₀ ratio in the LLC mesophase is constant to a mole ratio around 8 and then gradually increases at higher salt concentrations, where the extra surfactant molecules in the low Ltf concentrations are leached out from the media to compensate for a constant salt/surfactant ratio.

Furthermore, the salt-surfactant mixtures display mesophases in the following order V_1 (bicontinuous cubic), H_1 (normal hexagonal), and I_1 (micellar cubic), with a dominating hexagonal phase, with an increasing salt/surfactant mole ratio. Therefore, it is reasonable to conclude that the hexagonal phase of the Ltf-C₁₈E₁₀ system is metastable and undergoes a phase separation into micellar cubic phase and free surfactant at low $Ltf/C_{18}E_{10}$ mole ratios. Besides, the diffraction patterns in Figure 1c show some variations during the aging process; however, all diffraction lines are indexed to a micellar cubic phase, where the lines at 1.39, 1.70, 2.77, and 4.16°, 2θ , correspond to (200), (211), (400), and (600) planes, respectively. The line, corresponding to the (200) plane, remains in all of the samples but gradually loses its intensity and shifts to smaller angles due to increase in unit-cell size in the mesophase to accommodate the excess salt species. Interestingly, the mesophase in the $Lft-C_{18}E_{10}$ is cubic in all compositions, which is quite unusual for salt surfactant mesophases.

In addition, we recorded the ATR-FTIR spectra of a series of samples during solvent evaporation with different $Ltf/C_{18}E_{10}$ mole ratios. Figure 2a,b shows a series of spectra after aging the samples for 24 h, where the mesophase no longer undergoes changes. The peaks due to the Ltf species increase with an increasing salt concentration in the media. However, it is difficult to spectroscopically quantify the free surfactant in the media because the spectrum of the surfactant changes to the final spectrum upon the addition of as little as 0.1 salt/

surfactant to the media. This means that the initial conformation of the ethylene oxide units changes in the presence of a 0.1 salt per surfactant, and the rest is an assembly of the surfactant with the help of LiCF₃SO₃xH₂O. Notably, the Ltf peaks dominate the spectra. For instance, the sharp peak at 1035 cm⁻¹ is known to originate from the $\nu_s(SO_3)$ mode of the free triflate ion, and the tail on the high-energy site of this peak corresponds to the same mode in an ion pair. Figure 2b clearly shows that the triflate species are predominantly free ions. This is important for the ionic conductivity of the gel phase; see the latter. The intense peaks at 640, 1176, 1232, and 1255 cm⁻¹, due to $\delta(SO_3)$, $\nu_{as}(CF_3)$, $\nu_s(CF_3)$, and $\nu_{as}(SO_3)$, respectively, and a tail on the high-energy side of 1255 cm^{-1} , broken into two components at \sim 1272 and \sim 1300 cm⁻¹, originate from the asymmetric stretching mode of the SO₃ unit of the free and coordinated triflate ion, respectively.^{23,24}

Figure 2c-f shows the surfactant ν CH stretching and waterbending and -stretching regions of three samples in two different stages of the aging process. The fresh samples display weak water bending at $\sim 1649 \text{ cm}^{-1}$ and a broad waterstretching feature at 3440 cm⁻¹ with a shoulder around 3200 cm⁻¹, characteristic of the bulk water (hydrogen-bonded network of water). The ν CH stretching region displays peaks related to the surfactant molecules and ethanol. However, upon complete evaporation of ethanol, both the bending and stretching modes of the water display drastic changes; compare the spectra in Figure 2c-e. The bending mode gradually red shifts and becomes more intense and sharper, while stretching modes are blue-shifting, from 1649 to 1635 cm⁻¹ and 3440 to 3510 cm^{-1} , respectively, indicating that the water is becoming free or hydration-only water;²⁵ compare the spectra in Figure 2f. The bending mode also becomes sharper with an increasing Ltf in the media. The stretching to bending intensity ratio also displays drastic changes, from 2.5 to 1.0 increasing from 2 to $18Ltf/C_{18}E_{10}$ mole ratio, likely indicating the change in the bulk/hydration water ratio in the media. To support this observation and also to determine the water content of the samples, we monitored the weight change over a four-digit balance during ethanol evaporation of the solutions. Owing to the very fast evaporation of ethanol, the mass of the solution decays exponentially; see Figure 3a. After mesophase formation, the mass of the solution becomes constant. Indeed, the LLC mesophase stabilizes at around 2H₂O/Ltf for both 2-Ltf-C₁₈E₁₀



Figure 3. Variation of (a) the mass and (b) the conductivity of the samples with a $Ltf/C_{18}E_{10}$ mole ratio of between 2 and 18.



Figure 4. Application of Ltf based LLCs gel electrolytes for graphene optical modulators. (a) Schematic representation of a graphene-gold supercapacitor filled with Ltf salt-based liquid-crystalline gel electrolyte. The LLC system has $Ltf/C_{18}E_{10}$ mole ratio of 8. (b) Variation of normalized transmission of the graphene supercapacitor filled with Ltf salt-based liquid-crystalline gel electrolyte versus the wavelength of the incident light for various applied bias voltages. (c) Variation of Fermi energy level of graphene as a function of applied bias voltages for varying $Ltf/C_{18}E_{10}$ mole ratio in LLC gel electrolytes and typical ionic liquid used in graphene optical modulators.

and 4-Ltf- $C_{18}E_{10}$ and at around 1.3H₂O/Ltf for both 8-Ltf- $C_{18}E_{10}$ and 18-Ltf- $C_{18}E_{10}$.

Remarkably, the water/Ltf mole ratio decreases to almost 1.3 around 8 Lft/ $C_{18}E_{10}$ and remains constant above a mole ratio of 8. These observations are consistent with the behaviors observed in the XRD patterns as well as the discussions in the XRD Measurements section. Therefore, it is reasonable to conclude that the Ltf- $C_{18}E_{10}$ system is metastable up to an 8 Ltf/ $C_{18}E_{10}$ mole ratio and leaches out surfactant with some bulk-like water.

Additionally, we recorded the AC conductivity of a set of #-Lft- $C_{18}E_{10}$ during aging using an FTO substrate that was designed for this purpose.¹⁶ The AC ionic conductivity increases slowly in samples with a low Lft concentration and increases gradually after a mole ratio of 6; see Figure 3b. Notably, the Ltf-C₁₈E₁₀ mesophase displays the highest AC ionic conductivity among all lithium salt-surfactant systems investigated in the literature.^{12,22} The highest conductivity was reported for the 18-Lft-C₁₈E₁₀ to be 2.6 mS/cm. Notice that there is a factor of 3 compared with the polyelectrolytes of Lft using poly(ethylene oxide)s.²⁶⁻²⁸ There, the Lft has a limited solubility and also a stronger ion-ethylene oxide interaction that likely reduces the mobility of lithium ion in the media. However, in the lithium salt-surfactant systems, the salt is in the molten phase in the presence of a small amount of water with better ionic conductivity. It is likely that the lithium ion mobility is also enhanced in the confined space due to thermodynamic effects that need to be further investigated.^{29,30}

We employed the Ltf- $C_{18}E_{10}$ LLC mesophase as a gel electrolyte in a graphene optical modulator to evaluate the electro-optical performance of the LLC gels. For this purpose, five compositions were chosen with an Ltf/ $C_{18}E_{10}$ mole ratio of 4, 6, 8, 10, and 12. Figure 4a displays the schematic representation of the graphene-based supercapacitor structure used as an optical modulator. A gold film on a glass surface and graphene layer on an another glass surface form the capacitor structure (Figure S1). The electrolyte was placed between the capacitor plates by heating to a temperature higher than gel electrolyte's melting point (~75 °C) and then placing the liquid electrolyte on preheated graphene capacitor device; see Figure S1. It should be noted that the graphene–gold supercapacitor works by ion gating. Owing to the electric double-layer (EDL)

formation, the electric field is screened very close to the graphene surface (only a few nanometers) by the mobile ions in the medium. The fractional variation of the transmission of the incident light through the device as a function of the applied gate voltages is shown in Figure 4b and Figure S2. It should be noted that the ideal suspended graphene in air absorbs only $\Pi \alpha$ = 2.3% (α is the fine structure constant) of the incident light.³¹ When graphene is placed on a substrate such as a glass substrate in our case, the graphene absorbs 1.45% (corrected absorption = $\Pi \alpha (2/(1+n))$,² where *n* is the refractive index of BK7 glass at 600 nm, 1.5163) of the incident light,³² which is in close agreement with our observed variation (\sim 1.6%) in Figure 4b. This value should decrease with increasing the defect density on graphene. Because of the Pauli blocking, the graphene becomes transparent for incoming light at energies smaller than twice the Fermi level shift.³³ Note that the Pauli blocking happens when the final optical transition states are occupied because of the increase in the charge density on graphene. The observed Fermi level shift increases linearly with the applied gate voltages. The mole ratio of the $Ltf/C_{18}E_{10}$ gel electrolyte used in Figure 4b is 8. Also, a series of samples with varying composition of Ltf/C₁₈E₁₀ mole ratio were prepared and their electro-optical performances have been tested (Figure 4c). Remarkably, we observed that the $Ltf/C_{18}E_{10}$ mole ratio of 8 has the best performance in the graphene optical modulators. In addition, the fabricated device showed a similar performance after 3 months; see Figure S6. The capacitance and resistance of all compositions have been measured (see the Supporting Information, Figures S2–S6). In addition to the Ltf salt, we also prepared and used LiCl, LiBr, and LiNO3 salt-based LLC mesophases as electrolytes in graphene supercapacitors. However, we were only able to insert an LiCl-based electrolyte (due to high melting points of those mesophases) into the device, which performed very weakly when compared with the Ltf-based gel electrolytes; see Figure S2f. Figure 4c demonstrates the optical performance of all Ltf/C₁₈E₁₀ compositions and a commonly used ionic liquid in graphene supercapacitors; see Figure S7 as well. We observed that an Ltf/ $C_{18}E_{10}$ mole ratio of 8 displays the best performance in graphene optical modulators.

The Journal of Physical Chemistry C

CONCLUSIONS

A new LLC mesophase of Ltf-C₁₈E₁₀ has been prepared, characterized, and employed as a gel electrolyte in graphenebased optical modulators. A new LLC system consists of Ltf with a very small amount of water (as low as 1.3 water molecules per Ltf) and a nonionic surfactant ($C_{18}E_{10}$). The LLC mesophase forms over a broad range of Ltf/C₁₈E₁₀ mole ratios (2-18). The first step in the preparation is to obtain a clear ethanol solution of Ltf and $C_{18}E_{10}$. After obtaining the solution, the solvent, ethanol, is evaporated by spin coating over a glass substrate, and the gel phase is immediately formed. The mesophase is transparent in the visible region of the electromagnetic spectrum, mechanically flexible, and stable over a broad range of Ltf/C18E10 mole ratios. The LLC gel electrolyte displays good ionic conductivity and an excellent electro-optical performance in the graphene-based optical modulators. Maximum Fermi energy shift in graphene is observed in the $8Ltf/C_{18}E_{10}$. The LLC gel electrolyte has many advantages in terms of device fabrication, electrolyte leakage, electrochemical properties, etc. The results pave the way for the development of gel electrolytes, which can increase the Fermi energy level shift of graphene to allow for the realization of graphene-based optoelectronics in the visible region of the electromagnetic spectrum.

ASSOCIATED CONTENT

Supporting Information

The Supporting Information is available free of charge on the ACS Publications website at DOI: 10.1021/acs.jpcc.7b03622.

Schematic representation and photographs of the electrodes. More electrical and optical characterization data of the electrodes with gel electrolytes and ionic liquid. (PDF)

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Notes

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