Highly Proton Conductive Phosphoric Acid—Nonionic Surfactant Lyotropic Liquid Crystalline Mesophases and Application in Graphene Optical Modulators

Ebrima Tunkara,† Cemal Albayrak,† Emre O. Polat,‡ Coskun Kocabas,‡ and Ömer Dag*,†

†Department of Chemistry and ‡Department of Physics, Bilkent University, 06800, Ankara, Turkey

ABSTRACT Proton conducting gel electrolytes are very important components of clean energy devices. Phosphoric acid (PA, H₃PO₄·H₂O) is one of the best proton conductors, but needs to be incorporated into some matrix for real device applications, such as into lyotropic liquid crystalline mesophases (LLCMs). Herein, we show that PA and nonionic surfactant (NS, C₁₂H₂₅(OCH₂CH₂)₁₀OH, C₁₂E₁₀) molecules self-assemble into PANS—LLCMs and display high proton conductivity. The content of the PANS—LLCM can be as high 75% H₃PO₄·H₂O and 25% 10-lauryl ether (C₁₂H₂₅(OCH₂CH₂)₁₀OH, C₁₂E₁₀), and the mesophase follows the usual LLC trend, bicontinuous cubic (V₁)—normal hexagonal (H₁)—micelle cubic (I₁), by increasing the PA concentration in the media. The PANS—LLCMs are stable under ambient conditions, as well as at high (up to 130 °C) and low (−100 °C) temperatures with a high proton conductivity, in the range of 10⁻² to 10⁻⁶ S/cm. The mesophase becomes a mesostructured solid with decent proton conductivity below −100 °C. The mesophase can be used in many applications as a proton-conducting media as well as a phosphate source for the synthesis of various metal phosphates. As an application, we demonstrate a graphene-based optical modulator using supercapacitor structure formed by graphene electrodes and a PANS electrolyte. A PANS—LLC electrolyte-based supercapacitor enables efficient optical modulation of graphene electrodes over a range of wavelengths, from 500 nm to 2 μm, under ambient conditions.

KEYWORDS: phosphoric acid · lyotropic liquid crystals · proton conductor · gel electrolyte · graphene optical modulator

Assembling nanostructures, clusters, ions, and molecules into mesostructures, using surfactants, is an important process for the development of advanced functional materials. Specifically, organizing ionic conductors, acids, and bases into mesophases is beneficial in the advancement of new materials and technologies. There exist many chemicals for the self-assembly of a surfactant into lyotropic liquid crystalline (LLC) mesophases, such as water,¹⁻³ oil—water,⁴ aqueous solutions,⁵⁻⁷ ionic liquids,⁸⁻⁹ CO₂,¹⁰ and salts.¹¹⁻¹³ Furthermore, the LLC media with hydrophilic and hydrophobic domains offers a confined space for the synthesis of chemicals with unique properties.¹⁴⁻¹⁸ Only a few examples exist in the literature that use various acids and that investigate the influence of these acids on the existing LLC mesophases.¹⁹⁻²² There are also some examples in which the acid is the surfactant such as amino acids or carboxylic acids, which are used in the presence of water as the solvent in the LLC phase.²³⁻²⁵ Moreover, organizing acids in the LLC phases can be very beneficial for the synthesis of porous materials that require an acidic media.

To broaden the concept of the LLC phases, we have investigated using some salts as the solvents in the assembly of surfactants.¹¹⁻¹³ These salts have either low melting points or very high solubilities and act as solvents in the assembly of salt—surfactant mesophases, where the driving force is the soft-confinement effect (SCE).²⁶⁻²⁷ The small space in the hydrophilic domain of the LLC mesophases reduces the melting point, enhances the solubility of the salts, and keeps these highly concentrated solutions stable in the hydrophilic domains of the mesophases.¹²,¹³ Usually the solvent/solution—surfactant
interactions are strong enough to keep the salt species in the LLC phase at even very low temperatures (below liquid N2). Pure phosphoric acid is a liquid at temperatures above 42 °C and is also liquid at RT in the presence of as little as one water per H3PO4 and can be used as a solvent to form PA—nonionic surfactant LLC mesophases (PANS—LLCMs).

The hygroscopicity of compounds determines whether or not a compound can act as a solvent to assemble nonionic surfactants. The measure is known as the deliquescent relative humidity (DRH): above the DRH, salts spontaneously absorb water from their surroundings and dissolve. If the compound has a very low DRH value, it is likely to act as a solvent in the presence of a small amount of water. P2O5 is one of the most deliquescent compounds, and it immediately dissolves in any humidity. Furthermore, PA is an extremely useful acid in many applications, ranging from food to bone health. It is also one of the best known proton conductors and can be used as a solvent to form PA—nonionic surfactant LLC mesophases (PANS—LLCMs).

RESULTS AND DISCUSSION

In this investigation, we have used concentrated PA with 10-lauryl ether (C12H25(OCH2CH2)10OH, C12E10) without any additional water to form the PANS—LLCM phases. The molar and mass ratio of PA to C12E10 can be as high as 20 and 75%, respectively. The mesophases are bicontinuous cubic (Vid) at low, hexagonal (Hid, 2D or 3D) at intermediate, and micelle cubic (hid) at high PA concentrations. The PANS—LLCMs have been employed to control optical transmission of a graphene electrode in a supercapacitor geometry. PANS-based electrolytes enable us to operate graphene optical modulators in the near-IR—visible spectra.

The PANS samples diffract at small angles over a wide range of PA/C12E10 ratios, characteristic of the LLC mesophases. Figure 1 shows a series of XRD patterns from various compositions. The mesophase is semi-stable up to 2.0 PA/C12E10 and bicontinuous cubic with a unit cell parameter a of 127.1 Å. There is a phase transformation around a 2–3 PA/C12E10 mole ratio to a 2D hexagonal phase with a unit cell parameter a of 55.4 Å. The (110) line of the 2D hexagonal phase is also weakly observed upon destroying the orientation. The phase transformation continues by increasing the PA amount; it is 3D hexagonal at mole ratios around 4 to 5 and becomes cubic above a mole ratio of 6 PA/C12E10, with unit cell parameters a of 54.7 and c of 89.3 Å and a of 116.7 Å, respectively; see Figure 1.

Figure 1. XRD patterns of the PANS—LLCMs with a PA/C12E10 mole ratio of (a) 2, (b) 3, (c) 5, and (d) 6. Insets are the plots of d-spacing versus hkl relation of the corresponding structure.

Figure 2. FT-IR spectra of the PANS—LLCMs with a PA/C12E10 mole ratio of (a) 3, 6, 11, and 19 and (b) FT-IR spectral changes of the PANS—LLCMs of a 4 PA/C12E10 mole ratio over time.

Note also that the hexagonal phases are birefringent and display a characteristic focal conic fan texture with a typical domain size of 10—100 μm; see inset in Figure 1b.

We also monitored the room-temperature behavior of the PANS—LLCMs using FT-IR spectroscopy; see Figures 2 and S1. Figure 2a displays a set of FT-IR spectra of PANS—LLCMs with an increasing PA/C12E10 mole ratio. Clearly, the peaks related to the concentrated PA dominate the spectra with an increasing PA/C12E10 mole ratio, which closely resembles the spectrum of concentrated PA. These spectra show that the PA molecules form a network of hydrogen bonds with each other, water, and surfactant molecules in the mesophase; see Scheme 1. The peaks at 491 and 1020 and the broad peak around 1200 cm−1 are due to PA bending and stretching modes, respectively. The OH region is spread over a broad frequency range, indicating a strong hydrogen-bonding network with different strengths. The broad peak at around 1685 cm−1 is due to the H2O bending mode; the likely origin of such broadening and blue shift is that the water species are involved in the network and are partially protonated. The peaks at 2370, 3000, and 3500 cm−1 are due to various stretching modes of PA with some contribution from the stretching mode of water in the PA—H2O...
network. The stability and the response of PANS–LLCMs to atmospheric conditions and to high humidity conditions (25–70%) were also monitored over time using the samples spread over a Si wafer and with FT-IR spectroscopy; see Figures 2b and S1. However, above 15 PA/C12E10 mole ratio the mesophase is not stable to high humidity. Figure 2b clearly shows that there is almost no change in the spectra over 2 weeks under ambient laboratory conditions and only very little change in the extremely high PA containing samples, Figure S1. We also monitored PA itself over the Si wafer and noticed that within a few minutes it adsorbs so much water such that the sample flows from the silicon surface. This indicates that PANS–LLCMs do not demonstrate much response to atmospheric changes and are stable over very long periods.

The PANS–LLCMs are also stable over a very broad temperature range (−100 to 130 °C), but they depend on the PA/C12E10 mole ratio; the PANS–LLCM of low PA/C12E10 mole ratio has been monitored under a POM using a temperature-controlled hot plate. Figure S2 shows a set of POM images recorded between RT and −150 °C. Clearly, the sample never loses its birefringence at any temperature. We also checked the isotropization or melting point of the mesophase, which increases with an increasing PA concentration up to 14 mole ratios and gradually decreases at higher concentrations, and becomes liquid at RT in the samples with 18–20 mole ratios. Moreover, these samples are gel-like at lower temperatures.

The ac proton conductivity measurements were carried out using two different homemade cells; see Figure S3 using impedance spectroscopy. See Supporting Information for details. Briefly, the conductivity increases with an increasing PA concentration, Figure 3a. The increase in the PA concentration induces conformational and phase changes and enables the hydrophilic domains to accommodate more PA molecules. Such a compact arrangement in the cubic phase likely decreases the intermolecular distance between neighboring PA molecules and, hence, makes the hopping of protons more feasible. In the Grotthuss pathway, the conduction is facilitated by the hopping migration of protons, which require the close proximity of the ions involved, while in the vehicle pathway, the transportation of the proton is not affected by the absence of an infinite chain of hydrogen-bonded molecules. It requires only “vehicles” such as water, with high self-diffusion constants, to facilitate the proton mobility. Therefore, if we assume the Grotthuss pathway, then the higher composition PANSs, which are more viscous, should definitely be more conductive. Recently, Dag et al. reported that from the lithium salt–surfactant mesophases the less viscous samples are more conductive because of the high ionic mobility associated with the more liquid-like gels. Therefore, while the LLCM–hydrated metal salts obeyed the “vehicle” mechanism of ion conductivity, the lower resistivity in the highly viscous PANS gels is based on the fact that they obey the Grotthuss conduction mechanism.

The proton conductivity has been measured for all samples, but the temperature-dependent measurements were carried out for only five samples: PA/C12E10 mole ratio of 5, 10, 16, 18, and 20, Figure 3b. The results showed that the gels remained conductive over a wide range of temperatures, depending on the compositions. The conductivity increases with an increasing temperature until a certain value (above the melting points), then starts decreasing again, showing either the complete evaporation of water from the samples or the collapse of the mesophase caused by melting. This behavior was observed in all compositions but at different temperatures, which led to the conclusion that the vaporization temperature was affected by the viscosity of the composition. For example, the more viscous, low-concentrated samples remain conductive even at temperatures above 130 °C, while the more fluid-like samples were resistive above 100 °C. In the low-temperature regions, it was also realized that the more concentrated (less viscous) samples showed reasonable conductivity (7.6 × 10⁻⁵ S/cm) even at temperatures below −80 °C while the lower concentrated samples show only fair conductivity at temperatures around −30 °C.

It is already known that the increase in temperature causes the breakage of hydrogen bonds and, at the
same time, enhances the conductivity of the materials. Therefore, the conductivity of the PANS gels, which we assume to be largely dependent on hydrogen-bonded networks, may be at odds with this already accepted belief. However, this could be explained when we consider the different conductivity mechanisms and the possibility of changing from one mechanism to the other. When the gels completely melt, the decrease in the activation energy coupled with the fast mobility of the “vehicles” will enhance the proton conductivity even more. Therefore, the vehicle’s mechanism, which requires higher activation energy, will be the preferred mode of conduction in such a situation. This is what led to the observed rise in conductivity and non-Arrhenius behavior as a function of temperature in the viscous gels. This behavior continues until the gel-like liquid starts losing water molecules to the atmosphere and causes a “V” shape in the higher temperature regions; see Figure 3b. On the other hand, when the temperature was reduced to temperatures much lower than 0 °C, the resistance sharply increased. One of the factors that might be responsible for this sharp change is the decrease in the mobility of the carriers at temperatures below the freezing point of the gel or the formation of cracks (upon freezing) in the samples. In either case, semicircular arcs, showing bulk impedance of the sample, were observed in the high-frequency regions of all the samples that were completely frozen. The temperature-dependent conductivity plots show a non-Arrhenius behavior. This is known for PA in various media. The conductivity is Arrhenius below the glass transition temperatures, and the conduction takes place via a hopping mechanism (Grotthuss mechanism). Two different proton conductivity mechanisms (namely “Grotthuss” and “vehicle” mechanisms) are responsible for the non-Arrhenius behavior at higher temperatures.

We also employed the PANS—LLCMs as electrolytes in a graphene-based optical modulator and compared the performance of PANS with a well-known ionic liquid (N,N-diethyl-N-methyl-N-(2-methoxyethyl)-ammonium bis(trifluoromethanesulfonyl)imide). The graphene optical modulator is a supercapacitor formed by two graphene electrodes and an electrolyte between them (Figure 4a). The applied voltage between the graphene electrodes polarizes the electrolytes, which yields electrostatic doping on the graphene layers. Due to the low density of the states, the Fermi energy of the graphene shifts significantly under the electrostatic doping and results in the modulation of the optical transmittance owing to the Pauli blocking (see the inset in Figure 4b). The key challenge for the graphene modulator is to activate large doping concentrations with low applied voltages. The PANS electrolyte provides a very efficient device operation. The modulation of the optical transmittance is shown in Figure 4b. Transmittance in the visible and near-IR can be controlled by 1.7% (limited by the absorption of single-layer graphene) with relatively low voltages (<3 V). We compared the extracted Fermi energy of three PANS electrolyte concentrations with the ionic liquid electrolytes (Figure 4c). PANS (PA/C12E10 = 10) provides the best results, with Fermi energies up to 1.5 eV, which yield an optical modulation up to 500 nm. Note that the Fermi energy (E_F) has been measured from the transmittance spectrum, which shows a step-like function with a cut-off wavelength at 2E_F. Another advantage of the PANS over the ionic liquid is the high ionic mobility. Figure 4d shows the frequency-dependent capacitance of the modulator at 0 V. PANS electrolytes yield relatively larger capacitance at kilohertz frequencies (see Figures S5 and S6 for details of the measurements). The device performance of the optical modulators has been compared using three different PANS. Figure 5a–c shows the optical transmittance through the modulator for the PA/C12E10 ratios of 3, 10, and 20. Figure 5d shows the variation of the optical transmittance at 800 nm for the electrolytes.

**CONCLUSION**

In summary, PA and 10-lauryl ether form stable LLC mesophases over a very broad concentration range. The mesophases follow a general phase transition of V1—H1—I1 with increasing the PA content.
The PANS–LLCMs have a high proton conductivity, which is based on the Grotthuss mechanism (proton-hopping) at low temperatures and is based on both Grotthuss and vehicle mechanisms at higher temperature, and display non-Arrhenius behavior with increasing temperature. As an application, we fabricated graphene supercapacitors using PANS–LLCM electrolytes, which operate as optical modulators in the visible and near-IR spectra. We observed that PANS provides better device performance over ionic liquids in terms of operation wavelength and frequencies owing to the large ionic concentration and high proton conductivity.

METHODS
Preparation of PANS–LLCMs. A mixture of 7.000 g of water, 1.000 g of 10-lauryl ether, and phosphoric acid (for 19 different solutions the PA weights are 0.368, 0.552, 0.736, 0.921, 1.105, 1.289, 1.473, 1.657, 1.841, 2.025, 2.209, 2.393, 2.577, 2.762, 2.946, 3.139, 3.314, 3.498, and 3.682 g, corresponding to PA/C12E10 mole ratios of 2 to 20 with an increment of 1, respectively) in a closed glass vial was stirred for about 5 h. The homogenized solutions were spin-coated at a reasonable speed (500–700 rpm) on the surfaces of substrates (either glasses or silicon wafers). The gel phases (for conductivity, POM, and isotope-labeling experiments) were prepared by adding 1.000 g of 10-lauryl ether to the required quantities of PA (exact values listed above). The mixtures were made airtight through the use of Teflon tape. All the vials containing different PA/C12E10 ratios were put in a shaking water bath. The temperature of the bath was increased above the melting point of the surfactant (40–50 °C). This process took almost 24 h and resulted in the formation of highly viscous gels.

Preparation of Supercapacitors. Graphene layers used in the capacitors were synthesized by chemical vapor deposition on 25 μm thick copper foils (Alfa Aesar Cu foil (item #13382)) at 1050 °C under 10 Torr chamber pressure. Partial pressures of CH4 and H2 gases were set as 3 and 7 Torr, and corresponding rate of flows were 40 and 80 sccm, respectively. We transferred 1 × 2 cm2 graphene layers on glass microscope slides by using an s1813 photore sist as a mechanical support for graphene. After drying the photoresist at 70 °C for 12 h in the oven, the Cu foils were completely etched in the dense FeCl3 solution in 2 h. Graphene holding the photoresist was refloved at 110 °C to release the graphene layer on the microscope slide. The graphene-transferred microscope slides were put together by a 100 μm thick spacer. The gap between the two graphene electrodes was filled with a gel electrolyte or a commercially available liquid electrolyte to compare the electrolyte performances.

Instrumentation. The XRD patterns were recorded using a Rigaku Miniflex diffractometer equipped with a high-power Cu Kα source operating at 30 kV/15 mA and a wavelength of 1.5405 Å. The measurements for both the films and gel samples were performed over a broad range (450 to 1100 nm) by using a Bruker Vertex 70 V FT-IR spectrometer equipped with a Si photodiode. The bias voltage was applied by a source measure unit (Keithley 2400), and the leakage current was also monitored during the transmission measurements to observe the proper device operation. An HP 4194A precision LCR meter, which is programmed to measure the parameters of a serially connected capacitor–resistor network model, was used in capacitance and resistance measurements of the graphene supercapacitors.

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

Supporting Information Available: Composition- and time-dependent FT-IR spectra, temperature-dependent POM images, setups for conductivity measurements, Nyquist plots, and extra optical and electrical characteristics of the graphene supercapacitors. This material is available free of charge via the Internet at http://pubs.acs.org.

Acknowledgment. Authors thank TUBITAK under the project number 113Z730 for financial support. E.T. thanks IDB for the M.Sc. scholarship. O.D. is a member of Science Academy, Istanbul, Turkey.

REFERENCES AND NOTES
4. Alexandridis, P.; Olsson, U.; Lindman, B. A Record Nine Different Phases (Four Cubic, Two Hexagonal, and


