Novel Luminescent Ionic Adducts Based on Pyrene-1-sulfonate

Maria Angela Castriciano, Paola Cardiano, Enza Fazio, Placido Giuseppe Mineo, Angelo Nicosia, Roberto Zagami, Mariachiara Trapani, Luigi Monsù Scolaro, and Sandra Lo Schiavo

1. INTRODUCTION

Ionic liquids (ILs), defined as salts melting below 100 °C, have attracted extensive attention in the last 2 decades because of their custom-tailored physicochemical properties, such as low vapor pressure, wide liquid range, excellent chemical, thermal, and electrochemical stability, dispersant capabilities, dielectric constants, extraction capabilities, and so on.1–11 As such, ILs are highly versatile media replacing conventional organic solvents and electrolytes, finding applications in many strategic fields such as catalysis, electrochemical devices, separation processes, synthesis, nanochemistry, photonics, and, more recently, biotechnology.4–10 Anchoring of IL units to proper substrates such as polymer networks and/or nanomaterials is a successful synthetic methodology to access IL-based materials, usually referred to as supported ILs (SILs), with higher processing performance and even new functions.11–17 Among others, by promoting ionic self-assembly processes, ILs and SILs have shown to be ideal precursors leading to advanced ionic supramolecular assemblies.18–21

ILs with intrinsic functions, usually defined as task-specific ILs, are easily obtained by introducing active ions in IL structures.22–24 Thanks to the high disposal of organic emitter ions, this approach is a useful means to access novel soft phospholuminescent materials, alternative to rare-earth fluorophores, for application in photonics, sensing, optoelectronic devices, dye-sensitized solar cells, and so on.12,25–29 By using this strategy, ILs with improved physico-chemical properties and processing capabilities, such as solubility, photostability,
high quantum yields, and so on, differing significantly from those of the related salts, have been obtained.\textsuperscript{30--35} Further, systematic studies also showed that the photoactivity of ILs constitutes, to some extent, a tunable, structurally controlled IL function.

Because of the high relevance of the so-called “POSS nanotechnology” (POSS = polyhedral oligomeric silsesquioxanes) in the context of materials science, we have devoted our attention to the development of POSS-based ILs. Briefly, POSS are cage-like hybrid units with empirical formula \((\text{SiO}_{1.5})_{x}\), consisting of an inner inorganic core tethered with organic substituents widely used as efficient nanofillers inducing improved thermal, mechanical, and oxidation stability. Interestingly, when POSS is used as a scaffold for covalent binding of photoluminescent moieties, it experiences, among others, a dispersing action, which results in a significant improvement of quantum efficiency.\textsuperscript{38--42} In the course of our studies on POSS tetraalkylammonium-based ILs,\textsuperscript{38--40} we have reported on the synthesis and photochemical characterization of 1:4 POSS@TPPS ILs based on \textit{meso}-tetrakis-(4-sulfonatophenyl)porphyrin tetraanion (TPPS) and the POSS monofunctionalized cation, trimethylpropylammonium hepta-(isoctyl)octasilsesquioxane.\textsuperscript{41} The synthesis, based on a simple biphasic chloroform/water process, as well as the resulting optical properties, is highly dependent on the amphiphilic nature of the bulky cation. The latter acts as a highly dispersant agent in chlorinated solvents, reducing the well-known aggregation tendency of the TPPS tetra-anion, both in solution and in the solid state. Therefore, long emission lifetimes were observed for both the TPPS anions and TPPS J-aggregates. Analogous to porphyrins, pyrene derivatives constitute a class of attractive and thoroughly investigated solvatochromic fluorophores widely applied in optoelectronics, biochemistry, and sensor fields. These compounds are characterized by long excited-state lifetime and, in concentrated solution and in the solid state, lead to characteristic excimers because of self-association phenomena.\textsuperscript{42--53} Furthermore, pyrene acts as a probe for the local environment polarity because the intensity ratio of the first to third vibronic emission bands (I/III ratio) is very sensitive to the solvent polarity.\textsuperscript{54} On the basis of these considerations, pyrene sulfonate anions were chosen for the development of novel photoluminescent ILs.

In the present study, we report on the synthesis and characterization of a series of ion pairs based on monofunctionalized pyrene-1-sulfonate (PS). Because of their high versatility and applications, besides trimethylpropylammonium hepta-(isoctyl)octasilsesquioxane, conventional imidazolium cations, featured by “moderately” long alkyl chains, have been also considered as PS counterions (Scheme 1). Vinyl-imidazolium-based ILs are commonly used and they find a plethora of applications in various fields. It is well known that their properties can be opportunely tuned by varying the alkyl chain length. In addition, we chose to use vinyl-imidazolium derivatives to achieve potentially cross-linkable systems, thus tuning the IL properties. Furthermore, it has been reported that the presence of a vinyl substituent on the imidazolium ring provides additional hydrogen bonding interactions between the cations and anions whose strength decreases with increasing anion size.\textsuperscript{21} The new 1:1 ionic adducts were characterized by matrix-assisted laser desorption/ionization-time of flight (MALDI-TOF) mass spectrometry, thermogravimetric analysis (TGA), differential scanning calorimetry (DSC), and a series of spectroscopic techniques (NMR, UV–vis, steady-state, and time-resolved fluorescence spectroscopy). We anticipate that while the POSS–pyrene adduct is classifiable as an IL, the imidazolium ones, featuring melting points above 100 °C, have to be considered as simple ion pairs.

The combination of spectroscopic and light scattering data support the monomeric nature of the POSS–PS adducts in solution. On the contrary, imidazolium derivatives display a general attitude to self-organize in supramolecular structures through cation interactions, with the polycyclic anion acting as a nucleation center.

2. RESULTS AND DISCUSSION

2.1. Synthesis and Characterization. As described in the Experimental Section, PS POSS–ammonium and imidazolium derivatives (Scheme 1) were synthesized in high yields according to simple metathesis reactions. All new adducts have been obtained as pale yellow solids, except for POSS–PS, which appears very sticky, consistent with the nature of the cation. The characterization of the new ionic adducts was carried out by elemental analyses, conventional spectroscopic techniques (UV–vis, static and time-resolved fluorescence emission, and NMR spectroscopy), and MALDI–TOF spectrometry. Karl Fischer titrations reveal no significant level of water. Nevertheless, all samples were dried under vacuum overnight at 50 °C.

The \(^1\)H NMR spectra of the new ionic adducts display a signal pattern consistent with the presence of an asymmetric pyrene group between \(\delta 8.0–9.5\) ppm, which well integrates 1:1 with those of the cation counterpart. The presence of the PS polycyclic anion does not produce other significant changes with respect to those of the parent IL halides: the only exception concerns the up-field shifts observed for the NC(H) N imidazolium protons that, as it is well known, are very sensitive to the nature of the anion. MALDI-TOF mass spectra of all new ion pairs (see Supporting Information) show the presence of a single signal corresponding to the molecular mass peak of the ionic species.

The thermal behavior of the new ionic PS adducts has been investigated by DSC and TGA analyses (Table 1). As reported
in Table 1 and Figure 1, the occurrence of only a reversible glass transition at −4.2 °C and no melting point under 100 °C (DSC data) indicate that POSS–PS may be defined as an IL. Accordingly, imidazolium–PS must be considered as simple ion pair couples.

With respect to POSS–PS, imidazolium derivatives afford more complex and diversified DSC responses. Figure 1 (upper) reports the heating course for all imidazolium samples, collected at 10 °C min⁻¹. The curve of 1-vinyl-3-hexylimidazolium (VEIm)–PS features a broad and weak endothermic event at 22.2 °C, followed by two distinct exothermic peaks at 42.7 and 79.9 °C, pointing to the occurrence of amorphous-to-crystal and solid-to-solid transitions, respectively, together with a broad endothermic event at 124.5 °C (T onset). Actually, the shape of this last peak is not sharp, suggesting overlapped melting events. DSC thermograms of VDIm–PS present two very narrow endothermic events (120.9 and 135.7 °C), very likely associated with the melting of two distinct crystalline species with different arrangements or packing.

Finally, the DSC scan of 1-methyl-3-decylimidazolium (MDIm)–PS, bearing a methyl group in place of vinyl on the imidazolium ring, features a weak exothermic process (T onset = 113.7 °C), immediately followed by a melting event at 132 °C. The observed heating transition phase profiles are not reversible. In the corresponding cooling curves (Figure 1, lower), only a liquid-to-solid process is observed for all the samples. This exothermic process is well defined for both MDIm– and VDIm–PS (T m = 114.0 and 104.8 °C, respectively), while for VEIm–PS, it shows unresolved multipeaks with T onset at 98 °C. As expected, by lowering the DSC heating/cooling scan rate to 2 °C min⁻¹ (see Supporting Information), different DSC thermograms have been obtained. In the heating scan, the melting transition for MDIm–PS is featured by two narrow peaks at 132.1 and 135.4 °C, very likely arising from two distinct crystal-to-liquid phase transitions. The melting process for VDIm–PS is detected as a broad signal with a T onset of 133.43 °C and T max of 138.0 °C, which is preceded by a solid-to-crystal transition. By cooling, an unresolved double exothermal peak is observed for MDIm–PS at 114.0 °C, while for VDIm–PS, a crystallization peak appears at 104.8 °C. Conversely, VEIm–PS affords a heating curve displaying an endothermal peak at 125.7 °C and a cooling one featuring a broad exo-multipeak with T onset at 98.0 and T max at 88.5 °C. The thermal behavior of imidazolium ILs/salts has been widely explored, and the role played by the nature of cations and/or anions, alkyl chain lengths on the aromatic ring, and intermolecular forces and symmetry in determining glass, melting, and meso- and polymorphic-phase transitions has been demonstrated. On the basis of our experimental findings, it is reasonable to suppose that as solids, Imi–PS can be organized in supramolecular structures, via the alkyl chain, which can undergo “order–disorder” or vice versa phase transitions, thus affording nonreversible heating/cooling cycles and different DSC scanning profiles at different heating rates.

On the other hand, recent reports have described the role played by pyrene moieties in promoting and controlling the polymorphic superstructures of asymmetric dendritic biphenyl-based supramolecules. Our DSC data suggest that (i) the thermal transition phase behavior of POSS–PS is still determined by the bulky amphiphilic nature of the trimethylpropylammonium hepta(isocyt) octasilesquioxane cation and (ii) the substitution of bromide with a PS anion produces significant modifications on imidazolium-ILs, in terms of the melting point increase and the overall DSC thermogram profiles. Although the DSC technique is currently used for the determination of thermodynamic parameters of ILs/salts, it is not able to give satisfactory information on molecular and supramolecular polymorphic
structures. Further investigation of the thermal behavior for these imidazolium–PS ionic adducts goes beyond the aim of the present study. The thermal stability of the new adducts was determined by TG analyses (Figure 2, Table 1) in the temperature range between 50 and 800 °C, under a nitrogen flow (60 mL min⁻¹) and heating rate of 10 °C min⁻¹. The TGA profile of POSS–PS proceeds, essentially, in a large temperature range for which two temperatures of maximum volatilization (338.8 and 465.2 °C) are observed. The imidazolium–PS adducts afford very similar thermograms, as the mass losses take place in a single step. As their thermal stability is concerned, TGA data suggest that the presence of the vinyl substituent on the imidazolium ring induces a lower stability. Accordingly, VDIm–PS displays T_onset and T_max values significantly lower than those observed for MDIm–PS.

2.2. Photophysical Behavior in Solution. All the samples displayed a similar behavior in terms of good solubility in solvents with different polarities [tetrahydrofuran (THF), CHCl₃, and MeOH] up to millimolar concentration, where they are stable for months, as confirmed by the invariance of the UV–vis and fluorescence emission spectra. The photophysical properties and aggregation behavior of these adducts have been investigated in chloroform and methanol solution by various spectroscopic and light scattering techniques. Pyrene-1-sulfonate–tetra-n-butyl ammonium salt (TBA–PS) has been used as a reference. The UV–vis and fluorescence emission spectra in chloroform and methanol are similar for all the investigated samples. As an example, Figure 3 reports the spectroscopic features of MDIm–PS. Absorption bands with a characteristic vibronic fine structure, between 300 and 350 nm (upper graph), together with a structured emission band centered at ca. 378 nm (lower graph) are typical of the pyrene chromophore. The observed ipsochromic shift of the absorption bands in methanol with respect to chloroform (Δλ = −3 nm), together with intensity ratio modulation of the first to third vibronic emission bands, is ascribable to the change in solvent polarity. Beer’s law is obeyed up to millimolar concentrations and no additional nonstructured band at longer wavelengths, characteristic of excimer emission, has been observed for all the species. To confirm the absence of the excimer in solution, emission spectra at different sample concentrations and excitation spectra at different emission wavelengths have been collected (see Supporting Information). In any case, no evidence for the presence of excimers has been obtained. The fluorescence emission decays, in aerated solutions, show a biexponential profile with comparable long-lifetime values from 4.6 to 4.8 ns and a short one from 1.4 for MDIm to 2.1 ns ca. for POSS. As an example, Figure 4 shows a typical decay trace, while lifetimes and relative amplitude values are shown in Table 2.

The two emission lifetime values are consistently shorter than those reported for PS sodium salt (PS–Na) in aqueous solution but very similar to those obtained for TBA–PS in apolar solvent. This experimental evidence underlines that the presence of oxygen could be responsible for this discrepancy. The biexponential decay curves observed for these PS adducts are clearly indicative of two different species in solution.
Concentrations in solution of a chloroform and methanol solutions. DLS data collected in dynamic radii monodisperse population of spheric aggregates with hydrodynamic derivatization in CHCl₃ and MeOH at different concentrations of PS–imidazolium derivatives in CHCl₃.

| sample           | $r_1$/ns  | $r_2$/ns  | $A_1$ | $A_2$ | $r(t)$/ns |
|------------------|-----------|-----------|-------|-------|-----------|
| PS–Na            | 3.2 ± 0.02| 54.2 ± 0.3| 10    | 90    | 1.04 ± 0.03|
| TBA–PS           | 1.7 ± 0.04| 4.6 ± 0.02| 40    | 60    | 0.97 ± 0.05|
| MDIm–PS          | 1.4 ± 0.05| 4.7 ± 0.02| 32    | 68    | 1.02 ± 0.05|
| VDIm–PS          | 1.6 ± 0.05| 4.6 ± 0.02| 35    | 65    | 1.01 ± 0.08|
| VEIm–PS          | 1.8 ± 0.05| 4.7 ± 0.02| 32    | 68    | 1.05 ± 0.05|
| POSS–PS          | 2.1 ± 0.03| 4.8 ± 0.01| 43    | 57    |           |
| GF/VDIm–PS       | 1.6 ± 0.03| 3.7 ± 0.04| 14    | 86    |           |

*10 µM, $T = 298 K$, $\lambda_{ex} = 390$ nm, $\lambda_{em} = 450$ nm. *H₂O.

The fluorescence anisotropy decay [$r(t)$], which allows us to gain information on the formation of supramolecular structures via the rotational diffusion of the fluorophores, gives a rotational correlation time of about 1 ns, which is not fully compatible with the size of the single molecule (see data in Table 2). Further, the rotational correlation time suggests that the free rotation of each single fluorophore is partially inhibited.

In order to investigate the aggregation behavior in solution, dynamic light scattering (DLS) experiments were performed at different concentrations of PS–imidazolium derivatives in chloroform and methanol solutions. DLS data collected in Table 3 are in line with the presence in solution of a monodisperse population of spheric aggregates with hydrodynamic radii $R_H$ depending on the solvent polarity and adduct concentration. In chloroform, only some aggregate structures are detectable at low concentrations (1 µM), while larger ones are well identifiable at highest concentrations (100 µM). In the more polar methanol, nanometer-sized aggregates are already detectable at very low concentrations (1 µM). This behavior could be ascribed to assembling phenomena involving imidazolium cations. Comparative DLS experiments, performed on MDIm, VDIm, and VEIm bromide precursors, indicate that these compounds have no tendency to self-aggregate, independent of the alkyl chain length on the imidazolium ring. The absence of resonant light scattering peaks in the PS absorption region (see Supporting Information) allows us to exclude the presence of extended PS aggregates resulting from the strong electronic coupling of adjacent chromophores. However, it does not exclude the formation of dimer or small oligomers, which could be responsible for the observed spectroscopic behavior. Hence, on the basis of these DLS and photophysical investigations, we can speculate on the formation of supramolecular species obtained by the interaction between PS anion as monomers, dimers, or small oligomers embedded in nanostructured aggregates, mainly because of countercation supramolecular self-assembly with a consequent consistent reduction of the fluorophore mobility. We can conclude that extended stacking is not observed, presumably owing to the great “edge” effect due to the alkyl chains of countercations.

2.3. Interaction of PS Adducts with Graphite Flakes.

Polycyclic aromatic compounds, based on pyrene, perylene, and so forth, in a simple ionic or amphiphilic properly functionalized form constitute useful species for a supramolecular approach in graphite (GR) exfoliation to graphene as well as for noncovalent modification of graphitic surfaces. Among others, various substituted pyrene salts have been successfully employed in shear-assisted exfoliation of natural GR as well as of GR flakes (GFs). On the other hand, thanks to their ionic nature and their tunable surface...
tension, ILs are an ideal “solvent” for GR exfoliation and stabilization of graphene suspensions.

Stable GFs/PS-adduct THF suspensions have been obtained by using a sonication-assisted liquid phase procedure as described in the Experimental Section. These suspensions have been analyzed by UV−vis and Raman spectroscopy, the latter being one of the most powerful tools to investigate graphitic surfaces and edge modifications. All the GFs@PS-adduct composites afford comparable electronic and Raman spectra; hence, the discussion will be limited to the GFs@VDIM−PS sample.

According to Figure 5a, the Raman spectrum of GFs is consistent with a graphitic system bearing no significant structural defects, featuring a disordered, related low-intensity D peak at 1352 cm$^{-1}$, a sharp G band at 1578 cm$^{-1}$, ascribed to $E_g$ vibrational mode, and the 2D one centered at around 2718 cm$^{-1}$ and characterized by two subbands, namely, 2D$_1$ and 2D$_2$.

In the GFs/VDIM−PS Raman spectrum (Figure 5b), the GF pre-existing bands show a different intensity ratio, while new ones appear. In particular, the D band intensity increases, the G band is slightly split at a higher wavenumber, and the contribution D′ is now detectable at 1621 cm$^{-1}$. At the same time, the 2D pattern is broader and shifted to lower energy with the 2D$_1$ component (at around 2690 cm$^{-1}$) more pronounced. The intense band in the range 900−1050 cm$^{-1}$ is attributable to the PS moiety.

The UV−vis absorption spectra of VDIm−PS and GFs@VDIM−PS in THF solution are reported in Figure 6. By comparison, the modification induced by the presence of GFs on the VDIm−PS pattern is clearly evident. Even the fluorescence emission properties are modified (Table 3). The emission decay of GFs@VDIM−PS shows a biexponential profile with lifetimes of 3.7 and 1.6 ns and with relative amplitudes of 86 and 14%, respectively. The longest lifetime is shorter with respect to that of the VDIm−PS precursor (4.6 ns), while the relative amplitude is largely increased.

Even if the supramolecular interactions between the PS anion and GFs occur, they are not sufficient to afford an efficient exfoliation to graphene. For the sake of comparison, we have also explored the capabilities of halide IL precursors in GF processing. As an example, imidazolium bromide in THF solutions produces significant exfoliation of GFs, see Raman and scanning transmission electron microscopy (STEM) analyses in Supporting Information]. This finding suggests that the PS poly cyclic system is responsible for noncovalent functionalization of the GF surface and that the nature of cations does not play any synergetic role in the GR exfoliation process.

3. CONCLUSIONS

As part of our general interest regarding the design and synthesis of functional and SILs, here, we have reported on the PS anion attitude in generating fluorescent ILs. Amphiphilic trimethylpropylammonium hepta(isooctyl)-octasilsesquioxane and conventional imidazolium have been chosen as counterions. The experimental evidence shows that among the series of synthesized PS ion pairs, only the POSS derivative exhibits an IL behavior. Different from IL halide precursors, all imidazolium−PS derivatives display a crystalline nature. The PS anion, which does not undergo self-assembly processes, seems to play a main role in promoting the formation of imidazolium polymorphic structures, which are responsible for their complex transition phases. DLS investigations confirm the presence in solution of nanometer-sized species whose dimension is dependent on the solvent polarity and sample concentration; meanwhile, spectroscopic data indicate the absence of the extended electronically coupled chromophores in these supramolecular aggregates. These findings suggest, once more, that the presence of bulky surfactant cations is demanding in the design of luminescent-based PS ILs and more in general of ILs involving highly delocalized anions. Finally, we have also explored the capabilities of the new compounds in GF processing. On the basis of Raman data, we suggest that simple supramolecular interactions are operating between the PS anion and GF surface. These noncovalent forces are responsible for the stabilization observed in THF dispersion.

4. EXPERIMENTAL SECTION

4.1. Materials and Methods. All the chemicals and solvents employed, unless otherwise stated, were purchased from Sigma-Aldrich; aminopropylisobutyl POSS and aminopropyloisoctyl POSS were purchased from Hybrid Plastics; all these compounds were used as received. 1-Methyl-3- hexylimidazolium bromide, VEIm bromide, MDIm bromide, and 1-vinyl-3-decylimidazolium bromide were obtained according to a published procedure. Trimethylpropylammonium hepta(isobutyl) octasil sesquioxane iodide and trimethylpropylammonium hepta(isooctyl) octasil sesquioxane iodide were synthesized following a protocol already reported.

4.2. Synthesis. 4.2.1. Trimethylpropylammonium hepta(isooctyl)octasilsesquioxane Pyrene-sulfonate (POSS−PS). A suspension of 1-pyrenesulfonic acid sodium salt (0.1 g, 0.33 mmol) in 20 mL of deionized water was added dropwise to 0.430 g of trimethylpropylammoniumhepta(isooctyl) octasil sesquioxane iodide (0.30 mmol) dispersed in 10 mL of deionized water. The reaction mixture was left at 75 °C under vigorous stirring for 24 h. After decantation, the product separated as a sticky yellow solid; it was washed three times with water and then dried in a Rotavapor at 80 °C. Yield: 62%. 1H NMR (CDCl$_3$, δ, ppm): 0.66 (br, 16H), 0.87 (m, 65H), 0.99 (d, 21H), 1.18 (br, 9H), 3.31 (m, 2H), 3.38 (s, 9H), 8.21−8.00 (m, 7H), 8.73 (d, 1H), 9.35 (d, 1H). 29Si[1H] NMR (CDCl$_3$, δ, ppm): −67.34, −68.05, −69.04, −70.78. Elemental analysis: Calcd for C$_{78}$H$_{143}$NO$_{15}$SSi$_8$: C, 86.86%; H, 9.06%; N, 0.88%. Found: C, 86.87%; H, 9.09%; N,
0.91%. MALDI-TOF (m/z): 1591.3 (detected at m/z 2900.5 as [M]^2POSS adduct).

4.2.2. 1-Vinyl-3-hexylimidazolium PS. VEIm−PS was obtained as a pale yellow solid following a synthetic procedure similar to the previous one, using 0.19 g PS was isolated as a waxy pale yellow solid following a protocol as for VEIm−PS, and VEIm−PS; DSC curves of MDIm−PS, VDIm−PS, and Raman spectrum of CHCl3: (λem/nm) 348 (ε = 3.89 × 10^4 M⁻¹ cm⁻¹), 258, 268, 278, 317; 331. Fluorescence emission (CHCl3): (λem/nm) 340, (λem/nm) 378, 397,417. MALDI-TOF (m/z): 460.8 (detected at m/z 639.2 as [M]^2POSS adduct).

4.2.3. 1-Methyl-3-decylimidazolium PS. MDIm−PS was obtained as a waxy pale yellow solid following a synthetic procedure similar to that for MEIm−PS, using 0.32 g of 1-pyrenesulfonic acid sodium salt and 0.78 g of VEIm bromide (0.68 mmol) in deionized water. Yield: 76%. 1H NMR (CDCl3, δ, ppm): 0.86 (m, 3H), 1.16 (m, 14H), 1.65 (m, 2H), 3.98 (s, 3H), 4.03 (m, 2H), 7.01 (m, 1H), 7.11 (m, 1H), 8.21−7.97 (m, 7H), 8.75 (d, 1H), 9.34 (d, 1H), 9.94 (s, 1H). Elemental analysis: Calculated for C27H28N2O3S: C, 70.41%; H, 6.13%; N, 6.08%. Found: C, 70.56%; H, 6.12%; N, 6.06%. UV−vis (CHCl3): (λmax/ min) 348 (ε = 3.89 × 10^4 M⁻¹ cm⁻¹), 258, 268, 278, 317, 331. Fluorescence emission (CHCl3): (λem/nm) 340, (λem/nm) 378, 397,417. MALDI-TOF (m/z): 505.0 (detected at m/z 727.4 as [M]^2POSS adduct).

4.2.4. 1-Vinyl-3-decylimidazolium PS (VDIm−PS). VDIm−PS was isolated as a waxy pale yellow solid following a synthetic procedure similar to the previous one, using 0.19 g of 1-pyrenesulfonic acid sodium salt and 0.78 g of VEIm bromide (0.68 mmol) in deionized water. Yield: 76%. 1H NMR (CDCl3, δ, ppm): 0.86 (m, 3H), 1.16 (m, 14H), 1.65 (m, 2H), 3.98 (s, 3H), 4.03 (m, 2H), 7.01 (m, 1H), 7.11 (m, 1H), 8.21−7.97 (m, 7H), 8.75 (d, 1H), 9.34 (d, 1H), 9.94 (s, 1H). Elemental analysis: Calculated for C27H28N2O3S: C, 70.41%; H, 6.13%; N, 6.08%. Found: C, 70.56%; H, 6.12%; N, 6.06%. UV−vis (CHCl3): (λmax/ min) 348 (ε = 3.89 × 10^4 M⁻¹ cm⁻¹), 258, 268, 278, 317; 331. Fluorescence emission (CHCl3): (λem/nm) 340, (λem/nm) 378, 397,417. MALDI-TOF (m/z): 505.0 (detected at m/z 727.4 as [M]^2POSS adduct).

4.2.5. Exfoliation/Functionalization of GFs. In a general procedure, GFs (15 mg) and PS-based ionic adducts (30 mg) were sonicated for 24 h at room temperature then left to stand for 12 h, and finally centrifuged at 1300 rpm for 30 min to precipitate excess GF powder. The obtained light gray supernatant suspension has been used for analysis. By following a similar procedure, a specimen based on GFs and VDIm−Br was prepared for comparison.

4.3. Instruments. 1H and 29Si{1H} NMR solution spectra were obtained in deuterated solvents at room temperature on a Bruker AMX R-300 spectrometer operating at 300.13 and 59.62 MHz for 1H and 29Si nuclei, respectively. Chemical shifts (δ, ppm) were referred to SiMe4 TGA were performed by means of PerkinElmer Pyris TGA7 in the temperature range between 50 and 800 °C, under a nitrogen flow (60 mL min⁻¹) and heating rate of 10 °C min⁻¹. DSC measurements were carried out by using a TA Q20 instrument equipped with a refrigerant cooling system with a heating rate of 2 and 10 °C min⁻¹, in the −80 to +140 °C temperature range and under an anhydrous N2 atmosphere (60 mL min⁻¹). MALDI-TOF mass spectra were acquired using Voyager DE (PerSeptive Biosystems) using an acquisition protocol reported elsewhere.70−72 trans-2-[3-(4-tert-butylphenyl)-2-methyl-2-propenylidene]-malononitrile (DCTB) was used as the matrix. MALDI-TOF mass spectrometer calibration and average molecular mass determination were performed as reported in previous cases.73,74 Because of the isotopic composition, molecular species are detected in the mass spectra as clusters of peaks; to simplify their assignments, m/z values reported in the text are referred to the first peak of each cluster, corresponding to the ion containing the most abundant isotope of each element present. Micro-Raman spectra have been excited by the 532 nm diode laser wavelength, analyzed by a monochromator equipped with a 1800 line/mm grating, and collected by a Peltier-cooled charge-coupled device (CCD) sensor. STEM observations were carried out with a Zeiss-Gemini 2 electron microscope, operating at an accelerating voltage of 150 kV and equipped with a MultiScan CCD camera. STEM samples were prepared by depositing one drop of the solution on a 400 nm mesh carbon support sputter-coated with chromium and left to dry at room temperature for 3 h. UV−vis spectra were obtained using a Hewlett-Packard mod. 8453 diode array spectrophotometer. Static and time-resolved fluorescence emission measurements were performed on a Jobin Yvon-Spex FluoroMax 4 spectrophluorimeter using the time-correlated single-photon counting technique. A NanoLED (λ = 390 nm) has been used as the excitation source. Emission spectra were not corrected for the absorbance of the samples. Resonance light scattering (RLS) experiments were performed on a Jasco mod. FP-750 spectrophluorimeter, adopting a synchronous scan protocol with a right angle geometry.75 RLS spectra were not corrected for the inner-filter effect of the samples. Hydrodynamic particle sizes and size distributions were measured by DLS and carried out at 25 °C by a Zetasizer Nano-ZS (Malvern Instruments) equipped with a 633 nm He−Ne laser using backscattering detection. Each DLS sample was measured several times, and the results were averaged. IL stock solutions, obtained by dissolving the adducts in CHCl3 and/or MeOH, were stored in the dark and used within a week from preparation. The range of concentration (1 × 10⁻⁴ to 5 × 10⁻⁴ M) used in our experiments was determined spectrophotometrically by using the molar extinction coefficient at the Soret maximum.

**ASSOCIATED CONTENT**

5 Supporting Information

The Supporting Information is available free of charge on the ACS Publications website at DOI: 10.1021/acsomega.8b02961.

Positive MALDI spectra of POSS−PS, VDI−PS, VEIm−PS, and MDIm−PS; DSC curves of MDIm−PS, VDIm−PS, and VEIm−PS; fluorescence emission, extinction, and RLS spectra of VDIm−PS; and Raman spectrum and STEM image of GFs@VDIm−Br (PDF)
AUTHOR INFORMATION

Corresponding Authors
*E-mail: castriciano@pa.ismn.cnr.it. Phone: +39 090 3974108 (M.A.C.).
*E-mail: sloschia@unime.it (S.L.S.).

ORCID
Maria Angela Castriciano: 0000-0002-1514-8820
Paola Cardiano: 0000-0002-1380-5380
Placido Giuseppe Mineo: 0000-0003-3382-9272
Luigi Monsù Scolaro: 0000-0002-9742-9190

Author Contributions
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