Passing Current through Electrically Conducting Lyotropic Liquid Crystals and Micelles Assembled from Hybrid Surfactants with $\pi$-Conjugated Tail and Polyoxometalate Head

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Supporting Information

ABSTRACT: The solvent-mediated ability for molecularly encoded self-assembly into states of higher order (micelles, lyotropic liquid crystals) embodies the basis for many applications of surfactants in science and society. Surfactants are used frequently in recipes for nanoparticle synthesis. Because ordinary surfactants comprise insulating constituents (alkyl groups as side-chains and charged organic heads), such nanostructures are wrapped in an electrically inactive barrier, and this is a large disadvantage for future developments in nanotechnology. Implications of micelles with electrically conducting walls made from either “metallic” or “semiconducting” surfactants are huge, also in other areas such as nanoelectrocatalysis or micellar energy storage. We cross this frontier by replacing not only the hydrophilic chain but also the hydrophilic head by electronically conducting entities. We report the synthesis of surfactants with oligo para-phenylene-ethynylene as a $\pi$-conjugated side-chain attached to a redox-active, inorganic polyoxometalate cluster as charged head. It is proven that electronic communication between head and tail takes place. Hybridization on the molecular level leads to the emergence of advanced surfactant features such as semiconductor properties ($E_{\text{gap}} = 2.6$ eV) in soft lyotropic systems (micelles, liquid crystals).

KEYWORDS: surfactants, liquid crystals, polyoxometalates, organic–inorganic hybrids, molecular conductors

Molecular architectonics, the assembly of molecular entities into functional patterns coupling their properties to the nano-, then to the meso-, and finally to the macroscopic world, is key for implementing future technologies. Eventually, the spontaneous formation of such organized structures takes place as an intrinsic property of a system containing discrete constituents, a process termed self-assembly. A special class of self-assembling materials are liquid crystals (LCs), because they unify structural order and a high degree of mobility. This leads to a plethora of properties as also stated in the seminal article published by Tscherke in 2013. Liquid crystals are formed by certain molecular compounds, and one can distinguish between thermotropic and lyotropic LCs.

Rising interest was devoted to the self-assembly of $\pi$-conjugated systems because of their large potential in (supra)molecular electronics. A prominent application is for instance in field-effect transistors. Several papers have also reported on the semiconducting features of LC phases formed by these molecules. This year (2016) Kim and co-workers have summarized the potential of LCs based on graphene oxide. Most of the described systems belong to thermotropic phases. A molecular system with lyotropic (solvent-triggered self-assembly) and amphiphilic properties in combination with electronic conductivity could feature very interesting properties, for example, micelles as nanoreactors with electrically active and transmittive shells. Another important aspect is that surfactants located on the surfaces of nanoparticles lead to an insulating layer. This is a large disadvantage for numerous applications of nanoparticles requiring electronic transport.

Little is known about electrically conducting surfactants or amphiphiles in general. Li and co-workers presented in an interesting study a molecular hybrid of fullerene attached to a polythiophene oligomer. They showed some amphiphilic...
properties.\textsuperscript{12} Hecht and co-workers achieved a \textit{meta}-phenylene ethynylene derivative bound to an oligo-ethylene oxide chain as a hydrophilic constituent of the amphiphile. The authors reported intriguing, helical self-assembly.\textsuperscript{13} For an amphiphile, or finally a surfactant, appointed for molecular electronics, it becomes clear that the isolating character of typical, hydrophilic headgroups such as oligo-ethylene-oxide or ammonium is a large disadvantage. Therefore, a class of surfactants comprising a hydrophobic and electrically conducting chain and a (hydrophilic) headgroup, which is also capable of conducting electrons, would represent a major advance.

Recently, organic–inorganic hybrid surfactants with heads containing transition metal species were receiving attention because of their ability to self-assemble could be combined with added properties characteristic for coordination compounds (magnetic properties, catalytic activity, \textit{etc.}).\textsuperscript{14} Our group could identify a surfactant class with a pure inorganic head, namely, a lacunary polyoxometalate (POM) cluster \([\text{PW}_{11}\text{O}_{39}]\), and conventional alkyl chains as hydrophobic moieties. From our previous work we know the polytungstate head can be reduced (for example by electrochemical methods),\textsuperscript{18} and, in addition, reduced POMs belong to compounds with partially delocalized electronic systems (Robin–Day classification II).\textsuperscript{26} Thus, \([\text{PW}_{11}\text{O}_{39}]\) is a promising candidate for the desired, electrically conducting headgroup. Obviously the alkyl chain needs to be exchanged by a conducting entity, \textit{e.g.}, a \(\pi\)-conjugated system. Others have already shown the attachment of POM species to \(\pi\)-systems such as dyes (for example, perylene) can be done in general, but their target was not the synthesis of surfactant headgroups such as oligo-ethylene-oxide or ammonium is a becomes clear that the isolating character of typical, hydrophilic

RESULTS AND DISCUSSION

\textbf{Surfactant Preparation.} The target compound (3), also noted as \([\text{PW}_{11}\text{O}_{39}]\)TPE, a lacunary \([\text{PW}_{11}\text{O}_{39}]\) polytungstate cluster attached to two \textit{para}-phenylene ethynylene trimers via siloxane bridges, was synthesized as depicted in Scheme 1.

The isotope lines of the ESIMS signal centered at \(m/z = 1239 \text{ g mol}^{-1}\) and recorded in negative ion mode are separated by 0.33 unit, affirming the charge of the ion is \(=3\). Thus, the resulting mass of the molecular species \((m \times z = 3717 \text{ g mol}^{-1})\) matches nicely the nonfragmented molecular ion \(3\), and as a result, there is a perfect match with the theoretically expected isotope pattern (Figure 1c). In the \({}^{183}\text{W}\) NMR spectrum shown in Figure 1b one sees the characteristic 2:2:1:2:2:2 pattern of the lacunary \([\text{PW}_{11}\text{O}_{39}]\) POM cluster bound to two organo-siloxane units. Thus, one can infer the molecular architecture is intact. Also in the \({}^{31}\text{P}\) NMR spectroscopy (Figure S-2b) one sees only one signal (\(\delta = -13.94), \text{ which is characteristic for the central } \text{PO}_{4}^{2-} \text{ unit in } [\text{PW}_{11}\text{O}_{39}] \text{ and proves the high purity of the compound. FT-IR spectroscopy (Figure S-2a) confirms the latter results, and it can be seen that also the CC triple bonds are still present (\(\tilde{\nu} = 2150 \text{ cm}^{-1}\)). The presence of the characteristic Si–O–Si vibration (\(\tilde{\nu} = 1110 \text{ cm}^{-1}\)) is also in full agreement with the proposed structure of the surfactant. Therefore, the molecular structure of the target compound is proven unambiguously.

\textbf{Investigation of Self-Organization.} Similar to the alkyl-modified \([\text{PW}_{11}\text{O}_{39}]^{\text{2-}}\) surfactants (for example \((\text{CH}_3)_{14}\text{CH}_2\text{SiO}_{0.5})_2[\text{PW}_{11}\text{O}_{39}]^{2-}\) (4) as a reference; all reference systems used in this study are summarized in Supporting Information Figure S-3),\textsuperscript{15–18} we also expect that compound 3 has amphiphilic properties because of its dipolar character considering the hydrophilic and charged headgroup and (two) attached, hydrophobic hydrocarbon chains. Water is added to 3, and for concentrations \(c > 1 \text{ g/L}\) one still observes a sediment remaining in the test vial. This indicates the solubility of 3 is lower compared to classical organic surfactants, \textit{e.g.}, SDS (sodium dodecyl sulfate), and also the alkyl analogue 4. However, even at \(c = 0.1 \text{ g/L}\) we could observe the formation of self-assembled structures in water (see Figure 2). In particle size distribution functions obtained from dynamic light scattering (DLS) one sees two signals at a hydrodynamic diameter \(D_H = 4.9\) and 18 nm (Figure 2a).

The critical micelle concentration (CMC) of surfactants such as SDS (CMC = 2.3 g/L) or polysorbate 80 (CMC = 0.16 g/L).
is typically larger. Attempts to determine the CMC of PW11TPE precisely were not successful, because ring tensiometry fails for all surfactants with highly charged PW11 heads25 and other methods (such as DLS) are no longer sensitive to such low concentrations (<0.1 g/L). The overall lower solubility and lower critical aggregation concentration are assigned to the presence of the \( \pi \)-conjugated chain, which could induce other, stronger intermolecular interactions in the hydrophobic block such as \( \pi-\pi \) bonding. For further clarification, we have performed small-angle X-ray scattering (SAXS) measurements. Considering the Kratky plot of the SAXS data (see Supporting Information Figure S-4), the shape of the observed aggregates is close to a rigid-rod morphology, indicated by the linear increase of the curve. Further, the dimension of the aggregates is \( D_{G1} \approx 6 \) nm and \( D_{G2} \approx 20 \) nm according to the two different slopes obtained from the Guinier analysis of the data (Figure 2b), which fits nicely with the values obtained from DLS. Images received from transmission electron microscopy (TEM) analysis (Figure 2c) are consistent with the latter results. The impression of “hollow structures” is typical for self-assembled structures resulting from [PW11O39]\(^{3-}\) surfactants due to the large difference in electronic density between the headgroup region and the organic tail.\(^{15,17}\) Although there is a large tendency for film formation, the size distribution of the objects is narrow and the objects are slightly elongated with dimensions of \( \sim 15-20 \) nm. Compared to alkyl-modified surfactants, for example the reference system 4, there are some important differences. \( D_H = 4.9 \) nm is obviously smaller than the double extension of PW11TPE (see Figure 1a), and this speaks for a substantial interdigitation of the hydrocarbon chains. Because the hydrocarbon chain can adopt only a stretched conformation because of the triple bonds, interdigitation is problematic since intersection becomes almost unavoidable for any classical shape of a micelle. However, density functional calculations show that the two phenylene ethynylene chains do not prefer a parallel orientation, but there is an angle between them of \( \sim 60^\circ \) (see Supporting Information S-5). Considering this specific geometry of the amphiphile 3, we propose the special structure of the micellar aggregates shown in Figure 2d. Our suggestion is in line with the crystal structure of a lacunary Dawson polyoxometalate cluster bound to a pyrene moiety.23 The model also accounts for the elongated shape of the aggregates, which could be a result of the additional \( \pi-\pi \) interactions of the phenylene ethynylene moieties. It is important to note that all structures involving charged surfactants contain substantial amounts of counterions. For surfactants with polyoxometalate heads this aspect is crucial because of the high negative charge. We have recently explored polyoxotungstate surfactants with charges up to \( -5 \), and we saw that not only the choice of cation is important but also surfactants with charges higher than \( -3 \) exhibit unusual self-assembly features because of the inevitable electrostatic repulsion.25 Here, we have not varied the counterion. Sodium compounds were used in all cases. We assume that hydrated Na\(^+\) remains in close proximity to the
negative headgroup, but the exact position cannot be resolved as for other systems known in the literature.\textsuperscript{26–28}

The self-assembly at high concentration of PW\textsubscript{11}TPE (lyotropic LC) was studied using SAXS and TEM given in Figure 3. The water content in the samples was determined by thermogravimetric analysis and was found to be 4.7%. At low angles, two diffraction signals are present, which fit to a hexagonally ordered cylindrical liquid crystal phase (\(a = 4.99\) nm) (space group \(P_{6nmm}\)). In addition, there is one signal at higher \(q\) values (\(c = 1.14\) nm), which we assign to a dense packing of the \([\text{PW}_{11}\text{O}_{39}]\) headgroups located at the surface of the cylindrical aggregates (see Figure 3a).

However, the internal order is not high, which is signaled by the significant width of the SAXS signals. The latter is confirmed by TEM investigations, for which we could only find a disordered, wormlike phase (Figure 3b). Compared to the alkyl derivative as a reference (PW\textsubscript{11}C\textsubscript{16}),\textsuperscript{15} performing electron microscopy is intricate because PW\textsubscript{11}TPE undergoes radiation damage much more easily.

**Investigation of Electronic and Electrical Properties.** Now that the amphiphilic properties of PW\textsubscript{11}TPE (3) have been examined and some self-assembled structures have been introduced, the question needs to be answered, if there is any electronic communication between the redox-active \([\text{PW}_{11}\text{O}_{39}]\) headgroup and the \(\pi\)-conjugated chain(s). We studied the electronic properties of 3 in solution first. As the solubility of PW\textsubscript{11}TPE is limited in water, electrical (cyclic voltammetry (CV)) and electronic (UV/vis and photoluminescence (PL)) characterization was carried out in organic solvents for better comparison to references (PW\textsubscript{11}C\textsubscript{16} and TPE) and to meet the required minimal concentrations for the analytical techniques. Figure 4a shows cyclic voltammetry data of PW\textsubscript{11}TPE in comparison to the alkyl analogue PW\textsubscript{11}C\textsubscript{16}.

Both compounds show two reduction/oxidation waves due to the redox pair W(VI)/W(V). Two effects can be seen. The reversibility of the redox process is limited for PW\textsubscript{11}C\textsubscript{16} as documented by the second oxidation wave in comparison to the corresponding reduction wave. Interestingly, for PW\textsubscript{11}TPE the redox reversibility has increased significantly. Further, the reduction waves have shifted to lower potentials (−840 to −750 mV), meaning the transfer of one electron to the headgroup is eased.

It can be reasoned there is clearly an effect of the \(\pi\)-conjugated chain attached to the \([\text{PW}_{11}\text{O}_{39}]\) headgroup. Photoluminescence measurements also show an influence of the attached \(\pi\)-conjugated chain (Figure 4b). Whereas there is no or only limited signal for PW\textsubscript{11}C\textsubscript{16} (2) and PW\textsubscript{11}C\textsubscript{16} (4), the presence of the TPE chain leads to a noticeable fluorescence signal in the VIS region. Attaching the \([\text{PW}_{11}\text{O}_{39}]\) moiety to the TPE chain induces a red-shift of the fluorescence by \(\Delta\lambda_{\text{max}} = 18\) nm. From fluorescence lifetime measurements (Figure 4c) one sees there is clearly a biexponential decay for PW\textsubscript{11}TPE and the related relaxation constants are markedly different (see fits given in Supporting Information Figure S-6). Because the electronic system of \([\text{PW}_{11}\text{O}_{39}]\) is very different from the organic TPE chain, it is expected that disparate decay constants exist. Because excitation is based on the adsorption band of TPE, the occurrence of both decays is an indication of the successful transfer of the excited state from the chain to the headgroup. The bare TPE chain (for better comparison also fitted by a biexponential curve) compares well to the slower decay rate of PW\textsubscript{11}TPE, and the second component is much faster. Since it is known that fluorescence lifetimes are influenced by the presence of heavy atoms, the latter results can be interpreted as an additional evidence for the delocalization of the photogenerated, excited state from the \(\pi\)-conjugated chain to the polyoxometalate headgroup containing the heavy W atoms. We wanted to gain further, independent confirmation of the latter by using spectroelectrochemical (SEC) IR measurements (see Supporting Information Figure S-7). We expected a shift of the C≡C stretching vibration because of the occupation of antibonding orbitals, when an electron of a reduced \([\text{PW}_{11}\text{O}_{39}]\) headgroup would eventually hop to the \(\pi\)-conjugated chain. Clearly there is no shift, and as a result, one has to assume the transfer of electrons from the headgroup to the chain in solution is at least aggravated. Several factors can be responsible for this: First, one has to consider the reduction of conjugated \(\pi\)-systems is in general not favorable.\textsuperscript{29} Then, in solution rotation of the chain
about the Si–Csp2 bond is possible, and therefore the chances for overlap between the head and the π-system are reduced.

Therefore, it makes sense to investigate the described solid-state structure of 3 next. As expected, it can be seen that “bare” TPE is a yellow compound with an adsorption maximum at \( \lambda_{\text{max}} = 399 \) nm (\( E_h = 3.1 \) eV). The “bare” [PW11O39] also absorbs in the UV (Figure 5). Considering a “particle in a box model”, the significant red-shift of the absorption band of PW11OTPE compared to TPE can be explained by an extension of the electronic system. Further, one sees the band of 3 is significantly broader, and this is indicative for a bigger dispersion of the density of states (DOS) function. This and the fact that the optical properties of 3 in solution differ so much suggest that PW11TPE in its solid state has less molecular character and is gaining semiconducting properties. As a result, charge carriers can be delocalized over dimensions larger than just the molecular scale. This theory is confirmed by PL measurements shown in Supporting Information Figure S-8.

While pure TPE (5) is characterized by several distinct fluorescence features, all signals are quantitatively quenched for 3. We explain this by the increased delocalization of the excited state also to the headgroup region and fluorescence quenching because of the high atomic number of W, in addition to the
known “self-quenching” effect, due to excimer formation in high-concentration samples. 30

Final proof that PW11TPE truly is a semiconductor comes from current–voltage measurements and from testing the materials as a photoconductor (Figure 6). One sees that PW11TPE shows the typical I/V behavior of semiconductors. The asymmetry in the graph (Figure 6a) is presumably due to the possible redox process (reduction of the head; see above). When PW11TPE is irradiated with light, because of its semiconducting character, excitons (electron–hole pairs) are formed, and this increases the electrical conductivity of the material. Therefore, we can note an increase in current that quickly ceases when the light is turned off. Interestingly, afterward the resistance seems to be slightly higher than before but then recovers over time. When considering the results from PL and photosensor experiments together, it gets obvious that very interesting semiconducting properties arose from the presented hybrid material in LC phase. The determined band gap of 2.6 eV is very similar to established semiconductors such as ZnSe (2.7 eV), which opens the door for manifold applications.

CONCLUSIONS

In the current paper not only do we show that π-conjugated systems can be used for constructing soft semiconductors via thermotropic liquid crystals, but the presented organic–inorganic hybrid amphiphile allows entering the world of lyotropic, soft semiconductors. Because the set of analytical techniques is different for the micellar state and the LLC phase, it is very hard to compare the results directly. But it was shown that the type of aggregation also has an influence on the degree of electronic delocalization. When there is less conformational freedom of the π-conjugated chain with respect to the polyoxometalate head (in the solid state), delocalization is improved. It has also been demonstrated that in condensed phases intermolecular charge transfer between adjacent surfactants is possible and that this contributes to the semiconductor features.

When imagining possible applications of our materials, a first idea is to relate to classical semiconductors with similar gap energy (e.g., ZnSe). 31,32 However, compared to semiconductor quantum dots, a soft semiconductor like that presented here is fundamentally different. Therefore, our system should not be seen as an alternative to inorganic nanoparticles. It is much more promising to identify possibilities benefiting from the features of the presented system. These are the surfactant characteristics and the presence of the polyoxometalate group. Tempting ideas are to use conducting surfactants for surface stabilization of nanoparticles, enabling direct interparticle charge transfer, or to explore the option for the generation of self-organizing photovoltaic films. In combination with the idea of electrically conducting micelles and considering the catalytic activity of heteropolyoxometalates we want to approach innovative routes in electrocatalysis. When redox-active compounds are confined in micelles with conductive walls, one can also imagine finding possible applications in energy storage by charge storage and transport via reduction and oxidation. However, before these perspectives can be realized, most importantly the solubility in water has to be improved, for example by attaching side-chains to the π-backbone. In addition, more advanced physical measurements are planned to understand the mechanism of charge generation and transport in these systems.

METHODS

General Information. All experiments involving Pd complexes were carried out under a nitrogen atmosphere using standard Schlenk techniques. Solvents were dried and distilled prior to use. Unless otherwise specified, reagents were used as received without further purification. Refer to Supporting Information Figure S-1 regarding numbering of compounds.

Synthesis of ((4-iodophenyl)ethynyl)trimethylsilane (ii). A solution of 13.2 g (40 mmol) of 1,4-diodobenzene (i), 1 mol % Pd(PPh3)4, 2 mol % CuI, and 50 mmol of trimethylsilylacetylene (TMSA) in 90 mL of tetrahydrofuran (THF)/NEt3 (2:1, v/v) was prepared. After stirring for 20 h at 50 °C the solvent was evaporated and the residue was extracted with pentane. After evaporation of the solvent, the crude product was purified by column chromatography on silica gel and pentane as eluent. 1H NMR (400 MHz, CDCl3): δ (ppm) = 7.72–7.54 (m, 2H), 7.21–7.10 (m, 2H), 0.25, (s, 9H, TMS).

General Procedure for the Pd-Catalyzed Coupling of Terminal Alkynes and (ii). A solution of (ii), 1 mol % Pd(PPh3)4, 2 mol % CuI, and 1.05 equiv of terminal alkyne were prepared in THF/NEt3 (2:1, v/v). After stirring for 20 h at 50 °C the solvent was evaporated and the residue was extracted with pentane. After evaporation of the solvent, the crude product was purified by column chromatography on silica gel and pentane as eluent.

Trimethylsilyl-protected Tri-iso-propylsilylacetylenyl Phenylacetylene (TMS-iii): colorless oil; yield 85%; 1H NMR (400 MHz, CDCl3) δ (ppm) = 7.39 (s, 4H, Ph), 1.13 (m, 21H, TIPS), 0.25 (s, 9H, SiMe3).
Trimethylsilyl-protected Triiso-propylisilylacetylenyl bis-phenyl-acetylene (TMS-i:)(ppm): yellowish powder; yield 91%; \(^{1}H\) NMR (400 MHz, CDCl\(_3\)) \(\delta\) ppm = 7.45 (m, 8H, both Ph), 1.14 (m, 21H, TIPS), 0.26 (s, 9H, SiMe\(_3\)).

**General Procedure for the Selective Cleavage of the Trimethylsilyl Group.** TMS-protected alkyne was dissolved in 240 mL of CH\(_2\)Cl\(_2\)/MeOH (1:2, v/v), and 15 equiv of K\(_2\)CO\(_3\) was added. After stirring the resulting dispersion for 2 h, 240 mL of water was added. After separation of the phases, the organic solvent was removed. The obtained product was used without further purification.

(iii): white powder; yield 92%; \(^{1}H\) NMR (400 MHz, CDCl\(_3\)) \(\delta\) ppm = 7.42 (s, 4H, Ph) and 7.45 (s, 4H, both Ph), 3.15 (s, 1H, C=C), 1.13 (m, 21H, TIPS).

Preparation of Si-TPE (1; see Scheme 1) by Hydrosilylation of (v). To a degassed solution of 300 mg (0.6 mmol, 1.0 equiv) of (v) in 30 mL of CH\(_2\)Cl\(_2\) were added 102 mg of triethoxysilane (0.6 mmol, TMS-Si(OEt)\(_3\)). After stirring for 48 h the solvent was removed. The resulting dark yellow residue 1 was used without further purification.

\(^{29}Si\) NMR (79 MHz, CDCl\(_3\)) \(\delta\) ppm = -1.51.

**Synthesis of the Hybrid Polyoxometalate Species PW\(_{11}\)-TPE (3).** The lacunary polyoxotungstate K\(_{7}\)PW\(_{11}\)O\(_{39}\) was synthesized according to known literature. \(^{1}H\) NMR (400 MHz, DMSO) \(\delta\) ppm = 8.00 (s, 55H, aromatic protons), 3.13 (s, 36H, NMe\(_4\)), 1.09 (broad, 42H, TIPS); \(^{13}C\) NMR (162 MHz, DMSO) \(\delta\) ppm = 102.10 (6 siloxy groups), 81.35 (6 aromatic C), 54.76 (6 alkyl groups), 56.38 (6 alkoxy groups).

**Ion Exchange with Na-PW\(_{11}\)-TPE.** Cations were exchanged with Na\(^+\) by slow filtration of a 5 mg/mL solution of TMA-POM in acetonitrile through a column packed with Amberlite-IR120-Na. The success of the exchange was controlled via \(^{1}H\) NMR (no peak at \(\delta = 3.23\) tin any more).

**Preparation of High-Concentration Phases.** A dispersion of 100 mg of PW\(_{11}\)-TPE in water was prepared. Acetonitrile was added until full homogenization had taken place. Due to its lower boiling point (82 °C) compared to water, it is removed in a second step slowly under vacuum. The content of water can be controlled by the evaporation time, and its content was investigated using thermogravimetric analysis (TGA).

**Analytical Methods.** NMR measurements (\(^{1}H\), \(^{13}C\), \(^{29}Si\), \(^{31}P\)) were performed on a Varian Unity INOVA 400 spectrometer. The \(^{1}H\) NMR spectra were recorded on a Bruker Avance III 600 MHz spectrometer with 10 mm NMR tubes. ESIMS data were acquired on a Bruker microtolf II system. The solutions were injected directly into the evaporation chamber. SAXS was acquired on a Bruker NanoStar system equipped with pinhole collimation and Cu K\(_{α}\) radiation. The samples were placed between X-ray transparent foils and were measured in an evacuated chamber. Liquid samples were sealed in 1 mm Mark tubes made of soda lime glass. TEM was acquired on a Zeiss Libra 120 system and a JEOL JEM-2200FS. The dry sample was placed directly on carbon-coated copper grids. IR spectroscopy was performed on a PerkinElmer 100 system. UV/vis spectroscopy was performed on a Varian Cary 100. Dynamic light scattering was measured on a Viscotek 802 DLS machine. Photosensor measurements were performed on 3 mm × 3 mm sensor substrates from Umweltsensortechnik and measured with a Zahnem IM6 potentiostat. A solution of Na-PW\(_{11}\)-TPE in acetonitrile/water (1:1) solution was subsequently dropped onto the substrate and allowed to dry. Fluorescence spectroscopy was performed on a FluorTime 3000. All experiments were performed with \(\lambda_{ex} = 405\) nm. Because signals were too noisy in pure water due to the low concentration of PW\(_{11}\)-TPE, solutions with higher concentration (\(c = 5\) g/L) were prepared using THF. Cyclic voltammetry was measured with an Epsilon-potentiostat (BASI) (reference electrode: Ag/Ag\(^{+}\) wire; counter electrode: Pt wire; scan rate: 100 mV/s). Because signals were too noisy in pure water due to the low concentration of PW\(_{11}\)-TPE, solutions with higher concentration (\(c = 1\) g/L) were prepared using dimethyl sulfoxide (DMSO). TGA was recorded using a Netzsch TG 203 instrument.

**ASSOCIATED CONTENT**

3 Supporting Information

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**Author Contributions**  
A.K. has performed the experiments. S.P. has designed the research and wrote the paper. All authors have given approval to the final version of the manuscript.

**Notes**  
The authors declare no competing financial interest.

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