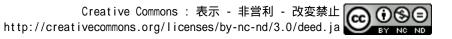
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Aggregation-Induced Emission of Water-Soluble Tetraphenylethene Derivatives at Polarized Liquid|Liquid Interfaces

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ABSTRACT

Aggregation-induced emission (AIE) behavior of water-soluble tetraphenylethene (TPE) derivatives bearing carboxy and sulfo groups was studied at polarized liquid liquid interfaces. The aggregation behavior of TPE derivatives in solution and at the water 1,2-dichloroethane (DCE) interface was highly dependent on their ionizable functional groups. Spectroelectrochemical analysis elucidated that the TPE derivatives were transferred across the interface accompanied by the adsorption process at the interface. The ion transfer and interfacial AIE features of TPEs responded reversibly to the externally applied potential, indicating no rigid crystalline structure formation in the interfacial region. The red-shift measured in intense interfacial emission spectra demonstrated that the carboxylate derivatives formed their J-aggregates specifically at the polarized water|DCE interface, while the aggregation processes with distinguishable emission properties took place in both interfacial region and organic solution in the sulfonate derivative system. The AIE features were also investigated at a glycerophospholipid-adsorbed interface as a model of biomembrane surface. The aggregation process of TPE derivatives was significantly modified through the interaction with phospholipid layers which stimulate the interfacial AIE process of tetra-anionic TPEs.

KEYWORDS: Aggregation-induced emission (AIE); Interface between two immiscible electrolyte solutions (ITIES); Tetraphenylethene (TPE); Polarization-modulation total internal reflection fluorescence (PM-TIRF); Potential-modulated fluorescence (PMF)

INTRODUCTION

A variety of organic luminescent molecules have been developed as spectrophotometric reagents for trace detection, bioimaging probes, sensitizers for photo-energy conversion, hosts/dopants for organic light-emitting devices and so on. The conventional aromatic fluorescent dyes with π conjugated system show a concentration quenching effect often referred as aggregation-caused quenching (ACQ) with π - π stacking interaction.¹⁻³ ACQ of fluorescent dyes impedes their effective applications on surfaces and at interfaces, where an enrichment of dye molecules with a certain surface-activity takes place spontaneously. Recent developments of aggregation-induced emission (AIE) system demonstrate promising applications in functional materials, optoelectronic and biomedical purposes.⁴⁻⁷ Tang and his coworkers have reported effective AIE features of a series of propeller shape compounds such as hexaphenylsiole (HPS) and tetraphenylethene (TPE) derivatives which undergo nonradiative relaxation in their monomer forms.⁸⁻¹⁰ These AIE luminogens (AIEgens) exhibit emissive features in the aggregated state because of the restriction of intramolecular motion (RIM), including rotation and vibration of the multiple phenyl groups.¹¹, ¹²

An interface between two immiscible electrolyte solutions (ITIES) has been utilized as a twodimensional reaction field for the formation of nanomaterials and 2D supramolecular architectures based on specific adsorption and phase transfer of reactants,¹³⁻¹⁶ and as a model of biomembrane surface to evaluate mass transport and pharmacokinetics in biomimetic systems.¹⁷⁻¹⁹ Various dye species and nanomaterials show self-aggregation/assemble features at polarized liquid|liquid interfaces, where the extent of aggregation of charged species can reversibly be controlled as a function of externally applied potential.²⁰⁻²⁴ The potential-driven aggregation accompanied by a colorimetric response is favorable for biomembrane imaging with a membrane potentialsensitivity. The membrane potential of the living cell plays a crucial role in physiological processes, including nerve impulse transmission and mass-transfer process of charged species moving into or out of the cell. A variety of voltage-sensitive dyes have been developed and used as a molecular probe in biochemical applications.²⁵⁻³⁰ Those dyes, however, exhibit emission properties mostly in the ACQ mechanism, which is not ideal to evaluate membrane potential with high-sensitivity in fluorescence imaging. Some AIEgens demonstrate their abilities as a membrane potential indicator for organic tissues as well as a photosensitizer for cancer phototherapy with selective staining of specific cells.³¹⁻³⁵ The adsorption and aggregation behavior of AIEgen at liquid|liquid interfaces, especially at a phospholipid-adsorbed biomimetic interface, under electrochemical control are thus important to elucidate their membrane potential-sensitive mechanism. It has been reported that surface-active AIEgens show the adsorption-promoted aggregation and emission induced by efficient lateral interaction of adsorbed molecules at liquid phase boundaries and on solid surfaces, ³⁶⁻³⁸ but no detailed analysis of the AIE effect at the liquid|liquid interface.

In this study, the interfacial aggregation and AIE features of anionic TPE derivatives were investigated water|1,2-dichloroethane (DCE) interface by at the surface-sensitive spectroelectrochemical techniques, i.e., potential-modulated fluorescence (PMF) spectroscopy and polarization-modulation total internal reflection (PM-TIRF) spectroscopy.³⁹⁻⁴¹ The analysis of acmodulated optical signals in the potential- and light polarization-modulation spectroscopies effectively improves the selectivity to the interfacial optical signals in comparison to the conventional differential voltfluorometry and TIRF spectroscopy.⁴²⁻⁴⁴ PMF spectroscopy is a powerful tool to elucidate a potential-driven interfacial process of charged species, while PM-TIRF spectroscopy allowed us to characterize the adsorption species oriented at the interface. The aggregation behavior of TPE derivatives in solution and at ITIES were highly dependent on their ionizable functional groups, and their interfacial AIE behavior responded to the externally applied potential. The AIE features of TPE derivatives were also examined at a biomimetic phospholipid-adsorbed interface.

EXPERIMENTAL SECTION

Reagents. Tetrakis(4-sulfophenyl)ethene (H₄TPETS) and its lithium salt were synthesized from tetraphenylethene (TPE) (Sigma-Aldrich, 98%) according to a literature method.^{45, 46} TPE was stirred in conc. H₂SO₄ at 388 K for 4 hrs and then the resulting H₄TPETS was obtained as precipitate by adding the reaction mixture dropwise into an ice-cooled ethyl acetate. The precipitate was washed with ethyl acetate and dried in vacuum at 353 K. The H₄TPETS recrystallized from methanol-ethyl acetate mixture was dissolved in 0.10 mol dm⁻³ LiOH aqueous solution and the evaporation of water left a readily water-soluble TPETS lithium salt (cf. ¹H NMR data in Supporting Information: S1). Tetrakis(4-carboxyphenyl)ethene (H₄TPETC) and 1,2di(4-carboxyphenyl)-1,2-diphenylethene (H₂TPEDC) were purchased from Strem Chemicals (99%) and Sigma-Aldrich (98%), respectively and used without further purification. The aqueous solutions were prepared with purified water by a Milli-Q system (Millipore, Direct-Q3UV). All other reagents were of the highest grade available. The pH condition of the aqueous solution was adjusted by the addition of an adequate amount of H₂SO₄ (pH \leq 3.2) or 2.0×10⁻³ mol dm⁻³ buffers, i.e. (pH 5.6) CH₃COOH/CH₃COOLi, (pH 7) LiH2PO4/LiOH, and ($8.8 \le pH$) H3BO3/LiOH. The composition of the electrochemical cells is illustrated in Figure 1. The purified water and 1,2dichloroethane (DCE) (Nacalai Tesque, HPLC grade, \geq 99.7%) as an organic solvent were saturated with each other and used for solution preparation. The supporting electrolytes were

for the aqueous 1.0×10^{-2} mol dm^{-3} Li₂SO₄ phase and 5.0×10^{-3} mol dm^{-3} bis(triphenylphosphoranylidene)ammonium tetrakis(pentafluorophenyl)borate (BTPPATPFB) for respectively. **BTPPATPFB** metathesis the organic phase, was prepared by of bis(triphenylphosphoranylidene)ammonium chloride (BTPPACl) (Aldrich, 97%) and lithium tetrakis(pentafluorophenyl)borate ethyl ether complex (TCI) in a 4:1 mixture of methanol and water, followed by recrystallization from acetone. A neutral glycerophospholipid, 1,2-dimyristoylsnglycero-3-phosphocholine (DMPC) (TCI, $\geq 97\%$), was added to the organic phase at 2.0×10⁻⁵ mol dm⁻³ for the biomimetic water|DCE interface.

Ag		$1.0 \times 10^{-3} \text{ mol } \text{dm}^{-3}$ BTPPACl $1.0 \times 10^{-2} \text{ mol } \text{dm}^{-3}$ LiCl (aq)	BTPPATPFB	TPE derivative buffer 1.0×10^{-2} mol dm ⁻³ Li ₂ SO ₄ (aq)	Ag ₂ SO ₄	Ag
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Figure 1. Composition of the electrochemical cell.

Apparatus. Ion transfer voltammetry at the water|DCE interface was performed in a fourelectrode spectroelectrochemical cell with the interfacial area of 0.50 cm^{2,47} Cyclic and ac voltammograms were measured by a four-electrode potentiostat (Hokuto Denko, HA-1010mM1A) with a lock-in amplifier (NF, LI5640). The Galvani potential difference ($\Delta_0^w \phi \equiv \phi^w - \phi^0$) was estimated by taking the formal transfer potential ($\Delta_0^w \phi^o$) of tetramethylammonium ion as 0.160 V.⁴⁸ In the spectroelectrochemical measurements, the water|DCE interface was illuminated in total internal reflection (TIR) with the angle of incidence (ψ) at ca. 75° by a cw laser diode at 376 nm (Coherent, OBIS 375LX-50). The laser power was attenuated to 10 mW to avoid photobleaching of sample solutions. A Fresnel rhomb half-wave plate (Sigma Koki, FRH-102) was used to select the linear polarization of the fundamental excitation beam. The fluorescence signal from the interfacial region was detected perpendicularly to the interface by an optical fiber and a monochromator equipped with a photomultiplier tube (Shimadzu, SPG-120S). The absorption and emission spectra of the TPE derivatives in solution were measured by a JASCO V-750 UV-vis spectrophotometer and a JASCO FP-8300 spectrofluorometer, respectively. All experiments were carried out in a thermostated room at 298±2 K.

PMF Analysis. The PMF study was carried out by analyzing the fluorescence signal from the interfacial region as a function of ac potential modulation. The amplitude of the ac potential modulation $(\Delta_o^w \phi_{ac})$ was 20 mV at 1 Hz and the linear sweep rate of $\Delta_o^w \phi_{dc}$ was 5 mV s⁻¹ in negative sweep.

$$\Delta_{\rm o}^{\rm w}\phi = \Delta_{\rm o}^{\rm w}\phi_{\rm dc} + \Delta_{\rm o}^{\rm w}\phi_{\rm ac}\exp(\mathrm{i}\omega t) \tag{1}$$

where i is an imaginary number and ω is the angular frequency. The real (ΔF_{re}) and imaginary (ΔF_{im}) components of PMF signals were analyzed by a digital lock-in amplifier (NF LI5640). The PMF signals arise only from the polarized interfacial region and unmodulated dc fluorescence signals from the bulk solution phase is negligible. The phase shift of PMF signal provides us to elucidate the charge transfer mechanism including ion transfer and adsorption at either side of the interface. Further details of the analytical procedure are described elsewhere.³⁹

PM-TIRF Analysis. The linear light polarization of an excitation beam was modulated at 13 Hz between p- and s-polarizations (i.e. parallel and perpendicular to the plane of incidence) by a half-wave liquid crystal retarder (LCR) thermostated at 318 K (Thorlabs, LCC1111T-A, LCC25/TC200). The polarization modulation efficiency of LCR was estimated as $P_m = 0.85$ at 376 nm.^{40, 41} The PM-TIRF signal (ΔF^{p-s}) in the case of $P_m \neq 1$ value is expressed by

$$\Delta F^{p-s} = F^p_m - F^s_m = (2P_m - 1)(F^p - F^s)$$
⁽²⁾

where F_m^p and F_m^s are the modulated fluorescence intensities measured with p- and s-polarized excitation modes, F^p and F^s are the fluorescence intensities arisen from the species oriented at the

interface under p- and s-polarized excitations. The modulated fluorescence signal from the interfacial region was analyzed by a digital lock-in amplifier (NF, LI5640) as a function of periodic polarization modulation of the incident beam. The PM-TIRF spectrum was measured at a fixed potential and the potential dependence of ΔF^{p-s} was recorded at 2 mV s⁻¹.

RESULS AND DISCUSSION

AIE Activity of Water-Soluble TPE Derivatives in Solution. The fluorescence titration experiments of TPE derivatives in aqueous solution were performed in order to investigate their fundamental AIE features. Their solubilities are drastically decreased under acidic conditions and some precipitates were formed as a result of the charge neutralization through the protonation of carboxylates. The emission intensities measured in the TPETC and TPEDC systems were enhanced, respectively, at pH < 6 and pH < 4.5, indicating AIE activity of protonated species in acidic aqueous solutions (**Figure 2**). In addition, the progressive red shifts of absorption and emission bands are indicative of the formation of emissive aggregates of H₄TPETC and H₂TPEDC (**Supporting Information: S2**). In alkaline aqueous solution, the anionic forms of TPETC^{4–} and TPEDC^{2–}, indicated no emissive properties. TPETS^{4–} with highly acidic sulfo groups is unprotonated within the examined pH conditions and the intermolecular electrostatic repulsion prevents the formation of self-aggregates. Accordingly, the fluorescence intensity observed for the transparent solution of TPETS^{4–} was negligibly small at 1 < pH < 9, demonstrating that TPETS is substantially AIE-inactive in aqueous solution.

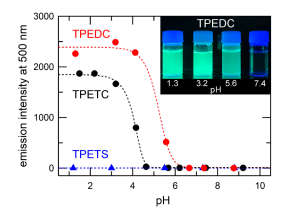


Figure 2. pH dependences of the emission intensity at 500 nm of TPE derivatives in aqueous solution. The red, black, and blue symbols depict TPEDC, TPETC, and TPETS, respectively. The excitation wavelength was 376 nm and the concentration of the TPE derivatives was 2.0×10^{-5} mol dm⁻³. **Inset**: Typical fluorescence image of TPEDC in aqueous solution under UV irradiation (365 nm).

The self-aggregation behavior of ionic species is generally influenced by solvent properties and the binary mixture system was employed to evaluate solvent effects on AIE features of anionic TPEs. A low polarity solvent, THF is miscible with water, thus the solvent effects arising from hydration-solvation, polarity and other physicochemical properties can be adjusted by a mixing volume ratio of water to THF. For instance, the relative permittivity (c_r) of water-THF mixture is approximately proportional to the volume fraction of water (f_w / vol%) between those of water ($c_{r,w}$ = 78.5) and THF ($c_{r,THF}$ = 7.6).⁴⁹ The water-THF mixtures with various water fractions were prepared by adding THF to the aqueous solution of TPE derivatives buffered at given pHs (**Supporting Information: S3**). All the examined TPE derivatives exist as nonemissive anionic monomer forms in alkaline aqueous solution (f_w = 100 vol%), whereas the emission intensities for tetraanionic TPE derivatives, TPETC⁴⁻ and TPETS⁴⁻, were drastically increased in low water fraction ($f_w \le 5$ vol%). These results coincide with previous reports for TPETS in the water-organic solvent mixture systems where the AIE-activity of TPETS with blue-shifted emission in water-THF mixtures is correlated with the self-assembly of nanostructured aggregates.⁴⁵ Bhosale and his co-workers reported rod-like nanostructures at $f_w \le 5$ vol% but the morphology of resultant emissive aggregates highly depends on the experimental conditions.⁴⁶ In contrast, the fluorescence from noncharged acid forms, H₄TPETC and H₂TPEDC, with AIE activity at pH 3 was totally quenched at $f_w < 80$ vol%. These neutral species were readily soluble and nonemissive in THF. The results obtained in the homogeneous solution system demonstrate that the charged and neutral species of the water-soluble TPE derivatives tend to form AIE-active self-aggregates, respectively, in low and high polarity media.

Ion Transfer Across the Water |DCE Interface. The ion transfer voltammograms of TPEDC²⁻ and TPETC⁴⁻ were measured under alkaline conditions since their protonated species affect the voltammetric response under acidic conditions, while those of TPETS⁴⁻ were obtained at pH 7.0. The ion transfer responses of $TPEDC^{2-}$ and $TPETC^{4-}$ were observed close to the negative edge of the polarizable potential window, i.e., $\Delta_o^w \phi_{TPEDC^2}^{\circ'} = -0.30 \text{ V}$ and $\Delta_o^w \phi_{TPETC^4}^{\circ'} = -0.41 \text{ V}$ (Figures **3a** and **3b**). TPEDC²⁻ exhibited well-defined quasi-reversible voltammetric responses for dianion with the peak separation of ca.30 mV and the diffusion coefficient in water was estimated as $D_{\text{TPEDC}^{2-}}^{\text{w}} = 2.2 \times 10^{-6} \text{ cm}^2 \text{ s}^{-1}$. Those parameters could not be obtained quantitatively for TPETC⁴⁻ owing to a significant overlap with the supporting electrolyte transfer. TPETS⁴⁻ exhibited complicate voltammetric waves at $\Delta_o^w \phi_{TPETS^4}^{o'} = -0.33$ V, where the negative current peak in CVs for the transfer of TPETS⁴⁻ from water to DCE was accompanied by a post-transfer response corresponding to a sharp admittance peak at -0.39 V (Figure 3c). These voltammetric responses could be related to the additional interfacial process of TPETS⁴⁻. On the other hand, the positive transfer current due to the back transfer to water was obtained as a well-defined diffusion-limited regime. In comparison with the base electrolyte system, the admittances measured in the watersoluble TPE systems were slightly increased at negative potentials, indicating that the adsorption of the charged TPE derivatives at the water|DCE interface.

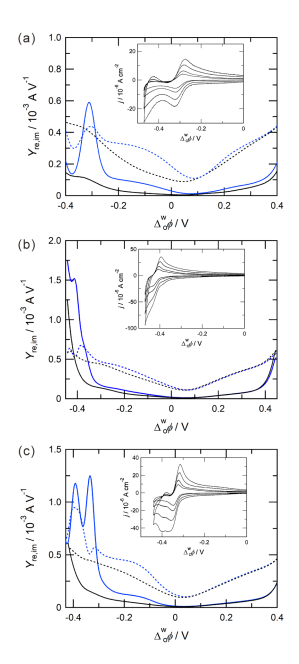


Figure 3. Ac voltammograms measured at the water|DCE interface for (a) TPEDC^{2–} at pH 8.8, (b) TPETC^{4–} at pH 8.9, and (c) TPETS^{4–} at pH 7.0. The solid and dashed lines depict the real (Y_{re}) and imaginary (Y_{im}) components of the admittance, respectively. The blue and black lines refer to the presence and absence of 5.0×10^{-5} mol dm⁻³ TPE derivatives. The ac potential modulation and the linear sweep rate were 10 mV at 7 Hz and 5 mV s⁻¹, respectively. **Insets:** Typical CVs measured at 10, 20, 50, 100 mV s⁻¹.

The PMF data measured for TPEDC²⁻ were shown in **Figure 4a**, in which $\Delta F_{\rm re}$ and $\Delta F_{\rm im}$ signals maximized around $\Delta_0^{\rm w} \phi_{\rm TPEDC^{2-}}^{\circ'}$ were observed with negative and positive signs in agreement with theoretical eqs. for ion transfer responses ($\Delta F_{\rm t,re}$ and $\Delta F_{\rm t,im}$).^{39, 50}

$$\Delta F_{t,re} = \frac{4.606\varepsilon \Phi_{f} I_{0}}{zFS \cos\psi} \left[\frac{\Delta_{o}^{W} \phi_{ac} \sigma \omega^{-3/2}}{(R_{ct} + \sigma \omega^{-1/2})^{2} + (\sigma \omega^{-1/2})^{2}} \right]$$
(3)

$$\Delta F_{t,im} = -\frac{4.606\varepsilon \Phi_{f} I_{0}}{zFS \cos\psi} \left[\frac{\Delta_{o}^{w} \phi_{ac} (R_{ct} + \sigma \omega^{-1/2}) \omega^{-1}}{(R_{ct} + \sigma \omega^{-1/2})^{2} + (\sigma \omega^{-1/2})^{2}} \right]$$
(4)

where ε is the molar absorption coefficient, $\Phi_{\rm f}$ is the fluorescence quantum yield, I_0 is the intensity of excitation light, F is Faraday constant, and S is the interfacial area. $R_{\rm ct}$ and σ are the charge transfer resistance and the Warburg term, respectively. Eqs.(3) and (4) provide negative $\Delta F_{\rm t,re}$ and positive $\Delta F_{\rm t,im}$ values for TPEDC²⁻ (z = -2). On the other hand, the adsorption responses of TPEDC²⁻ with $\Delta F_{\rm a,re} < 0$ and $\Delta F_{\rm a,im} > 0$ were obtained in the wide potential region ($\Delta_0^{\rm w} \phi_{\rm TPEDC^{2-}}^{\circ \circ} < \Delta_0^{\rm w} \phi < 0.25$ V), indicating a weak potential-dependence of the adsorption process at the aqueous side of the interface (eqs. (5) and (6)).

$$\Delta F_{a,re} = \frac{2.303 \varepsilon \Phi I_0 \Gamma Sz F}{RT} \left[\frac{b \Delta_0^w \phi_{ac} (k_{a,dc}^w c_0^w \alpha (1 - \theta_{dc}) - k_{d,dc}^w (\alpha - 1) \theta_{dc}) (k_{a,dc}^w c_0^w + k_{d,dc}^w)}{(k_{a,dc}^w c_0^w + k_{d,dc}^w)^2 + \omega^2} \right]$$
(5)

$$\Delta F_{a,im} = -\frac{2.303\varepsilon \Phi I_0 \Gamma Sz F}{RT} \left[\frac{b\Delta_o^w \phi_{ac} (k_{a,dc}^w c_0^w \alpha (1 - \theta_{dc}) - k_{d,dc}^w (\alpha - 1)\theta_{dc})\omega}{\left(k_{a,dc}^w c_0^w + k_{d,dc}^w\right)^2 + \omega^2} \right]$$
(6)

where Γ , α , $c_0^{\rm w}$ and $\theta_{\rm dc}$ are the saturated interfacial concentration, overall transfer coefficient for adsorption process, the bulk aqueous concentration and the dc surface coverage, respectively. $b\Delta_{\rm o}^{\rm w}\phi_{\rm ac}$ is a portion of the Galvani potential difference for the adsorption process. $k_{\rm a,dc}^{\rm w}$ and $k_{\rm d,dc}^{\rm w}$ are the dc components of the adsorption and desorption rate constants at given potentials, respectively. The PMF responses for TPEDC²⁻ seem to those of simple monomeric fluorescent dye but it should be noted that the monomeric TPETC exhibited only a feeble fluorescence in both aqueous and DCE solutions under present alkaline conditions, i.e. AIE-inactive in the bulk phases. Those PMF signals measured in the interfacial region are therefore associated with the interfacial AIE of the adsorbed species. The symmetrically charged TPETC⁴⁻ and TPETS⁴⁻ also exhibited significant enhancements of PMF intensities, respectively, at $\Delta_o^w \phi < \Delta_o^w \phi_{TPETC^4}^{\circ'}$ and $\Delta_o^w \phi < \Delta_o^w \phi < \Delta_o^{\circ'} \phi_{TPETC^4}^{\circ'}$ $\Delta_{o}^{w}\phi_{TPETS^{4}}^{\circ'}$ (Figures 4b and 4c), indicating their interfacial AIE features. The PMF results for TPETS⁴⁻, in particular, indicated apparent interfacial AIE activity maximized around -0.39 V in negative potential sweep, which is more than 60 mV negatively shifted relative to $\Delta_0^w \phi_{\text{TPETS}^4}^{\circ'} =$ -0.33 V. The delayed PMF enhancement would result from the aggregation after the accumulation of TPETS monomers at the interface. The TPETS monomer is AIE-inactive in the aqueous phase but it has a small AIE activity in the organic phase, where the emission maximum in DCE ($\varepsilon_{r,DCE}$ = 10.4) at 440 nm is located in a short-wavelength region compared to that in water at 466 nm as discussed below. Although the intense emission of TPETS in THF seems to associate with insoluble nanorod or particle formation,^{45,46} the voltammetric data showed the ion transfer features of tetravalent TPETS anions in a quasi-reversible diffusion-limited regime under present conditions (cf. Figure 3). The emission observed in the bulk organic phase therefore could arise from a readily dissociable self-aggregate of TPETS without forming a rigid crystalline structure.⁴⁶ It is also noteworthy that relatively large PMF intensity in the s-polarized light excitation mode was linked to a molecular orientation of fluorescent species at the interface.

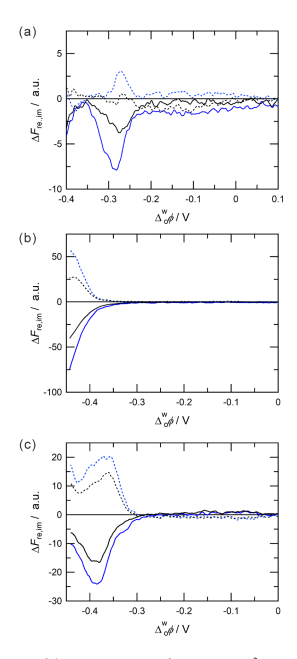


Figure 4. Potential dependences of the PMF responses for (a) TPEDC^{2–} at pH 8.8, (b) TPETC^{4–} at pH 8.9, and (c) TPETS^{4–} at pH 7.0. The solid and dashed lines depict the real (ΔF_{re}) and imaginary (ΔF_{im}) components of PMF, respectively. The blue and black lines refer to PMF measured, respectively, by the *s*- and *p*-polarized excitation beams. The concentration of the TPE derivatives was 5.0×10⁻⁵ mol dm⁻³. The ac potential modulation and the linear sweep rate were 20 mV at 1 Hz and 2 mV s⁻¹, respectively.

Characterization of Interfacial Aggregation States. The polarization-modulation fluorescence signals arise only from interfacial species adsorbed with a certain orientation, and undesirable optical signals from randomly oriented bulk species are, in principle, cancel out in the PM-TIRF experiments.⁴⁰ The intense PM-TIRF signals with negative sign ($\Delta F^{p-s} < 0$) were obtained for each TPE derivative (**Figure 5**). The negative ΔF^{p-s} signal is indicative of the excitation dipole moment of interfacial species relatively in parallel to the interface in accordance with the PMF results under the *s*-polarized excitation. The PM-TIRF responses showed a significant potential-dependence, where the ΔF^{p-s} magnitude was reversibly changed as a function of $\Delta_o^w \phi$ and enhanced around $\Delta_o^w \phi^{ot}$. The potential dependences of ΔF^{p-s} for TPETC and TPETS were not synchronized strictly with the voltammetric responses but keep in phase with the delayed PMF responses. As a result, the potential-dependent PM-TIRF responses measured for the interfacial AIE process coincide approximately with the PMF data (cf. **Figure 4**).

The PM-TIRF spectrum considered as emission spectrum of the interfacial species allows us to characterize the adsorption and aggregation states of the interfacial species at given potentials. **Figure 5b** shows the PM-TIRF spectra measured for TPEDC at the interface. As summarized in **Table 1**, TPEDC excited by a cw laser at 376 nm emits only a feeble fluorescence around 478 nm and 441 nm, respectively, in alkaline aqueous and organic phases. The intense PM-TIRF spectra therefore associate with the interfacial aggregates. The PM-TIRF spectra observed prior to the TPEDC^{2–} transfer from water to DCE ($\Delta_0^w \phi \ge -0.19$ V) have an intermediate emission maximum at 462 nm between those of bulk aqueous and organic species. The spectral features of adsorption species often reflect their specific physicochemical properties, and it may result from hydration/solvation state and polarity modified in the interfacial region. TPETC with $\Delta_0^w \phi_{\text{TPETC}^+}^{\circ\circ} = -0.41$ V indicated similar spectral changes at -0.42 V (**Supporting Information: S4**). In addition, the PM-TIRF spectra in the TPEDC system indicated apparent red-shift of the emission maximum from 462 nm at $\Delta_0^w \phi \ge -0.19$ V to 482 nm at -0.34 V, indicating a gradual formation of the J-type aggregates at the polarized interface around $\Delta_0^w \phi_{TPEDC^2}^{\circ, \circ}$.^{11, 51} TPETS also showed the red-shift of PM-TIRF spectrum at $\Delta_0^w \phi < \Delta_0^w \phi_{TPETS^4}^{\circ, \circ}$, in agreement with the J-aggregation after the interfacial accumulation of monomers (**Figure 5c** and **Table 1**). The PM-TIRF analysis demonstrated that the carboxylate and sulfonate derivatives specifically form AIE-active Jaggregates in the interfacial region by applying appropriate potentials.

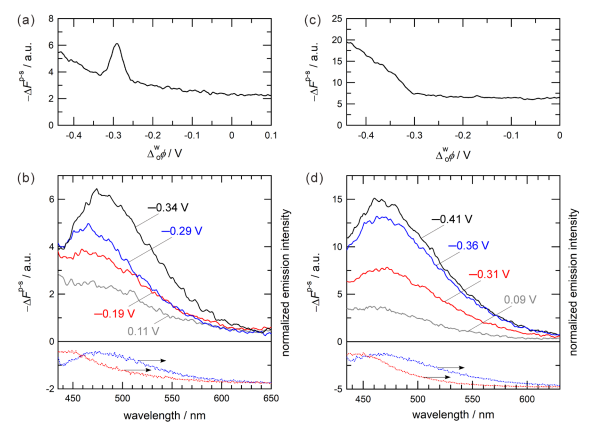


Figure 5. Potential dependences of PM-TIRF intensity at 460 nm and PM-TIRF spectra for **(a,b)** TPEDC^{2–} and **(c,d)** TPETS^{4–} at the water|DCE interface. The dotted lines refer to the normalized emission spectra measured in the aqueous (blue) and organic (red) phases. The concentrations of the TPE derivatives was 5.0×10^{-5} mol dm⁻³ in the aqueous phase (pH 9.0 for TPEDC^{2–} and pH 7.1 for TPETS^{4–}). **(a,c)** The potential sweep rate was 2 mV s⁻¹ in negative sweep.

	TPEDC system		TPETC system		TPETS system	
	$\Delta_{\rm o}^{\rm W} \phi / {\rm V}$	$\lambda_{\rm em,max}$ / nm	$\Delta_{\rm o}^{\rm w} \phi / { m V}$	$\lambda_{em,max}$ / nm	$\Delta_{\rm o}^{\rm w} \phi / {\rm V}$	$\lambda_{\rm em,max}$ / nm
	0.11	462	0.10	461	0.09	455
interface	-0.19	462	-0.35	458	-0.31	467
(PM-TIRF)	-0.29	465	-0.42	464	-0.36	469
	-0.34	482	-0.47	466	-0.41	469
aqueous phase	(pH 9.0)	478	(pH 9.0)	469	(pH 7.1)	466
organic phase		441		453		440

Table 1. Emission maxima of TPE derivatives at the water DCE interface and in solutions^a

^{*a*} The emission maximum wavelengths ($\lambda_{em,max}$) were measured with a cw laser at 376 nm.

AIE Behavior at Biomimetic Interfaces. The potential-induced interfacial AIE reaction found in the present study suggests the potential ability of simple water-soluble TPE derivatives to visualize the dynamic polarization of biomembranes or organic tissues. The DMPC-adsorbed water|DCE interface was employed for evaluating the AIE activity of anionic TPEs on a biomembrane surface. In all the examined systems, typical voltammetric features for phospholipid-adsorbed interfaces were observed, in which the capacitive current was reduced by the interfacial adsorption of neutral DMPC molecules and the facilitated lithium ion transfer appeared close to the positive edge of the potential window ($0.2 \text{ V} < \Delta_o^w \phi$) (Supporting Information: S5).⁵²⁻⁵⁴ The ion transfer responses of TPEDC²⁻ at $\Delta_o^w \phi_{TPEDC^{2-}}^{ov}$ were hardly affected by the DMPC layer but PM-TIRF results clarified the specific interaction between TPEDC and DMPC at the interface. As shown in Figure 6, the maximum intensity of negative ΔF^{p-s} signals was observed around -0.35 V, which is 50 mV more negative potential than $\Delta_o^w \phi_{TPEDC^{2-}}^{ov}$ at the bare water|DCE interface. The voltammetric and PM-TIRF results demonstrate that the potential-dependence of the interfacial AIE process of TPEDC is modified on the DMPC layer even though $\Delta_o^w \phi_{TPEDC^{2-}}^{ov}$ associated with the Gibbs energy of transfer between water and DCE (i.e. $\Delta G_{\text{tr.TPEDC}^2}^{w \to o} = -2F\Delta_o^w \phi_{\text{TPEDC}^2}^{\circ \circ}$) remained unchanged in the presence of DMPC. The emission maximum of the PM-TIRF spectrum at -0.34 V was measured at 470 nm, which is the intermediate value between the aqueous and organic phases, suggesting that the AIE mechanism of TPEDC²⁻ observed at the bare interface is effectively modified on the membrane surface (**Table 2**). In addition, no AIE was observed in the bulk organic phase including free DMPC monomers. The intense emission from the interfacial region in the biomimetic system could be associated with the direct RIM mechanism of TPEDC in the DMPC layer.^{11, 12}

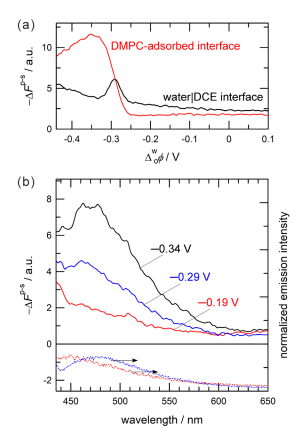


Figure 6. Potential dependences of (a) PM-TIRF intensity at 460 nm and (b) PM-TIRF spectra for TPEDC²⁻ at the DMPC-adsorbed interface. The dotted lines refer to the normalized emission spectra measured in the aqueous (blue) and organic (red) phases. The concentrations of TPEDC and DMPC were 5.0×10^{-5} mol dm⁻³ in the aqueous phase (pH 9.0) and 2.0×10^{-5} mol dm⁻³ in the organic phase, respectively. (a) The potential sweep rate was 2 mV s⁻¹ in negative sweep.

	TPEDC system		TPETC system		TPETS system	
	$\Delta_{\rm o}^{\rm w} \phi / { m V}$	$\lambda_{\rm em,max}$ / nm	$\Delta_{\rm o}^{\rm w} \phi / { m V}$	$\lambda_{\rm em,max}$ / nm	$\Delta_{\rm o}^{\rm W} \phi / { m V}$	$\lambda_{\rm em,max}$ / nm
					0.09	473
interface	-0.19	_	-0.35	464	-0.31	475
(PM-TIRF)	-0.29	460	-0.42	479	-0.36	484
	-0.34	470	-0.47	479	-0.41	484
aqueous phase	(pH 9.0)	478	(pH 9.0)	469	(pH 7.1)	466
organic phase		441		457		468

Table 2. Emission maxima of TPE derivatives at the DMPC-adsorbed interface and in solutions^a

^{*a*} The emission maximum wavelengths ($\lambda_{em,max}$) were measured with a cw laser at 376 nm.

In the symmetrically charged TPETC^{4–} and TPETS^{4–} systems, the ion transfer responses in ac voltammograms were weakened and negatively shifted at the biomimetic interface (**Supporting Information: S5**). The slow kinetic process across the DMPC layer could be responsible for the small voltammetric responses. On the other hand, the large red-shift of emission maxima in the PM-TIRF spectrum is associated with the promotion of J-aggregation on the DMPC layer (**Figure 7** and **Table 2**). In addition, the emission spectrum of TPETS measured in the organic phase containing free DMPC molecules showed a weak emission with a 28 nm red-shifted maximum at 468 nm, indicating the effective interaction between the TPETS anions and DMPC in the bulk organic phase. These results indicate that the DMPC layer stimulates the interfacial AIE process of tetra-anionic TPEs, in which the interaction with DMPC was more effective for TPETS.

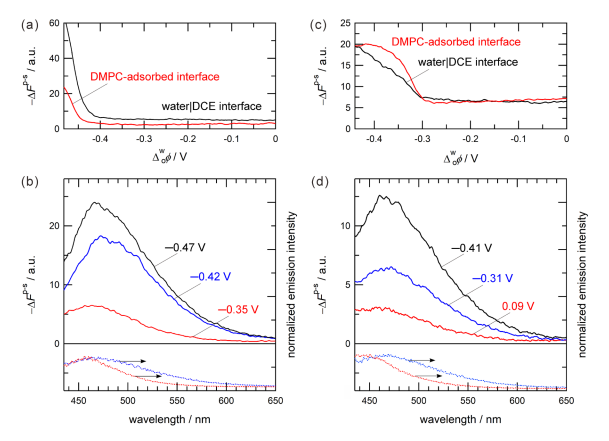


Figure 7. Potential dependences of PM-TIRF intensity at 460 nm and PM-TIRF spectra for **(a,b)** TPETC^{4–} and **(c,d)** TPETS^{4–} at the DMPC-adsorbed interface. The dotted lines refer to the normalized emission spectra measured in the aqueous (blue) and organic (red) phases. The concentrations of the TPE derivatives and DMPC were 5.0×10^{-5} mol dm⁻³ in the aqueous phase (pH 9.0 for TPETC^{4–} and pH 7.0 for TPETS^{4–}) and 2.0×10^{-5} mol dm⁻³ in the organic phase, respectively. **(a,c)** The potential sweep rate was 2 mV s⁻¹ in negative sweep.

Conclusions

The interfacial AIE behavior of the anionic TPE derivatives examined in this study was reversibly modulated depending on the externally applied potential. The anionic carboxylate derivatives, TPEDC^{2–} and TPETC^{4–}, exhibited the interface-sensitive AIE feature but no emission from the bulk aqueous and organic phases. On the other hand, TPETS^{4–} formed the aggregates with distinguishable emission properties at the interface and in the organic solution. Those specific AIE

responses show the advantage of the TPE-based AIEgens for sensitive-imaging of liquid interfaces and surfaces of micelles and polymers often buried in optical responses arising from bulk solution phases. The potential-induced AIE behavior observed for simple water-soluble TPE derivatives at the phospholipid-adsorbed interface is also important to develop the membrane-potential sensitive probe in biomedical applications, especially, noncovalent labeling of liposomes and exosomes.

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Supporting Information.

¹H NMR data for TPETS, UV-vis absorption and emission spectra in aqueous solutions, AIE responses in THF-water mixtures, PM-TIRF data for TPETC, ac voltammograms at biomimetic interfaces.

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