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Luminescence of triarylphosphines and their application to detection of elemental chlorine in aqueous solution†

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The concept of aggregation induced emission has been utilized to raise luminescence of triarylphosphines for the first time. Appreciable emission in the aggregated state as well as the Lewis basic character of these compounds were capitalized upon to demonstrate the functional utility as fluorescent chemosensors for elemental chlorine, Cl_2 , in aqueous solution.

Among phosphorus compounds, phosphines¹ have widely been employed in biology, nature and chemical industries as antioxidants, insecticides, herbicides, flame-retardants, catalysts, surfactants, nerve agents, energy transporters, semiconductors, dopants, plastics and so on. Curiously, the luminescence characteristics of phosphines have not been reported to date.¹,² The possible reasons for this are the free rotation of the substituents in the phosphine scaffold, and inversion of the lone-pair of electrons on the phosphorus atom.³ These procedures readily extinguish the emission of any fluorophore through electron transfer quenching⁴ and\or depopulation of the excited states *via* non-radiative mechanisms by internal conversions.⁵

Almost all non-radiative decay processes can be overcome readily by adopting suitable experimental conditions. ^{4,5} For example, the amino-functionalized anthracenes are weakly emissive in acetonitrile solution, however their photoluminescence quantum yields ($\Phi_{\rm PL}$) are raised by more than 200-fold by protonation of the amino-group/s. ⁴ Likewise, tetraphenylethene and its derivatives are found to be faintly emissive or nonemissive in dilute solutions, at room temperature, due to the active intramolecular motions of the multiple aryl rotors with respect to the central olefinic double bond. The $\Phi_{\rm PL}$ value has been shown to be increased by 5–150-fold by restriction of

the rotation of aryl rings through aggregation 5,6 or encapsulation. 7

As luminescent materials^{2,4-6,8} find many applications in dyes, optical sensors, molecular electronics, nonlinear optics, light emitting diodes, photovoltaic cells, field-effect transistors, etc., the present study was aimed at exploring the luminescence characteristics of triarylphosphines (TPPs, Chart 1) in particular. The interest in TPPs was spurred by the fact that they are available readily in pure form,1 insensitive to water and oxygen under ambient conditions, 1,2 fluorescent in the solid state as revealed by emission with a hand-held UV lamp, and should be expected to exhibit aggregation behaviour in polar solvents due to their hydrophobic nature. In addition, TPPs are powerful nucleophiles, and may form complexes with suitable analytes leading to changes in photophysics. Herein, I report the absorption and photoluminescence (PL) properties of three TPPs (Chart 1) in anhydrous as well as aqueous tetrahydrofuran (THF) and in the solid state. Also, the study accounts for the sensing activities of these TPPs for the detection of elemental chlorine. To the best of my knowledge, emission features of phosphine derivatives have not been heretofore investigated in detail.1,2,9

The optical properties of TPPs (Chart 1) were investigated by UV-vis absorption and fluorescence spectroscopy and the details are summarized in Table 1.

Fig. 1 shows the absorption spectrum of **TPHP** (1 \times 10⁻⁴ M) in THF and the water–THF mixture containing different water fractions from 10 to 95%. Independent of the medium very similar wavelengths of absorption (λ_{ab}) were observed, however, profile shapes were found to be broader and the absorption tails

Chart 1 Chemical structure of propeller-shaped TPPs investigated.

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Table 1 Optical properties of TPHP. TOTP and TPTP in THF, water-THF mixture and solid states

				$\lambda_{\mathrm{em}}^{f}(\mathrm{nm})$						
Luminogen	$\lambda_{ab,s}^{a}$ (nm)	$\lambda_{ab,a}^{b}$ (nm)	E_{g}^{c} (eV)	Soln ^g	Aggr. ^h	Frozen ⁱ	Film ^j	$\Phi_{\mathrm{PL,s}}{}^{k}\left(\% ight)$	$\Phi_{\mathrm{PL,a}}{}^{l}\left(\% ight)$	$d_{\rm H}^{m}$ (nm)
ТРНР	234, 255, 260, 266, 272	234, 261, 266, 273	$4.42,^d 4.30^e$	293	304	303	308	4	32	440
TOTP	229, 264, 271, 279	227, 264 272, 279	$4.32,^d 4.26^e$	299	308	308	309	7	28	301
TPTP	234, 258, 264, 271, 275	233, 263, 269, 275	$4.38,^d 4.28^e$	294	303	305	304	5	35	373

 $[^]a$ $\lambda_{\mathrm{ab,s}}$ —absorption wavelengths in THF. b $\lambda_{\mathrm{ab,a}}$ —absorption wavelengths in the THF-water (5:95, v/v) mixture. c E_{g} band gap (eV) estimated from the onset of absorption spectra. d In THF. e In the THF-water (5:95, v/v) mixture. f λ_{em} —maximum emission wavelength measured at the excitation wavelength of 240 \pm 5 nm. g In THF. h In the THF-water (5:95, v/v) mixture. i In methylcyclohexane rigid glass at 77 K. f In the film spin-casted on quartz substrate, Φ . $\Phi_{PL,S}$ —fluorescence quantum yield in THF estimated using biphenyll $\Phi_{PL} = 18\%$ in cyclohexane at $\Phi_{ex} = 233$ nm) as the standard, Φ . $^{l}\Phi_{PL,a}$ —fluorescence quantum yield in THF-water (5:95, v/v) estimated using biphenyl¹¹ ($\Phi_{PL}=18\%$ in cyclohexane at $\lambda_{ex}=233$ nm) as the standard. m Hydrodynamic radius (d_H) measured in the THF-water (5 : 95, v/v) mixture. soln; solution, aggr.; aggregate.

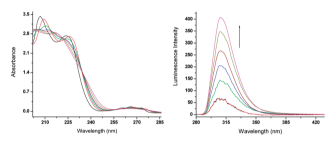


Fig. 1 UV-vis (left) and PL (right) spectra of TPHP (0.1 mM) measured in THF (black) and THF-water mixtures, v/v; 9:1 (red), 8:2 (green), 6:4 (blue), 4:6 (wine), 2:8 (dark yellow), 0.5:9.5 (magenta) under ambient conditions

extended to the long wavelength region in the aqueous-THF mixture. Similar phenomena were noticed from TOTP and TPTP as well, indicating that luminogens aggregated into nanoparticles in the aqueous-THF mixture as the Mie effect of nanoparticles is known to cause such tail levelling-off in the absorption spectra. 5,6,10 From the onset of the absorption profile, the band gap was estimated to be 4.3-4.42 eV, 4.26-4.32 eV and 4.28-4.38 eV, respectively, for TPHP, TOTP and TPTP.

The PL spectrum of **TPHP** $(1 \times 10^{-4} \text{ M})$ in THF and water-THF mixture containing different water fractions from 10 to 95% is shown in Fig. 1. Analogous spectra for TOTP and TPTP are provided in Fig. S1 and S2, ESI.† Notice that all of the compounds were weakly luminescent in THF, but the emission intensity gradually intensified with the increasing water content and reached a maximum value at the water content of 95% in the THF-water mixture. The emission intensities of TPHP, TOTP and TPTP, respectively, in the THF-water (5:95, v/v) mixture are 8-, 4- and 7-fold higher than those found in anhydrous-THF. To have a quantitative picture the $\Phi_{\rm PL}$ was estimated, with respect to biphenyl $\Phi_{\rm PL}=18\%$ in cyclohexane at $\lambda_{ex} = 233$ nm) as the standard, to the amounts of 4%, 7% and 5% (Table 1) in anhydrous-THF and 32%, 28% and 35% (Table 1) in the THF-water (5: 95, v/v) mixture, respectively, for **TPHP**, **TOTP** and **TPTP**. In THF, all TPPs are soluble completely (not a suspension) and as a result they undergo internal conversions (motions/rotations) without any constraint leading to poor luminescence activities.5,6 In aqueous-THF solution, however, TPPs aggregate themselves into lumps (Fig. 1) due to

their hydrophobic nature.^{1,2} As a consequence, non-radiative relaxations that happen by internal conversion are ceased.5-7 It was confirmed by low temperature- and solid state PL measurements as well, vide infra.

The internal conversions of the flexible system are insignificant at low temperature as well as in the condensed phase.2,12 Thus PL emission of TPPs has been investigated at a liquid N₂ temperature (77 K) and in the solid thin film. In both of these experiments, all compounds (Chart 1) show intense emission with $\lambda_{\rm em}$ centred between 303 and 309 nm (Fig. S3, ESI†). So it is reasonable to state that the intense emission (Table 1) detected from TPPs in the aqueous medium is by the effect of aggregation, through self-assembly, which facilitates radiative transitions. 5,6 However, the detailed mechanism for the aggregation induced emission of triarylphosphine is not identified. Doing that may help to enable phosphines to practical applications including in light emitting devices.6b Therefore the structure of TPPs shown in Chart 1 are novel aggregation induced emitters as analogues to well-known tetraphenylethenes6 that show promising emission in the condensed phase than in solutions.

A dynamic light scattering experiment (DLS) was used to investigate the size of the aggregates of TPHP, TOTP and TPTP assembled in the THF-water (5:95, v/v) mixture. A typical unimodal distribution was observed with an average particle size, in terms of the hydrodynamic radius (d_H) , of 440 nm, 301 nm and 373 nm (Fig. 2, S4 and S5, ESI†), respectively, for TPHP, TOTP and TPTP. Indeed particles were found to be relatively stable, as $d_{\rm H}$ values were almost unchanged over a week.

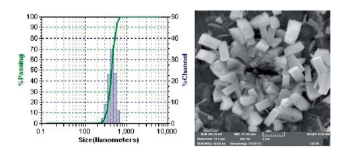


Fig. 2 Typical DLS profile (left) and the SEM image (right) of TPHP in the THF-water mixture (5:95, v/v) measured under ambient conditions.

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The scanning electron microscopy (SEM) was used to explore the morphology of particles of TPHP, a representative example, organized in the THF-water (5:95, v/v) mixture. Interestingly **TPHP** aggregated into brick like structures (Fig. 2 and S6, ESI†) with an average size of 450-500 nm similar to that noticed from DLS analysis.

The appreciable emission of the aggregate state as well as the nucleophilic nature of the phosphorus atom of TPPs encouraged exploring them as probes for sensor applications. Thus, the aqueous solution13 of all compounds (Chart 1) was titrated with various di- and tri-valent (Mn³⁺, Fe³⁺, Bi³⁺) transition metal cations as well as Cl2 and CO2 gases, which are having detrimental effects on humans and the environment.14 Chlorine gas (Cl₂) readily affected the PL emission intensities of TPPs significantly, whereas, neither CO₂ nor metal cations impacted the photophysics of the triarylphosphine luminogens under ambient conditions. Thus, the emission response against the quantity of Cl₂ was investigated intricately to gain insight into the interactions between Cl2 and TPPs (Chart 1) as inhaling or taking Cl₂ above 30 ppm causes eyes irritation, lung damage and aggravates the respiratory system by reaction with cells.14,15

As depicted in Fig. 3, the initial emission of nanoaggregates of TPHP (0.1 mM) in the THF-water mixture (5:95, v/v) weakened gradually with the addition of chlorine-water (ca. 2.3 mM), and attains a constant value (no further enhancement or loss) at 0.41 µM concentration of Cl2. The plots between changes in fluorescence intensity (I_{PL}) at 300 nm versus [Cl₂] are shown in Fig. 3. The limit of detection (LOD) and limit of quantitation (LOQ) were found at 0.015 \pm 0.002 μ M (\pm SD, n=3) and 0.052 \pm 0.006 μ M (\pm SD, n=10), respectively, suggesting the high sensitivity of the luminogen towards Cl₂ detection. A similar behaviour was observed from TPTP as well, c.f. Fig. 3.16 Phosphines are known to react with halogens, alkyl halides, etc., yielding respective phosphonium salts.17 Equally, TPHP and TPTP afforded the corresponding phosphonium salts, Ar₃PCl⁺Cl⁻, with the application of Cl₂ as examined by ³¹P{¹H} NMR spectroscopy analysis (Fig. S7, ESI†). The ³¹P{¹H} NMR spectrum of the parent **TPHP** resonates at -6.2 ppm, and was shifted to +67.4 ppm (deshielded) after reaction with Cl₂. The genuine dissolution of the phosphonium salt of the luminogen (Chart 1) in aqueous medium tone up the non-radiative transitions or weaken the PL emission intensity (Fig. 3).5,6 Thus the TPPs (Chart 1) investigated herein are novel fluorescence chemodosimeters for elemental chlorine.

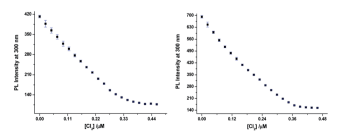


Fig. 3 Changes of fluorescence intensity of TPHP (left)/TPTP (right) at 300 nm as a function of [Cl₂].

In summary, the optical and sensor properties of three TPPs were studied in solution and solid states. Independent of their structure all are fairly luminescent in anhydrous THF, nevertheless, but exhibit moderate to good emission in THF-water mixtures at room temperature. The detailed analysis shows that, in anhydrous solution, all luminogens undergo internal conversion that depopulate the excited state through nonradiative pathways, whereas, in aqueous-THF, they aggregate themselves into nanoparticles of size between 300 and 440 nm, established by DLS and SEM analyses, which facilitate radiative transitions. It was verified by low temperature- and solid state photoluminescence experiments. In a remarkable manner, the luminescence of nanoaggregates of TPPs is sensitive to the elemental chlorine. Since such a character is not readily feasible either from classical luminophores² such as naphthalene, anthracene, pyrene, etc., or with familiar aggregation induced emitters⁵ such as triphenylethylenes, tetraphenylethylenes, siloles and cyano distyrylbenzenes, suggesting an advantage of electron rich phosphorous atoms in the phosphine scaffolds reported here. Therefore the present investigations may serve as the basis for future studies aimed at detection and quantification of Cl2 generated in neutrophils and design of novel functional materials based on phosphines for applications to solid state light emitting devices and molecular electronics.

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