Amine-assisted solubilization of unsubstituted zinc phthalocyananine for film deposition purposes†

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Typical zinc phthalocyanines (ZnPc) exhibit poor solubility in common solvents and, hence, are processed into thin films mostly from the vapor phase. The present work discloses how these limitations can be effectively overcome. Specifically, highly concentrated molecular solutions of unsubstituted ZnPc are prepared by combining a weakly structured ZnPc polymorph with binary liquid systems composed of a π-accepting solvent and a simple nitrogenous base, such as ammonia or tertiary aliphatic amine. The amine-assisted solubilization of ZnPc is rationalized by quantitative analysis of optical spectra and electrostatic potential maps of the dye molecule. A volatile aminoalcohol is proposed in order to rationally modify the habit of ZnPc crystalites and concurrently to produce uniform deposition of the crystallites by drop-casting the dye solutions onto a glass substrate. Finally, a versatile algorithm for wet-processed ZnPc films is declared.

Introduction

Nowadays, metallophthalocyanines (MPC) are applied to a wide range of industrial areas. Their intrinsic feature is the π-stacking propensity, which allows one to fabricate poly-crystalline thin films for molecular electronic devices. In general, an MPC film may be deposited from either vapor phase or solution. The former way is a technological mainstream, allowing one to obtain uniform films over large areas. Alternatively, wet-processing techniques of film fabrication are considered as more promising because of versatility and cost-efficiency.

To prepare an orderly MPC film from solution, a number of serious physicochemical obstacles should be overcome. Poor solubility of typical MPC in common solvents is a key problem being tackled by appending solubilizing substituents to the macroheterocycle. However, such a practice significantly raises a price of MPC and, thus, contracts the field of large-scale applications. Another problem is related to morphology of wet-processed MPC films. When high-grade polycrystalline films are demanded, the size, shape and spacing of nucleating dye islands should be controlled concurrently, while the coffee-ring effect, which inevitably follows the solvent evaporation from colloidal solutions, must be fully suppressed. Besides, a wet-processable dye is expected to adhere to solid substrates, forming a wetting layer as a basis for the vertical film growth. When the latter requirement is not met, the film formation is obstructed regardless of how efficiently a spreading solvent dissolves MPC. No wonder that polycrystalline films of unsubstituted MPC were obtained so far by neither solvent evaporationnor even Langmuir–Blodgett (LB) technique. Recently we examined Langmuir layers of lightly substituted CuPc and revealed a curious behavior of the subphase water that favored the vertical build-up of LB films but badly spoiled the film morphology. So, ammonia, being a usual component of liquid colorant formulations, was proposed as a milder promoter of MPC film growth. Other promising additives seem to be aminoalcohols that were recently applied for the same purpose.

The aforesaid reasons urged us to get down to unsubstituted ZnPc, a low-cost but highly functional dye for solar energetics. This dye is poorly soluble in common organic solvents but can be solubilized, to a certain degree, by nitrogenous bases. In particular, the work seeks to perform a dual task, namely, to develop ZnPc solutions applicable for efficient film deposition and, then, to get mechanistic insights into the amine-assisted solubilization of the dye.

Experimental

Materials

Unsubstituted ZnPc was synthesized according to the slightly modified procedure from phthalonitrile and zinc acetate in the presence of a small amount of N,N-dimethylacetamide. To obtain a dye sample for dissolution experiments, the synthesized ZnPc crystals were converted into the dry sulfate salt followed by regeneration with ammonia in absolute methanol. Organic solvents supplied by various manufactures are listed in Table S1.† They were used as received except for cyclopentanone that was doubly distilled to collect a narrow fraction boiling at
(130.5–131.0) °C. All aliphatic amines were obtained from Sigma-Aldrich and thoroughly distilled just before using.

**Procedures**

Saturation concentrations \(C_s\) of ZnPc were determined by spectrophotometry with pyridine as analytical medium. Light absorbance was measured at the analytical wavelength of 608 nm where the sharp vibrational satellite of the Q-band is located. An error in the \(C_s\) measurements was \(\pm 3\%\). Quantum chemical calculations were performed by the DFT method with the software package Gaussian09. In all calculations, the Becke3–Lee–Yang–Parr correlation functional (B3LYP) with the 6-31G(d,p) basis set was used. Optimized molecular structures were visualized by means of the software package GaussView 5.0.

**Instrumentation**

Wide-angle X-ray scattering data were collected on a Bruker D8 Advance apparatus using the Cu-Kα radiation. Optical spectra were measured on a PC-controlled spectrophotometer Specord M400 (Carl Zeiss). Optical microscopy images of the ZnPc films were taken with an Altami Polar 312 microscope (Altami, Russia).

**Results and discussion**

**Preparation of weakly structured ZnPc samples**

To minimize the effect of crystallinity on dye dissolution, weakly structured ZnPc samples were prepared by converting the synthesized crystals into the dry sulfate salt followed by regeneration with either aqueous or methanolic solutions of ammonia. So, two different ZnPc samples were obtained.

The former sample consists of transparent dark-cyan lamellas, 1–10 μm in length, and shows the optical spectrum similar to that of the initial crystals (Fig. 1, curves 1 and 3). In contrast, the latter sample is a fine amorphous powder that can be easily rubbed into clear green films. Its spectrum exhibits an intense peak at 695 nm (Fig. 1, curve 2), thus drastically differing from spectra of all known MPc polymorphs.

Significant structural differences between the lamellar and powdery samples were revealed by X-ray scattering measurements (Fig. 2).

In the former case, the diffraction curve shows only one peak in the small angle region, which corresponds to the 001 peak of the conventional \(\beta\)-phase. An analogous peak for the powdery sample is shifted to higher angles by ca. 0.5° but, in addition, there is a pair of weak peaks located between 8° and 9° at the position where the 20–1 peak of the \(\beta\)-phase is expected.

The 001 lattice plane of the conventional \(\beta\)-phase is known to be formed by horizontally extended stacks of the tilted MPc molecules, the tilt angle relative to the stacking axis being ca. 45°. At the same time, the 20–1 plane evolves optionally when molecular stacks are allowed to align slightly obliquely toward the 001 plane (see Fig. 4 in ref. 21).

Summarizing the data above allows one to classify the lamellar sample as a low-dimensional kind of the \(\beta\)-phase, while the powdery sample may be considered as an individual polymorph, in which the ZnPc molecules are tilted relative to the stacking axis a bit stronger than in the \(\beta\)-phase. Earlier a similar structure was observed in LB films of tert-butyl-substituted CuPc and termed a “o-phase” (i.e., overshifted).

**Dissolution and deposition of ZnPc**

Among non-specific organic solvents suitable for dissolution of unsubstituted ZnPc, the most efficient one is tetrahydrofuran (THF). However, drop-casting the saturated ZnPc solution in THF allows one to produce only needle-shaped microcrystals arranged in a coffee-ring pattern. So, as a matter of plain logic, regard was given to binary liquid systems composed of a volatile solvent and an additive performing at least two functions, namely, the solubilization of dye materials and the levelling of solid deposits.

As a first practical step, the \(C_s\) parameter for the powdery ZnPc sample in neat solvents was measured. Because no symptoms of dye dissolution in methanol (Met), acetone (Ace), chlorobenzene (CIB), and benzonitrile were visually detected, \(C_s\) in these solvents was roughly estimated as \(5 \times 10^{-7}\) mol kg\(^{-1}\) at best. In turn, \(C_s\) in THF was reliably measured as \(1.6 \times 10^{-4}\) mol.

**Fig. 1** Normalized optical spectra of the solid ZnPc samples: (1) lamellar; (2) powdery; (3) needle-shaped (as-synthesized).

**Fig. 2** X-Ray diffraction patterns of the lamellar (1) and powdery (2) ZnPc samples. Main peaks of the conventional \(\beta\)-phase are shown by sticks with Miller indices.
kg$^{-1}$, *i.e.*, nearly five times as low as $C_S$ reported in ref. 5. Thus, the dye sample under investigation contained no solubilizing impurities that could distort the solution behavior of ZnPc.

When ammonia was bubbled into suspensions of ZnPc in Ace, cyclopentanone (CyP), and ClB, the liquids instantly became deep blue. Optical spectra of the obtained solutions (Fig. 3) exhibit both well resolved bands and zero baselines, clearly indicating that these solutions are truly molecular and, hence, suitable for controlled crystallization of ZnPc. Numerical data on solubility of ZnPc in ammonia-saturated solvents are given in Table S2.$^\dagger$

The $C_S$ parameter was greatly increased by ammonia in Ace, CyP, and ClB, but slightly in THF. No effects in Met and other fatty alcohols were observed. Noteworthy, ammonia salts out ZnPc from solution in triethylamine (TEA), indicating that two amines compete for the dye molecule and, hence, interact with ZnPc by one and the same solvation mechanism.

Then, a number of volatile tertiary amines were tested as solubilizing additives (Fig. 4).

Because of different solubility of ZnPc in neat amines, the solubilizing effect is expressed here as logarithm of the ratio between $C_S$ values measured empirically and calculated in accordance with the additive principle. This effect, being equal to zero when no solubilization occurs, is expected to reach at least three units for really processable solutions. Fig. 5, exemplifying the solubilization of ZnPc by triethylamine (TEA), indicates that CyP deserves most attention as a spreading solvent for film deposition.

Notably, the most efficient solubilization occurs in low-volatile aromatics, such as acetophenone and benzonitrile, allowing one to prepare molecular solutions with $C_S > 1.0 \times 10^{-2}$ mol kg$^{-1}$.

In order to select amines applicable for film deposition, saturated solutions of ZnPc in the binary CyP-based systems were tested by placing one microdroplet onto hydrophilic glass followed by visual inspection of a solid deposit. The CyP–ammonia system was discarded because of high reactivity of ammonia towards ketones,$^{24}$ but promising results were obtained with some aminoalcohols.

For instance, when 3-dimethylamino-1-propanol (DMAP) was used as an additive, the dye deposit consisted of non-agglomerated prismatic crystallites with a low aspect ratio of ca. 1.5 and only a few of rod-like ones (Fig. 6). Ubiquitous needle-shaped particles, which had been detrimental for orderly film formation,$^{25}$ were fully lacking. Even severe annealing the crystallites did not shatter the prismatic habit. Moreover, no symptoms of the coffee-ring effect were observed. An overall uniformity of this deposit allows considering it as a primitive solid film grown up in the Vollmer–Weber (*i.e.*, island) mode.$^{26}$ Curiously, the needle-shaped habit of ZnPc crystallites was not eliminated by using $N,N$-diethylylaminoethanol, *i.e.*, aminoalcohol closely similar to DMAP in structure and volatility (see Fig. S3 and S4$^\dagger$).

**Scientific insight into the solubilization phenomenon**

Optical spectroscopy is a handy tool to study interactions of ZnPc with organic bases.$^{14}$ A key feature of the ZnPc spectrum in solution is a complex profile of the Soret band that spans the near UV region. This band has an intricate structure composed

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Fig. 3  Optical spectra of unsubstituted ZnPc in ammonia-saturated ChB (1) and Ace (2). The spectra are shown as measured. Experimental conditions: fresh ZnPc-saturated solutions; optical path, 0.1 cm; slit width, 1 nm.

Fig. 4  Saturation concentrations ($C_S$) of ZnPc in the binary CyP–amine systems at 20 °C. Dots show $C_S$ for the powdery sample (the o-phase); squares show $C_S$ for the lamellar sample (the low-dimensional β-phase). Abbreviations: TEA, triethylamine; NMM, $N$-methylmorpholine; DEAE, $N,N$-diethylylaminoethanol.

Fig. 5  Solubilization of ZnPc by TEA in organic solvents at 20 °C. The solubilizing effect is determined as a logarithmic ratio between the measured and calculated $C_S$ values at the solvent/TEA ratio of 6 : 1, v/v (0.08–0.10 mole fraction of TEA).
of two spectral envelopes (B1 and B2) and an individual band (B3) located at ca. 305 nm. After a basic molecule is axially coordinated to ZnPc, the band profile suffers changes and becomes available for reliable deconvolution. In this work, two kinds of the band profile were observed, asymmetrical and split (Fig. 7, top). The asymmetrical profile, recorded at low concentrations of organic bases, consists of three peaks belonging to the B1 envelope and two peaks being a part of the B2 one. When neat amine, such as TEA, is used as a solvent, the Soret band shows clear splitting at ca. 340 nm because an extra peak pops up in the B2 envelope (Fig. 7, bottom). The dual mode of the ZnPc–TEA interaction indicates the successive formation of two axial 1:1 and 1:2 complexes, thus being in agreement with the thermogravimetric data on solid ZnPc–amine crystals.

To rationalize the Soret band splitting, the orbital energy diagram for ZnPc is to be considered. According to the quantum chemical study, the B2 envelope consists of three rather than two \( \pi - \pi^* \) peaks, but one of them is rather weak (to be exact, 8% of the whole B2 envelope) because of meager population of the binding 5a2u orbital localized on the peripheral benzene rings (see Table 3 in ref. 28). Our analysis indicates that the intensity of the extra peak is negligible for the 1:1 complex and amounts to 15% of the B2 envelope for the 1:2 one. Thus, the asymmetric profile in Fig. 7 reflects the primary donation of the electron density from TEA to the macroheterocycle (further MHC), whereas the band splitting indicates the subsequent redistribution of the electron density onto the molecular periphery.

Finally, a series of DFT calculations was performed to complete the discussion above: the ZnPc molecule was optimized and thereupon the interactions of ZnPc with small molecules were examined (Tables S6 and S7†). The obtained results are in general agreement with earlier ones. Besides, the heat of interaction between typical Lewis acids and amines in inert media is known to range from 50 to 130 kJ mol\(^{-1}\) and the calculated ZnPc–ammonia interaction energy fits smoothly in this interval. At the same time, a serious discrepancy between the data acquired by the DFT and semi-empirical methods should be noted (cf. Fig. 8 below and Fig. 2 in ref. 31).

In order to understand how the electron density in the MHC is affected by small organic molecules, three-dimensional maps of molecular electrostatic potential (MEP\(^\text{32}\)) for ZnPc were generated. Fig. 9 shows a key result.

Upon coordination of one ammonia molecule, the MHC undergoes the dome-like distortion accompanied by overall levelling of the MEP surface: the central MEP peak disappears and peripheral cavities grow shallow. Another evident consequence of the axial coordination is electronic disparity of the upper and lower planes of the MHC.

Any MPC molecule may be considered as a combination of three kinds of particular molecular fragments: meso-nitrogen.
Conclusions

This contribution outlines a novel way to easily manipulate unsubstituted ZnPc for film deposition purposes. A key feature of the proposed approach is the use of a multifunctional amine additive, which is capable of solubilizing ZnPc in volatile solvents, modifying the habit of deposited dye crystallites, and producing uniform deposition of the crystallites on a solid substrate. The amine-assisted solubilization of ZnPc is explained by complex redistribution of the π-electron density triggered by a single axial ligand.
Conflicts of interest

There are no conflicts to declare.

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