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Probing the supramolecular features via $\pi-\pi$ interaction of a di-iminopyrene-di-benzo-18-crown-6-ether compound: experimental and theoretical study†

Adina Coroaba, pa Dragos-Lucian Isac, *a Cristina Al-Matarneh, a Tudor Vasiliu, a Sorin-Alexandru Ibanescu, a Radu Zonda, a Rodinel Ardeleanu, a Andrei Neamtu, ab Daniel Timpu, Alina Nicolescu, Francesca Mocci, at Stelian S. Maier, ae Aatto Laaksonen, af Marc Jean Médard Abadie ag and Mariana Pinteala

A novel DPyDB-C=N-18C6 compound was synthesised by linking a pyrene moiety to each phenyl group of dibenzo-18-crown-6-ether, the crown ether, through -HC=N- bonds and characterized by FTIR, 1H-NMR, ¹³C-NMR, TGA, and DSC techniques. The quantitative ¹³C-NMR analysis revealed the presence of two position isomers. The electronic structure of the DPyDB-C=N-18C6 molecule was characterized by UV-vis and fluorescence spectroscopies in four solvents with different polarities to observe particular behavior of isomers, as well as to demonstrate a possible non-bonding chemical association (such as ground- and excited-state associations, namely, to probe if there were forming dimers/excimers). The interpretation of the electronic structure was realized through QM calculations. The TD-CAM-B3LYP functional, at the 6-311+G(d,p) basis set, indicated the presence of predominant $\pi \to \pi^*$ and mixed π $\rightarrow \pi^* + n \rightarrow \pi^*$ transitions, in line with the UV-vis experimental data. Even though DPyDB-C=N-18C6 computational studies revealed a π -extended conjugation effect with predominantly $\pi \to \pi^*$ transitions, thorough fluorescence analysis was observed a weak emission, as an effect of PET and ACQ. In particular, the WAXD analysis of powder and thin films obtained from n-hexane, 1,2-dichloroethane, and ethanol indicated an amorphous organization, whereas from toluene a smectic ordering was obtained. These results were correlated with MD simulation, and it was observed that the molecular geometry of DPyDB-C=N-18C6 molecule played a defining role in the pyrene stacking arrangement.

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^aCentre of Advanced Research in Bionanoconjugates and Biopolymers, "Petru Poni" Institute of Macromolecular Chemistry, 41A Grigore Ghica Voda Alley, 700487 Iasi, Romania. E-mail: pinteala@icmpp.ro; isac.dragos@icmpp.ro

1 Introduction

Supramolecular chemistry has seen significant development over last decades, earning a lead role in the scientific research. Whilst in classical chemistry the focus is mainly on covalent bonds, in supramolecular chemistry the spotlight is on reversible interactions. The concept of self-assembly is defined by the spontaneous formation or development of some systems without exterior intervention, other than environmental conditions. Controlling this phenomenon is of great interest in the synthesis of materials with complex properties at the nano- and micro-scale. The control over self-assembly process has proved to be even more essential in developing supramolecular structures with biological functions, in understanding natural systems and for their synthetic implementation.1-3 The self-assembly of molecules is mainly based on noncovalent interactions between different parts of the molecules. This involves attraction and repulsive forces such as ionion interactions, van der Waals interactions, dipole-dipole or iondipole interactions, coordinative bonds, hydrogen bonds, and π π stacking.4-8

^bCenter of Fundamental Research and Experimental Development in Translational Medicine, Regional Institute of Oncology, General Henri Mathias Berthelot St, 700483 Iasi, Romania

Physics of Polymers and Polymeric Materials, "Petru Poni" Institute of Macromolecular Chemistry, 41A Grigore Ghica Voda Alley, 700487 Iasi, Romania

^dDepartment of Chemical and Geological Sciences, University of Cagliari, I-09042 Monserrato, Italy

[&]quot;Polymers Research Center, "Gheorghe Asachi" Technical University of Iasi, 73 Mangeron Blvd., 700050 Iasi, Romania

Department of Materials and Environmental Chemistry, Division of Physical Chemistry, Arrhenius Laboratory, Stockholm University, SE-106 91 Stockholm, Sweden *Faculy of Science, University of Montpellier, Place E. Bataillon, 34095 Montpellier, France

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Paper

The functionality of a supramolecular structure is endowed by some of the chemical groups belonging to the constituent molecules, which are capable of selectively interacting with various entities in the proximal environment. It is already known that crown ethers selectively interact with alkaline ions, based on the correlation between the size of the crown ether cycle and the ionic radius. Paper Exploiting this uniqueness, crown ether derivatives are often used as sensors, in it is uniqueness, crown or carriers for metallic ions. For some of the mentioned functions, it is useful to integrate functionalized crown ethers in supramolecular structures.

A particular way to create supramolecular structures is to use π -conjugated molecules. Studying charge transfer interactions, Koshkakaryan et al.22 developed a stable supramolecular assembly containing intercalated crown-ethers and naphthalene diimide-based molecules in 1:2 ratio, when naphthalene groups were situated in- and out-side of the crown ether ring. However, the use of π -conjugated molecules as binders represents an attractive way to build supramolecular structures. For this purpose, pyrene is a good choice, thanks to its propensity to self-aggregate by π -stacking, which can be visualized resorting to fluorescence emission spectroscopy. In this respect, pyrene excimers are often used as fluorescence markers for studying the systems' microheterogeneity. Pinteala et al.23 investigated the conformational changes and the aggregation hydrophobic-hydrophilic copolymers, using pyrene as a dye. They recorded the spectra of pyrene molecules captured in the space nearby copolymers and evaluated the local hydrophobicity of the microenvironment by evaluating the intensity ratio between the first and the third vibronic band in pyrene fluorescence emission spectra. Szajdzinska-Pietek et al.24 used pyrene and a pyrene derivate, 1,3-bis(1-pyrenyl)propane, to continue the study of the same copolymer.²³ According to them, the self-assembly of the chains leads to the formation of hydrophobic domains, determining an increase in the local concentration of fluorescent probes due to the π -stacking, a phenomenon revealed by excimer formation at a longer wavelengths (in the range of 420-550 nm). Moreover, they observed that the dyeing effect of the pyrene derivative is more sensitive and stronger than that of pyrene. Because of its characteristic photophysical behavior such as its prominent and typical absorption bands and fluorescence emission, 25,26 pyrene was involved in numerous supramolecular studies. The first two excited states of pyrene compounds, 1La and 1Lb, $\pi \rightarrow$ π^* , corresponding to $S_0 \to S_1$ and $S_0 \to S_2$ excitations, proved to have the main role in the photophysical processes involving such molecular systems.25,27-30 The band's intensity variation (including their appearance or disappearance), together with the wavelength shifts in UV-vis and fluorescence spectra varies from one pyrene derivative to another. The studies indicate that pyrene compounds are able to form dimers (by the weak association in the ground state), excimers (by the strong association in the excited states), and/or π - π stacking aggregates.^{25,29,31-37} Computational approaches such as density functional theory (DFT), time-dependent density functional theory (TD-DFT), and molecular dynamics (MD) simulations facilitate the understanding of the nature of electronic structure in the ground and

excited states (presence of $\pi \to \pi^*$ transitions described by molecular orbitals), and of the supramolecular interactions (π - π stacking associations) of pyrene derivatives. The presence of specific functional groups such as imine bond (-HC=N-), phenyl ring, or crown ethers in the moiety of pyrene units is able to induce new electron transitions (such as $n \to \pi^*$ ones³⁸), or different association interactions.

The published studies that address the π - π interaction in the pyrene-crown ether systems mainly refer to the changes in the molecular geometry of the assembly as a result of the binding of two pyrene units, in different chemical environments. Jarolímová *et al.*³⁹ studied crown ethers of different sizes, functionalized with pyreneamide derivatives, as potassium ion-selective nanosensors. The complexation of the cation with the crown ether ring determines the molecule unfolding, and consequently, the disappearance of the excimer emission in an on-off manner, which makes the compound usable as a chemosensor.³⁹

The present work reports the synthesis of di-iminopyrene-dibenzo-18-crown-6-ether (named as **DPyDB-C=N-18C6** in the present paper), a novel compound including crown ether and pyrene moieties linked by –HC=N- bonds (Fig. 1). The aim of the present study was to prove, based on photophysical properties measurements, the formation of a potential supramolecular arrangement mediated by inter- and intramolecular interactions between the **DPyDB-C=N-18C6** molecules. The experimental results were correlated with the molecular dynamics and with quantum mechanics calculations. For photophysical investigations, solvents with different polarities (*n*-hexane, toluene, 1,2-dichloroethane, and ethanol) were chosen.

2 Experimental section

2.1 Reagents and solutions

All reagents and solvents were purchased from Sigma-Aldrich (Merck KGaA, Darmstadt, Germany) and used without further purification.

2.2 Instrumentation and measurements

Analytical thin-layer chromatography was performed with commercial silica gel plates 60 F254 (Merck Darmstadt, Germany) and visualized with UV light ($\lambda_{max} = 254$ or 365 nm). The nuclear magnetic resonance (NMR) spectra were recorded using a Bruker Avance Neo 400 MHz spectrometer (Bruker, Germany). Chemical shifts were reported in delta (δ) units, part per million (ppm). The following abbreviations were used to designate chemical shift multiplicities: s = singlet, d = doublet, t = singlet

Fig. 1 Structural formula of di-iminopyrene-di-benzo-18-crown-6-ether.

triplet, q = quartet, m = multiplet, as = apparent singlet. The Fourier-transform infrared (FTIR) spectra were recorded using a Bruker VERTEX 70 FT-IR spectrometer (Bruker, Germany) at ambient temperature, in potassium bromide (KBr) pellets. The thermogravimetric analysis (TGA) was performed using STA 449F1 Jupiter equipment (Netzsch, Germany). The measurements were made in the temperature range 20-700 °C under nitrogen flow (50 mL min⁻¹) at a heating rate of 10 °C min⁻¹. Differential scanning calorimetry (DSC) measurements were conducted using a DSC 200 F3 Maia device (Netzsch, Germany). A mass of 15 mg of each sample was heated in a pierced and sealed aluminum crucible at a heating rate of 10 °C min⁻¹ under a nitrogen stream at a flow rate of 50 mL min⁻¹. The temperature against heat flow was recorded. The device was calibrated using indium according to standard procedures. Wide-angle X-ray diffraction (WAXD) analyses were performed using a D8 ADVANCE diffractometer (Bruker AXS, Germany) with Cu-K α radiation ($\lambda = 0.1541$ nm), a parallel beam with a Göbel mirror, and a dynamic scintillation detector. All the samples were investigated for a 2θ diffraction angle ranging from 0.8° to 60°, in a step scan mode of 2 s per step, with a step size of 0.02°, at 40 kV and 35 mA, at room temperature. The calculation of 2θ values, the interplanar distance (d), and peak heights were carried out using the Bruker "DIFFRAC-PLUS Evaluation - EVA" software. The Bruker "TOPAS" software was used for crystallite size data processing. UV-vis measurements were performed using a Lambda 35 device (PerkinElmer, USA). The absorption spectra were recorded in the 200-700 nm range for identical sample volumes (3 mL) with the following parameters: slit width 1 nm, scan speed 480 nm min⁻¹, and 1 nm data interval. The spectra of the samples were recorded at room temperature using 1 cm path length quartz cuvettes. Fluorescence measurements were carried out using a FluoroMax-4 spectrophotometer (Horiba, Japan). The emission spectra were recorded using an excitation wavelength of 360 nm. The concentrations of the solution obtained were determined by Quartz Crystal Microbalance with Dissipation (QCM-D) using a QSense Pro instrument (Biolin Scientific, Sweden). QSX 301 Gold sensors, acquired from Scientific Biolin Scientific AB, Sweden, were cleaned by placing them in a 5:1:1 mixture of Milli-Q water, ammonia (25%), and hydrogen peroxide (30%) at 75 °C for 5 minutes. The sensors were then rinsed with Milli-Q water and dried with nitrogen gas. The resonance frequency of a clean QCM sensor (f_0) was measured in air. Then, on gold QCM chips, a known volume of the solutions (100 µL for ethanol and toluene and 150 µL for n-hexane and 1,2-dichloroethane) with unknown concentrations were added drop by drop and dried under a nitrogen stream. The sensors were mounted again in the instrument and the resonance frequency after deposition (f_1) was measured also in the air atmosphere.

2.3 Synthesis of di-iminopyrene-dibenzo-18-crown-6-ether (DPvDB-C=N-18C6)

To a solution of diaminodibenzo-18-crown-6-ether (200 mg, 0.51 mmol) in ethanol (20 mL), acetic acid (200 μ L, 2.80 mmol,

4 equiv.) and 1-pyrenecarboxaldehyde (260 mg, 1.15 mmol, 2.1 equiv.) were added. The resulting mixture was stirred at room temperature for 24 h. The crude product was filtrated and then purified by centrifugation for separation from dichloromethane and ethyl ether. Yellow solid, yield 80%, IR: 3035, 2922, 2868, 1909, 1508, 1276, 1263, 1130, 1055 cm $^{-1}$. $^1\text{H-NMR}$ (400 MHz, CDCl $_3$): $\delta=4.14$ (as, 8H), 4.35 (dd, 8H), 7.02 (d, 4H), 7.08 (s, 2H), 8.08 (d, 2H), 8.15 (d, 2H), 8.18 (d, 2H), 8.28 (m, 8H), 8.77 (d, 2H), 9.05 (d, 2H), 9.54 (s, 2H). $^{13}\text{C-NMR}$ (100 MHz, DMSO): $\delta=69.5$, 106.6, 113.1, 114.6, 123.1, 123.7, 124.3, 124.6, 125.6, 126.4, 126.9, 127.1, 127.9, 129.2, 130.3, 131.2, 131.4, 133.1, 145.3, 147.5, 149.0, 157.0.

2.4 Molecular dynamics (MD) simulations

The behavior of the molecular system in different solvents and the ability of the target compound to establish intermolecular interactions between di-iminopyrene-dibenzo-18-crown-6-ether and polar solvents (using ethanol as an explicit model) were investigated by molecular dynamics (MD) simulations. All MD simulations were done using the GROMACS 2018 program.⁴⁰ The initial structure of the molecules was built using the AVO-GADRO software.41 The molecules were parameterized using the following protocol. The GAUSSIAN 16 software⁴² and RED-Tools⁴³ software were used to calculate the partial atomic charge distribution and to optimize the molecular geometry of the compounds. The method used for calculations was HF/6-31G* with the Connoly surface algorithm. The force field used to describe the system was AMBER force field. The parameters for bonds, angles, and dihedrals were calculated using the Ambertools18 software44 and the GAFF parameters. All simulations were carried out using the Beskow supercomputer from the KTH Royal Institute of Technology, PDC Center for highperformance computing.

The details of the simulations are presented in Table 1. The simulations were performed with the NPT ensemble at $T=293~\rm K$ and P=1 atm under isotropic conditions. The temperature was controlled using the Noose–Hoover thermostat and the pressure was controlled using the Parinello–Rahman barostat. The equations of motions were integrated with a time step of 2.5 fs. Before starting each simulation, energy minimization was performed using the steepest descent algorithm. Periodic boundary conditions (PBCs) were used for all simulations. The Van der Waals interactions were selected by the particle mesh Ewald (PME) method with a cut-off of 1 nm.

2.5 Quantum mechanics calculations

In this work, the quantum mechanics calculations were performed using the Gaussian (G16) package⁴² by density functional theory (DFT) and time-dependent density functional theory (TD-DFT) in order to investigate both the ground and excited states. The optimization of the calculations using the DFT method was employed with the PBE0/6-311+G(d,p) level of theory. The equilibrium geometry has been first optimized to the ground-state level. A conformational search was performed to identify if the equilibrium geometry corresponds to the local minimum of the potential energy surface. Then, the Hessian

40 ns

Ions system

Number of crown ether molecules/
System

Number of crown ether molecules/
Number of solvent molecules

Number of solvent molecules

Simulation box size (nm^3) duration

DPyDB-C=N-18C6 system

80 molecules (40 molecules of 23 278 ethanol 13.363 800 ns

Water

Table 1 Details of the molecular dynamics simulations

frequencies have been computed for the equilibrium geometry and no eigenvalues have been obtained, indicating that the molecular geometry structure corresponds to the local minimum on the potential energy surface.

each isomer)/0/0/0/0 10 molecules/10/10/10/40

Excited states and theoretical absorption spectra were computed with TD-CAM-B3LYP functional and 6-311+G(d,p) basis sets. To obtain a comparison between theoretical and experimental results, as well as a good estimation of the first electronic transitions, ten states were included in the TD-DFT calculations. Thus, the TD-DFT calculations at three different levels of theory, namely, TD-CAM-B3LYP/6-311+G(d,p), TD-PBE0/6-311+G(d,p), and TD-B3LYP/6-311+G(d,p) have been further used for comparison. All TD-DFT calculations were performed in gas phase and solvent environments with different polarities: n-hexane ($\varepsilon = 1.9$), toluene ($\varepsilon = 2.34$), 1,2-dichloroethane ($\varepsilon = 10.1$), and ethanol ($\varepsilon = 24.3$) in order to simulate the experimental UV-vis absorption spectra. The polarized continuum model (PCM)⁴⁵ was used to mimic the solvent's environment.

Both density functionals, PBE0 (ref. 46 and 47) and CAM-B3LYP,48 were used in the present theoretical calculations to explore the electronic structure of di-iminopyrene-dibenzo-18crown-6-ether. Moreover, PBE0, TD-BE0, and TD-CAM-B3LYP functionals were used in the theoretical calculations to explore the electronic structure of organic49,50 and inorganic51,52 compounds providing reliable geometries and good estimated values of the transition energies. Likewise, the TD-B3LYP/6-311+G(d,p) method was also included to study the theoretical electronic absorption spectra. This supplementary level of theory was employed because this method gives good results regarding the optoelectronic properties of π -conjugated compounds⁵³ and of 3-methoxy-2-((pyren-2yl-imino)methyl) phenol.38 Quantum mechanics calculations were carried out on the Tetralith supercomputer from the KTH Royal Institute of Technology, PDC Center for High-Performance Computing.

3 Results and discussions

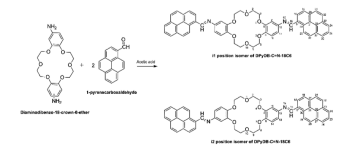
3.1. Synthesis and physicochemical characterization of diminopyrene-dibenzo-18-crown-6-ether

3.1.1 Synthesis of DPyDB-C=N-18C6. Di-iminopyrene-dibenzo-18-crown-6-ether was prepared by the direct reaction between the amino groups of diaminodibenzo-18-crown-6-ether and the aldehyde group of 1-pyrenecarboxaldehyde, in an acidic medium, with the formation of imine bonds between the two precursors, as presented in Scheme 1. Diaminodibenzo-18-crown-6-ether was synthesized as described by Ardeleanu

et al.,⁵⁴ as a mixture of two position isomers (noted as **i1** and **i2**) in regard to the amino position on the phenyl ring, and was used without further separation.

3.1.2 Fourier-transform infrared (FTIR) spectroscopy. The FTIR spectra of **DPyDB-C=N-18C6** (Fig. S1†) showed absorption bands between 2922–2868 and 1130–1055 cm⁻¹, characteristic for O-C-C-O and C-O-C ether bonds from the crown ether, and two intense absorption bands at 1610–1583 cm⁻¹, characteristic of the in-phase stretching vibrations of the newly formed -HC=N-imine bond. It was also noticed that the disappearance of the intense absorption bands at 2712, 1797, and 1678 cm⁻¹, characteristic of 1-pyrenecarboxaldehyde, is one more proof that the reaction took place.

3.1.3 Nuclear magnetic resonance (NMR) spectroscopy. In the ¹H-NMR spectrum of **DPyDB-C=N-18C6** (Fig. S2†), the signals at $\delta = 4.14$ –4.35 ppm have been assigned to aliphatic atoms from the crown ether structure. Protons H18 give a signal at $\delta = 9.05$ ppm, and protons H19 at $\delta = 8.77$ ppm, under the influence of the electron-withdrawing neighbors. The most deshelled proton is the one corresponding to the imine, H16, at 9.54 ppm. The disappearance of the characteristic protons for the amino group in the crown ether (4.68 ppm) and of the aldehyde group in 1-pyrenecarboxaldehyde (10.7 ppm) further confirm the structure of DPyDB-C=N-18C6. 13C-NMR spectra (Fig. S3†) showed signals in accordance with the proposed structure, with emphasis on the two different signals for the aliphatic zone corresponding to the atoms from the crown ether structure at $\delta = 69.5$ ppm, and a signal at $\delta = 157.0$ ppm corresponding to the C16 imine carbon. The quantitative ¹³C-NMR study (Fig. S4†) for diaminodibenzo-18-crown-6-ether revealed an approximatively 1:1 ratio between the two isomers, and it was assumed that the final compound, DPyDB-C=N-18C6, has the same isomer ratio as the starting compound.



Scheme 1 Synthetic of the preparation of di-iminopyrene-dibenzo-18-crown-6-ether.

3.1.4 Thermal analysis. Fig. S5† shows the thermogravimetric (TGA) and differential thermogravimetric (DTG) curves of DPyDB-C=N-18C6. Two thermal decomposition stages with a maximum weight loss in the range of 340-568 °C are present. The first stage was assigned to the loss of solvent molecules (ethanol and water) (1.07; 1.46 wt%). The second step, with initial temperatures between 340 and 420 °C, corresponds to the loss of 29.4% weight (crown ether moiety and 13.8% corresponding to the benzo amino structure at 420-568 °C). The more stable remaining structural entities thermally decompose in the domain 568-700 °C, leaving a residue that corresponds to the pyrene moiety.

Fig. S6† reveals that the differential scanning calorimetry (DSC) curves corresponding to the heating and cooling runs of the studied sample. The heating curve presented a peak at 185 °C, which is rather a heat anomaly or expresses a complex phase transition, 55 before sample degradation. No defined melting point was registered on the cooling curve, suggesting the amorphous character of the product.

3.2 The ability of supramolecular arrangement investigated by wide-angle X-ray diffraction (WAXD) analysis and simulated using molecular dynamics (MD)

3.2.1 WAXD analysis. To better understand the structure of DPyDB-C=N-18C6, WAXD analysis was performed on the product powder, revealing an amorphous material. Moreover, WAXD was used to determine whether the solvent can influence the supramolecular arrangement of the compound. In this respect, solvents with different polarities were used: n-hexane (ε = 1.9), toluene (ε = 2.4), 1,2-dichloroethane (ε = 10.1), and ethanol ($\varepsilon = 24.3$). Four highly concentrated solutions of **DPyDB-C=N-18C6** were prepared in each solvent and were left to rest for 24 hours. The resulting precipitates were collected, dropwise deposited on silicon wafer substrates, and left to dry by slow evaporation. The obtained films were then subjected to WAXD analysis. The results revealed that n-hexane, 1,2-dichloroethane, and ethanol solvents do not influence the supramolecular arrangement of the compound, the same amorphous character being put in evidence. However, the film obtained from toluene showed a crystalline character, as presented in Fig. 2 and Table 2. The broad peaks obtained at 2θ values of 12.4° and 22.8° belong to the silicon wafer substrate. The full width at half maximum (FWHM) of the peaks and the crystallite sizes were calculated using the Debye-Scherrer equation using the highest peak P1 as the reference; the following values were respectively obtained: 0.127 for FWHM and 69.74 nm for crystallite size. The scattering vector ratio (Table 2, q/reciprocal lattice vector) of approximatively 1:2:3:4 indicates a smectic ordering.⁵⁶ Based on this fact, it can be assumed that π - π , multipole, induced-multipole, and dispersion force interactions between toluene molecules and pyrene units57 lead to a potential supramolecular arrangement of DPyDB-C=N-18C6.

3.2.2 Molecular dynamics (MD) simulations. In order to investigate the aggregation capability of di-iminopyrene-dibenzo-18-crown-6-ether, molecular dynamic simulations of the aggregation process in ethanol were performed. Taking into

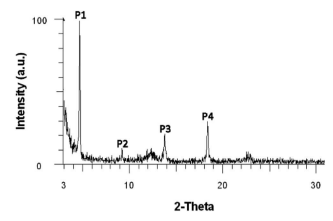


Fig. 2 Wide-angle X-ray diffraction (WAXD) diffractogram of the DPyDB-C=N-18C6 film obtained from the toluene solution.

consideration the quantitative ¹³C-NMR results, both position isomers were included in the MD simulations. Fig. 3 shows the initial conformation of the simulations.

The visual analysis of the simulation trajectory showed that the molecules started to aggregate as soon as the simulation was initiated. The fast aggregation was due to the high concentration simulated, which was much higher than the experimental one. The reason for the need to work under high-concentration conditions is that the simulation of the experimental concentration is impossible in a feasible amount of time, due to the high number of solvent molecules, which must be considered (108 solvent molecules). Therefore, to put in evidence the intermolecular interactions that govern the aggregation process, a system that would aggregate and reach equilibrium in a reasonable amount of time must be chosen.

Under the simulation conditions, **DPyDB-C=N-18C6** formed a single aggregate that includes 80 molecules (40 molecules of each position isomer). Fig. 4 depicts the final structure of the aggregate of **DPyDB-C=N-18C6**, with a detail of the interaction between molecules.

Visual analysis of the simulated aggregate shows an amorphous structure (Fig. 4A), in good agreement with the WAXD findings. Despite the high disorder degree of the aggregate, it can be seen that pyrene units are still capable of forming intermolecular π – π stacks between three or four near-by molecules. The results of MD computation indicated the existence of an almost isoenergetic structure, as a slipped-parallel aggregate, with interaction distances of about 3.45 and 3.28 Å. The ordered pyrene units are developed only at one side of the

Table 2 Angular positions of the peaks, interplanar distances, and scattering vector (q) data obtained from WAXD analysis

Peak	Angle (2-theta°)	d value (Å)	$q\ (\mathring{ m A}^{-1})$	q ratio
P1	4.626	19.09	0.328430	1.00
P2	9.191	9.61	0.652008	1.99
P3	13.753	6.43	0.974339	2.97
P4	18.365	4.83	1.298631	3.95

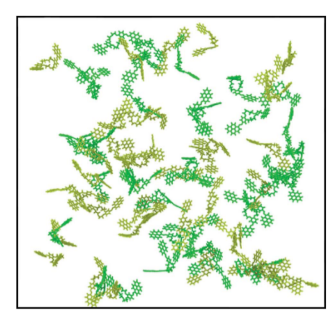


Fig. 3 Starting structures after the energy minimization for the DPyDB-C=N-18C6 compound. In yellowish green is the i1 position isomer and in green is the i2 position isomer. Ethanol molecules are hidden, for clarity.

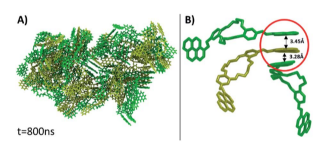


Fig. 4 (A) Final structure of the aggregate of DPyDB-C=N-18C6 molecules, resulting after 800 ns of simulation. (B) Detail of a one-sided stack of pyrene (circled in red). Ethanol molecules are hidden, for clarity.

molecule, the pyrene moieties on the other side being either not-aggregated or involved in stacks with pyrene groups belonging to totally different molecules (Fig. 4B). Due to the high degree of disorder, there was no way to determine whether the presence of two isomers has any effect on the final structure of the aggregate.

The inability of the compound to align its both pyrene cycles with the two pyrenes of the same partner molecule is due to the dihedral angle in –C–HC=N–C–, which includes the imine –HC=N– bond. It induces a twist between the planes of pyrenes and phenyl cycles respectively, which is propagated towards the plane of the second pyrene moiety. Another motivation of the inability to form more systematic intermolecular π – π stacks is the presence of a C–H bond (held by a –HC=N– unit) in the vicinity of a lone pair (belonging to the nitrogen atom of the same unit), which may induce a hindrance repulsion between neighboring molecules. In order to confirm the existence of the

twisted dihedral angle (-C-HC=N-C-), quantum mechanics (QM) molecular geometry optimization, presented later in the paper (see Section 3.3.2), was performed. The computational results also evidenced that hydrogen interactions were formed between nitrogen atoms of imine bonds (-HC=N-) and the -O-H moiety of ethanol. A similar interaction was also observed between the -OH group of ethanol with the oxygen atom of the crown ether segment in **DPyDB-C=N-18C6** molecules. The hydroxyl group of ethanol plays the role of hydrogen bond donor, while the nitrogen atoms belonging to imine groups, together with the oxygen atoms held by the crown ether, act as hydrogen bond acceptors.

3.3. Photophysical properties: experimental and theoretical studies

3.3.1 Experimental studies

3.3.1.1 UV-vis absorption spectra. To demonstrate the photophysical properties DPvDB-C=N-18C6, the UV-vis absorption of its solutions in the same solvents was examined. Regardless of the solvent, the compound presented low solubility, possibly due to an aggregation effect that occurs as a consequence of the inter- and intra-molecular interactions. In order to determine the concentrations of the compound in the investigated solutions, quartz crystal microbalance with dissipation (QCM-D) measurements were performed. In this respect, based on the QCM-D-measured change in frequency $\Delta f = f_1 - f_0$, the areal-mass difference (Δm) was calculated using the Sauerbrey equation: $^{58,59}\Delta m = -C\Delta f/n$, where $C(17.7 \text{ ng} (\text{cm}^{-2} \text{Hz})^{-1})$ represents a numerical constant related to the properties of quartz, and n (=7 in our case) is the harmonic overtone number which was considered. The results were normalized taking into account the volume of the solution, leading to the concentrations presented in Table S1.†

The experimental UV-vis spectra of **DPyDB-C=N-18C6**, at different concentrations in solvents of different polarities (*n*-hexane, toluene, 1,2-dichloroethane, and ethanol) are presented

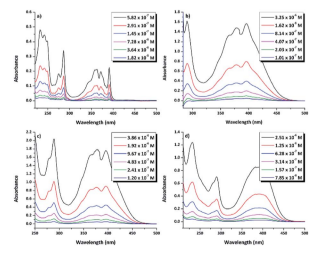


Fig. 5 UV-vis absorption spectra of di-iminopyrene-dibenzo-18-crown-6-ether in: (a) n-hexane, (b) toluene, (c) 1,2-dicholoroethane, and (d) ethanol.

in Fig. 5. The vibronic effects, related to the presence of both pyrene and phenyl rings, appear in two ranges: 200–300 nm and 310–475 nm. The bands in the first range may be assigned to $\pi \to \pi^*$ transitions, while the broad band in the second one could be ascribed to a mixture of $\pi \to \pi^*$ and n $\to \pi^*$ electronic transitions. Remarkably, the structure observed in 310–475 nm range in *n*-hexane was lost in the polar solvents. For each solvent, the spectral shape does not change with the increasing concentration of the solute and does not reveal any new peaks at longer wavelengths.

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3.3.1.2 Fluorescence spectroscopy. The photophysical properties of emission for the DPyDB-C=N-18C6 compound have been investigated by fluorescence spectroscopy at 360 nm excitation wavelength, in the same four solvents as for the UVvis and WAXD analyses (Fig. 6a). Comparing the profiles of the emission spectra, it was found that DPyDB-C=N-18C6 has a weak fluorescence emission, regardless of the chosen solvent. However, a slight increase in fluorescence emission can be seen in 1,2-dichloroethane, due to a different organization of the entire molecular system, interactions between the solvent and the molecules, as well as the increase in the stability of pyrenetype functional groups in this solvent.60 This is also sustained by the QM calculations when it was found that the values of the oscillator strength (f) for the $\pi \to \pi^*$ transitions are higher in this solvent (Table S2†). A weak fluorescence in solution could have many possible reasons, such as aggregation-caused quenching (ACQ) effect,61,62 concentration quenching, photoinduced electron transfer (PET), energy transfer, and a double bond in the imine groups. 62-64 According to references, 38,62-64 the most probable explanation for the weak fluorescence in our system may be mainly related to a PET effect, which arises from the imine nitrogen atom to the large π -conjugation system of the pyrene moiety. Other explanations could be an ACQ effect, concentration quenching, as well as the presence of both pyrene units on the sides of the crown ether that leads to a quenching of the fluorescence emission. The weak fluorescence emission was confirmed by a low photoluminescence quantum yield (PLQY), similarly to the results obtained for other pyrene derivatives. 62-64 Table 3 resumes the PLQY values for the DPyDB-C=N-18C6 compound in each solvent. The exception was the measurement of the quantum yield in toluene, where a higher value was obtained as compared to the other solvents, maybe due to the interactions that develop between toluene

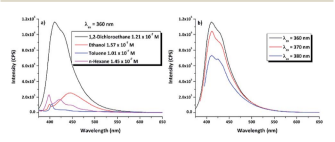


Fig. 6 (a) Fluorescence emission spectra of DPyDB-C=N-18C6 at $\lambda_{\rm ex}=360$ nm, in 1,2-dichloroethane, ethanol, toluene, and n-hexane. (b) Fluorescence emission spectra of DPyDB-C=N-18C6 in 1,2-dichloroethane, at $\lambda_{\rm ex}=360$ nm, 370 nm, and 380 nm.

Table 3 Absolute PLQY of DPyDB-C=N-18C6 in n-hexane, toluene, 1,2-dichloroethane, and ethanol

Solvent	Photoluminescence quantum yield (PLQY)	
<i>n</i> -Hexane	0.39%	
Toluene	0.99%	
Ethanol	0.31%	
1,2-Dichloroethane	0.56%	

molecules and the pyrene units.⁵⁷ Moreover, when the excitation wavelength was increased toward the visible region, there was a decrease in the intensity of the emission spectra regardless of the chosen solvent (Fig. 6b). This fact could be attributed to the low-lying $n \to \pi^*$ electronic transition.

3.3.2. Theoretical studies – quantum mechanics calculations

3.3.2.1 Ground state. In order to explain the inability of **DPyDB-C=N-18C6** to form good intermolecular π - π stacks revealed by MD simulations, quantum mechanics (QM) calculations in the ground state (GS) were performed. Moreover, the equilibrium geometries of both position isomers, predicted by quantitative ¹³C-NMR were studied to investigate if there are differences in the energy, dipole moments, as well as the intramolecular parameters (such as dihedral and valence angles, and bond length).

All equilibrium geometries of both position isomers (i1 and i2) reached after the completion of QM calculations using the PBE0/6-311+G(d,p) method are depicted in Fig. 7 and 8. DFT calculations were carried out in the ground state, considering the singlet state S₀ as a fundamental level. The conformational effect has been studied by rotating the plane of the pyrene moiety following a clockwise direction, while the plane of the phenyl moiety was orientated counter clockwise, by small steps of 10°. The conformational effect of the studied compounds was estimated along the potential energy surface (PES). The motions of both fragments (pyrene and phenyl rings) were performed keeping the plane of double bond -HC=N- fixed (Fig. S7†). The scan procedure was operated for a single plane, the one in which were included the pyrene moiety, the double bond fragment -HC=N-, the phenyl ring, and a few atoms belonging to the crown ether fragment (not the all them). The crown ether

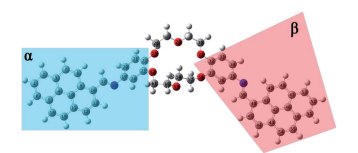


Fig. 7 Equilibrium geometries into the GS for the i1 isomer computed using the PBE0/6-311+G(d,p) method.

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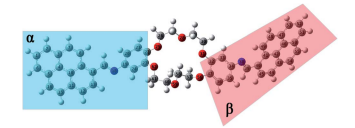


Fig. 8 Equilibrium geometries into the GS for the i2 isomer computed using the PBE0/6-311+G(d,p) method.

fragment was reduced because this sequence can have multiple conformational effects, and to be close to the local minimum given by the QM calculations, a cut of atoms has been made. The new molecular geometry obtained for these scans was previously optimized by the DFT – PBE0/6-311+G(d,p) method, which suffered a conformational search (Fig. S7 and S8 \dagger). The computational results after PES scan indicate that the equilibrium geometry from the GS minimum has the same orientations of sequences as the entire system considered in our study (Fig. S8 \dagger).

The DFT-PBE0/6-311+G(d,p) method indicates that the i2 molecular system was stable in energy as compared to i1 (around $0.42 \text{ kcal mol}^{-1}$ for i1), according to the data in Table S3.† The calculated dipole moment was of 4.59 D for i1 and of about 4.52 D for i2 (see Table S3†). Small differences in the energy stability of molecular systems, as well as into the values of dipole moments indicated that these compounds are hard to be isolated from the solution.

The structural parameters of these compounds, such as dihedral and valence angles, and bond lengths, were extracted after the DFT-PBE0/6-311+G(d,p) calculations. For easier comparison, both position isomers have been separated into two major regions, marked further in text with α and β symbols (Fig. 7 and 8). The α -section (including the pyrene unit, central linking –HC=N– bond, and phenyl fragment) is evidenced with blue color. The β -section (which includes the second phenyl moiety, second central bridge double bond, and second pyrene aromatic ring) is shown in a red color frame.

Based on the values resulting after the optimization using the PBE0/6-311+G(d,p) method performed for GS of i1, a slight variation in the C9–C14–C17=N18 dihedral angle around 4.2° (tending to planarity) was obtained in the case of α -section. On the contrary, the value of the C36–C35–N38=C39 dihedral angle in the β -section was 33.3°. The dihedral angles of fragment sequences are linking the pyrene and phenyl units (C14–C17=N18–C19 and C35–N38=C39–C40, respectively) lost to planarity. The C14–C17=N18–C19 dihedral angle from the α -section has a value around 176.2°, while the C35–N38=C39–C40 dihedral angle from the β -section was orientated with -177.5° . The C17=N18–C19–C24 dihedral angle has a value of around 145.4°, being twisted from planarity with 34.6°. The orientation of the N38=C39–C40–C41 dihedral angle tends to be close to planarity (around 3.6°) (Table S4†).

The optimization results in the case of i2 isomer showed that the C9–C14–C17=N18 dihedral angle (from $\alpha\text{-section})$ is close to planarity (175.2°), while the C35–C34–N38=C39 dihedral angle (from the β -section) is twisted with 39.4°. The values in Table S4† indicated that both bridge dihedral angles, which are linking α - and β -sections (C14–C17=N18–C19 and C34–N38=C39–C40), are close to planarity, having values around 176.2° and -177.1° , respectively. In the case of the β -section, the N38=C39–C40–C41 dihedral angle has a small value of about 4.8°. Therefore, the orientations of dihedral angles have the same trends for both i1 and i2 compounds.

To estimate the interatomic distance values, it was considered that the –HC=N– bond length represents the reference value for both **i1** and **i2** isomers. The computational results resumed in Table S5† indicate that, in the case of **i1** isomer, the –C17=N18– bond length in the α -section is slightly higher (with 0.001 Å) than the –N38=C39– length in the β -section. The same trend resulted in the case of the –HC=N– bonds in the **i2** isomer. No significant differences in the values were observed by comparing the –HC=N– central bridge from **i1** with the same bond belonging to **i2**. In this case, both compounds were isoelectronic, and differences appear at the position isomers.

The results of the PBE0/6-311+G(d,p) calculation in Table S5† predict that the adjacent bonds have different lengths in the α -section, compared with those in the β -section, for both **i1** and **i2** isomers. The intramolecular bond length variations appear due to the extended conjugation system. In the case of **i1**, the length of the adjacent bonds positioned in the α -section was higher in value than the β -section. On the contrary, in compound **i2**, the results are reversed (Table S5†).

Regarding the valence angles, all computational data results are presented in Table S6.† Analysing the obtained results, it can be asserted that the values for **i1** and **i2** isomers were comparable. Moreover, the computational data from Table S6† indicate a slight difference in valence angle values between the two positions. However, considering the valence angle values, as well as the values of central double bonds, both **i1** and **i2** isomers, with respect to the trigonal plane geometry, are in the sp² hybridization state.

In conclusion, it can be affirmed that the molecular geometry is not influenced by variations in bond length and valence angles. Moreover, because the dihedral angles between the pyrene unit and the phenyl aromatic ring of the crown ether (-C-HC=NC-) can vary up to 40° , the assembly in the π - π stacking can be hindered (as discussed in the MD simulations) (as shown in Section 3.2.2).

3.3.2.2. Excitation electronic transitions. The assignments of the excitation electronic transitions from the experimental spectra have been described and correlated with the TD-DFT calculations. The theoretical absorption spectra were calculated for both i1 and i2 position isomers. The equilibrium geometries optimized with the PBE0/6-311+G(d,p) level of theory were used in the TD-DFT computations. The TD-DFT absorption spectrum for each isomer was calculated in the gas phase and then into the implicit solvents: *n*-hexane, toluene, 1,2-dichloroethane, and ethanol, respectively. TD-DFT calculations were made using the PBE0/6-311+G(d,p), TD-calculations were made using the PBE0/6-311+G(d,p), TD-

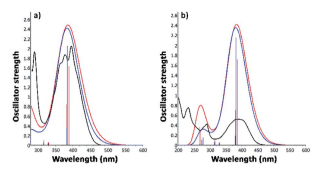


Fig. 9 Electronic absorption spectra of i1 and i2 position isomers, determined by the computational TD-CAM-B3LYP method in toluene (a) and ethanol (b), compared with the experimental UV-vis measurements.

CAM-B3LYP/6-311+G(d,p), TD-B3LYP/6-311+G(d,p) methods. The theoretical data such as values of absorption band maxima (λ_{abs}) and the oscillator strength (f) obtained from TD-DFT calculations are given in Table S2.† The results reveal that the TD-CAM-B3LYP/6-311+G(d,p) method predicts results closer to the experimental data, especially for the wavelengths located in the visible region. Moreover, a good correlation was obtained of electronic theoretical absorption spectra calculated at the TD-CAM-B3LYP/6-311+G(d,p) level of theory in toluene and ethanol solvents as compared to the experimental data (Fig. 9). However, TD-PBE0/6-311+G(d,p) and TD-B3LYP/6-311+G(d,p)

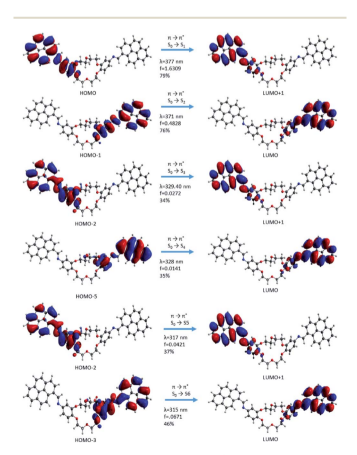


Fig. 10 Molecular orbital representation of the i1 isomer using TD-CAMB3LYp/6-31+G(d,p) in the gas phase.

overestimated the results in comparison with the experimental measurements (Table S2†).

Experimental and TD calculations indicated a strong intensity of the absorption band in the visible region. Comparing the TD calculations results with the experimental measurements illustrated in Fig. 9, no significant shift ranges of the maximum absorption wavelengths were observed. Moreover, the results presented in Fig. 9 indicated that both isomers can absorb at the same wavelength. The oscillator strength values (f) of the i2 isomer calculated with TD-CAM-B3LYP in all solvents (Table S2 \dagger) are higher than those of the i1 isomer, suggesting that the i2 isomer has a slightly hyperchromic effect.

Molecular orbitals corresponding to the excitation electronic transitions involved have been extracted from the TD-CAMB3LYP calculation in the gas phase. The surfaces of molecular orbitals were drawn at the 0.02 a.u. iso level, and they describe the electron density in both the GS and the excited state. The excitation character for the first six transitions of i1 and i2 isomers has been selected for representations (Fig. 10 and 11). The first six transitions have been selected for representation because they give the most intense signals in the electronic absorption spectra. The rest of the transitions have lower values of intensity due to the oscillator strength; hence these transitions are not expected to influence the theoretical absorption band structure. All first six excited state transitions represented in Fig. 10 and 11 suggested $\pi \to \pi^*$ and mixed π $\rightarrow \pi^*$ and n $\rightarrow \pi^*$ characters, which are in good agreement with the experimental absorption data. The mixed excited state transitions appear as an effect of energy levels being near

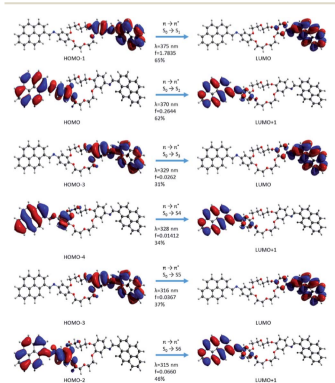


Fig. 11 Molecular orbital representation of the i2 isomer using TD-CAMB3LYp/6-31+G(d,p) in the gas phase.

degenerate, as well as due to the presence of intramolecular interactions (existence of electron density transfer). The electron densities in all cases were localized on the pyrene sequence.

In the case of the i1 isomer, the first transition was $\pi \to \pi^*$ $(S_0 \rightarrow S_1)$ and the electron density was localized on the pyrenephenyl and the imine bond (-HC=N-) sequences from the α section (Fig. 7). The molecular orbital that characterizes this π $\rightarrow \pi^*$ transition is HOMO \rightarrow LUMO+1. The photophysical process during this transition includes a slight delocalization of electron density from the phenyl ring (in the ground state, (π) up to antibonding orbitals of pyrene, as well as on the -HC=Nbond and partially on the phenyl bond corresponding to the excited state (π^*) . This delocalization suggests that the compound has a π -extended conjugation during the photophysical process. The second transition $\pi \to \pi^* (S_0 \to S_2)$ occurred when the process was localized on the sequences from the β-section. The molecular frontier was localized here between HOMO-1 \rightarrow LUMO (Fig. 10). The third transition has a mixed character, $\pi \to \pi^*$ and $n \to \pi^*$, but with predominantly $\pi \to \pi^*$ excitation. This type of transition confirms that the lone pair electrons from the nitrogen atom (belonging to the -HC=N- bond) can participate in the electron density conjugation effect. Only the first two excited state transitions have a major contribution (up to 75%, $S_0 \rightarrow S_1$ and $S_0 \rightarrow S_2$) and a higher oscillator strength value, which indicates that these are predominant in the absorption electronic spectra. The other transitions have small contribution and a small oscillator strength (f) value.

In the case of the i2 isomer, the first transition occurs from the phenyl-HC=N-pyrene sequences and corresponds to the occupied orbital π (HOMO-1) \rightarrow unoccupied orbital π^* (LUMO) in the same fragments. This excitation takes place in the β -section contrary to the situation from the i1 isomer (Fig. 10 and 11). Similar to the i1 isomer, the i2 isomer showed a π -extended electron density conjugation during the photophysical process. The π -extended electron density conjugation appears as an effect of an intramolecular interaction between phenyl, -HC=N-, and pyrene moieties. The second transition includes an π -extended conjugation during the photophysical process of group moieties from the α -section and the nature of this excitation was π (HOMO-1) $\rightarrow \pi^*$ (LUMO) and $S_0 \rightarrow S_2$ (Fig. 11). Both first $(S_0 \rightarrow S_1)$ and second transitions $(S_0 \rightarrow S_2)$ have a major contribution in the absorption electronic spectra because the oscillator strength values (f) were higher than the

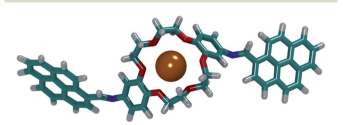


Fig. 12 Example of a complex formed between DPyDB-C=N-18C6 and a K^+ ion. C atoms are shown in teal, O atoms in red, N atoms in blue, K^+ atoms in orange and H atoms in white.

rest of the transitions. The other transitions have small oscillator strength values, and the contributions are less than 46%.

In summary, both pyrene units from the α - and β -sections have a major contribution to observed transitions, but only $S_0 \rightarrow S_1$ and $S_0 \rightarrow S_2$ transitions act with a major contribution on the absorption electronic spectra. The nature of transitions, predicted by the TD-DFT calculations, was also observed in other pyrene derivative studies. 25,29,30,37,65 In the case of the DPvDB-C=N-18C6 compound, no transfer of electron density between pyrene-pyrene fragments (a transfer from pyrene moiety from the α-section to the pyrene unit belonging to the β -section) was observed. Possibly, this potential situation was stopped by the presence of the crown ether sequence, which interrupts the extended conjugation. It can be assumed that if the electronic density was localized on pyrene units from the α - and β -sections, clear π - π stacking interactions would have been observed. The electronic density was localized only on one side of the crown ether sequence (α - or β -section) and this led to a perturbation, forming a well-ordered intermolecular π - π stack.

3.3.2.2.1. MD simulations of ion selectivity. Envisioning a chemo-sensor application for our novel compound was performed a molecular dynamic simulation of 10 DPyDB-C=N-18C6 molecules in a mixed ion solution to test the ion selectivity. The molecules were distributed randomly in an 8 \times 8 \times 8 nm box. Then, 10 of each Na⁺, K⁺ and Ca²⁺ ions and 40 Cl⁻ ions were added, and the entire system was solvated with water. To prevent the aggregation of the DPyDB-C=N-18C6 compound, a position restraint was applied to one of the C atoms from the crown ether. This restrained position allowed the molecule to twist and turn freely in the solution, but at the same time preventing diffusion in the solvent. Two simulations for 40 ns each were performed, starting from two different randomly dispersed structures in order to investigate what ions interact with DPyDB-C=N-18C6 and to eliminate any bias that may have been induced by the initial position of the simulated molecules. The final structures obtained after the simulation are presented in Fig. S9.†

By visually analysing both simulations, it was observed that the **DPyDB-C=N-18C6** molecules interacted only with K⁺ ions, forming stable complexes. Fig. 12 depicts one of the **DPyDB-C=N-18C6** molecule with a K⁺ ion trapped in the crown ether.

These simulations point towards a successful usage of the **DPyDB-C=N-18C6** molecules as a chemo-sensor selective for K⁺ ions. A detailed study of the complexation of **DPyDB-C=N-18C6** with different cations and the effect that the complexation has on the photophysical properties will be detailed in a future study.

4 Conclusions

In this work, a novel compound was synthesised by linking two pyrene moieties to diaminodibenzo-18-crown-6-ether through –HC=N– bonds, resulting in **DPyDB–C=N–18C6** product. The synthesis was confirmed by FTIR, ¹H-NMR, ¹³C-NMR, TGA, and DSC techniques. The quantitative ¹³C-NMR analysis revealed the presence of the two position isomers. WAXD technique and

MD simulations were used to evaluate the possibility of the supramolecular arrangement of DPyDB-C=N-18C6. Both experimental and theoretical methods showed that the compound forms an aggregate with weak π - π interactions. The MD simulation results revealed the existence of almost isoenergetic structures as a slipped-parallel aggregate with an interaction distance around 3.45 and 3.28 Å. Through the QM calculations, it was observed that -C-HC=N-C- dihedral angle, which links the pyrene units to the phenyl rings of the crown ether, was twisted around 40°. This situation explains the inability to align both pyrenes from the same DPvDB-C=N-18C6 molecule with the pyrenes from another molecule, as observed from the MD simulation results. This incapacity is also governed by the presence of a C-H bond and lone pairs of nitrogen from the -HC=N- bond, inducing a hindrance repulsion between individual molecules. Furthermore, the DPvDB-C=N-18C6 compound was investigated from the point of view of its photophysical properties. In order to understand the electronic structure, the UV-vis and fluorescence experiments together with the theoretical studies (OM calculations) were performed in solvents with different polarities (n-hexane, toluene, 1,2-dichloroethane, and ethanol). From the results, we observed the presence of $\pi \to \pi^*$ and $n \to \pi^*$ transitions, among which the $\pi \to \pi^*$ transition due to π -extended conjugation is predominant. By contrast, through fluorescence analysis, a weak emission was observed and could be explained by photoinduced electron transfer (PET) and aggregationcaused quenching (ACQ) effects. The frontier molecular orbitals into the ground state demonstrated that the electronic density of DPyDB-C=N-18C6 was localized only on one pyrene-CH=N-phenyl sequence of the crown ether, leading to a perturbation of a well-ordered intermolecular π - π stacking.

Author contributions

A. C., D.-L. I., and M. P. conceived and planned the experiments; C. A.-M., R. Z., and R. A. performed the synthesis of diminiopyrene-dibenzo-18-crown-6-ether compound; C. A.-M., S.-A. I., D. T., and A. N. performed the physico-chemical characterization of the di-iminopyrene-dibenzo-18-crown-6-ether compound; A. C. performed the photophysical properties measurements; T. V. performed the MD simulations; D.-L. I. performed the QM calculations; A. C., D.-L. I., C. A.-M., T. V., and M. P. wrote and revised the manuscript; F. M., A. L., M. J. M. A., and M. P. supervised the experimental and theoretical studies. All authors have been involved in checking the data, and have given approval to the final version of the manuscript.

Conflicts of interest

There are no conflicts to declare.

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