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Original Citation:

Availability:

This version is available <http://hdl.handle.net/2318/1521785> since 2017-06-27T16:55:43Z

Published version:

DOI:10.1021/acs.jpcc.5b04717

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This is the author's final version of the contribution published as:

Gigli, Lara; Arletti, Rossella; Vitillo, Jenny G.; Alberto, Gabriele; Martra, Gianmario; Devaux, André; Vezzalini, Giovanna. Thionine Dye Confined in Zeolite L: Synthesis Location and Optical Properties. JOURNAL OF PHYSICAL CHEMISTRY. C, NANOMATERIALS AND INTERFACES. 119 (28) pp: 16156-16165.
DOI: 10.1021/acs.jpcc.5b04717

The publisher's version is available at:
<http://pubs.acs.org/doi/pdf/10.1021/acs.jpcc.5b04717>

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Thionine Dye Confined in Zeolite L: Synthesis Location and Optical Properties

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1 Introduction

The physicochemical process of light conversion is under extensive investigation in order to develop more efficient devices for artificial photosynthesis, (1-3) water splitting, photovoltaic applications,(4) and targeting therapeutic agents.(5-7) The photoprocesses of these systems are induced by nanostructured functional materials, such as zeolites light sensitized by dye molecules adsorbed into their pores allowing for the supramolecular organization of the photoactive species. Neutral and cationic dye molecules were successfully incorporated into various zeolites with suitable channel dimensions, such as AlPO₄-5,(8, 9) zeolite Y,(10) and zeolite L(11, 12) as well as mesoporous materials like MCM-41.(13) Zeolite L (ZL) presents itself as an ideal host matrix because its arrays of parallel channels impose severe space restrictions and geometrical constraints to any inserted guest species, leading to very high concentrations of well-oriented dye molecules. Although dye–zeolite systems have been extensively reported in the literature(5-7, 14-17) and investigated experimentally—mainly by UV/vis spectroscopy, fluorescence microscopy, and IR/Raman spectroscopy(18-21)—as well as theoretically,(22-24) the influence of the zeolite pore systems on the dye arrangement and the potential molecules aggregation has not been yet fully investigated from a structural point of view by X-ray powder diffraction (XRPD).

Many factors influence the supramolecular organization of guest molecules, such as their size, shape, charge, and concentration, as well as the Si/Al ratio of the host framework and the nature/amount of the extra-framework species. To date, only a few X-ray powder and single crystal diffraction studies endeavored a detailed structural characterization(25-28) of dye–zeolite host–guest systems. Hennessy et al.(25) characterized the arrangement of the cationic dye methyl viologen (MV²⁺) in zeolite L. They combined X-ray powder diffraction, IR, and Raman spectroscopy and molecular modeling to localize the molecule in the channels, determined the amount of loaded dye, and proposed a slightly inclined orientation of the molecule with respect to the channel axis. Gigli et al.(29) used XRPD, IR spectroscopy, and molecular modeling(22-24) to determine the structure of zeolite L (ZL hereafter) loaded with high amounts of the neutral dye fluorenone. The theoretical and experimental results revealed a very densely packed dye arrangement, confirming the stability of the composite derived by the strong interaction between the dyes C=O group and the K⁺ ions present in the channels.(24, 29)

The family of phenothiazine derivatives (thionine = Th hereafter, methylene blue = MB hereafter, azure A and azure B) attracted considerable attention over the years due to their appropriate biological, chemical, photochemical, and photophysical properties(30) which led them to be used as a model for phototherapeutic agents in photodynamic therapy (PDT) as well as for dye-sensitized solar energy converter.(31, 32) Calzaferri et al.(33) performed first kinetics experiments on the incorporation of Th, MB, and ethylene blue into ZL. They demonstrated the successful incorporation of Th into the zeolite channels, estimating that the dye can exchange at most 7.5% of the K⁺ cations sited in the channel. Later studies on the aggregation properties of Th molecules within zeolites indicated that the channels cannot accommodate Th dimers due to space-filling effect and that the composite showed intense fluorescence.(34) The same authors reported that thionine forms aggregates with the co-guest dye molecule phenanthrene in the cavities of zeolite Y.(34)

The nature of the aggregation of thionine and other related dyes (methylene blue, oxazine 170, Nile blue A, acridine orange, pyronin-Y, and cresyl violet) within zeolites is controlled by different factors, chief among them the presence of water. Karuppanan et al.(35) studied the different optical behavior of Th and MB encapsulated into zeolites Y, ZSM-5, and MCM-41 in absence and presence of TiO₂ nanoparticles by applying absorption as well as picosecond and femtosecond time-resolved fluorescence techniques. They stated that the size, acidity, and hydrophilicity of the nanochannels and nanocavities of the specific zeolite influence the aggregation and the emission lifetime of Th. For example, Th encapsulated in TiO₂ loaded ZSM-5, in the absence of water, is present in two excited states species undergoing excited state interconversion, while Th

encapsulated in the large pore of the MCM-41 does not show this behavior, probably due to the presence of water molecules.

Knowledge of the geometrical arrangement of dye molecules and their chemical environment in the zeolite pores is a fundamental tool to understand the functionality of the host–guest systems. It also helps in elucidating their thermodynamic and kinetic stability, which is central to increasing the efficiency of the systems for energy transfer processes.

Simoncic et al.(26, 36) reported two single crystal X-ray studies on Th and MB incorporated into mordenite–Na. A slightly tilted arrangement of the Th molecules in the channels favors strong host–guest interactions between C, N, and S atoms of the dyes with the framework oxygen atoms. On the other hand, MB molecules were located in two partially occupied sites: one oriented nearly upright and the other slight inclined within the 12-member channel. Molecular mechanics studies on the orientation and dynamic diffusion of Th incorporated in mordenite were also performed by Deore et al.(37) The authors report that Th molecules during the diffusion along the main channel of mordenite assume a centered and upright position, even if one does not consider water molecules. Once the diffusion is completed, the dye molecules bind to the channel wall in the orientation reported by Armbruster et al.(26)

In this study, we report a detailed structural characterization of the organization of the cationic dye Th inserted into ZL channels, which is carried out by XRPD and augmented with FT-IR spectroscopy for the determination of the host–guest interactions as well as by absorption UV–vis and photoluminescence spectroscopies for the investigation of the electronic states of the guest.

2.1 Materials

The potassium ZL (K_{8.46}Al_{8.35}Si_{27.53}O₇₂·17.91H₂O)—LTL-framework type,(38) Si/Al ratio 2.9, space group P6/mmm, a = 18.3795(4), c = 7.5281(2), V = 2202.4(1) (Table 1)—used for this study was purchased from Tosoh Corporation (Japan). The chemical formula was obtained by X-ray fluorescence (XRF). Zeolite L presents large circular 12-ring (12MR) channels of size 7.4 × 7.8 Å(39–41) and smaller elliptical 8-ring (8MR) channels of size 1.9 × 5.6 Å, both running along the c-axis (Figure 1). The main channels are connected to the parallel 8MR ones by nonplanar boat-shaped 8-membered rings.

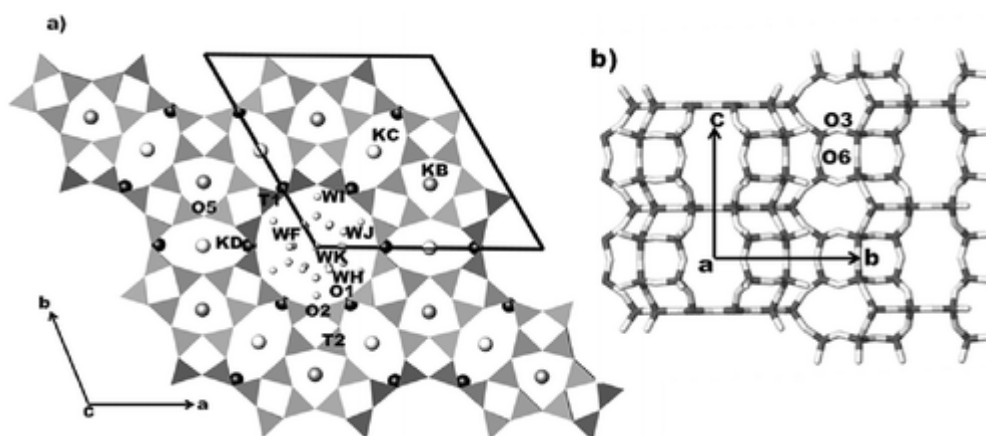


Figure 1. Projection of as-synthesized zeolite L structure and its unit cell along [001] (a) and along [110] (b). Light gray: water sites; dark gray: K sites.

Thionine acetate salt (C₁₂H₁₀N₃S⁺, 90%), was purchased from Sigma-Aldrich. Thionine, a metachromatic dye, is a flat molecule bearing a positive charge on the sulfur atom. Its dimensions, considering van der Waals radii, are 7.2 × 15 Å, and its structure is reported in Figure 2.

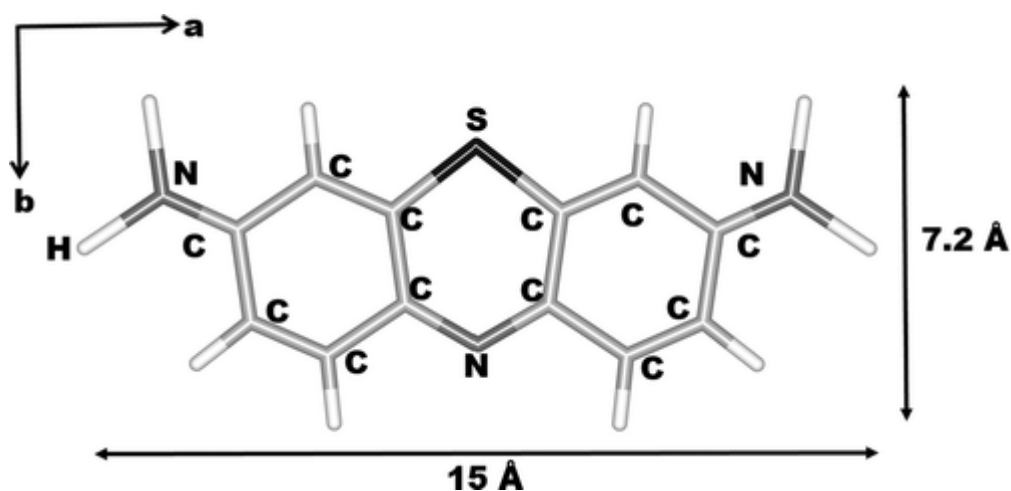


Figure 2. Structure and dimensions of thionine molecule.

2.2 Samples Preparation

Thionine incorporation was carried out by ion exchange, following the procedure described in refs 26 and 33. In a typical experiment, 60 mg of ZL powder was dispersed in either 6.3 or 15.7 mL of a 3.3×10^{-4} M aqueous thionine acetate solution at 80 °C for 3 days in a sealed ampule. After ion exchange, the blue powder was extensively washed: first three times with distilled water and then three times with ethanol to remove molecules adsorbed on the ZL surface. In this way two composites with different Th loading (labeled ZL/0.15Th and ZL/0.27Th) were obtained. Chemical, structural, and physical properties of the samples were determined by using the methods listed in the following section.

3.1 Thermogravimetric Analyses and Evolved Gas Mass Spectrometry (TGA-MSEGA)

TGA-MSEGA analysis of as-synthesized ZL, pure Th, and the two ZL/Th composites were performed on a Seiko SSC 520 thermal analyzer equipped with a quadrupole mass spectrometer (ESS, GeneSys Quadstar 422) using the following experimental conditions: heating rate 10 °C/min, temperature range RT–900 °C, air flux of 100 μ L/min. The gas emitted during the thermal reactions was monitored in order to allow for the unambiguous identification of the species responsible of the weight loss observed in the TGA curve. Gas analyses were carried out in multiple ion detection (MID) mode, following the intensity changes of eight species ($m/z = 18$ (H₂O), 44 and 45 (CO₂), 45.98 (NO₂), 52 (C₄H₄), 64 (SO₂), 76 (N₂O₃), and 78 (C₆H₆)) vs temperature. Before starting MID analysis, background subtraction was applied to set the zero point conditions.

3.2 IR-ATR Spectroscopy

Fourier transformed infrared spectra in air (2 cm^{-1} resolution, average on 256 scans) were collected in attenuated total reflection (ATR) mode on loose powder pressed on the internal reflection element (diamond) of a Bruker OPTIK platinum ATR accessory placed in a Bruker Vertex70 instrument (DTGS detector). The relative intensity of the bands over the spectral region considered was corrected for the different decay in the sample of the evanescent wave as a function of the incident wavelength, by dividing the original band intensities by the corresponding absorption wavelength. Atmospheric carbon dioxide and moisture signals have been subtracted from all the spectra by applying the Atmospheric Correction tools, as implemented in the Opus 6.5 software.

3.3 XRPD

XRPD patterns of the as-synthesized ZL, ZL/Th composites, and the pure dye were collected at the SNBL (BM01a) beamline of ESRF (European Synchrotron Radiation Facility) in transmission geometry, with a fixed

wavelength of 0.6825 Å. The powder samples were loaded and packed in a 0.3 mm boron capillary, mounted on a standard goniometric head, and spun during data collection. The bidimensional diffraction patterns were recorded on a PILATUS3M Series detector (pixel dimension 172 μm) at a fixed distance of 193 mm from the sample. One-dimensional diffraction patterns were obtained in the 2θ range 0–40° by integrating the two-dimensional images with the program FIT2D.(42) Structural refinements were performed by full profile Rietveld analysis using the GSAS package(43) with EXPGUI interface.(44) Notwithstanding the length of Th molecule (15 Å)—almost corresponding to the double of the ZL uc length along the c-axis (7.5 Å)—the diffraction data did not reveal the presence of superstructure and symmetry change in the ZL/Th composites. The refinements were performed in the parental ZL space group P6/mmm. The framework and potassium atom coordinates reported in ref 41 for the room temperature refinement were used as starting model, and the extraframework atoms were derived by inspection of the Fourier difference maps. The Bragg peak profile was modeled using a pseudo-Voigt function with 0.01% cutoff peak intensity. The background curve was fitted using a Chebyshev polynomial with 23 variable coefficients. The 2θ–zero shift was accurately refined into the data set pattern. The scale factor and unit-cell parameters were allowed to vary for all cycles. The refined structural parameters for each data histogram were the following: same thermal displacement value for all tetrahedral atoms, a second value for all framework oxygen atoms, and a third one for the Th molecule atoms. The thermal parameters of K sites and oxygen atoms of water molecules were allowed to vary. Occupancy factors and isotropic thermal displacement factors were refined in alternate refinement cycles. Soft constraints were imposed on Si–O (1.63 Å), C–C (1.39–1.48 Å), S–C (1.63–1.74 Å) and C–N (1.26–1.33 Å) distances during the first stages of the refinements and set free in the last cycles with tolerance values of 0.03 Å. The occupancy factors of Th sites were allowed to vary in the first refinement cycles and were successively constrained to an average value.

Table 1. Experimental and Refinement Parameters for ZL As-Synthesized as Well as ZL/0.15Th and ZL/0.27Th Composites

samples	ZL (ref 29)	ZL/0.15Th	ZL/0.27Th
space group	<i>P6/mmm</i>	<i>P6/mmm</i>	<i>P6/mmm</i>
<i>a</i> (Å)	18.3795(4)	18.3816(3)	18.3866(2)
<i>c</i> (Å)	7.5281(2)	7.5283(2)	7.5281(2)
<i>V</i> (Å ³)	2202.4(1)	2202.9(1)	2204.0(1)
<i>R_p</i> (%)	2.5	2.6	2.7

samples	ZL (ref 29)	ZL/0.15Th	ZL/0.27Th
R_{wp} (%)	3.6	3.6	3.7
RF^{*2} (%)	6.6	6.5	7.6
no. of variables	64	89	89
no. of observations	2211	2225	2223
no. of reflections	936	955	955

The final atomic positions and thermal parameters for the two composites (ZL/0.15Th and ZL/0.27Th) are given in Table S1, while the interatomic distances are reported in Table S2.

3.4 UV–Vis Absorption Spectroscopy

UV–vis–NIR measurements of the ZL/Th samples were carried out in the diffuse reflection mode (DR) on a Varian Cary 5000 spectrometer equipped with an integrating sphere whose inner surface was coated with Spectralon. The spectra were acquired in air with loose powders diluted in pure ZL (1:50 mass ratio) placed inside the standard powder cell of the instrument. The absorption spectrum of thionine acetate was collected in transmission by using a 10–5 M solution in ethanol in a Suprasil cuvette (optical path length: 10 mm). Ethanol was chosen for this measurement, as it is the only solvent guaranteeing the solubilization of dye molecules in monomeric form at this concentration.

3.5 Photoluminescence Spectroscopy

Photoemission/excitation steady-state spectra and fluorescence lifetimes of the ZL/Th samples used above were recorded on a Horiba Jobin Yvon Fluorolog 3 TCSPC spectrofluorometer.

The samples were placed in a Suprasil quartz cuvette with an optical path length of 2 mm. The spectrometer was equipped with a 450 W xenon lamp and a Hamamatsu R928 photomultiplier. A 10–7 M solution of thionine acetate in ethanol was used to record the emission spectra of the thionine monomer in solution (in Suprasil cuvette, optical path length: 10 mm). Fluorescence lifetimes were measured using a time-correlated single photon counting (TCSPC) technique (Horiba Jobin Yvon), with a pulsed laser diode as excitation source at 560 nm (NanoLed, Horiba) and a pulse rate of 1 MHz in a 90° arrangement to a TBX-4 detector. The detector was set to 636 nm, and the aperture of the band-pass was set to its maximum value (24.9 nm) due

to the well-known weak fluorescence intensity of thionine samples.(45, 46) The instrument was set in the Reverse TAC mode, where the first detected photon represented the start signal for the time-to-amplitude converter (TAC), and the excitation pulse triggered the stop signal. DAS6 decay analysis software was used for lifetime calculation.

4.1 TGA-MSEGA

TG and DTG spectra (Figure 3) clearly indicate intercalation of Th molecules into ZL channels. The amount of dye loaded in the two composites, as evaluated by the weight loss resulted in 0.15 and 0.27 Th molecules per uc (Table 2). Calzaferri et al.(33) reported that thionine can replace at most about 7.5% of the potassium in the ZL main channel (KD site), which corresponds to 0.27 molecule per unit cell. This is consistent with our data and evidence that the ZL/0.27Th sample corresponds to the maximum possible dye loading (Table 2).

Figure 3 shows the TG (a) and DTG curves (b) as a function of temperature for the as-synthesized ZL, pure crystalline Th, and ZL/Th composites. The thermal behavior of the as-synthesized ZL—whose total weight loss is about 11.9%—was described in detail in ref 41. In both composites the Th release occurs in one single, wide step in the temperature range of 320–590 °C. The mass spectrometry results showed that Th is thermally decomposed to CO₂, NO₂, and SO₂ species. The weight losses are resumed and interpreted for the two composites in Table 2.

The water weight losses (10% and 9.5% observed in the composites ZL/0.15Th and ZL/0.27Th, respectively, corresponding to 15.7 and 14.9 molecules per uc) are lower than that observed in ZL (18 molecules per uc) (Figure 3 and Table 2). This can be attributed to the Th molecules taking up space in the ZL channels.

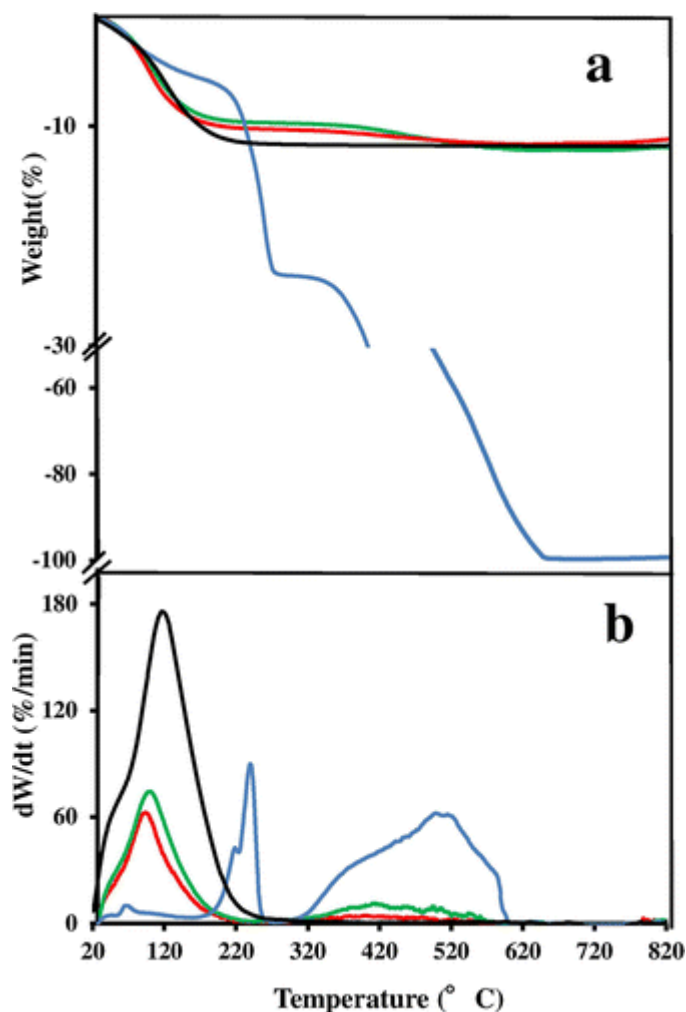


Figure 3. TG (a) and DTG curves (b) for the as-synthesized ZL (black solid curve), pure Th (blue curve), and the composites ZL/0.15Th (red line) and ZL/0.27Th (green line).

Table 2. Temperatures of Water and Th Weight Loss as Well as Number of Molecules (Water and Th), Determined by TGA-MSEGA, Compared with Those Obtained by the Rietveld Refinements for the ZL/Th Composites

sample	water loss T (°C)	water (wt %)	water molecules (puc) (TGA)	water molecules (puc) (structure refinement)	Th loss T (°C)	Th (wt %)	Th molecules (puc) (TGA)	Th molecules (puc) (structure refinement)
ZL(29)	110	11.9	18	18				
ZL/0.15Th102	100	15.7	15.8	15.8	320–590	1.5	0.15	0.19

sample	water loss T ($^{\circ}\text{C}$)	water (wt %)	water molecules (puc) (TGA)	water molecules (puc) (structure refinement)	Th loss T ($^{\circ}\text{C}$)	Th (wt %)	Th molecules (puc) (TGA)	Th molecules (puc) (structure refinement)
ZL/0.27Th100		9.5	14.9	14.8	320–590			

Nevertheless, the amount of water left and the amount of encapsulated Th are not exactly inversely correlated, indicating that thionine, entering the channels, removes only a limited amount of the original water.

4.2 IR-ATR Spectroscopy

Figure 4 shows the IR-ATR spectra of ZL/0.15Th (a, green) and ZL/0.27Th (b, blue) composites as well as of solid thionine acetate (c, black curve) and ZL (d, light gray curve) for comparison purposes. A summary of the assignment of the various signals is listed in Table S3. The spectra of the composites and bare zeolite were normalized with respect to the zeolite framework peak at 1001 cm^{-1} ; thus, differences in relative intensity among the signals due to water and dye molecules in the zeolite channels can be ascribed to actual differences in their amount and/or their decadic absorption coefficients. Basically, the spectra of the composites are close to a superposition of those of the bare ZL and of thionine ions, but some significant spectral features deserve a comment:

(i) Bands due to acetate anions (in particular, those in region of the spectra not occupied by zeolite or thionine cation modes such as νCH_3 at ca. 2950 cm^{-1} and $\nu\text{asymCOO}^-$ at 1544 cm^{-1}) are not present in the spectra of ZL/Th composites, indicating that the negative zeolite framework is acting as counteranion for the encapsulated Th cations.

(ii) The peak at 1608 cm^{-1} in thionine acetate, due to a C–C stretching mode of the aromatic rings, shifts to 1604 cm^{-1} in the composites and exhibits an increase in relative intensity with respect to the other dye modes. A similar behavior was previously reported for methyl blue after adsorption on titanates upon degassing and was considered associated with the direct interaction of π -orbitals with oxidic surfaces.⁽⁴⁷⁾ On the other hand, the apparent reversing in the relative intensity of the bands at 1385 and 1320 cm^{-1} in the solid thionine acetate and in the ZL/Th composites is due to the coexistence of signals related to thionine ($\nu\text{C–N}$ in the ring) and to acetate (νsCOO^-), the latter being no more important in the composites.

(iii) In the composite spectra, bands due to νNH_2 stretching modes, located at 3352 and 3235 cm^{-1} , exhibit a higher relative intensity. This should result from an increase in the polarization of the N–H bonds, suggesting the $-\text{NH}_2$ groups act as hydrogen-bond donors.

(iv) The broad signal due to the νOH of water molecules in the composites, spread over the $3600\text{--}3000\text{ cm}^{-1}$ range, appears similar in intensity for the two materials (slightly lower in the case of ZL/0.15Th) but weaker than for the bare ZL, in agreement with the lower content in H_2O molecules as indicated by TGA measurements. The opposite occurs for the $\delta\text{H}_2\text{O}$ signal: the main difference occurs again between bare ZL and ZL/0.15 Th, with respect to this latter and ZL/0.27 Th. The band is also slightly shifted to higher frequency (1645 cm^{-1}). The rationalization of this effect is not straightforward, but it seems reasonable to associate it to an interaction among water and dye molecules hosted in the zeolite channels (see Figure 4).

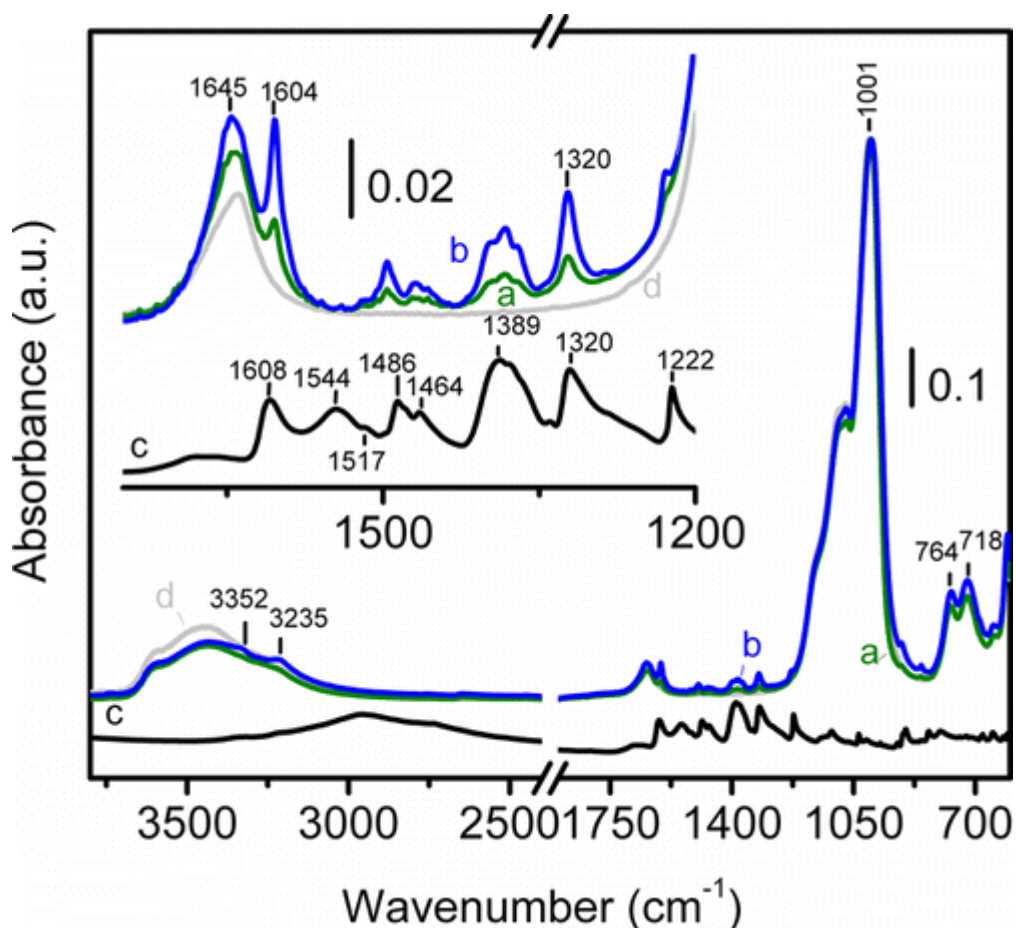


Figure 4. IR-ATR spectra of the ZL/0.15Th (a, green line) and ZL/0.27Th (b, blue line) samples. The inset shows a magnification of the 1750–1200 cm^{-1} region. The spectra for the solid thionine acetate (c, black line) and ZL (d, light gray line) in air are also reported for comparison.

4.3 Structure Refinement

The comparison between the XRPD patterns collected from ZL (see ref 29) and from the ZL/Th composites (Figure S1) shows slight differences in the intensity of some diffraction peaks, particularly in the low 2θ angle region, which is the most sensitive to the extraframework content and distribution. These differences are consistent with the Th penetration and in agreement with the TGA and ATR-IR spectroscopy results. The unit cell dimensions of both composites show an expansion along the a-axis with respect to the unloaded ZL (Table 1). This can be attributed to the penetration of the Th molecules, inducing a decrease of the ellipticity of the 8-membered rings perpendicular to [110] (Table 3). A small decrease of the ditrigonality of the D6R after the adsorption of Th is also observed (Table 3).

Table 3. Ellipticity of the 12MR and 8MR Channels Parallel to [001] and of 8MR Channel Parallel to [110] and Shape of the 6-Rings of the D6R before (ZL) and after the Adsorption of Thionine (ZL/0.15Th and ZL/0.27Th Composites)

ellipticity								
	12MR	8MR c	8MR⊥[110]	D6R			IDa	
	(O2–O2/O1–O1)(O1–O1/O5–O5)(O1–O1/O6–O6)O3–O3O5–O5O3–O5–O3O5–O3–O5							
ZL	1.03	1.70	1.61	3.84	4.97	94.83	144.64	1.29
ZL/0.15Th	1.04	1.72	1.64	3.83	4.95	94.94	143.92	1.29
ZL/0.27Th	1.04	1.67	1.32	3.81	4.83	96.55	142.17	1.27

a

ID = ditrignonalization index of the 6R forming the D6R, given by the ratio between the longest and the shortest axis of the 6R.

The structural refinements of the ZL/0.15Th and ZL/0.27Th composites gave similar results and are hence described here together. The configuration of zeolite L—with its high 6-fold symmetry—and the low amount of dyes consisting of light atoms made molecule localization difficult. However, although the Fourier differences for both composites map showed many residual electronic density peaks, it was possible to define the whole geometry of the molecule and localize it in the channel. In both composite materials, the Th molecule is aligned parallel to the 12MR channel axis and is sited on the mirror planes parallel to the c-axis. In Figure 5 only one of the six possible orientations of the molecule and the organization of the water molecules are represented for the two samples. An amount of 0.19 and 0.30 Th molecules per uc trapped in the zeolite porosities were estimated from the structural analysis for the ZL/0.15Th and ZL/0.27Th composites, respectively. The variation in the occupancy factor of the KD site—the only one accessible to the Th molecule—decreases consistently with the Th–K exchange (Table S1), in agreement with that determined for the two composites by TGA (Table 2).

The water content of ZL corresponds to 18 molecules distributed over five extraframework sites located in the main 12MR channel, labeled WF, WH, WI, WJ, and WK (Figure 1), as reported by Gigli et al.(41) These sites are partially occupied and weakly bound to the framework. The distribution of the residual water molecules in the two composites differs significantly from that of the unloaded ZL. In these composites the water molecules are located in the 12MR channel in four partially occupied sites. Two of them, labeled WH and WF, correspond to those present in ZL, while other two, W3 and W4, are new. As a whole, 15.8 molecules/uc in ZL/0.15Th and 14.8 molecules/uc in ZL/0.27Th composite were found, in good agreement with the TGA results (Table 3).

In the study of Simoncic et al.(26) of the Th–mordenite–Na system, strong interactions were found between the S, N, and C atoms of the Th molecules and the mordenite framework oxygen atoms, favored by a slightly inclined arrangement of the dye molecule along the channel axis.

In our study, the structural data (Table S2) indicates short distances between the C, S, and N atoms and water molecule sites suggesting water-mediated Th–ZL interactions.(53) In fact, N2–W4 (2.24–2.67, and 2.58–2.79 for ZL/0.15Th and ZL/0.27Th, respectively) and N2–WH distances (3.01 and 3.05 for ZL/0.15Th and ZL/0.27Th, respectively) suggest that the NH₂ groups act as hydrogen-bond donors, consistent with the results from the IR-ATR spectroscopy. Other strong interactions are found between S and WH (2.57 and 2.55 for ZL/0.15Th and ZL/0.27Th, respectively) and WF (2.95 and 2.93 for ZL/0.15Th and ZL/0.27Th, respectively) water molecules (see Table S1), in agreement with the changes in the vibrational features of H₂O molecules in the composites.

Water molecules in the ZL pores interact strongly with the dye and form a kind of solvent–matrix tube shaped around the Th molecule (Figure 5), favoring an upright arrangement of the molecules inside the 12MR channels.

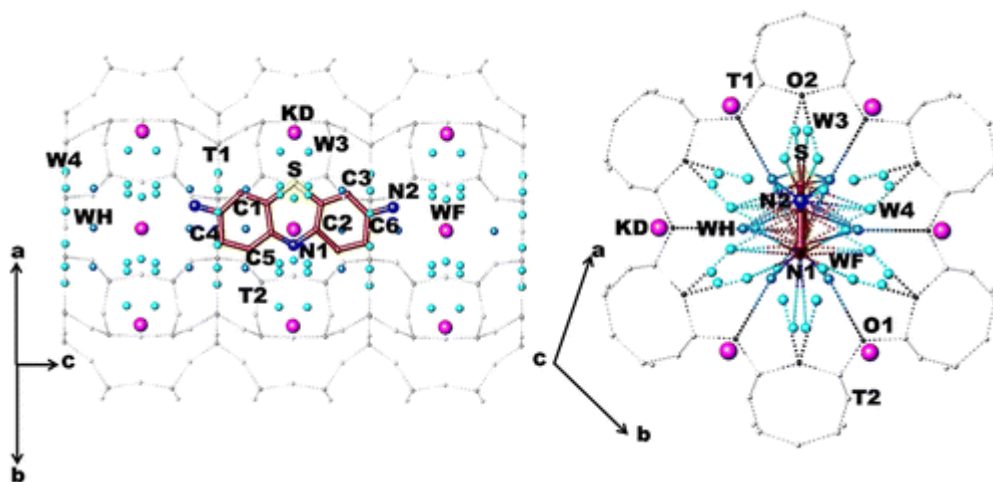


Figure 5. Arrangement of the thionine molecule and of the water–matrix surrounding the dye in the ZL/0.15Th and ZL/0.27Th composites viewed along [110] (a) and [001] (b), as obtained from structural refinements. Large fuxia spheres: K atoms; small light blue and turquoise spheres: water molecules (W). The distances between 2.5 and 3.2 Å are drawn as dashed. The position of only one thionine molecule is represented for sake of clarity.

4.4DR–UV–Vis Absorption Spectroscopy

DR–UV–vis spectra of the ZL/Th composites are reported in Figure 6. Both materials were diluted with pure ZL powder (1:50, mass ratio; see Experimental Section for details) in order to obtain signals, which, once converted by applying the Kubelka–Munk function, remained in the linear range of the relationship between intensity and concentration (KM values ≤ 1).

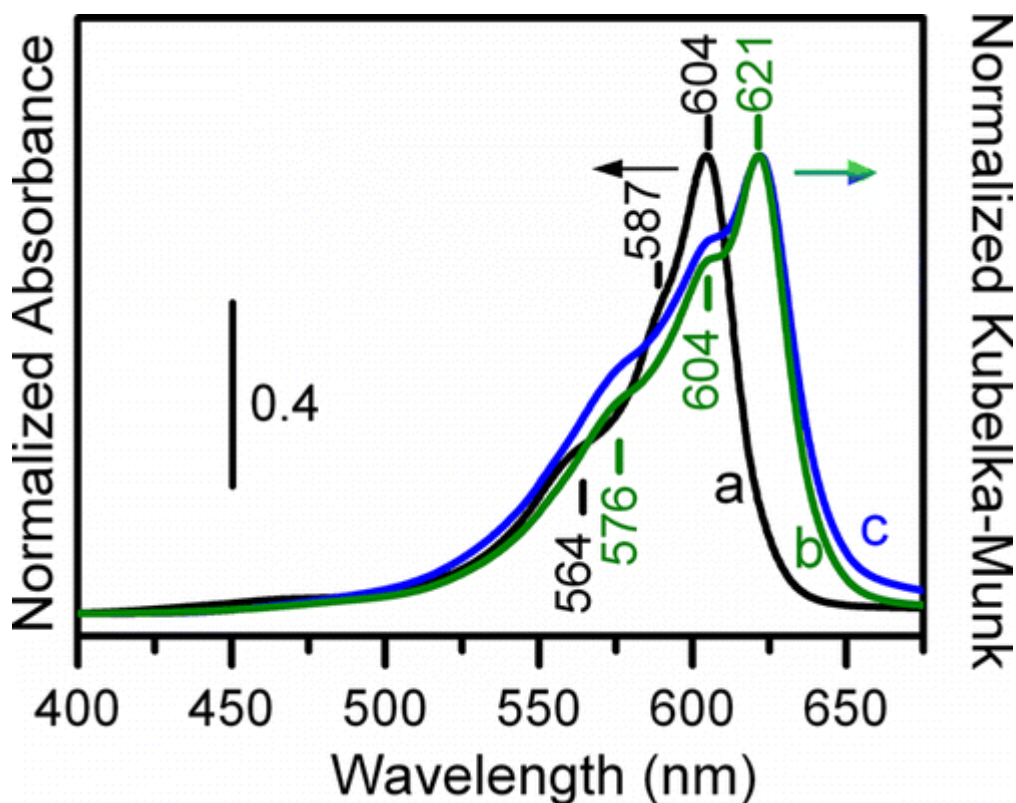


Figure 6. Electronic absorption spectra of (a) thionine acetate as 10^{-5} M ethanol solution (black curve), collected in the transmission mode; (b) ZL/0.15Th (green curve); (c) ZL/0.27Th (blue curve); the last two spectra were collected in the diffuse reflectance mode.

The spectrum of thionine acetate in ethanol solution (a, black curve) exhibits the typical profile expected for the monomeric form, with a main peak at 604 nm and two hypsochromic shoulders at ca. 587 and 564 nm, due to $0 \leftarrow 0$, $1 \leftarrow 0$, and $2 \leftarrow 0$ $\pi^* \leftarrow \pi$ transitions.⁽⁵⁴⁾ Looking at the ZL/0.15Th and ZL/0.27Th composites (b, green curve, and c, blue curve, respectively), the absorption spectra appear almost coincident each other in position and shape, indicating Th molecules basically experience the same environment independently of the loading, which actually is rather low in both cases. Interestingly, the spectral profile of both composites exhibits a similar shape with respect to Th molecules in solution but appeared red-shifted by 17 nm, indicating that the entrapment in the host affected the electronic states of the dye guest. This similarity in shape and the absence of a specific absorption peak at ca. 550 nm typical for thionine dimers (H aggregates)—found in the spectra of Th composites with zeolite with larger pores and higher cation content (ref 35)—indicates that in this case the Th molecules are present in a monomeric form in the ZL channels. Moreover, the better resolved vibronic structure in the ZL/Th spectra indicates the occurrence of some freedom restriction of dye molecules hosted in the zeolite channels. The red-shift of absorption maxima, which is related to the decrease of the energy gap between ground and excited electronic states, might be due to a modification of molecular planarity.⁽²⁸⁾ The deviation from planarity is difficult to evidence from by the XRPD structural refinement, since the electron density map shows an average situation and thus the Th molecule appears located on the mirror plane. As an alternative explanation, in the case of different hybrid systems containing cationic dyes, it has been observed that hydrogen bonds play a relevant role in the stabilization of both ground and excited electronic states.⁽⁵⁵⁾ In this respect, XRD and FT-IR data showed a strong interaction among water and Th molecules hosted inside the channels; the redistribution of the charge in the electronic excited state might then increase the hydrogen bond donor character of $-\text{NH}_2$ groups of Th molecules due to the interaction with water, stabilizing the LUMO and reducing the HOMO–LUMO energy gap.

4.5 Photoluminescence Spectroscopy

In the final part of the investigation, the photophysical behavior of Th molecules hosted in the zeolite channels was investigated. The steady-state photoluminescence spectra are displayed in Figure 7. When comparing the emission spectra of Th in solution (a, black curve) to the ZL/0.15Th and ZL/0.27Th composites (b, green curve, and c, blue curve, respectively), a red-shift of the latter two emission maxima was observed, in agreement with the analogous shift of the absorption spectra. Moreover, the photoemission profiles of Th in composites appeared much broader on the longer wavelength side with respect Th in solution, and this band broadening appears significantly larger than what was observed in the corresponding absorption spectra.

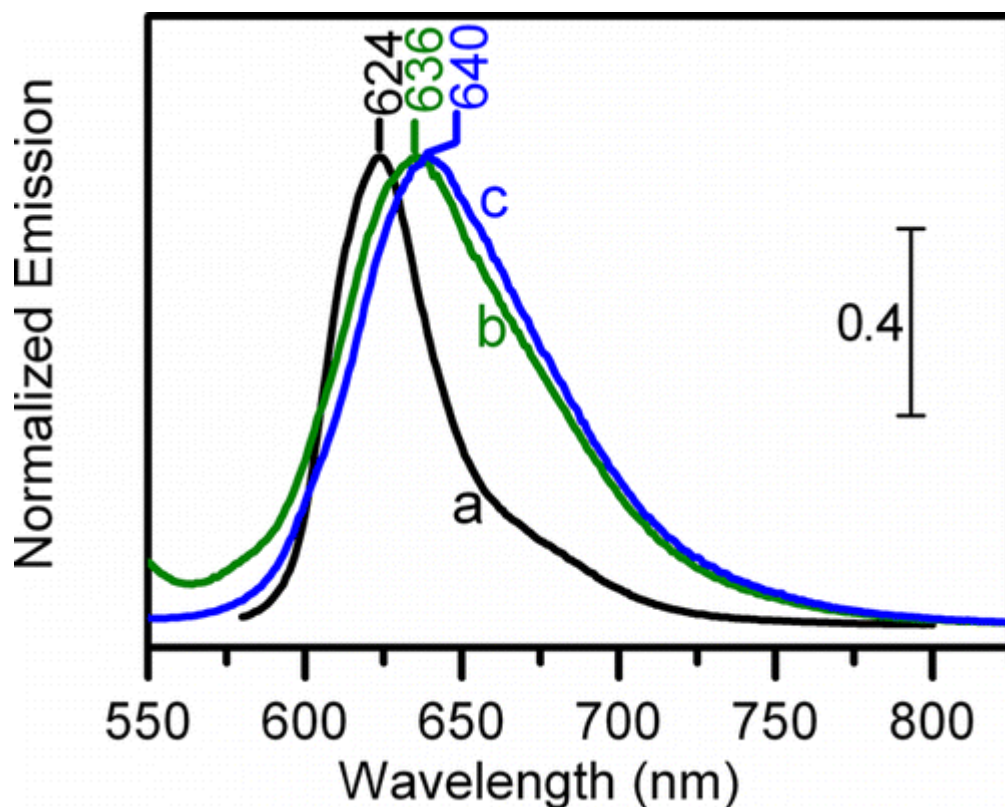


Figure 7. Normalized emission ($\lambda_{exc} = 530$ nm) spectra of (a) thionine acetate as a 10^{-7} M ethanol solution (black curve), (b) ZL/0.15Th (green curve), and (c) ZL/0.27Th (blue curve); these two spectra were collected on samples diluted with bare ZL (1:50, mass ratio).

Photoluminescence spectra of both ZL/Th composites were recorded on samples diluted in pure ZL until a 1:50 mass ratio, in order to avoid signals distortions due to interparticle self-absorption effects; however, due to the high local molar thionine concentration (<0.05 mol/L), the occurrence of intraparticle self-absorption cannot be excluded,⁽¹⁴⁾ likely explaining the slight red-shift observed by increasing Th loading. If the case, the occurrence of this phenomenon should prevent recording the true photoemission profile of photoluminescent Th molecules in the zeolite channels. Anyway, the photoluminescence spectra (or at least the part unaffected by self-absorption) of Th in composites appeared slightly broader than those the molecules in solutions, suggesting that the entrapment in the zeolite channel might have resulted in a modification of the shape of the potential well for the electronic ground state, with consequent changes in the probability of the radiative decays toward excited vibrational levels of this electronic state. Nevertheless, a modification only of the ground state is unlikely. However, changes in the shape of the potential well of the electronic excited state should have produced changes in the shape of the absorption spectra, and this was not the case. Hence, another possibility is that photoluminescent Th molecules in composites are only a small fraction of the total amount present in the zeolite channel, exhibiting a peculiar absorption profile included in the overall absorption spectrum. On this basis, the excitation spectra of Th in solution and of the

composites were collected and compared with the correspondent absorption spectra (Figure 8, panels A and B, respectively).

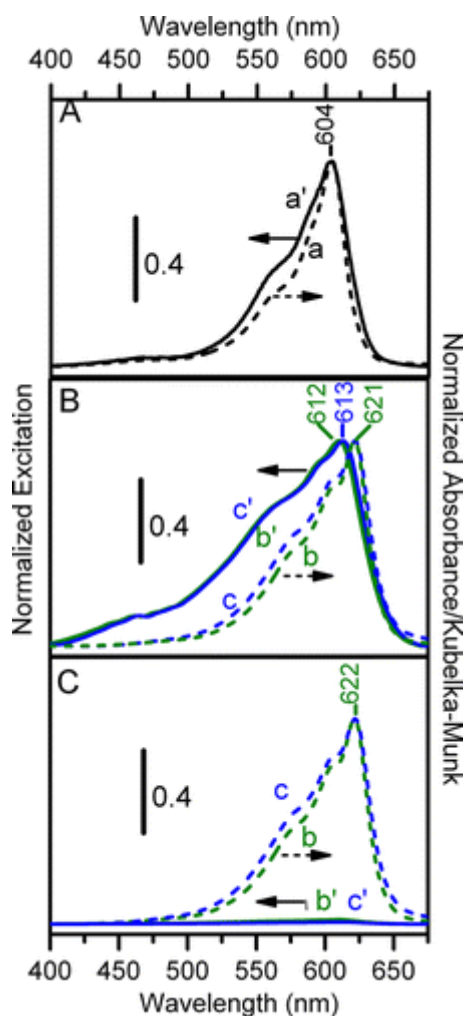


Figure 8. (A) Normalized absorption (a, dashed line) and excitation (a'; solid line; $\lambda_{em} = 700$ nm) spectra of thionine acetate in an ethanol solution. (B) Normalized absorption (dashed lines) and excitation (solid lines, almost coincident for the two composites) spectra of the ZL/0.15Th (b and b', green curves) and ZL/0.27Th (c and c', blue curves) samples diluted in bare ZL (1:50 mass ratio). (C) Comparison of absorption (dashed lines) and excitation (solid lines) spectra of ZL/0.15Th (b and b', green curves) and ZL/0.27Th (c and c', blue curves). Excitation spectra were scaled to the same height as the absorption spectrum.

The absorption profiles of the photoluminescent molecules were acquired in this way. Minor differences were observed for Th in solution (panel A, curves a, a'), whereas the excitation spectra of Th in the composites (panel B, curves b', c') appeared slightly blue-shifted and significantly broader toward shorter wavelengths than the corresponding absorption profiles (panel B, curves b, c), confirming that changes in the shape of the potential well for the electronic excited state also occurred for Th photoluminescent molecules entrapped in the zeolite. It must now be considered that the absorption profile of these molecules, i.e., their excitation spectrum, should be included in the overall absorption profile of all Th molecules present in the zeolite channels. Thus, the intensities of the excitation spectra of ZL/0.15Th and ZL/0.27Th were properly scaled in order to be included under the corresponding absorption curves in the 400–680 nm range. The results are shown in Figure 8C: it appears that photoluminescent Th molecules in composites contribute in a very minor extent to the absorption spectra of these materials. More precisely, on the basis of the ratio among integrated areas, it can be proposed that actually photoluminescent dyes represent ca. 5% of the total amount of Th molecules in the zeolite channels. The fact that the overwhelming part of entrapped dye

molecules appeared quenched can be the consequence of the interaction of their π -orbitals with the hosting environment, as proposed on the basis of the IR spectra (see above). Hence, the small fraction of photoluminescent molecules might correspond to dye molecules hosted in defect positions of the zeolite, where this interaction could be weaker. Moreover, time-resolved measurements (Table 4) indicated that more than 90% of photoluminescent Th molecules in the composite exhibit a lifetime equivalent to Th molecules in solutions, suggesting they are weakly influenced by the host. Conversely, the complementary fraction of entrapped photoluminescent Th molecules exhibit significantly longer lifetimes, likely resulting from an entrapment in peculiar locations allowing severe constraints toward external rotational degree of freedom of the guest molecules.

The intrinsic limits of the diffraction methods—showing the average structure—combined with the low thionine loading hinder the localization of small fractions of molecules potentially located in structural defect sites.

Table 4. Photoemission Decay Times (τ_F) of Thionine in Ethanol Solution (10^{-7} M) and of ZL/0.15Th and ZL/0.27Th Composites

sample	τ_{0F} (ns)	n_1 (%)	τ_{1F} (ns)	n_2 (%)
thionine in solution	0.65	100		
ZL/0.15Th	0.65	93	1.85	7
ZL/0.27Th	0.68	97	1.95	3

It has been demonstrated in the literature that the encapsulation of dye molecules within a zeolite matrix can improve their optical properties such as photostability, luminescence quantum yields, and lifetimes. This study was driven by the need to understand whether the encapsulation of thionine with different loadings in ZL, along with its molecular arrangement, could affect its emitting properties.

The spectroscopic and structural results confirmed a maximum possible incorporation of thionine of 0.27 molecules—present as monomers—per unit cell.

Most of the Th molecules are weakly influenced by the host and exhibit a lifetime equivalent to Th molecules in solutions; on the other hand, a small fraction of molecules, entrapped in peculiar locations, which hinder the external rotational degree of freedom of the Th, show longer lifetimes.

Observed and calculated diffraction patterns and final difference curve from Rietveld refinements of the pure ZL and of the ZL/0.15Th and ZL/0.27Th composites; atomic coordinates, occupancy factors, and thermal displacement parameters for the ZL/0.15Th, ZL/0.27Th composites; extraframework bond distances for

ZL/0.15Th and ZL/0.27Th, composites; the assignment of the various ATR-IR signals of pure compound Th and inside the ZL and bare ZL. The Supporting Information is available free of charge on the ACS Publications website at DOI: 10.1021/acs.jpcc.5b04717.

The authors declare no competing financial interest.

The BM01 beamline at the European Synchrotron Radiation Facility and Dr. Vladimir Dmitriev are acknowledged for allocation of experimental beamtime. Dr. Simona Bigi is acknowledged for the help in TGA-MSEGA analyses. This work was supported by the Italian MIUR, within the framework of the project: FIRB, Futuro in Ricerca “Impose Pressure and Change Technology” (RBFR12CLQD); INFOCHEM project PRIN 2010CX2TLM_006.

Reference

This article references 55 other publications.

1. Calzaferri, G. Nanochannels: Hosts for the Supramolecular Organization of Molecules and Complexes *Langmuir* 2012, 28, 6216– 6231, DOI: 10.1021/la3000872
2. Calzaferri, G.; Lutkouskaya, K. Mimicking the Antenna System of Green Plants *Photochem. Photobiol. Sci.* 2008, 7, 879– 910, DOI: 10.1039/b804682b
3. Calzaferri, G.; Méallet-Renault, R.; Brühwiler, D.; Pansu, R.; Dolamic, I.; Dienel, T.; Adler, P.; Li, H.; Kunzmann, A. Designing Dye-Nanochannel Antenna Hybrid Materials for Light Harvesting, Transport and Trapping *ChemPhysChem* 2011, 12, 580– 594, DOI: 10.1002/cphc.201000947
4. Gust, D.; Moore, T. A.; Moore, A. L. Mimicking Photosynthetic Solar Energy Transduction *Acc. Chem. Res.* 2001, 34, 40– 48, DOI: 10.1021/ar9801301
5. Tsotsalas, M. M.; Kopka, K.; Luppi, G.; Wagner, S.; Law, M. P.; Schafers, M.; De Cola, L. Encapsulating (111) In Nanocontainers for Scintigraphic Imaging: Synthesis, Characterization, and in Vivo Biodistribution *ACS Nano* 2010, 4, 342– 348, DOI: 10.1021/nn901166u
6. Li, Z.; Luppi, G.; Geiger, A.; Josel, H.-P.; De Cola, L. Bioconjugated Fluorescent Zeolite L Nanocrystals as Labels in Protein Microarrays *Small* 2011, 7, 3193– 3201, DOI: 10.1002/smll.201100959
7. El-Gindi, J.; Benson, K.; De Cola, L.; Galla, H.-J.; Kehr, N. S. Cell Adhesion Behaviour on Enantiomerically Functionalized Zeolite L Monolayers *Angew. Chem., Int. Ed.* 2012, 51, 3716– 3720, DOI: 10.1002/anie.201109144
8. Caro, J.; Marlow, F.; Wüst, M. Chromophore–Zeolite Composites: The Organizing Role of Molecular-Sieves *Adv. Mater.* 1994, 6, 413– 416, DOI: 10.1002/adma.19940060517
9. Ihlein, G.; Schuth, F.; Kraus, O.; Vietze, U.; Laeri, F. Alignment of a Laser Dye in the Channels of the AlPO₄-5 Molecular Sieve *Adv. Mater.* 1998, 10, 1117– 1119, DOI: 10.1002/(SICI)1521-4095(199810)10:14<1117::AID-ADMA1117>3.0.CO;2-W
10. Hoppe, R.; Schulz-Ekloff, G.; Wöhrle, D.; Kirschhock, C.; Fuess, H.; Uytterhoeven, L.; Schoonheydt, R. Incorporation of Methylene Blue in NaY Zeolite at Crystallographically Defined Positions *Adv. Mater.* 1995, 7, 61– 64, DOI: 10.1002/adma.19950070114
11. Calzaferri, G.; Pauchard, M.; Maas, H.; Huber, S.; Khatyr, A.; Schaafsma, T. J. Photonic Antenna System for Light Harvesting, Transport and Trapping *J. Mater. Chem.* 2002, 12, 1– 13, DOI: 10.1039/b106141k

12. Mahato, R. N.; Lülff, H.; Siekman, M. H.; Kersten, S. P.; Bobbert, P. A.; de Jong, M. P.; De Cola, L.; van der Wiel, W. G. Ultrahigh Magnetoresistance at Room Temperature in Molecular Wires *Science* 2013, 341, 257–260, DOI: 10.1126/science.1237242
13. Onida, B.; Bonelli, B.; Lucco-Borlera, M.; Flora, L.; Otero Arean, C.; Garrone, E. Spectroscopic Properties of Dye-Loaded Mesoporous Silicas of the Structural Type MCM-41 *Stud. Surf. Sci. Catal.* 2001, 135, 364–364
14. Devaux, A.; Calzaferri, G.; Miletto, I.; Cao, P.; Belser, P.; Brühwiler, D.; Khorev, O.; Häner, R.; Kunzmann, A. Self-Absorption and Luminescence Quantum Yields of Dye–Zeolite L Composites *J. Phys. Chem. C* 2013, 117, 23034–23047, DOI: 10.1021/jp408556g
15. Calzaferri, G.; Huber, S.; Maas, H.; Minkowski, C. Host-Guest Antenna Materials *Angew. Chem., Int. Ed.* 2003, 42, 3732–3758, DOI: 10.1002/anie.200300570
16. Calzaferri, G.; Devaux, A. In *Supramolecular Photochemistry*; Ramamurthy, V.; Inoue, Y., Eds.; Wiley & Sons: New York, 2011.
17. Fois, E.; Devaux, A.; Belser, P.; Brühwiler, D.; Calzaferri, G. Host–Guest Interactions and Orientation of Dyes in the One-Dimensional Channels of Zeolite L *Langmuir* 2013, 29, 9188–9198, DOI: 10.1021/la400579w
18. Calzaferri, G. Organic–Inorganic Composites as Photonic Antenna *Chimia* 2001, 55, 1009–1013
19. Maas, H.; Calzaferri, G. Constructing Dye-zeolite Photonic Nanodevices *Spectrum* 2003, 16, 18–23 [CAS]
20. Devaux, A.; Minkowski, C.; Calzaferri, G. Electronic and Vibrational Properties of Fluorenone in the Channels of Zeolite L *Chem. - Eur. J.* 2004, 10, 2391–2408, DOI: 10.1002/chem.200305673
21. Megelski, S.; Lieb, A.; Pauchard, M.; Drechsler, A.; Glaus, S.; Debus, C.; Meixner, A. Orientation of Fluorescent Dyes in the Nano Channels of Zeolite L. *J. Phys. Chem. B* 2001, 105, 25–35, DOI: 10.1021/jp002582c
22. Fois, E.; Tabacchi, G.; Devaux, A.; Belser, P.; Brühwiler, D.; Calzaferri, G. Host–Guest Interactions and Orientation of Dyes in the One-Dimensional Channels of Zeolite L *Langmuir* 2013, 29, 9188–9198, DOI: 10.1021/la400579w
23. Fois, E.; Tabacchi, G.; Calzaferri, G. Orientation and Order of Xanthene Dyes in the One-Dimensional Channels of Zeolite L: Bridging the Gap between Experimental Data and Molecular Behavior *J. Phys. Chem. C* 2012, 116, 16784–16799, DOI: 10.1021/jp304962w
24. Fois, E.; Tabacchi, G.; Calzaferri, G. Interactions, Behavior and Stability of Fluorenone Inside Zeolite Nanochannels *J. Phys. Chem. C* 2010, 114, 10572–10579, DOI: 10.1021/jp101635p
25. Hennessy, B.; Megelski, S.; Marcolli, C.; Shklover, V.; Bärlocher, Ch.; Calzaferri, G. Characterization of Methyl Viologen in the Channels of Zeolite L. *J. Phys. Chem. B* 1999, 103, 3340–3351, DOI: 10.1021/jp984573y
26. Simoncic, P.; Armbruster, T.; Pattison, P. Cationic Thionin Blue in the Channels of Zeolite Mordenite *J. Phys. Chem. B* 2004, 108, 17352–17360, DOI: 10.1021/jp047288p
27. Van Koningsveld, H.; Tuinstra, F.; Van Bekkum, H.; Jansen, J. C. Single Crystal Structure Analysis of a High-loaded Complex of H-ZSM-5 with Para-dichlorobenzene *Acta Crystallogr., Sect. B: Struct. Sci.* 1989, 45, 423–431, DOI: 10.1107/S0108768189004519
28. Ehrl, M.; Kindervater, H. W.; Deeg, F. W.; Braeuchle, C.; Hoppe, R. Optical Spectroscopy of Thiazine and Oxazine Dyes in the Cages of Hydrated and Dehydrated Faujasite-Type Zeolites: Molecular Dynamics in a Nanostructured Environment *J. Phys. Chem.* 1994, 98, 11756–11763, DOI: 10.1021/j100096a021

29. Gigli, L.; Arletti, R.; Tabacchi, G.; Fois, E.; Vitillo, J. G.; Martra, G.; Agostini, G.; Quartieri, S.; Vezzalini, G. Close-Packed Dye Molecules in Zeolite Channels Self-Assemble into Supramolecular Nanoladders. *J. Phys. Chem. C* 2014, 118, 15732– 15743, DOI: 10.1021/jp505600e
30. Moreira, L. M.; Lyon, J. P.; Romani, A. P.; Severino, D.; Rodrigues, M. R.; de Oliveira, H. P. M. Phenothiazinium Dyes as Photosensitizers (PS) in Photodynamic Therapy (PDT): Spectroscopic Properties and Photochemical Mechanisms; INTECH Open Access Publisher: 2012.
31. Mellish, K. J.; Cox, R. D.; Vernon, D. I.; Griffiths, J.; Brown, S. B. In Vitro Photodynamic Activity of a Series of Methylene Blue Analogues. *Photochem. Photobiol.* 2002, 75, 392– 7, DOI: 10.1562/0031-8655(2002)075<0392:IVPAOA>2.0.CO;2
32. Verbunt, P. P. C.; Debije, M. G. Progress in Luminescent Solar Concentrator Research: Solar Energy for the Built Environment; World Renewable Energy Congress: Sweden, 2011.
33. Calzaferri, G.; Gfeller, N. Thionine in the Cages of Zeolite L. *J. Phys. Chem.* 1992, 96, 3428– 3435, DOI: 10.1021/j100187a047
34. Ramamurthy, V.; Sanderson, D. R.; Eaton, D. F. Control of Dye Assembly within Zeolites: Role of Water. *J. Am. Chem. Soc.* 1993, 115, 10438– 10439, DOI: 10.1021/ja00075a101
35. Senthilkumar, K.; Paul, P.; Selvaraju, C.; Natarajan, P. Preparation, Characterization, and Photophysical Study of Thiazine Dyes within the Nanotubes and Nanocavities of Silicate Host: Influence of Titanium Dioxide Nanoparticle on the Protonation and Aggregation of Dyes. *J. Phys. Chem. C* 2010, 114, 7085– 7094, DOI: 10.1021/jp912267q
36. Simoncic, P.; Armbruster, T. Cationic Methylene Blue Incorporated into Zeolite Mordenite-Na: a Single Crystal X-ray Study. *Microporous Mesoporous Mater.* 2005, 81, 87– 95, DOI: 10.1016/j.micromeso.2005.01.019
37. Deore, S.; Simoncic, P.; Navrotsky, A. Molecular Mechanics Studies of Thionin Blue in Zeolite Mordenite. *Microporous Mesoporous Mater.* 2008, 109, 342– 349, DOI: 10.1016/j.micromeso.2007.05.016
38. Baerlocher, Ch.; McCusker, L. B.; Olson, D. H. Atlas of Zeolite Framework Types, 6th ed.; Elsevier: Amsterdam, 2007
39. Barrer, R. M.; Villiger, H. Z. The Crystal Structure of the Synthetic Zeolite L. *Z. Kristallogr.* 1969, 128, 352– 370, DOI: 10.1524/zkri.1969.128.3-6.352
40. Newsam, J. M. Structures of Dehydrated Potassium Zeolite L at 298 and 78K and at 78K Containing Sorbed Perdeuteriobenzene. *J. Phys. Chem.* 1989, 93, 7689– 7694, DOI: 10.1021/j100359a031
41. Gigli, L.; Arletti, R.; Quartieri, S.; Di Renzo, F.; Vezzalini, G. The High Thermal Stability of the Synthetic Zeolite KL: Dehydration Mechanism by in Situ SR-XRPD Experiments. *Microporous Mesoporous Mater.* 2013, 177, 8– 16, DOI: 10.1016/j.micromeso.2013.04.015
42. Hammersley, A. P.; Svensson, S. O.; Hanfland, M.; Fitch, A. N.; Häusermann, D. Two Dimensional Detector Software: From Real Detector to Idealized Image or Two Theta Scan. *High Pressure Res.* 1996, 14, 235– 248, DOI: 10.1080/08957959608201408
43. Larson, A. C.; Von Dreele, R. B. General Structure Analysis System "GSAS"; Los Alamos National Laboratory Report; Los Alamos, 1994; LAUR 86-748.
44. Toby, B. H. J. EXP GUI, a Graphical User Interface for GSAS. *J. Appl. Crystallogr.* 2001, 34, 210– 213, DOI: 10.1107/S0021889801002242

45. Mackay, R. A.; Lai, W. C. Energy Transfer From Rhodamine. 6G to Thionine in a Anionic Microemulsion *Colloids Surf., A* 2005, 254, 115– 123, DOI: 10.1016/j.colsurfa.2004.04.062
46. Rodriguez-Serrano, A.; Daza, M. C.; Doerr, M.; Marianc, C. M. A Quantum Chemical Investigation of the Electronic Structure of Thionine Photochem. *Photobiol. Sci.* 2012, 11, 397– 408, DOI: 10.1039/C1PP05267E
47. Horváth, E.; Ribic, P. R.; Hashemi, F.; Forróa, L.; Magrez, A. Dye Metachromasy on Titanate Nanowires: Sensing Humidity with Reversible Molecular Dimerization *J. Mater. Chem.* 2012, 22, 8778– 8784, DOI: 10.1039/c2jm16443d
48. Colthup, N. B.; Daly, L. H.; Wiberley, S. E. *Introduction to Infrared and Raman Spectroscopy*, 2nd ed.; Academic Press: New York, 1975.
49. Nakamoto, K. *Infrared and Raman Spectra of Inorganic and Coordination Compounds*, 4th ed.; Wiley: New York, 1986.
50. Hutchinson, K.; Heste, R. E. Raman Spectroscopic Studies of a Thionine-Modified Electrode *J. Chem. Soc., Faraday Trans. 1* 1984, 80, 2053– 2071, DOI: 10.1039/f19848002053
51. Bellamy, L. J. *The Infra-red Spectra of Complex Molecules*, 3rd ed.; Chapman and Hall: London, 1975
52. Jacobs, W. P. J. H.; van Wolput, J. M. C.; van Santen, R. A. An in Situ Fourier Transform Infrared Study of Zeolitic Vibrations: Dehydration, Deammoniation, and Reammoniation of Ion-Exchanged Y Zeolites *Zeolites* 1993, 13, 170– 182, DOI: 10.1016/S0144-2449(05)80274-X
53. Steiner, T. Water Molecules which Apparently Accept no Hydrogen Bonds Are Systematically Involved in C-H...O Interactions *Acta Crystallogr., Sect. D: Biol. Crystallogr.* 1995, 51, 93– 97, DOI: 10.1107/S09074444994007614
54. Lai, W. C.; Dixit, N. S.; Mackay, R. A. Formation of H Aggregates of Thionine Dye in Water *J. Phys. Chem.* 1984, 88, 5364– 5368, DOI: 10.1021/j150666a051
55. Epelde-Elezcano, N.; Duque-Redondo, E.; Martínez-Martínez, V.; Manzano, H.; López-Arbeloa, I. Preparation, Photophysical Characterization, and Modeling of LDS722/Laponite 2D-Ordered Hybrid Films *Langmuir* 2014, 30, 10112– 10117, DOI: 10.1021/la502081c
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