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Paul Legentil, Geneviève Chadeyron, Sandrine Therias, Nicolas Chopin,  
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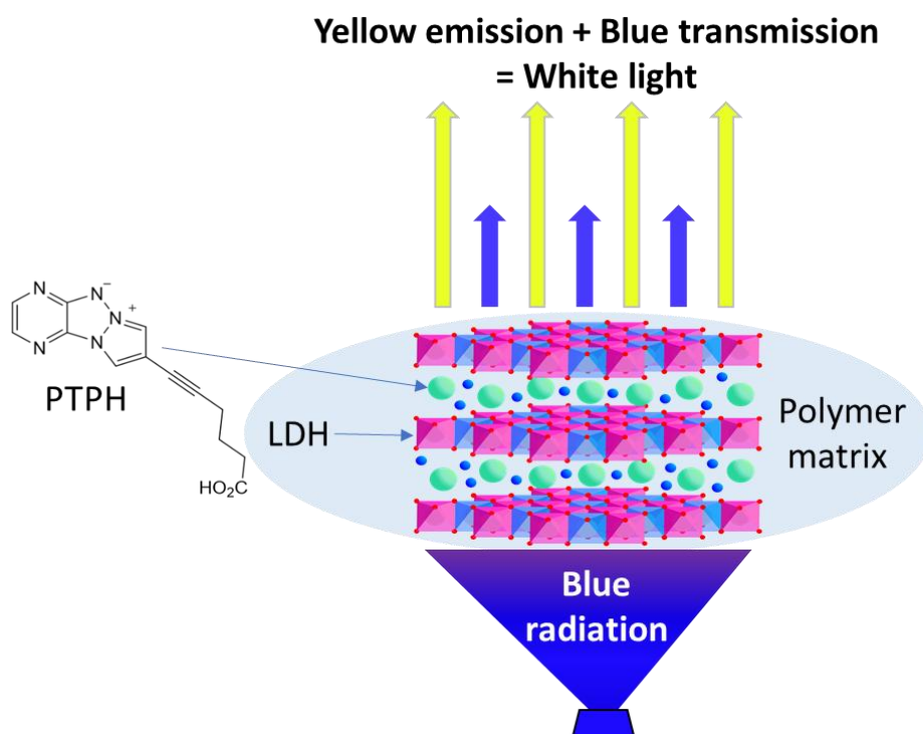
1 Luminescent N-heterocycles based molecular backbone  
2 interleaved into LDH and polymer: for an extensive study

3  
4 Paul Legentil<sup>a</sup>, Geneviève Chadeyron<sup>a</sup>, Sandrine Therias<sup>a</sup>, Nicolas Chopin<sup>b</sup>, Doina Sirbu<sup>b</sup>,  
5 Franck Suzenet<sup>b</sup>, Fabrice Leroux<sup>a,\*</sup>

6  
7 <sup>a</sup>Université Clermont Auvergne, Institut de Chimie de Clermont-Ferrand, UMR CNRS 6296, BP  
8 10448, F-63000 Clermont-Ferrand, France

9 <sup>b</sup>Université d'Orléans, Institut de Chimie Organique et Analytique - ICOA UMR CNRS 7311,  
10 Rue de Chartres, F-45100 Orléans, France

Graphical Abstract



11

## Highlights

- 12 • Pyrazino-1,3a,6a-triazapentalene dye type structure was intercalated into LDH by co-  
13 precipitation method
- 14 • Absolute photoluminescent quantum yields of pristine dye and LDH nanohybrid were  
15 compared
- 16 • Luminescent composite films were prepared with the LDH nanohybrid filler
- 17 • Optical properties of the composite films were characterized for potential lighting  
18 devices

## 19 Keywords

20 Layered double hydroxides; Dye; Intercalation; Fluorescence; Composite films.

## 21 Abstract

22 A pyrazino-1,3a,6a-triazapentalene dye-based molecule was intercalated by direct co-  
23 precipitation method into the galleries of negatively charged layered double hydroxide (LDH)  
24 to form a new luminescent LDH filler. The materials of cation composition  $Zn/Al = 2$  lead to  
25 well-defined organic inorganic assemblies. The structure of the hybrid materials was  
26 characterized by X-ray diffraction (XRD), Fourier-transform infrared spectra (FTIR) and UV-  
27 visible spectra. Optical properties of the materials, at both solid state and slurry state, were  
28 also recorded and absolute photoluminescent quantum yields (PLQY) were determined  
29 showing that the nanohybrid LDH slurry exhibited higher luminescence properties under UV  
30 excitation than the corresponding LDH powder. LDH fillers were used to prepare composite  
31 films by dispersion in both hydrophobic or hydrophilic polymer matrix such as poly(dimethyl  
32 siloxane) (PDMS) and poly(ethylene oxide) (PEO), respectively. Composite films were

33 characterized by Small Angle X-ray Scattering (SAXS) and PLQY were recorded in order to  
34 compare the behaviour of the nanohybrid LDH filler in both polymer matrices. It was found  
35 that the aggregation of particles is still observed by SAXS.

## 36 1. Introduction

37 It is a long-time tradition to use/mix clay minerals with dye molecules, this in particular for  
38 coloring used in Egyptian, Maya time, as beautifully exemplified with the famous «blue  
39 Maya» as well as for the dye immobilization for water treatment. Both domains were  
40 extensively covered with book chapters and reviews(Schoonheydt et al., 2018; Sen, 2017)  
41 and are beyond the scope of the present study focusing on advanced sustainable  
42 development in lighting devices.

43 It is known that optical properties may inform on the location and accommodation of the  
44 guest organic molecule into or onto the clay mineral as recently reported for laponite and  
45 fluorescent dye Safranin simply aggregated or built as Langmuir-Blodgett film (Saha et al.,  
46 2017).

47 As far as clay mineral and organic dye molecule are concerned, polymer may also play a role  
48 in dispersion the aggregated assembly as in polyamide fibers for clothes application using  
49 montmorillonite (Mmt) (Almasian et al., 2019) or layered double hydroxide (LDH) with azo-  
50 dye (Hajibeygi and Omid-Ghallemohamadi, 2017). Interesting work is also focused on very  
51 specific application as those regarding active food packaging polymer, where fluorescent  
52 hemicyanine dye-modified Mmt is solvent cast with chitosan to yield bio-nanocomposite  
53 films (Mekhzoum et al., 2016).

54 Concerning LDH, their structure is referred to the natural hydrotalcite mineral, and described  
55 with the ideal formula,  $[M^{II}_{1-x}M^{III}_x(OH)_2] [A^{m-}_{x/m} \cdot nH_2O]$ , where  $M^{II}$  and  $M^{III}$  are metal cations,

56  $A^{m-}$  the interlayered anions, the  $x$  value determines the charge density of the LDH and  $n$  mol  
57 of water molecules per formula weight are also located. Indeed, the isomorphic substitution  
58 of  $M^{II}$  with  $M^{III}$  cations in the brucite-like sheet gives rise to positive electrical charges  
59 balanced by exchangeable anions ( $A^{m-}$ ) leading to the well-known anion exchange/uptake  
60 LDH ability.

61 To date, pioneer works conducted by the group of U. Costantino were at first performed to  
62 better understand the host / guest interaction and sometimes to address the location of the  
63 organic dye, adsorption versus intercalation by scrutinizing the emission fluorescence and  
64 the associated emitting state energy coming from the possible interactions between the  
65 excited species of the dye molecule and the neighboring unexcited LDH host structure. This  
66 was exemplified with azoic dye methyl orange (MO) into LDH ( $Zn_2Al$  cation composition) for  
67 which the fluorescence of MO interleaved into LDH was found close to MO itself but shifted  
68 at higher energies (Costantino et al., 1999), with the xanthene dye fluorescein surface-  
69 immobilized or intercalated into LDH ( $Zn_2Al$ ), where a dual approach using computer  
70 modeling and diffuse reflectance experiments helped in describing guest arrangement  
71 (Costantino et al., 2000). Their approach was extended to phenolphthalein  $Zn_2Al$  hybrid  
72 materials for which space-resolved fluorescence imaged the dyes distribution and their  
73 interactions with LDH platelets (Latterini et al., 2002), to a series of chromophores,  
74 coumarin-3-carboxylic acid (3-CCA), 9-anthracenecarboxylic acid (9-ACA), 4-benzoylbenzoic  
75 acid (4-BBA) and 2-naphthalenesulfonic acid (2-NSA) (Aloisi et al., 2002), as well as covering  
76 the emerging aspect of fluorescent filler mostly for hydrophobic polymers with anti-  
77 microbial and photo-catalytic applications (Costantino et al., 2013) or simply for mechanical  
78 reinforcement (Marangoni et al., 2008).

79 After this gain in understanding the interaction between LDH and organic dye molecules,  
80 more recent study focus merely on applicative properties such as emphasizing on the  
81 tunable photoluminescent film using both fluorescein (FLU) and 1-heptanesulfonic acid  
82 sodium (HES) / Zn<sub>2</sub>Al hybrid materials for the electro-catalysis of dopamine (Shi et al., 2010).  
83 A step further concerns multicolor light-emitting ultrathin films that were built from Mg<sub>2</sub>Al  
84 LDH hosting photo-functional anions such as bis(N-methylacridinium)@polyvinylsulfonate  
85 ion pairs and derivatives of poly(p-phenylene), poly(phenylenevinylene), and  
86 poly(thiophene) showing a well-defined multicolor polarized fluorescence in association to  
87 high polarization anisotropy thus making them of interest in light displays and optoelectronic  
88 devices (Yan et al., 2011). In the same vein, flexible and robust free-standing films including  
89 LDH platelets disposed between poly(vinyl alcohol) (PVA) and a styrylbiphenyl derivative  
90 (BTBS) via layer-by-layer (LBL) were found to exhibit luminescent anisotropy (Dou et al.,  
91 2013).

92 Again, facing new concerns for possible development of optical devices such as to be  
93 economic, environmental friendly, rare-earth-free, and to address the life cycle assessment  
94 and shelf-life and durability of use, all these prompt us to further study hybrid pigments.

95 These last years, a lot of interest has been carried out to design organic molecules with  
96 optical properties for sustainable alternatives of inorganic luminophores, usually composed  
97 with rare-earth elements. A lot of organic molecules exhibit very high luminescence property  
98 such as N-Heterocyclic compounds and find applications in wide range such as lighting  
99 devices (Weber et al., 2015; Zhang et al., 2008), cell imaging sensors (Gogoi and Sen Sarma,  
100 2015; Mahajan et al., 2019; Volpi et al., 2018) or even dye-sensitized solar cells (Lu et al.,  
101 2018; Wu et al., 2019). Particularly, pyrazino-1,3a,6a-triazapentalene type organic backbone

102 has been identified and substituents are possibly attached to tune the optical properties  
103 (antenna effect, shift of the absorption/emission spectra) (Sirbu et al., 2019). In terms of  
104 tethering to any host structure, such substituents are the functional groups. Indeed, many  
105 researches have proven that both the alkyl chain length and the nature of functional groups  
106 could subtly and significantly impact the intermolecular interactions and molecular packing  
107 and consequently the optical properties of the dye at solid state or in concentrated solutions  
108 (Davis et al., 2008; Kumar et al., 2014). The present study is focus on a dye with a pyrazino-  
109 1,3a,6a-triazapentalene based molecular backbone and 5-hexynoic acid as substituent, this  
110 dye hereafter called compound "PTPH" in the following. The functionalized antenna of PTPH  
111 allows its intercalation within a LDH matrix thanks to the bonds occurring between the  
112 hydroxyl layers and the carboxylic acid function. Other substituents are also possible such as  
113 formic acid and benzoic acid but the PTPH is the only one which presents luminescence  
114 property under solid state. Organic molecules at solid state exhibiting luminescence are very  
115 rare. Usual dyes such as fluorescein, eosin or rhodamine form aggregates at solid state  
116 responsible of the quenching of luminescence (De et al., 2005; Math et al., 2006).

117 Thus, the dye PTPH has been incorporated into a  $Zn_2Al$  LDH matrix by one pot  
118 coprecipitation method. The nanohybrid LDH-PTPH was characterised by X-Ray diffraction  
119 and FT-IR spectroscopy. The photophysical properties of the pristine dye PTPH and hybrid  
120 LDH-PTPH are also scrutinized by UV-visible absorption and fluorescence emission as well as  
121 by recording the absolute photoluminescence quantum yield (PLQY). Consequently,  
122 composite thin films are prepared by dispersing the LDH-PTPH filler into silicon-type polymer  
123 matrix or poly(ethylene oxide). The photophysical properties of the resulting films are also  
124 determined in correlation with their morphology analysed by SAXS.

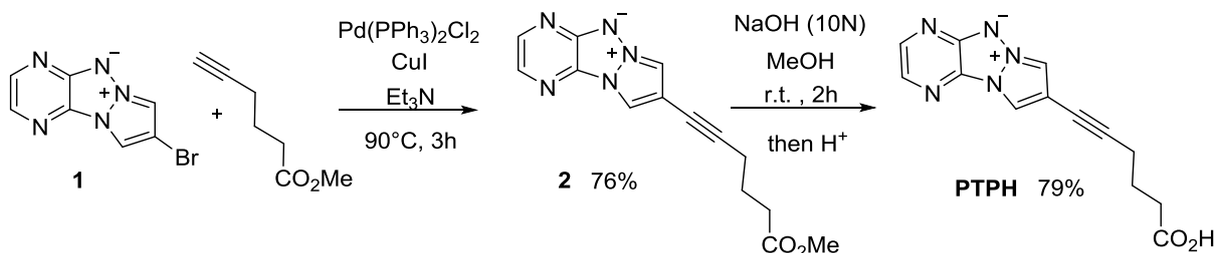
## 125 2. Materials and Methods

### 126 2.1. Chemicals

127 The reagents  $\text{Zn}(\text{NO}_3)_2 \cdot 6\text{H}_2\text{O}$  (purity 99.9+%),  $\text{Al}(\text{NO}_3)_3 \cdot 9\text{H}_2\text{O}$  (purity 99.9+%) and NaOH  
128 (97%) were obtained from Sigma Aldrich. The two-component silicon polymer, Bluesil RTV  
129 141 part A and part B, were supplied by Elkem. The poly(ethylene oxide) ( $M_w$  100,000  $\text{g}\cdot\text{mol}^{-1}$ )  
130 <sup>1</sup>) was supplied by Scientific Polymer Products.

### 131 2.2. Synthesis of PTPH

132 The N-heterocycles dye called PTPH was synthesized starting from the recently reported  
133 bromo derivative **1** (Sirbu et al., 2019). For this purpose, compound **1** was first engaged in a  
134 Sonogashira cross-coupling reaction with the methyl-5-hexynoate. The resulting ester **2** was  
135 then hydrolysed under basic conditions to afford the desired PTPH (Figure 1).



136

137

Figure 1 : Synthesis of PTPH

### 138 2.3. Synthesis of dye co-intercalated in $\text{Zn}_2\text{Al}$ LDH

139 The hybrid layered double hydroxide phase,  $\text{Zn}_2\text{Al}$ -Dye, was prepared by the co-precipitation  
140 method. The synthesis of  $[\text{Zn}_2\text{Al}_1(\text{OH})_6]^+[\text{NO}_3^-]_{1-x}[\text{PTPH}^-]_x \cdot m\text{H}_2\text{O}$ , where  $\text{PTPH}^-$  is the anionic  
141 dye, was performed using the target dye in 50 mL of deionized water. 50 mL of an aqueous  
142 solution of  $\text{Zn}^{2+}$  (1.6 mmol) and  $\text{Al}^{+3}$  (0.8 mmol) was added dropwise during 3 hours under  
143 magnetic stirring. For instance, to prepare the sample LDH-PTPH  $[\text{Zn}_2\text{Al}_1(\text{OH})_6]^+[\text{NO}_3^-]$   
144  $]_{0.60}[\text{PTPH}^-]_{0.4} \cdot m\text{H}_2\text{O}$ , 0.32 mmol of the dye was used. The pH was maintained at 9 by adding

145 0.25 M NaOH during all the addition process. The co-precipitation was performed under  
146 nitrogen at 20°C. The mixture was centrifuged at 5 000 rpm during 10 minutes; the  
147 sedimented solid on the bottom of the flask was washed several times with deionized water  
148 until to obtain a clear and transparent supernatant. A paste was recovered and stored at  
149 5°C. A small amount was dried over night at room temperature to obtain a powder for  
150 further characterizations and to determine the proportion of the dry extract as well as the  
151 uptake of PTPH in LDH-(NO<sub>3</sub><sup>-</sup>/PTPH<sup>-</sup>).

#### 152 2.4. Fabrications of composites thin film Silicon/LDH

153 The LDH hybrid powder was used to elaborate the polymer/hybrid material composite with  
154 different loadings. The two-component silicone polymer (silicone Bluesil-RTV 141 A&B) was  
155 composed of a viscous liquid called part A cured by polyaddition reaction with a catalyser,  
156 part B. The silicone film was prepared following the process: a mixture of the part A and 10  
157 phr of part B was prepared, homogenised with mechanical mixer (Planetary Centrifugal  
158 vacuum Mixer “Thinky Mixer”) during 10 minutes at 500 rpm then pressed in a three rolls  
159 mill Exakt 80E. Next, the final composite film was prepared by casting onto a Teflon surface  
160 using an Elcometer 4340 automatic film applicator. The blade knife height was fixed at 200  
161 µm and the casting speed was 30 mm/s. This two-component silicone polymer was cured at  
162 80°C for 2 hours. The film thickness was measured using an Elcometer 456 coating thickness  
163 gauge.

#### 164 2.5. Fabrications of composites thin film PEO/LDH

165 The LDH hybrid paste obtained after the last cleaning of LDH-PTPH synthesis was used to  
166 elaborate the polymer/hybrid material composite. Hydrophilic poly(ethylene oxide) (PEO)  
167 with Mw = 100,000 g.mol<sup>-1</sup> was used. 1.0 g of PEO powder was dispersed into 10 mL of

168 deionized water under magnetic stirring during 2 hours. 0.4 g of the LDH-PTPH paste was  
169 added to 5.0 g of the previous mixture and then homogenised during 1 hour. The final  
170 mixture was poured directly onto a Teflon surface and dried overnight to obtain the  
171 composite film PEO-LDH-PTPH.

## 172 2.6. Techniques of characterization

### 173 2.6.1. X-ray diffraction

174 LDH powders were characterised by X-Ray Diffraction, the patterns were measured with a  
175 Philips X-Pert Pro diffractometer operating with Cu-K $\alpha$  radiation ( $\lambda=1.5418 \text{ \AA}$ ). The data were  
176 collected in a  $2\theta$  range from  $5^\circ$  to  $70^\circ$  with a step size of  $1^\circ/\text{min}$ .

### 177 2.6.2. SAXS

178 Small Angle X-ray Scattering (SAXS) experiments were performed on an Empyrean  
179 Panalytical equipment with a ScatterX78 using a  $\theta/\theta$  goniometer and Cu anode (45 kV and 40  
180 mA). An elliptic W/Si focusing X-ray mirror for Cu radiation was used, and divergence fixed  
181 slit of  $1/32^\circ$ . The distance from incident beam to sample was 140 mm. The detector  
182 GaliPIX3D was at a distance of 240 mm from the sample. SAXS curves were recorded in  
183 continuous scan mode. The background (no sample) was removed in each case.

### 184 2.6.3. Fourier transformed-infrared spectroscopy

185 The infrared spectra were recorded with a Nicolet 5700-FTIR spectrometer with Omnic  
186 software. Spectra were obtained using a summation of 32 scans and a resolution of  $4 \text{ cm}^{-1}$ .  
187 Powders of LDH and fluorescein were studied by using the KBr pellet technique.

#### 188 2.6.4. UV-Visible absorption

189 The UV-visible absorption spectra of the samples were recorded in the wavelength range of  
190 200 to 800 nm with a UV-vis spectrophotometer (SP-3000 Plus) equipped with an  
191 integrating sphere with UV-Probe software.

#### 192 2.6.5. Luminescence

193 Quantum yields efficiencies were measured using an integrating sphere C9920-02G PL-QY  
194 measurement system from Hamamatsu Photonics. The setup consisted of a 150 W  
195 monochromatized Xe lamp, an integrating sphere (Spectralon coating,  $\varnothing=3.3$  in.) and a high  
196 sensibility CCD camera. Excitation and blue-excited emission spectra were recorded with a  
197 Jobin-Yvon set-up consisting of a Xe lamp operating at 400 W coupled with two  
198 monochromators (Triax 550 and Triax 180) and a cryogenically cold charge coupled device  
199 (CCD) camera (Jobin-Yvon Symphony LN2 series) for emission spectra and Hamamatsu 980  
200 photomultiplier for excitation ones. Excitation spectra were corrected for instrument  
201 response and Xe lamp intensity using sodium salicylate. The resolution of the system was  
202 better than 0.1 nm in both emission and excitation configurations. Luminescence decays  
203 were recorded using a second-harmonic generation on a pulsed Nd:YAG OPO Ekspla NT342A  
204 laser (3–5 ns pulse duration, 10 Hz, 5  $\text{cm}^{-1}$  line width, 0.3 mJ - 20 mJ in the UV-blue). The  
205 emitted photons were detected at right angle from the excitation and analysed through  
206 Edinburgh FLS980 spectrometer (Czerny-Turner monochromator, 300 mm focal length, 1200  
207 groove  $\text{mm}^{-1}$  grating and minimum band-pass of 0.1 nm) equipped with Hamamatsu R928P  
208 PMT (200–850 nm range).

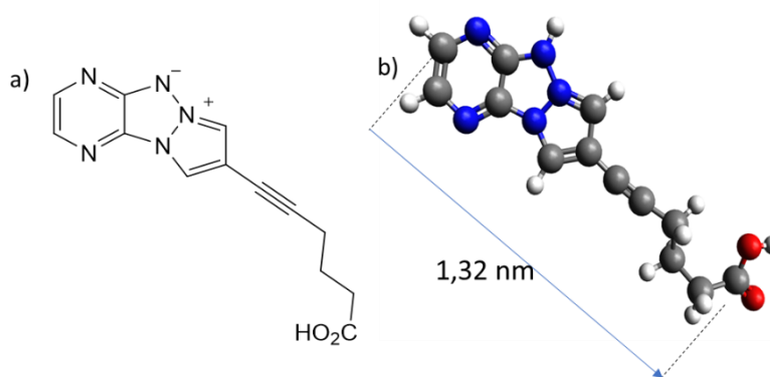
#### 209 2.6.6. Scanning electron microscopy (SEM)

210 Scanning electron microscopy was performed using a ZEISS Supra 55 VP scanning electron  
211 microscope in high vacuum, at 20 kV using a back-scatter electron detector (QBSD).

## 212 3. Results and Discussion

### 213 3.1. Characterisation of the pristine dyes and the LDH-Dye

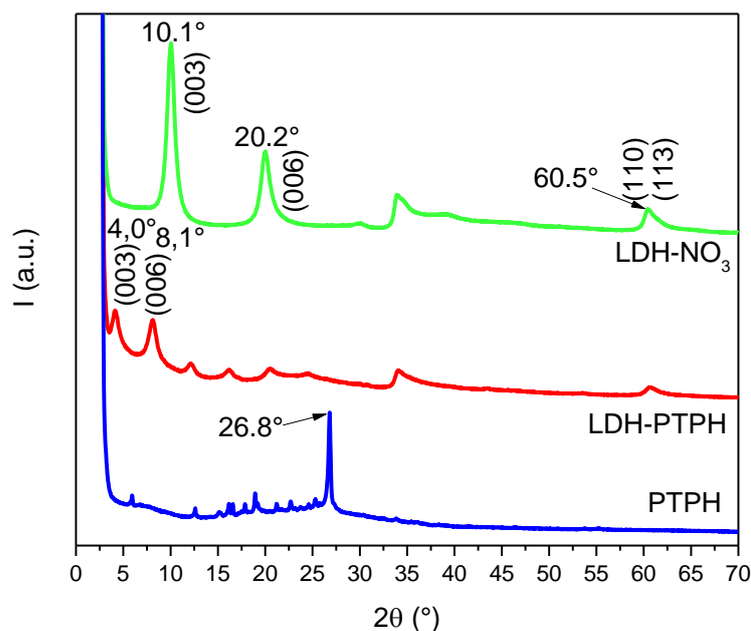
214 The organic molecule PTPH is represented in Figure 2 a). It is composed of a pyrazino-  
215 1,3a,6a-triazapentalene molecular backbone with a 4-hexynoic acid substituent chain.  
216 Carboxylic acid group is attached to allow the intercalation of the dye molecules between  
217 the LDH sheets (Yan et al., 2014). The sample LDH-PTPH powder was synthesized by  
218 coprecipitation method. The XRD patterns of LDH-PTPH powder as well as pristine PTPH  
219 powder and reference LDH-NO<sub>3</sub> powder where nitrate anions are intercalated in a Zn<sub>2</sub>Al LDH  
220 matrix, are represented in the Figure 3. Diffraction planes (00 $l$ ) are the fingerprints of the  
221 layered structure of LDH. For the reference sample LDH-NO<sub>3</sub>, the positions of  $2\theta = 10.1^\circ$  and  
222  $20.2^\circ$  (i.e. 0.78 nm and 0.39 nm) are characteristic of a nitrate LDH phase and assigned to  
223 (003) and (006) diffraction planes, respectively (Mahjoubi et al., 2017). The basal spacing  
224 given by the Scherrer equation (Scherrer, 1912) is 0.78 nm for a (003) diffraction plan at a  
225 position of  $2\theta = 10.1^\circ$ . (110) and (113) diffraction plans are partially overlapped around  $2\theta =$   
226  $60.5^\circ$  leading to a the metal-to-metal distance within the cationic hydroxide layers of a =  
227  $2d_{(110)} = 3.1 \text{ \AA}$ , this in agreement with a ratio Zn : Al of 2 : 1 (Vial et al., 2006).



228

229 *Figure 2: a) Molecular structure of the studied dye: a pyrazino-1,3a,6a-triazapentalene type structure; b) 3D molecular*  
230 *structure of the dye a with calculated size of the molecule*

231 The diffraction plan (003) and (006) of the XRD pattern of LDH-PTPH are observed at  
232 lower  $2\theta$ , the peak positions are  $4.0^\circ$  and  $8.1^\circ$  meaning that the basal spacing (2.21 nm) is  
233 larger than for the LDH-NO<sub>3</sub> (I. Khan and O'Hare, 2002), thus indicating that the large  
234 molecules PTPH are intercalated into the LDH host. Moreover, the geometry of the dye was  
235 optimized with Avogadro software and the length of the molecule is estimated to be 1.32  
236 nm (Figure 2 b)). The basal spacing agrees well the accommodation of the interleaved dye  
237 PTPH. The XRD pattern of the pristine sample PTPH is also represented, a narrow intense  
238 peak is observed at  $2\theta = 26.8^\circ$  which is not observed for the XRD pattern of LDH-PTPH that  
239 excludes the structural packing of free PTPH molecules onto the sample. The mean  
240 crystallite size (L) was calculated according to the Scherrer equation  $L = k\lambda/(B\cos\theta)$ , where B  
241 is the full width at half maximum (FWHM) of the diffraction peak (003) and k is the shape  
242 factor of the average crystallite and fixed to  $k = 0.9$  (assuming a spherical shape) (Cullity,  
243 1978). A coherence length  $L_c$  along the stacking direction of 17 nm was found for the LDH-  
244 PTPH powder meaning that LDH aggregates are composed in average by less than ten  
245 stacked layers.



246

247 *Figure 3: X-Ray Diffractogram (XRD) of the pristine dye PTPH and intercalated in a Zn<sub>2</sub>Al LDH matrix LDH-PTPH as well as a*  
 248 *reference Zn<sub>2</sub>Al LDH matrix with nitrate anions intercalated: LDH-NO<sub>3</sub>*

249 The FTIR spectra of PTPH and LDH-PTPH recorded between 400 and 4000 cm<sup>-1</sup> (Figure 4).

250 Several characteristic bands are observed for the pristine powder PTPH at 1300 cm<sup>-1</sup> (C-N

251 aromatic), 1515 cm<sup>-1</sup> (C=C aromatic), 1715 cm<sup>-1</sup> (stretching bond C=O from carboxylic acid)

252 and around 3100 cm<sup>-1</sup> (C-H aromatic). These bands are also visible in the spectra of the LDH-

253 PTPH powder that confirms the presence of PTPH in the sample. However, the IR band at

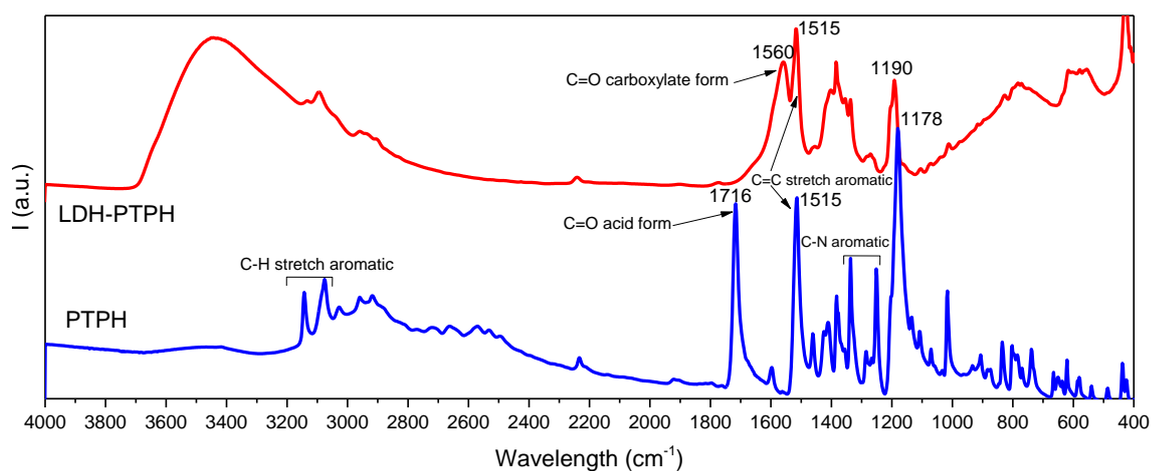
254 1715 cm<sup>-1</sup> (stretching bond C-O from carboxylic acid) disappears from the LDH-PTPH IR

255 spectra while a new band appears around 1560 cm<sup>-1</sup> which is characteristic of stretching

256 bond C=O from carboxylate. Thus, XRD data as well as FTIR spectra confirm the intercalation

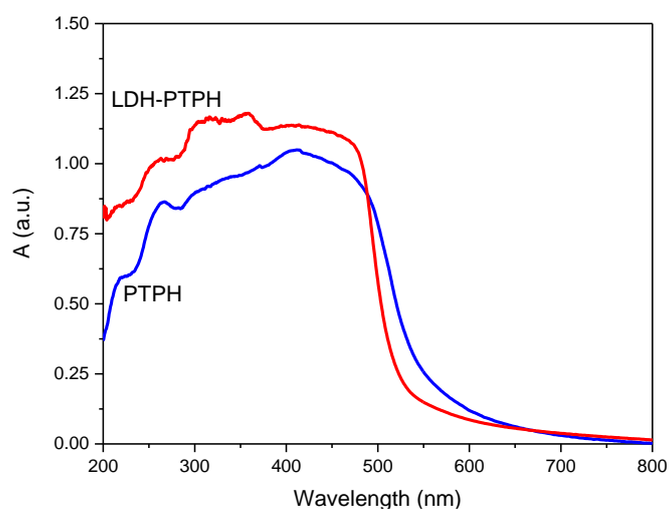
257 and its interaction, respectively, of the molecule PTPH under carboxylate form into the LDH

258 matrix.



259  
260 *Figure 4: FTIR spectra of the pristine dye PTPH and intercalated in a Zn<sub>2</sub>Al LDH matrix LDH-PTPH recorded into KBr pellets*

261 The UV-Vis absorption spectra of PTPH and LDH-PTPH powders recorded into KBr pellets  
262 (Figure 5). Both powders present a large absorbance domain between 200 and 500 nm  
263 without a specific absorbance band. The intercalation of PTPH in the LDH matrix does not  
264 affect its own absorbance properties.

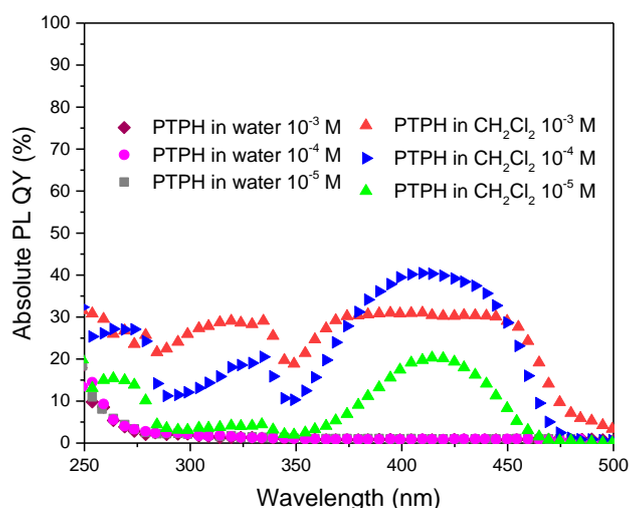


265  
266 *Figure 5: UV-visible spectra of the pristine dye PTPH and intercalated in a Zn<sub>2</sub>Al LDH matrix LDH-PTPH recorded into KBr*  
267 *pellets*

### 268 3.2. Photoluminescence of the pristine dye and the LDH-Dye

269 The absolute PLQY of the dye recorded between 250 and 500 nm for the compound PTPH in  
270 aqueous solution and in dichloromethane solution at three concentrations ( $10^{-3}$ ,  $10^{-4}$  and  $10^{-5}$   
271 M) is displayed in Figure 6. Thanks to the carboxylate group and the polarity of the dye,  
272 PTPH is soluble in water. It is also soluble in organic polar solvent such as dichloromethane

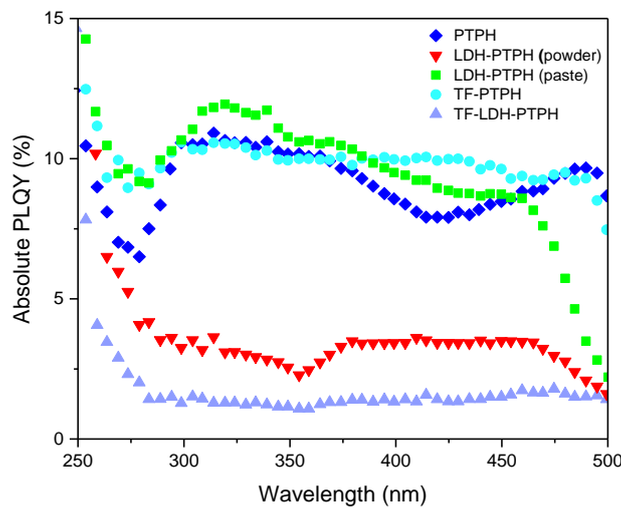
273 or DMSO. The absolute PLQY of PTPH in water is silent; the quenching of luminescence  
 274 properties occurs whatever the concentration. On the other hand, PTPH in dichloromethane  
 275 exhibits high absolute PLQY. In a high diluted system ( $10^{-5}$  M), PTPH shows absolute PLQY  
 276 between 340 and 465 nm, the absolute PLQY<sub>max</sub> is 20.3 % for  $\lambda_{exc} = 315$  nm, whereas the  
 277 absolute PLQY reaches 40.3 % for  $10^{-4}$  M concentration. At  $10^{-3}$  M, compound PTPH displays  
 278 a saturation step between 370 and 450 nm due to high concentration.



279  
 280 *Figure 6: Absolute PLQY in function of the wavelength for solution of PTPH in water and in dichloromethane at three*  
 281 *different concentration:  $10^{-3}$ ,  $10^{-4}$  and  $10^{-5}$  M*

282 The absolute photoluminescence quantum yield (PLQY) of the pristine powder PTPH is  
 283 illustrated on the Figure 7. The PLQY has been recorded between 250 and 500 nm as if the  
 284 dye was associated with an UV-LED. The PLQY varies between 6 and 11 % and the maximum  
 285 is reached at  $\lambda_{exc} = 315$  nm. The values are weak for a potential lighting application but in fact  
 286 they are very interesting for pristine solid organic dye. Indeed, most of the organic dyes at  
 287 the solid state such as fluorescein or eosin do not present any luminescent properties at  
 288 all.(De et al., 2005; De and Kundu, 2011; Yaguchi et al., 2012). The absolute PLQYs of LDH-  
 289 PTPH under powder and paste form are also reported on the Figure 6. As reported in  
 290 Experimental section, the paste is obtained after the washing/centrifugation process and  
 291 before the drying process. The absolute PLQY of the LDH-PTPH powder is almost constant as

292 a function of the wavelength (around 2 %) and are much weaker than the absolute PLQY of  
 293 the pristine powder PTPH. Expectedly, the dispersion of the dye into LDH matrix would allow  
 294 to reproduce a solid state mixed into a solvated system as in a liquid solution to increase the  
 295 absolute PLQY. However, the desired effect is not occurring as for the fluorescein(Legentil et  
 296 al., 2019). The absolute PLQY of the LDH-PTPH paste is higher than the absolute PLQY of the  
 297 powder, the absolute PLQY<sub>max</sub> is 11.9 % for  $\lambda_{exc} = 320$  nm. Counter intuitively, the paste,  
 298 which is just the powder dispersed in a small quantity of water, displays higher absolute  
 299 PLQY than the powder, while the luminescence is quenched for PTPH dispersed in water.

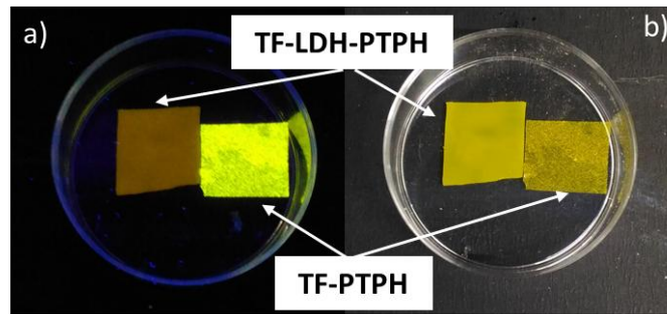


300  
 301 *Figure 7: Absolute PLQY of powders PTPH and LDH- PTPH as well as paste LDH-PTPH and both composite thin films TF-PTPH*  
 302 *and TF-LDH-PTPH*

### 303 3.3. Characterisation of the thin film

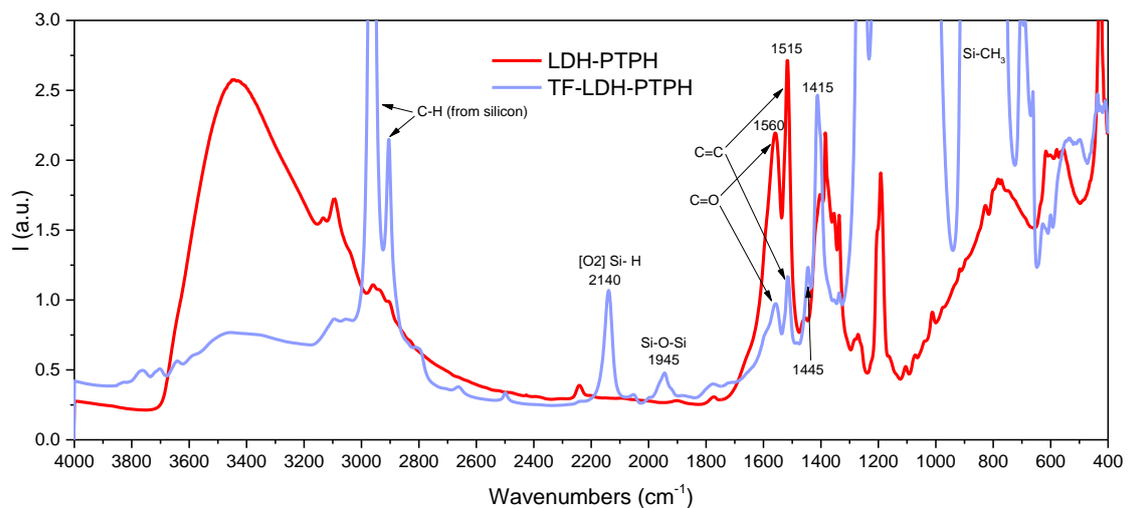
304 Two composite thin films with a silicon matrix (PDMS) was prepared (Deng et al., 2016). The  
 305 first one, TF-PTPH, was prepared with pristine powder PTPH with a loading rate of 5 wt. %  
 306 whereas the second one, TF-LDH-PTPH, was elaborated with the LDH-PTPH powder with a  
 307 loading rate of 20 %<sub>w</sub> to get around 5 %<sub>w</sub> of PTPH. Both films under UV radiation and under  
 308 daylight are illustrated in the Figure 8. First of all, one may observe that the dispersion of  
 309 PTPH molecule in the polymer matrix is quite inhomogeneous, as small aggregates are visible  
 310 in the thin film TF-PTPH leading to a granular aspect. On the contrary, the dispersion in TF-

311 LDH-PTPH is homogeneous without any aggregates. Thus, LDH matrix has a real additional  
312 value in dispersing of the dye PTPH into a polymer matrix. However, TF-PTPH seems to be  
313 more luminescent than TF-LDH-PTPH under UV radiation where luminescence is completely  
314 turned-off.



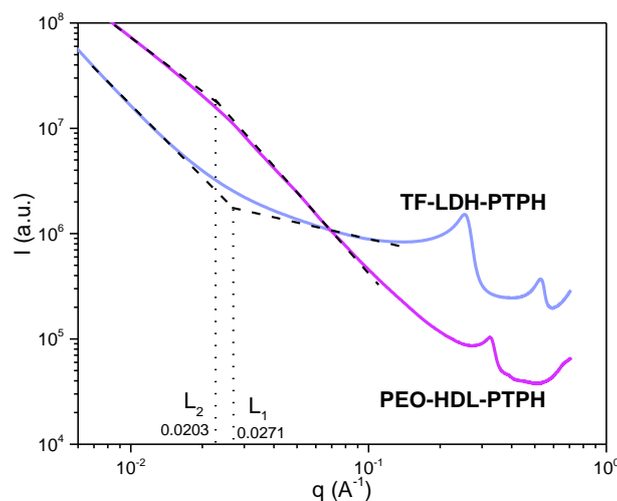
315  
316 *Figure 8: Composite thin films of TF-LDH-PTPH and TF-PTPH under daylight and UV irradiation ( $\lambda=265$  nm)*

317 The FTIR spectra of LDH-PTPH and TF-LDH-PTPH recorded between 400 and 4000  $\text{cm}^{-1}$   
318 (Figure 9). For TF-LDH-PTPH, several IR bands can be attributed to the PDMS polymer (2880-  
319 2960, 2140, 1945, 1445, 1415  $\text{cm}^{-1}$ ) (Bodas and Khan-Malek, 2006). Only two bands, relating  
320 to compound PTPH, at 1515 et 1560  $\text{cm}^{-1}$  are observed. Thus, the dispersion in the matrix  
321 polymer does not modify or degrade the LDH-PTPH powder.



322  
323 *Figure 9: FTIR spectra of powder LDH-PTPH recorded into KBr pellets and the composite thin film TF-LDH-PTPH between 400*  
324 *and 4000  $\text{cm}^{-1}$*

325 The small-angle X-ray scattering data of both composite films TF-LDH-PTPH and reference  
 326 silicon film are presented in Figure 10. Scattered intensity  $I$  is plotted versus the scatter  
 327 vector  $q$ , where  $q = 4\pi\sin(\theta)/\lambda$ . For TF-LDH-PTPH, a characteristic LDH pattern is observed  
 328 (Schaefer and Justice, 2007; Seftel et al., 2015). Two diffraction peaks are observed at 0.26  
 329 and  $0.53 \text{ \AA}^{-1}$  corresponding to distances of 2.42 and 1.18 nm, the first one is the basal  
 330 spacing of the LDH matrix. For PEO-HDL-PTPH, only one diffraction peak is observed at  $0.33$   
 331  $\text{\AA}^{-1}$  corresponding to a basal spacing of 1.90 nm. These values are closed and match with  
 332 data obtained from X-ray diffractogram of the powder LDH-PTPH (2.21 nm). According with  
 333 other SAXS investigations (Jarzębski et al., 1997; Tokudome et al., 2016; Wu et al., 2007), the  
 334 mean size ( $L$ ) of secondary aggregated particles, i.e. LDH pellets, can be calculated from the  
 335 scatter vector found between the slope modifications of the curve.  $L_2 = 27.3 \text{ nm}$  and  $L_1 = 23.1$   
 336 nm were measured, respectively for TF-LDH-PTPH and for PEO-HDL-PTPH. These values are  
 337 slightly higher than the size ( $L_c = 17 \text{ nm}$ ) obtained through the Scherrer equation from the X-  
 338 ray diffractogram of the powder HDL-PTPH. It could be explained by a modification of the  
 339 environment into the basal spacing or a swelling effect due to the polymer matrices.



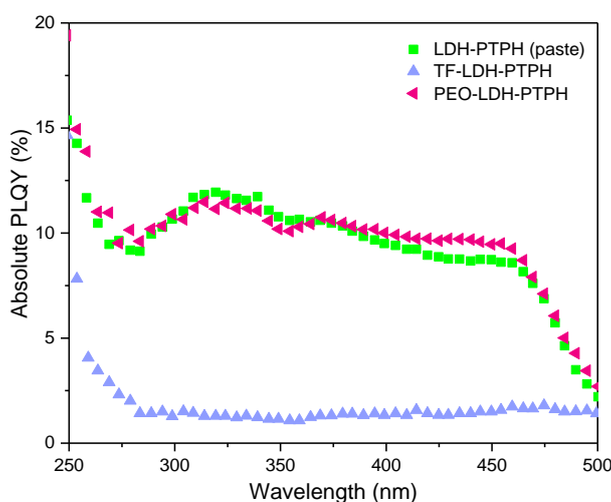
340

341 *Figure 10: Small-angle X Ray scattering (SAXS) of thin composite film TF-LDH-PTPH and a reference silicon film*

### 342 3.4. Photoluminescence of the thin film

343 The absolute PLQY of both composite films TF-LDH-PTPH and TF-PTPH are presented on  
344 Figure 7. The PLQY of TF-PTPH are close to those of the pristine powder PTPH with a  
345 maximum of 11 % for  $\lambda_{\text{exc}} = 315$  nm. The PLQY of the composite film TF-LDH-PTPH is  
346 extremely weak, between 1 and 2 %. The luminescence of the thin film is barely observed  
347 under UV irradiation. These results show that the intercalation of the molecule PTPH into  
348 LDH host is not suitable for the luminescence properties, this is also the case after dispersion  
349 into a PDMS-type polymer.

350 A hydrophilic polymer PEO is then used to prepare a second composite film with the LDH-  
351 PTPH paste which presents decent photoluminescence property. The absolute PLQY of the  
352 film PEO-LDH-PTPH were presented on Figure 11. The PLQY between 250 and 500 nm are  
353 perfectly matching with the PLQY of LDH-PTPH paste. The photoluminescence property of  
354 the paste is preserved through its dispersion into a hydrophilic polymer such as PEO, then  
355 yielding a luminescent composite film. Further studies are needed to unravel the unexpected  
356 behaviour of the observed stable luminescence in such medium.



357  
358 *Figure 11: Absolute PLQY of composite films PEO-LDH-PTPH and TF-LDH-PTPH as well as paste LDH-PTPH*

## 359 4. Conclusion

360 The study demonstrates that the organic molecule PTPH (pyrazino-1,3a,6a-triazapentalene  
361 type structure) exhibits efficient photoluminescence property for an organic molecule at  
362 solid state. The cumbersome organic guest PTPH is well intercalated into LDH matrix  $Zn_2Al$   
363 using a coprecipitation method, enough ensconced to get a hybrid luminescent material.  
364 Counter intuitively, the LDH-PTPH powder does not emit any radiation under blue/UV  
365 excitation (absolute PLQY lower than 2 %) whereas the LDH-PTPH paste produces  
366 luminescence with an absolute PLQY<sub>max</sub> of 11.9 % for  $\lambda_{exc} = 320$  nm. The dispersion of the  
367 LDH-PTPH powder in a polymer (PDMS) occurs readily thanks to the compatible LDH host  
368 matrix. However, such dispersion is not able to produce luminescence enough. When  
369 hydrophilic polymer such as poly(ethylene oxide) is combined with the LDH-PTPH paste, the  
370 absolute PLQY is similar to LDH-PTPH paste values, reaching an absolute PLQY<sub>max</sub> of 11.5 %  
371 for  $\lambda_{exc} = 315$  nm, and showing that the continuous phase; i.e. polymer, may play a role at  
372 nanoscale.

## 373 Acknowledgement

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## Supporting information

504

### 505 **Materials and methods**

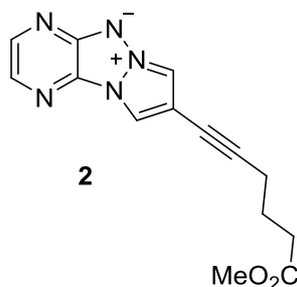
506 Nuclear Magnetic Resonance (NMR) spectra were recorded on a 250 or 400 MHz  
507 Bruker NMR spectrometers in CDCl<sub>3</sub> or DMSO. All chemical shift values are reported in parts  
508 per million (ppm) with coupling constant (J) values reported in Hz. All spectra were  
509 referenced to the CDCl<sub>3</sub> residual solvent peak CHCl<sub>3</sub> ( $\delta$  = 7.26 ppm) for <sup>1</sup>H NMR and the CDCl<sub>3</sub>  
510 solvent peak ( $\delta$  = 77.16 ppm) for <sup>13</sup>C NMR. The notation of signals is: Proton:  $\delta$  chemical shift  
511 in ppm (multiplicity, J value(s), number of protons). Carbon:  $\delta$  chemical shift in ppm.  
512 Fluorine:  $\delta$  chemical shift in ppm. Splitting patterns are assigned s = singlet, b = broad, d =  
513 doublet, td = triplet of doublet, dt = doublet of triplet, t = triplet, q = quartet, app= apparent.

514

515 All reaction were monitored by thin-layer chromatography (TLC) analysis using silica  
516 gel (60 F254). Solvents, unless otherwise stated, were purchased in reagent grade or  
517 anhydrous quality and used as received. Reagents were either purchased directly from  
518 commercial suppliers or prepared according to literature procedures. Yields of all the  
519 compounds refer to isolated compounds. Chromatography: Separations were carried out on  
520 Silica gel 60, (40-63  $\mu$ , 60 Å) purchased from Sigma Aldrich. High Resolution Mass  
521 Spectrometry (HRMS) were recorded on Maxis Bruker 4G. Melting points (mp [C°] ) were  
522 taken on open capillary tubes using a Electrothermal IA 9100 apparatus.

523 Reactants were purchased from commercial suppliers and were used without further  
524 purification. Compound **1** was synthesized following reported synthesis (Sirbu et al., 2019).

## 525 Experimental procedures



526

527 In a round bottom flask, the bromopyridazinotriazapentalene **1** (Sirbu et al., 2019) (1.0 g,  
528 4.20 mmol) was solubilized in Et<sub>3</sub>N (10 mL) with methyl-5-hexynoate (1.060 g, 8.40 mmol, 2  
529 eq.) and Pd(PPh<sub>3</sub>)<sub>2</sub>Cl<sub>2</sub> (294.8 mg, 0.42 mmol, 10 mol%). The mixture was stirred at room  
530 temperature for 20 min and then CuI (72.4 mg, 0.38 mmol, 9 mol%) was added. The mixture  
531 was stirred at 90°C for 3h. Solvent was removed under vacuum and crude compound purified  
532 by column chromatography (DCM/AcOEt 1/1) to give pure **2** (900 mg, 76%) as a yellow solid.

533 Pf: 99°C

534 IR  $\nu(\text{cm}^{-1})$ : 3138, 3078, 2959, 1721, 1507, 1424, 1382, 1368, 1336, 1256, 1186.

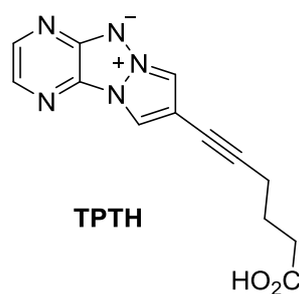
535 <sup>1</sup>H NMR (400 MHz, Chloroform-d)  $\delta$  8.35 (d,  $J = 2.7$  Hz, 1H), 8.03 (s, 1H), 7.83 (d,  $J = 2.7$  Hz,  
536 1H), 7.78 (s, 1H), 3.63 (s, 3H), 2.45 (td,  $J = 7.2, 5.4$  Hz, 4H), 1.89 (p,  $J = 7.2$  Hz, 2H).

537 <sup>13</sup>C NMR (101 MHz, CDCl<sub>3</sub>)  $\delta$  173.27, 152.43, 143.33, 130.43, 128.94, 111.86, 110.52, 107.83,  
538 93.28, 70.19, 51.66, 32.81, 23.49, 18.86.

539 HRMS (ESI)  $m/z$ : calculated for C<sub>14</sub>H<sub>14</sub>N<sub>5</sub>O<sub>2</sub> [M+H]<sup>+</sup> 284.1142; found 284.1141.

540

541



542

543 In a round bottom flask, **2** (500 mg, 1.77 mmol) was solubilized in MeOH (4 mL) with NaOH  
544 10N (1 mL, 10 mmol, 5.6 eq.). The mixture was stirred at room temperature for 2h. Solvents  
545 were removed under vacuum and the crude mixture solubilized in DCM and water (1/1).  
546 Aqueous phase was extracted and washed with DCM (3 x 20 mL) and then acidified with HCl  
547 2N until pH 4-5. Organic phase was extracted with DCM, washed with water and  
548 concentrated under vacuum to give pure **TPTH** (377 mg, 1.40 mmol, 79%).

549 *Pf*: ~175°C (decomp.)

550 IR  $\nu(\text{cm}^{-1})$ : 3143, 3074, 2960, 1715, 1512, 1382, 1336, 1250, 1176, 1016.

551  $^1\text{H}$  NMR (400 MHz, DMSO- $d_6$ )  $\delta$  12.14 (s, 1H), 8.85 (s, 1H), 8.58 (s, 1H), 8.36 (d,  $J = 2.7$  Hz,  
552 1H), 7.90 (d,  $J = 2.7$  Hz, 1H), 2.55 – 2.46 (m, 2H), 2.38 (t,  $J = 7.3$  Hz, 2H), 1.78 (p,  $J = 7.2$  Hz,  
553 2H).

554  $^{13}\text{C}$  NMR (101 MHz, DMSO)  $\delta$  173.96, 152.08, 142.65, 129.59, 128.92, 113.90, 111.37,  
555 106.68, 92.99, 70.80, 32.60, 23.46, 18.16.

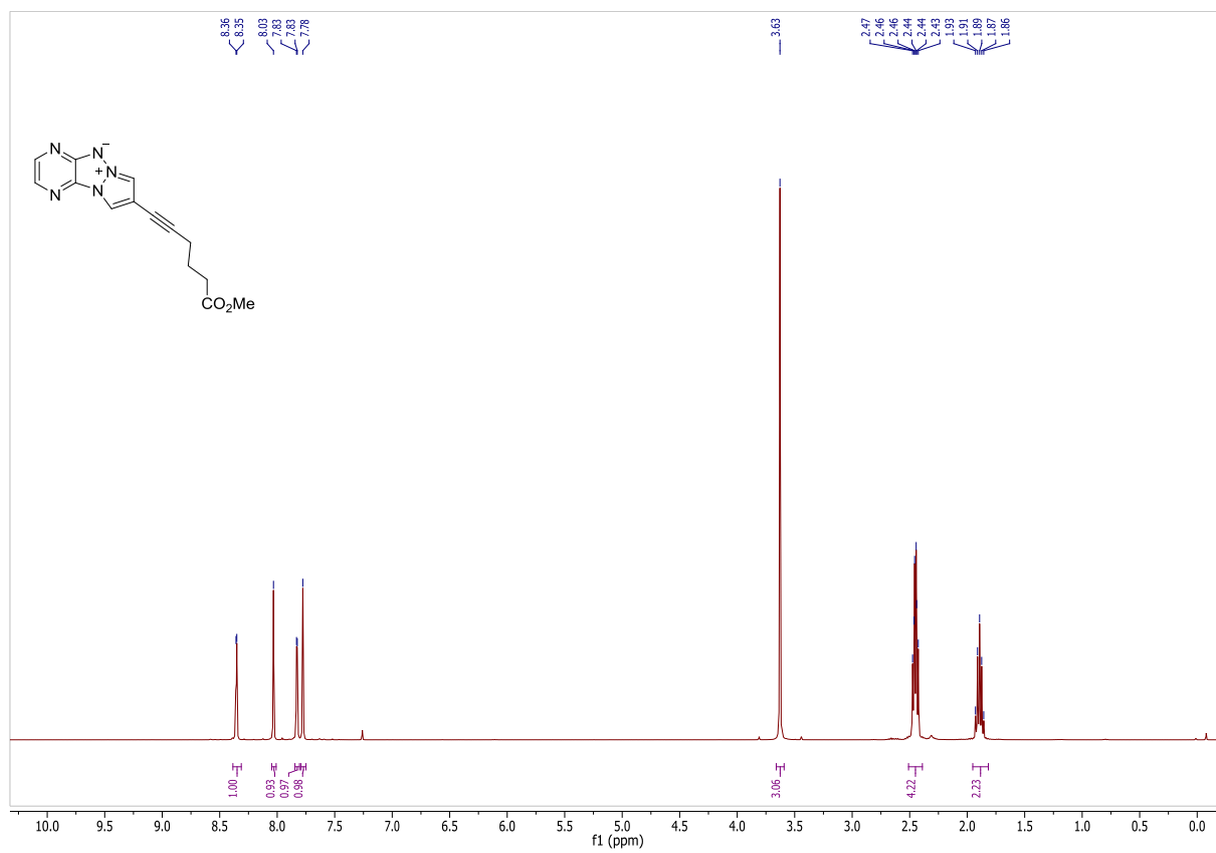
556 HRMS (ESI)  $m/z$ : calculated for  $\text{C}_{13}\text{H}_{12}\text{N}_5\text{O}_2$   $[\text{M}+\text{H}]^+$  270.0985; found 270.0984.

557

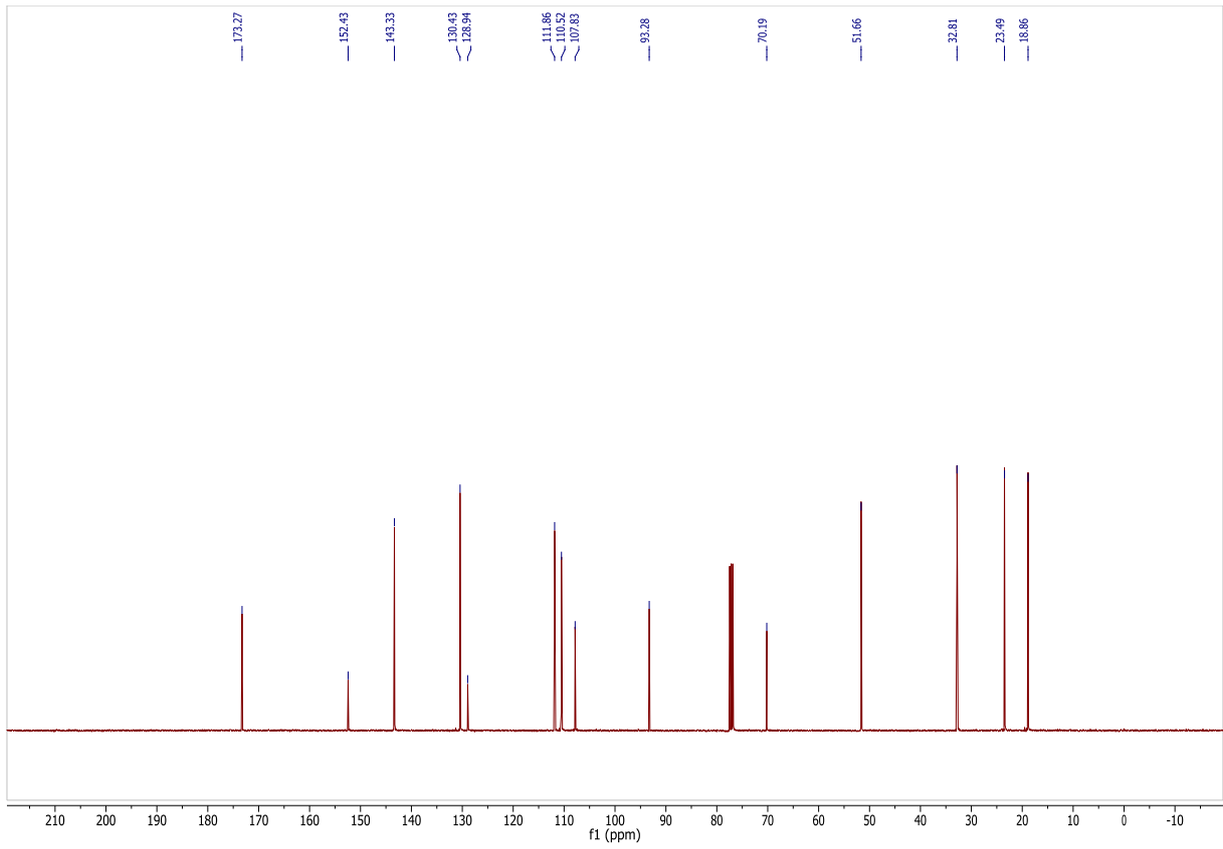
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560 NMR spectra

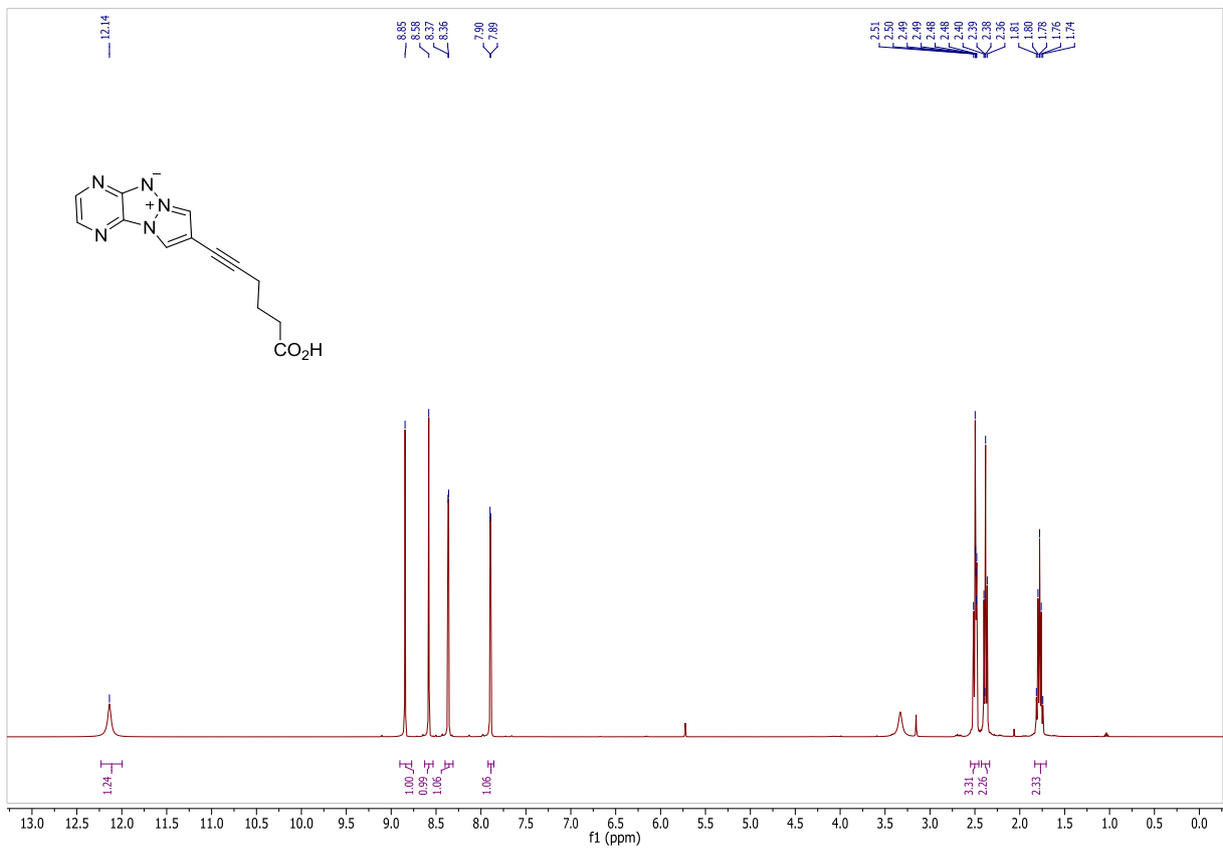


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