

# **Saponite-anthocyanin pigments: Slipping between the sheets**

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# **To cite this version:**

Luciano Clécio Brandão Lima, Fabrícia Castro-Silva, Edson Cavalcanti Silva-Filho, Maria Gardennia Fonseca, Maguy Jaber. Saponite-anthocyanin pigments: Slipping between the sheets. Microporous and Mesoporous Materials, 2020, 300, pp.110148.  $10.1016/j.micromeso.2020.110148$ . hal-02887318

# **HAL Id: hal-02887318 <https://hal.sorbonne-universite.fr/hal-02887318>**

Submitted on 2 Jul 2020

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## **ABSTRACT**

#### 

16 The present work describes the synthesis and characterization of hybrid materials based on β-cyclodextrin (β-CD) and cetyltrimethylammonium bromide 18 (CTAB) intercalated into saponite (SAP) and a commercially available **powder** of anthocyanin dye, Crystal Red Grape (RG). The interactions between the organic dye guest and the organo-clay host were investigated by X-ray diffraction, 21 thermogravimetric analysis, transmission electronic microscopy and  $^{13}C$  solid state nuclear magnetic resonance. The results support complex formation between β-CD and CTAB, their intercalation into the clay interlayer spaces or surface loading, and their 24 interaction with RG (CTAB\_SAP-RG, β-CD\_SAP-RG and β-CD+CTAB\_SAP-RG). The hybrid pigments formed exhibit different colors, enhanced stability against visible 26 light irradiation and basic pH conditions. These **hybrid pigments** are environmentally friendly and can be promising candidates in different application fields.

**KEYWORDS:** Saponite, Natural dye, Adsorption, Hybrid pigment, Color change.

- **1. Introduction**
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 The whole spectrum of colors obtained from natural sources has been employed and improved since the ancient civilization, but their use declined during the development of organic chemistry in  $19<sup>th</sup>$  century, with the "boom" of synthetic dyes and pigments to get away from the limitations of natural dyes, which are subject to the growing seasons of the plants or the life-cycles of the insects [1,2].

 This context has lead the researches into natural dyes to take main focus to historical, archaeological and cultural heritage aspects. However, in the last few decades, the efforts of producing dyes providing long-lasting coloring effect comes at the cost of complex organic structures and hazardous effects, reviving the age-old quest of humankind for natural resources, since the conventional uses of synthetic dyes have posed serious threat to global environment. In this way, the demand for eco-friendly products is switching the trend of using synthetic dyes over the use of naturally occurring colorants [3,4].

 The anthocyanins are the natural pigments that humanity and animals have most consumed from the beginning of time. Anthocyanins are therefore a highly desirable substitute for synthetic food colors, being regarded to be non-toxic and strongly associated to the disease prevention activities found in their sources by opposition to the synthetic food pigments [5–7].

 Anthocyanins are responsible for almost all nuances of blue, red, or purple pigments found in plants, especially flowers, fruits, and tubers. They are water-soluble glycosylated polyhydroxy and polymethoxy derivatives of 2-phenylbenzopyrylium 54 (flavylium) salts that have a common  $C_6-C_3-C_6$  structure consisting of two aromatic rings linked through an oxygenated heterocycle [8,9]. These molecules have an antioxidant and free-radical scavenging properties which promote their recognized anti- inflammatory, antitumor, anti-mutagenic, antiulcer, antiangiogenic, and antiaging activities in addition to its diabetes prevention.

 Despite their use in several fields due to their mentioned properties, anthocyanins have limitations due to their photo- and chemical-stabilities [10]. Anthocyanin dyes are unstable at external environmental conditions such as light, pH, oxygen, temperature [11].

 Therefore, biohybrid compounds that combine these biomolecules and an inorganic counterpart such as clay minerals have been widely investigated to reverse  these limitations by complexation in the host materials [12–14]. This field of research has grown from studies about the first and most famous example of biohybrid, Maya blue, which is composed of a natural clay mineral (palygorskite) and indigo dye of biological origin [1,15,16].

 The strategy to achieve the same stability known to Maya blue has been leading the scientists to try to enhance other organic dyes for several applications. Concerning the studies about anthocyanins, montmorillonite has been investigated as inorganic host materials in recent years [12,17,18], followed by palygorskite [19,20], sepiolite [21], laponite [22] and saponite [23]. However, no study explored the stability enhancement for anthocyanin after adsorption on modified clay.

 Among the many different types of clay minerals, saponite is a 2:1 trioctahedral 76 phyllosilicate belonging to smectite group, which shows interesting properties due to their intercalation/exfoliation characteristics, large specific surface area, surface acidity, cation exchange capacity, thermal stability, biocompatible, and non-toxic properties [24].

 The structure of saponite is composed of a magnesium octahedral sheet sandwiched between two silicon tetrahedral sheets via sharing oxygen. Substitution of  $S_1^{3+}$  cations by  $Al^{3+}$  cations in the tetrahedral layer confers a negative charge generally compensated by alkaline or alkaline earth cations located in the interlayer space. The 84 general formula of saponite can be expressed as  $M_{x}^{+}[Si_{4-x}Al_{X}][Mg_{3}]O_{10}(OH)_{2} \cdot nH_{2}O$ , 85 where M is the exchangeable interlayer cation, x  $(0.2 \le x \le 1.2)$  is the fraction of aluminum present in Si-O tetrahedral sheets, and n is the number of water molecules [25,26].

 Due to its negatively charged surface, saponite can be modified through the replacement of the interlayer inorganic cations by organic ones. Structures with organic cations, for example with quaternary ammonium cations such as hexadecyltrimethylammonium bromide (CTAB), have been extensively studied to provide surfactant-modified clays which can acquire hydrophobic and organophilic characteristics via the interlayer exchange process. The characteristics of this composite make it a promising candidate for the incorporation of dyes and aromatic compounds [13,27–30].

 Moreover, surface modification of clay minerals with biopolymers such as starch, cellulose, chitosan and cyclodextrin was also reported for several applications [31–34]. It can be noticed that the interaction between polysaccharides and

 anthocyanins has shown to be especially important to improve the stability of the latter [35,36]. Among the wide range of biopolymers, cyclodextrin has gained prominence due to its ability to form non-covalent inclusion complexes due to the adaptable hydrophobic tridimensional cavity. Cyclodextrin (CDs) are cyclic D-glucopyranose oligomers, which the most common are knows to have six, seven, or eight glucose units 104 linked by 1,4-α-glucosidic bonds, termed as  $\alpha$ -,  $\beta$ -, and γ-CDs, respectively [37,38].

 In this work, we report on the synthesis and characterization of hybrid pigments based on saponite, as well as its equivalent modified with β-cyclodextrin (β-CD) and 107 hexadecyltrimethylammonium bromide (CTAB), and a commercially available powdered of anthocyanin dye, Crystal Red Grape (RG). The interactions and structural characterizations were also carried out by different techniques to study the chemical-and photo-stabilities of the obtained pigments.

#### **2. Materials and methods**

# **2.1. Materials**

 [Anthocyanin](https://www.sciencedirect.com/topics/chemistry/anthocyanin) source was a Crystal Red Grape (RG) donated by San Joaquin Valley Concentrates (Fresno, CA, USA). Hexadecyltrimethylammonium bromide (CTAB), β-cyclodextrin (β-CD), citric acid, sodium citrate, sodium hydroxide, hydrochloric acid, and other applied chemicals were purchased from Aldrich or Sigma-Aldrich, all with an analytical grade and used without any previous purification.

- **2.2. Saponite synthesis**
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 For the synthesis of sodium saponite, the reagents were mixed in the following 123 order according to the following theoretical formula:  $\text{Na}_{0.3}(\text{Si}_{3.7}\text{Al}_{0.3})\text{Mg}_{3}\text{O}_{10}(\text{OH})_{2}$ : deionized water, hydrofluoric acid (40%, wt), sodium acetate (99%, wt), magnesium acetate tetrehydrate (99%, wt), basic aluminum acetate (19%, wt) and silica (Aerosil 130). The resulted hydrogels were aged under stirring at room temperature for 2 h and 127 then were autoclaved at 300 °C, 90 bars for 6 h. The autoclaves were cooled to room temperature and the products were washed thoroughly with distilled water and 129 centrifuged. The solids were then dried at 50 °C for 24 h [26].

#### **2.3. Synthesis of inclusion complex β-CD+CTAB.**

 The inclusion complex formed by reaction of β-CD with CTAB was prepared according to a method previously described (Yei et al., 2005) with modifications. Solutions of CTAB (5.59 mmol) and β-CD (16.8 mmol) in water (80 mL) were 136 prepared and mixed at room temperature. The mixture was stirred at 70 °C for 8 h and then it was left to stand at room temperature overnight. The mixture became turbid and the complex was obtained as a white crystalline precipitate. The precipitated product was collected after centrifugation and dried at 70 °C. The complex was then washed several times with water to remove any uncomplexed β-CD and CTAB. Finally, the 141 powder was dried at 60 °C for 24 h.

#### **2.4. Synthesis of hybrids organo-clay composites**

 Following the method described in [39] with modifications, a suspension of saponite (SAP, 5 g) in distilled water (250 mL) was stirred overnight in a 500 mL flask. The organic molecules (β-CD, CTAB or β-CD+CTAB inclusion complex; 2 g) were dissolved in 10 mL of 1 N HCl solution and added dropwise at room temperature to the stirred aqueous solutions of SAP. After stirring the mixture for 3 h, the white precipitate was collected by centrifugation, washed with water until no bromide ion could be 151 detected by an aqueous AgNO<sub>3</sub> solution, and then dried at 60  $\degree$ C for 24 h. The O-SAP 152 materials obtained are named β-CD\_SAP, CTAB\_SAP and β-CD+CTAB\_SAP.

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- **2.5. Adsorption studies**
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156 Anthocyanin powder (RG) was dissolved in  $0.020$  mol $L^{-1}$  citric acid buffer 157 solution (sodium citrate, citric acid at pH 3.0) to prepare a 5000 mg $L^{-1}$  solution that was used as stock solution.

# **2.5.1 Effect of contact time on the adsorption**

 The kinetics adsorption of anthocyanin on the materials were carried out as 163 follows: 20.0 mg of saponite (SAP) was added to 20.0 mL of 1000.0 mg $\cdot$ L<sup>-1</sup> dye solution and reacted for a time range of 2-240 minutes under mechanical stirring at 150 rpm and 25 ºC at pH 3. The samples were then centrifuged and equilibrium dye 166 concentrations were determined by UV-visible spectrophotometer (Model: Ocean 167 optics, HR2000) at 525 nm.

168 The quantity (moles) of the dye fixed in the adsorbate  $(a_{\alpha})$  was determined by 169 the Eq. $(1)$ ,

$$
q_e = \frac{(C_o - C_e)^* V}{m} \tag{1}
$$

171 where  $C_0$  and  $C_e$  are the initial and equilibrium concentrations in solution (mg $\cdot L^{-1}$ ), 172 m is the mass of the material in g and V is the volume of the solution used in mL.

173 Experimental results were adjusted to the pseudo-first [40], pseudo-second order 174 [41] and Elovich [42] adsorption kinetic models as described the following Eqs. (2)– 175 (4):

176

178

170

177 Pseudo-first order: 
$$
\ln(q_{e, \exp} - q_t) = \ln q_{e, \text{cal}} - K_1 t
$$
 (2)

Pseudo-second order: 
$$
\frac{t}{q_t} = \frac{1}{K_2 q_{e, cal}} + \frac{1}{q_{e, cal}} t
$$
(3)

Elovich: 179

$$
q_t = \beta(\ln \alpha \beta) + \beta(\ln t) \tag{4}
$$

180 Where  $q_e$  and  $q_t mg \cdot g^{-1}$  are the adsorption capacities at the equilibrium and in a 181 time t (min) respectively. In the Eq. (4)  $\alpha$  (mg⋅g<sup>-1</sup> min<sup>-1</sup>) and  $\beta$  (g⋅mg<sup>-1</sup>) are the initial 182 adsorption rates and Elovich constant related to the extent of surface coverage and also 183 to the activation energy involved in chemisorption, respectively.

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# 185 **2.5.2 Effect of initial concentration on the adsorption**

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 To investigate the influence of anthocyanin concentration on adsorption, 20 mg of saponite (SAP) was added to 20 mL of dye solution at the different initial dye 189 concentration at the 100 to 5000 mg $\cdot$ L<sup>-1</sup>. The mixture was stirred for 1 h at room temperature and then centrifuged to separate the adsorbent. The concentration of dye in supernatant was analyzed using UV-visible spectrophotometer as described in previous subsection.

193 Experimental results were adjusted to Langmuir [43], Freundlich [44] and 194 Temkin [45] models following the Eqs. (5)–(7):

196 Langmuir: 
$$
\frac{C_e}{q_e} = \frac{1}{K_L q_m} + \frac{C_e}{q_m}
$$
 (5)

197 Freundlich: 
$$
\ln q_e = \frac{1}{n_F} \ln C_e + \ln K_F
$$
 (6)

198 Temkin: 
$$
q_e = \frac{1}{n_T} \ln K_T + \frac{1}{n_T} \ln C_e
$$
 (7)

199 Where C<sub>e</sub> (mg<sup>-</sup>L<sup>-1</sup>) is the dye equilibrium concentration,  $q_e$  (mg<sup>-</sup>g<sup>-1</sup>) is the dye adsorbed 200 amount on solid/liquid interface,  $q_{max}$  (mg·g<sup>-1</sup>) is the maximum removal to form a 201 monolayer of the dye on surface,  $K_L$  (L $\cdot$ mg<sup>-1</sup>) is the Langmuir constant.

202 In the Freundlich model  $K_F$  (mg-g<sup>-1</sup>) (mg- $L^{-1}$ )<sup>-1/n</sup> and n<sub>F</sub> are the Freundlich 203 constant and a factor which are related to capacity and intensity of the adsorption, 204 respectively. For Temkin model,  $n_T$  is the constant related to adsorption energy 205 (J·mol<sup>-1</sup>), K<sub>T</sub> (L mg<sup>-1</sup>) is the Temkin isotherm constant, R is the gas constant (8.314) 206  $\text{J} \cdot \text{mol}^{-1} \cdot \text{K}$ ) and T is the temperature (K).

207

## 208 **2.6 Synthesis of the hybrid pigments**

209

210 1 g of SAP or each O-SAP were dispersed in 100 mL of anthocyanin (RG) citric 211 acid buffer solution at pH 3 (the RG concentration was 100 mg $L^{-1}$ ) and was left under 212 stirring for 1 h. The samples were then centrifuged and dried at 50  $\degree$ C for overnight. The 213 hybrid pigments samples are named SAP-RG,  $\beta$ -CD SAP-RG, CTAB SAP-RG and  $\beta$ -214 CD+CTAB SAP-RG.

215

# 216 **2.7. Characterizations.**

217

218 X-ray diffraction were recorded using D8 Advance Bruker-AXS Powder X-ray 219 diffractometer with CuKa radiation ( $\lambda = 1.5405$  Å). XRD patterns were performed 220 between 4-70° (2 $\theta$ ) with scan rate of 0.5 deg<sup>-</sup>min<sup>-1</sup>.

221 Infrared analyzes were performed on Agilent Cary 630 FTIR spectrometer using 222 an Agilent diamond Attenuated Total Reflectance (ATR) technique mode, with a 223 spectral resolution > 2 cm<sup>-1</sup> and 32 scans. Spectra were acquired by Microlab FTIR 224 Software (Agilent Technologies) between 4000 and  $650 \text{ cm}^{-1}$ .

 Thermogravimetric analyses were carried out using a TA Instrument SDT Q600 226 analyzer. The heating rate was of  $5^{\circ}C \cdot \text{min}^{-1}$  from 25 $^{\circ}C$  to 1000 $^{\circ}C$ , under dry air flow 227 of  $10 \text{ mL} \cdot \text{min}^{-1}$ , and using alumina pan.

 TEM study of the samples was performed on a JEOL 2010 microscope, 200 kV LaB<sub>6</sub> coupled Orius camera, from Gatan Company. Samples in the form of bulk powders were suspended in ethanol and then deposited on 400 mesh copper grids covered with an ultrathin carbon membrane of 2–3 nm thickness.

 MAS NMR spectra were obtained on a Bruker Avance III spectrometer 233 equipped with a 4 mm H-X MAS probe, operating at frequency of 500.17 MHz  $(^1H)$ , 234 125.77 MHz  $(^{13}C)$ . Chemical shifts were calibrated using the carboxyl signal of 235 adamantane (38.52 ppm) for  ${}^{13}C$ .

236 The <sup>13</sup>C Cross-Polarization spectra were acquired with a MAS rate of 14 kHz, a 237 ramp-CP contact time of 1 m s and a 1 s recycle delay and with a  ${}^{1}H$  decoupling spinal. Over an acquisition time of 40 m s, the number of scans to obtain the spectra depends on the S/N obtained for each sample. Spectra were processed with a zero-filling factor of 2 and with an exponential decay corresponding to a 25 Hz line broadening in the transformed spectra. Only spectra with the same line broadening are directly compared.

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# **2.8. Chemical and photo-stabilities of the hybrid pigments**

 Chemical stability of anthocyanin molecules loaded on the matrix was verified by exposure of the hybrid pigments to basic and acidic conditions in a desiccator 248 containing aqueous NH<sub>4</sub>OH or HCl. At first, the sample was exposed to NH<sub>3</sub> atmosphere. After exposure to basic environment, the samples were transferred into a 250 desiccator filled with HCl atmosphere. The sample was exposed to  $NH_3$  and HCl atmospheres, sequentially and repeatedly [23].

 The photo-stabilities were evaluated by exposure of solid pigments to white light irradiation for 192 h, using a LED lamp set to provide 100 Klx of illumination intensity, in which this time is equivalent to approximately 30 years of exposure in ambient light conditions. To study the effects of irradiation under oxidant and inert atmospheres, the pressed pigments (into pellets) were placed in a desiccator filled with air or nitrogen.

 The absorbance, reflectance and CIE (Commission Internationale de L'Eclairage) parameters were obtained from an Ocean Optics Halogen and Deuterium Light Source HL-2000-FHSA device as incident light beam and ocean optics USB4000 detector for acquisition. Ocean Optics QP400-1-UV-VIS fiberglass was used to link these devices. For each acquisition, the optimum signal was obtain with an average of 100 scans. The diffuse reflectance (R) converted into equivalent absorption coefficient F(R) using Kubelka–Munk equation (Eq. (1)) [46].

- 
- 
- 

 $F(R) = \frac{(1-R)^2}{2R}$ 265  $F(R) = \frac{(1 - R)}{2R}$  (1)

 The "Commission Internationale of l'Eclairage" CIE 1976 color space system was applied to evaluate the color of the pigments. Measurements were done on pressed 269 pellets samples as function of  $L^*$ ,  $a^*$  and  $b^*$  coordinates. The differences of colors between unexposed and exposed samples were calculated by  $\sqrt{((\Delta L^*)^2 + (\Delta a^*)^2 + (\Delta b^*)^2)}$  equation.

## **3. Results and Discussion**

#### **3.1 X-Ray Diffraction**

276 XRD patterns of the pure CTA $\overline{B}$  shows typical reflections at 6.83 $^{\circ}$ , 10.23 $^{\circ}$ , 13.63°, 16.90°, 20.55°, 21.47°, 23.83°, 37.95°, 40.18 (JCPDS 00-030-1746) (Fig. 1A.). Characteristic reflections of β-CD are observed in 4.53°, 9.04°, 12.70°, 13.58°, 14.75°, 17.94°, 18.86°, 21.28°, 22.83°, 24.38°, 25.78°, 27.18°, 32.04° and 34.92° in agreement with previous works [47,48]. The XRD pattern of the product resulting from the mixture 281 of cyclodextrin host and CTAB ( $\beta$ -CD+CTAB inclusion complex) is not a simple mixture of the two precursors but a new crystalline phase [37]. The differences observed between the patterns of their precursors were: i) the absence of the sharp and intense 284 reflections of β-CD and CTAB (i.e.  $4.53^\circ$ ,  $9.04^\circ$  for β-CD and  $16.90^\circ$ , for CTAB), ii) other ones shifted (i.e. 14.75°, 18.86° for β-CD and 13.63°, 20.55°, 21.47° for CTAB) and iii) presence of medium intense reflections after complexation (i.e. 14.75°, 18.86° for β-CD and 13.63°, 20.55°, 21.47° for CTAB), due to the encapsulation of the surfactant into the nano-hydrophobic cavities of β-CD [38,39].



291 **Fig. 1.** X-ray diffractograms of (A) CTAB, β-CD and β-CD+CTAB; (B) SAP,

292 CTAB SAP, β-CD SAP and β-CD+CTAB SAP; (C) SAP-RG, CTAB SAP-RG, β-

293 CD SAP-RG and β-CD+CTAB SAP-RG.

294

295 The XRD patterns of the sodium saponite  $(Fig. 1B)$  showed typical (hkl) reflections of the clay and confirmed the success of the synthesis. The reflection at 7.66° (2θ) was associated to the basal spacing, *d(001)* of 1.16 nm, which can be correlated with a thickness of monohydrated sodium saponite layer [49].

299 After loading of the organic molecules in SAP, the initial basal spacing increased 300 to 1.62 nm in CTAB\_SAP, 1.42 nm in β-CD\_SAP and 1.46 nm in β-CD+CTAB\_SAP 301 ( $\overline{Fig.}$  1B).

302 Different scenarios can be considered: for CTAB SAP sample, the process 303 involves cation exchange between  $CTA^+$  and  $Na^+$ . The orientation of the organic chain 304 in the interlayer space was suggested considering the dimensions of  $CTA<sup>+</sup>$  moieties 305 (Fig. 2) and the free interlayer space, which value was  $0.65$  nm for CTAB SAP, corresponding to the difference between the SAP layer thickness and the basal spacing 1.62 nm. Therefore, the results are consistent with a monolayer arrangement of the organic molecules, where the ammonium groups tethered to the clay surface, and the alkyl chains lay parallel to the layers [50].

310 In the β-CD\_SAP sample, since the  $d_{001}$  didn't increase significantly considering the size of β-CD, one plausible hypothesis is the adsorption of the organic part on the surface of the layers and maybe on the edges via hydrogen bonding.

313 For β-CD+CTAB\_SAP, a partial intercalation of the β-CD+CTAB complex can be proposed by ion exchange reaction between interlayer sodium cations of the SAP and positively charged portion of the inclusion complex.

 For SAP and β-CD-SAP samples loaded with anthocyanin, the (001) reflexion was broader with a slightly increase of *d(001)* values, suggesting maybe a heterogeneity 318 in the layer stacking due to dye intercalation. For CTAB\_SAP after dye loading, the (001) reflexion was also broader but with a slight decrease of *d(001*) values. In this case, a 320 partial release of of CTA<sup>+</sup> intercalated that convert the anthocyanin from the flavylium  $\cdot$  cation  $(AH_2^+)$  to quinoidal base form  $(AH)$  can occur, attenuating the electrostatic 322 repulsions by forming the pair CTA+/AH on the SAP surface or into interlayer space. This can also explain the blue color of the pigment and the heterogeneity in the layer stacking. However, no change was observed in the *d<sup>001</sup>* values of β-CD+CTAB-SAP 325 samples after dye loading suggesting RG loading on the surfaces. Fig. 2 illustrates these all possibilities.





 **Fig. 2.** Schematic representation of the (A) inclusion complex β-CD+CTAB and (B) hybrid formation.

## **3.2 Morphological analysis**

# TEM micrographs showed layers with alternate dark and bright fringes allowing the measurement of interplanar distances.

338 For pristine SAP, an interlayer distance of  $1.19\pm0.11$  nm was obtained (Fig. 3A). 339 In O-SAP samples, the interlayer distance increased to  $1.57\pm0.19$  nm,  $1.41\pm0.13$  nm and 340 1.43±0.13 nm for CTAB SAP, β-CD SAP and β-CD+CTAB\_SAP respectively (Fig. 3 C;E and G). After loading the anthocyanin molecules, the *d<sup>001</sup>* spacing were 1.39±0.19, 342 1.49±0.20, 1.51±0.19 and 1.43±0.19 nm for SAP-RG, CTAB\_SAP-RG, β-CD\_SAP-RG 343 and  $\beta$ -CD+CTAB SAP-RG, respectively (Fig. 3 B;D;F and G). The values corroborated with the XRD data.





 **Fig. 3.** TEM images of (A) SAP; (B) SAP-RG; (C) CTAB\_SAP; (D) CTAB\_SAP-RG; 347 (E) β-CD\_SAP; (F) β-CD\_SAP-RG; (G) β-CD+CTAB\_SAP and (H) β-348 CD+CTAB\_SAP-RG.

350 **3.3 FTIR**

351

352 The infrared spectra of pure β-CD indicated its main absorptions at 3280 and 2925 353 cm<sup>-1</sup> (Fig. 4A) attributed to *v*(O−H) and *v*(C−H), as well as the bands at 1152, 1077, 354 1023, and 940 cm<sup>-1</sup> (Fig. 4B) assigned to  $\delta$ (O−H), *ν*(C−C), and the α-1,4 linkage 355 skeletal vibration, respectively [51,52].

356 The main bands associated to CTAB occurred at 3016, 2916 and 2847  $cm^{-1}$  and 357 were attributed to the  $N(CH_3)$ <sub>3</sub> asymmetric vibration, C-H and CH<sub>2</sub> asymmetric and 358 symmetric stretching, respectively. Other bands in the range  $1460-1488$  cm<sup>-1</sup> were 359 assigned to  $[N(CH_3)_3]$  and  $(CH_2)$  bending modes, while C-N vibrations were detected at 360 911 cm<sup>-1</sup> and 963 [53,54].

361 The formation of the  $(\beta$ -CD)-(CTAB) complex attenuated the surfactant 362 absorptions, due to the seven repeating units in the β-CD, the spectrum of the inclusion 363 complex was largely dominated by the bands of the host portion, but even so, the 364 spectra in Fig.  $4A$  and Fig.  $4B$  display some differences in comparison to spectra of 365 their precursors that suggest the inclusion of CTAB into the β-CD cavity [51]. The band 366 at 3280  $\text{cm}^{-1}$  assigned to O-H stretching may correspond to the new hydrogen bonds in 367 the complexes. The region between 2800-2980 cm<sup>-1</sup> is attributed to  $v(C-H)$  in β-368 CD+CTAB: it shows the contributions of the (CH) group in  $\beta$ -CD, shifted from 2925 369 cm<sup>-1</sup> to 2919 cm<sup>-1</sup>, and also a weak band at 2852 cm<sup>-1</sup> related to the contribution of 370 (CH<sub>2</sub>) stretching of CTA<sup> $+$ </sup>, which appears as a shoulder probably due to the attenuation providing by the inclusion in the  $\beta$ -CD cavity. The bands at 1460-1488 cm<sup>-1</sup> 371 372 disappeared also suggesting the inclusion of the aliphatic chains in β-CD cavity, since 373 these bands are **mostly** associated to the  $(CH<sub>2</sub>)$  bending [55].

 $374$  In Fig.  $4C$  are shown the spectra of pristine SAP and the organoclays formed by 375 intercalation of the surfactant, cyclodextrin and the inclusion complex represented by 376 the CTAB\_SAP,  $\beta$ -CD\_SAP,  $\beta$ -CD+CTAB\_SAP respectively.

377 For the SAP sample, the sharp and weak band at 3677 cm<sup>-1</sup> and the region 378 between  $3000-3600$  cm<sup>-1</sup> are assigned to -OH vibrations modes. The band at 1633 cm<sup>-1</sup> 379 is attributed to –OH water bending. The band at 980 cm<sup>-1</sup> was assigned to the Si-O-Si 380 stretching [4,56].

381 After organomodification (Fig. 4C), the changes in CTAB\_SAP spectra that 382 confirm the presence of organic molecules are the new bands at 1478, 2847 and 2925,

 $cm^{-1}$ , which were assign to the characteristic vibrations of CTAB alkyl chain. Concerning the β-CD\_SAP spectrum, the different shape of the OH stretching and bending modes compared to the ones present in the pristine SAP suggests the contributions of OH groups brought from polysaccharide. A weak band at about 2924  $\cdot$  cm<sup>-1</sup> is attributed to v(CH). Finally, the  $\beta$ -CD+CTAB\_SAP sample shows more intense bands at 2853 and 2925 attributed to the CTAB alkyl chain in the β-CD+CTAB inclusion complex, in addition to the similar characteristics observed for β-CD\_SAP. The Fig. 4D presents the spectra of the natural dye (RG) and the hybrid pigments obtained. In the RG spectrum, the main bands related to the aromatic and phenolic 392 structure of the anthocyanin can be seen in the following regions :  $1250-1370$  cm<sup>-1</sup> and 393 1350-1530 cm<sup>-1</sup> attributed to C-O and C-C modes, respectively [17,18], 1550-1700 394 cm<sup>-1</sup> assigned to O-H bending and C=O stretching; and 3000-3700 cm<sup>-1</sup> associated with 395 stretching vibrations of O-H.

396 The presence of anthocyanin in the hybrids can be verified by the distinguished 397 signal of –OH stretching and bending modes due to the contributions of anthocyanin 398 bands and the appearance of the corresponding peaks in the region of 1320-1450 cm<sup>-1</sup>, 399 emerging in the FTIR spectrum of the hybrids.





401 **Fig. 4.** FTIR spectra of (A-B) CTAB, β-CD and the inclusion complex β-CD+CTAB; 402 (C) SAP and the organo-clay formed after incorporate CTAB, β-CD and β-CD+CTAB 403 molecules; (D) hybrid pigments formed after adsorption of anthocyanin dye molecule in 404 SAP, CTAB\_SAP,  $\beta$ -CD\_SAP and  $\beta$ -CD+CTAB\_SAP.

405

# **3.4 <sup>13</sup>** 406 **C MAS NMR**

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408 <sup>13</sup>C MAS NMR spectra of β-CD+CTA**B** and their precursors are presented in Fig. 409 5A. The signals of  $CTAB$  occur in the chemical shift range of 10–70 ppm. The 410 resonances at 16 ppm, 24 ppm and the complex signals that **appear** at 27-38 ppm are 411 assigned to the terminal methyl group, the final methylene near to terminal methyl and 412 the internal methylenes of the alkyl chain, respectively. The resonances at 55 ppm and 413 63 ppm are related to the N-CH<sub>3</sub> of the head-group and the first methylene group 414 directly bonded to nitrogen, respectively [57]. The spectrum of  $\beta$ -CD shows the 415 resonances of C1, C4 and C6 as multiple peaks that are spread over the chemical shift 416 range of 100-107 ppm, 80-86 ppm and 57-65 ppm, respectively. It occurs due to 417 asymmetric glucopyranosyl conformations. The chemical shift range of 68-77 ppm is 418 related to the signals of C2, C5 and C3 [58].



**Fig.** 5. <sup>13</sup>C MAS NMR spectra of (A) β-CD+CTA**B** and their precursors, (B) RG and 422 SAP-RG, (C) CTAB\_SAP-RG,  $\beta$ -CD\_SAP-RG and  $\beta$ -CD+CTAB\_SAP-RG.

423

424 The spectra of  $\beta$ -CD+CTAB complex (Fig. 5A) shows differences in comparison 425 with the precursors. The resonances related to  $β$ -CD carbons appeared without 426 significant shifts but as sharp singlet peaks after complexation, which implies that the β-427 CD unit adopts a symmetric cyclic conformation with the inclusion of CTAB alkyl 428 chain. Changes in chemical shift confirm the interaction, as occurred in  $C(4)$  and  $C(6)$ 429 glucopyranosyl monomer signals. For the CTAB signals, the peak assigned to terminal 430 methyl group is absent, the peaks of methylene shifted from 24 ppm to 23 ppm and the 431 complex signals related to internal methylenes shifted from the region of 27-38 ppm to 432 25-34 ppm. These changes indicate the inclusion of alkyl chain into β-CD hydrophobic 433 cavity [39].

434 The <sup>13</sup>C MAS NMR spectra of the RG and the RG-SAP are presented in  $\frac{Fig. 5B}{9}$ , 435 C. In the RG spectrum: the signal at 56 ppm is attributed to the  $OCH_3$  groups bounded 436 to the aglycone portion in  $C(3')$  and  $C(5')$  and the signals between 58-85 ppm are 437 assigned to the chemical shifts of the glycosidic portions. The signals above 90 ppm are 438 attributed to the aromatic carbons in flavylium cation, as indicated in  $\frac{Fig. 5B}{[59,60]}$ .

439 After incorporation of RG in SAP, any peaks in  $^{13}$ C MAS NMR spectrum of the 440 dye molecule was observed in the hybrid probably dominated by the other 441 contributions. On the other hand, three signals at 45 ppm, 76 ppm and 179 ppm were 442 assigned to the citric acid, which was also incorporated during the adsorption process. 443 The same behavior is observed for CTAB SAP-RG, β-CD SAP-RG and β-444 CD+CTAB SAP-RG  $(Fig. 5C)$  hybrids that showed peaks assigned to the organic 445 precursors and also three signals of the citric acid (CA).

446

## 447 **3.5 Thermal analyses**

448

449 In Fig. 6A, the first step of degradation at  $25{\text -}238$  °C (mass loss about 8%) for raw 450 SAP was assigned to the loss of physisorbed water and dehydration of interlayer 451 cations. The second event at 640-859 °C is due to dehydroxylation of SAP with a 452 weight loss of 3% [24,27]. DTG of SAP-RG shows an additionnal weight loss of 7% in 453 the region 210-623  $\degree$ C assigned to the degradation of the incorporated anthocyanin.

454 The weight losses in the second thermal event of O-SAP samples before the RG 455 adsorption ( $\overline{Fig.}$  6B-D, Table 1) were 2%, 6% and 30% in β-CD SAP, β-456 CD+CTAB SAP and CTAB SAP respectively. After dye loading, the organic mass 457 losses increases to 6%, 8% and 32% for β-CD\_SAP-RG, β-CD+CTAB\_SAP-RG and 458 CTAB SAP-RG, respectively.







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-CD\_SAP and  $\beta$ -CD\_SAP-RG; (D)  $\beta$ -CD+CTAB\_SAP and  $\beta$ -CD+CTAB\_SAP-RG.

463

<b>Sample</b>	<b>Temperature</b>	<b>Mass loss</b>
	$(^0C)$	(%)
<b>SAP</b>	25-238	8.0
	640-859	3.5
SAP-RG	25-210	13.0
	210-623	7.2
	623-878	3.6
CTAB_SAP	25-117	2.4
	117-710	29.5
	710-878	2.9
CTA <mark>B</mark> _SAP-RG	25-125	2.4
	125-702	32.2
	702-888	2.5
$\beta$ -CD_SAP	25-174	15.6
	174-319	2.5
	685-864	2.9
$\beta$ -CD_SAP-RG	25-182	9.3
	182-600	5.9
	600-885	3.6
β-CD+CTAB_SAP	25-180	9.6
	180-670	6.1
	670-892	3.1
$\beta$ -CD+CTAB_SAP-	25-181	6.5
RG		
	181-667	7.9
	667-885	3.0

465 **Table 1.** Temperature range and percentages of mass loss observed in DTG curves.

467 **3.6 Adsorption studies**

468



470

471 Adsorption kinetics studies were carried out and are depicted in Fig. 7.



 **Fig. 7.** Influence of the contact time on the RG adsorption on SAP; CTAB\_SAP; β-475 CD SAP; β-CD+CTAB SAP samples.

 The adsorption was fast during the first few minutes, mainly for raw saponite. The equilibrium times were reached after 10 min and resulted in dye adsorption capacity 479 close to 910 mg $\cdot$ g<sup>-1</sup> for SAP. For organo-saponite samples, the equilibrium times were 480 achieved after 30, 50 and 60 min for CTAB\_SAP, β-CD\_SAP and β-CD+CTAB\_SAP respectively, with lower maximum dye adsorption capacity than the pristine saponite. It is an indicative of the presence of different adsorption sites on organo-saponites.

 In order to better understand the adsorption kinetics, the pseudo-first-order, pseudo-second-order and Elovich models were applied to experimental data and the resulted kinetic parameters of the fittings are presented in Table 2.

 All systems were well-fitted to the pseudo-second order model, since that higher 487 R<sup>2</sup> coefficient and low difference between the  $q_{e,exp}$  and  $q_{e,theor}$  were observed. Same results were obtained in other works [27,61].





# 495 **3.6.2 Effect of the initial dye concentration**





 **Fig. 8.** Influence of the RG initial concentration on the adsorption process of SAP; 502 CTA**B**\_SAP; β-CD\_SAP; β-CD+CTA**B**\_SAP.

 The saturation occurred at high RG initial concentration and resulted in high 505 adsorption capacity for SAP-RG,  $(3500 \text{ mg} \cdot L^{-1} \text{ RG}$  initial concentration and adsorption 506 capacity about 2170.0  $mg-g^{-1}$ ). However for organo-saponite samples, equilibrium 507 occurred at 3000.0, 1000.0 and 2500 mg $L^{-1}$  of RG initial concentrations and showed 508 adsorption capacities close to 1230.0, 640 and 470 mg·g<sup>-1</sup> for β-CD\_SAP, β-509 CD+CTAB SAP and CTAB SAP respectively.

 The equilibrium data were adjusted to Langmuir, Freundlich and Temkin models, and the resulting parameters are summarized in Table 3. The values of isotherm correlation coefficients revealed that the experimental data were well-fitted to Langmuir 513 ( $\mathbb{R}^2$  closer to 1) compared with other models. Additionally, the R<sub>L</sub> values calculated 514 from the Langmuir equation suggested that the adsorption was favorable  $(0 < R_L < 1)$  in all cases. Adsorption sites in each sample were uniformly distributed on the surface and a monolayer adsorption took place [65].

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522 **Table 3.** Parameters obtained by fitting the experimental data to Langmuir, Freundlich

523 and Temkin models in the RG adsorption by the organo-saponites.

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## 525 **3.7 Chemical and photo-stabilities of the hybrid pigments**

## 526 **3.7.1 Spectra changes**

527

 The UV-Vis spectra of RG in citric acid solution buffered at pH 3 showed an 529 absorption at 525 nm ( $\frac{Fig. 9A}{P}$ ). The majority of anthocyanin molecules present in RG are malvidin-3,5-diglucoside, therefore the results are in concordance with the literature, that shows absorption around 520 nm for acid solution of anthocyanin [11,12,66].

532 The Fig. 9B shows the possible structural transformations of anthocyanin molecules in aqueous solution. The flavylium is the most stable form of anthocyanin that appears redish and turns to blue after conversion to quinoidal base when pH increases and finally to non- colored chalcone via hemicetal form.

536 Comparing the spectra of RG solution with those of hybrid pigments, a red shift 537 occurred from 525 to 539 nm for SAP-RG, β-CD\_SAP-RG, β-CD+CTA $B$ \_SAP-RG 538 samples and 554 nm for CTAB SAP-RG. The pH was adjusted by the addition of citric 539 acid buffer solution during the hybrid pigment formation. Shifts in the absorptions can

540 be related to electrostatic interactions between the clay mineral layer and the dye and/or 541 by intramolecular interaction between the organic moieties in the respective hybrids.

542 The shift and presence of a broad band in the spectrum of CTAB SAP-RG may 543 indicate the deprotonation of the flavylium cations into deprotonate quinoidal base  $AH<sub>2</sub>$  $544 \rightarrow AH$ , which can be stabilized by electrostatic interaction with the positively 545 charged surfactant in this hybrid system. This behavior may explain the blue color of 546 the hybrid, while the other ones were weakly red.





548 **Fig. 9.** (A) Diffuse reflectance UV–Vis spectra of hybrid pigments and the spectrum of 549 RG citric acid buffer solution pH 3; (B) schematic representation of possible 550 anthocyanin structures in different pHs in aqueous solution.

551

# 552 **3.7.2 Color changes in acid/basic environments**

553

 The hybrid pigments were exposed to acidic or basic atmosphere prepared in a desiccator, in which the samples were submitted to HCl or NH4OH atmosphere. The results upon exposure to acidic and basic vapors were monitored by visible absorption spectroscopy and also by visual changes in their photographs, these results are present 558 in  $\overline{\text{Fig.}}}10$ .



560 **Fig. 10.** Spectra changes after exposure to acidic and basic environments for (A) SAP-561 RG; (B) CTAB SAP-RG; (C) β-CD SAP-RG and (D) β-CD+CTAB SAP-RG. (E) 562 Digital photographs of the color changes of hybrid pigments upon exposure.

564 Colors changes from red to blue after exposure to basic atmosphere  $(NH_3$  from 565 aqueous NH4OH) were observed in the SAP-RG, β-CD\_SAP-RG and β-566 CD+CTAB\_SAP-RG. The process is reversible, as shown in Fig. 10E. The exposure 567 time to change the color was about 10 min. Similar results were obtained in the 568 literature.

569 In  $Fig. 10(A, C and D)$ , the absorption band have a redshift after exposure to basic atmosphere, and return to the same wavenumber, after exposure to basic atmosphere. After several cycles; the same observations on the spectra were noticed. The color change was reversible and repeatable for at least two cycles. The color changes are still observed after the second cycle of exposure, although strong acid and base conditions degraded the pigments.

575 The contribution of quinoidal base in the anthocyanin molecule in CTAB\_SAP-576 RG was more pronounced than in the other hybrids and explains its initial blue color. 577 After the first exposure to basic vapor, the right shift observed in  $\frac{Fig. 10B}{}$  is due to the  conversion of remaining flavylium cations to quinoidal base, which causes a change 579 only in nuance of blue (Fig. 10E). Hybrids being exposed to basic conditions were again submitted to acidic environment, the color change to red and their spectra were similar to the other hybrids and indicated that the anthocyanin molecules became in flavylium cation form. The behavior of color change in the following cycles was also similar to the others hybrid pigments [18,20,23,32].

- The decrease in relative absorbance is 80.4%, 73.8%, 83.2% and 84.3% for SAP-585 RG, CTAB SAP-RG, β-CD SAP-RG and β-CD+CTAB SAP-RG respectively. These data indicate that the CTAB\_SAP-RG has a slightly better chemical stability than the other pigments, since it has lowest decrease of relative absorbance after four cycles of 588 acid basic exposure.
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### **3.7.3 Photostability**

 The photostability of the hybrid pigments were studied under LED light exposure 593 for 192 h under air and nitrogen atmosphere. Fig. 11 depicts the UV-V is spectra.

 The progressive decreasing in the absorbance at maximum wavelength occurred without differences in the spectral patterns, which indicated no simultaneous reaction that could form different derivatives chromophore species during the light exposure. Degradation were most pronounced in the process carried out under air for all hybrids. This behavior is expected since the photodegradation is often considered in relation to 599 the oxidation caused by the evolution of singlet oxygen  ${}^{1}O_{2}$  [12]. The order of fading of 600 the hybrid pigments under air followed, CTAB SAP-RG (75%) >  $\beta$ -CD+CTAB SAP-601 RG (59%) >  $\beta$ -CD\_SAP-RG (37%) > SAP-RG (34%).

602 On the other hand, the **degree** of degradation were about 18% (based on  $A/A0 =$  0.82) for all hybrids under nitrogen atmosphere. The time exposure of the samples confirmed that the RG degradation process was less pronounced. Under nitrogen, reactive species are more difficult to form but the oxidation process remains possible due to the residual water molecules present in the pigments.



609 **Fig.** 11. Relative absorbance at  $\lambda_{\text{max}}$  and evolution of spectral patterns during light 610 exposure for 192 h under air and nitrogen atmosphere for (A), (B) and (C) SAP-RG; 611 (D), (E) and (F) CTAB\_SAP-RG; (G), (H) and (I)  $\beta$ -CD\_SAP-RG; (J), (K) and (L)  $\beta$ -612 CD+CTAB SAP-RG.

- 613 The colors of solid hybrid pigments were evaluated before and after light exposure
- 614 during 192 h (30 years of exposure in a museum). Measurements of  $L^*a^*b^*$  parameters
- 615 were carried out at different light irradiation times  $(Fig. 12)$ .



 **Fig. 12.** Photo-ageing of pigments followed by color change observations over 192 h of light exposure under (A) air and (B) nitrogen atmosphere.

 In solution, the color of RG changes from red in acid conditions to blue in neutral to weak alkaline conditions. However, the difference in color presented by the hybrids results from the different interactions between the host and the guest since the effect of pH on the color was controlled with the citric acid buffer solution. The color for SAP- RG is probably due to the intercalation of RG molecule in the interlayer space of SAP, which stabilizes the red color of flavylium cations. While adsorption in the 628 CTAB SAP-RG sample probably induced stabilization of the RG quinoidal base, leading to a blue color for the hybrid. The color for β-CD\_SAP-RG is close to the one observed in SAP-RG and corroborates with the proposal of intercalation of the RG 631 molecules in hybrids. Finally, the color for  $\beta$ -CD+CTAB SAP-RG was between the 632 colors of β-CD SAP-RG and CTAB SAP-RG, which is suggestive of a minor 633 stabilization of RG quinoidal base than in  $CTAB$  SAP-RG.

 The ΔE\* value is related to total color difference and is indicative of the light 635 stability of the pigments. Under air (Fig. 12A), the  $\Delta E^*$  values of the pigments increased gradually with the increase of the ageing time. Finally, the values reached 637 10.5 $\pm$ 0.9 for SAP-RG, 13.6 $\pm$ 0.6 for β-CD\_SAP-RG, 18.7 $\pm$ 0.2 for β-CD+CTAB\_SAP-638 RG and  $19.3\pm 2.5$  for CTAB SAP-RG after 192 h of light exposure. Lower  $\Delta E^*$ 639 variation in SAP-RG and β-CD\_SAP-RG under air is probably due to the protection of 640 the dye in the interlayer space of the saponite creating an oxygen hindering and 641 stabilizating the pigments under irradiation. Under nitrogen atmosphere,  $\Delta E^*$  values 642 were lower after 192 h of light exposure, the  $\Delta E^*$  values were 2.7 $\pm$ 0.2, 3.3 $\pm$ 0.2, 4.8 $\pm$ 0.5 643 and 5.7 $\pm$ 0.9 for CTAB\_SAP-RG, SAP-RG,  $\beta$ -CD+CTAB\_SAP-RG and  $\beta$ -CD\_SAP-644 RG, respectively. In addition, when the photostabilities under air are compared to 645 photostabilities under nitrogen, the most significant differences were observed for β-646 CD+CTAB SAP-RG and CTAB SAP-RG, which corroborate with the greater 647 exposure of RG molecules to oxygen attack on air atmosphere for these hybrids. The 648  $\Delta E^*$  values are summarized in Table 4.

649

650 **Table 4.** Total color differences (**ΔE\*)** for each sample after 192 h of light exposure 651 under air and nitrogen atmosphere.

<b>Sample</b>	$\Delta E^*$ in air	$\Delta E^*$ in $N_2$
SAP-RG	$10.5 \pm 0.9$	$3.3 \pm 0.2$
CTAB SAP-RG	$19.3 \pm 2.5$	$2.7 \pm 0.2$
$\beta$ -CD_SAP-RG	$13.6 \pm 0.6$	$5.7 \pm 0.9$
$\beta$ -CD+CTAB SAP-RG	$18.7 \pm 0.2$	$4.8 \pm 0.5$

#### 652

#### 653 **4 Conclusion**

654

655 Hybrid pigments were successfully prepared by incorporating anthocyanin into 656 unmodified and modified clay mineral based on synthetic SAP and CTAB and β-CD.

 The hybrid pigments exhibit different colors function of their host-guest interactions. The presence of the quinoidal base form of anthocyanin molecule in 659 CTAB SAP-RG sample is probably more pronounced than in the other hybrids, which explains their initial blue color. The enhanced stability against visible light irradiation and basic environment conditions were brought by the electrostatic interaction between the dye and the respective host materials. The intercalation of the dye molecules into the interlayer spaces of saponite in β-CD\_SAP-RG and SAP-RG protect the dyes from oxygen avoiding oxidation or the formation of other possible reactive oxygen species. 665 Reversibility in color upon exposure to acidic and basic atmosphere is an evidence for a  possible application of the obtained pigments as a sensor to atmospheric acidity, which can be exploited in several cycles if applied in less extreme pH conditions. These dyed materials are environmentally friendly and can be promising candidates in different application fields

# **Acknowledgments**

 We acknowledge the financial support from the CAPES/COFEBUB (Project n° 835/15), CAPES and National Council for Scientific and Technological Development (CNPq, Brazil) for financial support (Grant [310921/2017-1,](javascript:abrirPrestacao() M.G.F, 307460/2016-9, E.C.S.F.). The authors thank the Île-de-France region and CNRS for funding.

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