

Modification, hybridization and applications of saponite: An overview

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1	An Overview of Modification, Hybridization and Applications of Saponite
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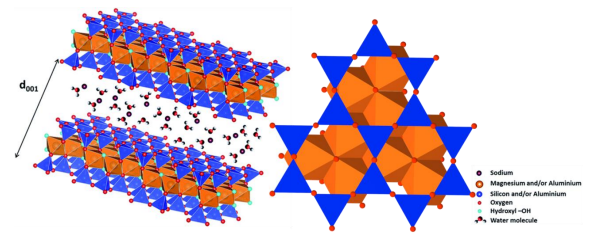
26 Abstract

27 Modification of saponite (Sap) by surface engineering and intercalation chemistry introduces guest 28 species into the structure of Sap and enhances the functionalities of the resultant Sap-based hybrids or 29 composites. This review summarizes and evaluates latest scientific advances in the strategies for 30 surface engineering, intercalation and hybridization of Sap, the insights into the relevant mechanisms, 31 and the properties and applications of the resultant Sap-based materials. Studies have indicated that Sap 32 can be inorganically modified by acid activation, inorganic cation exchange, pillaring, and adsorption. 33 The methods of preparing organo-saponite (OSap) hybrids can be categorized as follows: 1) 34 exchanging the inorganic cations in the interlayer space of Sap with organic cations; 2) covalent 35 grafting of organic moieties or groups onto the surface of Sap; 3) intercalating polymer into the 36 interlayer space of Sap by solution intercalation, and melt mixing or in situ polymerization. 37 Organic-inorganic modified Sap can be made through the reactions between organic species and 38 inorganic-modified Sap, or by the combination of inorganic species with organic-modified Sap. Modified Sap exhibits exceptional thermal stability, surface acidity, optical effects and adsorption. As 39 40 such, the modified Sap can be used for optical materials, adsorbents, catalysts and clay/polymer 41 nanocomposites (CPN). Literature survey suggests that future studies should place emphasis on 42 optimizing and scaling up the modification of Sap, probing the thermodynamics, kinetics and 43 mechanisms of the modification of Sap, endowing Sap with novel functionalities, and accordingly 44 advancing the practical applications of the resultant Sap-based materials.

Keywords: 45 Saponite; Modification; Hybrid; Adsorbent; Catalyst; Saponite/polymer 46 nanocomposite. 47 48 49 50 51 52 53 54 55

56 **1. Introduction**

57 Saponite (Sap) is a 2:1 trioctahedral clay mineral and belongs to the smectite group. Typically, a layer of Sap is composed of two tetrahedral -O-Si-O- sheets sandwiching a central octahedral -O-Mg-O-58 sheet (Fig. 1). A small amount of isomorphous substitution of Si⁴⁺ by Al³⁺ (and possibly other trivalent 59 cations such as Fe³⁺) in the tetrahedral sheet causes the layer to be negatively charged. This charge is 60 compensated by exchangeable interlayer cations including Na⁺, NH₄⁺, K⁺, Li⁺, and Mg²⁺ (M^{z+}). In this 61 context, the ideal structural formula of Sap can be described as $M_{x/z}^{z+}[Mg_6][Si_{8-x}Al_x]O_{20}(OH)_4 \cdot nH_2O$, 62 63 where M^{z+} represents the interlayer cations and x can range from approximately 0.4 to 1.2. The 64 majority or even all the layer charge of Sap is located on the tetrahedral sheet and this has an important influence in the properties of Sap. Nevertheless, some studies have suggested that Mg²⁺ in the 65 octahedral sheet can be substituted by trivalent cations such as Al³⁺. If Mg²⁺ cations are replaced by 66 Al^{3+} cations at a ratio of $3Mg^{2+}/2Al^{3+}$, the substitution in the octahedral sheet does create additional 67 positive charges in the layer; If at a ratio of $1Mg^{2+}/1Al^{3+}$, positive charges are created in the layer 68 (Brigatti et al., 2013). The positive charges can partly compensate the negative charges of the 69 tetrahedral sheet. Mg²⁺ in the octahedral sheet can also be substituted by other divalent cations (Ni²⁺. 70 $\operatorname{Co}^{2+}_{-}$ Fe², etc.) with no consequence on the layer charge. 71



73 Fig. 1. (A) Structure of saponite, two TO₄ (T =Si⁴⁺ and/or Al³⁺) tetrahedral sheets on each side of an octahedral sheet 74 occupied by magnesium and/or aluminium cations, forming 2D layers (Figure is made by using the VESTA software) 75 (Adapted and Reprinted from Lainé, et al., 2017, Copyright 2017, with permission from Royal Society of Chemistry under the 76 terms of Creative Commons CC By License; Momma and Izumi, 2011). Each TOT layer has a thickness of roughly 9Å. 77 Compensating cations (here sodium) are present in interlayer space and are surrounded by water molecules. The door 78 value obtained from X-ray diffraction experiments enables determining the thickness of the interlayer space, and hence, 79 the number of water layers in it (Lainé et al., 2017). (B) Top view of saponite structure (Figure is made by using the 3D 80 Max software).

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82 Saponite has a large specific surface area (SSA), surface acidity and cation exchange capacity
83 (CEC). In addition, it is distinct from dioctahedral smectites such as montmorillonite (Mt) and
84 nontronite. For example, Sap has a higher thermal stability than Mt (Casagrande et al., 2005; Vogels et

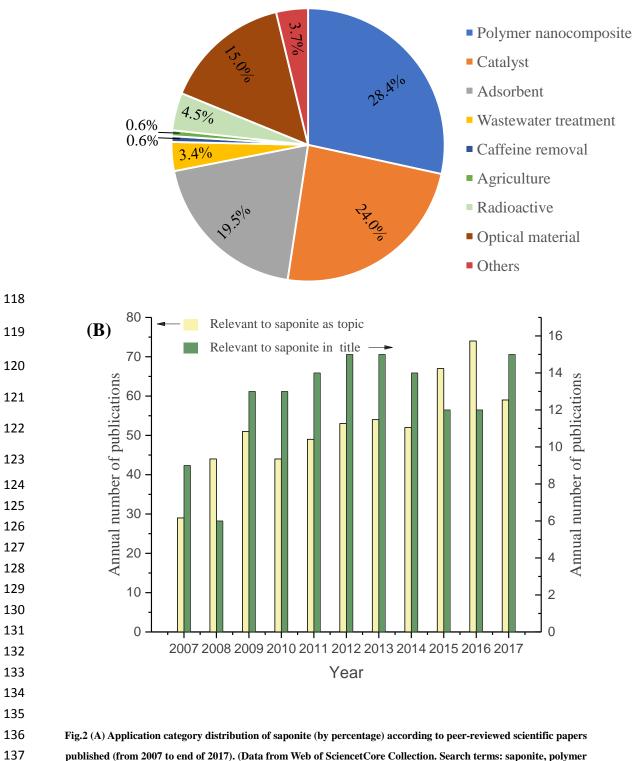
85 al., 2005; Tao et al., 2016); the size of Sap particles (~50 nm) is generally smaller than that of Mt (~300 86 nm) (Ebina and Mizukami, 2007; Paul and Robeson, 2008). In addition, Sap is easier to delaminate and 87 to be exfoliated into nanoplatelets, or even individual nanolayers in water (Takagi et al., 2013). 88 However, naturally occurring Sap is scarce. Due to the presence of impurities in the Sap deposits in 89 nature, laborious and time-consuming purification is required to obtain relatively pure Sap (Carrado et 90 al., 2006). Moreover, the chemical composition of natural Sap can be extremely variable, depending on 91 the geological genesis process (Utracki et al., 2007). This defects limits the use of natural Sap in 92 catalysis (Gómez-Sanz et al., 2017), adsorption (Franco et al., 2016a) and optics (Tsukamoto et al., 93 2016b). Hence, Sap-like solids with well-controlled chemical composition are synthesized and such 94 solid products are often called synthetic Sap. Generally, typical approach to the synthesis of Sap are 95 hydrothermal synthesis (Suquet et al., 1977; Kloprogge, 1999; Zhang, et al., 2009; Jaber et al., 2013; 96 Baldermann et al., 2014; Carniato et al., 2014; Zhang et al., 2017). The process can be modified to be 97 microwave-assisted synthesis (Trujillano et al., 2010; Gebretsadik et al., 2014), and sol-gel synthesis 98 (Garade et al., 2011; Garciano et al., 2014).

99 The synthesis of Sap is relatively mature, at least on a laboratory scale. However, either natural 100 Sap or synthetic Sap is rarely used directly. Under many circumstances the Sap needs modification, 101 including surface engineering, intercalation and hybridization of Sap. Taking advantage of the 102 swellability of Sap (Ferrage et al., 2010; Dazas et al., 2015; Ferrage, 2016), researchers can introduce 103 functional guest molecules into the structure of Sap and thus tune porosity, acidity and other physical 104 and chemical properties. Many types of Sap-based hybrids or composites have been successfully 105 prepared by the intercalation of bulky organic (Kurokawa et al., 2014; Marcal et al., 2015; Sas et al., 106 2017) and inorganic (Franco et al., 2016b; Miyagawa et al., 2017) cationic species, neutral polar 107 molecules (Marcal et al., 2015) and polymers (Mishchenko et al., 2016; Eguchi et al., 2017) into the 108 interlayer space of Sap. Covalent grafting of a moiety of organic molecules onto the surface of Sap 109 proves feasible. Accordingly, the resultant Sap-based materials have been examined for a wide range of 110 applications. Increasing studies show that such materials can be used as adsorbents (Marcal et al., 2015; 111 Sato et al., 2016), catalysts and catalyst supports (Carniato et al., 2014; Fatimah et al., 2016). They can 112 also be used as a functional additive in clay/polymer nanocomposites (CPN) (Wu et al., 2014; Wang et 113 al., 2015) (Fig. 2). Strikingly, Sap has been used as a support for the immobilization of luminescent

114 entities, rendering the resultant nanocomposites with particularities in optical imaging (Tsukamoto et

115 al., 2016b) and optoelectronic applications (Nanan et al., 2015).





nanocomposite, catalyst, adsorbent, wastewater treatment, caffeine removal, agriculture, radioactive, optical material) (B)
 Annual number of peer-reviewed papers published (from 2007 to 2017) relevant to the topics of saponite and the title
 contain saponite. (Data from Web of SciencetCore Collection. Search terms: saponite).

141 **2** Inorganic modification and hybridization of saponite

142 The inorganic modification of Sap is mainly achieved by the exchange of cations M^{z+} in the interlayer space of Sap with external inorganic cations or the generation of oxides in the form of pillars 143 in the interlayer space of Sap. The frequently-used methods include acid activation (M^{z+} exchanges 144 with H^+), inorganic cation exchange (exchange with inorganic metal cations), pillaring (M^{z+} first 145 146 exchanged with inorganic metal hydroxycations, followed usually by calcination to form oxide pillars), 147 and the hybridization of metal nanoparticles with Sap to form nanocomposite. After such inorganic modifications, the interlayer spacing, the thermal stability and the surface acidity of Sap are usually 148 149 increased. Thus the resultant Sap-based materials are commonly used as catalysts and adsorbents.

150 **2.1 Acid activation and inorganic cation exchange**

151 Acid activation refers to the process treating Sap with mineral acids such as sulphuric acid or hydrochloric acid. The final texture and structure of the treated Sap is dependent upon the 152 153 concentration of the acid used in the process. The product can be H^+ - Sap because merely M^{z+} cations in the interlayer space of Sap are exchanged by H⁺. It can also be smaller particulate H⁺-Sap if part of 154 155 Sap particles has been dissolved or damaged by the acid. It is worth noting that the dissolution of Sap is 156 much easier than that of Mt. Hence, the final product can even be an amorphous silica (Komadel and 157 Madejova, 2013). In the former two cases, the H⁺- Sap usually exhibits increased surface acidity, SSA 158 and pore volume (Komadel and Madejova, 2013; Franco et al., 2016b). A recent study shows that the microwave irradiation as heating in the acid activation of Sap can remarkably reduce the time for acid 159 160 activation (Franco et al., 2016b). Similar to acid-activated Mt (also known as bleaching earth), such 161 acid-activated Sap can be used as adsorbents (Ugochukwu et al., 2014a; Ugochukwu and Fialips, 2017a) and as catalysts (De Stefanis et al., 2013) or catalyst supports (Gebretsadik et al., 2016, 2017b). 162

163 The inherent nature of acid activation of Sap is a cation exchange reaction. As such, the 164 exchangeable cations M^{z+} in the interlayer space of Sap are replaced by H^+ ions. Nevertheless, under 165 many circumstances, the dissolution of part of the octahedral and tetrahedral sheets often occurs simultaneously (Kaviratna and Pinnavaia, 1994; Komadel and Madejová, 2013). The dissolution of the whole layer is also possible if the acid is strong enough. As a result, due to the attack by the proton H^{+,} the structural cations are released from the layers, leading the breakage of the layers (Kaviratna and Pinnavaia, 1994; Komadel and Madejová, 2013). Such attack prefers to be at the edge of the layer which is the broken-bond plane, but it can also take place on the oxygen plane.

171 When Sap is treated by mild acid and harsh conditions usually caused by the acid and temperature, 172 the morphology, structure, and texture of the resultant acid-treated Sap appears different (Bisio et al., 173 2008; Guidotti et al., 2009). When Sap is treated under mild conditions, usually the amount of Brønsted 174 acid sites increases without little alteration of the structure of Sap. In contrast, when Sap is treated 175 under harsh conditions, the cation exchange reaction is also accompanied by progressive dealumination 176 of the framework (layer) of Sap. Accordingly, a significant amount of porous amorphous silica can be 177 produced. Though microporosity and SSA are largely increased, the number of Brønsted acid sites 178 significantly decreases.

In addition to acid activation, the original cations M^{z+} in the interlayer space of Sap can also be exchanged with other inorganic cations, such as Li⁺, Cu²⁺, Ni⁺ (Vicente et al., 2011), Zn²⁺ (Intachai et al., 2017), and Fe³⁺ (Franco et al., 2016a). In particular, Cu²⁺, Ni⁺ and Fe³⁺ can be converted into zero-valent metal nanoparticles by appropriate reduction reaction. In this context, two-dimensional Sap acts as a nanoreactor with limited space for generating metal nanoparticles and such metal nanoparticles are -free of organic protectants (Miyagawa et al., 2017).

185 2.1 Pillaring, supporting and hybridization

186 A typical pillaring procedure of Sap involves the introduction of pre-prepared bulky inorganic 187 metal polyhydroxycations or polyoxocation into the interlayer space of Sap by a cation exchange reaction or the electrostatic attraction, following by calcination, which usually cause dehydration, 188 189 dehydroxylation and oxidation of the intercalated species (Fig. 3A) Hence, the polyoxocations are 190 converted to metal oxide clusters called pillars commonly (Gil et al., 2000). If the solid is calcined 191 under a flow of H₂, a reduction reaction occurs. Namely, the layered structure of Sap is pillared by the 192 metal oxide clusters. The pillars stably expand the 2:1 layers of Sap, prevent its collapse, expose their 193 internal surface and make the pillared Sap with micro- to mesoporosity (Ding et al., 2001; Vicente et al.,

194 2013). In addition, the pillars bring additional acid sites or redox sites and thus provide catalytic active
195 species for specific catalysis and adsorption (Table 1). Upon the pillaring procedures, the precursors
196 and the chemical compositions of pillars, the interlayer spacing and the porosity of the pillared Sap
197 vary.

- 198
- 199 (A)

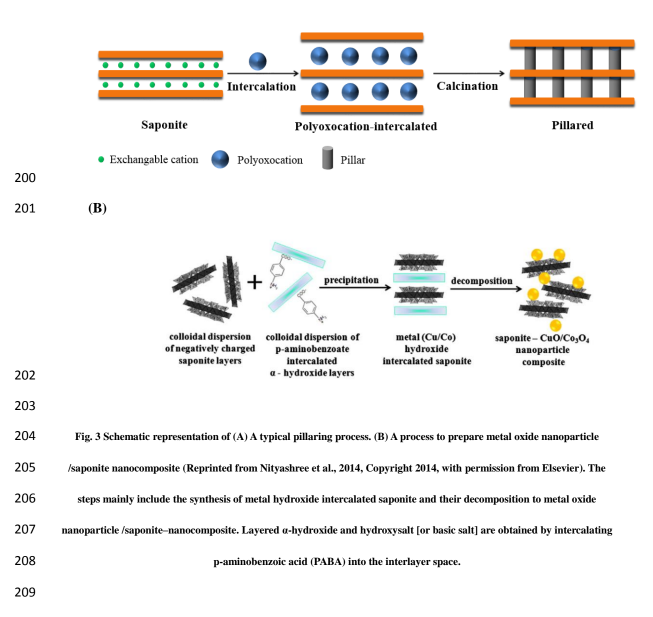


Table 1. Basal spacing (d₀₀₁), pore volume, surface area and applications of metal pillared saponite.

Pillaring		d ₀₀₁ *	$\mathbf{V_p}^*$	S _{BET} *	Applications	Ref.
agent	Amount	(Å)	(cm ³ /g)	(m ² /g)		

Al ₁₃	5mmol/g	17.8	0.188	177			
Cs- Al ₁₃	0.25 wt% ^{a)}	18.0	0.167	95	Catalyst for the synthesis of N-alkyl pyrazoles.	Velasco et al. (2011)	
Cs- Al ₁₃	1.25 wt%		0.156	111			
Al ₁₃	5mmol/g	18.0	-	274	Catalyst for acetalation of pentaerithritol	Kannan et al. (2011)	
Al	-	17.6	0.120	134			
Fe/Al	Al: 12 mmol/g Fe: 1	18.2	0.130	162	Catalyst for the cracking of polyethylene	De Stefanis et al. (2013)	
Zr	mmol/g		0.052	233	Catalyst for the isomula cal	Estimals at al	
Zr Ni/Zr	- 1 wt%	14.3	0.053 0.042	255 199	Catalyst for the isopulegol hydrogenation	Fatimah et al. (2015)	
Al	10mmol/g	17.0 16.6 ^{b)}	- 0.042 ¹	- 191. ¹	Photocatalytic phenol	Fatimah and	
Al-Ti	1.2 wt%	-	0.019 0.019^{1}	132 276 ¹	photo-oxidation in aqueous solution	Wijaya (2015)	
Fe	2 mmol/g		0.160	127	Adsorption of heavy metal	Franco et al.	
Al	20 mmol/g	12.8	0.180	178	cations from water	(2016a)	
	2 wt%		0.1441	53			
Magnetite	4 wt%		0.1455	56	Adsorption of dyes from	Makarchuk et	
	7 wt%	-	0.1464	53	aqueous solution	al. (2016)	
	10 wt%		0.3058	69			
Zr	5 wt%	13.4	0.0054	177	Catalyst for the conversion of	Fatimah et al.	
S-Zr	J wt70	15.7	0.0016	190	citronellal into menthol	(2016)	

212 *For clarity, consistency, and easy comparison, some original data have been rounded off.

a) Saponite pillared with Al and then the pillared solid doped with Cs. b) Samples prepared by using microwave irradiation as calcination procedure instead of conventional calcination method. S_{BET} : specific surface area calculated according to the BET equation; V_p = total pore volume; S-Zr: sulphated zirconium pillared saponite.

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As the precursors of the pillars, also called pillaring agents, can be metallic cations, metallic oxide sol, polyhydroxycations or polyoxocations (**Table 1**). $[Al_{13}O_4(OH)_{24}(H_2O)_{12}]^{7+}$ usually abbreviated as Al₁₃ is the most widely studied (Trujillano et al., 2009; Kannan et al., 2011; Velasco et al. , 2011). Accordingly, the Al-pillared saponite is well-documented. This is mainly because the preparation of Al₁₃ is very mature and its structure is better understood than that of other cations. In addition, Al₂O₃ 223 pillars is thought to possess solid acidity. Particularly, thermal resistance of Al-pillared Sap is higher 224 than that of Al-pillared Mt (Chevalier et al., 1994; Lambert et al., 1994; Bergaoui et al., 1995a,b). Pillars made from polycations which formed by the hydrolysis of Zr⁴⁺, Ti⁴⁺, Fe³⁺, or Ga³⁺ (Malla and 225 Komarneni, 1993; Vicente et al., 2008; Fatimah et al., 2015; Fatimah and Wijaya, 2015; Franco et al., 226 227 2016a; Makarchuk et al., 2016; Gebretsadik et al., 2017b) has also been reported. It is worth mentioned 228 that Ni²⁺ cation does not form polycations upon hydrolysis. In general, the basal spacing and the pore size of 229 the resultant pillared Sap are inherently decided by the size of the oligomers and accordingly the 230 products after calcination (Vicente et al., 2008; Franco et al., 2016a). In the two-dimensional network 231 of pillared Sap, interlayer spacing is controlled by the height of the pillars, whereas both the distance 232 between the pillars and the height of the pillars known as inter-pillar distance, determines the pore size. 233 The inter-pillar distance also reflects the density of pillars in the interlayer space of Sap. Of great 234 challenge is the interactions between guest pillars and the layers of the Sap. For Al-pillared Sap, the 235 -O-Al-O- on the Al₂O₃ pillars could form covalent bonds with -Al-O and -Si-O on the layer of Sap. 236 However, for many other elements with 4f/5f orbitals, the interactions remain unclear. Possibly 237 theoretical modeling can be conducive to getting more insights into this issue (Bian et al., 2015; Dong 238 et al., 2016). Besides, it has not yet to find an effective way to measure the size and charge of each 239 individual pillar accurately.

240 Though the chemical composition and structure of the metallic cation or its polycation used in the 241 pillaring process play a pivotal role in the characteristics and properties of the pillars in the pillared Sap, 242 other important factors such as the concentration of the pillaring agent (Khumchoo et al., 2016; 243 Makarchuk et al., 2016), the drying method and the calcination temperature (Vicente et al., 2008) have 244 significant effect on the pillars. For instance, recently, Fatimah and Wijaya (2015) found that during 245 the calcination procedure, when microwave irradiation for 15min was used, the Al-pillared Sap with a 246 higher SSA was obtained, compared with the Al-pillared Sap by conventional slow heating for 4h. 247 Inherently, under the two different calcination methods, the conversion of polycations would be 248 different. As conventionally to prepare pillared Sap need a long procedure, rapid microwave irradiation 249 for heating or calcination can shorten the process and much increase productivity.

Saponite can also be used as a matrix or support for adsorption of metallic nanoparticles or oxides
to produce nanocomposite. To this end, there are such ways as impregnation and deposition. Also the
Sap can be pre-modified judiciously if necessary. For example, in the case of Zn, it is deposited as

253 oxide on a cetyltrimethylammonium cation (CTA⁺)-intercalated Sap (Khumchoo et al., 2016) while in 254 the case of Ni, this metal is supported on Sap (Gebretsadik et al., 2017b). Notably, Sap easily 255 delaminates or can be exfoliated in water. By spontaneous flocculation of delaminated Sap, the 256 delaminated or exfoliated Sap can then be used to host other layered nanomaterials (e.g. Co, Cu 257 hydroxide) to form composites (Nityashree et al., 2014). Upon calcination, the metal hydroxide therein 258 decomposes to form metal oxide nanoparticle/Sap nanocomposite in which the nanoparticles get 259 uniformly distributed in Sap matrix (Fig. 3B). The particle size of the metal oxide nanoparticles formed 260 in the nanocomposite could be altered by varying factors such as the temperature, atmosphere and 261 duration of decomposition. If the nanoparticles are chosen to be nanosized magnetite, then a magnetic 262 nanocomposite can be achieved (Makarchuk et al., 2016).

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3 Organic modification of saponite

264 Positively charged organic cations can be intercalated into the interlayer space of Sap by cation 265 exchange or adsorption to produce OSap (Zhen and Wang, 2016; Tominaga et al., 2017). The inorganic 266 cations in the interlayer space of Sap, that compensate the negatively charged layer, can be exchanged 267 with positively charged organic cations. Such cations can be formed from surfactants, dye or 268 organometallic complexes in water (Kurokawa et al., 2014; Seki et al., 2015; Sas et al., 2017). The 269 organic cations are not only intercalated in the interlayer space of Sap, but also adsorbed on the surface 270 of Sap to form an electric double layer. Such organic modification of Sap is easily achieved. Studies 271 have demonstrated that during the synthesis process of Sap, OSap can be synthesized in a one-pot way 272 by the direct adding organic cations into the inorganic gel for synthesizing Sap (Bisio et al., 2011; 273 Kurokawa et al., 2014). The surface of Sap is usually hydrophilic and the inorganic cations in the 274 interlayer space tend to be hydrated (Ferrage, 2016). When such places are occupied by organic species 275 after organic modification, the surface is then covered by organic carbon chains or groups, thus the 276 hydrophilic Sap are changed into hydrophobic OSap. Accordingly, the polarity is altered (de Paiva et al., 277 2008). Besides, a wide range of organic cations, groups or molecules can be used and can perform a 278 variety of functions such as in optical properties (Wu et al., 2015), adsorption (Seki et al., 2010, 2015) 279 and catalysis (Goto et al., 2016). In addition, the inorganic layers of Sap play an active role in 280 stabilizing and protecting the intercalated organic species. Typically, their thermal stability or

281 heat-resistance remarkably increased (Bisio et al., 2011).

282 3.1 Organic cations

283 By taking advantage of cation exchangeability of Sap, organic cations can be incorporated into the 284 interlayer space of Sap by a cation exchange reaction. Consequently, organic cations replace the 285 original inorganic cation in the interlayer space of Sap to compensate the negative electric charge in the 286 layer of Sap. Such organic cations are commonly moieties of cationic surfactants such as 287 cetyltrimethylammonium (CTA), benzylammonium (BA), and didecyldimethylammonium (DDDMA) 288 cations. In particular, cationic surfactants have been successfully used as organic modifiers of Mt to 289 produce organo- Mt in industry (Lagaly and Weiss, 1969; Lagaly, 1981, 1986; Yu et al., 2014). In 290 addition, over the last decade, quaternary fulvic acid (QFA), and aminopropyl isobutyl 291 titanosilsesquioxane (Ti-NH₃POSS) have also been used to modify Sap (Table 2).

292 After organic modification, the surface of the Sap becomes hydrophobic from hydrophilic 293 (Ugochukwu et al., 2014b; Ugochukwu and Fialips, 2017b). Depending on the layer charge density 294 (LCD) of Sap, and the chain length or the stereoscopic structure and geometric size of organic cations, 295 the extent to which the inorganic cations are replaced by the organic cations, the organic moieties 296 appear to have several types of arrangements in the interlayer space of Sap. In the interlayer space of 297 Sap, the organic cations may lie flat as a monolayer (Okada et al., 2014) or bilayer (Marcal et al., 2015) 298 pseudo-trilayers or multilayers (Yu et al., 2014). Clearly, the arrangements of organic cations, along 299 with the length of their carbon chains, decide the degree of expansion of the Sap. For example, for 300 BA-intercalated Sap, when BA⁺ cations occur in the form of a monomolecular layer, the cations should 301 be aligned parallel to the oxgen plane of the Sap layer. As a result, the interlayer spacing of BA-Sap can 302 be increased a little by a few Angstroms from the basal spacing 1.26 nm of original Sap (Okada et al., 303 2014). By contrast, for CTA-intercalated Sap, when an arrangement of bilayer CTA⁺ ions is achieved, 304 the interlayer spacing of Sap increases remarkably by several Angstroms to a few nanometers 305 (Marcal et al., 2015). In this case, the ammonium groups occur like being tethered to the surface Sap 306 layer, and the alkyl chains lay almost vertical to the layers. Moreover, the intercalated organic cations 307 may undergo a conformational change when the thickness of the intercalated molecule is larger than 308 the interlayer spacing of Sap. This conformational change depends on the extent of loadings of organic 309 species and the layer charge density of Sap (Tominaga et al., 2017).

Modifier	Amounts (modifier/Sap)	Basal spacing (nm)	S _{BET} ^{#,} * (m ² /g)	Interlayer* space (Δd) (nm)	Ref.
Ti-NH ₃ POSS	-	-	-	2.60	Carniato et al. (2009,2011)
CONH	110% CEC	1.51	144	0.55	Seki et al. (2010, 2015)
CTAB	200% CEC	1.43 ^{a)} 1.35 ^{b)}	-	-	Bisio et al. (2011)
BA	50,100% CEC,	1.31, 1.30	_	0.35, 0.34	Okada et al. (2014)
CONH	200% CEC	1.51		-	
Arquad 2HT-75	500% CEC	3.60			
Octadecylamine	50% CEC	1.42	-		Albeniz et al. (2014)
OTAB	200% CEC	1.92		-	
DDDMA	35% CEC	1.40	330(EGME) °	-	Ugochukwu et al. (2014b, 2017b)
CTAB	-	1.84	0.1	0.88	Marçal et al. (2015)
QFA	50 wt%	1.57	-	-	Zhen and Wang, (2016)
BP	10-27% CEC	-	-	0.51-0.55	Tominaga et al. (2017)

311 Table 2. Characteristics of some organic modified saponite (Sap).

312 # The specific surface area (SSA) of the samples was determined by BET equation. ♀ The SSA of the samples was determined

by the EGME (Ethylene glycol monoethyl ether) method (Carter et al., 1965) * For clarity, consistency, and easy comparison,

some original data have been rounded off.

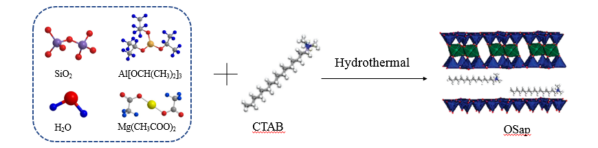
315 ^{a)} One-pot synthesized CTA-Sap. ^{b)} Post synthesized CTA-Sap.

Ti-NH₃POSS: aminopropyl isobutyl titanosilsesquioxane; CONH: neostigmine; CTAB: cetyltrimethylammonium bromide; BA:
 benzylammonium; OTAB: trimethyloctadecylammonium bromide; DDDMA: didecyldimethylammonium; QFA: quaternary

318 fulvic acid; BP: biphenyl derivative.

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A one-pot strategy for producing OSap has been developed as an alternative to the post-modification method of Sap (Roelofs and Berben, 2006; Bisio et al., 2011). This new method can intercalate organic cations into the interlayer space of Sap by the direct introduction of organic cations in the gel used for synthesizing Sap during the process (**Fig. 4**). By this methodology the procedure of producing OSap is significantly simplified and no longer time-consuming. Moreover, a low amount of surfactant is needed in such a method that thereby lowers the costs of producing OSap. Besides, the distribution of surfactant molecules could be more homogeneous in the OSap (Bisio et al., 2011)



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Fig. 4 Schematic drawing showing one-pot synthesis of organo-saponite (OSap) (Bisio et al., 2011). CTAB: 330 cetyltrimethylammonium bromide.

331

332 3.2 Dye molecule

333 Compared with Mt, Sap is more easy delaminated completely in water to form a transparent 334 dispersion. Such optical transparency, together with high absorption capacity of Sap, make it a very 335 good two-dimensional inorganic host for many organic dye molecules with optical functions 336 (Tsukamoto et al., 2016b), in particular fluorescence. The interlayer space and the layer charge can 337 provide a static confinement to the fluorescent dyes and is conducive to better distribution and 338 assembly of the dye molecules, thereby preventing fluorescence from quenching. A variety of dye 339 molecules, including porphyrin dyes (Takagi et al., 2013; Konno et al., 2014), cyanine dyes (Takagi et 340 al., 2013; Czímerová et al., 2017; Matejdes et al., 2017), acetylene derivatives (Suzuki et al., 2014), 341 methylene blue (MB) (Donauerová et al., 2015), triphenylbenzene derivatives (Tokieda et al., 2017), 342 biphenyl derivative (BP) (Tominaga et al., 2017), laser dyes rhodamine 6G (R6G), oxazine 4 (Ox4) 343 (Belušáková et al., 2017), and anthocyanin (Ogawa et al., 2017) have been successfully introduced onto 344 the surface or into the interlayer space of Sap. The driving forces involve cation exchange reactions, electrostatic attraction, hydrophobic interaction, van der Waals force, physical adsorption and 345 346 chemisorption between dye and Sap.

347 Organic dye cations generally tend to aggregate on the surface of Sap, due largely to the hydrophobic interaction and van der Waals force between organic species (Konno et al., 2014; 348 349 Donauerová et al., 2015; Matejdes et al., 2017; Sas et al., 2017; Tokieda et al., 2017). The aggregates 350 are categorized into two types: H-aggregates and J-aggregates. The former more frequently occurs and 351 is based on a sandwich-type intermolecular association and absorb the light of higher energy; the latter 352 appears less frequent and are formed by a head-to-tail intermolecular association. The types of the

353 aggregates are related to the amount of adsorbed dye cations. At higher ratio of dye/Sap, a large amount of H-aggregates tends to form whereas at the lower ratio of dye/Sap, J-aggregates are preferentially 354 355 formed (Boháč et al., 2016). The amount of the J-aggregates also increased with the aging of hybrid 356 colloids and is influenced by the solvents (Czímerová et al., 2017). In reverse, segregation of dye 357 cations on the surface of Sap often occurs (Konno et al., 2014). Segregation behavior depends 358 primarily on the chemical structure of the dye cations. Such aggregation drastically reduces the 359 efficiency of the intermolecular photochemical reactions because the functional dye cations are 360 separated from each other in the interlayer space of Sap or on the surface of Sap.

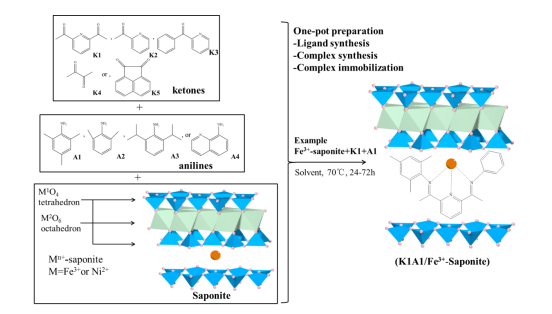
361 The structure of dye assembly on Sap can be effectively controlled by the electrostatic host-guest 362 interaction (Takagi et al., 2013). It includes the intermolecular distance, the molecular orientation, the 363 segregation, aggregation and integration, and the strength of the immobilization. In this context, a 364 size-matching effect have been proposed. In addition, the arrangement of organic fluorescent dyes can 365 be also affected by the layer charge density of Sap (Wu et al., 2015). Hence, it can be tuned by 366 changing the layer charge of Sap. According to a recent study by molecular dynamic simulation, (Wu et 367 al., 2015) as to bis-N-methylacridinium (BNMA)-intercalated Sap, when the layer charge density 368 increased, the electrostatic attraction between the layer of Sap and BNMA became stronger, and 369 BNMA in the interlayer space of Sap tended to arrange in a monolayer parallel to the layer. The parallel 370 orientation suppressed remarkably the molecular aggregation, preserving the photoactivity of dye 371 molecules.

372 3.3 Organometallic complex

373 Some organometallic complexes can be introduced into the interlayer space of Sap and interact 374 with the anionic layers of Sap. The confinement of organometallic complexes in the interlayer space of 375 Sap can regulate the molecular structure and the relevant arrangements, and increase the stability of the 376 complexes (Pimchan et al., 2014). Much importantly, the intercalation of organometallic complexes 377 into the interlayer space of Sap brings additional functions caused by the organometallic complexes. In 378 other words, the organometallic complex-Sap hybrids can act as catalysts (Pimchan et al., 2014; 379 Kurokawa et al., 2014; Goto and Ogawa, 2016), catalyst support (Goto and Ogawa, 2015) or optical 380 materials (Sato et al., 2014a; Hosokawa and Mochida, 2015; Tamura et al., 2015; Eguchi et al., 2017). 381 Up to now, many organometallic complexes, including bis(8-hydroxyquinoline)zinc(II) complex 382 (Znq₂) (Pimchan et al., 2014), iridium(III) complexes (Sato et al., 2014a; Tamura et al., 2015), nickel(II)

383 complexes with diketonato and diamine ligands (Hosokawa and Mochida, 2015), tris(2,20-bipyridine) $([Ru(bpy)_3]^{2+})$ (Goto and 384 ruthenium(II) complex Ogawa, 2016), and iron(II)-based 385 metallo-supramolecular complex polymer (polyFe) (Eguchi et al., 2017) have been used to modify Sap 386 to produce organometallic complex-Sap hybrids. Typically, the intercalation of an organometallic 387 complex in the interlayer space of Sap is achieved by exchanging the inorganic cations in the interlayer 388 space of Sap with cationic organometallic complexes. Recently, a new process to produce 389 organometallic complexes -Sap hybrids have been developed by Kurokawa et al. (2014) (Fig. 5). In 390 this process, transition-metal complex/Sap hybrids have been in situ synthesized successfully, instead the complexes should be pre-synthesized usually. The Fe³⁺- and Ni²⁺-exchanged Sap acted as an acid 391 392 catalyst and promoted ligand formation from an aniline derivative and a ketone derivative. The formed 393 ligand then simultaneously coordinated to the metal ions located in the interlayer space of Sap to form 394 bis(imino)pyridineiron(III), α -diiminenickel(II), and iminopyridinenickel(II) complexes, respectively. 395 Such methodology tactically combines catalysis, ligand synthesis, complex synthesis and 396 immobilization together in a one-pot process and can be regarded as a more efficient and cleaner 397 approach to producing organometallic complex -Sap hybrids.

398





400 Fig. 5 One-pot preparation of transition-metal complex/saponite hybrid (Adapted and reprinted from Kurokawa et

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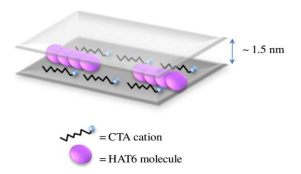
al., 2014, Copyright 2014, with permission from Elsevier).

403 **3.4 Grafting organic moiety and silylation**

404 Beside cations, neutral organic molecules can also be intercalated into the interlayer space of Sap. 405 Generally, cation exchange method cannot be used to introduce neutral organic molecules into Sap. 406 Instead, in the case of neutral organic molecules, the possible chemical interactions including covalent 407 bonds, hydrogen bonding, co-ordination, ion-dipole interaction, charge transfer, acid-base reactions, 408 and van der Waals forces can be used (Marcal et al., 2015; Tao et al., 2016). In particular, in grafting of 409 organic moiety onto Sap, covalent bonds between surface -OH groups of Sap and reactive organic 410 groups form. As such, the reactive hydrophobic organic groups durably immobilized on Sap, 411 preventing the organic moiety from leaching (He et al., 2005; Marcal et al., 2015; Tao et al., 2016). By 412 contrast, when organic cation-intercalated Sap is dispersed in solution, the long alkyl chain of organic 413 cations can leach into the medium. In addition to grafting, neutral organic compounds can also form 414 complexes with the interlayer cations of Sap (Kurokawa et al., 2014).

415 In the context of grafting organic moiety onto Sap, silylation, has captured particular attention. 416 The hydroxyl groups (Si-OH-) on the external surface and/or edges of Sap can react with the silvl 417 groups (R₃Si) of silanes to form covalent bonds (Marcal et al., 2015). In particular, the broken edge of Sap possesses active Si-OH, Mg-OH and Al-OH groups (Wu et al, 2015; Honda et al, 2016; Tao et al. 418 419 2016). Namely, the active groups of Sap readily react with Si-O groups of silane and then produce 420 silylated Sap hybrids (Herrera et al., 2004, 2005; He et al., 2014). After the edge of Sap is silylated with 421 properly-chosen silane, then it is possible for the Sap particles to be linked by silane (Zhou and Keeling 422 2013).

423



424

425 Fig. 6 Schematic drawing of the structure of CTA-HAT6-Saponite hybrid (Reprinted Nanan et al., 2015, Copyright 2015,

426 with permission from Elsevier). CTA: Cetyltrimethylammonium cation, and HAT6: Hexakis(hexyloxy) triphenylene (a

triphenylene-based liquid crystal molecule,)

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429 Alternatively, Sap can be firstly intercalated by organic cations, then the OSap enables neutral 430 organic molecules to be attracted into the interlayer space by hydrophobic interaction. For example, 431 Ishida et al. (2013) successfully prepared a supramolecular host-guest hybrid composed of a cationic 432 organic cavitand (octaamine, OAm, host), two neutral aromatic molecules (pyrene and thracene, 433 guests), and Sap. Due to strong Coulombic attraction between the ammonium groups of OAm and the 434 anionic layer of Sap, the OAm was anchored to the layer of Sap. OAm can then form capsular 435 assembly with organic neutral guest molecules. Namely, the cationic organic cavitand OAm enabled 436 neutral organic molecules to be intercalated into the layer space of Sap. More recently, Nanan et al. 437 (2015) successfully prepared a functional hybrid of a triphenylene-based liquid crystal molecule, 438 hexakis(hexyloxy) triphenylene, and Sap, by the reaction between colloidal Sap in the presence of a 439 cationic surfactant, cetyltrimethylammonium (CTA⁺) cation, and hexakis(hexyloxy) triphenylene 440 (HAT6) (Fig. 6). Such findings mean that one-pot facile process for introducing cationic and neutral 441 organic species into the interlayer space of Sap is a lot feasible.

442 **4 Saponite/polymer nanocomposites**

The idea of Sap/polymer nanocomposites might be borrowed from the Mt /polymer nanocomposites (Lambert and Bergaya, 2013). Similarly, the preparation of Sap/polymer nanocomposites is also achieved by solution intercalation (Jairam et al., 2013; Shin et al., 2013; Yang et al., 2013; Chang et al., 2014; Wu et al., 2014; Wang et al., 2015; Xi et al., 2015; Ju and Chang, 2016; Eguchi et al., 2017; Sas et al., 2017), melt mixing or melt intercalation (Zhen et al., 2012; Kitajima et al., 2013; Zhen and Wang, 2016) or *in situ* polymerization (Jairam et al., 2013; Nakamura and Ogawa, 2013; Tong and Deng, 2013; Bandla et al., 2017) (**Table 3**).

451 Table 5. Preparation and properties of typical sapointe/polymer nanocomposites	451	Table 3. Preparation and properties of typical saponite/polymer nanocomposites.
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Saponite (Sap)	Polymer	Method	Conditions	Properties	Ref.
now Con	PVA	malt mining	plasticizer: water and formamide; melt	coexistence of intercalated and	Zhen et al.
raw Sap	PVA	melt mixing	temperature: 115, 135, 125, and 110°C.	partially exfoliated layers of	(2012)

				Sap.	
QFA- Sap	PLA	melt mixing	plasticizer: ATBC; melt temperature: 170,180°C.	flocculated structure and partially intercalated morphology of the composite.	Zhen and Wang (2016)
synthetic Sap	QAL	solution intercalation	Lignin content: 30 wt%; solvent: water; stirred overnight; dried in vacuum at room temperature.	30.35±0.15 wt% of lignin was absorbed with Sap.	Jairam et al. (2013)
Sap	PVA	solution intercalation	PVA content: 0-10 wt%; solvent: water; vigorous stirring for 2 h at 80°C; dried at 80°C for 24 h.	thickness of resulting films: 54-63µm.	Shin et al. (2013)
synthetic	PVA	solution	Sap content: 0-10 wt%;	thickness of PVA hybrid films:	
Sap		intercalation	solvent: water; vigorous stirring for 3h; dried at 80°C for 24 h.	20–31µm.	Chang et al. (2014)
synthetic Sap	cellulose	solution intercalation	cellulose content: 4 wt%; solvent: LiOH/urea water solution; stirred at 1200 rpm for 10 min; air-dried	the longitudinal directions of cellulose and Sap oriented parallel to the film surface.	Yang et al. (2013)
synthetic Sap	TOCN	solution intercalation	at ambient temperature. Sap content: 0–50 wt%; solvent: water; stirring for 1 h; dried at 40°C for 3 d.	thickness of resulting films: 5.0–11.7μm.	Wu et al. (2014)
HTCC– Sap	PLA	solution intercalation	content of HTCC-Sap: 0.3-5 wt%; solvent: trichloromethane; vigorous stirring in a water bath at 60°C.	thickness of resulting films: about 0.5 mm.	Xi et al. (2015)
QFA–Sap	PLA	solution intercalation	content of QFA-Sap: 0.3-5 wt%; solvent: trichloromethane; vigorous stirring at 60°C for 4 h;	thickness of resulting films: about 0.5 mm.	Wang et al. (2015)
Sap	PAA	solution intercalation	Sap content: 0 to 40 wt%; solvent: DMAc; vigorous stirring at 30°C for 1 h; ultrasonication 6 times for 5 min each; heating at 80°C for 1 h; at 100,130, and 160°C for 30 min.	thickness of resulting films: 68-71μm.	Ju and Chang (2016)
synthetic Sap	RhPEG	solution intercalation	polymer content: 0.05–0.25 wt%; solvent: water; films were prepared using vacuum filtration technique.	-	Sas et al. (2017)
synthetic Sap	Poly- (NIPAm)	in situ polymerization	Sap content: 11, 23 and 46 wt%; monomer: NIPAm; cross-linking agent: 14.8 mmol L ⁻¹ KPS solution; react for 4 h at 70°C under nitrogen atmosphere;	Well-defined spherical particles of poly(NIPAm) hydrogel with the particle size of 800–1300 nm.	Nakamura and Ogawa (2013)
QAL–Sap	PSBA	in situ polymerization	co-stabilizer: hexadecane, AIBN; monomers: styrene, butyl acrylate; at temperature of 80±2°C for 6 h under continuous magnetic stirring; drying at 60°C for 12 h.	miniemulsions were only stable in lignin–Sap nanohybrid concentrations from 1.6 to 5.2 wt %.	Jairam et al. (2013)

VBTAC- Sap	PS	in situ polymerization	Sap content: 4–30 wt%; co-stabilizer: hexadecane and AIBN; monomer: styrene; degassing with N ₂ at RT for 30 min, and increasing temperature to 80±2°C keeping>6h under continuous mechanical stirring (600 rpm).	asymmetric polystyrene nanoparticles (hemispherical or truncated particles) with a size about 100–500 nm were synthesized.	Tong et al. (2013)
		in situ	Sap content: 1–3 wt%; monomer: AAm;	The 3 wt% Sap hydrogel	Bandla et
raw Sap	PAAm		solvent: water; initiator: potassium	showed the highest swelling	al., (2017)
		Polymentation	persulfate, accelerator: TEMEDA	capacity than other hydrogels.	un, (2017)

AAm: acrylamide; AIBN: 2, 2 azoisobutyronitrile; ATBC: acetyl tributyl citrate; DMAc: N, N-Dimethylacetamide; HTCC:
N-(2-hydroxyl) propyl-3-trimethyl ammonium chitosan chloride; KPS: potassium persulfate; NIPAm: N-Isopropylacrylamide;
PAA: poly(amic acid); PAAm: poly acrylamide; PLA: Poly(lactic acid); poly(NIPAm): poly(N-isopropylacrylamide); PSBA:
polystyrene co-butyl acrylate; PS: polystyrene; PVA: Poly(vinyl alcohol); QAL: quaternary ammonium lignin; QFA:
quaternary fulvic acid; TEMEDA: N,N,N¹,N¹-tetramethyl ethylenediamine; TOCN: 2,2,6,6-tetramethylpiperidinyl-1-oxyl
(TEMPO)-oxidized cellulose nanofibrils; RhPEG: rhodamine B (RhB) modified polyethylene glycol (PEG); VBTAC:
(ar-Vinylbenzyl) trimethylammonium chloride.

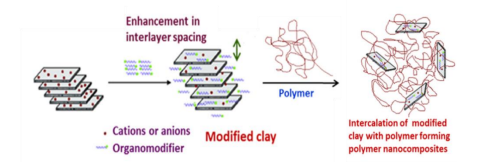
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460 Melt mixing is the simplest method since it is easily operated, economic and environmental friendly (Shen et al., 2002; Pandey et al., 2005; Zou et al., 2008). During melt mixing, Sap is firstly 461 mixed with thermoplastic polymers such as poly(vinyl alcohol) (PVA) and poly(lactic acid) (PLA) 462 463 (Zhen et al., 2012; Zhen and Wang, 2016). The mixture is then heated to the softening point of the 464 polymer in the presence of a plasticizer. With the mechanic force under heating, the Sap can be 465 exfoliated and dispersed in the polymer matrix to yield a CPN. Comparatively, solution intercalation is 466 a liquid-state process that brings about a good molecular level of mixture of Sap nanolayers and the polymer molecules (Zou et al., 2008). During this process, the soluble polymers (e.g. PVA, PLA, 467 468 poly(amic acid) (PAA), polyethylene glycol (PEG)) or the dispersion of macromolecules (e.g. lignin 469 and cellulose) in the solution are expected to be introduced into the interlayer space of Sap. Ideally, the 470 polymers are soluble in a certain solvent such as water, aqueous alkali/urea solutions, trichloromethane, 471 N, N-Dimethylacetamide (DMAc), etc. (Jairam et al., 2013; Shin et al., 2013; Yang et al., 2013; Chang et al., 2014; Wu et al., 2014; Wang et al., 2015; Xi et al., 2015; Ju and Chang, 2016; Eguchi et al., 2017; 472 473 Sas et al., 2017). However, the solution intercalation on an industry scale could be problematic because 474 of the use of cost-intensive solvent and its recovery (Zou et al., 2008). In in situ polymerization process, 475 Sap is firstly dispersed into a monomer(s) solution followed by direct polymerization. In this way, the 476 CPN are formed in situ (Jairam et al., 2013; Nakamura and Ogawa, 2013; Tong and Deng, 2013; 477 Bandla et al., 2017).

478 Whatever the process is used, an inherent issue is that ever, Sap is hydrophilic whereas polymers 479 are mostly hydrophobic. This limits the compatibility between Sap layers and polymer chains (Jairam 480 et al., 2013; Tong et al., 2013). Micro-phase separation often occurs in the CPN owing to the absence 481 of interfacial interaction between Sap layers and polymer chains. As a result, the intercalation and 482 uniform dispersion of Sap in the polymer matrix appear difficult (Wang et al., 2015; Zhen et al., 2016). 483 Therefore, to solve these issues, it is necessary to modify the surface of Sap with organic species to 484 increase the hydrophobicity of the surface of Sap nanolayers (Wu et al., 2014; Wang et al., 2015). In 485 addition, the bulky organic species can expand the interlayer space of Sap, allowing easier intercalation 486 of the polymer chains into the interlayer space of Sap (Kotal and Bhowmick; 2015) (Fig. 7).

487 Alternatively, the modification of polymers, instead of modification of Sap, can make the polymer 488 chains partly hydrophilic, thereby getting compatible with Sap so that polymer chains can well 489 penetrate into the interlayer space of Sap to produce CPN (Wu et al., 2014; Xi et al., 2015; Sas et al., 490 2017). For example, recently, Sas and co-workers (2017) produced a water-soluble and 491 positively-charged polymer (RhPEG) polyethylene glycol (PEG) with a reactive rhodamine B (RhB) 492 fluorophore and then RhPEG was successfully intercalated into the interlayer space of Sap by solution 493 intercalation with a vacuum filtration technique. In addition, certain polymer molecules or their 494 derivatives can be used to modify Sap first to produce OSap and such OSap is then used with other 495 polymers to produce CPN (Jairam et al., 2013; Xi et al., 2015)

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Fig. 7. Schematic drawing of modification of saponite followed by intercalation of polymer to form clay/polymer
 nanocomposites (CPN) (Reprinted Kotal and Bhowmick, 2015, Copyright 2015, with permission from Elsevier).

500

501Recently, bionanocomposites have attracted much attention (Shchipunov et al., 2012a,b; Migal, et502al., 2016). The conventional methods to produce CPN cannot be directly used to produce

503 Sap/biopolymer nanocomposites. Firstly, melt intercalation does not work since the majority of 504 biopolymers suffers from degradation upon heating. Secondly, the *in situ* polymerization, which starts 505 polymerization from monomers in dispersion of Sap, can also not be used, because under such a 506 circumstance the synthesis of biomacromolecules such as polysaccharides and proteins seems 507 impossible (Shchipunov et al., 2012a,b). In addition, Sap/biopolymer nanocomposites cannot be made 508 by a conventional solution intercalation because the flocculation or the precipitation caused by 509 cooperative electrostatic interactions between biomacromolecules and Sap nanoparticles, for example 510 the oppositely charged polysaccharide and Sap nanoparticles, usually occur (Shchipunov et al., 2009).

511 Chitosan, natural cationic polysaccharide containing free amino groups, have been intercalated 512 into Sap (Shchipunov and Postnova, 2010; Shchipunov et al., 2012a,b; Budnyak et al., 2016; Migal et 513 al., 2016). Shchipunov et al. (2009) demonstrated an effective way to achieve monolithic chitosan /Sap 514 hydrogel by shifting the pH of the reaction solution). The procedure to form Sap/chitosan hydrogel 515 nanocomposites includes several steps (Fig. 8A): (1) dispersing of chitosan in an aqueous dispersion of 516 Sap nanoparticles at a pH value corresponding to the neutral state of chitosan, (2) gradually acidifying 517 the dispersion medium to make chitosan macromolecules positively charged, and (3) forming network 518 structure via electrostatic attraction between Sap nanoparticles and chitosan macromolecules. These 519 physical cross-linked, a three-dimensional network exhibited a gradual increase in the viscosity and 520 finally got hydrogellation in later studies have also suggested that in such a hydrogel, part of the 521 protonated amino groups of chitosan were electrostatically cross-linked to the oppositely charged 522 surface of delaminated Sap nanoparticles in an aqueous solution (Shchipunov et al., 2012b; Budnyak et 523 al., 2014). Recently, Budnyak et al. (2016) revealed that the interaction between chitosan with Sap 524 involves both the electrostatic attraction and hydrogen bonding (Fig. 8B). It is noteworthy that this 525 method and the chitosan/Sap system can also be used to fabricate CPN films, similar to the structure of 526 mollusk shells (Shchipunov et al., 2012a).

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528 (A)

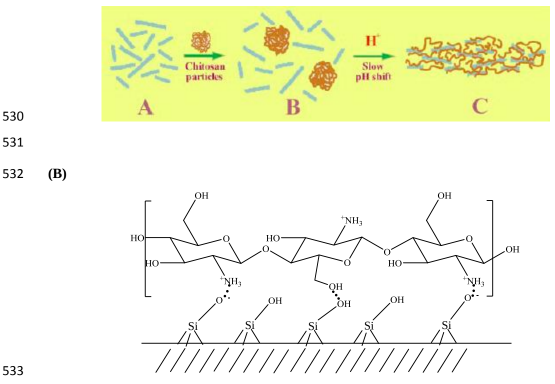


Fig. 8 (A) Schematic drawing of main stages of formation of monolithic hydrogel by chitosan and saponite (Reprinted from Shchipunov et al., 2009, with permission from The Royal Society of Chemistry). (B) Schematic diagram showing that the interactions between chitosan and saponite involves the electrostatic interaction as well as hydrogen bonding (Reprinted from Budnyak et al., 2016, Copyright 2016, with permission from Springer under the terms of the Creative Commons Attribution 4.0 International License).

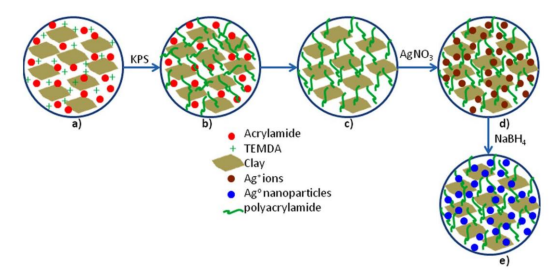
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540 **5 Organic - inorganic hybridization**

Organic-inorganic modified Sap can be prepared through the reaction between organic species and inorganic-modified Sap, or by combining inorganic species with organic-modified Sap. The organic-inorganic modification of Sap can be achieved by the *in situ* polymerization of polymers in pillared Sap (Zhen et al., 2014) or pillared organic Sap (Albeniz et al., 2014; Zhen and Sun, 2014; Zhen and Zheng, 2016) and the formation of metal nanoparticles in organically modified Sap (Venkatachalam et al., 2013; Khumchoo et al., 2015; Bandla et al., 2016).

547 The pillaring treatment of Sap can improve cation exchange and enlarge the interlayer space of 548 Sap, hence allowing the smooth insertion of polymer chains into the interlayer space of Sap during the 549 preparation of CPN by *in situ* polymerization (Zhen et al., 2014). However, the interfacial compatibility between pillared Sap and the polymer is very low and the organic modification of pillared Sap can be used to solve this problem efficiently (Zhen and Sun, 2014; Zhen and Zheng, 2016). The pillared and organic pillared Sap help in eliminating the toxic effects in the *in situ* polymerization process of polymers and facilitate the green polymerization of polymers. Moreover, pillared and organic pillared Sap can accelerate the crystallization process of polymer matrixes, hence improving their thermal stability.

In addition, the organically modified Sap can be used to stabilize and tune the size and 556 557 morphology of metal nanoparticles in the interlayer space. The interaction of the organic species and 558 Sap layers can affect the properties of the metal nanoparticles (Venkatachalam et al., 2013; Khumchoo 559 et al., 2015; Bandla et al., 2016). For example, Khumchoo et al. (2015) found that the 560 photoluminescence intensities and energy of zinc oxide in CTA-Sap were controlled by the host-guest 561 interactions of Sap and CTA. Bandla et al. (2016) presented a new process of developing silver 562 nanoparticles via the reduction of silver ions with NaBH₄ in aqueous solutions, using Sap/poly 563 acrylamide (PAAm) nanocomposite hydrogels (Fig. 9). The AgO nanoparticle-containing hydrogel 564 exhibited a strong antibacterial activity against two bacteria. These agents can easily find applications 565 in wound and burn dressings. The incorporated organic compound not only entered the interlayer space of Sap, but also coated on the structural unit layer of the outer surface of Sap. In such 566 inorganic-organic hybrids, the nanospace of Sap can be tuned by spatially controlling the number and 567 568 size (molecular structure) of the organic moieties, which affects their spatial distribution (Okada et al., 569 2014; Marcal et al., 2015).



572 Fig. 9 Schematic representation of formation of AgO nanoparticle hydrogel (Reprinted from Bandla et al., 2017,

573 Copyright 2017, with permission from John Wiley & Sons). KPS: potassium persulfate; TMEDA: N,N,N',N'-tetramethyl

574 ethylenediamine.

575 **6. Applications**

Earlier, Sap and its modified hybrids are often considered to be used as adsorbents (Marcal et al., 576 577 2015; Seki et al., 2015; Sato et al., 2016), catalysts and catalyst supports (Carniato et al., 2014; Fatimah 578 et al., 2016). Over the last decades, increasing studies aim at using Sap as functional additives in CPN 579 (Wu et al., 2014; Wang et al., 2015). It involves improving mechanical properties, gas permeability, 580 degradation, thermal stability and flame retardancy of the CPN. More recently, Sap has also been used 581 as a support for the immobilization of luminescent entities to prevent fluorescence from quenching by suppressing dye molecular aggregation and promoting luminescent energy transfer between donor and 582 583 acceptor dye pairs inside the interlayer space of Sap. In addition, dye molecules may exhibit unique 584 optical imaging (Tsukamoto et al., 2016b) and optoelectronic properties (Nanan et al., 2015).

585 **6.1 Optical materials**

586 Among clay minerals, Sap could be the most ideal host materials for the immobilization of 587 luminescent entities. Firstly, Sap easily delaminates to single nanolayers in water. Secondly, such an 588 aqueous dispersion of Sap is transparent in the range of UV-visible light. Thirdly, Sap itself is not 589 photo or redox active (Tsukamoto et al., 2016b). Fourthly, many luminescent organic dye molecules or 590 cations can be facilely introduced into the interlayer space of Sap to produce clay-based 591 nanocomposites. In this way, the aggregation of luminescent entities can be controlled so their 592 luminescent intensity can be stabilized or enhanced. In addition, the luminescent energy transfer 593 between donor and acceptor dye pairs in the interlayer space of Sap can be promoted. Thus the 594 resultant clay-based nanocomposites exhibit unique optical properties.

595 During fabricating luminescent materials, the aggregation of luminescent dye molecules often 596 occurs. The formation of irregular aggregates, such as H-aggregates, which are sandwich-type 597 structured molecular assemblies (Donauerová et al., 2015), significantly decreases the excited-state 598 lifetime of luminescent dye molecules. Moreover, irregular aggregates could lead to fluorescence 599 quenching or lower photoactivity (Takagi et al., 2013; Bujdák et al., 2016). Hence, suppressing the 600 aggregation of luminescent dye molecules is a critical issue. Saponite has a rigid structure and can 601 provide a confined microenvironment to luminescent molecules on a nanoscale. Hence, in the 602 interlayer space of Sap, J-aggregates with a more open geometry in which the luminescent dye 603 molecules better contact with the surface of Sap become favorable (Epelde-Elezcano et al., 2016). A 604 good dye distribution in the interlayer space of Sap can be achieved reducing molecular aggregation. 605 The net negative charge at the external tetrahedral sheet in the films of Sap permits strong electrostatic 606 Sap-dye interactions in a dye molecule/Sap hybrid. These interactions are conducive to suppressing dye 607 molecule aggregation. They can also increase luminous intensity of the dye molecules and extend the 608 fluorescence lifetime (Wu et al., 2015; Epelde-Elezcano et al., 2016; Boháč et al., 2016; Bujdák et al., 609 2016). In addition, in a confined microenvironment of the interlayer space of Sap, the dye molecules 610 have to occur in a parallel orientation with respect to the surface of the Sap nanoparticles (Tominaga et 611 al., 2017) (Fig. 10). Moreover, dye molecules in the interlayer space of Sap may undergo a 612 conformational change when the thickness of the intercalated molecule is larger than the interlayer 613 spacing of the Sap. Such intercalation renders a pseudo uniaxial pressure to an organic molecule 614 under ordinary temperature and pressure. In a high hydrostatic pressure field, the organic molecule 615 favors a planar conformation. Intercalating the fluorescence dye into the interlayer space of Sap 616 represents a promising progress in developing inorganic-organic luminescent hybrid films of high 617 performance.

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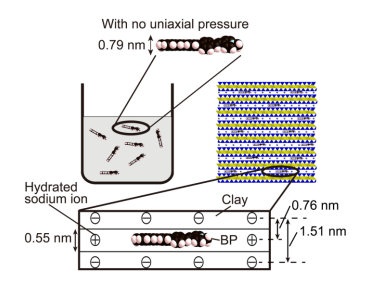


Fig. 10 Schematic representation of molecular confinement of biphenyl derivative (BP) in solution and in a hybrid film fabricated at 27% CEC. Sodium ions and anion sites in the smectite are indicated by circles that contain plus and minus signs, respectively (Reprinted from Tominaga et al., 2017, Copyright 2017, with permission from Springer Nature under the terms of Creative Commons CC By License).

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626 Aggregation of luminescent dye molecule can be further tuned by the co-adsorption of 627 alkylammonium cations because they compete with dye cations for adsorption sites, thereby isolating 628 each dye molecule and suppressing the aggregation (Belusăkova et al., 2015; Donauerová et al., 2015; 629 Epelde-Elezcano et al., 2016). In other words, co-intercalation of alkylammonium cations in the 630 interlayer space of Sap allows potential fluorescent J-aggregates rather than undesirable H-aggregates 631 (Epelde-Elezcano et al., 2016). It is noteworthy that the layer charge of Sap and the solvents also remarkably affect the aggregation of luminescent dye molecule in the interlayer space, hence 632 633 influencing the properties of the Sap/dye hybrids (Wu et al., 2015; Czímerová et al., 2017).

634 Luminescent energy transfer is the dynamic process and occurs via the excited state of a donor dye 635 molecule (Calzaferri, 2012; Verma and Ghosh, 2012; McLaurin et al., 2013). The process plays a 636 critical role in natural and artificial photochemical processes such as photosynthesis, photo-sensitized 637 molecular transformation and solar cells (Gust et al., 2009; Hasobe, 2013). Intercalating luminescent 638 cationic dye species in the confined interlayer space of Sap can promote Förster resonance energy 639 transfer (FRET) between donor and acceptor dye pairs (Ishida et al., 2013; Olivero et al., 2014; Sato et 640 al., 2014b; Belušáková et al., 2015; Goto et al., 2015; Fujimura et al., 2016; Tsukamoto et al., 2016a; 641 Belušáková et al., 2017). In addition, FRET is sensitive to surface concentration, which could be easily 642 controlled by an appropriate selection of the dye/Sap ratio (Belušáková et al., 2017). Promoting FRET 643 between donor and acceptor dye pairs in the interlayer space of Sap provide a new way for developing 644 new artificial light harvesting systems.

645

646 (A)

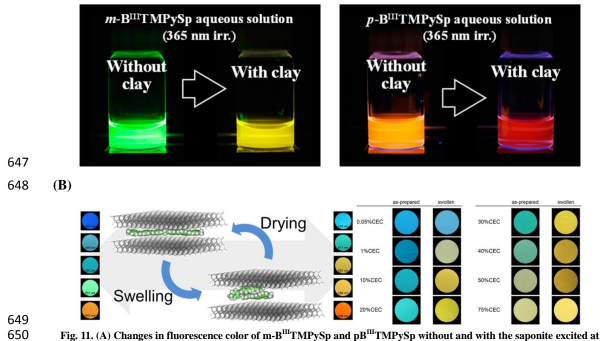


Fig. 11. (A) Changes in fluorescence color of m-B^{III}TMPySp and pB^{III}TMPySp without and with the saponite excited at
365 nm in water. [subporphyrin] = 2.7 × 10⁻⁶ M (0.4% versus CEC of the saponite) (Reprinted with permission from
Tsukamoto et al., 2016b. Copyright (2016) American Chemical Society). (B) Fluorescence colors of saponite–DPDP
hybrid films with various %CEC loading. Left and right columns correspond to the as-prepared and swollen states,
respectively. DPTP:4,4'-[4,1-phenylenedi-2,1-ethynediyl]bis[1-methylpyridinium] diiodide (Adapted and reprinted with
permission from Tominaga et al., 2016. Copyright (2016) American Chemical Society)).

656

657 Luminescent dye-Sap complexes can act as highly sensitive sensors and can be used in artificial 658 photosynthesis systems (Tominaga et al., 2016; Tsukamoto et al., 2016b; Ogawa et al., 2017). For example, Tsukamoto et al. (2016b) intercalated two types of meso-substituted +3-charged subporphyrin 659 derivatives having m-methylpyridinium and p-methylpyridinium (m-B^{III}TMPySp and p-B^{III}TMPySp) 660 661 into the interlayer space of Sap. The photo- or optical properties, with and without the anionic layers of Sap of the two subporphyrin derivatives proved to be different. The absorption and fluorescence 662 spectra of subporphyrin-Sap complexes shifted to longer wavelengths because of luminescent 663 molecules lying flat on the surface of Sap. Changes in fluorescence color of m-B^{III}TMPySp and 664 pB^{III}TMPySp without and with the Sap excited at 365 nm in water can be observed (Fig. 11A). 665 Tominaga et al. (2016) prepared chromic hybrid films consisting of fluorescent organic molecules with 666 667 planar π -conjugated systems and synthetic Sap. The swelling and drying of Sap can reversibly 668 switching fluorescence color (Fig. 11B). In the hybrid films, the organic molecules formed an excimer in the swollen interlayer space of Sap, whereas they were a monomer in the dried interlayer space. The 669 670 emission wavelengths of the excimer and monomer were different, leading to the switching fluorescence color of the hybrid films. Formation and deconstruction of the excimer can be controlled under a mild external stimulus. Moreover, the color switching can be repeated for many cycles. The findings imply that both the confined microenvironment by the interlayer space of the Sap and the reversible swellability can be used for switching optical properties of the resultant hybrids and such intelligent methodology and mentality can be extended to make many other smart materials with switchable properties.

677 6.2 Adsorbents

678 Saponite can be used as cation exchangers and adsorbents to remove radioactive ions in nuclear wastes (Srinivasan, 2011; Sato et al., 2016; Adraa et al., 2017; Sato and Hunger, 2017). It can also be 679 680 applied to separation of heavy metals from wastewater and fixation of the hazardous cations in soil 681 (Budnyak et al., 2016; Franco et al., 2016a; Petra et al., 2017). After organic modification of Sap, the 682 affinity of OSap for organic species is increased and thus can be used to remove dyes and pigments 683 from wastewater (Nakamura and Ogawa, 2013; Makarchuk et al., 2016; Tangaraj et al., 2017). It can 684 also be used to remove carcinogens from foods and food additives (Seki and Ogawa, 2010; Carraro et 685 al., 2014; Marcal et al., 2015; Okada et al., 2015; Seki et al., 2015).

686

6.2.1 Adsorption of metallic cations

687 Heavy metal pollution is a severe problem and threatens ecological systems and human health. 688 Typically, nowadays with the increasing relevant mining and uses of the heavy metal resources. Some 689 waters, groundwater and soil have been polluted by Cd^{2+} , Cu^{2+} , Hg^{2+} , and $Pb^{2+}(Cruz-Guzmán et al.,$ 690 2012; Yang et al., 2016; Zhu et al., 2016). Numerous methods for handling the pollution of the heavy691 metals have been reported in the literature, but little is satisfactory, in view of the high expenses and692 low efficiency (Padilla-Ortega et al., 2016).

693 Over last few years, the use of inorganically pillared, organically modified, chemical and 694 mechanochemical activated Sap has been studied to adsorb and fix heavy metal cations. Franco et al. 695 (2016a) evaluated the effectiveness of adsorption of heavy metal cations (As^{3+} , Cd^{2+} , Cr^{3+} , Cu^{2+} , Hg^{2+} , 696 Ni²⁺, Pb²⁺ and Zn²⁺) at very low concentrations (10–100 ppb) by natural Sap and Fe-pillared and 697 Al-pillared Sap, aiming at purifying the drinking water. Natural Sap and pillared Saps showed notably

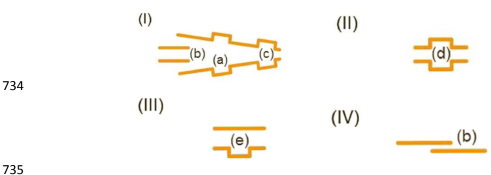
higher adsorption capacities of those cations except Hg^{2+} than carbon used as a reference. In particular, 698 both natural and pillared Sap appeared to be highly efficient to adsorb Cr^{3+} . The adsorption capacity is 699 700 in the order: Fe- pillared Sap> natural Sap> Al-pillared Sap. Recently studies have shown that 701 Sap/chitosan nanocomposites can act as efficient adsorbents for the removal of heavy metals from 702 polluted water. The adsorption behavior is influenced by the pH value of the medium, contact time, and 703 initial metal ion concentration (Budnyak et al., 2016). For such nanocomposite, in addition to the 704 interlayer space of Sap to adsorb the cations, the adsorption can occur through complexation of aqua, 705 acetic, or bi-ligand complexes of the cations with amino groups of chitosan and such formation of 706 complexes heavily depends on the pH value of the medium, namely, acidic basic and neutral one.

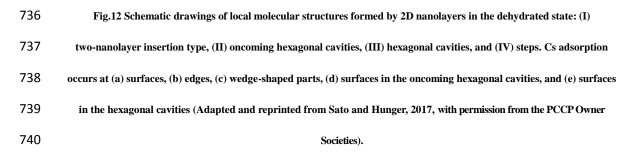
707 Besides chemical modification, mechanochemical activation by high-energy ball milling or 708 grinding have recently been demonstrated to activate Sap for effective fixation of heavy metal cations. 709 Comparatively speaking, mechanochemical activation is a relatively low-cost, effective, and green 710 method to practical application. Such activated Sap can be produced effectively in a planetary ball mill, 711 where the impact energy arising from the ball to ball and ball to wall collisions is transferred to the Sap 712 powder. Such mechanochemical activation can substantially produce fresh and active surfaces for Sap 713 particles. As a result, the surfaces are active for adsorbing heavy metal cations (Vdović et al., 2010). 714 Owing to the existence of many -OH groups on the surface of Sap, the adsorbed metal cations can 715 form insoluble hydroxides on the surface of Sap. Hence the cations are well fixed. More recently, Petra et al. (2017) disclosed that the adsorption for Cu^{2+} by activated Sap was more efficient than for Ni²⁺. 716 717 Nevertheless, for producing effective Sap adsorbents for fixation of heavy metals, tedious screening for 718 the types of metal cations, condition and the modification of Sap remain necessary and challenging.

719 Earlier studies have revealed that radioactive Cs firmly stays on Sap in soil or water (Kim and 720 Kirkpatrick, 1997; Mcbride, 1997; Sato et al., 2016; Dzene et al. 2017). 2D nanolayered Sap can 721 form several types of structures and provide several types of sites for adsorption (Fig. 12) (Sato and 722 Hunger, 2017). The adsorption can occur at the external surfaces (oxygen plane), the broken-bond 723 edges, the wedge-shaped parts, the surfaces in the oncoming hexagonal cavities, and the surfaces in the 724 hexagonal cavities. Consequently, at different sites, the amount and the strength of adsorption appear 725 different due to the different geometrical confinement, physisorption, and chemical bonds. The edges 726 can significantly contribute to the Cs adsorption, along the oxygen planes. Both the oxygen planes 727 and the edges of Sap should be used as the active sites for adsorbing and decontaminating radioactive

728 Cs. Such subtle analysis and findings have much deepened into the insights into the adsorption of Sap 729 for radioactive elements. Finally, it should be pointed out that few studies are reported on the 730 adsorption of radioactive U, Th and Pu ions from nuclear wastes. These elements are more practically 731 used for nuclear energy, so such studies could be more needed (Bian, et al., 2015).

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- 733



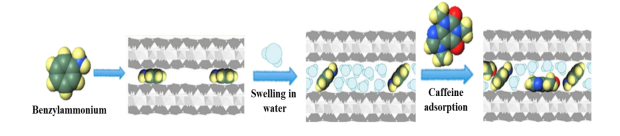


741

742 6.2.2 Removal of carcinogens and caffeine

743 In addition to cation species, the adsorption of nonionic species is also possible through ion-dipole 744 interactions and hydrogen bonding (Theng, 1974). When the cations in the interlayer space of Sap are 745 replaced with organic cations, the surface of the resultant Sap become active for the adsorption of 746 various ionic and nonionic organic species (Ogawa, and Kuroda, 1997). The applications of Sap based 747 on the adsorption is now expanded to the selective adsorption and removal of carcinogens and additives 748 from foods and plants. 2-phenylphenol (also known as 2-hydroxybiphenyl, 2PP) is an antifungal agent 749 used for oranges. On-site removal of 2PP by organically-modified Sap can improve the safety of fresh 750 orange juice (Seki and Ogawa, 2010). Even in the presence of sucrose in the starting aqueous solution, 751 2PP can also be selectively adsorbed on organically-modified Sap. Interestingly, Seki et al. (2015) 752 found that the removal of 2PP by CONH-modified Sap (CONH-Sap) from aqueous sucrose solution

- was even more efficient than that from the solution without sucrose. Monolayer 2PP adsorption to
 CONH-Sap reached 0.067 g/g in the presence of sucrose and 0.062 g/g in the absence of sucrose. The
 basic reason is ascribed to the difference in the 2PP solubility in water and in aqueous sucrose solution.
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- 757



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Fig. 13 Schematic drawing of the process of preparation of benzylammonium (BA)-saponite and the adsorption of
 caffeine over water dispersed BA-saponite (Reprinted and adapted with permission from Okada et al., 2015, Copyright (2015)
 American Chemical Society).

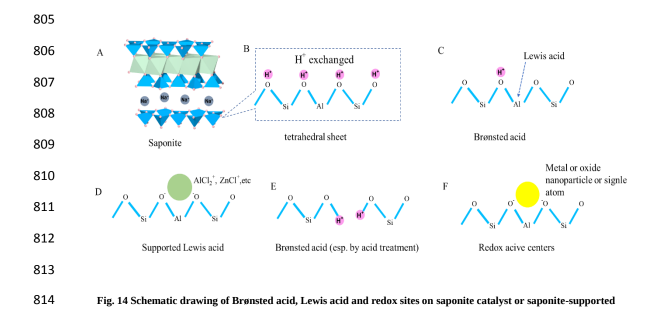
762

763 The large consumption of coffee worldwide has made caffeine a chemical marker or indicator of 764 water pollution. Microorganisms in wastewater usually cannot metabolize the compound effectively. 765 Recently, OSap can act as an efficient adsorbent of caffeine. Adsorption capacity depends primarily on 766 the intercalated organic cations. For example, Okada et al. (2014) have clearly demonstrated the 767 enhanced adsorption of caffeine in water on benzylammonium (BA)-modified Sap (BA-Sap, 768 respectively) (Fig. 13). By contrast, CONH-Sap adsorbed less caffeine than raw Sap and BA-Sap. It 769 was suggested that BA enlarges the siloxane surface area of Sap and make it more available for the 770 adsorption of caffeine. The interlayer space of BA-Sap is further expanded significantly in water by 771 the intercalation of water molecules into it, making the nanospaces suitable for accommodating 772 caffeine molecules. By interacting with BA, the siloxane in the layers of the Sap plays an important 773 role in caffeine adsorption. Marcal et al. (2015) found that CTAB- and APTS-modified synthetic Sap 774 are also efficient absorbents for the adsorption of caffeine from aqueous solutions. The kinetic studies 775 the adsorption fitted into the pseudo-second-order equation, implying that chemical adsorption 776 controlled the process. Comparatively, APTS-Sap exhibited a high affinity for caffeine and possessed a 777 maximum adsorption capacity of 80.54 mg/g after 4 h. The above studies indicated that Sap-based materials are promising candidates for the selective adsorption and removal caffeine from aqueoussolution.

780 6.2.3 Removal of dyes and pigments from wastewater

Dyes and pigments are widely used in the textiles, paper, plastics, leather, and cosmetic industries for coloring products. Most dyes and pigments are toxic and not biodegradable, so they need to be removed from wastewater in industry (Hashemian et al., 2015). A wide range of physicochemical and biological methods for removing dyes and pigments from wastewaters have been studied. The adsorption process proves more effective (Bhatnagar and Sillanpää, 2010). In recent years, there has been increasing interest in using OSap to adsorb organic molecules of dyes and pigments (Rafatullah et al., 2010; Nityashree et al., 2014).

788 As always, the organic modification of Sap increases its affinity for dyes. The OSap can also be 789 made into hydrogel for adsorption. For example, Nakamura and Ogawa (2013) examined the 790 adsorption of a cationic dye (rhodamine 6G) on the spherical particle of a Sap-containing 791 poly(N-isopropylacrylamide) hydrogel. The cation adsorption capacity of the hybrid hydrogel particles 792 was mainly determined by the cation exchange capacity of Sap. Tangaraj et al. (2017) investigated the 793 adsorption capacities of CTA-Sap with three fluorescent dyes, namely Rhodamine 640 perchlorate 794 rhodamine (Rho), sulforhodamine B (SR), and Kiton red 620 (KR). The edges and faces of Sap 795 particles was found responsible for the adsorption of dyes. The adsorption isotherms fitted well into the 796 non-linear Langmuir isotherm model. Most of the dye molecules can be released in aqueous solution. 797 Another issue is that after adsorption, the separation of Sap particles from the aqueous solution are 798 considerable difficulty due to their easy delaminate and high dispersion. To overcome this, a suggested 799 method is to endow saponite particles with magnetic properties (Chen et al., 2016). As a result, 800 magnetized Sap can be separated by the simple procedure of magnetic separation and removed from 801 the water (Xu et al., 2012; Giakisikli and Anthemidis, 2013). Moreover, magnetite/Sap nanocomposites 802 can possess a microporous and mesoporous structure and thus have enhanced adsorption performance 803 in comparison to Sap alone and magnetite alone (Makarchuk et al., 2016).



catalyst.

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Over the past few decades, many strategies have been developed to produce clay minerals-based catalysts (Zhou, 2011). In the layer of Sap, isomorphous substitution of some Si(IV) by Al(III) in the tetrahedral sheet not only make the layer to be negatively charged but also creates Lewis acidity. The exchangeable cations in the interlayer space of Sap, typically Na⁺, K⁺, and Ca²⁺, which compensate the layer charge, can be exchanged with H⁺, thereby creating Brønsted acidity (**Fig. 14B**).

822 more Brønsted acid sites can be created through H⁺ coordination or silanol group –Si-O-H⁺ (Fig. 14C).

Besides.

823 Such acidity can be enhanced by proper acid treatment of Sap to partly remove Al and create more

824 edges or broken bonds (Fig. 14E). Many studies have proved that Al in the tetrahedral sheet exhibits

825 Lewis acidity. In addition, two-dimensional nanolayer possesses the large surface and the surface can

826 be used to support Lewis acid reagents like AlCl₃, ZnCl₂ and FeCl₃ (Fig. 14D). Such Sap prove to be an

827 efficient catalyst for acid-catalyzed organic reactions, for example the synthesis of β -amino ketones via

828 Mannich-type reactions under mild conditions (Gómez-Sanz et al., 2017). In addition, the synthesis of

829 Sap can be easily adapted to introduce some metal or metal oxide centers in the octahedral or

830 tetrahedral sheet of Sap so that the M-Sap are able to catalyze redox reactions (Fig. 14F). Moreover,

831 due to its easily and controllably expandable interlayer space with large surface area, Sap can also be

832 used as catalyst supports. (Table 4)

Catalyst	Reaction	S _{BET} (m²/g)	T (K)	t (h)	Conversion (%)	Yield (%)	Ref.
Cs (1.25 wt%)/Al ₁₃ -pillared Sap	Synthesis of N-alkyl pyrazoles	47	353	1	100	100	Velasco et al. (2011)
Ni (20 wt%)/Sap	Hydrogenation of styrene oxide	RT	344	1	100	100	Vicente et al. (2011)
MgFe-Sap NiMgAl-Sap	Epoxidation of (Z)-cyclooctene by hydrogen peroxide	467 334	RT	48	7.5 8.8	7.5 8.8	Trujillano et al. (2011)
Acid activated V(0.05mmol/g) / Sap ¹	Oxidative dehydrogenation of propene into coke	-	773	0.5	21	-	Ostinelli et al. (2012)
Zirconia-cluster (1mmol/g)/ Sap	Isomerization of 1-butene	410	573	2.5	82	-	Kameshima et al. (2013)
Nb ^V /Sap	Oxidative abatement of CEES	-	298	8	98	71.5*	Carniato et al. (2014)
TiO ₂ (0.42 g/L) /[Ru(bpy) ₃] ²⁺ -Sap	Oxidation of benzene to phenol	-	VIS	5	72	69	Goto and Ogawa (2016)
ZnO/Sap	Photocatalytic decomposition of Methylene Blue	168	UV	3	93	-	Khumchoo et al. (2016)
Sulphated zirconium pillared Sap	Conversion of citronellal	190*	MW	10 min	100	-	Fatimah et al. (2016)
Cu (5 wt%)/Na-Sap	Diesel soot combustion	-	673	-	100	99	Gebretsadik et al. (2016)
Ni (40 wt%)/H-Sap	Glycidol hydrogenolysis to 1,3-propanediol	229	453	1	100	29	Gebretsadik et al. (2017b)

Table 4. Catalytic activity of saponite (Sap) and Sap supported catalysts.

Re (7 wt%) modified Ni (40 wt%)/ acid Sap	Glycidol hydrogenolysis to 1,3-propanediol	331	393	4	98	46.1	Gebretsadik et al. (2017a)
Raw Sap	Synthesis of β-amino ketones by Mannich reaction	131	318	5	80	-	Gómez-Sanz et al. (2017)

* For clarity, consistency, and easy comparison, some original data have been rounded off.

836 T: Temperature; t: Time; VIS: Visible light irradiation; UV: UV irradiation; MW: Microwave irradiation; RT: Room temperature;

837 CEES: (2-chloroethyl)ethyl sulfide.

838

839 The incorporation of transition elements, such as Ni (Trujillano et al., 2011), Fe (Trujillano et al., 840 2011), V (Ostinelli et al., 2012), and Nb (Carniato et al., 2014), into the framework of Sap, including 841 acid activated Sap, allows the combination of high surface area and porosity of the Sap with the solid 842 acidic and redox catalytic active centers. Theoretically, the transition metal atoms can be incorporated 843 in the octahedral or tetrahedral sheets of Sap. When transition elements are introduced into the 844 tetrahedral sheet of Sap, the co-existence of active transition metal cations, for example V (Ostinelli et 845 al., 2012) and Nb cations (Carniato et al., 2014), and the tetrahedral silicon cations, a synergetic effect 846 of the redox properties of the transition metals and the Brønsted acid sites makes the transition 847 metal-containing Sap an effective catalyst (Ostinelli et al., 2012; Carniato et al., 2014). Recent studies 848 demonstrated that the transition metal-containing Sap can be used to catalyze the epoxidation of 849 (Z)-cyclooctene by hydrogen peroxide (Trujillano et al., 2011), the conversion of propene into coke 850 (Ostinelli et al., 2012) and the oxidative abatement of chemical warfare agents (Carniato et al., 2014). 851 Moreover, the active cations in the framework of Sap could be more stable and leaching of the active 852 metallic cations during catalytic reaction in a liquid phase can be mitigated to some extent (Trujillano 853 et al., 2011). Pillaring treatments of Sap can increase the SSA, the pore volume and the surface acidity 854 and re-dox catalysis of Sap. In particular, pillars in the pillared Sap can be designed and made with 855 additional specific catalytic performances (De Stefanis et al., 2013; Zhen et al., 2014; Zhen and Zheng, 856 2016). In pillared Sap, the major sources for Brønsted and Lewis sites are associated with the lattice 857 hydroxyls of Sap and the pillars in the interlayer space of Sap. For example, Kameshima et al. (2013) 858 successfully prepared zirconia-cluster pillared Sap and revealed that both the amounts of Brønsted and 859 Lewis acids remarkably increased after pillaring Sap with zirconia-cluster. The zirconia-cluster was a 860 main source for generating Lewis acidity besides the tetra-coordinated Al in the tetrahedral sheet of Sap while Brønsted acidity was provided by H⁺ coordination and silanol group generated by removal of
some Al from the tetrahedral sheet of Sap (Fig. 14E). Recently, regarding sulphated zirconia-pillared
Sap, Fatimah et al. (2016) discovered that the sulphate anions on the surface can increase the number
and strength of the Brönsted acid sites of zirconia-pillared Sap.

865 Saponite, acid activated Sap, pillared Sap, and other related hybrids can also be used as a catalyst 866 support. The support offers expandable interlayer space, high SSA, surface acidity, and good thermal 867 stability (Garciano et al., 2014; Gebretsadik et al., 2016; Khumchoo et al., 2016). In particular, in view 868 of synthesis, swellability, surface acidity, and thermal stability, Sap has an advantage over Mt. Such 869 Sap-supported catalysts can catalyze a wide range of reactions. Recent examples are Cs/Al-pillared Sap 870 catalysts for the synthesis of N-alkyl pyrazoles (Velasco et al., 2011), Ni/Sap catalysts for the 871 hydrogenation of styrene oxide (Vicente et al., 2011), Co-Mo/Sap catalysts for upgrading squalene 872 (C₃₀H₅₀) (Garciano et al., 2014), TiO₂/Al-pillared Sap catalysts for photocatalytic phenol 873 photo-oxidation (Fatimah and Wijaya, 2015), Ni/Zr-pillared Sap catalysts for hydrogenation of isopulegol (Fatimah et al., 2015), TiO₂ /[Ru(bpy)₃]²⁺-Sap catalysts for photocatalytic oxidation of 874 875 benzene to phenol (Goto and Ogawa, 2015, 2016), ZnO/Sap catalysts for photodegradation of 876 Methylene Blue (Khumchoo et al., 2016), Cu/Sap catalysts for soot combustion (Gebretsadik et al., 877 2016), and Ni (Gebretsadik et al., 2017b) and MO_x-modified nickel-copper catalysts (Gebretsadik et al., 878 2017a) supported on H-Sap for glycidol hydrogenolysis (Table 4). Saponite-supported catalysts are 879 much stable even under irradiation and acid conditions (Khumchoo et al., 2016). In addition, a more 880 recent study form Gebretsadik et al. (2017b) indicated that the acidity of H-Sap is conducive to the 881 dispersion and reduction of NiO catalyst on the surface of H-Sap. The NiO and Ni particles could play 882 a role in changing the amount and strength of surface acid sites of the Ni /Sap (Gebretsadik et al., 883 2017a). As proved by many reactions, there are indeed interactions between Sap support and metal 884 atoms or metal oxides and the two can have a synergistic catalytic effect. However, the details remain 885 unclear.

886 **6.4 Polymer additives**

Recently, Sap and organo-Sap has been used as reinforcing fillers in CPN (Table 3). Basically, it
has been considered to possess the same functions as Mt nanolayers. By contrast, Sap could be more

easily dispersed in the polymer matrix. Compared with pure polymers, CPN exhibit remarkable
improvement in mechanical strength, Young's modulus, gas barrier properties, thermal stability, and
water resistance (Zhen et al., 2012; Jairam et al., 2013; Yang et al., 2013; Zhen and Wang, 2016). The
enhancement of those properties of the CPN is commonly ascribed to the existence of well-dispersed
Sap nanolayers with a high aspect ratio in the CPN (Chang et al., 2014; Zhen and Wang, 2016).

894 In addition, Sap can act as a heterogeneous nucleating agent and accelerate the crystallization rate 895 of polymers (Xi et al., 2015; Zhen and Wang, 2016). When the crystallization is improved, it also helps 896 improve the thermal stability of the CPN (Zhen et al., 2014; Zhen and Zheng, 2016). Furthermore, 897 during the *in situ* polymerization process, Sap can act as a green catalyst which catalyzes the 898 polymerization reactions. Besides, Sap as adsorbents can contribute to adsorbing and eliminating the 899 toxic chemicals during the polymerization process (Zhen et al., 2014; Zhen and Zheng, 2016). More 900 recently, the CPN have been endowed with antibacterial properties. (Xi et al., 2015; Bandla et al., 901 2017). This broadens the functions of Sap and the application of the CPN.

902 It is noteworthy that the amount of Sap in the polymer matrix has a significant effect on the 903 properties of the resultant Sap/polymer nanocomposite. In many cases, the addition of a small amount 904 of Sap is sufficient to improve the thermal and mechanical properties and gas barrier. For example, 905 Chang et al. (2014) found that for even 3-10 wt.% Sap in the PVA matrix, the CPN exhibited much 906 higher transition temperature than pure PVA. Ju and Chang (2016) found that upon the addition of Sap 907 into PI matrix up to a critical loading amount of Sap (20 wt.%), the glass transition temperatures (T_{o}) and the initial thermal degradation temperature (Ti_{D}^{i}) of the Sap/PI hybrid films were remarkably 908 909 improved. Nevertheless, the coefficient of thermal expansion (CTE) decreased. If Sap was added more 910 than the critical content, the thermal properties worsened. Wu et al. (2014) found that the Sap/TOCN 911 nanocomposite with 10 wt.% Sap exhibited the Young's modulus of 14.5 GPa, tensile strength of 425 912 MPa, and strain-to-failure of 10.2%. Due to the addition of rigid Sap, the resistance to fracture of the 913 Sap/TOCN nanocomposite increased more than sevenfold when the loading amount of Sap is increased 914 from 0 to 10 wt.%. Finally, it is worth mentioning that Sap is not always just an additive for polymers. 915 In some cases, both the polymer and Sap are main components of the CPN, in particular as it is in the 916 hybrid films (Zhou et al., 2011). In addition, Sap can be the main component and the polymers could be 917 used with a lower amount to improve some functions (Shin et al., 2013).

918 **7. Concluding remarks and future work**

919 Last decade has witnessed great ad advances in modification and hybridization spanning from 920 simple interaction and surface engineering to fabrication of nanocomposite and hierarchical assembly. 921 The researchers have smartly developed many strategies for introducing functional inorganic, organic 922 and organic-inorganic guests into the interlayer space of Sap. In addition to acid activation, inorganic 923 cation exchange, pillaring and catalyst supporting, the nanoscale interlayer space of Sap is increasingly 924 being considered as a nanoreactor to produce a functional metal or metal oxide nanoparticle/Sap hybrid 925 with magnetic, selective adsorptive and catalytic properties. Besides OSap and CPN, organically 926 modified Sap and polymers can be made into functional films and such materials show promising 927 applications in optics, biological and medical materials.

Undoubtedly, the great strides in the approaches to the modification and hybridization of Sap, along with the upsurge of inorganic nanotechnology and the discovery of functional organic molecules, provide many opportunities to use Sap for producing advanced materials. Meanwhile, previous studies have highlighted some challenges and problems which need laddering, as prudently and briefly remarked as follows:

933 (1) To the best of our knowledge, Sap is a scarce natural resource and is not as abundant as Mt; At present, the synthesis, modification and applications of Sap or the related studies is still limited on a 934 935 laboratory scale. In order to scale up the modification and application of Sap, some critical issues need 936 to be resolved. For example, a cleaner, efficient and profitable process of synthetic Sap should be first 937 developed. For the modification of Sap, the wastewater and the toxic solvents need handling properly. In most cases, successful modification of Sap is usually a multiple-step process, which is 938 939 time-consuming and cost-intensive. A good alternative way could be to integrate the synthesis with the 940 modification of Sap. The one-pot reaction significantly shortens the process and simplifies the 941 procedure, thereby lowering the cost. A few studies have shed new light on this technology. However, it 942 is unknown if many other organic species can be introduced into the Sap in this way. Alternatively, a 943 supercritical continuous flow process could worth be applied to synthesize and modify Sap. Such 944 intensified process can improve mixing of reactants, mass and heat transfer and accordingly shorten the 945 reaction and improve productivity.

946 (2) Although the methods of modifying Sap have been greatly expanded, few studies provide the 947 mechanisms, thermodynamics, and kinetics. Theoretically, molecular dynamics simulations and 948 experimental cryo-electron microscopy (cryo-EM) and other scientific tools are helping throw new 949 light on the mechanisms. In order to get a better understanding of the modification and scale up the 950 relevant process, detailed investigations and data for engineering are of basic necessity.

951 (3) There is a need to put Sap and Sap-based materials onto the practice in industry. A critical issue is 952 to modify Sap with functional species with the practical applications. As Sap is a good player in the 953 hydrogel system, a combination of magnetic nanoparticles with Sap to produce magnetic 954 nanoparticles/Sap hydrogel can bring exceptional properties in magnetism, rheology, adsorption, 955 catalysis, and biocompatibility. Thus, the clay-based nanocomposites have potential to be applied in 956 electromagnetic devices, magnetorheological fluids/ferrofluids, magnetic adsorbents, catalysts, 957 biomaterials, biomedicines or highly simulated robots. In addition, there are various choices of 958 functional guest species that exhibit exceptional biological, electrical, and optical properties and could 959 to be introduced into Sap to produce new functional materials.

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- 972 973
- 974 References 975
- Adraa, K. E., Georgelin, T., Lambert, J. F., Jaber, F., Tielens, F., Jaber, M., 2017. Cysteine-montmorillonite composites for heavy metal cation complexation: a combined experimental and theoretical study. Chemical Engineering Journal, 314, 406-417.
- Albeniz, S., Vicente, M. A., Trujillano, R., Korili, S. A., Gil, A., 2014. Synthesis and characterization of organosaponites.
 Thermal behavior of their poly (vinyl chloride) nanocomposites. Applied Clay Science, 99, 72-82.
- Baldermann, A., Dohrmann, R., Kaufhold, S., Nickel, C., Letofsky-Papst, I., Dietzel, M., 2014. The Fe-Mg-saponite solid solution series-a hydrothermal synthesis study. Clay Minerals, 49(3), 391-415.

Bandla, M., Abbavaram, B. R., Kokkarachedu, V., Sadiku, R. E., 2017. Silver nanoparticles incorporated within intercalated
 clay/polymer nanocomposite hydrogels for antibacterial studies. Polymer Composites, 38(S1), E16-E23

Belušáková, S., Lang, K., Bujdák, J., 2015. Hybrid systems based on layered silicate and organic dyes for cascade energy transfer.
 The Journal of Physical Chemistry C, 119(38), 21784-21794.

⁹⁸⁶ Belušáková, S., Martínez-Martínez, V., Arbeloa, I. L., Bujdak, J., 2017. Resonance Energy Transfer between Dye Molecules in

- 987 Colloids of a Layered Silicate. The Effect of Dye Surface Concentration. The Journal of Physical Chemistry C, 121(15), 988 8300-8309
- 989 Bergaoui, L., Lambert, J. F., Franck, R., Suquet, H., Robert, J. L., 1995a. Al-pillared saponites. part 3. - effect of parent clay layer 990 charge on the intercalation. Journal of the Chemical Society Faraday Transactions, 91(14), 2229-2239.
- 991 Bergaoui, L., Lambert, J. F., Vicenterodriguez, M. A., Michot, L. J., Villieras, F., 1995b. Porosity of synthetic saponites with 992 variable layer charge pillared by Al₁₃ polycations. Langmuir, 11(8), 2849-2852.
- 993 Bhatnagar, A., Sillanpää, M., 2010. Utilization of agro-industrial and municipal waste materials as potential adsorbents for water 994 treatment-a review. Chemical Engineering Journal, 157(2), 277-296.
- 995 Bian, L., Dong, F. Q., Song, M. X., Dong, H. L., Li, W. M., & Duan, T., Xu, J. B., Zhang, X. Y., 2015. DFT and two-dimensional
- 996 correlation analysis methods for evaluating the Pu³⁺-Pu⁴⁺ electronic transition of plutonium-doped zircon. Journal of Hazardous 997 Materials, 294(8), 47-56.
- 998 Bian, L., Song, M. X., Dong, F. Q., Duan, T., Xu, J. B., Li, W. M., Zhang, X. Y., 2015. DFT and two-dimensional correlation 999 analysis for evaluating the oxygen defect mechanism of low- density 4f (or 5f) elements interacting with Ca-Mt. RSC Advances,
- 1000 5, 28601-28610.Bisio, C., Carniato, F., Paul, G., Gatti, G., Boccaleri, E., Marchese, L., 2011. One-pot synthesis and 1001 physicochemical properties of an organo-modified saponite clay. Langmuir, 27(11), 7250-7257.
- 1002 Bisio, C., Gatti, G., Boccaleri, E., Marchese, L., Bertinetti, L., Coluccia, S., 2008. On the acidity of saponite materials: a 1003 combined HRTEM, FTIR, and solid-state NMR study. Langmuir, 24(6), 2808-2819.
- 1004 Boháč, P., Czímerová, A., Bujdák, J., 2016. Enhanced luminescence of 3, 3'-diethyl-2, 2'-thiacyanine cations adsorbed on saponite particles. Applied Clay Science, 127, 64-69. 1005
- 1006 1007 Brigatti, M. F., Galan, E., Theng, B. K. G., 2013, Structure and Mineralogy of Clay Minerals. In: Bergaya, F., Lagaly, G. (Eds.), Developments in Clay Science, Vol. 5, Handbook of Clay Science, Second Edition. Amsterdam, Elsevier, pp. 21-81
- 1008 Budnyak, T. M., Yanovska, E. S., Kichkiruk, O. Y., Sternik, D., Tertykh, V. A., 2016. Natural minerals coated by biopolymer 1009 chitosan: synthesis, physicochemical, and adsorption properties. Nanoscale Research Letters, 11(1), 492-503.
- 1010 Budnyak, T., Tertykh, V., Yanovska, E., 2014. Chitosan immobilized on silica surface for wastewater treatment. Materials 1011 Science, 20(2), 177-182.
- 1012 Bujdák, J., Ratulovská, J., Donauerová, A., Bujdáková, H., 2016. Hybrid materials based on luminescent alkaloid berberine and 1013 saponite. Journal of Nanoscience and Nanotechnology, 16(8), 7801-7804.
- 1014 Calzaferri, G., 2012. Nanochannels: hosts for the supramolecular organization of molecules and complexes. Langmuir 28, 1015 6216-6231.
- 1016 Carniato, F., Bisio, C., Gatti, G., Boccaleri, E., Bertinetti, L., Coluccia, S., Monticelli, O., Marchese, L., 2009. 1017 Titanosilsesquioxanes embedded in synthetic clay as a hybrid material for polymer science. Angewandte Chemie, 121(33), 1018 6175-6177
- 1019 Carniato, F., Bisio, C., Gatti, G., Guidotti, M., Sordelli, L., Marchese, L., 2011. Organic-Inorganic Hybrid Saponites Obtained by 1020 Intercalation of Titano - Silsesquioxane. Chemistry-An Asian Journal, 6(3), 914-921.
- 1021 Carniato, F., Bisio, C., Psaro, R., Marchese, L., Guidotti, M., 2014. Niobium (V) saponite clay for the catalytic oxidative 1022 abatement of chemical warfare agents. Angewandte Chemie International Edition, 53(38), 10095-10098.
- 1023 Carrado K.A., Decarreau A., Petit S., Bergaya F., Lagaly G., 2006. Synthetic clay minerals and purification of natural clays, 1024 1025 Chapitre 4. In: Bergaya F., Theng B., Lagaly G. (Eds.), Developments in Clay Science, Vol. 1, Handbook of Clay Science, Elsevier, pp. 115-139.
- 1026 Carraro, A., De Giacomo, A., Giannossi, M. L., Medici, L., Muscarella, M., Palazzo, L., Quaranta, V., Summa, V., Tateo, F., 2014. 1027 Clay minerals as adsorbents of aflatoxin M 1 from contaminated milk and effects on milk quality. Applied Clay Science, 88, 1028 92-99.
- 1029 Carter, D.L., Heilman, M.D., Gonzalez, C.L., 1965. Ethylene glycol monoethyl ether for determining surface area of silicate
- 1030 minerals. J. Soil Sci. 100, 356-360.
- 1031 Casagrande, M., Storaro, L., Lenarda, M., Rossini, S., 2005. Solid acid catalysts from clays: Oligomerization of 1-pentene on 1032 Al-pillared smectites. Catalysis Communications, 6(8), 568-572.
- 1033 Chang, J. H., Ham, M., Kim, J. C., 2014. Comparison of Properties of Poly (vinyl alcohol) Nanocomposites Containing Two 1034 Different Clays. Journal of nanoscience and nanotechnology, 14(11), 8783-8791.
- 1035 Chen, L., Zhou, C.H., Fiore, S., Tong, D.S., Zhang, H., Li, C.S., Ji S.F., Yu, W.H., 2016. Functional magnetic nanoparticle/clay 1036 mineral nanocomposites: preparation, magnetism and versatile applications Applied Clay Science. 127-128, 143-163
- 1037 Chevalier, S., Franck, R., Suquet, H., Lambert, J. F., Barthomeuf, D., 1994. Al-pillared saponites. part 1.- IR studies. Journal of
- 1038 the Chemical Society Faraday Transactions, 90(4), 667-674.
- 1039 Crini, G., Badot, P. M., 2008. Application of chitosan, a natural aminopolysaccharide, for dye removal from aqueous solutions by 1040 adsorption processes using batch studies: a review of recent literature. Progress in polymer science, 33(4), 399-447.
- 1041 Cruz-Guzmán, M., Celis, R., Hermosín, M. C., Koskinen, W. C., Nater, E. A., Cornejo, J., 2012. Heavy metal adsorption by
- 1042 1043 montmorillonites modified with natural organic cations. Soil Science Society of America Journal, 70(1), 215-221. Czímerová, A., Sas, S., Čeklovský, A., 2017. Unique solvatochromism of cyanine/saponite hybrid systems. Applied Clay Science,
- 1044 198-203. 1045 Dazas, B., Lanson, B., Delville, A., Robert, J.L., Komarmeni, S., Michot, L.J., Ferrage E., 2015. Influence of tetrahedral layer 1046 charge on the organization of interlayer water and ions in synthetic Na-saturated smectites. Journal of Physical Chemistry C, 119,
- 1047 4158-4172. 1048 De Paiva, L.B., Morales, A.R., Valenzuela Díaz, F.R., 2008. Organoclays: properties, preparation and applications. Applied Clay 1049 Science, 42, 8-24.
- 1050 De Stefanis, A., Cafarelli, P., Gallese, F., Borsella, E., Nana, A., Perez, G., 2013. Catalytic pyrolysis of polyethylene: A 1051 1052 comparison between pillared and restructured clays. Journal of Analytical and Applied Pyrolysis, 104, 479-484.
- Ding, Z., Kloprogge, J. T., Frost, R. L., Lu, G. Q., Zhu, H. Y., 2001. Porous clays and pillared clays-based catalysts. Part 2: a 1053 1054 review of the catalytic and molecular sieve applications. Journal of Porous Materials, 8(4), 273-293.
- Donauerová, A., Bujdák, J., Smolinská, M., Bujdáková, H., 2015. Photophysical and antibacterial properties of complex systems

- 1055 based on smectite, a cationic surfactant and methylene blue. Journal of Photochemistry & Photobiology B Biology, 151, 135-141.
- 1056 Dong, F. Q., Bian, L., Song, M. X., Li, W. M., Duan, T., 2016. Computational investigation on the $f^n \rightarrow f^{n-1}d$ effect on the
- 1057 electronic transitions of clinoptilolite. Applied Clay Science, 119,74-81.
- 1058 Dzene, L., Verron, H., Delville, A., Michot, L.J., Robert, J.L., Tertre, E., Hubert, F., Ferrage, E., 2017. Influence of tetrahedral 1059 layer charge on the fixation of cesium in synthetic smectite. Journal of Physical Chemistry C, 121, 23422-23435.
- 1060 Ebina, T., Mizukami, F., 2007. Flexible transparent clay films with heat - resistant and high gas - barrier properties. Advanced 1061 Materials, 19(18), 2450-2453.
- 1062 Eguchi, M., Momotake, M., Inoue, F., Oshima, T., Maeda, K., Higuchi, M., 2017. Inert Layered Silicate Improves the 1063 1064 Electrochemical Responses of a Metal Complex Polymer. ACS Applied Materials & Interfaces, 9(40), 35498-35503.
- Epelde-Elezcano, N., Martínez-Martínez, V., Duque-Redondo, E., Temiño, I., Manzano, H., López-Arbeloa, I., 2016. Strategies 1065 for modulating the luminescence properties of pyronin Y dye-clay films: an experimental and theoretical study. Physical 1066 1067 Chemistry Chemical Physics, 18(12), 8730-8738.
- Fatimah, I., Rubiyanto, D., Huda, T., 2015. Preparation and Characterization of Ni/Zr-Saponite as Catalyst in Catalytic Hydrogen 1068 Transfer Reaction of Isopulegol. In Materials Science Forum (Vol. 827, pp. 311-316). Trans Tech Publications.
- 1069 Fatimah, I., Rubiyanto, D., Huda, T., Zuhrufa, Z., Yudha, S. P., Kartika, N. C., 2016. Novel sulphated zirconia pillared clay 1070 nanoparticles as catalyst in microwave assisted conversion of citronellal. Materials Technology, 31(4), 222-228.
- 1071 Fatimah, I., Wijaya, K., 2015. Microwave assisted preparation of TiO₂/Al-pillared saponite for photocatalytic phenol 1072 photo-oxidation in aqueous solution. Arabian Journal of Chemistry, 8(2), 228-232.
- 1073 Ferrage, E., 2016. Investigation of the interlayer organization of water and ions in smectite from the combined use of diffraction 1074 1075 experiments and molecular simulations. A review of methodology, applications, and perspectives. Clays and Clay Minerals, 64, 346-371.
- 1076 Ferrage, E., Lanson, B., Michot, L.J., Robert J.L., 2010. Hydration properties and interlayer organization of water and ions in 1077 1078 synthetic Na-smectite with tetrahedral layer charge. Part 1. Results from X-ray diffraction profile modeling. Journal of Physical Chemistry C, 114, 4515-4526.
- 1079 Franco, F., Benítez-Guerrero, M., Gonzalez-Triviño, I., Pérez-Recuerda, R., Assiego, C., Cifuentes-Melchor, J., Pascual-Cosp, J., 1080 2016a. Low-cost aluminum and iron oxides supported on dioctahedral and trioctahedral smectites: a comparative study of the 1081 effectiveness on the heavy metal adsorption from water. Applied Clay Science, 119, 321-332.
- 1082 Franco, F., Pozo, M., Cecilia, J. A., Benítez-Guerrero, M., Lorente, M., 2016b. Effectiveness of microwave assisted acid 1083 treatment on dioctahedral and trioctahedral smectites. The influence of octahedral composition. Applied Clay Science, 120, 1084 70-80.
- 1085 Fujimura, T., Ramasamy, E., Ishida, Y., Shimada, T., Takagi, S., Ramamurthy, V., 2016. Sequential energy and electron transfer in 1086 a three-component system aligned on a clay nanosheet. Physical Chemistry Chemical Physics, 18(7), 5404-5411.
- 1087 Garade, A. C., Biradar, N. S., Joshi, S. M., Kshirsagar, V. S., Jha, R. K., Rode, C. V., 2011. Liquid phase oxidation of p-vanillyl 1088 alcohol over synthetic Co-saponite catalyst. Applied Clay Science, 53(2), 157-163.
- 1089 Garciano II, L. O., Tran, N. H., Kannangara, G. K., Milev, A. S., Wilson, M. A., Volk, H., 2014. Developing saponite supported 1090 cobalt-molybdenum catalysts for upgrading squalene, a hydrocarbon from the microalgae Botryococcus braunii. Chemical 1091 Engineering Science, 107, 302-310.
- 1092 Gebrestadik, F. B., Llorca, J., Salagre, P., Cesteros, Y., 2017a. Hydrogenolysis of glycidol as an alternative route to obtain 1093 1,3-propanediol selectively using Mox -modified nickel-copper catalysts supported on acid mesoporous saponite. Chemcatchem, 1094 9(19), 3670-3680.
- 1095 Gebretsadik, F. B., Cesteros, Y., Salagre, P., Giménez-Mañogil, J., Garcia-Garcia, A., Bueno-López, A., 2016, Potential of 1096 1097 Cu-saponite catalysts for soot combustion. Catalysis Science & Technology, 6(2), 507-514.
- Gebretsadik, F. B., Ruiz-Martinez, J., Salagre, P., Cesteros, Y., 2017b. Glycidol hydrogenolysis on a cheap mesoporous acid 1098 saponite supported Ni catalyst as alternative approach to 1, 3-propanediol synthesis. Applied Catalysis A: General, 538, 91-98.
- 1099 1100 Gebretsadik, F. B., Salagre, P., Cesteros, Y., 2014. Use of polymer as template in microwave synthesis of saponite. Study of several factors of influence. Applied Clay Science, 87, 170-178.
- 1101 Giakisikli, G., Anthemidis, A. N., 2013. Magnetic materials as sorbents for metal/metalloid preconcentration and/or separation. A 1102 1103 review. Analytica chimica acta, 789, 1-16.
- Gil, A., Gandia, L. M., Vicente, M. A., 2000. Recent advances in the synthesis and catalytic applications of pillared clays. 1104 Catalysis Reviews, 42(1-2), 145-212.
- 1105 Gómez-Sanz, F., Morales-Vargas, M. V., González-Rodríguez, B., Rojas-Cervantes, M. L., Pérez-Mayoral, E., 2017. Acid clay 1106 minerals as eco-friendly and cheap catalysts for the synthesis of β-amino ketones by Mannich reaction. Applied Clay Science, 1107 1108 143, 250-257.
- Goto, T., Ogawa, M., 2015. Visible-light-responsive photocatalytic flow reactor composed of titania film photosensitized by 1109 metal complex-clay hybrid. Acs Applied Materials & Interfaces, 7(23), 12631-12634.
- 1110 Goto, T., Ogawa, M., 2016. Efficient photocatalytic oxidation of benzene to phenol by metal complex-clay/TiO2 hybrid 1111 photocatalyst. Rsc Advances, 6(28), 23794-23797.
- 1112 Guidotti, M., Psaro, R., Ravasio, N., Sgobba, M., Carniato, F., Bisio, C., Gatti, G., Marchese, L., 2009. An efficient ring opening 1113 reaction of methyl epoxystearate promoted by synthetic acid saponite clays. Green Chemistry, 11(8), 1173-1178.
- 1114 Gust, D., Moore, T.A., Moore, A.L., 2009. Solar fuels via artificial photosynthesis. Acc Chem Res, 42(12), 1890-1898.
- 1115 Hashemian, S., Dehghanpor, A., Moghahed, M., 2015. Cu_{0.5}Mn_{0.5} Fe₂O₄ nano spinels as potential sorbent for adsorption of 1116 brilliant green. Journal of Industrial and Engineering Chemistry, 24, 308-314.
- 1117 Hasobe, T., 2013. Porphyrin-based supramolecular nanoarchitectures for solar energy conversion. Journal of Physical Chemistry 1118 Letters, 4(11), 1771-1780.
- 1119 1120 He, H., Duchet, J., Galy, J., Gerard, J. F., 2005. Grafting of swelling clay materials with 3-aminopropyltriethoxysilane. Journal of colloid and interface science, 288(1), 171-176.
- 1121 1122 He, H.P., Li, T., Tao, Q., Chen, T.H., Zhang, D., Zhu, J.X., Yuan, P., Zhu, R.L., 2014. Aluminum ion occupancy in the structure of synthetic saponites: effect on crystallinity. American Mineralogist. 99(1), 109-116.
- 1123 Herrera, N.N., Letoffe, J.M., Putaux, J.L., David, L., Bourgeat-Lami, E., 2004. Aqueous dispersions of silane-functionalized laponite clay platelets. A first step toward the elaboration of water-based polymer/clay nanocomposites. Langmuir 20, 1124 1125 1564-1571.
- 1126 Honda, M., Shimoyama, I., Okamoto, Y., Baba, Y., Suzuki, S., & Yaita, T. 2016. X-ray absorption fine structure at the cesium L₃
- 1127 absorption edge for cesium sorbed in clay minerals. Journal of Physical Chemistry C, 120(10), 5534-5538.

- 1128 Hosokawa, H., Mochida, T., 2015. Colorimetric Humidity and Solvent Recognition Based on a Cation-Exchange Clay Mineral 1129 Incorporating Nickel (II)-Chelate Complexes. Langmuir, 31(47), 13048-13053.
- 1130
- Intachai, S., Khaorapapong, N., Ogawa, M., 2017. Hydrothermal synthesis of zinc selenide in smectites. Applied Clay Science, 1131 135, 45-51,
- 1132 1133 Ishida, Y., Kulasekharan, R., Shimada, T., Takagi, S., Ramamurthy, V., 2013. Efficient Singlet-Singlet Energy Transfer in a Novel Host-Guest Assembly Composed of an Organic Cavitand, Aromatic Molecules, and a Clay Nanosheet. Langmuir, 29(6), 1134 1748-1753.
- 1135 Jaber, M., Komarneni, S., Zhou, C.H., 2013. Chapter 7.2, Synthesis of clay minerals. In: Bergaya, F., Lagaly, G. (Eds.), 1136 Developments in Clay Science, Vol. 5, Handbook of Clay Science(second Edition), Part A. Amsterdam, Elsevier
- 1137 Jairam, S., Tong, Z., Wang, L., Welt, B., 2013. Encapsulation of a biobased lignin-saponite nanohybrid into polystyrene co-butyl 1138 acrylate (PSBA) latex via miniemulsion polymerization. ACS Sustainable Chemistry & Engineering, 1(12), 1630-1637.
- 1139 Ju, J., Chang, J. H., 2016. Comparison of the properties of colorless polyimide nanocomposites containing saponite or 1140 organically modified hectorite. Journal of Thermoplastic Composite Materials, 29(4), 558-576.
- 1141 Kameshima, Y., Kageyama, K., Mizunuma, Y., Komatsu, T., Isobe, T., Nakajima, A., Okada, K., 2013. Preparation and solid acidity of zirconia-cluster/saponite composites. Journal- Ceramic Society Japan, 121(1409), 49-53. 1142
- 1143 1144 Kannan, V., Sreekumar, K., Gil, A., Vicente, M. A., 2011. Acetalation of pentaerithritol catalyzed by an Al-pillared saponite. Catalysis letters, 141(8), 1118-1122.
- 1145 Kaviratna, H., Pinnavaia, T. J. (1994). Acid hydrolysis of octahedral Mg²⁺ sites in 2: 1 layered silicates: An assessment of edge 1146 attack and gallery access mechanisms. Clays and Clay Minerals, 42(6), 717-723.
- 1147 Khumchoo, N., Khaorapapong, N., Ogawa, M., 2015. Formation of zinc oxide particles in cetyltrimethylammonium-smectites. 1148 Applied Clay Science, 105-106, 236-242.
- 1149 Khumchoo, N., Khaorapapong, N., Ontam, A., Intachai, S., Ogawa, M., 2016. Efficient Photodegradation of Organics in Acidic 1150 Solution by ZnO-Smectite Hybrids. European Journal of Inorganic Chemistry, 2016(19), 3157-3162.
- Kim, Y., Kirkpatrick, R. J., 1997. ²³Na and ¹³³Cs NMR study of cation adsorption on mineral surfaces: local environments, 1151 1152 dynamics, and effects of mixed cations. Geochimica Et Cosmochimica Acta, 61(24), 5199-5208.
- 1153 Kitajima, S., Bertasi, F., Vezzù, K., Negro, E., Tominaga, Y., Di Noto, V., 2013. Dielectric relaxations and conduction 1154 1155 mechanisms in polyether-clay composite polymer electrolytes under high carbon dioxide pressure. Physical Chemistry Chemical Physics, 15(39), 16626-16633.
- 1156 Kloprogge, J. T., S. Komarneni and J. E. Amonette, 1999. Synthesis of Smectite Clay Minerals: A Critical Review, Clays Clay 1157 Miner. 47:529-554.
- 1158 Komadel, P., Madejová, J., 2013. Acid activation of clay minerals. In: Bergaya, F., Lagaly, G. (Eds.), Developments in Clay 1159 Science, Vol. 5, Handbook of Clay Science, Second Edition. Amsterdam, Elsevier, pp. 385-409.
- 1160 Konno, S., Fujimura, T., Otani, Y., Shimada, T., Inoue, H., Takagi, S., 2014. Microstructures of the porphyrin/viologen monolayer 1161 on the clay surface: segregation or integration.. Journal of Physical Chemistry C, 118(35), 20504-20510.
- 1162 Kotal, M., Bhowmick, A. K., 2015. Polymer nanocomposites from modified clays: Recent advances and challenges. Progress in 1163 Polymer Science, 51, 127-187.
- 1164 Kurokawa, H., Hayasaka, M., Yamamoto, K., Sakuragi, T., Ohshima, M. A., Miura, H., 2014. Self-assembled heterogeneous late 1165 transition-metal catalysts for ethylene polymerization; New approach to simple preparation of iron and nickel complexes 1166 immobilized in clay mineral interlayer. Catalysis Communications, 47, 13-17.
- Lagaly, G., 1981. Characterization of clays by organic compounds. Clay Minerals. 16, 1-21. 1167
- 1168 Lagaly, G., 1986. Interaction of alkyamines with differents types of layered compounds. Solid State Ionics 22, 43-51.
- 1169 Lagaly, G., Weiss, A., 1969. Determination of the layer charge in mica-type layer silicates. In Proceedings of the International 1170 Clay Conference, Tokyo (Vol. 1, pp. 61-80). Israel University Press, Jerusalem.
- 1171 Lainé, M., Balan, E., Allard, T., Paineau, E., Jeunesse, P., Mostafavi, M., Robert, J.-L., Le Caër, S., 2017. Reaction mechanisms
- 1172 in swelling clays under ionizing radiation: influence of the water amount and of the nature of the clay mineral. RSC Advances, 1173 7(1), 526-534.
- 1174 Lambert, J. F., Chevalier, S., Franck, R., Suquet, H., Barthomeuf, D., 1994. Al-pillared saponites. part 2.-NMR studies. Journal of 1175 the Chemical Society Faraday Transactions, 90(4), 675-682.
- 1176 Lambert and Bergaya, 2013. Smectite Polymer Nanocomposites (Chapter 13.1). In: Bergaya and Lagaly, Eds. Handbook of Clay 1177 Science Vol. 5A, Developments of Clay Science, Elsevier
- 1178 Makarchuk, O. V., Dontsova, T. A., Astrelin, I. M., 2016. Magnetic nanocomposites as efficient sorption materials for removing 1179 dves from aqueous solutions. Nanoscale Research Letters, 11(1), 161-168
- 1180 Malla, P. B. and S. Komarneni, "Properties and Characterization of Al₂O₃ and SiO₂-TiO₂ Pillared Saponite as Affected by 1181 Pillaring", Clays Clay Miner. 41: 472-483 (1993).
- 1182 Marcal, L., de Faria, E. H., Nassar, E. J., Trujillano, R., Martin, N., Vicente, M. A., Rives, V., Gil, A., Korili, S.A., Ciuffi, K. J., 1183 2015. Organically modified saponites: SAXS study of swelling and application in caffeine removal. ACS applied materials &
- 1184 interfaces, 7(20), 10853-10862. 1185 Matejdes, M., Himeno, D., Suzuki, Y., Kawamata, J., 2017. Controlled formation of pseudoisocyanine J-aggregates in the
- 1186 interlayer space of synthetic saponite. Applied Clay Science, 140, 119-123.
- 1187 Mcbride, M. B., 1994. Environmental chemistry of soils. Environmental chemistry of soils. Oxford University Press, 1-146.
- 1188 McLaurin, E.J., Bradshaw, L.R., Gamelin, D.R., 2013. Dual-emitting nanoscale temperature sensors. Chem. Mater. 25, 1189 1283-1292.
- 1190 1191 Migal', E. A., Mishchenko, M. D., Ozheredov, I. A., Postnova, I. V., Sapozhnikov, D. A., Shkurinov, A. P., Shchipunov, Y. A., 2016. A terahertz spectroscopic study of chitosan-based bionanocomposites containing clay nanoparticles. Colloid Journal, 78(2), 1192 189-195
- 1193 Miyagawa, M., Shibusawa, A., Maeda, K., Tashiro, A., Sugai, T., Tanaka, H., 2017. Diameter-controlled Cu nanoparticles on 1194 saponite and preparation of film by using spontaneous phase separation. RSC Advances, 7(66), 41896-41902.
- 1195 Momma, K., Izumi, F., 2011. VESTA 3 for three-dimensional visualization of crystal, volumetric and morphology data. Journal 1196 of Applied Crystallography, 44(6), 1272-1276.
- 1197 Nakamura, T., Ogawa, M., 2013. Adsorption of cationic dyes within spherical particles of poly (N-isopropylacrylamide) hydrogel 1198 containing smectite. Applied Clay Science, 83, 469-473.
- 1199 Nanan, S., Khumchoo, N., Intachai, S., Khaorapapong, N., 2015. A hybrid of hexakis (hexyloxy) triphenylene and synthetic

- 1200 saponite. Applied Clay Science, 114, 407-411.
- 1201 1202 Nityashree, N., Gautam, U. K., Rajamathi, M., 2014. Synthesis and thermal decomposition of metal hydroxide intercalated saponite. Applied Clay Science, 87, 163-169.
- 1203 Ogawa, M., Kuroda, K., 1997. Preparation of inorganic-organic nanocomposites through intercalation of organoammonium ions into layered silicates. Bulletin of the Chemical Society of Japan, 70(11), 2593-2618.
- 1204 1205 Ogawa, M., Takee, R., Okabe, Y., Seki, Y., 2017. Bio-geo hybrid pigment; clay-anthocyanin complex which changes color 1206 1207 depending on the atmosphere. Dyes & Pigments, 139, 561-565.
- Okada, T., Oguchi, J., Yamamoto, K. I., Shiono, T., Fujita, M., Iiyama, T., 2015. Organoclays in water cause expansion that 1208 facilitates caffeine adsorption. Langmuir, 31(1), 180-187.
- 1209 Olaya, A., Moreno, S., Molina, R., 2009. Synthesis of pillared clays with aluminum by means of concentrated suspensions and 1210 1211 microwave radiation. Catalysis Communications, 10(5), 697-701.
- Olivero, F., Carniato, F., Bisio, C., Marchese, L., 2014. Promotion of Förster resonance energy transfer in a saponite clay 1212 containing luminescent polyhedral oligomeric silsesquioxane and rhodamine dye. Chemistry-An Asian Journal, 9(1), 158-165.
- 1213 1214 Ostinelli, L., Recchia, S., Bisio, C., Carniato, F., Guidotti, M., Marchese, L., Psaro, R., 2012. Acid/vanadium - containing saponite for the conversion of propene into coke: potential flame - retardant filler for nanocomposite materials. Chemistry-An 1215 Asian Journal, 7(10), 2394-2402.
- 1216 1217 Padilla-Ortega, E., Darder, M., Aranda, P., Figueredo Gouveia, R., Leyva-Ramos, R., RuizHitzky, E., 2016. Ultrasound assisted preparation of chitosan-vermiculite bionanocomposite foams for cadmium uptake. Applied Clay Science. 130, 40-49.
- 1218 Pandey, J. K., Reddy, K. R., Kumar, A. P., Singh, R. P., 2005. An overview on the degradability of polymer nanocomposites. 1219 1220 Polymer degradation and stability, 88(2), 234-250.
- Paul, D. R., Robeson, L. M., 2008. Polymer nanotechnology: nanocomposites. Polymer, 49(15), 3187-3204.
- 1221 Petra, L., Billik, P., Melichová, Z., Komadel, P., 2017. Mechanochemically activated saponite as materials for Cu²⁺, and Ni²⁺, 1222 1223 removal from aqueous solutions. Applied Clay Science, 143, 22-28.
- Pigatto, G., Lodi, A., Finocchio, E., Palma, M. S., Converti, A., 2013. Chitin as biosorbent for phenol removal from aqueous 1223 1224 1225 1226 1227 1228 1229 solution: Equilibrium, kinetic and thermodynamic studies. Chemical Engineering and Processing: Process Intensification, 70, 131-139
- Pimchan, P., Khaorapapong, N., Ogawa, M., 2014. The effect of cetyltrimethylammonium ion and type of smectites on the luminescence efficiency of bis(8-hydroxyquinoline)zinc(ii) complex. Applied Clay Science, 101, 223-228.
- Rafatullah, M., Sulaiman, O., Hashim, R., Ahmad, A., 2010. Adsorption of methylene blue on low-cost adsorbents: a review. Journal of hazardous materials, 177(1), 70-80.
- 1230 1231 Rodriguez, M. V., González, J. D. D. L., Munoz, M. B., 1995. Preparation of microporous solids by acid treatment of a saponite. Microporous Materials, 4(4), 251-264.
- 1232 1233 1234 1235 Roelofs, J. C. A. A., Berben, P. H., 2006. Preparation and performance of synthetic organoclays. Applied clay science, 33(1), 13-20.
- Sas, S., Danko, M., Bizovská, V., Lang, K., Bujdák, J., 2017. Highly luminescent hybrid materials based on smectites with polyethylene glycol modified with rhodamine fluorophore. Applied Clay Science, 138, 25-33.
- 1236 Sato, H., Ochi, M., Kato, M., Tamura, K., Yamagishi, A., 2014a. Energy transfer in hybrid langmuir-blodgett films of iridium 1237 complexes and synthetic saponite: dependence of transfer efficiency on interlayer distance. New Journal of Chemistry, 38(12), 5715-5720.
- 1238 1239 1240 Sato, H., Tamura, K., Taniguchi, M., Yamagishi, A., 2014b. Efficient energy transfer of cationic iridium (III) complexes on the surface of a colloidal clay. Applied Clay Science, 97, 84-90.
- 1241 1242 Sato, K., Fujimoto, K., Dai, W., Hunger, M., 2016. Quantitative Elucidation of Cs Adsorption Sites in Clays: Toward Sophisticated Decontamination of Radioactive Cs. The Journal of Physical Chemistry C, 120(2), 1270-1274.
- 1243 Sato, K., Hunger, M., 2017. Molecular studies of Cs adsorption sites in inorganic layered materials: the influence of solution 1244 1245 concentration. Physical Chemistry Chemical Physics, 19(28), 18481-18486.
- Seki, Y., Ide, Y., Okada, T., Ogawa, M. 2015. Concentration of 2-phenylphenol by organoclays from aqueous sucrose solution. 1246 1247 Applied Clay Science, 109, 64-67.
- Seki, Y., Ogawa, M., 2010. The removal of 2-phenylphenol from aqueous solution by adsorption onto organoclays. Bulletin of 1248 the Chemical Society of Japan, 83(6), 712-715.
- 1249 Shchipunov, Y. A., Sarin, S. A., Silant'ev, V. E., Postnova, I. V., 2012a. Self-organization in the chitosan-clay nanoparticles 1250 1251 system regulated through polysaccharide macromolecule charging. 2. Films. Colloid Journal, 74(5), 636-644.
- Shchipunov, Y. A., Silant'ev, V. E., Postnova, I. V., 2012b. Self-organization in the chitosan-clay nanoparticles system regulated through polysaccharide macromolecule charging. 1. Hydrogels. Colloid Journal, 74(5), 627-635.
- 1252 1253 1254 Shchipunov, Y., Ivanova, N., Silant'ev, V., 2009. Bionanocomposites formed by in situ charged chitosan with clay. Green Chemistry, 11(11), 1758-1761.
- 1255 1255 1256 1257 1258 1259 Shchipunov, Y., Postnova, I., 2010. Water-soluble polyelectrolyte complexes of oppositely charged polysaccharides. Composite Interfaces, 16(4-6), 251-279.
- Shen, Z., Simon, G. P., Cheng, Y. B., 2002. Comparison of solution intercalation and melt intercalation of polymer-clay nanocomposites. Polymer, 43(15), 4251-4260.
- Shin, J., Kim, J. C., Chang, J. H., 2013. Flexible clay hybrid films with various poly (vinyl alcohol) contents: Thermal properties, 1260 morphology, optical transparency, and gas permeability. Macromolecular Research, 21(12), 1349-1354.
- 1261 Srinivasan, R., 2011. Advances in application of natural clay and its composites in removal of biological, organic, and inorganic 1262 contaminants from drinking water. Advances in Materials Science & Engineering, 2011(1), 1-5.
- 1263 Suquet, H., Iiyama, J.T., Kodama, H., Pezerat, H., 1977. Synthesis and swelling properties of saponites with increasing layer 1264 charge. Clays and Clay Mienrals, 25, 231-242.
- 1265 Suquet, H., Malard, C., Copin, E., Pezerat, H., 1981. Variation du paramètre b et de la distance basale dout dans une série de 1266 saponites à charge croissante: I Etats hydratés. Clay Minerals, 16(1), 53-67.
- 1267 Suzuki, Y., Sugihara, H., Satomi, K., Tominaga, M., Mochida, S., Kawamata, J., 2014. Two-photon absorption properties of an 1268 acetylene derivative confined in the interlayer space of a smectite. Applied Clay Science, 96(26), 116-119.
- 1269 1270 Takagi, S., Shimada, T., Ishida, Y., Fujimura, T., Masui, D., Tachibana, H., Eguchi, M., Inoue, H., 2013. Size-matching effect on inorganic nanosheets: control of distance, alignment, and orientation of molecular adsorption as a bottom-up methodology for 1271 nanomaterials. Langmuir, 29(7), 2108-2119.
- 1272 1273 Tamura, K., Yamagishi, A., Kitazawa, T., Sato, H., 2015. Harvesting light energy by iridium(iii) complexes on a clay surface. Physical Chemistry Chemical Physics, 17(28), 18288.
- 1274 Tangaraj, V., Janot, J. M., Jaber, M., Bechelany, M., Balme, S., 2017. Adsorption and photophysical properties of fluorescent

- 1275 dyes over montmorillonite and saponite modified by surfactant. Chemosphere, 184, 1355-1361.
- 1276 1277 Tao, Q., Fang, Y., Li, T., Zhang, D., Chen, M.Y., Ji, S.C., He, H.P., Komarneni, S., Zhang, H.B., Dong, Y., Noh, Y. D., 2016. Silylation of saponite with 3-aminopropyltriethoxysilane. Applied Clay Science, 132, 133-139.
- 1278 Theng, B. K. G., 1974. The Chemistry of Clay-Organic Reactions. The chemistry of clay-organic reactions. Hilger.
- 1279 1280 Tokieda, D., Tsukamoto, T., Ishida, Y., Ichihara, H., Shimada, T., Takagi, S., 2017. Unique fluorescence behavior of dyes on the clay minerals surface: surface fixation induced emission (s-fie). Journal of Photochemistry & Photobiology A Chemistry, 339, 1281 1282 67-79.
- Tominaga, M., Nishioka, Y., Tani, S., Suzuki, Y., Kawamata, J., 2017. Tunable high-pressure field operating on a cationic 1283 biphenyl derivative intercalated in clay minerals. Scientific Reports, 7(1), 7651.
- 1284 Tominaga, M., Oniki, Y., Mochida, S., Kasatani, K., Tani, S., Suzuki, Y., Kawamata, J., 2016. Clay-Organic Hybrid Films 1285 1286 Exhibiting Reversible Fluorescent Color Switching Induced by Swelling and Drying of a Clay Mineral. The Journal of Physical Chemistry C, 120(41), 23813-23822.
- 1287 Tong, Z., Deng, Y., 2013. The formation of asymmetric polystyrene/saponite composite nanoparticles via miniemulsion 1288 1289 polymerization. Journal of Applied Polymer Science, 127(5), 3916-3922.
- Trujillano, R., Rico, E., Vicente, M. A., Herrero, M., Rives, V., 2010. Microwave radiation and mechanical grinding as new ways 1290 1291 for preparation of saponite-like materials. Applied Clay Science, 48(1), 32-38.
- Trujillano, R., Rico, E., Vicente, M. A., Rives, V., Ciuffi, K. J., Cestari, A., Gil, A., Korili, S. A., 2011. Rapid microwave-assisted 1292 synthesis of saponites and their use as oxidation catalysts. Applied Clay Science, 53(2), 326-330.
- 1293 1294 Trujillano, R., Vicente, M. A., Rives, V., Korili, S. A., Gil, A., Ciuffi, K. J., Nassar, E. J., 2009. Preparation, alumina-pillaring and oxidation catalytic performances of synthetic Ni-saponite. Microporous and Mesoporous Materials, 117(1), 309-316.
- 1295 Tsukamoto, T., Ramasamy, E., Shimada, T., Takagi, S., Ramamurthy, V., 2016a. Supramolecular surface photochemistry: cascade 1296 energy transfer between encapsulated dyes aligned on a clay nanosheet surface. Langmuir, 32(12), 2920-2927.
- 1297 Tsukamoto, T., Shimada, T., Takagi, S., 2016b. Photophysical Properties and Adsorption Behaviors of Novel Tri-Cationic Boron 1298 1299 (III) Subporphyrin on Anionic Clay Surface. ACS applied materials & interfaces, 8(11), 7522-7528.
- Ugochukwu, U. C., Fialips, C. I., 2017a. Crude oil polycyclic aromatic hydrocarbons removal via clay-microbe-oil interactions: 1300 Effect of acid activated clay minerals. Chemosphere, 178, 65-72.
- 1301 1302 Ugochukwu, U. C., Fialips, C. I., 2017b. Removal of crude oil polycyclic aromatic hydrocarbons via organoclay-microbe-oil interactions. Chemosphere, 174, 28-38.
- 1303 Ugochukwu, U. C., Jones, M. D., Head, I. M., Manning, D. A., Fialips, C. I., 2014a. Effect of acid activated clay minerals on 1304 biodegradation of crude oil hydrocarbons. International Biodeterioration & Biodegradation, 88, 185-191.
- 1305 Ugochukwu, U. C., Manning, D. A.C., Fialips, C. I., 2014b. Microbial degradation of crude oil hydrocarbons on organoclay 1306 minerals. Journal of Environmental Management, 144, 197-202.
- 1307 1308 Utracki, L. A., Sepehr, M., Boccaleri, E., 2007. Synthetic, layered nano-particles for polymeric nanocomposites (PNC's). Polymers for Advanced Technologies, 18(1), 1-37.
- 1309 Vdović, N., Jurina, I., Škapin, S.D., Sondi, I., 2010. The surface properties of clay minerals modified by intensive dry milling -1310 revisited. Applied Clay Science. 48, 575-580.
- 1311 Velasco, J., Pérez-Mayoral, E., Mata, G., Rojas-Cervantes, M. L., Vicente-Rodríguez, M. A., 2011. Cesium-saponites as excellent 1312 1313 environmental-friendly catalysts for the synthesis of N-alkyl pyrazoles. Applied Clay Science, 54(2), 125-131.
- Venkatachalam, S., Hayashi, H., Ebina, T., Nakamura, T., Nanjo, H., 2013. Optoelectronic properties of nanostructured ZnO thin 1314 films prepared on glass and transparent flexible clay substrates by hydrothermal method. Japanese Journal of Applied Physics, 1315 1316 52(5), 492-494.

Verma, S., Ghosh, H.N., 2012. Exciton energy and charge transfer in porphyrin aggregate/semiconductor (TiO₂) composites. J. 1317 Phys. Chem. Lett. 3, 1877-1884.

- 1318 1319 Vicente, I., Salagre, P., Cesteros, Y., 2011. Ni nanoparticles supported on microwave-synthesised saponite for the hydrogenation of styrene oxide. Applied Clay Science, 53(2), 212-219.
- Vicente, M. A., Belver, C., Sychev, M., Prihod'ko, R., Gil, A., 2008. Relationship between the surface properties and the catalytic 1320 performance of Al-, Ga-, and AlGa-pillared saponites. Industrial & Engineering Chemistry Research, 48(1), 406-414. 1321
- 1322 Vicente, M. A., Gil, A., Bergaya, F., 2013. Pillared Clays and Clays Minerals. In: Bergaya, F., Lagaly, G. (Eds.), Developments in 1323 Clay Science, Vol. 5, Handbook of Clay Science, Second Edition. Amsterdam, Elsevier, pp. 523-557.
- 1324 Vogels, R. J. M. J., Kloprogge, J. T., Geus, J. W., 2005. Synthesis and characterization of saponite clays. American Mineralogist, 90(5-6), 931-944.
- 1325 1326 1327 Wang, W., Zhen, W., Bian, S., Xi, X., 2015. Structure and properties of quaternary fulvic acid-intercalated saponite/poly (lactic acid) nanocomposites. Applied Clay Science, 109, 136-142.
- 1328 Wu, C. N., Yang, Q., Takeuchi, M., Saito, T., Isogai, A., 2014. Highly tough and transparent layered composites of nanocellulose 1329 and synthetic silicate. Nanoscale, 6(1), 392-399.
- 1330 Wu, L., Lv, G., Liu, M., Li, Z., Liao, L., Pan, C., 2015. Adjusting the Layer Charges of Host Phyllosilicates To Prevent 1331 Luminescence Quenching of Fluorescence Dyes. The Journal of Physical Chemistry C, 119(39), 22625-22631.
- 1332 Wu, L. M., Liao, L. B., Lv, G. C., 2015. Influence of interlayer cations on organic intercalation of montmorillonite. Journal
- 1333 Colloid & Interface Science, 454, 1-7.
- 1334 Wysokowski, M., Klapiszewski, Ł., Moszyński, D., Bartczak, P., Szatkowski, T., Majchrzak, I., Siwinska-Stefanska, K., 1335 Bazhenov, V.V., Jesionowski, T., 2014. Modification of chitin with kraft lignin and development of new biosorbents for removal 1336 of cadmium (II) and nickel (II) ions. Marine drugs, 12(4), 2245-2268.
- 1337 Wysokowski, M., Petrenko, I., Stelling, A. L., Stawski, D., Jesionowski, T., Ehrlich, H., 2015. Poriferan chitin as a versatile 1338 1339 template for extreme biomimetics. Polymers, 7(2), 235-265.
- Xi, X., Zhen, W., Bian, S., 2015. Preparation and properties of polylactic acid/N-(2-hydroxyl) propyl-3-trimethyl ammonium 1340 chitosan chloride-intercalated saponite nanocomposites. Iranian Polymer Journal, 24(3), 243-252.
- 1341 Xu, P., Zeng, G. M., Huang, D. L., Feng, C. L., Hu, S., Zhao, M. H., Lai, C., Wei, Z., Huang, C., Xie, G.X., Liu, Z. F., 2012. Use 1342 of iron oxide nanomaterials in wastewater treatment: a review. Science of the Total Environment, 424, 1-10.
- 1343 Yang, F., Sun, S., Chena, X., Chang, Y., Zha, F., Lei, Z., 2016. Mg-Al layered double hydroxides modified clay adsorbents for 1344 efficient removal of Pb²⁺, Cu²⁺ and Ni²⁺ from water. Applied Clay Science, 123, 134–140.
- 1345 Yang, Q., Saito, T., Isogai, A., 2013. Transparent, flexible, and high - strength regenerated cellulose/saponite nanocomposite 1346 films with high gas barrier properties. Journal of Applied Polymer Science, 130(5), 3168-3174.
- 1347 Yu, W.H., Ren, Q.Q., Tong, D.S., Zhou, C.H., Wang, H., 2014. Clean production of CTAB montmorillonite: formation

- 1348 mechanism and swelling behavior in xylene. Appl. Clay Sci. 97–98, 222–234.
- 1349 Zhang, C. Q., He, H. P., Tao, Q., Ji, S. C., Li, S.Y., Ma, L. Y., Su, X. L., Zhu, J. X., 2017. Metal occupancy and its influence on thermal stability of synthetic saponites. Applied Clay Science, 135, 282-288.
- 1351 Zhang, D., Zhou, C.H., Lin, C.X., Tong, D.S., Yu, W.H., 2010. Synthesis of clay minerals. Appl. Clay Sci. 50, 1–11.
- 1352Zhen, W., Li, J., Xu, Y., 2014. In situ intercalation green polymerization, characterization, and kinetics of poly(lactic acid)/zinc1353oxide pillared saponite nanocomposites. Polymer Composites, 35(6), 1023-1030.
- Zhen, W., Lu, C., Liang, M., 2012. Structure and properties of thermoplastic saponite/poly (vinyl alcohol) nanocomposites.
 Applied Clay Science, 57, 64-70.
- 1356 Zhen, W., Sun, J., 2014. Properties, structure and crystallization of poly lactic acid/zinc oxide pillared organic saponite
 1357 nanocomposites. Polymer Korea, 38(3), 299-306.
- 1358 Zhen, W., Wang, W., 2016. Structure, properties and rheological behavior of thermoplastic poly (lactic acid)/quaternary fulvic
 1359 acid-intercalated saponite nanocomposites. Polymer Bulletin, 73(4), 1015-1035.
- **1360** Zhen, W., Zheng, Y., 2016. Synthesis, characterization, and thermal stability of poly (lactic acid)/zinc oxide pillared organic
- 1361 saponite nanocomposites via ring opening polymerization of D,L-lactide. Polymers for Advanced Technologies, 27(5),1362 606-614.
- 1363 Zhou, C.H., 2011. An overview on strategies towards clay-based designer catalysts for green and sustainable catalysis. Appl.
 1364 Clay Sci. 53, 87–96.
- 1365 Zhou, C.H., Keeling, J.L., 2013. Fundamental and applied research on clay minerals: from climate and environment to1366 nanotechnology. Appl. Clay Sci. 74, 3–9.
- 1367 Zhou, C.H., Shen, Z.F., Liu, L.H., Liu, S.M., 2011a. Preparation and functionality of claycontaining films. J. Mater. Chem. 21, 1368 15132–15153.
- 1369 Zhou, C.H., Xia, X., Lin, C.X., Tong, D.S., Beltramini, J., Zhu, R., Chena, Q., Zhou, Q., Xi, Y., Zhu, J., He, H., 2016. Adsorbents
 1370 based on montmorillonite for contaminant removal from water: a review. Applied Clay Science, 123, 239–258.
- ased on montholine for containing tentova noin water, a review. Applied Clay Science, 123, 237–236.
- Zou, H., Wu, S., Shen, J., 2008. Polymer/silica nanocomposites: preparation, characterization, properties, and applications.
 Chemical Review, 108(9), 3893-3957.
- 1373