Trapping at the Solid–Gas Interface: Selective Adsorption of Naphthalene by Montmorillonite Intercalated with a Fe(III)–Phenanthroline Complex

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ABSTRACT: In this study, stable hybrid materials (Mt–Fe(III)Phen), made by the μ-oxo Fe(III)–phenanthroline complex [(OH₂)₃(Phen)-FeOFe(Phen)(OH₂)]⁴⁺ (Fe(III)Phen) intercalated in different amounts into montmorillonite (Mt), were used as a trap for immobilizing gaseous benzene and naphthalene and their mono chloro-derivatives at 25 and 50 °C. The entrapping process was studied through elemental analysis, magic angle spinning NMR spectroscopy, thermal analysis, and evolved gas mass spectrometry. Naphthalene and 1-chloronaphthalene were found to be immobilized in large amount at both temperatures. Molecular modeling allowed designing of the structure of the interlayer in the presence of the immobilized aromatic molecules. Adsorption is affected by the amount of the Fe complex hosted in the interlayer of the entrapping hybrid materials. On the contrary, under the same conditions, benzene and chlorobenzene were not adsorbed. Thermal desorption of naphthalenes was obtained under mild conditions, and immobilization was found to be reversible at least for 20 adsorption/desorption cycles.

1. INTRODUCTION

Monocyclic and polycyclic aromatic molecules are produced by several industrial and human activities, including the combustion processes of vehicles. Many efforts have been devoted to the production of materials able to remove the pollutants through immobilization processes or catalytic degradation. Several are presently available, but most lack selectivity. In addition, they are successfully operated mainly in solution. Only a few materials, such as zeolite HZSM-5† and diatomite-based ceramic monoliths coated with silicalite-1,‡ trap effectively aromatic molecules in the gas phase, and a number of challenges related to selectivity are to be tackled, which are complicated by the low partial pressure of these compounds in the flue gas. Natural and engineered zeolites are extensively employed as catalysts in a variety of reactions involving benzene and naphthalene. For example, zeolites are commercial catalysts currently utilized in refineries to remove benzene from gasoline by alkylation with olefins (BenzOUT process). Other reactions of benzene are catalyzed by zeolites, such as self-alkylation and hydrogenation, oxidation to obtain phenol, and hydrogenation to produce cyclohexane. Immobilization followed by a catalytic reaction implies that the target aromatic species is transformed into different final molecules and the overall process is irreversible, thus desorption does not restore the starting species. Simple adsorption, instead, allows the aromatic molecule to retain its structure and the recovery of the chemical compound. For example, activated carbon is largely employed as an industrial adsorbent for gases because of its effectiveness and low cost. However, this material shows many drawbacks as the limited thermal stability and poor versatility. More importantly, the adsorbed organic gas molecules are confined in the slitlike micropores, and this could result in a difficult desorption and regeneration. Moreover, in some cases, immobilization partly occurs via chemical adsorption, which changes the functional groups and the structure of activated carbon. For example, Guo et al. found that after chlorobenzene adsorption, the lactone and quinone groups on activated carbon decreased, and the structure of activated carbon slightly changed. In addition, activated carbon, also after specific treatments, is poorly selective.

Here, we have investigated the effectiveness of a hybrid material made by the intercalation of the μ-oxo di-fac-(triaqua-(1,10-phenanthroline-k²N,N’)-iron(III))⁴⁺, [(OH₂)₃(Phen)-FeOFe(Phen)(OH₂)]⁴⁺ complex (Fe(III)Phen) onto montmorillonite (Mt–Fe(III)Phen hereafter) to immobilize some selected paradigmatic aromatic molecules from the gas phase: benzene, chlorobenzene, naphthalene, and 1-chloronaphthalene...
lene. In previous papers, this hybrid material (Mt–Fe(III)-Phen) was characterized, unraveling that the Fe(III)Phen complex has a structuring effect on the interlayer because of the presence and orientation of extended aromatic regions of the ligand;\textsuperscript{10,11} then this hybrid material was proven effective in trapping volatile sulfur compounds, aliphatic thiols and H\textsubscript{2}S.\textsuperscript{12,13} In this case, the entrapping mechanism for thiols and H\textsubscript{2}S is controlled by the redox and coordination properties of the metal center. In the present work, on the other hand, we investigated the ability of the interlayer structured by the Fe(III)Phen complex, which bears two coordinated planar phenanthroline ligands almost parallel to each other,\textsuperscript{10} to capture specific aromatic compounds, thanks to the polycyclic nature of the Phen ligand.

Here, benzene, chlorobenzene, naphthalene, and 1-chloronaphthalene trapping by Mt–Fe(III)Phen was studied to understand if a relationship exists between the entrapping efficiency and the size of the aromatic moiety and to test the effect of a halogen substitution. Removal of such pollutants from the environment would be a result of great social and industrial significance.\textsuperscript{14}

2. EXPERIMENTAL SECTION

2.1. Materials. Mt STx-1a from the Clay Minerals Society (The Clay Minerals Society, Source Clays Repository, University of Missouri, Columbia, MO) was characterized elsewhere.\textsuperscript{15,16} It was used “as received”, avoiding any purification or exchange process intended to obtain a homogeneous counterion, as efficiency of this clay mineral in the adsorption of the Fe(III)Phen complex was already proved when used “as it is”.\textsuperscript{10,12,13}

All the chemicals used for preparation of the adsorbent material were of analytical grade (purity > 99%) and purchased from Carlo Erba (Fe\textsubscript{2}(SO\textsubscript{4})\textsubscript{3} 8H\textsubscript{2}O and NaOH pellets) or from Sigma-Aldrich (1,10-phenanthroline C\textsubscript{12}H\textsubscript{8}N\textsubscript{2}, Phen). The aromatic molecules tested for adsorption in the gas phase were as follows: benzene 99.9% (AnalaR, analytical reagent, benz hereafter), chlorobenzene ≥ 99.5% (Fluka, Clbenz hereafter), naphthalene scintillation grade (BDH Laboratory reagents, naph hereafter), and 1-chloronaphthalene 90% (Aldrich Chemie, Clnaph hereafter).

2.2. Methods. 2.2.1. Preparation of Mt Intercalated with Different Amounts of the Fe(III)Phen Complex: Mt–Fe(III)-Phen Materials. The solutions of [(OH\textsubscript{2})\textsubscript{2}(Phen)FeOFe(Phen)(OH\textsubscript{2})\textsubscript{2}]\textsuperscript{4+} (Fe(III)Phen) were prepared by dissolving Fe\textsubscript{2}(SO\textsubscript{4})\textsubscript{3} 8H\textsubscript{2}O in aqueous Phen solutions (typically 2 ± 12 mM) in order to have a 1:1 Fe(III)/Phen molar ratio.\textsuperscript{10} The Fe(III)Phen solutions (20 mL) in the 1–6 mM range of concentration were mixed with 100 mg of Mt to form the corresponding suspensions. The suspensions were shaken at 250 rpm in an orbital incubator (Stuart Scientific Orbital Incubator SI50) at 20 °C for 30 min using a Haake k20 thermocryostat and then kept in the thermostated incubator. After the solid–liquid separation, the supernatant was removed with a pipette. The solids were washed several times with distilled water and dried at 30 °C. They have different levels of interlayer saturation by the filling of complex molecules, and all constitute the Mt–Fe(III)Phen adsorbed materials. Quantification of the amount of the intercalated complex was obtained by UV–vis measurements (JASCO V-570 Spectrophotometer) performed on the different supernatants, previously centrifuged at 14 000 rpm (Thermo mod. Esprreso) for 1 min to remove residual clay traces. \(q_{\text{complex}}\) is defined as the content in the iron complex of the Mt–Fe(III)Phen material, expressed as moles of Fe(III)Phen per 100 g of Mt–Fe(III)Phen; \(q_{\text{complex}}\) ranges from 0.005 to 0.038 (saturation limit of Mt) moles of Fe(III)Phen/100 g Mt–Fe(III)-Phen..\textsuperscript{10,12,13}

2.2.2. Immobilization of Aromatic Compounds in the Gas Phase on Mt–Fe(III)Phen at Different \(q_{\text{complex}}\) Values. Mt–Fe(III)Phen prepared with \(q_{\text{complex}}\) ranging from 0.005 to 0.038 moles of Fe(III)Phen/100 g Mt–Fe(III)Phen was exposed to the vapors of aromatic compounds benz, Clbenz, naph, and Clnaph at 25 and 50 °C for 1 month. The Mt–Fe(III)Phen adsorbent material (100 mg) was finely dispersed in a glass Petri dish (diameter 5 cm) and placed in a sealed glass box in the dark. The aromatic compound (benz, Clbenz, naph) was placed in a beaker inside the same glass box, which was closed with a sealing cap. The glass boxes were thermostated at 25 and 50 °C for 1 month. In this way, we can confidently assume that at each temperature, the box was always saturated with the vapor of the aromatic compound at the vapor pressure (ref 17, Table S1, page S2). After 1 month, the elemental analyses were performed on samples to find the maximum amount of the adsorbed aromatic compounds. Hereafter, these materials after exposure to benz, Clbenz, naph, and Clnaph at 25 and 50 °C will be indicated as Mt–Fe(III)Phen–benz\textsubscript{25}, Mt–Fe(III)Phen–benz\textsubscript{50}, Mt–Fe(III)–Phen–Clbenz\textsubscript{25}, Mt–Fe(III)–Phen–Clbenz\textsubscript{50}, Mt–Fe(III)–Phen–naph\textsubscript{25}, Mt–Fe(III)–Phen–naph\textsubscript{50}, Mt–Fe(III)–Phen–Clnaph\textsubscript{25}, and Mt–Fe(III)–Phen–Clnaph\textsubscript{50}. In addition, for the material prepared with the amount of the complex which gives the maximum adsorption capacity, the “best” suffix was introduced: Mt–Fe(III)Phen–benz\textsubscript{best25}, Mt–Fe(III)–Phen–benz\textsubscript{best50}, Mt–Fe(III)–Phen–Clbenz\textsubscript{best25}, Mt–Fe(III)–Phen–Clbenz\textsubscript{best50}, Mt–Fe(III)–Phen–naph\textsubscript{best25}, Mt–Fe(III)–Phen–naph\textsubscript{best50},Mt–Fe(III)–Phen–Clnaph\textsubscript{best25}, and Mt–Fe(III)–Phen–Clnaph\textsubscript{best50}.

2.2.3. Kinetics of Immobilization of the Aromatic Compounds. The kinetics of immobilization was investigated on Mt–Fe(III)Phen showing the maximum immobilization capacity of naph and Clnaph at 25 and 50 °C, respectively (\(q_{\text{complex}}\) = 0.022 and 0.025 moles of Fe(III)Phen/100 g of Mt–Fe(III)Phen), which will be indicated as Mt–Fe(III)Phen–naph\textsubscript{25} and Mt–Fe(III)Phen–naph\textsubscript{50} respectively (\(q_{\text{complex}}\) = 0.022 and 0.025 moles of Fe(III)Phen/100 g of Mt–Fe(III)Phen). Kinetics of immobilization was investigated on Mt–Fe(III)Phen showing the maximum immobilization capacity of naph and Clnaph at 25 and 50 °C, respectively (\(q_{\text{complex}}\) = 0.022 and 0.025 moles of Fe(III)Phen/100 g of Mt–Fe(III)Phen), which will be indicated as Mt–Fe(III)Phen–naph\textsubscript{25} and Mt–Fe(III)Phen–naph\textsubscript{50}. Batches of Mt–Fe(III)Phen–naph\textsubscript{25} and Mt–Fe(III)Phen–naph\textsubscript{50} samples were prepared and exposed to the vapors of naph and Clnaph at 25 and 50 °C, respectively, as a function of time in the range 1 h to 30 days, using the same procedure described in the previous paragraph. The elemental analyses performed on samples at different times allowed to investigate the kinetics of immobilization of the aromatic compounds.

2.2.4. Desorption of Aromatic Compounds from Mt–Fe(III)Phen–naph\textsubscript{best25} and Mt–Fe(III)Phen–Clnaph\textsubscript{best25} at Different Temperature Values; Adsorption/Desorption Cycles. Batches of Mt–Fe(III)Phen–naph\textsubscript{best25} and Mt–Fe(III)Phen–Clnaph\textsubscript{best25} were treated at different temperature values: 160, 180, and 200 °C for \(t = 10\) min; elemental analysis was performed on the samples after every minute of treatment to measure the residual amount of aromatic compounds as a function of time. Samples were completely desorbed at \(T = 200\) °C, and then they underwent 20 adsorption/desorption cycles (adsorption at 25 °C for 400 h and desorption at 200 °C); the content in the aromatic compound was checked during the 20 cycles at different times by elemental analysis.
2.2.5. Elemental Analysis. The elemental analyses (C, N, S) of the samples before and after exposure to the vapors of aromatic compounds were performed by a Carlo Erba elemental analyzer (model 1106) and used to calculate the amount of the adsorbed aromatic compound.

2.2.6. Thermogravimetric Measurements and Analysis of the Evolved Gases with Mass Spectrometry. Thermogravimetric analyses (TGAs) on the samples before and after exposure to the vapors of the aromatic compounds were made with a Seiko SSC 5200 thermal analyzer equipped with a quadrupole mass spectrometer (ESS, GeneSysQuad 422) to characterize the gases evolved during the thermal processes (mass spectrometry evolved gas analysis, MS-EGA). Gas sampling by the spectrometer was via an inert, fused silicon capillary system, heated to prevent the condensation of gases. Gas analyses were performed in multiple ion detection (MID) mode to determine the nature of the evolved chemical species with temperature (or time). Background subtraction was used to obtain the point zero conditions before starting MID analysis. Measurements were performed on each air-dried sample under the following experimental conditions: heating rate: 20 °C/min; heating range: 25−1200 °C; data measurement: every 0.5 s; purging gas: ultrapure helium, flow rate: 100 μL/min. Mass analyses were carried out in multiple ion detection mode measuring the m/z ratios 17 and 18 for H2O, 28 and 44 for CO2, 30 for NO and NO2, 34 for H2S, 46 for SO2, and 48, 64, and 66 for combustion gases (mass spectrometry evolved gas analysis, MS-EGA). Gas sampling by the spectrometer was via an inert, fused silicon capillary system, heated to prevent the condensation of gases. Gas analyses were performed in multiple ion detection (MID) mode to determine the nature of the evolved chemical species with temperature (or time). Background subtraction was used to obtain the point zero conditions before starting MID analysis. Measurements were performed on each air-dried sample under the following experimental conditions: heating rate: 20 °C/min; heating range: 25−1200 °C; data measurement: every 0.5 s; purging gas: ultrapure helium, flow rate: 100 μL/min. Mass analyses were carried out in multiple ion detection mode measuring the m/z ratios 17 and 18 for H2O, 28 and 44 for CO2, 30 for NO and NO2, 34 for H2S, 46 for SO2, and 48, 64, and 66 for SO2 (m/z is the dimensionless ratio between the mass number and the charge of an ion); secondary electron multiplier (SEM) and FARADAY detector at 1100 V were employed with 1 s of integration time on each measured mass. To avoid differences in relative humidity, samples were equilibrated for 15 min inside the oven at 25 °C using a 100 μL/min flow of ultrapure helium.

2.2.7. NMR Measurements. NMR spectra were acquired at 300 K using an AVANCE III HD 600 Bruker spectrometer equipped with a 2.5 mm H/X CPMAS probe operating at 600.13 and 150.90 MHz for 1H and 13C, respectively. Samples were packed into 2.5 mm zirconia rotors and spun at the magic angle. 1H NMR spectra were obtained at 30 or 33 kHz magic angle spinning (MAS) rate, using DEPTH sequence in order to remove baseline distortions, at 125 kHz spectral width, 10 s relaxation delay, 2.3 μs 90° pulse, 4k data points, and 32 scans. The empty rotor 1H spectrum was subtracted to compensate for background effects.

Cross-polarization-MAS (CP-MAS) 13C NMR spectra were obtained at 16 kHz MAS rate, using the standard Bruker CP sequence with 139 kHz spectral width, 1 s relaxation delay, 2.3 μs 90° 1H pulse, radio frequency field strength of about 62 kHz for Hartmann−Hahn match, 0.2 ms contact time, 4k data points, and 28k−32k scans. All chemical shifts were referenced by adjusting the spectrometer field to the value corresponding to 38.48 ppm chemical shift for the deshielded line of the adamantane 13C NMR signal, as previously reported.19

2.2.8. Molecular Modeling. A 3 × 2 × 1 supercell of Mt with two Fe(III)Phen complexes with eight water molecules per Fe complex, viewed from (010) (a) and (100) (b) planes. The Fe, H, O, N, C, Si, Al, and Mg atoms are in purple, white, red, blue, gray, ochre, pink, and green colors, respectively.

On the other hand, taking into account the amount of water (2.96%, see after) found experimentally by TGA in Mt−Fe(III)Phen showing the maximum immobilization capacity of naphth by adjusting the spectrometer field to the value corresponding to 38.48 ppm chemical shift for the deshielded line of the adamantane 13C NMR signal, as previously reported.19

A double supercell was generated, being a 3 × 4 × 1 of Mt and two Fe(III)Phen complexes. One of these complexes was rotated in order to place both phenanthroline moieties oriented to the same side maintaining all rings parallel to the clay surface, creating a hydrophobic channel. The distances between the phenanthroline complexes are 7.56−8.25 and 6.58 Å in the b and a axes directions, respectively (Figure 1).
where $\Delta E$ (adsorption energy) is the energy involved in the adsorption of the molecule(s) of adsorbate (naphthalene) in the double supercell of Mt/Fe(III)Phen; $E_{\text{adsorption complex}}$ is the total energy of the double supercell of Mt/Fe(III)Phen hosting the molecule(s) of the adsorbate (naphthalene); $E_{\text{adsorbate}}$ is the total energy of the molecule(s) of the isolated adsorbate species (naphthalene); and $E_{\text{host}}$ is the total energy of the double supercell of Mt/Fe(III)Phen alone.

3. RESULTS

3.1. Selective Trapping of Aromatic Gas Phases by Mt−Fe(III)Phen. Figure 2 shows the amount of benz, Clbenz, naph, and Clnaph immobilized by Mt−Fe(III)Phen ($q_{\text{arom}}$, expressed as moles of immobilized species per 100 g of Mt−Fe(III)Phen) as a function of $q_{\text{complex}}$ (moles of Fe(III)Phen per 100 g of Mt−Fe(III)Phen). (naph = triangle, Clnaph = circle, benz = diamond, Clbenz = square).

![Figure 2. Number of moles of aromatic compounds immobilized at 25 (open symbols) and 50 °C (filled symbols) per 100 g of Mt−Fe(III)Phen ($q_{\text{arom}}$) as a function of $q_{\text{complex}}$ (moles of Fe(III)Phen per 100 g of Mt−Fe(III)Phen). (naph = triangle, Clnaph = circle, benz = diamond, Clbenz = square).](image)

The maximum immobilization capacity ($q_{\text{arom}}^{\text{max}}$) strongly increases with temperature. In fact, the $q_{\text{arom}}^{\text{max}}$ value for naph ranges from 0.035 mol/100 g Mt−FePhen at 25 °C to 0.060 mol/100 g Mt−FePhen at 50 °C and for Clnaph from 0.043 mol/100 g Mt−FePhen at 25 °C to 0.067 mol/100 g Mt−FePhen at 50 °C. These entrapment performances are remarkable because they were achieved at rather low vapor pressures of adsorbates. At $q_{\text{complex}}$ values higher than that corresponding to $q_{\text{arom}}^{\text{max}}$, the entrapment efficiency clearly decreases and, at the saturation limit in Fe(III)Phen ($q_{\text{complex}} = 0.038$), for naph and Clnaph, the immobilization ability drops at about 60% (67%) and 35% (27%) of the $q_{\text{arom}}^{\text{max}}$ at 25 °C (50 °C). Because of the low efficiency of Mt−Fe(III)Phen in the immobilization of benz and Clbenz, data on these two aromatic gaseous phases were omitted.

3.2. Temperature Dependence of the Trapping Kinetics for Naphthalene and 1-Chloronaphthalene. The adsorbing rates of naph and Clnaph on Mt−Fe(III)Phen were obtained by measuring the amounts of the aromatic compounds immobilized by Mt−Fe(III)Phenbest25 and Mt−Fe(III)Phenbest50 (see Experimental Section) as a function of time at both temperatures (Figure 3). Upon exposure to naph and Clnaph, the content in the aromatic compound inside Mt−Fe(III)Phenbest25 and Mt−Fe(III)Phenbest50 at 25 °C (open symbols) and by Mt−Fe(III)Phenbest50 at 50 °C (filled symbols), $q_{\text{arom}}$, is the number of moles of immobilized aromatic molecule per 100 g of Mt−Fe(III)Phen. The Fe(III)Phen adsorbing materials are prepared with $q_{\text{complex}} = 0.022$ at 25 °C and $q_{\text{complex}} = 0.025$ at 50 °C (see Experimental Section).

![Figure 3. Entrapping kinetics of naph (triangle) and Clnaph (circle) by Mt−Fe(III)Phenbest25 at 25 °C (open symbols) and by Mt−Fe(III)Phenbest50 at 50 °C (filled symbols). $q_{\text{arom}}$ is the number of moles of immobilized aromatic molecule per 100 g of Mt−Fe(III)Phen. The Mt−Fe(III)Phen adsorbing materials are prepared with $q_{\text{complex}} = 0.022$ at 25 °C and $q_{\text{complex}} = 0.025$ at 50 °C (see Experimental Section).](image)

and Clnaph, the content in the aromatic compound inside Mt−Fe(III)Phenbest25 and Mt−Fe(III)Phenbest50 (calculated by the carbon content exceeding that of Fe(III)Phen) strongly increases with time to reach a constant value after about 48 h at 25 °C and after about 24 h at 50 °C for both naph and Clnaph (Figure 3). These data show that the process takes place in a single step and the entrapment process is fast, in particular at 50 °C, according to the increase of the partial pressure and the adsorption rate constant with temperature. The amount of trapped gas phase at 50 °C is about twice higher than at 25 °C.

3.3. Thermogravimetric Measurements and Regeneration/Reversibility Tests on Mt−Fe(III)Phen−naphbest25 and Mt−Fe(III)Phen−Clnaphbest25. Mt−Fe(III)Phenbest25 saturated in naph and Clnaph at 25 °C (Mt−
Fe(III)Phen–naphbest25 and Mt–Fe(III)Phen–Cl(naphbest25) were characterized by TGA and MS-EGA, respectively, to determine the temperature of desorption and the nature of the released species (Figures 4 and 5). The thermal behavior of Mt–Fe(III)Phen alone was characterized previously. After exposure to the aromatic molecules, major changes in TGA [(and its first derivative curve, differential TGA (DTGA)] were observed in the temperature range 110–300 °C (Figures 4 and 5). In particular, the DTGA profiles of Mt–Fe(III)Phen–naphbest25 and Mt–Fe(III)Phen–Cl(naphbest25) show a well-defined peak at $T = 70$ °C and a shoulder at about 135 °C that are related to the dehydration of the interlayer (H$_2$O release, $m/z = 18$), already observed for Mt–Fe(III)Phen alone. A further noticeable thermal effect appears at about 220 °C for Mt–Fe(III)Phen–naphbest25 and at 205 °C (the peak also shows a well-defined shoulder at 240 °C) for Mt–Fe(III)Phen–Cl(naphbest25). This is not observed for Mt–Fe(III)Phen and can be related to the release of the undecomposed aromatic molecule (naph release, $m/z = 128$), between 150 and 290 °C; Cl(naph) release, $m/z = 162$ and 127 after the loss of Cl, between 160 and 340 °C. At temperatures higher than 350 °C, the DTGA profiles closely resemble that obtained for Mt–Fe(III)Phen alone and correspond to the thermal decomposition of the iron complex and, afterward, to Mt dehydroxylation. The analysis of the thermogravimetric profiles coupled with the MS-EGA shows that within the temperature range 20–350 °C, the mass loss due to dehydration is partially overlapped to the desorption of the aromatic molecules. In the DTGA curves, however, the peaks corresponding to the two processes are well resolved, allowing assignment of the first process (at $T$ about 70 °C) to the emission of the interlayer water and the second mass loss (at about 200–220 °C) to that of the aromatic compounds. In Mt–Fe(III)Phen, the loss of interlayer water was previously quantified being about 5.46%; the contributions in mass loss due to water and aromatic molecule can be evaluated by TGA: for Mt–Fe(III)Phen–naphbest25, these contributions are 2.96 ± 0.26% of H$_2$O and 4.12 ± 0.14% of naphthalene, while for Mt–Fe(III)Phen–Cl(naphbest25) they are 3.16 ± 0.24% of H$_2$O and 6.21 ± 0.48% of 1-chloronaphthalene. The contents in the aromatic compound determined by TGA (0.033 ± 0.002 mol/100 g Mt–Fe(III)Phen for naph and 0.039 ± 0.003 mol/100 g Mt–Fe(III)Phen for Cl(naph)) are in good agreement with those calculated by elemental analyses (0.035 ± 0.003 mol/100 g Mt–Fe(III)Phen for naph and 0.043 ± 0.002 mol/100 g Mt–Fe(III)Phen for Cl(naph) both at 25 °C). The release at relatively low temperature of the intact aromatic compound indicates that Mt–Fe(III)Phen can be used several times as the adsorbent material. Mt–Fe(III)Phen–naphbest25 and Mt–Fe(III)Phen–Cl(naphbest25) were heated at 160, 180, and 200 °C to optimize the conditions for material regeneration, in particular, to determine the lowest temperature at which a complete and quick removal of the aromatic compound is possible. In fact, Figures 4 and 5 show that the maximum rates of the release of the aromatic compounds are reached at 220 and 205 °C for naph and Cl(naph), respectively, but the release processes already start at about 160 °C.

The changes in the content of aromatic molecules at the different temperature values versus time are reported in Figure 6 for Mt–Fe(III)Phen–naphbest25 and Mt–Fe(III)Phen–Cl(naphbest25). The rate of desorption is dependent on the

Figure 4. Thermogravimetric curves (TGA and DTGA) and mass analysis of the evolved gases as a function of temperature recorded at $m/z = 18$ (H$_2$O), 30 (NO and NO$_2$), 44 (CO$_2$), and 128 (naph) of Mt–Fe(III)Phen–naphbest25; $q_{\text{complex}} = 0.022$ moles of Fe(III)Phen per 100 g Mt–Fe(III)Phen corresponding to the maximum immobilization capacity at 25 °C.
treating temperature; at all the temperatures, however, complete desorption can be obtained. In particular, at 200 °C after 10 min a completely desorption of naph and Clnaph occurs from Mt\textsuperscript{−}Fe(III)Phen\textsuperscript{−}naph\textsuperscript{best25} and Mt\textsuperscript{−}Fe(III)Phen\textsuperscript{−}Clnaph\textsuperscript{best25}. The thermal treatment at \(T \leq 200\) °C is fully reversible for at least 20 cycles, that is, the hybrid material can thermally release and, afterward, uptake the aromatic compound again in the same amount and with a similar time course (Figure 7).

### 3.4. \textsuperscript{1}H and \textsuperscript{13}C CP-MAS NMR Characterization of Mt–Fe(III)Phen Exposed to Naphthalene and 1-Chloronaphthalene.

\textsuperscript{1}H NMR spectra (Figure 8) of the samples exposed to naph (Mt–Fe(III)Phen–naph\textsuperscript{best50} Figure 8b) and Clnaph (Mt–Fe(III)Phen–Clnaph\textsuperscript{best50} Figure 8a) confirm the presence of the aromatic molecules immobilized inside Mt–Fe(III)Phen\textsuperscript{best50}. In fact, together with signals of structural hydroxyl protons at 2.8 and 1.4 ppm, a signal at 7.4 ppm appears after exposure to naph, whereas a resonance at 7.4 and a shoulder at 8.2 ppm are observed in the case of Clnaph. These chemical shifts parallel those observed in solution for the two host molecules.\textsuperscript{22} Both spectra are quite different from that of Mt–Fe(III)Phen\textsuperscript{best50} (Figure 8c). The same holds for \textsuperscript{13}C CP-MAS NMR spectra (Figure 9), which are both dominated by a signal at 126.6 and 127.5 ppm for the samples exposed to naph and Clnaph, respectively.

The comparison between the \textsuperscript{1}H NMR spectral profiles of Mt–Fe(III)Phen–Clnaph\textsuperscript{best50} and Mt–Fe(III)Phen–naph\textsuperscript{best50} (inset Figure 8) shows that Clnaph signals at 8.2 and 7.4 ppm slightly exceed that of naph at 7.4 ppm. Deconvolution of \textsuperscript{1}H NMR spectra allows to derive a Clnaph/naph molar ratio of about 1.14, in fair agreement with that
obtained by elemental analysis (from Figure 2, at 50 °C: $q_{\text{Claph}}^\text{max} / q_{\text{naph}}^\text{max} = 0.067/0.060 = 1.12$). Structural hydroxyl protons are found to be slightly shielded with respect to those of Mt−Fe(III)Phen−$\text{Claph}_{\text{best25}}$ (3.2 and 1.9 ppm) and even to those of Mt itself (3.0 and 1.7 ppm). 19

3.5. Computational Simulation of the Interlayer of Mt−Fe(III)Phen Saturated with Naphthalene. The modeling of the adsorbate gaseous phases into Mt−Fe(III)Phen was reported only for naphthalene. The adsorbate molecules, naphthalene and naphthalene dimer, were included in a cubic periodical box of $20 \times 20 \times 20$ Å. In the naphthalene dimer, both aromatic rings were parallel and with the same orientation at a distance of 4 Å. The optimization of this dimer maintained parallel both aromatic rings with one ring rotated 90° with respect to another.

By placing one naphthalene molecule between both phenanthroline complexes in a parallel disposition with respect to the phenanthroline rings and the (001) clay surface, the optimization yielded a structure with the naphthalene molecule inclined between both phenanthroline complexes and not intercalated inside the phenanthroline rings (Figure 10a). The shortest distances between the naphthalene H atoms and the phenanthroline C atoms are at 3.5 Å. The adsorption energy was $-23.75$ kcal/mol, indicating that this adsorption is energetically favorable.

Two naphthalene molecules at 3.6 Å in a parallel disposition were also placed in a similar initial position as above, parallel to the clay surface. The optimized structure showed that both adsorbates remained parallel at 3.5 Å and inclined with respect to the phenanthroline complexes at 2.4−3.3 Å between naphthalene and phenanthroline rings (Figure 10b). Starting with the naphthalene molecules placed separately, one close to the clay surface and another in the center of the interlayer space, the optimization yielded the same above structure. The adsorption energy was $-45.4$ kcal/mol, indicating also a highly favorable process.

Besides, another initial configuration was considered, placing both naphthalene molecules in a perpendicular disposition with respect to the mineral surface. The optimization of this structure showed that the adsorbates remained in the same perpendicular disposition with respect to the phenanthroline complexes at 2.4−3.3 Å between naphthalene and phenanthroline rings (Figure 10c). The naphthalene rings remained parallel at 3.45 Å. The distances between the naphthalene and phenanthroline rings are at 2.28−2.70 Å. Besides, a certain interaction between the naphthalene H atom and the mineral surface basal O atoms is observed at 2.6 Å. The adsorption energy was $-52.16$ kcal/mol. These short intermolecular distances explain the higher adsorption energy of this model.
1.5 molecules of naphthalene per Fe complex. Then, we included three naphthalene molecules in a parallel disposition and placed them perpendicularly with respect to the (001) clay surface (Figure 10d). After the optimization, all naphthalene molecules remained parallel with inter-ring distances of 3.66–3.70 Å and perpendicular to the (001) plane of the clay mineral. The shortest distances between the naphthalene rings and the phenanthroline rings are 2.84–2.98 Å. In the same way, this trimer of naphthalene was optimized in a 20 × 20 × 20 Å box maintaining all aromatic rings parallel with the middle ring rotated 90° with respect to others. This rotation was not observed when this trimer is intercalated in the clay interlayer space. The adsorption energy was ΔE = −72.89 kcal/mol, indicating an energetically favorable adsorption process according to the above results.

4. DISCUSSION

Mt–Fe(III)Phen is able to rapidly immobilize a large amount of naph and Clnaph but shows an extremely low affinity toward benz and its chloro-derivative. Previous studies have shown that Mt adsorbs Fe(III)Phen in excess (twice the cation exchange capacity, CEC) and induces a structuring of the interlayer as a function of the amount of the immobilized complex. The interaction of Mt–Fe(III)Phen with aromatic species in the gas phase is likely to be controlled by the presence and disposition of the aromatic moiety of the iron complex mainly through the shaping of the interlayer volume with hydrophobic channels and holes having suitable dimensions for guest molecules. If the holes are large enough, π-stacking interactions could be established between adjacent naphthalene rings. Molecules with large aromatic regions, such as naphthalenes, yield stronger intermolecular stacking interactions than benzene and its derivatives, which are also more volatile. These reasons would explain the larger affinity of Mt–Fe(III)Phen for naphthalenes than benzenes: maybe shape and dimensions of the array made by three naphthalene molecules, with strong π interactions, fit better with the available hydrophobic nanospaces than the benzene-type ones. This selectivity is a step ahead with respect to classical organoclays, which, albeit showing highly hydrophobic interlayers, are not selective.

This view is supported by the observation that the amount of adsorbed molecules (q_arom) as a function of the composition of the adsorbent hybrid material (q_complex) shows a maximum. In fact, the amount of immobilized aromatic derivative is controlled by two opposite contributions: the available volume in the interlayer and the amount of aromatic moieties with which the adsorbate can interact. The peak of maximum adsorption corresponds to a slightly higher amount of the stoichiometric proportion related with the CEC capacity, named semi-saturated sample in previous work. This excess of Fe complex acts as a pillaring agent taking into account previous X-ray diffraction profiles, where the d(001) spacing increased with a partial excess of Fe complex. This corroborates previous molecular modeling work where these excess Fe complexes have a perpendicular disposition with respect to the interlayer surface, acting as pillars in the interlayer space. These pillars do not avoid the entrance of aromatics and the increase of d(001) spacing and internal free volume facilitates a slightly higher adsorption capacity.

The maximum adsorption of naph and Clnaph at 50 °C is observed at a q_complex value higher than that determined at 25 °C. This result is probably related to the fact that temperature disfavors entropically and enthalpically the adsorption process, which is exothermic. Clnaph is immobilized to a larger extent
than naph at both temperatures, despite the larger vapor pressure of the latter compound, especially at 50 °C. This effect could be related to the dipole induced by Cl in the Clnaph molecule. The entrapping efficiency for naph and Clnaph is larger at 50 °C than at 25 °C. High temperature, although in general disfavors the adsorption processes, increases the vapor pressure of the aromatic molecule, which enhances the entrapment. However, this increase of vapor pressure with the temperature is much higher than the increase of adsorption efficiency (e.g., for Clnaph, $P_{25°C}/P_{35°C} = 6.8$, while $d_{50°C}/d_{25°C} = 1.6$). This can indicate that the intercalation of aromatics decreases the entropy, that is, the adsorption has a negative entropy contribution, explaining the lower increase in desorption efficiency with higher temperature with respect to the higher increase in vapor pressure.

The entrapping performances, achieved at rather low vapor pressures of adsorbates, are similar or even higher than those obtained using other well-known high-performance adsorbing materials such as synthetic zeolites, active carbons, or organoclays. In organoclays, the intercalation of alkyl chains in the interlayer imparts the material with the ability to bind exogenous molecules through not specific hydrophobic interactions. The material proposed here exploits the structuring effect that the iron complex has on the interlayer to efficiently bind exogenous aromatic hydrocarbons.

In addition, the immobilization rate is also high at 25 °C and the hybrid material also proved to be stable over time after treatment at 200 °C and reusable. Desorption is also fast at 160 °C, and the released species is the pristine species. This allows a quantitative recovery of the aromatic species and adsorbent solid.

5. CONCLUSIONS

We have shown that Mt–Fe(III)Phen, a hybrid material easily obtained by fast intercalation of the Fe(III)–Phen complex into Mt, is able to efficiently trap naphthalene and 1-chloronaphthalene at the solid/gas interface. This property is related to the strong interactions of these species with the aromatic moiety of the iron complex, providing a stable immobilization. This process occurs at room temperature and its characteristics can be modified changing the composition (iron complex content) of the hybrid material. In fact, the efficiency of Mt–Fe(III)Phen as naphthalenes trap is controlled by the saturation level of the complex, which modulates structuration, hydrophobicity, and availability of the aromatic moiety inside the interlayer. In particular, structuration of the interlayer probably results in the formation of a network of strongly hydrophobic channels lined by the phenanthroline moieties of the Fe(III)Phen aggregates. The immobilization is fast and fully reversible: the entrapment material can be thermally regenerated and brought back to its full immobilizing capacity quickly and in mild conditions. The high adsorption efficiency of Mt–Fe(III)Phen is mainly determined by the unique nanostructure and tailorable properties of the interlayer, which make it suitable for the capture of various aromatic molecules which have a strong environmental impact. Because the mechanism that controls naphthalene immobilization probably involves the formation of stacking interactions between molecules entrapped in hydrophobic interlayer spaces, the structure and the extent of the hydrophobic channels are probably the determinants of the observed entrapping activity of Mt–Fe(III)Phen.

The already proved entrapping ability of Mt–Fe(III)Phen toward thiols and H2S has been here extended to the aromatic molecules of environmental interest. The novelty of the present work is the exploitation of the aromatic moiety of Fe(III)Phen, which is able to structure the interlayer to form hydrophobic channels which allow the specific entrapment of naphthalene and 1-chloronaphthalene. We are also convinced that it is possible to expand the opportunities offered by this approach employing other ligands with affinity for aromatic moieties and extending the research to the study of the immobilization of other aromatics.

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