Halloysite nanotubes filled with salicylic acid and sodium diclofenac: effects of vacuum pumping on loading and release properties

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Abstract
In this work, we investigated the effects of the vacuum pumping on both the loading efficiencies and the release kinetics of halloysite nanotubes filled with drug molecules dissolved in ethanol. As model drugs, salicylic acid and sodium diclofenac were selected. For comparison, the loading of the drug molecules was conducted on platy kaolinite to explore the key role of the hollow tubular morphology on the filling mechanism of halloysite. The effects of the pressure conditions used in the loading protocol were interpreted and discussed on the basis of the thermodynamic results provided by Knudsen thermogravimetry, which demonstrated the ethanol confinement inside the halloysite cavity. Several techniques (TEM, FTIR spectroscopy, DLS and ζ-potential experiments) were employed to characterize the drug filled nanoclays. Besides, release kinetics of the drugs were studied and interpreted according to the loading mechanism. This work represents a further step for the development of nanotubular carriers with tunable release feature based on the loading protocol and drug localization into the carrier.

Graphic abstract
The filling efficiency of halloysite nanotubes is enhanced by the reduction of the pressure conditions used in the loading protocol.

Keywords Halloysite nanotubes · Sustained release · Clay nanoparticles · Drug loading · Knudsen thermogravimetry
**Introduction**

In the very last decades, we observed a growing interest of the scientific community in the study of clays, which were employed in numerous applications. The reason for their rapid diffusion has to be found in their wide variety of different and tunable features, including the morphology, size, interfacial chemistry [1–7]. Moreover, they are natural occurring building blocks that can be used as raw components for the design of new green and eco-sustainable materials [8–13]. Nowadays, the latter aspect appears to be the most important and attractive factor since it deals with some environmental needs which demand great efforts especially from researchers and technologists [14–16]. Halloysite is a very promising aluminosilicate whose unit formula is Al₂Si₂O₅(OH)₄·2H₂O and whose most intriguing feature is its distinctive shape [17, 18]. The dimensions of halloysite nanotubes (HNTs) are influenced by their natural geological deposit [19]. Compared to the carbon nanotubes (CNTs), halloysite nanotubes are easily available and abundant worldwide. Consequently, the price of HNTs is more than 100 times lower with respect to that of CNTs. Moreover, HNTs are biocompatible nanomaterials, while CNTs can be toxic to the human health reducing their potential applications in biotechnological and pharmaceutical fields. Based on the European Union Standards, the amount of heavy metals in halloysite is much lower than the maximum limits on harmful species [1]. Similar to carbon nanotubes, HNTs can be easily rendered conductive by metallization and/or coating with polymers [20, 21]. In addition, both HNTs and CNTs are effective fillers for the mechanical reinforcement of polymeric matrix driving to the preparation of smart nanocomposites [20]. According to the previous considerations, HNTs are considered a green alternative nanofiller with a broad range of technological applications. Basically, the nanotubes length is from 200 nm to 2 µm, their internal diameter is 15–50 nm, whereas the external diameter is 50–200 nm [22]. Furthermore, outer and internal surfaces possess opposite charges because of their different chemistry (Al–OH and Si–O–Si groups, respectively) [23]. Thus, the inner (Al) surface is positively charged, while the outer (Si) surface is negatively charged [24]. Consequently, this unique property can be exploited for the surface modification with organic molecules (e.g., polymers, biopolymers or surfactants), through both electrostatic and van der Waals interactions, thus leading to the design of hybrid and bio-hybrid systems that find applications in a broad range of applications [25–34]. In general, electrostatic interactions are convenient to target the interaction/adsorption site for halloysite nanotubes [35–38]. Most interestingly, halloysite nanotubes can be loaded with negatively charged active species, such as biological molecules, anti-corrosion compounds and antioxidants that can be protected inside the inner lumen of the nanotubes [39–41]. This one is a very important property, which endows the preparation of new smart nanocarriers for loading, storage and sustained release of chemical agents [42–45]. Moreover, halloysite exhibits low toxicity [32, 46, 47]. To the light of these aspects, thermo-responsive nanocarriers were prepared by the functionalization of HNTs with charged PNIPAAMs for the controlled delivery of drugs into biological systems [48]. Additionally, the design of new hybrid and biohybrid materials was carried out using drug-loaded halloysite for health applications as antimicrobial patches or anti-inflammatory drugs-releasing tablets [49–52]. Similarly, the nanotubes were used for the encapsulation of corrosion inhibitors with the aim to coat and protect metal surfaces [53–55]. Furthermore, HNTs were loaded with antioxidant species to preserve food by designing packaging nanomaterials [56] and, finally, clay nanotubes were used for cultural heritage treatment by loading anti-acid species [57]. Recent reviews and research articles evidenced that halloysite is a support for the development of antimicrobial nanocomposites [58–62]. The control of the loading mechanism is crucial to extend the HNTs applications as drug delivery systems and nanocontainers [63]. In this regard, the confinement effect of a liquid into a very narrow space inside of the nanoparticles plays a crucial role [64]. Similar observations can be considered for halloysite nanotubes. First, Price et al. proved that by subjecting the drug/clay dispersion to a vacuum pumping step, where the pressure is lower than the standard atmospheric value, the drug loading efficiency is greatly enhanced for comparison with encapsulation protocols carried out without any particular pressure control [65]. Hence, in our previous study, we demonstrated the confinement of water within the inner lumen of halloysite and this finding was shown to be deeply correlated with the driving mechanism of nanotubes filling and to the optimization of the loading efficiency. Accordingly, the vacuum pumping allows the overall pressure to approach the vapor pressure of the solvent, which evaporates faster, thus leaving empty space to be filled by the guest molecules that are forced to enter and gather inside [66]. Literature reports several studies on the loading of halloysite from organic solvents by using the vacuum assisted procedure [43, 67, 68]. Nevertheless, the physico-chemical insights of this mechanism have not been studied. In this work, we moved forward by investigating the confinement effect of an organic solvent, namely ethanol, to understand its influence on the loading mechanism, on the drug localization within the clay and on the releasing kinetics. Indeed, this seems to be the most important aspect for all the applications we discussed above, concerning the use of halloysite as controlled and tunable delivery system.
Experimental

Materials

Kaolinite (Kao), diclofenac sodium salt (DF) and absolute ethanol (99.9%) are Sigma Aldrich products. Salicylic acid (SA) is from Fluka. Halloysite nanotubes (HNTs) are from I-Minerals Inc. mined in the geological deposit of Latah County with physico-chemical properties detailed elsewhere [22]. As reported elsewhere [22], the investigated halloysite possesses a basal spacing of 7.58 Å.

Drug loading protocols

The encapsulation of salicylic acid and diclofenac sodium salt within the halloysite cavity was performed by ethanol suspensions following two protocols. First, we prepared drug solutions in ethanol by magnetically stirring for 2 h at 20 °C. The drugs concentration was at the maximum of their solubility (saturated solutions). Halloysite was added keeping a HNTs/drug ratio of 2:1. The HNTs/drug suspensions were subjected to ultrasonication for 5 min. Afterwards, the dispersions were divided into two aliquots following two different protocols: (i) the dispersion was transferred to a vacuum jar for 30 min allowing to reduce the pressure conditions \((P = 0.01 \text{ atm})\) and to remove ethanol. Then, the vacuum was broken for 10 min. The cyclic vacuum pumping step was repeated three times. (ii) The dispersion was kept under stirring for 2 h at \(P = 1 \text{ atm}\). In both the two protocols, room temperature conditions were considered.

The same procedure was repeated using kaolinite to investigate the influence of the vacuum conditions on the filling process by altering the nanoclay morphological features. It should be noted that both drugs are highly soluble in ethanol. In particular, the solubilities in ethanol of salicylic acid and sodium diclofenac are 2.35 [69] and 0.5202 mol L\(^{-1}\) [70], respectively. These values are significantly larger compared to the corresponding molar solubilities in water (0.0159 [69] and 0.0674 mol L\(^{-1}\) [70] for salicylic acid and sodium diclofenac, respectively). Accordingly, loading protocols based on ethanol dispersions could enhance the amounts of drugs encapsulated within the clay nanotubes with respect to those expected using aqueous suspensions. Furthermore, both ethanol and water are largely employed for the filling of halloysite nanotubes with guest molecules as evidenced by recent reviews [21, 47] and research articles [40, 49, 66].

Drug release kinetics

The study of the drug release kinetics was carried out via UV–VIS spectrophotometry. Since the main purpose is to target the molecules accumulation within HNTs and to study the crucial effects on the delivery properties, water was used as releasing medium. Therefore, 10 mg of loaded nanoclays (HNTs or Kao) was placed into water and the drugs release was studied as reported elsewhere [49]. It should be pointed out that the absorbance values at 276 and 296 nm were considered for diclofenac sodium salt and salicylic acid, respectively. Releasing profiles were determined using the following equation [71]:

\[
C'_n = C_n + \left( \frac{V}{V_0} \right) \cdot \sum_i \left( C_i \right),
\]

where \(C_n\) and \(C'_n\) refer to the drug concentrations in water before and after the correction, respectively. On the other hand, \(V\) is the volume for each measured aliquot, while \(V_0\) represents the full volume for each sample. The same procedure was followed for the samples whose drug loading was performed using both the two protocols described in the previous paragraph. Drug release tests were conducted three times. As concerns the dissolution of pure diclofenac and salicylic acid, the whole amounts of pure drugs could be detected in solution already at the first experimental point.

Methods

A Q5000 IR apparatus (TA Instruments) was employed for thermogravimetry (TG). The measurements were performed under inert atmosphere using different \(\text{NO}_2\) flows for the balance and the sample (10 and 25 cm\(^3\) min\(^{-1}\), respectively). The samples were heated at 20 °C min\(^{-1}\) from room temperature to 700 °C. The apparatus was calibrated using the Curie temperature of standards [72]. The drug loading amounts were calculated by the quantitative analysis of TG curves as detailed in Supplementary Material [73]. Knudsen thermogravimetry (KTG) measurements were conducted at 30 °C using Knudsen cells with a hole (diameter of 20 μm) on their top surface. Ethanol dispersions of both nanoclays (concentration of 50 wt%) were investigated by KTG measurements to explore the solvent evaporation process. According to literature [66], ethanol activity coefficients were calculated. The evaporation rate can be expressed as:

\[
dm/dt = K_F \cdot (P_{\text{int}} - P_{\text{ext}}),
\]

where \(K_F\) is a constant related to the volatile gas, \(P_{\text{int}}\) is the partial pressure of the solvent vapor inside the cell and \(P_{\text{ext}}\) is the external pressure. It should be noted that \(K_F\) refers to a given gas escaping through a Knudsen orifice with specific geometrical size and at a specified temperature. Based on the fundamentals of KTG, the thermodynamic activity of ethanol (\(\alpha_{\text{EtOH}}\)) can be estimated by the following equation:

\[
(dm/dt)_n/(dm/dt)_{\text{EtOH}} \approx (P_{\text{int}})_n/(P_{\text{int}})_{\text{EtOH}} = \alpha_{\text{EtOH}}.
\]
being \((\text{dm/dt})_n\) and \((\text{dm/dt})_{\text{EtOH}}\) the mass loss rates for the nanoclay ethanol dispersion and pure ethanol, respectively.

A Zetasizer NANOZS (Malvern Instruments) instrument was used to conduct both dynamic light scattering (DLS) and \(\zeta\)-potential experiments, which were carried out at 25 °C. The analyses of DLS curves provided the apparent hydrodynamic radius (Rh) of pure and loaded clay nanotubes, while \(\zeta\)-potential data were helpful to monitor the effect of the drug loading on the HNTs surface charge.

TEM micrographs were acquired with a Jeol JEM 2100 microscope operating at 200 kV. A drop of each dispersion has been deposited in the grid, which was dried prior to the acquisition of TEM images. A 3 mm nickel grid holey carbon coated (Taab) was selected for TEM analyses. It should be noted that no coating was applied to the samples. TEM images of both pure and loaded HNTs were analyzed using ImageJ software [74] for the determination of the corresponding averaged gray-scale profiles.

Kinetics of drug releases were studied by UV–Vis spectrophotometer (Specord S600 Analytik Jena).

Fourier transform infrared (FTIR) spectra were determined using a Frontier FTIR spectrometer (PerkinElmer) at 25 °C in the wavenumber range between 500 and 4000 cm\(^{-1}\). KBr pellets were employed for FTIR analyses.

Results and discussion

Drugs loading onto halloysite nanotubes: mechanism and structure

The amounts of organic molecules loaded into the nanoclays were determined through thermogravimetry taking into account the equilibrium moisture contents of all materials (before and after the drug loading). The moisture content data are reported in Supplementary Material (Table 1S). Compared to those of pristine clay minerals, the thermogravimetric curves of the hybrid materials evidenced a significant mass loss due to the decomposition of the organic moiety at ca. 250 and 180 °C for DF and SA, respectively (Fig. 1S in Supplementary Material). The pressure conditions induced a relevant effect on the loading efficiency of HNTs. In particular, the reduced pressure enhanced the filling efficacy for HNTs (the loading increases by factors 4 and 6 for DF and SA, respectively); whilst for Kao, the dependence on the preparation condition is less crucial although the reduced pressure generated an increase up to a factor 2 in the loading amount (Fig. 1).

The successful loading of salicylic acid into the nanotubes was confirmed by FTIR spectroscopy, which evidenced the presence of the characteristic signals of the drug (such as vibrations of \(\text{C}=\text{C}\) and \(\text{C}=\text{O}^+\) groups at 1485 and 1582 cm\(^{-1}\), respectively [51]) in the HNTs/SA hybrids.

Morphological investigations were carried out by TEM on halloysite nanotubes before and after drugs loading under different pressure conditions. Figure 2 shows the images for pristine nanotubes as well as for HNTs/DF and HNTs/SA samples prepared with the different protocols. For a better visualization of the status of the HNTs cavity, an averaged gray-scale profile was plotted along the perpendicular direction of the main nanotube axis for all samples (see insets in Fig. 2) to have semi-quantitative additional insights about the effectiveness of the drug loading procedure. The empty cavity of ca. 10 nm with a wall thickness of 20 nm is evidenced by the gray-scale contrast. It is suggested that a lower pressure increases the filling of the halloysite lumen. The cavity appears full filled when the loading is conducted under reduced pressure for both DF and SA. These findings are a semi-qualitative proof of the TG data, which evidenced increases of the drug loading at \(P=0.01\) atm. These aspects are in agreement with DLS measurements, since no change in the hydrodynamic radius can be observed after the loading of the drug inside the clay. As examples, the distribution curves of the intensity as a function of the hydrodynamic radius for HNTs and HNTs/DF samples are presented in Supplementary Material (Fig. 2S). The influence of the pressure on the loading efficiency and drug localization in halloysite nanotubes is correlated with the different volatility of confined and bulk solvents. Namely, the solvent into the lumen possesses a larger vapor pressure compared to the bulk one and, consequently, a flux of ethanol into the halloysite lumen is required to compensate the different evaporation velocity. Consequently, drug molecules

![Fig. 1 Loading amounts for nanoclay/drug systems prepared at different pressure conditions](image-url)
precipitate and their accumulation within the HNTs cavity can be observed.

Furthermore, the influence of the drugs loading on the HNTs surface charge was explored. To this purpose, $\zeta$-potential measurements were performed. As shown in Supplementary Material (Table 2S), the $\zeta$-potential of halloysite is not largely altered by the encapsulation of anionic drugs within the HNTs cavity. Contrary to these results, literature \[21, 34, 48\] reports that the loading of anionic species from aqueous suspensions generates substantial variations of the HNTs $\zeta$-potential as a consequence of the electrostatic attractions with the halloysite inner surface, which is positively charged in a wide pH range. This discrepancy is related to the influence of the solvent polarity (and the dielectric constant) on the acid/base dissociation equilibria. In general, the dissociation constants in ethanol are lower than those in water.
and, consequently, the formation of ions occurs in a smaller extent. On this basis, HNTs/drugs electrostatic interactions are reduced in ethanol compared to those in water.

**Drugs release controlled by the loading protocol**

Halloysite nanotubes have been deeply investigated for applications in drug delivery. However, the possibility to target the drug localization within the nanomaterial is a promising result for attaining tunable release based on different preparation protocols, which would extend the applicability of these systems. On this basis, drug release experiments were performed for SA and DF embedded into the both nanoclays (halloysite and kaolinite) under the two different pressure conditions \( (P = 0.01 \text{ atm} \text{ and } P = 1 \text{ atm}) \). As a general result, the percentage of released drug showed a monotonic increase with the time reaching a constant value that corresponds to the quantitative release of the loaded material (Figs. 3, 4). These trends are typically observed for small molecules released by nanoclays and can be interpreted by considering the drug/nanomaterial interactions as well as the potential intercalation and/or localization of the active molecules in the porous support [75]. It was observed that the loading protocol generates relevant effects on the release profile for halloysite nanotubes based systems (Figs. 3, 4). In particular, the nanotubes loaded at reduced pressure showed slower release kinetics than the corresponding systems obtained at ambient pressure. These results reflect the drugs localization within the HNTs cavity. It should be noted that the incorporation into the halloysite lumen generates a sustained release profile due to confinement and tortuosity effects that are less relevant when the drug is mainly outside the nanotubes. To validate this hypothesis, the release experiments for kaolinite/drug hybrids were considered. Interestingly, the release from kaolinite based systems was not influenced by the loading protocol (Figs. 3, 4). These results are in agreement with the morphological planar structure of kaolinite that, differently than halloysite, does not possess a cavity to be preferentially filled under reduced pressure conditions. For a clear quantitative evaluation, the time necessary for the release of 50% \( (t_{50\%}) \) of the payload is reported in Table 1 for all the systems. It should be noted that the \( t_{50\%} \) values were calculated by considering the maxima of the UV–Vis absorbance as 100% of drug releases, which are in agreement with the loading results obtained from thermogravimetric analysis. Based on \( t_{50\%} \) values, one

![Fig. 3 Release kinetics of sodium diclofenac salt loaded at \( P = 1 \text{ atm} \) (red squares) and \( P = 0.01 \text{ atm} \) (black circles)](image1)

![Fig. 4 Release kinetics of salicylic acid loaded at \( P = 1 \text{ atm} \) (red squares) and \( P = 0.01 \text{ atm} \) (black circles)](image2)

**Table 1** Drug release kinetics data

|            | \( t_{50\%} \) min |
|------------|-------------------|
|            | \( P = 0.01 \text{ atm} \) | \( P = 1 \text{ atm} \) |
| **HNTs**   |                   |
| DF         | 270 ± 30          | 4.7 ± 0.2 |
| SA         | 570 ± 60          | 135 ± 3  |
| **Kaolinite** |                  |
| DF         | 160 ± 20          | 163 ± 12 |
| SA         | 425 ± 20          | 330 ± 20 |
can state that the preparation protocol can tune the release time even by an order of magnitude in the case of halloysite nanotubes; whilst its effect is almost negligible for flat kaolinite clay mineral.

A general summary on the effect of the loading protocol is sketched in Fig. 5.

**Ethanol evaporation in clay nanostructures: mechanism for drugs loading**

The influence of the pressure conditions on the loading of anionic guest molecules was reported in aqueous media and it was strictly correlated with the water confinement [66]. Due to its confinement, water molecules exhibit a faster evaporation if they are located into the halloysite cavity. As a consequence, a flux of the solution driven by capillarity occurs within the HNTs cavity, in which the drug molecules accumulate once the water evaporation takes place. Such a process is certainly facilitated under low-pressure conditions due to the increased evaporation rate. It should be noted that the loading of anionic drugs in aqueous solvent can be favored by the electrostatic interactions. Namely, anionic compounds can interact with the HNTs inner surface, which possesses positive charges. On the contrary, the loading of anionic species in ethanol is slightly affected by electrostatic attractions as demonstrated by the ζ-potential data of HNTs/SA and HNTs/DF hybrids (see Table 2S in Supplementary Material). Therefore, the loading mechanism by ethanol suspensions can be interpreted by considering the solvent confinement within halloysite cavity.

To verify this interpretation, the ethanol evaporation rate in the presence of halloysite nanotubes and platy kaolinite was explored by KTG experiments. The results are discussed in terms of relative vapor pressure after the normalization by the bulk ethanol evaporation ($P/P_°$) as a function of the ethanol/clay mass ratio ($R_{\text{EtOH:Clay}}$).

Figure 6 shows that $P/P_° > 1$ was detected for halloysite suspensions with $R_{\text{EtOH:Clay}}$ ranging between $6 \times 10^{-3}$ and $6 \times 10^{-2}$, while this peculiarity was not observed in kaolinite dispersions. On this basis, we can assert that a fraction of ethanol with a faster evaporation rate than that of pure solvent was induced by the presence of halloysite nanotubes. According to the Gibbs Thomson effect, the latter represents the thermodynamic proof of the ethanol confinement within the HNTs cavity. Based on geometric considerations and the bulk densities, halloysite nanotubes should be totally filled with ethanol for $R_{\text{EtOH:Clay}}$ equals to ca. $5 \times 10^{-2}$. This result is consistent with the composition range ($6 \times 10^{-3} < R_{\text{EtOH:Clay}} < 6 \times 10^{-2}$) in which the ethanol confinement occurs. The Knudsen thermogravimetric data (Fig. 6) allowed us to interpret the loading results (Fig. 1) by taking into account that the replacement of the ethanol solution inside the halloysite lumen is facilitated under low-pressure conditions. This finding is related to the enhancement of ethanol volatilization rate once the solvent vapor pressure is approaching. Consequently, the amounts of drug molecules entrapped into the cavity can be increased by vacuum pumping operation.
From the data measured in the low $R_{\text{EtOH}:\text{clay}}$ range (corresponding to $P/P^o$ values between 0.05 and 0.95), one can investigate the ethanol isothermal desorption from the nano-clays surfaces. Similar to water/clay systems, the Guggenheim–Anderson–de Boer (GAB) model [76] can be used as fitting model, which is expressed by the following equation:

$$R^o (g_{\text{ethanol}}/g_{\text{clay}}) = \frac{K \times C \times P / P^o}{(1 - K \times P / P^o) \cdot (1 - K \times P / P^o + C \times K \times P / P^o)},$$

being $R^o$ the ethanol:clay mass ratio for the monolayer coverage, while $C$ is a constant related to the monolayer adsorption. Moreover, $K$ is a coefficient that depends on the formation of multilayers during the adsorption process.

It should be noted that the GAB model reveals successful in the fitting of ethanol/HNTs and ethanol/kaolinite desorption isotherms. As an example, Supplementary Material (Fig. 4S) shows the fitting by GAB model for the ethanol desorption from HNTs. The adsorption parameters obtained by GAB method are provided in Table 2. Both clays provided $C > 2$ and, therefore, the ethanol desorption isotherms can be classified as type III [76]. The equilibrium constant value for the multilayer formation is ca. one order of magnitude larger than the corresponding constant for monolayer. This result indicates that the ethanol–ethanol interactions are favored compared to those between ethanol and nanoclays [77].

Despite the equivalent surface chemistry, the monolayer saturation was reached at different the ethanol:clay mass ratios for halloysite and kaolinite. In particular, halloysite can adsorb an amount of ethanol that is ca. 30 times larger than kaolinite. This result cannot be ascribed to the larger specific surface of HNT because the specific surface ratio between HNT and Kao is ca. 3.5 [66] and, therefore, it reflects a denser ethanol layer on the halloysite nanotubes than in flat kaolinite.

![Fig. 6](https://example.com/fig6.png)

**Fig. 6** Relative vapor pressure for ethanol as a function of the ethanol to clay mass ratio for kaolinite (red broken line) and halloysite (full blue line) system

| Table 2 Parameters for the ethanol desorption isotherms |
|---------------------------------|-----------------|-----------------|
| $R^o (g_{\text{ethanol}}/g_{\text{clay}})$ | $K$ | $C$ |
| HNTs | $(1.80 \pm 0.09) \times 10^{-3}$ | $0.714 \pm 0.016$ | $5.7 \pm 0.6$ |
| Kaolinite | $(6.23 \pm 0.03) \times 10^{-5}$ | $1.008 \pm 0.001$ | $50.2 \pm 0.9$ |

From the data measured in the low $R_{\text{EtOH}:\text{clay}}$ range (corresponding to $P/P^o$ values between 0.05 and 0.95), one can investigate the ethanol isothermal desorption from the nanoclays surfaces. Similar to water/clay systems, the Guggenheim–Anderson–de Boer (GAB) model [76] can be used as fitting model, which is expressed by the following equation:

Conclusions

This work explores the filling of halloysite nanotubes with drug molecules (salicylic acid and sodium diclofenac) dissolved in ethanol solvent. The influence of the pressure conditions on the loading efficiencies and the release kinetics was investigated by thermodynamic and structural viewpoints. We observed that the vacuum pumping in/out operation determines an increase of the loading amounts of both drugs, which exhibited a slower release kinetics with respect to the corresponding loaded nanotubes prepared at environment pressure. These results were explained on the basis of the Knudsen thermogravimetry experiments, which proved the ethanol confinement within the halloysite cavity as a consequence of the Gibbs Thomson effect. Compared to the bulk solvent, the confined ethanol evidenced a faster evaporation rate that generates a flux of the drug solution within the nanotubes’ cavity, and, consequently, a payload of the guest molecules. This process is enhanced at reduced pressure conditions driving to increase the filling efficiency of halloysite nanotubes as confirmed by morphological investigations. According to these considerations, the loadings and the release properties were not significantly affected by pressure conditions for platy kaolinite filled with salicylic acid and sodium diclofenac. It should be noted that the drugs encapsulation slightly affected the halloysite surface charge ruling out the presence of relevant electrostatic interactions between halloysite inner surface and the guest molecules.

The present work represents a fundamental step for development of halloysite-based nanocarriers with tunable drug loading and release kinetics.

**Supplementary Information** The online version contains supplementary material available at [https://doi.org/10.1007/s40097-021-00391-z](https://doi.org/10.1007/s40097-021-00391-z).
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Compliance with ethical standards

Conflict of interest The authors declare that there is no conflict of interest.

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