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## RESEARCH ARTICLE

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### Key Points:

- Clay content and clay mineralogy significantly affect the induction time of methane hydrate formation as well as the hydrate morphology
- Illite-rich clays reduce the cage occupancy of hydrates and thus the resulting gas storage capacity (GSC)
- Microscale analyses underpins the need to characterize microscopic properties of hydrates to estimate GSC

### Supporting Information:

Supporting Information may be found in the online version of this article.

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## Influence of Clay-Containing Sediments on Methane Hydrate Formation: Impacts on Kinetic Behavior and Gas Storage Capacity

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**Abstract** On Earth, natural hydrates are mostly encountered in clay-rich sediments. Yet their formation processes in such matrices remain poorly understood. Achieving an in-depth understanding of how methane hydrates accumulate on continental margins is key to accurately assess (a) their role in sustaining the development of some chemosynthetic communities at cold seeps, (b) their potential in terms of energy resources and geohazards, and (c) the fate of the methane releases, a powerful greenhouse gas, in this changing climate. This study investigated the formation of methane hydrates and their gas storage capacity (GSC) in clay-rich sediments. A set of hydrate experiments were performed in matrices composed of sand, illite-rich clay, and montmorillonite-rich clay at different proportions aiming to determine the role of mineralogy on hydrate formation processes. The experiments demonstrate that a clay content of 10% in a partially water saturated sand/clay mixture increases the induction time by ~60%, irrespective of the nature of the clay used. The increase in water saturation in the two matrices promotes hydrate formation. Micro-Raman spectroscopic analyses reveal that increasing the clay content leads to a decrease in the hydrate small-cage occupancy, with an impact on the storage capacity. Finally, the analyses of collected natural samples from the Black Sea (off Romania) enable us to estimate the GSC of the deposit. Our estimates is different from previous ones, and supports the importance of coupling multiscale properties, from the microscale to the geological scale, to accurately assess the total amount of methane hosts in hydrate deposits worldwide.

**Plain Language Summary** Natural gas hydrates are amongst the largest methane reservoirs on Earth. They are sensitive to temperature increase. Societal and environmental concerns surrounding natural gas hydrates pertain to their decomposition, and in particular to the amount of gas they may release and its fate: can it trigger geohazards, or jeopardize the development of unique chemosynthetic communities encountered on continental margins? In-depth knowledge of hydrate formation processes and properties are essential to provide reliable answers to these questions. The majority of hydrate deposits are characterized by clay-rich sediments. We led a comprehensive study coupling fine microstructural analyses of both natural and synthetic hydrate samples with macroscale laboratory experiments within different matrices to show how clays can affect their formation kinetics and storage capacity. We found that even a small amount of clay can significantly change the formation kinetics of hydrates, and the matrix mineralogy affects their storage capacity. This study is crucial with a view to accurately assessing the amount of methane trapped in hydrate deposits, and for improved prediction of the consequences of their decomposition on the environment in a changing climate.

## 1. Introduction

Gas hydrates are crystalline compounds in which gas molecules are trapped in cavities called cages formed by water molecules (Desmedt et al., 2012; Ruffine et al., 2018; Sloan & Koh, 2007). The most naturally occurring form of hydrates corresponds to deposits where methane (CH<sub>4</sub>) is overwhelmingly present (more than 99%-mol) (Kvenvolden, 1998). On Earth, they are widely distributed in both the permafrost regions and marine sediments along the continental margins as their formation is governed by low-temperature and high-pressure conditions (Koh et al., 2012; You et al., 2019). Accordingly, they are very sensitive to climate change. Their decomposition on continental margins may affect both seabed stability (Sultan et al., 2004) and the development of autochthonous chemosynthetic communities (Levin, 2005; MacDonald et al., 2003), whereas in permafrost, part of the

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methane released is transferred to the atmosphere and making these deposits a potential contributor to global warming (Dickens et al., 1997; Kennett et al., 2003; Ruppel & Kessler, 2017).

Natural gas hydrates are well known for their high gas-storage capacity, making them the most important methane reservoir on earth, with estimates ranging from  $3\text{--}5 \times 10^{15}$  (Boswell & Collett, 2011; Milkov, 2004; Wallmann et al., 2012) to  $3 \times 10^{18} \text{ m}^3$  (Trofimuk et al., 1975). This huge amount of methane is regarded as an opportunity for countries with limited energy resources to meet their internal and external energy demands (Hancock et al., 2005; Yamamoto et al., 2019). Furthermore, the high gas-storage capacity of hydrates opens a route to innovative development of (coupled)-processes related to seawater desalinization and/or gas storage, with potential applications in carbon capture, utilization and storage (CCUS) (Linga et al., 2007) and hydrogen storage (Nakayama et al., 2010).

Although natural gas hydrates have been investigated for decades, their formation and decomposition processes are still poorly understood, limiting our knowledge on their growth and distribution mechanisms in porous matrices in the natural environment. It is still unclear how sediment mineralogy quantitatively affects the growth kinetics and final distribution of gas hydrates. Yet, such properties are key to a quantitative insight into how fluid flows within a hydrate deposit; which is relevant in understanding community distribution over a hydrate deposit, assessing slope and seafloor stability, and developing reliable technologies for methane production. Besides, the kinetic behavior of hydrate formation is influenced by a variety of parameters associated with sediments such as particle and pore sizes, mineralogy, water saturation, and pore water chemistry (Malagar et al., 2019). These parameters affect the induction time (Benmesbah et al., 2020; S. Wang et al., 2016), the rate of hydrate growth and the water-hydrate conversion rate (Babu et al., 2013; Linga et al., 2009; Loh et al., 2012; Mekala et al., 2014). Therefore, to obtain in-depth knowledge on formation kinetics and the distribution of hydrates in porous matrices, it is important to investigate the influence of each of the aforementioned parameters separately. However, such an endeavor will lead to a huge amount of experiments to carry out. Therefore, the selection of key parameters to investigate is crucial. Moreover, most of natural gas hydrate deposits are hosted by silty and clayey sediments (Boswell & Collett, 2011; Boswell et al., 2020) and the mechanisms of hydrate formation in such fine-grained sediments are still poorly investigated at the lab scale. This is partly due to the difficulty in forming hydrate in poorly permeable clays (Chuvilin et al., 2002; Lei & Santamarina, 2018). From a mineralogy point of view, clays are hydrated aluminosilicates with a complex chemical and crystalline structure characterized by a stack of layers forming thin plate-like particles (Grim, 1962), whereas from a geotechnical ones, they represent sediment particles with sizes smaller than 2 or 4  $\mu\text{m}$  (Das, 2008). These features result in a large specific surface area and small pores size (Das, 2008; Kumari & Mohan, 2021; Shmulovich et al., 1994), and gives rise to important physico-chemical interactions in the presence of water such as adsorption, structural swelling and ionic exchanges, which can significantly affect the hydrate formation processes.

There are few studies what have focused on the influence of clay on the kinetics of hydrate formation in clay-rich matrices. Among them, Zeng et al. (2022), by studying methane hydrate formation in mixtures of silica sand and montmorillonite, found that at low montmorillonite content (10–25 wt %) the induction time is shortened, and it significantly increases when the montmorillonite content reaches 40 wt %. However, Zhang et al. (2017) showed that the addition of 10 wt % bentonite to a sand-clay matrix increases the induction time from 560 to 2,600 min and decreases the average hydrate growth rate by  $\sim 50\%$ . This inhibiting effect was attributed to restricted contact between gas and water resulting from the presence of fine clay particles between sandy grains. Likewise, Kumar et al. (2015) studied methane hydrate formation in different mixtures of sand and bentonite at various clay contents (25, 50, and 75 wt %) in a 500  $\text{cm}^3$  cylindrical reactor using the excess gas method. They found that even 25% of bentonite reduces the rate of hydrate formation by a factor of 4 compared to a matrix composed exclusively of sand. The lowest water to hydrate conversion rate was achieved for a sand/clay ratio of 25/75 wt %. They also investigated the influence of water saturation and noted that an increase in this parameter from 50% to 100% drastically reduced the rate of hydrate growth by a factor of 7 in a sand/clay mixture with a ratio 25/75 wt %. This has been related to better water-gas contact in a partially water-saturated sediment.

As mentioned above, there is a large discrepancy on the amount of methane trapped in these deposits, and very little attention has been paid to the potential influence of the nature of the host sediments on their structural characteristics, particularly on the occupancy of the large ( $\theta_{LC}$ ) and small ( $\theta_{SC}$ ) hydrate cages (Sloan & Koh, 2007). The cage occupancy is crucial in calculating the gas storage capacity (GSC), corresponding to the hydrate volumetric expansion factor (EF) at standard temperature and pressure (Boswell & Collett, 2011; Meroy & Sinayuc, 2016). Accurate determination of this parameter will help in the reduction of uncertainties on the estimates of methane trapped within the hydrates. This is essential, not only in determining their potential as an energy resource or to

assess seafloor/slope stability, but also to better constrain and predict their possible impact on climate change. Liu et al. (2008) carried out a study on hydrate structure by Raman spectroscopy, and they showed that the hydrate cage occupancy ( $\theta_{LC} \sim 96.5\%$  and  $\theta_{LC} \sim 95\%$ ) is similar in bulk phase and in silica sands with particle sizes ranging from 53–75 to 150–180  $\mu\text{m}$ . The physico-chemical interactions between water molecules and clays are directly related to their specific nature (surface area, interlayer cations, structural organization), and can in turn significantly affect host-guest interactions during hydrate formation (Davidson et al., 1987; Seol et al., 2010). For this reason, Yeon et al. (2011) investigated methane occupancy of natural hydrate in Na-montmorillonite using solid state  $^{13}\text{C}$  MAS NMR and found that sodium ( $\text{Na}^+$ ) cations were inserted into small cages (SC). This leads to values of relative cage occupancy ratios ( $\theta_{LC}/\theta_{SC}$ ) as high as 2.4, thus far higher than the unity when small and large cages are completely filled. Such an occupancy of SC by  $\text{Na}^+$  was observed in bulk propane hydrate by Seol et al. (2012). Their results obtained in sand and clay illustrate the need for further experiments to better assess how and to what extent clayey sediments may affect cage occupancy.

The main goal of this work is to improve our understanding on the formation kinetics of methane hydrates in the presence of complex matrices containing sand, illite, and montmorillonite; and to assess how the sediment mineralogy affect hydrate storage capacity. The two aforementioned clays are characterized by very different chemical structures and swelling properties, and are commonly encountered in marine sediments (Fütterer, 2006), such as the hydrate-bearing sediments of the Western Black Sea (Chazallon et al., 2020; Ruffine et al., 2021). First, we investigated the effects of the nature and fraction of clays on the induction time and hydrate distribution. Different water saturations were applied in order to assess the importance of water content on hydrate formation. Second, Raman micro-spectroscopy has been performed on the synthesized hydrates to determine their cage occupancy and calculate their GSC. Finally, we compare the results with those obtained on the natural hydrate specimens collected on the Romanian margin of the Black Sea, and discuss the implications of the results regarding the estimation of the amount of  $\text{CH}_4$  stored in the natural hydrate deposits of the Black Sea.

## 2. Materials and Methods

### 2.1. Materials

We selected Fontainebleau (FB) silica sand (Laboratoires Humeau, France), illite-rich clay (also called ILL in the following text) supplied by Argiletz Laboratoires (France) and Montmorillonite-rich clay K10 (called MMT hereafter) from Sigma Aldrich, as the porous media. Table 1 summarizes the semi-quantitative mineralogical analysis performed by X-ray diffraction (XRD). It shows that the matrix ILL contains  $\sim 60\%$  clay, 24% quartz, while MMT is composed of 66% clay and 20% quartz.

Illite belongs to the 2:1 clay family in which the elementary aluminosilicate sheet is characterized by an octahedral (aluminum) layer sandwiched between two tetrahedral (silica) layers. The substitution of one  $\text{Si}^{4+}$  ion out of four by an  $\text{Al}^{3+}$  ion in the tetrahedral layers leads to a negative charge of the sheet which is compensated by the presence of  $\text{K}^+$  cations in the interlayer space (Weil & Brady, 2017). The strong interactions between these anhydrous cations ( $\text{K}^+$ ) confer a non-expansive behavior to illite in the presence of water. Montmorillonite belongs to the smectite group and it is also a 2:1 clay, like illite. They differ from each other mainly in the nature of interlayer counter-ions such as  $\text{Na}^+$ ,  $\text{Ca}^{2+}$ ,  $\text{Mg}^{2+}$  or  $\text{Li}^+$  (Odom & Low, 1978; Weil & Brady, 2017), resulting in a larger internal specific surface area and a higher cation-exchange capacity relative to illite. The swelling capacity of the MMT K10 used in this study is reduced due to a chemical treatment applied by suppliers.

Fontainebleau Sand is 100% quartz, characterized by grain sizes between 80 and 450  $\mu\text{m}$  (Figure 1). The clay matrices are composed of elongated platy grains (Figures 1a and 1b), with particle sizes ranging from 0.3 to 80  $\mu\text{m}$  (mean size of 5  $\mu\text{m}$ ) and from 0.6 to 100  $\mu\text{m}$  (mean diameter of 12.7  $\mu\text{m}$ ) for ILL and MMT, respectively (Figure 1c).

We used deionized water (18  $\text{M}\Omega$  cm at 25°C) degassed by ebullition prior to the experiment and methane with a purity of 99.995% supplied by Air Liquid.

### 2.2. Laboratory Methane Hydrate Formation in a High-Pressure Cell

#### 2.2.1. Experimental Apparatus

The high-pressure apparatus used was designed to simulate upward gas transport as encountered at seeps in natural sedimentary environments. It allows the synthesis of small hydrate-bearing cores, while studying selected

**Table 1**  
Main Properties of the Matrices Used in This Study

| Property                              | Illite-rich clay (ILL)     | Montmorillonite-rich clay (MMT) | Sand FB                    | Characterization method   |
|---------------------------------------|----------------------------|---------------------------------|----------------------------|---|
| Particle size ( $D_{\text{mean}}$ )   | 5.3 $\mu\text{m}$          | 12.7 $\mu\text{m}$              | 212 $\mu\text{m}$          | Malvern Mastersizer 3,000 laser diffraction   |
| Coefficient of uniformity ( $C_u$ )   | 4.53                       | 5.93                            | 1.55                       | $C_u = D_{60}/D_{10}$   |
| Roundness ( $R$ )                     | –                          | –                               | 0.4                        |   |
| Specific surface ( $S_s$ )            | 66.4 $\text{m}^2/\text{g}$ | 240.8 $\text{m}^2/\text{g}$     | 0,01 $\text{m}^2/\text{g}$ | ILL and MMT: $\text{N}_2$ BET adsorption measurements<br>Sand: $S_s = 6/(G_s \times \rho_w \times D_{50})$ (Santamarina et al., 2001) |
| Global mineralogy                     |                            |                                 |                            | Sand: from supplier specifications  |
| Quartz                                | 29%                        | 20%                             | 100%                       |   |
| Clays                                 | 56%                        | 66%                             |                            |   |
| Calcite                               | 13%                        | –                               |                            | LL and MMT: X-ray diffraction using a D8 Advance BRÜKER-AXS   |
| Gypsum                                | 1%                         | –                               |                            |   |
| Plagioclase                           | –                          | 14%                             |                            |   |
| Clay fraction mineralogy <sup>a</sup> |                            |                                 |                            | X-ray diffraction after separation of the clay fraction from the other components of the matrix                                       |
| Smectite                              | <5%                        | 82%                             |                            |   |
| Illite                                | 77%                        | 15%                             |                            |   |
| Kaolinite                             | 17%                        | <5%                             |                            |   |
| Chlorite                              | <5%                        | <5%                             |                            |   |

<sup>a</sup>Clay minerals have been separated from the rest of the matrix before XRD analyses, by a series of preparations and chemical treatments (Grim, 1962).

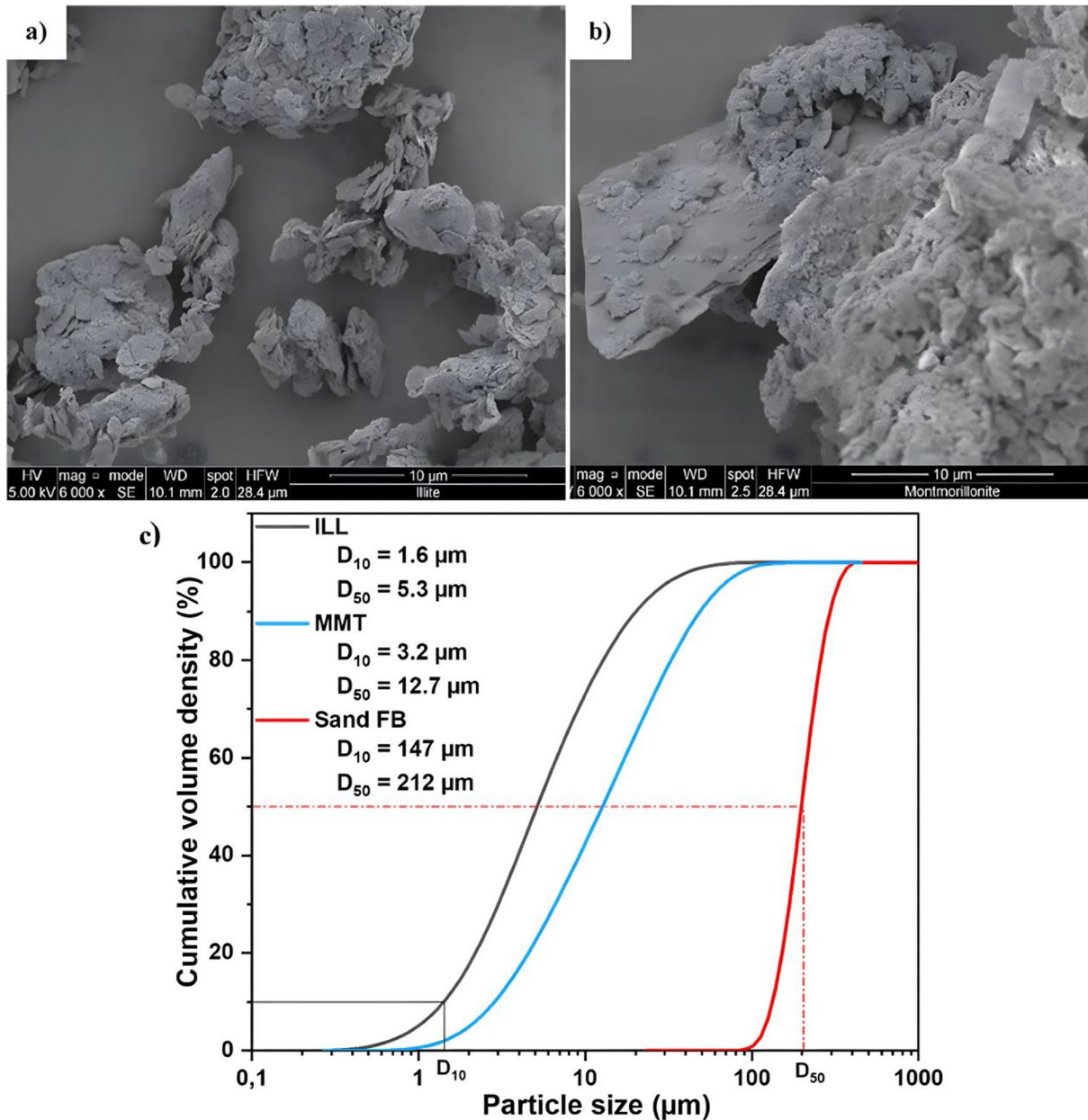
physicochemical parameters of the systems and/or key environmental factors that control their formation and decomposition processes.

Its detailed description is provided in Ruffine (2015) and its schematic diagram is shown in Figure 2. Briefly, it consists of a stainless steel high-pressure cylindrical cell (316-L) manufactured by TOP-Industrie (France). The cell measures 17 cm in height and 6 cm internal diameter with a volume of ~508 mL, and can be used at pressures up to 22 MPa and temperatures ranging from 253 to 373 K. Two flanges located at the top and the bottom hermetically seal the cell. Thermal regulation is ensured by fluid circulation using a thermostatic bath, Ministat 230 (Huber), through a jacket housing the cell. A 15-cm-long thermocouple type K with an uncertainty of  $\pm 0.2$  K, inserted in the upper flange allows for temperature monitoring during experiments. Pressure is measured by a WIKA P-31 pressure transducer located at the bottom of the cell with an uncertainty of  $\pm 0.02$  MPa. These two sensors are connected to a Eurotherm Chessel 6100E data logger that allows continuous data acquisition. Water and methane injection are controlled by a set of high-pressure mass flow and pressure regulators (Bronkhorst). The Bronkhorst Flow DDE, Flow View and Flow Plot softwares provide instrument control and data recording. Thus, the flow rate of water can be adjusted between 0.05 and 2.5  $\text{g min}^{-1}$  with an accuracy of 1% of the full scale (FS), and gas-flow rate from 10 to 500  $\text{mL min}^{-1}$  with an accuracy of 0.1% of FS.

### 2.2.2. Experimental Procedure

We prepared matrices of pure sand, clay matrix/sand (ILL/sand/and MMT/sand) with clay content up to 60%, and ILL/MMT at ~60% clay. The mixtures were prepared by weighing, taking into account the fact that the ILL and MMT clay matrices are not entirely composed of clay minerals (Table 1). Thus, the proportions of each mixture component (ILL/sand, MMT/sand, and ILL/MMT) were determined according to the desired clay content. The mass ratios of clay matrix/sand and the corresponding clay contents are given in Table 2. It is therefore necessary to make the link between the clay content (fraction of clay minerals) in a mixture, and the proportion of the clay matrix (MMT or ILL) in that mixture.

The prepared matrix was either introduced by gravity directly into the high-pressure cell or into a cylindrical aluminum cell that is inserted afterward in the high-pressure cell (Figure S1 in Supporting Information S1). The

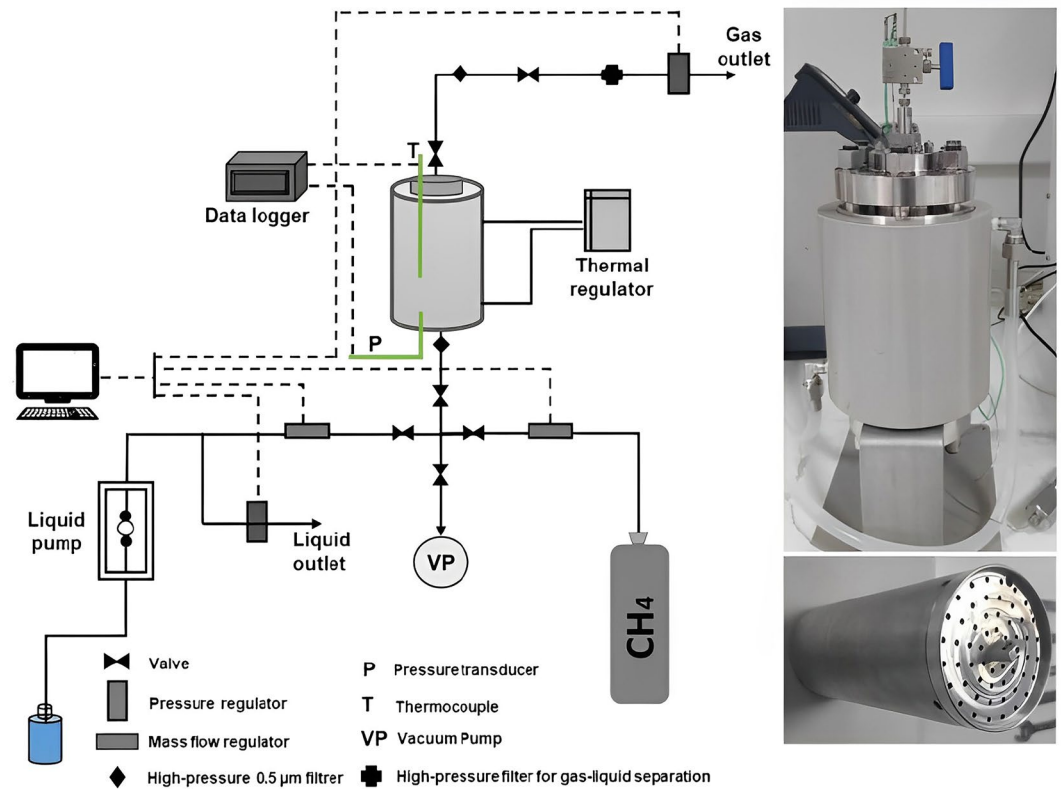


**Figure 1.** Scanning Electron Microscopy images showing aggregated platy particles of (a) ILL and (b) MMT. (c). Particle Size Distribution of Fontainebleau sand and the clay-rich matrices (ILL and MMT) determined by a Malvern Mastersizer 3,000 laser. (a)  $D_{10}$  and  $D_{50}$  ( $D_{mean}$ ) are grain sizes below which 10% and 50% of all particles are found, respectively.

latter was used when it was necessary to recover the hydrate-bearing core for visual inspection. The matrix was unconsolidated and therefore subjected to very low or practically no effective stress (self-weight).

Prior to any experiment, the apparatus was vacuumed by applying pressure of about  $10^{-6}$  MPa in order to remove any residual air pockets from the matrix. Water was injected at 293.15 K with constant flowrate (2.5 and  $0.5 \text{ g min}^{-1}$  for pure sand and clay-rich matrices, respectively) in the center of the upper flange of the cell for optimal distribution in the matrix. The system was then left at 293.15 K for 24 hr to ensure no further gravitational water movement in the matrix.

Prior to gas injection, cell temperature was set at 277 K before gas supply. Once thermal stability was reached (4 hr minimum), methane was injected through the bottom of the cell (Figure 2) at a constant flowrate of



**Figure 2.** Schematic diagram of the apparatus used to synthesize hydrate-bearing cores. The images to the left depict the high-pressure cell (top) and the aluminum cell (bottom).

57 mL min<sup>-1</sup>, until the system reached the gas cylinder pressure (~11 MPa). During methane injection, the temperature, as well as the upstream and downstream pressures of the cell, were recorded every 10 s. Hydrate formation was detected from a pressure drop (illustrated by a change in pressure slope when monitoring the evolution of pressure as a function of time) associated with a sudden increase in temperature due to the exothermic nature of hydrate formation (Figure 3). The gas injection was stopped when the pressure remained stable for at least 15 hr to ensure completion of hydrate formation.

### 2.2.3. Design the Experiments

Three series of experiments were performed:

1. The first series was dedicated to the investigation of the effect of clay content on hydrate formation in porous media. Experiments were carried out with partially saturated sand-clay mixtures with various clay content and at a constant volume of water ( $V_w$ ). The latter was calculated as follows:

$$V_w = S_w \times V_v \quad (1)$$

$$V_v = V_{\text{Cell}} - V_s - 2 \times V_{\text{ps}} \quad (2)$$

Where  $V_{\text{cell}}$  (mL) is the aluminum cell effective volume, the volume of voids  $V_v$  (mL),  $V_{\text{ps}}$  (mL) is the volume of the porous stone, and  $V_s$  (mL) is the sand volume computed from the sand grain density ( $\rho_s = 2.65 \text{ g/cm}^3$ ).

Equation 2 is only used to determine the volume of voids in sand. Since sand and clay-rich matrices do not have the same density and porosity, water saturation in the presence of clay was calculated from their estimated porosity ( $n$ ), using geometric soil parameters (Holtz et al., 2011):

$$e \times S_w = G_s \times \omega \quad (3)$$

**Table 2**  
Mineralogical Composition and Properties of the Different Clay Matrix/Sand Mixtures Used for the Experiments

|          | Mixtures    |             |                         | Mineralogy                |                            |          | Hydraulic conductivity                |  |
|----------|-------------|-------------|-------------------------|---------------------------|----------------------------|----------|---------------------------------------|--|
|          | Clay matrix | Sand FB (%) | Clay (%)                | Quartz (%)                | Calcite (%)                | Porosity | $K$ (m s <sup>-1</sup> ) <sup>a</sup> |  |
| Sand     | 0           | 100         | —                       | 100                       | —                          | 0.42     | $2 \times 10^{-4}$                    |  |
| ILL/Sand | 17.9        | 82.1        | 10.0                    | 87.3                      | 2.3                        | 0.47     | $4.80 \times 10^{-9}$                 |  |
|          | 35.7        | 64.3        | 20.0                    | 74.6                      | 4.6                        | 0.52     | $4.36 \times 10^{-10}$                |  |
| MMT/Sand | 71.4        | 28.6        | 40.0                    | 49.3                      | 9.3                        | 0.58     | $2.37 \times 10^{-10}$                |  |
|          | 100         | 0.0         | ~60                     | 29.0                      | 13.0                       | 0.63     | $2.23 \times 10^{-10}$                |  |
| MMT/Sand | 15.2        | 84.8        | 10.0                    | 87.6                      | —                          | 0.48     | $2.08 \times 10^{-8}$                 |  |
|          | 90.9        | 9.1         | ~60                     | 25.5                      | —                          | 0.74     | $1.08 \times 10^{-8}$                 |  |
|          | ILL (%)     | MMT (%)     | Illite (%) <sup>b</sup> | Smectite (%) <sup>b</sup> | Kaolinite (%) <sup>b</sup> |          |                                       |  |
| MMT/ILL  | 90.0        | 10.0        | 70.8                    | 10.9                      | 15.5                       | —        | 0.63                                  |  |
|          | 70.0        | 30.0        | 58.4                    | 26.7                      | 12.5                       | —        | 0.66                                  |  |
|          | 47.0        | 53.0        | 44.1                    | 44.9                      | 9.1                        | —        | 0.69                                  |  |

<sup>a</sup>For sand,  $K$  is obtained by Hazen's equation:  $K = CH \times D_{10}^2$  (Holtz et al., 2011). For clay matrix/sand mixtures, values were obtained by oedometer tests conducted in the laboratory according to the standard method D5856-15 of ASTM International (ASTM, 2007). <sup>b</sup>The proportions given here represent the resulting clay fraction mineralogy calculated from the composition of the original materials (Table 1).

$$e = \frac{n}{1 - n} \quad (4)$$

Porosity is derived from Equations 3 and 4:

$$n = \frac{\omega}{\omega + \frac{1}{G_s}} \text{ for } S_w = 100\% \quad (5)$$

Where  $e$  is the void ratio,  $G_s$  (2.66) is the specific gravity of soil solids and  $\omega$  (%) is the water content, obtained by the ratio between the pore water weight and the solid grains weight. A detailed description of this procedure is given in the Supporting Information S1. Calculated porosity values are reported in Table 2. These values are used to obtain the volume of voids, which then allows to calculate water saturation using Equation 1.

- The second set of experiments aimed to jointly examine the effect of water saturation and clay type on hydrate formation. They were performed only with the aforementioned clay matrices, at ~60% of clay (Table 2), but with different water saturation varying from 40% to 82% for ILL and from 35% to 100% for the MMT-rich matrix.
- The purpose of the third set was to study the kinetic behavior of hydrates in the presence of different mixtures of the two types of clay. Experiments were performed at 75% water saturation with three MMT/ILL mixtures with mass ratio of 10/90, 30/70, and 47/53 (Table 2), respectively.

After each experiment, the cell was quickly depressurized before recovering the hydrate core. The core was cut longitudinally immediately after recovery for hydrate morphology observation, and then stored in liquid nitrogen at 77 K. Selected cross sub-sections were made later to collect specific samples for micro-Raman spectroscopy analysis.

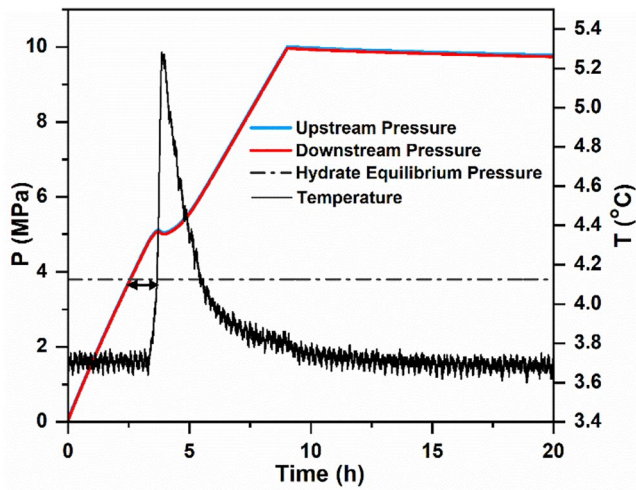
### 2.3. Recovery and Sampling of Natural Gas Hydrate From the Western Black Sea

A 6-m-long hydrate-bearing core, labeled GAS2-CS05, was recovered at 734 m water depth and ~9°C during the Ghass2 Cruise (2021) in the Romanian sector of the Western Black Sea. The core was cut into sections of 1 m in length and opened lengthwise to collect hydrate specimens with a sampling resolution of ~10–30 cm (Table 3). The samples were preserved in liquid nitrogen for future analysis by micro-Raman Spectroscopy. Sediment samples were also collected near the hydrates within the core for subsequent mineralogy analysis and particle size distribution (Table 3). Overall, the sediment is composed on average of ~60% clay, ~23% quartz, and ~8.5% calcite. However, clay mineralogy varies with depth, with the upper 2.4 m dominated by illite (~62%) and a smectite-rich lower part (~53%).

### 2.4. Micro-Raman Spectroscopy Analysis

#### 2.4.1. Apparatus and Experimental Procedure

Micro-Raman analyses were performed using a Labram HR Evolution spectrometer (Horiba, Jobin Yvon, France) coupled with a confocal microscope allowing to accurately investigate a specimen over a micrometric area. A 405 nm wavelength solid laser produces the excitation source and the beam is focused on the sample thanks to a ×50 long working distance objective. The confocal pinhole is set to 300 μm and the collected Raman



**Figure 3.** Example of typical pressure and temperature profiles during hydrate formation. (←→) Represents induction time. Upstream and downstream pressures are measured at the bottom and top of the core, respectively. Hydrate equilibrium pressure in bulk phase at 277 K is 3.8 (dashed black line) (Sloan & Koh, 2007).

respectively (Sum et al., 1997; Tulk et al., 2006). Since the number of large cages is three times that of SC in the SI hydrate structure, the relative cage occupancy of large to SC is expressed as follows:

$$\frac{\theta_{LC}}{\theta_{SC}} = \frac{A_{LC}}{3A_{SC}} \times \frac{\sigma_{SC}}{\sigma_{LC}} \quad (6)$$

Where  $\theta_{LC}$  and  $\theta_{SC}$  are the absolute cage occupancies in large and SC, respectively;  $A_{LC}$  and  $A_{SC}$  are the Raman peak integrated intensities (areas) of the large and SC, respectively. These areas are determined by peak deconvolution using the Gauss/Lorentz function after a suitable baseline correction with the software Labspec 5.  $\sigma$  represents the Raman scattering cross section for each cage and  $\sigma_{LC}/\sigma_{SC}$  is 0.977 for  $\text{CH}_4$  hydrate (Qin & Kuhs, 2013).

To calculate absolute cage occupancies, the relative cage occupancy obtained from micro-Raman measurements is coupled with the well-known Van der Waals and Platteeuw thermodynamic theory (Waals & Platteeuw, 1958), expressing the chemical potential of water in the SI hydrate:

$$\mu_w(h) - \mu_w(h_0) = -\frac{RT}{23} 3 \ln(1 - \theta_{LC}) + \ln(1 - \theta_{SC}) \quad (7)$$

Where  $\mu_w(h)$  is the chemical potential of water in a hypothetical empty lattice, and  $\mu_w(h_0)$  is the chemical potential of water in a reference state. In the equilibrium state:

$$\mu_w(h) - \mu_w(h_0) = \Delta\mu_w^0 \quad (8)$$

And  $\Delta\mu_w^0$  is the difference between the chemical potential of water in the empty gas-hydrate lattice and the stable-ice lattice, whose commonly used value is  $1297 \text{ J mol}^{-1}$  (Davidson et al., 1986). The Van der Waals and Platteeuw theory assumes that the free energy of the hydrate is independent of cavity occupation, that one hydrate cavity contains only one guest molecule, and that the interactions between guest molecules are negligible.

The absolute occupancies of the cages are obtained by combining Equations 6 and 7. Thus, the gas storage capacity (GSC in  $\text{m}^3$ ) or EF, which corresponds to the volume of methane that can be released by the dissociation of a unit volume of hydrate in standard conditions ( $0^\circ\text{C}$  and 1 atm), is calculated as defined in Mery and Sinayuc (Mery & Sinayuc, 2016):

**Table 3**  
Hydrates Samples Depth, Average Mineralogy, and Mean Particle Size of Their Associated Sediments

| Hydrates samples | Associated sediments |                                    |                            |
|------------------|----------------------|------------------------------------|----------------------------|
|                  | Global mineralogy    | Clay mineralogy                    | $D_{50}$ ( $\mu\text{m}$ ) |
| 0.39–2.40        | Clays (~65%)         | Illite (~62%)                      | ~4.5                       |
|                  | Quartz (~19%)        | Smectite (~11%)                    |                            |
|                  | Calcite (~7%)        | Kaolinite (~20%)<br>Chlorite (~7%) |                            |
| 2.50–5.80        | Clays (~57%)         | Illite (~28%)                      | ~27.6                      |
|                  | Quartz (~26.5%)      | Smectite (~53%)                    |                            |
|                  | Calcite (~5%)        | Kaolinite (~14%)                   |                            |
|                  |                      | Chlorite (~5%)                     |                            |



$$GSC = EF = \frac{M_{CH_4}}{M_{CH_4} + N \times M_{H_2O}} \times \frac{\rho_H \times V_H}{\rho_{CH_4}} \quad (9)$$

And

$$N = \frac{23}{3\theta_{LC} + \theta_{SC}} \quad (10)$$

$N$  is the hydration number of  $CH_4$  hydrate,  $M_{CH_4}$  and  $M_{H_2O}$  ( $g \text{ mol}^{-1}$ ) are the molar weights of  $CH_4$  and  $H_2O$ , respectively;  $V_H$  is the unit hydrate volume ( $1 \text{ m}^3$ );  $\rho_H$  and  $\rho_{CH_4}$  are the densities of  $CH_4$  hydrate ( $910 \text{ kg m}^{-3}$ )  $CH_4$  gas ( $0.717935 \text{ kg m}^{-3}$ ), respectively, in standard conditions.

### 3. Results and Discussion

#### 3.1. Hydrate Morphology and Structural Load Carrying Fraction

Both the hydrate-bearing cores synthesized from the series of laboratory experiments in the different matrices composed of pure sand, ILL/sand, MMT/sand, MMT/ILL mixtures with different clay contents (10%~60%) and the natural hydrate samples, were visually inspected in order to describe the hydrate morphology (Holland et al., 2008; Terzariol et al., 2020, 2021).

Figure 4 shows selected photographic images of the morphology of hydrates formed in different matrices. Hydrates in the sandy matrix are widely distributed along the cores. Cross sections (Figure 4a) show that hydrates are concentrated in the central part of the core. These areas of the cores exhibited a more pronounced resistance to splitting. Similar experimental observations of hydrates appearing as cement between sand grains were made by Ruffine (2015) and Klapproth et al. (2006). In clay/sand matrices with clay content of 10% (Figures 4b and 4c), morphology is similar to that observed in sand, but the distribution is more heterogeneous. In the 17.9% ILL/82.1% sand mixture, hydrates are more concentrated in the lower part of the core (Figure 4b), whereas they are more widely distributed along the core when the clay used is MMT (white parts, Figure 4c).

A different morphology is observed in the 35.7% ILL/64.4% sand mixtures (Figure 4d). The hydrates exhibit a grain-displacive behavior and are localized in the central and lower parts of the cores. They are massive and are arranged as veins with thickness varying from 1 to 3 mm. In the 71.4% ILL/28.6% sand matrix, hydrates also displace particles to form dispersed nodules (Figure 4e). They form a network of interconnected fine millimeter-thick veins (Figure 4e). This morphology, resulting from grain displacement, is increasingly pronounced in the more enriched clayey matrix and in the MMT/ILL mixtures, with the thickest hydrate veins observed in our experiments (Figures 4f and 4g). The photographic images of the Black Sea natural hydrates show that the morphologies observed on our clay-rich synthetic hydrates are consistent with natural hydrate-bearing sediments. Hydrates fill parallel fractures in the Black Sea clayey sediments (Figures 4g and 4i).

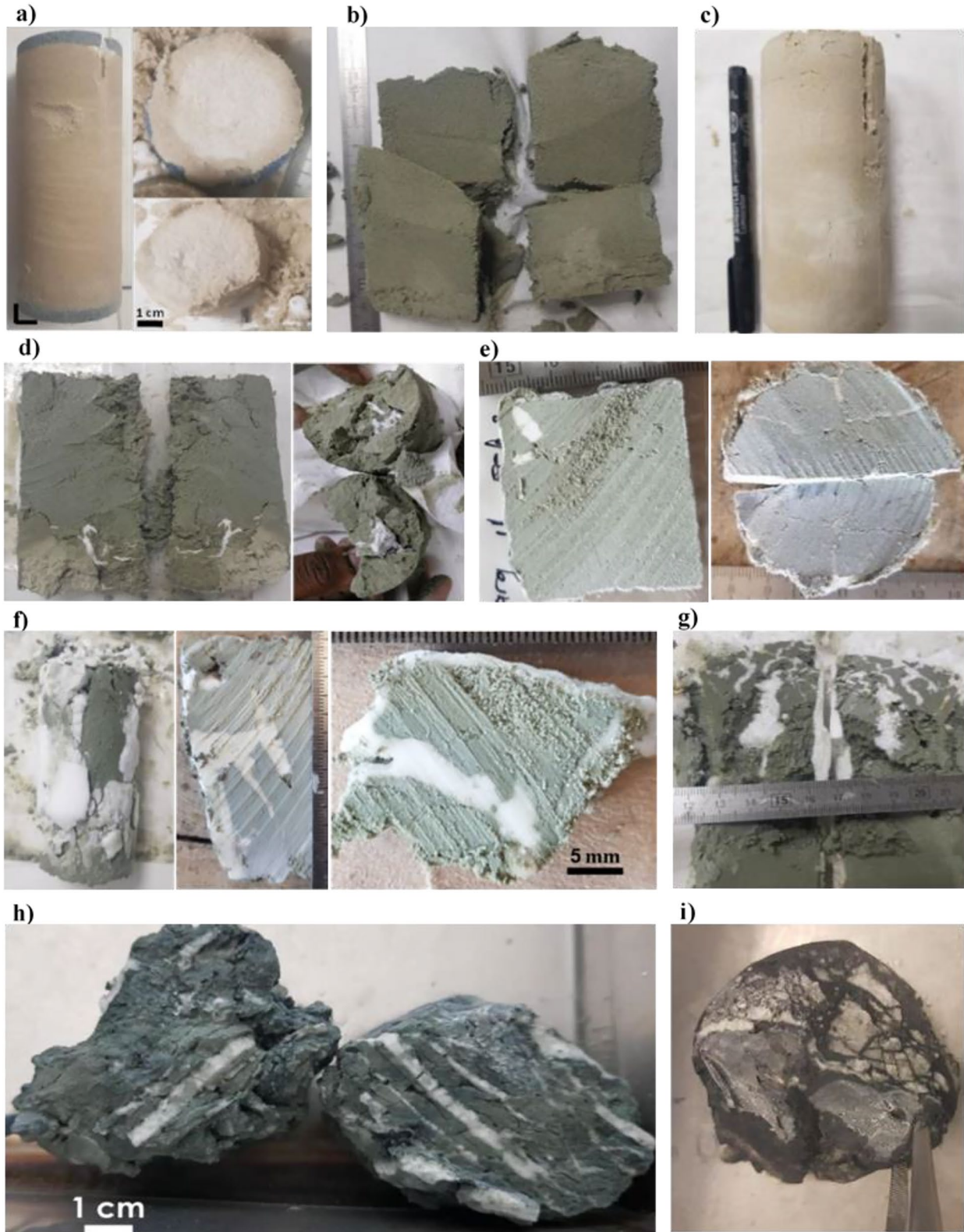
Sediment properties control hydrate morphology. A pore-invasive and grain-displacive pore habit can be anticipated from the ratio of pore-throat capillarity  $\Delta u$  and effective stress  $\sigma'$  (Dai et al., 2012; Terzariol et al., 2020, 2021). Capillary pressure is estimated by:

$$\Delta u = \frac{4T_s}{r} \quad (11)$$

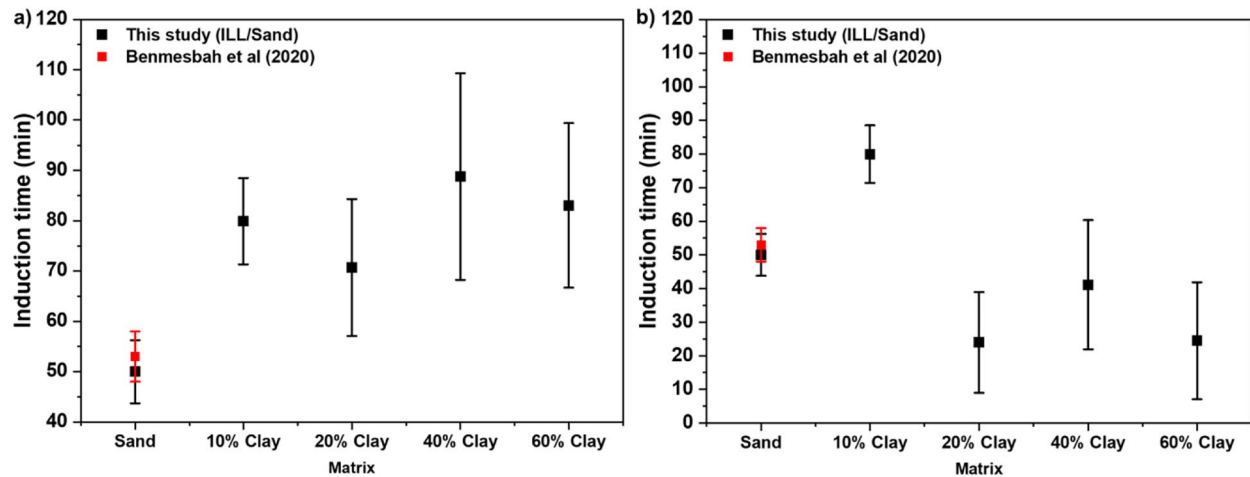
Where  $T_s$  ( $\sim 0.039 \text{ N/m}$ ) is methane hydrate/water interfacial tension;  $r$  is the mean pore size and can be approximated for sand by  $(\sqrt{2} - 1)D_{10}$ . In the case of clayey matrices (ILL and MMT), this pore size can be estimated from specific surface ( $S_s$ ), mineral density ( $\rho_m$ ), and void ( $e$ ) (equations and explanation in Terzariol et al. (2021)):

$$d_p = \frac{2e}{S_s \rho_m} \quad (12)$$

A first-degree approximation of the capillarity in our test sediments, shows that  $\Delta u = 3 \text{ kPa}$  in sand FB, while in ILL and MMT, this value reaches 9 and 12 MPa respectively. Thus, it can be expected that sand FB might form pore-filling hydrate even in the absence of applied vertical loads. On the other hand, ILL and MMT will favor grain-displacive hydrate formation as observed in our experiments.



**Figure 4.** Photographic images showing the morphology of synthetic hydrate-bearing sediment (a–g) and natural specimens (h, j): (a) Disseminated hydrates in sand. (b, c) Disseminated hydrates in sand. (b, c) Disseminated hydrates in clay matrix: (b) 17.9% ILL/82.1% sable, (c) 15.2% MMT/84.8% sand. (d) Hydrate veins and nodules in clay matrix (35.7% ILL/64.3% sand). (e) Nodules and vein network in clay (71.4% ILL/28.6% sand). (f) Thick veins and hydrates surrounding the matrix (100% ILL). (g) Network of hydrate veins of variable thickness in a MMT/ILL matrix. (h) Natural sample collected at 5.35 mbsf showing hydrate-filling fractures. (i) Massive hydrate sample collected at 3.2 mbsf in the Romanian sector of the Western Black Sea.



**Figure 5.** Average values of induction time for hydrate formation in matrices with different clay content and at constant water volume (not constant water saturation). (a) Measured induction time from bulk hydrate equilibrium pressure. (b) Corrected induction from capillarity effect. Bars represent the standard deviation. Experiments of Benmesbah et al. (2020) were carried out in the same conditions as the present study (sandy matrix, and same water saturation and methane injection flowrate).

In the case of a fines (i.e., particle size  $\leq 74 \mu\text{m}$ )/coarse mix sample, we needed to determine which sediment fraction carries the mechanical load. The ILL and MMT matrices are both fines since they have  $D_{99.85}$  and  $D_{97}$  smaller than  $74 \mu\text{m}$  (Figure 1c). In other words, the proportion of either the ILL or MMT matrix in a mixture corresponds to the fines content. The Revised Soil Classification System (RSCS) (Park & Santamarina, 2017) allows us to estimate the fines fraction thresholds (FF) from compacity (for sands) and effective stress dependent void ratios (for fines). An ILL/sand FB and MMT/sand FB mixtures show a mechanical transition zone of  $\text{FF} = 15\% - 29\%$  and  $8\% - 20\%$ , respectively, displaying good agreement with the fine content at which the change in morphology occurs in our synthesized samples. Regarding fluid flow, these thresholds are  $\text{FF} = 8\%$  and  $2.5\%$  for ILL/sand and MMT/sand, respectively.

### 3.2. Kinetics of Hydrate Formation

#### 3.2.1. Influence of Clay Content on Induction Time

The induction time is defined as being the elapsed time between the equilibrium pressure of methane hydrate at experiment temperature and the pressure drop associated with the onset of hydrate formation (Figure 3). Figure 5a summarize the measured induction times for the first series of experiments performed at constant injected water volume in sand and clay/sand mixtures (experiments 1\_1 to 5\_5 in Table 4). The average induction time for hydrate formation within the sand is  $50 \pm 6$  min, which is close to the value of  $53 \pm 5$  min obtained by Benmesbah et al. (2020) in a previous study using the same apparatus and operating conditions.

As shown in Figure 5a, once clay was added to sand, even in a small amount, the induction time significantly shifted to higher values. The addition of 10% of clay to a sandy matrix increases the induction time from 50 min (without any clay) to  $80 \pm 8$  min, for example, an increase of  $\sim 60\%$ . Our experiments indicate that the variation of induction time is relatively small when varying the amount of clay in the ILL/sand matrix. Despite data scattering (Table 4), likely related to the stochastic nature of hydrate nucleation (Bagherzadeh et al., 2011; Linga et al., 2009; Mekala et al., 2014), the values of induction time remains relatively constant for matrices with 20%, 40%, and  $\sim 60\%$  of clay, with values of  $71 \pm 13$ ,  $89 \pm 20$ , and  $83 \pm 16$  min, respectively. Therefore, our results suggest that clay acts as a kinetic inhibitor for this VW. This is in agreement with previous conclusions of Zhang et al. (2017) who also show that the addition of 10 wt% of bentonite clay to sand, at 55% of water saturation, significantly increases the induction time, which then remains almost unchanged even for clay content up to 30 wt%.

It is well known that gas flow is a key factor controlling hydrate formation in sediment (Xu & Ruppel, 1999; You et al., 2019). As mentioned by Park and Santamarina (Park & Santamarina, 2017), the presence of fines, at even a low content, is enough to drastically affect fluid flow in sand/clay mixtures. In the present work, we have estimated this fines fraction  $\text{FF} = 8\%$  for the ILL/sand mixtures. Thus, the addition of ILL ( $D_{99.7} < 74 \mu\text{m}$ ) fills the voids between sand particles, leading to a reduction in matrix permeability, and therefore strongly hindering gas migration to the hydrate formation front. This is partially supported by hydraulic conductivity ( $K$ ),

**Table 4**  
*Summary of Measured Kinetic Parameters From Different Series of Hydrate Formation Experiments*

| Mixtures | Experiments | Clay content<br>(ratio clay<br>matrix/sand) | Water<br>saturation (%) | Measured<br>induction<br>time (min) | Pressure at the<br>onset of hydrate<br>formation (MPa) | Amplitude of<br>the thermal<br>anomalies $\Delta T$<br>(°C) | Load<br>carrying<br>fraction | Void ratio<br>for fines<br>at low<br>effective<br>stress | Pore size diameter<br>( $\mu\text{m}$ ) | Corrected<br>induction<br>time from<br>capillarity |
|----------|-------------|---|-------------------------|-------------------------------------|--|---|------------------------------|--|---|--|
| Sand     | 1_1         | 0% clay (0/100)                             | 56.0                    | 55.8                                | 5.10   | 1.10  | Sand                         | N/A  | 60.8                                    | 55.8   |
|          | 1_2         |   | 56.7                    | 50.5                                | 4.78   | 0.81  | Sand                         | N/A  | 60.8                                    | 50.5   |
|          | 1_3         |   | 56.0                    | 43.3                                | 4.71   | 0.73  | Sand                         | N/A  | 60.8                                    | 43.3   |
| ILL/Sand | 2_1         | 10% clay<br>(17.9/82.1)                     | 52.0                    | 83.3                                | 5.22   | 1.66  | Sand                         | N/A  | 60.8                                    | 83.3   |
|          | 2_2         |   | 52.4                    | 70.2                                | 5.12   | 1.58  | Sand                         | N/A  | 60.8                                    | 70.2   |
|          | 2_3         |   | 52.4                    | 86.3                                | 5.71   | 1.66  | Sand                         | N/A  | 60.8                                    | 86.3   |
|          | 3_1         | 20% clay<br>(35.7/63.4)                     | 49.0                    | 81.3                                | 5.50   | 0.61  | ILL                          | 1.52   | 0.017                                   | 34.0   |
|          | 3_2         |   | 49.0                    | 75.3                                | 5.34   | 1.69  | ILL                          | 1.52   | 0.017                                   | 29.3   |
|          | 3_3         |   | 49.0                    | 55.3                                | 4.82   | 1.42  | ILL                          | 1.52   | 0.017                                   | 6.0  |
|          | 4_1         | 40% clay<br>(71.4/28.6)                     | 45.0                    | 72.5                                | 4.47   | 0.44  | ILL                          | 1.52   | 0.017                                   | —  |
|          | 4_2         |   | 45.0                    | 82.0                                | 5.22   | 1.16  | ILL                          | 1.52   | 0.017                                   | 27.5   |
|          | 4_3         |   | 45.0                    | 111.8                               | 5.63   | 1.27  | ILL                          | 1.52   | 0.017                                   | 54.7   |
|          | 5_1         | 60% clay<br>(100/0)                         | 41.3                    | 99.7                                | 5.45   | 0.63  | ILL                          | 1.52   | 0.017                                   | 42.7   |
|          | 5_2         |   | 41.2                    | 95.2                                | 5.38   | 1.36  | ILL                          | 1.52   | 0.017                                   | 39.0   |
|          | 5_3         |   | 41.0                    | 65.0                                | 4.87   | 1.25  | ILL                          | 1.52   | 0.017                                   | 7.2  |
|          | 5_4         |   | 41.0                    | 66.3                                | 4.81   | 1.53  | ILL                          | 1.52   | 0.017                                   | 5.5  |
|          | 5_5         |   | 41.0                    | 88.4                                | 5.20   | 1.13  | ILL                          | 1.52   | 0.017                                   | 28.2   |
|          | 6_1         | 60% clay<br>(100/0)                         | 55.0                    | 56.7                                | 4.84   | 0.54  | ILL                          | 1.52   | 0.017                                   | 5.3  |
|          | 6_2         |   | 55.0                    | 53.0                                | 4.82   | 0.48  | ILL                          | 1.52   | 0.017                                   | 3.7  |
|          | 6_3         |   | 55.0                    | 67.2                                | 4.98   | 0.57  | ILL                          | 1.52   | 0.017                                   | 13.0   |
|          | 7_1         | 60% clay<br>(100/0)                         | 62.0                    | 44.0                                | 4.57   | 0.40  | ILL                          | 1.52   | 0.017                                   | —  |
|          | 7_2         |   | 62.0                    | 60.0                                | 4.91   | 0.91  | ILL                          | 1.52   | 0.017                                   | 9.7  |
|          | 7_3         |   | 62.0                    | 39.0                                | 4.51   | 0.46  | ILL                          | 1.52   | 0.017                                   | —  |
|          | 8_1         | 60% clay<br>(100/0)                         | 82.0                    | 59.7                                | 4.98   | 0.31  | ILL                          | 1.52   | 0.017                                   | 11.7   |
|          | 8_2         |   | 82.0                    | 52.2                                | 4.91   | 0.30  | ILL                          | 1.52   | 0.017                                   | 7.7  |
|          | 8_3         |   | 82.0                    | 52.2                                | 4.92   | 0.67  | ILL                          | 1.52   | 0.017                                   | 9.0  |

Table 4  
Continued

| Mixtures | Experiments | Clay content (ratio clay matrix/sand) | Water saturation (%) | Measured induction time (min) | Pressure at the onset of hydrate formation (MPa) | Amplitude of the thermal anomalies $\Delta T$ (°C) | Load carrying fraction | Void ratio for fines at low effective stress | Pore size diameter ( $\mu\text{m}$ ) | Corrected induction time from capillarity |
|----------|-------------|---------------------------------------|----------------------|-------------------------------|--|--|------------------------|--|--------------------------------------|---|
| MMT/Sand | 9           | 10% clay (15.2/84.8)                  | 52.0                 | 90.0                          | 5.87   | 4.09   | Sand                   | N/A  | 60.8                                 | 90.0                                      |
|          | 10          | 60% clay (91/9)                       | 36.0                 | 125.2                         | 5.93   | 3.01   | MMT                    | 4.04   | 0.013                                | 51.2                                      |
|          | 11_1        | 60% clay (91/9)                       | 55.0                 | 132.5                         | 6.22   | 4.11   | MMT                    | 4.04   | 0.013                                | 65.3                                      |
|          | 11_2        | 60% clay (91/9)                       | 55.0                 | 100.0                         | 5.67   | 3.75   | MMT                    | 4.04   | 0.013                                | 34.0                                      |
|          | 12_1        | 60% clay (91/9)                       | 75.5                 | 60.5                          | 5.13   | 2.40   | MMT                    | 4.04   | 0.013                                | 5.5                                       |
|          | 12_2        | 60% clay (91/9)                       | 75.0                 | 83.7                          | 5.68   | 2.87   | MMT                    | 4.04   | 0.013                                | 29.2                                      |
| MMT/ILL  | 13          | 60% clay (91/9)                       | 100                  | 38.7                          | 4.95   | 0.35   | MMT                    | 4.04   | 0.013                                | –   |
|          | 14          | ~60% (10 MMT/90 ILL)                  | 75.0                 | 89.2                          | 5.83   | 0.20   | ILL                    | 1.52   | 0.017                                | 52.0                                      |
|          | 15_1        | ~60% clay (30 MMT/70 ILL)             | 75.0                 | 85.1                          | 5.67   | 0.18   | ILL                    | 1.52   | 0.017                                | 40.8                                      |
|          | 15_2        | ~60% clay (30 MMT/70 ILL)             | 75.0                 | 92.7                          | 5.76   | 0.16   | ILL                    | 1.52   | 0.017                                | 50.0                                      |
|          | 16_1        | ~60% clay (53 MMT/47 ILL)             | 75.0                 | 126.0                         | 6.83   | 0.22   | MMT                    | 4.04   | 0.013                                | 69.2                                      |
|          | 16_2        | ~60% clay (53 MMT/47 ILL)             | 75.0                 | 135.0                         | 6.73   | 0.35   | MMT                    | 4.04   | 0.013                                | 79.7                                      |

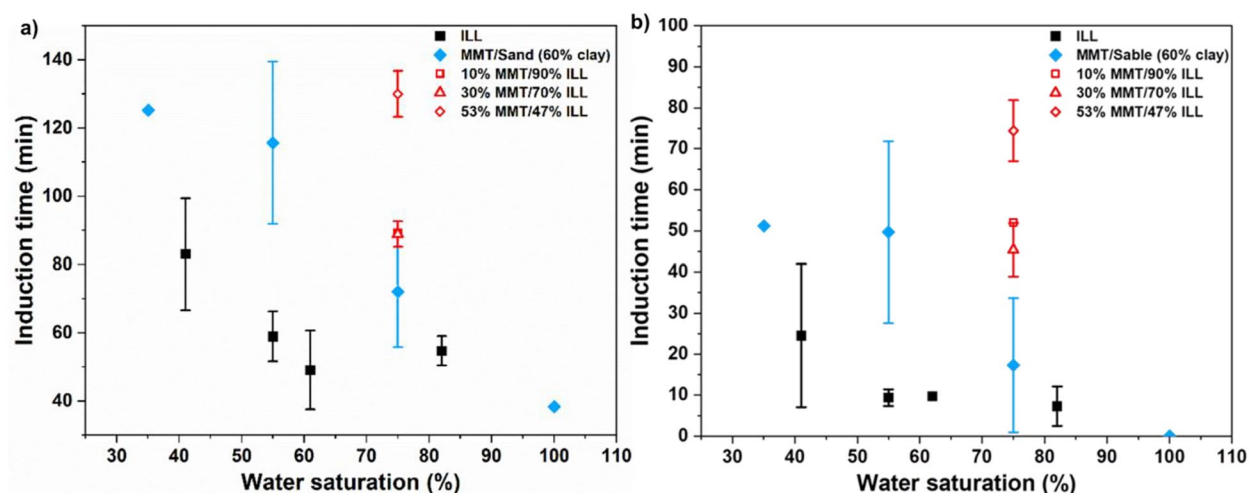
proportional to the intrinsic permeability of the matrices used here.  $K$  for sand is  $2 \times 10^{-4} \text{ m s}^{-1}$ , whereas the values for the mixtures ILL/sand with 10%, 20%, 40%, and ~60% clay are  $4.80 \times 10^{-9}$ ,  $4.36 \times 10^{-10}$ ,  $2.37 \times 10^{-10}$ , and  $2.23 \times 10^{-10} \text{ m s}^{-1}$ , respectively. A difference of five orders of magnitude is noted between the hydraulic conductivity of the sandy matrix and the mixture with 10% clay (17.9% ILL). Considering our gas injection procedure, this large difference could be one plausible explanation for the longer time observed for hydrate formation in 10% clay matrix. Such behavior may explain the fact that the induction time remains almost constant for the matrices ILL/sand, irrespective of the clay content, since the values of their permeabilities are very close.

To take account for capillarity effect inherently related to small pore size in clayey sediments, we corrected the induction time discussed above and measured based on the bulk hydrate equilibrium pressure. This correction requires to determine the methane hydrate equilibrium pressures associated with the different mixtures porous media (sand, ILL, MMT, and their different mixtures) at the experimental temperature (~277 K). In fact, the conclusions of several studies have shown the thermodynamic inhibiting effect of small pore size, which shifts the hydrate phase boundary of hydrates toward lower temperatures and/or higher pressures (Handa & Stupin, 1992; Østergaard et al., 2002; Uchida et al., 2002). The temperature depression ( $\Delta T_{\text{pore}}$ ) related to this effect is calculated using Equation 5 in Lee et al. (2008) and first introduced by Clennell et al. (1999):

$$\Delta T_{\text{pore}} = \frac{2\gamma_{\text{Hw}}T_{\text{bulk}}\cos\theta}{\rho_{\text{H}}\Delta H r} \quad (13)$$

Where  $\gamma_{\text{Hw}}$  (27 mJ m<sup>-2</sup>) (Clennell et al., 1999) is the hydrate-water interfacial tension,  $T_{\text{bulk}}$  (K) is the methane hydrate equilibrium temperature in bulk phase,  $\theta$  the contact angle between the pore wall and hydrate,  $\rho_{\text{H}}$  (kg m<sup>-3</sup>) is the hydrate mass density,  $\Delta H$  (kJ kg<sup>-1</sup>) the heat of hydrate dissociation, and  $r$  is the pore size of the sediment. Due to the lack of pore size distribution, this latter parameter were estimated in this study for each mixture from the load fraction carrying (RSCS) as suggested in Terzariol et al. (2020). As explained in Section 3.1, the operation consists in defining which fraction of a clay matrix/sand matrix mixture controls its mechanical behavior. The results of the load carrying fraction reported in Table 4, were obtained from the fines fraction thresholds previously calculated: 29% at low effective stress for ILL/sand and MMT/sand, respectively. As a result, the mixture 17.9 ILL/82.1 sand (with 10% clay) for example, is sand-controlled while the others are ILL-controlled. The mean pore diameters of these load carrying fractions sand, ILL and MMT are 60.8, 0.017, and 0.013  $\mu\text{m}$ , respectively.

The application of Equation 13 to these pore sizes allowed to determine the hydrate equilibrium pressures at the experimental temperature (~277K), whose values are ~38 bar (similar to the bulk phase) for sand, ~4.8 for ILL and ~5.1 MPa for MMT. Based on the changed equilibrium conditions, the corrected induction times in mixtures affected by capillarity were found to be lower, than those previously obtained by considering a bulk equilibrium pressure of methane hydrate. As seen in Table 4 and Figure 5b, the trends of the evolution of the induction time in the ILL/sand are modified. Values remained unchanged for sand and 10% clay matrix where hydrate formation is not subjected to the capillarity pressure because of the large pore size (60.8  $\mu\text{m}$ ). Thus, it can still be observed that the induction time increases with the addition of 10% clay to sand. However, it decreases when the clay



**Figure 6.** Induction time as a function of the initial water saturation in illite and montmorillonite-rich clay. (a) Measured induction time from bulk hydrate equilibrium pressure. (b) Corrected induction time from capillarity effect. There is no addition of sand to the MMT/ILL matrices. Note that the experiments at 41% water saturation for ILL are also represented in Figure 5 at 60% clay for a constant amount of water.

content reaches 20% (i.e., when the load-carrying fraction is the clay matrix ILL) and remains relatively constant with further increase of the clay content. Indeed, applying the capillarity correction shifts the equilibrium pressure to higher values (from  $\sim 3.8$  in bulk phase to 4.8 MPa). The latter becomes located closer to the hydrate formation incipient pressure (e.g., the pressure drop) that occurs well after reaching the equilibrium pressure of methane hydrate within the cell. Thus, mechanically, the induction time is shortened. However, this result with the increase in clay content is unexpected since capillarity should inhibit hydrate formation. Nevertheless, this trend could be explained by the small size of clay particles and their high specific surface area compared to sand. As emphasized in previous studies, this specific surface area of small grains provides a greater number of hydrate nucleation sites and thus reduce the induction time (Chong et al., 2016; Heeschen et al., 2016; Mekala et al., 2014).

These results are not in agreement with those of Zeng et al. (2022) and Zhang et al. (2017) who rather observed an increase in induction time in a sand-clay mixture with increasing clay content. However, none of these studies take into account the effect of capillary pressure on hydrate equilibrium conditions in the determination of induction time. All these elements discussed highlight the complexity of hydrate formation mechanisms in the presence of clay, including gas transport, the effect of grain size and capillarity driven by small pores.

### 3.2.2. Influence of the Initial Water Saturation and Nature of Clay on Hydrate Formation Kinetics in Clay-Rich Matrices

This section is dedicated to the results obtained after the series of experiments (2) and (3) in the experimental section (experiments 5\_1 to 8\_3 and 10 to 16\_2 in Table 4). Figures 6a and 6b show respectively the measured and corrected (from capillarity) induction time of hydrate formation in clay-rich matrices (ILL and MMT) at different initial water saturations. It can be observed that the trends of the recalculated induction times are broadly the same as those measured according to bulk hydrate equilibrium conditions. However, previous work showed that capillary-induced depressions in equilibrium temperatures ( $\Delta T_{\text{pore}}$ ) is reduced with increasing water saturation (Uchida et al., 2004; Vavra et al., 1992). Therefore, it becomes less relevant to apply the correction only based on the theoretical pore diameter explained in Section 3.2.1. From our data set, it seems difficult to take quantitatively into account the variation of the capillary pressure with the water saturation. The following discussion, regarding the evolution of the induction as function of water saturation in ILL and MMT, is based on the initial measured values (Figure 6a), although it remains valid for Figure 6b.

The experiments performed with the ILL matrix show that induction time decreases significantly from  $83 \pm 16$  to  $49 \pm 11$  min when water saturation increases from 40% to  $\sim 62\%$  and then remains relatively steady with a further increase in water saturation; whereas a continuous decrease in induction is observed for experiments performed in the MMT-rich matrix (Figure 6b). Induction time is reduced by a factor of 3.2 when water saturation increases

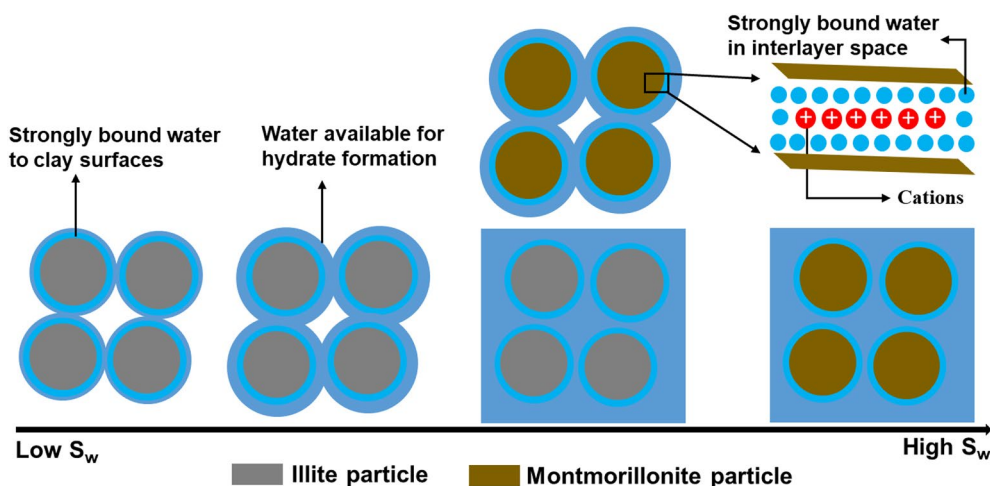


Figure 7. Schematic representation of water distribution in illite and montmorillonite as a function of water saturation.

from 35% to 100%. It is well known that clays have a high water adsorption capacity owing to their high specific surface area (Kumari & Mohan, 2021; Martin, 1960). In addition, clay minerals have a strong adsorption regime at low water content (Hatch et al., 2012; Schuttlefield et al., 2007), which consequently reduces the amount of water available for gas/water contact at low water saturation. Additionally, the inherent capillarity pressure associated to the small pores of ILL (0.017) and MMT (0.013  $\mu\text{m}$ ) is more significant at low water saturation and thus affecting the hydrate formation behavior.

Furthermore, the trends observed in Figure 6 indicate that hydrate formation occurs faster in ILL than in MMT-rich matrix for low to medium water saturation levels ( $S_w \leq 55\%$ ). Thus, the average induction time is almost twice higher for hydrate formation in montmorillonite than in illite for a saturation of 55%. This could be attributed to the different capillarity pressures and different water absorption capacities and mechanisms related to the different properties and mineral structures of these two types of clay. First, in agreement with its larger specific surface (240.8  $\text{m}^2/\text{g}$ , Table 1), MMT has a higher adsorption capacity than ILL (66.4  $\text{m}^2/\text{g}$ ). In addition, with its structural ability to swell, water uptake in montmorillonite is done by adsorption not only on the external surfaces of the clay particles but also and especially in interlayer spaces with hydration of exchangeable cations (Cases et al., 1992), whereas adsorption in illite only occurs on external surfaces (Figure 7) (Dettmann, 1958). These additional sites for bound-water in montmorillonite strongly affect water availability and could therefore delay hydrate formation mainly at low to medium water saturation. As water content increases, water fills the pores of the matrix (Cases et al., 1992; Hatch et al., 2012; H. Wang et al., 2020) and capillarity decreases (Vavra et al., 1992), making free water directly available for gas/water contact to promote hydrate formation. As shown in Figure 7, this pore filling process occurs at lower water saturation for illite with respect to montmorillonite (Hatch et al., 2012; Sun et al., 2021) because of the strongly bound water in its interlayer spaces; and this could explain the higher induction times in MMT-rich matrix than in ILL at 55% water saturation, the constant induction times in ILL from 55% water saturation and the shorter induction times obtained at higher water saturations in MMT-rich matrix.

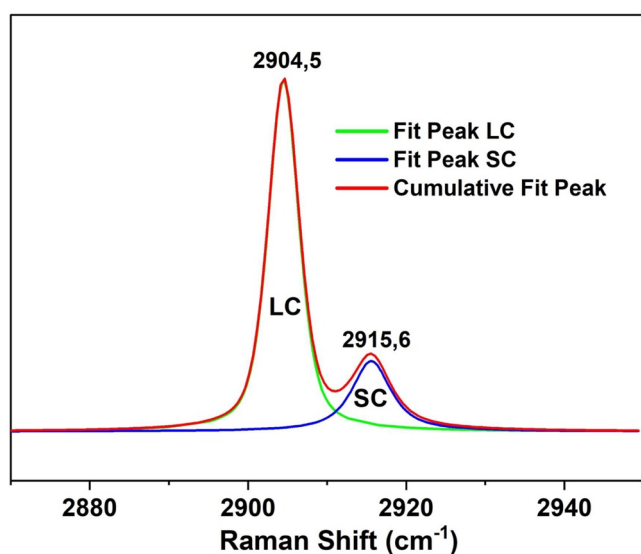


Figure 8. Typical Raman spectra of methane hydrate from 2,860 to 2,950  $\text{cm}^{-1}$  showing the signatures of  $\text{CH}_4$  trapped in LC (2,904) and small cages (2,915  $\text{cm}^{-1}$ ) and their deconvolution thanks to the Gauss-Lorentz (Voigt) function of the Labspec 5 software.

These observations are also corroborated by pressure variation (Table 4) at the onset of hydrate formation from one clay to another with the increase in water saturation. The pressure required for hydrate formation in ILL at 55%  $S_w$  ranges from 4.81 to 4.98 MPa (Experiments 6\_1 to 6\_3) while it is in the range of 5.67–6.22 MPa (Experiments 11\_1 and 11\_2) in the MMT-rich matrix. These higher formation pressures in the MMT-rich matrix are not

**Table 5**  
*Relative Cage Occupancies, Absolute Cage Occupancies, Hydration Numbers and Gas Storage Capacities Determined for Each Matrix*

|          | Matrix          | $\theta_{LC}/\theta_{SC}$ | $\theta_{LC}$ (%) | $\theta_{SC}$ (%) | $N$         | GSC or EF (m <sup>3</sup> ) |
|----------|-----------------|---------------------------|-------------------|-------------------|-------------|-----------------------------|
|          | Bulk hydrate    | 1.10 ± 0.04               | 97.1 ± 0.3        | 88.5 ± 3.2        | 6.06 ± 0.04 | 163.9 ± 1.0                 |
| Sand     | Sand            | 1.14 ± 0.06               | 97.4 ± 0.3        | 85.2 ± 4.0        | 6.09 ± 0.05 | 162.9 ± 1.4                 |
| ILL/Sand | 20% Clay        | 1.23 ± 0.07               | 97.4 ± 0.3        | 79.8 ± 4.3        | 6.17 ± 0.06 | 159.9 ± 1.5                 |
|          | 40% Clay        | 1.24 ± 0.08               | 97.8 ± 0.3        | 78.6 ± 4.9        | 6.18 ± 0.07 | 159.6 ± 1.6                 |
|          | 60% Clay        | 1.29 ± 0.07               | 97.8 ± 0.2        | 76.4 ± 4.7        | 6.22 ± 0.06 | 158.7 ± 1.5                 |
| MMT/Sand | 10% Clay        | 1.14 ± 0.04               | 97.8 ± 0.3        | 85.2 ± 3.4        | 6.09 ± 0.03 | 162.9 ± 1.0                 |
|          | 60% Clay        | 1.18 ± 0.04               | 97.6 ± 0.1        | 81.5 ± 2.6        | 6.13 ± 0.03 | 160.8 ± 0.9                 |
| MMT/ILL  | 10% MMT/90% ILL | 1.26 ± 0.05               | 97.7 ± 0.2        | 77.9 ± 3.7        | 6.19 ± 0.05 | 159.4 ± 1.2                 |
|          | 30% MMT/70% ILL | 1.22 ± 0.05               | 97.7 ± 0.2        | 80.1 ± 3.5        | 6.16 ± 0.05 | 160.3 ± 1.1                 |
|          | 53% MMT/47% ILL | 1.19 ± 0.06               | 97.6 ± 0.3        | 81.7 ± 3.9        | 6.13 ± 0.05 | 160.8 ± 1.4                 |

only due to capillarity in small pores but also to the low activity of strongly bound-water as mentioned in previous studies (Clennell et al., 1999; Sun et al., 2021). It is therefore logical that higher pressure levels for hydrate formation onset in MMT matrix at  $S_w \leq 55\%$  lead to longer induction times. Similarly, the hydrate formation pressure in the MMT-rich matrix decreases when increasing water saturation from 35% to 100%, resulting in the decrease in induction time. These results are in perfect agreement with the studies of Uchida et al. (2004) and Yakushev (2019), who show that the inhibiting thermodynamic effect (i.e., higher pressure or lower temperature required for hydrate formation) of bentonite clay, associated with the low activity of the strongly adsorbed water and capillarity, vanishes when water saturation reaches 90%. It is important to point out that the MMT used in the present study underwent chemical treatment leading to the leaching of some of the interlayer cations, and its clay fraction is not exclusively composed of smectite (Table 1). This results in a decrease in the swelling capacity compared to an untreated montmorillonite for which the specific surface area is greater and can reach up to 800 m<sup>2</sup>/g (Newman, 1983). Thus, we can reasonably speculate that the difference found in the induction time between the two clay used here, and also as a function of water saturation, would have been much more significant.

Previous studies carried out in pure sand did not highlight a clear relationship between induction time and initial water saturation (Benmesbah et al., 2020). Interestingly, induction times from 55% to 82% and at 100% water saturation in ILL and MMT-rich matrix, respectively, are in the same range or lower than the times obtained in pure sand at 55% water saturation. This is related to the fact that the sandy matrix is directly under a pore filling regime at this level of water saturation. However, the induction time for the MMT-rich matrix at  $S_w \leq 55\%$  (Figure 6) is 2.5 higher than for pure sand (Figures 5 and 6). These different kinetic behaviors of hydrate formation in sand and clay (in the present study) at different water saturations are clearly linked to the enhanced water/sediment interactions in clays (especially water adsorption on clay particle surfaces and strong binding between water and interlayer cations).

From the results of this set of experiments, we chose a  $S_w$  of 75% as an optimum value to synthesize hydrates in different MMT/ILL mixtures (Table 4, Figure 6). Unexpectedly, the values are higher than those obtained in the MMT-rich matrix at 75%  $S_w$ . On the other hand, it is worth highlighting that hydrate formation is delayed when the montmorillonite proportion in the matrix is increased from 30% MMT (26.7% smectite) to 53% MMT (44.9% smectite), with the average induction time increasing from 89 to 130.6 min and the average pressure at the hydrate formation onset increasing from 5.71 to 6.83 MPa.

### 3.3. Effect of Clay on Hydrate Cage Occupancy and Storage Capacity

From the large number of micro-Raman analyses performed, the processing of spectral signatures of CH<sub>4</sub> hydrates (Figure 8) allowed us to describe the variation of the relative cage occupancy, the absolute occupancies of both large and SC, and thus to infer the GSC as a function of the mineralogy of the matrix (Table 5, Figure 9). Since hydrates were formed by pore filling in sand and as veins/lenses in clay-rich matrices (Figure 4),



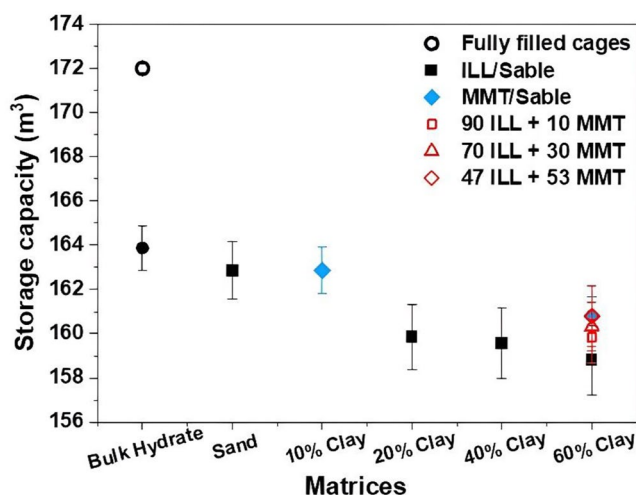


Figure 9. Gas storage capacity of hydrate as a function of matrix mineralogy.

measurements were carried out on sand + hydrate powders and from vein sampling for hydrate-bearing sandy matrix and hydrate-bearing clayey matrices, respectively.

The average relative occupancy obtained for bulk hydrates in this study is  $1.10 \pm 0.04$ , which corresponds to an absolute occupancy of  $97.1\% \pm 0.3\%$  of large cages and  $88.5\% \pm 3.2\%$  of SC. These values are in good agreement with those obtained for bulk methane hydrate ( $\theta_{LC}/\theta_{SC}$  ranging from 1.05 to 1.12) in previous studies (Qin & Kuhs, 2013; Sum et al., 1997).

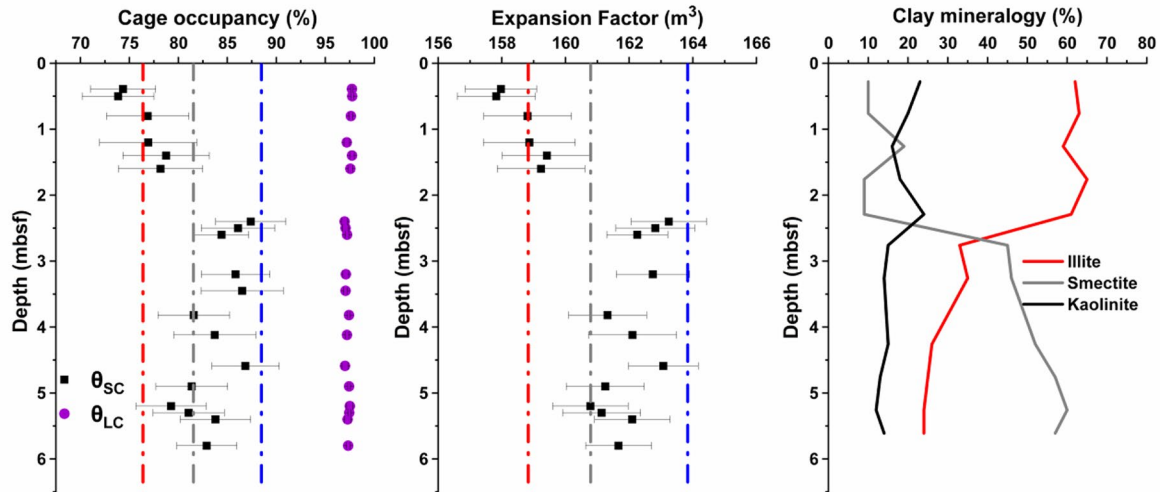
Table 5 shows that the large cages are almost fully occupied with  $\theta_{LC} \sim 98\%$  regardless of the mineralogy of the matrix used. However, the SC are partially filled with occupancy fractions decreasing (e.g., increase in the relative cage occupancy) with increasing clay content especially for the ILL/sand mixture. The absolute occupancy of the SC of the hydrate formed in sand ( $\theta_{SC} = 85.2\% \pm 4.0\%$ ) is relatively close to that of the bulk hydrate, while the average value is  $76.4\% \pm 4.7\%$  for the hydrate formed in the ILL/sand matrix (at  $\sim 60\%$  clay), for example, a deviation of  $\sim 12\%$  (maximum of  $\sim 21\%$ ) compared to the bulk hydrate. This difference is less significant when the clay used is MMT since small cage occupancy is  $85.2\% \pm 3.4\%$  (similar

to sand) and  $81.5\% \pm 2.6\%$  for MMT/sand mixtures with 10% and 60% clay, respectively. Our results indicate that the presence of clay affects the hydrate cage occupancy and it seems to be related to the nature of the clay (Table 5). Conversely (Yeon et al., 2011), found anomalous cage occupancy of natural hydrates disseminated in Na-montmorillonite with  $\theta_{LC}/\theta_{SC}$  up to  $\sim 2.40$ . They show that the mobile interlayer  $\text{Na}^+$  cations are inserted into SC during hydrate formation and compete with methane, thereby reducing the absolute occupancy of SC. An analogy might be made with the  $\text{K}^+$  ions present in the interlayer space of illite. However, this is unlikely because of the strong interaction between this anhydrous and non-exchangeable cation and the illite sheets that reduce their mobility in the presence of water (which does not enter into the interlayer space) (Dettmann, 1958; Odom & Low, 1978). By increasing the pressure from 3.5 to 15 MPa during bulk methane hydrate formation, Qin and Kuhs found a very slight change in the relative cage occupancy from 1.15 to 1.10, probably related to pressure-dependent change of the relative size of the large and SC (Qin & Kuhs, 2013). Our samples were synthesized under the same temperature and pressure conditions, the only parameter modified during the experiments was the mineralogy of the matrix used. However, a clear explanation cannot be given at this stage on the decrease in absolute occupancy of SC with increasing illite content.

This decrease in the filling rate of SC induces a small decrease in the hydrate storage capacity (Figure 9) ranging from  $163.9 \pm 1.0$  for the bulk hydrate to  $158.7 \text{ m}^3 \pm 1.5$  in the ILL/sand mixture with 60% clay, that is, a deviation of  $\sim 3\%$ . Considering only the minimum and maximum values for bulk hydrate and the hydrate-bearing ILL respectively, a maximum deviation of  $\sim 4.5\%$  is reached. From our experiments, the increase in clay content does not significantly affect the GSC of hydrate. However, given the very large volumes of methane stored in hydrate deposits (around  $10^{15} \text{ m}^3$ ), these findings could affect the overall assessment of the amount of methane stored. For comparison, when the cages are completely filled (i.e.,  $\theta_{LC} = \theta_{LC} 100\%$ ), the hydrate storage capacity is  $172 \text{ m}^3$  and the values that were commonly used in the estimates of methane-in-place volumes in hydrates are 180 (Klauda & Sandler, 2005), 170 (Gornitz & Fung, 1994), or  $164 \text{ m}^3$  (K. A. Kvenvolden, 1998). All these values are higher than our values calculated from the micro-Raman spectroscopy analyses.

#### 4. Application to the Black Sea Hydrate Deposits

The Black Sea is the largest isolated sea in the world with an area of  $4.23 \times 10^5 \text{ km}^2$ . It represents the largest anoxic basin on Earth (Overmann & Manske, 2006), and it is characterized by the occurrence of gas hydrates from the slope sediments to the deep sea (Ker et al., 2019; Klaucke et al., 2006; Popescu et al., 2006; Riboulot et al., 2018). In the Romanian sector of the Black Sea, hydrates are encountered within clay-rich sediments ( $\sim 60\%$  clays, Table 3) characterized by a relatively diverse mineralogy (illite, smectite, and kaolinite), with depth-variable mineral proportion (Ballas et al., 2018; Ruffine et al., 2021). We performed deep micro-structural characterization of the hydrate-bearing sediments as a function of depth on the collected core GAS-CS05 from the western Black Sea.



**Figure 10.** Absolute cage occupancy (left), gas storage capacity (middle), and clay mineralogy (right) of the natural hydrate-bearing sediment samples as a function of depth. The dash dotted lines in the left and middle figures represent the average small cage occupancy and the storage capacity obtained from the synthetic hydrates in the ILL matrix with 77% illite (red), the 53% MMT/47% ILL matrix with ~45% smectite (gray) and the synthetic bulk hydrate (blue).

As for synthetic hydrates, micro-Raman spectroscopy performed on the natural specimens has allowed to obtain the variation of cage occupancy, which in turn enabled to estimate the GSC as summarized in Figure 10. We observed that the large cages are almost fully occupied with  $\theta_{LC} \sim 98\%$  along the entire length of the core. However, small cage occupancy varies with depth, highlighting two zones (Figure 10a). The first zone is located at less than 2.4 m below sea floor (mbsf) and is characterized by average values of  $\theta_{SC}$  varying between  $\sim 74\%$  and  $\sim 78\%$ , which correspond to an average storage capacity between 157.8 and 159.4  $m^3$ . The hydrate samples collected at depth below 2.4 mbsf exhibit more SC occupied by methane. Thus, hydrates have a  $\theta_{SC}$  ranging between 79.3% and 87.4%, and a resulting storage capacity of around 160.8 and 163.2  $m^3$ , respectively. Interestingly, the horizon located at  $\sim 2.4$  mbsf corresponds to the change in clay mineralogy (Figure 10c), with an illite-rich upper zone ( $\sim 62\%$  illite) and a lower zone dominated by the presence of smectite ( $\sim 53\%$ ). Our results show a clear link between hydrate cage occupancy and sediment mineralogy: the SC are less occupied in the illite-rich layer compared to the smectite-rich layer. These results are in good agreement with those determined from our investigations from the synthetic cores discussed in the first part of this paper, as shown in Figures 10a and 10b. These figures also present the average values of small cage occupancy and gas storage capacities obtained on the hydrates synthesized in the ILL matrix, the 53% MMT/47% ILL matrix and the bulk synthetic hydrates.

Several estimates of the amount of  $CH_4$  bound as hydrates in the Black Sea (Klauda & Sandler, 2003; Mery & Sinayuc, 2016; Vasilev & Dimitrov, 2003) have been made in the last two decades using different storage capacities. Using an EF of 180  $m^3$ , Klauda and Sandler calculated an amount of methane bound in Black Sea hydrates of  $85 \times 10^{13} m^3$  (Klauda & Sandler, 2003). A total volume of  $71.8 \times 10^{12} m^3$  was estimated by Mery and Sinayuc using an EF value of 165.6  $m^3$  (Mery & Sinayuc, 2016). The difference between these two estimates is mainly related to the methods used to derive hydrate saturation in sediments. However, the Black Sea sediments consist on average of  $\sim 60\%$  clay, with illite being the dominant clay mineral ( $>51\%$ ) (Ballas et al., 2018; Huvaj & Huff, 2016; Ruffine et al., 2021; Stoffers & Müller, 1972). Applying to their estimates the average storage capacity of the hydrate in the illite-rich layer (158.7  $m^3$ ), determined from our analysis, the total volume estimated is  $\sim 11.8\%$  (i.e.,  $10^{14}$ ) and  $\sim 4.6\%$  (i.e.,  $3.4 \times 10^{12} m^3$ ) less than from those determined by Klauda and Sandler, and Mery and Sinayuc, respectively.

More recently, it has been shown for the Romanian sector of the Black Sea, that salinization triggered the destabilization of shallow gas hydrates, releasing methane that contributes to the current bubble emissions widely observed at the slope area (Riboulot et al., 2018). The authors estimated that, at the scale of the Black Sea, this destabilization could release in the long term up to  $\sim 2.1 \times 10^{11} m^3$  of  $CH_4$  (EF  $\sim 165 m^3$ ) into the water column, with potential impact on marine ecosystems and climate if the gas reaches the atmosphere. We obtained a differ-

ence of  $\sim 3.83\%$  ( $8 \times 10^9 \text{ m}^3$ ) using a storage capacity of  $158.7 \text{ m}^3$ . Even if important uncertainties on some parameters inherent to the deposits such as the hydrate saturation in sediments (Vasilev & Dimitrov, 2002) remain to achieve better estimates of methane trapped in hydrate on Earth, it appears from our study that integrating microscale properties such as cage occupancy and the matrix mineralogy is essential.

## 5. Conclusions

Experiments of methane hydrate formation were conducted in matrices consisting of mixtures of sand, illite and montmorillonite, with an aim to investigating how matrix mineralogy affects the formation processes. Our results showed that mineralogy significantly affects the formation kinetics and GSC of hydrates. Indeed, clay behaves as a kinetic inhibitor in comparison with sand, as adding only 10% from a sand matrix increases the average induction time by more than 60%. It was also shown that the incipient hydrate formation occurs much faster in illite-rich matrix than in the montmorillonite at low to medium water saturation (35%–55%  $S_w$ ). This has been attributed to the combined effect of capillarity and the low activity of the strongly bound water in montmorillonite, which significantly reduces the amount of water available for hydrate formation. The effect of water adsorption and capillarity vanished when increasing water saturation in the clayey matrices, leading to a decrease in the induction time at higher water saturations.

Micro-Raman spectroscopy analysis on the synthesized hydrates revealed that increasing the clay content up to  $\sim 60\%$ , in illite in particular, decreases the average small cage occupancy by  $\sim 12\%$  compared to the bulk hydrate. This variation mechanically leads to a decrease in the average GSC that reaches a value of  $\sim 158.7 \text{ m}^3$ , lower than those generally used to estimate the amount of hydrate-bound gas. These results on synthetic hydrates were found to be consistent with those obtained from natural hydrate samples collected in the Romanian sector of the Black Sea, and therefore allowed us to revise the estimate of methane trapped in this deposit. Overall, our study shows the importance of considering the effect of the mineralogy of the matrix on hydrate formation kinetics and GSC; key properties to better predict the amount of methane potentially released from hydrate decomposition, either by climate change or anthropogenic operations for natural gas production.

## Data Availability Statement

The data of sediments characterization, pressure, and temperature from high pressure experiments for hydrate synthesis, and of hydrate cage occupancy and storage capacity, are available in Mendeley Data (<https://data.mendeley.com/>, Agnissan, 2023).

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