The impact of mineral compositions on hydrate morphology evolution and phase transition hysteresis in natural clayey silts

Hang Bian a,b, Xuwen Qin c,d,e, Jinsheng Sun f,g, Wanjing Luo a,**, Cheng Lu h,b,d,e, Jian Zhu h, Chao Ma d,e, Yingfang Zhou a,i,*

a School of Energy Resources, China University of Geosciences, Beijing, 100083, China
b Center of Oil & Natural Gas Resource Exploration, China Geological Survey, Beijing, 100083, China
c China Geological Survey, Beijing, 100083, China
d Guangzhou Marine Geological Survey, China Geological Survey, Guangzhou, 510075, China
e National Engineering Research Center of Gas Hydrate Exploration and Development, Guangzhou, 511458, China
f China University of Petroleum (East China), Qingdao, 266580, China
g CNPC Engineering Technology R&D Company Limited, Beijing, 102206, China
h iRock Technologies, Beijing, 100094, China
i School of Engineering, University of Aberdeen, UK

ARTICLE INFO

Handling Editor: Wojciech Stanek

Keywords:
Clayey silt
Hydrate phase transition
Micro-scale properties
Micro-scale imaging

ABSTRACT

Natural gas hydrate is a clean and high-efficient energy resource, and more than 90% of its reserves are contained in fine-grained (typically clayey silts) sediment. In this work, for the first time, the micro-scale imaging is performed to explore hydrate phase transition, morphology evolution and fundamental characteristics (mineral compositions, pore structures and seepage capacity) in clayey silt sediments. The results indicate that clayey silts formation properties strongly depend on dominant minerals component in the sediments. The clay-rich clayey silt possesses more microcapillary interstice with smaller permeability than that in quartz-rich sediment. Hydrates generally occur as microfracture-filling (veins) and grain-displacing (nodulus) in sand-rich clayey silt. While they occur in the form of fracture-filling (vein) and foraminifera-filling in clay-rich clayey silt sediments. Biological fossils (especially foraminifera) provides potential space for hydrate aggregates. But hydrate formed inside it depends on the structures of fossils, the mineral components and pore structure of surrounding matrix. The hysteresis between hydrate formation and decomposition found to be more significant in clay-rich clayey silt sediment than quartz-rich sediment. And this hysteresis inside foraminifera is much more serious compared with that in matrix. In addition, dispersed pore-filling hydrates forms during decomposition, which has adverse effect on continuous gas production.

1. Introduction

Under the background of economic globalization, the global energy consumption is projected to increase in the coming years [1]. However, carbon emission from the conventional fossil fuels has led to increasing environmental concern and global warming, which becomes an imminent consideration [2]. NGH is crystalline solid, in which guest gas molecules are trapped in host water cavities [3]. NGH is widely distributed in continental permafrost and marine sediments, and it is considered as an environmental-friendly alternative energy resource compared with other fossil fuels [4]. The estimated global natural gas hydrate reserve is twice of that in combined conventional fossil fuels [3]. The promising prospect has arisen great interests of energy consuming countries in recent years and many of them have carried out geological explorations [5] or pilot field tests [6]. According to statistics, hydrate in clayey silt sediments contains more than 90% of the total NGH resources in the world [7,8]. Typical clayey-silt hydrate reservoirs are globally distributed, including the Ulleung Basin in Korea [9], the northern part of the South China Sea [10], the Gulf of Mexico [11,12], the Cascadia Margin [13], the offshore Peru [14,15] and the K-G Basin in India [16], etc.

Thoughtful understanding of hydrate formation and decomposition
process, such as hydrate morphology, hydrate phase transition and the evolution of them can provide strong theoretical support for better NGH development. The attenuation of P- and S-waves and change of conductivity during seismic exploration and resource assessment are directly governed by hydrate structure and saturation in NGH reservoir [17–22]. In addition, the physical properties of hydrate-bearing sediments, such as thermal conductivity [23], intrinsic permeability [24] and mechanical strength [25] are also dominated by the specific distribution of hydrates in porous medium.

With the recent development of imaging technologies, micro-CT has been widely used as a nondestructive instrument to observe the inner structure of hydrate-bearing samples [26]. Numerous scholars have conducted research on hydrate formation and decomposition process based on CT technology. For hydrate formation, most of the scholars [27–30] found that hydrates preferentially developed at gas-water interface. Hydrates possess irregular microstructures [30], and the nucleation and distribution of hydrates are heterogeneous at grain scale as well as sample scale [28,31,32]. The distribution and migration of gas and water phases dominate the subsequent growth direction [31] and surface morphology of hydrates [28,31]. Specifically, Lei et al. [33] indicated that hydrate tended to adopt round and smooth surfaces when in contact with water, while exhibited angular interfaces when in contact with gas. Moreover, gas occurrence types can also alter hydrate occurrences [34].

For hydrate decomposition, some scholars indicated that hydrate decomposition started at the hydrate-gas interface, not at the contacts with quartz grains [6,35]. Gupta et al. [36] confirmed the characteristic of radial dissociation of hydrates. The different changing rate of hydrate surface area at different decomposition stages suggests that hydrate morphology will change during decomposition [37]. Some scholars indicated that this hydrate morphology transition is caused by the change of the distribution of decomposed water [35] and the limited diffusion of gas in water phase [38]. Moreover, hydrate re-formation was likely to occur during initial decomposition process [35]. And water saturation was the dominant factor of hydrate re-formation site [39]. Meanwhile the hydrate re-formation may promote homogeneous distribution of hydrate in pores [6].

The distinction between different hydrate habits (cementing or non-cementing) was initially deduced from the effect of hydrate morphologies on elastic wave velocity [40]. Several idealistic theoretical models (including pore-filling, grain-cementing, grain-coating, load-bearing and patchy) for pore habits of laboratory-synthesized hydrates in porous medium have been proposed in previous literatures [41–45]. Some researchers have conducted the studies on hydrate pore habits evolution based on micro-CT technology. Results show that different hydrate formation conditions (such as excess-gas and excess-water) can lead to different evolution rules of hydrate habits [31,33]. Mass and heat transfer during hydrate formation can also lead to habit transformation.

However, it is worth noting that the above-mentioned studies were conducted in large size (hundreds to thousands of microns) glass beads or quartz sand, which are the experimental substitutes of coarse-grained hydrate sediments. These materials are enormously different from the natural fine-grained clayey silt hydrate sediments in mineral composition, particle size distribution, types of porous space and seepage capacity. The natural clayey silt hydrate sediments possess typical characteristics of tight reservoirs, including extremely small pore size, low porosity and ultra-low permeability [46]. Numerous field coring data indicates that hydrates in coarse-grain sediments basically show the same hydrate occurrence as those observed in large-grained glass beads or quartz sands in laboratory tests. While the hydrate occurrences in fine-grained sediments (typically clayey silts) mainly show in grain-displacing occurrence (including fracture-filling/veins, chunks/-nodes and lenses types), which are significantly different from those in coarse-grained sediments [47,48]. The essence of the difference is that further hydrate growth mostly proceeds by invading adjacent pores in coarse-grained sediment when hydrate growth is restricted by mineral particles; While in fine-grained sediment, further hydrate growth is mainly achieved by replacing particles to enlarge the occupied pores [41]. It is mainly caused by the relative balance between the effective stress and the hydrate-water capillarity in sediments with difference particle sizes. Capillary force prevails in fine-grained sediment, so that hydrate is easier to replace particles than to invade into small throats. The pore throat size of coarse sediment is much larger and the capillary force is insignificant. Therefore, hydrate is more inclined to invade the adjacent pores under high effective stress [41].

By far, there has been limited lab study on hydrates in natural clayey silt at micro-scale. Wu et al. observed hydrate occurrences in silty sediment and discussed mechanical properties of hydrate-bearing clayey silt by self-developed CT triaxial apparatus [49]. Liu et al. [50] conducted nano-CT scanning on samples from Shenhu area in the South China Sea and found that clayey silts are usually rich in foraminifera, which increases pore space of sediments. Liu et al. [51] performed X-CT scanning on natural hydrate-bearing clayey silt pressure-core at low-temperature and high-pressure condition. Results indicated that there were only bulk hydrates formed in clayey silt without bio-fossils, and no dispersed hydrates were found in matrix pores. Li et al. [52] synthesized hydrate in foraminifera and demonstrated that liquid and gas migrated into the shells through through-holes and hydrates directly contacted inner wall of foraminifera. At present, there is few research on evolution of hydrate occurrences in fine-grained sediments, and even less research on the characteristics of hydrate phase transition in such sediments.

The overall aim of this work is to investigate the characteristics of hydrate phase transition and morphology evolution in clayey silts. Two representative clayey silts with different dominant minerals collected from the hydrate reservoir in the South China Sea are selected by quantitative characterization through QEMSCAN technology. Quantitative characterizations of particle size, pore space and seepage capacity are performed based on the obtained multi-scale spatial structures of samples by SEM and micro-CT scanning techniques. Similar to the previous study [53], xenon is utilized as a substitute of methane to generate hydrate in this study. Hydrate phase transition experiments in natural clayey silts are successfully performed by using stepwise pressurization and depressurization methods for the first time. The pore-scale distribution and morphology evolution of hydrate at typical states during formation and decomposition are captured. The decisive effect of mineral compositions on the phase transition properties and morphology evolutions has been discussed contrastively. The results of this research are the theoretical basis of studies on evolution characteristics of seepage capacity and mechanical properties during hydrate phase transition in clayey silts. In addition, the results possesses guiding significance to the production tests of clayey silt hydrate-bearing reservoirs.

2. Sample preparation and methodology

2.1. Dry scan of clayey silt samples

As observed in the field, the contents of silt and clay vary significantly in different fine-grained NGH sediments. For example, the overall sand content is lower in the K-G basin of India, but some high-concentrated sand regions can still be observed [54]. In Ulleung Basin, mineral contents of the sandy clay sediment are different from those in the clay-dominant fine-grained sediments [55]; and there are many microfossils dispersing in clay minerals in the clay-dominant sediments. In the South China Sea, the sand content in clayey silt sediment of Qiong Dongnan area is higher than that of Shenhu area [56,57]. While the clay content of the sediments in the offshore Peru, the Gulf of Mexico and the Cascadia Margin are generally dominant [41]. Previous work also suggests classifying the fine-grained NGH sediments into three main lithofacies with distinct grain size distributions (type-A-silty clay to clayey silt, Type B-clayey silt and Type C-sandy silt to silty sand), indicating
that grain size increases with the decrease of clay content [58]. And thus, two typical samples (clay-rich and sand-rich clayey silt natural hydrate sediments) were selected as example to analyze the hydrate occurrence evolution in clayey silt hydrate reservoir. Before phase transition experiments, the two samples are scanned by several methods to provide basic images for the subsequent qualitative and quantitative characterization of physical properties including mineralogy, pore space and seepage capacity. The specific preparation process of SEM samples is detailed in Appendix A.

**MAPS imaging.** Scanning electron microscope (SEM) is an instrument for high-resolution imaging of materials [59]. The electrons interact and scatter as impacting various minerals. SEM images can be obtained by scanning the surface of specimen and mapping the intensity of corresponding scattered electrons and other energy signals [60]. The primary electrons reflected by elastic collisions, also known as back-scattered electrons (BSE), is one of the main parts of the imaging signal in SEM. The electrons are converted into video signals through the detector and the SEM image can be formed after amplification and brightness adjustment.

MAPS is a modular automated processing system with automatic imaging control function. It can seamlessly stitch thousands of high-resolution SEM images together to generate a complete, multi-scale SEM image of the area of interest. In this study, MAPS images of the target areas are acquired by stitching plenty of consecutive SEM images with a resolution of 5 μm and pixel number of 2048 × 1768.

**QEMSCAN imaging.** QEMSCAN (Quantitative Evaluation of Minerals by Scanning electron microscopy) is a comprehensive automatic mineral detection method. QEMSCAN technology combines energy-dispersive spectroscopy (EDS) and backscattered electron (BSE) intensity from a SEM and identifies minerals by comparison with a predefined mineral database [61]. The specific descriptions are referred in the research of Gottlieb et al. [62]. In this work, a 15 kV electron beam is used to obtain 2D SEM images for quantitative analysis of minerals in a high vacuum condition. The scanning resolution is 5 μm, and the scanning area is 8 mm × 8 mm. A series of data processing is then performed, specifically including integrating several mineral phases into mineral aggregates, decomposing mixed spectra (boundary phase processing), image filtering, and particle classification.

**FIB-SEM imaging.** FIB-SEM (Focused Ion Beam Scanning Electron Microscopy) is the combination of SEM and focused ion beam [63,64]. Firstly, the 3D slice range and thickness (number of slices) are set. After scanned by electron beam, the sample surface is etched by the large focused ion beam and then smoothed by the small one. Subsequently, SEM scans the new sample surface [65]. This cycle continues until the sample is cut to a predetermined depth. Finally, plenty of continuous 2D images are reconstructed into 3D FIB-SEM images using reconstruction algorithm.

**Micro-CT imaging.** 3D internal space of samples can also be tested by micro-computed tomography (micro-CT), which is a non-destructive technology. The high-pressure chamber is fixed on the 3D detection platform, and rotates 360° for scanning [53]. The radiation emitted from X-ray source is attenuated by various phases and then received by the detector. To optimize image quality, each sample is scanned under 120 kV. By converting the attenuated X-rays into digital and electrical signals, the attenuation coefficients are acquired, and 3D gray-scale images of the samples can be subsequently obtained by 3D reconstruction algorithm [53].

### 2.2. Hydrate phase transition experiment

**Experimental preparations.** A cylinder with a length of 4 mm and a diameter of 5 mm are sampled from each of the two pieces for hydrate phase transition experiment. Xenon gas (purity of 99.999%) and the in-situ seawater produced by thawing frozen samples serve as raw materials of hydrate formation. Both methane hydrate and xenon hydrate are classified as Type I structure hydrate [66]. The similarities of the two hydrates in various aspects have been verified, including lattice structure, phase equilibrium thermodynamic, growth kinetics and mechanical properties [67–69]. In addition, the linear attenuation coefficient of xenon hydrate is much higher than that of methane hydrate [66,70], which can significantly enhance adsorption of X-rays and thus improve imaging contrast. Serious risks related to the operations with potentially explosive gas under high pressure can be greatly alleviated, since the stability pressure of xenon hydrate is much lower. In brief, xenon hydrate is a convincing substitute of methane hydrate for experimental study. Moreover, according to the latest in-situ NMR and well logging data achieved during natural gas hydrate drilling program in the Shenhu Area [71], pore space of clayey-silt hydrate reservoir is mainly filled with capillary irreducible water, followed by clay irreducible water, with almost no movable water. Thus, excess-gas method is used here to generate connate water condition in the clayey-silt hydrate reservoir.

The experimental setup (see Fig. 1) is an integrated test system composed of experimental apparatuses and micro-CT facility. Specific parameters of experimental apparatus were introduced in detail in the previous work [53]. At preparation stage, micro-CT operation is tested. The scanning resolution of sample A (clay-rich) is 5.04 μm, while it is 2.54 μm for sample B (sand-rich). Then, xenon gas is charged and discharged for several times in the chamber (loading sample) to exhaust the air.

**Experimental procedures.** Fig. 2 depicts pressure-temperature paths over the entire experiment with temporal distribution of micro-CT scans of clayey silts. Temperature is controlled at approximately 285.15 K (12 °C) throughout the process. Stepwise pressurization method is utilized for hydrate formation and stepwise depressurization is used during decomposition. As shown in previous studies, most of the clayey silt is unconsolidated with a labile structure [72–76]. The microstructure and fluid distribution will be influenced if a larger pressure gradient is applied, and thus a relatively small pressure gradient should be applied to avoid potential damage. Preliminary tests for hydrate formation and decomposition with different pressure gradients were carried out. Based on the results of a series of preliminary tests, the pressure gradient for hydrate formation, which can not only greatly maintain the original porous space, but also control the formation process within an appropriate period. Moreover, small gradients are also used to capture the initial pressure of hydrate phase transition for analyzing the hysteresis phenomenon (Section 3.4).

Multiple CT scans are conducted within phase transition process. The CT scan of ‘Initial State’ is obtained before injecting gas into the chamber, and the initial scanning time is set as time ‘0’. In the stage of pressure-induced hydrate formation, xenon gas is injected continuously into the chamber with initial pressure equal to 0.1 MPa and then the pressure is increased stepwise with an interval of 0.1 MPa until hydrate is formed in the sample. Hydrate formation starts when pressure increased to 1.1 MPa in sample A, to 1.0 MPa in Sample B; then second CT scan, ‘P-1’, is performed immediately. Subsequently, xenon gas continues to be injected into the chamber until the pressure reaches 1.5 MPa to ensure sufficient hydrate formation. CT scan ‘P-2’ is performed after 7 days (168 h) at this pressure-temperature state to obtain the initial fluid-hydrate-gas-solid distribution before hydrate decomposi-
tion. The pressure is then reduced stepwise with an interval of 0.1 MPa to study hydrate decomposition process. The first CT scan of hydrate decomposition process, ‘DP-1’, is conducted 24 h after initial hydrate decomposition (the pressure at this point of sample A is 0.6 MPa and that of sample B is 0.3 MPa). The pressure continuously to drop by 0.1 MPa, and CT scan ‘DP-2’ is performed after keeping at this state for 24 h. This process is repeated until hydrates disappear thoroughly, and then the pressure is released as the atmosphere pressure.

### 2.3. Micro-scale characterization and modeling based on images

**Image postprocessing.** Equipment operating conditions (such as current, voltage, etc.) and instability of external conditions (such as
temperature and pressure) during imaging process will affect sensors and circuit elements, and therefore generate image noise. In this work, the non-local means filter \cite{77} is utilized for 2D and 3D image denoising while preserving the original information of the image greatly. The watershed algorithm \cite{78} is then applied for phase segmentation based on the atomic number contrast of SEM images (including 2D BSE and 3D FIB-SEM images) or on the relative radiodensity of various materials to X-rays in micro-CT images. Thus, the resulting segmented gray-scale images can be used for further micro-scale characterizations and simulations.

**Minerals characterization.** Through enlarging 2D MAPs images, different types of mineral grains and pore space structures at whatever positions can be clearly observed. Based on Qemscan images, minerals content of samples can be quantitatively analyzed. Therefore, not only mineralogical information, but also grain morphology, mineral embedded characteristics and elemental occurrences of the clayey silt samples can be obtained.

**Particle size distribution.** Minerals can be separately extracted and marked based on 2D MAPS images with high resolution. The single particle size can be calculated with the method of equivalent diameter, as shown in Eq (1),

$$E_qD = \sqrt{\frac{4 \times S_{area}}{\pi}}$$

where $E_qD$ denotes equivalent circular diameter, $S_{area}$ denotes the surface area of the single particle. Finally, the particle size distribution can be statistically summarized.

**Pore space characterization.** Pore space characterization based on 2D MAPS images is to acquire the equivalent aperture of pores through central axis method, after which the continuous distribution of multi-scale pore size can be obtained by statistical analysis.

While based on 3D images, porosity of clayey silt samples can be calculated as the fraction of pore voxels over the total voxels. Pore size distributions are obtained by NMR simulations. NMR relaxations are simulated using the Random Walker algorithm \cite{79,80}. Throat size distributions are obtained by performing mercury injection simulations in 3D digital models. Moreover, tortuosity of the pore space can be obtained by simulating Brownian Motion of diffusive particles (random walkers) through pore space using the Random Walker Method \cite{81}. These methods have been described in detail in previous study \cite{53} and will not be repeated here.

**Intrinsic permeability simulation.** In this study, computational fluid dynamics (CFD) simulations are performed in 3D digital models to obtain absolute permeability in one certain direction. The fluid flow in porous space is assumed to be extremely slow and incompressible. The linear Stokes Equations are solved by the finite difference method (FDM). Specific introduction was described in detail in previous work \cite{53}.

3. Results and discussions

3.1. Fundamental characteristics of clayey silts

Fig. 3 shows the backscattering SEM images of sample A and sample B with magnifying parts indicating dominant minerals and various types of pore space. The Fig. indicates that particles in both samples are randomly arranged and unconsolidated. Particle size of sample A is much finer than that of sample B. In sample A, the proportions of carbonate (basically in the form of biological fossils) and clay are relatively high. Other minerals (such as quartz, mica and feldspar, etc.) are almost tightly surrounded by clay minerals. The porous space in sample A can be classified as microfractures, biological fossil pores (mainly foraminifera), interparticle pores, and clay intercrystalline pores. Among them, biological fossil pores and clay intercrystalline pores comprise a large proportion of total porous space.

As for sample B, porous space occurs mainly in the form of large pores, interparticle macropores and clay intercrystalline pores. The proportion of biological fossil pores is much less than that of sample A.
Particles are relatively loose-packed. Clay minerals do not totally fill the pore space among large-grained minerals, but mostly surround these minerals.

**Minerals.** Occurrences of particles with different shapes and dimensions in the two samples suggest complicated compositions and heterogeneous distributions of clayey silt minerals. Fig. 4 demonstrates quantitative characterization of minerals in both samples. Moreover, minerals are classified into several main types (including carbonate, clay, quartz, mica, feldspar, pyrite and others) and their proportions are shown in Fig. 4 (C). For sample A, the ratio of carbonate (mainly representing in biological fossils) is the highest at 37.03%; the second is clay, whose content is more than a quarter of the whole; and quartz accounts for approximately 18%. While for sample B, the most abundant mineral is quartz (44.43%), and both clay and carbonate occupy approximately 16%.

According to Fig. 5, particle size range of sample A is relatively wide. The first peak of the distribution curve (A) occurs at the particle size of approximately 4 μm, which is contributed mainly by clay minerals. While the second peak occurs at the particle size of 450 μm, representing the abundant biological fossils (such as foraminifera). The distribution curve of sample B peaks when particle size reaches 40 μm, which is approximately 10 times larger than that of sample A (excluding biological fossils).

**Pore space and seepage capacity.** Firstly, the large-area pore size distributions of samples (Fig. 6) are obtained based on MAPS images. The radii of pores in sample A covers a wide range from submicron to submillimeter. Among them, 83.4% of the pores are submicron-scale pores. The curve peaks at 18.1% when the pore radius is 0.25 μm. Submillimeter pores are intragranular pores of biological fossils, accounting for approximately 1%. The distribution curve of pore radius in sample B peaks at 10.1% when the pore radius equals 1 μm. There is a low peak (1%) between 0.1 μm and 0.2 μm in pore radius. The number of pores with radius between 1 μm and 10 μm accounts for 55.4% of the total, and the proportion of submicron pores occupies 32.6%, which denotes that more than half of the pores in sample B are macropores or intergranular pores.

Larger-sized micron-scale and millimeter-scale pores increase the total porosity of clayey silts and provide potential space for hydrate.
aggregates. However, plenty of submicron-scale or even nanoscale clay intercrystalline pores in silty matrix is the main factor affecting hydrate formation and seepage characteristics, which cannot be well identified at the resolution of MAPS images. Therefore, FIB-SEM scanning with a resolution of 65 nm and micro-CT scanning with a resolution of 0.95 μm are conducted respectively on silty matrix of sample A and B. Previous research shows that the selection of typical REV can promote analysis efficiency (accuracy) [82–84]. Fig. 7 shows the respective REV model of silty matrix of the two samples. Relative threshold segmentation process and gray distribution curves have been shown in Appendix B. Through size independence verification (detailed in Appendix C), the side length of the REV from clay-rich clayey silt matrix is determined to be 300 pixels, while that of the REV from quartz-rich clayey silt matrix is 500 pixels. Simulated physical parameters of pore space and seepage capacity of two sample matrix are listed in Table 1. Pore-throat size distribution curves are depicted in Fig. 8.

Table 1 demonstrates that porosity values of the silty matrix in the two samples are similar. The effective porosity accounts for more than 95% of the respective total porosity, indicating that pores in the two silty matrixes are well-connected. However, due to the high content of fine-grained minerals (mainly clay minerals) in sample A, the mean pore radius and mean throat radius are both less than 0.1 μm, which are less than one tenth of those in sample B. Combined with Table 2 [85], it can be seen from Fig. 8 that more than 70% of pores and 85% of throats in sample A matrix are categorized as microcapillary interstice. While nearly 90% of pores and more than 85% of throats in sample B matrix are classified as capillary interstice.

Moreover, tortuosity of the pore space inside silty matrix of sample A is higher and the fluid flow should be more complicated. Simulation results confirm that although porosity values of the two samples are similar, their seepage capacities differ significantly. According to Chen et al. [86], silty matrix of sample A with the permeability of far less than 0.1 md belongs to tight reservoir. While permeability of that in sample B is more than 680 times that in sample A, and sample B matrix belongs to low-permeability reservoir.

3.2. Hydrate occurrence in clayey silts

Micro-CT images of the clay-rich clayey silt (sample A) at typical states are listed in Fig. 9. Some microfractures are randomly distributed, which makes the structure highly heterogeneous. Three of them are circled with yellow dashed box and denoted as A, B and C, respectively. Hydrate formation in clay-rich clayey silt. At the early stage, the injected gas with the increasing pressure cannot flow into the matrix due to the ultra-low permeability of the dense clay-dominated silty matrix. Thus, xenon gas accumulates in fracture A, which is in good connectivity with the external space. Meanwhile, hydrates form rapidly along fracture sidewall. Since hydrates in fracture A are in direct and sufficient contact with gas, the water in matrix gradually diffuses to the surface of hydrate layer [87] for further hydrate formation. Both the outward migration of internal liquid and the accumulation of injected gas cause local compression of the unconsolidated sediment. As a result, fracture A

Fig. 3. MAPS images of Sample (A) and (B) with magnifying parts showing main types of minerals and pore space.
is transformed into a penetrating fracture with increasing aperture (shown in the state P-1 and P-2), providing more space for hydrates. In this case, hydrates would fill the entire fracture if gas and water could be sufficient.

As for fracture B in a deeper position, the fracture aperture is similar with that of fracture A at initial state. Gas gradually invades forward till fracture B is connected with fracture C. In the meanwhile, a certain amount of hydrate is formed in B and a thin hydrate layer occurs along sidewall of C. However, the upper matrix continuously moves downward, which hinders gas from entering fracture B and impedes subsequent hydrate formation. As a result, hydrate in fracture B only shows a slight increase during later formation and the occurrence of hydrates appears as fracture-filling.

While for fracture C, the aperture is the widest at the initial state. However, due to the poor connectivity between fracture C and gas, it narrows once the pressurized gas is injected into the sample. It is not until the limited gas in fracture B breaks into fracture C that a thin hydrate layer is formed on the sidewall of C. Subsequently, hydrate formation in C is terminated due to gas supply disruption. Fracture C is continuously compressed until the previously formed hydrates fill its space.

In addition, it is difficult for gas to enter inside matrix pores due to the ultra-high capillary force caused by both microcapillary interstice and the water film surrounding clay particles. On the other hand, hydrate formation continuously extracts water from the matrix to the previously formed hydrates (microfracture-filling and foraminifera-filling in this study) and the matrix becomes compacted at the meanwhile [88]. Thus, although the direct verification is limited by the resolution of micro-CT, it should be hard to form dispersed pore-filling (embedded) hydrates in such capillary and microcapillary interstice of

Fig. 4. (A) Qemscan image of sample A; (B) Qemscan image of sample B; (C) Proportions of main mineral categories in the two clayey silts.
clay-dominated silty matrix. Hydrate mainly exists in the form of fracture-filling and foraminifera-filling. 

Hydrate decomposition in clay-rich clayey silt. In the initial stage of depressurization (from P-2 to DP-1), hydrates in fractures remain unchanged, while those in the gap between the sample and the chamber decrease obviously. From DP-1 to DP-2, the uniformly distributed 

Table 1
Porous space and seepage characteristics of the matrix part in clayey silt samples.

<table>
<thead>
<tr>
<th>Sample</th>
<th>Porosity [%]</th>
<th>Effective Porosity [%]</th>
<th>Permeability [md]</th>
<th>Tortuosity</th>
<th>Mean Pore Radius [μm]</th>
<th>Mean Throat Radius [μm]</th>
</tr>
</thead>
<tbody>
<tr>
<td>A</td>
<td>16.46</td>
<td>15.78</td>
<td>0.0126</td>
<td>13.55</td>
<td>0.0998</td>
<td>0.0917</td>
</tr>
<tr>
<td>B</td>
<td>19.67</td>
<td>18.98</td>
<td>8.68</td>
<td>9.1</td>
<td>2.06</td>
<td>1.38</td>
</tr>
</tbody>
</table>

Fig. 5. Particle size distribution of sample (A) and (B) based on MAPS images.

Fig. 6. Pore size distribution curves based on MAPS images of sample (A) and (B).

Fig. 7. The respective REV model of silty matrix in sample (A) and (B).
hydrate layer on top of fracture A decomposes into discontinuous particles, indicating inhomogeneous hydrate decomposition. Meanwhile, the thickness of hydrate layer at the bottom slightly increases, and the volume of hydrate aggregation at the lowest position becomes larger. It is analyzed that the decomposed water of top hydrates in fracture A migrates to the bottom, and then re-form hydrates with the trapped decomposed gas [60] under local high pressure and local reduced temperature. During this period, hydrates in fracture B and C still remain undecomposed. At the final state (DP-3), hydrates in fracture A, B and C completely decomposed; however, the structure of fractures cannot recover to the initial state. Specifically, the apertures of the fractures are narrowed compared with that at the initial state, fracture A remains penetrated and fracture B is prolonged accordingly. The gap between the matrix above fracture A and the chamber wall obviously shrinks, which indicates an expansion of the above matrix. It is speculated that the decomposed water absorbed into the matrix by the massive clay minerals leads to this phenomenon. However, specific distribution of decomposed water cannot be identified due to the resolution limitation of the current micro-CT technology.

**Hydrate formation in quartz-rich clayey silt.** Fig. 10 shows Micro-CT images of quartz-rich clayey silt (sample B) during hydrate phase transition with amplifying images of two local areas. At the initial state, sample B is loosely packed with intensively-distributed microfractures and pores inside, and no dominant channel exists between the sample and the chamber wall. In the early stage of hydrate formation (P-1), gas invades microfractures and hydrate forms at the meanwhile. From the local amplifying image at P-1, it can be seen that some microfractures (red dashed box) are completely filled with hydrates. Some hydrates displace sediment grains and occur in the form of nodulus. While other hydrate particles are scattered (blue dashed box) in microfractures and gradually connect with each other with the increasing hydrate saturation. As a result, the dispersed spheroid and discoid hydrate particles transform into vein hydrates with a greater aspect ratio. This phenomenon has also been found in the research of Wu et al. [49].

The sample is continuously squeezed downward at the same time as gas injection. The pore space where hydrates have not yet formed is compressed till the surrounding particles are tightly packed; while pores and microfractures partially occupied by hydrates are squeezed till the hydrates can support the compressed porous space. This phenomenon is consistent with the previous research that hydrate will act as the skeleton in clayey silt sediment with the increasing hydrate saturation [41, 89]. And hydrate formation in fine-grained sediment can increase the strength of sediment to some extent. Top surface of the sample gradually becomes concave with pressurization, causing further compression of matrix porous space. The previously formed hydrate deforms to some extent. Besides a few microfractures and pores well-connected with the sample-chamber gap or top gas source, most of the new-generated hydrates from P-1 to P-2 are located in the gap. This phenomenon should be caused by water migration from matrix to the gap during compression. Finally, hydrates in quartz-dominated clayey silt mainly show in

<table>
<thead>
<tr>
<th>Pore categories</th>
<th>Size range [μm]</th>
<th>Flow Characteristics</th>
</tr>
</thead>
<tbody>
<tr>
<td>Supercapillary interstice</td>
<td>Φ &gt; 500 or f_w &gt; 250</td>
<td>Fluids can flow freely under gravity with a high flow rate or even a vortex.</td>
</tr>
<tr>
<td>Capillary interstice</td>
<td>0.2 &lt; Φ &lt; 500 or 0.1 &lt; f_w &lt; 250</td>
<td>Fluids cannot flow freely due to the capillary force. Liquids can flow inside the pores only if the external force is greater than the capillary resistance.</td>
</tr>
<tr>
<td>Microcapillary interstice</td>
<td>Φ &lt; 0.2 or f_w &lt; 0.1</td>
<td>The attractive force between the fluid and the surrounding medium molecules is always extremely large, and an ultra-high pressure gradient is required for fluids migration.</td>
</tr>
</tbody>
</table>

**Fig. 8.** Pore-throat size distributions of silty matrix in sample (A) and (B) based on 3D pore space models.
microfracture-filling (vein) and grain-displacing (nodulus) forms.

**Hydrate decomposition in quartz-rich clayey silt.** In the initial stage of depressurization (DP-1), decomposition starts in the chamber-sample gap and the microfractures well-connected to the external porous space. Dispersed hydrates (in yellow circles) that not present during formation occur in matrix at state DP-1. After pressure drops to DP-2 and maintains for 24 h, the dispersed hydrates do not disappear, but show a significant increase. It is analyzed that hydrates in the pore space well-connected with the outside gradually decompose with pressure decrease, throats connecting with the internal pore space become wider. Therefore, pressure is gradually transmitted to the interior of matrix, causing further hydrate decomposition. During this period, the
support cementation is significantly lost with decreasing hydrate saturation, which leads to the decrease of hardness and failure strength of clayey silt sediment [90]. The matrix gradually migrates downwards. Meanwhile, hydrates in the chamber-sample gap decompose and the sample laterally extends until it is in close contact with the chamber wall. As a result, the well-connected pore space and the dominant migration channels (chamber-sample gap) for decomposed gas and water disappear. As the matrix pores in sample B belong to the category of capillary interstice, molecular gravitational forces are always exerted between the liquid particles and between the liquid and the pore walls. High capillary force prevents internal fluids from flowing freely. The local temperature decrease caused by hydrate decomposition and the local pressure increase caused by the accumulation of decomposed gas and water lead to hydrate reformation. The reformed hydrates appear in dispersed pore-filling habits embedded in the silty matrix. The embedded type of hydrate occurrence has also been found in the research of Wu et al. [49] and Dai et al. [41]. Pressure drops further to DP-2, fracture-filling hydrates continuously decompose. The decomposed gas and water flow into matrix pores, and are still trapped under molecular gravitational force. And the conduction of external pressure drop is still hindered by high capillary force. Thus, more dispersed pore-filling hydrates occur due to local temperature decrease and pressure increase. At the final stage of depressurization, matrix gradually becomes loosely packed and hydrates decompose completely. Abundant microfractures and matrix pores reappear. However, the inside structure is significantly different from that at the initial state.

In conclusion, for clay-dominated clayey silt reservoir, fine matrix particles are densely packed. Under the ultra-high capillary force and the strong adsorption force of clay minerals on water, it is difficult for gas to enter the microcapillary interstice in matrix. In general, gas can invade forward through microfractures and the formed hydrates are in the shape of fracture-filling. As for the initial fractures with large aperture or the penetrating fractures, hydrates grow in layer on fracture sidewalls and are gradually thickened. Hydrates would fill the fractures if gas and water could be sufficient. For both cases, hydrates continuously squeeze the surrounding matrix and widen fractures during growth, leading to the damage of porous space where no hydrate formed initially. Hydrates mainly exist in the form of microfracture-filling (vein) and foraminifera-filling. While the quartz-dominated clayey silt reservoir is rich in microfractures and matrix pores. Due to the high content of pores above submicron level, the connectivity between internal porous space and gas is better. Hydrates are in the form of grain-displacing (nodulus) and microfracture-filling (vein).

During hydrate decomposition, clay-dominated clayey silt migrates downward under gravity due to the loss of support from hydrates. Consequently, sediment strength decreases, fractures are narrowed. Finally, matrix structure is significantly different from that at initial state. Likewise, hydrates dissociation in quartz-dominated clayey silt starts from the locations that is well-connected with the external porous space. During decomposition, the disappearance of dominant channels (microfractures) as well as the accumulation of decomposed water and gas can lead to an increase of pore pressure and a decrease in temperature, which can cause re-formation of dispersed hydrates in matrix pores. There still be a large amount of microfractures and matrix pores after complete hydrate decomposition. However, the aperture of most fractures is narrowed and the matrix structure differs from that at initial state.

The results possess theoretical importance for practical development of clayey silt hydrate reservoirs. Targeted reconstruction methods should be implemented for reservoirs with different mineral compositions and hydrate occurrences. Appropriate reconstruction can largely ensure productivity by reducing the transformation from fracture-filling to dispersed pore-filling (embedded) hydrate and thus the discontinuity of production in quartz-rich reservoirs, or alleviating the shrinkage or even disappearance of dominant channels for decomposed gas and water in clay-rich reservoir. Moreover, for achieving long-term development of clayey silt hydrates, thermal stimulation should be taken as a parallel measure to weaken hydrate re-formation and its adverse impacts.

Error analysis between in-situ reservoir conditions and laboratory conditions. The results shown in this section are based on xenon hydrate phase transition tests at pore scale in laboratory. Due to the better stability of xenon hydrate at lower pressure, the experiment pressure is much lower than the actual reservoir pressure (13.5 MPa) [91].

For the microstructure of clayey silt sediment, Wu et al. [49] indicate that the void ratio of clayey silt sediment can significantly decrease with the increase of formation pressure. The actual higher pore pressure can enhance cementing bonds, the ability to resist deformation [92], and also increase the elastic modulus for the pore fluid. Therefore, the failure strength of hydrate-bearing sediment is higher, and the failure volumetric strain is lower at the actual reservoir condition. As for hydrate microstructure, Lee et al. [93] demonstrate that the geometry of grain-displacing hydrate formed by filling existing fractures are mostly governed by the geometry of fracture, whereas those formed by heaving mechanisms are governed by in-situ stress conditions and geometric properties. Assuming an unlimited amount of methane and an unlimited time for hydrate formation, the overburden pressure is the key factor for controlling the thickness of hydrate. There is a negative relation between the thickness of hydrate and the overburden stress [94, 95]. Thus, the actual hydrate thickness should be theoretically smaller than that of hydrates formed in laboratory.

3.3. Hydrate in foraminifera

Fig. 11 shows 3D distribution of foraminifera in sample A. The enrichment of foraminifera highly increases the volume of intragranular pores. According to the previous research, the high abundance of foraminifera in clayey silt is favorable for the storage and enrichment of hydrates [96,97]. Fig. 12 illustrates that the hard shell of foraminifera can keep their internal space from being compressed when unconsolidated clayey silt is deformed during hydrate formation. Some foraminifera (e.g., in the red circle in Fig. 9) are well-connected with external gas source; therefore, hydrates can be easily generated inside them. As for other foraminifera (e.g., in the blue circle in Fig. 9), clay minerals tightly wrap their outer shells due to sample deformation during pressurization. The blockage of gas channels connecting with their inner space prevents hydrate formation in these foraminifera. Therefore, whether hydrate can form inside foraminifera depends not only on their structure [98], but also on the types of minerals and matrix structure around them.

Two representative examples of hydrate phase transition inside foraminifera are listed in Fig. 12. Two pieces of hydrates in foraminifera A show as pore-filling morphology initially. Then, significant phenomenon

![Fig. 11. 3D distributions of foraminifera containing in clayey silt sample A.](image-url)
of hydrate Ostwald Ripening [99], which is generated by Gibbs-Thomson effect, is observed during process from P-1 to DP-1. That is, two pieces of hydrates merge together and gradually grow into a shape of regular cube. It should be noted that hydrate is only generated in the leftmost foraminifer. The highlighted material in the lower right circular pore should be pyrite aggregates. Likewise, there are two pieces of hydrates generated in foraminifer B. However, Ostwald Ripening phenomenon has not been observed as the two chambers in the shell are separated. Two pieces of hydrates grow as regular cubes in respective space before being confined by fossil walls. Foraminifera-filling hydrate tends to adhere to the inner walls of the shell. If time and/or gas and water source was sufficient, hydrates would fill the entire cavity. Moreover, this type of hydrate is completely surrounded by foraminifera shells, which cannot facilitate the cementation of clayey silt matrix during formation.

In addition, hydrate volume in both foraminifera A and B did not reach the highest point at pressure peak (P-2). During depressurization stage from P-2 to DP-2, hydrates inside foraminifera still continue to increase even though partial hydrates in matrix have already decomposed. In summary, foraminifera-filling hydrates in clayey silt reservoir show more significant decomposition hysteresis.

3.4. Hydrate phase transition hysteresis

During hydrate phase transition, the initial formation pressure (P-1) is found to be inconsistent with the initial decomposition pressure (DP-1) at the same temperature. As shown in Fig. 13, the pressure of P-1 of sample A is 1.1 MPa, while the pressure at DP-1 is 0.6 MPa. As for sample B, P-1 is 1.0 MPa and DP-1 is 0.3 MPa. In other words, hydrate shows the phenomenon of 'difficult to synthesize, difficult to decompose' in clayey silt. In Fig. 12, the hydrate initial formation (P-1), initial decomposition (DP-1) and final decomposition (DP-3) state points in both samples are projected on the xenon hydrate phase diagram, which is acquired in pure Xe system [100].

**Hydrate formation hysteresis.** Fig. 14 shows that initial formation pressures in two samples are different, but both of them are much higher than the equilibrium pressure at the same temperature. This phenomenon is supposed to be a composite effect of hydrate nucleation condition...
and phase boundary shift.

Firstly, when it comes to the hydrate nucleation conditions, Sloan indicated that both the gas phase and the water phase are disordered at the molecular level, whereas hydrate crystals are intrinsically ordered [3]. As entropy favors disorder over order, the initial hydrate formation is hindered by a long, metastable period (induction period). During this period, overpressure (or supercooling) as a driving force is indispensable for rearranging the disorderly gas and liquid water into the orderly hydrate crystal structure till the critical size for spontaneous growth. As shown in Fig. 15, hydrate nucleation and growth have direct analogies in crystallization processes such as the precipitation of salt from solution [3]. Mullin [101] demonstrated that (1) the right side of the equilibrium line AB is the stable region, in which crystallization is impossible; (2) the area between AB and CD is the metastable (supersaturated) region, in which spontaneous crystallization cannot occur, but if any seed exists in the metastable solution, the seed will grow; (3) the left side of CD is the unstable region, in which spontaneous crystallization can take place. A direct analogy is shown in Fig. 15 (b) for hydrates when the concentration axis of Fig. 15 (a) is replaced by a logarithmic pressure axis [3]. The AB and CD lines in Fig. 15 (b) possess the same meaning as in (a) by analogy. To the left of line CD in (b), hydrate nucleation will readily occur due to the high driving force [3]. In summary, a high driving force (overpressure in this study) is required for initial spontaneous hydrate formation in a system without hydrate seeds.

Secondly, various factors of practical clayey silt hydrate reservoirs can lead to phase boundary shift. Capillary effect of small pores can shift thermodynamic phase boundary toward lower temperature (or higher pressure), which is referred to as the Gibbs-Thomson effect [102]. Specifically, Liu et al. [103] proposed that capillary effect of pores in fine-grained quartz sand (particle size <35 μm) is so significant that temperature can be reduced by 0.5–1.5 K at the same pressure. Clennell [104] indicated that fine-grained sediments inhibit hydrate formation until the temperature is 0.5–3 °C below the equilibrium value. Clennell [104] also pointed out that strong adsorption of water by clay-rich sediments can reduce water activity and thus inhibit hydrate formation, especially at high gas saturation. Moreover, Yang et al. [105] demonstrated that saline ions in aqueous solution possess ionic effect, which can destroy ionization equilibrium, alter the equilibrium constant of hydrated ions and thus shift the phase boundary to lower temperature and higher pressure. In summary, hydrate nucleation conditions and phase boundary shift caused by various factors make the initial hydrate formation pressure significantly higher than the equilibrium value.

Furthermore, the initial hydrate formation pressure (P-1) in sample A is higher than that in sample B. According to the information in Table 2, an ultra-high pressure gradient is required for gas entering the microcapillary interstices in sample A matrix. Moreover, it is difficult for water to flow freely as the rich clay possesses strong adsorption to water. Therefore, hydrate formation can only start when gas and water are in contact with each other through a high driving force (overpressure) and a slow diffusion process. While for sample B, microfractures are intensively distributed and sizes of matrix pores are larger. Low content of clay minerals possesses weaker adsorption to water and smaller pressure gradient is required for fluid flow in porous space. That is, the driving force required for initial hydrate formation in sample B is lower than in sample A.

**Hydrate decomposition hysteresis.** Hydrate decomposition should be easier and faster than hydrate nucleation, as hydrate decomposition is an entropy-increasing process. That is, ordered hydrate crystals develop into disordered gas and water. According to Sloan [3], the theoretical P-T point at which hydrates completely decompose should be infinitely approaching phase boundary. However, both the DP-3 points in Fig. 14 are beneath the phase equilibrium line. In particular, the initial decomposition pressure (DP-1) of the hydrate in sample B is located in the stable region, that is, hydrates have not start to decompose until the system pressure drops to the stable region. It seems to be contradictory with the upper shift of phase boundary during hydrate precipitation.
formation. In some literatures, this unexpected high stability of hydrate at conditions out of thermodynamic equilibrium is also called ‘self-preservation’ effect [106,107].

According to the results in section 3.1, matrix pores of sample B are mostly capillary interstices, and hydrates mainly occur as grain-displacing (nodulus) and microfracture-filling (vein) types. Hydrate formation and creep deformation of clayey silt can lead to a dramatical reduction in porous space. The ultra-high capillary force of narrow throats hinders pressure conduction into pores. Moreover, the pore space of in-situ clay-silt hydrate reservoir is mainly filled by capillary-bound water, followed by clay-bound water, with almost no movable water [71]. Therefore, hydrates remain unchanged until the system pressure drops to a point where the internal and external pressure difference can overcome high capillary force of matrix pores and water adhesion force by clay minerals. While for sample A, hydrates occur as fracture-filling type (except those in foraminifera). The conduction of system pressure is not limited by capillary force. Decomposed water and gas can transport without the hindrance of matrix pores. Thus, initial system pressure required for complete hydrate decomposition in samplehydrate decomposition pressure (DP-1) in sample A is higher than that in sample B, that is, hydrate in sample A is easier to decompose.

As shown in Fig. 12, foraminifera-filling hydrate is the other main type of hydrate occurrence in sample A. In the early stage of depressurization, clay minerals wrapped around foraminifera restrict pressure conduction. While in later stage, clay minerals hindered the output of decomposed water and gas. According to the results in section 3.3, foraminifera-filling hydrates show a significant decomposition hysteresis. Therefore, the final decomposition pressure (DP-3) of hydrate in sample A is above the phase boundary. For sample B, the thin throats hinder transport of decomposed gas and water, which inhibits further hydrate decomposition. Subsequently, the low temperature caused by decomposition and the high pressure caused by accumulation of gas and water result in hydrate re-formation in pores with smaller size. Thus, the system pressure required for complete hydrate decomposition in sample B is lower than that in sample A.

Different hydrate occurrences and mineral compositions of reservoirs can lead to different hysteresis degrees of hydrate phase transition. Targeted depressurization schemes should be adopted for hydrate reservoirs under different conditions. Reasonable pressure depletion together with monitoring system can largely avoid geological hazards potentially triggered by strength reduction during decomposition. The factors and relevant sensitivity analysis of hydrate decomposition hysteresis in reservoirs with inhomogeneous pore distribution and large differences in mineral contents need to be investigated in detail in the future.

4. Conclusions

In this work, hydrate phase transition experiments are performed in two clayey silts collected at different locations of hydrate reservoir in the South China Sea. The basic characteristics, including mineral compositions and porous space, are qualitatively and quantitatively analyzed. The main conclusions are as follows.

1. In clay-rich clayey silt, more than 70% of the matrix pores (except biological fossils) are microcapillary interstice. And the matrix is classified as tight reservoir (k ≈ 0.01 mD). For quartz-rich clayey silt, more than 90% of the matrix pores are capillary interstice and the matrix is low permeable (k < 0.03 mD).

2. For clay-rich clayey silt, hydrates occur in the form of microfracture-filling (vein) and foraminifera-filling. While hydrates mainly exist in the form of microfracture-filling (vein) and grain-displacing (nodulus) in quartz-rich clayey silt

3. Biological fossils (especially foraminifera) are mainly found in clay-rich clayey silt sample. Although the abundant foraminifera can provide favorable porous space for hydrate formation, the through-holes on some foraminifera shells are tightly blocked by clay minerals, which hinders hydrate formation inside.

4. During hydrate formation in clayey silts, the clay minerals, the saline ions, the high content of fine grains (Gibbs-Thomson effect) and the high driving force (overpressure in this study) required for a critical-size hydrate core formation induced by entropy reduction, can all lead to the shift of phase boundary toward low-temperature and high-pressure condition. This phenomenon of hydrate formation hysteresis is more obvious in the clay-rich sample.

5. The impeded pressure conduction can lead to decomposition hysteresis. Pressure difference must continuously increase for initial decomposition until it can overcome the high adhesion of water by clay minerals and the high capillary force of matrix pores and throats. Hydrates in biological fossils present more remarkable decomposition hysteresis.

6. During hydrate decomposition, silty matrix gradually moves downward due to decreasing strength. For clay-rich sample, the dominant channels for decomposed water and gas will be compressed and affect productivity. While in quartz-rich reservoir, capillary interstitium hinders migration of decomposed gas and water, resulting in the dispersed formation of secondary hydrate in silty matrix and the consequent discontinuous gas production.

Therefore, for clayey silt hydrate reservoirs with different mineral compositions and hydrate occurrences, both the reservoir stability and the decomposition hysteresis should be taken into consideration, and reasonable depressurization scheme should be adopted together with monitoring system. Targeted reconstruction methods should be used to impede the shrinkage or even disappearance of dominant channels and to avoid the secondary pore-filling (embedded) hydrates dispersed in fine capillary interstice. Moreover, in order to achieve long-term stable development of clayey silt hydrate reservoirs, thermal stimulation should be considered as parallel measure. The future work should focus on how the mechanical and fluid flow properties change with hydrate phase transition, which hope to be contributed to improved measures for impelling efficient development in fine-grained hydrate sediments.

Credit author statement

Hang Bian: Conceptualization, Methodology, Writing – original draft, Validation, Formal analysis; Xuwen Qin: Conceptualization, Investigation; Jinseng Sun: Conceptualization, Investigation; Wanjing Luo: Conceptualization, Writing – review & editing; Cheng Lu: Resources, Funding acquisition; Jian Zhu: Experiment design, Writing – review & editing; Chao Ma: Writing – review & editing, Investigation; Yingfang Zhou: Conceptualization, Writing – review & editing.

Declaration of competing interest

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

Data availability

Data will be made available on request.

Acknowledgements

The authors are grateful to the National Natural Science Foundation of China, China [51991365]; China Geological Survey Project, China [DD20211350]; Guangdong Major Project of Basic and Applied Basic Research, China [2020B0301030003]; Key Program of Marine Economy Development (Six Marine Industries) of Special Foundation of Department of Natural Resources of Guangdong Province, China [2021]56.
Appendix

A. SEM sample preparation

Firstly, the original sample was prepared precisely into a specified shape. Then, the prepared sample was dried in the oven for dry imaging. After that, the argon ion polishing instrument is utilized to polish the sample surface. The principle is that the ionized argon gas accelerates to the sample and the surface is polished by ion sputtering through momentum transfer during collision. As a result, the surfaces are smooth and parallel. And the carbon coater is finally used for protecting sample surface and improving the scanning effect of SEM.

B. Threshold segmentation and gray distribution curves

For efficient analysis, the original 16bit binary images are firstly transformed into 8bit images. Due to the great variety of minerals, the curves of various substances overlap each other. The integral gray value distribution curve of each silty matrix is shown in Fig. A.1 (A) and (B), respectively. The segmentation threshold of the two phases (solid phase and fluid phase) is determined by the watershed algorithm [108] in Avizo software. The segmented CT images are shown in the figures. The blue represents pore space and the yellow represents rock phase. The threshold value of clay-dominated clayey silt matrix is 129, and that of quartz-dominated clayey silt matrix is 69.

C. Size independence verification of the REVs shown in Fig. 7

In the silty matrix of the two clayey silt samples, 5 REVs at random positions are selected and the side length of REV (the minimum is 50 pixels) increases with an increment of 50 pixels. The maximum side length of REV in clay-dominated matrix is 500 pixels, while it is 800 pixels in quartz-dominated matrix. The curves of porosity versus side length of REV have been shown in the Fig. A.2 (A) and (B). With the increase of the side length of REV, the porosity gradually converges. The average and the variance of the porosity of REVs with different sizes are calculated. The hypothesis tests (T test and F test) are performed with the values of the maximum REVs, and the confidence interval is 95%. Detailed descriptions of the methods can refer to the research of Liu et al. [109]. Finally, the side length of the REV from clay-dominated clayey silt matrix is determined to be 300 pixels, while that of the REV from quartz-dominated clayey silt matrix is 500 pixels.
References


