1	Laboratory insights into the effect of sediment-hosted methane hydrate morpholog			
2	on elastic wave velocity from time-lapse 4D synchrotron X-ray computed			
3	tomography.			
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#### 16 Key Points

17 We observe the evolution of methane hydrate morphology in porous media by 4D X-ray CT

18 imaging and laboratory geophysical experiments.

19 X-ray CT shows that hydrate morphology evolves from an initial pore-floating, to pore-bridging,

20 to a final inter-pore hydrate framework.

We found anomalously low S-wave velocity probably caused by the presence of water filmsbetween hydrate and host grains.

## 23 Abstract

24 A better understanding of the effect of methane hydrate morphology and saturation on elastic 25 wave velocity of hydrate bearing sediments is needed for improved seafloor hydrate resource and 26 geohazard assessment. We conducted X-ray synchrotron time-lapse 4D imaging of methane 27 hydrate evolution in Leighton Buzzard sand, and compared the results to analogous hydrate 28 formation and dissociation experiments in Berea sandstone, on which we measured ultrasonic P-29 and S-wave velocity, and electrical resistivity. The imaging experiment showed that initially 30 hydrate envelops gas bubbles and methane escapes from these bubbles via rupture of hydrate 31 shells, leading to smaller bubbles. This process leads to a transition from pore-floating to pore-32 bridging hydrate morphology. Finally, pore-bridging hydrate coalesces with that from adjacent 33 pores creating an inter-pore hydrate framework that interlocks the sand grains. We also observed 34 isolated pockets of gas within hydrate. We observed distinct changes in gradient of P- and S-35 wave velocity increase with hydrate saturation. Informed by a theoretical model of idealized 36 hydrate morphology and its influence on elastic wave velocity, we were able to link velocity 37 changes to hydrate morphology progression from initial pore-floating, then pore-bridging, to an 38 inter-pore hydrate framework. The latter observation is the first evidence of this type of hydrate 39 morphology, and its measurable effect on velocity. We found anomalously low S-wave velocity 40 compared to the effective medium model, probably caused by the presence of a water film 41 between hydrate and mineral grains.

42

#### 43 1 Introduction

44 Gas hydrates are naturally occurring ice-like clathrate compounds that form when sufficient gas 45 (methane is the most common in nature) and water coexist under low temperatures and high 46 pressures, generally found in marine and permafrost environments (Kvenvolden, 1993). 47 Currently, seafloor gas hydrates are being considered as a viable alternative energy resource 48 (Boswell & Collett, 2011), and may have an important role in future climate change (Archer et 49 al., 2009), carbon dioxide sequestration (Jung et al., 2010) and continental slope stability (Sultan 50 et al., 2004). As such, it is important to obtain accurate estimates of the amount and distribution 51 of gas hydrates, largely reliant on geophysical remote sensing technologies and data 52 interpretation. Such estimates depend on knowledge of hydrate formation processes and how 53 they affect geophysical properties. In general, the presence of hydrate increases the seismic 54 velocity (Helgerud et al., 1999) and electrical resistivity (Edwards, 1997) of host sediments; this 55 depends on the amount of hydrate occupying the pore space (saturation) and hydrate 56 morphology, i.e., spatial distribution of the hydrate grains within the host sediment (e.g., Dai et 57 al., 2012; Ecker et al., 2000; Priest et al., 2005; Waite et al., 2009). In particular, geophysical 58 remote sensing methods use elastic wave velocity and electrical resistivity anomalies to quantify 59 hydrates in marine sediments, based on rock physics models that relate these anomalies to 60 hydrate content (e.g., Collett, 2001; Cook & Waite, 2018; Doveton, 2001; Ecker et al., 2000; 61 Edwards, 1997; Helgerud et al., 1999; Spangenberg, 2001).

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63 Accurate quantification of *in situ* methane hydrates is hampered by our limited understanding of 64 the effects of hydrate content, morphology and distribution on the geophysical properties of the 65 hydrate bearing sediments, along with sediment type, porosity, permeability, and pore fluid 66 salinity (e.g., Waite et al., 2009). These effects are difficult to understand unambiguously from 67 studies of natural samples alone because of spatial averaging. Moreover, using natural samples 68 for laboratory studies of geophysical and geomechanical properties is challenging because: (i) 69 coring is technically difficult and requires expensive drill ships with pressurised sampling 70 capability; and (ii) absolute preservation of in situ conditions is not possible currently (Tulk. 71 1999). However, controlled laboratory experiments on synthetic hydrate samples offer a viable 72 alternative to gain insights into the physical properties of hydrate-bearing sediments. Synthetic 73 hydrate samples allow exploration of potentially the full range of hydrate saturations and

74 morphologies for different sediment types, although laboratory methods have their own75 challenges.

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77 Notable insights have been gained from laboratory studies to date (e.g., Handa, 1990; Kerkar et 78 al., 2014; Priegnitz et al., 2015; Priest et al., 2009; Tohidi et al., 2001), but further research is 79 needed into the following areas: i) the causes of the commonly observed discrepancy between 80 hydrate saturation estimates from seismo-acoustic and electrical resistivity methods (Attias et 81 al., 2016; Goswami et al., 2015; Lee & Collett, 2006; Miyakawa et al., 2014; Sahoo et al., 2018) 82 (referred to here as the seismic-electrical discrepancy); and ii) the effect of methane hydrate 83 saturation and its spatial distribution in the host sediment on the seismo-acoustic velocity of 84 hydrate bearing sediments. Some studies associate the seismic-electrical discrepancy to the coexistence of gas and hydrate, as the presence of gas can reduce the seismic velocity but not the 85 electrical resistivity of the sediment (e.g., Goswami et al., 2015; Lee & Collett, 2006; Miyakawa 86 87 et al., 2014; Sahoo et al., 2018). This discrepancy could also be due to incorrect assumptions 88 about the morphology or distribution of hydrate within the pores.

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90 Natural hydrates commonly exist in several different morphologies (or habits) within the host 91 sediments. In this study, the term "morphology" refers to the spatial distribution of the hydrate 92 grains within the host sediment. Natural hydrate can be broadly divided into two main types 93 based on its morphology : sediment grain displacing or pore-fluid displacing hydrate (e.g., 94 Holland et al., 2008). Sediment grain displacing hydrate physically moves apart sediment grains, 95 forming solid hydrate volumes larger than the original sediment pore size; examples include 96 hydrate veins, layers, and lenses generally found in fine-grained sediments (e.g., Holland et al., 97 2008). By contrast, pore fluid displacing hydrate grows inside the intact structure of sediment pores. Most pore fluid displacing natural hydrate is observed in cores from coarse-grained silty 98 99 or sandy layers. For example, cores from NGHP1 (Collett et al., 2015) and IODP Expedition 100 311 (Riedel et al., 2010) showed pore fluid displacing hydrate in coarse-grained layers. Such 101 sandy units are often the targets for hydrate reservoirs of potential economic importance, and we 102 restrict this study to pore-fluid displacing hydrate.

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104 Pore-fluid displacing hydrate can be sub-divided into cementing or non-cementing morphologies 105 (Figure 1) based on whether hydrate grows adhering to sediment grains or floating in the pore 106 fluid inside the pore space (e.g. Ecker et al., 1998). The distinction between different pore-fluid 107 displacing hydrate morphologies were initially deduced from the effect of hydrate morphology 108 on elastic wave velocity (e.g. Ecker et al., 1998). . Cementing morphology occurs when hydrate 109 bonds the host mineral grain contacts (Ecker et al., 1998); the effect on elastic wave velocity was 110 conceptualized as either hydrate located exclusively at grain contacts (contact cementing) or 111 hydrate evenly coating mineral grains (grain coating), a proportion of which bonds grain contacts 112 (Ecker et al., 1998; Helgerud et al., 1999). Several studies (e.g., Chand et al., 2006; Priest et al., 113 2005) have deduced from elastic wave measurements that hydrate forms cement under excess 114 gas conditions by coating the mineral grains, with a fraction of the hydrate saturation acting as 115 cement. Formation of hydrate in cementing or non-cementing morphology also depends on the 116 sediment mineralogy; clay and sand interact with hydrate differently (Kumar et al., 2015; Sloan 117 & Koh, 2007). By contrast, non-cementing hydrate forms when hydrate grows away from the 118 sediment grain contacts (Ecker et al., 1998). The cementing morphology has a much greater 119 effect on the elastic properties of hydrate-bearing sediments than the non-cementing morphology 120 (e.g., Best et al., 2013; Ecker et al., 1998; Priest et al., 2009; Waite et al., 2004). However, the 121 non-cementing morphology is thought to dominate natural hydrate systems, and has been 122 sampled, or inferred, at locations such as Mallik, Mackenzie Delta (Uchida et al., 2000), the 123 Nankai Trough (e.g. Fujii et al., 2015), Alaminos Canyon, Gulf of Mexico (Boswell et al., 2009), 124 and Mount Elbert, Alaska North Slope (Stern et al., 2011). Useful summaries of observations of 125 hydrate morphologies at various sites around the world are given in Holland et al. (2008) and in 126 Dai et al. (2012).

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128 If non-cementing hydrate grows in the pore space without bridging neighbouring sediment 129 grains, then it is termed pore-floating (Hu et al., 2014) or pore-filling hydrate (i.e. hydrate may 130 be partially filling the pore, but not contacting more than one grain of the sand frame). In this 131 manuscript, we use the term "pore-floating" for such a hydrate morphology. If hydrate bridges 132 neighbouring sediment grains (i.e. contacts more than one grain in the sand frame) then this is 133 termed "frame-supporting" or "load-bearing" or "pore-bridging" hydrate. We will use the term 134 "pore-bridging" to describe this morphology, which has been reported for pore-floating hydrate

135 saturations greater than 25 - 40% (Hu et al., 2014; Priest et al., 2009; Waite et al., 2009). Priest et 136 al. (2009) deduced a pore-bridging morphology for methane hydrate formed in sand under excess 137 water conditions from observed increases in seismic velocity at hydrate saturations higher than 138 30%; for saturations of 10% and 15%, the seismic velocity was between those for pore-bridging 139 and pore-floating hydrate (Priest et al., 2009). (Yun et al., 2005) showed that Tetrahydrofuran 140 (THF) hydrate grows in the pore-floating morphology up to 40% hydrate saturation; for higher 141 saturations, the measured velocity was much higher than that predicted for the pore-floating 142 morphology, qualitatively consistent with a pore-bridging morphology. While cementing and 143 pore-bridging hydrate are both associated with an increase in the elastic moduli of the composite 144 sediment, pore-floating hydrate affects the elastic properties of the pore-fluid (e.g., Ecker et al., 145 2000).

Studies of gas hydrate using techniques like X-ray imaging have shown that gas hydrate often has a complex morphology. Recent studies in sands suggest that a thin film of water is present between the host mineral grains and the hydrate (Bonnefoy et al., 2005; Chaouachi et al., 2015; Sell et al., 2018; Tohidi et al., 2001). This water film should exist for both cementing and porebridging hydrate in sands.

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152 Recently, Sahoo et al. (2018) found hydrate formation does not take up all the methane gas or 153 water even if the system is under two phase water-hydrate stability conditions, leading to 154 coexisting gas, water and hydrate. Sahoo et al., (2018) deduced this coexistence of gas and 155 hydrate using thermodynamic calculations from pore pressure and temperature measurements. 156 The authors hypothesised that the dominant mechanism for coexisting gas is the formation of 157 hydrate films around methane gas bubbles. Also, co-existence of gas and hydrate in the gas 158 hydrate stability zone (GHSZ) has been inferred in natural sediments (e.g., Guerin et al., 1999; 159 Milkov et al., 2004; Lee and Collett, 2006; Miyakawa et al., 2014). Researchers have attributed 160 this coexistence to the following causes: (i) influx of gas into the GHSZ along fractures or faults (Gorman et al., 2002; Lee & Collett, 2006; Smith et al., 2014); (ii) local deviations from two 161 162 phase water-hydrate stability conditions (pressure-temperature-salinity) resulting in local hydrate 163 dissociation within the GHSZ (Guerin et al., 1999; Milkov et al., 2004); or (iii) hydrate 164 formation kinetics (Torres et al., 2004). Inclusions of gas within hydrate can also enable 165 coexistence of gas with hydrate in two phase water-hydrate stability conditions (e.g., Schicks et

166 al., 2006). Hydrate formation on the surface of gas bubbles results in isolation of the remaining 167 gas inside the hydrate shell from the pore water outside, leading to co-existing gas. Sahoo et al., 168 (2018) showed experimentally that coexisting gas can cause significant errors in hydrate 169 saturation calculations from electrical resistivity, as both hydrate and gas are resistive compared 170 to seawater. Coexistence of gas and hydrate can also cause uncertainty in local hydrate saturation 171 estimation from electrical resistivity. For example, Miyakawa et al., (2014) proposed that co-172 existing gas and hydrate leads to a velocity decrease with no corresponding decrease in 173 resistivity in the Kumano basin, Nankai, Japan. Other such discrepancies found in the literature 174 are listed in Table 1 of Sahoo et al. (2018).

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176 In this study, we set out to observe changes in geophysical properties during methane hydrate 177 growth in coarse-grained hydrate reservoir analogues (porous media), and to link them to 178 observed changes in hydrate morphology using time-lapse (4D) X-ray CT imaging of the pore 179 spaces. We also want to image the mechanism of co-existing gas and hydrate in two-phase 180 water-hydrate stability condition. To achieve this, we conducted two separate laboratory hydrate 181 formation and dissociation experiments, one on Berea sandstone to obtain ultrasonic P- and S-182 wave velocity and electrical resistivity variations with hydrate saturation, and another on 183 Leighton Buzzard sand to obtain 4D time-lapse images from synchrotron radiation X-ray 184 computed tomography (SR-XCT). We then used the effective medium rock physics model of 185 Marín-Moreno et al. (2017) to predict the effect of changing hydrate morphology on elastic wave 186 velocities based on previously conceived idealised hydrate morphologies (pore-floating and 187 pore-bridging). The X-ray imaging confirmed the existence of these idealised morphologies at 188 certain periods during hydrate formation, and provided the first known direct evidence for a third 189 morphology, here called inter-pore hydrate framework. This last morphology was inferred to 190 affect the elastic velocities, although not modelled. Also, lower than expected S-wave velocities 191 were attributed to the presence of a water film between the inter-pore hydrate framework and the 192 host porous medium..

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196 Overall, our results provide further evidence of how methane hydrate saturation relates to

197 hydrate morphology, of how this morphology influences elastic wave velocity and electrical

resistivity, two important geophysical parameters used in hydrate exploration, and of the

199 mechanism of coexisting gas and hydrate.

200

## 201 **2 Methods**

#### 202 **2.1 Porous media**

203 We used a cylindrical sample (4.97 cm diameter and 2.06 cm height) of Berea sandstone with a porosity of 0.22, permeability of 448 mD ( $4.48 \times 10^{-13}$  m<sup>2</sup>) as a stable, inert, and well-204 205 characterized porous medium for the ultrasonic experiment. The use of loose sand would have 206 been preferable as a seafloor hydrate analogue, but the available pulse-echo system was 207 configured for rock samples only. The grain size is about 100 µm, similar to observations by 208 other researchers (e.g., Minagawa et al., 2008). X-ray diffraction analysis of the sample Berea 209 rock by Han et al. (2015) showed 1.7% illite and 3.3% K feldspar in volume. For the synchrotron 210 imaging experiment, we weighed and tamped Leighton Buzzard sand (a mean grain size  $d_{50}$ 211 =100 um) directly into the cylindrical hydrate rig (2 mm diameter and 23 mm height) to obtain a 212 sample of 35% porosity (a typical permeability is several Darcies for such sand packs). We tried 213 to cut a 2 mm diameter Berea sandstone sample suitable for synchrotron imaging, but the Berea 214 disintegrated during the attempts. Lee, (2008) found that the permeability of gas hydrate bearing 215 sediment at the Mallik 5L-38 with hydrate saturation between 12% and 34% to be very similar to 216 the permeability of hydrate bearing Berea sandstone (Kleinberg et al., 2003), and maximum 217 hydrate saturation in our experiment was 26%. Therefore, we choose Leighton Buzzard sand for 218 the synchrotron experiments, which has a similar quartz mineralogy (although uncemented). We 219 used the same hydrate formation method in both experiments according to Section 2.2 (i.e 220 hydrate forming from gas bubbles, in an excess water environment, with a water wet sediment). 221 Given the similar mineralogies and grain shapes of the host porous samples, we assume there is 222 no significant difference in hydrate morphology evolution between experiments, although this 223 has not been verified. Each sample was firstly oven-dried at 60 °C before placing in their 224 respective experimental rigs. 225

## 227 2.2 Hydrate formation

We followed the method of Sahoo et al. (2018) and Waite et al. (2004) with high initial brine saturation (83.5% for ultrasonic and 90% for synchrotron samples, respectively) giving excess water conditions (Ellis, 2008; Priest et al., 2009). Our experimental setup with gas injected from the base of the sample represents gas hydrate systems with localized gas flow, such as at the base of the gas hydrate stability zone (GHSZ), or near to gas chimneys.

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234 An initial hydrostatic triaxial confining pressure of 10 MPa was applied to the Berea sample in 235 the ultrasonic rig to ensure the integrity of coupling between the sample and buffer rods 236 (confining pressure was provided by the reaction of the rigid sample container to applied pore pressure in the synchrotron sample). A pore fluid line vacuum (<1 Pa) was applied to each 237 238 sample to maximize the removed air from the pore space. Still under vacuum, 3.5 wt% NaCl 239 deionized and deaerated water solution was injected to partially fill the sample pore spaces 240 (83.5% for ultrasonic and 90% synchrotron samples). The partially saturated samples were left 241 under vacuum conditions for 3 days, to favor a homogeneous pore fluid distribution throughout 242 the sample by capillary forces. The brine imbibition and distribution was facilitated by brine 243 injection after an applied vacuum, and by the high wettability of the quartz grains. Thus, we 244 assume water vapor and any remaining air occupied the remaining pore space of each sample.

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246 In the ultrasonic experiments, methane gas was injected to achieve a pore fluid pressure of 11.9 247 MPa and, and the confining pressure was increased simultaneously to 21.9 MPa to maintain a 248 constant differential pressure of 10 MPa (confining minus pore pressure). The pore fluid system 249 was then sealed, keeping the pore fluid line between the sample and valve V<sub>A</sub> (Figure 2a) filled 250 with methane gas, which is free to move in and out of the sample as a result of potential pore 251 pressure variations Buoyancy could make the gas to accumulate in the upper part of the sample. 252 However, that was not the case in our experiment, as we did not identify internal reflections in 253 the P-wave signal (in addition to the top and base sample reflections; Supplementary Figure S1). 254 The initial gas and water distribution in the synchrotron rig was observed from the 3D imaging, 255 discussed in Section 3.1. Finally, hydrate was formed by cooling each system to fall within the

- gas hydrate stability conditions (Figure 3) and above the freezing point of water. After hydrateformation, dissociation was achieved by heating the system (Figure 3).
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259 The hydrate formation procedure was similar in both experiments with slight variations as 260 described below. Ultrasonic P- and S-wave velocities of Berea sandstone are known to be 261 sensitive to changes in differential pressure due to microcracks in mineral grains (e.g., Nur & 262 Simmons, 1969). Hence, a constant differential pressure was maintained in the ultrasonic rig to 263 rule out any such effects. This approach ensured that any change in velocity could be attributed 264 to changes in pore fluids and hydrate formation only. Four cycles of hydrate formation and 265 dissociation were completed in the ultrasonic rig; a differential pressure of 10 MPa was 266 maintained for cycles 1 and 2, and 55 MPa for cycles 3 and 4. Hydrate, gas and brine saturation 267 were calculated from the measured pressure and temperature changes of the system using a 268 thermodynamic method (Sahoo et al., 2018). In the synchrotron rig, no confining pressure was 269 applied, with a pressure cell (made from PEEK) providing rigid confinement. We applied 10 270 MPa of methane pressure directly through the injection inlet, filling the pore fluid pipe and sand 271 sample volume with a pre-calculated amount of brine solution. We then left the rig for three days 272 for the pore fluids to redistribute within the pore space. Only one cycle of hydrate formation and 273 dissociation was performed in the synchrotron rig.

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#### 275 **2.3 Ultrasonic velocity and electrical resistivity measurements**

276 We used a stainless steel high-pressure cell, designed to host 5 cm diameter rock samples under 277 hydrostatic confining pressure up to 65 MPa (Figure 2a) (Ellis, 2008). The cell was 278 instrumented to monitor pore fluid pressure, and the inner and outer (ambient) cell temperature. 279 The inner temperature sensor was placed on the outer surface of the rubber sleeve to indicate the 280 sample temperature. The inlet pore pressure pipe was connected via a three-way valve to a 281 vacuum pump, a methane gas cylinder (with pressure regulator) and a brine reservoir. A syringe 282 pump was used to inject brine into the sample in a controlled manner, while the temperature of 283 the system was regulated by a controlled cooling circuit.

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The inner cell was configured for ultrasonic pulse-echo measurements of P- and S-wave velocity ( $V_p$  and  $V_s$ ) with an accuracy  $\pm 0.3\%$  (Best et al., 1994). The inner rubber sleeve that prevents

direct contact between the mineral oil, used as confining fluid, and the rock sample is perforated by 16 electrodes for electrical resistivity measurements (North et al., 2013). Under typical operating conditions, the resistivity measurement error is  $\leq 0.1\%$  (at A/C frequencies 1 - 500 Hz) for samples in the electrical resistivity range 1 - 100  $\Omega$  m (North et al., 2013). This system does not allow simultaneous ultrasonic and electrical measurements because the ultrasonic system gives a ground path for the electrical system. The resistivity system took nearly one hour for each measurement, so we have fewer resistivity measurements.

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#### 295 2.4 Synchrotron X-ray CT imaging

296 We designed and manufactured a miniature cylindrical hydrate rig to fit the SRXCT stage at the 297 TOMCAT beamline, Swiss Light Source (SLS), Switzerland. It was manufactured from 298 monolithic PEEK plastic by precision lathe and drilling machines (see Figure 2b). The rig had an 299 internal diameter of 2 mm, a wall thickness of 0.8 mm and sample scan height of 10 mm (23 mm 300 total height). Internal and external thermocouples were installed below the scan zone to measure 301 the temperature throughout the experiment. In order to reduce the temperature within phase 302 boundary, cooled nitrogen gas was blown at 5L/min onto the sample using CryojetXL (Oxford 303 Instruments). The temperature of the jet was initially calibrated by varying discharge and  $N_2$  gas 304 temperature to provide stable 2°C sample internal temperature. The gas hydrate formation and 305 dissociation process was imaged through computed tomography (CT) using monochromatic X-306 rays from a synchrotron source (TOMCAT SLS). Beam energy of 21 keV, 81 mm propagation 307 distance, 200 ms exposure time (1501 projections over 180° sample rotation) with 1.25 x, 4 x and 308 10 x objectives were chosen after trial runs to obtain images at 1.625 µm, 0.625 µm and 0.325 309 µm voxel size respectively. The transmitted and refracted x-rays from the sample was converted 310 to visible light by LuAG:Ce scintillator, thereafter magnified and recorded by sensitive CCD 311 cameras (2560x2160 pixels). Figure2b shows details of the hydrate rig including the pore fluid 312 injection system and temperature control. The TOMCAT facility uses phase shifts of the X-rays 313 as they pass through the sample (Fitzgerald, 2000; Stampanoni et al., 2002). We used the phase 314 reconstruction algorithm described by Paganin et al. (2002). Reconstructed CT data were post-315 processed using ImageJ and visualized using Amira-Avizo® 3D software.

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317 We calibrated the grey scale values in the CT data to known standard densities (sand, brine, 318 hydrate and methane gas; e.g., Kneafsey et al., 2007). The grey scale intensity represents the 319 extent to which the X-ray signals are attenuated. Different material in the sample, attenuate the 320 X-ray signal to a different extent, resulting in different grey scale intensity. The grey scale 321 intensities are mainly dependent on density of the material and beam energy but are also slightly 322 influenced by other factors like atomic number, reconstruction algorithm (Koeberl, 2002; 323 Phillips & Lannutti, 1997). Following the approach of previous studies (e.g., Kneafsey et al., 324 2007, Jassonov et al., 2009), the grey scale values were calibrated with density of sand, brine and methane gas  $(2650 \text{ kg/m}^3, 1025 \text{ kg/m}^3, 18 \text{ kg/m}^3)$  using scans with no hydrate. This provides a 325 326 relation between gray scale intensity and density. The grey scale intensity range for hydrate was 327 derived by using this relationship with a hydrate density of 925 kg/m<sup>3</sup>. By choosing the grey 328 scale intensity range derived from associated material density (Figure 4c), we segmented the 2D 329 slices stack to obtain separate sand, brine, hydrate and methane gas as binary images file stacks 330 (e.g., Iassonov et al., 2009). We then used these individual stacks to estimate the volume of gas, 331 hydrate and water. Porosity was calculated by adding the volume of gas, water and hydrate. An 332 example of reconstructed 2D slice at two different hydrate formation time along with its grey 333 scale intensity profile drawn between two sand particles passing a methane bubble and brine 334 within the pore between these particles is shown in Figure 4. The density based boundary lines 335 (Figure 4c) clearly shows ability of this segmentation technique to distinguish different phases 336 especially between methane hydrate, gas (between 20-60 $\mu$  profile length) and brine (65-75 $\mu$ 337 profile length) within the pore space. This segmentation allows a range of grey scale for each 338 component (sand, brine, methane gas and hydrate) to be highlighted, for enhanced visual 339 contrast. For example, Rees et al. (2011) studied a natural gas hydrate bearing sediment sample 340 collected offshore India (NGHP-1) using this segmentation technique.

Each scan took around 10 mins. However, as hydrate formation is a relatively slow process
taking between 40-50 hours, and we had to trade-off between storage and analysis capacity and
frequency of scans able to capture the main changes during hydrate formation, we decided a scan
frequency of ~30 minutes. We increased the scan frequency to the maximum possible (~15
mins) during rapid hydrate formation, and reduced the scan frequency during other times (1-3
hours). See supplementary information for additional scans (Movie S1, Figure S2).

## 348 2.5 Rock Physics Model

349 We used the hydrate bearing effective sediment (HBES) model of Marín-Moreno et al. (2017) to 350 relate changes in velocity to changes in hydrate saturation and morphology (at least for pre-351 conceived, idealised morphologies of cementing, pore-floating hydrate discussed in Section 1). 352 HBES model calculates frequency dependent (from seismic to ultrasonic) P- and S-wave velocity 353 and attenuation of hydrate bearing sediment (Figure 5). It was derived from the previous HEG 354 (Hydrate Effective Grain) model of Best et al. (2013) which adapted extant static (zero 355 frequency, broadly equivalent to seismic frequencies) velocity models for hydrate morphologies 356 (cementing, pore-floating, pore-bridging) by Ecker et al. (1998) and Helgerud et al. (1999) for 357 the purpose of predicting attenuation and velocity dispersion, using model concepts developed 358 by Leurer & Brown (2008) and Leurer (1997) for clay-squirt flow attenuation in marine 359 sediments. The central idea is that hydrate can be treated as an effective medium of solid hydrate 360 with fluid inclusions (similar to clay assemblages in Leurer et al. (1997)). During the passing of 361 an elastic wave, the different elastic compliances of the porous medium host (e.g. sand grain 362 framework) and the porous hydrate grains creates local fluid pressure gradients between the 363 hydrate inclusions and the sand frame pores, leading to viscous fluid flow (squirt flow) of water, 364 and associated wave energy loss. The squirt flow element is embedded in the Biot-Stoll global 365 fluid flow model (Biot, 1956b, 1956a)(Biot, 1956b, 1956a)(Biot, 1956b, 1956a)(Biot, 1956b, 366 1956a)(Biot, 1956b, 1956a)(Biot, 1956b, 1956a)(Biot, 1956b, 1956a)(Biot, 1956b, 1956a)(Biot, 367 1956b, 1956a)(Biot, 1956b, 1956a)(Biot, 1956b, 1956a)(Biot, 1956a, 1956b) giving an effective 368 medium solution for frequency-dependent P- and S-wave velocity and attenuation in hydrate-369 bearing sediments and rocks, as a function of both hydrate content (saturation) and the specific 370 morphologies above.

The HBES model extended the HEG model to include additional loss mechanisms identified from the pore-scale hydrate morphology, and the coexistence of gas, water and hydrate. In addition to the HEG model squirt flow due to hydrate grain fluid inclusions described above (termed sub-micro squirt flow in the HBES model), the HBES model introduces another form of squirt flow (termed micro squirt flow) due to the low aspect ratio pores that are created during

hydrate formation between the hydrate grains and the sand frame pore walls (Figure 5). The
model incorporates gas bubbles resonance effects according to Smeulders and van Dongen
(1997) (Figure 5).

379 The HBES model was developed for hydrate growing in the sediment pore space, and do not 380 account for grain displacing hydrate. The HBES model accounted for pore-floating and 381 cementing (grain coating and contact cementing) morphologies. We adapted the HBES model 382 for pore-bridging hydrate using the approach of Ecker et al., (2000), which considers that pore 383 bridging hydrate reduces the porosity and affects the elastic properties of the solid phase. All the 384 HBES model input parameters are given in Table 1. As our Berea sandstone sample had 1.7 385 volume % illite and 3.4 volume % k-feldspar, we first used the Voigt-Reuss-Hill average to 386 calculate the bulk and shear moduli of the grains, which were then used as inputs to the HBES 387 model. The saturation of hydrate, gas and brine in the pore space was calculated from changes in 388 pore pressure and temperature using the thermodynamics approach of Sahoo et al. (2018). This 389 saturation calculation showed the presence of co-existing gas even at maximum hydrate 390 saturation, and hence we included the bubble resonance effect of the HBES model. The pore size 391 in our Berea sample varied from 11 µm to 73 µm, measured from SR-XCT at TOMCAT, SLS 392 Switzerland (Sahoo et al., 2018), and we choose to use 10 and 20 µm bubble radii in the HBES 393 model. Based on observations from the synchrotron images we expect that, initially, the bubble 394 would almost completely fill the pore, and with the formation of hydrate, the bubble size would 395 reduce. The aspect ratio and concentration of pores created during hydrate formation were 396 chosen based on the values used in Marín-Moreno et al. (2017). The concentration of inclusions 397 in hydrate was set to zero, as they have a negligible effect on V<sub>p</sub> in our measurement frequency 398 band (Marín-Moreno et al., 2017).

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### 400 **3.0 Results and Discussions**

401

#### 402 **3.1 Synchrotron imaging of hydrate formation.**

After carefully inspecting the extensive X-ray CT dataset (9 TB of data), we selected the images
that could best describe the key hydrate formation processes that we observed (Figure 6). The 3D
volume reconstructed time-lapse sequence in Figure 6 demonstrates the evolution of hydrate

406 morphology within the pore space. A full time-lapse sequence of 2D slices is shown in the 407 supplementary information (Movie S1, Figure S2). Hydrate grows initially as a shell around gas 408 bubbles scattered throughout the pore space, as also reported for methane hydrate formation in 409 water without sediments (e.g., Klapp et al., 2012). The gas is sparsely distributed as sub-410 spherical "bubbles" with a range of sizes; some gas bubbles almost completely fill the pores, 411 while others occupy part of the pore only (Figure 6a, b). Hydrate formation starts at the gas-412 water interface as expected. Hydrate films that develop on bubbles at some point seem to 413 rupture, and the trapped gas escapes (Figure 6c, d). Hydrate formation consumes methane, 414 reducing the pore fluid pressure, and is also an exothermic reaction. The resulting pressure drop 415 and temperature increase may lead to bubble expansion and/or rupture of the hydrate shell. Some 416 of the escaped gas forms smaller bubbles which later forms more hydrate (Figure 6 c, d). Hydrate 417 growing in adjacent pores then starts to coalesce as the pores are further filled with hydrate, 418 creating an inter-pore hydrate framework interlocking with the sand grain framework (Figure 6c, 419 d). The spherical shapes of bubbles distort due to further hydrate growth into the gas bubble. 420 Ultimately, the gas remains as isolated pockets surrounded by hydrate, while hydrate occupies 421 most of the pore space in patches throughout the volume (Figure 6e, f). Such distorted bubble 422 shapes and uneven, porous hydrate distribution could have an impact on elastic wave 423 propagation mechanisms (Section 3.3). The presence of such pockets of gas confirms the 424 hypothesis of co-existence of water, hydrate and methane gas proposed by Sahoo et al. (2018) 425 from electrical resistivity and thermodynamic calculations. A film of water is evident between 426 hydrate and sand even at maximum hydrate saturation (Figure 6e).

The highest hydrate content occurs where the porosity is highest, but the lowest hydrate content is not where the porosity is lowest (Figure 7). The depth variation of  $S_h$  increases as the average  $S_h$  increases (Figure 7); at 45h 10 min,  $S_h$  varies between 38 and 60%, about a mean saturation of 44%. Even though the gas was injected from below, the maximum gas hydrate saturation occurs towards the upper part of sample. We installed two thermocouples to measure internal (just below the scan zone) and external (room) temperature; we did not observe any temperature increase in the sample during hydrate formation.

We note the following caveats when comparing the results of synchrotron imaging to theultrasonic experiments in the sections below. Although the sample's porosity, permeability, and

436 size are different, it may not affect observed hydrate morphologies as described in Section 2.1. 437 The inlet pipe in the synchrotron imaging rig is in the center of the sample, while it is off-center 438 for the ultrasonic rig (Figure 2). The ratio of diameter of each sample to that of the pore fluid 439 inlet pipe is 24.5 for the ultrasonic rig and 15.7 for the imaging rig, respectively. These 440 differences might have affected the initial distribution of the gas and hydrate in the sample. The 441 off-center position of the inlet pipe in the ultrasonic rig might have caused a less homogenous 442 distribution of gas in the sample compared to the synchrotron rig. However, as hydrate formation 443 in both experiment was from gas bubbles in excess water condition, we expect the observed 444 hydrate formation morphologies to be similar in natural hydrate systems, especially for coarse-445 grained sediments that are fed by free gas from below. For example, porous hydrate have been 446 found in the southern summit of Hydrate Ridge (offshore Oregon, USA) which likely formed when methane hydrate film coated gas bubbles moved upwards within the sediments and 447 448 coalesced together (Suess et al., 2001). Such porous hydrates have also been found offshore 449 Nigeria (Sultan et al., 2014).

#### 451 **3.2** Ultrasonic velocity changes during hydrate formation and dissociation.

452 The cooling of the system to 5 °C generated a reduction in pore pressure as shown in Figure 3, 453 points ABC. The pressure drop can be explained mainly by hydrate formation, with some 454 contribution from methane gas contraction and increased gas solubility between AB. A slight 455 increase in temperature can also be seen in Figure 3a between points B and C, caused by the 456 exothermic reaction of hydrate formation (Hwang et al., 1990). Figure 8 shows rapidly 457 increasing P- and S-wave velocities (V<sub>p</sub> and V<sub>s</sub>, respectively) during the initial 10 hours of 458 hydrate formation, followed by a more gradual increase thereafter. Once hydrate formation 459 ceased, indicated by the end of the pore pressure decrease at about 260 hours (Figure 3b), the 460 system was left at that pressure and temperature for several hours to ensure complete hydrate formation. The asymptotic behavior of the pore pressure in Figure 3b, and of  $V_p$  and  $V_s$  in 461 462 Figures 8a,b is evidence that no further hydrate formation took place (e.g., Waite et al., 2004). 463 As the system was taken out of hydrate stability by heating, there was an increase in pore 464 pressure (trajectory CD in Figure 3) and a decrease in  $V_p$  and  $V_s$  (Figure 8 c, d).

465

466 The time taken to reach the cycle's maximum velocity, and associated maximum hydrate 467 saturation, is longer in the first cycle than in subsequent cycles, which may be due to a methane 468 hydrate formation memory effect (Ohmura et al., 2003; Sloan & Koh, 2007; Takeya et al., 2000). 469 The memory effect is related to survival of meta stable clathrate clusters of water after hydrate 470 dissociation making subsequent hydrate nucleation more likely (Rodger, 2006; Sloan & Koh, 471 2007). As some researchers dispute the memory effect(e.g., Buchanan et al., 2005), we offer an 472 additional explanation. We propose this time reduction may be due to an increase in the contact 473 area of the gas/brine interfaces after the first formation and dissociation cycle, associated with a 474 more even distribution of smaller gas bubbles, as observed in the synchrotron images. Smaller 475 spheres have a larger surface area per unit volume than larger ones, and this change could 476 increase the total reaction surface area.

477

#### 478 **3.3 Morphology of hydrate from ultrasonic velocity**

#### 479 **3.3.1** Changes in gradient of change of velocity during hydrate formation

480 The observed increases in  $V_p$  and  $V_s$  in Figure 8a & b depend on both hydrate saturation and

481 morphology. Here, we apply the hydrate bearing effective sediment (HBES) model of Marín-

482 Moreno et al. (2017) to our experimental results with the aim of relating changes in velocity to

- 483 changes in hydrate saturation and morphology.
- 484

485 Small increases in  $V_p$  and  $V_s$  for hydrate saturations (S<sub>h</sub>) up to about 5% are well represented by 486 implementing a pore-floating hydrate morphology in the HBES model (Figure 9). As hydrate 487 saturation increases above 5%, the observed  $V_p$  increases more steeply and approaches the pore-488 bridging morphology model results at about  $S_h = 15\%$ . It is generally accepted that when 489 saturation of pore-floating hydrate increases, it eventually starts bridging the pores (Priest et al., 490 2009; Waite et al., 2009). Above  $S_h = 15\%$ , the gradient of  $V_p$  becomes smaller, and diverges 491 below the predicted pore-bridging increase of the HBES model. This observation indicates that, 492 for saturations above 15%, only a small proportion of the newly formed hydrate is adding to 493 bridging of the pores. Similarly, V<sub>s</sub> continues to track the pore-floating HBES model up to S<sub>h</sub> of 494 5%. For  $S_h > 5\%$ , in contrast to  $V_p$ ,  $V_s$  falls below the HBES model for pore-bridging 495 morphology. A possible explanation is that when hydrate fills the pores in our experiments, it 496 does not make solid-solid bonds to the host sand grains, as assumed in the HBES pore-bridging 497 model. Instead, a thin, bound water layer may exist between the water wet sand grains and the 498 hydrate, as observed in our synchrotron images (Figure 6). The presence of the water film could 499 increase the Berea's frame bulk modulus in a similar manner to the pore-bridging model, but not 500 the frame shear modulus. Only when sufficient hydrate has grown to interlock the sand grains 501 would the frame shear modulus increase, and then still less than for solid-solid contacts. This 502 phenomenon could explain the rise of V<sub>s</sub> above the pore-floating model line, but below the pore-503 bridging model line.

504

505 There is a steep increase in  $V_s$  around  $S_h = 23\%$  in Figure 9b, with a less pronounced increase in 506  $V_p$  (Figure 9a). This behavior is also seen in Figure 8. The increase in Vs with hydrate formation 507 in cycles 1 and 3 shows two distinct segments, with an initial increase followed by a plateau,

followed by a renewed velocity increase up to a plateau at the maximum  $V_s$  (Figure 8b). Cycles 1

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and 3 can be considered as the first cycles at each differential pressure, 10 MPa for cycle 1 and

510 55 MPa for cycle 3. The steep increase in cycle 1 and 3 (at 39 hours for cycle 1, and 47 hours for

511 cycle 3, in Figure 8b), might occur when sufficient hydrate has grown to interlock extensively

512 the sediment. The sudden increase in  $V_s$  could indicate a threshold of interlocking when the rock

513 frame shear modulus is stiffened significantly. The HBES model does not consider this new

- 514 inter-pore hydrate framework morphology..
- 515 While we expect different hydrate formation/dissociation rates in Berea and Leighton Buzzard

516 sand (e.g. hydrate formation of about 80 & 45 hours respectively), we would expect the pore

517 scale morphological evolution to be similar in both the experiments (as discussed in Section 2.1).

518 The coalescence of hydrate from adjacent pores creating an inter-pore hydrate framework, which

519 interlocks the host grains (Figure 6) could be linked with this rapid increase in  $V_s$  seen in Figures

520 8 & 9. Such a steep increase is less prominent in  $V_p$  (Figure 9) suggesting that the increase in

521 bulk modulus is dominated by replacement of pore-fluid with pore-bridging hydrate, irrespective

522 of whether the hydrate significantly interlocks or not.

For cycle 3, electrical resistivity also shows a similar steep increase at  $S_h = 21\%$  in Figure 8. We do not have resistivity measurements between  $S_h$  of 1% and 21%, but it is clear that steep increase in velocity matches with that of resistivity (Figure 10). The steep increase in resistivity can be also be explained by coalescence of hydrate from adjacent pores creating an inter-pore hydrate framework. When hydrate from adjacent pores coalesces, they might block electrical current conduction paths, causing a rapid increase in resistivity. The gradient of increase in resistivity seems to decrease at higher hydrate saturations, reaching a maximum at  $S_h = 25\%$ .

530

531 A possible explanation for the absence of such behavior in cycles 2 and 4 could be a reduction in 532 gas bubble size that results in pore-bridging aggregates formed by smaller hydrate "grains". 533 Hydrate forms on gas bubble surfaces, and smaller gas bubbles would result in smaller hydrate 534 grains. With more hydrate formation, such hydrate grains would aggregate to eventually bridge 535 the pores and interlock the rock frame, as discussed earlier. For a given volume of hydrate, 536 smaller hydrate grains will form hydrate aggregates with more discontinuities than larger hydrate 537 grains, resulting in a smaller shear modulus. Hence, aggregation of smaller hydrate grains may 538 lead to a weaker effect on V<sub>s</sub>. As discussed earlier, hydrate dissociation can lead to more uniform

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539 distribution of methane gas and a reduction in bubble size. So it is possible that bubble size is

be lower in cycles 2 and 4 than in cycles 1 and 3. The patterns of change of  $V_s$  and  $V_p$  for first and

541 third cycles are similar, and the patterns are also similar between the second and fourth cycle

542 (Figure 8 a, b). As expected, the magnitude of changes is higher for lower differential pressures

543 (cycles 1 and 2).

544

545 While our experiments show transitions of the geophysical properties at specific hydrate 546 saturations in our experiments, it is likely that such transitions occur at different hydrate

547 saturations depending on sediment type and hydrate formation method. In Section 1, we

548 discussed the results of Priest et al., (2009) and Yun et al., (2005), which show similar transitions

549 at slightly different saturations. ( $S_h = 30\%$  - Priest et al., 2009;  $S_h = 40\%$  - Yun et al., 2005).

550

# 551 **3.3.2 Possible effect of water film on wave velocities**

552

553 The observation that  $V_p$  matches the pore-bridging HBES model but not  $V_s$ , likely due to the 554 presence of water films between hydrate and sand, adds another level of complexity to the effect 555 of hydrate morphology on elastic wave velocities. We no longer should view hydrate as pore-556 bridging in the sense of solid-solid contacts, as developed initially (e.g., Ecker et al., 2000; Priest 557 et al., 2009). We should account also for the presence of a water film between hydrate and 558 sediment surface as seen in Figure 6 and other studies (e.g., Bonnefoy et al., 2005; Chaouachi et 559 al., 2015; Sell et al., 2018; Tohidi et al., 2001). Gas hydrate bearing sediment should be viewed 560 as a three-phase system of interlocking solid hydrate and host grain frameworks separated by 561 water. Indeed, the presence of a water film between sand grains and hydrate is consistent with 562 the Leclaire et al. (1994) three phase Biot model adapted for hydrate by Guerin & Goldberg 563 (2005) and Carcione & Tinivella (2000). Best et al. (2013) found that this model gave 564 reasonable predictions of shear wave attenuation. This observation also implies that the hydrate cementing model concept may need to be revisited to include this water layer effect (e.g., 565 566 Chaouachi et al., 2015; Sell et al., 2018; Tohidi et al., 2001).

567

#### 568 3.3.3 Different maximum velocity in different hydrate formation cycles

570 We see higher maximum  $V_p$  and  $V_s$  for cycle 1 than for cycle 2, while those for cycle 3 and 4 are 571 similar (Figure 8). The percentage difference in maximum  $V_p$  between cycles 1 and 2 is about 572 double that of Vs. Different maximum velocities between cycles 1 and 2 can be explained by (i) 573 higher hydrate saturation in cycle 1 than in cycle 2, and/or (ii) reduction in bubble size in cycle 2. 574 In cycle 1, S<sub>h</sub> could not be calculated due to a malfunctioning pressure gauge, and in cycles 2, 3 and 4 the maximum  $S_h$  values were 23%, 26% and 25%, respectively. The HBES model shows 575 576 that a slight increase of pore-bridging hydrate equal to S<sub>h</sub> of 3 to 4% could account for the 577 observed discrepancies in maximum V<sub>p</sub> and V<sub>s</sub> between cycles 1 and 2 (Figure 9). A reduction in 578 bubble size can cause a similar effect. The latter mechanism is also consistent with a 579 redistribution of methane gas and a change in bubble size occurring during hydrate dissociation

580 and reformation, as discussed above.

- 581
- 582 **3.4 Effect of differential pressure**
- 583

584 The rate of change in V<sub>p</sub> and V<sub>s</sub> with S<sub>h</sub>, for S<sub>h</sub> of 10% to 15%, is much higher at a differential pressure of 10 MPa, than at 55 MPa (Figure 11). The onset of the rapid increase in  $V_p$  and  $V_s$ 585 586 occurs at a lower S<sub>h</sub> at 10 MPa than at 55 MPa. This behavior might be due to the presence of 587 microcracks at 10 MPa that are mostly closed at 55 MPa (e.g., Prasad and Manghnani, 1997). 588 When the microcracks are open, hydrate formation is likely to cause a much more dramatic 589 initial stiffening of the rock frame bulk moduli than when the microcracks are closed at higher 590 pressures, leading to a steeper increase in  $V_p$  and  $V_s$  because the frame moduli are initially 591 weaker. This effect is similar in magnitude to the normal velocity-pressure dependence reported 592 for Berea and similar sandstones with microcracks, where velocity increases more rapidly at 593 lower than at higher pressures (Eberhart-Phillips et al., 1989; Prasad & Manghnani, 1997). The effect of microcracks is evident as the initial (start of hydrate formation cycle)  $V_p$  and  $V_s$  for the 594 595 first and second cycles are lower than those for the third and fourth cycles (Figure 8). 596

597 4 Conclusions

598

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- 599 Estimates of hydrate content, and of hydrate related geohazards, from seismic data depend on our
- 600 understanding of the morphology and formation process of non-cementing hydrate in porous
- 601 media. From our experimental observations, we can conclude the following:

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Our time resolved/lapse (4D) SR-XCT images show that hydrate grows initially in a
 pore-floating morphology and transitions to a pore-bridging morphology. Then,
 eventually, it forms an inter-pore hydrate framework that interlocks with the sand grain
 framework, although separated by water films. To our knowledge, this is the first
 observation of such a methane hydrate morphology existing in a porous medium.

Construction
2. The SR-XCT images confirm the occurrence of a hydrate film around methane gas
bubbles, trapping gas inside, as the mechanism of co-existence of gas with hydrate under
hydrate stability conditions. SR-XCT images show that water films occur between
hydrate and sand when using methane and brine for hydrate formation. They also confirm
the previously inferred porous nature of hydrate. Using rock physics modeling, we were
able to link these morphological transitions to changes in the rate of increase of P- and Swave velocity with hydrate saturation.

The size of gas bubbles forming hydrate has a significant effect on velocities. The
presence of smaller gas bubbles can result in reaching maximum hydrate saturations
sooner than with large gas bubbles because there is an increase in surface reaction area.
Smaller gas bubbles also result in smaller hydrate grains, and when they aggregate the
number of discontinuities is larger than for larger hydrate grains, resulting in lower shear
modulus and velocity.

4. While P-wave velocities match the modeled velocity for pore-bridging hydrate, S-wave
velocities are higher than the pore-floating model and lower than the pore-bridging
model, likely due to presence of water films between hydrate and the rock frame. Both
ultrasonic velocities and imaging results indicate that hydrate-bearing sediment is a
system of interlocking solid hydrate and host grain frameworks separated by water films,
with isolated pockets of gas within the hydrate.

These observations are likely to be typical of natural hydrate-bearing sediments charged by gas from below. Such inter-pore hydrate framework and co-existing gas, if widespread in nature, should be considered when estimating *in situ* hydrate contents from elastic wave velocities.

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# 642 **References**

- Archer, D., Buffett, B., & Brovkin, V. (2009). Ocean methane hydrates as a slow tipping point in
  the global carbon cycle. *Proceedings of the National Academy of Sciences of the United States of America*, 106(49), 20596–601. https://doi.org/10.1073/pnas.0800885105
- Attias, E., Weitemeyer, K. A., Minshull, T. A., Best, A. I., Sinha, M., Jegen-Kulcsar, M., ...
  Berndt, C. (2016). Controlled-source electromagnetic and seismic delineation of subseafloor
  fluid flow structures in a gas hydrate province, offshore Norway. *Geophysical Journal*
- 649 *International*, 206(2), 1093–1110. https://doi.org/10.1093/gji/ggw188
- Best, A. I., Priest, J. A., Clayton, C. R. I., & Rees, E. V. L. (2013). The effect of methane hydrate
  morphology and water saturation on seismic wave attenuation in sand under shallow subseafloor conditions. *Earth and Planetary Science Letters*, *368*, 78–87.
  https://doi.org/10.1016/j.epsl.2013.02.033
- Best, A. I., Mccann, C., & Sothcott, J. (1994). The relationships between the velocities ,
  attenuations and petrophysical properties of reselvoi r sedimentary rocksl. *Geophysical Prospecting*, 151–178.
- Biot, M. A. (1956a). Theory of Propagation of Elastic Waves in a Fluid-Saturated Porous Solid.
  I. Low-Frequency Range. *The Journal of the Acoustical Society of America*, 28(2), 168–
  178. https://doi.org/10.1121/1.1908239
- Biot, M. A. (1956b). Theory of Propagation of Elastic Waves in a Fluid-Saturated Porous Solid.
  II. Higher Frequency Range. *The Journal of the Acoustical Society of America*, 28(2), 179–
  191. https://doi.org/10.1121/1.1908241
- Bonnefoy, O., Gruy, F., & Herri, J. M. (2005). Van der Waals interactions in systems involving
  gas hydrates. *Fluid Phase Equilibria*, 231(2), 176–187.
- 665 https://doi.org/10.1016/j.fluid.2005.02.004
- Boswell, R., & Collett, T. S. (2011). Current perspectives on gas hydrate resources. *Energy Environ. Sci.*, 4(4), 1206–1215. https://doi.org/10.1039/C0EE00203H
- Boswell, R., Shelander, D., Lee, M. W., Latham, T., Collett, T. S., Guerin, G., ... Goldberg, D.
  (2009). Occurrence of gas hydrate in Oligocene Frio sand: Alaminos Canyon Block 818:
  Northern Gulf of Mexico. *Marine and Petroleum Geology*, 26(8), 1499–1512.
  https://doi.org/10.1016/j.marpetgeo.2009.03.005
- Buchanan, P., Soper, A. K., Thompson, H., Westacott, R. E., Creek, J. L., Hobson, G., & Koh, C.
  A. (2005). Search for memory effects in methane hydrate: Structure of water before hydrate
  formation and after hydrate decomposition. *Journal of Chemical Physics*, *123*(16).
  https://doi.org/10.1063/1.2074927
- 676 Carcione, J. M., & Tinivella, U. (2000). Bottom-simulating reflectors: Seismic velocities and
   677 AVO effects. *Geophysics*, 65(1), 54–67. https://doi.org/10.1190/1.1444725
- Chand, S., Minshull, T. A., Priest, J. A., Best, A. I., Clayton, C. R. I., & Waite, W. F. (2006). An
  effective medium inversion algorithm for gas hydrate quantification and its application to
  laboratory and borehole measurements of gas hydrate-bearing sediments. *Geophysical Journal International*, *166*(2), 543–552. https://doi.org/10.1111/j.1365-246X.2006.03038.x
- 681 *Journal International*, 100(2), 345–352. https://doi.org/10.1111/j.1503-240X.2000.05058.2
   682 Chaouachi, M., Falenty, A., Sell, K., Enzmann, F., Kersten, M., Haberthür, D., & Kuhs, W. F.
   683 (2015). Microstructural evolution of gas hydrates in sedimentary matrices observed with
- 684 synchrotron X-ray computed tomographic microscopy. *Geochemistry, Geophysics,*
- 685 *Geosystems*, 16(6), 1711–1722. https://doi.org/10.1002/2015GC005811
- Collett, T. S. (2001). A Review of Well-Log Analysis Techniques Used to Assess Gas-Hydrate Bearing Reservoirs. In *Natural Gas Hydrates: Occurrence, Distribution, and Detection* (pp.

- 189–210). American Geophysical Union. https://doi.org/10.1029/GM124p0189
  Collett, T. S., Riedel, M., Boswell, R., Presley, J., Kumar, P., Sathe, A., ... Lall, M. V. (2015). *Indian National Gas Hydrate Program Expedition 01 report. Scientific Investigations Report.* Reston, VA. https://doi.org/10.3133/sir20125054
- 692 Cook, A. E., & Waite, W. F. (2018). Archie's Saturation Exponent for Natural Gas Hydrate in
   693 Coarse-Grained Reservoirs. *Journal of Geophysical Research: Solid Earth*.
   694 https://doi.org/10.1002/2017JB015138
- Dai, S., Santamarina, J. C., Waite, W. F., & Kneafsey, T. J. (2012). Hydrate morphology:
  Physical properties of sands with patchy hydrate saturation. *Journal of Geophysical Research*, *117*(B11), B11205. https://doi.org/10.1029/2012JB009667
- Daigle, H., Cook, A., & Malinverno, A. (2015). Permeability and porosity of hydrate-bearing
  sediments in the northern Gulf of Mexico. *Marine and Petroleum Geology*, 68, 551–564.
  https://doi.org/10.1016/j.marpetgeo.2015.10.004
- Doveton, J. H. (2001). All Models Are Wrong , but Some Models Are Useful : " Solving " the
   Simandoux Equation Prolog : The Archie Equation.
- Eberhart-Phillips, D., Han, D.-H., & Zoback, M. D. (1989). Empirical relationships among
   seismic velocity, effective pressure, porosity, and clay content in sandstone. *Geophysics*,
   54(1), 82–89. https://doi.org/10.1190/1.1442580
- Ecker, C., Dvorkin, J., & Nur, A. (1998). Sediments with gas hydrates: Internal structure from seismic AVO. *Geophysics*, 63(5), 1659–1669. https://doi.org/10.1190/1.1444462
- Ecker, C., Dvorkin, J., & Nur, A. (2000). Estimating the amount of hydrate and free gas from
  surface seismic. *Geophysics*, 65(2), 565–573. https://doi.org/10.1190/1.1820496
- Edwards, R. N. (1997). On the resource evaluation of marine gas hydrate deposits using sea-floor
  transient electric dipole-dipole methods. *Geophysics*, 62(1), 63–74.
  https://doi.org/10.1100/1.1444146
- 712 https://doi.org/10.1190/1.1444146
  713 Ellis, M. H. (2008). *Joint Seismic and Electrical Me*
- Ellis, M. H. (2008). Joint Seismic and Electrical Measurements of Gas Hydrates in Continental
   Margin Sediments. PhD thesis, University of Southampton. Retrieved from
   http://eprints.soton.ac.uk/id/eprint/63293
- Fitzgerald, R. (2000). Phase-Sensitive X-Ray Imaging. *Physics Today*, 53(7), 23–26.
   https://doi.org/10.1063/1.1292471
- Fujii, T., Suzuki, K., Takayama, T., Tamaki, M., Komatsu, Y., Konno, Y., ... Nagao, J. (2015).
  Geological setting and characterization of a methane hydrate reservoir distributed at the first offshore production test site on the Daini-Atsumi Knoll in the eastern Nankai Trough,
- Japan. Marine and Petroleum Geology, 66, 310–322.
- 722 https://doi.org/10.1016/j.marpetgeo.2015.02.037
- Gorman, A. R., Holbrook, W. S., Hornbach, M. J., Hackwith, K. L., Lizarralde, D., & Pecher, I.
  (2002). Migration of methane gas through the hydrate stability zone in a low-flux hydrate
  province. *Geology*, *30*(4), 327. https://doi.org/10.1130/0091-
- 726 7613(2002)030<0327:MOMGTT>2.0.CO;2
- Goswami, B. K., Weitemeyer, K. A., Minshull, T. A., Sinha, M. C., Westbrook, G. K., Chabert,
  A., ... Ker, S. (2015). A joint electromagnetic and seismic study of an active pockmark
  within the hydrate stability field at the Vestnesa Ridge, West Svalbard margin. *Journal of Coophysical Bacography Solid Earth*, 120(10), 6707, 6822
- 730 *Geophysical Research: Solid Earth*, 120(10), 6797–6822.
- 731 https://doi.org/10.1002/2015JB012344
- 732 Guerin, G., & Goldberg, D. (2005). Modeling of acoustic wave dissipation in gas hydrate-
- bearing sediments. *Geochemistry, Geophysics, Geosystems*, 6(7), 1–16.

734 https://doi.org/10.1029/2005GC000918

- Guerin, G., Goldberg, D., & Meltser, A. (1999). Characterization of in situ elastic properties of
   gas hydrate-bearing sediments on the Blake Ridge. *Journal of Geophysical Research: Solid Earth*, 104(B8), 17781–17795. https://doi.org/10.1029/1999JB900127
- Han, T., Best, A. I., Sothcott, J., North, L. J., & MacGregor, L. M. (2015). Relationships among
  low frequency (2Hz) electrical resistivity, porosity, clay content and permeability in
  reservoir sandstones. *Journal of Applied Geophysics*, *112*, 279–289.
- 741 https://doi.org/10.1016/j.jappgeo.2014.12.006
- Handa, Y. P. (1990). Effect of hydrostatic pressure and salinity on the stability of gas hydrates. *The Journal of Physical Chemistry*, 94(6), 2652–2657. https://doi.org/10.1021/j100369a077
- Helgerud, M. B., Dvorkin, J., Nur, A., Sakai, A., & Collett, T. S. (1999). Elastic-wave velocity in
  marine sediments with gas hydrates: Effective medium modeling. *Geophysical Research Letters*, 26(13), 2021–2024. https://doi.org/10.1029/1999GL900421
- Holland, M., Schultheiss, P., Roberts, J., & Druce, M. (2008). Observed Gas Hydrate
  Morphologies in Marine Sediments. In *6th International Conference on Gas Hydrates*.
- Hu, G., Ye, Y., Liu, C., Best, A. I., & Li, C. (2014). Gas Hydrate Distribution in Sediment Pore
  Space and Its Impact on Acoustic Properties of Hydrate-Bearing Sediments.
- Iassonov, P., Gebrenegus, T., & Tuller, M. (2009). Segmentation of X-ray computed tomography
  images of porous materials: A crucial step for characterization and quantitative analysis of
  pore structures. *Water Resources Research*, 45(9), 1–12.
  https://doi.org/10.1029/2009WR008087
- Jung, J. W., Espinoza, D. N., & Santamarina, J. C. (2010). Properties and phenomena relevant to
   CH 4 -CO 2 replacement in hydrate-bearing sediments. *Journal of Geophysical Research*,
   *115*(B10), B10102. https://doi.org/10.1029/2009JB000812
- Kerkar, P. B., Horvat, K., Jones, K. W., & Mahajan, D. (2014). Imaging methane hydrates
  growth dynamics in porous media using synchrotron X-ray computed tomography. *Geochemistry Geophysics Geosystems*, 15, 4759–4768.
- 761 https://doi.org/10.1002/2014GC005373
- Klapp, S. A., Enzmann, F., Walz, P., Huthwelker, T., Tuckermann, J., Schwarz, J. O., ... Brewer,
  P. G. (2012). Microstructure characteristics during hydrate formation and dissociation
  revealed by X-ray tomographic microscopy. *Geo-Marine Letters*, *32*(5–6), 555–562.
  https://doi.org/10.1007/s00367-012-0276-0
- Kleinberg, R. L., Flaum, C., Griffin, D. D., Brewer, P. G., Malby, G. E., Peltzer, E. T., &
  Yesinowski, J. P. (2003). Deep sea NMR: Methane hydrate growth habit in porous media
  and its relationship to hydraulic permeability, deposit accumulation, and submarine slope
  stability. *Journal of Geophysical Research: Solid Earth*, *108*(B10).
  https://doi.org/10.1029/2003JB002389
- Kneafsey, T. J., Tomutsa, L., Moridis, G. J., Seol, Y., Freifeld, B. M., Taylor, C. E., & Gupta, A.
  (2007). Methane hydrate formation and dissociation in a partially saturated core-scale sand
  sample. *Journal of Petroleum Science and Engineering*, 56(1–3), 108–126.
  https://doi.org/10.1016/j.petrol.2006.02.002
- Koeberl, C. (2002). High-resolution X-ray computed tomography of impactites. *Journal of Geophysical Research*, *107*, 1–9. https://doi.org/10.1029/2001JE001833
- Kumar, A., Sakpal, T., Roy, S., & Kumar, R. (2015). Methane hydrate formation in a test
   sediment of sand and clay at various levels of water saturation. *Canadian Journal of*
- 779 *Chemistry*, *93*(8), 874–881. https://doi.org/10.1139/cjc-2014-0537

780 Kvenvolden, K. A. (1993). Gas hydrates-geological perspective and global change. Reviews of 781 Geophysics, 31(2), 173. https://doi.org/10.1029/93RG00268 782 Leclaire, P., Cohen-Ténoudji, F., & Aguirre-Puente, J. (1994). Extension of Biot's theory of 783 wave propagation to frozen porous media. The Journal of the Acoustical Society of 784 America, 96(6), 3753-3768. https://doi.org/10.1121/1.411336 785 Lee, M. W. (2008). Models for Gas Hydrate-Bearing Sediments Inferred from Hydraulic 786 Permeability and Elastic Velocities. Retrieved from 787 https://pubs.usgs.gov/sir/2008/5219/pdf/SIR08-5219\_508.pdf 788 Lee, M. W., & Collett, T. S. (2006). Gas Hydrate and Free Gas Saturations Estimated from 789 Velocity Logs on Hydrate Ridge, offshore Oregon, U.S.A. In Proceedings of the Ocean 790 Drilling Program, 199 Scientific Results (Vol. 204, pp. 1–25). Ocean Drilling Program. 791 https://doi.org/10.2973/odp.proc.sr.204.103.2006 792 Leurer, K. C. (1997). Attenuation in fine-grained marine sediments: Extension of the Biot-Stoll 793 model by the "effective grain model" (EGM). GEOPHYSICS, 62(5), 1465-1479. 794 https://doi.org/10.1190/1.1444250 795 Leurer, K. C., & Brown, C. (2008). Acoustics of marine sediment under compaction: Binary 796 grain-size model and viscoelastic extension of Biot's theory. The Journal of the Acoustical 797 Society of America, 123(4), 1941–1951. https://doi.org/10.1121/1.2871839 798 Marín-Moreno, H., Sahoo, S. K., & Best, A. I. (2017). Theoretical modeling insights into elastic 799 wave attenuation mechanisms in marine sediments with pore-filling methane hydrate. 800 Journal of Geophysical Research: Solid Earth, 122(3), 1835–1847. 801 https://doi.org/10.1002/2016JB013577 802 Milkov, A. V., Dickens, G. R., Claypool, G. E., Lee, Y. J., Borowski, W. S., Torres, M. E., ... 803 Schultheiss, P. (2004). Co-existence of gas hydrate, free gas, and brine within the regional 804 gas hydrate stability zone at Hydrate Ridge (Oregon margin): Evidence from prolonged 805 degassing of a pressurized core. Earth and Planetary Science Letters, 222(3-4), 829-843. 806 https://doi.org/10.1016/j.epsl.2004.03.028 807 Millero, F. J., Chen, C.-T., Bradshaw, A., & Schleicher, K. (1980). A new high pressure equation 808 of state for seawater. Deep Sea Research Part A. Oceanographic Research Papers, 27(3–4), 809 255-264. https://doi.org/10.1016/0198-0149(80)90016-3 810 Minagawa, H., Nishikawa, Y., Ikeda, I., Miyazaki, K., Takahara, N., Sakamoto, Y., ... Nairta, H. 811 (2008). Characterization of sand sediment by pore size distribution and permeability using 812 proton nuclear magnetic resonance measurement. Journal of Geophysical Research: Solid 813 Earth, 113(7), 1-9. https://doi.org/10.1029/2007JB005403 814 Miyakawa, A., Saito, S., Yamada, Y., Tomaru, H., Kinoshita, M., & Tsuji, T. (2014). Gas 815 hydrate saturation at Site C0002, IODP Expeditions 314 and 315, in the Kumano Basin, 816 Nankai trough. Island Arc, 23(2), 142-156. https://doi.org/10.1111/iar.12064 817 North, L. J., Best, A. I., Sothcott, J., & MacGregor, L. (2013). Laboratory determination of the 818 full electrical resistivity tensor of heterogeneous carbonate rocks at elevated pressures. 819 Geophysical Prospecting, 61(2), 458-470. https://doi.org/10.1111/j.1365-820 2478.2012.01113.x 821 Nur, A., & Simmons, G. (1969). The effect of saturation on velocity in low porosity rocks. Earth 822 and Planetary Science Letters, 7(2), 183-193. https://doi.org/10.1016/0012-823 821X(69)90035-1 824 Ohmura, R., Ogawa, M., Yasuoka, K., & Mori, Y. H. (2003). Statistical Study of Clathrate-825 Hydrate Nucleation in a Water/Hydrochlorofluorocarbon System: Search for the Nature of

the "Memory Effect." The Journal of Physical Chemistry B, 107(22), 5289-5293. 826 827 https://doi.org/10.1021/jp027094e 828 Paganin, D., Mayo, S. C., Gureyev, T. E., Miller, P. R., & Wilkins, S. W. (2002). Simultaneous 829 phase and amplitude extraction from a single defocused image of a homogeneous object. 830 Journal of Microscopy, 206(1), 33–40. https://doi.org/10.1046/j.1365-2818.2002.01010.x 831 Phillips, D. H., & Lannutti, J. J. (1997). Measuring physical density with X-ray computed 832 tomography. NDT & E International, 30(6), 339–350. https://doi.org/10.1016/S0963-833 8695(97)00020-0 834 Prasad, M., & Manghnani, M. H. (1997). Effects of pore and differential pressure on 835 compressional wave velocity and quality factor in Berea and Michigan sandstones. 836 Geophysics, 62(4), 1163–1176. https://doi.org/10.1190/1.1444217 837 Priegnitz, M., Thaler, J., Spangenberg, E., Schicks, J. M., Schrötter, J., & Abendroth, S. (2015). 838 Characterizing electrical properties and permeability changes of hydrate bearing sediments 839 using ERT data. Geophysical Journal International, 202(3), 1599-1612. 840 https://doi.org/10.1093/gji/ggv245 841 Priest, J. A., Best, A. I., & Clayton, C. R. I. (2005). A laboratory investigation into the seismic 842 velocities of methane gas hydrate-bearing sand. Journal of Geophysical Research, 110(B4), 843 B04102. https://doi.org/10.1029/2004JB003259 844 Priest, J. A., Rees, E. V. L., & Clayton, C. R. I. (2009). Influence of gas hydrate morphology on 845 the seismic velocities of sands. Journal of Geophysical Research: Solid Earth, 114(B11), 846 B11205. https://doi.org/10.1029/2009JB006284 847 Reagan, M. T., & Moridis, G. J. (2008). Dynamic response of oceanic hydrate deposits to ocean 848 temperature change. Journal of Geophysical Research: Oceans, 113(12). 849 https://doi.org/10.1029/2008JC004938 850 Rees, E. V. L., Priest, J. A. J. A. J. A., & Clayton, C. R. I. (2011). The structure of methane gas 851 hydrate bearing sediments from the Krishna-Godavari Basin as seen from Micro-CT 852 scanning. Marine and Petroleum Geology, 28(7), 1283–1293. 853 https://doi.org/10.1016/j.marpetgeo.2011.03.015 854 Riedel, M., Collett, T. S., & Malone, M. (2010). Expedition 311 Synthesis: scientific findings. In 855 Proceedings of the IODP, 311 (Vol. 311). Integrated Ocean Drilling Program. 856 https://doi.org/10.2204/iodp.proc.311.213.2010 857 Rodger, P. M. (2006). Methane Hydrate: Melting and Memory. Annals of the New York Academy 858 of Sciences, 912(1), 474–482. https://doi.org/10.1111/j.1749-6632.2000.tb06802.x 859 Sahoo, S. K., Marín-Moreno, H., North, L. J., Falcon-Suarez, I., Madhusudhan, B. N., Best, A. 860 I., & Minshull, T. A. (2018). Presence and consequences of co-existing methane gas with 861 hydrate under two phase water-hydrate stability conditions. Journal of Geophysical 862 Research: Solid Earth. https://doi.org/10.1029/2018JB015598 863 Schicks, J. M., Naumann, R., Erzinger, J., Hester, K. C., Koh, C. A., & Sloan, E. D. (2006). 864 Phase transitions in mixed gas hydrates: Experimental observations versus calculated data. 865 Journal of Physical Chemistry B, 110(23), 11468–11474. https://doi.org/10.1021/jp0612580 866 Sell, K., Quintal, B., Kersten, M., & Saenger, E. H. (2018). Squirt flow due to interfacial water 867 films in hydrate bearing sediments. Solid Earth, 9(3), 699–711. https://doi.org/10.5194/se-9-699-2018 868 869 Setzmann, U., & Wagner, W. (1991). A New Equation of State and Tables of Thermodynamic 870 Properties for Methane Covering the Range from the Melting Line to 625 K at Pressures up 871 to 100 MPa. Journal of Physical and Chemical Reference Data, 20(6), 1061–1155.

872 https://doi.org/10.1063/1.555898

- Sloan, E. D., & Koh, C. A. (2007). *Clathrate hydrates of natural gases*. New York: CRC Press.
   Retrieved from https://www.crcpress.com/Clathrate-Hydrates-of-Natural-Gases-Third Edition/Sloan-Jr-Koh/p/book/9780849390784
- Smith, A. J., Flemings, P. B., Liu, X., & Darnell, K. (2014). The evolution of methane vents that
  pierce the hydrate stability zone in the world's oceans. *Journal of Geophysical Research B: Solid Earth*, *119*(8), 6337–6356. https://doi.org/10.1002/2013JB010686
- Spangenberg, E. (2001). Modeling of the influence of gas hydrate content on the electrical
  properties of porous sediments. *Journal of Geophysical Research: Solid Earth*, *106*(B4),
  6535–6548. https://doi.org/10.1029/2000JB900434
- Stampanoni, M., Borchert, G., Wyss, P., Abela, R., Patterson, B., Hunt, S., ... Rüegsegger, P.
  (2002). High resolution X-ray detector for synchrotron-based microtomography. *Nuclear Instruments and Methods in Physics Research Section A: Accelerators, Spectrometers, Detectors and Associated Equipment, 491*(1–2), 291–301. https://doi.org/10.1016/S01689002(02)01167-1
- Stern, L. A., Lorenson, T. D., & Pinkston, J. C. (2011). Gas hydrate characterization and grainscale imaging of recovered cores from the Mount Elbert Gas Hydrate Stratigraphic Test
  Well, Alaska North Slope. *Marine and Petroleum Geology*, 28(2), 394–403.
  https://doi.org/10.1016/j.marpetgeo.2009.08.003
- Suess, E., Torres, M. E., Bohrmann, G., Collier, R. W., Rickert, D., Goldfinger, C., ... Elver, M.
  (2001). Sea Floor Methane Hydrates at Hydrate Ridge, Cascadia Margin. In *Natural Gas Hydrates—Occurrence, Distribution and Detection* (pp. 87–98). American Geophysical
  Union. https://doi.org/10.1029/GM124p0087
- Sultan, N., Cochonat, P., Foucher, J.-P., & Mienert, J. (2004). Effect of gas hydrates melting on
  seafloor slope instability. *Marine Geology*, 213(1–4), 379–401.
  https://doi.org/10.1016/j.margac.2004.10.015
- 897 https://doi.org/10.1016/j.margeo.2004.10.015
- Sultan, N., Bohrmann, G., Ruffine, L., Pape, T., Riboulot, V., Colliat, J. L., ... Wei, J. (2014).
  Journal of Geophysical Research : Solid Earth. *Journal of Geophysical Research: Solid Earth*, *119*(December 2011), 2679–2694. https://doi.org/10.1029/2010JB007453.Pockmark
- Takeya, S., Hori, A., Hondoh, T., & Uchida, T. (2000). Freezing-memory effect of water on
  nucleation of CO<sub>2</sub> hydrate crystals. *The Journal of Physical Chemistry B*, 104(17), 4164–
  4168. https://doi.org/10.1021/jp993759+
- Tohidi, B., Agena, W. F., Clennell, M. Ben, Burgass, R. W., & Biderkab, A. B. (2001). Visual
  observation of gas-hydrate formation and dissociation in synthetic porous media by means
  of glass micromodels. *Geology*, 29(9), 867–870. https://doi.org/10.1130/00917613(2001)029
- Torres, M. E., Wallmann, K., Tréhu, A. M., Bohrmann, G., Borowski, W. S., & Tomaru, H.
  (2004). Gas hydrate growth, methane transport, and chloride enrichment at the southern
  summit of Hydrate Ridge, Cascadia margin off Oregon. *Earth and Planetary Science Letters*, 226(1–2), 225–241. https://doi.org/10.1016/j.epsl.2004.07.029
- Tulk, C. A. (1999). Storage and handling of natural gas hydrate. *Bulletin of the Geological Survey of Canada*, (544).
- 914 Uchida, T., Dallimore, S., & Mikami, J. (2000). Occurrences of Natural Gas Hydrates beneath
- 915 the Permafrost Zone in Mackenzie Delta: Visual and X-ray CT Imagery. *Annals of the New*
- 916 York Academy of Sciences, 912(1), 1021–1033. https://doi.org/10.1111/j.1749-
- 917 6632.2000.tb06857.x

- 918 Waite, W. F., Winters, W. J., & Mason, D. H. (2004). Methane hydrate formation in partially
- 919 water-satulated Ottawa sand. *American Mineralogist*, 89(July), 1202–1207.
  920 https://doi.org/10.2138/am-2004-8-906
- Waite, W. F., Santamarina, J. C., Cortes, D. D., Dugan, B., Espinoza, D. N., Germaine, J., ...
  Yun, T.-S. (2009). Physical properties of hydrate-bearing sediments. *Reviews of Geophysics*, 47(4), RG4003. https://doi.org/10.1029/2008RG000279
- 924 Yun, T. S., Francisca, F. M., Santamarina, J. C., & Ruppel, C. (2005). Compressional and shear
- wave velocities in uncemented sediment containing gas hydrate. *Geophysical Research Letters*, 32(10), L10609. https://doi.org/10.1029/2005GL022607
- 927

**Table 1**. Values used in the HBES model runs (Marín-Moreno et al., 2017). Marín-Moreno etal. (2017)

Parameter	Value	Units	Reference
Hydrate bulk modulus	$7.9 \mathrm{x10}^{9}$	Ра	(Best et al., 2013)
Hydrate shear modulus	$3.3 \times 10^9$	Pa	(Best et al., 2013)
Hydrate Poisson's ratio	0.32		
Methane bulk modulus	$K_{\rm CH4}(P_p,T)$	Pa	(Millero et al., 1980)
Methane density	$\rho_{\mathrm{CH4}}(P_p,T)$	kg m <sup>-3</sup>	(Millero et al., 1980)
Methane viscosity	$\mu_{\mathrm{CH4}}(P_p,T)$	Pa s	(Millero et al., 1980)
Methane irreducible saturation	0.02		(Reagan and Moridis, 2008)
Sand/Quartz grain bulk modulus	$36 \times 10^9$	Pa	(Ecker et al., 2000)
Sand/Quartz grain shear modulus	$45 \times 10^{9}$	Pa	(Ecker et al., 2000)
Sand/Quartz grain Poisson's ratio	0.062		
Sand/Quartz grain density	2650	kg m <sup>-3</sup>	(Ecker et al., 2000)
Sand/Quartz grain diameter	$1 \times 10^{-4}$	m	(Best et al., 2013)
Sand/Quartz grain coordination	85		(Eaker et al. 2000)
number	0.3		(Eckel et al., 2000)
Water bulk modulus	$K_{\rm W}(P_p,T)$	Pa	(Setzmann & Wagner, 1991)
Water density	$\rho_{\rm W}(P_p,T)$	kg m <sup>-3</sup>	(Setzmann & Wagner, 1991)
Water viscosity	$\mu_{\rm W}(P_p,T)$	Pa s	(Setzmann & Wagner, 1991)
Water irreducible saturation	0.2		(Reagan & Moridis, 2008)
Intrinsic permeability without	10 <sup>-13</sup>	$m^2$	(Datale et al. 2015)
hydrate	10	111	(Dargie et al., 2013)
Intrinsic permeability exponent for	3		
cementing hydrate	5		
Intrinsic permeability exponent for	2		
pore-filling hydrate			
Tortuosity	3		based on (Mavko et al., 2009)
van Genuchten's (1980) capillary	0.45		(Reagan & Moridis 2008)
pressure fitting parameter	0.15		(Reugun & Morrais, 2000)
van Genuchten's (1980) capillary	2000	Pa	(Reagan & Moridis 2008)
pressure gas entry parameter	2000	1 4	(Itelagan ee Morrano, 2000)
Critical porosity	0.36		(Mavko et al., 2009)
K Feldspar bulk modulus	$37.5 \times 10^9$	Pa	(Mavko et al., 2009)
K Feldspar shear modulus	15x10 <sup>9</sup>	Pa	(Mavko et al., 2009)
Illite bulk modulus	$62.21 \times 10^9$	Pa	(Mavko et al., 2009)
Illite shear modulus	$25.70 \text{ x} 10^9$	Pa	(Mavko et al., 2009)



Figure 1. Conceptual diagram showing different pore-fluid displacing hydrate morphologies. A: Contact cement, B: Grain coating cement, C: Pore-floating, D: Pore-bridging and E: Inter-pore hydrate framework.

933

**(a)** 

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Figure 2. Schematic representation of the experimental setup: a) ultrasonic rig; b) synchrotron rig.



Figure 3. Changes of a) pressure and temperature and b) pressure with time during methane hydrate formation and dissociation in the Berea sandstone. Only the second cycle of hydrate formation and dissociation is shown for clarity. The green line is the methane hydrate phase boundary for 35 g/L salinity, calculated using the approach of Tohidi et al. (1995). Blue dots represent cooling and red dots represent heating. In a) time is shown in hours (h). Trajectory ABC marks cooling of the system to 5 °C and hydrate formation. Trajectory CD shows hydrate dissociation.

(a) Time: 23h 30min

(b) Time: 45h 10min



(c)



Figure 4. 2D phase reconstructed CT slices from synchrotron imaging of the hydrate/sand sample after (a) 23hr 30m, (b) 45hr 10m. (c) Cross section through grey scale images indicating evolution of hydrate formation extracted at the same locations shown in (a) and (b) marked in yellow. Also shown are the grey levels of the four phases methane gas, methane hydrate, brine and sand obtained by density normalization.



Figure 5. Conceptual diagram showing different loss mechanisms considered in the Hydrate-Bearing Effective Sediment (HBES) model of Marín-Moreno et al. (2017) a) Biot's type global fluid flow, b) micro squirt flow c) sub-micro squirt flow due to inclusions of gas and water in hydrates and d) gas bubble resonance. Blue represents water, black is gas and white is hydrate. After Marín-Moreno et al., (2017).

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Figure 6. 3D Synchrotron radiation X-ray computed tomography images at different times during hydrate formation in sand. Red is gas, brown is sand, white is hydrate and blue is water. P1 and P2 marked in (a), (b) and (c) are two arbitrarily selected sand grains to aid visual comparison. Times are: 16hr 42 m (a and b), 23h 30m (c) and (d), 45h 10m (e) and (f). (a), (c) and (e) show all four phases while (b), (d) and (f) show only gas and hydrate for the same data volumes, respectively.





Figure 7. Changes in the samples' average areal distribution with depth of (a) hydrate and (b) gas at different times during hydrate formation in the X-ray CT analysis. Porosity distribution is also shown in (a) and (b). Dashed and dotted orange lines show the upper and lower error bounds for 16h 45m; errors bounds at other times are similar.



Figure 8. Changes in ultrasonic (648 kHz) P- and S-wave velocity ( $V_p$ ,  $V_s$ ) during hydrate formation (a, b) and dissociation (c, d). Differential pressure was 10 MPa for cycles 1 & 2, and 55 MPa for cycles 3 & 4. Although the cycles are continuous and sequential, zero time for a given cycle marks the beginning of cooling or heating for hydrate formation or dissociation, respectively.





Figure 9. Comparison of measured and modeled change in (a) P- and (b) S-wave velocity with hydrate formation. The experimental data is that of the third cycle of hydrate formation with a differential pressure of 55MPa. The modeled velocities were obtained using the HBES model (Marín-Moreno et al., 2017) with two bubble radii of  $2 \times 10^{-5}$  m and  $1 \times 10^{-5}$  m. The error in the experimental data is smaller than the symbol size.



Figure 10. Changes in electrical resistivity and (a) P- and (b) S-wave velocity with hydrate saturation during cycle 3. The error in the experimental data is smaller than the symbol size.



Figure 11. Changes in a) P- and b) S-wave velocity with hydrate saturation at differential pressures of 10 MPa (cycle 2) and 55 MPa (cycles 3 and 4). Saturations for cycle 1 are not shown because they could not be calculated due to a nonfunctional pressure gauge.