# Physical property changes in hydrate-bearing sediment due to depressurization and subsequent repressurization

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Abstract. Physical property measurements of sediment cores containing natural gas hydrate are typically performed on material exposed at least briefly to non-in situ conditions during recovery. To examine effects of a brief excursion from the gas-hydrate stability field, as can occur when pressure cores are transferred to pressurized storage vessels, we measured physical properties on laboratory-formed sand packs containing methane hydrate and methane pore gas. After depressurizing samples to atmospheric pressure, we repressurized them into the methane-hydrate stability field and remeasured their physical properties. Thermal conductivity, shear strength, acoustic compressional and shear wave amplitudes and speeds are compared between the original and depressurized/repressurized samples. X-ray computed tomography (CT) images track how the gas-hydrate distribution changes in the hydrate-cemented sands due to the depressurization/repressurization process. Because depressurization-induced property changes can be substantial and are not easily predicted, particularly in water-saturated, hydrate-bearing sediment, maintaining pressure and temperature conditions throughout the core recovery and measurement process is critical for using laboratory measurements to estimate in situ properties.

#### 1. Introduction

Pressures and temperatures in continental margin and permafrost sediments can stabilize gas hydrate, a crystalline solid in which hydrogen-bonded water molecules enclose individual guest molecules [Kvenvolden and Lorenson, 2001]. The most common guest molecule in naturally occurring gas hydrate is methane. Hydrate-bound methane represents a potential energy resource [Kerr, 2004; Ruppel, 2007], and may play

a role as a greenhouse gas in the global climate [Kvenvolden, 1993]. Sediment weakening caused by gas-hydrate dissociation has the potential to cause submarine slumps or slides, endangering seafloor infrastructure [Hovland and Gudmestad, 2001; Nixon and Grozic, 2007].

Interest in methane hydrates has motivated several large-scale field efforts to recover and study hydrate-bearing sediment from marine environments and beneath permafrost. Pressure coring systems represent a significant advance toward recovering pristine cores from which to infer *in-situ* physical property values. The Fugro and HYACE systems used aboard Ocean Drilling Program (ODP) Leg 204 [*Leg 204 Shipboard Scientific Party*, 2003] and International Ocean Drilling Program (IODP) Expedition 311 [*Expedition 311 Scientists*, 2006] are designed to retrieve and preserve meter-long core samples near their in situ hydrostatic stress. Although some physical property testing can be accomplished without ever releasing the hydrostatic stress [*Yun, et al.*, 2006], specialized measurements, such as triaxial shear-strength tests, are not yet possible without a brief hydrostatic stress release.

To accommodate these and other laboratory tests, pressure cores containing hydrate-bearing sediment have been rapidly subsectioned and transferred at one atmosphere to storage and transfer vessels, in which they are subsequently repressurized and stabilized with methane gas [Collett, et al., 2006; Expedition 311 Scientists, 2006]. This process takes approximately five minutes to complete (P. Schultheiss, personal communication, 2007).

Pressure cores containing hydrate-bearing sediment have also been subsectioned and transferred into storage and transfer vessels through ball valves, which allow the process

to be completed with almost no loss of hydrostatic pressure [Collett, et al., 2006; Winters, et al., 2005]. Although core manipulation and measurement systems can be designed and built to accommodate sample transfers, many existing laboratory systems have not been designed with this capability. For some measurements, pressurized transfer may never be possible. In such cases, samples must undergo a one-atmosphere transfer to the laboratory apparatus prior to repressurization and the measurement of physical properties. In this work, we focus primarily on effects from a single one-atmosphere transfer, though we also consider a case involving a second transfer, as would be required to transfer material from a storage vessel into some laboratory systems.

Rapidly venting a pressure core stored above 0°C to atmospheric pressure exposes the core to pressure well below the hydrate stability pressure, inducing hydrate dissociation and gas bubble formation. Repressurization with methane once the subsectioned sample is placed in a storage and transfer vessel stabilizes hydrate remaining in the sample, but also provides gas for additional hydrate formation. The net effect of such a transfer on the distribution of water, gas and hydrate within a core and the corresponding changes in the core's physical properties must be evaluated to determine whether measurements on disturbed core provide values representative of the undisturbed core.

To limit hydrate dissociation, cores have also been frozen prior to depressurization, or have been depressurized and subsequently stored in liquid nitrogen at atmospheric pressure. Though hydrate dissociates more slowly when depressurized below 0°C [Stern, et al., 2001], pore water expansion during the freezing process can generate a network of cracks that significantly disturb the sediment fabric. We discuss these effects in Section

5 with regard to recovering natural core material, but our laboratory work is focused on unfrozen material.

To examine the alteration of hydrate-bearing sediment that undergoes hydrate dissociation caused by sample depressurization above 0°C and subsequent hydrate formation following sample repressurization, we measured physical properties of methane hydrate-cemented sands formed in the laboratory. As discussed in Section 3, X-Ray computed tomography (CT) imagery, thermal conductivity, shear strength, acoustic compressional and shear wave speeds and amplitudes were measured first on undisturbed hydrate-cemented sand samples, and then again after the samples equilibrated following a brief exposure to atmospheric pressure. These results are presented in Section 4.

By construction, our laboratory samples were combinations of sand, methane hydrate, water and free methane gas, formed in the presence of free gas as discussed in Section 2. Naturally-occurring hydrate formed in the presence of free gas is thought to occur in the shallow sediments of actively venting regions such as the Cascadia margin [Bohrmann, et al., 1998], and at the base of the hydrate stability zone where gas can be recycled into the hydrate stability field [Guerin, et al., 1999; Yuan, et al., 1999].

The extent to which hydrate cements sediment grains is still debated, even with regard to studies of a single site [Guerin and Goldberg, 2005; Lee and Collett, 2005], but the strong intergranular bonding observed in our laboratory-formed samples is analogous to the bonding observed highly hydrate-saturated sands [Yun, et al., 2007]. Sands containing pore-space hydrate saturations exceeding 50% have been found in permafrost [Lee and Collett, 1999; Lee and Collett, 2005; Mount Elbert Science Team, 2007] and marine settings [Matsumoto, 2002; Smith, et al., 2006], and represent potential resource

targets. We discuss connections between our laboratory observations and the behavior of highly hydrate-saturated sands in Section 5.

In sediment with lower hydrate saturations, particularly in fine-grained sediment, a brief depressurization can alter sediment physical properties via gas bubble formation and effective stress loss [*Yun, et al.*, 2006] in addition to the hydrate redistribution we observe in our laboratory samples. These processes are discussed in Section 6.

## 2. Sample Preparation

We created samples containing quartz sand, methane hydrate and methane gas in the pore space. The hydrate-bearing sand samples were all formed according to a single procedure for the physical property measurements conducted at the U.S. Geological Survey facility in Massachusetts and the CT scanning experiments carried out at Lawrence Berkeley National Laboratory in California. Samples were not transferred between institutions.

Ottawa sand with a grain size range of 50-300 µm for the CT scanning and 250-500 µm for the physical property testing, was premixed with water, adding water in steps until the desired initial water content was reached. Samples were formed by tamping ~1cm lifts of the sand+water mixture into the measurement vessel. This packing process produced samples with ~38% porosity, as determined from the overall sample volume and mass of sand and water used. Sample-specific porosities and initial water saturations are given in Table 1. The choice of quartz grain size represents the on-site availability, not intrinsic requirements of either the CT or physical property measurement systems.

Samples were then pressurized with methane gas and cooled into the hydrate-stability field. Hydrate formation was allowed to continue until the formation rate became insignificant. Apparatus-specific measurements, described below, were used to determine when the sample was equilibrated, so the formation times differ between systems. Hydrate formed in this fashion within gas-rich, water-limited samples surrounds and cements sediment grains [Waite, et al., 2004].

The one-atmosphere transfer was simulated by depressurizing to one atmosphere, then repressurizing using methane gas. The sample was then allowed to re-equilibrate until the hydrate formation rate became insignificant. Timing for the depressurization and repressurization steps was guided by the five-minute transfer accomplished in the field. In practice however, depressurization and repressurization rates were limited by the complexity of the laboratory instrument used, and therefore varied from system to system as noted below.

## 3. Measurement Technique

Three separate systems were used to assess the effects of a brief depressurization on hydrate-bearing sands. Schematics for all three systems are given in Figure 1, with details of their use given below.

3.1. X-Ray Computed Tomography (CT): CT was used to map the initial hydrate distribution as well as the hydrate distribution within the cemented sand resulting from a simulated one-atmosphere transfer. A complete description of the CT imaging technique is given in Kneafsey et al. [2007]. To summarize, CT images were collected using a

modified Siemens HiQ medical X-ray computed tomography scanner (Fig. 1B), scanning 0.5 cm slices over the length of the sample (Fig. 1A). A CT image captures the sample density, with a resolution given by the voxel size. For scans shown here, the voxel size was 0.25 x 0.25 x 5 mm. The 5 cm diameter, 37 cm long cylindrical sample was jacketed in a rubber sleeve and surrounded by confining fluid in a pressure vessel. The pressure vessel itself was housed in a temperature-controlled PVC jacket and placed in an insulating box.

Methane hydrate was initially formed in sample CT 1 (see Table 1) at approximately 4°C and 5.2 MPa methane pore pressure. After 20 hours, the sample was scanned to obtain the initial gas-hydrate distribution. From mass balance, it was determined that ~67% of the initial pore water had converted to hydrate. The average pore-space hydrate saturation was 35 ± 4%, with hydrate saturations ranging from 26% near the sample perimeter to 46% near the sample's axis (Figure 2). The sample was then depressurized over the course of 10 minutes, held at atmospheric pressure for 6 minutes, then repressurized over a 10-minute period. The sample was out of the methane hydrate stability field for 19 minutes. CT scans were taken 1.66 and 24 hours after repressurization was completed. Measured density changes in the sample before and after the simulated one-atmosphere transfer reflect both the redistribution of methane hydrate and any additional hydrate growth from previously unreacted water.

3.2. Thermal Conductivity,  $\lambda$ : Thermal conductivity is a measure of how easily heat travels through a material. We measured  $\lambda$  using an axially-positioned probe in a 41-mm-diameter, 133 mm long cylindrical sample as described by Waite *et al.* [2006]. The

measurement vessel is shown schematically in Figure 1C. A resistive wire in the probe heats the sample, and  $\lambda$  is calculated from the temperature change measured by a thermistor in the probe. The uncertainty in our thermal conductivity measurements is conservatively estimated to be  $\pm 1\%$  [Waite, et al., 2006]. In a sand containing gas,  $\lambda$  is sensitive to the pore-space hydrate saturation [Waite, et al., 2002] and can be expected to change in response to a redistribution of hydrate around the thermistor.

Two pore-space hydrate saturations were tested for thermal conductivity changes during a simulated one-atmosphere core transfer. Characteristics of samples Thermal 1 and 2 are given in Table 1. Methane hydrate was formed at 5°C and 12 MPa methane pore pressure from initially partially water-saturated sand. The confining pressure balanced the pore pressure, meaning the samples experienced no effective stress. After 11 days, the measured  $\lambda$  showed no further change, and hydrate formation was deemed complete. Sample conditions were changed to 6.1°C and 12 MPa methane pore pressure and allowed to equilibrate overnight at the test conditions used in the acoustic and strength measurements described below.

The simulated one-atmosphere transfer was carried out by dropping the pore and confining pressures to atmospheric pressure over a 1-minute period. Three minutes later, the pore and confining pressures were increased to 12 MPa over 1 minute. To examine the effects of a second one-atmosphere transfer, as would occur if a core were transferred out of a storage vessel and into a laboratory system, a second depressurization/repressurization cycle was carried out, 7 days after the first simulated transfer.

3.3. Acoustic wave speed: Compressional and shear waves contain information about the rigidity of grain-to-grain contacts, which in turn is affected by the extent to which hydrate binds grains together [Helgerud, et al., 1999]. We used the Gas Hydrate And Sediment Test Laboratory Instrument (GHASTLI), described by Winters et al. [2000], to perform compressional and shear wave analysis on methane-hydrate-cemented sands containing two different pore-space hydrate saturations (GHASTLI 1, 1a, 2 and 2a in Table 1). As with the thermal property samples, the remaining pore space was filled with pressurized methane gas.

Cylindrical samples approximately 14 cm high and 7 cm in diameter were prepared as described above by tamping premixed, wet sand into a flexible liner, subsequently capped at each end by a metal endcap. Figure 1D shows a prepared sample, ready to be raised into the GHASTLI pressure chamber.

The endcaps house axially-positioned, 1 MHz compressional and shear wave transducers. Acoustic waves were produced in the top endcap and were detected by the transducer crystal in the bottom endcap. Wave speed was calculated by dividing the acoustic travel time through the sample by the sample length, as measured using a linear voltage displacement transducer. The acoustic wave speed uncertainty is  $\pm 0.5\%$  for compressional wave speeds,  $V_p$  [Waite, et al., 2004]. Because of uncertainties in choosing the shear wave arrival, the uncertainty in the shear wave speed,  $V_s$ , is conservatively estimated to be  $\pm 3\%$ .

In the GHASTLI pressure vessel, the sample's methane pore pressure, controlled by a syringe pump connected through a line in the endcap, was held at 12 MPa while the temperature was lowered from room temperature and held at 6.1°C. The independently-

controlled confining pressure was held via syringe pump at 12.25 MPa to impart an effective pressure of 250 kPa. As hydrate formed in the sample, the acoustic signal increased in amplitude and the measured wave speed increased [Waite, et al., 2004]. Hydrate formation was deemed complete when the acoustic amplitude and measured wave speed stabilized, 5 days after onset of hydrate formation for the low-saturation sample, 14 days for the higher-saturation sample.

To simulate a one-atmosphere core transfer, we reduced the pore pressure and confining pressure together over the course of 3 minutes, held them at atmospheric pressure for 1.5 minutes, and repressurized them over the course of 5.5 minutes. The samples were out of the methane-hydrate stability field for 7.5 minutes. They were then allowed to equilibrate at the original formation conditions until the acoustic signal restabilized, which took 2 days for the low hydrate saturation sample, and 3 days for the high hydrate saturation sample.

3.4. Shear Strength: Triaxial shear-strength measurements provide a means of assessing sediment stability. In hydrate-bearing sediment, strength loss resulting from hydrate dissociation can cause localized failures in drilling and seafloor pipeline applications [Hovland and Gudmestad, 2001; Moridis and Kowalsky, 2006], or regional-scale slope failure [Paull, et al., 2000]. Undrained shear strength was measured by using a ram to axially strain the sample while the confining pressure, pore pressure, and load supported by the sample were recorded. The ram extended at a constant 5.6 mm/hour, equivalent to an initial strain rate of 4.4 %/hour. We ran the shear-strength test in the

undrained configuration, meaning the pore space was isolated and the pore pressure changed as the sample length decreased.

Unlike the other property measurements described here, shear-strength measurements are destructive and therefore cannot be carried out on a single sample both before and after a simulated one-atmosphere transfer. We estimate the effect of a one-atmosphere transfer on shear strength by comparing the shear strength for samples GHASTLI 1 and 2, which undergo the transfer procedure, with nominally identical samples that do not undergo the simulated one-atmosphere transfer (GHASTLI 1a and 2a in Table 1). Shear-strength measurement uncertainties in hydrate-bearing sediment are difficult to quantify because shear strength depends sensitively on how sediment grains are packed, the extent to which they are interlocked, and their degree of cementation [Bowles, 1979] Difficulties in precisely replicating the distribution and sediment bonding by methane hydrate increases sample-to-sample variations in shear strength. Extensive work by Hyodo et al. [2005] on laboratory-formed mixtures of methane hydrate and sediment suggests uncertainties of ±15% are to be expected.

## 4. Laboratory results for gas-rich, methane hydrate-bearing sediment

In a one-atmosphere transfer, methane-hydrate-cemented sediment first undergoes a depressurization-induced dissociation that can disrupt or destroy the sediment fabric and radically alter a range of physical properties [Francisca, et al., 2005]. Subsequent hydrate formation following repressurization will not likely restore all of the sample's physical properties, particularly if the initial and final hydrate distributions differ.

Here we look at the net physical property changes resulting from a brief pressure excursion to one atmosphere. Conclusions are drawn by comparing measurements on a given sample before and after a brief depressurization, meaning sample-to-sample variations are not responsible for the observed property changes. The exception is the undrained shear strength,  $S_u$ , which as noted previously, requires a destructive test of two samples to evaluate the effects of a brief depressurization. Sample-to-sample variability increases the  $S_u$  analysis uncertainty relative to the other measured properties (see Table 2). Following the discussion of property changes resulting from a brief depressurization, we comment in Section 4.5 on the dependence of the sample's initial physical properties on hydrate saturation.

4.1. X-Ray Computed Tomography (CT): Density changes resulting from the simulated core retrieval process were observed in CT images taken 1.66 and ~24 hours after briefly depressurizing the sample. Figure 3A is a cross section of sample C1 showing the average density change 1.66 hr after the simulated core transfer, relative to the sample density after the initial hydrate formation. Figure 3B shows the average density change measured ~24 hr after the depressurization/repressurization procedure, relative to the sample density after the initial hydrate formation. The density changes are small enough that, in spite of the constant level of measurement noise, speckling is more apparent in the density difference profiles shown in Figure 3 than in the sample density profile shown in Figure 2.

Figure 3 shows the average over 56 of the 78 scanned sections of the sample. The remaining 22 sections contained serious scanning artifacts that rendered them unusable in

this study. In any given slice, measurement noise tends to obscure the hydrate-redistribution pattern, but a sensitivity analysis shows that density changes exceeding 0.01 g/cm<sup>3</sup> observed in regions of 0.04 cm<sup>2</sup> in the averaged cross section have nearly a 90% level of confidence and can be considered real. The white holes in the center top of the cross-section are from thermocouples embedded in the sample. The speckled pattern on the left side of the figures results from averaging scans down the length of a sample with a slightly non-uniform diameter.

As discussed in Section 4.5, capillary action can cause water to migrate during the initial hydrate formation, leading to the radial inhomogeneity observed in Figure 1 prior to the brief depressurization. The initial density distribution is not responsible for the density change pattern resulting from the brief depressurization, however. Because the density change profiles shown in Figure 3 are the density difference between the post-and pre-depressurization densities, the initial sample's inhomogeneities are removed. Both the density increase near the sample's central axis, and the observed decrease at the sample's perimeter represent sample inhomogeneity caused by the brief depressurization.

A density increase near the sample's central axis, combined with a density decrease observed at the sample perimeter, can be explained by a net inward migration of material. We propose the hydrate redistribution process illustrated in Figure 4 accounts for much of the observed density change. Figure 4A represents a cross section of an essentially uniform distribution of sand grains, methane hydrate and methane gas. Depressurizing the sample destabilizes the hydrate, which begins dissociating to methane gas and water (Fig. 4B). Hydrate dissociation is endothermic, and because the surrounding bath provides the most readily available heat source to fuel dissociation, dissociation initially

occurs most efficiently on the outer sample perimeter. Time-lapse CT imagery during hydrate dissociation shows the dissociation front propagates toward the central axis of the sample over time [*Kneafsey, et al.*, 2007].

The sample is repressurized with methane gas before the dissociation front reaches the sample's central axis. Hydrate persisting near the sample's central axis forms a surface for rapid growth of new hydrate following the repressurization step [Osegovic, et al., 2006]. Capillary forces draw water to the hydrate formation front [Gupta, et al., 2006; Kneafsey, et al., 2007] (Fig. 4C), reducing the water available for hydrate formation near the sample perimeter. Overall, the process increases the pore-space hydrate saturation near the sample's central axis, while reducing the hydrate saturation near the sample's perimeter (Figs. 3 and 4D).

Two additional hydrate formation processes contribute to the density increase near the sample's central axis, though as shown below, neither process can account for the density decrease at the sample perimeter. The two processes are: 1) new hydrate formation from water that had not yet formed hydrate just prior to the brief depressurization, and 2) hydrate formation from water that does not migrate during the brief depressurization.

Prior to the brief depressurization of sample C1, approximately 67% of the initial water was thought to have been converted to hydrate. Depressurization can crack hydrate layers that isolated portions of the remaining initial water volume from the surrounding methane gas [Kneafsey, et al., 2007]. Subsequent repressurization exposes methane gas to that previously unreacted water, allowing new hydrate growth. This growth is in addition to growth from water liberated via hydrate dissociation during the brief depressurization. The volume increase associated with converting a volume of water to

gas hydrate in sands at these pressures means low-density methane pore gas is replaced by higher-density hydrate, resulting in a net density increase.

Because the water/methane interface is an active hydrate growth zone, a portion of the available water likely forms hydrate in place, without first migrating toward the sample's central axis. Hydrate forming in place from water liberated during dissociation returns the local density to pre-depressurization levels, resulting in no net density change. Hydrate forming in place from previously unreacted water increases the sample density everywhere it occurs. The relative contribution of this mechanism to the observed density increase near the sample's central axis is not known, but this mechanism has been observed in CT scans of similar samples in which the pore gas pressure was changed, but remained within the methane-hydrate stability field [Kneafsey, et al., 2007]. Without water migration resulting from the brief depressurization, however, hydrate formation following the brief depressurization cannot cause the density decrease noted at the sample's perimeter in Figure 3.

We attribute the observed density changes induced by a brief depressurization to a combination of hydrate growth from water that has migrated toward the sample's central axis, in place hydrate re-growth from water liberated from hydrate during the depressurization, and new hydrate growth from previously unreacted water.

As we show in Sections 4.1-4.5, physical property changes resulting from a brief depressurization of gas-rich, hydrate-bearing sand vary with the initial hydrate content, the amount of unreacted water prior to depressurization, and pore-size dependent capillary forces. We therefore use the CT results to visualize hydrate redistribution, but

not to predict the magnitude of physical property changes occurring in other gas-rich, hydrate-bearing sands as a result of hydrate redistribution.

4.2. Effect on thermal conductivity measurements: Because we measure  $\lambda$  using an axially positioned probe, our measurement is most sensitive to changes occurring near the sample's central axis. The hydrate redistribution observed in the CT scans should increase the measured  $\lambda$  for two reasons: (1)  $\lambda$  of methane hydrate is ~0.62 W/m·K [Waite, et al., 2007], more than an order of magnitude larger than that of the methane gas (0.045 W/m·K [Vargaftik, et al., 1993]) being replaced by hydrate; (2) methane hydrate provides a more efficient thermal bridge between the high-thermal-conductivity sand grains than does methane gas [Waite, et al., 2002].

The 5.4% increase in thermal conductivity for a pore-space hydrate saturation,  $S_h$ , of ~20% is significantly less than the 13.8% increase observed for the  $S_h \approx 34\%$  case (Table 2). The dependence on initial hydrate saturation likely reflects differences not only in the extent of the hydrate migration toward the sample's central axis, but also in the amount of new hydrate growth from pore water that became isolated from the methane gas and did not form hydrate during the initial hydrate synthesis.

A second one-atmosphere transfer was carried out on both samples, causing an additional increase in  $\lambda$ . The effect was smaller than for the first cycle, resulting in an additional 3.6% increase for the low-hydrate-saturation sample and an additional 2.5% increase for the higher-hydrate-saturation sample. Hydrate redistribution following the first depressurization cycle reduces the hydrate saturation in the outer portions of the sample. Reducing the volume of hydrate near the outer surface of the sample, where

hydrate is most susceptible to alteration from a brief depressurization, mutes the impact of a subsequent brief depressurization on the sample's physical properties.

The impact of a one-atmosphere transfer on  $\lambda$  depends on the location at which  $\lambda$  is measured. If we had adopted a guarded-plate measurement scheme in which the sensor is placed on the sample surface rather than the interior [Cook and Leaist, 1983; Taylor, et al., 2007], our measurement sensitivity would have been highest in the zone of hydrate depletion. We would therefore have expected a decrease rather than increase in  $\lambda$ . The physical property variation with measurement location in a sample further complicates the estimation of in situ properties from property measurements of disturbed core material.

4.3. Effect on acoustic wave speed measurements: As with thermal conductivity measurements, our acoustic measurements are most sensitive to changes occurring along the sample's central axis because of the central location of our transducers in their endcaps. Increasing the hydrate saturation along the sample's axis is expected to more effectively cement sand grains, stiffen the sample, and thereby increase the measured wave speed [Dvorkin, et al., 2000] in the central portion of the sample where the transducers are most sensitive. As the sample stiffens with increasing hydrate cementation of the sand grains, acoustic transmission should be more efficient, increasing the observed acoustic amplitude [Waite, et al., 2004].

Figure 5 compares the acoustic signal produced by the shear wave crystals before and after the brief excursion from the hydrate stability field. Results from the higher of the two hydrate saturations tested are shown, but both samples display the same basic

waveform changes: (1) acoustic travel times generally decrease, meaning the associated wave speeds increase; (2) acoustic amplitudes generally increase, by 66% in the case of the shear wave shown in Figure 5.

As noted in the CT results, hydrate forms rapidly following repressurization. Even just 3 hours after repressurization, the waveform amplitude surpassed the predepressurization amplitude (blue curve in Fig. 5). The waveform required only 3 days to fully develop and equilibrate following repressurization (red curve in Fig. 5), as compared to the 14 days required for the initial waveform to grow and stabilize.

As shown in Table 2, the P-wave speed,  $V_p$ , for the GHASTLI 2 sample decreased by 2.4%. The correspondingly large 47% increase in shear wave speed,  $V_s$ , suggests hydrate redistribution and new hydrate growth significantly increased the grain-to-grain stiffness along the central axis of the sample. In the presence of free gas this stiffness increase could indicate enhanced intergranular cementation, but the corresponding P- wave speed and amplitude increase [*Lee*, 2004] is not observed. An enhanced S-wave speed response coupled with a muted P-wave speed response to an increased hydrate saturation has been observed in laboratory studies of THF hydrate-bearing sand [*Yun*, *et al.*, 2005]. Yun et al. [2005] show how such a response can be modeled assuming hydrate forms initially on the surface of grains, with subsequent growth into the pore space rather than at grain contacts. As discussed in Section 5, an insensitivity of  $V_p$  to a brief depressurization has also been observed in naturally-occurring, highly-cemented sands [*Winters*, *et al.*, 1999].

4.4. Effect on shear strength measurements: The hydrate saturation increase along the sample axis resulting from a brief depressurization has the potential to stiffen the sample

by strengthening bonds between sand grains, but that strengthening comes at the expense of cement lost from between grains near the sample perimeter during the brief depressurization. Despite changes in hydrate distribution, the average equilibrium concentration of gas hydrate within any cross section may therefore be similar before and after a brief depressurization.

Figure 6 compares the shear-strength response to axial strain in four different samples. The simulated one-atmosphere transfer appears to reduce the peak shear stress each sample can support, but even the peak stress decrease of 18% for the  $S_h \approx 20\%$  case is within the expected sample-to-sample variation of  $\pm 15\%$ .

Other shear characteristics do not correlate with the effects of hydrate redistribution. Prior to failure, for instance, the slope of shear stress versus strain is nearly independent of hydrate saturation, in agreement with measurements on methane-hydrate-bearing sediment by Hyodo et al. [2005], and that slope does not change dramatically as a result of hydrate redistribution. The shear-stress decrease following the peak stress also does not correlate with the hydrate redistribution, becoming more pronounced for the  $S_h \approx 20\%$  case, but less pronounced for the  $S_h \approx 43\%$  case. Based on our study, the shear-strength measurement changes are not significantly correlated with effects of hydrate redistribution in cemented, gas-rich sand samples. This lack of correlation complicates the assessment of in situ shear properties based on measurements of briefly depressurized samples.

4.5. Hydrate formation controls on the initial physical property measurements: In the same fashion that a brief depressurization induces water migration via capillary action

within a sample, capillary action causes water to migrate within a sample during the initial hydrate formation procedure [Kneafsey, et al., 2007]. As shown in Figure 2, an initially uniform sample of partially-water-saturated sand can form a slightly inhomogeneous hydrate-bearing sand. While it is possible that our higher-hydrate-saturation samples exhibit lower thermal conductivity and shear wave speed than their lower-hydrate-saturation counterparts because of differences in their formation-induced inhomogeneities, the inhomogeneities have not yet been quantified in terms of the extent to which they depend on initial water saturation, hydrate formation rate, and other factors.

# 5. Implications for highly hydrate-saturated sand

The weak effect a brief depressurization has on  $V_p$  in our laboratory-formed, hydrate-bearing sands has also been observed in a hydrate-bearing sand sample recovered beneath permafrost at the Mallik 2L-38 site in the MacKenzie Delta region of the Canadian Northwest Territories [Winters, et al., 1999]. In spite of obvious differences in sample character and handling, the Mallik sample provides a link between our laboratory results and expectations for highly hydrate-saturated sands.

The Mallik 2L-38 core contained pore space hydrate saturations of 70-74% [*Lee and Collett*, 1999]. Collected as a conventional core, the sample was depressurized during its recovery from depth. The sample also froze during retrieval through the overlying permafrost. The sample was stored in a methane-pressurized vessel and kept frozen during transport to the GHASTLI facility. The sample was then depressurized briefly below the freezing temperature before being transferred into GHASTLI at room temperature.

The core's transfer into GHASTLI is similar to the brief depressurization experienced by our laboratory-formed, hydrate-bearing sands. The Mallik sample was depressurized while it was frozen, which limits the hydrate dissociation rate [Stern, et al., 2001]. The sample partially thawed during the room-temperature transition into GHASTLI, however [Winters, et al., 1999]. Similar to Fig. 4, the sample was most susceptible to dissociation near its cylindrical surface prior to being repressurized into the methane hydrate stability field. In spite of the potential for sample disturbance during retrieval, transport and transfer into GHASTLI,  $V_p$  for the Mallik 2L-38 core measured in GHASTLI was in agreement with downhole logging results [Winters, et al., 1999].

A key similarity between our laboratory-formed, hydrate-cemented sands and the Mallik sample is the extent to which methane hydrate bonds sediment grains. Naturally occurring hydrate in sand is generally thought to be a load-bearing sediment component [Kleinberg, et al., 2005], rather than providing the strong intergranular bonding seen in out hydrate-cemented laboratory samples. Nevertheless, in laboratory studies of noncementing THF hydrate sediments with pore-space hydrate contents exceeding 50% appear to maintain intergranular bonding by hydrate during a reduction in effective stress, limiting the sample alteration during sample recovery as long as the sediment is returned to its in situ effective stress state prior to measuring the acoustic wave speeds [Lee, et al., 2008]. Pore space hydrate saturations exceeding 50% in sands have been found not only at the Mallik site, but also beneath permafrost at the Mt. Elbert site on the Alaskan North Slope [Mount Elbert Science Team, 2007], as well as in marine sands in the Nankai Trough offshore Japan [Matsumoto, 2002], and at the Tigershark well in the Gulf of Mexico [Smith, et al., 2006].

Our laboratory technique, forming methane hydrate from partially water-saturated sands, cements sand grains with hydrate, providing strong intergranular bonding. Following the primary hydrate formation stage,  $V_p$  in the 43% hydrate saturation sample is 3430 m/s (see Table 2), more than triple the value for the partially-water-saturated sand pack prior to hydrate formation [Waite, et al., 2004]. Even in even our 20% hydrate saturation sample,  $V_p$ , changes by less than 5% in response to a brief depressurization. This resilience in  $V_p$  is not expected in non-cementing hydrate cases for all hydrate saturation levels, however. For pore-space hydrate contents below ~40%, intergranular bonding by hydrate is less significant and effective stress changes can alter the sample [Yun, et al., 2007].

An important difference between our laboratory samples and the Mallik 2L-38 core is the temperature during depressurization. Our laboratory samples were above freezing prior to depressurization, while the Mallik core was frozen. Depressurization below the freezing point preserves hydrate by lowering the dissociation rate [*Stern, et al.*, 2001], but freezing also causes interstitial pore water to expand, potentially fracturing the sediment.

In highly hydrate-saturated sands such as the Mallik 2L-38 sample, consistent  $V_p$  results from laboratory and downhole logging measurements [Winters, et al., 1999] imply pore water volumes were low enough that pore-water freezing did not significantly alter the sample. Prior to depressurization, freezing high hydrate-content sands even from marine environments may preserve hydrate without significantly altering the host sediment fabric. Lower hydrate-content sediments, however, may contain enough water that expansion due to pore-water freezing does significantly disturb the host sediment fabric.

# 6. Implications for water-saturated, methane hydrate-bearing sediment

A critical difference between the gas-rich, hydrate-bearing sand samples discussed above and water-saturated hydrate-bearing core recovered in the field is the extent to which gas produced during a brief depressurization escapes the sample. Our gas-rich, hydrate-bearing sands were permeable, containing gas-filled pathways through which gas produced via hydrate dissociation could escape the sample during depressurization. In water-saturated sediment, particularly fine-grained sediment, such vent pathways are restricted. These restrictions lead to bubble formation and effective stress reduction [Yun, et al., 2006], both of which are exacerbated by methane hydrate dissociation. As discussed below, bubble formation and effective stress reduction, combined with hydrate redistribution, can lead to physical property changes that vary from sample to sample and are therefore difficult to correct for.

6.1. Bubble formation: Depressurizing water-saturated, hydrate-bearing sediment draws gas dissolved in the pore water out of solution in addition to producing gas via hydrate dissociation. Repressurization can force gas back into the dissolved phase, but the maximum quantity of dissolved gas is limited by the gas's solubility in water. In a sample containing gas produced from hydrate dissociation, the solubility limit can be exceeded, allowing hydrate-forming gas to remain in the gas phase immediately following repressurization. The persistence of gas bubbles following pressurization can lead to hydrate formation, because the bubble's gas-water interface provides an efficient hydrate growth surface [Freer, et al., 2001]. Because bubble location and volume depend

on the sample's lithology, pore-water methane saturation, and hydrate content, hydrate formed from bubbles will be distributed according to those sample-dependent characteristics. As shown in our laboratory samples, hydrate redistribution can alter the sample's physical properties.

Bubble growth itself can alter a sample's physical properties, attenuating compressional waves and reducing the measured wave speed relative to *in situ* values. The magnitude of the change depends on the extent of bubble formation, which itself depends on the sample's lithology, pore-water methane saturation, and hydrate content. Although hydrate dissociation can significantly increase the amount of free gas produced as a result of core depressurization, bubble growth is a concern for any water-saturated core with dissolved gas, regardless of whether hydrate is present.

By maintaining near *in-situ* hydrostatic pressure throughout the core-acquisition and property-measurement process, bubble formation and hydrate redistribution can be curtailed. Compressional wave-speed measurements made using the Instrumented Pressure Testing Chamber (IPTC) on clay-rich Gulf of Mexico sediment recovered and measured while maintaining near *in-situ* pressure are only 2% below the logging while drilling (LWD) results [*Yun, et al.*, 2006].

6.2. Loss of effective stress: Although current pressure-coring technology is able to preserve near-in situ hydrostatic pressure, the effective stress pushing sediment grains against each other is not maintained. Particularly in fine-grained sediment, bubble formation caused by hydrate dissociation can increase the pore pressure, further reducing the effective stress [Xu and Germanovich, 2006], weakening grain-to-grain contacts and

reducing the sediment frame stiffness [Francisca, et al., 2005]. As a result, the compressional and shear wave speeds as well as the sediment strength are all reduced [Yun, et al., 2006]. As described above, bubble formation varies from sample to sample, complicating efforts to predict or account for the ensuing degradation of the matrix stiffness. By maintaining the hydrostatic pressure, however, pressure coring can avoid the additional matrix weakening caused by bubble formation and expansion.

## 7. Conclusion

We have shown that in gas-rich, laboratory-formed methane-hydrate-cemented sands, even a brief depressurization to one atmosphere induces hydrate redistribution within the sample. This redistribution creates a sample with new physical properties that can differ significantly from those of the original sample. Measured changes depend on whether a given property is measured near the sample perimeter, which loses hydrate during the simulated core transfer, or near the sample center, which gains hydrate. Briefly dropping the pressure to one atmosphere a second time, to simulate the transfer of core from a storage vessel into a laboratory measurement apparatus, leads to continued hydrate redistribution, though to a lesser extent than the initial simulated transfer.

Estimating in situ properties from initially water-saturated recovered core that has experienced a depressurization cycle is further complicated by bubble formation and loss of effective stress, with hydrate dissociation exacerbating both processes. Property changes therefore vary not only based on which portion of the sample is being measured, but on sample-to-sample differences in initial hydrate saturation and lithology.

Unpredictable location- and sample-dependent property changes complicate the assessment of *in situ* properties based on recovered core material. Accurate *in situ* property estimates from recovered cores require careful preservation of the *in situ* stress state throughout the recovery and measurement process, even in non hydrate-bearing material.

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**Table 1.** Porosity,  $\phi$ , and pore-space hydrate saturation,  $S_h$ , of hydrate-bearing Ottawa sands in this study prior to the brief depressurization. Remaining pore space contains pressurized methane gas. Full conversion of pore water to methane hydrate is assumed for all samples except CT 1, which contains a 14.5% porespace saturation of residual water in addition to methane hydrate. GHASTLI samples 1a and 2a were not subjected to a brief depressurization. Uncertainties are  $\pm 4\%$ .

Sample	φ(%)	$S_h$ (%)	
CT 1	38	35	
Thermal 1	37	20	
Thermal 2	42	34	
GHASTLI 1	37	20	
GHASTLI 1a	38	19	
GHASTLI 2, 2a	37	43	

**Table 2.** Physical property changes in methane-hydrate-cemented sands resulting from a brief depressurization to one atmosphere, followed by a repressurization into the methane hydrate stability field. Measured properties include thermal conductivity,  $\lambda$ , compressional and shear wave speeds,  $V_p$  and  $V_s$  respectively, and triaxial undrained shear strength,  $S_u$ .

Property	Porosity (%)	$S_h$ (%)	Initial Value	Final Value	Change (%)	Uncertainty(%)
λ	37	20	1.67 W/m·K	1.76 W/m·K	5.4	±1
λ	42	34	$1.60 \text{ W/m} \cdot \text{K}$	1.82 W/m·K	13.8	±1
$V_p$	37	20	2830 m/s	2960 m/s	4.5	±0.5
$\hat{V_p}$	37	43	3430 m/s	3350 m/s	-2.4	±0.5
$V_s$	37	20	1580 m/s	1770 m/s	11.7	±3
$V_s$	37	43	1320 m/s	1945 m/s	47.0	±3
$S_u$	37/38†	20/19†	2.31 MPa	1.89 MPa	-18	±15
$S_u$	37	43	2.94 MPa	2.81 MPa	-4.5	±15

<sup>†</sup>The two samples required for undrained shear strength differed slightly in their initial porosity and hydrate saturation,  $S_h$  (see Table 1).

## **Figure Captions**

- Figure 1. A: Schematic of the X-Ray CT sample vessel. The sample is 37 cm long, by 5 cm in diameter. B: The CT sample vessel and surrounding temperature control bath are positioned in the X-Ray CT scanner using a translating table. C: Cutaway schematic of the thermal conductivity pressure vessel. Cylindrical hydrate samples 13.3 cm long by 4.1 cm in diameter are formed around a thermal probe (enlarged to show detail). D: View of a 14 cm long, 7 cm diameter sample prior to raising the sample up into the main pressure chamber of the Gas Hydrate And Sediment Test Laboratory Instrument (GHASTLI). Acoustic measurements are made using axially-positioned sensors in the top and bottom endcaps. Shear strength measurements are made by axially straining the sample using a ram brought into contact with the heat exchanger. The load supported by the sample during shear is measured by the load cell beneath the sample.
- Figure 2. Cross-section of a methane-hydrate-cemented quartz sand containing methane gas in the pore space. Figure represents the average of 56 CT scans along the length of the sample. The pore-space hydrate saturation ranges from 26 to 46%, with an average of  $35 \pm 4\%$ . Speckling in the center top portion of the scan is caused by thermocouples embedded in the sample.
- Figure 3. Density change in a methane-hydrate-cemented quartz sand as a result of a brief depressurization to one atmosphere and subsequent repressurization with methane gas. Figure represents the average of 56 CT scans along the length of the

sample. Hydrate reforms rapidly, measurably increasing the density near the central sample axis while decreasing the density near the sample surface within 1.66 hours of repressurization (A). The redistribution pattern becomes more accentuated ~24 hours after repressurization (B). White holes in the center top portion of both scans are thermocouples embedded in the sample.

Figure 4. Cross-sectional schematic (not to scale) of methane hydrate redistribution in a quartz sand (grey circles) during a depressurization and subsequent repressurization with methane gas: (A) Initially, methane hydrate (red) and methane gas (white) are fairly uniformly distributed; (B) Depressurization induces hydrate dissociation to water (dark blue) and methane gas. Dissociation front moves inward (black arrows) as heat is supplied from the surrounding bath; (C) Hydrate growth following repressurization is most efficient where hydrate persists, drawing water inward via capillary action (black arrows); (D) Final hydrate distribution is concentrated near the sample's central axis and depleted near the sample perimeter.

Figure 5. Hydrate redistribution resulting from a brief depressurization followed by repressurization changes certain aspects of the acoustic signal. The original shear-wave arrival near 110 ms travel time (dotted blue curve) increases in amplitude by ~66%, and travels 47% faster (solid red curve) as a result of the hydrate redistribution. A possible normal mode feature initially arriving near 80

μs (*M. Lee personal communication*, 2007), experiences a similar shift, but the small P wave arrival near 40 μs remains nearly unchanged.

Figure 6. Triaxial shear testing for hydrate-cemented sands that have (dotted curves) and have not (solid curves) experienced a brief depressurization. A brief depressurization does not significantly alter the shear stress dependence on strain prior to shear, but does reduce the peak shear stress. The stress drop following the peak stress does not correlate with whether samples have experienced a brief depressurization.













