Methane hydrate rock physics models for the Blake Outer Ridge

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ABSTRACT

Seismic analyses of methane hydrate data from the Blake Outer Ridge indicate high P-wave velocity and anomalously low S-wave velocity in sediments containing methane hydrates. In an attempt to explain this observed P-wave and S-wave velocity structure at the transition from gas to hydrates, the effect of different hydrate models on elastic moduli and velocities are explored. After construction of an initial gas model, the properties of the hydrates are estimated using the bound averaging method together with the Voigt and Reuss bounds for elastic moduli. The result suggests that the hydrates becomes part of the rock matrix and softens the pores by fracturing. The additional formation of ice is required to obtain the desired P- to S-wave velocity ratio in the hydrate bearing sediments, indicating temperature conditions around the freezing point of water.

INTRODUCTION

Methane hydrates are ice-like, crystalline lattices of water molecules in which gas molecules are trapped physically without the aid of direct chemical bonds. They form only under certain pressure and temperature conditions which limit their occurence to polar and deep oceanic regions. Due to the enormous amount of methane that is apparently sequestered within hydrate structures, they are likely to have a "greenhouse" effect on future global climate (Leggett, 1990), and might also be an important future energy source (Kvenvolden, 1993). Therefore, a good understanding of the characteristics and properties of the hydrate deposits is essential. However, only limited information is available from deep-sea drilling due to the high instability of the hydrate. Thus, information such as acoustic velocity and resistivity is mostly inferred from seismic reflection data (?Rowe and Gettrust, 1993; Singh et al., 1993; Ecker and Lumley, 1994). Although the properties of pure methane hydrate were determined in several lab measurements (Stoll and Bryan, 1979; Pearson et al., 1986), the properties of an in situ hydrate deposit is still quite controversial.

One of the first regions that was associated with the occurence of methane hydrates is on the eastern United States margin, offshore Florida, on the Blake Outer Ridge. Deep-sea drilling on the Blake Outer Ridge (Kvenvolden and Barnard, 1983) found high methane content below a subbottom depth of 50 m. Furthermore, approaches to estimate the amount of hydrate in this region (Lee et al., 1992a,b) suggested a significant amount of hydrate above the stability zone. Seismic investigations (Rowe and Gettrust, 1993; Ecker and Lumley, 1994)

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indicated a considerable increase of compressional wave velocity in the hydrate. Based on AVO amplitude responses and impedance estimations, Ecker and Lumley (1994) showed that the hydrate layer is characterized not only by high P-wave velocity but also by anomalously low S-wave velocity. The hydrate layer seemed to be underlain by a thick gas layer having lower P-velocity but considerably higher S-velocity than the hydrate.

This study is a first attempt to explore the effects of different hydrate structures on the elastic moduli and velocities, and compare them with those obtained by Ecker and Lumley (1994). In this way, a clear discrimination between different models is possible and should give a good insight into the possible hydrate structure at the Blake Outer Ridge.

In this paper I discuss the properties of several hydrate structures which were obtained by modeling a transition from gas saturated sediments to sediments containing methane hydrate. An initial gas saturated, shaly sand model is constructed by using deep-sea drilling information and the seismic velocity information given by Ecker and Lumley (1994). The bound averaging method for fluid substitutions is then used together with the Voigt and Reuss bounds to model the elastic moduli of hydrate layers with different mechanical structures. The resulting moduli are converted into velocities and compared with those obtained by the seismic study. Although the high clay content in the sediments in this area introduces a significant uncertainty in the moduli and velocity calculations, a clear trend in the behavior of the hydrate properties can be observed for the different structures.

POSSIBLE HYDRATE STRUCTURES

Based on the seismic analysis performed by Ecker and Lumley (1994) on data from the Blake Outer Ridge, it can be assumed that the high velocity hydrate layer (2.5 km/s) is underlain by a gas layer having a P-wave velocity of approximately 1.6 km/s. Furthermore, the study showed that the hydrate layer has to be characterized by anomalously low S-wave velocity of around 0.5 km/s which increases to 1.1 km/s in the gas saturated sediments, in order to explain the AVO data. The estimated velocities in the gas layer yield a Poisson's ratio of 0.1 which is consistent with free gas.

Several different hydrate structures can be imagined to explain this considerable increase in S-wave velocity and simultaneous decrease in P-wave velocity at the transition from hydrate bearing sediments to sediments containing free gas. The simplest model can be best described as a so-called "slush" model. As the gas/water pore fluid in the sediments reaches the base of the hydrate stability zone, it is transformed into a mixture of hydrate and water, i.e. a slush. This resembles a simple substitution of the pore filling while the properties of the actual rock matrix remain unchanged. The model assumes that the pressure-temperature conditions at the transition zone allow the formation of hydrate at temperatures above the freezing point of water. In the case of lower temperatures, part of the water might freeze in which case the gas/water pore filling is converted into a hydrate/ice/water slush which will affect the elastic properties in a different way.

Another possible way to model the transition from gas to hydrate is to assume that the hydrate becomes part of the rock matrix which would result in a decrease of the porosity of the composite. An important factor in this model is the influence of the methane solidification on the rock matrix. One possibility is that the hydrate formation and addition to the rock matrix causes fracturing of the sediments, thus causing some of the pores to become crack-like and compliant. On the other hand, there might be no change in the stiffness due to the freezing of the methane at all. Again, the temperature at the transition zone might be an important factor, also for the additional formation of ice which would reduce the porosity of the mixture even further.

The properties of hydrate, water, ice, and gas used to construct the different hydrate structures are shown in Table 1. The gas properties are valid for a pressure of 10MPa which is a realistic assumption for the depth of the gas-hydrate transition zone at the Blake Outer Ridge.

substance	K [MPa]	μ [MPa]	$\rho [\text{kg/m}^3]$	v_p [m/s]	v_s [m/s]
water	2250	0	1000	1500	0
pure hydrate	5600	2400	767	3450	1770
ice	6650	2550	916	3310	1670
gas	10	0	100	375	0

Table 1: Elastic properties of water, pure hydrate, ice and gas.

MODELING TECHNIQUE

In order to estimate the actual effects of the hydrate structures proposed in the preceding section on the elastic and acoustic properties of hydrate bearing sediments, I used the so-called Bound Averaging Method (BAM) by Marion (1990). This method estimates the changes in elastic moduli and velocities that result from substituting one pore-filling material with another.

The estimation of the moduli for a volume mixture of minerals and pore fluids requires the incorporation of the following parameters:

- the elastic moduli of all constituents of the solid phases
- the volume fractions of each solid constituent
- the elastic moduli of all constituents of the pore fluids
- the volume fractions of each fluid constituent
- the stiffness of the phase mixture

If the stiffness of the mixture is not known, only upper and lower bounds can be specified for the moduli. At any given composition of the phases, the effective modulus will fall between the bounds. Its precise value, however, can not be determined as it depends on the stiffness of the given phase state. A mixture containing stiff pores will be located along the upper bound while a mixture containing soft pores will be located along the lower bound. Strictly empirical bounds are the so-called Voigt and Reuss bounds which give the ratio of average stress and average strain within the composite. For the Voigt upper bound, the strain is assumed to be uniform everywhere. The Reuss lower bound assumes a state of uniform stress within the entire mixture.

For a compound of mineral and pore fluid, the Voigt upper bound is given by:

$$M_v = \phi M_{fluid} + (1 - \phi) M_{min} \tag{1}$$

The Reuss lower bound can be determined by:

$$M_r = \frac{\phi}{M_{fluid}} + \frac{(1 - \phi)}{M_{min}} \tag{2}$$

with: $\phi = \text{porosity} = \text{volume fraction of the pore fluid}$

 M_{min} = elastic modulus of the mineral

 M_{fluid} = elastic moduli of the pore fluid If the mineral is composed of several constituents, the effective moduli for the pure mineral without any pores are determined using the Voigt-Reuss-Hill average:

$$M_{vrh} = \frac{M_v + M_r}{2} \tag{3}$$

The Voigt and Reuss bounds are obtained by replacing the moduli and porosity in equations (1) and (2) above with the moduli of the individual constituents and its volume fractions, respectively, and summing over the number of constituents. The actual moduli of a multi-component pore fluid is obtained by just using the Reuss bound with the individual moduli and volume fractions, and summing over them.

Knowing the elastic modulus of a given mixture, the bound averaging method (BAM) assumes that the fractional vertical position of the modulus within the bounds, w, is a measure of the stiffness of the mixture. Presuming that w is independent of the pore filling properties, it should remain constant with any change in pore fluid. Thus, the elastic properties of a new composite can be determined from an initial composite. The realization of the BAM method is shown in Figure 1. First, the elastic modulus of an initial model with pore filling 1 and its resulting Voigt and Reuss bounds for different volume mixtures are determined. The stiffness parameter w of the initial state is given by the position of the elastic modulus between the bounds. The Voigt and Reuss bounds are then recalculated for a pore filling material 2. As w remains constant for the transition from filling 1 to filling 2, the elastic modulus of the new composite can be obtained by determining the modulus value at the fractional position w between the new bounds.

If the composite undergoes a change in porosity, an additional linear interpolation is required. Keeping the porosity constant, the initial state is converted using the BAM method.

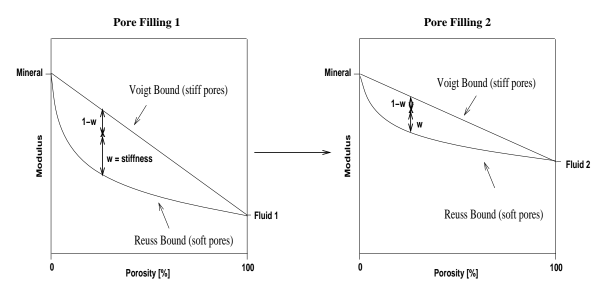


Figure 1: Bounding average method for fluid substitution. [NR]

Then, the porosity of this state is changed by linear interpolation between the upper and lower bounds. From the resulting position, the elastic modulus of the final state is obtained.

In case of an additional change in stiffness, as can be expected by substituting a fluid pore filling with a frozen one, the elastic modulus is determined in a similar way. The initial state is again transformed using the BAM method. After having determined the new position between the upper and lower bounds, the mixture can be softened by moving it closer to the Reuss bound, or stiffened by moving it closer to the Voigt bound. Having accounted for this change in the stiffness parameter, linear interpolation to a new porosity is possible.

The resulting elastic moduli are converted into velocity information using the following relationships between P-wave velocity, S-wave velocity, bulk modulus, shear modulus and density:

$$v_p = \sqrt{\frac{K + \frac{4}{3}\mu}{\rho}} \tag{4}$$

$$v_s = \sqrt{\frac{\mu}{\rho}}. (5)$$

The effective density of a two phase mixture is obtained by:

$$\rho = f_1 \rho_1 + f_2 \rho_2 \tag{6}$$

where f_i and ρ_i represent the volume fractions and densities of the individual phases.

CONSTRUCTION OF THE INITIAL MODEL

Results from deep sea drilling at the Blake Outer Ridge suggest high concentrations of clay within the sediments. The effects of clay on the elastic properties of shaly sandstones are still quite speculative and many different possible correlations exist between clay content and acoustic and elastic properties of sandstone. Due to the relatively high scattering in the possible properties of shaly sandstone, a large uncertainty is introduced into all calculations in this study.

Based on the velocity information given by Ecker and Lumley (1994), I assume the actual gas saturated sediments to have a P-wave velocity of 1.6 km/s and a S-wave velocity of approximately 1.1 km/s.

Using the given velocity and mineral information for the Blake Outer Ridge, a gas layer was constructed which could fulfill the velocity requirements. Using Han's (1986) empirical relationship for water saturated, clay-bearing sandstones,

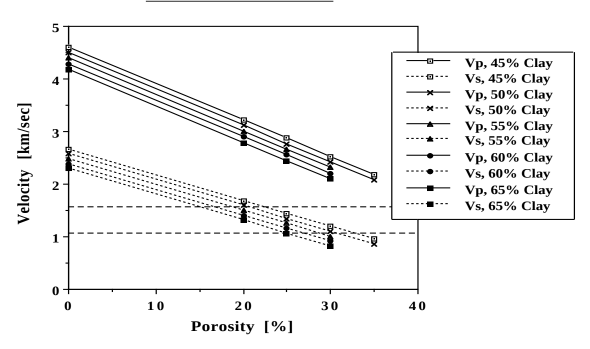
$$v_p = 5.59 - 6.93\phi - 2.18C \tag{7}$$

$$v_s = 3.52 - 4.91\phi - 1.89C \tag{8}$$

in which ϕ gives the porosity and C gives the clay content, the properties of water saturated, shaly sandstones were determined for clay concentrations between 45% and 65% and porosities between 0% and 45%. The water was then replaced with water-gas mixtures having different gas concentrations. The properties of water and gas that were used for this substitution were shown in Table 1. The resulting P- and S-wave velocities for the different mixtures were then compared with the given velocities. Figure 2 shows the P- and S-velocities versus porosity for different clay concentrations and two different saturations. Assuming no gas saturation in the sediments, the desired P- and S-wave velocities can not be obtained. Saturating the shaly sediments with a 40% gas - 60% water mixture decreases the P-wave velocities for all clay concentrations considerably. The S-wave velocities remain approximately constant. The P- to S-wave ratio, however, is for all settings slightly higher than the given one. This is caused by the use of Han's (1986) empirical relationship for saturated shaly sandstones, which might not resemble the realistic conditions. Nevertheless, it should give a reasonable approximation for this first modeling attempt.

Based on the determined velocities and P- to S-wave ratios for the different clay concentrations, I assumed the sediments to have around 55% clay concentration and a porosity of approximately 31%. As the P-wave velocity remains approximately constant after a gas saturation of only a few percent, the S-wave velocity, however, slightly increases with increasing gas concentration, I choose a 60% gas - 40% water saturation to improve the P- to S-wave ratio. This model yields a P-wave velocity of approximately 1.65 km/s and an S-wave velocity of approximately 1.0 km/s. The properties of the pure mineral and the water-gas mixture of this initial model are shown in Table 2. The bulk and shear moduli of the mineral were obtained by using the previously determined velocities for a sandstone containing 55% clay. As the density of sandstone does not vary much with clay concentration, an averaged density

100% Brine Saturated



40% Gas, 60% Brine Saturated

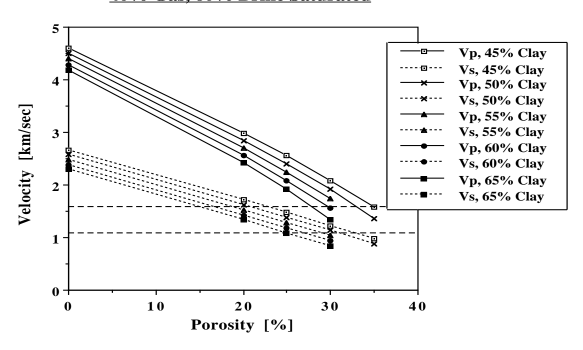


Figure 2: P- and S-wave velocities versus porosity for different clay concentrations. The upperfirst figure shows the velocities for 100% water saturated sandstone, the second one shows velocities for a saturation of a 60% gas - 40% water mixture. The two dotted lines in each figure represent the desired P- and S-velocities for the shaly sandstone. Christine2-velocity [NR]

substance	K [MPa]	μ [MPa]	$\rho [\text{kg/m}^3]$
mineral	27700	15400	2500
fluid	16.62	0	460

Table 2: Properties of the pure mineral and the fluid of the constructed gas layer.

of 2500 kg/m³ was assumed. The properties of the gas-water mixture were determined from the parameters listed in Table 1. The so obtained initial model serves as an input for the BAM method. It is shown in Figure 3. The modified Voigt bound in Figure 3 describes the expected

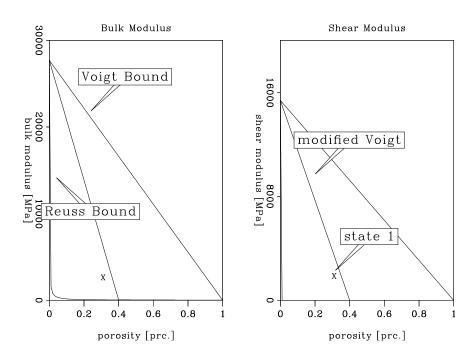


Figure 3: Voigt, Reuss and modified Voigt bound for the initial gas/water saturated sediment model. State 1 gives the actual bulk and shear modulus of this model. christine2-in1-ann [ER]

behavior of porous material for a porosity smaller than the critical porosity of the material (Nur et al., 1991). For porosities lower than the critical porosity, the mineral grains are load-bearing, while for greater porosities the sediments become a suspension. The critical porosity for shaly sand is assumed to be approximately 40%. The elastic moduli for the initial model, described as state 1 in the Figure, fall well within the load-bearing domain.

HYDRATE MODELING

1. SLUSH MODEL

Hydrate/water slush

In this first modeling approach, the 60% gas - 40% water saturation of the initial sediments is substituted with a 60% hydrate - 40% water mixture. The porosity and stiffness of the whole composite remains unchanged. Using the properties of hydrate and water specified in Table 1, the properties of the hydrate/water slush can be obtained from equations (2) and (6):

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K_{slush} = bulk modulus = 3510 MPa

\mu_{slush} = shear modulus = 0 MPa

\rho_{slush} = 860 kg/m<sup>3</sup>
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Although pure hydrate has a shear modulus of approximately 2400 MPa, the shear modulus of the hydrate-water pore filling is set to zero. It can be assumed that even for small amounts of water a slush has absolutely no shear strength. The resulting bulk and shear modulus of the hydrate model are compared with those of the initial gas model in Figure 4. Based on the

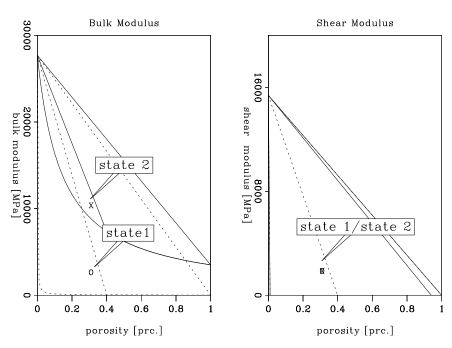


Figure 4: The solid curves show the Voigt and Reuss bound for sediments containing a hydrate/water slush. The dotted curves represent the bounds of the initial gas/water saturated sediments. State 1 gives the moduli of the initial state, while state 2 represents the slush state. christine2-final1-ann [ER]

increase in the bulk modulus of the pore filling by substituting gas/water with a hydrate/water slush, the upper and lower bounds for the bulk modulus are much narrower. The actual bulk

modulus of the new composite, marked as state 2 in Figure 4, is considerably higher than that of the initial composite, marked by state 1. This indicates a lower compressibility of the new mixture due to the additional strength of the hydrate. As the shear modulus of both pore fillings was assumed to be zero, the Voigt and Reuss bounds remain the same for the change in pore fluid. Therefore, the resulting shear modulus of the hydrate/water bearing sediments is identical with the one of the sediments containing gas and water. Both mixtures have low shear strength. Thus, replacing a gas/water pore filling with a hydrate/water slush increases the bulk modulus significantly but leaves the shear modulus constant. Using the bulk and shear moduli marked as state 2 in Figure 4, the actual P- and S-wave velocity can be determined to be:

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v_p = 2.60 \text{ km/s}
v_s = 0.97 \text{ km/s}
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The increase of the P-wave velocity from 1.65 km/s in the gas model to 2.6 km/s in the hydrate model resembles the increase observed in the seismic data. However, the decrease in the S-wave velocity which is based solely on the change in density as the shear properties remain constant for the pore-filling substitution, is negligibly small. Thus, a transition from gas/water to a hydrate/water slush seems unreasonable.

Hydrate/water/ice slush

In this model it is assumed that the temperature at the base of the hydrate stability zone allows the formation of ice. In this case, the gas/water mixture transforms into slush containing hydrate, ice, and a small amount of remaining water. Considering a 60% hydrate, 20% ice, and 20% water mixture, the following properties of the slush are calculated using equations (2) and (6):

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K_{slush} = 4420 \text{ MPa}

\mu_{slush} = 0 \text{ MPa}

\rho_{slush} = 843 \text{ kg/m}^3
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The shear modulus was again set to zero presuming zero shear strength of the slush. Therefore, regarding the preceding model, only a change in bulk modulus and thus P-wave velocity is expected. The result of the pore filling substitution can be seen in Figure 5. Due to the additional ice in the slush, the bulk modulus of this model increased even more than that in the previous one. As predicted, the shear modulus remained constant. Converting the moduli into velocities yields:

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v_p = 2.77 \text{ km/s}
v_s = 0.97 \text{ km/s}
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Comparing these velocities with those obtained with the hydrate/water slush model, it is obvious that the presence of ice increased the P-wave velocity further. However, as the addition of

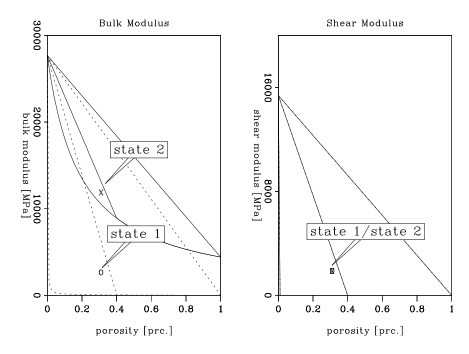


Figure 5: The solid lines give upper and lower bounds of the slush model. The dotted lines show those of the initial model for comparison. State 2 indicates the actual moduli of the sediments containing slush, state 1 gives the moduli of the gas layer. christine2-final2-ann [ER]

ice to the slush does not change the shear strength and leaves the density fairly unchanged, a hydrate/water/ice slush also does not result in a desired increase of the P-wave velocity and a decrease in S-wave velocity.

The slush model can be further modified by changes in the gas and thus hydrate content, but its characteristic of having no shear strength and only considerable small changes in density will always yield a relatively small decrease in the S-wave velocity. Consequently, a slush model is not a realistic representation of the hydrate structure at the Blake Outer Ridge.

2. Matrix Model

No change in stiffness

In this modeling approach the hydrate that forms when the gas comes in contact with the base of the hydrate stability zone becomes part of the mineral. The resulting composite is 100% water saturated and has a reduced porosity. The solidification of the methane is assumed not to change the stiffness of the rock. The initial model of 69% mineral and 31% pore fluid consisting of 60% gas and 40% water is thus transformed into 88% mineral and 12% pore fluid consisting of 100% water. In contrast to the slush model, both the properties of the pure mineral and the pore filling of the hydrate bearing sediments have changed. Using the parameters given in Table 1 and Table 2, and the equations for the Voigt bound, Reuss bound, and

Voigt-Reuss-Hill average, the properties of the pure mineral and the pore fluid are determined and shown in Table 3. A comparison with the properties of the initial model given in Table 2

substance	K [MPa]	μ [MPa]	$\rho [\text{kg/m}^3]$
mineral	18800	9650	2120
fluid	2250	0	1000

Table 3: Properties of the pure mineral and pore fluid of the hydrate model.

shows that the addition of hydrate to matrix has decreased both the mineral bulk modulus and the shear modulus considerably. The replacement of the gas/water pore filling with water leads furthermore to a significant increase of the fluid bulk modulus and density. Figure 6 shows these effects clearly. The resulting bulk modulus, marked as state 2 in Figure 6, is significantly

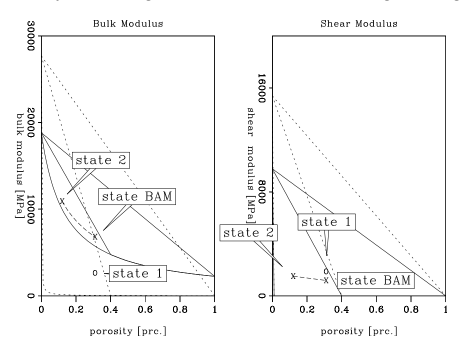


Figure 6: The solid lines represent the bounds for the hydrate model while the dotted lines give the bounds of the initial model. State 1 marks the actual moduli given for the initial model, state 2 those for the hydrate model. State BAM gives the moduli after the BAM method and before interpolation to the new porosity. christine2-final1a-ann [ER]

higher than the old bulk modulus, marked as state 1, indicating that the new sediment structure has much lower compressibility. State BAM represents the modulus obtained by transforming the gas model using the BAM method without changing the porosity. The shear modulus of the hydrate structure has slightly decreased with regard to the modulus of the gas structure. The velocities obtained from these elastic moduli are:

 $v_p = 2.55 \text{ km/s}$ $v_s = 0.87 \text{ km/s}$

Comparing these velocities with those of the initial gas model ($v_p = 1.6 \text{ km/s}$, $v_s = 1.0 \text{ km/s}$) shows that the P-wave velocity increased considerably while the S-wave velocity decreased slightly. This velocity trend resembles the one obtained from the seismic study by Ecker and Lumley (1994). However, although the increase in P-wave velocity is in good agreement with the actually observed one, the model did not succeed in reproducing an S-wave velocity ratio of approximately 2:1 from gas to hydrates. Thus, adding the hydrate to the rock matrix and hence decreasing the porosity of the resulting water-saturated composite predicts the correct velocity trend but its effect on the S-wave velocity is not significant enough.

Change in stiffness

Assuming that the hydrate becomes part of the rock matrix during its formation, fracturing of the composite is possible due to the solidification of the methane. This would introduce crack-like and compliant pores resulting in a softening of the hydrate bearing sediments. To realize this model, the initial gas model of 69% mineral and 31% gas/water mixture is again transformed into 88% mineral containing 22% hydrate and 12% water filling the pores. In contrast to the previous model in which no fracturing of the rock was assumed, the stiffness of the hydrate bearing sediments is changed by assuming that the fracturing causes a 50% softening of the rock. The properties of the pure mineral and the pore filling are not changed by a softening of the rock and are thus the same as in the previous model shown in Table 3. The result of this modeling approach can be seen in Figure 7.

The moduli obtained with this model are again marked as state 2 in Figure 7, and those obtained from the initial model are marked as state 1. "Softened" represents the position after application of the BAM method and 50% softening. As in the model without fracturing, the bulk modulus increased significantly with respect to the initial model. On the other hand, the shear modulus shows a more notable than before. The bulk and shear moduli of this hydrate model yield the following velocities in the hydrate bearing sediments:

$$v_p = 2.40 \text{ km/s}$$

 $v_s = 0.62 \text{ km/s}$

A comparison with those values obtained without softening the sediments shows that the additional fracturing caused the P-wave velocity to decrease slightly from 2.55 km/s to 2.4 km/s. An even larger effect can be observed on the S-wave velocity which decreased from 0.87 km/s to 0.62 km/s. This indicates that further softening of the pores would result in an even more decreased S-wave velocity of the hydrate structure. Apparently, a decrease similar to the one determined from the seismic data (1.1 km/s in the gas, 0.5 km/s in the hydrate) is possible in this way. However, further softening would also introduce a more pronounced decrease in the P-wave velocity compared to the one obtained without softening. Therefore, this model provides a good explanation for the observed anomalous decrease in S-wave velocity, but lacks the simultaneously large increase in P-wave velocity. Another possible hydrate structure has to exist which would leave the S-wave velocity decreased by fracturing approximately constant

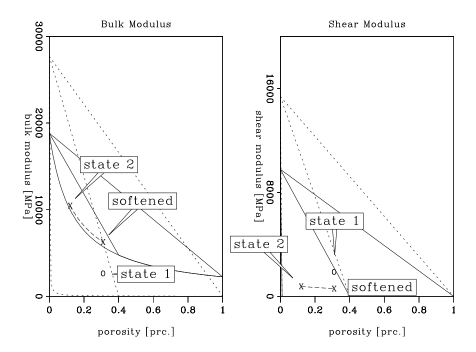


Figure 7: Solid lines represents the bounds for the hydrate model, and dotted lines represent the bounds for the gas model. State 1 marks the elastic moduli of the initial model, state 2 marks the final elastic moduli and "softened" describes the position after BAM and softening. christine2-final1b-ann [ER]

but would again increase the P-wave velocity. As shown in a previous section, the additional formation of ice can be expected to have such an effect. This assumption is investigated in the following hydrate model.

Change in stiffness and ice formation

This modeling approach explores the effects of additional ice formation at the gas-hydrate transition zone, caused by temperature conditions around the freezing point of water. During the formation of hydrate and ice, they become both part of the mineral and cause fracturing of the rock. Presuming that 50% of the water in the pore space freezes, a sediment model containing 94% mineral and 6% pores filled with water is constructed. The mineral is composed of 73% shaly sands, 20% hydrates and 7% ice. The initial model on the other hand contained 69% mineral and 31% pores filled with a gas/water mixture. Due to the additional formation of ice, the properties of the pure mineral have changed with regard to the previous model without ice. As the model still assumes 100% water saturation, the properties of the pore filling did not changed. The new mineral is characterized by:

 $K_{min} = 17800 \text{ MPa}$ $\mu_{min} = 9100 \text{ MPa}$

$$\rho_{min} = 2043 \text{ kg/m}^3$$

The addition of ice to the pure mineral has decreased its elastic properties even further. The effect of the ice formation on the elastic moduli can be seen in Figure 8. As before, state 1

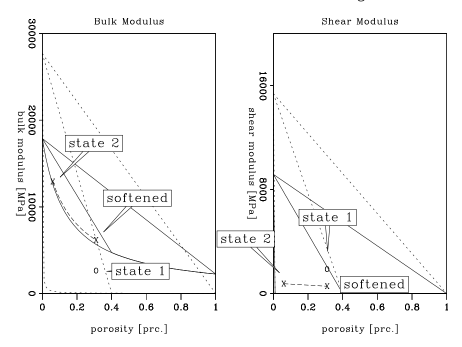


Figure 8: The solid curves represent the bounds for the hydrate sediments, while the dotted curves represent the bounds for the gas sediments. The actual moduli of the gas sediment are marks as state 1, those of the hydrate sediments as state 2. "Softened" describes the position after transformation of the gas layer moduli with the BAM method and softening of the rock. [christine2-final2b-ann] [ER]

represents the moduli of the gas/water saturated sediments while state 2 marks the moduli of the hydrate structure after interpolation to a porosity of 6%. "Softened" describes the position after transformation of the initial model using the BAM method and softening of the pores. It is obvious that this model results in the same velocity behavior as the previous model without ice. The bulk modulus has increased considerably while the shear modulus in the hydrate structure decreased. The actual velocities that can be determined are:

$$v_p = 2.65 \text{ km/s}$$

 $v_s = 0.62 \text{ km/s}$

A comparison of these values with those obtained without the presence of ice shows that the P-wave velocity has increased while the S-wave velocity remained constant. Figure 9 shows this behavior clearly. Consequently, the P-wave velocity which becomes too low by softening the pores can be increased again by the addition of ice to the mineral. As predicted, there is no notable effect on the S-wave velocity. The obtained velocities are in good agreement with those obtained by the seismic analysis ($v_p = 2.5 \text{ km/s}$, $v_s = 0.5 \text{ km/s}$) and an even better agreement

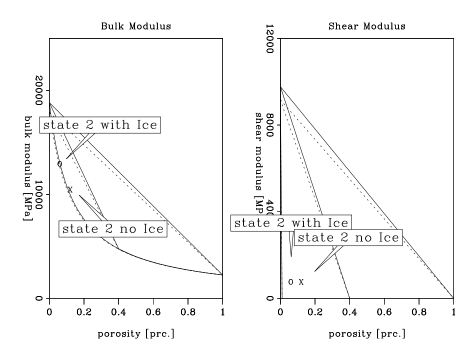


Figure 9: Comparison of the bulk and shear moduli obtained with and without ice. The solid lines represent the bounds for the absence of ice while the dotted lines give the bounds for the presence of ice in the mineral. christine2-comp-ann [ER]

could be achieved by further softening of the pores. Thus, assuming temperature conditions close to the freezing point of water and fracturing during the hydrate and ice formation seems to be a reasonable model for the hydrate structure at the Blake Outer Ridge.

However, a comparison of the actual numbers should be done very carefully due to the fact that the initial shaly sand model was determined only in first approximation. Based on the high clay content present in the sediments of the Blake Outer Ridge, a relatively high uncertainty is introduced into the elastic moduli and velocities of the initial gas structure. A different starting model might thus result in different absolute velocity numbers in the hydrate structures. Furthermore, it has to be considered that the velocities determined by Ecker and Lumley (1994) might also have some uncertainties.

CONCLUSIONS

The properties of different hydrate structures were evaluated in an attempt to explain the seismic velocities determined at the Blake Outer Ridge. All results were related to an initial gas/water saturated rock model containing 60% shaly sands and a porosity of 31%.

The analysis showed that the P-wave velocity in sediments containing hydrates increases significantly independently from the apparent hydrate structure. The S-wave velocity on the other hand is extremely dependent on the way the hydrate forms and affects the rock matrix. The assumption of a so-called "slush" model could not reproduce the required considerable

decrease in S-wave velocity at the transition from gas to hydrate. Presuming, however, that the hydrate becomes part of the mineral and causes softening of the pores due to induced fracturing, the S-wave velocity in the hydrate layer could be decreased significantly. Simultaneously, fracturing induced a slightly lower P-wave velocity in the hydrate structure. It was therefore necessary to assume temperature conditions close to the freezing point of water to allow the additional formation of ice. This resulted in an increase of the P-wave velocity while the S-wave velocity remained constant, creating the desired P- to S-wave ratio.

It can be concluded that a hydrate model which adds hydrate and ice to the mineral and softens the pores by fracturing seems to be a reasonable assumption for the hydrate bearing sediments of the Blake Outer Ridge based on the given initial model. The high clay concentration at the Blake Outer Ridge introduced, however, high uncertainties into this initial model due to the still quite speculative properties of clay. Thus, the effects of different clay concentration and properties as well as different gas concentration in the initial gas structure have to be evaluated further.

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