A two-dimensional model of the passive coastal margin deep sedimentary carbon and methane cycles

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Abstract

We present a new geologic-time and basin-spatial scale model of the continental margin methane cycle. The model, SpongeBOB, is used to simulate evolution of the carbon cycle in a passive sedimentary continental margin in response to changing oceanographic and geologic forcing over a time scale of 200 million years. The geochemistry of the sediment column is altered by the addition of vertical high-permeability channels intended to mimic the effects of heterogeneity in the real sediment column due to faults, and produces results consistent with measured pore-water tracers $SO_4^{\ 2-}$ and $^{129}I.$ Pore water DIC concentrations are consistent with chemical weathering (CaCO₃ formation from igneous rocks) at depth within the sediment column. The carbon isotopic composition of the DIC is consistent with a methane production efficiency from POC of 50%, which is somewhat lower than redox balance with the H/C of organic matter in the model. The hydrate inventory in the model is somewhat less sensitive to temperature than our previous results with a one-dimensional model, is quite sensitive to reasonable changes in POC, and extremely sensitive to the ability of methane bubbles to rise within the sediment column, and how far gas-phase methane can get through the sediment column before it redissolves when it reaches undersaturated conditions. Hydrate formation is also sensitive to deep respiration of migrating petroleum. Other phenomena which we simulated had only small impact on the hydrate inventory, including thermogenic methane production, and production / decomposition of dissolved organic carbon.

Introduction

Models of methane hydrate formation in sediments of the deep-sea have tended to follow the example of early diagenesis modeling in adopting a one-dimensional formulation [Burdige, 2011; Chatterjee et al., 2011; Davie and Buffett, 2001; Davie and Buffett, 2003a; Davie and Buffett, 2003b; Garg et al., 2008; He et al., 2011; Liu and Flemings, 2007; Xu, 2004]. For the top meter of the sediment column [Archer et al., 2002], the 1-D approximation is a good one, because vertical diffusion is far faster than any possible lateral pore fluid transport or diffusional effects. The methane cycle challenges a 1-D formulation, however, because the relevant chemical processes occur hundreds of meters below the sea floor, and clearly show the impact of pore fluid and gas phase migration in more than one dimension. One-dimensional models of the upper sediment require fluid flow imposed as a bottom boundary condition, and the hydrate inventory of the model is very sensitive to this parameter [Buffett and Archer, 2004]. The methane cycle is also impacted by time-dependence of the sediment and organic carbon deposition rates, which are governed by onshore-offshore processes, another benefit of incorporating the second dimension into the model formulation.

SpongeBOB is formulated in the onshore-offshore dimension laterally, and to bedrock in the vertical, to internalize the entire coastal margin carbon and methane cycles, of which the methane hydrates near the sea floor are just one manifestation. Many of the processes and mechanisms, such as all the different means of downslope sediment transport, or pathways for subsurface fluid flow, are impossible to represent accurately given our present state of knowledge, but by representing them as best we are able, we hope to gain some qualitative picture of the relative importances of the various driving factors and processes.

The model is set up to simulate an accumulating passive continental margin such as the East coast of the United States, which has been accumulating sediment since rifting of the Atlantic about 200 million years ago. One of the best-studied locations for methane hydrates is the Blake Ridge, a "drift deposit" ridge jutting out into the Atlantic [Borowski, 2004]. The Blake Ridge owes its existence to the hydrodynamics of particle sedimentation, presumably driven by existing topography and the energetic western boundary flow of the Atlantic. Although Blake Ridge is attached to the continental slope of the U.S., our model results (which only represent the slope, with no drift-deposit ridge) suggest that Blake Ridge might not be entirely typical of slope sediments in general, and thus might be deceptive jumping-off point for global extrapolation.

We gauge the sensitivities of this complex model by showing a variety of sensitivity model runs, varying critical parameters or turning process on and off. The characteristics of the runs will be described as they come up in the model formulation section, and they are summarized in Tables 1 and 2.

Model Formulation

Computational Grid

SpongeBOB is formulated on a free-form grid sigma-type vertical coordinate system in depth (Figure 1), in which the grid cells expand to fill the sediment column as it expands through time, stretching from bedrock to the sediment surface. As sediment accumulates at the sea surface each time step, sediment is also moved between subsurface grid cells in order to maintain the stretched grid. At the beginning of the simulation all grid cells contain enough solid material to fill a thickness of 1 meter. The initial porosity of the cells is "drained", such that the sediment load is carried entirely by the solid matrix with no excess pore fluid pressure to drive a Darcy flow. Sediment is deposited on the top face of the top cell, and a material flow rate is calculated for the bottom face of the top cell, such that the top cell keeps some of the new material for itself, but passes most of it on fill the subsurface cells below. The process continues through the sediment column, and the boxes all expand uniformly.

A few grid cells in the top of the domain are restricted, such that they cannot expand beyond a thickness of 200 meters, in order to maintain a minimum computational resolution in the region of the hydrate stability zone. After enough model time has passed that a restricted cell reaches its maximum thickness, it passes along the entire incoming advective flux from above into the cell below.

Isostacy

The sediment column elevation is governed by isostasy, the idea that the crust and all the material sitting on top of it are all floating in a denser fluid of mantle material [*Watts*, 2001]. Laterally, the model domain spans between continental crust and ocean crust, as can be seen in the transition in the thickness of the crust in Figure 1.

The effect of cooling with age on the elevation of the ocean crust is driven by the evolution of a time-dependent thermal boundary layer, extending through the crust into the mantle below. As the material in the crust and the mantle boundary layer cool, they contract and become denser. The thickness of the boundary layer is assumed to increase according to $dz = \operatorname{sqrt}(\kappa T)$ [Parson and Sclater, 1977]. The temperature profile with depth is assumed to be linear, allowing us find the temperature at the crust / mantle boundary by interpolation, from which to compute the average temperatures of the crust and the mantle boundary layer. The temperature at the base of the mantle boundary layer is taken to be 1600 K [Iamori et al., 1995]. The floating pile in the ocean crust endmember consists of the sediment column, the ocean crust, and the thermal boundary layer as it thickens and thermally contracts through time.

The continental crust is simpler because there is no time-dependent cooling. The crust is defined to be 13 km thick and its density is 3.0 g/cm³. Any sediment load also contributes to the mass of the column, and the equilibrium elevation of the bedrock.

The blending of domains is done by calculating pure end-member continental and ocean crust scenarios at each model grid point, and then scaling between them using an ad-hoc ocean fraction variable, ranging from 0 for continent to 1 for the ocean end member. The transition is set up with a horizontal length scale of 50 km. For each end member, continental and oceanic, the masses and heights of the column are computed. The two scenarios are blended together at this stage, taking a composite column mass that scales between the two end members, and a composite column height. Although this is not a mechanistic simulation of crustal dynamics at a passive margin, it is sufficient to produce a final envelope of sediment that resembles seismic sections of passive margins (Figure 4), as a driver for the sedimentary core of the model.

Sea level is defined as 4000 meters elevation above generic ocean crust. The hypothetical elevation of the top of the mantle, the sea in which the crustal material floats, is defined to be consistent with this.

The equilibrium heights for the top of bedrock are smoothed horizontally through the grid using a diffusion algorithm that achieves a spatial scale of a few 10's of kilometers (three passes with a diffusion coefficient of 10^6 m² yr⁻¹). Flexure and rigidity of the plates becomes important in accretionary margins, but for this study the lateral interaction in isostacy is rudimentary, merely setting the stage for the sediment column. The elevation of the bedrock evolves toward the isostatic equilibrium value with a time constant of 10^{-4} yr⁻¹.

Deposition

The goals of the sedimentation scheme are to create appropriate sedimentation rates and grain sizes, in adaptive response to a time-evolving submarine landscape. The existence of a shelf break, and the prograding clinoform pattern of sediment deposition, requires the existence of a region of fast sediment accumulation just off the shelf break, a sediment depocenter [*Pirmez et al.*, 1998]. Sediment input from the continent is also sorted by grain size, with largest grains depositing closer to shore and finer grain sediments offshore (Figure 2).

Offshore sediment transport and deposition in SpongeBOB is parameterized in an approach following Pirmez et al [1998]. Sediment comes into the model domain on the continental (left-hand) boundary, suspended in the ocean. It advects offshore with a sediment transport velocity that decreases inversely with water depth, as if the ocean margin were a giant river carrying suspended particles away from the continent in a plug flow. Suspended particles settle out following Stokes' law according to their grain size

$$w_{\text{sink}} = \rho_{\text{sediment}} R_{\text{particle}} g / (25*10^{-6} \rho_{\text{seawater}})$$

where R_{Particle} is the particle radius, ρ stands for density, and g is Earth's gravitation acceleration. Particles deposit only if the shear stress at the bed, calculated as

$$\tau_{\text{shear}} = 5 \, * \, 10^{\text{--}5} \, * \, \rho_{\text{seawater}} \, * \, v^2$$

is too low to lift the particles faster than they sink, determined as a critical shear cutoff

$$\tau_{\text{critical}} = \rho_{\text{seawater}} * w_{\text{sink}}$$

The sediment transport scheme is not mechanistically realistic, but is intended to serve as a realistic boundary condition driver for the sediment column core of the model. The sediment transport offshore velocity (0.1 m s⁻¹ in water depth depth of 10 m) and the resulting shear deposition scheme were tuned to produce a reasonable facsimile of a seismic section of the Atlantic margin (Figure 4).

The critical shear velocity depends on particle radius, as does the settling speed, effectively sorting the particles by size with offshore distance. The impact of water depth on the plug flow and hence the shear velocity allows changes in sea level to impact sedimentation and the evolution of the sediment column.

Slope-Driven Offshore Transport

Turbidites in abyssal sediments attest to the importance of slope failure to the margin sediment dynamics [*Meiburg and Kneller*, 2010]. These processes are handled as a function of the sea floor grade. If the grade exceeds a critical value, here taken to be 6%, the material that would deposit according to the shear / sinking speed criterion is instead

classified as "resuspended". Also, if the sea floor slope ever manages to exceed the critical slope, surface sediment is eroded into the resuspended pool. The resuspended material redeposits using the same settling speed / offshore transport criterion as used for the primary settling material, but resuspended material only begins to deposit where the sea floor slope is less than 1%, reflecting the difference between a turbidity flow which only stops on the plain as opposed to individual sinking particles assumed in the primary sedimenting material.

The particulate organic carbon (POC) concentration of the sediment is determined in the model as a function of water depth at the time of initial sediment deposition, as discussed under the heading Carbon Cycle. When material is resuspended from the sea floor, in contrast, its POC concentration is conserved and redeposited. Because the offshore transport is assumed to be turbidites, we assume it escapes the degradation that individual settling particles, the primary deposition, are subject to as they slowly sink through the water column. Resuspended organic carbon is assumed to be distributed among the grain size fractions of the resuspended material according to surface area, selectively concentrating (according to mass) the organic carbon into the smaller size fractions.

The sediment transport scheme is intended to be a utilitarian parameterization of sediment transport processes, rather than a mechanism-resolving model such as Sedflux [Syvitski, 2001]. The scheme as we have tuned it achieves the goal of a well-defined shelf break, driven by a strong sediment depocenter just off the shelf break, and response to sea level changes that mimics the Sedpak model [Strobel et al., 1989]. The accumulation rate at 2 km depth in the model is close to that of Blake Ridge, about 2 cm / kyr. Total sediment thicknesses and extents are comparable with a seismic image of the Atlantic margin from Kennett [1982] (Figure 4). Model sensitivity to these sediment transport processes was gauged by altering the critical sea floor slope in sensitivity simulation Lo Slope.

Sea Level Changes

Sedimentation is heavily impacted by eustatic changes in sea level, with highest rates close to shore but in water that is not too shallow. Sea level changes are imposed on the simulation as a 240 million year cycle, peaking at high sea level about 100 million years ago and declining since then, to generally and smoothly follow the larger-scale sea level changes in the Haq [1987] curve (Figure 3). The in organic carbon (POC) of the new sediment covaries with sea level in the model (see Carbon Cycle section). Two sets of simulations were done, one which neglected time-dependent sea level changes (Table 1), and another set which included these effects (Table 2). Results in this paper will be presented as time slices from various points in the simulations, but animations of the entire runs are available as links from the figure captions.

Pore Water Geophysics

Porosity and Excess Pressure

The pore fluid excess pressure, which drives the pore fluid flow, is calculated from a constitutive equation in which the excess pore fluid pressure can be calculated from the porosity and the mass load overhead [Flemings et al., 2002]. The idea is that there is a one-to-one relationship between the porosity of the sediment and the amount of solid load it can support. The compression of the bulk sediment by exclusion of pore water at hydrostatic pressure is governed by

$$\phi_{\text{hydrostatic}} = 0.38 \ e^{-\beta_1 \ S_{\text{supp}}} + 0.32 \ e^{-\beta_2 \ S_{\text{supp}}}$$
 (1)

where the β values are bulk compressibilities and S_{supp} is the solid load that can be supported by sediments of that porosity. Since pore fluids are relatively close to hydrostatic, the porosity profile in the sediment column is largely governed by this relationship. We find that to produce the relatively high porosity of surface sediments and the lower porosities of the deep sediment column requires two exponentials with different β values (Figure 5b). Coefficient β_1 is 0.0312 and produces compaction on a depth scale of about 10 km, representing ductile flows and recrystalization of mineral grains. The value of β_2 is 0.2, producing a compaction depth scale of about 2 km (Figure 5), representing the collapse of clays in the shallower sediment column. Because of the double exponential form of the constitutive equation, calculating S_{supp} from Φ is an iterative calculation

The excess pressure is then calculated as

$$P_{\text{excess}} = S_{\text{v}} - P_{\text{hydro}} - S_{\text{supp}}$$

where S_v is the total load of the sediment column from all phases, and P_{hydro} is the hydrostatic pressure (the load due to the fluid phase).

Temperature and Heat Flow

Subsurface temperatures are primarily governed by thermal conduction, a diffusional process, although fluid advection and latent heat also affect the temperature. The thermal conductivity and heat capacity are taken as a volume-weighted averages of the components: sediment, pore fluid, methane hydrate, and methane gas. A geothermal heat flux is imposed at the base of the sediment column of 70 mW m⁻², with sensitivity runs spanning from 50 to 120 mW m⁻².

Permeability

The vertical permeability k_y is parameterized as

$$k_h = \frac{\bar{r}^2 + \phi^3}{180 (1 - \phi)^2}$$

where \bar{r} is the mean grain radius and ϕ the porosity [Bear, 1972]. A factor of 100 anisotropy is imposed, following the example of [Daigle and Dugan, 2011], such that

$$k_{h} = 100 k_{y}$$

to parameterize for the effect of layers like sandy turbidites, that are unresolved in the model.

We show results of two model permeability formulations, one as presented so far, and the other with added vertical channels of enhanced permeability, such as faults, pipes, channels, and similar structures (cases *Base* and *No Channels*). The channels are situated every 5 grid cells throughout the model domain, and remain stationary as the continental margin progrades offshore through the simulation. The horizontal grid spacing is 3.15 km, and the vertical permeability in the channel is 10 times higher than given above. This is only a crude representation of a realistic flow regime through faults and permeable channels that seem to dominate subsurface flows [*Flemings et al.*, 2003]. Most of these features would be much smaller than the grid resolution of the model can represent more realistically, although some seismic "blowout zones", disturbance of the layered sedimentary structure due presumably to gas escape from the deep sediment column, do span several kilometers in horizontal extent [*Hill et al.*, 2004]. The intent is to gauge in a qualitative way the sense and potential magnitude of the impact of heterogeneity on the flow and chemical evolution of the sediment column.

The permeabilities and the excess pressures of the two simulations at the end of the simulations are shown in Figure 6.

Darcy flow

Fluid flow follows Darcy's law in response to the pore fluid excess pressure gradient [Athy, 1930], to find along- and across-grid velocities as

$$u_{_{Darcy,i\rightarrow i+1}} = \frac{k_{_{h,i}} + k_{_{h,i+1}}}{2\mu} \frac{P_{_{excess,i}} - P_{_{excess,i+1}}}{(\Delta x_{_i} + \Delta x_{_{i+1}})/2}$$

and

$$w_{\text{Darcy},j \rightarrow j+1} = \frac{k_{v,j}}{\mu} \frac{P_{\text{excess},j} - P_{\text{excess},j+1}}{(\Delta z_i + \Delta z_{i+1})/2}$$

where m is the dynamic viscosity, k the permeability, P_{excess} the excess pressure, and Δx and Δz the grid dimensions. All of these variables except the viscosity are defined on the two grid dimensions i and j. The velocities are defined on the cell faces.

A flow limiter in the vertical protects the calculation from numerically overstepping, for thin, permeable layers, by testing for the change in fluid pressure that would result from unimpeded Darcy flow each time step, as

$$\Delta P_{\rm excess} = \Delta \phi \cdot \frac{\partial S_{\rm supp}}{\partial \phi}$$

where the derivative of the porosity / S_{supp} relation ship (equation 1) is stored for each grid cell as part of the iterative solution of equation 1 for S_{supp} . If the vertical pressure gradient would change by more than 10% in a time step, a stand-in vertical velocity is calculated that limits the change to 10%, as

$$\Delta \phi_{\text{imposed}} = 0.1 \left(P_{\text{excess,j}} - P_{\text{excess,j+1}} \right) \left(\frac{\partial S_{\text{supp}}}{\partial \phi} \right)^{-1}$$

The idea is that a thin sandy layer would quickly drain, reaching zero excess pressure gradient, and the stand-in velocity is designed to achieve that end in a stable way even if the time step for Darcy flow would need to be much shorter. The 10% coefficient implies that it will take approximately 10 time steps (5 years) for the layer to relax to a drained condition (any faster leads to numerical instability). Most of the vertical flow in the model however is well-resolved in time, leaving the flow limiter for a few "emergency" grid points and times.

Total fluid flow including sedimentation and grid stretching

Darcy flow is defined relative to the solid grains. This is a different definition of fluid flow from the convenient and usual metric in sediment diagenesis models, which is defined relative to the sediment-water interface, which we will call w_{seafloor} . Formation of methane hydrate in models is very sensitive to the value of w_{seafloor} , upward carrying methane into the stability zone from below, or downward away from the stability zone [Buffett and Archer, 2004].

At the sediment surface, the two definitions of vertical velocity differ because the sediment surface is moving as solid and material accumulates. The burial of fluid in the pores between the sedimenting solid constitutes a vertical flow across the sea floor, when the reference frame is the moving sea floor itself. At the sedient surface,

$$W_{\text{seafloor}} = W_{\text{Darcy}} + \phi_0 \cdot \text{SedRate}$$

where SedRate is the bulk sedimentation rate and ϕ_0 the porosity of freshly sedimented material.

Beneath the sea floor, the calculation of w_{seafloor} requires two additional terms. One is the movement from one grid cell to another as the vertical grid in the model stretches to fill the expanding sediment column (explained above). The fluid component of this bookkeeping "flow" is calculated as the change in elevation of a grid boundary in a time step, multiplied by the porosity.

$$\mathbf{w}_{\text{bur}} = -\frac{\partial \mathbf{Z}_{\text{boundary}}}{\partial \mathbf{t}} \cdot \mathbf{\phi}$$

where $\partial z_{\text{boundary}}/\partial t$ represents the movement of the cell boundary.

For subsurface grid points, w_{seafloor} is calculated as

$$W_{\text{seafloor}} = W_{\text{Darcy}} + W_{\text{bur}} + \frac{d}{dt} (z_{\text{boundary}} - z_{\text{seafloor}})$$

The second term represents the relative movements of the seafloor and the depth of the subsurface grid cell in question. Like w_{bur} and w_{Darcy} , w_{seafloor} is defined on the cell faces. Numerically, the velocities are combined before any fluxes are calculated, using the net flow across a grid boundary within a given time step, rather than advecting material back and forth in multiple operations which would result in needless numerical mixing.

A subsurface value of $w_{seafloor}$ is defined relative to the moving sea floor, so that if a subsurface parcel were flowing upward just quickly enough to remain at a constant depth below the seafloor, its $w_{seafloor}$ would be defined as zero. Sections of w_{Darcy} and $w_{seafloor}$ are shown in Figure 7, and values at the sediment surface in Figure 8.

When vertical permeable channels are added to the simulation, the channels take most of the compaction flow. The horizontal anisotropy applied to the permeability allows the sediment columns in between the channels to vent fluid into the channels for expulsion. Interestingly for the formation of methane hydrate discussion to come, the channeled simulation finds strongly downward w_{seafloor} in the grid cells in between the channels. These cells have vented their upward Darcy flow into the permeable channels, leaving the burial flux of fluid with sedimentation to drive w_{seafloor} in the downward direction, where it brings methane-depleted fluid into the hydrate stability zone from above, rather than methane-rich fluid from below.

Carbon Cycle

Deposition

The carbon cycle of the deep biosphere is driven by the rain of particulate organic carbon (POC) to the sea floor. Organic carbon deposition and concentrations in surface sediments of the ocean depend on primary productivity in the ocean, runoff of terrestrial POC in some places, degradation in the water column, offshore transport, and degradation in the surface sediments which depends on the oxygen concentration of the overlying water and perhaps the temperature.

Rather than mechanistically simulating that complexity, the POC concentrations and characteristics of the sedimenting material in these simulations are determined as a simple function of water depth of initial deposition, and time, allowing us to control POC deposition as a boundary condition to the model rather than simulating it as a prognostic output. We used the POC parameterization to drive a scenario in which periods of high sea level have enhanced carbon preservation, a cartoonish representation of the transition from the Cretaceous to the present [Hunt et al., 2002].

Although the link between ocean oxygenation and POC deposition is not mechanistically clear or straightforward to model, we concoct and drive the model with an ocean oxygenation "index" that varies from 1 to 4, with 1 the most oxic and 4 anoxic. Ocean state 1 is called "Atlantic", and the sedimenting POC concentration profile is comparable to sediment surface POC concentration off Cape Hatteras today [*Mayer*, 2002]. In this state the POC concentration is at most 2% dry weight POC, with an H/C ratio of 0.7. The anoxic end-member buries up to 5% POC, with an H/C ratio of 2 (Figure 3b).

The ocean oxygenation index covaries with sea level and the deep ocean temperature, resulting in a scenario in which a warm high-sealevel high POC deposition ocean 100 million years ago transitions into the characteristics of the Atlantic ocean by the end of the simulation. For a given grid cell and time step, the POC concentrations are interpolated in depth, then interpolated in the ocean state, so that carbon deposition can vary smoothly in time, in synchrony with changes in eustatic sea level.

The impact of a persistent change in ocean oxic state on hydrate abundance was gauged in simulations with time-uniform forcing, called *Atlantic*, *Pacific*, *OMZ*, and *Anoxic*. In the time-dependent simulations, high POC concentrations accumulate in a time of high sea level in the simulation, which becomes a buried feature by the end of the simulation (Figure 9 left).

Biological Degradation

Biological degradation is taken to be a function of temperature (T, °C), sediment age, and dissolved methane concentration.

$$k_{resp} = 0.5 \text{ yr}^{-1} \cdot e^{-\frac{Ea}{RT}} \cdot H_{smooth} (50^{\circ}\text{C} - \text{T}, 10^{\circ}\text{C})^{6}$$
$$\cdot e^{-1.1 \log(\text{Age}) - 0.5} \cdot \left(\frac{35 \text{ mM}}{[\text{CH}_{4}] + 35 \text{ mM}} \right)$$

where k_{resp} is a first-order rate constant for POC degradation. The activation energy Ea is 70 kJ/mol, resulting in a tripling in the rate with a temperature increase of 10 °C. This is a relatively moderate temperature sensitivity; Burdige [2011] argues that it could be 200 kJ/mol, which would increase the reaction rate by a factor of 24 for an increase in temperature by 10 °C.

The second term represents inhibition of biological activity above a critical temperature of 50 °C. The formulation uses a smoothed "Heaviside" function to shut down the respiration reactions above a threshold temperature of 50 °C. This function is used several times in the SpongeBOB formulation, defined as

$$H_{\text{smooth}}(x, \Delta x) = \frac{1}{\pi} \left(0.5 \ \operatorname{arctan} \left(\frac{x}{\Delta x} \right) \right)$$

This function provides a smooth transition between the on and off states (1 and 0), rather than the discontinuous step function of the true Heaviside function. Raising the function to a power, as done here, tightens the transition, here intended to mimic a fairly strong shutdown of respiration rates when it gets too hot.

The third term represents the correlation between carbon age and its reactivity, mimicking the astonishingly wide correlation observed in nature [Hedges and Keil, 1995; Middelburg et al., 1997]. The correlation between age and reactivity does not imply a direct or simple causation, and "priming experiments" [Bianchi, 2011], demonstrate that the age parameterization cannot be universally true. But the correlation between age and reactivity captures many orders of magnitude of variation, and seems essential to account for [Wallmann et al., 2006].

The effect of the "metabolite inhibition" effect as proposed by Wallmann et al. [2006] is represented in the last term in the formulation. Wallmann included dissolved inorganic carbon concentration in addition to methane, but for the sake of model stability and development we took the simplifying liberty of excluding DIC inhibition of methanogenesis.

Respiration is concentrated in the top 2 km of the sediment column, limited by temperature (Figure 10). It is also generally focused offshore of the shelf break, except during periods of very high sea level, during which POC can accumulate in the shelf sediments. When sea level falls, the biologically available POC in the shelf sediment burns out, leaving most of the biological activity in slope sediments, and in particular in the sediment depocenter just off the shelf break.

Model sensitivity to the respiration formulation was gauged by varying the "labile fraction" of the POC (cases *Bio 10%*, *Base* (which is 50%), and *Bio 100*%).

Biological Methanogenesis

The respiring carbon is assumed to produce DIC and molecular hydrogen, a transient reducing agent. If sulfate (SO_4^{2-}) is present in the pore water, the hydrogen reduces and consumes it; otherwise, it reacts with CO_2 (dissolved inorganic carbon or DIC) to produce methane in reactions

$$CH_x + 2 H_2O \rightarrow CO_2 + (2+x/2) H_2$$

followed by CO₂ reduction

$$1/2 \text{ CO}_2 + 2 \text{ H}_2 \rightarrow 1/2 \text{ CH}_4 + \text{H}_2\text{O}$$

where the variable x denotes the relative concentration of POH / POC in the reacting organic matter.

The maximum efficiency with which the biologically produced CO_2 can be converted to CH_4 is determined by conservation of carbon, hydrogen, and oxygen from the organic matter to methane plus CO_2 , realizing that supplemental oxygen and hydrogen comes from H_2O . This maximum CH_4 production efficiency is

$$R_{CH_4} = \frac{1}{2} + \frac{1}{8} \cdot \frac{POH}{POC}$$

which works out to 62% if the H/C ratio of the reacting organic matter is 1. However, carbon isotopic systematics and data, described below, will restrict the efficiency of CH_4 production somewhat, as if about 20% of the H_2 reacts with oxidized mineral phases such as Fe_2O_3 rather than with CO_2 to make methane, leaving an overall methane production efficiency of about 50%.

Methane is oxidized in the presence of SO_4^{2-} by the anaerobic oxidation of methane (AOM) reaction. The reaction in nature is fast enough to permit just a very narrow depth range of overlap between the high- SO_4^{2-} pore waters in the surface sediment column and high- CH_4 waters below [*Burdige and Geiskes*, 1983; *Chatterjee et al.*, 2011; *Davie and Buffett*, 2001] f. The reaction in the model is based on whichever species is stoiciometrically limiting, with a effective rate constant of 5% yr⁻¹, unrealistically slow but numerically stable and able to eliminate simultaneous coexistence of SO_4^{2-} and CH_4 in any grid cell. SO_4^{2-} is only found in a few grid cells just inside the continental shelf break. The grid resolution of the model is too coarse to resolve the details of pore water SO_4^{2-} , so we leave an analysis of the behavior and impacts of SO_4^{2-} to future work.

Petroleum and DOC

The model produces petroleum based on chemical and thermal conditions. Petroleum formation in the real world is limited by the H/C ratio of the organic matter, proceeding if the H/C is greater than 1, and continuing until the ratio approaches 1. The temperature dependence of the rate constant is governed by an activation energy of 150 kJ mol⁻¹, with a pre-exponential constant of 1*10¹⁶/year. These parameters confine petroleum generation to a temperature window of 60-150 °C, as inferred from the distribution of oil [*Hunt*, 1995].

Petroleum genesis temperatures are too warm to allow bacterial consumption, but we gauge the potential impact of bacterial respiration of petroleum by assuming that 10% of the oil is upwardly mobile at a rate of 1 m per thousand years. If it reaches cool enough conditions it is consumed by bacterial respiration, following the temperature and solute concentration dependence (but not the POC age dependence) of bacterial POC degradation kinetics above. The rate of methane production by this process of "secondary respiration" is shown in Figure 10b to be deeper than POC respiration, which is more confined to the upper sediment column. The model also assumes "sloppy feeding", in which 10% of biological POC consumption goes to dissolved organic carbon (DOC). Simulations to determine the importance of these processes in the model are called *No Petro* and *No DOC*.

Thermal Methanogenesis

Methanogenesis from thermal degradation of organic carbon proceeds according to an Arrhenius equation with an activation energy of 230 kJ mol⁻¹ and a pre-exponential rate constant of 4 · 10²¹ yr⁻¹, which produces a lifetime of a million years at about 160 °C. In contrast to the biological rate of methane production, the rate constant for thermal methane generation is very well constrained and consistent in field data, driven by the hunt for petroleum [*Hunt*, 1995].

The rate of methane production is taken to be this rate constant multiplied by the concentration of particulate organic hydrogen, POH. The POC inventory decreases by 1/4 of the decrease in POH during methanogenesis. The distribution of the H/C ratio in the sediment column (Figure 9 right) is primarily driven by the effects of changing ocean chemistry (anoxia) and thermogenic degradation. The H/C ratio approaches zero as the carbon matures, and the decrease in H directly limits the production of methane in the model.

Methane Bubbles and Hydrate

The dissolved methane concentration is compared with the solubility of dissolved methane, relative to the two possible other phases of gas bubbles or hydrate, in Figure 11. Hydrate or bubbles both release or take up methane from the dissolved phase. Disequilibria in the three-phase methane system is restored toward equilibrium by conversion to or from bubbles or hydrate with a numerically-tractable but ad-hoc time

constant of 1000 years. This unrealistically slow time constant, a compromise that should doubtless be higher? for future detailed simulations, permits the dissolve methane concentration to reach unrealistically high supersaturation values ($\Omega = 1.2$ in Figure 11), but since dissolved methane is a transient intermediate species in the pathway POC to CH_4 to bubbles, the rate of bubble formation will be determined by methaneogenesis, and the dissolved methane concentration in the steady state will grow in to fulfill that balance, leaving only the question of whether the altered CH_4 concentration changes some other aspect of its dynamics, by diffusing more quickly for example.

When the precise depth of the stability boundary, determined by depth interpolation, falls within a grid cell, the non-gaseous CH₄ is allocated into bubbles and hydrate in proportion to the volumes of the two zones, and the concentrations of the phases, for purposes of advection and diffusion, are calculated using the partial grid cell volumes. This prevents a numerical artifact that otherwise appears as the depth of the stability boundary crosses through the coarse numerical grid of the model.

The methane cycle in the SpongeBOB requires gas-phase mobility in the sediment column. If bubbles are not allowed to migrate, gas completely fills some parts of the sediment column pore space (Figure 12c). Gas volume in the pore space of the model currently and unrealistically has no impact on the fluid pressure or flow, allowing unlimited gas accumulation to fill up to 200% of the original pore volume. That much methane simply has to go somewhere.

In a basin-scale sense, gas mobility comes as no surprise. The view from petroleum geology is that, on geologic time scales, sediment columns are permeable to gas, except under permafrost soils and evaporites, which overlay the largest gas accumulations in the world in Siberia and the Middle East, respectively. Other parts of the world, for example the North Sea, are vastly enriched in oil relative to the expected amount of gas were it retained [*Hunt et al.*, 2002]. On the smaller spatial scales of structural hydrate deposits such as at Hydrate Ridge [*Treguer et al.*, 1995], gas migration is also apparent. Gas migration is a fundamental part of the Liu and Flemings [2007] model to explain the seismic "wipeout zones" and observed methane emissions from the seafloor within the hydrate stability zone [*Hill et al.*, 2004].

In contrast, for modeling "stratigraphic" methane hydrate deposits typical of most of the continental margin, gas migration is typically neglected or assumed to be small [Burdige, 2011; Chatterjee et al., 2011; Davie and Buffett, 2001; Davie and Buffett, 2003a; Davie and Buffett, 2003b; Garg et al., 2008; He et al., 2011; Liu and Flemings, 2007; Xu, 2004]. These one-dimensional methane hydrate models span a domain a few hundred meters below the stability zone boundary and assume that the bubbles there are immobile. This is not a priori wrong; it could be that gas transport is only important in the structural hydrate deposits, more obviously associated with faults and channels. However, we do find that methane hydrate formation in SpongeBOB is strongly driven by upward migration of gas, the same gas flux as is required to evacuate excess gas from the deep sediment column.

Gas tends to form if the dissolved concentration of the gas exceeds the Henry's law solubility, a function of the pressure of the gas. Surface tension raises the pressure of the gas phase somewhat over that of the pore fluid, but the differences are small relative to the overall pressure, so the impact of excess gas pressures on methane solubility have been neglected in the model. Bubbles typically grow, in cohesive sediments, by fracturing the sediment matrix [Boudreau et al., 2005; Jain et al., 1995], requiring a further excess pressure that is a function of sediment properties such as fracture strength. With a continuous source of methane, dissolved methane concentrations can rise until Henry's law produces gas at whatever pressure is required to create the bubble.

It is another issue to understand the upward mobility of the gas, as opposed to its growth in place. Even at pressures near the base of the sediment column, the compressed gas is buoyant, manifesting as an upward body force that intensifies as the bubble rises and decompresses. One source of excess pressure at the top of a bubble is the hydrostatic pressure difference between the top and the bottom of the bubble [Jain and Juanes, 2009]. However, for a bubble several kilometers deep in the sediment column to break through the sediment column by the difference in hydrostatic pressure, it would have to several kilometers tall for bubble-top excess pressures to break through the sediment column. Presumably gas migration from the deep sediment resembles gas blasting through a pipe more than it does a long bubble contemplating rising. The gas can manufacture its own means of escape by building up to lithostatic pressure, possibly venting episodically.

Laboratory experiments find a critical gas saturation (pore volume fraction) of about 10% where gas begins to mobilize [Leas et al., 1950], and values of 1-2% are often assumed in models for natural gas in wells [Reagan and Moridis, 2007]. Higher gas saturations connect bubbles together, crossing the hydrostatic pressure gradient as well as providing a pathway for the gas to flow, which is important because gas is much less viscous than pore water, even when it is compressed. The standing stock of bubbles in the sediment column is another transient intermediate reservoir for methane in the pathway POC -> dissolved methane -> standing gas bubbles -> migrating gas. In the steady state, the standing gas bubble reservoir will find the concentration at which the methane fluxes balance. The essential balance is between methanogenesis and upward gas flux in the model. A bias in the bubble flux parameterization would result in an incorrect standing bubble concentration but no error in the bubble flux itself, unless the bubble concentration leads to some distortion of the methane budget, such as by burial of gas.

We have adopted a parameterization of bubble migration that depends on the bubble fraction in pores

$$CH_4$$
 loss = $H_{smooth} (F_b - 10\%, 5\%)^3$

where again the H_{smooth} is a smoothed version of a Heaviside function. The bubbles escape as their concentration approaches or exceeds a critical pore volume fraction of 10%. In practical terms, the bubbles in SpongeBOB only fill at most 1-4% of the pore

space, because the loss rate, however slow, is able to keep up with production, even when the saturation value is much less than the parameterized critical value.

Rather than resolve the rise of the gas through time, we instantaneously redistribute some of the CH₄ lost as migrating bubbles into the dissolved phase to grid cells above where the bubble first started moving. This approach is analogous to the treatment of sinking plankton in the water column of many ocean chemistry models, in which the sinking organic matter is assumed to reach some mean distance before instantaneously redissolving. The bubbles once in motion rise through the sediment until they encounter pore water that is undersaturated with respect to dissolved methane, at which point the upward flow is attenuated with height as

$$CH_4$$
 flux(z) = CH_4 flux(0) · $e^{-z/z_{scale}}$

where the value of the scale height parameter z_{scale} is taken to be 500 meters in the high-resolution run, but varied to 100 and 2000 m in two of the sensitivity runs (called $Bubb_100m$ and $Bubb_2k$). The methane cycle in the model in general, except for the standing bubble inventory, is not very sensitive to the bubble mobility parameterization, but the methane hydrate inventory is extremely sensitive to what happens to the gas after it begins moving.

The distribution of bubbles resulting from these assumptions is shown in Figure 9. A standing crop of bubbles is sustained by respiration in the upper sediment column, and there is a column of bubbles carrying methane from the thermogenic zone up toward the sediment surface. Thermogenic methane in the model systematically evades the hydrate stability zone, because the sediment column is only deep enough to reach high enough temperatures when the top sediment column has nearly reached the sea surface, so that there is in general no hydrate stability zone over thermal methane production in the model (Figure 9). This geophysical mechanism could lead to a general bias toward biogenic rather than thermogenic methane in deep margin hydrate deposits.

DIC and Subsurface Weathering

Measurements of the titration alkalinity of pore waters (a reasonable facsimile to DIC under pH-neutral conditions) from Blake Ridge reach fifty times higher than the ocean value within the top few hundred meters, but then either decrease to half that by the deepest samples measured, about 700 mbsf (Rodrigous). A similar decrease with depth was documented from the West African margin [Sivan et al., 2007].

In the model, if CO_2 is allowed to accumulate in deep sediment pore water without a compensating sink, the DIC concentration of the model increases monotonically to bedrock (Figure 13 b and c). The difference can be explained as a chemical weathering or carbonate formation reaction such as

anorthite + CO_2 -> kaolinite + $CaCO_3$

Throughout the upper sediment column, the equilibrium concentration of dissolved CO₂ based on this equilibrium is much lower than the measured concentrations. At low temperatures typical of the surface of the Earth or the upper sediment column this reaction is too slow to reach equilibrium, but at a high enough temperature the reaction proceeds quickly as the metamorphic decarbonation reaction. We treated this reaction as a first-order uptake in DIC. This neglects the impact of the alkalinity or pH of the solution on the CO₂ concentration, but the constancy of porewater alkalinity / DIC ratios [Sivan et al., 2007] and pH make this seem a reasonable assumption. The reaction as written will affect the concentration of calcium, Ca²⁺, but it is not calcium limited unless there is a shortage of unweathered igneous calcium-bearing rock. The model assumes equilibrium with CaCO₃, rather than try to resolve the porewater chemistry into its components alkalinity, calcium, and DIC.

The kinetics of DIC uptake are taken to be a function of temperature as

$$k_{_{urey}} = k_{_{urey_0}} e^{\frac{-Ea}{RT}}$$

where the activation energy Ea is 120 kJ mol⁻¹, increasing the rate constant by a factor of five in response to a temperature increase of 10 °C. The slowest pre-exponential multiplier we tried was 3*10¹³ yr⁻¹, which produces a reaction timescale of about 10⁸ years at 15 °C, so that essentially nothing happens in surface sediments, to a time constant of 1000 years at 100 °C. We also show profiles of model DIC concentration with faster uptake kinetics, by factors of 3 spanning two orders of magnitude.

The model profiles (Figure 13 b) are compared with alkalinity data for ODP sites 994 and 997 on the Blake Ridge [Rodger, 2000]. Both the peak concentration and the depth of the peak are best simulated by the model with an intermediate rate constant of the range we tried, $3 * 10^{14} \text{ yr}^{-1}$.

For all runs with nonzero Urey reaction rate constant, there is a drawdown in DIC concentration with depth. The similarity to measured concentrations lends strong support to the significance of this reaction to sediment pore water DIC. It is no surprise that carbonate formation occurs within the sediment column [Rodger, 2000], but it is interesting to note the impact that this process has on measured DIC concentrations.

Also in all weathering runs, the concentration of DIC in sediments warmer than 100 °C in the model is effectively controlled by equilibrium, determined from thermodynamic data for the phases in the weathering reaction. The highest DIC levels in the simulation are found in the hottest deepest sediments, due to the effect of temperature on the equilibrium condition, the beginnings of metamorphic decarbonation (Figure 13a). At temperatures between about 50 to 100 °C, the equilibrium DIC concentration is relatively low, but the reaction kinetics in the model are still fast enough for equilibrium to be reached, pulling DIC concentrations down in that depth range, and isolating the high DIC concentrations in the hottest sediments from reaching the sediment surface.

Carbon Isotopes

The model tracks parallel tracers representing the stable carbon isotopes in DIC and CH₄. Using methodology I learned from Ernst Maier-Riemer for ocean modeling [*Maier-Reimer et al.*, 1982], the isotopologs undergo the same advection, diffusion, and chemical reaction operators, but with offsets in the reaction rates corresponding to the fractionation factors. Rather than try to numerically resolve the large relative concentration differences between the isotopes, the isotopic standard ratio for all isotopic systems is defined to be 1, meaning equal concentration of both isotopes at 0 °/ $_{oo}$. The results, presented in °/ $_{oo}$ deviations from the standard ratio, are directly comparable with isotopic ratios of natural samples presented relative to their separately defined standard ratios (PDB for δ^{13} C, for example).

The isotopic boundary conditions are that the DIC in the ocean is assumed to be 0 $^{\circ}/_{\circ \circ}$, and organic matter is assumed -25 $^{\circ}/_{\circ \circ}$. Organic matter is respired to DIC without fractionation. CO₂ reduction carries a fractionation of -90 $^{\circ}/_{\circ \circ}$ at 0 $^{\circ}$ C, dropping to about -55 $^{\circ}/_{\circ \circ}$ at 50 $^{\circ}$ C [Whiticar and Faber, 1986]. Thermogenic methane is assumed to have the isotopic composition of organic matter. Diffusion and anaerobic oxidation of methane (AOM) also fractionate the carbon isotopes.

The isotopic composition of DIC in the sediment column (Figure 14) is lightened by degradation of organic matter, but then pulled heavier by extraction of isotopically light CH_4 . As a result, the isotopic composition of DIC or alkalinity in porewater samples starts near $0^{\circ}/_{\circ\circ}$ at the sediment surface, then trends light in the near subsurface due to respiration in the non-methanogenesis zone, then gets heavier, reaching a maximum value of about $+10^{\circ}/_{\circ\circ}$ (Figure 14). These features are mimicked by the model, but the degrading effect of the coarse grid resolution can clearly be seen in the broadening and attenuation of the subsurface negative isotopic excursion.

Sivan et al [2007] interpret the maximum $\delta^{13}C$ of the DIC as an indicator of the rate of methanogenesis, but it turns out to be more sensitive in the model to the relative partitioning of respired carbon between DIC and methane. If the respired carbon has an isotopic composition of -25 °/ $_{oo}$, and the fractionation between CH₄ and DIC at some temperature were 60 °/ $_{oo}$, for example, then a balance of about 50% methane versus DIC would tend to produce DIC at an isotopic composition of +10 °/ $_{oo}$. In time, this source composition dominates the pore water value. Model experiments did not show a sensitivity of $\delta^{13}C$ in DIC to the overall methaneogenesis rates, but it does respond to the relative CH₄ production efficiency. The best fit seems to require some loss of H₂ in the pathway, as if by reduction of mineral phases.

Iodine 129

Iodine 129 has been used as a tracer for the age of the degrading organic matter in a sediment column [Fehn et al., 2000]. Iodine 129 is produced by cosmic ray spallation in the atmosphere and it has a 22.8 million year e-folding lifetime. Iodine, radio and stable, is incorporated into biological material, and it is released to the pore water as the carbon

degrades biologically. Iodine released by older organic carbon will have less Iodine-129 because of radioactive decay. In the model, both stable and unstable iodine are carried with the pore flow and are subject to diffusion along with the rest of the dissolved tracers.

Fehn et al [2000] measured I^{129} ages 55 million years old in pore waters of solid sediment that was 1.8 to 6 million years old, a difference of about 50 million years, which they interpreted as indicative of 50 million year old carbon degrading.

Iodine 129 seems like a very rich tracer in the SpongeBOB model, indicative of the history of the accumulation of the sediment column and the expulsion of fluid from it (Figure 15). Variations in I¹²⁹ age arise from compaction-driven fluid flow. When fluid flow is prevented in the model, the iodine ages correspond to the sediment ages (Figure 15d). When fluid flow is enabled but without the permeable channels (Figure 15b), there are iodine ages 60 million years older than the solid ages at its maximum, located in mid-column sediments under the deep continental slope, along with I¹²⁹ ages less than the solid age in its minimum under the continental shelf. This pattern persists when permeable vertical conduits are available, with a superimposed pattern that the I¹²⁹ ages within the conduits are older than in the surrounding low-flow columns (Figure 15c). The old I¹²⁹ ages "grow in" as the continental margin ages (compare Figures 15a and c).

The iodine ages in the depth range of Blake Ridge in SpongeBOB never get as old as Fehn et al [2000] measured at Blake Ridge. The values in the conduits reach 20 million years old, while Fehn measured 50 million years. The model results suggest that the old I¹²⁹ ages measured at Blake Ridge probably have more to do with pore fluid flow than with the age of respiring organic carbon. They also suggest that Blake Ridge pore water geochemistry may be affected by focused subsurface flow, such as crudely approximated by the model, and thus would over-represent the abundance of methane hydrate along margin sediments overall (which cannot all be the subject of focused upward flow).

Model Sensitivities

Scenario Runs

In addition to the "Base HR" and "No Conduits HR" cases, a series of sensitivity simulations were done, at lower spatial resolution of only 10 grid points horizontally instead of 40. The setup, intent, and summary of the conclusions from the sensitivity runs are described in Tables 1 and 2. One suite of simulations was forced by time-invariant forcing, comprising sea level, ocean "oxic" state (POC deposition patterns), and temperature, with variations as described in Table 1. These allow a clearer diagnosis of the model sensitivity to these variables. Another suite of simulations imposes time-varying forcing shown in Figure 3, as described in Table 2. The time evolution of sea level, POC inventory, methanogenesis rates, and hydrate inventories are shown in Figure 16c for the two representative cases, the *Base* case, which including the sea level changes, and case *TO*, which does not.

Sensitivity to Ocean Temperature

The effect of ocean temperature on model hydrate inventory is shown in Figure 17a. These are from time-invariant suite of runs, with temperature a function of water depth only. The inventory of hydrate decreases roughly linearly with temperature, finally reaching zero hydrate with a warming of 8 °C. The active margin simulations (presented in the companion paper) have a similar temperature sensitivity to the passive case. The SpongeBOB model is significantly less sensitive to temperature than the Davie and Buffett [2001] one-dimensional model (Figure 17a). These are essentially equilibrium calculations, comparable to those of Buffett and Archer [2004].

The difference is probably due to the fundamentally different processes controlling vertical methane fluxes between the two models. The 1-D model relies on the transport of dissolved-phase methane into the stability zone. In a warmer ocean, the stability zone gets thinner, which increases the diffusive loss of dissolved methane in that model by requiring the diffusive gradient to be steeper. This had the impact in the Buffett and Archer [2004] simulations of restricting the formation of hydrate to regions of higher POC content, reducing the ocean's ultimate equilibrium inventory of methane nearly completely with 3 °C of warming. Methane trapping in the stability zone, when it is transported by bubbles in SpongeBOB, would also be affected by a change in the stability zone thickness, but the mechanism is different and so it is not surprising that the two models would have different temperature sensitivity.

Sensitivity to POC

The model sensitivity to POC deposition was gauged using time-invariant forcing, by varying the "ocean oxic state" variable through the spectrum from "Atlantic" (low POC concentrations in surface sediments, state=1) to "Pacific" (2), "OMZ" (3), and "Anoxic" (4). The range in sediment surface POC concentrations is intended to bracket something of the range in the ocean, and Figure 17b shows the sensitivity to that range. Changes in POC deposition have a much greater capacity to increase the hydrate inventory than changes in temperature could, because the ocean is near freezing and it can mostly warm from here.

The ability of high POC to overcome warm temperatures to produce hydrates in the geologic past [*Gu et al.*, 2011] is not supported by these model results. The time-dependent simulation has no hydrate through the warm, high-POC time 100 Myr ago (by coincidence year 100 million in the simulation) until the ocean cools after 150 Myr, whereas the time-independent simulation begins to accumulate hydrate after only 50 Myr (Figure 17c).

The sensitivity of the hydrate inventory to POC can also be seen in the model response to changes in the labile fraction of the sedimenting POC (Figure 17c). In practice, the respiration zone is thin enough that the POC is not often limited by depletion of the labile POC fraction, so the labile fraction acts like a scaling of an apparent bulk POC degradation rate constant. With the tight correlation between organic carbon age and

reaction rate [Middelburg et al., 1997], this range in effective POC degradation rate constants might optimistically be viewed as a range in the uncertainty in the POC degradation rates.

The sensitivity to biologically labile POC in particular shows that biogenic methane production is the important factor in the model. A model sensitivity run with thermogenic methane production disabled (called *No Therm*) corroborates this by the lack of much change in the hydrate inventory. The simulation *No Petro*, which disables secondary respiration of migrating petroleum, produces no hydrate, a strong effect to be sure. However, the absence of hydrate without petroleum migration in the model does not mean that petroleum is the whole story. The hydrate inventory is so responsive to reasonable changes primary POC respiration that were we to neglect secondary petroleum respiration in the model, the primary respiration could be increased enough to tip back over to the point of preserving hydrate.

Sensitivity to Bubble Transport

The hydrate inventory in the model depends critically on methane transport by rising bubbles through the sediment column, and redissolution if the bubbles enter an undersaturated condition. This mechanism is novel to SpongeBOB, not shared by the Davie and Buffett [2001] or most other 1-D methane hydrate accumulation models. If bubble transport is disabled, hydrate disappears from the results (Figure 16). Figure 17d shows the extreme model sensitivity to the redissolution height scale, which determines what fraction of the methane will be recaptured by redissolution rather than escape directly to the sea floor. When a scale height of 2 km is imposed, hydrate nearly disappears from the simulation. Shortening the height scale from 500 m to 100 m increases the hydrate inventory by a more than a factor of four. Clearly this is a parameter to which the model is very sensitive, and further work will be required to capture the complexities of the real world in a numerical model.

Pore fluid flow heterogeneity

Old iodine 129 ages in pore waters at Blake Ridge, relative to the age of the solids, can only be reproduced in the model as resulting from heterogeneity in the pore fluid flow field, as crudely mimicked in the code by the vertical high-permeability channels. This conclusion is reinforced by the heterogeneity in penetration depth for dissolved sulfate, SO_4^{2-} [Borowski et al., 1996], which seem to show a bimodal distribution, with some sites where the methane/sulfate boundary is just a few 10's of meters, and others where sulfate reaches hundreds of meters or more. The shallow sites correlate with the presence of methane hydrate, just like in the model where the hydrate is found. The importance of the channels justifies the imposition of an upward flow in the Davie and Buffett one-dimensional model [2001], which was interpreted as representing a zone of upward flow, analogous to the channels here.

Other model (in)sensitivities

Most of the other scenarios had a fairly minor impact on the hydrate inventory. Thermogenic methane is not a significant contributor to the pelagic hydrate methane pool. The permeable channels do not determine the solution, nor does sediment erosion and redeposition, or the value of the critical seafloor slope that drives it.

Conclusions

The SpongeBOB model treats the coastal margin methane cycle from a much larger perspective than previous models of methane hydrate accumulation in margin sediments. The model domain spans to bedrock, closing the budgets for pore fluid and carbon, and into the onshore – offshore dimension. When heterogeneities in the sediment permeability are introduced, the lateral dimension becomes an integral part of the dynamics, rather than the model being a sort of palisade of one-dimensional systems. Also, rates of sedimentation are governed by the evolution of the water depth overlying the sediment column, so that sedimentation variations in time are controlled by things going on in the lateral dimension. Its two-dimensional formulation therefore allows the model to span geologic time.

Bubble migration through the sediment column is an important part of the methane cycle in the model. Viewed from a basin perspective, it is no surprise that methane is lost from the deep sediment column. Another related but separate question is how much of the methane gas lost from the deep sediment column is recaptured to the dissolved phase.

The SpongeBOB model departs from most previous models of methane hydrate accumulation [Burdige, 2011; Chatterjee et al., 2011; Davie and Buffett, 2001; Davie and Buffett, 2003a; Davie and Buffett, 2003b; Garg et al., 2008; He et al., 2011; Liu and Flemings, 2007; Xu, 2004] by postulating that this upward bubble flux might play a role in the stratigraphic methane hydrate deposits on passive continental margins. It turns out that the model methane hydrate inventory depends very strongly on the scale height for methane redissolution within an undersaturated zone. The fate of methane gas within the sediment column clearly needs further attention and development.

Heterogeneity in the vertical flow field, simulated very crudely in SpongeBOB as vertical channels of increased permeability, affect chemical tracers such as iodine 129 and SO_4^{2-} , as well as focusing the formation of methane hydrate in the channels. Old I^{192} ages in hydrate-bearing sediments [*Fehn et al.*, 2000] and the observed correlation between SO_4^{2-} penetration depth and presence of hydrate [*Borowski et al.*, 1996] are consistent with the conclusion that hydrate distribution is steered by upward flow.

The abundance of methane hydrate in the model is poised on a knife edge, responding very strongly to changes in the degradation rate of organic matter, to gas transport and redissolution in the sediment column, and to ocean and subsurface temperatures. For this reason it is currently impossible to calculate the global inventory of methane hydrate from modeling first principles. The goal of the process-oriented model SpongeBOB is to

weigh the relative sensitivities of the hydrate inventory to potential changes in forcings in the past and future, and to identify processes that need to be understood better in the real world, such as gas transport.

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Table 1. Model scenarios with uniform sea level and POC deposition state (ocean oxic state = 3).

ТО	No temperature offset	Higher hydrate than the Base case because of higher POC rain in the last bits of the simulation
T2, T4 etc.	2, 4, 6, 8, and 10 °C temperature increases	Weaker temperature sensitivity than the 1-D models (Figure 17)
Atl, Pac, Anoxic	"Atlantic" (ocean state 1) "Pacific" (state 2), and "Anoxic" (state 4) POC rain forcings, shown in Figure 3. Default is state 3 in the T* runs.	Stronger POC dependence for this range of conditions than T dependence

Table 2. Simulations with a time-varying sea level and ocean oxic state (POC deposition) ${\bf P}$

		T
Simulation name	Characteristics	Salient results
Base	Base case	
Lores Base	Lower resolution version of the full base case for comparison with sensitivity runs	Nearly 50% less hydrate than the higher resolution case
No Conduits	Vertical permeable channels eliminated	Similar hydrate inventory to Lores Base
Labile 10%, Labile 100%	Changed the biologically-available POC fraction from 50% in base to 10% and 100%	Nearly linear dependence of hydrate inventory on rate constant.
Geotherm 120, 90, 50	Geothermal heat flux in mW m ⁻² relative to base of 70	No hydrate above 90
No Bubb Mig	Bubble migration disabled	Eliminates hydrate
Bubb Mig 100m, Bubb Mig 2km	Redissolution height scale for rising bubbles from 500 m in base to 100 m and 2km	Strong impact on hydrate inventory
Lo Slope	The maximum sea floor slope is taken to be 3% instead of 6% in base	No strong hydrate inventory impact
No Slides	Landslides disabled	No strong hydrate inventory impact
No Thermogen	Thermal methanogenesis disabled	No strong hydrate inventory impact
No Petroleum	Petrogenesis disabled	Eliminates hydrate
No DOC	DOC production and	No strong hydrate inventory

respiration disabled	impact	
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Figure Captions

- 1. SpongeBOB in isostatic equilibrium. Colors are temperatures on the computational grid of the model, traced horizontally by the black lines. The thickness of the underlying crust grades from thin ocean crust on the right side of the domain into a thicker (and less dense) continental crust on the left. The transition between ocean and continental crust is smoothed over a spatial scale of 25 km. Local isostacy applies, on a time scale of 10,000 years and with spatial smoothing over 20 km. An animation of the simulation is archived in the Supplemental Material and can be viewed at http://geosci.uchicago.edu/~archer/spongebob_passive/fig1.atlantic.movie.gif
- 2. Sediment transport model results. A) Mean grain size ($\phi = \log_{10}(mm)$) in the sediment column half-way through (left) and at the end of the simulation (right). B) Snapshots of the sedimentation rate of particles of various sizes. C) Snapshots of the total sedimentation rate, with a depocenter just off the shelf break. An animation of this figure is archived in the Supplemental Material and can be viewed at http://geosci.uchicago.edu/~archer/spongebob passive/fig2.atlantic.movie.gif
- 3. A) Sea level time variation with time imposed on the simulation, and corollary changes in the "ocean state", which drive POC deposition concentration patterns as in (B), where Atlantic and Pacific represent conditions typical of the East or Northwest coasts of United States, "OMZ" is stronger oxygen minimum conditions typical of the Eastern equatorial Pacific, and Anoxic refers to anoxic conditions. Concentrations of POC in ocean sediments vary through this range and beyond; the intent here is to allow for variations that span something like the observed range.
- 4. A) Sediment column constructed from a seismic section off Cape Cod, from Kennett [1982], with cutout (b) resized for comparison with model results (C).
- 5. Porosity at the end of the simulation, section (A) and profile (B) taken on the heavy solid line in A: (solid line) full model solution, (long dashes) "drained" porosity contribution from β_1 , (short grey dashes) sum of drained porosity contribution from β_1 + β_2 .
- 6. Log of the permeability from the end of the simulation, (A) from the Base case and (B) the No Chan case with the vertical permeable channels disabled and at lower horizontal resolution.
- 7. Vertical velocity relative to the moving sea floor (w_{seafloor}) for the Base (A) and No Channels (B) cases, from the end of the simulation. An animation of the simulation is archived in the Supplemental Material and can be viewed at http://geosci.uchicago.edu/~archer/spongebob_passive/w_seafloor.movie.gif
- 8. Rates of vertical flow at the sea floor. Solid lines are the Base scenario, dashed the No Channel run. (A) Darcy flows (w_{Darcy}) are upward, while (B) flows relative to the moving

- seafloor (w_{seafloor}) can be found in both directions. An animation of the simulation is archived in the Supplemental Material and can be viewed at http://geosci.uchicago.edu/~archer/spongebob_passive/w_seafloor_sf.movie.gif
- 9. POC concentrations (dry weight percent) (A and C) and H/C elemental ratio of organic matter (B and D). From half-way through (A and B) the simulation, at a period of high sea level, and (C and D) at the end of the simulation, when sea level is relatively low, as today. An animation of the simulation is archived in the Supplemental Material and can be viewed at

http://geosci.uchicago.edu/~archer/spongebob_passive/poc.movie.gif

- 10. Methane sources in the sediment column. (A) From biological respiration of POC. (B) From biological degradation of migrating petroleum. (C) From thermal alteration of POC. An animation of the simulation can be viewed at http://geosci.uchicago.edu/~archer/spongebob_passive/ch4_src.movie.gif
- 11. Concentration of dissolved methane plotted in moles / m³ (A and B), and relative to saturation (C and D). Base model run (A and C), and No Channels (B and D). An animation of the simulation is archived in the Supplemental Material and can be viewed at http://geosci.uchicago.edu/~archer/spongebob_passive/ch4.movie.gif
- 12. Bubbles (A, C, E) and hydrate (B, D, F) (volume percent of pore space). From the Base model (A and B), the No Channels model (C and D), and with bubble migration disabled (E and F). An animation of the simulation is archived in the Supplemental Material and can be viewed at

http://geosci.uchicago.edu/~archer/spongebob_passive/bubbles.movie.gif

- 13. DIC concentrations, moles/m³. (A) Base case section, (B) Base case profiles compared with data in red from Blake Ridge ODP sites 994 and 997 [Rodgiruez et al., 2000] and the West African margin Sivan et al [2007], and (C) with Urey reactions that consume CO₂ disabled (note change in color scale from A). An animation of the simulation is archived in the Supplemental Material and can be viewed at http://geosci.uchicago.edu/~archer/spongebob_passive/dic.movie.gif
- 14. Carbon isotopic composition of methane from half-way through (A), and at the end of the simulation (C). (B) is the δ^{13} C of DIC at the end of the simulation, compared with data Blake Ridge ODP sites 994 and 997 [Rodgiruez et al., 2000] and the West African margin Sivan et al [2007] (D). The model lacks the resolution to capture the isotopically light spike just below the sea floor, but the deep isotopic composition matches well. Animations of this plot is archived in the Supplemental Material and can be viewed at http://geosci.uchicago.edu/~archer/spongebob_passive/del13.movie.gif.
- 15. Iodine-129 porewater age sediment age, in millions of years. With permeable channels 100 100 Myr of model time, (A) and 200 Myr (B), without permeable channels (C), and (D) with pore fluid flow disabled, t=30 myr instead of 140 myr for the other

simulations. Animations of this plot is archived in the Supplemental Material and can be viewed at http://geosci.uchicago.edu/~archer/spongebob_passive/i129.movie.gif.

- 16. Results of model sensitivity runs. (A and B) Hydrate inventory at the end of the simulation, (A) for the time-uniform forcing runs and (B) for the time-varying forcing runs. (C) The time trajectory of hydrate inventory, Base (time-varying) vs. T0 (time-uniform).
- 17. Summary of model sensitivities. Y-axis is the hydrate inventory relative to the appropriate base case = 1. (A) Temperature, (B) ocean oxic state (see Figure 3), (C) geothermal heat flux, (D) percent labile fraction of depositing POC, and (E) migrating bubble redissolutution scale height.

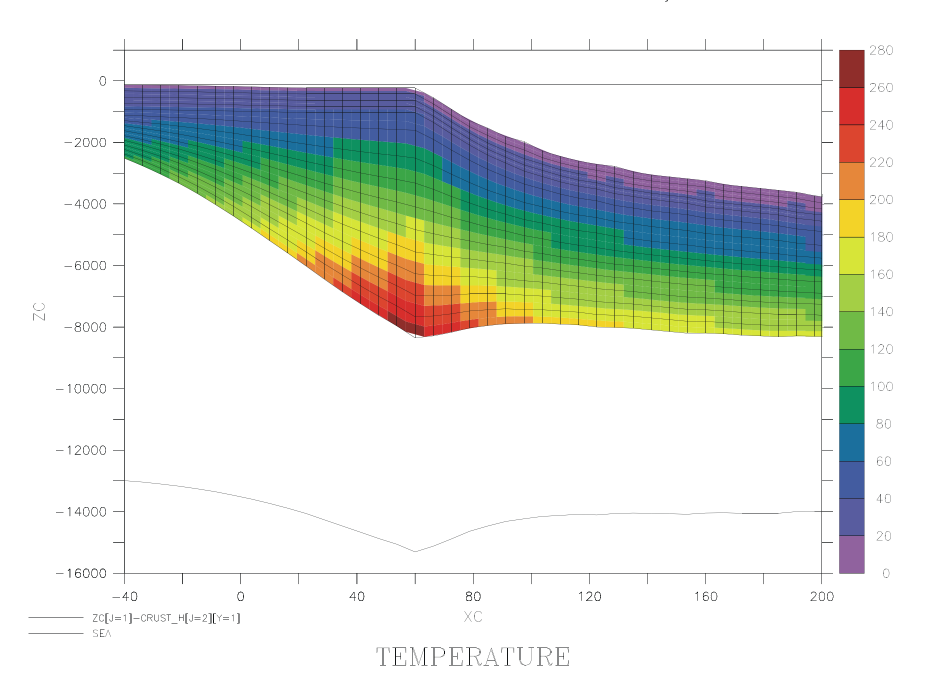
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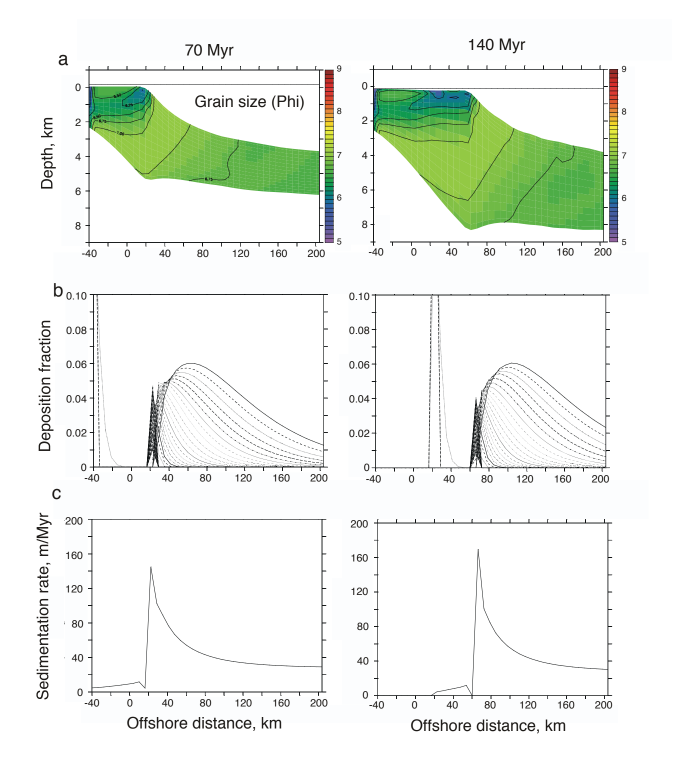
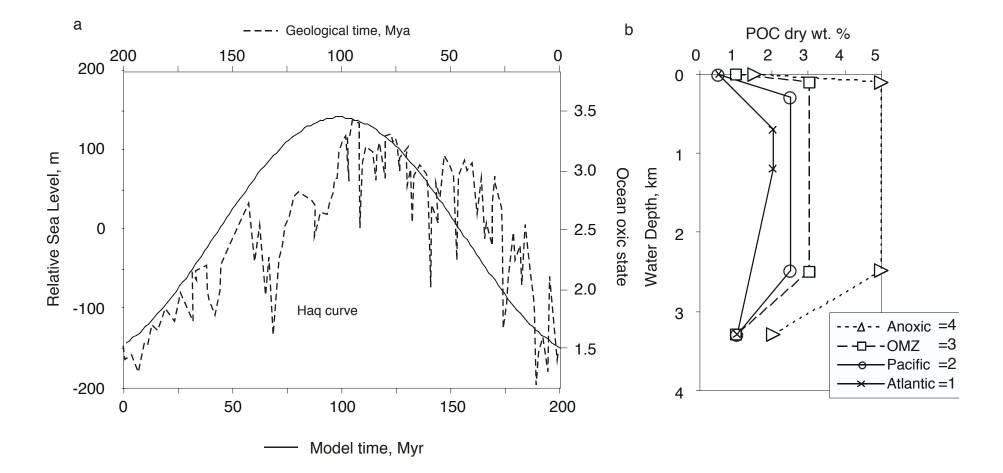


Figure 2





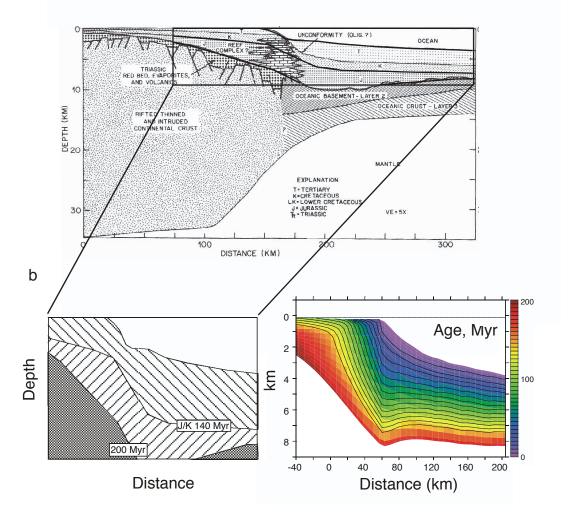
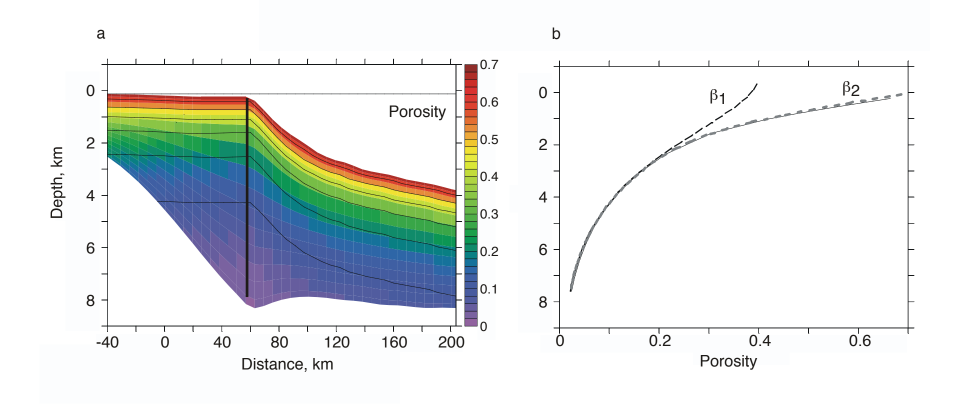
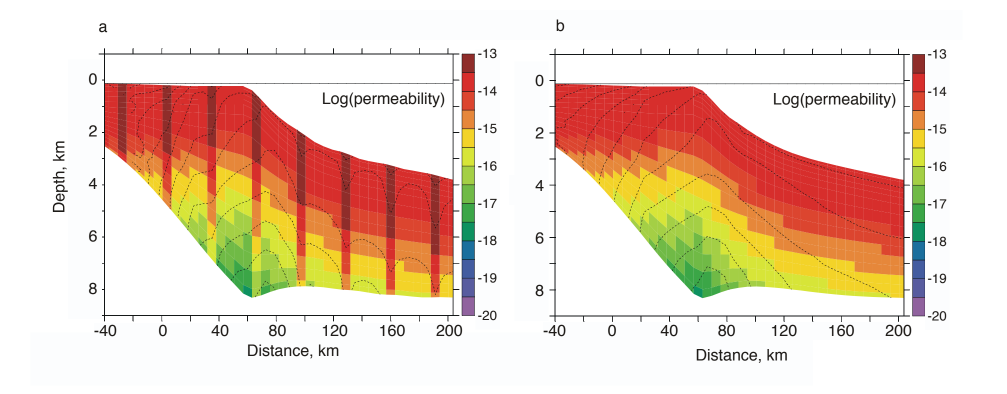
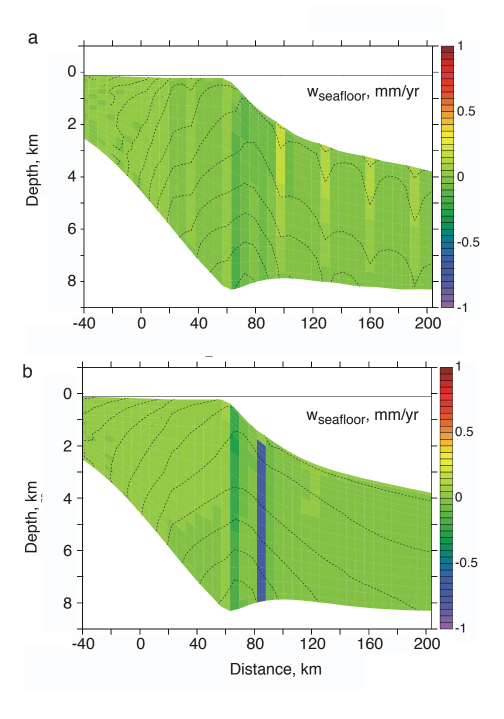
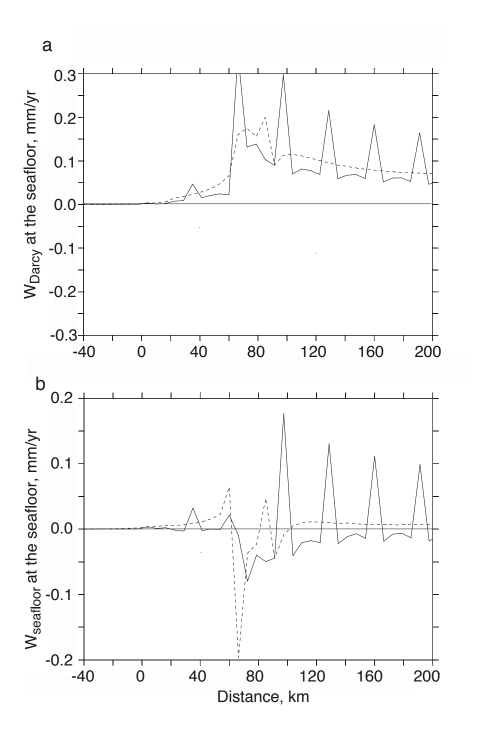


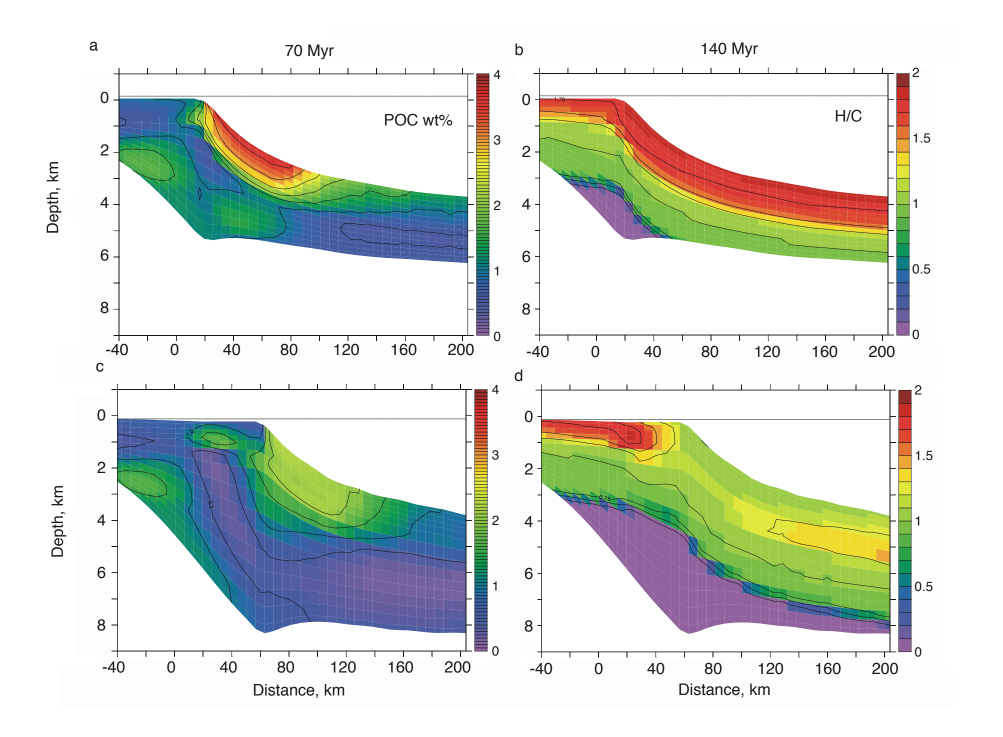
Figure 4

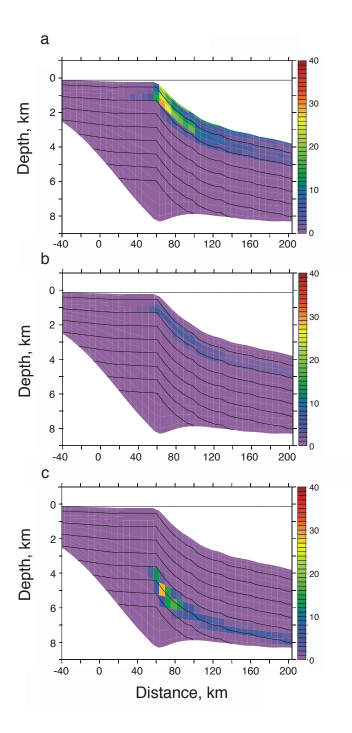












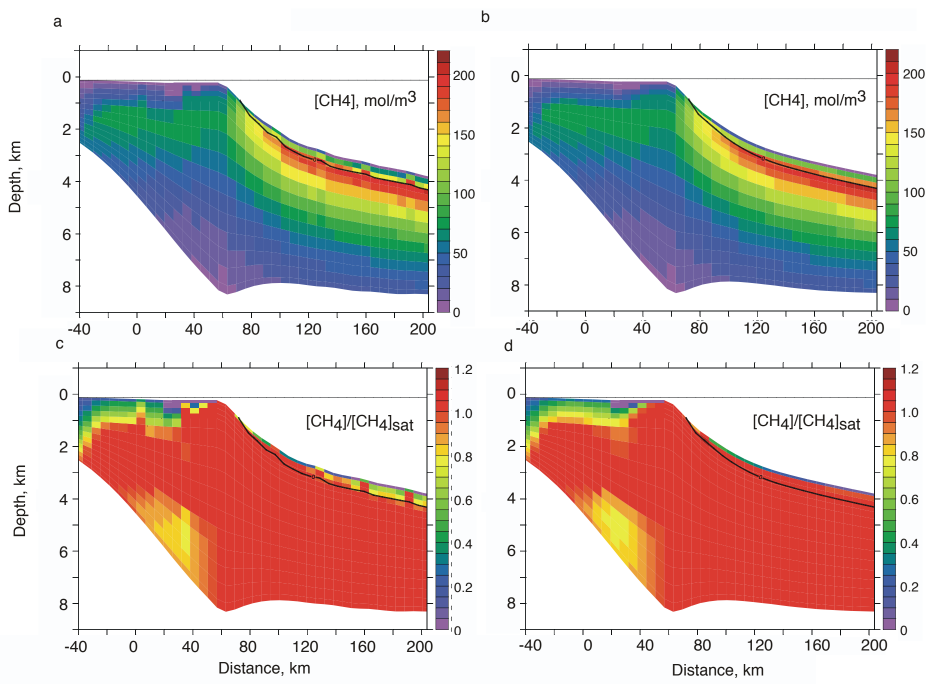
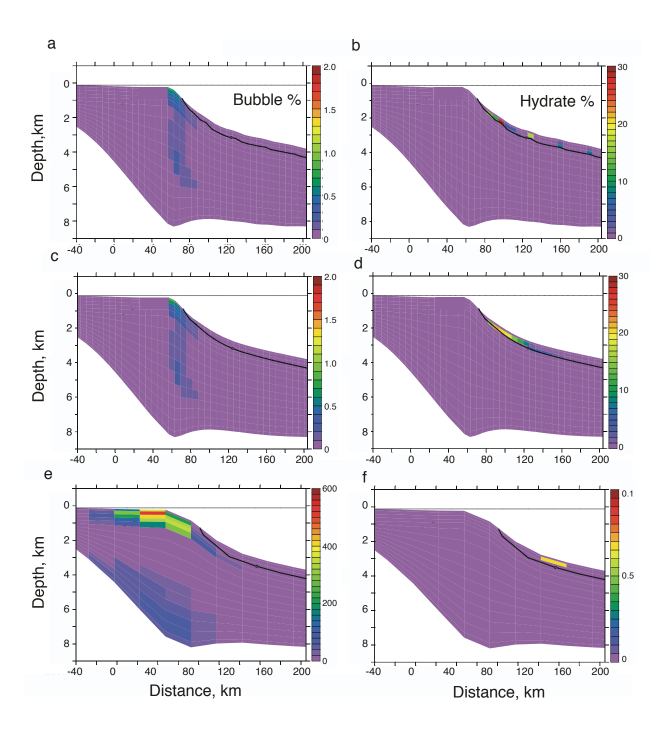
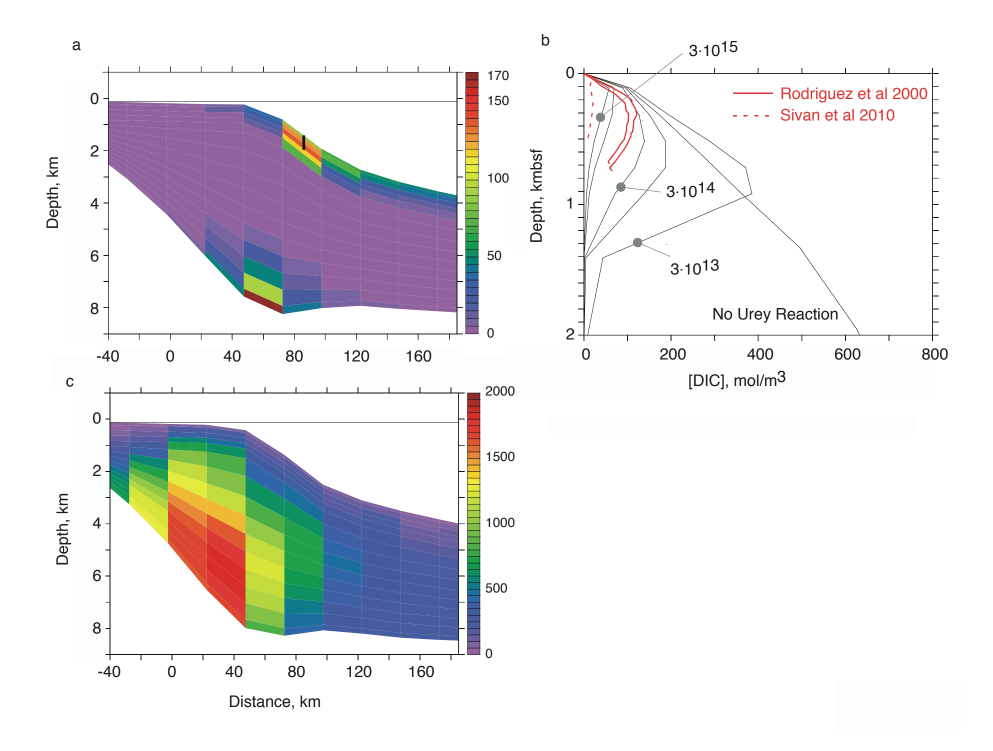
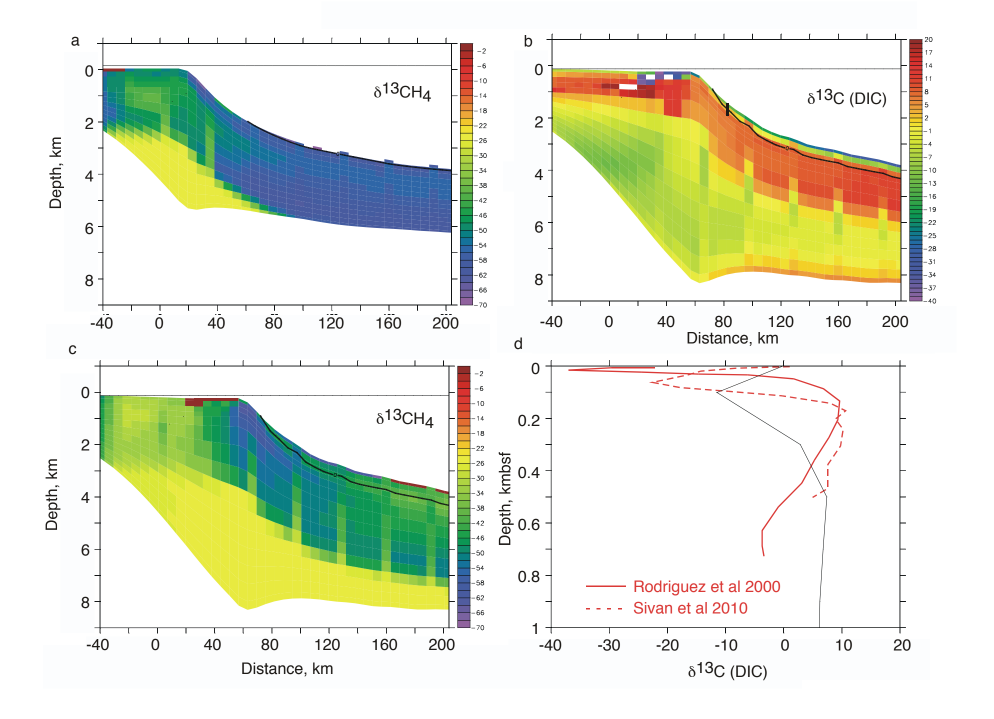


Figure 11. Concentration of dissolved methane plotted in moles / m3 (A and B), and relative to saturation (C and D). Base model run (A and C), and No Chimneys (B and D). An animation of the simulation can be viewed at http://geosci.uchicago.edu/~archer/spongebob_passive/ch4.movie.gif







129_I age-solid age, Myr

