Response to Anonymous Referee #1

Dear editor,

We would like to thank the first referee for his/her clearly positive comments that help us improve our manuscript. We replied to the comments of referee #1 in detail and explained how we have modified the manuscript for publication in Biogeosciences. Referee's comments are shown in black and our responses are shown in blue.

Anonymous Referee #1

«Overall comments» In general, I feel positive about the overall contribution of the paper. The topic is interesting and relevant to the goal of Biogeosciences. The approaches that the authors adopted are interdisciplinary and provide educative information to this topic. The data and interpretation are mostly convincing with several points that I request for further clarification (see comments below). The authors need to improve the presentation a lot as figure 2 is hard to read, some of the references cited are out of date, a few sentences are quite awkward to read, and Table 2 needs more polishing. I've included my detail suggestions for these technical issues in the pdf file. Despite these minor flaws, I strongly encourage to publish this paper after all of my concerns are addressed.

«Detail comments» My major concerns about the paper are as follow: for analytical and model approach-

1) In quite a few of samples, the sulfate concentrations are over seawater value (28mM). The authors explained this as dissolution of anhydrite. The alternative explanation will be re-oxidation of hydrogen sulfide in the porewater samples after they were collected. In the sampling procedure the authors described, I do not see any description such as flushing the porewater samples with N2 gas or fixing sulfide with Zn(OAc)2 solution to get rid of sulfide. Some clarification about how this is of concern should be addressed.

Response:

All of the samples were processed inside an anaerobic glove bag with N2 atmosphere. We added this information to the paper. However the reviewer is correct that it may be possible that some H2S is oxidized in the shallow subsurface of the cores due to oxygen penetration due to bioturbation.

Therefore, excess sulfate could potentially be from H2S oxidation and/or from sulfate input from groundwater. We note that in this specific site groundwater has been previously identified as a source of excess sulfate and excess Sr. Regardless we now include the option of H2S oxidation in the revised manuscript, whilst further noting that, since $R_{SD}$ is the net sulfate depletion, model results may underestimate the true $R_{POC}$ and $R_M$ due to sulfide oxidation. However, this is only relevant for very few cores in Group-1 and Group-2 where a shallow subsurface excess sulfate is observed. Although we mention this option for completion we believe that groundwater input is
a more likely source due to correlation between excess sulfate and excess Sr which has been previously described and is consistent with groundwater input.

Perry et al. (2002) identified dissolution of evaporites within the freshwater lens as the probable source of the excess $\text{SO}_4^{2-}$ found in some Yucatán groundwater by using the ratio between sulfate and chloride ($100\times(\text{SO}_4/\text{Cl})$). Ratios higher than seawater (average seawater is 10.3) are expected where gypsum/anhydrite dissolution is involved (Perry et al. 2002). The other indicator is $\text{Sr}/\text{Cl}$ ratio which in groundwater is invariably higher than the seawater value and indicates dissolution of celestite (from evaporite) and/or aragonite (Perry et al. 2002). The region east and south of Lake Chichancanab, Mexico, referred to as the Evaporite Region by Perry et al. (2002) is characterized by distinctive topography and the high-sulfate content of groundwater (Perry et al. 2002). The groundwater from the presumed source region, Lake Chichancanab, flows northward into the Celestún Estuary which can be recognized by the progressive decrease in the ratio $[\text{SO}_4/\text{Cl}]_{\text{groundwater}}/[\text{SO}_4/\text{Cl}]_{\text{seawater}}$ in water from southeast to northwest (Perry et al., 2009). These parameters in Celestún lagoon published in Young et al., (2008) are consistent with our interpretation that gypsum/anhydrite dissolution involved in the groundwater contributes to Celestún lagoon.

2) The authors modeled the system for 1 Myr to reach steady state. I wonder if this is a reasonable assumption to make in this case? From the high sedimentation rate (0.25- 0.35 cm/yr) of these cores, the age of the sediments investigated is not older than several years. Besides, this environment must be very dynamic with episodic input of water from different sources, bioturbation, and even sediment reworking. Why not simulate the system only to their real age, say 1-5 years? I believe this will significantly impact the results.

Response:

Yes, it's true that this is a very dynamic study area. Below in Fig R1 we show the example of modeling with 1 yr and 5 yr simulations for core 1CEL_Oct01. The results are the same as the 1 Myr simulations which means this dataset has reached steady state within 1 yr or less and using longer time scales does not make a difference.

Specifically we note that the long simulation time of 1 Myr is a default setting in this version of the model to ensure that the results for all cores are under steady state. However, we see the possible confusion. The 5 year simulation time now reported can be justified using the following equation (Boudreau, 1997):

$$t = \frac{L^2}{2 \times D_M}$$

where $t$ is time, $L$ is the distance involved in a typical diffusive movement (length of the model column) and $D_M$ is the molecular diffusion coefficient. The time for methane and sulfate diffusion over the length of modeled sediments (20 cm) is less than 0.6 yr ($D_M(\text{CH}_4) = 659$ (cm$^2$ yr$^{-1}$) and $D_M(\text{SO}_4^{2-}) = 382$ (cm$^2$ yr$^{-1}$) for e.g. core 1CEL_Oct01). Now we show the data using 5 years as the time needed for steady state and changed the text and figures accordingly. This makes no difference to the results.
Fig. R1: Depth profiles for modeled (lines) and measured/calculated (symbols) concentration of dissolved methane (dashed line; open circle), sulfate (solid line; solid circle) in the upper panel and sulfate depletion (solid line; solid circle), zero sulfate depletion (dashed line) and chloride (open circle) in the lower panel for core 1CEL_Oct01 for: (A) 1yr and (B) 5yr simulation times.

3) I find it difficult to understand the reactions described in the appendix:
   a. Page17931, line20-22: “Since AOM may play a minor role in the methane and sulfate rich sediment and RAOM was included in the net reaction rates of methane and sulfate this is justified.” I don’t understand at all what does this sentence mean. AOM should play an important role when you have abundant methane and sulfate isn’t? What is justified? By what?

   Response:

   We have now simplified the model set-up because, based on our data, we cannot accurately quantify the relative proportion of sulfate loss due to organoclastic sulfate reduction and AOM. The sentence pointed out by the reviewer was unclear and has been removed.

   b. Eq. A6: so you exclude entirely AOM when SO4-dep is positive? I thought SO4-dep>0 means active removal of sulfate? Not by AOM?

   Response:

   SO4-dep>0 means that active removal of sulfate is dominated by organoclastic sulfate reduction, although $R_{SD}$ includes $R_{SR}$ and $R_{AOM}$. Since we can see evidence for methane production along with sulfate reduction in many of our sites especially cores in Group-1 and Group-2, we use the rate derived from $[SO_4^{2-}_{dep}]$ profile ($R_{SD}$) to represent the rate of organoclastic sulfate reduction ($R_{SR}$).
In the sulfate reduction zone, we assume the co-occurrence of the following reactions:

(1) \[ \text{CH}_2\text{O} + 0.5\text{SO}_4 \rightarrow \text{H}_2\text{S} + \text{DIC} \quad (=R_{SR}) \]

(2) \[ \text{CH}_2\text{O} \rightarrow 0.5\text{DIC} + 0.5\text{CH}_4 \quad (=R_M) \]

(3) \[ 0.5\text{CH}_4 + 0.5\text{SO}_4 \rightarrow 0.5\text{H}_2\text{S} + 0.5\text{DIC} \quad (=R_{AOM}) \]

In the reaction stoichiometry, \( R_{SR} \) in reaction (1) is \( 0.5R_{POC} \). In terms of net sulfate reaction, \( R_{SD} = R_{SR} + R_{AOM} \). Since the 3 reactions are coupled in the sulfate reduction zone and reaction (2) + reaction (3) is equal to reaction (1), this means that \( R_{SD} = 0.5R_{POC} \). Hence, \( R_{AOM} \) is negligible compared with \( R_{SR} \) for \( R_{SD} \). We have revised the equations for \( R_{SD} = R_{SR} + R_{AOM} = R_{SR} = 0.5R_{POC} \). To estimate the fraction of organic matter degradation via methanogenesis \( (R_M) \) and organoclastic sulfate reduction \( (R_{SR}) \), Michaelis-Menten kinetic limitation term used for methanogenesis \( (R_M) \) and organoclastic sulfate reduction \( (R_{SR}) \) are expressed as:

\[
R_{SR} = 0.5 \cdot R_{POC} \cdot f_{SO_4^{2-}}
\]

\[
R_M = 0.5 \cdot R_{POC} \cdot (1 - f_{SO_4^{2-}})
\]

AOM may occur in these systems, but the data and model sensitivity results indicate that it is insufficient to prevent \( \text{CH}_4 \) escape to the bottom water, probably because of the abundant organic matter available for sulfate reducers to use instead of \( \text{CH}_4 \). We have revised this in the manuscript.

### d. Eq. A7
I understand you related \( R_{POC} \) to \( R_{SR} \) assuming all sulfate reduction is organoclastically. Again, is this a good assumption? What’s the role of AOM in sulfate reduction? I think you are right that organoclastic SR is important here but you need to explain this better.

**Response:**

AOM plays a minor role in sulfate reduction. Please see the response to comments (3).

### e. Eq. A12
How does \( R_{organic} \) differ from \( R_{POC} \)? How does the comparison of these two rates like? From table 2, I see them can be orders of magnitude different (e.g. \( 1\text{CH}_\text{Dec00} \)). Why?

**Response:**

The reason we simulate \( [\text{SO}_4^{2-}\text{dep}] \) profiles to derive rates of organic matter degradation, organoclastic sulfate reduction and methanogenesis is because measured organic matter contents in this area show evidence for a change in depositional regime over time (Gonneea et al., 2004,
and Fig. 4 in this version). Organic matter cannot therefore be used for accurate organic matter degradation calculations. To avoid the confusion between $R_{\text{organic}}$ and $R_{\text{POC}}$, we have removed the sampling and analytical methods, results, discussions and the model equation related to the measured organic contents from our manuscript and refer to measured organic matter contents from Gonnea et al (2004) and Eagle (2002, thesis).

This is stated more clearly in the revised manuscript.

4) Refer back to my comment (2), time scale of your model is really important. It determines the scale of your kinetic constants. For example, you use 0.01 1/yr for your $k_{\text{corg}}$. It may be a lot different if you only run the model for 5 years and. for scientific interpretation/discussion- I think the experiment and model results support most of the interpretation by the authors. I however feel that the authors should extend the discussion a bit more from the following prospects:
1) Maybe my biggest concern for the paper is the assumption of steady state. The authors should provide good reasons why they think this assumption is adequate as the system is so dynamic.

Response:

Please see the response to comment (2). We are now running the model for 5 years to reach steady state (see comment above). We use a steady state model because we do not have enough data to constrain a dynamic model such as regular monitoring of porewater sulfate, methane and chloride concentrations. The steady state model is still useful because we apply it to a wide range of profile types which represent the different conditions in the system and capture the system dynamics.

2) The authors presented tremendous amount of temporal/spatial porewater data in this paper but did not spend much effort in discussing these. The groping of data is based on the shape of profiles and thus their dominate reactions. Do these groups correspond to any particular location or season that might explain the such dominance in terms of biogeochemistry?

Response:

We have included additional discussion to show there is no relation between profile time and location or season and that the variability is a result of the system being heterogeneous and probably highly dynamic. We do not have sufficient data to differentiate between temporal and spatial trends, however as we show cores collected at the same time close to each other may differ and cores collected at the same sites during different times also differ from each other hence we believe that if we monitored any one site continuously it is likely that all profile types will be captured at one site.

3) Results from incubation experiments are one of the highlights in this paper but the authors only mentioned it briefly in 5.1 section. I wonder are the authors able to derive some rates from the experiments that can be compared with the rates estimated by modeling. Also, how do all these rates compared to other similar environments? I feel like the authors should put their results in a larger global context to reveal the significance of their data.
Response:

Yes, we are able to derive some rates from the experiments which have been added to Table 1 in the manuscript. An additional table (Table 3) includes rates estimated by modeling which can be used to compare with the rates from the experiments.

The maximum methane production rates listed in Table 1 from TMA, methanol and H₂ treatments are higher than the methane production rates from coastal freshwater and brackish wetland sediments which were measured using radiolabeled acetate and bicarbonate in slurries and reported in Segarra et al. (2013).

In addition to depth-integrated rates, Table 3 listed model derived maximum methanogenesis (Max-RM), sulfate reduction and AOM rates (Max-R_{SR}). Maximum methane production rates estimated from TMA, methanol and H₂ treatments of sediment slurry incubations (Table 1) are similar to values reported by model derived Max-RM at station 16CEL (Table 3) the site from which sediments were collected for sediment slurry incubations. Model derived Max-RM in some cores can reach to 1-2 orders of magnitude higher than rates derived from the sediment slurry incubations (e.g., cores 1CEL_Jul02, 1_1CH_Oct01, 2CEL_Oct01 and 14CEL_Dec00). Although our model results show that organoclastic sulfate reduction dominates organic matter degradation, model derived Max-RM are even higher than the maximum sulfate reduction rates in cores 1_1CH_Oct01 and 1_2CH_Oct01. Methanogenesis rates in this study area are more important than in other mangrove systems where methanogenesis is negligible (e.g., Thailand, Kristensen et al., 2000; Malaysia, Alongi et al., 2004; Australia, Kristensen and Alongi, 2006).

We include this in the discussion section of the manuscript.

4) The authors introduced the different seasons of this area and the potential impact to the sediment and porewater systems. However, I do not see further discussion about how their results reflect such seasonality. I feel a great pity that the authors did not translate the “numbers” they got from their modeling and experiments into something helpful to understand the spatial and temporal heterogeneity of the environment.

Response:

Model derived methane fluxes to the water column are listed in Table 2 (F_{methane (top)}) and reveal that fluxes (0.011-21 mmol CH₄ m⁻² d⁻¹) are similar or up to two orders of magnitude larger than fluxes reported for other mangrove systems in, e.g., Florida (0.02 mmol CH₄ m⁻² d⁻¹, Barber et al., 1988; Harriss et al., 1988), Australia (0.03-0.52 mmol CH₄ m⁻² d⁻¹, Kreuzwieser et al., 2003), and India (5.4-20.3 mmol CH₄ m⁻² d⁻¹, Purvaja and Ramesh, 2001). Our values as well as the depth-integrated rates (Fig. 2) show no relation between sampling time and location or season. Since all of the different types of methane depth profiles (group-1, group-2, etc) were found during each sampling trip, and no obvious trends in spatial and temporal distribution (seasons and sampling locations) were observed, model derived methane effluxes to the water column and the variability in the porewater methane concentrations and depth-integrated turnover rates suggest a very dynamic system with high methane production and efflux rates.

We have included this in the discussion section of the manuscript.
«Minor/technical comments»

1) My biggest comments on the technical part of the paper is its presentation. The lead author tend to use long sentences with many clauses. I would suggest split the long sentences into shorter ones which will be more understandable for readers who know nothing about modeling especially.

Response:

We have improved the English structure.

2) The authors also need to consider more recent literatures. When the hypothesis was built solely based on some 80’ and 90’ papers, it’s hard not to think there may be different views in the current research.

Response:

More recent literature has been included.

3) The Figure 2 is small and difficult to read. You need to figure out a different way to present these.

Response:

Fig. 2 was moved to supplementary material and replaced with a figure showing one typical profile per group. All figure qualities were improved.

4) I have a few comments for Table 2. You need to be more careful about the significant digits. I don’t think the model can give that many meaningful digits. The use of “F” at header row is confusing. I know you explain below but it is intuitionally awkward especially when you mixed the real fluxes with depth-integrated rates. The negative sulfate depletion rates and sulfate reduction rates are also awkward. It makes no physical sense unless you meant the reactions are reversible, which I think are not.

Response:

The table has been revised.
Response to Referee #2 (Dr. Pohlman)

Dear editor,

We would like to thank the second referee, Dr. Pohlman, for his support of our manuscript and for giving us comments to improve the manuscript. We replied to his comments in detail point by point and explained how we have modified the manuscript for publication in Biogeosciences. Dr. Pohlman's comments are shown in black and our responses are shown in blue.

General Comments:

The authors present sulfate, methane and chloride data from sediment cores collected from two coastal mangrove systems in the Yucatan Peninsula. The authors group the cores into 5 sets that generalize the sulfate and methane profile behavior. Because the analytical data are limited to concentration profiles of 3 constituents, they apply the Wallman et al. 2006 transport-reaction model to explain potential processes affecting the pore water geochemistry. An unusual and interesting observation is that methane and sulfate often coexist in the porewater, suggesting a non-competitive substrate (i.e., one used only by methanogens) allows methanogens to be active in the presence of sulfate reducers. A series of incubations that includes a treatment with the non-competitive substrates TMA and methanol demonstrates the microbial machinery and other factors required to produce methane from these substrates is present in the sediments from the investigated sites. The suggested implication is that mangrove ecosystems may be large methane emitters, provided the observations and model results accurately represent mangrove systems at large.

Although the diversity of data is limited, the authors do a commendable job of testing the hypothesis that non-competitive substrates accounted for the accumulation of methane in the sulfate reduction zone. The study does not provide definitive evidence that the process is active, as the only substrate-level data supporting its activity are from ex situ incubation experiments. The study should be used as motivation for tackling this specific question in greater detail in a mangrove ecosystem. It would appear others have observed the same effect in mangroves, but this appears to be the first to suggest a mechanism for the repeated observation. This is an important and interesting contribution. With moderate revisions, this reviewer recommends publication of this manuscript in Biogeosciences.

Specific Comments:

1. The grouping of the profiles helps to consolidate the data in a way that makes the application of the model more systematic. However, the authors have a tendency to overstate the certainty of their findings. For example, the model does not “illustrate” that methane is produced from DOM...it suggests production from these unmeasured carbon sources is possible. Also, shallow methane production does not necessarily promote high methane fluxes to the water column and atmosphere as the authors state. Although benign in intent, these statements being expressed definitely in the abstract may be misleading because they imply the conclusions are based on data. Be clear that the conclusion are based on modeling results and that no measurements regarding fluxes were obtained.
Response:

We have changed the wording used to be more consistent with our data and less definitive (e.g. change "illustrate" to "suggests" and change "promote" to "increase the likelihood").

We have also made changes throughout the manuscript in order to more clearly differentiate the modeling results from the field and laboratory measurements.

This reviewer recommends the authors provide a figure with generalized sulfate profiles (and methane, if applicable) for each group in Fig 2. Such a model (and a description of each group in the Fig 2 headings) would give the reader a better intuitive sense for the groupings.

Response:

Fig. 2 was moved to supplementary material and replaced with a figure showing one of the typical profiles per group.

2. Why would mangroves have such a high abundance of non-competitive substrates in comparison to other brackish systems?

Response:

This is a question we can only speculate about. Mangrove forests are known to be highly productive ecosystems with the capacity to release high concentrations of DOM to sediment porewaters (Kristensen et al., 2008). Litter from trees (leaves, propagules and twigs) and subsurface root growth provide further significant inputs of organic carbon to mangrove sediments which are unique for this type of system. We have now included these sentences in the manuscript.


3. Using the near surface methane gradients and modeled results, the authors should quantify the differing methane flux potentials for each environment rather than only speculating about the importance of this methane source.

Response:

Model derived methane fluxes to the water column are listed in Table 2 ($F_{methane\ (top)}$) and reveal fluxes (0.011-21 mmol CH$_4$ m$^{-2}$d$^{-1}$) that are similar or up to two orders of magnitude larger than fluxes reported for other mangrove systems in Florida (0.02 mmol CH$_4$ m$^{-2}$d$^{-1}$, Barber et al., 1988; Harriss et al., 1988), Australia (0.03-0.52 mmol CH$_4$ m$^{-2}$d$^{-1}$, Kreuzwieser et al., 2003), and India (5.4-20.3 mmol CH$_4$ m$^{-2}$d$^{-1}$, Purvaja and Ramesh, 2001). Since all of the different types of methane depth profiles (Group-1, Group-2, etc) were found during each sampling trip and no differences in spatial and temporal distribution (seasons and sampling locations) were
observed, model derived methane effluxes to the water column and the variability in the porewater methane concentrations suggest a very dynamic system with high methane production and efflux rates. We have included this in the discussion section of the manuscript.

4. The site description should include a description of where and why anhydrite might contribute excess sulfate. An alternate possibility not discussed is oxidation of sulfides. Total sulfides were not measured, so their potential contribution cannot be discussed. Perry and others have written much about why anhydrites and gypsum are found on the Yucatan platform. More details would make this argument more convincing. The evidence for contributions from anhydrite are not especially compelling. Basically, the authors state that there is anhydrite in the area, so that explains the excess sulfate. From looking at one of the Perry references, it is not clear that one would expect a groundwater contribution in the Chelem lagoon (inside the Chicxulub impact zone). More details would be helpful. Sr data would be even better, but that is not likely to be available and is not required.

Response:

Perry et al. (2002) identified dissolution of evaporites within the freshwater lens as the probable source of the excess $SO_4^{2-}$ found in some Yucatán groundwater by using the ratio between sulfate and chloride ($100 \times (SO_4/Cl)$). Ratios higher than seawater (average seawater is 10.3) are expected where gypsum/anhydrite dissolution is involved (Perry et al. 2002). The other indicator is Sr/Cl ratio which in groundwater is invariably higher than the seawater value and indicates dissolution of celestite (from evaporite) and/or aragonite (Perry et al. 2002). The region east and south of Lake Chichancanab, Mexico, referred to as the Evaporite Region by Perry et al. (2002) is characterized by distinctive topography and the high-sulfate content of groundwater (Perry et al. 2002). The groundwater from the presumed source region, Lake Chichancanab, flows northward into the Celestún Estuary which can be recognized by the progressive decrease in the ratio $[SO_4/Cl]_{groundwater}/[SO_4/Cl]_{seawater}$ in water from southeast to northwest (Perry et al., 2009). These parameters in Celestún lagoon published in Young et al., (2008) are consistent with our interpretation that gypsum/anhydrite dissolution involved in the groundwater contribute to Celestún lagoon.

Though there are no published $SO_4$ and Sr data for groundwater and surface water in Chelem lagoon, Perry et al., (2009) measured strontium concentrations greater than seawater in the saline groundwater of the Northern Yucatan Peninsula east of the Ring of Cenotes, and Chelem lagoon is located within this region. We included this in the discussion section of the manuscript.

5. Were the sediments dried and prepared for TOC analysis as part of this study, or Gonneea et al., 2004? The methods do not include the analysis. The results do not specify the origin of the data. Please clarify.

Response:

The original data are from Gonneea et al., (2004) and Eagle, (2002, master thesis). The TOC study utilized splits of the sediment cores collected for methane concentration analysis.
We have included both references in the manuscript.

6. Increasing OM content with depth? How is this? Suggestion of a changed depositional pattern not discussed.

Yes, organic matter profiles show a changed depositional pattern (Gonneea et al., 2004). Since this pattern can't be used for organic matter degradation calculations, we simulate [SO$_4^{2-}$$_{dep}$] profiles to derive organic matter degradation rates. To avoid the confusion between TOC analysis, TOC data expressions and reaction rates for $R_{organic}$ (Eq. A11) and $R_{POC}$ (Eq. A7) in this version, we have removed the sampling and analytical methods, results, discussions and the model equation related to the measured particulate organic contents from our manuscript and refer the measured organic matter contents to Gonneea et al (2004) and Eagle (2002, thesis).

7. Why would negative sulfate depletion be observed at the surface and not at depth if the source of the excess sulfate is from depth? See Core 7CH-Oct01.

Response:

It may be possible that some $H_2S$ is oxidized in the shallow subsurface of the cores due to oxygen penetration due to bioturbation.

Therefore, excess sulfate could potentially be from $H_2S$ oxidation and/or from sulfate input from groundwater. We note that in this specific site groundwater has been previously identified as a source of excess sulfate and excess Sr. We now include this option in the revised manuscript, whilst further noting that, since $R_{SD}$ is the net sulfate depletion, model results may underestimate the true $R_{POC}$ and $R_M$ due to sulfide oxidation. However, this is only relevant for very few cores in Group-1 and Group-2 where a shallow subsurface excess sulfate is observed. Although we mention this option for completion we believe that groundwater input is a more likely source due to correlation between excess sulfate and excess Sr which has been previously described and is consistent with groundwater input.

Technical Corrections:

17921, line 21: ‘porewater’
17923, line 7: delete ‘that’
17923, line 12: ‘sites’
17927, line 5: ‘inhibited’
17930, line 3: ‘atmosphere’
17920, line 20: ‘chloride’

Response:
These technical corrections have been revised in the manuscript.

Figures:
1. Put letters on the figure panels
   Response: The figure has been revised.

2. Some units on figures indecipherable (e.g., CH4 conc)
   Response: The figure has been revised.
A list of all relevant changes made in the manuscript:

Dear editor,

Based on both referees' comments, we have incorporated our responses in the revised manuscript. The relevant changes are as follows:

1. modifying model descriptions which include AOM and heterotrophic sulfate reduction ($R_{SR}$) into net sulfate depletion rates ($R_{SD}$), use $R_{SR}=R_{SD}$ and use 5 yrs for steady state simulations;

2. combining cores for Group-2 and Group-3 into one group, so there are four Groups of porewater profiles;

3. using the second modeling approach to simulate the original Group-2 data which has been combined with Group-3 in this version;

4. adding methanogenesis rates calculated from sediment slurry experiments in Table 1, adding maximum model-derived rates of methanogenesis and sulfate reduction in Table 3 and comparing both methanogenesis rates in the main text;

5. removing the sampling and analytical methods, results, discussions and the model equation related to the measured particulate organic contents from our manuscript and refer the measured organic matter contents to Gonneea et al (2004) and Eagle (2002), since organic matter profiles show a changed depositional pattern which can't be used for organic matter degradation calculations;

6. adding the sampling and analytical methods and results for porewater methane;

7. including a description of where and why anhydrite might contribute excess sulfate in the discussion section of the manuscript, though oxidation of sulfides is a possible source for excess sulfate;

8. including model derived methane fluxes to the water column listed in Table 2 ($F_{methane \, (top)}$) in the discussion section of the revised manuscript;

9. including additional discussion to show there is no relation between profile time and location or season and that the variability is a result of the system being heterogeneous and probably highly dynamic;

10. describing AOM plays a minor role in this study in the last paragraph;

11. showing one selected profile per group in Fig. 2 for illustration and presenting the other profiles for each group (9 cores for Group-1, 6 cores for Group-2, 2 cores for Group-3 and 3 cores for Group-4) in the Appendix (Fig. A1);

12. revising all Tables according to modified modeling approaches (there are no mark-up changes for Tables shown in this file);

13. improving English structures and presentations;

14. improving qualities of all figures.

15. Andrew Dale (GEOMAR) has been added as a co-author for his contribution and advice on the numerical modelling.
Methane and Sulfate Dynamics in Sediments from Mangrove-dominated Tropical Coastal Lagoons, Yucatán, Mexico

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Abstract

Porewater profiles in sediment cores from mangrove-dominated coastal lagoons (Celestún and Chelem) on the Yucatán Peninsula, Mexico, reveal the widespread coexistence of dissolved methane and sulfate. A numerical transport-reaction model suggests that methane in the upper sediments is produced in the sulfate reduction zone at rates ranging between 0.012 and 31 mmol m$^{-2}$ d$^{-1}$, concurrent with sulfate reduction rates between 1.1 and 24 mmol SO$_4^{2-}$ m$^{-2}$ d$^{-1}$.

The model also indicates that a significant fraction of methane is transported to the sulfate reduction zone from deeper zones within the sedimentary column, by rising bubbles and gas dissolution. Sediment slurry incubation experiments show that non-competitive substrates such as trimethylamine (TMA) and methanol can be utilized for microbial methanogenesis at the study sites. Our results suggest that a large fraction of the methane formed in the sediments escapes to the overlying water column. By combining field measurements with transport-reaction modeling, we are able to demonstrate that sediments in coastal lagoons within mangrove ecosystems are characterized by shallow methane production and accumulation depths, likely due to non-competitive substrate utilization in near-surface sediments and extensive bubble transport and dissolution; this may favor high methane emission rates.

I Introduction

Wetlands are the largest natural source of methane (CH$_4$) to the atmosphere, accounting for between 20-25% of the global atmospheric methane budget (Fung et al., 1991; Whalen 2005).

Methane produced in wetlands is primarily biogenic, arising from microbial activity in anaerobic sediments and soil. Since sulfate-reducing bacteria outcompete methanogens for common substrates (Oremland and Polcin, 1982), freshwater wetlands typically have much higher methane fluxes to the atmosphere than brackish to fully marine wetlands (Bartlett et al., 1987; Bartlett et al., 1985; Segarra et al., 2013). Marine and estuarine sediments are generally characterized by comparatively lower rates of methanogenesis with a methane production and accumulation zone located deeper within the sediment pile below the sulfate reduction zone (Holmer and Kristensen 1994; Martens and Val Klump 1984; Poulton et al., 2005; Segarra et al., 2013). In these marine or estuarine systems methane that diffuses upwards towards the sediment surface can be oxidized both anaerobically (AOM) and aerobically within the sediments and in the water column, reducing emissions to the atmosphere (Whalen, 2005).

Despite brackish to marine salinities, methane fluxes comparable to those measured in freshwater wetlands have been reported for coastal mangrove-dominated lagoon systems in...
sustained relatively high methane fluxes despite high sulfate concentrations. For example, if the microbial activity of sulfate reducers is high and sulfate replenishment from the overlying water is slow, sulfate may become depleted in the upper centimeters of the sediment, thus allowing methanogens to occur close to the sediment surface. Additionally, methanogens can co-exist with sulfate reducers when non-competitive substrates (those used only by methanogens and not by sulfate reducers) are available. Moreover, in these systems methane may migrate from deeper in the sediment to shallower depth and to the water column. Typically, a large percentage of the methane produced in sediments is oxidized prior to reaching the atmosphere, and in shallow-water systems, the oxidation takes place primarily in the sediments and not in the water column (Martens and Valklump 1980; Misch and Gosselink, 2000; Weston et al., 2011; Segarra et al., 2013, 2015). However, accumulation and transport of methane in gas bubbles reduces the exposure time of methane to oxidants such as oxygen and sulfate, allowing a large fraction of gas to escape the sediment (Barnes et al., 2006; Martens and Valklump, 1980).

The objective of this study was to examine porewater methane distribution within the sediments of two mangrove-dominated coastal lagoons in Mexico and relate them to sulfate concentrations of the sediments. We aim to gain a better understanding of the factors controlling the methane flux from coastal mangrove-dominated lagoon sediments. To this end, we applied a numerical transport-reaction model based on Wallmann et al. (2006) and Chuang et al. (2013) to simulate porewater methane and sulfate concentration profiles. We also performed sediment slurry incubation experiments to test the effect of competitive and non-competitive substrates on methanogenesis in the lagoon sediments. The results provide quantitative data on methane dynamics in coastal mangrove-dominated lagoon systems and highlight their importance as methane sources to the atmosphere.

**2 Study sites**

The mechanism of gas release from the sediment can also affect exposure time to oxidizing conditions in the sediment and may therefore have a significant effect on the total methane flux to the atmosphere. If enough methane builds up within the sediment, it can be released in the form of bubbles (ebullition), which can travel through the sediment and water column quickly, resulting in minimal oxidation.

In order to better understand the processes controlling methane flux from these sediments to the water column and atmosphere in these systems. By examining the spatial and temporal differences in porewater methane distributions at the mangrove lagoons, and relating them to sulfate concentrations and organic carbon content of the sediments, we can gain a better understanding of the factors controlling atmospheric methane flux from coastal mangrove ecosystems.
Fieldwork was conducted in two mangrove-dominated coastal lagoons located on the western Yucatán Peninsula, Mexico (Figure 1). The typical climatological pattern for this area consists of a dry season (March–May), a rainy season (June–October) during which the majority of the annual rainfall (>500mm) occurs, and the “nortes” season (November–February), which is characterized by moderate rainfall (20-60mm) and intermittent high wind speeds greater than 80 km hr⁻¹ (Herrera-Silveira, 1994).

Celestún Lagoon (20°52′N, 90°22′W) is long, narrow, and relatively shallow (average depth = 1.2 m). The inner and middle sections of the lagoon always have lower salinities than the section near the mouth due to year-round discharge of brackish groundwater from multiple submarine springs (Young et al., 2008). Salinity within the lagoon fluctuates seasonally, with salinity in the inner zone ranging from 8.9 to 18.2 during the course of this study, grading out to marine salinities at the mouth of the lagoon (Young et al., 2008). The lagoon is surrounded by 22.3 km² of well-developed mangrove forest, and has experienced relatively little disturbance from human development and/or pollution such as wastewater discharge (Herrera-Silveira et al., 1998). Sediments in Celestún consist primarily of autochthonous carbonate ooze.

Chelem Lagoon (21°15′N, 89°45′W), (average depth = 0.7 m), in contrast, receives very little groundwater input and the surrounding area has been heavily impacted by urban development. Salinity in Chelem ranges from brackish to hypersaline (24.8 - 40.3 during the study period), and vegetation surrounding the lagoon consists of scrub mangrove forest (Young et al., 2008). The construction of Yucalpeten Harbor in 1969 (Valdes and Real 1998) increased circulation and resulted in sandy marine sediments entering the lagoon. Sediments in Chelem deposited since 1969 consist of a heavily bioturbated mix of sands and autochthonous carbonate ooze, with a large number of shells of living and dead burrowing organisms (Valdes and Real 1998). In the following text, CEL and CH denote cores collected from Celestún Lagoon and Chelem Lagoon, respectively.

3 Sampling and analytical methods

3.1 Porewater solutes

Sediment cores were collected along lengthwise transects in both lagoons during the three different seasons; April 2000 (dry season), December 2000 (nortes season), and October 2001 (late rainy season). Duplicate samples (1_1CH_Oct01 and 1_2CH_Oct01) were collected at station 1CH in Chelem lagoon. Sediments were sampled using hand-held acrylic push cores (7 cm inner diameter) either 30 or 60 cm in length. The push cores had holes drilled along the side.
at 2 cm intervals, which were sealed with electrical tape prior to sampling. Subsamples for porewater methane analysis were collected in the field immediately after core collection from the holes along the sides of the push cores, using plastic 3 mL syringes with the needle attachment end removed. The sediment plugs from the syringes were immediately extruded into 20 mL glass Wheaton bottles and sealed with blue butyl stoppers and aluminum crimp caps. 3 mL of degassed Milli-Q water and 0.3 mL of saturated mercuric chloride (HgCl₂) solution were added to create a slurry and halt all biological activity within the sample. After subsampling, the cores were capped, the holes were resealed, and the cores were transported back to the lab for sectioning and porewater extraction. The cores were extruded and sliced into 2.5 cm depth intervals in an anaerobic glove bag under an N₂ atmosphere and transferred into centrifuge tubes for porewater extraction. Core length was measured immediately after collection and just prior to extrusion in order to correct for compaction during transport. Average compaction was 6% of the total core length, and never exceeded 20%.

Porewater for sulfate (SO₄²⁻) and chloride (Cl⁻) analysis was extracted by centrifuging all the sediment from each depth interval and filtering the porewater through sterile 0.20 µm syringe filters. Samples were kept frozen in 20 mL acid-cleaned glass scintillation vials until analysis. Porewater sulfate and chloride concentrations were measured by ion chromatography using a Dionex DX-500 IC equipped with an Ionpac AS9-HC column (4mm) and AG9-HC (4mm) guard column. The samples were diluted 5-fold with Milli-Q water prior to analysis in order to bring the sulfate and chloride within the appropriate analytical range for the ion chromatograph. Methane concentrations for all samples were measured on an SRI 310 Gas Chromatograph (GC) equipped with a flame ionization detector and an Alltech Haysep S 100/120 column (6’ x 1/8” x 0.085”). Helium was used as the carrier gas at a flow rate of 15 ml/min and the column and detector temperatures were maintained at 50 °C and 150 °C, respectively. Peak integration was performed using Peak Simple NT software. Methane gas standards were prepared by diluting 100% methane in helium, and five standards bracketing the range of sample concentrations were measured at the beginning, middle, and end of each set of analyses. Average standard error of repeat injections of standards throughout a sample run (between 2 to 6 hours of continuous analysis) was 1.8% (n=152). Porewater methane concentration in the sediment core subsamples was determined after vigorously shaking of the sealed serum bottles containing the sediment slurries to ensure complete mixing, followed by at least 3 minutes of standing equilibration time to ensure that the porewater methane was fully equilibrated with the...
headspace in the serum bottles. A small volume of headspace (0.25-0.5 µL) was drawn out of each serum bottle using a gas-tight syringe, and analyzed for methane concentration on the SRI 310 GC. The total volume of porewater in each sample was calculated using the difference between the total wet weight of the sediment minus the dry weight of the sediment, correcting for the added water and HgCl₂ solution.

3.2 Sediment slurry incubation experiments

Sediment slurry incubations were performed in order to examine changes in methane production over different time intervals and at different substrate concentrations (Table 1). Incubations consisted of three competitive substrates (H₂, acetate, formate), two non-competitive substrates (methanol, trimethylamine (TMA)), and four types of controls. The controls (preparation methods are described below) consisted of an un-amended sediment control under anaerobic conditions, an un-amended aerobic control (partial oxygen headspace), a killed control in which the sediment was autoclaved to kill all living organisms in the sediment, and a chemical control in which biological methanogenesis was inhibited through the addition of 2-bromoethanesulfonic acid (BES) to a final concentration of 40 mM within the slurry. Triplicate bottles were prepared for each condition (controls and substrate additions), and methane headspace concentrations were measured at 3-4 time intervals over the course of 29 days.

All the sediment slurries were prepared semi-anaerobically by homogenizing sediment in a blender with artificial seawater mixture in a 1:1 ratio under continuous flow of nitrogen gas. Large pieces of leaves, twigs, and shells were removed from the sediment prior to homogenization. 70 mL glass Wheaton bottles were flushed with nitrogen gas for 1 minute prior to the addition of the sediment slurry. 30 mL of slurry was then added to each bottle under continuous nitrogen flow, and the bottles were sealed using blue butyl rubber stoppers and aluminum crimp seals. Substrate additions were made by injecting the substrate solution into the bottle immediately after sealing the bottles, except for the H₂ gas treatment and the aerobic control. For the addition of H₂, the entire headspace of the bottles was flushed with 100% H₂ gas. After each headspace sampling the H₂ gas removed by microbial activity in the sediment was replaced by inserting a gas tight syringe filled with 100% H₂ gas into the bottles, and allowing the gas to be drawn into the bottles until equilibrium pressure was reached. The aerobic controls were prepared like the anaerobic, un-amended controls, except that 8 mL (20% of the total headspace) of 100% O₂ was added to the bottles immediately after they were sealed. In order to ensure that the sediment slurries remained aerobic, 100% O₂ was added to the bottles throughout the incubation period. The sediment slurries were kept at room temperate
Headspace samples (0.25 mL) were extracted from the bottles at each time interval using a gas-tight syringe. Methane concentrations were measured on an HP 5730A GC equipped with a flame ionization detector. GC calibration and creation of standard curves were based on successive dilutions of 100% methane. Analytical error was approximately 5% for methane concentrations below 10 ppm-v (446 nM), and less than 3% for methane concentrations above 10 ppm-v as determined by repeat analyses of standards and samples.

4 Results

4.1 Porewater concentrations of dissolved species

Representative porewater methane profiles were plotted alongside sulfate profiles in Fig. 2 and Fig. A1. Profiles were assigned to one of four profile-types based on the relation between methane and sulfate distributions down core (see below). Considerable spatial and temporal variability in porewater chemistry was observed with no systematic seasonal differences in concentration trends. For example, porewater methane concentrations varied by up to three orders of magnitude in both lagoons, even between sites in close proximity to each other (i.e. 1CEL and 2CEL, Oct01; 1CH and 2CH, Dec00), and at the same station sampled during different seasons (i.e. 2CEL Dec00, Oct01; 1CH Apr00 and Oct01). No consistent differences were evident between the stations at the sides of the lagoons and those located in the center of the lagoons, or between stations located in the inner zone of the lagoons and those located near the mouth. For instance, methane above calculated saturated concentrations (1.1 and 1.3 mM) was observed in cores 1CEL Jul02 (the inner zone of Celestún lagoon) and 14CEL Dec00 (near the mouth of the lagoon). This is particularly interesting because the mouth of the lagoon has much higher salinities than the inner zone (Young et al., 2008). The variability (both spatial and temporal) in the porewater methane concentrations and in the spatial and temporal distribution of profile types suggest a very dynamic system where both concentration and distribution patterns in the porewater vary constantly (spatially throughout the lagoons and temporally at distinct sites). Such variability is indicative of rapid methane production and efflux rates.

Porewater sulfate concentrations ranged from 0.21 to 35.3 mM in Celestún lagoon and from 4.13 mM to 33.5 mM in Chelem lagoon and show different trends (Fig. 2; Fig. A1). In many of the cores a negative relation between methane and sulfate was observed. Specifically, higher sulfate was associated with lower methane in cores located near the mouth of the lagoons (16CEL Jul02, 16CEL Oct01, 14CEL Oct01, 14CEL Jul02 and 5CH Apr00) and lower
The relationship between porewater salinity (represented by chloride concentration), methane, and sulfate concentrations was spatially and temporally variable (Fig. 3). Generally, higher sulfate concentrations were associated with higher chloride in cores located near the mouth of the lagoons and lower sulfate with lower chloride in the inner zone of the lagoons (Fig. 3A). Despite these general trends there are no clear consistent relationships between methane and chloride (Fig. 3B) and sulfate and methane (Fig. 3C) when the data was considered collectively. The lack of consistent trends suggests multiple processes impacting the distribution of methane and sulfate. These include physical processes, such as mixing and dilution with seawater or groundwater, and biological processes such as sulfate reduction, methanogenesis and methane oxidation. Brackish groundwater enters the Celestún lagoon through at least 30 subsurface discharge points (Young et al., 2008), and the chloride profiles suggest that some of this groundwater may seep through the sediments, resulting in localized decline in porewater salinities.

To account for mixing with seawater or freshwater and to extract information on the processes controlling the distribution of porewater solutes, the observed sulfate depletion ([SO$_4^{2-}_{\text{dep}}$]$_{\text{OBS}}$) relative to seawater was calculated as the difference between the expected sulfate concentration contributed from sulfate (based on porewater chloride concentration) and the measured sulfate concentration:

$$[\text{SO}_4^{2-}_{\text{dep}}]_{\text{OBS}} = \frac{[\text{SO}_4^{2-}]_{\text{SW}}}{[\text{Cl}]_{\text{SW}}} \times [\text{Cl}]_{\text{measured}} - [\text{SO}_4^{2-}]_{\text{measured}} \quad (1)$$

where 0.05171 is applied for the $\frac{[\text{SO}_4^{2-}]_{\text{SW}}}{[\text{Cl}]_{\text{SW}}}$ ratio (Pilson, 1998). Positive values indicate that sulfate has been removed from the porewater, most likely through sulfate reduction while negative values indicate an external source of sulfate not associated with chloride that is other than seawater, in this case the groundwater (see discussion below).

Based on the observed trends in sulfate depletion, when considered together with methane, four different porewater trends can be described, referred to as Groups 1 through 4 here (Fig. 2, Fig. A1). The majority of profiles fell into Group-1 (10 cores); these profiles showed positive sulfate depletion profiles (e.g. sulfate consumption or loss) with methane profiles mirroring the sulfate concentration profiles (methane production or input). The peaks for methane and sulfate depletion occurred at the same depth as the lowest measured sulfate concentrations. In Group-2
4.2 Sediment slurry incubation experiments

All the sediment slurries with added substrates showed an increase in headspace methane concentrations that was significantly greater than those observed with either the un-amended aerobic and anaerobic controls or the treated controls (Figure 4). The greatest increases in headspace methane concentration were seen with additions of the two noncompetitive substrates, TMA and methanol. The H₂ treatment showed the next highest methane production rate, followed by formate then acetate. Of the four control conditions, the un-amended anaerobic treatment had the highest increase in headspace methane concentration than the un-amended anaerobic treatment, although there was no detectable change in the headspace methane concentration in the aerobic treatment between 150 and 700 hours. Both the autoclaved and BES treatments did not show any changes in headspace methane concentration greater than the instrumental detection limits. The maximum methane production rates for each treatment are listed in Table 1.

5 Discussions

5.1 Co-existence of Methane and Sulfate in Sediments

Seawater transport into the sediment by diffusion and bioirrigation due to the activity of burrowing animals has clear effects on porewater solutes. These processes are a source of seawater sulfate and mask sulfate loss by microbial reduction. Although, as indicated above, considerable variability in porewater profile distribution trends was observed, and different profile types were found throughout the lagoons, certain trends were more common at distinct locations. Specifically, sites characterized by sulfate addition from input of seawater into the sediment slurry incubation experiments show strong relations with sulfate depletion profiles in

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sediment (cores in Group-4) were found primarily near the mouth of both lagoons where low
methane was associated with near-zero sulfate depletion. Negative sulfate depletion (Group-3),
on the other hand, which indicates the presence of porewater that is enriched in sulfate relative
to chlorine, was seen primarily in the middle zone of Celestún Lagoon where groundwater
springs rich in sulfate due to anhydrite dissolution are present, as reported by Perry et al. (2009;
2002). Positive sulfate depletion profiles co-occurring with methane (Groups 1 and 2) were
seen throughout the lagoons but mostly at sites in the inner zone of both lagoons, suggesting
significant sulfate reduction at rates higher than the replenishment from sulfate rich
groundwater or from the overlying seawater and a source of methane to the shallow sections of
the sediment.

It is surprising that at many sites particularly within Groups 1 and 2 in the inner zone of both
lagoons (1CEL, 2CEL, 3CEL and 1CH) high concentrations of methane and sulfate
cocurred at the same depth in the sediment. Co-existence of methanogenesis and sulfate
reduction is not normally observed because sulfate reduction is more energetically favorable
than methanogenesis, and sulfate reducers should outcompete methanogens for common
substrates such as hydrogen and acetate (Oremland and Polcin, 1982; Jørgensen and Kasten,
2006). Moreover, anaerobic oxidation of methane (AOM) coupled with sulfate reduction at the
base of the sulfate reducing zone should further deplete methane (Capone and Kiene, 1988;
Valentine and Reeburgh, 2000). There are several possible explanations for these observations.
First, the high methane concentrations measured in the sulfate rich porewater may be supplied
by a rapid non-diffusive mechanism from below the sulfate reduction zone (like rising gas
bubbles), limiting the exposure time to AOM. Second, methane may be produced in-situ at
these depths supported by a high abundance of competitive substrates in the sulfate reduction
zone hence sustaining both methanogenesis and sulfate reduction (Holmer and Kristensen,
1994). Third, methanogens may instead be able to thrive on various non-competitive substrates
(Oremland and Polcin, 1982; Wellsbury and Parkes, 2000; Lee et al., 2008; Taketani et al.,
2010). Indeed, use of non-competitive substrates by methanogens, including methanol,
trimethylamines and dimethylsulfide, has been reported for mangrove sediments, coastal
lagoons and continental shelf sediments (Ferdelman et al., 1997; Lyimo et al., 2000; Mohanraju
et al., 1997; Purvaja and Ramesh, 2001; Torres-Alvarado et al., 2013; Maltby et al., 2016). Our
slurry incubation experiments demonstrated that the methanogenic community at Celestún is
capable of using a wide range of substrates, including H₂, acetate, formate, methanol, and
trimethylamine (Fig. 4). Both methanol and trimethylamine are not utilized by sulfate reducers,
which could allow methanogens to thrive in the sulfate reduction zone (Fig. 4). The use of
non-competitive substrates by the methanogenic community has important implications for

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methane fluxes to the atmosphere as it allows for methane production at shallow depths in the sediment and reduces the potential for complete oxidation of methane. Although processes and trends similar to those described above have been reported for other mangrove sediments (e.g., Lee et al., 2008; Purvaja and Ramesh, 2001), the co-occurrence of sulfate and methane and related biogeochemical reactions in these reports remain qualitative in nature. In the following section, we use a transport-reaction model to better quantify the processes controlling methane fluxes from the sediments in these mangrove-dominated tropical coastal lagoons.

5.2 Model set-up and application to mangrove-dominated coastal lagoon sediments

In order to understand methane production and consumption and how these processes relate to sulfate dynamics in the lagoon sediments, we used two different approaches to simulate methane and sulfate porewater profiles.

In the first approach, a transport-reaction model was applied to profiles of Group-1 where methane and sulfate co-occur with no indication of groundwater sulfate input and where sulfate reduction that surpasses sulfate addition from seawater (Fig. 2; Fig. A1). Data in Group-1 have positive net sulfate depletion rates indicative of sulfate reduction. The sulfate depletion is seen within the zone where methane concentrations are high. In these cores the net sulfate depletion rates can be used to derive the minimum methanogenesis rates (see model details in the Appendix). Reactions considered in this first approach include organic matter degradation via heterotrophic sulfate reduction, methane production via methanogenesis and methane addition from gas bubble dissolution (Haeckel et al., 2004; Chuang et al., 2013).

A second approach (detailed in the Appendix) was used for simulating the profiles for Group-2, Group-3 and Group-4 which show no positive net sulfate depletion rates when integrated over the core length. These sites are affected by groundwater input or by considerable irrigation and input of seawater. Here, the link between sulfate and methane reactions is less clear and hard to quantify directly.

The following equation was solved to quantify the rates of reaction and transport of dissolved methane and sulfate in the upper 20 cm of the sediments in both approaches (Berner, 1980; Boudreau, 1997):

\[ \Phi \frac{\partial C}{\partial t} = \frac{\partial (\Phi \cdot D_s \frac{\partial C}{\partial x})}{\partial x} - \frac{\partial (\Phi \cdot C)}{\partial x} + \Phi \cdot R_s \]

(2)

where \( x \) is sediment depth, \( t \) is time, \( \Phi \) is porosity, \( D_s \) is the solute-specific diffusion...
Model derived **sulfate depletion and methane concentrations** are shown in Fig. 2 and Fig. A1. Modeled porewater data for Group-1 (the most common trend) show that methane generated from organic matter degradation within the upper sediments is a more important methane source than methane diffusing from below and gas bubble dissolution, as further seen in the results of 1CEL_Jul02 and the sensitivity analysis from 2CEL_Jul02 (Fig. 5A). In 1CEL_Jul02, for example, gas dissolution of methane transported from deeper sediments is not necessary to achieve a good model fit to the data, and in-situ methanogenesis alone can reproduce methane concentrations similar to the measured data even though methane concentrations are oversaturated (> 1.1 mM (in situ solubility)) (Fig. 2). In contrast, the modeled methane profile for 2CEL_Jul02 (black dashed line) arguably does require the inclusion of methane from gas dissolution (Р_diss) (Fig. 5A). In Fig. 5A, the gray dashed and solid lines represent only gas dissolution in the methane reaction terms (no methanogenesis within the modeled 20 cm column) using different gas dissolution constants (k_diss values are 0.2 yr⁻¹ and 0.5 yr⁻¹ respectively). The model results shown as the gray dashed line simulate the methane concentrations below 10 cm depth, whereas those shown by the gray solid line reproduce methane concentrations in the upper 5 cm, but neither reproduces the data throughout the whole core. Comparing results considering methanogenesis and gas dissolution (black solid line) and methanogenesis only (black dashed line), it is clear that both methanogenesis and some gas dissolution are needed for reproducing the methane distribution in core 2CEL_Jul02. This illustrates the complexity of controlling processes and the dynamic nature and resulting temporal variability in methane fluxes at this and the other sites in the lagoons.

### 5.3 Model derived depth-integrated turnover rates and fluxes

Table 2 lists the calculated depth-integrated turnover rates and fluxes for the individual cores.
For profiles in Group-1, methane sources include methanogenesis within the upper 20 cm and/or methane transported from deeper sections (>20 cm) via bubble transport and dissolution. Methane can be supported fully by methanogenesis without gas bubble dissolution within the modeled upper 20 cm in cores 1CEL_Dec00, 1CEL_Jul02, 1_1CH_Oct01 and 1_2CH_Oct01. Gas bubble dissolution and transport from deeper sediments contributes more methane than methanogenesis in cores 1CEL_Apr00, 1CEL_Oct01, 2CEL_Dec00 and 3CEL_Jul02.

Methane sinks include emissions to the water column or methane diffusion into deeper sediments (>20 cm) and oxidation. Our model shows that the major sink for methane, however, is emission to the water column accounting for over 90% of methane produced within the upper 20 cm (e.g. 1CEL_Apr00, 1CEL_Oct01, 3CEL_Apr00 and 3CEL_Jul02). Model derived methane fluxes to the water column are listed in Table 2 ($F_{\text{methane (top)}}$) and range from 0.012-20 mmol CH₄ m⁻² d⁻¹. These are similar to or up to two orders of magnitude larger than fluxes reported for other mangrove lagoon systems in Florida (0.02 mmol CH₄ m⁻² d⁻¹), Barber et al., 1988; Harriss et al., 1988), Australia (0.03-0.52 mmol CH₄ m⁻² d⁻¹), Kreuzwieser et al., 2003), and India (5.4-20.3 mmol CH₄ m⁻² d⁻¹), Purvaja and Ramesh, 2001). Since all methane depth profile types were observed throughout the year with no differences in spatial and temporal distribution (seasons and sampling locations), our results support the idea that methane fluxes in coastal mangrove lagoon systems respond very dynamically to environmental stimuli.

Sulfate sinks include heterotrophic sulfate reduction and AOM, although the model suggests that AOM plays a minor role compared to heterotrophic sulfate reduction. Sulfate reduction ranges from 1.1 to 24 mmol SO₄²⁻ m⁻² d⁻¹ and is the major sink for both sulfate and organic carbon in most cores. Sulfate reduction accounts for 2.2 to 48 mmol C m⁻² d⁻¹ of total anaerobic carbon respiration, which is in the same range of values listed in Kristensen et al. (2008) for most mangrove sediments.

Mangrove forests are known to be highly productive systems with the capacity to release high concentrations of dissolved organic matter (DOM) to surrounding sediments and porewaters (Kristensen et al., 2008). Tree litter and subsurface root growth provide further significant inputs of organic carbon to mangrove sediments which are unique for this type of system. The rate of organic matter mineralization ($R_{\text{org}}$; Eq. A6) derived from sulfate depletion ranges from 3.2 mmol C m⁻² d⁻¹ to 110 mmol C m⁻² d⁻¹. Although our modeling approach for determining degradation rates is not without uncertainty, it is more accurate than rates derived from down-core trends in organic matter content because of temporal variability in accumulation rates in this area (Gonneea et al., 2004). Particulate organic matter will also contain a high amount of refractory carbon that is not easy to quantify and separate from the bulk pool. The derived degradation rates likely represent the more labile particulate components and labile
DOM that was not considered (or measured) in this study. The high calculated organic carbon oxidation rates derived here are thus not unexpected since mangrove systems in general (e.g., Dittmar et al., 2006; Dittmar and Lara, 2001; Lee 1995; Odum and Heald, 1975) and the lagoons in Yucatan in particular are dominated by high concentrations of DOM, a large fraction of which is likely to be labile (Young et al., 2005).

Depth–integrated methane production or consumption rates ($R_{CH_4}$) and net sulfate inputs ($R_{SO_4}$) calculated from Eqs. (A9) and (A10) for cores in Group-2, Group-3 and Group-4 are listed in Table 2. The methane and sulfate net production/consumption rates ranged from -0.060 to +11 mmol CH4 m⁻² d⁻¹ and -69 to +21 mmol SO₄²⁻ m⁻² d⁻¹. (Negative values indicate net sulfate or methane consumptions while positive values indicate production or addition from external sources). Although sulfate depletion values for cores in Group-2 are positive (e.g. net sulfate reduction), sulfate concentrations at some depths of the porewater are relatively high, suggesting continuous sulfate input from deeper within the sediments or from seawater. Cores in Group-3 and Group-4 show negative or zero sulfate depletion that likely results from sulfate addition from groundwater (Group-3) or seawater (Group-4), thus prohibiting accurate calculation of sulfate reduction and methanogenesis rates. Although, in theory, H₂S oxidation is a more likely source for the excess sulfate, we believe that sulfate-rich groundwater input is a more likely source due to correlation between excess sulfate and excess Sr which has been previously described for groundwater in this region (Young et al., 2008). Perry et al. (2002) identified dissolution of evaporites within the freshwater lens at some Yucatan sites as a probable source of excess sulfate in groundwater using the sulfate-to-chloride ratio (100×[SO₄²⁻/Cl⁻]). Ratios higher than seawater (average seawater is 10.3) are expected where gypsum/anhydrite dissolution occurs (Perry et al. 2002). Another indicator is the Sr/Cl ratio which is invariably higher in the Yucatan groundwater than in seawater and indicates dissolution of celestite (from evaporite) and/or aragonite (Perry et al., 2002). The region east and south of Lake Chichancanab, referred to as the Evaporite Region by Perry et al. (2002), is characterized by distinctive topography and high sulfate groundwater concentrations (Perry et al., 2002). The groundwater from the Lake Chichancanab area flows northward into the Celestún Estuary which can be recognized by the progressive decrease in the ratio $\frac{Sr_{groundwater}}{Sr_{seawater}}$ in water from southeast to northwest (Perry et al., 2009). The Sr and sulfur trends for Celestún lagoon (Young et al., 2008) are consistent with our interpretation that gypsum/anhydrite dissolution in groundwater is the source of excess sulfate in the porewater of Group-3 in Celestún lagoon. Due to the impact of groundwater, our sulfate reduction and methanogenesis rates estimated using the model are minimum rates and independent rates of methanogenesis rates estimated using the model are minimum rates and independent rates of methanogenesis rates estimated using the model are minimum rates and independent rates of
groundwater discharge into each core are needed for obtaining more realistic estimates in these sites.

In addition to depth–integrated rates, Table 3 also includes maximum methanogenesis/methane production (Max-RM) and sulfate reduction/consumption (Max-RSR) rates solved by Eq. 2 in the model. Interestingly, the maximum methane production rates estimated from TMA, methanol and H₂ additions to sediments in the slurry incubations (Table 1) are similar to model derived Max-RM at station 16CEL (Table 3), which is the site from which sediments were collected for the slurry incubations. The rates in the TMA, methanol and H₂ treatments from the slurry incubations (Table 1) and in some of our stations are higher than methane production rates from previously reported coastal freshwater and brackish wetland sediments that were measured using radiolabeled acetate and bicarbonate in slurries (Segarra et al., 2013).

Modeled Max-RM in some cores were 1-2 orders of magnitude higher than rates derived from the sediment slurry incubations (e.g., cores 1CEL Jul02, 1 1CH Oct01, 2CEL Oct01 and 14CEL Dec00). Although heterotrophic sulfate reduction generally dominates organic matter degradation, Max-RM values are even higher than the maximum sulfate reduction rates in some cores (1 1CH Oct01, 1 2CH Oct01 and 1CH Dec00). Both the methanogenesis rates measured in the sediment slurry incubations and the modeled maximum methanogenesis rates in this study area were much higher than those reported for some mangrove systems (e.g., Thailand, Kristensen et al., 2000; Malaysia, Alongi et al., 2004; Australia, Kristensen and Alongi, 2006) but similar to other sites in India (Ramesh et al., 2007).

AOM is expected to play an important role in tropical porewaters with abundant methane and sulfate (Biswas et al., 2007). However, our model results and sensitivity analyses indicate that AOM is insufficient to prevent methane escape to the bottom water, probably because of the abundant organic matter available for sulfate reducers to use instead of methane. In our sensitivity tests (using core 1CEL Oct01 as an example), if AOM is allowed to be responsible for sulfate and methane consumption (no heterotrophic sulfate reduction and methanogenesis; \( R_{SO4} = R_{AOM} \)) then methane concentrations would decrease to negative values (gray solid lines in Fig. 5B), which is inconsistent with observations. Although based on our data it is not possible to accurately quantify the relative proportion of sulfate loss due to heterotrophic sulfate reduction and/or AOM, our model results suggest AOM plays a minor role in this setting.

Future investigations on the role of AOM in these dynamic mangrove-dominated tropical coastal lagoons are needed (e.g., Thalasso et al., 1997; Raghoebarsing et al., 2006; Lee et al., 2008; Kristensen et al., 2008; Beal et al., 2009; Silvan et al., 2011; Segarra et al., 2013).
6 Conclusions

The variable trends observed in porewater chemistry indicate a very dynamic system spatially and temporally throughout the year. This can be explained by physical processes such as mixing and dilution with seawater or groundwater, gas bubble rise and dissolution, and microbial processes which operate at different rates during different times at all sites. Although our modeling suggests that organic carbon degradation rates are dominated by heterotrophic sulfate reduction in these cores, methanogenesis both in shallow and deeper sediments is prevalent. The co-occurrence of methane and sulfate reduction (documented by sulfate depletion) in shallow sediments of non-competitive substrates and ample dissolved and labile organic matter in the shallow sediments as well as the contributions of methane from deeper sediment through gas rise and dissolution. Model results demonstrate that the largest sink for methane in these sediments is efflux to the water column. Build-up of methane at shallow depths may reduce the fraction of methane that is oxidized prior to entering the water column, thereby increasing the flux at the sediment-water interface. This shallow methane pool may also encourage methane flux through bubble release, which can result in a larger fraction of the methane reaching the atmosphere without being lost to oxidation. Specifically, the ability of the microbial community in these sediments to use non-competitive substrates may allow for methane production in the upper sections of the sediment potentially contributing to the higher than expected atmospheric methane flux measured from mangrove-dominated tropical coastal lagoons.

Appendix: Modeling procedure used in the evaluation of porewater observations from sediments in mangrove-dominated tropical coastal lagoons, Yucatán, Mexico

Details of the modeling procedure and parameters used are described here. The following reactions are considered in the model:

Heterotrophic sulfate reduction ($R_{SR}$):

\[ 2\text{CH}_2\text{O} + \text{SO}_4^{2-} \rightarrow 2\text{HCO}_3^- + \text{H}_2\text{S} \]  

\hspace{1cm} (R1)

Methanogenesis ($R_M$):

\[ 2\text{CH}_2\text{O} \rightarrow \text{CO}_2 + \text{CH}_4 \]  

\hspace{1cm} (R2)

Gas bubble dissolution ($R_{MB}$):

\[ \text{CH}_4(g) \rightarrow \text{CH}_4(aq) \]  

\hspace{1cm} (R3)

The net reaction terms ($R_c$ in Eq. 2) are given in Table A1, boundary conditions are listed in

- 16 -
Table A2, best-fit model parameters are given in Table A3 and model derived concentration profiles are shown in Fig. 2 and Fig. A1. 

In Eq. (2), sediment porosity decreases with depth due to steady-state compaction:

\[ \Phi = \Phi_f + (\Phi_0 - \Phi_f) \cdot e^{-D_v v} \]  

(A1)

where \( \Phi_f \) is the porosity below the depth of compaction (0.78 for Celestún and 0.83 for Chelem), \( \Phi_0 \) is porosity at the sediment surface (0.90 for Celestún and 0.89 for Chelem) and \( p \) (1/15 cm\(^{-1}\)) is the depth attenuation coefficient. These parameters were determined from the measured porosity data at each site or at a nearby site (Eagle, 2002). Under the assumption of steady state compaction, the burial of porewater was calculated as in (Wallmann et al. 2006):

\[ v = \frac{\Phi_f \cdot w_f}{\Phi} \]  

(A2)

where \( w_f \) is the sedimentation rate of compacted sediments calculated from excess \(^{210}\)Pb data (0.25 cm/yr for Celestún and 0.35 cm/yr for Chelem; Gonneea et al. (2004)). Sediment burial results in the downward movement of both sediment particles and porewater relative to the sediment water interface.

The sediment diffusion coefficient of each solute (\( D_s \)) was calculated according to Archie’s law considering the effect of tortuosity on diffusion (Boudreau 1997):

\[ D_s = \Phi^2 \cdot D_M \]  

(A4)

where \( D_M \) is the molecular diffusion coefficient at the in situ temperature, salinity and pressure (Table A1) calculated according to (Boudreau 1997). We used the same tortuosity coefficient (\( \Phi^2 \) corresponding to \( m = 3 \) in Archie’s law) as reported by Wallmann et al. (2006) for fine-grained sediments.

Since net sulfate consumption is observed in Groups 1 profiles (Fig. 2; Fig. A1), we use the following calculations to obtain net sulfate depletion rates (\( R_{SO_4} \) mmol SO\(_4\)\(^{2-}\) cm\(^{-1}\) yr\(^{-1}\)). \( R_{SO_4} \) is proportional to the difference between modeled (\( C(SO_4^{2-}icone text here: 

\[ (C(SO_4^{2-})_{obs}) \]  

-17
The corresponding kinetic constant is set to be high ($k_{SD} \geq 100$ $\text{y}^{-1}$) to ensure that simulated concentrations are very close to measured values. $R_{SD}$ implicitly includes $R_{SR}$ as well as anaerobic oxidation of methane ($R_{AOM}$):

$$\text{CH}_4 + \text{SO}_4^{2-} \rightarrow \text{HCO}_3^- + \text{HS}^- + \text{H}_2\text{O} \quad (R4)$$

The numerical modeling procedure outlined in Wallmann et al. (2006) is used as a basis to simulate the rate of sedimentary organic carbon degradation ($R_{POC}$) by sulfate reduction and methanogenesis. Since the measured organic matter content in both lagoons showed evidence for a change in depositional pattern over time (Gonneea et al., 2004 and Eagle, 2002), these measurements cannot be used for reliable organic matter degradation calculations. Hence, $R_{SD}$ (Eq. A5 below) was first calculated and then used to estimate $R_{POC}$ (Eq. A6) and subsequently to derive $R_M$ (Eq. A7). Here, we assume the three reactions (R1, R2 and R4) co-occur in the sulfate reduction zone such that the net reaction for methanogenesis and AOM (reactions R2+R4) is equal to carbon respiration by heterotrophic sulfate reduction (reaction R1). In other words, $R_{SD} = 0.5R_{POC}$.

To approximate the fraction of $R_{POC}$ due to $R_{SD}$ and $R_{SR}$, a Michaelis-Menten kinetic limitation term is applied to Eq. (A5-A7) (Wallmann et al., 2006):

$$R_{SR} = R_{SD} = 0.5 \cdot R_{POC} \cdot f_{SO_4^{2-}} \quad (A5)$$

$$R_{POC} = \frac{R_{SD}}{0.5 \cdot f_C \cdot f_{SO_4^{2-}}} \quad (A6)$$

$$R_M = 0.5 \cdot f_C \cdot R_{POC} \cdot (1 - f_{SO_4^{2-}}) \quad (A7)$$

where $f_{SO_4^{2-}} = \frac{c_{SO_4^{2-}}}{c_{SO_4^{2-}} + K_{SO_4}}$ is the Michaelis–Menten rate-limiting term for sulfate reduction.

At sites where methanogenesis was insufficient to simulate the measured methane data, methane was added as an external source by dissolution of gas bubbles (Chuang et al., 2013).
Gas bubbles were observed in the field. The rate of dissolution of the gas bubbles (R3) rising through the sediment (\(CH_4(g) \rightarrow CH_4\)) was also considered as (Haeckel et al., 2004):
sediment surface to 0.38 cm at depth. Since most of the porewater profiles were fitted directly, only a few years of simulation time (5 yr) was needed to achieve steady state. Mass balance was typically better than 99.9%.

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Table 1: Experimental conditions and sampling time intervals for methane headspace concentration analysis of sediment slurry incubations.

<table>
<thead>
<tr>
<th>Treatment</th>
<th>Initial concentration of treatment</th>
<th>Experiment length (days)</th>
<th>Number of measurements</th>
<th>Methane production rate (nmol CH₄ cm⁻³ slurry d⁻¹)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Controls</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>No amendment (anaerobic)</td>
<td>N₂ headspace</td>
<td>29</td>
<td>3</td>
<td>1.3 × 10⁻⁴ to 2.0 × 10⁻³</td>
</tr>
<tr>
<td>Autoclaved</td>
<td>N₂ headspace</td>
<td>29</td>
<td>3</td>
<td>0 to 2.6 × 10⁻¹</td>
</tr>
<tr>
<td>Aerobic- O₂ gas</td>
<td>16% O₂ headspace (0.36 mM)</td>
<td>29</td>
<td>3</td>
<td>5.7 × 10⁻⁴ to 3.5 × 10⁻³</td>
</tr>
<tr>
<td>BES</td>
<td>40 mM</td>
<td>29</td>
<td>3</td>
<td>0 to 1.3 × 10⁻⁴</td>
</tr>
<tr>
<td>Competitive substrates</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>H₂ gas</td>
<td>100% headspace (1.8 mM)</td>
<td>29</td>
<td>3</td>
<td>5.4 × 10⁻³ to 6.2</td>
</tr>
<tr>
<td>Acetate</td>
<td>10 mM</td>
<td>29</td>
<td>3</td>
<td>6.8 × 10⁻⁴ to 9.2 × 10⁻⁴</td>
</tr>
<tr>
<td>Formate</td>
<td>10 mM</td>
<td>29</td>
<td>3</td>
<td>6.9 × 10⁻⁴ to 1.6 × 10⁻⁴</td>
</tr>
<tr>
<td>Noncompetitive substrates</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Methanol</td>
<td>10 mM</td>
<td>29</td>
<td>4</td>
<td>2.0 × 10⁻² to 19</td>
</tr>
<tr>
<td>TMA</td>
<td>10 mM</td>
<td>29</td>
<td>4</td>
<td>5.4 × 10⁻⁴ to 40</td>
</tr>
</tbody>
</table>
Table 2. Model-derived depth–integrated turnover rates (mmol m\(^{-2}\) d\(^{-1}\)), dissolved methane fluxes to the water column (mmol m\(^{-2}\) d\(^{-1}\)) and contributions of methanogenesis to net methane production (%) and heterotrophic sulfate reduction to POC degradation (%). CEL and CH represent cores collected from Celestún Lagoon and Chelem Lagoon.

<table>
<thead>
<tr>
<th>Group</th>
<th>Length of model column (cm)</th>
<th>(R_{SO4}^{−})</th>
<th>(R_{R2}^{−})</th>
<th>(R_{POC}^{−})</th>
<th>(R_{R4}^{−})</th>
<th>(F_{methane}(top))</th>
<th>(F_{methane}(bottom))</th>
<th>(R_{R4}/(R_{R4}+R_{R3}))</th>
<th>(2R_{R4}/R_{POC})</th>
<th>(F_{POC})</th>
<th>(F_{CH4})</th>
</tr>
</thead>
<tbody>
<tr>
<td>Group 1</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>1CEL_Apr00</td>
<td>20</td>
<td>3.7</td>
<td>0.13</td>
<td>7.7</td>
<td>0.41</td>
<td>0.59</td>
<td>0.06</td>
<td>25%</td>
<td>97%</td>
<td></td>
<td></td>
</tr>
<tr>
<td>1CEL_Dec00</td>
<td>20</td>
<td>2.2</td>
<td>1.5</td>
<td>7.4</td>
<td>0</td>
<td>0.94</td>
<td>-0.60</td>
<td>100%</td>
<td>59%</td>
<td></td>
<td></td>
</tr>
<tr>
<td>1CEL_Oct01</td>
<td>20</td>
<td>6.2</td>
<td>0.12</td>
<td>13</td>
<td>0.32</td>
<td>0.40</td>
<td>-0.04</td>
<td>27%</td>
<td>98%</td>
<td></td>
<td></td>
</tr>
<tr>
<td>1CEL_Jul02</td>
<td>20</td>
<td>3.6</td>
<td>8.0</td>
<td>23</td>
<td>0</td>
<td>6.0</td>
<td>-1.98</td>
<td>100%</td>
<td>31%</td>
<td></td>
<td></td>
</tr>
<tr>
<td>2CEL_Dec00</td>
<td>20</td>
<td>1.1</td>
<td>0.05</td>
<td>2.3</td>
<td>0.76</td>
<td>0.54</td>
<td>-0.27</td>
<td>5.8%</td>
<td>96%</td>
<td></td>
<td></td>
</tr>
<tr>
<td>2CEL_Jul02</td>
<td>20</td>
<td>11</td>
<td>0.08</td>
<td>22</td>
<td>0.05</td>
<td>0.11</td>
<td>-0.02</td>
<td>63%</td>
<td>99%</td>
<td></td>
<td></td>
</tr>
<tr>
<td>3CEL_Apr00</td>
<td>20</td>
<td>1.3</td>
<td>0.29</td>
<td>3.2</td>
<td>0.24</td>
<td>0.68</td>
<td>0.15</td>
<td>55%</td>
<td>82%</td>
<td></td>
<td></td>
</tr>
<tr>
<td>3CEL_Jul02</td>
<td>20</td>
<td>7.1</td>
<td>0.24</td>
<td>15</td>
<td>2.2</td>
<td>3.0</td>
<td>0.63</td>
<td>10%</td>
<td>97%</td>
<td></td>
<td></td>
</tr>
<tr>
<td>1CEL_Oct01</td>
<td>20</td>
<td>13.75</td>
<td>24</td>
<td>31</td>
<td>110</td>
<td>0</td>
<td>11</td>
<td>-19.54</td>
<td>100%</td>
<td>42%</td>
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</tr>
<tr>
<td>1CEL_Oct01</td>
<td>20</td>
<td>3.0</td>
<td>26</td>
<td>58</td>
<td>0</td>
<td>20</td>
<td>-5.6</td>
<td>100%</td>
<td>10%</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Group 2</td>
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<tr>
<td>1CH_Dec00</td>
<td>20</td>
<td>0.52</td>
<td>-7.0</td>
<td>0</td>
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<td>4.5</td>
<td>7.8</td>
<td></td>
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<td>0</td>
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<td>3.2</td>
<td>-0</td>
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<td>-0</td>
<td>-0</td>
<td>0</td>
<td></td>
<td>6.9</td>
<td>-0</td>
<td></td>
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<td>11</td>
<td>-0.01</td>
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<td>11</td>
<td></td>
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<td>14CEL_Jul02</td>
<td>20</td>
<td>0.27</td>
<td>-0</td>
<td>0</td>
<td></td>
<td>3.7</td>
<td>0.27</td>
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<td>20</td>
<td>-0.047</td>
<td>0.013</td>
<td>0</td>
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<td>-1.8</td>
<td>-0.060</td>
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<td>Group 3</td>
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<td>-0.01</td>
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<td>14CEL_Dec00</td>
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<td>3.4</td>
<td>-0.13</td>
<td>0</td>
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<td>3.6</td>
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<td>0.088</td>
<td>-0.01</td>
<td>0</td>
<td></td>
<td>2.9</td>
<td>0.10</td>
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<td>Group 4</td>
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4 \(R_{SO4}^{−}\) is net sulfate depletion (mmol m\(^{-2}\) d\(^{-1}\) of SO\(_{4}^{−}\)). \(R_{R2}^{−}\) is heterotrophic sulfate reduction (mmol m\(^{-2}\) d\(^{-1}\) of SO\(_{4}^{−}\)). \(R_{R2}\) is methanogenesis rate (mmol m\(^{-2}\) d\(^{-1}\) of CH\(_{4}\)). \(R_{POC}\) is total POC mineralization (mmol m\(^{-2}\) d\(^{-1}\) of CH\(_{4}\)). \(F_{methane}(bottom)\) is methane flux across the sediment surface (mmol m\(^{-2}\) d\(^{-1}\) of CH\(_{4}\)). \(F_{methane}(top)\) is the methane flux to the water column and vice versa. \(F_{POC}\) is the methane flux out of the sediments (mmol m\(^{-2}\) d\(^{-1}\) of CH\(_{4}\)). Negative values in \(F_{methane}\) represent methane flux to deep sediments and vice versa. \(R_{SO4}^{−}\) is net sulfate input (mmol m\(^{-2}\) d\(^{-1}\) of SO\(_{4}^{−}\)) and \(R_{CH4}\) is net methane production (mmol m\(^{-2}\) d\(^{-1}\) of CH\(_{4}\)) for cores in group 2 to group 4. See Appendix for further model details.
Table 3: Maximum model-derived rates of methanogenesis and sulfate reduction for cores in Group-1 and maximum model-derived rates of methane production and sulfate consumption for cores in Group-2, Group-3 and Group-4. CEL and CH represent cores collected from Celestún Lagoon and Chelem Lagoon.

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<th>Max-R&lt;sub&gt;CH4&lt;/sub&gt; (nmol CH&lt;sub&gt;4&lt;/sub&gt; cm&lt;sup&gt;-3&lt;/sup&gt; d&lt;sup&gt;-1&lt;/sup&gt;)</th>
<th>Max-R&lt;sub&gt;SR&lt;/sub&gt; (nmol SO&lt;sub&gt;4&lt;/sub&gt;&lt;sup&gt;2-&lt;/sup&gt; cm&lt;sup&gt;-3&lt;/sup&gt; d&lt;sup&gt;-1&lt;/sup&gt;)</th>
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Table A1: Rate expressions applied in the differential equations ($R_c$ in Eq. (2))

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<td>CH$_4$</td>
<td>$+R_M + R_{MB}$</td>
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<tr>
<td>SO$_4^{2-\text{dep}}$</td>
<td>$+R_{SD}$</td>
<td>Group-1</td>
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<td>SO$_4^{2-}$</td>
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<td>Group-2, Group-3 and Group-4</td>
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### Table A2: Boundary conditions used in the model

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<th>$\text{CH}_4$ (top)</th>
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<th>$\text{SO}_4^{2-}$ dep (bottom)</th>
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Table A3: Imposed and best-fit parameters in each core

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<th>T (°C)</th>
<th>S (bar)</th>
<th>P (cm² yr⁻¹)</th>
<th>Dₑ(CH₄) (cm² yr⁻¹)</th>
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<th>$\Delta$SR</th>
<th>$k₀₂$ (mM yr⁻¹)</th>
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- Table continues...
Figure 1: Maps of (A) the Yucatán Peninsula with lagoon locations, (B) Celestún Lagoon and (C) Chelem Lagoon showing the sampling stations (circles) of sediment cores.
Figure 2: Depth profiles of modeled (lines) and measured (circles) and calculated (triangles) concentration of dissolved methane (dashed lines; open circles), sulfate (solid lines; solid circles) in the upper panel and sulfate depletion (solid lines; solid circles), zero sulfate depletion (dashed lines) and chloride concentration (gray circles) in the lower panel for each profile type (Groups 1-4, see text). One selected profile per group is shown here and the other profiles for each group (9 cores for Group-1, 6 cores for Group-2, 2 cores for Group-3 and 3 cores for Group-4) are presented in the Appendix (Fig. A1). CEL and CH represent cores collected from Celestún Lagoon and Chelem Lagoon.
Figure 3: Relationship between (A) $[\text{Cl}^-]$ and $[\text{SO}_4^{2-}]$, (B) $[\text{Cl}^-]$ and $[\text{CH}_4]$ and (C) relationship between $[\text{CH}_4]$ and $[\text{SO}_4^{2-}]$ in porewater samples.
Figure 4: (A) Headspace methane concentrations in sediment slurry incubations. (B) Expansion of (A), showing results for acetate, formate, and controls. (C) Expansion of (A), showing results for controls only. Error bars represent one standard deviation for triplicate sample bottles.
Figure 5: Model sensitivity analysis of methane concentrations for cores in Group 1 to the different processes controlling methane concentrations in porewaters. Black dashed lines denote the standard simulation results: \( CH_4 \) production rate = \( R_{\text{methanogenesis}} + R_{\text{M}} + R_{\text{MB}} \). \( R_{\text{methanogenesis}} \) is methanogenesis, \( R_{\text{MB}} \) is methane bubble dissolution, \( R_{\text{AOM}} \) is anaerobic oxidation of methane and \( R_{\text{SD}} \) is net sulfate depletion.
Figure A1: Depth profiles of modeled (lines), measured (circles) and calculated (triangles) concentration of dissolved methane (dashed lines; open circles), sulfate (solid lines; solid circles) in the upper panel and sulfate depletion (solid lines; solid triangles), zero sulfate depletion (dashed lines) and chloride (gray circles) in the lower panel for each profile type (Groups 1-4, see text). One selected profile per group is shown in Fig. 2 for illustration and here the rest of other profiles are shown (9 cores for Group-1, 6 cores for Group-2, 2 cores for Group-3 and 3 cores for Group-4). CEL and CH represent cores collected from Celestún Lagoon and Chelem Lagoon.
Figure A1: Continued.
Figure A1: Continued.
Figure A1: Continued.

Deleted: Figure 6: Sensitivity of methane concentrations for cores in group 1 and group 2 to the different processes: (A) $R_{CH4} = -R_{AOM} + R_{MB}$ (gray solid line; $k_{MB} = 1000 \text{ yr}^{-1}$), (B) $R_{CH4} = -R_{AOM} + R_{MB}$ (gray solid line; $k_{MB} = 0.5 \text{ yr}^{-1}$), $R_{CH4} = -R_{AOM} + R_{MB}$ (gray dashed line; $k_{MB} = 0.2 \text{ yr}^{-1}$), and $R_{CH4} = -R_{AOM} + R_{M}$ (black solid line), (C) $R_{CH4} = -R_{AOM} + R_{MB}$ (gray solid line), (D) $R_{CH4} = -R_{AOM}$ (gray solid line), (E) $R_{CH4} = -R_{AOM} + R_{MB}$ (gray solid line), (F) $R_{CH4} = -R_{AOM} + R_{MB}$ (gray solid line). Black dashed curves denote the standard simulation results: (A) $R_{CH4} = -R_{AOM} + R_{MB}$ and (B) $R_{CH4} = -R_{AOM} + R_{MB}$ ($k_{MB} = 0.1 \text{ yr}^{-1}$).