Author responses to:

Interactive comment on “Impacts of temperature and soil characteristics on methane production and oxidation in Arctic polygonal tundra” by Jianqiu Zheng et al.

We appreciate both referee’s reviewing our revised manuscript and are glad that our changes have adequately addressed Referee #1’s comments. And we have used the thoughtful comments from new Referee #3 to clarify methodology in the second revision of this paper. This document includes responses to Referee #3’s comments. Finally, we have attached a comparison of the revised manuscript with changes tracked from the originally submitted version.

Anonymous Referee #1

No comments submitted for this version of the manuscript.

Anonymous Referee #3

This manuscript reports effects of temperature and soil layers on CH4 production and oxidation rates in the arctic tundra with a soil incubation experiment. I have major comments regarding the incubation methods, which lead to the results and conclusions:

1) The authors introduced that low-centered polygons (LCPs) are wetter than flat-centered polygons (FCPs) and high-centered polygons (HCPs) (P4). Then why not taking samples from wetter LCPs?

We previously measured and observed high rates of methanogenesis and CO2 production in wet and saturated LCP samples (Roy Chowdhury et al., 2015). Field measurements of CH4 fluxes from surface soils demonstrated higher CH4 emissions from saturated LCPs than drier FCPs or HCPs, but steeper gradients in porewater methane concentrations in FCP/HCP features (e.g. Vaughn et al., 2016). Therefore, we focused our aerobic CH4 oxidation experiments on FCP samples that were expected to be more significant sources of CH4 oxidation activity. This rationale has been clarified in the second paragraph of the Introduction.

2) It is not clear if all soil samples are incubated anaerobically (closed lids)? If not, soil moisture would change over the 90 days of incubation, and thus changing the CH4 oxidation rates. If yes, the aerobic condition would be totally different from that in the field. Also, would the CH4 concentration in the air space be saturated and inhibit further production of CH4?

We modified methods section 2.3 to clarify that “All vials were sealed with butyl rubber septa and crimp sealed to prevent evaporation and gas exchange.” Soil moisture did not change in these sealed vials. The headspace gases for the incubation experiments were selected to match the soil’s redox properties and are described in section 2.3 and illustrated in Figure 1.
Due to the low solubility of CH\textsubscript{4} in water, it is unusual for CH\textsubscript{4} accumulation to inhibit methanogenesis in a liquid-gas system. In our incubations, CH\textsubscript{4} accumulated with a linear rate profile (Figure 3) showing no indication of product inhibition or even substrate limitation. For comparison, biogas from anaerobic digestors frequently reaches 60-70\% CH\textsubscript{4} concentration—orders of magnitude higher than our incubations.

3) P5. L15. Soil samples were homogenized. Then the soil structure was damaged, compared with that in the field. How does soil texture affect the CH\textsubscript{4} production and consumption?

The Roy Chowdhury et al. (2015) reference was added to section 2.3 to explain the homogenization process, along with a short description of the homogenization process and its effects on the soil structure. Our method using an oscillating power tool inside a glove box disrupts large, frozen soil clumps and removes gravel or litter, allowing us to place representative soil samples inside the serum vials. This method does not affect the soil microaggregate structure or expose the samples to oxidation, drying or significant warming that could disrupt anaerobic microbial processes. We recognize that incubation experiments do not reproduce the gradients and transport processes of soil pedons in the field (see below). The small soil sample volume used in our microcosms is unlikely to present diffusion constraints, and we assume equilibrium with headspace in our measurements. Our experimental set-up, therefore, could focus on the specific question of understanding the temperature sensitivity of the distinct processes of interest, i.e. methane production and potential oxidation.

4) How does the microbial community differ (in diversity and density) during the incubation from those in the field? This shift may change the Q10 function if the incubation results would be used to predict the field condition.

We agree that increased temperatures are likely to change microbial community composition, both in incubations and in the field (see for example Hoj et al., 2007, ISME J. 2:37 and Conrad et al. 2009, Environ. Microbiol. 11:1844). However, Q\textsubscript{10} relationships are strictly empirical representations of a processes’ temperature sensitivity. Microbial community changes, differences in gene expression and enzyme production, and fundamental Arrhenius enzyme kinetics are all grouped together in this term, along with any changes in sorption or transport rates. Current process-enabled models do not distinguish among these factors. To reduce the influence of microbial biomass changes on Q\textsubscript{10} calculations, we use a ratio of maximum activity rates, which are usually highest at the beginning of activity or immediately after a lag period (see Fig. 6 and our extended discussion in Roy Chowdhury et al., 2015).

Future manipulation experiments in the field and novel model structures and parameterization may provide a framework to quantify changes in microbial community composition from laboratory and field warming experiments and begin to differentiate mechanisms underlying Q\textsubscript{10} relationships. We added a short comment on this issue on pp. 12-13 of the revised manuscript.

Minor comments:
Title: remove “polygon”, which causes confusion to readers.

Done.
P2. L14. It is unclear here if the “source strength” means the carbon producing CO2 or CH4.

This sentence has been revised for clarity: “... a current estimate of net CH4 exchange from tundra to the atmosphere ranging widely from 8 to 29 Tg C yr\textsuperscript{-1}.”


We added a citation to the Le Mer and Roger (2001) review whose conceptual model of methane cycling in wetlands has been highly reproduced. A reference to the metatranscriptomic study by Kim and Liesack (2015) was also added to this paragraph.

P4. L13. Describe the diameter of the “SIPRE auger”.

We added the 3-inch inner diameter dimension of the core liner to this description. Additional details about this auger are published in Roy Chowdhury et al. (2015).

P12. L26. The diffusion should be discussed. This also relates to my previous comment that the soil structure and text determine the diffusion of gas fluxes (CO2 and CH4). Thus, the change in diffusivity may lead to changes in gas effluxes and the response to temperature.

This discussion section uses observed Q\textsubscript{10} values to develop an example simulation of competing methane oxidation and methanogenesis rates in the thin FCP transitional layer. It illustrates the impact of different process temperature sensitivities on potential CH4 fluxes and microbial biomass ratios in a warming environment.

A variety of transport processes in the soil column will certainly affect surface gas fluxes. However, our discovery that maximum rates of methanogenesis and methane oxidation occur in the same thin transitional layer of FCP soil enables this simple simulation that neglects diffusion. We added a corollary to this section reminding readers that future simulations will need to include a full range of gas transport processes in order to integrate soil horizons and model net surface gas fluxes.
Impacts of temperature and soil characteristics on methane production and oxidation in Arctic tundra

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Abstract

Rapid warming of Arctic ecosystems accelerates microbial decomposition of soil organic matter and leads to increased production of carbon dioxide (CO₂) and methane (CH₄). CH₄ oxidation potentially mitigates CH₄ emissions from permafrost regions, but it is still highly uncertain whether soils in high-latitude ecosystems will function as a net source or sink for CH₄ in response to rising temperature and associated hydrological changes. We investigated CH₄ production and oxidation potential in permafrost-affected soils from degraded ice-wedge polygons on the Barrow Environmental Observatory, Utqiaġvik (Barrow) Alaska, USA. Frozen soil cores from flat and high-centered polygons were sectioned into organic, transitional and permafrost layers, and incubated at -2, +4 and +8 °C to determine potential CH₄ production and oxidation rates. Significant CH₄ production was only observed from the suboxic transition layer and permafrost of flat-centered polygon soil. These two soil sections also exhibited highest CH₄ oxidation potential. Organic soils from relatively dry surface layers had the lowest CH₄ oxidation potential compared to saturated transition layer and permafrost, contradicting to our original assumptions. Low methanogenesis rates are due to low overall microbial activities measured as total anaerobic respiration and competing iron
reduction process. Our results suggest that CH$_4$ oxidation could offset CH$_4$ production and limit surface CH$_4$ emissions, in response to elevated temperature, and thus must be considered in model predictions of net CH$_4$ fluxes in Arctic polygonal tundra. Future changes in temperature and soil saturation conditions are likely to divert electron flow to alternative electron acceptors and significantly alter CH$_4$ production, which should also be considered in CH$_4$ models.

1 Introduction

Arctic ecosystems store vast amounts of organic carbon in active layer soils and permafrost (Hugelius et al., 2014; Shiklomanov et al., 2010). Rising temperatures, increased annual thaw depth, and a prolonged thaw season accelerate microbial degradation of this carbon reservoir (Shiklomanov et al., 2010; Schuur et al., 2015; Schuur et al., 2013). The potential carbon loss due to these direct effects is estimated to be 92±17 Pg carbon over the coming century (Schuur et al., 2015). The extent of soil organic matter (SOM) decomposition and partitioning between CO$_2$ and CH$_4$ emissions highly depend upon soil saturation conditions. Thawing of ground ice and ice-wedge degradation cause ground subsidence and significant changes in soil water saturation (Liljedahl et al., 2016) generating heterogeneous surface CH$_4$ fluxes (Schädel et al., 2016), with a current estimate of net CH$_4$ exchange from tundra to the atmosphere ranging widely from 8 to 29 Tg C yr$^{-1}$ (McGuire et al., 2012).

Understanding the factors that control CH$_4$ fluxes is key to reducing model uncertainties and predicting future climate feedbacks. The unique polygonal ground in Arctic coastal plain tundra creates natural gradients in hydrology, snow pack depth and density, and soil organic carbon storage that control CH$_4$ fluxes (Liljedahl et al., 2016; Lara et al., 2015). Thermal contraction processes create cracks in the tundra soil, which can fill with water that freezes to produce massive ground ice (French, 2007). This ice forms wedges that create the borders of three dominant polygon types, defined by their surface relief and subsurface hydrology: low-centered polygons (LCPs), flat-centered polygons (FCPs), and high-centered polygons (HCPs) (MacKay, 2000). Poorly drained LCPs are characterized by wet centers bordered by raised, relatively dry rims and wet troughs. FCPs lack the rims of LCPs and are drier (Wainwright et al., 2015). When ice wedges erode and water drains from the polygons, troughs subside and rims disappear to form drier HCPs. Methane emissions from wet and inundated LCP sites were 1-2 orders of magnitude larger than emissions from drier FCP and HCP sites (Vaughn et al., 2016; Sachs et al., 2010). Although a number of factors, including vegetation height and plant composition (von Fischer et al., 2010), soil inundation (Sturtevant et al., 2012), thaw depth (Sturtevant and Oechel, 2013; Grant et al., 2017), and season (Chang et al., 2014) were suggested as explanatory factors for CH$_4$ flux variations, the hundred-fold difference in CH$_4$ flux between polygon types could not be fully explained by variations in moisture or temperature (Sachs et al., 2010; Vaughn et al., 2016). Measurements of dissolved CH$_4$ concentrations in soil pore waters suggested a huge disconnect between >100 µM concentrations of dissolved CH$_4$ in the active layer below 20 cm and negligible dissolved CH$_4$ at 10 cm soil depth (~1 µM). This CH$_4$ gradient is particularly steep in FCPs.
δ¹³C-CH₄ data suggested CH₄ oxidation played an important role in mitigating CH₄ in soil porewater and limiting surface CH₄ emissions (Vaughn et al., 2016). However, CH₄ oxidation potential is rarely studied in permafrost-affected soils. Therefore, we investigated CH₄ cycling in FCP soil for this study.

Methane oxidation mitigates terrestrial CH₄ emission. Up to 90% of CH₄ produced in the soil is consumed in the upper dry layers of soil by aerobic CH₄-oxidizing bacteria (methanotrophs) before reaching the atmosphere (Le Mer and Roger, 2001). Methane oxidation rates are usually greatest in oxic, surficial soils, although methane oxidation is known to occur under oxygen-limiting conditions as well (Roslev and King, 1996). In the classical model of CH₄ oxidation profiles, there is usually a vertical gradient of decreasing O₂ concentration in the top cm of the soil column that is inversely correlated with an increasing gradient of CH₄ through the suboxic/anoxic active layer (Le Mer and Roger, 2001). The relative abundance of methanotrophs generally correlates with CH₄ oxidation activity at the soil surface, and methanogens are relatively abundant in the deeper layer where oxygen is limiting (Lee et al., 2015; Kim and Liesack, 2015). Methane oxidation potential in wetlands and peat bogs is highest near the water table, while most CH₄ is produced below the water table (Whalen and Reeburgh, 2000). In contrast, methanogenic and methanotrophic communities can overlap in the rhizosphere (Liebner et al., 2012; Knoblauch et al., 2015), where roots or Sphagnum create an oxic/anoxic interface providing a substantial amount of oxygen for methanotrophs (Laanbroek, 2010; Parmentier et al., 2011) and organic acid substrates for methanogenesis.

Soil CH₄ fluxes result from the net effect of microbial CH₄ production and oxidation, coupled with transport processes. The rates of CH₄ oxidation are mainly governed by the abundance and composition of methanotrophic microbial communities and environmental factors including CH₄ and O₂ availability, soil air-filled porosity and soil-water content (Preuss et al., 2013). Previous studies of boreal lakes and wetlands showed that CH₄ production is more sensitive to temperature changes than CH₄ oxidation, as CH₄ oxidation rates respond more strongly to CH₄ availability than temperature increase (Liikanen et al., 2002; Segers, 1998). Soils at the Barrow Environmental Observatory in Utqiaġvik (Barrow), Alaska experience a wide range of arctic temperatures, from -20 to +4°C (Shiklomanov et al., 2010). Soil respiration and methanogenesis continue at low temperatures close to 0 °C, even after the soil surface freezes trapping gas under ice. Therefore, substantial annual CH₄ and CO₂ emissions from the Alaskan Arctic occur during the spring thaw (Commane et al., 2017; Raz-Yaseef et al., 2017; Zona et al., 2016). However, it is unclear how accelerated warming in Arctic soils affects the opposing processes of CH₄ production and oxidation due to their nonlinear response to temperature changes (Treat et al., 2015).

In this study, we investigated the rates and temperature sensitivities of CH₄ production and oxidation from permafrost-affected soils in Utqiaġvik (Barrow), Alaska. Although various studies have identified significant and frequently correlated factors affecting CH₄ and CO₂ production in permafrost ecosystems upon thawing, the oxidation of CH₄ is not considered in most incubation studies (Treat et al., 2015). Here we used the natural geomorphic gradient of FCP and HCP soils to represent degraded polygon tundra soils with relatively oxic active layers that will potentially act as CH₄ sinks. Methane production and
oxidation assays were performed separately using anoxic or oxic incubations at three temperatures. We also measured additional mechanisms influencing CH₄ production, including accumulation of organic acids and competing anaerobic respiration through iron reduction (Lipson et al., 2010). Specifically, we tested the following hypotheses regarding CH₄ dynamics in FCP and HCP soils: (I) CH₄ production is localized in the more reduced subsurface while CH₄ oxidation occurs at the soil surface; (II) CH₄ production is more sensitive to temperature increase than CH₄ oxidation and will likely exceed CH₄ oxidation in wet areas in response to warming.

2 Materials and Methods

2.1 Site description and soil sampling

The study site is located at the Barrow Environmental Observatory (BEO), Utqiaġvik (Barrow), Alaska as part of the Intensive Study Site areas B (High-Centered Polygon, HCP) and C (Flat-Centered Polygon, FCP) of the Next Generation Ecosystem Experiments in the Arctic project. The centers of HCPs in area B were covered by lichens, moss and dry tundra graminoids, while centers of FCPs in area C hosted wet tundra graminoids, mosses and bare ground (Langford et al., 2016). Intact frozen soil cores from the centers of a water-saturated FCP (N 71° 16.791', W 156° 35.990') and a well-drained HCP (N 71° 16.757', W 156° 36.288') were collected with a modified SIPRE auger containing a sterilized liner (3-inch inner diameter), driven by a hydraulic drill during a field campaign in April 2012 (Herndon et al., 2015a; Herndon et al., 2015b; Roy Chowdhury et al., 2015). All samples were kept frozen during core retrieval, storage and shipment to Oak Ridge National Laboratory (Oak Ridge, TN). The frozen cores were stored at -20 °C until processing. The thaw depth measured in September 2012 in the HCP center was 40 cm, and thaw depths in the FCP varied from 41-47 cm.

The frozen soil cores were inspected and processed inside an anaerobic chamber (Coy Laboratories, MI. H₂ ≤ 2% and O₂ < 1 ppm). Both cores were sectioned into 10-cm segments, and each segment was inspected for evidences of roots and undecomposed organic matter. Soil Munsell color was recorded to qualitatively infer redoximorphic conditions. The soil core collected from the FCP center showed evidence of buried and discontinuous organic matter at approximately 50 cm depth, below the active, organic layer and above the permafrost. This transition layer was attributed to episodic thawing and cryoturbation (Schuur et al., 2008; Bockheim, 2007). Soil geochemical properties were measured for each 10-cm segment to assess soil geochemical depth profiles. Specifically, we measured soil gravimetric water content and soil pH, using the 1:2 soil slurry method within 1M KCl. Fe(II) concentrations were measured to characterize redox conditions in the soil segments. Briefly, a soil subsample (~2 g) was extracted with 0.1M KCl solution for 30 min in an anaerobic chamber. The extracts were filtered with 2 µM syringe filters and analyzed immediately using the colorimetric 1,10-phenanthroline method (Hach method 8146). Absorbance was determined at 510 nm using a DU 800 spectrophotometer (Beckman Coulter, CA).
2.2 Soil pore water gas measurements

Dissolved gas (CO\(_2\) and CH\(_4\)) concentrations in soil pore water were determined at each 10-cm depth-interval from the FCP and HCP cores. A 1:1 (w:v) soil slurry was prepared by mixing 10 g wet soil in 10 mL de-ionized and de-gassed water under anoxic conditions inside an anaerobic chamber. The samples were then placed in 15 mL crimp-sealed serum vials (Wheaton, NJ). Vials were inverted and shaken at 4 °C for 12 h to allow for exchange between the dissolved and soil gas phases. Then, ~5 mL of the aqueous phase was exchanged with Ar (99.9 % purity) using a Gastight syringe (Hamilton, NV), and samples were manually shaken vigorously for 5 min to allow for equilibration between the aqueous and gas phase. Subsequently, a 500-µL headspace sample was drawn and immediately analyzed with an SRI 8610C gas chromatograph using the method previously described (Roy Chowdhury et al., 2015). The detection limit for CH\(_4\) was 1 ppm v. Concentrations of CH\(_4\) and CO\(_2\) were corrected for dissolved gases based upon temperature and pH-dependent Henry’s Law constants (Sander, 2015).

2.3 Low temperature respiration experiments

To investigate the temperature response of CH\(_4\) production and overall organic carbon mineralization rates, samples of FCP and HCP soil cores were incubated at -2, +4 or +8 °C for approximately 90 days. Based on measured geochemical similarities, core segments were combined to represent the organic layer that comprises most of the active layer, transition layer (only present in FCP), and permafrost. Soils from each layer were homogenized inside an anaerobic chamber using sterile tools and equipment to establish microcosms (Roy Chowdhury et al., 2015). Our homogenization method using an oscillating power tool inside a glove box disrupts large, frozen soil clumps and removes gravel or litter, allowing us to place representative soil samples inside the serum vials. This method does not affect the soil microaggregate structure or expose the samples to oxidation, drying or significant warming that could disrupt anaerobic microbial processes. Gases for the microcosm headspaces were selected based on gravimetric water contents and concentrations of reduced Fe(II) (Howeler and Bouldin, 1971) to best represent field conditions. Thus, organic layer soils from both FCP and HCP cores were incubated under air, while the transition layer and permafrost of FCP and HCP cores were incubated under anoxic conditions with N\(_2\) headspaces. All vials were sealed with butyl rubber septa and crimp sealed to prevent evaporation and gas exchange. Anoxic microcosms were flushed with N\(_2\) three times after sealing to remove residual H\(_2\) and O\(_2\) from the headspace. It is important to note that at -2 °C, soil water remained unfrozen in these samples due to freezing point depression (Romanovsky and Osterkamp, 2000).

A subsample of the combined, homogenized soil (~10 g) was placed in a sterile 70-mL serum bottle sealed with a blue butyl stopper and aluminum crimp seal to form a soil microcosm. For each homogenized soil layer, 9 replicate microcosms were constructed at each incubation temperature (Figure 1). Headspace CO\(_2\) and CH\(_4\) concentrations were measured at 2 to 15 days’ intervals using gas chromatography (see Section 2.2). After 5, 10 or 20 days of incubation, three replicated microcosms were destructively sampled for methane oxidation assays (see section 2.4 and Figure 1) and additional soil geochemical analysis.
Subsamples from microcosms opened after 10, 20 and 90 days’ incubation were further processed for analysis of pH, Fe(II) and total organic acids. pH and Fe(II) concentrations were measured as described in section 2.1. Iron reduction rates were estimated by the changes in measured Fe(II) concentration. Organic acids were analyzed using NH₄HCO₃ extracts. Briefly, NH₄HCO₃ extractions (12 h) were centrifuged for 15 min at 6500 g, then the supernatants were filtered through 0.2-µm membrane filters before analysis. Filtered sample were analyzed for low-molecular-weight organic acids using a Dionex ICS-5000’ system (Thermo Fisher Scientific, MA) equipped with an IonPac AS11-HC column with a KOH mobile phase. Total organic acid concentrations (TOA) were calculated by summing molar C equivalents for each measured species, normalized per gram of dry soil.

2.4 Methane oxidation potential assay

Soil samples were incubated in oxic conditions supplemented with ample CH₄ substrate to measure CH₄ oxidation potential (Roy Chowdhury et al., 2014). Methanogenesis is expected to be negligible under these oxic conditions. The methane oxidation assays (MOAs) were constructed using both freshly thawed (labeled as 0 day) and pre-incubated (labeled as various days of pre-incubation) samples to account for potential delays in the overall microbial activities (Figure 1). Replicated samples (about 2 g) were slurried in a 1:1 (w:v) ratio with autoclaved de-ionized water in 26-mL serum bottles under ambient condition. A 1% CH₄ headspace was introduced into each crimp-sealed bottle by replacing 0.23 mL headspace with 99.99% CH₄ (Scott, Air Liquide). For freshly thawed samples, 9 replicates were constructed for each incubation temperature and each soil layer. For pre-incubated samples at the designated sampling day, 3 replicated incubations (~10 g per sample) were destructively sampled to construct 9 replicates of MOAs to incubate at the same incubation temperature they were pre-incubated (Figure 1).

Due to the limited number of shaking-incubators, only the 4 °C MOA from FCP and 8 °C MOA from HCP were shaken to minimize potential gas-liquid phase transfer limitations. Headspace CO₂ and CH₄ concentrations in MOAs were measured at 2 to 15 days’ time intervals using gas chromatography (section 2.2).

2.5 Rate estimation, temperature sensitivity and statistical analyses

Concentrations of CH₄ and CO₂ from soil microcosms were fitted with hyperbolic, sigmoidal, or linear functions (Roy Chowdhury et al., 2015). Rates of CO₂ production were calculated using derivatives of the best curve-fitting equations with parameters listed in Table S3. Methane oxidation rates were calculated from the loss of headspace CH₄, which were best fitted with simple linear regression. All rate calculations are reported on per gram soil dry weight basis. The temperature dependence was calculated using the conventional Q₁₀ relationship by taking the ratio of maximum production or oxidation rates at 8 and -2°C based on triplicate measurements.
Changes of soil physicochemical properties were evaluated with one-way ANOVA, Tukey’s Honest Significant Difference (HSD) test. The effect of soil layers (organic, transition layer, and permafrost) and incubation temperature (-2, 4 and 8 °C) were examined with Tukey’s HSD test. All curve fittings and statistical analyses are performed with R 3.4.0 (The R Foundation for Statistical Computing) and validated with Prism (GraphPad Software, ver. 7.0a).

2.6 Calculation of net CH₄ emission

To evaluate the net result of CH₄ production and oxidation and how this result changes in response to temperature increase, we applied a simplified model simulation. Representation of the CH₄ oxidation rate (Roxi) is based on Michaelis-Menten kinetics with linear dependence on the biomass of methanotrophs (Xu et al., 2015), while the CH₄ production rate (Rpro) is calculated from measurements directly:

\[ R_{\text{oxi}} = B_{\text{methanotrophs}} \cdot V_{\text{max,oxi}} \left( \frac{C_{\text{CH}_4}}{C_{\text{CH}_4} + K_m,\text{CH}_4} \right) \left( \frac{C_{\text{O}_2}}{C_{\text{O}_2} + K_m,\text{O}_2} \right) \]

\[ R_{\text{pro}} = B_{\text{methanogens}} \cdot V_{\text{measure,pro}} \]

where \( B_{\text{methanotrophs}} \) and \( B_{\text{methanogens}} \) represent the estimated biomass of methanotrophs and methanogens respectively. \( K_{m,\text{CH}_4} \) and \( K_{m,\text{O}_2} \) are the half saturation coefficients (mM) with respect to CH₄ and O₂ concentrations, respectively. Values of \( K_{m,\text{CH}_4} \) and \( K_{m,\text{O}_2} \) vary within different models. We started with \( K_{m,\text{CH}_4} = 0.005 \) and \( K_{m,\text{O}_2} = 0.02 \) (Riley et al., 2011) and further applied a wide uncertainty range of 0.0005-0.05, 0.002-0.2. The maximum CH₄ oxidation rate \( V_{\text{max,oxi}} \) and CH₄ production rate \( V_{\text{measure,pro}} \) were obtained from the incubations. Initial CH₄ and O₂ concentrations were calculated from soil porewater dissolved gas measurement and soil air-filled porosity estimations. With the above parameters, we estimated the biomass ratio of methanogens to methanotrophs \((B_{\text{methanogens}} / B_{\text{methanotrophs}})\) under both net CH₄ production and net CH₄ oxidation scenarios, and further evaluated how the biomass ratio would change in response to rising temperature to keep the soil as a net source or sink of CH₄.

3 Results

3.1 Soil attributes and pore water characteristics

Soil cores from FCP and HCP center positions showed distinct vertical profiles of soil moisture expressed as gravimetric water content (g g⁻¹ dry soil). The soil core from FCP was characterized by a wet surface within the top 10 cm below ground, a much drier organic layer between 10 to 40 cm, and a bottom layer below 40 cm with significantly higher water content. In the HCP core, soil moisture gradually increased from the top to the bottom (Figure 2). Soil bulk density (g cm⁻³) is negatively correlated
with gravimetric water content in both FCP and HCP along the depth profile ($R^2$=-0.93, and $R^2$=-0.86, respectively). A similar water distribution has been recorded by continuous field measurements of volumetric water content at the nearby NGEE_BRW_C soil pit monitoring site (http://permafrostwatch.org). Fe(II) concentration showed a strong positive correlation with gravimetric water content in both FCP and HCP cores ($R^2$=0.81, and $R^2$=0.91, respectively). The soil pH in FCP increased steadily with soil depth ($R^2=0.95$), with an average of 4.7. In the HCP soil core, soil pH varied by 1.5 pH unit, with an average of 5.4. Overall, soil moisture, Fe(II) concentration and pH increased with depth in the centers of both polygon types.

Dissolved CO$_2$ in soil pore water showed a similar general trend in both FCP and HCP cores. The concentration of dissolved CO$_2$ increased from 100 µM in the surface soil (0-10 cm) to approximately 950 µM at 30 to 40 cm depth in FCP, and it decreased below 40 cm. In HCP, approximately 400 µM dissolved CO$_2$ was measured in the surface soil. The concentration was 400-500 µM in the top 30 cm and also decreased significantly below 40 cm. A strong correlation between dissolved CO$_2$ concentration and soil bulk density was observed for both FCP and HCP ($R^2=0.91$, and $R^2=0.85$, respectively).

The highest dissolved CH$_4$ concentration (about 85µM) was found between 30 to 40 cm in soil pore water of FCP, approximately 10 times the CH$_4$ concentration measured from the top 10 cm and 2-4 times higher than the CH$_4$ concentration measured below 40 cm. In HCP, significant CH$_4$ accumulation in soil pore water was found below 50 cm of the HCP core, while no dissolved CH$_4$ was detected above 50 cm.

Soil cores of FCP and HCP were divided into organic, transitional and permafrost layers to facilitate incubation setup. The top 10 cm of both cores contained mostly plant material, litter and ice or snow: these sections contained little soil and were not studied further. The organic layers of FCP and HCP cores were both oxic, with low Fe(II) concentrations and minimal dissolved CH$_4$ in soil pore water, while deeper layers were more reduced with 5-7 fold higher Fe(II) concentrations and more dissolved CH$_4$ (Figures 1 and S1). Measured total carbon and nitrogen content showed distinct patterns in FCP and HCP cores (Table S1). The total carbon content of the FCP permafrost (31%) was nearly twice as large as the organic layer. The FCP transition layer contained much less carbon than the adjacent layers (20% of FCP permafrost), leading to a low C/N ratio of 16. For the HCP, the total carbon contents of organic and permafrost layers were 21% and 17%, respectively, significantly lower than that of the FCP permafrost. Inorganic carbon quantified as CO$_2$ released upon acid treatment was less than 0.001% for each layer of the FCP and HCP cores.

### 3.2 Temperature responses of CH$_4$ production and oxidation

Thawed FCP and HCP soil samples were incubated in microcosms at fixed temperatures to assess methanogenesis rates. CH$_4$ production was only observed in microcosms from the transition layer and permafrost of FCP, which were incubated under
anoxic conditions. CH₄ production started within 5 days after the anoxic incubations were set up (Figure 3). Cumulative CH₄ concentrations at all temperatures were best fitted with a linear model (Table S2). Soils from the transition layer yielded about 10 times more CH₄ than permafrost at same incubation temperatures. Transition layer soil showed a stronger temperature effect than permafrost. CH₄ production rate increased by 1.6 and 3.1 times as the temperature increased from -2 °C to 4 °C and 8 °C, respectively. Measurements of CH₄ concentrations in the headspace of HCP permafrost were mostly below the quantification limit of the gas chromatograph flame ionization detector (1ppm, Figure S2). Organic soils from both FCP and HCP were incubated with air to best represent the field condition, and methanogenesis was unlikely to occur under oxic conditions. Headspace O₂ was not completely consumed after 90 days incubation (calculations not shown).

Potential rates of aerobic CH₄ oxidation were measured in freshly thawed and pre-incubated soils to minimize total microbial growth limitations. Pre-incubated soils showed much higher CH₄ oxidation potential compared to corresponding freshly thawed soil in both FCP and HCP. In FCP soil, CH₄ oxidation potentials measured in soils from the transition layer and permafrost were significantly higher than those measured in organic soils (Figure 4a, 4b). Similarly, permafrost from HCP showed higher CH₄ oxidation potentials than organic soils (Figure 4c, 4d). Overall, CH₄ oxidation rates in HCP soils were 80-90% lower than rates from the equivalent FCP soil layers.

Rates of both CH₄ production and oxidation responded positively to temperature increase (Figure 5). In the transition layer of FCP, CH₄ production showed much higher temperature sensitivity than CH₄ oxidation from both freshly thawed and pre-incubated soils, with an estimated Q₁₀ value of 4.1. The Q₁₀ value of CH₄ oxidation was 2.0 from freshly thawed soils and only 1.1 in pre-incubated soils. Similarly, the Q₁₀ value for CH₄ oxidation in permafrost also dropped from 1.7 in freshly thawed soils to 1.0 in pre-incubated soils. However, CH₄ production in the permafrost responded slowly to temperature increase, with an estimated Q₁₀ value of 1.7. Overall, permafrost showed significantly lower CH₄ production rates than soils from the upper transition layer, and also had a much lower temperature sensitivity for CH₄ production. However, the measured CH₄ oxidation potentials were similar in both soils, with similar temperature responses.

3.4 Soil respiration in response to rising temperature

Total soil respiration was evaluated using CO₂ production to characterize the observed variation in CH₄ production. Soils in all of the microcosm incubations produced CO₂ by aerobic respiration under oxic conditions or by anaerobic respiration and fermentation under anoxic conditions. CO₂ production started immediately in microcosms of FCP samples, including organic soils that were incubated under oxic conditions and soils from transition layer and permafrost samples that were incubated under anoxic conditions (Figure 6). CO₂ accumulation was best modeled by a hyperbolic function, except the organic layer soil incubated at -2 or +4 °C where the best fit was a linear function (Table S3, Figure 6). This exception is likely due to continuous aerobic respiration at lower temperatures, indicating that substrate limitation was not reached within 90 days. The
transitional layer had the slowest CO₂ production rates and least carbon loss via CO₂, in contrast to its relatively high rate of methanogenesis.

CO₂ production in microcosms of HCP samples was significantly delayed with much lower production compared to FCP soils. The HCP cumulative CO₂ production profiles were best fitted with a sigmoidal model, compared to the hyperbolic model that best fit FCP data (Table S3). A prolonged delay in CO₂ accumulation was observed in both HCP organic and permafrost samples. CO₂ production started about 10 days after the microcosm setup in the organic layer and reached a maximum rate at 30 days (+4 and +8 °C) or 75 days (-2 °C). The delay was longer in permafrost incubations, with a rapid increase after 40 to 50 days followed by a plateau. Therefore, microorganisms mineralized more carbon from FCP soils than HCP soils, and CO₂ production began sooner in FCP than HCP soil incubations.

Temperature showed significant effects on CO₂ production from the organic and transitional layers of FCP during 90-day incubations (p<0.01 for each layer, Figure 6). FCP soils incubated at +8 °C produced substantially more CO₂ than those incubated at lower temperatures. In the permafrost layer of FCP, CO₂ production was significantly higher at +8 °C compared to 2 °C (p<0.05, ANOVA with Tukey’s multiple comparisons test). Anaerobic CO₂ production in FCP permafrost showed much higher temperature sensitivity than that from FCP transition layer (Q₁₀ = 2.3~3.3 and 1.2~1.3, respectively, Table S4).

3.5 Organic acids production and iron reduction in FCP soils

Organic acids were produced as intermediate metabolites during microbial degradation of organic matter, and they probably fueled methanogenesis and iron reduction. Specific organic acids were analyzed from soil extracts from FCP transitional and permafrost layers (Table S5). The dominant organic acids included formate, acetate, propionate, butyrate, and oxalate, consistent with previous analyses from LCP soils (Herndon et al., 2015a). Trace amount of lactate, pyruvate, and succinate were detected, with concentrations less than 0.05 µmol g⁻¹ soil. To compare the changes in organic acids over time, total organic carbon contained in formate, acetate, propionate, butyrate and oxalate products was calculated (T₉₀, µmol C g⁻¹, Figure 7a,b). Concentrations of dominant organic acids measured in the permafrost were approximately 10 times higher than those measured in the transition layer. The difference could be partly explained by much lower SOC content (5.8%) in the transitional layer than that in the permafrost (30.8%). T₉₀ increased by 23%, 70% and 65% at -2, 4 and 8 °C, respectively, in transitional soils during 90-day anoxic incubations. Permafrost initially contained a much higher concentration of organic acids, and T₉₀ increased by a lower percentage, by 1%, 15% and 25% at -2, 4 and 8 °C, respectively.

Significant increases in Fe(II) concentrations were observed from anoxic incubations of FCP samples over the incubation period (Figure 7c,d). In soils from the transition layer, Fe(II) concentrations stayed at a similar level during the first 20 days of anoxic incubation, and then increased significantly from ~50 µmol g⁻¹ to ~100 µmol g⁻¹ during the 20 to 90 days incubation.
period at -2, 4 and 8 °C. In the permafrost, the highest level of iron reduction within the first 20 days was observed in samples incubated at -2 °C. Between 20 and 90 days, the highest iron reduction rates were observed in samples incubated at 8 °C. The estimated Q10 values were 1.2 and 1.3 for transition layer and permafrost, respectively. If we assume that iron reduction is coupled to acetate oxidation to produce CO₂, stoichiometric calculations suggest that iron reduction could account for 96% and 70% of CO₂ produced in the transitional and permafrost layers at -2 °C. At 8 °C iron reduction could account for 74% and 61% of acetate oxidation in the transitional and permafrost layers.

4 Discussion

The widespread ice-wedge degradation in the Arctic causes morphological succession and hydrological changes in tundra ecosystems (Liljedahl et al., 2016). FCP and HCP features represent successively more degraded polygons. Despite clear geomorphological differences between polygon types that affect drainage, vegetation and snow cover, the frozen FCP and HCP organic layers share similar gravimetric water contents, pH, SOC, and Fe(II) concentrations. Permafrost from both polygons contains more water, dissolved CH₄, and Fe(II) than organic layer soils indicating more reducing environments. Concentrations of CH₄ measured in soil pore water from FCP and HCP cores increased with depth. These results are consistent with field measurements of CH₄ dissolved in soil water sampled from the sites during the thaw season (Herndon et al., 2015b). This CH₄ gradient suggests CH₄ oxidation in the upper organic layer. Therefore, we developed the following hypotheses for CH₄ production and oxidation, which were informed by previous studies of methane cycling in temperate ecosystems but untested in the Arctic. (1) CH₄ is produced in the more reduced subsurface, and consumed by methane oxidizers at the upper section of the soil column where O₂ is available. (2) Methane production has higher temperature sensitivity than CH₄ oxidation, and is likely to exceed the CH₄ consumption rate in wet areas in response warmer temperature.

Aerobic CH₄ oxidation is usually assumed to be limited by O₂ and CH₄ diffusion. Therefore, upper layers of soil, the rhizosphere and soil at the water table would be expected to have the highest CH₄ oxidation activities (Shukla et al., 2013; Gulledge et al., 1997). The abundance of the pmoA marker gene for CH₄ oxidation decreased with soil depth in HCP trough soils (Yang et al., 2017) and in permafrost-affected soils from the Canadian Arctic (Frank-Fahle et al., 2014), consistent with that conceptual model. Thus, the organic layers of FCP and HCP in the BEO tundra would have higher potential for CH₄ oxidation. However, the highest CH₄ oxidation potentials were observed in the transitional and permafrost layers of FCP, the only layers with active methanogenesis. This result suggests that most active methanotrophs are found in these deeper soil layers, where CH₄ is available as a C and energy source.

Our results demonstrated CH₄ oxidation might not be primarily O₂ diffusion-limited, but rather limited by the availability of CH₄ in this system. The highest CH₄ oxidation potentials were measured below the rhizosphere, in suboxic layers where CH₄...
has accumulated. Given that half-saturation constants for \( \text{CH}_4 \) and \( \text{O}_2 \) used in methanotrophy models vary over 1-2 orders of magnitude (Segers, 1998; Riley et al., 2011), aerobic \( \text{CH}_4 \) oxidation could occur throughout much of the soil column, as advective, diffusive or plant-mediated transport processes introduce \( \text{O}_2 \) into the soil. Others have observed deep soil \( \text{CH}_4 \) oxidation activity in peatlands (Hornibrook et al., 2009), fens (Cheema et al., 2015) and wet tundra (Barbier et al., 2012), often correlated with water table depth (Sundh et al., 1994).

The water table in the center of Barrow LCPs and FCPs varies somewhat during the thaw season but remains close to the surface (<10 cm below surface for most of the thaw season) (Liljedahl et al., 2015; Liljedahl et al., 2016). Precipitation balances evapotranspiration during the thaw season, with little lateral runoff (Dingman et al., 1980), and volumetric water contents remain constant for these features (http://permafrostwatch.org). In HCPs, the water table drops up to 20 cm below the surface following snowmelt (Liljedahl et al., 2015), and the soils have a lower volumetric water content. Due to limited drainage in the flat coastal plain, the frozen cores analyzed here are representative of field conditions for much of the thaw season. Water isotope analysis demonstrated that most water in the deep active layer comes from summer precipitation rather than seasonal ice melt (Throckmorton et al., 2016). Precipitation during September and October 2011 was above average for Barrow (http://climate.gi.alaska.edu/), suggesting a high water table and limited gas diffusion during the winter freeze-up before we collected soil cores in early 2012. As a result of these annual and transient changes in saturation, methanotrophs could colonize a broad range of the soil column below the rhizosphere. A comparison of methanogenesis and methane oxidation potential in peat bogs demonstrated that methanotrophs survived temporary exposure to anoxic conditions, suggesting these organisms can tolerate rapid changes in the water table and redox potential (Whalen and Reeburgh, 2000). These observations argue against hypotheses that \( \text{CH}_4 \) oxidation occurs primarily at the surface layer or at the water table interface.

The net effect of \( \text{CH}_4 \) production and oxidation in response to temperature change determines the sign of surface-atmosphere \( \text{CH}_4 \) flux. The 4.1-fold increase in \( \text{CH}_4 \) production in the transition layer due to a 10 °C rise in temperature (\( Q_{10} \)) was similar to the average value of 4.26 reported in a recent meta-analysis of permafrost-affected soils (Schädel et al., 2016). These values are substantially higher than the temperature sensitivity of \( \text{CH}_4 \) oxidation from both freshly thawed and pre-incubated samples (\( Q_{10} = 1 \) to 2). While both \( \text{CH}_4 \) production and oxidation respond positively to increased temperature, \( \text{CH}_4 \) production rates are predicted to increase more rapidly with higher temperature at this critical interface between the organic layer and permafrost. This difference in temperature sensitivity of \( \text{CH}_4 \) production and oxidation was also found in Arctic lakes (Lofton et al., 2014). \( \text{CH}_4 \) oxidation potential from freshly thawed FCP soils showed a temperature sensitivity coefficient (\( Q_{10} \)) between 1.7 and 2.0, which is consistent with reported values from peat (Segers, 1998). Higher \( Q_{10} \) values for \( \text{CH}_4 \) oxidation were reported in drier mineral Arctic cryosols with low organic carbon content (Jørgensen et al., 2015; Christiansen et al., 2015). \( Q_{10} \) relationships are strictly empirical representations of a processes’ temperature sensitivity. Microbial community changes, differences in gene expression and enzyme production, and enzyme kinetics are all grouped together in this term, along with
any changes in sorption or transport rates. Current process-enabled models do not distinguish among these factors, but future field experiments and model structures could provide a framework to differentiate mechanisms.

We simulated the net effect of CH₄ production and oxidation based on rate measurements from the transitional layer of FCP that exhibited the highest CH₄ oxidation potential. Methane oxidation rates are 14, 9, and 7 times the methanogenesis rate at -2, 4, and 8°C, respectively. It is quite likely that methanogenesis will outpace CH₄ oxidation under much warmer temperature if there is no change in soil water content. We introduced model representations to explore this possibility, as previous studies suggest CH₄ oxidation rate is strongly regulated by the CH₄ supply (Liikanen et al., 2002; Lofton et al., 2014). Excluding diffusion in this thin layer, we modeled the distribution of the active biomass ratio between methanogens and methanotrophs (B_methanotrophs/B_methanogens), which would cause Arctic soils to act as CH₄ sink or source in response to rising temperature (Figure 8). Given the large difference in the rates of CH₄ production and oxidation, CH₄ oxidation would easily exceed methane production even with an active biomass ratio B_methanotrophs/B_methanogens lower than 1. A much lower biomass ratio B_methanotrophs/B_methanogens (0.07-0.26) would make the soil a net source of CH₄. Studies of functional genes involved in methane production (mcrA) and oxidation (pmoA) in the active layers in the Western Canadian Arctic region suggested substantial variation in the pmoA/mcrA abundance ratio: the range is between 8.5×10⁻⁵ and 7.6×10⁻⁵ (Frank-Fahle et al., 2014). Thus, accurate prediction of soil CH₄ production requires quantification of methanogen and methanotroph populations as model constraints. These results suggest the importance of parameterizing the temperature response function and biomass growth function specifically for methanogenesis and methane oxidation in model simulations to determine if the rates of methanogenesis and methane oxidation offset each other. These results support the hypothesis that methanogenesis can be offset by high methane oxidation rates in degraded tundra, although comprehensive simulations of surface gas fluxes will need to account for gas transport through the full soil column.

The incubations of thawed FCP and HCP soils from all layers revealed substantial differences in the temporal dynamics of CO₂ production. An initial lag was observed from HCP samples at all temperatures, suggesting low initial microbial activity, which might be due to low initial microbial biomass or substrate limitation. In contrast, rapid CO₂ production in FCP soils was observed from both organic layer incubated under oxic conditions and transitional and permafrost layers incubated under anoxic conditions. This temporal pattern of rapid accumulation after thawing was also observed for anaerobic respiration from LCP soils and HCP trough soils (Roy Chowdhury et al., 2015; Yang et al., 2016).

Coupled iron reduction and organic carbon oxidation processes made substantial contributions to total anaerobic CO₂ production relative to fermentation and methanogenesis. Acetate was most abundant intermediate and exhibited the most dynamic concentration changes among individual organic acids measured from the soils. Using reaction stoichiometry for acetoclastic methanogenesis and anaerobic respiration through iron reduction (Istok et al., 2010), we estimated the amount of acetate being consumed by these parallel processes in soils from the transition layer and permafrost (Figure 9). In transition
layer soils, about half of available acetate was consumed by methanogenesis during the first 20 days of incubation (1.34 µmol g\(^{-1}\) out of 2.54 µmol g\(^{-1}\)), while Fe(III) reduction consumed approximately 23% of available acetate (0.60 µmol g\(^{-1}\) out of 2.54 µmol g\(^{-1}\)). In contrast, the partitioning of methanogenesis and Fe(III) reduction inverted during 20 to 90 days of anoxic incubation. Methanogenesis and Fe(III) reduction consumed 31% and 59% of available acetate, respectively. Permafrost contained a higher level of available acetate at the beginning of the incubation, and over 60% of the available acetate was rapidly consumed by Fe(III) reduction during the first 20 days of incubation. From 20 to 90 days of incubation, iron reduction still consumed over half of the depleted acetate in the permafrost. This estimate is consistent with our initial characterization of the FCP core, where the soils from the transition layer contained the highest dissolved CH\(_4\) concentrations, and permafrost was associated with significantly higher Fe(II) concentrations (Figure S1). If hydrogen or other organic anions such as formate or propionate were oxidized by methanogens or iron reducers, then estimated acetate production levels would decrease slightly. These simulations indicate that Fe(III) reduction is responsible for most of the acetate mineralized and CO\(_2\) produced in these soil incubations. As a significant anaerobic respiratory process in Arctic soils, Fe(III) reduction should also be included in CH\(_4\) models to better predict greenhouse gas production in response to a changing climate.

Based on these findings, we propose the following scheme of soil carbon biogeochemistry in the FCP (Figure 10): (1) Increasing temperature facilitates aerobic decomposition of organic carbon in the organic layer, and accelerates anaerobic carbon decomposition in the lower active layer and transition layer through fermentation, iron reduction, and methanogenesis to form CH\(_4\) and CO\(_2\). (2) CH\(_4\) produced in the transitional and permafrost layers is oxidized close to the site of production or transported to the atmosphere. (3) Fe(III) reduction is the primary anaerobic process responsible for the depletion of acetate, the major SOC decomposition intermediate. In this scheme, the oxic/anoxic interface could dynamically move in the soil column with changes in the water table and pore water distribution. Although the transitional layer contained much less carbon than the permafrost layer, the total carbon loss as the sum of CO\(_2\) and CH\(_4\) was comparable to that from permafrost. Laboratory measurements suggested that acetate accumulated in the organic layer could be transported into deeper layers to support iron reduction and methanogenesis (Yang et al., 2016). This transport might occur through vertical movement of dissolved organic compounds or mixing through cryoturbation (Drake et al., 2015). We will use results from these incubation experiments to structure and parameterize a thermodynamically based microbial growth model that improves simulations of anaerobic organic matter decomposition with CO\(_2\) and CH\(_4\) production.

5 Conclusions

Increased warming is predicted to accelerate the transition from wet LCPs to drier FCPs and HCPs in Arctic tundra, which probably function as potential CH\(_4\) sinks. This study demonstrated that CH\(_4\) oxidation capacity was tightly linked to CH\(_4\) availability. Thus, the zone of highest CH\(_4\) oxidation potential is at the suboxic area near the FCP transition layer and the upper
permafrost. The measured CH₄ oxidation potential is an order of magnitude higher than the methanogenesis rate. With higher CH₄ residence time in the soil column due to limited gas diffusion in the field, CH₄ oxidation could easily consume CH₄ produced in deep permafrost soil at warming temperature. Given that iron reduction-coupled respiration predominates anaerobic organic carbon decomposition, CO₂ is likely to remain the major form of carbon emission from degraded polygons.

This finding provides critical information about the dynamics of CH₄ production and oxidation with increasing temperature that need to be incorporated into Arctic terrestrial ecosystem models for better predictions.

Data availability

The dataset can be found in (Zheng et al., 2017).

Author contributions

DG, SW, and BG conceived and organized the research study; TRC, DG and SW collected core samples; IZ, TRC, ZY and performed experiments and acquired data; IZ, TRC and DG analyzed and interpreted data; JZ and DG drafted the manuscript. All authors contributed revisions to the manuscript and have given approval to the final version of the manuscript.

Competing interests

The authors declare no competing interests.

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Soil incubation and methane oxidation assay experimental design. Soils from the transitional layer of a FCP core were used here to illustrate how incubations and methane oxidation assays (MOAs) were constructed at a given temperature. Incubation replicates were destructively sampled after 10 or 20 days to set up MOA with pre-incubated soils. For example, 9 replicate microcosms were created for the FCP transition layer soil, incubated under anoxic conditions. CO₂ and CH₄ in the headspace were cumulatively measured. After 10 days, 3 were opened and subsampled for triplicate MOAs. After 20 days, another 3 were opened and subsampled for triplicate MOAs.
Figure 2. Depth profile of gravimetric water content, bulk density, Fe(II) concentration, pH and soil pore water dissolved CO$_2$ and CH$_4$ concentrations from FCP (blue) and HCP (red) cores. Replicate measurements were plotted as high and low values in each soil section as blue filled and empty circles (FCP) and red triangles (HCP). Trend lines were plotted based on the average values.
Figure 3. CH$_4$ production in soil microcosms from (a) transition layer, and (b) permafrost layers of FCP at indicated temperatures. The temporal profiles of CH$_4$ production were best fitted with the linear regression model, and the shaded area represents 95% confidence interval.
Figure 4. CH$_4$ oxidation potential measured from soils incubated at the indicated temperatures from (a) FCP after 0 days, (b) FCP after 20 days, (c) HCP after 5 days, and (d) HCP after 10 days. Error bars indicate ±1 standard deviation from three replicate incubations.
Figure 5. Temperature sensitivity of CH₄ production and oxidation measured from the transition and permafrost layers of FCP. Temperature responses of CH₄ oxidation rates from freshly thawed (O_transition and O₂0_transition) and pre-incubated (O20_transition and O20_permafrost) soils were both estimated.
Figure 6. Cumulative CO$_2$ production in soil microcosms from FCP and HCP samples at indicated temperatures.
Figure 7. Changes in total organic acids carbon (Top panels) and Fe(II) concentrations (Bottom panels) in soils from transition layer and permafrost of FCP during anoxic incubations. Total organic acids (µmol C g⁻¹) were calculated from the concentrations of individual organic acids (Table S5). Error bars for Fe(II) concentrations are ±1 standard deviation from three replicate incubations.
Figure 8. Simulations of active biomass ratio $\frac{B_{methanotrophs}}{B_{methanogens}}$ distribution for Arctic soils to act as CH$_4$ sink or source in response to rising temperature. Dissolved CH$_4$ and O$_2$ concentrations of 0.1 mM in soil pore water are assumed. Half saturation constants ($K_{m,CH_4}$ and $K_{m,O_2}$) represent the baseline value (Grey), high (Blue) and low(Red) range for sensitivity analysis. Biomass ratios below the curves indicate the system could be a CH$_4$ source, while biomass ratios above the curves suggest a CH$_4$ sink.
Figure 9. Changes in acetate concentrations associated with production (white bars) and consumption by iron reducing bacteria (red bars) or methanogens (blue bars) were estimated using stoichiometric calculations based on measurements of methane and Fe(II) during incubations from 0 to 20 days and from 20 to 90 days at the indicated incubation temperatures.
Figure 10. Conceptual model of aerobic and anaerobic soil organic carbon decomposition pathways and the release of CO$_2$ and CH$_4$ from a flat-centered polygon. Lines on the left marked as Fe(II), CO$_2$, and CH$_4$ represent measurements across the soil column.