

Year-round simulated methane emissions from a permafrost ecosystem in Northeast Siberia

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Abstract. Wetlands of northern high latitudes are ecosystems highly vulnerable to climate change. Some degradation effects include soil hydrologic changes due to permafrost thaw, formation of deeper active layers, and rising topsoil temperatures that accelerate the degradation of permafrost carbon and increase in CO₂ and CH₄ emissions. In this work we present 2 years of modeled year-round CH₄ emissions into the atmosphere from a Northeast Siberian region in the Russian Far East. We use a revisited version of the process-based JSBACH-methane model that includes four CH₄ transport pathways: plant-mediated transport, ebullition and molecular diffusion in the presence or absence of snow. The gas is emitted through wetlands represented by grid cell inundated areas simulated with a TOPMODEL approach. The magnitude of the summertime modeled CH₄ emissions is comparable to ground-based CH₄ fluxes measured with the eddy covariance technique and flux chambers in the same area of study, whereas wintertime modeled values are underestimated by 1 order of magnitude. In an annual balance, the most important mechanism for transport of methane into the atmosphere is through plants (61%). This is followed by ebullition ($\sim 35\%$), while summertime molecular diffusion is negligible (0.02%) compared to the diffusion through the snow during winter ($\sim 4\%$). We investigate the relationship between temporal changes in the CH₄ fluxes, soil temperature, and soil moisture content. Our results highlight the heterogeneity in CH₄ emissions at landscape scale and suggest that further improvements to the representation of largescale hydrological conditions in the model will facilitate a more process-oriented land surface scheme and better simulate CH₄ emissions under climate change. This is especially necessary at regional scales in Arctic ecosystems influenced by permafrost thaw.

1 Introduction

During the last 30 years, atmospheric temperatures at northern high latitudes have risen more than the global average (Schuur et al., 2015; Serreze et al., 2000). In consequence, many permafrost areas in these regions have experienced expedited thawing rates in recent years. Permafrost in northern high-latitude ecosystems contains twice as much carbon as the current carbon pool in the atmosphere and about half of the global soil organic carbon (Hugelius et al., 2014; Tarnocai et al., 2009). About two-thirds of the terrestrial Arctic is classified as wetlands (Liljedahl et al., 2016; Hugelius et al., 2014) and permafrost underlies most of them. Wetlands globally contribute about 25 % of the total CH₄ emissions (using bottom–up approaches between 2003 and 2012) from natural sources into the atmosphere. Nearly 4% of the total global emissions from top-down inversions correspond to emissions at latitudes $> 60^{\circ}$ N (Saunois and al., 2016). The degradation of freshly available carbon from permafrost thaw is expected to contribute strongly to a positive carbon-climate feedback in Arctic ecosystems (e.g., Beer, 2008). Changes in air temperature, surface topography, and projected shifts in precipitation in Arctic tundra ecosystems (Kattsov and Walsh, 2000; Lawrence et al., 2015) influence the soil hydrologic regime in permafrost areas. Also, thawing permafrost will induce changes in the surface wetness due to surface subsidence of ice-rich soils (Christensen et al., 2004; Helbig et al., 2017b). These changes will therefore also influence the magnitude of future emissions of CO2 and CH₄ into the atmosphere from Arctic terrestrial ecosystems (Hugelius et al., 2014; Lawrence et al., 2015; Schuur et al., 2008). Drier soil columns will enhance methane oxidation and increase CO₂ emissions (Kittler et al., 2016; Kwon et al., 2016; Lawrence et al., 2015; Liljedahl et al., 2016; Sturtevant et al., 2012), also leading to changes in plant community structure (Christensen et al., 2004; Kutzbach et al., 2004; Kwon et al., 2016). Thus, it is imperative to improve our understanding of the effects of climate change in permafrost wetlands, specifically their contribution to greenhouse gases in the atmosphere.

Freeze and thaw soil processes are critical mechanisms that modulate the seasonality of CH₄ emissions in permafrost ecosystems of the Arctic (Panikov and Dedysh, 2000). Most of the annual CH₄ emissions from Arctic wetlands take place during summer (growing season). In spring, the rising air and soil temperatures promote the melt of snow and ice in the soil, stimulating the microbial production of gas within the mostly anoxic active layer (i.e., the surface soil layer that thaws during summer and freezes again during autumn). During this season, episodic releases of large amounts of CH4 in the form of bursts have been evidenced in wetlands (e.g., Friborg et al., 1997; Song et al., 2012), peatlands (e.g., Tokida et al., 2007), and lakes (e.g., Jammet et al., 2015) of northern high latitudes. During late autumn, CH₄ emissions still take place when the active layer starts to freeze gradually from the top and ice begins to fill the soil pore spaces, i.e., the socalled zero curtain period. Through this period, the remaining CH₄ in the soil that was produced during the growing season or in the deeper warm soil layers is squeezed out of the soil. This remaining gas is emitted to the atmosphere via molecular diffusion, and via the "pressure pumping" phenomenon due to advection enhanced by wind (Bowling and Massman, 2011; Massman et al., 1997), through the forming layer of snow (Mastepanov et al., 2008, 2013; Zona et al., 2016). Previous studies have reported that CH₄ emissions during the cold season in Arctic tundra ecosystems account for up to 50 % of the total annual CH₄ flux released in the form of gas bursts (Mastepanov et al., 2008; Zona et al., 2016).

Soil and vegetation at northern high latitudes remain covered by snow during most of the year (October to May). Snow is an effective thermal insulator between the soil and the atmosphere, and it is a porous medium that allows the diffusive exchange of gases. Only a few observational efforts have previously been made to constrain gas fluxes through the snow in tundra and permafrost environments during the long and cold Arctic winter. CH_4 emissions have been measured using flux chambers and eddy covariance (EC) towers in various snow-covered areas, e.g., in boreal forest soils (Kim et al., 2007; Whalen and Reeburgh, 1992), boreal peat landscape bogs and fens (Helbig et al., 2017b; Panikov and Dedysh, 2000; Rinne et al., 2007; Smagin and Shnyrev, 2015), and subalpine soils (Mast et al., 1998; Wickland et al., 1999). Also, methane flux measurements are available in the Alaskan tundra (Zona et al., 2016) and in the Zackenberg valley in northern Greenland (Mastepanov et al., 2008; Pirk et al., 2016). In boreal peat bogs of West Siberia, cold season CH₄ emissions contribute 3.5 to 11 % of the annual CH₄ fluxes (Panikov and Dedysh, 2000). In other Arctic permafrost tundra ecosystems, however, winter CH₄ emissions were 1 to 2 orders of magnitude lower than the emissions during summer, and only accumulate in the snowpack in the presence of layers of ice blocking their exit route to the atmosphere (Pirk et al., 2016). Wickland et al. (1999) concluded that in snow-covered subalpine wetland soils, CH4 fluxes accounted for 25% of the annual fluxes, similarly to the recent results shown in a boreal peat landscape of northwestern Canada (Helbig et al., 2017b). However, there are still large uncertainties in cold season CH₄ emissions from wetlands and permafrost ecosystems of the Arctic tundra, particularly related to projected changes in vegetation phenology due to climate warming which might also lead to changes in snow cover; e.g., more shrubs will tend to hold more snow during winter (Blanc-Betes et al., 2016; Domine et al., 2015). A thicker snow layer will insulate the soil column more during autumn and winter, preserving the heat of the active layer after the preceding growing (zero curtain period) season. This will further impact the extent of subsequent wintertime CH₄ productions and emissions.

Numerical models have made much progress in better simulating the magnitude and temporal and spatial variability of CH₄ emissions in boreal regions. Methane models include the traditional theoretical and empirical approaches that describe the mechanistic understanding of the processes involved in the production, oxidation, and transport of CH₄ in terrestrial ecosystems (e.g., Grant, 1998; Riley et al., 2011; Walter and Heimann, 2000). Previous studies have also improved the scaling representation from plot to regional areas in specific locations, and also to global frameworks (Bohn et al., 2015; Lawrence et al., 2015; Melton et al., 2013; e.g., Riley et al., 2011; Ringeval et al., 2011; Tagesson et al., 2013; Wania et al., 2010). However, there are still many shortcomings in land surface models for boreal regions because a complex network of processes and a wide range of spatial and seasonal variation characterize these areas. These are particularly related to the inability to account for methane emissions and uptake in dry non-wetland areas, capturing shifts in vegetation cover (e.g., Chen et al., 2015), representing soil thawing and freezing cycles due to a poor soil thermal physics representation (e.g., Schuldt et al., 2013; Zhu et al., 2014), varying wetland extents and water tables (Bohn et al., 2015), accounting for microtopography effects on surface water and methane emissions dynamics (Cresto Aleina et al., 2013, 2016), upscaling to larger areas, or coupling to Earth system models (e.g., Mi et al., 2014; Xu et al., 2015). Also, the majority of models lack a snow scheme that interacts with gas transport and a descriptive representation of peat soils (Xu et al., 2016). Special challenges exist for regional-scale model simulations in wetland-dominated areas influenced by climate change, such as in Arctic permafrost ecosystems, mostly because of the lack of observational constraints sufficient to understand the processes in these areas and to evaluate model outputs.

The aim of this work is to analyze the performance of an improved process-based methane model, designed for Arctic tundra and wetlands underlain by permafrost, when applied to a regional domain in Northeast Siberia. Our intention is to evaluate the potential of a refined process-based methane model as a proof of concept, for its application to scales larger than site level. Also, a regional-scale application will allow the identification of spatial heterogeneities in CH₄ emissions in boreal regions. To address these objectives, we simulate year-round CH₄ emissions during 2014 and 2015 with the process-based JSBACH-methane model (Kaiser et al., 2017) in a region dominated by low-lying wetland areas and continuous permafrost in the Russian Far East. This model includes freeze and thaw soil cycles associated with explicit methane production, oxidation and transport. The latter takes place through distinct pathways: plant-mediated, ebullition, and diffusion. In this work, we use an improved version of the model that explicitly simulates CH₄ emissions into the atmosphere in the presence of snow during the nongrowing season, and also contains a revised representation of CH₄ transported by plants including the description of relevant features of vascular plants based on the volume of roots in the soil pore space. We present and analyze the year-round temporal variation of the CH₄ emissions and their relationship with the environmental controls at a regional (model domain) scale. The model performance was assessed by comparison of the simulated CH₄ emissions against year-round EC measurements and summertime chamber flux measurements in the same study area. Because temporal variation in the amount of inundated areas is essential for the estimation of CH₄ emissions from wetlands (Prigent et al., 2007), our model also includes a representation of inundated areas using a TOPMODEL approach. We evaluate the modeled horizontal extent of the inundated areas against the wetland area from a high-resolution remote sensing product.

2 Methods

2.1 Site description

The target region of this study is located in Northeast Siberia, Sakha Republic. The model domain is centered on the town of Chersky and to the west is dominated by low-lying wetland areas of the Kolyma River floodplain and to the east by dry upland tundra (Fig. 1a). This is a region of continuous permafrost and active layer depths that range between 20 and 180 cm. Winter spans from October to May, with daily air temperatures that remain well below the freezing point and average daily temperatures of about 13 °C during July (Dutta et al., 2006). In this region prevail dry climate conditions, with a mean annual precipitation of 218 mm (60% as snow and 40% as rain; Dutta et al., 2006). At the Kolyma River floodplain, the soil profile has a top layer of organic material (\sim 15–25 cm thick) that is located above alluvial mineral soils, i.e., silty clay (Kittler et al., 2016; Kwon et al., 2016). In this area, the vegetation is heterogeneous and representative of wet tussock tundra. There, the water-logged areas are covered by the tussock-forming sedges (Carex appendiculata and Carex lugens) and cotton grasses (Eriophorium angunstifolium) (Kwon et al., 2016). During the spring snowmelt (May and June), large sections of the Kolyma floodplain usually become inundated, and during summer, the extent of surface water recedes due to evapotranspiration and drainage to the river channels located nearby. However, most areas remain inundated throughout the year (Kwon et al., 2016) and microtopographic structures typical of polygonal tundra landscapes are sparse in this region. The eastern part of the model domain has more elevated slopes and drier soils with tundra vegetation dominated by grasslands and forests, i.e., dwarf evergreen and deciduous shrubs, Sphagnum mosses, lichens, and few trees (Dutta et al., 2006; Merbold et al., 2009). Loess soil deposits originating from the accumulation of aeolian and alluvial sediments characterize the soil in this region.

2.2 Model configuration

The model results presented in this work were obtained with a regional configuration in offline mode of the land surface component of the MPI-ESM (Max Planck Institute for Meteorology Earth System Model), the so-called Jena Scheme for Biosphere Atmosphere Coupling in Hamburg (JSBACH) model. We used a JSBACH version that has been extended from the version of the CMIP5 activity (e.g., Brovkin et al., 2013; Raddatz et al., 2007; Reick et al., 2013). Modifications include the addition of a multilayer hydrology scheme (Hagemann and Stacke, 2015) and the representation of permafrost physical processes (Ekici et al., 2014). The model domain covers an area of 7° in longitude (158 to 165° E) and 3° in latitude (66.5 to 69.5° N). Using a horizontal resolution of 0.5° (Fig. 1b), this results in a model domain with 14×6 equally spaced grid cells. The vertical structure in the model domain comprises 11 non-equidistant soil layers with thicknesses that increase from 6.5 cm at the top to 23.2 m at the bottom, reaching a maximum column depth of 40.5 m. This vertical refinement is necessary to achieve numerically stable solutions for the gas diffusion equation. In the model domain, the root zone is confined to the top five layers (max-



Figure 1. (a) Image from Copernicus Sentinel data (2016) showing the heterogeneous landscape in most of the model domain in Northeast Siberia, also showing the location of nearby cities and the floodplain; (b) geographical location of the model domain used in this study also depicted with the midpoints of the model grid cells (pink circles) and boundaries (dashed lines) underlain by a geoTIFF image (data from 1 to 10 July 2007) from the EAWS product. The boundaries of grid cells A and B are delimited with pink lines. The continuous black line delimits the 35 model grid cells used for the evaluation of modeled inundated areas against the EAWS product.

imum depth of 1.1 m) with maximum and mean root depths of 0.88 and 0.42 m, respectively. The soil ice content is restricted to the top six layers (maximum depth of 2.0 m), with the bedrock located from the sixth layer downwards.

In JSBACH, each grid cell has defined fractions for different types of vegetation that are assigned across a maximum of 11 non-equal tiles, hence a hospitable fraction to vegetation that represents the sub-grid-scale heterogeneity of vegetation cover. The remaining fraction of the grid cell where vegetation is not assigned is then associated with a land cover type that represents areas inhospitable to vegetation such as rocky surfaces and deserts (Reick et al., 2013).

In our model domain, only four land cover types were present (ordered by dominance in the model domain): (1) C3 grasses, (2) deciduous trees, (3) evergreen trees, and (4) deciduous shrubs (see Fig. S1 in the Supplement for the spatial distribution of the cover types in the model domain).

The model configuration contains the basic JSBACH modules with components from the Biosphere-Energy-Transfer-Hydrology model, BETHY (Knorr, 2000). The vegetation carbon is categorized into three groups: wood, green, and reserve. The Yasso07 soil carbon and decomposition model (Tuomi et al., 2009, 2011) takes care of the transport and decomposition of carbon into the soil. It simulates the breakdown of litter and soil organic matter based on measurements of soil carbon and litterbag experiments, and has been previously implemented in JSBACH (Goll et al., 2015; Thum et al., 2011). In Yasso07, soil litter is divided into three classes: non-woody, woody, and humus. The non-woody class is subdivided into four pools representing groups of chemical compounds with an independent decomposition rate determined by changes in air temperature and precipitation; thus, it has no relation to plant species (Goll et al., 2015; Tuomi et al., 2009).

Most of the CH₄ emissions into the atmosphere from Arctic terrestrial ecosystems are from wetland areas; thus, the representation of the wetland extent in CH₄ models is of relevance. We use a TOPMODEL (TOPographic MODEL) approach (Beven and Kirkby, 1979; Kleinen et al., 2012; Stocker et al., 2014) to determine the fraction of any grid cell that is inundated, implying it has a water table at or above the soil surface. We obtain a grid cell mean water table position from the soil hydrology scheme (Hagemann and Stacke, 2015) determined from the saturation state of the soil layers: the lowest soil layer that is not completely frozen or completely saturated contains the grid cell mean water table, with the exact location within the layer given as the layer fraction that is saturated. Details on the TOPMODEL scheme in JS-BACH are shown in Sect. 1 of the Supplement.

The position of the local water table depth z_i is used to define the grid cell wetland area (Eq. S1 in the Supplement); i.e., the grid cell wetland area is defined where $z_i \ge 0$, and it

is subject to a minimum CTI (compound topographic index) threshold value χ_{min_cti} that limits the maximum possible areas that can be flooded following the approach of Stocker et al. (2014), with lower values leading to larger wetland areas. In this configuration, the constant prescribed value of χ_{min_cti} and the exponential decay of transmissivity with depth *f* (Eq. S1) are tunable parameters of the TOPMODEL module used to expand or reduce the fraction of inundated surface areas in a model grid cell. Within the inundated fraction of 0.95 is assumed. The decomposition of soil organic matter is reduced to 35 % of the aerobic decomposition in line with Wania et al. (2010).

The CH₄ production and emission processes in the model are tightly linked to the volumetric soil porosity to allocate gas transport (Kaiser et al., 2017). This model configuration contains a permafrost module to explicitly simulate soil freeze and thaw processes coupled to the hydrological and thermal regimes in the soil column (Ekici et al., 2014). This is a relevant process in permafrost regions where changes in the soil ice content drive the seasonal changes in the volumetric soil pore space, and changes in the soil moisture ultimately determine whether the soil pores are filled with water or air.

2.2.1 Methane module

In this work, the JSBACH-methane configuration presented in Kaiser et al. (2017) underwent several modifications. Besides being coupled to TOPMODEL and the soil carbon Yasso07 components, the CH₄ module itself acquired several extensions: (i) a refined description of plant-mediated transport, (ii) allowance of gas transport via diffusion through the snow during the non-growing season, and (iii) change in the order of transport processes. Details on each of these changes are listed in Sect. 2 of the Supplement.

In the process-based JSBACH-methane module, the equilibrium between the concentrations in free atmosphere, soil air, and soil moisture is assumed for the initialization of the methane and oxygen concentrations in the soil. During each model time step, CH₄ is produced in the soil column depending on the soil hydrological conditions (i.e., ice content and soil moisture), soil temperatures, soil pore space, and the available decomposed carbon. The fraction of CH₄ produced from the total carbon decomposition under anaerobic conditions for mineral soils ($f_{CH4anox}$) is prescribed as 0.5 (i.e., 50 % of the anaerobically decomposed carbon is used to produce CH₄). Since this setting is highly uncertain, the model response to a range of $f_{CH4anox}$ values is tested in sensitivity experiments as part of this work.

The JSBACH-methane module contains two explicitly modeled CH_4 oxidation processes: bulk soil oxidation and rhizospheric oxidation of methane (plant oxidation). These oxidation pathways interact iteratively in the model with the methane transport processes, reducing the methane pool when oxidation takes place. Only part of the oxygen in the soil is available for methane oxidation, and this discrimination relates to the amount of carbon dioxide produced during heterotrophic respiration, which uses up to a maximum value of 40% of the total oxygen content in the soil. An additional 10% of the available oxygen is assumed to be unavailable because it is used in other processes such as respiration by microbes. This leads to only 50% of the total oxygen in the soil being available for CH₄ oxidation.

To facilitate the interaction between the CH₄ and TOP-MODEL modules, the ice-free pores of the soil column are prescribed at a saturation level of 95% in the fraction of the grid cell that was determined as inundated. This concept mimics the lateral distribution of water that creates water-logged conditions, depending on the topographic profile. However, the soil temperatures, ice content, and available carbon for CH₄ production are not changed in the model during this process. Thus, CH₄ emissions from a grid cell happen under a combination of soil temperatures, ice content, and available carbon decomposition characteristic of an unsaturated soil column on the one hand and ice-free soil pores with soil moisture at 95 % saturation on the other. Ultimately, the methane production, oxidation, and transport processes only take place in the saturated portion of the grid cell (Fig. S2). The transport of the gases to and from the atmosphere is distributed across four explicitly modeled transport processes: plant-mediated transport, ebullition, and molecular diffusion without snow and through the snow. The transport pathways follow a sequential order based on the expected priority, with their efficiency based upon prevailing soil moisture content (set to constant 95% saturation in the inundated areas) taking into account the ice-corrected volumetric soil porosity, which in turn depends on the soil temperature.

The *plant-mediated transport* in the model only takes place in areas with C3 grasses and follows Fick's first law, including the diffusion of gas between the roots of plants and the surrounding soil pores. In wetland ecosystems, many plants have developed an efficient aerenchyma system that functions as a transport mechanism of gases between the atmosphere and their roots. Plants need oxygen for metabolic processes and the root exodermis is an efficient barrier that keeps the oxygen inside the plant roots and, at the same time, slows down the diffusion of gas from the soil into the roots; thus, the gas flow is restricted by the thickness of the exodermis tissue. In the JSBACH-methane module, the root exodermis has a prescribed diffusivity value of 80 % of the total diffusivity of the same gas in water, for the gas transport from soil into the plant. *Ebullition* takes place when excess gas that has not been dissolved in the available soil pore liquid water forms bubbles that are rapidly transported upwards from their source in the deep soil layers through the water and into the atmosphere, successfully bypassing the oxic areas in the soil. Diffusion is the molecular transfer of gas from high to low concentration gradients between soil layers and

the atmosphere following Fick's second law. In this model version, diffusion is now also allowed to take place through a layer of snow using a simplified formulation that does not take into account the enhanced advection of gas in the snow-pack due to wind, i.e., pressure pumping.

Between the model time steps, the amount of gas is constant, whereas the gas concentrations change in relation to the varying ice-free pore space. Further details on how these schemes are included in the model are shown in the supplementary material, and for more details the reader is also referred to Kaiser et al. (2017).

2.2.2 Experimental setup and sensitivity experiments

The model was forced with the daily reanalysis atmospheric data from CRUNCEPv7 (the Climate Research Unit from the University of East Anglia, analysis of the National Centers for Environmental Prediction reanalysis atmospheric forcing version 7.0) from 1901 to 2015 with a spatial resolution of 0.5° (Viovy and Ciais, 2016). Prescribed annual means of atmospheric CO₂ values (https://www.esrl.noaa.gov/gmd/ ccgg/trends/global.html, las access: March 2018) were also used to drive the model. The model was spun up for 10000 years of simulation by repeating cycles of atmospheric data from 1901 to 1930 (\sim 330 cycles) to equilibrate the soil carbon pools and ensure pre-industrial steady state (Chadburn et al., 2017; McGuire et al., 2016). The total carbon (woody, green, and reserve) after spinup in the entire model domain showed little change over the last 500 years of the spinup period. The methane module was de-activated during this procedure. After that, simulations were initialized with reanalysis data from 1931 until 2015 (85 years). To allow equilibration of the soil carbon pools to the hydrology as well as equilibration of CH₄, a model adjustment period of 850 years (10 cycles using the 85 years of reanalysis data) was added. After this period, the subsequent output of the model was stored and used for data analysis. In this simulation, we used prescribed reference values for parameters in the TOPMODEL and methane modules that represent the control simulation. A description of the most relevant prescribed parameters and variables in the control simulation is outlined in Table 1.

To evaluate the robustness of the model and identify the parameters to which the model is most sensitive, a set of sensitivity experiments was done following a cost efficient parameter-permutation approach (Saltelli et al., 2000). Six model parameters that are prescribed in the model and are involved in the newly modified code for this model version were selected. These parameters are either not provided in the published literature, the published values are largely uncertain due to the nature of the method used to obtain these values, or the measured values cover a wide range of options characterizing different conditions in nature. The selected parameters are χ_{min_cti} for the evaluation of TOPMODEL, d_r and R_{fr} for the evaluation of plant-mediated transport, h_{snow}

and ϕ for evaluation of the transport via diffusion through the snow, and the fraction of anoxic decomposed carbon that becomes CH₄ ($f_{CH4anox}$) for the evaluation of the methane production. For each parameter, reference values from the control simulation were decreased or increased for one parameter at a time, by a fixed value (shown together with the results in Table 2), resulting in a total of 12 independent sensitivity simulations.

The values for the parameters $\chi_{\min \text{ cti}}$, d_{r} and R_{fr} and h_{snow} are highly uncertain. The first one is a parameter that is part of the TOPMODEL parameterization, whereas the rest are highly uncertain or absent in the published literature; therefore, we decided to choose extreme values with respect to their values in the control simulation. The selected values for ϕ and $f_{CH4anox}$ were kept within ranges reported in the literature. The snow porosity is derived from measurements of snow and ice, and ultimately controls the amount of gas that can diffuse through the snow layer. Different snow densities lead to different snow porosities: 330 kg m^{-3} ($\phi = 0.64$) for wind packed snow, 263 kg m^{-3} ($\phi = 0.71$) for settled snow and 128 kg m^{-3} ($\phi = 0.86$) for fresh damp new snow. These values were tested to reflect the effect of gas diffusion through less to more porous snow layers. All ϕ values were calculated with $\rho_{ice} = 910 \text{ kg m}^{-3}$.

The parameter $f_{CH4anox}$ is highly uncertain in the literature. In our model, a setting of $f_{CH4anox} = 1.0$ would imply that all of the decomposed soil carbon would become CH₄ under anaerobic conditions. The value used in the reference simulation is 0.5. In the context of the sensitivity experiments, we decrease $f_{CH4anox}$ to 0.1 (i.e., 10% of the decomposed carbon will become CH₄ and 90% will be oxidized), and to 0.3 (i.e., 30% of the decomposed carbon will become CH₄ and 70% will be oxidized).

Each sensitivity simulation consisted of a re-initialization from the conditions in the control simulation from the last time step on 31 December 1999. This was to allow the model to adjust to the parameter change for 13 years before the year of result analysis (i.e., 2014). In order to keep consistency in the treatment of our simulations, the same reinitialization procedure was done for a reference simulation by re-initializing the control simulation from the restart conditions on 31 December 1999, as in the sensitivity experiments, but without changing any parameter (i.e., maintaining the same parameters as in the control simulation). The results from the sensitivity experiments were compared to the results from the reference simulation. The temporal resolution of all the model simulations is 30 min, with hourly output averaged for analysis into daily and monthly values.

2.3 Observational data

2.3.1 Wetland product

Methane emissions to the atmosphere in the model occur largely from areas with a water table at or above the sur-

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Parameter	Description	Value	Unit
Xmin cti	Threshold to define maximum areas that can be flooded in a grid cell (TOPMODEL)	12	_
f = 1	Exponential decay of transmissivity with depth (TOPMODEL)	2	_
$d_{ m r}$	Root diameter	2	mm
r	Resistance factor of root exodermis	0.8	_
hexo	Thickness of exodermis	0.06	mm
$R_{\rm fr}$	Principal fraction of the pore-free soil volume occupied by roots	40	%
ϕ	Porosity of snow	0.64	_
$h_{\rm snow}$	Snow depth threshold	5	cm
f _{CH4anox}	Fraction of anoxic decomposed carbon that becomes CH ₄	0.5	-
$D_{\rm air}^{\rm CH_4}$	Diffusion coefficient of CH ₄ in free air at 0 °C and 1 atm	$1.95 imes 10^{-5}$	${\rm m}^2{\rm s}^{-1}$
$D_{\rm air}^{\rm O_2}$	Diffusion coefficient of O_2 in free air at 0 °C and 1 atm	1.82×10^{-5}	${\rm m}^2{\rm s}^{-1}$
$\rho_{\rm ice}$	Ice density	910	$\mathrm{kg}\mathrm{m}^{-3}$
ρ_{snow}	Snow density (together with ρ_{ice} leads to $\phi = 0.64$ and $\tau = 0.77$)	330	$\mathrm{kg}\mathrm{m}^{-3}$

Table 1. Summary of the most relevant prescribed parameters in the JSBACH-methane control and reference simulations.

Table 2. Results from sensitivity experiments (the specific descriptions of the parameters listed below are given in Table 1). Statistical *p*-values are given for the experiments whose results significantly differ from the results in the reference simulation.

Variable	Value	Unit	Annual mean of total CH ₄ / (mg CH ₄ m ⁻² d ⁻¹)
Xmin_cti	11 12 ¹ 13	_	$\begin{array}{c} 4.2\pm 5.0^2\\ 6.2\pm 7.3\\ 9.2\pm 10.7^2\end{array}$
$d_{ m r}$	2 ¹ 5 8	mm	$\begin{array}{c} 6.2 \pm 7.3 \\ 6.2 \pm 7.3 \\ 6.2 \pm 7.3 \end{array}$
R _{fr}	$0.2 \\ 0.4^1 \\ 0.6$	_	6.2 ± 7.3 6.2 ± 7.3 6.2 ± 7.3
φ	0.64^{1} 0.71 0.86	_	$6.2 \pm 7.3 \\ 6.2 \pm 7.3 \\ 6.2 \pm 7.3$
h _{snow}		cm	6.2 ± 7.3 6.2 ± 7.3 6.2 ± 7.3
f _{CH4anox}	$0.1 \\ 0.3 \\ 0.5^1$	-	$\begin{array}{c} 1.2 \pm 1.4^2 \\ 3.7 \pm 4.3^2 \\ 6.2 \pm 7.3 \end{array}$

¹ Parameter value in reference simulation; ² significant at p < 0.001.

face. These fractions of "inundated" areas in each model grid cell represent the horizontal extent of wetlands (including lakes, peatlands, or temporally inundated areas). As described in Sect. 2.2, in our study the inundated fraction for each grid cell is estimated through the TOPMODEL approach. For evaluation, we compared the spatial distribution of the inundated areas per grid cell to the wetland extent remote sensing product from ENVISAT ASAR (the European Space Agency's ENVISAT with an Advanced Synthetic Aperture Radar operating in Wide Swath mode C-band). The ENVISAT ASAR WS-wetland product (EAWS) was tested for operational monitoring in northern Russia, where smallscale ponds and an overall high soil moisture level are common surface features (Reschke et al., 2012). The backscatter of the EAWS product for high latitudes has a higher spatial and temporal resolution (150 m and 2 to 3 days, respectively) than other commonly used wetland products (e.g., Prigent et al., 2007), which have spatial resolutions on the order of kilometers. Thus, the EAWS product is able to capture small water bodies like tundra ponds and wetland patches that remain almost unchanged throughout the year and are associated with permafrost areas. The spatial coverage of the EAWS product includes most of northern Russia and is subdivided into 10 mosaics, each with different coverage areas. It is freely available as GeoTIFF images, each representing a 10-day mean in a wetland map during July and August in 2007 (i.e., 1-10, 11-20, 21-31 July, 1-10, 11-20, and 21-31 August, all in 2007; Reschke et al., 2012).

For the evaluation of the modeled wetland extent, each 10-day-mean image of the EAWS product was mapped to the same grid of georeferenced rectangular cells of the JS-BACH domain. From the total 84 model grid cells, 35 model grid cells fall into the area coverage of the EAWS images (Fig. 1b). The wetland fraction from the EAWS remote sensing product (w_{rs}) in percentage was calculated as the ratio of pixels flagged as wetland (ID = 1) to the total number of pixels contained in the grid cell. In the model, the spatial wetland fraction (w_{mod}) is represented as the fraction of the total grid cell area that is inundated (i.e., with a water table at or above the soil surface). To facilitate a direct comparison against w_{rs} , the w_{mod} values from the control simulation

were averaged to the same 10-day mean in 2007 as the remote sensing data.

2.3.2 Chamber measurements

To evaluate the performance of the methane model, we compared the total modeled methane fluxes (F_{mod}) to the total methane fluxes measured with flux chambers (F_{ch}) in the Kolyma River floodplain (Fig. 1b; see also Kwon et al., 2016). In this study, chamber fluxes from an undisturbed control area were considered for model evaluation purposes. The chamber flux measurements were done during the early to mid-growing season (15 June to 20 August) in 2014. As additional ancillary variables, water table depth, vegetation cover, and soil temperature were also measured. For further details on the gas measurements, calculations, and discussion of the chamber flux results, the reader is referred to Kwon et al. (2016).

The surface area of each chamber along the control transect is 0.36 m²; therefore, even all 10 chambers combined can only represent a very small fraction of the surface area of a single model grid cell $(2.5 \times 10^9 \text{ m}^2)$. However, since both F_{mod} and F_{ch} are normalized to a unit area (CH₄ m⁻² day⁻¹), it is possible to directly compare F_{ch} to F_{mod} . For the model evaluation exercise, we extracted the daily F_{mod} corresponding to the same dates of the chamber flux measurements, and only the emissions from that model grid cell where the chamber plots were geographically positioned (grid cell A, Fig. 1b). We also show the results from an adjacent grid cell (grid cell B, Fig. 1b) to demonstrate the spatial heterogeneity between the model grid cells for a region close to the chamber flux measurements. This specific second grid cell was chosen to highlight the fact that even areas that appear similar in overall ecosystem structure can produce deviating CH4 flux rates, for example influenced by environmental factors such as soil depth, inundation fractions, or C3 grass coverage.

Due to the heterogeneous topographic characteristics in the study site, the microsites of the chamber plots within the control area include water-saturated (average water table during the growing season > 10 cm below the surface, observed in eight chamber plots) and unsaturated characteristics (dry soil conditions, i.e., water table < 10 cm below the surface, observed in two chamber plots; Kwon et al., 2016).

Thus, the total F_{ch} from the chamber plots was averaged separately for the wet plots (F_{ch_wet}) and for the dry plots (F_{ch_dry}). This heterogeneity in the data finds its equivalent in the model grid cell heterogeneity as estimated by TOP-MODEL, where on average only a portion of the grid cell area is inundated and the rest remains dry during a specific period of time. The modeled methane emissions correspond exclusively to the portion of the grid cell with near-water saturated soils. Similarly, the chamber flux measurements evidenced predominant emissions of this gas in plots with wet soils, whereas the emissions in dry plots were negligible (see Appendix A3). Thus, to obtain the total F_{ch} , the chamber flux measurements, and to account for emissions predominantly from wet soils, F_{ch_wet} and F_{ch_dry} were scaled to the daily inundated fractions w_{mod} for the corresponding model grid cell A. More details are presented in Appendix A:

$$F_{\rm ch} = F_{\rm ch_wet} \cdot w_{\rm mod} + F_{\rm ch_dry} \cdot (1 - w_{\rm mod}). \tag{1}$$

At two of the chamber sites, temperature sensors (hereinafter referred to as redox systems) continuously recorded the soil temperature profile at three soil depths (4, 16, and 64 cm). The redox systems are located in a site dominated by dry soils and a site dominated by wet soils, and thus these temperature measurements reflect the important influence of soil water levels on the soil thermal regime across the seasons.

2.3.3 Eddy covariance measurements

The model results were also compared to ecosystem-scale methane fluxes measured by an EC tower situated in the Chersky floodplain near the northern end of the chamber plot transect in a control area (Fig. 1 and Tower 2 at 68.62° N and 161.35° E in Fig. 1 of Kittler et al., 2016). The observation height is at 5.11 m.a.g.l., and fluxes are available at 30 min intervals. For details on the instrumental setup, raw data collection, and EC data post-processing, the reader is referred to Kittler et al. (2016, 2017). The analysis of uncertainties in the EC data is presented in Appendix A. The field of view ("footprint area") of an EC system with the given sensor height above the ground normally extends up to several hundred meters in the main wind direction at any given time, changing with atmospheric turbulence conditions (Fig. 1 of Kittler et al., 2016). The position of the EC tower falls within the area of model grid cell A (shown in Fig. 1b) and far away from the grid cell borders; thus, it is assumed that all the CH₄ fluxes measured with the EC system fall within the area of grid cell A. To improve this comparison due to the difference in spatial scale between the EC footprint and model grid cell areas, for the former we analyzed the vegetation composition within the footprint using the highest-resolution land cover maps based on WorldView-2 remote sensing imagery. For this analysis, we aggregated vegetation classes to differentiate between areas of predominant wet soils or wetlands (dominated by the cotton grass *Eriophorum angustifolium*) and dry soils (dominated by shrubs and the tussock Carex appendiculata). We then compared the extent of the wetlands to the inundated fractional area of the model grid cell considered as the corresponding model wet area. It has been recently shown in the literature that the type of vegetation in tundra landscapes is a good indicator of the spatial distribution and variation of CH₄ fluxes (Davidson et al., 2017), and it is also expected that the majority of the CH₄ fluxes are emitted from wetlands in tundra ecosystems (Helbig et al., 2017a). About 26 % of the fluxes measured by the EC tower were emitted from wetland areas within the footprint, i.e., from wet soils with cotton grasses. Within the entire model



Figure 2. Latitudinal distribution of the difference between the fractions of the grid cell inundated areas simulated with TOPMODEL in JSBACH-methane (w_{mod}) and the inundated areas estimated from the EAWS product (w_{rs}) for the same grid cells (1–10 August 2007). The inset figure is the mean spatial distribution of the fraction of inundated areas in the model domain for 1–10 August 2007. Grid cells with inundated areas <1 % are not shown.

grid cell A, the inundated fraction is between 17.7 and 19.9 % (10-day-mean values in summer months) during the summer of 2014, while C3 grasses cover 33.3 % of the area (with no explicit separation between cotton grasses and tussocks). To investigate the EC methane fluxes for a smaller wetland area similar to that one in the model grid cell, it is possible to linearly scale the 10-day-mean EC methane fluxes to the in-undated fraction from the model. Results of this scaling approach for fluxes in summer 2014 are shown in Appendix A.

3 Results

3.1 Evaluation of inundated areas

Within the context of this analysis, fractions of inundation are given as the percentage of the total grid cell area that holds water at or above the surface. The first comparison between remote sensing (w_{rs}) and simulated (w_{mod}) wetland extents, using an initial TOPMODEL configuration, showed that the model mostly overestimated the extent of inundated fractions. For example, in the predominantly wet sections north of the model domain (>68.5° N), the averaged $w_{\rm rs}$ is 9%, whereas $w_{\rm mod}$ was simulated at 15%. However, in drier areas (< 68.5° N) $w_{\rm rs}$ is on average 1.2%, whereas the model did not predict inundation in those grid cells. Since modeled methane emissions only take place in the inundated areas of a grid cell, it was necessary to modify the prescribed TOPMODEL parameters to improve w_{mod} towards w_{rs} . To achieve this, the initially prescribed maximum threshold for inundation ($\chi_{min cti}$) was modified in a similar fashion to the sensitivity experiments through a step change in the parameter value and subsequent analysis of results, until the horizontal extent of inundated areas in the model decreased compared to the results of the initial configuration. Changes in this value have an effect only on wet areas. In Fig. 2 the latitudinal distribution of the percent difference between w_{mod} and $w_{\rm rs}$ for 1–10 August 2007 after parameter adjustment (i.e., $\chi_{min_cti} = 12$) is depicted. We show only the results corresponding to one EAWS image because the results are similar for the other five available GeoTIFF images. The distribution of modeled grid cell inundated areas during the same period of time is shown in the inset of Fig. 2: model grid cells with more than 1% of inundated area are found from the northwestern to southeastern parts of the model domain, and also include some grid cells in the western and northern parts. The spatial distribution of the modeled inundated areas throughout the year does not vary considerably because the inundated fraction in the model takes into account the accumulation of liquid and frozen water. However, the fraction of inundation within each grid cell varies in relation to drier or wetter conditions. After parameter adjustment through step change tests, the comparison between w_{mod} and $w_{\rm rs}$ resulted in a mean difference of -1 ± 8 % and a median of 2% integrated over all six 10-day-mean periods. However, some outlier values result in considerable single-pixel differences between $w_{\rm mod}$ and $w_{\rm rs}$, ranging from +19% in the southernmost areas (<68.5° N) to -23 % in the northernmost areas (>69° N). The best agreement between model and EAWS product is observed between latitudes 68.5 and 69° N (Fig. 2).

During the process of optimization between w_{mod} and w_{rs} , the parameter f was not modified because this would influence both the inundated and dry areas of a grid cell. The best value for $\chi_{\text{min_cti}}$ that resulted in a closer agreement between w_{mod} and w_{rs} is applied in the configuration of the control and reference simulations of this work. We exemplify the effect on the modeled methane emissions due to changes in the $\chi_{\text{min_cti}}$ value, and include this parameter in the sensitivity experiments shown in the following section.

3.2 Sensitivity experiments

We investigated the impact of different values for selected model parameters (shown in column 2 of Table 2) on the individual transport processes and total CH₄ emissions. We compared the results from the reference simulation at daily resolution in 2014 to six pairs of sensitivity experiments (Fig. 3, with pairs of sensitivity experiments shown as panels within rows a through f). The annual mean model domain CH₄ emissions for each experiment are also summarized in Table 2. From all the sensitivity experiments, a significant difference in model output between the reference simulation and the simulations with modified settings was found only for parameters χ_{min_cti} and $f_{CH4anox}$ (for both variables n = 365 and p < 0.01 after a nonparametric Wilcoxon ranksum test; see also Fig. 3, all panels in row a and f).



Figure 3. Results from the sensitivity experiments for the six selected parameters described in Table 2. Daily methane emissions for the individual transport pathways and total methane emissions are shown. The inset figures in some of the panels are zooms to periods of time where larger differences between signals are depicted.

The prescribed threshold parameter χ_{min_cti} in the TOP-MODEL module sets the maximum possible area in the grid cell that can be flooded. A higher χ_{min_cti} value leads to a larger wetland extent in already inundated areas within the model grid cell. Consequently, our results show that a change in χ_{min_cti} has a large effect on the CH₄ emissions: describing $\chi_{min_cti} = 13$ leads to nearly 1.5 times higher CH₄ emissions during summer and autumn compared to the results using the reference value of 12, and about 2 times higher than the results with the lower χ_{min_cti} test value of 11 (Fig. 3, row a). The effect of varying χ_{min_cti} in the resulting model mean inundated fraction is shown in Fig. S3 of the Supplement. With the higher χ_{min_cti} value (i.e., 13), the annual average of the inundated fraction in the model domain (0.054) increases by 54 %, whereas with the lower χ_{min_cti} value (i.e., 11) the annual average inundated fraction in the model domain (0.024) decreases by 35 %, both with respect to the annual average of the inundated fraction in the model domain from the reference simulation (0.0367).

The f_{CH4anox} parameter is a prescribed fixed value used to define the fraction from the total decomposed soil organic matter that will be allocated for CH₄ production (i.e., anoxic carbon mineralization), with the rest becoming CO₂. As in many other land surface models, only mineral soils are considered in our model configuration (limitation further discussed below in Sect. 4.3). For a f_{CH4anox} value of 0.5 (control) the resulting mean summertime CH₄ emissions in the model domain were 5 times higher than the emissions with lower $f_{CH4anox}$ values (0.1 or 0.3) (Fig. 3, row f).

The remaining parameters tested in this sensitivity test (Table 2) show that with the chosen values, the simulated CH₄ emissions are not significantly different from each other. The d_r parameter associated with the plant-mediated transport pathway shows that the difference between the simulated CH₄ emissions is not statistically significant through the year between fine plant roots of 2 mm (as defined in the reference simulation) and thicker roots of 8 mm (Fig. 3, row b). A small variation can be noted for ebullition and diffusion mostly during July (Fig. 3, columns 1 and 2 of row b). The difference in emissions due to an increase in the soil root volume from 20 to 60 % is also not statistically significant (Fig. 3, row c).

For the selected parameters associated with the emissions of CH₄ through the snow, the porosity of snow $\phi = 0.64$ used in the reference simulation (ρ_{snow} of 330 kg m⁻³ for wind packed snow) has a mean tortuosity $\tau = 0.77$, calculated with Eq. S6 (Supplement). The tortuosity value decreases with denser snow, and thus for $\phi = 0.71$ corresponds to $\tau = 0.79$ ($\rho_{snow} = 263$ kg m⁻³ for aged settled snow), whereas $\phi =$ 0.86 means $\tau = 0.85$ (ρ_{snow} of 128 kg m⁻³ for fresh damp new snow). Our sensitivity results from these experiments show that the differences between the winter CH₄ emissions through a layer of fresh damp snow, or through a wind packed snow layer, are not statistically significant (Fig. 3, row d).

Finally, the fixed limiting snow depth, which discriminates between ordinary CH₄ transport via diffusion and diffusion through the snow, was also tested. In the reference simulation, this switch happens at a fixed $h_{\text{snow}} \ge 5 \text{ cm}$. In the sensitivity experiments, we decreased h_{snow} to 3 and 1 cm. The results show that differences between the individual and total CH_4 emissions through various h_{snow} values are not statistically significant (Fig. 3, row e). A time shift is seen however in the CH₄ emissions from mid-October until mid-November (Fig. 3, column 4 of row e), with larger emissions through snow taking place earlier if h_{snow} is thinner. Nevertheless, this temporal shift in the CH₄ emissions through the snow does not influence the total CH₄ emissions. The regionally aggregated CH₄ transport via ebullition, diffusion, and plants during the same months is reduced as h_{snow} becomes thinner, thus compensating for the shift in the emissions in the presence of snow and maintaining a mass balance in the annual total emissions.

3.3 Evaluation of modeled emissions with eddy covariance and chamber measurements

3.3.1 Evaluation of year-round modeled total CH₄ emissions

The methane emissions from EC measurements used here to evaluate the modeled CH_4 emissions at grid cell scale span from April 2014 until September 2015, while data from chamber measurements are restricted to the period of June to August 2014. The modeled total CH₄ emissions used for this comparison correspond to the grid cell where both the EC tower and the chambers are geographically located (grid cell A, Fig. 1b). Also, we show the modeled total CH₄ emissions from the neighboring grid cell to the west (grid cell B, Fig. 1b) and further discuss their association with environmental variables and prescribed parameters in the model. The EC data presented here have been subject to a thorough quality check, and gap-filling was subsequently applied to produce a continuous time series (also see Appendix A for details on uncertainty analysis). For the year-round data evaluation it is not possible to apply the suggested linear scaling approach between the EC flux and model flux data, based on the vegetation type as an indicator of wetland areas in the EC footprint and the inundated fraction predicted in the model (Sect. 2.3.3 and Appendix A). This is due to the lack of year-round vegetation coverage from remote sensing data that would otherwise allow one to obtain a temporal varying wetness area for the EC footprint. However, the results shown in Appendix A from the suggested scaling approach for data in summer 2014 serve as a demonstration that (1) the areas with wet soils within the EC footprint and the model grid cell, translated into the areas where the majority of the CH₄ emissions take place, show only minor differences, and that (2) the offset between methane fluxes from EC and from the model can be largely attributed to these differences in the extent of wetland areas. In the course of this paper we will consider the EC fluxes as representative of the processes within the entire model grid cell, therefore allowing a direct comparison to the modeled CH₄ fluxes.

Despite the large spatial scale of the modeled emissions, the monthly means of the CH₄ emissions from the EC and chamber measurements agree well with the monthly model results for grid cells A and B and for 2014 and 2015. Positive correlations between the measured and modeled CH₄ fluxes, with correlation coefficients (R^2) higher than 0.95, are observed in all comparisons except for the correlation between the chamber measurements and the results from grid cell B ($R^2 = 0.85$; Fig. 4). Fig. 5a and b display box plots of the monthly mean CH₄ emissions for summer months (June, July, and August) from each data set: both grid cells (A and B), EC data for 2014 and 2015, and chamber flux measurements for 2014 only. In 2014, the median of the CH₄ emissions from grid cell B is consistently higher than the rest of the other compared data sets, and this is followed by the EC fluxes (Fig. 5a). The same is observed in 2015 (except for the lack of chamber flux measurements during that year; Fig. 5b). During both years, the median of the modeled CH₄ emissions from grid cell A is generally lower than the rest of the compared data sets (Fig. 5a and b).

The time series of the monthly mean CH₄ emissions from model grid cells A and B is compared with the observational data sets in Fig. 5c. The shaded area around the mean values is 1 standard deviation calculated from the daily values; thus, it represents the range of variability in the emissions



Figure 4. Comparison between modeled CH₄ emissions and flux measurements by chambers and EC in the Chersky floodplain: correlation between results for model grid cells A and B and measurements during 2014 and 2015 (the light grey line is the 1 : 1 line).

within each month. To analyze the contribution of uncertainties in the daily variability of the EC data, random (due to, e.g., turbulent sampling or instrument error) and systematic (e.g., instrument calibration or drift) errors in this data set were assessed. The uncertainties in the EC data are given as error bars in the monthly averages in Fig. 5c and account on average for 0.35 ± 0.22 mg CH₄ m⁻² d⁻¹ of the monthly emissions. These uncertainties are smaller than the spread of the daily variability. A summary of the methods followed to account for these errors is presented in Appendix A.

For the 2 years of analysis, modeled CH₄ emissions of grid cell A underestimate the EC monthly values by 4.7 ± 8.1 mg CH₄ m⁻² d⁻¹. However, the modeled values from grid cell B are higher for most of 2014 and 2015 by as much as 6.1 ± 10.5 mg CH₄ m⁻² d⁻¹. During winter, the model CH₄ emissions from both grid cells are on average 3.7 mg CH₄ m⁻² d⁻¹ lower than the EC measurements (Fig. 5c). The modeled CH₄ emissions from both grid cells show large interannual variability. This is evidenced in Fig. 6, which compares the standard deviation of the monthly fluxes between the 2 years of analysis. In the model results, particularly grid cell B shows large interannual variability in summer months (10.9 mg CH₄ m⁻² d⁻¹ in June and 5.6 mg CH₄ m⁻² d⁻¹ in July).

Methane emissions from the chamber flux measurements are lower than the model results of both grid cells for June and July 2014 (on average by 16.6 mg CH₄ m⁻² d⁻¹ in June and 24.3 mg CH₄ m⁻² d⁻¹ in July), but also than the EC flux

data (on average by 15.3 mg CH₄ and 25.1 mg CH₄ m⁻² d⁻¹ in June and July, respectively). However, the results from grid cell A are in closer agreement with the chamber flux measurements, with the chamber data showing larger emissions by 8.2 mg CH₄ m⁻² d⁻¹ compared to results from grid cell A during August 2014. The shaded areas of the EC data show the largest spread in daily variability of all of the presented data sets, particularly in summer months. Thus, despite the disagreement between monthly mean values, there is an obvious overlap in the shaded areas between all data sets during 2014, while a larger disagreement is observed only during the summer of 2015 between results from grid cell B and EC data (Fig. 5c).

The root mean square error between the daily CH₄ fluxes from grid cell A and the observations, normalized to the mean of the measurements (NRMSE = RMSE/mean (CH₄)_{obs} × 100), is on average < 30 % from June to October for both years, while for spring and winter it is on average 80 %, with the maximum NRMSE during May in 2014 (107 %) and 2015 (104 %). Thus, the large variation of the measured daily fluxes in summer leads to a lower error when compared to the summertime modeled fluxes, whereas the lower magnitude and variation in wintertime fluxes leads to a larger error between modeled values and the observations being captured.



Figure 5. Box plot for summer (JJA) methane emissions from model grid cells A and B, eddy covariance and chamber flux measurements for (a) 2014 and (b) 2015 (without chamber flux measurements). The central horizontal line on each box is the median for each data set and whiskers are the minimum and maximum values; (c) time series of monthly CH_4 emissions for 2014 and 2015 for grid cells A and B in the model, from eddy covariance as well as chamber flux measurements. Shaded areas depict 1 standard deviation of the monthly mean of each data set calculated from the daily resolution model output. Error bars in the EC fluxes are the uncertainty of the monthly averages of the gap-filled and quality checked signal.

3.3.2 Relationship between soil temperatures and CH₄ emissions

To examine the relationship between soil temperatures and CH_4 emissions, we first compared the modeled and measured soil temperature profiles. The temporal evolution of the vertical profiles of daily soil temperatures, measured with the redox systems, is shown in Fig. 7a for the wet plot and in Fig. 7b for the dry plot in 2015. The measured soil temperatures are soil temperatures.

Difference of standard deviation of monthly CH₄ flux between 2014 and 2015



Figure 6. Comparison of the standard deviation of the monthly fluxes between 2014 and 2015.

tures were only available from August to December in 2014, and behaved similarly in 2015 for the same months. The temperature values measured with the sensors at 4, 16, and 64 cm depth were linearly interpolated every 2 cm through the vertical soil column to construct the soil temperature profiles shown in Fig. 7a and b. For comparison, the modeled vertical profiles of the daily soil temperatures in 2015 for the top four soil layers (bottom depth of 3, 12, 29, and 58 cm) in grid cells A and B were also linearly interpolated every 2 cm (Fig. 7c and d). During winter and spring, the measured soil temperatures are not lower than -16 °C, while the modeled temperature values are as low as -26 °C within extended sections of the period from December to May. The measured values in the dry plot show abrupt temperature changes during the transition between freezing conditions ($<0^{\circ}C$) and warmer conditions (>0°C) during mid-December and mid-May. This abrupt change is also seen in the wet plots with freezing conditions remaining for a shorter period of time; i.e., the change to and from warmer temperatures takes place only from the end of January until mid-May. Also, generally colder temperatures are observed in the top part of the soil column and gradually extend to deeper soil layers as the season progresses. In contrast, although the modeled soil temperatures reach lower values during winter, a smoother transition of temperature is evidenced from freezing to warmer conditions in spring, and to freezing conditions again in autumn. In the model results, the soil temperature remains homogeneous along the vertical profile.

Measured temperatures above freezing conditions occur from mid-June until the end of September. As summer progresses, warmer soil temperatures extend from the surface to deeper soil layers (Fig. 7a and b). In the dry plot, however, the warmer conditions remain only in the top 16 cm of the soil column (Fig. 7b) due to lower soil moisture content and lower thermal conductivity compared to the wet plot. The model is able to capture the timing of the seasonal transition from spring to summer at the end of May, the duration of the summer conditions, and the magnitude of the tempera-



Figure 7. Hovmöller diagrams showing the time evolution of the vertical profiles of daily soil temperature during 2015 from eddy covariance fluxes measured (**a**) at the wet plot and (**b**) at the dry plot and from the model data (**c**) grid cell A and (**d**) grid cell B. The data were interpolated linearly from the depths where data are available (4, 16, and 64 cm in the sensors of redox systems and 3, 12, 29, and 58 cm in the model).

ture values. For grid cells A and B, the summer temperature profiles are more similar to the wet than to the dry plot. The average measured soil temperature in the range of the sensor's depths (top 64 cm) during summer (June, July, and August 2015) in the dry plot was 2.1 °C, while in the wet plot for the same period the average measured soil temperature was 4.7 °C; in the model, the average soil temperature in the top 58 cm is about 4.9 °C during the summer of 2015. The modeled warm soil temperatures $(>5 \circ C)$ reach deeper soil layers in summer; however, this is not observed in the measured data. This could simply be due to the coarse vertical resolution of the data because of the large gap between sensors (from 16 to 64 cm depth). Thus, to evaluate the extent of the warm soil temperatures depicted in the model, this portion of the soil column needs to be better resolved vertically by the measurements.

A larger disagreement between measured and modeled soil temperatures, however, occurs during the transition from

autumn to winter. The measured temperatures remain around -3 °C in the top 64 cm from October to mid-December, until they change abruptly to around < -10 °C in the dry plot during mid-December (Fig. 7b), and in the wet plot towards the end of January (Fig. 7a). In contrast, the model results show a gradual transition between the seasons, with decreasing soil temperatures to values < 0 °C starting in mid-October (Fig. 7c and d).

To investigate the effect on the CH_4 emissions due to the abrupt changes in the measured soil temperatures, we plotted the soil temperature at 12 cm (for model data) and at 16 cm (measured values for the wet plot) against the total CH_4 emissions for grid cells A and B and from EC measurements in 2014 (Fig. 8). The modeled soil temperatures represent the entire grid cell conditions, whereas the CH_4 emissions are only from the saturated and inundated portions of the grid cell (Fig. S2). Despite this disagreement, CH_4 processes in the model follow the seasonal variation in soil temperature.



Figure 8. Model soil temperatures at 12 cm depth and measured values at 16 cm depth measured at a wet site, against the total methane emissions for grid cells A and B in 2015.

This relationship, however, can only be analyzed on a qualitative basis. A positive non-linear correlation between soil temperatures and CH₄ emissions is observed in all comparisons. Fitted polynomial curves are plotted on top of each data set. During 2015 (and 2014, data not shown), the CH₄ emissions measured with EC drop faster with the changes in temperature until freezing conditions. Between -3 and -14 °C, little variation in the lowest CH₄ emissions is observed, whereas the change in modeled CH₄ emissions with respect to changes in soil temperature is more gradual within that range of sub-zero temperatures. Lower methane emissions in the model compared to those in the EC data take place in winter and are associated with even lower soil temperatures than the ones registered by the sensors in the redox systems.

3.4 Year-round modeled methane emissions

Domain means of the seasonal courses of CH_4 emissions from the different CH_4 transport pathways in 2014 and 2015 from the reference simulation, as well as the daily mean snow depth, are shown in Fig. 9. The results show a distinct seasonality for each of the individual methane emission pathways. Overall, the lowest CH_4 emissions occur between November and May. During these months, the timing of the CH_4 emissions through the snow is largely modulated by the changes in the snow depth, and accordingly takes place predominantly in spring and autumn. The methane emissions via plants, ordinary molecular diffusion, and ebullition are mostly restricted to the period May through mid-November in areas when and where h_{snow} does not exceed 5 cm (or is absent). The magnitude of the CH₄ emissions through molecular diffusion is the least relevant among the four modeled transport pathways.

3.4.1 Summertime CH₄ transport pathways

From May to mid-November, CH₄ emissions take place only in the grid cells with inundated areas, with the highest flux rates simulated for the center west of the domain (Fig. S5). During this period h_{snow} is either absent or does not exceed 5 cm. Ebullition precedes the emissions through plants during late March 2014 and during early April 2015. In both years, the mean of the CH₄ emissions in the model domain through ebullition rises steadily, followed by a short but pronounced decrease to $3.5 \text{ mg CH}_4 \text{ m}^{-2} \text{ d}^{-1}$; the ebullition of CH₄ rises again to reach its maximum during midsummer, with a similar magnitude in both years (7.2 mg $CH_4 m^{-2} d^{-1}$). This maximum is achieved later in 2014 (9 August) than in 2015 (16 July) and during the same days as the peak of maximum gas transported through plants. The domain means of the CH₄ emissions through plants reached their maximum value of $15 \text{ mg CH}_4 \text{ m}^{-2} \text{ d}^{-1}$ in both years. Similar to ebullition, right after the summer maximum of emissions, the emissions through plants start to decrease until mid-November. In contrast to 2014, a shoulder is shaped in the emissions through ebullition and plants, and to a lesser extent in diffusion, during the end of August to mid-September in 2015, indicating the continuation of CH₄ emissions even as the soil starts to freeze (Fig. 9). Annually, CH₄



Figure 9. Year-round mean simulated CH_4 emissions of the model domain through different pathways and domain mean snow depth for (a) 2014 and (b) 2015.

transported by plants is the dominant pathway, contributing 61 % to the total domain mean annual CH₄ emissions. These emissions take place from May to mid-November in areas where $h_{\text{snow}} < 5$ cm, and are restricted to areas where C3 grasses are present (Fig. S1 and the panels of the first column in Fig. S5). The gas transported through ebullition is 33.9 % in 2014 and 35.7 % in 2015 of the total annual CH₄ emissions.

Methane emissions through ordinary molecular diffusion also take place if $h_{\text{snow}} < 5 \text{ cm}$ in the inundated portion of the grid cells (panels in the third column of Fig. S5). In the absence of snow during summer and early autumn, CH₄ emissions via diffusion in the model domain average about 2.9×10^{-3} mg CH₄ m⁻² d⁻¹ (similar in 2014 and 2015), while during late autumn, winter and early spring the emissions via this pathway are only possible if $h_{\text{snow}} \ge 5$ cm. For those few grid cells with $h_{\text{snow}} < 5$ cm during the nongrowing season (November to May), the CH₄ emitted via molecular diffusion is 2 to 3 orders of magnitude lower (mean of 3.4×10^{-5} mg CH₄ m⁻² d⁻¹ for both years) than during the growing season (June to September) (Fig. 9). Methane transported via molecular diffusion during the growing season contributes only 0.02 % to the total CH₄ annual budget.

3.4.2 Impact of snow on the winter and seasonal variation of CH₄ emissions

During early spring, late autumn, and winter, methane emissions take place through a layer of snow $\geq 5 \text{ cm}$ deep. The mean maximum accumulation of snow in the model domain takes place in spring: earlier in 2014 (0.23 m on 21 March) than in 2015 (0.17 m on 8 March). The spatial distribution of the spring snow depths in 2014 and 2015 (Fig. S4a and b) shows deeper snow layers in the dryer southwestern part of the model domain. On average, the layer of snow starts to melt rapidly at the beginning of May in 2014 and at the end of April in 2015, reaching total snowmelt by 2 June 2014 and 27 May 2015 (Fig. 9). The average CH₄ emissions through the snow in the entire model domain during January and February are $0.17 \text{ mg} \text{ CH}_4 \text{ m}^{-2} \text{ d}^{-1}$ in 2014 and 0.12 mg $CH_4 m^{-2} d^{-1}$ in 2015. The CH_4 emissions fluctuate through the winter, and these changes are related to changes in the thickness of the snow cover. During the rapid snowmelt period in spring (March, April, and May), the daily domain average CH₄ emissions into the atmosphere through the snow increase (Fig. 9), with domain mean average spring CH₄ emissions of 0.65 and 0.43 mg $CH_4 m^{-2} d^{-1}$ in 2014 and 2015, respectively. The maximum domain mean daily emissions of CH₄ outside the growing season are modeled during May, with 1.66 mg $CH_4 m^{-2} d^{-1}$ in 2014 and 0.96 mg $CH_4 m^{-2} d^{-1}$ in 2015, and these take place predominantly in the central part of the model domain (panels in the fourth column of Fig. S5). In the entire model domain, the emissions of CH₄ through snow contribute 4.7 and 2.7 % to the total mean annual CH₄ emissions for 2014 and 2015, respectively. Although deeper spring snow layers are modeled in 2014 than in spring 2015 (Fig. 9) in the areas where CH_4 is emitted into the atmosphere (Fig. S4a and b), the total methane emissions through snow from January to mid-May amount to \sim 70 mg CH₄ m⁻² in 2014, and only 66 % of that value in 2015 ($\sim 46 \text{ mg CH}_4 \text{ m}^{-2}$; Fig. 9).

Integrated over the model domain during autumn, the snow starts to accumulate later in 2014 (9 October 2014) than in 2015 (30 September 2015), and the snow layer becomes rapidly deeper until December at a similar accumulation rate for both years (Fig. 9). As the snow accumulates, the emissions via ebullition and plants decline, but diffusion through snow rises as soon as the snow depth reaches 5 cm in some grid cells. From November to December, the mean CH₄ emissions through the snow in the domain amount to 37.3 mg CH₄ m⁻² in 2014, and 33 % less in 2015 (12.4 mg CH₄ m⁻²). The modeled CH₄ emissions through the snow only consider the ordinary molecular diffusion of CH₄ between the soil and the atmosphere, and the pressure pumping



Figure 10. Mean daily ancillary variables and CH_4 emissions in the model domain in 2014 and 2015: (a) total CH_4 emissions, (b) mean soil temperature in the root zone (top five soil layers), (c) domain mean relative soil moisture content in the top five soil layers, (d) domain mean relative soil ice content in the top five soil layers, and (e) inundated fraction of the grid cell.

effects due to advection of gas by wind are not taken into account.

At the grid cell level, in Fig. 10 we show the CH₄ emissions through the snow from the EC measurements and those from grid cells A and B simulated by the JSBACH model, all at daily resolution. The time series of daily emissions are shown from the beginning of October 2014 to the end of April 2015 (Fig. 10a) and in October 2015 (Fig. 10c). The difference between the model methane emissions for grid cells A and B and EC data is shown in Fig. 10b and d for the same cold season periods. Comparable to the EC measurements, the winter emissions in the model drop abruptly at the end of October 2014, remaining low until March 2015. During October 2014, the model CH₄ emissions in grid cell B are higher, while the emissions from grid cell A are more similar to the EC measurements (Fig. 10a). This is also found in the first half of October 2015 (Fig. 10c). However, during this month the EC measurements show no clear trend, while the model CH₄ emissions show a decreasing trend over time. During most of the winter in 2014/2015 (i.e., from November 2014 until April 2015), the modeled CH₄ emissions from grid cells A and B remain lower than the EC measurements by on average $2.8 \text{ mg CH}_4 \text{ m}^{-2} \text{ d}^{-1}$. During January, February and March in 2015 the mean model CH₄ emissions for grid cells A and B are $0.4 \text{ mg CH}_4 \text{ m}^{-2} \text{ d}^{-1}$,

while the EC data show persistently higher values averaging 3.8 mg CH₄ m⁻² d⁻¹ for the same months (Fig. 10a). Model emissions start rising (2.0 mg CH₄ m⁻² d⁻¹) to values similar to those in the EC data (2.8 mg CH₄ m⁻² d⁻¹) only in mid-April.

To investigate whether the CH₄ emissions from the model during the entire wintertime are equivalent to the total winter emissions measured by EC, we calculated the cumulative sum of the modeled CH₄ emissions and EC from October 2014 to March 2015. The uncertainty as the standard deviation of the monthly cumulative fluxes is shown in error bars for each data set (Fig. 10e). Our results show that, despite a higher earlier release of methane in grid cell A, the modeled total emissions released during that winter are not equivalent to those from the EC measurements, with the latter providing evidence of larger total CH₄ emissions in winter than predicted by the model. The cumulative uncertainties are also larger in the eddy covariance data, and this is due to the large daily variability compared to the model results. In our model, the emissions through the snowpack only take into account the molecular diffusion of gas, whereas the advection of gas due to wind as an additional transport pathway is not included.



Figure 11. Time series of the daily mean of methane emissions through snow from EC measurements and model data for grid cells A and B during (a) October 2014 to March 2015 and (c) October 2015; the difference between grid cells A and B and the EC data are shown in panels (b) and (d) for the same period of time; (e) cumulative CH_4 emissions for the period from the end of autumn in 2014 until the end of spring in 2015 for the same data sets. Error bars in each data set are the standard deviation of the monthly accumulated fluxes.

3.4.3 Impact of environmental controls on CH₄ flux seasonality

Several systematic interannual differences between the timing and magnitude of the individual CH₄ transport pathways in 2014 and 2015 were found in the model results. These include, e.g., the maxima of the individual emissions, which occur a few days later in 2014 than in 2015. To improve the interpretation of the temporal variability of CH₄ emissions through the different pathways, we analyze the temporal changes in soil temperatures within the root zone (top five soil layers) as simulated by the model. It is important to note that because of the current structure of the model, the depicted soil temperature in Fig. 11b reflects the average conditions of the entire grid cell, and not only the inundated portion with saturated soils where CH₄ emissions take place. Still, the analysis of the temporal changes in the mean grid cell soil temperatures gives an indication of the nature and magnitude of the seasonal changes that indirectly control the CH₄ emissions. The gradient of temperatures in the root zone for the entire domain between spring and summer is steeper in 2015 than in 2014 (Fig. 11b). The maximum soil temperatures are similar in both years (8.7 °C); however, this maximum was reached at the beginning of August in 2014, while in 2015 the maximum was reached at the beginning of July and remained high throughout August. During the rest of the year, the mean soil temperatures were 2 °C higher in 2014 compared to 2015 (-4.5 and -6.5 °C, respectively). The mean changes in temperature in the top five soil layers reflect the changes in the air temperature as given in the atmospheric forcing data. According to the mean air temperature in the model domain, the summer of 2014 was colder than the summer of 2015 (by up to 10 °C for individual days during June, Fig. S6a). This leads to delayed warming of the soil, later high CH₄ production, and thus a later release of CH₄ into the atmosphere during summer in 2014 than in 2015, as shown in Fig. 11a. These findings are in good agreement with those recently presented in Helbig et al. (2017b). In a comparison of meteorological records of air temperature between 2013 and 2016 in northwestern Canada, the authors found that the coldest May of those years took place in 2014. As a result, during that year a shift in air temperature influenced the soil temperature, and with it the year-to-year methane fluxes, especially during spring.

Figure 11c depicts the model domain mean relative soil moisture content in the top five soil layers for 2014 and 2015. As with soil temperature, the soil moisture reflects the average conditions of the entire grid cell and not just those in the inundated portion where the soil moisture is set to nearly saturation levels. Although these values are not linked to the

area of the grid cell where CH₄ is transported and emitted, we can still show the temporal changes in soil moisture content in the non-saturated portion of the grid cell between years and seasons. These changes can be linked to changes in precipitation patterns (Fig. S6b) and soil temperatures. According to the mean precipitation from the CRU-NCEP reanalysis data, more precipitation fell in the model domain during early July in 2014 compared to the same period in 2015 (Fig. S6b). This led to the top five soil layers becoming wetter on average in 2014 (Fig. 11c) and potentially allowed higher thermal capacity in the soil during that period. In contrast, more precipitation fell during most of August and September 2015 than for the same periods in 2014 (Fig. S6b), leading to an increase in the relative moisture content towards the end of summer and early autumn (Fig. 11c). These changes in soil moisture influence the soil temperature at the grid cell scale, and thus the soil temperature feedbacks to the CH₄ processes. Therefore, it is possible to indirectly relate the effects of changes in grid-cell-scale soil moisture to the changes in the modeled CH₄ emissions.

The mean relative soil ice content in the top five layers of the model domain (Fig. 11d) was higher in the winter and spring of 2014 than in 2015, and this is a general observation for the entire domain. However, the air temperatures from the reanalysis data during that period were on average higher in 2014 than in 2015 (Fig. S6a). The ice content decreases at a fast rate during June in both years; however, the complete loss of ice in the soil is reached earlier in June of 2015 than in 2014, and this is a reflection of colder temperatures in June 2014, delaying the complete melt of the more abundant ice in the soil during that year relative to the same month in 2015 (Fig. S6a). The soil ice content feeds back to the modeled available pore space for CH₄ production; thus, the ice content changes in the soil can be indirectly linked to the CH₄ emissions. The earlier reduction of ice content in the soil during June 2015 might have contributed to the earlier release of methane during that month, via ebullition, compared to 2014 (Fig. 9). The lower air and soil temperatures at the beginning of autumn in 2015 (Fig. S6a) led to higher ice content in the soil during October 2015 compared to 2014 (Fig. 11d). The soil temperatures remain warmer in autumn of 2014, enabling more CH₄ to be emitted during November 2014 when the snow starts to accumulate, in contrast to 2015 (Figs. 9 and 11a).

4 Discussion

4.1 Sensitivity experiments

Through the model sensitivity experiments we identified that changes to the values of the parameters χ_{min_cti} and $f_{CH4anox}$ caused statistically significant differences in the total CH₄ emissions (p < 0.001). A significant increase in CH₄ emissions with increasing inundated surface area (TOPMODEL

parameter $\chi_{min cti}$) highlights the importance of this approach in regulating the extent of the grid cell inundated areas. However, further investigations and improvements in the TOPMODEL approach, as well as a better integration into the hydrology scheme of JSBACH, are needed in order to better constrain the modeled CH₄ emissions with JSBACH. The results of our sensitivity experiments also provided evidence that the magnitude of the simulated CH₄ emissions responds strongly to changes in the parameter values of the fraction of anaerobic decomposed soil organic matter that becomes methane, $f_{CH4anox}$. In soil systems where fermentation and methanogenesis are exclusive processes, i.e., without the presence of alternative pathways for respiration via terminal electron acceptors by other microbial groups that ultimately can suppress the production of CH_4 , the CO_2 : CH_4 ratio after anaerobic carbon mineralization is normally 1:1 (Conrad, 1999); i.e., $f_{CH4anox} = 0.5$. We used this value in the reference simulation because it was previously reported in the literature as characteristic of water-saturated polygon centers (Preuss et al., 2013), and it is similar to the value reported for unsaturated zones in boreal bogs (Whalen and Reeburgh, 2000). However, in wetland areas, CH₄ is still subject to oxidation after its production and the CO₂ : CH₄ ratio is expected to increase and to vary among types of wetlands (Bridgham et al., 2013). Thus, although the value of $f_{CH4anox}$ determines the fraction of CH₄ produced under anoxic conditions, this CH₄ still can undergo oxidation before it is emitted into the atmosphere. Furthermore, $f_{CH4anox}$ can be theoretically related to the fraction of CH₄ that is left after oxidation and before it is emitted into the atmosphere ($f_{ox} = 1$ $f_{CH4anox left}$). Values of f_{ox} have been previously reported as ranging between 0.6 and 0.7 for sites with vascular plants. On the other hand, it can be nearly equal to 1 in sites with, for example, a layer of Sphagnum moss, where the majority of the produced CH₄ is oxidized, or in bottom soils in pond centers where slow molecular diffusion of CH₄ takes place through the water (Knoblauch et al., 2016). Under the latter conditions, f_{ox} can be approximated to >0.9 (i.e., >90 % of the produced CH₄ is oxidized before it is emitted into the atmosphere). This value has been estimated in polygonal ponds without vascular plants, empirically supporting the relevance of CH₄ oxidation below the water table in these types of environments (Knoblauch et al., 2016). A lower CH₄ oxidation fraction occurs in the presence of vascular plants that are effective at bypassing the aerobic areas in the soil. Under these conditions, $f_{CH4anox_left}$ can increase moderately from 0.2 to 0.4 (i.e., $f_{\rm ox}$ is from 0.6 to 0.8, meaning that 60 to 80% of the produced CH₄ is oxidized in the soil column). Although current estimates for $f_{CH4anox_left}$ from laboratory and on-site experiments are still scarce, they mostly agree that those are lower than our reference value of 0.5. This is expected because $f_{CH4anox}$ left excludes the portion of CH₄ that is oxidized directly after production, whereas $f_{CH4anox}$ is only the initially produced CH₄. Still, our modeled CH₄ emissions might benefit from prescribing a spatially variable f_{CH4anox} value linked to the distribution of vascular plants and soil wetness in the model domain.

As for the rest of the selected parameters for the sensitivity exercise, no significant differences were observed in the modeled CH₄ emissions for the individual pathways or the total flux. Specifically, varying the diameter of roots from finer to thicker, and varying the amount of available soil volume occupied by roots, did not cause significant differences in modeled CH₄ emissions with the new formulation of the plant transport in the JSBACH-methane model. These results suggest that the revisited and simplified formulation for plant-mediated transport of gas allows a reduction in the uncertainties of methane transported through this pathway, which previously relied on predefined plant root characteristics that are often not available from observational studies. Instead, we define the volume in the soil that is occupied by roots.

The lack of sensitivity in the CH₄ emissions to most of the selected parameters might ultimately be due to the explicit restriction of gas transport via diffusive processes modeled by Fick's first law (plant transport and molecular diffusion through snow) that was set in the model. The role of this restriction is to limit the diffusion of gas once the concentration gradient between two interfaces equals zero; i.e., it reaches equilibrium. Thus, this restriction takes place when the concentration gradient between, e.g., the gas in the soil pore spaces and within the plant's roots (for plant-mediated transport) or between the gas in the soil pore spaces and the atmosphere above the snow layer (for diffusion of gas through the snow) equals zero. Because the transport pathways can occur in parallel (except for diffusion with and without a snow layer), or emissions can be shifted in time, the modeled total CH₄ emissions may not be influenced by the set of parameters tested here. Finally, changes in the threshold depth of snow that limit the diffusion of gas through this layer revealed some differences in the partitioning of the methane flux into the four transport pathways. These differences indicate that a thinner threshold depth favors the other three transport pathways. However, the resulting total CH₄ emissions with the three tested snow threshold depths were not statistically different.

4.2 Year-round model methane emissions

We simulated for the first time year-round methane emissions in a Northeast Siberian region centered on the city of Chersky. Our results showcase the ability of the improved JSBACH-methane model to reproduce seasonality in the CH₄ emissions when compared to fluxes measured by EC and chambers in a study site near Chersky. The different transport pathways in this process-based model play an important role in defining the timing of the year-round emissions since they are closely linked to the soil physical state and expected share in the total emissions. During the growing season, plant-mediated transport dominated the emissions, contributing about 61.4 % in 2014 and 61.7 % in 2015 of the total annual CH₄ emissions, followed by ebullition (33.9 and 35.7 %) and molecular diffusion during summer when snow is not hindering the emissions (0.02 % for both years). These patterns agree well with the findings presented in Kwon et al. (2016) for the CH₄ emissions measured with chambers at the Chersky floodplain, and by Kutzbach et al. (2004) and Knoblauch et al. (2016) in the Lena River Delta. In these works it is shown that the dominant CH₄ transport pathway in tundra wetland ecosystems (about 70–90 % of the total annual emissions) is diffusion through the aerenchyma structures of the plants when they are present. Methane emissions during the non-growing season contributed 4.7 and 2.7 % of the annual methane emissions in 2014 and 2015, respectively.

As for the methane oxidation, the bulk soil oxidation accounts for about 1 % of the total methane production during the growing season at grid cell scale, and only about 0.6 % for rhizospheric CH₄ oxidation (results not shown). This leads to most of the methane that is produced in the soil being emitted into the atmosphere through the different transport pathways. Past observational and laboratory studies have estimated the methane oxidation in boreal and tundra soils. Whalen and Reeburgh (2000) showed that about 55 % of the CH₄ diffusing from saturated boreal soils was oxidized while reaching the surface. Through bottle incubations, Knoblauch et al. (2016) measured the volumetric CH₄ oxidation potential of soil and moss samples collected from ponds of the Lena Delta. The authors found that the fraction of produced CH₄ that is oxidized before it is emitted was between 61 and 78 %using a stable isotope approach. In samples from pond areas without vascular plants, the fraction increased up to 90 % of the total produced CH₄ following a potential methanogenesis approach, and from diffusive CH4 fluxes into the bottom water this was between 63 and 94 %. Berestovskaya et al. (2005) measured CH₄ oxidation rates of different soil samples from the Russian Arctic tundra and found that generally the rates of methane oxidation exceeded the rates of methane production, especially at temperatures of 5 °C. For this to happen, methane-oxidizing bacteria rapidly consume the methane released from the freshly thawed tundra soils and the methane already deposited in the unfrozen soil, and this takes place even before methanogens produce new methane. Based on these scarce observations in boreal soils, the oxidation processes in our model are still far off and need to be revisited in order to improve the contribution of the methane oxidation processes to the total methane emissions.

The JSBACH-methane model does not explicitly consider specific mechanisms related to the carbon decomposition and thaw in Arctic permafrost and wetland ecosystems, such as CH₄ production in the soil from root exudates (Knoblauch et al., 2016; Ström et al., 2012), vertical transport of soil organic matter and its vertically resolved decomposition (Braakhekke et al., 2011, 2013; Koven et al., 2015), and microbial community dynamics (McCalley et al., 2014) involved in anoxic CH₄ oxidation or the production of CH₄ in anaerobic microsites confined in oxic soils. Although these processes might contribute substantially to the dynamics of CH₄, research on these processes in soil-permafrost and wetland environments is still lacking or poorly understood, with controversial results so far (Bridgham et al., 2013).

At the grid cell scale, the characteristics defined in the model input parameters exert an important influence on the spatial heterogeneity and temporal variability of the modeled environmental controls and CH₄ emissions. For example, in the model domain the soil depths range between 0.1 and 10.6 m (Fig. S4c; grid cell A is 0.89 m and grid cell B is 10.6 m), whereas the depth of the root zone is from 0.1 to 0.89 m (Fig. S4d; grid cell A is 0.72 m and grid cell B is 0.67 m). Also, the cover fraction of vegetation differs among grid cells, and in this model, the coverage of C3 grasses is particularly relevant for CH₄ emissions through the plant roots, e.g., 33.3 % in the area of grid cell A and 91.6 % in grid cell B (Fig. S1). Finally, grid cell A has lower soil moisture and soil ice content relative to the pore volume in the top five soil layers, and a larger inundated area, compared to grid cell B (Fig. S8a, b, and d). These differences predominantly explain the shift in the dominant growing season CH₄ transport pathways and seasonal changes between grid cells (Fig. S8e and f).

To further demonstrate the heterogeneity in the modeled total CH₄ emissions, we show in Fig. S5 the time series of the daily CH₄ fluxes in 2014 and 2015, for nine model grid cells (grid cell A and the eight grid cells surrounding it, which includes grid cell B). In this area of the domain, the range of the mean emissions is between 24 and 75 mg CH₄ m⁻² d⁻¹, with similar values between years 2014 and 2015. To further analyze the spatial heterogeneity, more is discussed below in the context of spatial distribution of CH₄ fluxes in the entire model domain (Fig. S9).

The modeled CH₄ emissions represent fluxes from exclusively inundated areas (water table at or above the surface); thus, emissions from areas with a water table below the surface are neglected. In Fig. A3b (Appendix A) are shown summertime CH₄ fluxes (June to August 2014) measured with chambers in the Chersky floodplain (Kwon et al., 2016), plotted against the water table in the chamber microsite at the time of the flux measurements. CH₄ oxidation predominantly exceeded production in dry microsites (water tables were below the surface up to about 10 cm), and this was evidenced by a small uptake of CH₄ (on average 3 mg CH₄ m⁻² d⁻¹). The fluxes in these dry plots were almost negligible during the growing season. Thus, the modeled fluxes of CH₄ represent the majority of the emissions in this tundra ecosystem.

Our results show a good agreement between the modeled CH_4 emissions (at the grid cell scale) and measured CH_4 emissions with EC and chambers. Overall, the modeled yearround and measured methane emissions at daily temporal resolution are on the same order of magnitude, and both fall within their monthly range of variability. In both the EC footprint area and model grid cell area, the methane emissions are not spatially homogeneous, but bound to the distribution of wetland (inundated) areas, which are also linked to the type of vegetation. This was demonstrated for the summer of 2014, where EC CH₄ fluxes are in closer agreement with the model methane emissions after a linear scaling approach of the wet soil areas in the EC footprint.

In the model, CH₄ emissions integrated in our study region were on average 22.5 mg $CH_4 m^{-2} d^{-1}$ during the growing season of 2014 and 2015. These modeled values are also in good agreement with measurements in other Arctic wetland areas influenced by permafrost using eddy towers, chambers, and more recently airborne techniques. Kutzbach et al. (2004) reported CH₄ emissions of 28 mg $CH_4 m^{-2} d^{-1}$ measured with chambers during the onset of the growing season from a polygon center of the wet tundra in the Lena Delta. For a variety of locations in polygons of the same region, Sachs et al. (2010) reported mean summer methane emissions of about 55 mg $CH_4 m^{-2} d^{-1}$. Knoblauch et al. (2016) presented mean summer fluxes of $46 \text{ mg CH}_4 \text{ m}^{-2} \text{ d}^{-1}$ also measured with chambers at the margins of ponds also in the Lena Delta. Larger summer methane emission values have been reported elsewhere, e.g., from automatic chambers at the Zackenberg research station, with maximum emissions of about 168 mg CH_4 m⁻² d⁻¹ at the onset of the growing season (Mastepanov et al., 2008). Merbold et al. (2009) reported CH₄ emissions of $\sim 600 \text{ mg}$ $CH_4 m^{-2} d^{-1}$ measured by chambers at the peak of the growing season (August) in 2005 in the Chersky floodplain.

At the lower end of the observational data, Wille et al. (2008) measured CH₄ emissions of about 30 mg CH₄ m⁻² d⁻¹ during mid-summer in the Lena Delta. The authors argued that the measured values were generally lower than other estimates and that the main controlling factors of their measurements were low soil temperatures and the influence of atmospheric turbulence during their period of study. Rinne et al. (2007) reported CH₄ fluxes of about 84 mg CH₄ m⁻² d⁻¹ measured using EC at a boreal fen in southern Finland. Eddy covariance CH₄ fluxes measured in the Alaskan tundra showed a larger range of values, with an average of 32 mg CH₄ m⁻² d⁻¹ during the onset of the growing season (Zona et al., 2016). Finally, airborne measurements of CH₄ emissions from wetlands in Alaska were estimated to be about 56 mg CH₄ m⁻² d⁻¹ (Chang et al., 2014).

4.3 Representation of inundated fractions of the grid cell

In this model version, we incorporated the TOPMODEL approach to explicitly model the distribution of inundated areas according to the topography profile. Although this is still only a robust approximation, the implementation of this approach enabled the representation of wetlands in the highly heterogeneous landscape of Northeast Siberia, which is not possible with the traditional hydrology scheme of JSBACH

because it does not allow standing water (Kaiser et al., 2017; Hagemann and Stacke, 2015). In contrast to standard remote sensing products, the highly resolved product of Reschke et al. (2012) is the best data available so far for our application. After TOPMODEL parameter adjustments, large differences still remain between w_{mod} and w_{rs} ; however, most of these can be attributed to uncertainties due to the model technique to simulate wetlands and its horizontal resolution.

In the model, the fraction of the grid cell that becomes inundated refers to the area where the water table lies at or above the soil surface and varies with changes in available solid (snow) and liquid water (rain) leading to a 15-day time step. However, in nature, most wetlands have periods of the year when no visible standing water is above the surface, and the water table is located a few centimeters below the surface (Bridgham et al., 2013; Kwon et al., 2016). Previous studies have demonstrated the dependence of CH₄ emissions on the location of the water table in tundra ecosystems (e.g., Helbig et al., 2017a; Kwon et al., 2016; Merbold et al., 2009; Sturtevant et al., 2012; Zona et al., 2009) and, as shown before, the work by Kwon et al. (2016) revealed that CH₄ fluxes measured by chambers in the Chersky floodplain are significantly influenced by the location of the water table at the plot scale. Larger CH₄ emissions were measured in sites where the water table was at or above the surface compared to drier sites. With the TOPMODEL approach it is not possible to characterize the location of inland water bodies (i.e., lakes), and the explicit location of peatlands is also not taken into account because the model only considers mineral soils. This separation would help to identify the inundated portions of land with more or less relative input of organic carbon to better localize the methane emissions. The lack of organic layer representation in the model is mainly due to the difficulties of coupling sub-grid-scale hydrology and carbon cycles in a holistic manner. The model however considers the amount of carbon that is available in the soil, based on a soil carbon and litterbag approach, and that one for decomposition and production of methane.

Future important advancements in our model are necessary in the context of a process-based representation of peatland extent as well as the CH_4 balance in non-inundated areas; currently, these are not taken into account in our study, contrary to Kaiser et al. (2017) for a site-level study. This is especially relevant for the applicability of this model to other regions where uptake of methane in dry areas might play a substantial role (e.g., Flessa et al., 2008; Jørgensen et al., 2015).

4.4 Impact of the revised model structure

The model reproduces well the observed temporal trends in the CH_4 emissions, and patterns can be linked to changes in the environmental controls. However, integrating the TOP-MODEL approach into JSBACH led to a decoupling of some physical soil state variables. Soil moisture content, soil ice content, and soil temperature influence the heat capacity of the soil and the ice content (i.e., soil freeze and thaw processes), and control the accumulation of gas, microbial activity, diffusion rates of gases, and amount of oxygen in the soil (Sturtevant et al., 2012; Wickland et al., 1999; Pirk et al., 2016). In the JSBACH-methane model, soil moisture of the ice-free soil pores in the inundated part of the grid cell was set to 95 % saturation, for purposes of justifying inundation in the TOPMODEL approach. Although the temperature and ice conditions in the soil are not influenced by this change, this leads to a missing link in terms of the distribution of soil water to soil ice or soil moisture. However, a direct connection between each of these physical variables and the CH₄ processes is definitively present.

Between data years, the soil temperature in the summer of 2014 was lower from mid-June than at the same period during 2015, leading to a phase lag in the maximum summer CH₄ emissions of nearly a month earlier in 2015. However, during the autumn of 2015, the soil was colder than in 2014, with more soil ice content and less methane production during this period of the year. This translates into lower CH₄ emissions into the atmosphere from November 2015 until the end of the year than during the same period in 2014. Because seasonal changes in soil wetness must be taken into account for modeling year-round gas emissions in permafrost Arctic tundra environments (Pirk et al., 2016), the JSBACH-methane version used for this work requires further improvements to better integrate the TOPMODEL approach using a fully mechanistic thermal and hydrology scheme at landscape scale able to interact with inundated area fractions at grid cell scale (Stacke and Hagemann, 2012).

4.5 Simulation of grid cell soil temperature

Large uncertainties in the simulation of CH₄ emissions from northern wetlands with models come from limitations in the representation of freezing and thawing soil processes, snow layer dynamics, and the robust mapping of the distribution of wetlands (Bridgham et al., 2013). Kaiser et al. (2017) reported that the process-based JSBACH-methane module considers the effects of permafrost thawing and freezing, and thus also the seasonal changes in the physical state of the soil on CH₄ processes. However, our analysis of the soil temperature profiles showed that during the cold season the simulated soil temperatures are nearly 10 °C lower than the values measured on site. Moreover, they gradually increase through spring and summer to reach values similar to the measurements. In contrast, the soil temperature seasonal cycle observed in the Chersky floodplain shows strong links to thawing and freezing processes (Göckede et al., 2017). These differences could be related to a negative bias in soil moisture content at the grid cell scale - which is driven by non-inundated areas - used to calculate the soil thermal regime. This limits the validity of the soil thermal properties as well as changes in latent heat. In addition, the carbon decomposition scheme used in this model version is driven by precipitation and atmospheric temperature. Therefore, actual changes in the soil temperature regime and wetness are not fully linked to the carbon dynamics. Finally, in addition to snow, near-surface vegetation in tundra environments (e.g., mosses and lichens) is also an effective thermal insulator of soil (Porada et al., 2016), regulating high surface temperatures in summer and cold temperatures during winter, and should be taken into account in a next version of the land surface model.

4.6 Role of non-growing season CH₄ emissions

In this work, we present, to our knowledge, the first simulated CH₄ emissions during the non-growing season with a land surface model at a regional scale. Our results show that changes in the snow layer depth control the temporal variation of the molecular diffusion of CH₄ through the snow. Our sensitivity studies corroborate that setting a thinner layer of snow as a threshold depth to switch to the CH₄ emission process during the cold season only promotes some changes in the partitioning of the methane flux among the four transport pathways. For example, a thinner snow layer promotes an earlier release of CH₄ than was otherwise emitted during late summer with a thicker layer of snow. However, the magnitude of the emissions through the snow is also determined by the amount of CH₄ that is produced, calculated from the decomposed carbon that is driven only by air temperature and precipitation in the carbon decomposition module. Changes in the physical properties of the snowpack (i.e., porosity and density) defined in the model have no clear effect on the timing of the emissions through the snow; this may lead to the conclusion that our choice of values for the capacity of snow to transport CH₄ was large enough. The physical restriction of gas transport via diffusive processes modeled by Fick's first law ensures that only physically possible rates of gas transport are being modeled.

In the recent work by Pirk et al. (2016), the authors demonstrated that the fluxes of CH₄ through the snowpack of permafrost Arctic wetlands during wintertime reflected a continuous emission of low amounts of gas still being produced in the soil, rather than solely the release of gas stored in the soil that was produced during the preceding growing season. These observations are in agreement with those from Mast et al. (1998), where the authors reported evidence of microbial activity throughout winter in subalpine soils permitted by the insulating effect of the snow layer. The results of Pirk et al. (2016) showed that there was no apparent sink or source of CH₄ within the snowpack, and their measurements captured a linear concentration gradient through the snow (Pirk et al., 2016). This observation validates the application of Fick's first law for diffusion of fluxes through the snow during winter, as applied in our model configuration. However, our formulation does not take into account the "pressure pumping" process reported by Massman et al. (1997) and Bowling and Massman (2011) that is related to the persistent advection of gas enhanced by wind through the snowpack. Based on isotopic analysis of CO₂ through the snowpack of a mountain forest, Bowling and Massman (2011) found that in the presence of wind, the pressure pumping effect contributed up to 11% of the total emissions during winter.

Our comparison at a grid cell scale to wintertime fluxes measured from EC at the Chersky floodplain (from January until March 2015 on average $3.8 \text{ mg CH}_4 \text{ m}^{-2} \text{ d}^{-1}$) shows that the modeled CH₄ emissions during this season (0.4 mg $CH_4 m^{-2} d^{-1}$) are consistently lower by about 1 order of magnitude. The measured EC fluxes are similar to other measurements with other methods from earlier studies. The work by Panikov and Dedysh (2000) showed winter methane emissions measured by chambers of about $5.0 \text{ mg CH}_4 \text{ m}^{-2} \text{ d}^{-1}$ from boreal peat bogs in western Siberia in mid-February. Pirk et al. (2016) measured CH₄ fluxes above the snowpack of about 2.4 mg $CH_4 m^{-2} d^{-1}$. In subalpine soils covered with snow, Mast et al. (1998) reported average winter CH₄ emissions of 4.4 mg CH₄ m⁻² d⁻¹ in moist soils calculated from samples collected through the snowpack. However, our modeled winter CH₄ emissions are comparable to those reported by Smagin and Shnyrev (2015) of about $0.6 \text{ mg CH}_4 \text{ m}^{-2} \text{ d}^{-1}$ measured by chambers during the coldest months of the year (February and March) in environments with different soil wetness in a West Siberian bog landscape.

Moreover, it is important to note that our results represent average values of a grid cell with a $0.5^{\circ} \times 0.5^{\circ}$ horizontal resolution, whereas measurements represent a much smaller spatial scale. Integrating the latter into the grid cell level must lead to an overestimation of the emission values at the grid cell level.

Other works have reported large CH_4 emissions from dry areas during the non-growing season. Using EC measurements, Zona et al. (2016) showed large fluxes from dry areas of the Alaskan tundra during the zero curtain period. Also, the findings of Mastepanov et al. (2013) imply that a portion of the active layer still remains free of ice during late autumn, and moisture and temperature changes are limited by the low thermal conductivity and heat capacity of dry soils. In the model, the consistently lower CH_4 emissions during winter can be explained by a low bias in soil temperature, leading to a low bias also in methanogenesis and larger oxidation within the topsoil.

5 Conclusion and outlook

The refined configuration of the JSBACH-methane model presented in this study has the ability to represent grid-cell-scale year-round CH₄ emissions at a comparable magnitude to those measured by chambers and EC in the same study area. The model was successfully applied to a regional domain in a floodplain of Northeast Siberia underlain by permafrost. The seasonal transition of the four CH₄ transport

pathways is mainly controlled by changes in the soil temperature, and only indirectly linked to soil moisture. The majority of the annual emissions take place through vascular plants. Given the relatively large scale of the model regional domain and uncertainties in the methane fluxes associated with forcing the model with reanalysis data and evaluating net emissions at the local level, it is difficult to quantify the emissions through the individual emission pathways. However, this explicit representation is necessary in processbased modeling, particularly for forecast regarding Arctic methane emissions under future climate scenarios.

The findings of this study demonstrate that to improve the understanding of the interannual variability of CH₄ fluxes from wetlands in boreal permafrost areas, to improve process model evaluation, and to improve the contribution of the individual emission pathways, more highly resolved temporal observational data are required, especially of year-round CH₄ EC fluxes and soil temperatures which are generally scarce and challenging for boreal and tundra areas. This is particularly important for improving modeling CH₄ emissions through snow, which in our model show a low bias when compared to EC measurements.

Finally, our model will greatly benefit from further improvements for regional simulations, which will also contribute to advancing the application to a global scale. In summary, the following model improvements are suggested: (1) a descriptive scheme for snow layer dynamics may benefit the simulation of wintertime CH₄ emissions, including pressure pumping effects due to advection of gas enhanced by wind; (2) improvements to prescribed model parameters such as soil depth until bedrock and initial soil moisture saturation, which are normally obtained from global-scale configurations of JSBACH; (3) an improved connection between the TOPMODEL approach for simulating the inundated fractions in a model grid cell, and soil state variables such as soil moisture, soil temperature, and ice content. This in turn might lead to improvements in the soil thermal properties for dry versus wet areas, and to the representation of noninundated areas to understand the dynamics of sources and sinks of CH₄. This might be alleviated if sub-grid-scale heterogeneity is included in future model developments. (4) Finally, improving the temporal transitions and seasonality of the water table levels will help to better constrain the surface heterogeneity of hydrologic responses to permafrost thaw and the spatial distribution of carbon decomposition.

Code and data availability. The JSBACH land surface model used in this study is intellectual property of the Max Planck Society for the Advancement of Science, Germany. The JSBACH source code is distributed under the Software License Agreement of the Max Planck Institute for Meteorology and can be accessed on personal request. The steps to gain access are explained under the following link: http://www.mpimet.mpg.de/en/science/models/ license/, last access: April 2018. The EC data set is available through the European Fluxes Database Cluster (site code: RU-Ch2). The chamber flux data are available upon request to Mathias Göckede (mgoeck@bgc-jena.mpg.de). The model data is also available upon request to Karel Castro Morales (kcastro@bgcjena.mpg.de).

Appendix A: Details on the in situ flux observation program

A1 Uncertainty assessment in EC flux data

The uncertainty analysis for the EC flux data followed procedures well established in the literature (Aubinet et al., 2012), and was split into random and systematic errors. The largest sources of random errors are associated with the turbulent sampling and instrument issues. These errors were quantified for each 30 min flux value through the TK3 flux processing software (Mauder and Foken, 2015). Errors related to footprint uncertainties were not quantified, since there are no major transitions in biome types within the core areas of the flux footprints.



Figure A1. Accumulated source-weight function for the EC tower in a control area within the Chersky study site, based on data from the growing season (mid-June–mid-September) in 2014. Solid white isolines indicate the 80, 60, 40, and 20% levels; the dashed line is the 10% level. Background colors indicate aggregated land cover classes based on WorldView-2 data.

Systematic errors can be introduced by unmet theoretical assumptions and methodological challenges, as well as by instrument calibration and data processing issues. To minimize this error, the instruments in the Chersky area were maintained and calibrated on a regular basis. Data intercomparison with a second EC tower located 600 m away from the tower that is the source of the data presented here yielded no systematic offset in the frequency distributions of wind speed, sonic temperature, and methane mixing ratios between the two towers. The TK3 software package contains all the required conversions and corrections for the flux data pro-

cessing, and yielded good agreement in a comparison with EddyPro (Fratini and Mauder, 2014). To avoid methodological issues that may bias flux data results, we employed a rigid post-processing quality control and flagging system scheme based on well-established analyses for stationarity and welldeveloped turbulence (Foken and Wichura, 1996), followed by additional tests to flag implausible data points in the resulting flux time series. Further details on this analysis are presented in Kittler et al. (2017).

No u^* -threshold was applied to the flux data set, since we determined that the stationarity of the signal and integral turbulence characteristics are also for nighttime conditions. This information facilitates identification of data sets with regular turbulent exchange, also during stable stratification, therefore producing fewer gaps compared to a bulk exclusion of data during stable nighttime stratification through the u^* filter method. After filtering out low-quality fluxes, the data coverage of methane fluxes was 86 % during the growing season and 67 % during the winter from the original full 30 min flux data set (Kittler et al., 2017). To produce a continuous flux record for quantification of long-term CH₄ budgets, the remaining gaps in the data were filled by averaging the existing flux data within a moving window of 10-day length centered on the gap. Uncertainties for gap-filled values were quantified as a standard deviation within the corresponding window, similar to the definition of gap-filling uncertainties for the CO₂ flux via the marginal distribution sampling routine of Reichstein et al. (2005).

To produce aggregated uncertainty values for longer time periods, we applied the procedures suggested in Rannik et al. (2016). All random errors were combined by considering them as independent variables that normally decrease with the length of the averaging period. Averaged over 2014 and 2015, the CH₄ flux uncertainty based on the 30 min data is 7.4 ± 8.3 nmol m⁻² s⁻¹, a result comparable to the 4.7 ± 3.8 nmol m⁻² s⁻¹ reported for a fen ecosystem by Jammet et al. (2017).

A2 Source-weight function of the EC flux data and scaling for model flux evaluation

We conducted a source-weight analysis (i.e., footprint analysis) to determine the fractional contribution of different land cover types within the field of view of the EC flux tower. Source-weight functions for each 30 min flux measurement were computed based on the Lagrangian stochastic footprint model of Rannik et al. (2003). Footprints were accumulated, analyzed, and interpreted using an approach presented by Göckede et al. (2006, 2008). We projected these footprints onto a land cover map from WorldView-2 with 2 m horizontal resolution (Fig. A1). In the context of the presented study, we aggregated the originally identified 22 land cover classes into 9 classes to concentrate on the dominant elements of the vegetation community structure.



Figure A2. (a) Non-scaled 10-day mean EC methane fluxes representing emissions from 26 % of a wet area in the footprint between June and August 2014 (black line), the 10-day mean EC methane fluxes scaled to the 10-day mean inundated fraction (IF) from the model for the same period of time (red line), and 10-day mean model methane emissions for grid cell A, which imply emissions from the IF from the model (blue line). Error bars in all lines are 1 standard deviation of the 10-day mean flux values; (b) shows the 10-day mean IF from the model used to scale the EC fluxes (blue line) and the constant wetness percentage of the footprint area calculated from the vegetation coverage remote sensing images (i.e., 26 %).

Since the EC tower is situated on a slightly elevated patch of tundra, tussocks and shrubs featuring various levels of wetness (red and orange colors in Fig. A1) dominate the immediate surroundings. Even though inundated parts of the study area, in this case identified by the prevalence of the cotton grass Eriophorum angustifolium (blueish colors in Fig. A1), dominate the area encircled by the 10% isoline that is used here to mark the boundary of the cumulative footprint area, they are mostly present in the outer reaches, therefore combining just about 26 % of the total flux signal sampled by the eddy system. Another 31 % is contributed by wet to moist tussock tundra with some shrubs. Overall coverage fractions within the major wetness categories remain approximately constant between tower footprint and two larger regions covered within the same WorldView-2 data set, indicating that this composition of wetness levels is typical for the Kolyma floodplain ecosystems analyzed within the context of this study. Furthermore, C3 grasses cover 33.3 % of model grid cell A, whereas the inundated fraction in that same grid cell ranges between 17.7 % and 19.9 %, calculated as a 10-day mean during June, July and August 2014. Thus, to improve the comparison between EC and model CH₄ fluxes, we corrected the 10-day EC mean fluxes (related to 26% of the footprint wet cotton grass area) through a linear scaling to obtain the fluxes from a smaller wet area, i.e., corresponding to the 10-day-mean inundated fraction of the model grid cell. The results of this scaling exercise are shown in Fig. A2. The non-scaled 10-day EC fluxes for the period of analysis are on average 65 mg CH₄ m⁻² d⁻¹, and the scaled fluxes decreased on average by 24% (mean 49.6 mg CH₄ m⁻² d⁻¹) after a correction by considering a smaller wet area within the footprint, reaching a magnitude similar to the 10-day-mean fluxes from the model (48.6 mg CH₄ m⁻² d⁻¹). This exercise emphasizes that wetness is the dominant control for total methane emissions in these ecosystems.

A3 Flux chamber observations

As shown in the study of Kwon et al. (2016), in the Chersky site were located two transects of 10 permanently installed PVC collars for flux chamber measurements. With distances of approximately 25 m between individual microsites, both



Figure A3. Daily methane flux rates (a) aggregated from flux chamber measurements within the growing season of 2014. Measurements are separated into drained (one wet microsite, nine dry microsites) and control (eight wet microsites, two dry microsites) transects; (b) flux rates against the water table at each microsite. Dry plots had a water table at or below the surface (up to 10 cm), whereas wet plots had a water table at or above the surface.

transects cover a distance of ~ 225 m within a drained and control section in this area. Site locations were selected quasi-randomly to reflect the dominant microsite characteristics (e.g., vegetation composition, wetness level) that were observed at each of the target locations. With a chamber footprint of 60 cm × 60 cm, this technique allowed study of microsites with rather homogeneous environmental conditions, as compared to the EC fluxes with often heterogeneous footprint areas. Details on the chamber program, overall methane flux rates observed, and functional relationships with, e.g., soil temperature, vegetation, and wetness levels are provided in Kwon et al. (2017).

Figure A3a displays average flux rates for wet and dry microsites observed within drained and control transects during sampling campaigns in summer 2014 (Kwon et al., 2016). These results demonstrate that methane release rates were virtually zero in the absence of standing water. At some of the dry microsites, defined by having the water table below the surface (on average up to 10 cm), slightly negative CH₄ flux rates were predominantly observed (mean of 3 mg CH₄ m⁻² d⁻¹) and almost negligible emissions, indicating the oxidation of methane (uptake) under highly aerobic conditions. Thus, the methane emissions in this tussock tundra ecosystem of Northeast Siberia take place predominantly in wet areas.

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