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Divergent patterns of experimental and model-derived permafrost ecosystem carbon dynamics in response to Arctic warming

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Abstract

In the last few decades, temperatures in the Arctic have increased twice as much as the rest of the globe. As permafrost thaws in response to this warming, large amounts of soil organic matter may become vulnerable to decomposition. Microbial decomposition will release carbon (C) from permafrost soils, however, warmer conditions could also lead to enhanced plant growth and C uptake. Field and modeling studies show high uncertainty in soil and plant responses to climate change but there have been few studies that reconcile field and model data to understand differences and reduce uncertainty. Here, we evaluate gross primary productivity (GPP), ecosystem respiration (R_{eco}), and net ecosystem C exchange (NEE) from eight years of experimental soil warming in moist acidic tundra against equivalent fluxes from the Community Land Model during simulations parameterized to reflect the field conditions associated with this manipulative field experiment. Over the eight-year experimental period, soil temperatures and thaw depths increased with warming in field observations and model simulations. However, the field and model results do not agree on warming effects on water table depth; warming created wetter soils in the field and drier soils in the models. In the field, initial increases in growing season GPP, Reco, and NEE to experimentally-induced permafrost thaw created a higher C sink capacity in the first years followed by a stronger C source in years six through eight. In contrast, both models predicted linear increases in GPP, Reco, and NEE with warming. The divergence of model results from field experiments reveals the role subsidence, hydrology, and nutrient cycling play in influencing the C flux responses to permafrost thaw, a complexity that the models are not structurally able to predict, and highlight challenges associated with projecting C cycle dynamics across the Arctic.

1. Introduction

Permafrost soils contain 1460–1600 Pg (10^{15} g) carbon (C), a stock roughly double in size to the C presently in the Earth's atmosphere (Hugelius *et al* 2014, Schuur

et al 2015, Strauss *et al* 2017). Continuously frozen soils keep the C stocks buried in permafrost, which is widespread across an area that covers 24% of the terrestrial Northern hemisphere (Brown *et al* 2002). Over the last few decades, the Arctic has been

experiencing rapid increases in temperature (Huang et al 2017, Richter-Menge et al 2017), and frozen ground conditions are projected to become less stable in the coming decades (Romanovsky et al 2010, 2012). Warming and drying threaten permafrost in upland ecosystems as a result of increased natural drainage as the water table descends (Hinzman et al 2005, Liljedahl et al 2016). In low lying areas the reverse occurs, and ground subsidence caused by permafrost thaw can create anoxia as the topography and the underlying permafrost prevent runoff. Once thawed, organic matter is vulnerable to decomposition, and large amounts of C from permafrost ecosystems are projected to be released to the atmosphere in the form of carbon dioxide (CO₂) and methane (e.g. Schädel et al 2014, Schuur et al 2015). Any additional release of C to the atmosphere has the potential to amplify climate change and the permafrost C feedback, although the magnitude of the response is highly uncertain.

The representation of permafrost C dynamics in process-based Earth system models remains a crucial challenge in projecting potential C cycle-climate feedbacks (McGuire et al 2018). Manipulative experiments provide opportunities to inform these longterm projections about the magnitude, and underlying mechanisms of permafrost C dynamics in a changing Arctic (Oberbauer et al 1992, Mauritz et al 2017, Christiansen et al 2018). Chamber based field CO₂ flux measurements represent the balance of gross primary productivity (GPP) and ecosystem respiration (R_{eco}) and document gains and losses of C in response to environmental change. Experimental soil warming in the field can be achieved by installing snow fences that passively warm the soil during winter months (Dorrepaal et al 2009, Natali et al 2011, Johansson et al 2013, Leffler et al 2016, Christiansen et al 2018). This type of passive warming is distinct from air warming and active soil warming techniques, which are not covered in this manuscript.

While field-based experiments provide detailed information on process-based mechanisms and smallscale variations in C dynamics in response to warming, they are limited in space and time and rarely extend beyond a decade. Process-based models have been used for circumpolar projections of near-surface permafrost degradation (Lawrence and Slater 2005, Lawrence et al 2012, Slater and Lawrence 2013, Koven et al 2013a) and C dynamics under various scenarios of future warming (Koven et al 2011, Schaefer et al 2011, MacDougall et al 2012, Schaphoff et al 2013, Schneider von Deimling et al 2012, 2015, Koven et al 2015b, Burke et al 2012, 2017). Differences in modeled permafrost extent and C dynamics among models are large and disagree on estimation of present-day permafrost extent, the size of the permafrost C pool, and the timing and amount of permafrost C release and plant C uptake. A recent model intercomparison project assessed permafrost degradation, changes in soil



and vegetation C, and whole ecosystem C for low and high emissions pathways (Representation Concentration Pathways, RCP4.5 and RCP8.5, (McGuire *et al* 2018)). For most models, net loss of ecosystem C would not occur until after 2100 as vegetation gain would offset soil C losses until then. By 2300, soil C losses exceeded gains in vegetation C but variation among models was large. The lack of agreement across models is striking and implies that new ways to evaluate and improve the underlying biogeophysical and biogeochemical processes in these models is required.

One method towards improving model performance is to utilize field experimental data to examine model responses to environmental perturbations. Here, we use eight years of field-based measurements of CO₂ dynamics from a unique snow-fence experimental field warming study, in which permafrost is experimentally degraded in an upland tundra site, and compare it with Community Land Model (CLM) simulations that replicate the field experiment. Using results from grid cells that broadly represent active layer thickness (ALT) and productivity in control plots, we evaluate the modeled CO2 response to warming and compare it to equivalent results from the field experiment. The objectives of this study are to: (1) compare the magnitude and pattern of environmental variables in CLM4.5 and 5.0 relative to experimental warming in the field; (2) evaluate annual and seasonal responses of ecosystem CO₂ dynamics that are simulated by the model after eight years of warming, and (3) identify key drivers responsible for variation and bias between field and model responses.

2. Methods

2.1. Field experiment

In this field-to-model comparison, we used eight years of data (data citations are provided in the supplementary material, available online at stacks.iop.org/ERL/13/ 105002/mmedia) from an experimental field warming study that was established in the fall of 2008 (Natali et al 2011). The Carbon in Permafrost Experimental Heating Research site is situated in upland moist acidic tundra in the Eight Mile Lake Watershed $(-149.23^{\circ}W)$, 63.88°N, 670 m.a.s.l.) and is completely underlain by permafrost (Osterkamp et al 2009, Schuur et al 2009). The area is characterized by degrading permafrost and a mean annual air temperature of -0.94 °C from 1977 to 2015 (Healy and McKinley Stations, Western Regional Climate Center and NOAA National Centers for Environmental Information). The vegetation is dominated by the graminoid Eriophorum vaginatum, mosses, vascular plants, and deciduous shrubs (Schuur et al 2007, Natali et al 2011, Deane-Coe et al 2015, Salmon et al 2016). Experimental warming is imposed by installing six snow fences that accumulate snow, insulate the ground and increase surface and deep soil temperatures during the winter and into the spring. The unique

feature of this experimental setup is the springtime removal of the additional snow prior to melt out. Snow removal from the warmed plots prevents a delayed start of the growing season and avoids additional water input. Details of the experimental design can be found in Natali *et al* (2011); numerous publications address the effects of warming and permafrost degradation on aboveground biomass (Deane-Coe *et al* 2015, Salmon *et al* 2016), nitrogen (N) availability (Salmon *et al* 2016), vegetation phenology (Natali *et al* 2012), and gaseous C fluxes (Natali *et al* 2014, Mauritz *et al* 2017).

Net ecosystem exchange (NEE) was measured between May and September using an automated chamber system, and CO₂ concentrations were measured continuously using an infrared gas analyzer (LI-820, LICOR Corp., Lincoln Nebraska). We obtained net NEE values by gap-filling using a hyperbolic light-response curve, and Reco was modeled using an Arrhenius-type temperature response and soil temperature at 10 cm. Gap-filling was used to fill in missing data and to estimate Reco under low light conditions (when photosynthetically active radiation was $<5 \,\mu$ mol m² s⁻¹; details can be found in Mauritz *et al* (2017)). GPP was calculated as $GPP = NEE - R_{eco}$, where positive NEE values denote a CO₂ sink. Annual NEE was estimated by using the average of two nongrowing season Reco models developed by Webb et al (2016) for the non-growing season months (October-April).

Aboveground plant biomass during the first five years of the field experiment is calculated using a point intercept method and site-specific allometries based on harvests performed at this site (Salmon *et al* 2016). No direct measurements of aboveground biomass were performed for the last three years of the field experiment.

2.2. Model experiments

Two versions of CLM were used for the model simulations, CLM4.5 (Oleson et al 2013) and CLM5.0 (http://www.cesm.ucar.edu/models/cesm2/land/). CLM4.5 includes processes that permit simulation of critical thermal, hydrologic, and biogeochemical processes that are relevant for permafrost and its response to climate change (Koven et al 2013b, Lawrence and Slater 2005, Lawrence et al 2008, Riley et al 2011, Swenson et al 2012). CLM5.0 includes numerous changes and additional processes, including, critically for this comparison, a reformulation of the coupling between C and N cycles that in principle permits plants to uptake N during periods when they are not photosynthesizing. For each model, a control and snow-fence simulation was performed at 0.5 degree resolution for Alaska and Western Canada (Southwest bound: 58°15'N, 179°45'W; Northeast bound: 84°45'N, 95°15'W, figure S1) with transient atmospheric forcing from the GSWP3 version 1 climate reanalysis http:// hydro.iis.u-tokyo.ac.jp/GSWP3/. Simulations were spun up to steady state conditions by cycling over



atmospheric conditions from 1901-1920. Subsequently, transient control simulations followed the typical land only protocols over the 20th century (Oleson et al 2013, Koven et al 2013b). In the snow-fence simulation, the effects of the field snow-fence on ground temperature were replicated by decreasing the snow thermal conductivity by 50%, which effectively increases the snow insulation and warms the ground without affecting the water budget of the model. The reduced snow thermal conductivity was used for the period September through March. On April 1, the snow conductivity was reverted to its standard value, following the snow removal in the field experiment. To best match the vegetation characteristics at the field site, all of the grid cells in the domain used the same Arctic C3 grass plant functional type (PFT), which is the CLM PFT that represents tundra.

Analyses centered on edaphic conditions and C fluxes during the first eight years of warming. We focused data analyses and model evaluations on model grid cells that experience similar environmental and biological conditions as the field site location (figure S1). Details on grid cell selection can be found in the supplement.

2.3. Variables

We selected variables from all model simulations that were equivalent to variables measured in the field experiment (table 1). Model output is monthly and the measurement frequency for field data varies from halfhourly (soil temperatures and C fluxes, table 1) to multiple times a week (thaw depth, water table depth, and leaf area index) to once a year (e.g. snow depth, aboveground biomass, and leaf chemistry). For seasonal variation, we averaged monthly values over eight years and for trends over time we averaged values over all months for a given year (soil temperature) or only over the growing season months (water table depth). Cumulative CO₂ fluxes were calculated by summing half-hourly data to monthly values to match model output. Field and model GPP, Reco, and NEE were additionally summed to growing season periods. We also estimated an annual cumulative NEE value that accounts for respiration during the non-growing season (October-April, details are provided in the field data section).

3. Results

3.1. Environmental variables

During the experimental period (2009–2016), mean annual soil temperature at 10 cm depth in the field control plots was 0.41 ± 0.24 °C (figure 1(a)). Modeled annual soil temperatures at 10 cm in the control simulations were colder (-3.79 ± 0.19 and -5.29 ± 0.2 in CLM4.5 and CLM5.0 respectively), which was to be expected given that the selected grid cells were further north than the field site location

Table 1. Environmental variables, carbon dynamics, and plant related variables measured in the field experiment and extracted from model simulations.

	Variable ^a	Description	CLM variable	Unit ^a		
				Field	Model	$Frequency (field)^{b}$
Environmental	Soil temperature	10 cm depth	TSOI_10 CM,	°C	К	Half-hourly
	Thaw depth	Active layer thickness	ALT	cm	m	Multiple times per week during growing season
	Water table depth		ZWT_PERCH	cm	m	Multiple times per week during growing seasor
	Snow depth	Snow height	SNOWDP	m	m	$1 \mathrm{x} \mathrm{yr}^{-1}$ (March or April)
Carbon fluxes	GPP	Gross primary productivity	GPP	$g CO_2$ - $C m^{-2} half$ -hour ⁻¹	$g CO_2 - C m^{-2} s^{-1}$	Half-hourly
	R _{eco}	Ecosystem respiration	ER	gCO_2 -C m ⁻² half-hour ⁻¹	gCO_2 -Cm ⁻² s ⁻¹	Half-hourly
	NEE	Net ecosystem exchange	NEE	gCO_2 -C m ⁻² half-hour ⁻¹	gCO_2 -Cm ⁻² s ⁻¹	Half-hourly
Plant related	Biomass ^c	Aboveground peak biomass	LEAFC	$dry g m^{-2}$	gCm^{-2}	$1 \mathrm{x} \mathrm{yr}^{-1} (\mathrm{July})$
	Foliar carbon		LEAFC	%	gCm^{-2}	$1 \mathrm{x} \mathrm{yr}^{-1}$ (July)
	Foliar nitrogen		LEAFN	%	gNm^{-2}	$1 \mathrm{x} \mathrm{yr}^{-1}$ (July)
	LAI ^d	Leaf area index	TLAI	Unitless	Unitless	\sim biweekly during growing season

^a Units for field and model data were corrected for analyses to match each other.

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^b In the field, measurements were taken each year from 2009 to 2016 except for biomass, which was only measured from 2009 to 2013, and LAI, which was measured in 2011 and from 2013 to 2016. Model data are available for each month and year.

^c Above ground peak biomass in the model was calculated as leaf C multiplied by 2 assuming plant biomass consists of 50% C. ^d LAI estimated for field was based on parameters and equations from Street *et al* (2007). LAI = $a^*e^{(b*NDVI)}$; a = 0.0148, b = 6.192, parameters and equation from Street *et al* (2007), NDVI measured in the field.





Figure 1. Environmental variables for eight years of warming for field and model data. Mean annual values for soil temperature at 10 cm depth are shown in (a), active layer thickness in (e), growing season water table depth in (i). Mean monthly values averaged over eight years are shown in panels (c), (g), and (k). Note that (e) and (f) show active layer thickness and (g) and (h) thaw depth. Bar graphs show the difference between warming minus control for annual data in (b), maximum in (j), and monthly in (d), (h), and (l). For field and modeled data, black dots and lines represent control and red represents warming. Values are means of six field replicates (snow fences), 75 grid cells in CLM4.5, and 42 grid cells in CLM5.0. Error bars are standard errors in the field and standard deviation in the modeled data. Shaded areas show standard deviated for modeled data (light grey for control and light red for the warming treatment). Note different axis scaling between annual and seasonal panels.

(figure S1). The warming treatment increased soil temperature by 0.83 ± 0.13 °C in the field and 1.32 ± 0.07 in CLM4.5 and 1.40 ± 0.08 °C in CLM5.0 (figure 1(b), table S1). Soil warming occurred during winter months in both the field and models

and was strongest in March (figures 1(c) and (d), table S2).

At the beginning of the field experiment, ALT in control plots was 0.55 ± 0.01 (figure 1(e), table S1) and similar in the models as defined by the

geographical masking criteria. With each year of warming, ALT increased linearly but to a lesser extent in the models than in the field (figure 1(f)). Over the course of the growing season, field and model thaw depth showed similar trends (figure 1(g), table S2) although the difference in thaw depth between warming and control was greater in the field (figure 1(h)).

Averaged over eight years, mean growing season water table depth in the field was closer to the surface in the warmed plots $(-14.6 \pm 2.7 \text{ cm})$ than in the control plots $(-19.7 \pm 1.6 \text{ cm})$, and the warmed plots became wetter each year (figure 1(i)). In the models, control water table depth varied little over the eight years and was similar in magnitude to the field $(-12.6 \pm 0.5 \text{ cm} \text{ in CLM4.5}, \text{ and } -16.1 \pm 0.9 \text{ in CLM5.0}; figure 1(i), error represents variation across years) but unlike in the field, the warming treatment dried the soils out. After snow melt in May, modeled water table depth was similar to the field and then dropped each month until September with warmed soils in the models drying out more than the modeled control (figures 1(k) and (l), table S2).$

3.2. Carbon fluxes

Cumulative growing season (May-September) GPP increased in control plots over the experimental period in the field (204–492 g C m⁻² yr⁻¹, figure 2(a), table S1). Warmed plots showed stronger increases in GPP in the first four years of warming (2009-2013) compared to control plots and then declined in the second half of the experimental period. The largest relative difference between warming and control in the field was in 2011 with 57% higher GPP values in the warming treatment whereas after eight years GPP was 8% lower in warmed plot compared to control (figure 2(b), table S1). In the first year of simulations, modeled GPP was similar to field measurements as defined by the selection of grid cells. Warming had a gradual but smaller effect on GPP in both models than in the field. After three years of warming (2011), when the highest difference was found in GPP in the field, CLM4.5 showed an increase in GPP of 8% with warming and CLM5.0 an increase of 2%. Interannual patterns of growing season Reco in the field and models were comparable to GPP (figure 2(e) and (f), table S1). Seasonal GPP and Reco showed similar fluctuations in field and modeled data with peak fluxes occurring in July (figure 2(c) and (g), table S2).

Cumulative growing season NEE in field controls showed C uptake (7–84 g C m⁻² yr⁻¹) during the eight years of the experiment. Modeled controls started as small growing season C sources (CLM4.5–18 g C m⁻² yr⁻¹; CLM5.0–21 g C m⁻² yr⁻¹) and then became C sinks in the second year. In the field, the difference in growing season NEE between warmed and control plots increased until year five (2013: relative increase of +55%) whereas in year eight, NEE in warmed plots had declined to -105% of NEE in control plots (figure 2(i) and (j)).



In contrast, the models exhibited a gradual increase and linear NEE response to warming.

When accounting for non-growing season CO_2 dynamics, the field site was a net C source each year in both control and warmed plots (negative NEE, figure 2(m), table S1). Warmed plots were a smaller CO_2 source in years two to five in the field experiment, and in year six (2014) and onward, the warmed plots became a larger CO_2 source than the control plots (2016, control: -58 and warming: $-156 \text{ g C m}^{-2} \text{ yr}^{-1}$). Generally, modeled cumulative NEE for control and warming fluctuated around zero with a minimal difference between control and warming indicating that the ecosystem may be C neutral over the years (figure 2(m)).

3.3. Biomass and plant tissue responses

In the control plots, peak biomass in July ranged from 420 g dry weight m^{-2} in the first year to 582 g dry weight m^{-2} in year five of the experiment (2013). Though aboveground biomass increased in all plots during the first five years of the experiment, increases in the warmed plots were greater and by year five, warmed plots had 25% more aboveground biomass than controls (table S1). Aboveground peak biomass in the model is leaf C multiplied by two, assuming plant biomass consists of 50% C. In both models, peak biomass was less than half of what was estimated in the field (CLM4.5: 199 g dry weight m^{-2} in the first year, CLM5.0: 113 g dry weight m^{-2}) and CLM5.0 showed consistently lower values than CLM4.5 (table S1).

4. Discussion

Experimental warming increased soil temperature and ALT in both the field and models but this caused saturated soils in the field and drier soils in the models. These different responses in environmental variables to soil warming led to higher increases in GPP, R_{eco} , and NEE in the field in the first five years of warming and a decline in all CO₂ flux components after five years. In both models, warming caused a gradual but weaker increase in GPP, R_{eco} , and NEE that continued throughout the experimental period. These nonlinear patterns in CO₂ fluxes in the field experiment indicated that soil warming and permafrost thaw created water dependent dynamics that may increase the C source strength of the ecosystem, a trend that was not captured by either of the models.

4.1. Observed and modeled biogeophysical variables Arctic landscapes are very heterogeneous and soil warming can lead to warmer and wetter soils but also to warmer and drier soils as water table depth responses can be patchy. Throughout the field experimental period, we observed increases in annual soil temperatures and ALT in the control plots, which presumably is in response to the ongoing climate





change that is already occurring (Richter-Menge *et al* 2017). Modeled soil temperature at 10 cm was much colder during winter months than in the field, which led to colder annual temperatures in the models. The selection of grid cells was limited to those that had similar ALT and GPP values as in the field in year one and did not account for accurate initial soil temperatures. This approach ensured that the volume

of available soil C matched field conditions and matching GPP ensured comparable vegetation dynamics. Adding soil temperature to the mask for grid cell selection resulted in only a few grid cells in CLM4.5 and none in CLM5.0. It is unclear whether the mismatch between model mean soil temperatures and active layer predictions is due to model physics or whether the site-level environmental meteorological



conditions are not representative of the regional-scale atmospheric dynamics used to drive the model.

Over the course of the eight years of experimental warming, permafrost on the plot scale has resulted in wetting and drying as expected with widespread thaw at the landscape level (Osterkamp et al 2009). Ground ice content in the field accounts for $54\% (\pm 6\%)$ up to a depth of 1 m at the beginning of the experiment (in 2009) and cryogenic structures are porous ice closer to the surface and ice lenses occurring further down the profile. Total ice content may still be underestimated as sampling of soil cores excludes massive ice wedges. Not only did the model predict drier surface soils with warming as opposed to the field, it also predicted the water table depth to move further away from the surface during the growing season and at a higher rate in the warming treatment. The divergence in the models from the field is most likely due to the loss of ground ice in the field causing the ground to subside and the surface getting closer to the water table, a feature that is not represented in the model versions included in our analysis. Lee et al (2014) incorporated excess ice into soil layers of CLM4.5 and then projected the amount of surface subsidence that could occur. However, the subsidence that was calculated was only characteristic and the model hydrologic state was not strongly affected by the inclusion of excess ice. Thaw-induced soil drying as a result of increased drainage was also observed in previous modeling efforts (Lawrence et al 2015, Chadburn et al 2017). Ground surface subsidence generally exhibits a heterogeneous pattern depending on the amount of ground ice and high precipitation can enhance subsidence and move the water table closer to the surface (Mauritz et al 2017). The effects of subsidence have thus far not been considered in models meaning that we likely are missing an important dynamic in permafrost transitions.

An additional factor that could potentially contribute to the divergence in water table dynamics between field observations and model simulations could be discrepancies in soil hydraulic properties between field and model. The soil properties at the site are described as Aeolian silt but detailed soil characterizations are not available. In CLM, soil properties are specified in each grid cell based on the Harmonized World Soil Database. If the specified deep soil properties in the selected CLM grid cells do not reflect those at the site, the runoff and soil drying response to warming could differ. Without more detailed characterization of deep soil properties at the site, it is difficult to probe this question in detail, though we should note that we are effectively sampling CLM for several different soil types through our grid cell filtering method. The differences in water table depth between observed and modeled data demonstrates that accurate representation and prediction of near-surface hydrology in the Arctic remains a challenge in models complicating comparisons between field manipulation experiments and models.

4.2. Factors contributing to variable CO₂ dynamics with warming in the field

For accurate simulation of CO_2 dynamics to soil warming, we need to consider which environmental and biological factors are linked to the increase in growing season CO_2 flux components in the first five years of the experiment followed by a decrease. Growing season fluxes provide the best resolution of field CO_2 fluxes as non-growing season CO_2 dynamics rely on models and gap-filling and our understanding of non-growing season fluxes remains limited. In contrast, growing season GPP is directly linked to photosynthesis and can be easily compared between field and models.

The GPP response as observed in the field could be driven by two primary factors: active layer deepening and subsequent increase in N availability. Deepening of the active layer with soil warming expands the rooting zone and increases the space for vertical root distribution where plants access nutrients and water (Iversen et al 2014). Thaw of deep permafrost soil not only increases the potential rooting zone for tundra plants but also releases biologically available N that is accessible to support further plant growth (Keuper et al 2012, Salmon et al 2016, Keuper et al 2017, Wild et al 2018). Plant growth in moist acidic tundra is often N-limited (Shaver et al 1992, Chapin et al 1995, Mack et al 2004, Sistla et al 2012) and at the field experiment higher aboveground plant biomass during the initial thaw stages has been linked to increasing N availability (Salmon et al 2016). In addition, thaw and changing N availability may promote changes in species composition, and field manipulations suggest there will be a shift to greater shrub dominance with warming over longer time scales (e.g. Chapin et al 1995, Sistla et al 2013, DeMarco et al 2014). At the field site however, the observed increase in GPP during the first five years of warming induced by thaw was driven by productivity of the already dominant sedge, E. vaginatum (Salmon et al 2016), which is consistent with the transient response observed in Sweden by Johansson et al (2013).

The decline in GPP, R_{eco} , and NEE in the field after five years of warming demonstrates that a deepening of the ALT and increasing N availability cannot drive unlimited GPP increases. Initial thaw might allow a rapid response of existing dominant species but over longer timeframes different drivers may dominate. The loss of ground ice resulting in soil subsidence and a water table position closer to the surface may favor plant species capable of dealing with more water-logged conditions. Unpublished data from the latest field season (year nine of the field experiment, Taylor *et al* 2017) show that *E. vaginatum* declines in wet areas possibly explaining the decline in GPP which turns the ecosystem into an annual C source with continued warming.



4.3. Model divergence from field observations

Increases in GPP in response to warming are strongly attenuated in both models compared to the field experiment. Both ALT and N limitation can be identified as main causes for the weaker increase in GPP with warming in the models. ALT increased less than 15% in both models in year eight of soil warming compared to a 55% increase in the field which added more depth for root growth, water uptake and nutrient acquisition in the field. Smaller increases in ALT with warming in the model are the result of colder winter temperatures in the model compared to the field site. Further, N limitation of Arctic plant productivity may not be well represented in either of the models. This is indicated in CLM4.5 by a lack of change in the fraction of potential GPP (FPG) with warming over time as well as a very weak response in net N mineralization with warming (table S3). In CLM5.0, nutrient limitation shows almost no change with warming and increases in net N mineralization with warming were less than 10% over the years (table S3). Additional N availability with depth was shown to have a small effect on plant productivity in CLM4.5 (Koven et al 2015a), which further emphasizes insufficient representation of N limitation in CLM4.5. Nutrient cycling along with aboveground and belowground coupling and microbial dynamics were also identified as inadequately represented processes for high latitude ecosystems in CLM4.5 (Bouskill et al 2014). Modifications to the plant N cycle in CLM5.0 aimed to improve the realisms of vegetation N cycling (Shi et al 2015, Ali et al 2016, Ghimire et al 2016); (http://www.cesm. ucar.edu/models/cesm2/land/). While aspects of the terrestrial N cycle are improved with these developments, results here show a surprising low level of N limitation in arctic ecosystems simulated by both versions of CLM (table S3). Thus, although warming stimulates modest changes in N mineralization rates, given low plant nutrient limitation, this elicits negligible changes in plant productivity. This suggests that aspects of the soil biogeochemical code and considerations of plant-microbial competition for N deserve further revisions in the CLM. Improving nutrient limitation to better capture future changes in the C balance has also been identified for other models (Chadburn et al 2017) and provides a key process that can improve model performance.

5. Conclusion

Models are an important tool to project future C dynamics in the Arctic, as well as globally. To obtain meaningful projections for the global C cycle, we need to ensure that the relevant processes in models are represented and validated against field observations. In this study, the models are able to capture some but not all aspects of an experimental warming. Both versions of CLM predict that the response to warming is gradual and increases in GPP, R_{eco} , and NEE are small with warming, and that short-term ecosystem C gains give way to medium-term C neutrality. This is in contrast to the field observations, which showed that initial increases in C uptake subsequently reverse to C losses, such that ecosystem C losses exceeded C gains after eight years of experimental warming. Short-term responses as observed in the first five years of the field experiment are not indicative of longer-term responses requiring manipulative experiments to continue beyond a decade.

This field and model comparison provides insights into the underlying causes for model divergence of ecosystem responses to soil warming on a decade-long timescale. The divergences between field and model responses to warming demonstrate the importance of thaw-induced changes in surface hydrology and N limitation on C dynamics and that it will be critical to accurately represent these processes in future model projections.

In this data-model comparison, we have identified important steps for model improvement and as a next step it will be crucial to expand site level comparisons between observational and model data to global simulations. The complexity of interacting processes in models will need to be carefully evaluated and accounting for the spatial heterogeneity in tundra landscapes and the proportion of wet and dry surface soils on the circumpolar scale will be challenging but important elements to consider.

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References

Ali A A *et al* 2016 A global scale mechanistic model of photosynthetic capacity (LUNA V1.0) *Geosci. Model Dev.* 9 587–606

Bouskill N J, Riley W J and Tang J Y 2014 Meta-analysis of highlatitude nitrogen-addition and warming studies implies ecological mechanisms overlooked by land models *Biogeosciences* 11 6969–83

Brown J, Ferrians O J J, Heginbottom J A and Melnikov E S 2002 Circum-Arctic Map of Permafrost and Ground-Ice Conditions, version 2nd edn (National Snow and Ice Data Center) https:// nsidc.org/data/ggd318

Burke E J, Ekici A, Huang Y, Chadburn S E, Huntingford C, Ciais P, Friedlingstein P, Peng S and Krinner G 2017 Quantifying uncertainties of permafrost carbon–climate feedbacks *Biogeosciences* 14 3051–66

Burke E J, Hartley I P and Jones C D 2012 Uncertainties in the global temperature change caused by carbon release from permafrost thawing *Cryosphere* 6 1063–76

- Chadburn S E *et al* 2017 Carbon stocks and fluxes in the high latitudes: using site-level data to evaluate Earth system models *Biogeosciences* 14 5143–69
- Chapin F S I, Shaver G R, Giblin A E, Nadelhoffer K J and Laundre J A 1995 Responses of arctic tundra to experimental and observed changes in Climate *Ecology* 76 694–711

Christiansen C T, Lafreniére M J, Henry G H R and Grogan P 2018 Long-term deepened snow promotes tundra evergreen shrub growth and summertime ecosystem net CO₂ gain, but reduces soil carbon and nutrient pools *Glob. Change Biol.* **24** 3508–25

Deane-Coe K, Mauritz M, Celis G, Salmon V, Crummer K, Natali S and Schuur E G 2015 Experimental warming alters productivity and isotopic signatures of tundra mosses *Ecosystems* 18 1070–82

DeMarco J, Mack M C, Bret-Harte S M, Burton M and Shaver G R 2014 Long-term experimental warming and nutrient additions increase productivity in tall deciduous shrub tundra *Ecosphere* 5 art72

Dorrepaal E, Toet S, van Logtestijn R S P, Swart E, van de Weg M J, Callaghan T V and Aerts R 2009 Carbon respiration from subsurface peat accelerated by climate warming in the subarctic *Nature* **460** 616–U79

Ghimire B, Riley W J, Koven C D, Mu M and Randerson J T 2016 Representing leaf and root physiological traits in CLM improves global carbon and nitrogen cycling predictions *J. Adv. Model. Earth Syst.* **8** 598–613

Hinzman L D *et al* 2005 Evidence and implications of recent climate change in northern Alaska and other arctic regions *Clim. Change* 72 251–98

Huang J *et al* 2017 Recently amplified arctic warming has contributed to a continual global warming trend *Nat. Clim. Change* **7** 875–9

Hugelius G *et al* 2014 Estimated stocks of circumpolar permafrost carbon with quantified uncertainty ranges and identified data gaps *Biogeosciences* **11** 6573–93

Iversen C M, Sloan V L, Sullivan P F, Euskirchen E S, McGuire D A, Norby R J, Walker A P, Warren J M and Wullschleger S D 2014 The unseen iceberg: plant roots in arctic tundra New Phytologist 205 34–58

Johansson M, Callaghan T V, Bosiö J, Åkerman J H, Jackowicz-Korczynski M and Christense T 2013 Rapid responses of permafrost and vegetation to experimentally increased snow cover in sub-arctic Sweden *Environ. Res. Lett.* 8 35025 Keuper F, Bodegom P M, Dorrepaal E, Weedon J T, Hal J, Logtestijn R S and Aerts R 2012 A frozen feast: thawing permafrost increases plant-available nitrogen in subarctic peatlands *Glob. Change Biol.* 18 1998–2007

Keuper F, Dorrepaal E, van Bodegom P M, van Logtestijn R, Venhuizen G, van Hal J and Aerts R 2017 Experimentally increased nutrient availability at the permafrost thaw front selectively enhances biomass production of deep-rooting subarctic peatland species *Glob. Change Biol.* 23 4257–66

Koven C D, Riley W J and Stern A 2013a Analysis of permafrost thermal dynamics and response to climate change in the CMIP5 earth system models *J. Clim.* **26** 1877–900

Koven C D, Riley W J, Subin Z M, Tang J Y, Torn M S, Collins W D, Bonan G B, Lawrence D M and Swenson S C 2013b The effect of vertically resolved soil biogeochemistry and alternate soil C and N models on C dynamics of CLM4 *Biogeosciences* 10 7109–31

Koven C D, Ringeval B, Friedlingstein P, Ciais P, Cadule P, Khvorostyanov D, Krinner G and Tarnocai C 2011 Permafrost carbon-climate feedbacks accelerate global warming *Proc. Natl Acad. Sci.* **108** 14769–74

Koven C D, Lawrence D M and Riley W J 2015a Permafrost carbon —climate feedback is sensitive to deep soil carbon decomposability but not deep soil nitrogen dynamics *Proc. Natl Acad. Sci.* **112** 3752–7

Koven C D *et al* 2015b A simplified, data-constrained approach to estimate the permafrost carbon–climate feedback *Phil. Trans. R. Soc.* A **373** 20140423

Lawrence D M, Slater A G, Romanovsky V E and Nicolsky D J 2008 Sensitivity of a model projection of near-surface permafrost degradation to soil column depth and representation of soil organic matter J. Geophys. Res.-Earth Surf. **113** F02011

Lawrence D M, Koven C D, Swenson S C, Riley W J and Slater A G 2015 Permafrost thaw and resulting soil moisture changes regulate projected high-latitude CO₂ and CH₄ emissions *Environ. Res. Lett.* **10** 94011

Lawrence D M and Slater A G 2005 A projection of severe nearsurface permafrost degradation during the 21st century *Geophys. Res. Lett.* **32** L24401

Lawrence D M, Slater A G and Swenson S C 2012 Simulation of present-day and future permafrost and seasonally frozen ground conditions in CCSM4 J. Clim. 25 2207–25

Lee H, Swenson S C, Slater A G and Lawrence D M 2014 Effects of excess ground ice on projections of permafrost in a warming climate *Environ. Res. Lett.* **9** 124006

Leffler A J, Klein E S, Oberbauer S F and Welker J M 2016 Coupled long-term summer warming and deeper snow alters species composition and stimulates gross primary productivity in tussock tundra *Oecologia* 181 287–97

Liljedahl A K *et al* 2016 Pan-arctic ice-wedge degradation in warming permafrost and its influence on tundra hydrology *Nat. Geosci.* **9** 312–8

MacDougall A H, Avis C A and Weaver A J 2012 Significant contribution to climate warming from the permafrost carbon feedback *Nat. Geosci.* 5719–21

Mack M C, Schuur E A G, Bret-Harte M S, Shaver G R and Chapin F S 2004 Ecosystem carbon storage in arctic tundra reduced by long-term nutrient fertilization *Nature* **431** 440–3

Mauritz M, Bracho R, Celis G, Hutchings J, Natali S M, Pegoraro E, Salmon V G, Schädel C, Webb E E and Schuur E A G 2017 Nonlinear CO₂flux response to 7 years of experimentally induced permafrost thaw *Glob. Change Biol.* **23** 3646–66

McGuire A D *et al* 2018 The dependence of the evolution of carbon dynamics in the northern permafrost region on the trajectory of climate change *Proc. Natl Acad. Sci.* **115** 3882–7

Natali S M, Schuur E A G, Trucco C, Hicks Pries C E, Crummer K G and Baron Lopez A F 2011 Effects of experimental warming of air, soil and permafrost on carbon balance in Alaskan tundra *Glob. Change Biol.* **17** 1394–407

Natali S M, Schuur E A G and Rubin R L 2012 Increased plant productivity in Alaskan tundra as a result of experimental warming of soil and permafrost *J. Ecol.* **100** 488–98



Natali S M, Schuur E A G, Webb E E, Hicks Pries C E, Crummer K G, Pries C E H and Crummer K G 2014 Permafrost degradation stimulates carbon loss from experimentally warmed tundra *Ecology* **95** 602–8

Oleson K W *et al* 2013 Technical Description of version 4.5 of the Community Land Model (CLM) *Ncar Tech. Note* NCAR/ TN-503+STR National Center for Atmospheric Research (https://doi.org/10.5065/D6RR1W7M)

Oberbauer S F, Gillespie C T, Cheng W, Gebauer R, Serra A S and Tenhunen J D 1992 Environmental-effects on CO₂ efflux from riparian tundra in the northern foothills of the brooks range, Alaska, USA *Oecologia* **92** 568–77

Osterkamp T E, Jorgenson M T, Schuur E A G, Shur Y L, Kanevskiy M Z, Vogel J G and Tumskoy V E 2009 Physical and ecological changes associated with warming permafrost and thermokarst in Interior Alaska *Permafrost Periglacial Process.* 20 235–56

Richter-Menge J, Overland J E, Mathis J T and Osborne E 2017 Arctic Report Card 2017 https://arctic.noaa.gov/Report-Card/Report-Card-2017

Riley W J, Subin Z M, Lawrence DM, Swenson S C, Torn M S, Meng L, Mahowald N M and Hess P 2011 Barriers to predicting changes in global terrestrial methane fluxes: analyses using CLM4Me, a methane biogeochemistry model integrated in CESM *Biogeosci.* **8** 1925–53

Romanovsky V E, Smith S L and Christiansen H H 2010 Permafrost thermal state in the polar Northern Hemisphere during the international polar year 2007–2009: a synthesis *Permafrost and Periglacial Process.* **21**106–16

Romanovsky V, Smith S L, Christiansen H H, Shiklomanov N I, Streletskiy D A, Drozdov D S, Oberman N G, Kholodov A L and Marchenko S S 2012 *Permafrost in Arctic Report Card 2012* NOAA ftp://ftp.oar.noaa.gov/arctic/ documents/ArcticReportCard_full_report2012.pdf

Salmon V G, Soucy P, Mauritz M, Celis G, Natali S M, Mack M C and Schuur E A G 2016 Nitrogen availability increases in a tundra ecosystem during five years of experimental permafrost thaw *Glob. Change Biol.* **22** 1927–41

Schädel C, Schuur E A G, Bracho R, Elberling B, Knoblauch C, Lee H, Luo Y, Shaver G R and Turetsky M R 2014 Circumpolar assessment of permafrost C quality and its vulnerability over time using long-term incubation data *Glob. Change Biol.* 20 641–52

Schaefer K, Zhang T, Bruhwiler L and Barrett A P 2011 Amount and timing of permafrost carbon release in response to climate warming *Tellus* B 63 165–80

Schaphoff S, Heyder U, Ostberg S, Gerten D, Heinke J and Lucht W 2013 Contribution of permafrost soils to the global carbon budget *Environ. Res. Lett.* **8** 14026

Schneider von Deimling T, Meinshausen M, Levermann A, Huber V, Frieler K, Lawrence D M and Brovkin V 2012 Estimating the near-surface permafrost-carbon feedback on global warming *Biogeosciences* **9** 649–65

Schneider von Deimling T, Grosse G, Strauss J, Schirrmeister L, Morgenstern A, Schaphoff S, Meinshausen M and Boike J 2015 Observation-based modelling of permafrost carbon fluxes with accounting for deep carbon deposits and thermokarst activity *Biogeosciences* **12** 3469–88

- Schuur E A G, Crummer K G, Vogel J G and Mack M C 2007 Plant species composition and productivity following permafrost thaw and thermokarst in Alaskan tundra *Ecosystems* 10 280–92
- Schuur E A G, Vogel J G, Crummer K G, Lee H, Sickman J O and Osterkamp T E 2009 The effect of permafrost thaw on old carbon release and net carbon exchange from tundra *Nature* **459** 556–9
- Schuur E A G *et al* 2015 Climate change and the permafrost carbon feedback *Nature* 520 171–9
- Shaver G R, Billings W D, Chapin F S, Giblin A E, Nadelhoffer K J, Oechel W C and Rastetter E B 1992 Global Change and the Carbon Balance of Arctic Ecosystems Carbon/nutrient interactions should act as major constraints on changes in global terrestrial carbon cycling *BioScience* 42 433–41
- Shi M, Fisher J B, Brzostek E R and Phillips R P 2015 Carbon cost of plant nitrogen acquisition: global carbon cycle impact from an improved plant nitrogen cycle in the community land model *Glob. Change Biol.* **22** 1299–314
- Sistla S A, Asao S and Schimel J P 2012 Detecting microbial N-limitation in tussock tundra soil: implications for Arctic soil organic carbon cycling *Soil Biol. Biochem.* **55** 78–84
- Sistla S A, Moore J C, Simpson R T, Gough L, Shaver G R and Schimel J P 2013 Long-term warming restructures Arctic tundra without changing net soil carbon storage *Nature* 497 615–8
- Slater A G and Lawrence D M 2013 Diagnosing present and future permafrost from climate models J. Clim. 26 5608–23

Strauss J *et al* 2017 Deep yedoma permafrost: a synthesis of depositional characteristics and carbon vulnerability *Earth-Sci. Rev.* **172** 75–85

- Street L E, Shaver G R, Williams M and van Wijk M 2007 What is the relationship between changes in canopy leaf area and changes in photosynthetic CO2 flux in arctic ecosystems? *J. Ecol.* **95** 139–50
- Swenson S C, Lawrence D M and Lee H 2012 Improved simulation of the terrestrial hydrological cycle in permafrost regions by the Community Land Model J. Adv. Model. Earth Syst. **4** M08002

Taylor M A, Schuur E A G, Mauritz M, Pegoraro E, Salmon V G and Natali S M 2017 Eight Mile Lake Research Watershed, Carbon in Permafrost Experimental heating research (CiPEHR): aboveground plant biomass, 2009–2017 Bonanza Creek LTER - University of Alaska Fairbanks. BNZ:501 https://doi.org/10.6073/pasta/ 1af376985d83cd7e01c61b67abfa9f91

Webb E E, Schuur E A G, Natali S M, Oken K L, Bracho R, Krapek J P, Risk D and Nickerson N R 2016 Increased wintertime CO₂ loss as a result of sustained tundra warming J. Geophys. Res.: Biogeosci. 121 249–65

Wild B *et al* 2018 Amino acid production exceeds plant nitrogen demand in Siberian tundra *Environ. Res. Lett.* **13** 34002