Permafrost carbon-climate feedbacks accelerate global warming

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Permafrost soils contain enormous amounts of organic carbon, which could act as a positive feedback to global climate change due to enhanced respiration rates with warming. We have used a terrestrial ecosystem model that includes permafrost carbon dynamics, inhibition of respiration in frozen soil layers, vertical mixing of soil carbon from surface to permafrost layers, and CH4 emissions from flooded areas, and which better matches new circumpolar inventories of soil carbon stocks, to explore the potential for carbon-climate feedbacks at high latitudes. Contrary to model results for the Intergovernmental Panel on Climate Change Fourth Assessment Report (IPCC AR4), when permafrost processes are included, terrestrial ecosystems north of 60°N could shift from being a sink to a source of CO₂ by the end of the 21st century when forced by a Special Report on Emissions Scenarios (SRES) A2 climate change scenario. Between 1860 and 2100, the model response to combined CO₂ fertilization and climate change changes from a sink of 68 Pg to a 27 + -7 Pg sink to 4 + -18 Pg source, depending on the processes and parameter values used. The integrated change in carbon due to climate change shifts from near zero, which is within the range of previous model estimates, to a climate-induced loss of carbon by ecosystems in the range of 25 + -3 to 85 +-16 Pg C, depending on processes included in the model, with a best estimate of a 62 + -7 Pg C loss. Methane emissions from high-latitude regions are calculated to increase from 34 Tg CH₄/y to 41-70 Tg CH₄/y, with increases due to CO₂ fertilization, permafrost thaw, and warming-induced increased CH₄ flux densities partially offset by a reduction in wetland extent.

carbon cycle | land surface models | cryosphere | soil organic matter | active layer

oreal and Arctic terrestrial ecosystems are particularly sensi-Brive to future warming (1). These cold regions are crucial to the global carbon cycle because they are rich in soil organic carbon, which has built up in frozen soils, litter, and peat layers. Laboratory incubation experiments (2) and field studies (3) suggest that this old carbon could be lost rapidly through decomposition in response to warming. In particular, the slow burial of soil carbon below the base of seasonally thawed surface layers (the active layer) into deeper permafrost layers has led over tens of millennia to the formation of an enormous stock. This carbon stock is presently not actively cycling, but might become available for respiration if frozen soils thaw. Estimates of the total northern carbon pool are 495 Pg for the top meter of soils, 1,024 Pg to 3 m, and an additional 648 Pg for deeper carbon stored in yedoma (frozen, carbon-rich sediments) and alluvial deposits (4). Such a huge permafrost carbon pool, formed during the Pleistocene and Holocene, exists because decomposition is strongly inhibited in frozen soils, thus allowing old, otherwise labile carbon to persist and accumulate slowly to the present.

In the recent Coupled Carbon-Climate Change Model Intercomparison Project (C^4 MIP) (5)—which formed the estimate for the strength of the carbon-climate feedback for the Intergovernmental Panel on Climate Change Fourth Assessment Report (IPCC AR4) (6, 7)—and other studies (e.g., ref. 8) that examine the effects of CO₂ fertilization and climate change on the net carbon balance of terrestrial and ocean ecosystems, most terrestrial biosphere models predicted an enhanced carbon sink due to warming in high latitudes (Fig. 1D) (9), through longer growing seasons and enhanced productivity that offsets the warminginduced increase in heterotrophic respiration. However, none of these coupled models accounted for carbon vulnerable to decomposition when permafrost thaws. Models that have considered permafrost carbon losses calculate total emissions of CO₂ from permafrost carbon from 7–17 Pg by 2100 (10) to 190 + -64 Pg by 2200 (11). In addition to frozen soil carbon, northern wetlands are a strong source of methane (CH_4) to the atmosphere, averaging 35–45 Tg CH_4/y (12, 13), and this methane source is sensitive to changes in permafrost, wetlands hydrology, and ecosystem productivity. None of the models of C⁴MIP accounted for the climate feedbacks of natural CH₄ sources, even though CH₄ is a very efficient greenhouse gas [global warming potential (GWP) = 25] on 100 y timescale) (14).

Model

We selected the ORCHIDEE model as a representative land component of the C⁴MIP models, and designed four separate sets of simulation experiments to explore the sensitivity of the northern high-latitude CO₂ and CH₄ balance to the inclusion of critical soil carbon processes (Table S1). Typically, soil carbon models have used either a single bulk vertically integrated soil pool, though (10) adapted this approach to high latitudes by normalizing the carbon of this single pool relative to the thickness of the active layer. Here, in all cases, we use a fully vertically discretized soil carbon module, recently developed (15), where decomposition rates are calculated for each soil level, to dynamically model the steep vertical gradient in soil carbon residence time that occurs at the permafrost table in permafrost-affected soils (Fig. S1). In addition, the model soil physics has been improved to more realistically capture the effects of organic matter on active layer thickness (15).

The four experiments explored here are (i) control, in which soil carbon is vertically resolved but no additional processes are added; (ii) freeze, inhibition of decomposition in seasonally frozen soil layers, but no soil carbon in permafrost soil layers; (iii) permafrost, inclusion of permafrost carbon through vertical

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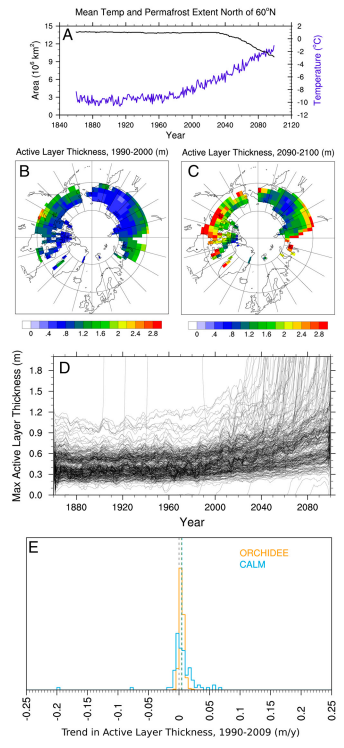


Fig. 1. Change in permafrost extent and properties over the model simulation period, for the region $60^{\circ}N-90^{\circ}N$. (A) Black line, permafrost extent (to 50 m); blue line, mean annual temperature for the high-latitude terrestrial region. (B) Active layer thickness (maximum depth of seasonally thawed soils), 1990–2000. (C) Active layer thickness, 2090–2100. Blank grid cells in (B–C) are those where we do not calculate permafrost within the top 50 m. (D) Trends in active layer thickness for all permafrost grid cells in the model. (E) A histogram of modeled and observed [CALM, (25)] active layer thickness trends (m/y) based on regression over the period 1990–2009.

mixing and soil organic insulation (15); and (iv) heat, inclusion of microbial heat release by decomposing microbes to the soil thermal budget (16).

In addition to the CO_2 balance, we model the climate response of CH_4 natural emissions by both deep permafrost layers and wetlands. For deep permafrost, we incorporate in ORCHIDEE the detailed process-based model of (16), in which (*i*) methanogenesis can occur in oxygen-poor deep permafrost horizons, and methanotrophy in the aerated upper soil profile; (*ii*) soil gas (O_2 and CH_4) diffusion is calculated to trigger methanotrophy vs. aerobic decomposition; and (*iii*) heat release due to exothermic decomposition reactions (decomposition, methanogenesis, and methanotrophy) can be included in the soil thermal budget.

For CH₄ emissions by wetlands in regions outside permafrost areas and in upper soil layers of permafrost regions, we use the wetland-CH₄ enabled version of ORCHIDEE (17, 18), in which wetland extent (saturated soil fraction) is calculated prognostically using the TOPMODEL (19, 20) subgrid approach, and methane emission rates are calculated for a given wetland extent, following an approach similar to Walter et al. (21). We model the temperature sensitivity of methanogenesis using a Q_{10} of 3, relative to an initial location-dependent mean annual temperature T_{mean} , based on a site-level optimization (17). We calculate two separate sets of wetland CH_4 fluxes, one allowing the base T_{mean} to change with changing climate, and the other where T_{mean} remain fixed, to bracket the uncertainty associated with possible microbial decomposition adaptation to warming. Wetland simulations are also calculated with separate biochemical CO₂ fertilization effect alone, and with the combined fertilization and climate effect of CO2. We then add these CH4 flux distributions across the high latitudes to the deep permafrost CH₄ emissions calculated from the permafrost model, to obtain total high-latitude natural CH₄ emissions.

In each simulation experiment a new mechanism is added to test its effect on the modeled CO₂ balance. The control case uses the standard ORCHIDEE soil carbon temperature sensitivity to respiration, with a classic temperature sensitivity Q_{10} of two (ref. 5). In the freeze experiment, we inhibit soil carbon decomposition by seasonal freezing (different sensitivity functions of low frozen respiration rate to warming were tested; see SI Text and Fig. S2). In the control and freeze cases, there is no vertical movement of soil carbon; thus no permafrost carbon stocks exist in these simulations. In the permafrost experiment, we add an initial permafrost carbon pool beneath the active layer by including thermal insulation by soil carbon and cryoturbation as in ref. 15; this mixing leads to the downward movement and burial of soil carbon from seasonally thawed soil layers into the upper permafrost (to ~3 m, Fig. S3), allowing a realistic model initialization. In addition, the specific very thick permafrost loess deposits in yedoma areas are initialized prior to the 10,000-y model equilibration with uniform carbon concentrations below the active layer to match observed carbon stocks (4) to include the presence of this relic frozen-but-labile Pleistocene carbon, mainly over Eastern Siberia. Finally, in the heating experiment, the soil thermal budget of the model accounts for the exothermic heat released by decomposition, exactly as described by ref. 16. We estimate uncertainty of each process using an ensemble of runs and varying key parameters over a given range.

We perform all model simulations over the period 1860-2100. For each experiment, we calculate a control run with preindustrial CO₂ levels and climate, a CO₂-only run with increasing CO₂ but fixed climate, and a CO₂+ climate run where both CO₂ concentration and climate vary. We calculate the effect of CO₂ (Fig. 2*C*) as the difference between the CO₂-only and the control runs, and the effect of climate change (Fig. 2*D*) as the difference between the CO₂+ climate and the CO₂-only runs. For all experiments, we run ORCHIDEE offline, so that each experiment is forced by the same meteorology. The model is forced by climate fields constructed as a base climatology (22, 23) plus anomalies relative to a climatological period 1961–1990 of the Institut Simon Pierre Laplace Climate Model 4 climate system model

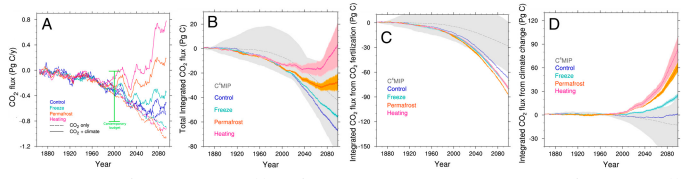


Fig. 2. Change in carbon fluxes over the model run. (A) Mean fluxes over modeled period. Contemporary budget estimate from McGuire et al. (1) (B) integrated changes. (C) Integrated changes in carbon balance due to rising CO_2 concentration alone. (D) Integrated change in carbon balance due to climate change alone (difference between CO_2 -only and CO_2 +climate change).

(24) for prescribed greenhouse gas-forced historical and future [Special Report on Emissions Scenarios (SRES) A2].

Results and Discussion

For each experiment, the initial equilibrium soil carbon stocks differ as a result of the processes included (Fig. S4 and Table S1), with a large increase in high-latitude soil C stocks (from ~ 200 Pg to ~ 500 Pg C in the top 3 m of soil) from permafrost processes, leading to better agreement with soil carbon observations (4) in freeze and permafrost, however, a substantial underestimate of initial carbon stocks still exists because we do not model the buildup of peatlands or organic soils.

We run the ORCHIDEE model fitted with these processes added in a transient climate change scenario. The modeled climate response leads to significant warming at high latitudes (Fig. 1), with mean high-latitude surface soil temperature rising approximately 8 C by 2100—much larger than the global mean—and permafrost extent (within the top 3 m) reduced by 30%. In addition, where permafrost does still exist at 2100, the active layer is deepened, with consequent thawing of previously frozen carbon. The changes in permafrost properties have a lag with respect to surface warming, and changes in active layer depth over the observed period (1990–2009) are small (mean 0.5 cm/y) and agree well with observed changes in active layer thickness (25), which we calculate by linear regression of all circumpolar active layer monitoring network (CALM) sites poleward of 60°N.

The modeled carbon fluxes of the region north of 60° N (Fig. 2) change as a result of both the effect of CO₂ fertilization on photo-

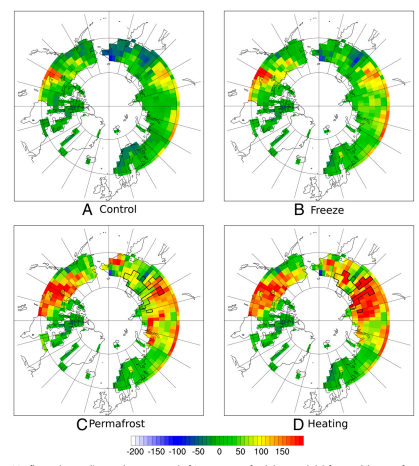


Fig. 3. Spatial patterns of net CO₂ fluxes due to climate change at end of 21st century, for (A) control, (B) freeze, (C) permafrost, and (D) heating experiments. Units are in gC/m2/y. Outlined cells are initialized as containing deep yedoma carbon.

synthesis, and the warming due to climate change. In all experiments, the effect of CO₂ fertilization is to increase vegetation productivity and thus create a carbon sink of 69-88 Pg relative to the control, whereas that of climate change is a net loss of carbon relative to the CO₂-only case, whose magnitude differs strongly between experiments. We also show the model range for the C⁴MIP experiments in Fig. 2 C and D, with a cumulative sink ranging from 0 to 60 Pg (mean 29 Pg) due to CO₂ fertilization alone, and from a sink of 77 Pg to a source of 20 Pg (mean 14 Pg sink) due to warming. The uncertainty evidenced by the large spread between the C4MIP models relates to their different parameterizations, their initial carbon storage as well as their remaining climate drifts (due to low-frequency variance and initial model disequilibrium), and associated drifts in the carbon fluxes. ORCHIDEE shows a very high sensitivity to CO₂ fertilization at high latitudes; this high sensitivity is likely due to a number of biases in the model: (i) there is no limitation by N in the model, and thus increases in CO₂ directly allow increases in productivity; and (ii) the baseline productivity of ORCHIDEE at high latitudes is higher than other models (26, 27), thus a proportionally similar change in the productivity leads to a larger gross change. These issues are also evident in the CH₄ emissions, which show a high sensitivity to CO₂ fertilization through substrate availability and local hydrologic feedbacks (18). Future work to integrate a dynamic N cycle and improve soil hydrology should reduce these biases. The sensitivity to CO₂ fertilization increases further in the model experiments because the longer turnover times of soil carbon with permafrost processes lead to a greater capacity for changes in productivity to translate to changes in storage.

In the control case given the ORCHIDEE model, the effect of warming is to lead to a large increase in vegetation productivity through longer growing seasons (+37 d over 1990-2100) that offsets the increase in heterotrophic respiration during the 21st century. Thus, this simulation gives only a small loss due to warming of 1 Pg C by 2100, a result within the range of C⁴MIP models (5), as seen in Fig. 2D. In the freeze experiment, the larger initial soil carbon stocks and higher effective temperature sensitivity of decomposition lead to a cumulative source due to warming of 25(+-3) Pg, which occurs mainly in the spring and fall (Fig. S5) due to a lengthened unfrozen soil carbon decomposing season (to a mean of 165 d relative to 130 d in 1990). The permafrost experiment gives an even larger cumulative source of CO₂ of 62(+-6) Pg due to warming over the 21st century. This carbon source is caused by partial decomposition of the old permafrost carbon pool, with the largest changes in the summer. Lastly, in the heating experiment, the extensive thawing of permafrost carbon stocks is accelerated by soil microbes releasing heat within the bottom of the active layer, which leads to a cumulative carbon loss due to warming of 85(+ - 16) Pg. We note that the warminginduced carbon loss also begins earlier in the experiment with microbial heating (Fig. 2A), leading to a contemporary highlatitude carbon sink that is small, but within the range of regional estimates (1). In the permafrost case, which does not include the heating term, Arctic ecosystems shift from a CO₂ fertilizationdriven sink to a climate change-driven source before 2100.

Fig. 3 shows the spatial distribution of the climate-induced CO_2 flux anomalies for each of the model experiments during the period 2090–2100. The control case shows widespread sink, which is partially attenuated in the freeze case. In the permafrost and heating cases, the region becomes a net source, with CO_2 emissions highest in regions that lie at the margins of the current permafrost zone, where permafrost is lost or the active layer substantially deepened in the future. Large carbon losses are seen in central Canada for the permafrost experiment, where substantial permafrost stocks exist that are vulnerable to warming. The effect of the microbial heat release in the heating experiment is particularly strong in Eastern Siberia, where it leads to more rapid permafrost degradation and associated carbon loss than is calcu-

lated in the permafrost experiment. The yedoma carbon stocks do not substantially contribute to the CO_2 or CH_4 fluxes in the permafrost case, because they are located in the coldest regions of Siberia, which are the most stable with respect to warming and thus have not thawed to depth by 2100 in this simulation.

Fig. 4 and Table S2 show the CH_4 balance for the permafrost and heating cases, also accounting for CH_4 emissions from wetlands. The effect of CO_2 fertilization is to increase the productivity of wetland plants and thus the methanogenesis substrate, leading to increased CH_4 emissions, to 71–74 Tg/y from 34 Tg/y in the early 20th century; this model estimate is subject to the same biases as for the larger carbon cycle, and has large uncer-

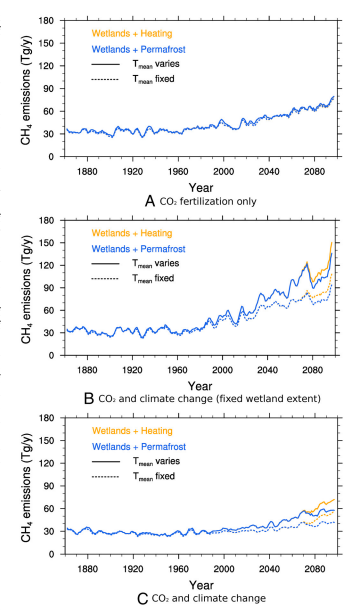


Fig. 4. CH₄ fluxes from high-latitude region over model runs (Tg CH₄). (A) CH₄ emissions under CO₂ fertilization alone; (B) CH₄ emissions under combined CO₂ increase and climate change, but holding wetland extent fixed; (C) CH₄ emissions under full climate change experiment with CO₂, climate, and its effect on wetland extent all varying. For each case, two separate wetland CH₄ experiments were carried out, with the reference temperature for methanogenesis, T_{mean} , remaining fixed or changing with climate. In addition, two separate permafrost CH₄ experiments were carried out, corresponding to the permafrost and heating experiments for the carbon balance.

tainties. Including warming as well, but holding wetland extent fixed, leads to enhanced emissions of 84-120 Tg/y, with the large value for the runs in which T_{mean} is held fixed. However, warming leads in our wetland hydrology model to a reduction of wetland area due to increased evapotranspiration, leading to less summer inundation and thus less CH4 emission, for an increase to only 41–57 Tg/y. A similar shrinking of Arctic lakes has already been observed (28, 29), however this term is a large source of uncertainty in the CH₄ model. In the permafrost simulation, the deep permafrost carbon stores that could serve as the basis for extra methane emissions (16) are thawed only partially and in their upper layers in the time frame considered, thus not leading to large upland permafrost CH₄ emissions. Therefore, the change in CH₄ emissions is almost entirely realized from changes of wetland areas and flux intensity. By contrast, in the heating simulation, a fraction of 0-30% of deep permafrost thaws by the self-heating feedback that is described by ref. 16, leading to extramethanogenesis because of the deeper yedoma permafrost carbon that is decomposed. This switch on of deep permafrost methanogenesis leads to an additional methane source of up to 14 Tg CH_4/y , 40% of the current total high-latitude CH_4 natural source (gas hydrates nonmodeled) although with large uncertainties. Using a CH₄ GWP of 25 and summing the changes to the integrated CO₂ and CH₄ budgets over the scenario with fixed methanogenesis T_{mean} leads to a change in the high-latitude GWP of -63 PgC-equivalent for the control case and -22 PgCequivalent for the permafrost case. However, climate change alone induces an increase in GWP of the region of 47 Pg Cequivalent for the permafrost case.

The version of ORCHIDEE used here for testing the sensitivity of high-latitude CO₂ and CH₄ fluxes to warming does not include C-N interactions, which may affect both the CO₂-fertilization and climate response to plant growth (30, 31). In particular, mineralization of nitrogen from thawing permafrost soil organic matter could lead to both enhanced plant growth and decomposition, with an uncertain sign on the net carbon balance response to the added N (32, 33). Inclusion of these interactions in ORCHIDEE without permafrost representation (18) leads to almost cancellation of the high-latitude carbon sink due to CO₂ fertilization. By contrast, when including C-N interactions and warming, the balance at high latitudes between increased growth and respiration is only shifted slightly. Including C-N interactions in our simulations should strongly reduce the CO₂induced sink potential of high-latitude ecosystems, turning all of our simulation experiments into carbon sources by 2100; however, the uncertainty associated with the warming-induced increase in N mineralization is unresolved here. Finally, several other processes, not modeled here, could also affect the highlatitude CO₂ balance, including northerly expansion of the boreal forest (34), changes to the fire regime (10, 35), or other disturbance mechanisms.

We attempted to incorporate in this study some of the latest mechanistic understanding about the mechanisms controlling soil CO_2 respiration and wetland CH_4 emissions, but uncertainties remain large, due to incomplete understanding of biogeochem-

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ical and physical processes and our ability to encapsulate them in large-scale models. In particular, small-scale hydrological effects (36) and interactions between warming and hydrological processes are only crudely represented in the current generation of terrestrial biosphere models. Fundamental processes such as thermokarst erosion (37) or the effects of drying on peatland CO_2 emissions (e.g., ref. 38) are lacking here, causing uncertainty on future high-latitude carbon-climate feedbacks. In addition, large uncertainty arises from our ability to model wetland dynamics or the microbial processes that govern CH_4 emissions, and in particular how the complicated dynamics of permafrost thaw would affect these processes.

The control of changes in the carbon balance of terrestrial regions by production vs. decomposition has been explored by a number of authors, with differing estimates of whether vegetation or soil changes have the largest overall effect on carbon storage changes (39–41). These results demonstrate that with the inclusion of two well-observed mechanisms: the relative inhibition of respiration by soil freezing (42) and the vertical motion in Arctic soils that buries old but labile carbon in deeper permafrost horizons, which can be remobilized by warming (3), the high-latitude terrestrial carbon response to warming can tip from near equilibrium to a sustained source of CO_2 by the mid-21st century. We repeat that uncertainties on these estimates of CO_2 and CH_4 balance are large, due to the complexity of high-latitude ecosystems vs. the simplified process treatment used here.

The 61 Pg C reduction in cumulative carbon fluxes at 2100 between our permafrost and control cases imply that when taking frozen soil processes into account, climate change can lead to a large reduction of the carbon sinks in high-latitude. About one third (24 Pg) of this climate-induced carbon loss is due to seasonally frozen soil carbon, the rest being due to permafrost processes. The modeling studies included in the IPCC AR4 (6, 7) inferred that tropical ecosystems would act as a climate change-induced carbon source, mid- and high-latitude ecosystems could be regions where climate change would enhance carbon storage; we show here that including the vast permafrost carbon pool in models leads to a qualitatively different result, in which high latitudes act as future CO₂ and CH₄ sources, leaving only the mid latitudes as potential climate regulators. We note as well that significant permafrost stocks exist and a steep loss continues at 2100, so that beyond the time horizon considered here there is still a potential for enormous carbon losses from high-latitude soils to continue.

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