

Tundra soil carbon is vulnerable to rapid microbial decomposition under climate warming

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Microbial decomposition of soil carbon in high-latitude tundra underlain with permafrost is one of the most important, but poorly understood, potential positive feedbacks of greenhouse gas emissions from terrestrial ecosystems into the atmosphere in a warmer world^{1–4}. Using integrated metagenomic technologies, we showed that the microbial functional community structure in the active layer of tundra soil was significantly altered after only 1.5 years of warming, a rapid response demonstrating the high sensitivity of this ecosystem to warming. The abundances of microbial functional genes involved in both aerobic and anaerobic carbon decomposition were also markedly increased by this short-term warming. Consistent with this, ecosystem respiration (R_{eco}) increased up to 38%. In addition, warming enhanced genes involved in nutrient cycling, which very likely contributed to an observed increase (30%) in gross primary productivity (GPP). However, the GPP increase did not offset the extra R_{eco} , resulting in significantly more net carbon loss in warmed plots compared with control plots. Altogether, our results demonstrate the vulnerability of active-layer soil carbon in this permafrost-based tundra ecosystem to climate warming and the importance of microbial communities in mediating such vulnerability.

Permafrost, defined as 'subsurface earth materials remaining below 0 °C for two consecutive years¹', is a unique characteristic of polar regions and high mountains. In permafrost-underlain high-latitude tundra, plant-derived carbon has accumulated over hundreds to thousands of years because low temperatures and saturated soils reduce microbial decomposition of soil organic C (refs 5,6). As a result, nearly 50% of the global soil organic C is stored in Northern Hemisphere permafrost and the active-layer soils above, although they cover only 16% of the global terrestrial area⁷. High-latitude tundra has long been recognized as being highly responsive to climate change⁸. Recent accelerated warming in the northern high-latitude region⁹ has resulted in rapid permafrost degradation, and studies suggest that permafrost could decline by 30–70% by the end of the twenty-first century^{10,11}. During permafrost degradation, frozen soil becomes biologically active,

with microbial decomposition resulting in massive ecosystem C loss, which is likely to dominate the overall net C exchange in permafrost regions¹. Although plant responses to climate warming in the active layer of the tundra soil have been intensively studied^{12–14}, microbial responses have not been examined until very recently^{4,15–18}.

Although various observational studies have documented the responses of tundra ecosystems to natural warming¹⁹, and some incubation studies revealed microbial community changes on permafrost thaw in laboratory settings^{16,18}, very few studies examined microbial responses to climate warming in tundra ecosystems in the field. As field experimental warming can directly examine the impacts of temperature increases on the microbial community *in situ*¹³, an ecosystem warming experiment, Carbon in Permafrost Experimental Heating Research (CIPEHR), was established in September, 2008, in Interior Alaska. The experiment is located in typical moist acidic tussock tundra²⁰, a dominant tundra type, on permafrost that is close to the freezing point and thus especially vulnerable to thaw in a warming climate²¹. In this experiment, snow fences (that is, increased snow pack for insulation) were used in the soil warming treatment to increase soil temperature, coupled with early spring snow removal to control snow-water equivalents in both warmed and control plots. Soil warming and control treatments were arranged in six replicates, providing sufficient statistical power. This is the first warming experiment to degrade surface permafrost without delaying spring snow melt¹⁴. To understand how vulnerable the active layer of the tundra soil is to climate warming, a total of 12 subsurface soil samples from a representative depth of 15–25 cm were collected from both warmed and control plots after short-term (1.5 years) warming for geochemical and microbial analyses. These samples represented active-layer soil that freezes in winter and thaws in the growing season, and were within the organic horizon along the depth profile. As soil microbial community structure is tightly linked to changes in the aboveground plant community and soil environmental conditions²², we predicted that short-term warming would result in selective microbial growth, which would be seen as a shift in the active-layer microbial community structure and

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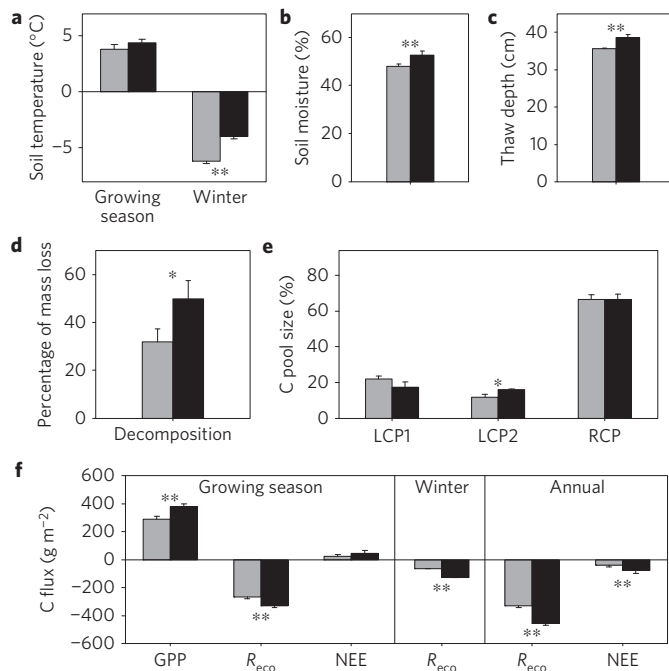


Figure 1 | Warming effects on soil variables and ecosystem C fluxes. Grey bars represent control plots and black bars represent warmed treatment plots. **a**, Soil temperature in both growing season (May to September 2010) and wintertime (December 2009 to March 2010) averaged across 5, 10, 20 and 40 cm. **b**, Soil moisture. **c**, Maximum thaw depth. **d**, Standard cellulose filter paper decomposition rate (mass loss) in the field. **e**, Proportion of soil C pools in total organic C, including labile C pool 1 (LCP1, mainly polysaccharides) and 2 (LCP2, mostly cellulose), and recalcitrant C pool (RCP). **f**, Growing season (May to September 2010), wintertime (October 2009 to April 2010) and annual ecosystem C fluxes, which were estimated on the basis of the C amount from CO₂ emissions. GPP, gross primary productivity; R_{eco} , ecosystem respiration; NEE, net ecosystem C exchange. Positive values indicate C sink, and negative values represent C source. Error bars represent standard error of the mean. The differences between warmed and control plots were tested using two-tailed *t* tests, indicated by ** when $p < 0.05$, or * when $p < 0.10$. Panels **a–d,f** were reanalysed from previously published data^{12,23,31}.

accompanying gene content, especially in those populations and traits important to both aerobic and anaerobic C decomposition and nutrient cycling. Consequently, soil C in this tundra ecosystem would be highly vulnerable to climate warming.

Short-term soil warming altered several environmental attributes (for example, plant, soil microclimate and soil properties) of the tundra^{12,14,23}. First, soil temperature (5–40 cm) increased by 2.3 °C (from –6.2 °C to –4.0 °C) in response to warming in wintertime and by 0.6 °C (from 3.8 °C to 4.4 °C) during the growing season in 2010 (Fig. 1a), which led to a substantial surface permafrost thaw as indicated by an increased thaw depth (8.8%, $p < 0.001$; Fig. 1c). Similarly, soil moisture increased in response to warming (over 10%, $p = 0.03$; Fig. 1b). Second, GPP increased (30.3%, $p = 0.02$; Fig. 1f), mainly owing to enhanced growth of graminoids (57.5% increase in biomass, $p = 0.05$). Warming also extended the growing season length through earlier bud break and delayed senescence¹². In addition, the percentage of mostly cellulose fraction of the labile C pool²⁴ in total soil organic C was higher (36.1%, $p = 0.06$) in warmed than control soils (Fig. 1e). The C amount of this pool under warming tended to increase as well, but not statistically significantly (Supplementary Fig. 5). Together, these results indicated that environmental attributes of the tundra soil were altered rapidly by short-term warming.

Table 1 | Significance tests on the effects of warming on the microbial community functional structure detected by GeoChip hybridization.

	MRPP		ANOSIM		Adonis	
	δ	p	R	p	F	p
Euclidean	69.02	0.04	0.38	0.02	1.69	0.04
Horn	0.06	0.04	0.36	0.02	1.20	0.23
Bray	0.09	0.03	0.30	0.02	1.28	0.21

Three different permutation tests were performed, including the multiple response permutation procedure (MRPP), analysis of similarity (ANOSIM) and permutational multivariate analysis of variance (Adonis), calculated with Euclidean, Horn or Bray distance. Bold values indicate $p < 0.05$.

The observed alterations in the soil microclimate (temperature, moisture, thawing depth), soil C, and GPP in response to warming would be expected to cause significant changes in the microbial communities in the active layer of tundra soil. Consistent with this expectation, the microbial community functional gene structure was markedly different between warmed and control plots as revealed by the detrended correspondence analysis of the GeoChip data (Supplementary Fig. 1a), indicating increases in certain genes and possibly the organisms that host these. Three different non-parametric multivariate statistical tests (ANOSIM, Adonis and MRPP) showed that the functional community structure differed substantially between warmed and control plots (Table 1). However, significant differences in the overall community structure were not detected with 16S ribosomal RNA gene-based amplicon and shotgun metagenomic sequencing approaches (Supplementary Fig. 1b,c). This is most likely due to the high heterogeneity of soil environments, low taxonomic resolution of the experimental approaches, and/or high noise associated with random sampling²⁵. Canonical correspondence analysis revealed that soil temperature, moisture and plant GPP were the main significant variables related to the microbial community functional structure ($F = 1.68$, $p = 0.005$; Supplementary Fig. 2a). This is also consistent with our central hypothesis that warming-induced changes in plant productivity and soil microclimate significantly alter the soil microbial community structure. In addition, soil community DNAs were shotgun sequenced and a total of 3.24 billion raw sequences were obtained for these samples (Supplementary Table 2). Although the overall metagenome structures were not separable into warmed versus control groups (Supplementary Fig. 1c, Supplementary Fig. 3 and Supplementary Table 1), a small portion (7.4%) of total subsystems, genes associated with microbial physiological attributes and ecosystem processes, were significantly different between warmed and control plots ($p < 0.05$; Fig. 2d). In particular, warmed plots were enriched in genes associated with pathways related to labile C utilization (Supplementary Table 3). Our above results indicated that the microbial communities in the active layer of tundra soil were responsive to warming.

Soil warming also significantly impacted a number of microbial functional and phylogenetic groups important for C decomposition. First, more than half (54.5%) of the detected C decomposition genes were increased by warming according to GeoChip signal intensities ($p < 0.05$; Fig. 2a), including those involved in degrading starch (for example, *amyA* encoding α -amylase), hemicellulose (for example, *ara* encoding arabinofuranosidase), cellulose (for example, cellobiase), chitin (for example, endochitinase), aromatics (for example, *vdh* encoding vanillin dehydrogenase) and lignin (for example, glyoxal oxidase, phenol oxidase). Also, the total fungal functional gene intensity detected by GeoChip was more abundant in warmed plots than control plots (4.7%, $p < 0.001$; Supplementary Fig. 4a). Increases of the genes involved in recalcitrant C decomposition (Fig. 2a) suggest the

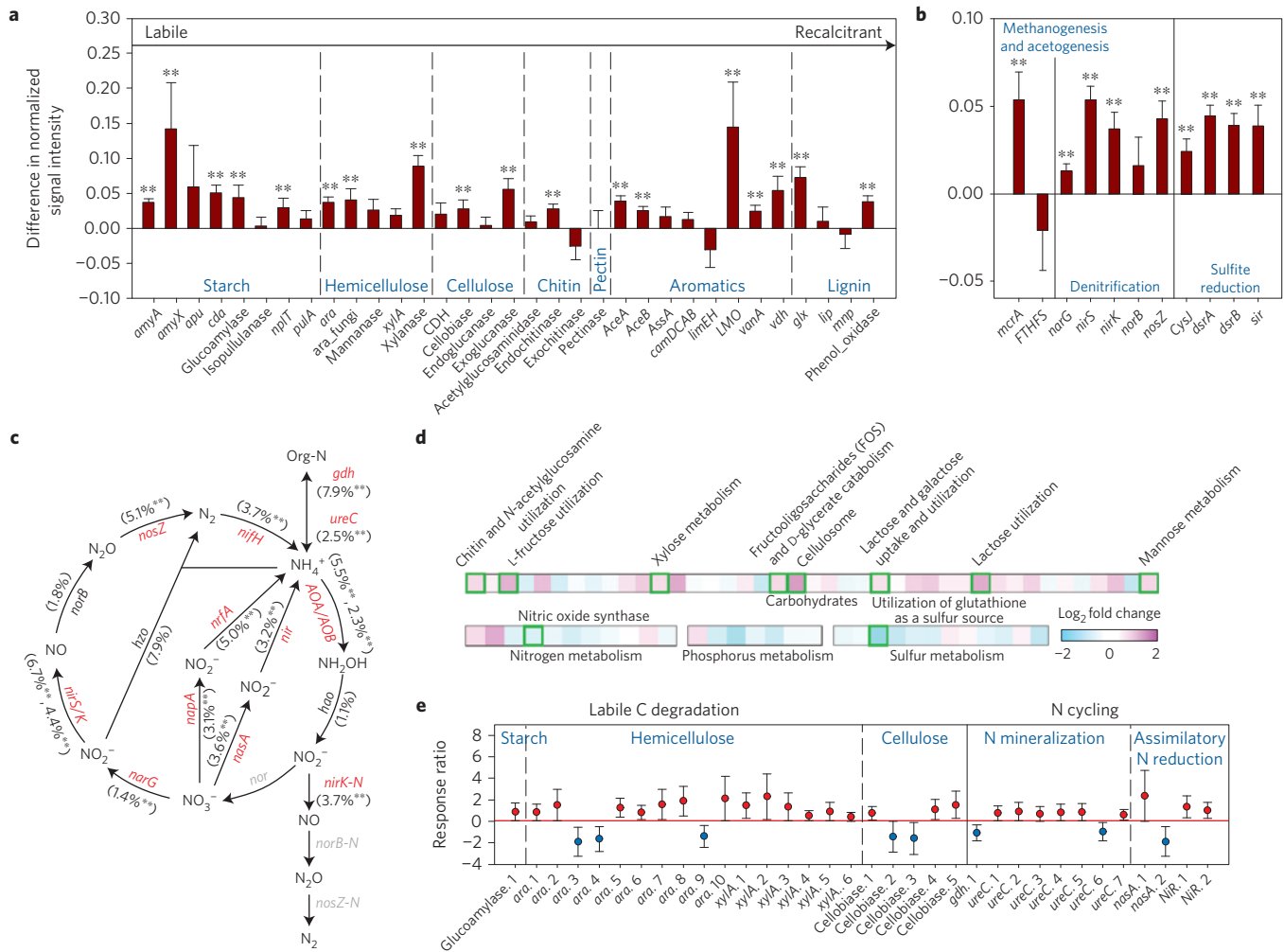


Figure 2 | Warming effects on functional genes involved in biogeochemical cycling processes. **a**, C degradation from GeoChip data. The targeted substrates were arranged in order from labile to recalcitrant. **b**, Anaerobic processes from GeoChip data. **c**, N processes from GeoChip data. The percentage change in N gene abundance in response to warming is indicated in parenthesis. Genes where change in abundance was significant ($p < 0.05$) are labelled in red. Grey-coloured genes were not targeted by the version of GeoChip used here, not detected or not applicable. **d**, Abundance of subsystems involved in C, N, phosphorus and sulfur cycling from metagenomic shotgun sequence data. Changes in subsystems are indicated as fold change (\log_2 (warmed/control)) in abundance. Significant differences between warmed and control plots are highlighted with green squares. **e**, Response ratios showing significant changes in abundance of gene clusters involved in C and N cycling from the metagenomic shotgun sequence data. These gene clusters were identified by searching the shotgun sequence data sets using GeoChip genes as queries. Each cluster on the x axis represents a group of sequences among which the similarities are $\leq 95\%$. The GenBank GI numbers of the representative sequences for the gene clusters are listed in Supplementary Table 8. Error bars indicate 95% confidence intervals of abundance differences between warmed and control groups. The full names of the genes in this figure are listed in Supplementary Table 7.

possible degradation of old recalcitrant C and thus a potential positive feedback to climate warming. In addition, shotgun metagenome sequence data revealed that a substantial portion (19.5%, 8 of 41) of C degradation pathways was increased by warming ($p < 0.05$; Fig. 2d, Supplementary Table 3), including those for cellulose, mannose metabolism, carbohydrate hydrolases, fructooligosaccharides and raffinose utilization, lactose and galactose uptake and utilization, L-fructose utilization, xylose utilization, chitin utilization and N-acetylglucosamine utilization. More specifically, many individual genes involved in labile C degradation (for example, starch, hemicellulose and cellulose), which were identified from metagenome sequences using GeoChip genes as queries, were increased (95% confidence interval; Fig. 2e). Overall, as these functional genes directly participate in aerobic C degradation, their higher abundance could enhance C decomposition and hence contribute to positive climate feedback.

The potential for accelerated C decomposition was supported by several independent pieces of evidence. First, the cellulose decomposition rate measured by adding external cellulose substrate was higher under warming (Fig. 1d). Also, both winter R_{eco} , derived almost exclusively from heterotrophic soil respiration, and growing season R_{eco} , including both autotrophic (from plants) and heterotrophic soil respiration, increased with warming (100% and 24%, $p < 0.05$; Fig. 1f; ref. 23). In addition, strong correlations were observed between both growing season and wintertime R_{eco} and the functional gene groups involved in degrading almost all C compounds targeted by GeoChip, including starch, hemicellulose, cellulose, chitin, aromatics and lignin (Supplementary Table 5), suggesting that changes in abundance of these genes could be important in mediating R_{eco} (ref. 26).

Permafrost thawing induced by warming often increases soil water content and creates a mosaic of flooded areas interspersed

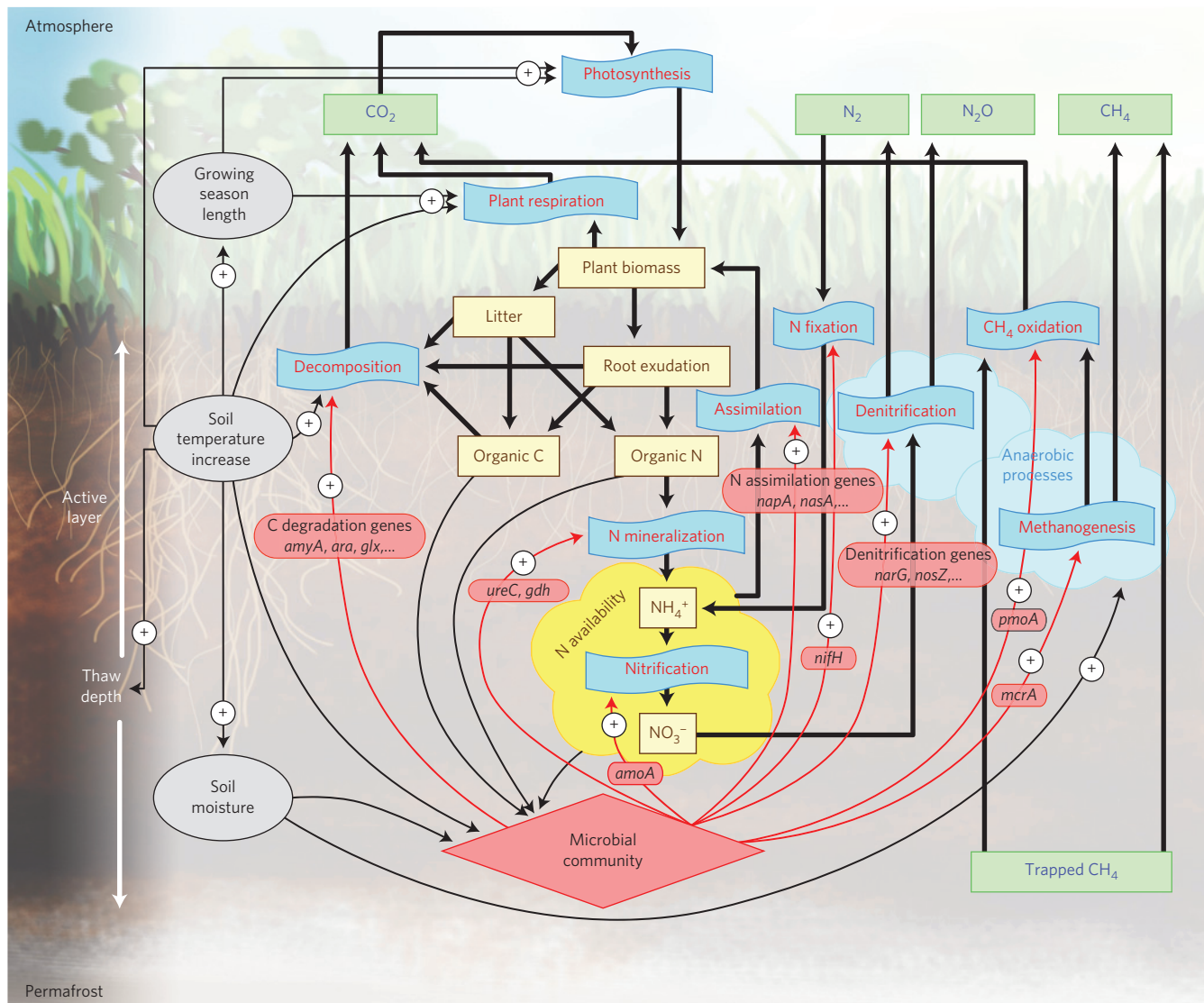


Figure 3 | A conceptual model of the impact of warming on the active layer of tundra ecosystem processes. Greenhouse gas pools are represented by green square frames, material pools by yellow square frames, and biological processes by frames in the shape of blue punched tape. Material flows are indicated by thicker black arrows. Impacts of environmental attributes (for example, soil temperature) and microbial community are marked by narrow arrows in black and red, respectively, and labelled with a '+' if increases in gene abundance were observed in this study.

within dry areas, which may potentially enhance anaerobic C decomposition^{1,18}. As water is heterogeneously distributed spatially and temporally, oxygen would also be expected to be unevenly distributed in the soil¹⁴. GeoChip hybridization-based analysis revealed that genes involved in several important anaerobic respiration processes, such as denitrification, methanogenesis and sulfate reduction, were increased by warming ($p < 0.05$; Fig. 2b). Shotgun metagenome sequence analysis also revealed that the pathway for denitrification (marginally) increased in response to warming ($p = 0.08$; Supplementary Table 6). These results were consistent with laboratory incubation studies, where methanogenic pathways were increased within several days after permafrost thaw^{16,18}. Although some upland permafrost areas were observed to be CH_4 sinks²⁶, for our studied site, significantly increased CH_4 emission rates after several years of warming have been reported²⁷. As CH_4 and N_2O have 28 and 265 times the warming potential of CO_2 per mole²⁸, respectively, anaerobic microbial responses are most likely of considerable importance.

Estimates show that methane emission in northern wetlands, including tundra, accounts for 25% of the global methane

release from natural sources²⁹. GeoChip analysis revealed that warming increased the gene encoding methyl coenzyme M reductase A (*mcrA*), a key enzyme in methanogenesis ($p < 0.01$; Supplementary Fig. 4b). Although 16S rRNA sequence analysis detected two methanogens (*Methanobacteria* and *Methanomicrobia*; Supplementary Fig. 4d), the relative abundance between warmed and control plots was not significantly different for either methanogen. Warming, however, resulted in a greater abundance of *pmoA*, a gene encoding particulate methane monooxygenase subunit A ($p = 0.01$; Supplementary Fig. 4b), suggesting that more of the methane produced could be oxidized in the aerobic upper soil horizon at the warmed plots. Similar findings were reported in recent studies on both incubated permafrost soils¹⁶ and active-layer samples *in situ*^{17,26}.

Warming also increased genes involved in N cycling (Fig. 2c,e), microbial phosphorus utilization and sulfur metabolism (Supplementary Fig. 4c). Most (82.4%) of the GeoChip-detected functional genes involved in N cycling were increased in response to warming ($p < 0.05$; Fig. 2c), consistent with the previous finding that warming enhances nutrient cycling³. For example,

the abundance of N₂-fixing bacteria was higher in response to warming ($p < 0.05$; Fig. 2c), and two bacterial classes (Opitutae and Deltaproteobacteria) detected by PCR amplification of the *nifH* gene had higher abundance in warmed samples ($p < 0.05$; Supplementary Fig. 4e). Also, the abundance of key genes (for example, *gdh* and *ureC*) in N mineralization was higher in warmed than control soil ($p < 0.05$ or 95% confidence interval; Fig. 2c,e). In addition, warming seemed to increase nitrification and denitrification processes, as indicated by increased *nirK* and *amoA* genes from GeoChip data (Fig. 2c). The increase in *amoA* could potentially lead to higher nitrate concentrations, which is also supported by the greater abundance of genes for various reductive processes that use nitrate as an electron acceptor, such as *narG*, *nirS/nirK* and *nosZ* for denitrification, *napA* and *nrfA* for dissimilatory nitrate reduction to ammonium, and *nasA*, *nirA* and *nirB* for assimilatory nitrate reduction (Fig. 2c,e). Microbial phosphorus utilization genes (phytase and *ppx*) and eight of the eleven detected sulfur metabolic genes had higher abundance in warmed than in control plots (Supplementary Fig. 4c). Although the significant increase in abundance of the genes involved in nutrient-cycling processes observed in warmed plots may potentially enhance the rates of nutrient cycling, more in-depth studies are necessary to determine the rates and extent of stimulation of different nutrient-cycling processes.

The increased abundance of N cycling genes (particularly those involved in N mineralization, N fixation and nitrification) and other nutrient-cycling genes could increase nutrient (especially N) availability in soil, which is important for ecosystem C dynamics because N is a limiting factor for plant growth in most tundra ecosystems¹². That warming enhanced plant N uptake is supported by the observation that from 2009 to 2010 plant foliar N mass increased in warmed plots (35%, $p < 0.01$), but remained unchanged in control plots¹². The enhanced plant N uptake could in turn affect GPP, which increased in response to warming ($p < 0.05$; Fig. 1f). Moreover, almost all genes involved in N cycling (18 of 19) and most C degradation genes (22 of 33) showed significant correlations with GPP ($p < 0.05$; Supplementary Table 5). Increased N₂ fixation, mineralization and nitrification could counteract the potential higher N loss from soil due to increased plant N uptake, denitrification and nitrate leaching. As a net result, the soil N availability seemed not to be affected by warming¹⁴.

In summary, our results highlight the importance of microbial-community-mediated feedbacks of the active layer to warming, as illustrated in a conceptual model (Fig. 3). In response to warming, deeper thaw depth increased the amount of C accessible for decomposition. Within the active layer, soil C is also more vulnerable to degradation through the following mechanisms: first, short-term soil warming altered the active-layer microbial community structure, demonstrating rapid responses by these communities; annual R_{eco} released 127 g more C m⁻² from warmed plots compared with controls, resulting in 38.6% more C loss from soil. Also, warming increased the abundance of functional genes involved in anaerobic processes, which could lead to a greater positive feedback by releasing more CO₂, CH₄ and N₂O (Fig. 3). In contrast, the potentially higher nutrient availability resulting from the increased abundance of nutrient-cycling genes would also stimulate plant growth (Fig. 3). In this study, increased GPP did not completely offset the C loss from the warming-induced R_{eco} increase. The net C loss from warmed plots doubled in 2010 (Fig. 1f), and was estimated to increase more in an actual climate warming scenario²³. However, it should be noted that the experimental results reported in this study were derived from the active layer of the Alaskan tundra soil. To generalize whether the results observed are applicable to permafrost requires further analyses with actual permafrost.

Overall, whether the tundra soil acts as a C source or sink depends on plant and microbial responses to climate warming.

Our results indicate that the soil C is highly vulnerable to climate warming and this vulnerability is determined by a set of complex microbial feedbacks to the temperature increase. Improved predictions by ecosystem models to climate warming³⁰ may be possible through better assessment of microbial functional capacities and their responses.

Methods

Methods and any associated references are available in the [online version of the paper](#).

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Author contributions

All authors contributed intellectual input and assistance to this study and manuscript preparation. J.Z., E.A.G.S., Y.L., J.M.T. and K.T.K. developed the original concepts. K.X., M.M.Y., Z.J.S., L.W., Z.H., Y.Q., Y.D., J.D.V.N., Q.W. and C.L. contributed reagents and data analysis. R.B. handled all soils processing and subsampling for microbial analysis. S.N. provided key field data. M.M.Y. and L.C. did sequencing and GeoChip hybridization. K.X., M.M.Y. and J.Z. performed data analysis and integration. K.X., M.M.Y. and J.Z. wrote the paper with help from E.A.G.S., K.T.K., Y.L., J.R.C. and J.M.T.

Additional information

Supplementary information is available in the [online version of the paper](#). Reprints and permissions information is available online at www.nature.com/reprints. Correspondence and requests for materials should be addressed to J.Z.

Competing financial interests

The authors declare no competing financial interests.

Methods

Site description and sampling. The Carbon in Permafrost Experimental Heating Research (CIPEHR) site was established in September 2008 at a moist acidic tundra area of Interior Alaska near the Denali National Park in the Eight Mile Lake region (63° 52' 59" N, 149° 13' 32" W). The experimental plots were located in the discontinuous permafrost region where permafrost thaw has been observed in the past several decades. Experimental design and site description were described in detail previously¹⁴. Briefly, three experimental blocks were located approximately 100 m away from each other. In each block, two snow fences were erected in the winter of each year (October to April) about 5 m apart. The soil warming treatment plots were located 5 m back from the leeward side of the snow fences, and the paired control plots were at the windward side of the snow fences. Soil temperature was increased in the warmed plots owing to thicker snow cover on the soil surface and lower wind strength. Snow fences were removed in the spring before snow melt to provide uniform hydraulic conditions in both winter warming and control treatments. From 1976 to 2009, mean monthly temperature in the field ranged from -16 °C in December to 15 °C in July, with an annual mean temperature of -1.0 °C. The average annual precipitation was 378 mm. Only C₃ plant species were observed in this area. Dominant species include *Eriophorum vaginatum*, *Vaccinium uliginosum*, some other vascular species, non-vascular feather moss and lichen. In the experimental plots, soil from the ground surface to a depth of 45–65 cm, depending on sampling cores, was rich in organic carbon (C) materials; below that depth was mineral soil with a mixture of glacial till and windblown loess. The active-layer depth was about 50 cm.

Twelve soil cores, six from treatment and six from control plots, were taken using electric drills in destructive sampling plots at the six snow fences in the beginning of the 2010 growing season (May), one and one-half years after the initiation of the winter warming treatment. Our analysis provides a snapshot of the soil microbial community response to early stage soil warming. The 15–25 cm depth soil fractions were analysed in this study.

Environmental and soil chemical measurements. Thaw depth was measured weekly during the growing season (May to September 2010) using a metal depth probe¹⁴. The thaw depth data presented in this study were the average values for the 2010 growing season.

Constantan–copper thermocouples and CR1000 data loggers (Campbell Scientific) were used to measure and record soil temperature and moisture content at 5, 10, 20 and 40 cm every half hour in flux bases installed in each plot¹⁴. The soil temperatures in Fig. 1 were reanalysed from previously published data¹², which represent either growing season or wintertime (December 2009 to March 2010) temperatures averaged over 5–40 cm soil depth. To represent the microclimate of the soil where and when the microbial communities were sampled, the soil temperature data used in canonical correspondence analysis (CCA; Supplementary Fig. 2) were the average values at 20 cm depth from December 2009 to May 2010. Volumetric water content from the soil surface to 15 cm depth was measured using site-calibrated Campbell CS616 water content reflectometer probes¹⁴. Soil moisture data presented in this study were averaged over the 2010 growing season.

To prepare soils for microbial and chemical analyses, visible roots and stones were removed by metal forceps. To measure soil C and nitrogen (N), soil samples (5 g) were dried at 70 °C until constant weight, ground to powder, encapsulated in silver foil and fumigated with HCl for 24 h at room temperature to remove soil inorganic C (carbonates). Soil C and N concentrations were analysed in the Colorado Plateau Stable Isotope Laboratory at the Northern Arizona University on a DELTA V Advantage isotope ratio mass spectrometer (Thermo Fisher Scientific), configured through a Finnigan CONFLO III (Thermo Fisher Scientific) and using a Carlo Erba NC2100 elemental analyser (CE Elantech). The total organic C (TOC) and soil N content of each sample were calculated as the percentage mass of C or N (ref. 32).

To measure soil C pools, soil samples were processed with a two-step hydrolysis procedure to separate the labile and recalcitrant C pools²⁴. First, 5 N H₂SO₄ was used to hydrolyse dried soil at 105 °C for 30 min, from which the hydrolysate and wash-offs were collected after centrifugation as labile pool 1, containing mainly polysaccharides. Second, the residue was then shaken continuously overnight at room temperature with 26 N H₂SO₄, followed by hydrolysis at 105 °C for 3 h with acid diluted to 2 N. The hydrolysate and wash-offs were recovered as labile pool 2, containing mostly cellulose. The recalcitrant C pool consisted of the remaining organic C. The organic C in labile pools 1 and 2 was analysed using a Shimadzu TOC-V CPH PC-Controlled TOC analyser (Shimadzu Corporation) and the organic C in the recalcitrant C pool was analysed using a PerkinElmer Optima 2000DV ICP-OES spectrometer (PerkinElmer) in the Environmental and Agricultural Testing Service laboratory at North Carolina State University.

Aboveground plant communities. Aboveground plant community investigations were conducted as described previously^{12,14}. In brief, aboveground biomass and net primary productivity (ANPP) were determined by a non-destructive point-frame method using a 60 × 60 cm point frame with a grid size of 8 × 8 cm (ref. 33). At

each of the 49 intersecting grid points, a metal rod (1 mm diameter) was placed vertically through the plant canopy. Species identity and tissue type (leaf, stem or fruit) were recorded for every 'hit' with the rod. Aboveground live biomass for each vascular plant species, moss and lichen was estimated by applying allometric equations developed for this site to the average number of point-frame 'hits' per plot¹⁴. Vascular plant ANPP was estimated as the sum of the current year's apical growth (leaves, stems, flowers and fruits) and secondary growth. The ratio of biomass between each tissue type and total plant was determined from destructive harvesting of a site adjacent to CIPEHR (ref. 35). Secondary growth was evaluated using growth rates determined from tussock tundra at Toolik Lake, Alaska³⁵. Moss NPP was measured by the cranked wire method, which measures vertical growth of moss using a stainless-steel reference wire inserted at the moss surface^{34,36}. Three to five cranked wires were placed in four moss types in each treatment at all fences to measure the growth from mid-May to mid-September. Feather moss NPP was estimated as the product of linear growth per stem, stem density, biomass per unit stem growth and percentage cover. Allometric equations developed for the Eight Mile Lake watershed³⁴ as well as percentage cover were used to convert the vertical growth of other types of moss into biomass. Moss NPP was the sum of all types of moss NPP. Current year's fully formed green leaves from six vascular plants found across plots were collected at peak biomass (mid-July) for measuring foliar N and at the end of the growing season (late September) for senescent N (ref. 14). At least three leaves from two to three individuals in each plot were collected each time. Leaves were dried at 60 °C, finely ground, and analysed on a continuous flow isotope ratio mass spectrometer (Thermo Fisher Scientific) coupled with a Costech elemental analyser.

Decomposition. Weighed cellulose filter papers (Fisher brand P8 09-802-1B) were placed into fibreglass mesh bags and placed vertically at 0–10 cm in the field soils in September 2009 and collected in September 2010. The bags were rinsed and dried at 60 °C for weighing. The percentage of mass loss was calculated to represent decomposition rate.

Ecosystem C flow. Ecosystem C flux measurements were described previously^{14,23}. Growing season net ecosystem exchange (NEE) and ecosystem respiration (R_{eco}) were measured from May to September 2010 using an automated CO₂ flux system coupled to the flux chambers¹⁴. R_{eco} was determined with night measurements. Gross primary productivity (GPP) was estimated as the difference between NEE and R_{eco} (values are positive for C flowing from atmosphere to terrain and vice versa). Winter respiration was estimated using a parameterized winter respiration model, adjusted using in-plot winter respiration measurements in March and April 2009 using an infrared gas analyser in a portable CO₂ flux system. In winter, there was no photosynthetic activity and R_{eco} represents mainly microbial respiration. The C flux data used for analysis in this study were reanalysed from previous published data sets²³.

Soil DNA extraction. Soil DNA was extracted using a PowerMax Soil DNA Isolation Kit (MO BIO), and the quality was assessed based of spectrometry absorbance at wavelengths of 230 nm, 260 nm and 280 nm (ratios of absorbance at 260/280 nm around 1.8, and 260/230 nm >1.7) detected by a NanoDrop ND-1000 spectrophotometer (NanoDrop Technologies). Then it was quantified with Pico Green using a FLUOstar OPTIMA fluorescence plate reader (BMG LabTec) before being used for gene array labelling and sequencing library preparation. Detailed protocols for soil microbial community analysis (for example, amplicon sequencing, shotgun sequencing and GeoChip) are provided as Supplementary Information.

GeoChip analysis. GeoChip 4.2 is a comprehensive gene array containing 107,950 probes designed for covering 792 functional gene families from 11 major functional categories including C, N, phosphorus and sulfur cycling^{37,38}. One microgram of DNA from each sample was mixed with random primers and denatured before dNTP, fluorescent dye Cy-3 dUTP and DNA polymerase were added for labelling at 37 °C for 6 h, followed by heating at 95 °C for 3 min. Labelled DNA was purified and dried up. For hybridization, DNA was resuspended in hybridization solution containing a sample tracking control, formamide, SSC, SDS, a Cy3-labelled alignment oligonucleotide, a Cy5-labelled alignment oligonucleotide and a Cy5-labelled common oligonucleotide reference standard target. After denaturing, the mixtures were deposited onto the glass microarray and hybridized at 42 °C for 16 h. Then the arrays were washed and dried, and scanned by an MS 200 Microarray Scanner (NimbleGen) at 532 nm and 635 nm. NimbleScan software version 2.5 (NimbleGen) was used to grid and process the images to transform them into signal intensity. The raw signals from NimbleScan were submitted to the Microarray Data Manager on our website (<http://iee.ou.edu/microarray>), cleaned, normalized and analysed using the data analysis pipeline. Briefly, spot signal-to-noise ratio and minimum intensity cutoff were used as standard to remove unreliable spots. Both the universal standard and functional gene spot intensities are used to normalize the signals among arrays. Data were log

transformed after cleaning and normalization. A total of 48,188 functional gene probes were detected across all samples in this study.

Illumina MiSeq sequencing of 16S rRNA gene amplicons. DNAs were amplified for the V4 region of 16S rRNA genes using primer set 515F and 806R, and sequenced in one run on a MiSeq using 2 × 150 pair end format³⁹. Raw sequences were assembled using RDP's paired-end reads Assembler. Any assembled sequences with any ambiguous bases ('N') were discarded. Then, 5.28% of the remaining reads were identified as chimaeras using Uchime⁴⁰ and removed. The remaining sequences were clustered into operational taxonomic units (OTUs) using Uclust⁴¹ at 97% identity, and randomly resampled to the depth of 42,684 reads per sample. Representative sequences chosen by Uclust from each OTU were annotated taxonomically using the RDP Classifier⁴² with the confidence cutoff 0.5. Finally, 512,208 sequences in 23,677 OTUs were obtained.

454 pyrosequencing of *nifH* gene amplicons. *nifH* genes from the DNA samples were amplified using the primer pair *nifH* Poly F (5'-TGCGAYCCSAARGCBGACTC-3') and Poly R (5'-ATSGCCATCATYTCRCGGA-3') and sequenced on the 454 GSFLX Titanium platform at Macrogen^{43–45}. After trimming primers, the sequences were cleaned using LUCY (ref. 46). Sequences with 'N', those containing frameshift(s) detected by FrameBot⁴⁷, and those identified as chimaeras by Uchime⁴⁰ according to the Zehr *nifH* Database⁴⁸ were all removed from downstream analyses. The remaining 162,523 sequences were clustered into 2,643 non-singleton OTUs using CD-HIT (ref. 49) at 0.95 identity, which was an arbitrary but strict enough cutoff to identify different species according to previous studies on *nifH* and other N-cycling-related genes^{50–52}. Finally, 2,643 non-singleton OTUs were normalized to relative abundance (scale all of the sample sequence numbers to the largest one) for statistical analyses. The representative sequence for each OTU was assigned taxonomic information using the FrameBot⁴⁷ nearest-neighbour match with an identity cutoff of 0.5.

Shotgun metagenome sequencing analysis. Each soil metagenome was prepared using the TruSeq Kit and sequenced at the Los Alamos National Laboratory Genome Facility using the Illumina HiSeq 2000 in one flow cell lane with a 2 × 150 bp paired-end kit³⁹. A total of 3.24 billion reads were generated from the 12 samples, with both phylogenetic and functional information extracted. After data processing, it was found that one of the samples from the control (C1) did not produce enough useful sequence during shotgun sequencing and thus this sample was removed from all subsequent analyses. For phylogenetic analysis, the metagenome reads were trimmed⁵³ and searched against representative OTU sequences from GreenGenes⁵⁴ using BLAT (ref. 55). Paired reads that both matched GreenGenes reference sequences were identified as 16S reads and were extracted for further analyses. These 16S reads were subsequently searched against the 99%-clustered GreenGenes OTU sequences. The reads were assigned to the taxon that was the lowest common ancestor of the two reads in a pair. The 797,898 reads were assigned to 23,167 OTUs in total. For functional subsystem analysis, 25 million reads were randomly resampled from each sample. Open reading frames were predicted on non-16S encoding reads using FragGeneScan⁵⁶. The translated amino acid sequences were then searched against the M5NR database⁵⁷ using BLAT. Reads matching genes incorporated in the SEED database⁵⁸ were assigned to the corresponding best-matched subsystem(s). The numbers of assigned reads were taken as a proxy of abundance of the SEED subsystem(s). An approach combining re-sampling techniques, the DESeq package⁵⁹, and binomial testing with adjusted *p* values⁶⁰ was then applied to identify significantly differentially abundant subsystems (pathways) under warming versus control plots, as described previously⁶¹.

Annotating shotgun sequences on the basis of GeoChip genes. An ecological functional gene-oriented metagenomic analysis pipeline (EcoFun-MAP) has been developed to fish out sequence reads of important environmental functional genes from shotgun metagenome sequence data. EcoFun-MAP is a method designed for annotating metagenomic sequences by comparing them with functional genes used to fabricate GeoChip. In the preparation of the reference databases, keyword queries were submitted to the NCBI (ref. 62) online GenBank for 308 functional genes to retrieve candidate reference sequences, from which 5 to 200 distinct representative sequences from each gene were manually selected functional gene seed sequences (FGSSs). The selected FGSSs were aligned using both global and local algorithms in ClustalW (ref. 63), and the resulting alignments were used as input for another program HMMBUILD (ref. 64) to build both global and local HMMER (ref. 64) models (FGSS-HMM). Next, the candidate reference sequences for each gene were searched back against corresponding FGSS-HMM using HMMSEARCH (ref. 64). The output sequences, termed functional gene reference sequences (FGRSs), were clustered into OTUs for each gene using CD-HIT at the similarity threshold of 95%. In addition, BLAST databases were constructed on the FGRSs with MAKEBLASTDB (ref. 65). To this end, two reference databases

involved in the method were established: FGSS-HMM and FGRS-BLAST. For annotation, sequences from HighSeq were resampled to the minimal number of reads in a sample, and were quality trimmed by Btrim⁶⁶. All trimmed nucleotide sequences were translated into protein sequences using FragGeneScan⁵⁶. HMMSEARCH was used for annotating the predicted protein sequences with the FGSS-HMM database, and both global and local model hits were counted as valid results. Also, all FGSS-HMM confirmed sequences were compared together against the FGRS-BLAST database with BLASTN (ref. 65). Only best hits (Rank No. 1 in BLAST results) between probes and sequences were kept as final processing results. The web-based pipeline application of EcoFun-MAP can be accessed with request.

Statistical analysis. Statistical analyses were carried out using R software version 2.15.1 using the package vegan (v.2.3-2; ref. 67) when not specified. Detrended correspondence analysis⁶⁸ was performed to visualize the overall microbial community composition among samples. Three complementary non-parametric multivariate analyses, non-parametric multivariate analysis of variance (Adonis; ref. 69), analysis of similarity (ANOSIM; ref. 70), and the multi-response permutation procedure (MRPP; ref. 71), were used to test the differences in soil microbial communities between warming and control treatments. CCA (ref. 72) was performed to determine the linkage between environmental variables and microbial community composition. For selecting environmental variables, those containing redundant information were reduced to minimum number, keeping only the variables that had significant impacts. Also, the final sets of variables should have the variance inflation factors all <20. Finally, soil temperature, soil moisture and GPP remained in the CCA model (Supplementary Fig. 2a) of GeoChip data. Labile C pool 1 and 2 (%), and soil N content (%) were selected for 16S rRNA gene-based analysis (Supplementary Fig. 2b). The significance of the CCA model was tested by analysis of variance (ANOVA). According to CCA results, variation partitioning analysis was performed to determine the contribution of each individual variable or groups of variables to total variations in soil microbial community compositions. CCA was also used to determine correlations between abundance of subcategories of functional genes and the individual environmental variables (Supplementary Table 5). Two-tailed *t*-tests were performed to examine whether the differences between warming and control treatments were significant on the basis of several important biotic and abiotic variables (that is, soil C contents, aboveground biomass and total bacteria, archaea and fungi abundance) using Microsoft Excel 2010. ANOVA (ref. 73) was performed to test the treatment effect on the abundance of each functional gene involved in C and N cycling for GeoChip or relative abundances of OTUs of certain genus or phylum groups. In addition to the warming treatment effect, the probe OTU also factored into the model for partitioning the variance of probes within each functional gene. Response ratio was used to compute the effects of warming on functional genes relevant to GeoChip probes from shotgun sequences using the formula described in ref. 74.

Data availability. Raw shotgun metagenome, 16S rRNA and *nifH* amplicon gene sequences are available in the European Nucleotide Archive (<http://www.ebi.ac.uk/ena>) under study no. PRJEB10725. GeoChip raw and normalized signal intensities can be accessed through the URL ftp://129.15.40.240:8187/nclimate2940/Raw_GeoChip_Data.txt and ftp://129.15.40.240:8187/nclimate2940/Normalized_GeoChip_Data.csv.

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