Central metabolic responses of microorganisms to years and decades of soil warming

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Abstract

Microbial physiological responses to long-term warming are poorly understood. Here we applied metatranscriptomics to investigate how microorganisms react to medium-term (8 years) and long-term (>5 decades) subarctic grassland soil warming of +6 °C.

Decades, but not years, of warming induced changes in relative abundances of eukaryotic, prokaryotic, and viral transcripts and reduced functional richness. However, irrespective of the duration of warming, we observed a community-wide upregulation of central (carbon) metabolisms and cell replication in the warmed soils, whereas essential energy metabolism and protein biosynthesis complexes and pathways were downregulated. This coincided with a decrease of microbial biomass and lower soil substrate concentrations (e.g. dissolved organic carbon and phosphorus).

We conclude that permanently accelerated reaction rates at higher temperatures facilitate a downregulation of energy metabolism and protein biosynthesis, potentially freeing energy and matter for substrate acquisition and growth. This resource allocation seems to be a common response in microorganisms and allows sustaining high metabolic activities and replication rates even after decades of soil warming.
Introduction

Global temperatures and atmospheric carbon dioxide (CO$_2$) levels have increased steadily over the last 100 years$^{1,2}$. The terrestrial carbon (C) cycle feedback to the climate system represents a major uncertainty in the prediction of future temperatures$^3$. Soil microorganisms, including Bacteria, Archaea, Fungi, and protists, are responsible for the turnover of soil organic matter (SOM) and the subsequent release of CO$_2$ from soils to the atmosphere$^2$. Higher temperatures commonly lead to higher microbial activities, so global warming should accelerate the decomposition of SOM to CO$_2$$^4$.

On the other hand, SOM consists largely of microbial necromass and warming may stimulate microbial growth and thus necromass production$^{5,6}$, promoting SOM formation. Whether soils will ultimately act as C sinks or sources, thus depend on how microorganisms respond to long-term warming. Nevertheless, microbial responses to global warming are currently poorly represented in Earth system models$^7$, which can, to some extent, be attributed to the challenges associated with studying and quantifying microbial responses to environmental change in complex soil environments$^4$.

Microbial responses to long-term soil warming may include i) quantitative and compositional changes of microbial communities, ii) physiological adjustments of individual microorganisms, including changes in growth and resource use, via transcriptional and translational regulation, iii) shifts in microbial interactions and emergent properties of the community, and iv) microbial adaptations by genomic rearrangements and evolutionary changes of the genetic code.

In one of the few truly long-term warming studies (Harvard Forest Warming Experiment: 26 y, +5 °C, mid-latitude forest soil)$^8$, the observed warming effects on the microbial community included a decrease in fungal biomarkers and abundance, a decrease in microbial biomass, a community shift toward Gram-positive Bacteria, and an increase in bacterial evenness and abundance of Bacteria with low copy numbers of ribosomal RNA (rRNA) operons. A meta-analysis of 25 in situ soil warming experiments (1–15 y, +0.5–5.5 °C, various soil ecosystems) found initial increases in soil respiration
(46 ± 8%) due to warming followed by significant decreases over time. However, less than half of the studies estimated microbial biomass, and two thirds of the studies that quantified changes reported decreases in microbial biomass with warming. Increased rates of SOM degradation and soil respiration followed by a return to pre-warming rates within a few years were repeatedly observed. This pattern is often explained by an “acclimation of soil respiration”; a shift in the response of respiration to ongoing warming that leads to different sensitivities of soil respiration to temperature (and references therein). Alternatively, a return to pre-warming states can be explained by the depletion of easily degradable substrates.

It has recently been shown that natural geothermal activity can enable the study of soil warming on decadal time scales. We here make use of the longest in situ soil warming experiment worldwide, ForHot, in which a subarctic grassland site has been exposed to soil warming for decades (>50 y). More recent soil temperature gradients emerged nearby after an earthquake in 2008. The effects of long-term and short- to medium-term soil warming on abiotic and biotic properties and processes at these sites have been described in a range of publications. For example, Walker et al., Marañón-Jiménez et al., and Poeplau et al. identified considerable soil environmental changes in the warmed grassland plots, including reductions in topsoil C and nitrogen (N) pools by about 40% and decreased soil aggregation. These changes were accompanied by lower soil microbial biomass but higher soil respiration rates per unit of microbial biomass, contradicting the concept of a physiological acclimation of microorganisms. Amplicon sequencing of rRNA genes, on the other hand, did not indicate warming-induced changes of the microbial community composition at genus and operational taxonomic unit (OTU) level. Recently a meta-analysis on 128 measured variables at this site, including a broad variety of biotic and abiotic soil properties, pools, and processes was published, reporting a systemic overreaction to 5–8 years versus decades of warming. However, details of underlying microbial responses influencing the ecosystem-scaled responses remained largely unexplored.
Here we aimed at providing the first in-depth functional analysis of soil microbial responses to warming using a metatranscriptomics approach, i.e. high-throughput shotgun-sequencing of total RNA. In contrast to DNA based methods (e.g., metagenomics or amplicon sequencing), which can only show genetic potential and are often restricted to specific microbial groups, metatranscriptomics allows the comprehensive study of the soil microbiome, i.e., the entire active soil microbial community (Bacteria, Archaea, Eukaryotes, and viruses) and their functions, by studying expressed genes (messenger RNA (mRNA) and rRNA)\(^{21}\). We analysed 16 soil microbiomes. In situ microbial gene expression profiles of long-term warmed soils (LTW) exposed to +6 °C above ambient (\(E_t\)) for >50 y and medium-term warmed soils (MTW) exposed to +6 °C for 8 y were compared to ambient controls (\(A_t\)). Additionally, we measured dissolved and total soil C, N and phosphorous (P) concentrations and estimated microbial biomass. Our main objective was to elucidate if and how soil microorganisms alter their cellular activities and functions in response to warming.

**Results**

**Warming effects on soil physicochemical and biological properties**

Warmed soils differed from their ambient counterparts (Fig. 1b); total, dissolved organic, and microbial C, N, and P concentrations were lower in the warmed soils, mirroring previous studies that also report lower substrate concentrations and microbial biomass contents in the warmed soils at the same site\(^{15-18}\). Soil pH ranged from 4.4 to 6.0 and was slightly higher in the warmed soils (\(P < 0.05, n = 16, P_{corr} > 0.1, \text{Supplementary Table 1, 2}\)).
Figure 1. Grassland samples and warming-induced differences on physicochemical and biological properties. 

- **a** Schematic representation of the study sites, their development over time, and the metatranscriptomics samples (see Methods for details).
- **b** Trend charts (see Methods) indicating differences in DNA and RNA concentrations (per unit of soil), contents of dissolved organic C, N, and P (DOC, DON, and DOP, respectively), total C, N, and P (C\textsubscript{tot}, N\textsubscript{tot}, and P\textsubscript{tot}, respectively), microbial C, N, and P (C\textsubscript{MO}, N\textsubscript{MO}, and P\textsubscript{MO}, respectively), RNA:DNA ratio, and soil and microbial C:N ratios (see Supplementary Tables 1 and 2 for absolute values and significant differences, respectively).
- **c** DNA and RNA content per unit of microbial biomass and correlations between microbial biomass and DNA and RNA content per unit of soil (Supplementary Table 3).

DNA and RNA concentrations per unit of microbial C (C\textsubscript{MO}) had opposite trends, with higher DNA but lower RNA concentrations in the warmed soils, irrespective of the warming duration (Fig. 1c). While the RNA content per g soil positively correlated with C\textsubscript{MO} (r\textsubscript{s}(16) = 0.81, P\textsubscript{corr} < 0.001), the DNA content did not (r\textsubscript{s}(16) = 0.12, P\textsubscript{corr} = 0.664) (Fig. 1c). C\textsubscript{MO} was likewise positively correlated with total soil C, N, and P, dissolved organic C and P, microbial N and P, and water content (Supplementary Table 3).

To investigate what structural and functional changes in the soil microbiomes were associated with these substantial differences in microbial biomass and soil chemistry we performed metatranscriptomics.

**Compositional changes of the functional microbial communities across all domains of life**

Illumina paired-end sequencing produced an average of 6.69 × 10\textsuperscript{6} mRNA reads per sample (Supplementary Table 4). Bacteria dominated the mRNA transcript pools (93.36–98.52%), followed by Eukaryota (1.28–6.45%), Archaea (0.5–1.6%), and viruses (0.2–1.9%). Overall, more than 1,000
different families were detected, and mean family richness was lower in the heated soils, albeit not significant (Supplementary Fig. 1a). An NMDS analysis of mRNA reads assigned at family level indicated no clear overall separation between Aₜ and Eₜ samples and a greater variability between MTW replicates than LTW replicates (Supplementary Fig. 2a). Thus, we analysed the detailed taxonomic assignments of LTW and MTW separately (Fig. 2). In the long-term warmed soils, but not in the medium-term warmed soils, several taxa showed warming-induced differences in relative transcript abundances (Fig. 2a, Supplementary Table 6). Those that may be affected by warming include phylogenetic groups across all domains of life and included Fungi, protists, and Chloroflexi, which showed lower mean relative abundances in LTW-Eₜ and Deltaproteobacteria, Planctomycetes, Verrucomicrobia, as well as viruses, which showed higher mean relative abundances in LTW-Eₜ compared to LTW-Aₜ (Fig. 2a, b). A taxonomic analysis of rRNA reads corroborated the bacterial mRNA profiles, but the mean relative abundances of Eukaryotes differed between the rRNA datasets and the mRNA datasets (Supplementary Fig. 3).

We further screened all taxonomic ranks down to family level (Fig. 2c). Five taxa showed lower relative expression levels in LTW-Eₜ than LTW-Aₜ, and 17 taxa showed higher relative expression levels in LTW-Eₜ than LTW-Aₜ (Fig. 2c). Differential gene expression analyses underpinned these results, as many of the reported taxa were significantly different. In contrast, none of the six taxa that displayed higher relative expression levels in MTW-Eₜ than MTW-Aₜ was significantly differential expressed (Fig. 2c). We repeated the analysis on the bacterial fraction of the mRNAs to test whether Eukaryota abundances influenced the bacterial patterns and found almost identical warming-induced abundance patterns (Supplementary Fig. 4). We also attempted to increase the taxonomic resolution by assembling the metatranscriptome reads (Supplementary Methods). Our approach using rnaSpades²², however, did not provide a sufficient number of long mRNA contigs (<10% of functionally annotated mRNA contigs were longer than the unassembled reads).
Fig. 2. Taxonomic annotation of mRNA reads and warming-induced abundance patterns. a Sankey plots showing both the fraction of mRNA reads (mean over all LTW and MTW replicates, respectively) assigned to domains and the composition and relative abundances of eukaryotic and bacterial mRNAs. The depicted groups account for 100% of eukaryotic and bacterial mRNAs, respectively. All bacterial phyla with a relative abundance of ≥1% in at least one of the sampled soil groups are depicted; the remaining phyla are summed (other Bacteria). Potential warming-induced differences in mRNA abundances are indicated with + (t-tests, n = 8, P < 0.05, Pcorr < 0.1; Supplementary Table 6). b Log2-fold changes of mean relative abundances between At and Er of the taxa listed left-hand. c Exploratory analysis showing warming-induced taxon abundance patterns. Only taxa with higher or lower relative abundances in all four replicates of one group relative to their counterparts are depicted (4/4 filter, see Methods). Bacterial taxa are shown at family level; higher taxonomic levels are only shown if no family belonging to these higher levels passed the filter. Higher bacterial taxonomic levels are accordingly not shown if any family belonging to these levels passed the filter. Potential warming-induced differences in mRNA abundances are indicated (differential gene expression analysis, n = 8, Pcorr < 0.05 (*), Pcorr < 0.1 (+); Supplementary Table 7).
We subsequently analysed functional mRNA annotations to identify how soil warming influences the transcription of genes involved in central metabolic functions and cellular processes.

**Warming-induced gene expression profiles**

An average of $2.66 \times 10^6$ mRNA reads per sample (Supplementary Table 4) was assigned to a molecular function (KO number) defined in the KEGG Orthology database. KEGG offers a hierarchical structure with four layers, hereafter referred to as KEGG1 (i.e. *Metabolism*, *Genetic* and *Environmental information processing*, and *Cellular processes*), KEGG2 (see Fig. 3), KEGG3 (i.e. pathways and functional complexes), and KO (molecular functions, i.e. enzymes and enzyme subunits). Significantly fewer unique functions (KOs) were detected in LTW-E than LTW-A (t-tests, $n = 8$, $P < 0.05$), but not between MTW-E and MTW-A (Supplementary Fig. 1b). PERMANOVA analyses further revealed a significant effect of warming on expressed functions (KOs), and the functions expressed by different microbial families (Supplementary Fig. 2c, d; Supplementary Table 5).

To identify the nature of this functional response we explored in more detail the functional assignments to KEGG1 categories *Metabolism* (65.4%), *Genetic information processing* (16.8%), *Cellular processes* (9.0%), and *Environmental information processing* (8.9%) (Fig. 3). Transcripts for *Metabolism* and major *Metabolism* subcategories such as *Carbohydrate metabolism*, *Lipid metabolism*, and *Nucleotide metabolism* had higher relative abundances in the warmed soils, while *Genetic information processing* subcategories such as *Transcription* and *Translation* had lower relative abundances (Fig. 3, Supplementary Table 8). Transcript patterns for *Cellular processes* and *Environmental information processing* were not consistent.
Fig. 3. Functional annotation of mRNA reads. Sankey diagram showing the mean relative abundances of KEGG1 and KEGG2 categories over all samples. The heatmap depicts the relative abundances of all KEGG2 categories with mean relative abundances >1%; samples are sorted by soil temperature (\(A_T\), ambient soil temperature; \(E_T\), +6 °C). The heatmap colour code indicates the KEGG1 affiliation of the KEGG2 categories. Potential warming-induced differences in mRNA abundances are indicated (differential gene expression analysis, \(n = 16\), \(P_{corr} < 0.01 (**), P_{corr} < 0.05 (*)\), \(P_{corr} < 0.1 (+)\); Supplementary Table 8).

**Community-wide shifts in gene expression of central metabolisms and cellular functions**

Since the gene expression patterns observed within the broad functional categories suggested functional temperature-dependencies (Fig. 3) we screened all KEGG3 categories to identify which specific metabolic pathways and functional complexes were the basis for these overall patterns. One third of all functionally annotated mRNAs were affiliated with KEGG3 categories that exhibited warming-induced abundance patterns similar to those in Fig. 3 (Supplementary Fig. 5). Out of 55 KEGG3 categories with a visually distinct pattern, 13 showed lower relative expression levels in \(E_T\) than \(A_T\), while 42 KEGG3 categories showed higher relative expression levels in \(E_T\) than \(A_T\). We further investigated the most abundant of these 55 KEGG3 categories, each making up >1% of total annotated mRNA reads. Eight of these abundant categories showed higher relative expression levels...
in E7 and encompassed central C metabolisms and metabolic pathways associated with amino acids and nucleotides, while five showed higher relative expression levels in A7 and included protein biosynthesis related complexes and Oxidative phosphorylation (i.e. ATP formation) (Fig. 4a). This change in the transcript abundances for the most central metabolisms in organismal function was observed for multiple taxa (Fig. 4b). In the long-term warmed soils, four phyla, eight classes, four orders, and four families, representing 33.9%, 38.1%, 11.1%, and 1.8% of the LTW metatranscriptomes, respectively, displayed this pattern, confirming that this is a taxonomically broad type of warming response. This was supported by the analysis of the medium-term warmed soils, where seven phyla, 14 classes, seven orders, and eleven families, representing 87.2%, 65.0%, 24.1%, and 6.0% of the MTW metatranscriptomes, respectively, displayed the same expression pattern. It should be noted that the low percentages at family level is due to an average of only 22.1% of the annotated mRNA reads being assigned to a family-level taxon. Interestingly, the pattern occurred in taxa (Fig. 4b taxa in bold) regardless of whether these taxa became more active (higher relative abundance of mRNA and rRNA) or less active with warming (Fig. 2, Supplementary Fig. 3), indicating that the physiology of these community members had been altered in a similar way. Furthermore, our extensive screening revealed that on average this was true for nearly two thirds (63.7%) of the taxa present in LTW and MTW (Fig. 4c, Supplementary Fig. 6). However, the percentage of taxa expressing the pattern varied between 43 and 95% depending on the KEGG category and the warming duration and was highest for the KEGG3 categories related to RNA, protein and energy metabolisms.
Fig. 4. Overall and taxon-specific warming-induced transcript abundance patterns. a Pathways and complexes with mean relative abundances ≥1% that passed the differential abundance-pattern filter (13/16-filter, see Methods). Colour code indicates the corresponding KEGG1 category. Differential gene expression analysis results are indicated next to the KEGG 3 categories (n = 16, \( P_{\text{corr}} < 0.05 \) (*), \( P_{\text{corr}} < 0.1 \) (+); Supplementary Table 8). b Taxa reflecting the observed warming-induced abundance patterns presented in (a). All taxa with a mean relative abundance ≥1% in LTW-\( \Delta T \), LTW-\( \Sigma T \), MTW-\( \Delta T \), or MTW-\( \Sigma T \) and not more than two deviations from the overall pattern are depicted. Numbers in square brackets next to the taxonomic ranks are the sums of the relative abundances of the individual taxa represented below (grey circles). Colour code indicates the mean relative abundance of a KEGG3 category: orange, higher expression levels in \( \Sigma T \); blue, higher expression levels in \( \Delta T \). Pale, dark, and intermediate colours indicate how strong and widely distributed a pattern was across LTW and MTW soils, respectively (see legend “Expression levels”). See (a) for KEGG3 abbreviations. c Table listing the percentage of all abundant taxa (i.e. taxa with >1‰ relative abundance) within a taxonomic rank that featured the overall warming-induced abundance pattern of a specific KEGG3 category (see Supplementary Fig. 6 for details on the taxa). Numbers in square brackets next to the taxonomic ranks give the total number of abundant taxa within the taxonomic rank. Bold numbers indicate pathways and complexes with warming-induced differential abundance patterns observed in more than two thirds of all abundant taxa within a taxonomic rank.
Warming effects on microbial growth, energy metabolism, protein biosynthesis and C metabolisms

In addition to Glycine, serine & threonine metabolism and Amino sugar & nucleotide sugar metabolism (Fig. 4), the less abundant amino acid metabolisms Phenylalanine, tyrosine & tryptophan biosynthesis, D-Alanine, D-Glutamine & D-glutamate metabolism, and Lysine biosynthesis showed higher relative transcript abundances in the warmed soils (Supplementary Fig. 5). Higher relative abundances of Pyrimidine, and Purine metabolism transcripts were also seen (Fig. 4). This indicated an upregulation of the production of major building-blocks of nucleic acids and proteins in the warmed soils, prompting us to investigate warming effects on bacterial growth (cell division), energy metabolism, and protein biosynthesis in detail. The eukaryotic fraction was excluded from the further analyses due to the large variations between LTW and MTW (Fig. 2).

**Fig. 5. Warming-induced abundance profiles of bacterial growth-related transcripts.** Boxplots showing relative transcript abundances of the KEGG3 categories Peptidoglycan biosynthesis, Glycerolipid metabolism, and DNA replication, and FtsZ in the bacterial fractions of the LTW-A1, LTW-E1, LTW-A2, and LTW-E2 metatranscriptomes. The depicted KEGG3 categories are involved in the build-up of new cell walls and cell membranes and responsible for the duplication of genomic DNA, which precedes bacterial cell division; and FtsZ represents a key enzyme in bacterial cell division (see schematic drawings above the boxplots). P-values ($P_{\text{corr}}$) indicating significant differences are displayed above boxplot-pairs (Supplementary Table 9).

We observed trends of warming-induced transcriptional responses in a broad set of genes fundamental to bacterial cell replication (Fig. 5, Supplementary Table 9). In LTW, we observed
increased relative transcript abundances for Peptidoglycan biosynthesis, DNA replication and FtsZ in E, albeit insignificant. However, in MTW-E, relative transcript abundances for Peptidoglycan biosynthesis, Glycerolipid metabolism, and DNA replication were significantly higher in MTW-E than MTW-A (t-tests, n = 8, $P_{corr} < 0.05$, Fig. 5), while the trend for FtsZ transcripts was similar but insignificant.

While the relative abundances of cell replication-related transcripts increased with warming, transcripts for energy conservation (Oxidative phosphorylation) showed the opposite trend. All four enzyme complexes of the membrane-bound prokaryotic electron transport chain and ATP synthase (producing ATP, the energy currency of cells) showed lower mean relative transcript abundances in E (Fig. 5a). In contrast, transcripts of enzymes providing inorganic phosphate ($P_i$) to the ATP synthase, especially polyphosphate kinase, showed higher mean relative abundances in E than A.

Similar to Oxidative phosphorylation (i.e. ATP formation), transcripts for multiple complexes related to protein biosynthesis showed a trend towards lower relative abundances in E (Fig. 6b). While insignificant after correcting for multiple testing ($P_{corr} > 0.05$, Supplementary Table 10), such trends were also observed for RNA polymerase, ribosomal proteins, ribosomal RNA, the main molecular chaperones GroEL and DnaK (involved in protein folding), and the Sec-dependent pathway, the dominant protein export pathway present in the metatranscriptomes. Contrary to all other categories of the protein biosynthesis machinery, transcripts for RNA degrading complexes (RNA degradosomes) showed higher relative abundances in E than A (Fig. 6b).

We also investigated in detail the gene expression profiles for central C metabolisms. While the transcript abundances of KEGG3 categories Pyruvate metabolism, Glycolysis/Gluconeogenesis, Methane metabolism, and C fixation pathways in prokaryotes pointed towards higher expression levels in the warmed soils (Fig. 4), these differences were not significant. This suggests that while a shift in community metabolism is indicated, this might not be reflected consistently in all pathway steps and responsible bacteria that contribute to the above mentioned KEGG3 categories.
Fig. 6. Warming-induced abundance profiles of transcripts related to bacterial energy metabolism and protein biosynthesis. Boxplots show the relative transcript abundances of enzymes and enzyme complexes involved in (a) membrane-bound electron transport and ATP synthesis (oxidative phosphorylation) and (b) protein biosynthesis in the LTW-ATL, LTW-ET, LTW-ATL, and LTW-ETL metatranscriptomes (see Supplementary Data 1 for a list of KOs summarised in the boxplots). Schematic representations of the enzymes and enzyme complexes are provided above each boxplot and are based on the KEGG pathway drawings. Membrane-bound complexes are embedded in a lipid bilayer. Potential warming-induced differences are indicated by p-values (uncorrected) < 0.05 (*) and corrected p-values < 0.1 (+) (t-tests, n = 8, Supplementary Table 10). GroEL and DnaK are part of the KEGG category RNA degradation, thus skewing its overall warming-induced abundance pattern towards lower transcript abundances in ET than AT as seen in (Fig. 4a).

Finally, we examined, within all KEGG3 categories, the differential expression of single functions (KO). Notably, the observed patterns, including those not significant at the KEGG3 level, were confirmed by multiple functions (KO) that were differentially expressed and significant after multiple testing.
correction (Supplementary data S2). For functions related to protein biosynthesis and oxidative
phosphorylation (ATP formation) the patterns were particularly strong and supportive of the above
described trends. Less, although substantial, support was found for central carbon and amino acid
metabolism patterns.

**Discussion**

Here we used metatranscriptomics to elucidate how soil microorganisms change their gene
expression of central metabolic functions and cellular processes in response to warming. We showed
that long-term (>50 y), but not medium-term (8 y), soil warming (+6 °C) resulted in a significant,
albeit small, decrease of unique molecular functions encoded by soil Bacteria, Archaea, Fungi, and
microbial Eukaryotes. However, irrespectively of the duration of warming, physiological responses to
warming as shown by the transcriptional tuning of central metabolisms for C, energy, protein
biosynthesis, and growth, were consistent and common across many community members.

The apparent overexpression of genes involved in **Pyruvate metabolism**, **Glycolysis/Gluconeogenesis**, **Methane metabolism**, and **C fixation pathways in prokaryotes** indicate that long-term as well as
medium-term warming affect the central cellular C metabolism of soil microorganisms. However,
inconsistencies in the relative transcript abundances of subjacent KOs indicated that the perceived
upregulation of these metabolisms does not involve all associated enzymes, reflecting either how
metabolic pathway efficiency can be controlled by only a few rate-limiting steps\(^4\) or that the
patterns created by the distinct up-regulation of pathways in some organisms are diluted by other
organisms that share pathway steps but not the response pattern.

In contrast, the reduction in relative transcript abundances of oxidative phosphorylation complexes,
RNA polymerases, protein processing enzymes, and ribosomal proteins were consistent across
multiple enzyme subunits and pathway steps. A downregulation of the protein biosynthesis
machinery is not immediately recognizable as a strategy to counteract substrate limitation or maximize growth rates. However, a reduction of ribosomes may be biochemically favourable under the warming conditions. Ribosomes are macromolecular complexes that are the sites of protein synthesis (translation). They consist of ribosomal proteins and rRNAs and are present in tens of thousands of copies per bacterial cell. Thus, ribosomes can comprise over one third of the dry cell mass and rRNAs can account for >90% of the total cellular RNA content. Starving *Escherichia coli* and *Salmonella* spp. cells reduce their ribosomal content, suggesting that a downregulation of the translation machinery, which accounts for up to 40% of total cellular proteins, is metabolically favorable in nutrient-limited ecosystems. Furthermore, carefully tuning concentrations of abundant proteins such as ribosomal proteins can also free resources for accelerating other reactions. We not only demonstrated lower relative transcript abundances of ribosomal proteins and protein biosynthesis-related enzymes in the warmed soils, but our results also showed lower microbial biomass per gram of soil dry weight, and a decrease in RNA per gram dry soil that correlated significantly with the biomass. A significant decrease in microbial biomass per gram of soil dry weight in the warmed soils has been observed previously at the same site. Furthermore, the transcriptional patterns for protein biosynthesis were consistent for many community members, regardless of their relative abundances of total mRNAs and rRNAs in the long- and medium-term warmed soils and the controls. Thus, a downregulation of protein biosynthesis-related enzymes and a reduction of ribosomes might be a common response of microorganisms, especially bacteria, to ecosystem warming and partly responsible for the lower microbial biomass in the warmed soils. It seems contradictory that we observed, at the same time, higher relative transcript abundances for enzymes related to cell division and biosynthesis of amino acid, the building-blocks of proteins. However, a reduced ribosomal content is not necessarily linked to lower protein synthesis rates or lower growth rates. It is long known that increased temperatures accelerate mRNA synthesis and the protein synthesis rate per ribosome (peptide chain elongation rate) in the model organism *Escherichia coli*. Conversely, it was indicated more recently that lower growth rates induced by slower reaction rates...
are associated with an increased content of ribosomal proteins\textsuperscript{30,33}. Likewise, an acceleration of reaction rates in the membrane-bound electron transport chain and ATP synthesis (\textit{Oxidative phosphorylation}) despite lower expression might also have been driven by temperature and might be indicated by the higher relative transcript abundances of enzymes (i.e. polyphosphate kinase) providing $P_i$ to the ATP synthase in the warmed soils. Since the translation machinery accounts for up to 40\% of total cellular proteins\textsuperscript{30} and protein biosynthesis is the most costly type of macromolecular synthesis\textsuperscript{34–36} the reduced number of ribosomes would furthermore allow conservation and reallocation of energy (ATP) from the generation of ribosomal proteins to the synthesis of metabolic proteins.

Accelerated reaction rates due to increased temperatures presumably allowed microorganisms to reduce their ribosomal content and their machinery for oxidative phosphorylation, liberating energy and matter for substrate acquisition and growth. Through this transition, soil warming may have led to metabolically more active microbial cells driven by temperature and a different partitioning of cellular resources. Previous observations of higher biomass-specific growth, respiration, organic C uptake, and turnover rates in the same long-term warmed soils\textsuperscript{16} corroborates this interpretation. These results are clearly not consistent with the suggested thermal acclimation of soil microorganisms used to explain a return to pre-warming SOM degradation rates after a few years of warming\textsuperscript{11,12} (and references therein). Rather, the apparent thermal acclimation in soil respiration rates might be driven by a reduction in microbial biomass caused by reduced carbon and nutrient concentrations, as previously suggested\textsuperscript{16}. However, our results point at a different type of microbial acclimation to warming where the physiological adjustments allow the microbial community members to maintain a high activity even after decades of warming, despite more limiting conditions.

The microbial functional responses were more pronounced and widespread in the microbial communities in the medium-term warmed soils than in the long-term warmed soils reflecting the
recently reported systemic overreaction to years versus decades of warming in the same soils\(^{15}\). The authors proposed that an initial acceleration of biotic activity due to warming led to rapid decreases in C, N, and P pools within the first years after the onset of warming, followed by a decrease of microbial and fungal biomass and system stabilisation within decades. Our taxonomic analyses of the mRNA transcript pools extended our insight into this proposed soil warming chronology; long-term, but not medium-term, soil warming resulted in differential abundances. Relative mRNA transcript abundances of several bacterial taxa, fungi, protists, and viruses were affected by long-term warming. This indicated shifts in trophic interactions (e.g. from top-down to bottom-up control of the prokaryotic community) and possibly a reduced importance of fungi, with repercussion on organic matter decomposition. Reduced Fungi:Bacteria ratios have been described previously in long- and short-term warming experiments of forest and grassland soils\(^{8,37}\) and Fungi appear to be more abundant and active at lower temperatures\(^4,38\) and in soils with low pH\(^{39,40}\). Besides the higher temperatures and slightly higher pH in the warmed ForHot grassland soils, the lower concentrations of various soil C, N and P compounds and decreased soil aggregate sizes\(^{19}\) may explain the lower relative abundances and diversity of fungal mRNAs. The few and non-significant taxonomic differences between ambient and medium-term warmed soils suggest that while the soil microorganisms respond functionally, there is little effect on microbial community structure. However, we cannot exclude that the apparent lack of taxonomic response after eight years resulted from the high variation between the biological replicates. Our results somewhat contrasts previous studies on the ForHot grassland sites that reported generally little taxonomic shifts with warming\(^{16,20}\). However, these studies applied DNA-based methods (i.e., 16S gene and ITS amplicon sequencing) targeting the potential microbial community, including active cells, dormant cells, and relic DNA. These methods are also prone to biases from PCR and primers. In contrast, metatranscriptomics is PCR-free and deploys random hexamer-primers targeting prokaryotic, eukaryotic, and viral transcripts in parallel, thus allowing a comprehensive analysis of the active soil microbiome. Furthermore, our analyses did demonstrate a stronger effect of warming on microbial functions.
(KOs) than on taxonomy. This functional response was similar in medium-term and long-term warmed soils. Thus, our study provides evidence for a decoupling of microbial community structure and functions, as recently suggested from several ecosystems including soil$^{41-44}$. How warming affects the microbial control of the global C cycle is a key question to better understand the soil-climate feedbacks, an answer to which is urgently needed$^{4,45}$. Here, for the first time in a comprehensive study of the transcriptional response of microorganisms across all domains of life, we show that downregulation of energy metabolism and protein biosynthesis is central in the microbial soil warming response that allow microorganisms to maintain high metabolic activities and cell division rates even after decades of warming. We suggest that accelerated biochemical reaction rates due to higher temperatures, have a positive feedback on central metabolisms via increased relative transcript abundances, if not constraint by reduced substrate concentrations, further accelerating microbial decomposition of SOM and the subsequent release of CO$_2$ to the atmosphere.
Methods

Grassland sites and soil sampling

Soil samples were collected from a natural grassland near Hveragerði (64°00′N, 21°11′W), Iceland, in July 2016. The grassland is part of the ForHot experiment\textsuperscript{14} and features two sites, each consisting of replicated soil temperature gradients that were formed by natural geothermal activity. One site has experienced warming for >50 y, possibly since before 1708\textsuperscript{14}. Geothermal activities may have varied over time, but warming has been stable in the area since at least 1963, and the warming intensities of the temperature gradients have not varied since detailed measurements began\textsuperscript{14–16}. The second site recently developed similar temperature gradients after an earthquake in 2008. We collected 16 samples that were later analysed in detail; i.e. four samples of long-term warmed soils (LTW) exposed to elevated temperatures (+6 °C above ambient, LTW-E\textsubscript{T}) and four corresponding controls at ambient temperatures (LTW-A\textsubscript{T}) as well as four samples of medium-term (8 y) warmed soils (MTW) exposed to +6 °C above ambient temperatures (MTW-E\textsubscript{T}) and four corresponding controls (MTW-A\textsubscript{T}). The grassland soils are classified as Silandic Andosols, and both sites are dominated by Agrostis capillaris with varying undergrowth and vascular pant and moss cover\textsuperscript{14}. We took soil samples (0–10 cm depth) from ambient and +6 °C plots of four replicate blocks at one time point. Samples were immediately frozen in liquid N\textsubscript{2} for subsequent nucleic acid extraction and metatranscriptomics after sieving to 2 mm.

Physicochemical soil properties

Fresh aliquots of each soil sample were extracted with either 1 M KCl or 0.5 M NaHCO\textsubscript{3} solution for 30 min at room temperature or fumigated with chloroform for 48 h and subsequently extracted using the same extractants. Various soil C and N compounds and soil P compounds (Supplementary Table 1) were quantified in the KCl and NaHCO\textsubscript{3} extracts, respectively, using standard procedures\textsuperscript{46}. C\textsubscript{MO}, N\textsubscript{MO}, and P\textsubscript{MO} were calculated as the differences between DOC, TDN, and TDP contents in the fumigated and non-fumigated extracts. Total C and N contents and stable isotope ratios were
analysed in dried soil aliquots using an elemental analyser coupled to an isotope ratio mass spectrometer (EA-IRMS; EA1110 coupled via a ConFlo III interface to a DeltaPLUS IRMS, Thermo Fisher Scientific). Soil pH was determined from sieved soil samples in 0.05 M CaCl₂ solution and gravimetric water content was determined.

**Nucleic acid extractions and sequencing**

We extracted total nucleic acids from 16 sieved soil samples (Supplementary Table 7), i.e. four replicates from each of the four sampled soil groups (LTW-A₁, LTW-E₁, MTW-A₁, MTW-E₁), representing two warming intensities (A₁, +0 °C, and E₁, +6 °C) of >50 y warmed soils (LTW) and 8 y warmed soils (MTW). These samples were selected because 6 °C above ambient is within the predicted and most severe range of (soil) warming over the next 60–100 y⁴⁷,⁴⁸. Each sample was extracted in triplicate, as previously described⁴⁹. Briefly, a phosphate buffer, a detergent solution containing CTAB, and TE saturated phenol (pH 8) were added to 0.3 g of soil in a lysis matrix E tube (MP Biomedicals) containing silica beads and shaken vigorously for 30 s (6.5 m s⁻¹) in a FastPrep machine (MP Biomedicals). After centrifugation, the aqueous supernatant was retained. This procedure was repeated two more times using fresh buffer, detergent, and phenol. The three supernatants were then pooled, followed by phenol:chloroform:isoamylalcohol (25:24:1) and chloroform:isoamylalcohol (24:1) extraction and precipitation of the nucleic acids using PEG 8000. The nucleic acids were treated with DNase (RQ1, Promega) before the metatranscriptomes were generated. RNA quantity and quality were assessed by automated agarose gel electrophoresis (Bioanalyzer 2100, Agilent), a NanoDrop™ spectrophotometer (ND-1000, Peqlab), and a RiboGreen™ assay kit (Thermo Fisher Scientific). Total RNA was amplified using the MessageAmp II-Bacteria Kit (Ambion Life Technologies) with an input of 100 ng RNA, following the kit protocol, except the RNA was linearly amplified for 14 h. The three technical replicates of each of the 16 samples were pooled prior to the amplification. Overlapping paired-end 125-bp reads were sequenced using the HighSeq2500 sequencer (Illumina) at the Vienna Biocenter Core Facilities, Vienna, Austria.
Sequence data pre-processing

We performed the following pre-processing steps and the majority of the subsequent analyses using the Life Science Compute Cluster (LiSC) run by CUBE (Division of Computational Systems Biology), Department of Microbiology and Ecosystem Science, University of Vienna, Austria. PEAR v.0.9.10 (Paired-End reAd mergeR, default settings) was used to merge the raw paired-end reads. We subsequently used SortMeRNA v.2.1 to filter out non-rRNA reads from the total RNA reads. The non-rRNA reads were quality filtered (-min_len 180 -max_len 240 -min_qual_mean 30 -ns_max_n 5 -trim_tail_right 15 -trim_tail_left 15) using prinseq-lite v.9.20.4. A second filtering step was performed to obtain mRNA reads and reduce the size of the dataset for later analyses. All non-rRNA reads that aligned to any sequence in the NCBI nr database (as of September 2018) using a non-conservative DIAMOND blastx search (v.0.9.18, -k 1 --min-score 52.5) of the same data set against the KEGG database as of February 2019). We, therefore, used a bit score of 52.5 rather than an e-value of 0.0001 for all following analyses to obtain a better comparability between different database searches, independent of increasing database sizes. We performed a DIAMOND blastx search (v.0.9.25, -k 25 --min-score 52.5) against the NCBI nr database (as of October 2019) to obtain taxonomic information for the mRNA reads. We used the program blast2lca, the standalone implementation of the MEGAN (v.6.11.1) LCA (lowest common ancestor)-assignment algorithm, to obtain one taxonomic assignment for each read based on the 25 nr database hits. The following parameters were used to obtain the LCAs: -ms 50 -me 0.01 -top 5 -mid 0.0 (mapping file: prot_acc2tax-Jul2019X1.abin). Subsamples of 200,000 rRNA reads were taxonomically annotated as previously described.

Functional and taxonomic annotation

We functionally annotated the total mRNA reads (Supplementary Table 7) by aligning them to the KEGG database (as of April 2019) using a DIAMOND blastx search (v.0.9.18, -k 1 --min-score 52.5). A prior analysis indicated that at a bit score of 52.5 equalled an e-value of ≤0.0001 (DIAMOND blastx of the same data set against the KEGG database as of February 2019). We, therefore, used a bit score of 52.5 rather than an e-value of 0.0001 for all following analyses to obtain a better comparability between different database searches, independent of increasing database sizes. We performed a DIAMOND blastx search (v.0.9.25, -k 25 --min-score 52.5) against the NCBI nr database (as of October 2019) to obtain taxonomic information for the mRNA reads. We used the program blast2lca, the standalone implementation of the MEGAN (v.6.11.1) LCA (lowest common ancestor)-assignment algorithm, to obtain one taxonomic assignment for each read based on the 25 nr database hits. The following parameters were used to obtain the LCAs: -ms 50 -me 0.01 -top 5 -mid 0.0 (mapping file: prot_acc2tax-Jul2019X1.abin). Subsamples of 200,000 rRNA reads were taxonomically annotated as previously described.
Data analysis

We used R56 to analyse the data, perform statistical tests, and graphically display the results (Rstudio version 1.1.456) including the R packages ggalluvial version 0.10.0 (https://CRAN.R-project.org/package=ggalluvial), ggplot2 version 3.2.0 (https://CRAN.R-project.org/package=ggplot2), vegan version 2.5-6 (https://CRAN.R-project.org/package=vegan), and edgeR version 3.32.0 (https://bioconductor.org/packages/edgeR). Further details on the R packages can be found in the respective sections. Functionally annotated and taxonomically classified mRNA datasets were merged. Functionally annotated mRNA reads that lacked a taxonomic classification were tagged with “no match”. The resulting data were normalised and filtered.

Filtering and normalisation. First, we normalised the data to the library size and transferred them to permille (‰). Second, only functions (KOs, KEGG database entries) present in all four replicates of at least one group (LTW-ATai, LTW-ETai, MTW-ATei, or MTW-ETei) were kept (removing 0.6‰ of the data). Third, low abundant functions were removed by filtering out KOs with a total relative abundance (sum of all 16 datasets) below <0.1‰ (removing 4.6‰ of the data remaining after the previous step), which equalled a mean relative abundance of a specific function per sample of <0.00625‰ (mean relative abundance [‰] = 0.1‰/16). A second dataset containing all taxonomically classified mRNA reads (without functional annotation) was likewise normalised and transferred. The taxonomy-only dataset was filtered by removing all families (taxonomic strings from domain to family) that were not present in all replicates of at least one group (LTW-ATai, LTW-ETai, MTW-ATei, or MTW-ETei) (removing 0.07‰ of the data); no second filter was applied. Viral reads were summarised prior to filtering depending on the subsequent analyses.

Trend charts. Trend charts (e.g. Figure 1C) were created by calculating the means of AT and ET replicates, respectively (LTW and MTW separate and combined) and comparing them to the mean over all samples (AT and ET combined, LTW and MTW separate and combined) which was set to 1. The shapes of the individual wedges reflect the magnitude of the differences. Wedges are coloured alternating (brown and grey) to ease discriminability. Dark brown and dark grey indicate values >1
(i.e. above the over-all mean) while light brown and light grey indicate values <1 (i.e. below the over-
all mean).

**Heatmaps.** Heatmaps were generated using the `geom_tile` function of the ggplot2 R package. Before plotting, normalised data were transformed by z-scoring, either over all 16 samples or separately for TLW and MTW. Two explorative filters were subsequently applied for selecting patterns of interest.

Abundance patterns of taxa present in LTW and MTW, respectively, were subject to a stringent filter (referred to as 4/4-filter); only taxa with higher or lower relative abundances in all four replicates of one group relative to their counterparts passed the filter threshold (e.g. a taxon passes the filter if the four highest values are found in LTW-A and the four lowest in LTW-E). A less stringent filter (referred to as 13/16-filter) was applied when the relative abundances of KEGG functions between A and E across all samples were compared (e.g. Figure 4). Patterns were retained only if: i) at least six samples of one temperature group (A or E) were higher than the third highest sample of the other temperature group (E or A) and at least seven samples of one temperature group (E or A) were lower than the third lowest sample of the other temperature group (A or E) or ii) at least seven samples of one temperature group (A or E) were higher than the third highest sample of the other temperature group (E or A) and at least six samples of one temperature group (E or A) were lower than the third lowest sample of the other temperature group (A or E). Therefore, the critical threshold to pass the 13/16-filter lies at 80% consensus with the most stringent warming-associated distribution (i.e. the eight highest relative transcript abundances are found in one temperature group and the eight lowest in the opposite temperature group).

**Alluvial plots.** Alluvial plots (Sankey diagrams) were created using the R package ggalluvial. Individual Sankey diagrams were manually merged if more than two levels were shown.

**Boxplots.** Boxplots were generated using `geom_boxplot` (R package ggplot2).

**Non-metric multidimensional scaling (NMDS).** NMDS was used to obtain ordination plots depicting (dis)similarities between the microbial functions and microbial community structures of the samples. We used the `metaMDS` function implemented in the R package vegan, two dimensions, and a
maximum of 10,000 random starts in search of a stable solution. The sequencing data were normalised and filtered as described above prior to the NMDS analyses. GUSTA ME (GUide to STatistical Analysis in Microbial Ecology)\textsuperscript{37} was consulted for selecting the appropriate dissimilarity index (i.e. “canberra”).

**Statistics and post hoc analyses.** The basic R function *t.test* was used to perform two-sided Student’s *t*-Tests to identify significant differences between A\textsubscript{T} and E\textsubscript{T} (of LTW and MTW, respectively, or across LTW and MTW). Obtained p-values were corrected for multiple testing (Benjamini-Hochberg procedure, basic R function *p.adjust*). Corrected p-values (P\textsubscript{corr}, q-values) < 0.05 were considered to indicate significant differences. We deliberately chose a parametric test (decreasing the chance of making a type II error) combined with multiple-testing adjustment and considered corrected p-values < 0.1 to indicate a temperature-dependent trend, reflecting the explorative nature\textsuperscript{58} of our study.

The basic R function *cor.test* was used to identify associations between microbial biomass and RNA content, DNA content, and various C, N, and P concentrations by applying Spearman’s rank correlation (two-sided). The ggplot function *geom\_smooth* was used to indicate correlations (method = *lm*). Taxonomic (family) and functional (KO) richness was estimated from raw read counts (families and KOs not present in all four replicates of at least one group (LTW-A\textsubscript{T}, LTW-E\textsubscript{T}, MTW-A\textsubscript{T}, or MTW-E\textsubscript{T}) were considered as noise and excluded) using the vegan function *rarefy*. Log2-fold changes in transcript abundance between temperature group (GOA\textsubscript{T}, GOE\textsubscript{T}, GNA\textsubscript{T}, and GNE\textsubscript{T}) means were calculated in R: log2(mean E\textsubscript{T}/mean A\textsubscript{T}). Permutational multivariate analysis of variance (PERMANOVA, vegan function *adonis*) was used to identify the effect of warming (A\textsubscript{T} and E\textsubscript{T}) and warming duration (LTW and MTW), respectively, on the distribution of samples (see NMDS plots) by physicochemical soil properties (including microbial biomass measures, DNA, and RNA concentrations) or gene expression. In all cases, 10,000 permutations were calculated, and the dissimilarity indices were the same as those used in the NMDS analyses (see above). Differential gene expression analyses on taxonomic and functionally annotated datasets were performed using edgeR (function *glmQLFTest*). Raw read counts of families or functions (KOs, KEGG database entries)
present in all four replicates of at least one group (LTW-A\textsubscript{T}, LTW-E\textsubscript{T}, MTW-A\textsubscript{T}, or MTW-E\textsubscript{T}) were used as input data. Low abundant taxa (kingdoms, phyla, classes orders, and families <1\%), low abundant KEGG categories (KEGG 1, KEGG 2, and KEGG 3 <1\%), and low abundant functions (KO <0.5\%) were excluded from the analyses and the default trimmed mean of M-values normalization (TMM) method was used to normalize the data.

Adobe Illustrator (CC 23.0.2.) was used for final figure editing.
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Author contributions

AR, ATT, TU, and AS conceived the study. BDS, AR, IJ, and JPe established the experimental sites. AR and JPr collected the samples. ATT, JPr, and JS processed the samples in the lab. AS, ATT, and MBD analysed the sequencing data. AS created the figures. AS and ATT wrote the manuscript with inputs from all authors.

Competing interests

The authors declare no competing interests.

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Additional information

Supplementary information, figures, and tables are available. The raw sequence data are available at the NCBI Sequence Read Archive (SRA); BioProject ID: PRJNA663238, accession numbers SAMN16124403–SAMN16124422.