Atmospheric trace gases support primary production in Antarctic desert surface soil

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Cultivation-independent surveys have shown that the desert soils of Antarctica harbour surprisingly rich microbial communities1–3. Given that phototroph abundance varies across these Antarctic soils2,4, an enduring question is what supports life in those communities with low photosynthetic capacity3,5. Here we provide evidence that atmospheric trace gases are the primary energy sources of two Antarctic surface soil communities. We reconstructed 23 draft genomes from metagenomic reads, including genomes from the candidate bacterial phyla WPS-2 and AD3. The dominant community members encoded and expressed high-affinity hydrogenases, carbon monoxide dehydrogenases, and a RuBisCO lineage known to support chemosynthetic carbon fixation6,7. Soil microcosms aerobically scavenged atmospheric H2 and CO at rates sufficient to sustain their theoretical maintenance energy and mediated substantial levels of chemosynthetic but not photosynthetic CO2 fixation. We propose that atmospheric H2, CO2 and CO provide dependable sources of energy and carbon to support these communities, which suggests that atmospheric energy sources can provide an alternative basis for ecosystem function to solar or geological energy sources8,9. Although more extensive sampling is required to verify whether this process is widespread in terrestrial Antarctica and other oligotrophic habitats, our results provide new understanding of the minimal nutritional requirements for life and open the possibility that atmospheric gases support life on other planets.

Terrestrial Antarctica is among the most extreme environments on Earth. Its inhabitants experience the cumulative stresses of freezing temperatures, limited carbon, nitrogen and water availability, strong UV radiation, and frequent freeze–thaw cycles2,10,11. Although it was once believed that these conditions restrict life, we now know that the continent hosts a surprising diversity of macrofauna and microorganisms12,13. Surveys indicate that the phylum-level composition of microbial communities in Antarctic soils is similar to those of temperate soils3, but Antarctic communities are highly specialized at the species level and strongly structured by physicochemical factors1,3,10. In many Antarctic soils, microorganisms are thought to live in dormant states2, with metabolic energy directed towards cell maintenance rather than growth3. However, it is unclear how these communities obtain the energy and carbon needed for maintenance, given that these soils are often low in organic carbon and contain few classical primary producers2,3. Cyanobacteria and algae are known primary producers in such ecosystems2,4, but multiple sites have now been described at which phototrophs are present in low abundance or restricted to lithic niches5,12,14. These findings suggest that unidentified energy and carbon sources may also support ecosystem function. Here, we used shotgun metagenomics and biochemical measurements to clarify the basis of primary production in two such Antarctic sites. Three surface soils were initially sampled from Robinson Ridge (Wilkes Land)11 (Extended Data Fig. 1). Physicochemical analysis showed that the soils were low in total organic carbon (0.13 – 0.24%), nitrogen (0.012 – 0.023%), and moisture (4.0 – 5.6%) content (Supplementary Table 1). We used shotgun DNA sequencing to produce a 264-MB metagenome from these soils, comprising 208,233 predicted genes (Supplementary Table 2) and 451 predicted 16S/18S rRNA operational taxonomic units (OTUs) (Supplementary Table 3). The communities were dominated by Actinobacteria, Chloroflexi, Proteobacteria, Acidobacteria, and two candidate phyla, AD3 and WPS-2 (Fig. 1a). Identified phototrophs were limited to Cyanobacteria, with an average relative abundance of 0.28% (Fig. 1a). Consistent with other studies5,11,14, the low photosynthetic capacity and organic carbon content of Robinson Ridge soils contrasts with the inferred bacterial diversity, suggesting that alternative energy sources support these communities. To gain a deeper understanding of ecosystem function, we applied differential coverage binning25, which has been used to construct genomes of uncultivated inhabitants of diverse environments15–18. We constructed 23 draft genomes from the Robinson Ridge metagenome, including two WPS-2 and three AD3 genomes (Fig. 1b; Supplementary Table 4). These candidate phyla affiliate with the Terrabacteria superphylum19 and are sister lineages to Chloroflexi and Armatimonadetes19 (Extended Data Fig. 2).

We subsequently analysed the metabolic potential of the metagenome and derived genomes (Extended Data Fig. 3). All 23 sequenced organisms harboured terminal oxidase genes to sustain aerobic respiration, consistent with their aerated habitat, and genes for the oxidation of organic carbon compounds were also abundant (Supplementary Tables 5, 6). Genes supporting CO2 fixation through the Calvin–Benson–Bassham (CBB) cycle were unexpectedly widespread, being found in the genomes of Actinobacteria, WPS-2, and AD3 (Fig. 1b, Supplementary Table 5). Phylogenetic analysis revealed that these genomes encoded the type IE rubulose-1,5-bisphosphate carboxylase (RuBisCO) (rbcLIE) (Fig. 2b, Extended Data Fig. 4a, b); this lineage is known to support hydrognotrophic growth of Actinobacteria, but is absent from phototrophs. Instead, enzymes responsible for the aerobic respiration of molecular H220–22 and CO23,24 were encoded in multiple genomes (Fig. 1b, Supplementary Table 5). The high-affinity lineages identified—namely the structural genes encoding group 1h [NiFe]-hydrogenases (hylL and hylS) (Fig. 2a, Extended Data Fig. 5a) and type I [MoCu]-carbon monoxide dehydrogenases (coxL, coxS and coxM) (Extended Data Fig. 6a)—support scavenging of H2 and CO at the trace concentrations found in the atmosphere21,23. We also identified determinants of methanotrophy, ammonia oxidation, nitrogen cycling, and psychothermogenesis (Supplementary Tables 7, 8, 9).

Collectively, the metagenome data suggest that Antarctic surface soil microbial communities have the capacity to scavenge H2, CO2 and...
CO from the atmosphere for use as energy and carbon sources. We confirmed this theory by detecting both the expression and activity of the enzymes catalysing these processes in Robinson Ridge soils. Reverse transcription (RT)–PCR confirmed the expression of the genes encoding type IE RuBisCO (rbcL1E), high-affinity hydrogenase (hhyl), and carbon monoxide dehydrogenases (coxL) (Extended Data Fig. 7). Gas chromatography experiments showed that the soil communities aerobiocally oxidized atmospheric H2 and CO (threshold of 190 parts per billion by volume) H2 and 20 parts per billion by volume CO; rate of 3.49 nmol atmospheric H2 per hour per gram dry weight and 0.42 nmol atmospheric CO per hour per gram dry weight (Fig. 3a; Extended Data Fig. 6b; Supplementary Table 10), with H2 oxidation also measurable at −12 °C (Extended Data Fig. 5b). However, we did not observe atmospheric methane oxidation, despite recovering the genome of a putative alphaproteobacterial methanotroph. We next measured the capacity of the soils to fix carbon by tracing assimilation of 14C-labelled CO2. Although basal levels of CO2 fixation were variable (Extended Data Fig. 4c, d), the addition of H2 stimulated CO2 fixation by an average of twofold (Fig. 3b). By contrast, light stimulation had no consistent effect (Fig. 3b).

To ensure that the use of H2, CO2 and CO from the atmosphere as energy and carbon sources was not an isolated phenomenon, we sampled another Antarctic site: Adams Flat (Princess Elizabeth Land) (Extended Data Fig. 1). We collected three soils (Supplementary Table 1) with low organic carbon content and high Actinobacteria–low Cyanobacteria communities (Extended Data Fig. 8). Enzymes responsible for trace gas scavenging were expressed and active in these soils (Fig. 3a; Extended Data Figs 6b, 7; Supplementary Table 10), and H2 supplementation stimulated carbon fixation by an average of eightfold (P = 0.0039) (Fig. 3b). In further support of the theory that trace gas scavenging operates in a range of Antarctic desert sites, analysis of public metagenomes revealed that hhyl, coxlE, and rbcL1E are relatively abundant in the McMurdo Dry Valleys region (Extended Data Fig. 9).

On the basis of these findings, we propose that the two Antarctic sites sampled harbour largely dormant microbial communities that conserve energy by oxidation of atmospheric trace gases (Fig. 4). Biochemical studies showed that soils oxidize atmospheric H2 and CO at rates that are theoretically sufficient to sustain the energy needs of their microbial communities; assuming 1.4 × 1014 cells per C-mol of biomass (the amount of biomass that contains 1 mol carbon) and a maintenance energy of 1.68 kJ per C-mol biomass per hour at 10 °C, the rates of atmospheric H2 and CO oxidation observed can support 5.5 × 107 bacteria per gram of Robinson Ridge soil and 8.0 × 107 bacteria per gram of Adams Flat soil (Supplementary Table 10). Consistently, pure culture studies have shown that trace gas scavenging supports the persistence of diverse heterotrophic aerobes during organic carbon starvation20,22,25. Atmospheric H2 and CO are dependable energy sources
for dormant bacteria as they are strong reductants, diffuse through cell membranes, and occur at low concentrations throughout the troposphere\textsuperscript{24,27,28}.

Moreover, we propose that the primary producers in these communities are bacteria from the phyla Actinobacteria, AD3, and WPS-2 that generate biomass by consuming atmospheric $\text{H}_2$, $\text{CO}_2$ and $\text{CO}$. Of the draft genomes harbouring group 1h $[\text{NiFe}]$-hydrogenase genes, 55\% also contained type IE RuBisCO genes (Fig. 2a, b; Supplementary Table 5), suggesting that some bacteria generate biomass through $\text{H}_2$-driven CO fixation and others exclusively scavenge $\text{H}_2$ for energy acquisition. The metagenomes suggest that minor community members may also assimilate carbon through the serine pathway (methanotrophs), the 3-hydroxypropionate pathway (ammonia oxidizers), and oxygenic photosynthesis (cyanobacteria). On the basis of the high completeness, minimal contamination, and monophyly of their genomes (Supplementary Table 4), we propose the names \textit{Candidatus Dormibacteraeota} (desert bacterial phylum) and \textit{Candidatus Eremiobacteraeota} (dormant bacterial phylum) for candidate phyla WPS-2 and AD3, respectively (Supplementary Information). The type genera \textit{Candidatus Eremiobacteraeota} and \textit{Candidatus Dormibacteraeota} are also proposed, based on two of the obtained genome sequences (Ga011786, Ga0137693), as per recent recommendations\textsuperscript{29}.

Overall, the microbial community structure of the two Antarctic sites sampled appears to be shaped by selection for bacteria that can persist in these physically extreme, chemically deprived environments. Microbial phototrophs are in low abundance, probably restricted by climatic conditions such as low water and nutrient availability, summer radiation, and winter darkness\textsuperscript{30}. Instead, most inhabitants are inferred to be dormant heterotrophic aerobic bacteria that support energy generation and, in some cases, carbon fixation by oxidising atmospheric trace gases. In oligotrophic Antarctic soils, there is evidence that aeolian processes drive some ecosystem recruitment from less hostile microenvironments (for example, hypolithons) and niche processes subsequently select for the most functionally specialized microorganisms\textsuperscript{30}; hence, the ability to scavenge trace gases is likely to be an important survival-determining trait that may influence ecosystem succession. Given the concurrent findings from Robinson Ridge, Adams Flat, and the public McMurdo Dry Valley metagenomes, trace gas scavenging may be a general mechanism in Antarctic desert soils. However, broader surveys are now needed to determine the distribution, significance, and role of this process in primary production and ecosystem succession in terrestrial Antarctica, particularly in comparison to phototrophy. Indeed, there are diverse Antarctic sites where phototrophs are dominant, including hypolithons and high-latitude or moist soils\textsuperscript{4,4}. It is likely that both phototrophy and gas scavenging co-occur in many Antarctic sites, with the balance of these processes shifting depending on physicochemical factors. It will also be of interest to determine whether trace gas scavenging supports primary production in other oligotrophic ecosystems, for example the hyperarid deserts of Atacama, where Actinobacteria harbouring genes for trace gas scavenging are also dominant\textsuperscript{17}. Whereas most ecosystems are driven by solar or geologically derived energy, primary production in these Antarctic desert surface soils appears to be supported by atmospheric trace gases.

**Online Content** Methods, along with any additional Extended Data display items and Source Data, are available in the online version of the paper; references unique to these sections appear only in the online paper.

### Figure 3 Trace gas oxidation and chemosynthetic carbon fixation in Antarctic desert soils.
A. Gas chromatography measurements of oxidation of atmospheric $\text{H}_2$ (mixing ratio, 0.53 parts per million by volume (p.p.m.v.)) at $10\,^\circ\text{C}$. Values shown for two biologically independent soil samples (mean of technical triplicates). B. Ratios of $^{14}\text{C}$-labelled CO$_2$ assimilation following stimulation with light and/or $\text{H}_2$, Results shown for two biologically independent Robinson Ridge soils (technical quadruplicate) and three biologically independent Adams Flat soils (technical triplicate). Centre values show medians; boxes show upper and lower quartiles; whiskers show maximum and minimum values. Where sample size was appropriate, statistical significance between paired technical replicates was tested using a two-tailed Wilcoxon signed-rank test.

### Figure 4 Proposed metabolic capacity of the Robinson Ridge soil.
Depicted are members of the five microbial phyla that are predicted to persist by trace gas scavenging. These phyla are variably capable of aerobic respiration of atmospheric $\text{H}_2$ (all phyla) or $\text{CO}$ (Actinobacteria, Dormibacteraeota), and chemosynthetic fixation of atmospheric $\text{CO}_2$ (Actinobacteria, Dormibacteraeota, Eremiobacteraeota). It remains to be determined whether atmospheric $\text{H}_2$ oxidation supports NAD(P)$^+$ reduction through direct coupling or reverse electron transport, although direct coupling remains thermodynamically favourable at $10\,^\circ\text{C}$ (Gibbs free energy of reaction ($\Delta G$) = $-19\,\text{kJ}\,\text{mol}^{-1}$). The community is also predicted to respire the small amount of soil organic carbon present as it becomes bioavailable.
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Supplementary Information is available in the online version of the paper.

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METHODS

No statistical methods were used to predetermine sample size. The experiments were not randomized and the investigators were not blinded to allocation during experiments and outcome assessment.

**Site description and sample collection.** This study focused on two coastal ice-free sites in different regions of eastern Antarctica. Robinson Ridge (∼66.367739, 110.585262), located 10 km south of Casey station in the Windmill Islands coast of Wilkes Land (Extended Data Fig. 1a, b), is part of a pristine polar desert (Extended Data Fig. 1c). Adams Flat (∼68.5502, 78.02106) is located 242 km from Davis station in the Vestfold Hills regions of Princess Elizabeth Land (Extended Data Fig. 1d). Both sites are devoid of vascular plants, but harbour a limited diversity of macroflora, with tardigrades and nematodes shown to be present in Robinson Ridge4. Three surface soil samples (100 g) from the top 10 cm of the soil profile were collected in December 2005 for a previous study on Robinson Ridge1 and January 2014 for Adams Flat. For both sites, samples were collected along a spatially explicit sampling design comprised of three 300-m-long transects11,33 separated by 2-m distances from each other. As these soils were included in the Australian Antarctic Division’s polar soil archive, samples were sieved down to 63 μm, aliquoted into 5–25-g subsamples, and stored at −80 °C until analysis.

**Physicochemical analysis.** Latitude, longitude, slope, aspect and elevation were recorded for each sample taken using ARC GIS software (ESRI). For all soils, detailed physical and chemical data were analysed in-house using standard procedures. Total carbon, nitrogen, and phosphorus, pH, water-holding capacity, grain size, and conductivity were recorded1. In addition, extractable ions were measured (Cl−, Br−, NO3 −, NO2 −, PO4 3−, SO4 2−, and NH4 +) and X-ray fluorescence elemental analysis was conducted (SiO2, TiO2, Al2O3, Fe2O3, MnO, MgO, CaO, Na2O, K2O, P2O5, SO3, and Cl). The soils were low in total organic carbon content (average 0.17% for Robinson Ridge, 0.09% for Adams Flat) and moisture measured as dry matter fraction (average 4.4% for Robinson Ridge, 0.42% for Adams Flat) (Supplementary Table 1).

**Community DNA extraction.** Total community DNA was extracted from the six soil samples for microbial community profiling and functional metagenomic analysis. In all cases, DNA was extracted from 0.25–0.3 g of each sample in technical triplicate using the FastDNA SPIN Kit for Soil (MP Biomedicals). All DNA extracts were quantified and DNA lystate quality was evaluated using automated ribosomal intergenic spacer analysis (ARISA) as described21.

**Metagenome sequencing, assembly, and binning.** For the three Robinson Ridge samples, DNA was extracted in triplicate and used for shotgun metagenome sequencing. Metagenome libraries were prepared using the Nextera DNA Library Preparation Kit (Illumina) and sequenced using three-fifths of an Illumina HiSeq2000 flowcell lane at the Institute for Molecular Biosciences (University of Queensland). The raw reads (2 × 100-bp reads, 14.9 Gb) were processed using Trimmomatic33 for adaptor removal and quality filtering, and BBMap to merge the paired-end reads. FastQ files were assembled and annotated using the assembly algorithm in CLC Genomics Workbench v8 (CLC Bio). The genome sequences were then quality controlled and in silico predicted using RDP 14.7. All contigs were binned on the basis of differential coverage profiles, k-mer frequencies, and GC content using GroupM15 and MetaBAT9. Population genome bins obtained with GroupM were further refined using the GroupM refine function. The genome completeness and contamination were estimated with CheckM28 by calculating the presence of lineage-specific single-copy marker genes. The two sets of population genomes from the GroupM and MetaBAT binning were compared using RefineM to identify possible duplicates. Twenty-three genomes (>30% completeness, <10% contamination) were selected for further analysis and accounted for 56–65% of total reads obtained from the three soil samples. Of these, 11 genomes were 90% complete and less than 5% contaminated (Supplementary Table 4).

**Community analysis.** We determined the microbial community structure of the three soil samples from Robinson Ridge and three from Adams Flat. For both sites, the bacterial and archaeal community structure was determined by 16S rRNA gene amplicon sequencing as described previously21,28. Community alpha diversity was inferred from the amplicon sequencing data by calculating observed richness, Chao1, and the Shannon index (H′) as described21. For the Robinson Ridge samples only, bacterial, archaeal, and eukaryote community structure was also determined by retrieving 16S and 18S rRNA genes from the metagenomes. Community composition profiles were generated with Community by mapping reads onto a database of bacterial, archaeal and eukaryotic 16S and 18S rRNA genes (that is, the SILVA29 and GreenGenes48 databases clustered at 97% similarity). To infer the phylogeny of WPS-2 and AD3, a genome tree was generated using 38 universal conserved marker genes41 from 4,624 bacterial and archaeal genomes retrieved from the Integrated Microbial Genomes database (IMG)42 together with the recovered population genomes. A concatenated alignment of the marker genes was used to generate the genome tree with FastTree43 and the tree was visualized in iTOL44.

**Functional analysis.** The population genomes derived from Robinson Ridge were functionally annotated using Prokka46 and the KEGG Orthology database (Kyoto Encyclopedia of Genes and Genomes)47. Genes specifically involved in carbo- hydrate metabolism were identified using dbCAN, an HHM-based database for carbohydrate-active enzyme (CAZY)48 annotation. Potential secondary metabolic biosynthesis clusters were identified using antiSMASH48. [NiFe]-hydrogenase, [MoCo]-carbon monoxide dehydrogenase, and RuBiSCO enzymes encoded within the metagenomes were classified by constructing phylogenetic trees of their catalytic subunits14,49,50. The derived protein sequences encoding the catalytic subunits of these enzymes were aligned with reference sequences reported in previous studies21,49,50, using ClustalX49, Evolutionary relationships were visualized on phylogenetic trees constructed with MEGA70 using the maximum-likelihood method. Trees were bootstrapped using 500 replicates and rooted with suitable outgroup sequences. The relative abundance of hzyL, coxL, and rbclE in McMurdo Dry Valley samples was determined by retrieving ten public metagenomes through the Joint Genome Institute (JGI IDs 104803, 106469, 35851, 3300002548, 3300012042, 3300012045, 3300012185, 3300012188, 3300012257, and 3300012678). The genes of interest were retrieved from the downloaded metagenomes by BLAST using the hzyL, coxL, and rbclE gene sequences identified from the Robinson Ridge metagenome as queries. The identity of the retrieved type I coxl and type II ecbL large subunit genes was confirmed by constructing phylogenetic trees as described above, whereas genes encoding group I h [NiFe]-hydrogenase large subunits were identified using HyDB50. The relative abundance of the three genes was compared with those in five public forest metagenomes (JGI IDs 66726, 69782, 92543, and RuBiSCO) for sequences sequenced using Illumina MiSeq PE50 technology. The MGG4 genome was generated using RT-PCR. For expression analysis, total RNA was extracted from 2 g from each of the six soil samples using the MoBio PowerSoil Total RNA Isolation Kit (MO BIO). All RNA extracts were quantified using the RNA Analysis Kit (Agilent) and cDNA was synthesized using Maxima First Strand cDNA Synthesis Kit (Thermo Fisher Scientific). RT–PCR was used to confirm the expression of the group 1 h [NiFe]-hydrogenase large subunit gene hzyL (primers NiFe-2444: 5′-GGGATCTGGGCGGACAAA-3′; NiFe-568r: 5′-TCTCCGGCGGTGGACGCCTC-3′) and the type I [MoCo]-carbon monoxide dehydrogenase large subunit gene coxl (primers type-1-1288F: 5′-TSSKYYACSGCGWSSSTTA-3′; type-1-1540r: 5′-TAYGAYWWWGGYRAYTA-3′) using previously described degenerate primers, mixtures, and thermal cycler conditions21,54. Expression of the type I RuBiSCO large subunit gene (rbclE) was confirmed using specifically designed primers based on the sequences obtained from Robinson Ridge (rbclE1F: 5′-GGCABGTSTGTYYTGACGASCA-3′; rbclE1R: 5′-TGGACAACBCCRAVARCTTGGC-3′). For this gene, the RT–PCR mixtures were comprised of 1× Promega reaction buffer (Promega), 0.4 mM deoxynucleotide triphosphates, 0.4 mM of each primer, 0.1 mg ml−1 bovine serum albumin, 1.25 U Go-Taq polymerase (Promega), 1 μl diluted cDNA, and nuclease-free water to a final volume of 25 μl. A standard PCR protocol was used for the type I IE RuBiSCO large subunit gene as follows: 95 °C for 5 min; 30 cycles of denaturing at 95 °C for 30 s, annealing at 55 °C for 30 s and extension at 72 °C for 30 s; and a final extension of 72 °C for 5 min. Owing to the high degeneracy of coxl PCR primers, multiple bands were produced after thermal cycling. To confirm the identity of the PCR products, bands were separated by gel electrophoresis, the band of the correct size (780 bp) was excised aseptically, and its DNA was extracted using the Zymoclean Gel DNA Recovery Kit (Zymol). For hydrogenase and RuBiSCO, purified PCR products were cloned and transformed using the pGEM-T Easy Vector System (Promega). The purified coxl PCR products and positive hydrogenase and RuBiSCO clones were sequenced using Sanger sequencing by the Macrogen Inc. (Seoul, South Korea). The identity of all sequenced clones was confirmed using NCBI BLAST against the nucleotide database.

**Gas chromatography analysis.** We used gas chromatography to determine whether the microbial communities in the collected soil samples scavenged atmospheric H2, CO, or CH4. Soil samples of 1 g were placed into sterile 114–ml serum bottles and sealed with butyl rubber stoppers. H2, CO, or CH4 gas was then added to achieve final initial headspace concentrations of ~60 p.p.m.v., ~5 p.p.m.v., or ~30 p.p.m.v., respectively. Soil incubation temperatures were monitored throughout using a 511 single input digital thermometer (Fluke). Headspace (1 ml) were sampled in triplicate using a gas sampler (VICI Precision Sampling). H2 and CO partial pressures were measured using a PPI Gas Analyser (Peak Performer) equipped with a reducing compound photometer (RCP: H2, CO), flame ionizing detector (FID: CH4), Unibeads 15 60/80 column, and Molecular Sieve 13X 60/80 column, as previously described20. Samples were calibrated against H2, CO, and CH4 standards
prepared in advance. The experiments each used two biologically independent Robinson Ridge and two biologically independent Adams Flat soil samples. Gas chromatography traces were recorded of technical triplicates for all H₂ measurements and technical duplicates for all CO measurements. Two negative controls, namely heat-killed soils (1 g; 121°C, 15 psi, 20 min) and serum bottles without soil, were monitored in parallel to regular sampling to ensure that the observed reduction of H₂ and CO concentrations was a consequence of biological oxidation. The theoretical bacterial population sustained by trace gas scavenging (N) was calculated by the methods of Conrad²⁸ based on the observed trace gas oxidation rate (d), the Gibbs free energy of gas oxidation (ΔG), and the theoretical maintenance energy of the population (m). ΔG was calculated to be −200.9 kJ mol⁻¹ for the reaction H₂ + 0.5 O₂ → H₂O and −236.2 kJ mol⁻¹ for the reaction CO + 0.5 O₂ → CO₂ using the Nernst equation (10°C, ambient gas concentrations). m was estimated to be 1.68 kJ per C-mol biomass per hour using the Tijhuis equation ⁵⁵.

Carbon fixation analysis. The soil samples were incubated with radiolabelled carbon dioxide (¹⁴CO₂) under different conditions to determine whether their microbial communities could mediate chemosynthetic or photosynthetic CO₂ fixation. Each of the soil samples (0.25 g) were added to sterile 4-ml glass vials, sealed with rubber septum lids, and stored on ice before down-stream analysis. Gaseous ¹⁴CO₂ (1% v/v) was generated by mixing 75 μl radiolabelled sodium bicarbonate solution (NaH¹⁴CO₃, Perkin Elmer, 53.1 mCi nmol⁻¹) with 75 μl 10% HCl solution in a sealed 4-ml glass vial and incubating for 2 h at room temperature. To each sample, 160 μl of ¹⁴CO₂ (1% v/v) gas was added using a gas-tight syringe (1 ml, SGE Analytical Science), obtaining initial headspace mixing ratios of 100 p.p.m.v. H₂ and 400 p.p.m.v. ¹⁴CO₂. Both syringes (1 ml, SGE Analytical Science) were run with soils that were heat-killed for 12 h at 100°C using a cooled vacuum drying oven (Memmert GmbH). Scintillation counts from heat-killed controls were subtracted and the amount of ¹⁴CO₂ fixed per sample was calculated on the basis of the reported specific radioactivity of the original bicarbonate solution. A control experiment validated that light stimulation caused a 15-fold increase in CO₂ fixation over the dark-incubated samples in a phototroph-containing soil community collected from Mitchell Peninsula in December 2005.

Data availability. The raw shotgun sequencing datasets generated during the current study have been deposited in the National Center for Biotechnology Information (NCBI) Sequence Read Archive (RR samples 1, 2 and 3 under accession numbers SRR5223441, SRR5223442 and SRR5223443, respectively). The metagene data have been deposited into the IMG-M web portal (https://img.jgi.doe.gov/cgi-bin/main/m/main.cgi) under accession number 3300009490, and the scaffolds of all 23 draft genomes have been deposited into the IMG-M web portal under accession numbers 2667527203, 2667527204 and 2698536762–2698536743.

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Extended Data Figure 1 | Field sites sampled in this study. a, Map of Antarctica highlighting Casey station in the Windmill Islands and Davis station in the Vestfold Hills, eastern Antarctica. b, The location of Robinson Ridge with respect to Casey station. c, Photo of sampling site at Robinson Ridge within the Windmill Islands region. d, Photo of sampling site in Adams Flat within the Vestfold Hills region. Map courtesy of D. Wilkins; Adams Flat photo courtesy of T. Mooney.
Extended Data Figure 2 | A concatenated genome tree showing phylogeny of the sequenced genomes retrieved from the Robinson Ridge metagenomes. The phyla retrieved are coloured green. The phylogeny of the three retrieved genomes of candidate phylum Dormibacteraeota (AD3) and the two retrieved genomes of candidate phylum Eremiobacteraeota (WPS-2) is shown (coloured pink and blue, respectively). Also shown is their affiliation with the superphylum Terrabacteria (highlighted in yellow).
Extended Data Figure 3 | Distribution of genes in the Robinson Ridge metagenome according to the KEGG classification system. The most abundant metabolic categories identified were those associated with carbohydrate (18%), amino acid (17%), and energy metabolism (8%).
Extended Data Figure 4 | See next page for caption.
Extended Data Figure 4 | Genetic organization, amino acid phylogeny, and inferred activity of RuBisCO enzymes in sampled soils.

a, Organization of type IE RuBisCO genes retrieved from Robinson Ridge. The genes included in Bin 12: Dormibacterota (AD3, top) and Bin 22: Eremiobacterota (WPS-2, bottom) are shown. *rbcS*, RuBisCO small subunit; *rbcL*, RuBisCO large subunit; *cbbX*, RuBisCO expression protein; *cbbP*, phosphoribulokinase; *cbbR*, transcriptional regulator protein; *cbbT*, transketolase.

b, Expanded phylogenetic tree of RuBisCO large subunit genes retrieved from Robinson Ridge metagenomes. The phylogenetic tree was generated by the maximum-likelihood method and robustness was tested with 500 replicates. The majority of sequences within the metagenome and bins encode type IE RuBisCO (chemosynthesis-type), except for one of the unbinned sequences, which encodes a type IA RuBisCO (photosynthesis-type).

c, Box plots showing amount of 14C-labelled CO2 assimilated by Robinson Ridge and Adams Flat soil communities following light or H2 stimulation. Results are shown for two biologically independent Robinson Ridge soils (technical quadruplicate) and three biologically independent Adams Flat soils (technical triplicate). Centre values show medians. Maxima, upper and lower quartile, and minima values are also shown. Where sample size is appropriate, statistical significance between paired technical replicates was tested using a two-tailed Wilcoxon signed-rank test.

d, Amount of 14C-labelled CO2 assimilated for each individual Robinson Ridge and Adams Flat soil tested. Soils are named as in Supplementary Table 1 and results are shown for technical quadruplicates (Robinson Ridge) and technical triplicates (Adams Flat). Centre values show means, upper values show maxima, and lower values show minima. Although a variable level of basal CO2 fixation was observed, these results show that H2 addition consistently stimulated CO2 fixation.
Extended Data Figure 5 | Amino acid phylogeny and inferred activity of hydrogenase enzymes in sampled soils. a, Expanded phylogenetic tree of [NiFe]-hydrogenase large subunit genes retrieved from the Robinson Ridge metagenome data. The phylogenetic tree was generated by the maximum-likelihood method and robustness was tested with 500 replicates. Hydrogenase genes were common within the metagenome and bins. They clustered exclusively with the group 1h [NiFe]-hydrogenases, suggesting that they mediate atmospheric H₂ scavenging. b, H₂ oxidation activities of Robinson Ridge soils incubated at 10 °C, 4 °C, and −12 °C. Gas chromatography traces show H₂ oxidation of soils in modified ambient air headspaces. Results are shown from one soil sample (Soil 1) following averaging of results from technical triplicates.

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Extended Data Figure 6 | Amino acid phylogeny and inferred activity of carbon monoxide dehydrogenase enzymes in sampled soils.

a, Phylogenetic tree of type I [MoCu]-hydrogenase large subunit genes retrieved from the Robinson Ridge metagenome data. The phylogenetic tree was generated by the maximum-likelihood method and robustness was tested with 500 replicates. Multiple Actinobacteria and AD3 genomes encoded this subunit, suggesting that they may mediate atmospheric CO scavenging.

b, Gas chromatography results showing oxidation of atmospheric CO (mixing ratio 0.10 p.p.m.v.) mediated by the soil communities at 10 °C in modified ambient air headspaces. For both the Robinson Ridge and Adams Flat datasets, values are shown for two biologically independent soils following averaging of technical duplicates.
Presence and expression of trace gas scavenging genes in Robinson Ridge and Adams Flat soil samples.

The presence and expression of the genes encoding the group 1h [NiFe]-hydrogenase large subunit (hhyL) and type IE RuBisCO large subunit (rbcL1E) was confirmed by agarose gel electrophoresis. Lane M shows the DNA ladder (Bioline 100-bp Molecular Weight Marker). Lanes 1 and 5 show amplifications from community DNA extracted from the soil samples. Lanes 2 and 6 show amplifications of cDNA derived from reverse transcription of community RNA extracted from the soil samples. No template negative controls for both the PCR (lanes 3, 7) and RT–PCR (lanes 4, 8) are also shown. Independent PCR and RT–PCR reactions were conducted twice for both Robinson Ridge and Adams Flat soil samples, with similar results. For gel source data, see Supplementary Fig. 1. For carbon monoxide dehydrogenase, the bands obtained from agarose gel electrophoresis were blurred owing to the degeneracy of the primer set. Sequencing of the bands confirmed the expression of carbon monoxide dehydrogenase genes homologous to GenBank accession numbers EU888210 (79%), EU888266 (79%), and KJ468250 (88%).
Extended Data Figure 8 | Microbial community structure of Adams Flat soil samples. Community structure was determined by targeted amplicon sequencing of bacterial 16S rRNA genes from three biologically independent soil samples. All samples show a high abundance of potential trace gas scavengers from the phylum Actinobacteria and a low abundance of phototrophs from the phylum Cyanobacteria.
Extended Data Figure 9 | Comparison of the relative abundance of genes associated with trace gas scavenging in public metagenomes. The relative abundance of the genes *hhyL*, *coxL*, and *rbcL1E* was compared between McMurdo Dry Valley and forest metagenomes publicly available through JGI. Results are shown for ten biologically independent McMurdo Dry Valley metagenomes and five biologically independent forest metagenomes. Centre values show medians; boxes show upper and lower quartiles; whiskers show maxima and minima. Statistical significance was tested using a one-tailed Student's *t*-test.
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▶ Experimental design

1. Sample size
   Describe how sample size was determined.
   No statistical methods were used to predetermine sample sizes. Three biologically independent soil samples were collected from the two sampling sites, Robinson Ridge (December 2005) and Adams Flat (January 2014). For both sites, samples were collected along a spatially explicit sampling design comprised of three 300 m long transects separated by two meter distances from each other. This sample size was sufficient to perform the in-depth metagenome analyses and supporting biochemical assays needed to confirm that novel primary production processes operating in these soils. However, the relatively limited sample size meant that we were unable to confirm whether trace gas scavenging is a widespread mechanism operating in Antarctic soils. Due to the very limited amounts of Robinson Ridge soils that remained following metagenome and physicochemical analysis, we used sample sizes lower than three for some gas chromatography and carbon fixation experiments.

2. Data exclusions
   Describe any data exclusions.
   No exclusions.

3. Replication
   Describe whether the experimental findings were reliably reproduced.
   All attempts at replication were successful. For carbon fixation studies (Figure 3b, Extended Data Figure 4), we observed that there was a large and unexplained variation in basal levels of CO2 that was fixed between each of the five soils tested. However, the core finding that H2 addition stimulated carbon fixation was reliably reproduced. H2 addition stimulated all 17 sample pairs in the dark condition and 15 of the 17 sample pairs under light illumination.

4. Randomization
   Describe how samples/organisms/participants were allocated into experimental groups.
   Samples were not randomized for the experiments.

5. Blinding
   Describe whether the investigators were blinded to group allocation during data collection and/or analysis.
   Blinding was not applicable to use in this study as we were investigating a new process and only two sites were examined.

Note: all studies involving animals and/or human research participants must disclose whether blinding and randomization were used.
6. Statistical parameters

For all figures and tables that use statistical methods, confirm that the following items are present in relevant figure legends (or in the Methods section if additional space is needed).

| n/a | Confirmed |
|-----|-----------|
| ☑   | ☑         |

- The exact sample size (n) for each experimental group/condition, given as a discrete number and unit of measurement (animals, litters, cultures, etc.).
- A description of how samples were collected, noting whether measurements were taken from distinct samples or whether the same sample was measured repeatedly.
- A statement indicating how many times each experiment was replicated.
- The statistical test(s) used and whether they are one- or two-sided (note: only common tests should be described solely by name; more complex techniques should be described in the Methods section).
- A description of any assumptions or corrections, such as an adjustment for multiple comparisons.
- The test results (e.g. P values) given as exact values whenever possible and with confidence intervals noted.
- A clear description of statistics including central tendency (e.g. median, mean) and variation (e.g. standard deviation, interquartile range).
- Clearly defined error bars.

See the web collection on statistics for biologists for further resources and guidance.

7. Software

Describe the software used to analyze the data in this study.

- ARC GIS 10.1, Trimmomatic 0.36, CLC Genomics Workbench v8, BamM 1.4.1, GroopM 0.3, MetaBAT 0.25.2, CheckM v0.9.4, RefineM 0.0.20, CommunityM 1.0.6, FastTree 2.1, iTOL v3.6, dbCAN 5.0, ClustalX 2.0, MEGA7, Antismash 2.0, Graphpad Prism 7, BBMap 35.66, Prokka 1.12.

For manuscripts utilizing custom algorithms or software that are central to the paper but not yet described in the published literature, software must be made available to editors and reviewers upon request. We strongly encourage code deposition in a community repository (e.g. GitHub). Nature Methods guidance for providing algorithms and software for publication provides further information on this topic.

8. Materials and reagents

Policy information about availability of materials

8. Materials availability

Indicate whether there are restrictions on availability of unique materials or if these materials are only available for distribution by a for-profit company.

All materials are available. However, only limited quantities remain of the Antarctic soil samples used for this study.

9. Antibodies

Describe the antibodies used and how they were validated for use in the system under study (i.e. assay and species).

Not applicable.

10. Eukaryotic cell lines

a. State the source of each eukaryotic cell line used.

Not applicable.

b. Describe the method of cell line authentication used.

Not applicable.

c. Report whether the cell lines were tested for mycoplasma contamination.

Not applicable.

d. If any of the cell lines used are listed in the database of commonly misidentified cell lines maintained by ICLAC, provide a scientific rationale for their use.

Not applicable.
Animals and human research participants

Policy information about studies involving animals; when reporting animal research, follow the ARRIVE guidelines

11. Description of research animals

Provide details on animals and/or animal-derived materials used in the study.

Not applicable.

Policy information about studies involving human research participants

12. Description of human research participants

Describe the covariate-relevant population characteristics of the human research participants.

Not applicable.