Grazing weakens competitive interactions between active methanotrophs and nitrifiers modulating greenhouse-gas emissions in grassland soils

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Grassland soils serve as a biological sink and source of the potent greenhouse gases (GHG) methane (CH4) and nitrous oxide (N2O). The underlying mechanisms responsible for those GHG emissions, specifically, the relationships between methane- and ammonia-oxidizing microorganisms in grazed grassland soils are still poorly understood. Here, we characterized the effects of grazing on in situ GHG emissions and elucidated the putative interactions between the active microbes involving in methane oxidation and nitrification activity in grassland soils. Grazing significantly decreases CH4 uptake while it increases N2O emissions basing on 14-month in situ measurement. DNA-based stable isotope probing (SIP) incubation experiment shows that grazing decreases both methane oxidation and nitrification processes and decreases the diversity of active methanotrophs and nitrifiers, and subsequently weakens the putative competition between active methanotrophs and nitrifiers in grassland soils. These results constitute a major advance in our understanding of putative relationships between methane- and ammonia-oxidizing microorganisms and subsequent effects on nitrification and methane oxidation, which contribute to a better prediction and modeling of future balance of GHG emissions and active microbial communities in grazed grassland ecosystems.

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INTRODUCTION
Methane (CH4), as the second most potent greenhouse gas (GHG) after carbon dioxide (CO2), is considered to be responsible for ~20% of the anthropogenic global warming effect [1]. Its atmospheric concentration has doubled from about 700 ppb at pre-industrial times to the current concentration of 1850.5 ppb [2]. Aerobic oxidation of CH4 in soils by methane-oxidizing bacteria (MOB), also known as methanotrophs, represents the largest biological sink for atmospheric CH4 [3]. Methanotrophs have the unique ability to grow on CH4 as their sole source of carbon and energy. They are ubiquitous in the environment and play a major role in the removal of the greenhouse gas methane from the atmosphere if released into the atmosphere [4].

The key step of aerobic methane oxidation, the initial oxidation of CH4 to methanol, is catalyzed by the methane monooxygenase which exists either as a particulate, membrane-bound form (pMMO), or a soluble, cytosolic form [5]. It is noticeable that pMMO of MOB and ammonia monooxygenase (AMO) of ammonia-oxidizing bacteria (AOB) are homologous members [6], which are grouped into the copper-containing membrane-bound monooxygenase (CuMMO) family [7]. The AMO of ammonia-oxidizing archaea (AOA) is also a CuMMO, but it is a phylogenetically distant one from those of MOB and AOB [8]. Traditionally, ammonia (NH3) was converted into nitrite by AOA/AOB and further oxidized into nitrate by nitrite-oxidizing bacteria (NOB). The evolutionary links between MOB and AOB, together with the similar molecular structure of their substrates (NH3 and CH4, respectively), lead to functional similarities enabling them to oxidize both NH3 and CH4, although neither AOB nor MOB are capable of growing on the alternative substrate [9–11].

Methanotrophs and ammonia oxidizers are considered as strong competitors for N to maintain their growth and activity [12, 13]. However, response and activity of methanotrophs to N levels remains unsolved and is still debated. Previous studies either demonstrated that ammonium could stimulate [12, 14–17], suppress [18, 19], or exert little impact on [20, 21] the global methanotrophic CH4 sink. Therefore, there are some gaps of knowledge concerning the mechanisms controlling the relationships between methane oxidation and N levels. Additionally, N levels are particularly important in controlling niche specialization of the ecology and evolution of AOA and AOB [14, 22]. AOA generally dominates ammonia oxidation in N-limited soils while...
AOB dominates ammonia oxidation in N-rich environments [12, 14, 22]. It is thus hypothesized that ammonium-assimilating MOB decreases nitrification rate by competing for N with AOA and AOB, especially in N-limited soils.

Grasslands are known as important biological sinks of CH4 and sources of N2O and grassland soils are often viewed as important ecosystems influencing global environmental change through their strong capacity to produce, store, and cycle C and N substrates [23, 24]. It is reported that grassland ecosystems contain more than one-third of above- and below-ground C reserves because grasslands cover nearly half of the Earth’s land surface [25]. Notably, management practices of grasslands can alter the exchange of C and N between atmosphere and soil, aboveground and belowground biomass. Livestock grazing, for example, is the most important and pervasive practice in grasslands. Increasing stocking rates have been implemented to meet the increasing food demands of a growing population, but grazing modifies C and N cycling in grassland soils [26]. Grazing has also been shown to be an important factor regulating the emission and uptake of greenhouse gases [27] with moderately-grazed grassland having high CH4 uptake, while heavy grazing suppresses soil CH4 uptake and stimulates N2O emissions [28, 29].

Analysis of the active methanotrophs in such ecosystems has shown that grazing decreases the abundance and diversity of the active methanotrophs in steppe soils [30]. Grazing also significantly decreases nitrification activity, and different grazing intensity leads to diverse active communities with AOA and AOB focused on effects of grazing on nitri-fication or methane oxidation, but very little is known about their relationships in grassland ecosystem as well as the responsible microbial communities.

In this study, we firstly investigated in situ greenhouse gas emissions in ungrazed and grazed grasslands for 14-months (August 2014 to October 2015) using closed static chambers. Then a controlled incubation experiment was conducted in the laboratory to understand microbial relationships modulating greenhouse gas emissions in grassland by performing DNA–SIP and Illumina HiSeq 16S ribosomal RNA (rRNA) gene sequencing. The present study aimed to (1) characterize the impact of long-term grazing on active methanotrophs and nitrifiers, (2) examine the mechanisms underlying the effects of N level on CH4 and N2O oxidation, and (3) clarify the relationships occurring between important players of the methane and ammonia oxidation in grazed grassland soils. We hypothesized that grazing would suppress the competitions between methane- and ammonia-oxidizing microorganisms by decreasing their diversities and activities. Elucidating those putative relationships is crucial for understanding the methane-nitrogen cycling and its effects on global climate change.

MATERIALS AND METHODS

Site description and soil sampling

The experimental site was situated at the Inner Mongolia Grassland Ecosystem Research Station (IMGERS, 43°37’S, 116°43’E) on the Xilingol steppe of the Xilin River basin, bounded by the west side of the Daxing-An Mountain. This ecoregion possesses a semi-arid continental climate with mean annual temperature of −0.4 °C, and 348 mm of precipitation which was distributed unevenly across the seasons, falling mostly during June–August. The vegetation type was Leymus chinensis with some Stipa grandis and Telespiza aquarescas. The soil was classified as Calcic Chernozem according to ISSS Working Group R8, 1998.

Two adjacent long-term field plots were established at the Research Station, one ungrazed and enclosed since 1983 (“Ungrazed”) and the other subjected to free grazing (“Grazed”). Both sites were derived from a paddock that had uniform fertility and slope. At each site, a transect of 60 m × 400 m was established with five equal-sized replicate plots of dimension 60 m × 70 m. A 10-m-wide buffer strip was established between each plot to avoid interactions and allow for sampling. Soil samples were collected from the upper 10-cm layer from five random locations within each plot following an “S” sampling pattern using a 5-cm diameter soil auger in August 2016. The five samples from each plot were grouped into a single composite sample, packed with ice packs, and transported to the laboratory. Soil samples were then passed through a 2-mm sieve. Subsamples were air-dried for physicochemical analysis. The remaining fresh soils were used for the incubation experiment.

Soil pH was measured using a pH meter (Mettler-Toledo, Switzerland) after shaking the soil at a soil-water ratio of 1:2.5. Gravimetric soil moisture content was analyzed by oven-drying at 105 ± 2 °C for 24 h. Soil bulk density was measured using the volumetric method according to Lampurlanes and Cantero-Martinez [32]. Soil organic matter was determined by dichromate digestion [33]. Total C and N were determined by dry combustion in a Vario Max CNS analyzer (Elementar Instrument, Mt. Laurel, NJ). Olsen P was extracted by 0.5 M NaHCO3 (pH = 8.5) for 0.5 h and determined with the molybdenum blue method [34]. Available potassium was extracted by 1 M ammonium acetate and determined by flame emission spectrophotometry. Exchangeable NH4+ and NO3– were extracted with 1 M KCl for 1 h and determined by a flow injection analyzer (SAN+−+, Skalar, Holland).

In situ CH4 and N2O measurements

To elucidate the effects of grazing on N2O and CH4 emissions from grassland soils, we collected gas samples using closed-chamber greenhouse-gas flux method from August 2014 to October 2015. For gas sampling, a static chamber (50 cm diameter) was installed in each of the five replicated plots into the ungrazed and grazed soils (in close proximity to the soil sampling points) resulting in ten chambers. Gas samples (40 ml) were collected five times during the in-situ field study, in May (early pasture-growing season, 2015 only), August (peak pasture-growing season), and October (non-growing season) in both 2014 and 2015. Gas samples were collected 0 and 40 min after chamber closure using a 20-ml syringe and injected into preevacuated 20-ml glass bottles and samples were collected between 10 and 12 a.m. The concentrations of N2O and CH4 in gas samples were analyzed using a gas chromatograph (Shimadzu GC-2010 Plus, Japan).

Construction and sampling of soil microcosms

Fresh soil (equivalent to 6.0 g dry weight) was pre-incubated at 40% field moisture capacity in a 120-ml serum bottle for ten days at 25 °C in darkness before the incubation experiment. Microcosms were established using seven treatments in triplicate, see details in Table S1 including (i) 13C–CH4 without urea addition (U0 + 13C14CH4); (ii) Low 13C-urea with 13CO2–CO2, corresponding to 20 mg 13C-urea-N kg−1 soil; (iii) High 13C-urea with 13CO2 (U100 + 13CO2, corresponding to 100 mg 13C-urea-N kg−1 soil); (iv) Low 13C-urea with 12CO2–CO2 and 13C–CH4 (U20 + 12CO2–13CH4); (v) High 13C-urea with 12CO2–CO2 and 13C–CH4 (U100 + 12CO2–13CH4); (vi) Low 13C-urea with 12CO2–CO2, corresponding to 100 mg 13C-urea-N kg−1 soil; (vii) Low 13C-urea with 12CO2–CO2 and 13C–CH4 (U20 + 12CO2–13CH4); (viii) High 13C-urea with 12CO2–CO2 and 13C–CH4 (U100 + 12CO2–13CH4).

Microcosms supplemented with 13C–CH4 or 13C–urea microcosms were used to monitor the active methanotrophs and nitrifiers, respectively, while the microcosms supplemented with 13C–urea and 13C–CH4 were used to monitor the putative relationships between active methanotrophs and nitrifiers. The application rate of U20 was to mimic the annual total N deposition in the grassland ecosystems. The annual total inorganic N deposition to Inner Mongolia grassland regions ranged from 4 to 20 kg N ha−1 year−1 [35], which is equivalent to 3–15 mg N kg−1 dry soil, assuming an effective soil depth of 15 cm. The treatment of U100 was implemented to simulate N input accompanied with livestock excrements, as up to 300 kg N ha−1 y−1 can be returned to grassland soils in the form of livestock urine [36]. Urea was added weekly by dropple additive of freshly made urea solution, as degradation of urea into NH4+–N would occur very quickly in soils, to achieve 60% field capacity to establish a substrate-rich environment for nitrifying communities. For the U0 + 13CH4 treatment, distilled water was added instead of urea solution to achieve 60% field capacity. Microcosms were flushed before any supplementation of urea or methane with synthetic air (20% O2, 80% N2) for 1 min to maintainoxic conditions. The bottles were then sealed with rubber stoppers and aluminum caps and 5% and 1% (v/v) of labeled or unlabeled CO2 and CH4 were injected into each corresponding microcosm through the rubber septum. The unlabeled and labeled urea and CO2 (99 atom% 13C) were purchased from the Shanghai Engineering Research Center of Stable Isotopes (Shanghai, China). The unlabeled and labeled CH4 (99 atom%
% $^{13}$C were purchased from Sigma-Aldrich (St Louis, MO, USA). The microcosms were incubated at 25 °C in the dark for 21 days. For the $^{13}$CH$_4$ microcosms, the concentrations of CH$_4$ were measured daily by gas chromatography (Shimadzu GC-2010 Plus, Japan), and $^{13}$CH$_4$ or $^{12}$CH$_4$ was then renewed to maintain 1% CH$_4$ in the microcosms. The measurement time and dynamic consumption of CH$_4$ are detailed in Table S2 and Fig. S1. Time and dynamic consumption of CH$_4$ for the comprehensive analysis of the microbial community [39]. The estimated absolute corresponding cell numbers was essential for the comprehensive analysis of the microbial community [39–41].

RESULTS

Greenhouse-gas emissions, methane, and ammonia oxidation activity

The in situ GHG measurements revealed that total N$_2$O emission was 0.134 kg ha$^{-1}$ in the grazed soils over the two years, and was significantly lower than that in the grazed soils (0.212 kg ha$^{-1}$).

Quantitative PCR (qPCR) of the amoA and pmoA genes

The qPCR assays targeting bacterial and archaeal amoA genes, as well as the pmoA genes of total DNA extracts and DNA gradient fractions, were carried out in triplicate with LightCycler 480 (Roche Applied Science). The primers and PCR conditions used are detailed in Table S3. Each qPCR was performed in a 20-µl reaction mixture containing 10 µl SYBR Premix Ex Taq (TaKaRa, Dalian, China), 0.5 µM of each primer and 1 µl of DNA template (10–100 ng) and an appropriate amount of milli-Q water to make a total volume of 20 µl. Melting curve analysis was performed at the end of each real-time PCR run to confirm PCR product specificity, by measuring fluorescence continuously with the temperature increasing from 50 °C to 99 °C. High efficiencies ranging between 80.5 and 108.3% were obtained for amplification of the functional genes, with $R^2$ values ranging between 0.990 and 0.998.

Illumina HiSeq sequencing and phylogenetic analysis

High-throughput sequencing of the V4 region of the 16S rRNA gene (amplified using the universal 515F-907R primer set) was used to assess the microbial community composition, including methane-oxidizers and nitrifiers. The HiSeq sequencing was carried out using the total DNA extracted from soil microcosms on day 0 and day 21 and the fractions 3–12 of the labeled $^{13}$CO$_2$ and $^{15}$CH$_4$ and control $^{13}$CO$_2$ and $^{12}$CH$_4$ microcosms at Day 21 (Tables S4, S5). Purified amplicons were pooled in equimolar and paired-end sequenced using an Illumina HiSeq 2500 (PE250) platform. The raw fastq files were demultiplexed, quality-filtered by Trimmomatic and merged by FLASH [37] according to the following criteria: (i) the reads were truncated at any site receiving an average quality score <20 over a 50-bp sliding window; (ii) sequences whose overlap was longer than 10 bp were merged according to their overlap sequence; (iii) sequences of each sample were separated according to barcodes (exactly matching) and primers (allowing two nucleotide mismatching), and reads containing ambiguous bases were removed. UPARSE was used to cluster cleaned sequences into operational taxonomic units (OTUs) at 97% similarity threshold. One representative sequence of each OTU was selected to determine taxonomic identification by RDP Classifier algorithm against the Silva (SSU128) 16S rRNA database. The 16S rRNA genes affiliated with aerobic MOB were selected based on two phyla: Proteobacteria and Verrucomicrobia. Of which, aerobic Proteobacterial MOB was divided into two major groups mainly based on phylogeny type I (Gammaproteobacteria) and type II (Alphaproteobacteria). Type I MOB harboring the family Methylococaceae, which were further divided into type Ia (Methylococcales), type Ib (Methylochloroccales) and type Ic (Methylovelularaceae). Type II MOB included the family Methylophagaceae (including genera Methylophaga and Methylosinus) and Beijerinckiaceae (including genera Methylocella, Methylocaldum, and Methyloferula). MmoX, a subunit of soluble methane monoxygenase, was not accounted for in the 16S sequence analysis. One phylum (Thaumarchaeota) was suggested for AOA. Comparative 16S rRNA sequence analyses of cultured AOB revealed that members of this physiological group are confined to two monophyletic lineages within β- and γ-proteobacteria. The former lineage includes the genera Nitrosomonas and Nitrosospira and the latter the genera Nitroscococcus, Nitrososphaeraceae and Nitrosoglobus. All reads classified as belonging to the genera Nitroscococcus, Nitrosococcus, Nitrosopina, Nitrobroder, and Nitrosira were used for NOB phylogenetic analysis.

A representative sequence was used from each OTU of the 16S rRNA for phylogenetic analysis. Phylogenetic analysis of archaeal and bacterial amoA genes, NOB, as well as pmoA genes of total DNA and of 13C-labeled DNA, was then conducted by Molecular Evolutionary Genetic Analysis software (MEGA6.0) with 1000-fold bootstrap support. The tree topology was checked by the neighbor-joining algorithm and the minimum evolution method [38]. The entire dataset of 16S rRNA gene reads of total and fractionated DNA was deposited in NCBI's Sequence Read Archive under BioProject accession numbers PRJNA594293 and PRJNA594236, respectively.

Since it is insufficient to reveal the changes of actual taxon abundances based only on those of relative abundances by high-throughput sequencing analysis [39, 40], the estimated absolute abundance (EAA) of each microbial genus considering both the corresponding relative abundance and the absolute corresponding cell numbers was essential for the network visualization was generated by Gephi 0.9.2 [42].

Statistical analysis

One-way analysis of variance followed by Duncan’s multiple range test was used to check for the significant differences between treatments for abundance of functional genes, nitrification potential, and methane-oxidizing potential using SPSS software (version 20). The independent samples t test was performed with SPSS 20 (IBM) to check for the significant differences between grazed and ungrazed soils for soil physicochemical properties and α-diversity indices. $P < 0.05$ was considered to be statistically significant. The α-diversity indices of active methane-oxidizing bacteria and nitrifying communities were calculated using the “vegan” packages in the R environment. Network analysis was performed by CoNet of cytosecpe 3.6.1 based on Spearman’s correlation for labeled communities of methane-oxidizers and nitrifiers in microcosms incubated at low and high levels of urea with or without methane supplementation in ungrazed and grazed soils. A co-occurrence event was validated when the Spearman’s coefficient was greater than 0.65 and the $P$ value was lower than 0.01. Nodes in the microbiome network represents individual microbial taxa (OTUs), and edges corresponded to the pairwise correlations between nodes. The co-occurrence networks were visualized by Gephi 0.9.2 with Community Detection plugin.
(P < 0.01) (Fig. 1). Total CH₄ emission was −3.7 kg ha⁻¹ in the ungrazed soils, and was significantly lower than that in grazed soils (−2.6 kg ha⁻¹).

In the incubated soils for 21 days in presence of CH₄, a total of 37, 51, and 24 µmol CH₄ g⁻¹ dry soil was oxidized in ungrazed soils, while 6, 11, and 3 µmol CH₄ g⁻¹ dry soil was oxidized in grazed soils under supplementation of 0, 20, and 100 µg N g⁻¹ of urea (Fig. 2a). Accordingly, the methane oxidation was significantly stimulated in the U₂₀ + CH₄ treatments but significantly suppressed in the U₁₀₀ + CH₄ treatments compared to U₀ + CH₄ soils during the incubation period for both ungrazed and grazed soils. The inconsistency in methane addition resulted from different methane oxidation abilities in these treatments, as we keep 1% CH₄ in the headspace of the microcosms. The inconsistency in methane addition just indicated different functional activity of methanotrophs in these treatments. The less amount of methane addition in U₁₀₀-treated soils compared with that in U₀ and U₂₀-treated soils was ascribed to the fact that high N levels (U₁₀₀) significantly depressed methane oxidation activity.

In both ungrazed and grazed incubated soils, net nitrification rate was higher in soil incubated with higher concentrations of urea (Fig. 2b), with a maximum of 180 and 136 µg NO₃⁻ g⁻¹ dry soil in U₁₀₀-supplemented soils in ungrazed and grazed soils, respectively. The nitrate concentration in the soils increased during the incubation period regardless of treatments, and ammonium decreased conversely over time with a sharper decrease during the first 7-days of the incubation (Fig. S2). Methane supplementation did not impact nitrification rate, apart in U₂₀-supplemented ungrazed soils, for which nitrification was reduced from 89 to 41 µg g⁻¹ following methane addition (Fig. 2b).

**Abundance of methylotrophic and nitrifying communities**

The abundance of pmoA-related organisms increased during the incubation under low urea supplementation (U₂₀) in both ungrazed and grazed soils, but decreased dramatically for the high urea supplementation treatment (U₁₀₀) compared to the control treatment (Fig. 3a; Table S6). A similar trend was observed for the relative abundance of methylotrophs estimated by 16S rRNA gene sequencing in microcosms incubated for 21 days (Fig. 3b; Table S6). The relative abundance of 16S rRNA genes affiliated with methane-oxidizing bacteria in both ungrazed and grazed soils was significantly stimulated by the low rate of urea addition (20 µg N g⁻¹), and decreased following the addition of 100 µg urea-N g⁻¹ compared to those without urea addition.

The abundance of bacterial amoA gene increased significantly over time in both ungrazed and grazed soils supplemented with high urea (Fig. 3c; Table S7). After incubation, the bacterial amoA gene abundance was significantly higher in U₁₀₀- than in U₂₀-treated soils. Methane addition did not affect the bacterial amoA gene abundance. The dynamic changes of the relative abundance of 16S rRNA genes affiliated with AOB were similar to the variations of bacterial amoA genes copy numbers under urea and methane treatments in both ungrazed and grazed soils (Fig. 3e; Table S7). By comparison, irrespective of methane addition, the abundance as
also occurred towards the heavy fractions (1.732 g ml\(^{-1}\)) of the AOB gene abundance (a), the relative abundance of targeted 16S rRNA genes affiliating with methane-oxidizing bacteria (b), the amoA gene copy numbers in total DNA (c), relative frequencies of targeted 16S rRNA genes affiliating with ammonia-oxidizing bacteria (d) in the ungrazed and grazed soils after 21-day incubation were represented. The error bars represent standard errors of three replicates for the U0 + CH\(_4\) treatments, and 6 replicates for the U20 and U100 treatments (\(^{13}\)C-labeled treatments) (a–c). The error bars represent standard errors of six replicates for the U20 and U100 treatments (\(^{13}\)C-labeled treatments) (d). Different lower-case letters indicate significant differences among different treatments. Asterisks indicate significant difference between ungrazing and grazing under the same urea treatment. d.w.s refers to dry weight of soil.

well as the relative abundance of *Thaumarchaeota* (including all known AOA) decreased significantly during the incubation period with urea addition in both ungrazed and grazed soils (Fig. S3).

In ungrazed soils, the relative abundance of targeted 16S rRNA genes affiliated with NOB increased over time in U20, U100, and U100 + CH\(_4\) microcosms, but remained constant in U20 + CH\(_4\) microcosms (Fig. S4a). On the contrary, in grazed soils, the relative abundance of NOB only increased in U100-treated microcosms (Fig. S4b).

**Active methane-oxidizing bacteria and nitrifying communities**

Isopycnic gradient centrifugation was conducted on the total DNA from each treatment to detect which putative autotrophic methanotrophs and ammonia-oxidizers were incorporating \(^{13}\)C-CH\(_4\) or \(^{13}\)C-CO\(_2\) during the 21-day incubation (Fig. 4). The buoyant density in the 15 fractions ranged from 1.676 to 1.786 g ml\(^{-1}\) from the top to the bottom of the ultracentrifugation tube. The abundances of *pmoA* and *amoA* genes in the 15 fractions were determined by specific qPCR assays (Figs. 4 and S5).

In ungrazed soils, the copy numbers of *pmoA* genes in all \(^{13}\)C-spiked microcosms peaked in heavy fractions (1.738–1.778 g ml\(^{-1}\)), while those in the presence of \(^{12}\)CH\(_4\) were detected in fractions with lower buoyant densities (between 1.680 and 1.738 g ml\(^{-1}\)) (Fig. 4a–c). In grazed soils, a shift of DNA buoyant density also occurred towards the heavy fractions (1.732–1.770 g ml\(^{-1}\)) in microcosms with \(^{13}\)CH\(_4\) under U0 and U20 treatments, but to a lesser extent than in the ungrazed soils. In addition, the distributions of *pmoA* gene abundance in microcosms with \(^{12}\)CH\(_4\) and microcosms under U100 + \(^{13}\)CH\(_4\) treatments were mainly in lighter fractions (1.698–1.732 g ml\(^{-1}\)) with no peak reported in the heavy fractions (Fig. 4f–h). Sequencing of the V3–V4 region of the 16S rRNA gene generated 1,124,000 and 1,088,000 high-quality reads in ungrazed and grazed soils, respectively (Table S4). The labeling of MOB was further supported by distribution of relative abundance of methanotrophs across the whole buoyant density (Fig. S6).

The \(^{13}\)C-labeling of the AOB populations was stronger in the U100 than U20 treatments in both ungrazed and grazed soils irrespectively of methane addition treatment (Fig. 4d, e and i, j). For the \(^{12}\)CO\(_2\) controls (U20 + \(^{13}\)CO\(_2\) + \(^{12}\)CH\(_4\) and U100 + \(^{13}\)CO\(_2\) + \(^{12}\)CH\(_4\)), the AOB *amoA* gene relative abundance was distributed in the light fractions (1.705–1.738 g ml\(^{-1}\)). A small but detectable shift of DNA buoyant density was detected in U20 + \(^{13}\)CO\(_2\) microcosms with the relative abundance of AOB peaking in the heavy fractions (1.732–1.742 g ml\(^{-1}\)) (Fig. 4d and i). A major shift into the heavier fractions was detected for bacterial *amoA* genes in U100 + \(^{13}\)CO\(_2\) microcosms, with a buoyant density of 1.750 g ml\(^{-1}\) (Fig. 4e and j). In contrast, no significant difference of migration between the \(^{12}\)CO\(_2\) and \(^{13}\)CO\(_2\) treatments was observed for the archaeal *amoA* gene (Fig. S5). The relative frequency of the 16S rRNA gene sequences affiliated with ammonia oxidizers across the whole buoyant density gradient of DNA fractions suggested the similar labeling trend to that of *amoA* gene abundance in the "heavy" DNA fractions (Fig. S7). Combined with the results that AOA *amoA* genes abundances decreased during incubation (Fig. S3), suggested that AOA was not important for ammonia oxidation in the grassland soil tested. Therefore, the possible bias against *Thaumarchaeota* due to primers used in the present study exerted little effect on the results.

In both ungrazed and grazed soils, a higher proportion of 16S rRNA genes affiliating to NOB were labeled in the \(^{13}\)CO\(_2\) than in the \(^{12}\)CO\(_2\) treatments, with more labeling happening in the absence of \(^{13}\)CH\(_4\) apart from the grazed microcosms with highest urea addition treatment (Fig. S8).

**Diversity of active methane-oxidizing bacteria and nitrifying communities**

There were obvious dissimilarities (P < 0.05) in diversity of active methane-oxidizing bacteria and nitrifying communities for the Shannon’s and Simpson’s indexes, respectively (Fig. S5). The Shannon’s and Simpson’s indexes of active MOB and AOB were significantly higher in ungrazed soils than those in grazed soils, while the variation of those of NOB was not significant between ungrazed and grazed soils. Grazing, therefore, significantly decreased the community diversity of active ammonia oxidizers and methanotrophs.

**Phylogenetic analysis of active MOB and AOB**

The taxonomic analysis focused on the organisms present in the "heavy fractions" in the \(^{13}\)CO\(_2\)/\(^{12}\)CH\(_4\) microcosms, as these AOB, NOB and MOB were actively growing (Figs. 4; S5–S8).

The majority of \(^{13}\)C-labeled MOB in ungrazed soils mainly belonged to *Methylobacter*, with some *Methylocaldum* and USCa, and AOB mainly grouped with *Nitrosospira*, with some
Nitrosomonas, and Nitrosococcus (Figs. 6a, c and S9). By contrast, the 13C-labeled MOB and AOB in grazed soils were almost exclusively affiliated with Methylobacter and Nitrosospiro, respectively (Figs. 6b, d and 9). In ungrazed soils, over 98% of active MOB sequences were derived from Methylobacter species in low and high urea amended soils, while 59.9% of active MOB were affiliated with Nitrosomonas in no urea treated soils (Fig. 6a).

The active AOB community grouped with Nitrosospira was significantly decreased from 67.0% and 71.1% in low and high urea treated soils without methane addition, respectively, to 65.5% and 62.6% in microcosms with methane addition (Fig. 6c).

In stark contrast, the active AOB classified into Nitrosomonas was significantly increased by methane addition in both low and high N soils. Irrespective of methane addition, the active AOB community classifying as Nitrosococcus was suppressed by high urea addition.

Absolute abundance of different active microbial genera affiliated with MOB and AOB

The estimated absolute abundances (EAA) of the major active genera affiliated with active MOB and AOB were performed based on the combined analysis of specific qPCR and Illumina HiSeq sequencing analysis (Fig. 7). The EAA results showed that Methylobacter lineage of MOB was the dominant active MOB in the grassland ecosystems (Fig. 7a, b). The Methylocaldum and USCα lineages of MOB were exclusively observed in ungrazed soils without urea addition (Fig. 7a). In both ungrazed soils and in grazed soils, the addition of U20 significantly increased while U100 decreased the absolute abundance of active Methylobacter lineage of MOB (Fig. 7a, b). The active bacterial amoA genes fell within Nitrosospira, Nitrosomonas, and Nitrosococcus, with Nitrosospira accounting for the largest percentage among labeled microcosms (Fig. 7c, d). In ungrazed soils, the absolute abundance of the three lineages of AOB was higher in U100 than U20 microcosms (Fig. 7c).

In grazed soils, the absolute abundance of Nitrosospira was 3.25-fold higher in U100 compared to U20 (Fig. 7d).

**Network description**

Network analysis based on co-occurring patterns of AOB, NOB, and MOB in "heavy fractions" in all microcosms was implemented to investigate potential relationships between the three functional guilds in ungrazed and grazed soils (Fig. 8). The resulting topological properties commonly used in network analysis in terms of the numbers of nodes and edges, average connectivity, and average clustering coefficient were detailed in Table S8. Positive and strong correlations (P < 0.05) were observed between active AOB and NOB phylotypes, while negative correlations (P < 0.05) were discerned between active MOB and the two above-mentioned nitrifiers in both ungrazed and grazed soils. In ungrazed soils, OTU 2, which fell within the Methylobacter and was the most abundant phylotype of active MOB (more than 80% of active MOB sequences), exhibited strong negative correlations with OTU 9, OTU 466, and OTU 703, which were affiliated with Nitrosospira, Nitrosococcus, and Nitrosomonas, respectively (Fig. 8a).

Furthermore, in grazed soils, a strong and negative relationship was detected between Methylobacter and Nitrosospira (Fig. 8b).

**DISCUSSION**

**Grazing reduced CH₄ uptake and increased N₂O emissions**

Grazing, associated with trampling, excreta patches and grazing grass, greatly alters plant growth, soil properties, and nutrient
transformations, especially affecting soil C/N cycling [31, 43, 44]. A large number of studies investigating the effects of grazing on GHGs emissions have proposed that grazing caused a significant decline in soil CH4 uptake [21, 43, 45]. It was reported that soil compaction [46–48] and drought or waterlogging stress [49] decreased methane oxidation and increased methane production. The inhibition effects of grazing on CH4 consumption in the present study could thus be mediated by soil bulk density and moisture, which were appreciably increased and decreased under grazing, respectively, as evidenced by the strong relationships between CH4 uptake and bulk density ($r = -0.962$) and soil moisture ($r = 0.966$) (Table 1). Intriguingly, the effect of grazing on N2O emissions remains controversial. The stimulating effects of grazing on N2O emissions in the present study were supported by previous studies [50–52], while contrasting results were also reported [45]. However, the N2O emissions observed under the grazing treatment are highly dependent on the season, with most of the annual N2O emissions being reported from semi-arid grasslands experiencing soil freeze-thaw cycles during the spring season, while grazing decreases N2O emissions during the spring thaw period [45, 53], and our present measurements were taken during the grass growing season from May to October annually within 14-months. Therefore, GHGs emissions from grazed grassland soils are a result of multiple factors, including the soil properties, climate conditions, and anthropogenic activities. The quantitative environmental impact of grazing on the global nitrogen cycle and on greenhouse gas emission needs to be fully considered and quantified to provide comprehensive recommendations for future grassland management.

**Communities and activity of methanotrophs and nitrifiers**

The significant higher Shannon’s and Simpson’s indexes in ungrazed soils revealed that diversity of active MOB and AOB in ungrazed soils were higher than those in grazed soils. We may therefore conclude that grazing decreases the community diversity of active nitrifiers and methanotrophs in grassland soils. This could be the direct reasons why grazing suppressed CH4 oxidation and nitrification activity (Fig. 2).

The strong positive correlation ($r = 0.841$, $P < 0.05$) between AOB amoA gene abundance and nitrification activity, together with the results of DNA-SIP, indicate that AOB rather than AOA dominated microbial ammonia oxidation in the grassland soils. The predominant role of AOB in ammonia oxidation is consistent with previous studies showing that AOB generally dominates ammonia oxidation in N fertilized neutral pH agricultural soils [14, 54–57], while AOA may dominate ammonia oxidation in acidic agricultural soils [58, 59] or in unfertilized grassland soils [60]. Surprisingly, albeit the low nitrification activity, grazing appreciably increased N2O emissions as well as AOB amoA gene copy numbers and absolute abundance of active AOB affiliated with *Nitrosospira* (Figs. 1, 2, 6, 7). Apart from the soil conditions caused by grazing favored for N2O emissions, the significant increase of active AOB affiliated with *Nitrosospira* could also partially explain the distinct increase of N2O emission in grazed soils, as *Nitrosospira* were considered as the main contributors to the N2O emissions [61]. The active AOB in grazed soils were all affiliated with *Nitrosospira* while those in ungrazed soils were grouped into *Nitrosospira* and *Nitrosomonas* (Fig. 6). *Nitrosospira* strains were previously shown to outcompete *Nitrosomonas* in microcosms [62] while *Nitrosomonas* has a greater nitrification.

**Fig. 5** Alpha-diversity indices of active methanotrophs and nitrifiers in grassland soils. Alpha-diversity measurements of Shannon index (a, c, e) and Simpson index (b, d, f) of active MOB (a, b), AOB (c, d) and NOB (e, f) in ungrazed and grazed soils. Different letters indicate significant differences ($P < 0.05$) based on the analysis of variance.
activity potential than *Nitrosospira* [63]. These findings might explain the unexpected increase of active AOB abundance with low nitrification activity in grazed soils. A field in situ experiment revealed that excreta depositions considerably stimulated N$_2$O emissions from nitrifer denitrification dominated by AOB in grassland ecosystems [44, 64]. Combined with the results that AOB dominated nitriification, we proposed that the high N$_2$O emissions in the grazed soils resulted from nitrifer denitrification possessed by *Nitrosospira* AOB in the present study. Although we failed to explore the activity of comammox in the grassland soils, comammox were found to lack NOR homologs and cannot produce N$_2$O via nitrifier denitrification [65]. Meanwhile, comammox yielded N$_2$O at levels that are comparable to AOA but much lower than AOB via nitrification [66]. Therefore, these results suggest that grazing tends to convert major N$_2$O-generating processes from nitrification to nitrifier denitrification, which was dominated by *Nitrosospira* lineage, rather than AOB in grazed soils. Nevertheless, more studies are needed to quantify the contribution of nitrification and nitrifier denitrification to N$_2$O emissions in grazing grassland in future.

**Nitrogen-triggered relationships between CH$_4$ oxidation and nitrification**

The methane oxidation was increased and decreased under low and high urea addition, respectively, compared to the non-urea microcosm in both ungrazed and grazed soils (Fig. 2a). Similarly, both the pmoA copy numbers and the relative frequency of targeted 16S rRNA affiliated with methanotrophs were suppressed by the high urea addition (Fig. 3). It has been reported that the response of the methanotrophs to different N levels is inconsistent, which is traditionally ascribed to the inherent characteristics of the methanotroph composition or the N load-tested [67]. In fact, the mineral N is essential for methanotrophs to form biomass [68]. That is why urea addition is usually observed to stimulate methane oxidation and MOB growth in soil [14–16]. The grassland soils in our present study had low fertility with low mineral N levels (Table 1), so the addition of 20 µg urea-N g$^{-1}$ increased the abundance of MOB communities (Fig. 3), resulting in the higher methane-oxidizing activity than in the absence of urea (Fig. 2a). By contrast, the application of 100 µg urea-N g$^{-1}$ reduced both

**Fig. 6**  Phylogenetic tree of the 13C-labeled 16S rRNA genes affiliated with MOB and AOB from the labeled microcosms after incubation for 21 days. Phylogenetic analysis of the 16S rRNA genes affiliated with MOB (a, b) and AOB (c, d) in 13C-labeled DNA. The designation “HF” indicates the 13C-DNA in the active fraction after the ultracentrifugation of the total DNA extract from the labeled microcosms. The designation “U0 + CH$_4$” indicates OTU-1 contains 13,312 reads with > 97% sequence similarity, accounting for 31.8% of the total AOB 16S rRNA gene reads in the 13C-DNA from the CH$_4$-treated soil microcosms. The scale bars represent 2% nucleic acid sequences divergence for the 16SrRNA genes in ungrazed (a) and grazed (b) soils, respectively. The designation “HF” indicates the 13C-DNA in the active fraction after the ultracentrifugation of the total DNA extract from the labeled microcosms. The designation “U20 + CO$_2$” indicates OTU-1 contains 1096 reads with > 97% sequence similarity, accounting for 65.5% of the total bacterial AOB 16S rRNA gene reads in the 13C-DNA from the U20 + CO$_2$-treated soil microcosms. The scale bars represent 2% nucleic acid sequences divergence for the 16SrRNA genes in ungrazed (c) and grazed (d) soils, respectively.
Table 1. Physicochemical properties of the grassland soils used in this study.

| Soil     | pH | Soil moisture (%) | Bulk density (g cm⁻³) | Soil organic matter (g kg⁻¹) | Total N (g kg⁻¹) | Total C (g kg⁻¹) | Olsen P (mg kg⁻¹) | Available K (mg kg⁻¹) | NO₃⁻-N (mg kg⁻¹) | NH₄⁺-N (mg kg⁻¹) |
|----------|----|-------------------|-----------------------|-----------------------------|-----------------|-----------------|-----------------|---------------------|----------------|----------------|
| Ungrazed | 7.19a | 14.3a              | 1.00b                 | 28.0a                       | 1.63a           | 17.0a           | 5.4b            | 199b                | 42.5b           | 3.5b           |
| Grazed   | 7.04b | 7.6b               | 1.54a                 | 14.7b                       | 0.80b           | 9.8b            | 9.3a            | 230a                | 54.6a           | 4.9a           |

Values are means of triplicate. Different lowercase letters indicate significant differences between two soil types at P < 0.05 based on the analysis of variance.
methane oxidation and MOB growth in the grassland soils compared to no and low urea amendments (Figs. 2a and 3). High N fertilization was shown to reduce methane oxidation in some unfertilized arable land [69], while no inhibition impact was observed in paddy soils receiving annual N fertilization [14, 15]. This discrepancy might be attributed to inherent properties of the soils used (soil texture, soil N, and organic matter content), as those two studies were all carried out in paddy soils which received N fertilization annually [14, 15], whereas our present experiment was conducted in grassland soils under oligotrophic conditions (Table 1). Indeed, ammonia likely acted as the competitive inhibitor for MMO under the high urea treatment, as previously demonstrated [11]. Together, these results suggested that N was a trigger unlocking putative competition between methane and ammonia oxidation in global carbon and nitrogen cycling.

Putative competition between active nitrifiers and MOB

This study observed negative correlations between *Methylcobacter*-like MOB and *Nitrosospira* lineage of AOB, while strong and positive correlations between AOB and NOB in both ungrazed and grazed soils (Fig. 8). *Methylcobacter* was the most responsive MOB in all the microcosms regardless of urea levels (Figs. 6a, b, 7a, b), indicating the competitive life strategy of *Methylcobacter* [67]. Similarly, *Nitrosospira* species dominated AOB communities and were ubiquitous in all treatments (Figs. 6c, d, 7c, d). Strong positive links could be attributed to niche overlap and cross-feeding, while negative relationships could be attributed to competition and amensalism [70]. The strong negative correlations between *Methylcobacter*-like MOB and *Nitrosospora* lineage of AOB suggest the putative competitive relationships between these two critical players of soil C and N cycling competing for N or even for oxygen in the soils [14]. The strong and positive correlations between AOB and NOB suggested putative syntrophic relationships between those nitrifiers, as the nitrite produced by AOB is a substrate for NOB [71].

In the ungrazed control soils, strong negative relationships occurred between *Methylcobacter* and both *Nitrosospora* and *Nitrocoecus* (Fig. 8a). In the grazed soils, the putative competitive relationships between MOB and AOB exclusively occurred between *Methylcobacter* and *Nitrosospira* (Fig. 8b). These results indicated that the network in ungrazed soils incorporated a substantially higher number of significant correlations than that in grazed soils. Previous study pointed out that grazing weakened the correlations between soil micro-food webs and ecosystem functions (soil C and N mineralization) [72]. To the best of our knowledge, the present study is the first work to reveal the impact of grazing on putative competition between active methanotrophs and nitrifiers. We did not directly analyze the interactions of one group upon another, and between the taxonomic groups and process rates. In addition, some important controls (e.g., UO + 12CH4, UO + 13CO2, UO + 12CO2) should be taken into consideration for further studies targeting the active methanotrophs and nitrifiers under N gradient. Nonetheless, our study unraveled the putative competitive relationships between active MOB and AOB based on the abundance, activity, composition, and network analysis of the functional genes. These results suggest that grazing decreases methane and ammonia oxidation activity, and diversity of nitrifiers as well as methanotrophs, and subsequently weakens the putative competitive relationships between methanotrophs and nitrifiers in grassland soils.

CONCLUSIONS

In this study, we linked the activity, composition, and potential relationships between active methanotrophs and nitrifying communities using stable-isotope techniques. *Methylcobacter* lineage of MOB and *Nitrosospira* lineage of AOB were the dominating active methane and ammonia oxidizing microorganisms, respectively, in ungrazed and grazed soils, and *Nitrocoecus* and *Nitrosporomans* lineages of AOB were also involved in ammonia oxidation in ungrazed soils. The diversities of active MOB and AOB communities were higher in ungrazed than in grazed soils. And the network of co-occurring active phylootypes of MOB, AOB, and NOB was also more complex in ungrazed than in grazed soils, while the methane and ammonia oxidation activities were higher in ungrazed than in grazed soils. Therefore, these results suggest that grazing decreased diversity of active microbes mediated in C/N cycling, thus suppressing both methane oxidation and nitrification activity, weakening their putative competitive relationships, and thereby increasing methane emissions. These studies will help to understand the complex biotransformation processes of C and N, which are important for development of management practices to mitigate greenhouse-gas emissions in grassland soils.

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AUTHOR CONTRIBUTIONS
HP, YL, and CG-R conceived and designed the project. HP performed the research. YWL and CYL assisted with sample collection. HP and HJF analyzed the data and created the figures. HP wrote the manuscript. YL, ZJJ, YPZG, QCZ, CXT, JMX, HJD, and CG-R revised the paper. All authors discussed and interpreted the data and contributed to the manuscript. All authors read and approved the final manuscript.

COMPETING INTERESTS
The authors declare no competing interests.

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