Adaptive Multi-Paddock Grazing Lowers Soil Greenhouse Gas Emission Potential by Altering Extracellular Enzyme Activity

Bharat M. Shrestha 1,†, Edward W. Bork 2, Scott X. Chang 1,*, Cameron N. Carlyle 2, Zilong Ma 1, Timm F. Döbert 3, Dauren Kaliaskar 1 and Mark S. Boyce 3

1 Department of Renewable Resources, University of Alberta, Edmonton, AB T6G 2E3, Canada; bshresth@ualberta.ca (B.M.S); zilong@ualberta.ca (Z.M.); kaliaska@ualberta.ca (D.K.)
2 Department of Agricultural, Food and Nutritional Science, University of Alberta, Edmonton, AB T6G 2P5, Canada; ebork@ualberta.ca (E.W.B.); cameron.carlyle@ualberta.ca (C.N.C.)
3 Department of Biological Sciences, University of Alberta, Edmonton, AB T6G 2R3, Canada;
doebert@ualberta.ca (T.F.D.); boyce@ualberta.ca (M.S.B.)
* Correspondence: sxchang@ualberta.ca; Tel.: +1-780-492-6375
† Current address: Agriculture and Agri-Food Canada, Beaverlodge Research Farm, Beaverlodge, AB T0H 0C0, Canada.

Received: 18 September 2020; Accepted: 10 November 2020; Published: 13 November 2020

Abstract: Adaptive multi-paddock (AMP) grazing is a form of rotational grazing in which small paddocks are grazed with high densities of livestock for short periods, with long recovery periods prior to regrazing. We compared the fluxes of greenhouse gases (GHGs), including carbon dioxide (CO₂), methane (CH₄), and nitrous oxide (N₂O), from soils of AMP-grazed grasslands to paired neighboring non-AMP-grazed grasslands across a climatic gradient in Alberta, Canada. We further tested GHG responses to changes in temperature (5°C vs. 25°C) and moisture levels (permanent wilting point (PWP), 40% of field capacity (0.4FC), or field capacity (FC)) in a 102-day laboratory incubation experiment. Extracellular enzyme activities (EEA), microbial biomass C (MBC) and N (MBN), and available-N were also measured on days 1, 13, and 102 of the incubation to evaluate biological associations with GHGs. The 102-day cumulative fluxes of CO₂, N₂O, and CH₄ were affected by both temperature and moisture content (p < 0.001). While cumulative fluxes of N₂O were independent of the grazing system, CH₄ uptake was 1.5 times greater in soils from AMP-grazed than non-AMP-grazed grasslands (p < 0.001). There was an interaction of the grazing system by temperature (p < 0.05) on CO₂ flux, with AMP soils emitting 17% more CO₂ than non-AMP soils at 5°C, but 18% less at 25°C. The temperature sensitivity (Q₁₀) of CO₂ fluxes increased with soil moisture level (i.e., PWP < 0.4FC ≤ FC). Structural equation modelling indicated that the grazing system had no direct effect on CH₄ fluxes, but had an effect on CH₄ fluxes on days 1 and 13, indicating that CH₄ uptake increased in association with AMP grazing. Increasing soil moisture level increased fluxes of GHGs—directly and indirectly—by influencing EEAs. Irrespective of the grazing system, the MBC was an indirect driver of CO₂ emissions and CH₄ uptake through its effects on soil EEAs. The relationships of N-acetyl-β-glucosaminidase and β-glucosidase to N₂O fluxes were subtle on day 1, and independent thereafter. AMP grazing indirectly affected N₂O fluxes by influencing N-acetyl-β-glucosaminidase on day 13. We conclude that AMP grazing has the potential to mitigate the impact of a warmer soil on GHG emissions by consuming more CH₄ compared to non-AMP grazing in northern temperate grasslands, presumably by altering biogeochemical properties and processes.

Keywords: climate change; grazing management; GHG mitigation; northern temperate grasslands; soil incubation; soil moisture; temperature
1. Introduction

Grasslands cover more than 30% of terrestrial land globally and generate important ecological services, including enhancing food security by providing forage for more than 1.8 billion livestock and holding 33% of the terrestrial carbon (C) stock [1]. However, grasslands can affect global climate change by being a sink or source of greenhouse gases (GHGs), including carbon dioxide (CO$_2$), methane (CH$_4$), and nitrous oxide (N$_2$O), depending on several drivers, including their responses to management [2,3]. The identification of grazing management practices that achieve GHG reductions within grasslands [4,5] would have direct social and policy implications for land use and management [6]. Although GHG emissions from grasslands are responsive to grazing [7–9], grasslands vary in their response, depending on which specific grazing practices are used [9]. Moreover, under future climate change scenarios, increased temperature and altered soil moisture may interact with grazing to affect GHG dynamics, because fluxes of GHG vary with temperature and moisture level [10–12] depending on the defoliation level [7].

Adaptive multi-paddock (AMP) grazing is a form of rotational grazing in which grasslands are sub-divided into many small paddocks and high densities of animals graze for short periods within each paddock, with long rest periods between successive grazing events to facilitate vegetation recovery [13,14]. The AMP grazing system has been reported to improve physical, chemical, and biological soil properties compared with areas subject to continuous grazing [15], in which cattle graze throughout the growing season in the same area. In particular, vegetation recovery is suggested to be more rapid in AMP-grazed areas due to enhanced control over the extent of defoliation and more uniform impact of animals on grassland soils, including the physical impact on soils through concentrated animal activity (“herd effect”) [16] and distribution of excreta [13]. Ultimately, these practices have been suggested as a means to increase the fixation of atmospheric C into plant biomass, and bolster soil organic C (SOC) [13] and macro-nutrients [17], improve soil function and health, and mitigate climate change through increased C storage [18,19]. There are multiple ways AMP grazing may alter vegetation and soils leading to altered C cycling. High stocking densities can increase soil compaction [20], affecting bulk density, penetration resistance, and water infiltration [21] slowing decomposition processes [22]. Trampling can increase fine litter, which is typically a source of more recalcitrant C, incorporation into soils [19], and can subsequently alter microbial abundance and composition [23]. Furthermore, AMP grazing may alter the chemical composition of vegetation by favoring grazing-tolerant plants that tend to have more recalcitrant carbon [24] or, in contrast, by encouraging plant regrowth, which tends to be less recalcitrant [15]. However, the benefits of AMP grazing in boosting productivity, maintaining grassland function, and enhancing C sequestration remain unclear and are sources of debate [25,26]. An understanding is therefore needed of the mechanisms regulating C and nutrient cycling under AMP grazing compared to non-AMP systems and to parse out the role of AMP grazing on GHG emissions. Measurements of GHGs in the field are influenced simultaneously by variation in environment and management regimes. Thus, controlled incubation studies are one approach to specifically test the grazing legacy effects of soil microbes in altering pathways of nutrient cycling in soils [27], and thereby to formulate grazing management strategies to mitigate atmospheric GHGs [9].

Soil extracellular enzyme activity (EEA) regulates soil organic matter (SOM) decomposition and nutrient cycling [28], thereby influencing GHG fluxes from the soil. Plants and microbes release enzymes during the decomposition process of SOM, catalyzing the decomposition of target molecules [29]. As different enzymes can be responsible for decomposing a single biopolymer in SOM, several enzymes need to be measured simultaneously to fully understand the role of EEAs in the cycling of C and nutrients such as nitrogen (N). Soil EEAs can change in response to biotic (e.g., vegetation, faunal influences) and abiotic conditions of soils (e.g., temperature and moisture) that affect soil physical and chemical properties [30,31]. Thus, measurement of EEAs in grassland soil under different microclimatic conditions is necessary to help understand the role of the grazing-induced environment in affecting GHG fluxes from the soil. While effects of temperature and moisture on GHG emissions have been widely studied in both the field [7,32] and in laboratory conditions [33,34],
none have examined the specific influence of AMP grazing and the role of associated EEAs in regulating GHG fluxes.

The goal of this study was to determine whether AMP grazing alters potential GHG fluxes from grassland soils relative to neighboring conventional grazing practices (hereafter “non-AMP”), and to test whether the grazing system has altered the sensitivity of GHG fluxes to changes in soil temperature and moisture. We conducted a laboratory incubation experiment to test the a priori influence of grazing system on GHG emissions and associated microbial activity. An additional objective was to explore the effects of grazing regime, temperature, and moisture conditions on microbes and EEAs, and resultant GHG fluxes in grassland soils.

2. Materials and Methods

2.1. Study Sites and Soil Collection

Grasslands under AMP grazing were initially identified by a voluntary selection process from a pool of attendees at a series of rancher workshops held across Alberta and more widely advertised using local producer groups. Prospective candidates responded to a series of questions concerning their grazing management activities using an online self-registration system. To qualify as an AMP ranch, ranches had to meet criteria regarding the number of paddocks used per herd (>10), the minimum size of the ranch (>65 ha), the frequency of cattle rotation, and the use of flexible adjustment of stocking density in response to climatic variation across the region. These conditions were subsequently verified via phone interviews and field visits. Additionally, we required that AMP ranches had used this system for at least 10 years, that no cultivation/seeding was done in the last 10 years, and that each AMP ranch had a neighboring ranch (within 10 km) with a similar cultivation history and ecosite conditions (landform and soil type) supporting cattle grazing. Moreover, AMP ranches were considered for study only if a portion of the grasslands therein were free of bale grazing and available for soil sampling to avoid confounding effects of additional C and nutrient inputs from hay.

Grasslands from a total of 11 pairs of ranches were selected across south-central Alberta, Canada, for this study. Each pair comprised of AMP grasslands and their neighboring non-AMP grasslands, where the latter utilized conventional grazing management. The 11 pairs represented a broad agro-climatic (i.e., soil and vegetation) gradient across northern temperate grasslands of Alberta, Canada. Selected ranches were, in order of declining aridity, situated within the Mixed grass, Aspen parkland, Foothills fescue, and Boreal transition regions. Soils coinciding with these natural regions were Orthic Brown Chernozems (Mixed grass), Orthic Black to Eluviated Black Chernozems (Foothills and Parkland), and Dark Gray Chernozems to Gray Luvisols (Boreal). Soil organic matter content ranges from 2.5 to 3.4% in Brown, 3.5 to 5.5% in Gray, and 5.5 to 8.5% in Black Chernozem soils [35]. The 30-yr normal (1984–2014) mean annual precipitation (MAP) ranged from 332.3 to 533.3 mm, with mean annual temperatures (MAT) ranging from 2.0 to 4.1 °C. The annual heat moisture (AHM, \[AHM = (MAT + 10)/(MAP/1000)\]) index, an index of aridity and moisture limitations on ecosystem productivity [36,37], ranged from 24.3 in moist areas to 44.1 in arid areas. AHM is a useful climatic variable because it accounts for both changes in moisture and temperature [38].

Six sampling points were randomly selected within a representative grassland area of 10 ha on each of the 22 studied ranches. Two mineral soil cores (3.8 cm diameter, 15 cm deep) from each sampling location were collected in the last week of August 2017. After overlying litter and mulch were removed, six mineral soil samples from each ranch were combined, bagged, and placed in a cooler to transport to the University of Alberta, where they were stored at 4 °C until processed during the second week of November.

2.2. Soil Processing and Characterization

Soil moisture content was determined from 20 g sub-samples by weighing them fresh, drying at 105 °C for 27 h to a constant mass, and then reweighing. Bulk soil was air-dried and sieved through a
2 mm screen, with all coarse fragments removed, including rocks, roots, and litter. Sub-samples of air-dried soils were ground to 0.1 mm size with a ball mill (Retsch MM200 Mixer Mill, Thomas Scientific, Swedesboro, NJ, USA) and then analyzed for total C and N by dry combustion using an automated elemental analyzer (Elementar Analysensysteme GmbH, Langenselbold, Germany). Soil pH was measured with a 1:5 (w:v) mix of soil:water [39], bulk density was determined using the core method [40], and the texture was determined using the hydrometer method [41].

2.3. Soil Preparation, Incubation, Gas Sampling, and Analysis

Water holding capacity of sieved soils at different matric potentials was determined using the pressure-plate method [42]. Sub-samples of air-dried soils were first placed in O-rings on ceramic porous plates and saturated for 24 h. Saturated soils were then pressurized at 0.1, 0.5, 1.0, 5.0, and 15.0 bars for 72 h, after which the moisture content at each pressure level was quantified by drying at 105 °C for 27 h to a constant mass and reweighing. Water content at 15 bar was considered the permanent wilting point (PWP), while 0.1 bar was the field capacity (FC) of sandy soils [43] and 0.33 bar the FC of clayey soils [42]. Water content at 0.33 bar was estimated by linear extrapolation of water contents at 0.1, 0.5, and 1.0 bar. The moisture content of air-dried sieved soil was also determined following the oven-dry method (described above) to help maintain the desired soil moisture level throughout the subsequent incubation experiment.

For each grassland investigated, 100 g of oven-dry equivalent air-dried soil was placed in each of six 500 mL Mason jars for the incubation experiment. Sufficient water was added (with a dilute 0.005 M CaSO\textsubscript{4} to protect micro-aggregates from disruption) to bring these soils to a moisture level of either FC, 40% FC, or PWP [44]. One set of Mason jars with soil from each moisture treatment was placed in an incubator at 5 °C, while the other set was placed in another incubator at 25 °C. The tops of all jars were covered with perforated aluminum foil for five consecutive days to stabilize microbial activity. On the fifth day (collection day 0), initial GHG samples were collected from the headspace air of the jars immediately after closing them using a lid equipped with a rubber septum. Soils were further incubated for 24 h with the lids closed, and then headspace samples were collected again to determine the change in GHG concentrations. Subsequent sampling of GHGs occurred on days 1, 2, 4, 7, 10, 13, 18, 23, 28, 35, 42, 52, 62, 72, 82, 92, and 102. The change in gas concentration between the 0 and 24 h headspace samples on each sampling day was used to calculate daily GHG flux per unit dry mass of soil. Soil moisture levels were maintained throughout the incubation period by tracking water loss by weighing the jars and replenishing the water at least 3 days prior to each gas sampling event. Headspace air samples were collected with an air-tight 20 mL syringe (Norm-Ject, Henke Sass Wolf, Tuttinglen, Germany) and injected into 12 mL pre-vacuumed soda glass Isomass Exetainers (Labco Limited, Lampeter, Wales, UK).

Greenhouse gas samples were analyzed with a Varian CP 3800 gas chromatograph (Varian Canada, Mississauga, Canada) containing three detectors. A thermal conductivity detector (TCD) and flame ionization detector (FID) simultaneously determined the concentration of CO\textsubscript{2} and CH\textsubscript{4}, respectively [45], while the electron capture detector (ECD) determined the concentration of N\textsubscript{2}O [46]. Standard curves were generated using mixtures of gases at standard concentrations of CO\textsubscript{2} (360 ppm), CH\textsubscript{4} (1.6 ppm) and N\textsubscript{2}O (1.0 ppm) (Praxair, Mississauga, Ontario, Canada) and used to calculate the headspace concentrations of respective gases.

2.4. Measurements of Microbial Activities and Soil Parameters

A parallel set of soils at the same moisture level were prepared by placing 50 g of oven-dry equivalent air-dried soil in 200 mL conical flasks for measuring extracellular enzyme activities (EEAs), microbial biomass C (MBC) and N (MBN), and reactive N (available-N), on day 1 (start), day 13, and day 102 (end) of the incubation period.

Activities of select extracellular enzymes involved in C (xylosidase: Xylo, β-glucosidase: BG, cellobiosidase: Cello) and N (N-acetyl-β glucosaminidase: NAG) cycling in soil were
analyzed. To assess the EEA, a standard fluorometric method was used with 96-well microplates (see Sinsabaugh et al. [47]) with acetate buffer solution (pH 5.0). One gram of fresh soil and 125 mL of buffer were mixed to make a soil solution and 200 µL of the solution was pipetted into each well of the microplate. Depending on the enzyme type, microplates with soil solutions and enzyme substrates were incubated for three (BG, NAG), four (Xylo), or seven hours (Cello) at 25 °C. After incubation, microplates were read on a Biotek Synergy HT (BioTek Instruments, Inc., Winooski, VT, USA) with 360 nm excitation and 460 nm emission [48]. Substrates used in this experiment were 4-MUF-β-D-glucopyranoside, 4-MUF-β-D-cellobioside, 4-MUF-β-D-xyloside, and 4-MUF-N-acetyl-β-glucosaminide.

Soil MBC and MBN were analyzed by the chloroform fumigation-extraction method [49, 50]. For fumigation, 10 g of moist soil sample was fumigated with chloroform in a desiccator for 24 h. Soil extracts were obtained by mixing 10 g of moist soil with 50 mL of 0.5 mol L\(^{-1}\) K\(_2\)SO\(_4\) solution, shaking for 1 h in a reciprocating shaker (250 rpm), and filtering through Q\(_2\) filter papers. Soil extractions were analyzed for MBC and MBN by a TOC-V analyzer connected to a TN module (Shimadzu Corporation, Kyoto, Japan). The MBC and MBN were calculated as the difference between the C and N extracted from fumigated and non-fumigated soil samples, respectively.

Soil NO\(_3\)\(^-\) and NH\(_4\)\(^+\) were determined using the colorimetric method in soil solution. The vanadium oxidation method was used for NO\(_3\)\(^-\) [51], and the indophenol blue method was used for NH\(_4\)\(^+\) [52] and analyzed on a spectrophotometer (GENESYS\textsuperscript{TM} 10S UV-Vis Spectrophotometer, ThermoFisher Scientific, Ottawa, Canada). The sum of NH\(_4\)\(^+\)-N and NO\(_3\)\(^-\)-N was expressed as total available N (avail-N). The MBC, MBN, and avail-N on each sampling day were calculated per unit mass of soil (mg kg\(^{-1}\) soil).

2.5. Data Preparation

All GHG concentrations were converted to gas fluxes per unit mass of dry soil using the following Equation (1), modified after Lang et al. [34]:

\[
R = \frac{\rho \times \Delta c 	imes V \times 273}{W \times \Delta t \times (273 + T)} \times 24
\]

where,
- \(R\) = flux of GHGs, specifically CO\(_2\) (mg CO\(_2\)-C kg\(^{-1}\) day\(^{-1}\)), N\(_2\)O (µg N\(_2\)O–N kg\(^{-1}\) day\(^{-1}\)), and CH\(_4\) (µg CH\(_4\)-C kg\(^{-1}\) day\(^{-1}\)),
- \(\rho\) = density of N\(_2\)O, CO\(_2\), or CH\(_4\) in a standard state,
- \(\Delta c\) = change in gas concentration between incubation times of \(t_1\) (0 h) and \(t_2\) (~24 h) (ppbv h\(^{-1}\) or ppmv h\(^{-1}\)),
- \(V\) = volume of the Mason jar (mL),
- \(T\) = incubation temperature (°C),
- \(W\) = dry weight of soil (kg), and
- \(\Delta t\) = time difference (h) between GHG measurements (\(t_2 - t_1\)).

The temperature sensitivity of CO\(_2\) flux (\(Q_{10}\)) was calculated using the following Equation (2) [53, 54]

\[
Q_{10} = \left(\frac{R_2}{R_1}\right)^{(T_2 - T_1)/10}
\]

where,
- \(R_2\) and \(R_1\) are the cumulative CO\(_2\) emissions measured at temperatures \(T_2\) and \(T_1\), respectively, and \(T_2 > T_1\).

The proportion of total mineralized SOC as CO\(_2\) was calculated as follows:

\[
% C_{min} = \frac{\sum_{i=1}^{102} (CO_2 - C) \times 100}{Total SOC in soil sample}
\]
where,

- \( \% C_{\text{min}} \) is the proportion of total organic carbon mineralized as CO\(_2\) throughout the incubation experiment,
- CO\(_2\)-C is the cumulative sum of SOC mineralized as CO\(_2\) during the 102-day incubation period.

The net flux of cumulative GHGs over the entire experimental period was calculated as follows [55]:

\[
\text{Net GHG flux} = \text{CO}_2 + (\text{CH}_4 \times 28) + (\text{N}_2\text{O} \times 265)
\]  

(4)

where,

- Net GHG flux is the sum of all GHGs (mg CO\(_2\)-e kg\(^{-1}\)), and 28 and 265 are the global warming potential of CH\(_4\) and N\(_2\)O, respectively, compared to CO\(_2\) given a 100 y life span of trace gases [55].

Resulting EEA rates were expressed in \( \mu \text{mol h}^{-1} \text{g}^{-1} \) of oven-dry soil using the following equation [47].

\[
\text{Enzyme activity} \left( \mu\text{mol g soil}^{-1}\text{h}^{-1} \right) = \frac{\text{Signal} \times 125 \text{ mL}}{1000 \times \text{Time(h)} \times 0.2 \text{ mL} \times W / (1 + M)}
\]  

(5)

where,

- Enzyme activity = EEA rate \( (\mu\text{mol h}^{-1} \text{g}^{-1} \text{dry soil}) \),
- \( W \) = the fresh weight of soil in g, and
- \( M \) = the moisture content of soil.

\[
\text{Signal} \ (\text{nmol}) = \frac{(\text{Assay}/E_c \times Q_c) - (\text{Substrate}/E_c) - (\text{Soil}/E_c \times Q_c) + (\text{Buffer}/E_c \times Q_c)}{E_c \times Q_c}
\]  

(6)

where,

- \( E_c \) is the emission coefficient, and
- \( Q_c \) is the quench coefficient.

2.6. Statistical Analyses

Normality and homogeneity of variance for all data were tested with Shapiro–Wilk tests in univariate analyses. A log transformation was applied to the total 102-day cumulative flux of CO\(_2\) and N\(_2\)O, and a cube root transformation was applied to CH\(_4\) flux. However, non-transformed values are presented for ease of data interpretation. The fixed effects of the grazing system, temperature, and moisture level were then analyzed on cumulative GHGs using a 3-way factorial mixed model ANOVA using transformed data (where appropriate) for a split-split plot experimental design. Grazing is the whole plot factor, incubation temperatures the sub-plot (i.e., incubation chambers) factor, and moisture level the sub-sub plot factor. The grazing system, soil temperature, and moisture were fixed effects, with blocked ranch pairs as random effects. Temperature sensitivity \( (Q_{10}) \) of CO\(_2\) flux was analyzed in a two-way mixed model with the grazing system and soil moisture as fixed effects, and ranch pair as random effects. All effects were evaluated at the 5% level of significance. All analyses were performed using the software RStudio version 1.2.5033 [56].

We used structural equation modelling (SEM) to evaluate the relationships among grazing, soil temperature, soil moisture, MBC, individual EEAs, and soil GHG emissions. We developed a conceptual SEM model (Figure S1), hypothesizing direct effects of MBC and EEA on GHG fluxes, MBC on EEA, and grazing, soil temperature, and soil moisture effects on MBC, EEA, and GHG fluxes. The categorical variables “grazing”, “soil temperature” and “soil moisture” were coded as 0 for non-AMP and 1 for AMP; 0 for 5 °C, and 1 for 25 °C; 0 for PWP, 1 for 0.4FC, and 2 for FC, respectively. Following Grace [57], we assessed the conceptual model (full model) vs. reduced models by the goodness-of-fit statistics and used akaike information criterion (AIC) to select the final model among alternative models. The final model had the lowest AIC value. We conducted SEM analysis using the “piecewiseSEM” package in R software [58].
3. Results

3.1. Basic Soil Properties

Soil physical and chemical properties, including bulk density, texture, moisture content, pH, SOC, and N, were not affected by the grazing system (Table S1). However, SOC \((p = 0.023)\) and soil available-N \((p = 0.05)\) varied with the geographic location of the grasslands studied, as represented by AHM (data not shown). More arid grasslands (higher AHM index) were associated with lower SOC \((r = -0.62; \ p < 0.001)\) and available-N \((r = -0.54; \ p < 0.001)\), and a higher C:N ratio \((r = 0.86; \ p < 0.001)\).

3.2. Effects of Grazing, Soil Temperature, and Moisture on Cumulative GHG Fluxes

Total emissions of CO\(_2\) during the entire incubation period were affected by soil temperature and moisture \((p < 0.001)\), with a further interaction of grazing \(\times\) soil temperature (Table 1). At 5 °C, AMP soils emitted 17% more CO\(_2\) compared to non-AMP soils, while at 25 °C, AMP soils emitted 18% less CO\(_2\) than non-AMP soils (Figure 1a). Fluxes of CO\(_2\) from soils at FC were 2.1 and 2.7 times greater compared to soils at PWP and at 0.4FC, respectively; notably, the responses to moisture remained similar in both grazing systems (Figure 1e).

Table 1. Summary of ANOVA, degree of freedom, F- and \(p\)-values for the cumulative flux of greenhouse gases (GHGs) during the 102-day long incubation period. \(p\)-values in bold are \(\leq 0.05\).

| Response Variable | Fixed Effect | Degree of Freedom \(^v\) | F-Value | \(p\)-Value |
|-------------------|--------------|---------------------------|----------|-----------|
| Carbon dioxide (CO\(_2\)) | Grazing (G) | 1, 10 | 0.41 | 0.534 |
| | Temperature (T) | 1, 100 | 555.63 | \(<0.001\) |
| | Moisture (M) | 2, 100 | 120.61 | \(<0.001\) |
| | G \(\times\) T | 1, 100 | 10.06 | 0.020 |
| | G \(\times\) M | 2, 100 | 0.64 | 0.528 |
| | T \(\times\) M | 2, 100 | 1.13 | 0.326 |
| | G \(\times\) M \(\times\) T | 2, 100 | 0.10 | 0.905 |
| Nitrous oxide (N\(_2\)O) | G | 1, 10 | 0.33 | 0.578 |
| | T | 1, 100 | 47.69 | \(<0.001\) |
| | M | 2, 100 | 45.21 | \(<0.001\) |
| | G \(\times\) T | 1, 100 | 3.06 | 0.084 |
| | G \(\times\) M | 2, 100 | 0.42 | 0.660 |
| | T \(\times\) M | 2, 100 | 0.37 | 0.689 |
| | G \(\times\) M \(\times\) T | 2, 100 | 0.44 | 0.645 |
| Methane (CH\(_4\)) | G | 1, 10 | 6.81 | 0.026 |
| | T | 1, 100 | 44.88 | \(<0.001\) |
| | M | 2, 100 | 24.65 | \(<0.001\) |
| | G \(\times\) T | 1, 100 | 0.26 | 0.609 |
| | G \(\times\) M | 2, 100 | 0.68 | 0.510 |
| | T \(\times\) M | 2, 100 | 0.08 | 0.927 |
| | G \(\times\) M \(\times\) T | 2, 100 | 0.16 | 0.852 |
| Net GHG \(^§\) | G | 1, 10 | 0.39 | 0.547 |
| | T | 1, 100 | 581.50 | \(<0.001\) |
| | M | 2, 100 | 122.82 | \(<0.001\) |
| | G \(\times\) T | 1, 100 | 10.55 | 0.002 |
| | G \(\times\) M | 2, 100 | 0.64 | 0.528 |
| | T \(\times\) M | 2, 100 | 1.04 | 0.358 |
| | G \(\times\) M \(\times\) T | 2, 100 | 0.11 | 0.898 |
| Q\(_{10}\) (CO\(_2\)) \(^‡\) | G | 1, 10 | 12.19 | 0.006 |
| | M | 2, 40 | 7.27 | 0.002 |
| | G \(\times\) M | 2, 40 | 0.68 | 0.514 |
| C\(_{\text{min}}\)/SOC \(^†\) | G | 1, 10 | 0.95 | 0.354 |
| | T | 1, 100 | 436.89 | \(<0.001\) |
| | M | 2, 100 | 110.22 | \(<0.001\) |
| | G \(\times\) T | 1, 100 | 4.36 | 0.039 |
| | G \(\times\) M | 2, 100 | 1.09 | 0.190 |
| | T \(\times\) M | 2, 100 | 35.35 | \(<0.001\) |
| | G \(\times\) M \(\times\) T | 2, 100 | 0.35 | 0.703 |

\(^v\) Degrees of freedom (numerator, denominator); \(^§\) Net GHG = Resultant flux of greenhouse gases; \(^‡\) Q\(_{10}\) = Temperature sensitivity of cumulative CO\(_2\) flux; \(^†\) C\(_{\text{min}}\)/SOC = Percentage of soil organic C (SOC) mineralized as CO\(_2\)-C.
Figure 1. Left panels show effects of grazing and temperature on total fluxes of GHGs (mean ± SE) in a 102-day incubation experiment on soils from either adaptive multi-paddock (AMP) or conventional (non-AMP) grazing systems: (a) CO$_2$-C, (b) N$_2$O-N, (c) CH$_4$-C, and (d) net GHG flux. Right panels show effects of grazing and moisture on total flux of GHGs: (e) CO$_2$-C, (f) N$_2$O-N, (g) CH$_4$-C, and (h) net GHG flux. Negative values show the consumption of CH$_4$ in the soil. Statistics in the inset box show an interaction between grazing and temperature on (a) CO$_2$ flux and (d) net GHG flux ($p < 0.05$).

Cumulative fluxes of N$_2$O during the incubation period were independent of grazing ($p \geq 0.58$), but remained strongly influenced by soil temperature and moisture ($p < 0.001$, Table 1). Soils at 25 °C produced 3.4 times more N$_2$O than soils at 5 °C (Figure 1b). Mean N$_2$O flux from soils at FC were 1.5 and 3.1 times higher than in soils at PWP and 0.4FC, respectively (Figure 1f).

The cumulative uptake of CH$_4$ during the incubation period was affected by the grazing system, soil temperature, and moisture (Table 1). Overall, CH$_4$ uptake was 2.6-fold greater in AMP soils (52.4 ± 6.4 µg CH$_4$-C kg$^{-1}$) in comparison to non-AMP soils (20.4 ± 2.5 µg CH$_4$-C kg$^{-1}$) (Figure 1c). Similarly, soils at 25 °C had 2.9-fold greater uptake of CH$_4$ compared to soils incubated at 5 °C (Figure 1c). Furthermore, uptake of CH$_4$ was 2.6-fold greater in soils at FC than soil at PWP ($p < 0.001$), and the latter, in turn, was double ($p < 0.001$) that in soils at 0.4FC (Figure 1g).
Net GHG emissions (CO₂-e) throughout the incubation varied with soil temperature and the interaction of grazing with temperature (Table 1). Net GHG emissions were 15% lower in non-AMP than in AMP systems at 5 °C, but were 22% higher in non-AMP soils at 25 °C compared to the AMP soils (Figure 1d). Net GHG emissions were also affected by soil moisture (Table 1). Soils at FC emitted 3.1 and 3.7 times more net GHGs (p < 0.05) in comparison to soils at either PWP or 0.4FC, while the latter did not differ from one another (Figure 1g).

The temperature sensitivity (Q₁₀) of CO₂ flux within these grassland soils differed between the AMP and non-AMP grazing systems (Table 1), and were consistently greater (p < 0.001) from non-AMP soils (2.76 ± 0.11) compared to AMP soils (2.13 ± 0.08). Additionally, Q₁₀ values increased with soil moisture as follows: PWP (2.25 ± 0.11) < 0.4FC (2.43 ± 0.14) ≤ FC (2.67 ± 0.14) (p < 0.05 for all comparisons).

3.3. Proportion of SOC Mineralized as CO₂

The proportion of SOC mineralized as CO₂ during the incubation period was affected by soil temperature, moisture, and by an interaction of grazing × temperature, and temperature × moisture (Table 1). At 5 °C, the proportion of SOC mineralized as CO₂-C was <0.5%, but at 25 °C reached as high as 2.9% (Figure 2a). Within the AMP soils, SOC mineralization was 3.7 times greater at 25 °C than at 5 °C, while in non-AMP soils this increase was 5.6 times (Figure 2a). Increasing soil moisture and soil temperature both increased the proportion of SOC mineralized (Figure 2b), though high temperature (25 °C) and high moisture (i.e., FC) in combination increased CO₂-C mineralization by as much as 17+ fold relative to soils at low temperature where moisture was below FC (Figure 2b).

3.4. Factors Affecting GHG Fluxes

The use of SEM evaluating CO₂, CH₄, and N₂O responses to the grazing regime, temperature, and moisture conditions during incubation, as well as the associated MBC and EEA levels, revealed marked differences among GHGs, and the sampling times during incubation (Figure 3; Table S2).
Figure 3. Structural equation models showing relationships between temperature, moisture, soil microbial properties, various extracellular enzyme activities, and fluxes of greenhouse gases (a) CO$_2$, (b) CH$_4$ and (c) N$_2$O on either day 1 (top row), day 13 (middle row) and day 102 (bottom row) of the soil incubation. The green arrows depict a positive relationship, while red arrows depict a negative relationship on the response variable. Different thickness of arrows indicates the relative strength of the relationship between variables. Refer to text for enzyme names abbreviated in the figures.
Emissions of CO$_2$ from soil were consistently positively influenced by increasing soil moisture, and even more so by greater soil temperatures (Table S3), throughout all three sampling periods—namely days 1, 13, and 102 (Figure 3a). In contrast, the grazing system had no impact on CO$_2$ emissions, either directly, or indirectly by moderating MBC. Instead, MBC was largely decoupled from soil temperature and moisture (Table S4), with the exception of soil temperature on day 13, at which time higher temperatures reduced MBC (Figure 3a; Table S5). Soil EEAs had a strong positive response to both increasing MBC and higher soil moisture, particularly on day 13, but also on day 102, and to a lesser extent on day 1 (Table S6; Table S7).

Unlike CO$_2$, CH$_4$ fluxes directly declined (i.e., uptake increased) in response to higher soil temperature during the first two sampling periods, and also decreased more within soils subject to AMP rather than non-AMP grazing at these times (Figure 3b). By the end of the incubation period, however, no direct grazing effect remained, while CH$_4$ flux increased at that point with soil temperature and declined with greater moisture (Figure 3b). CH$_4$ fluxes were also closely coupled with MBC, but only via the indirect influence of MBC on EEA. On day 13, CH$_4$ flux increased with greater Cello EEA, and decreased with greater BG EEA (Figure 3b). On day 102, Xylo EEA was the only enzyme that was linked to soil CH$_4$ flux, with higher Xylo EEA associated with decreasing CH$_4$ flux (Figure 3b).

Based on the SEM, fluxes of N$_2$O were not affected by the grazing regime either directly or indirectly, with the lone effect of grazing being an increase in NAG EEA in soil arising from AMP grazing (Figure 3c). The latter, however, did not link further to N$_2$O flux. Overall, significant relationships evident between the grazing systems and associated soil N$_2$O fluxes were limited to a consistent positive effect of soil temperature throughout the incubation period, and a positive effect of soil moisture at the start of the incubation (Figure 3c).

4. Discussion

4.1. Effects of Grazing Systems, Soil Temperature and Moisture on GHG Flux

We purposely conducted this study within an incubation environment to control extraneous sources of environmental variation common under field conditions. Furthermore, by using a paired design in which grasslands were on similar ecosites, we controlled many external physical factors (e.g., soil texture, SOM, etc.), and thereby isolated the influence of inherent differences in soil chemical and biological parameters on soil GHG fluxes [59] generated by the grazing treatments. In our study, by varying only soil temperature and moisture levels during the incubation, we were able to isolate soil-based influences on GHG fluxes, which in turn, were hypothesized to arise due to differences in prior grazing practices.

Our results showed that CH$_4$ oxidation was higher in grasslands subject to AMP grazing rather than non-AMP grazing, and that fluxes of N$_2$O did not vary in relation to grazing, while the flux of CO$_2$ depended on soil temperature. Increases in soil temperature from 5 °C to 25 °C led to 2+ fold increases in N$_2$O flux and 3+ fold increases in CO$_2$ flux, but simultaneous increases in CH$_4$ uptake. A similar pattern occurred with increases in soil moisture: soils at FC had marked increases in CO$_2$ and N$_2$O flux, as well as increased CH$_4$ uptake. Overall, net GHG fluxes strongly paralleled those associated with CO$_2$, which is not surprising given that the latter represented the primary contributor to net GHG emissions (~98%). Collectively, these findings corroborate the notion that mineralization of soil C increases with increasing temperature and moisture [33,60]. The emission of CO$_2$, which is a key indicator of mineralizable C, is well known to be directly influenced by soil temperature and moisture [33,34,61–64]. For example, the low flux of GHGs in soils at PWP compared to other moisture levels shows that dry soils inhibit microbial activity, leading to decreased respiration [65]. An increasing flux of N$_2$O with higher moisture may also reflect increased microbial activity involved in the formation of N$_2$O [65].

While grazing effects were not as marked as soil temperature and moisture, distinct patterns were nevertheless evident in GHGs relative to whether soils originated from AMP- or non-AMP-grazed
grasslands, and these differences were more likely to manifest in soils subject to a higher temperature or elevated moisture. Under warm and moist conditions, soils under AMP grazing had lower emissions of CO$_2$ and increased uptake of CH$_4$ compared to soils from non-AMP grasslands. Soils from AMP systems were generally a better sink for CH$_4$ than soils from non-AMP systems within each temperature level during the entire experimental period, as determined by net GHG values. These results are indicative of a smaller GHG footprint in grasslands subject to AMP grazing and in line with findings of other studies such as CH$_4$ sink capacity of the Northern Great Plains pastures [9] and lower GHG emissions from AMP-grazed than feedlot beef productions [66]. Another field experiment showed that AMP grazing led to a greater CH$_4$ sink capacity compared to moderate and heavy continuous grazing [67]. Although the mechanisms accounting for these differences remain unknown, below, we discuss potential causes based on our lab-incubation study.

Soils from non-AMP grasslands may have more labile C that microbes can rapidly mineralize, particularly at high temperatures, leading to elevated CO$_2$ fluxes. In contrast, AMP soils may contain elevated recalcitrant material from the incorporation of above-ground litter, induced by intensive animal activity (i.e., hoof action) [68], which in turn, could slow down decay [8].

Fluxes of N$_2$O result from nitrification and denitrification processes in the soil [69]. Water content is crucial in these processes as it transports energy required by soil microorganisms to function [70]. A positive relationship between soil moisture and N$_2$O emissions has been reported in an incubation study on mixed grassland soils [61], and is consistent with our current findings. In addition to moisture, soil temperature and aerobic conditions affect N$_2$O flux, with the former two factors explaining almost 90% of the variation [70]. In a warming and defoliation field experiment conducted in the same northern temperate grasslands encompassed by our study, N$_2$O flux increased with increasing temperature and more severe defoliation [7].

Soil compactness, together with organic C and N turnover induced by grazing, determines the abundance and diversity of active methanotrophs, as well as CH$_4$ oxidation in grassland soils [23]. Under normal conditions, N-deposition reduces the CH$_4$ sink capacity of soils [71]. Given the similar physical and chemical soil properties among our paired grasslands, the specific mechanism for the increased CH$_4$ uptake in AMP soils is not clear but may be related to differences in soil microbes. However, our results show that increasing temperature facilitates CH$_4$ uptake across all moisture levels in soils from both grazing systems, confirming the favorable temperature response of methanotrophs [12,72].

Grassland soils are well-documented CH$_4$ sinks due to methanotrophy. The identity of methanotroph communities (Type I or II) responsible for CH$_4$ oxidation depends on soil properties, including SOM and the synthesis of secondary chemicals during the SOM mineralization process, as well as SOC, N cycling, and soil pH [73]. In simulated global change experiments, Type II methanotrophs decreased with increased precipitation and temperature [74]. As our study did not test these factors, further investigation is warranted to determine which type of methanotrophs are specifically involved in the increased CH$_4$ oxidation within AMP soils.

4.2. SOC Mineralization and the Temperature Sensitivity of CO$_2$ Emissions

Our results showed that SOC mineralization as CO$_2$-C was more than 4-fold higher at 25°C compared to 5°C, with a further increase in non-AMP grasslands at high temperatures. We found that $Q_{10}$ increased as moisture increased within incubated soils from both the AMP and non-AMP grasslands. Importantly, $Q_{10}$ was consistently greater in non-AMP soils in comparison to AMP soils, which suggests a greater risk of C loss exists within soils grazed under conventional grazing.

The $Q_{10}$ value of grasslands found here (2.4 ± 0.1) falls within the range of globally observed values (2.5 ± 2.0), and temperature-normalized values (2.0 ± 1.7) for grasslands [59]. It also corroborates earlier findings that values of $Q_{10}$ generally increase with soil moisture levels, where at the same temperature, soils with greater moisture have higher CO$_2$ emissions [64]. Moreover, we found a strong synergistic effect of greater moisture and temperature in elevating $Q_{10}$ values, highlighting
the potential impact of simultaneous changes in both these climatic factors on future grassland soil C storage. While grassland managers are unlikely to seek lower soil moisture due to its fundamental importance in regulating plant (and forage) growth, it does highlight the crucial role that soil water plays in the decomposition of grassland SOM, with implications for various scenarios of global climate change [64,75]. Additionally, it reinforces the importance of adopting grazing management practices that limit soil temperatures, such as the retention of adequate insulating litter [76–78].

There may be several mechanisms accounting for this difference in temperature sensitivity between grazing systems. For example, as mentioned above, relative to non-AMP soils, AMP soils may contain more complex (recalcitrant) compounds [79] due to the so-called “herd effect”, that incorporates coarse standing litter and debris into the surface soil [68]. Should this be the case, the available pool of soil C in AMP soils may contain more structural plant materials such as lignin that are found in lower abundance within roots [80]. Grazing is known to influence nutrient cycling by modifying litter breakdown within species as well as the soil environment for decomposition [27]. Yet another explanation is that the microbial communities in soils between the grazing systems may differ, in part due to the differences in plant chemistry, but also differences in the microenvironment. The non-AMP-grazed areas are often grazed for longer periods of time with less rest, and this could lead to lower levels of insulating litter on the soil surface. Litter, in turn, is widely known to be important for regulating soil temperature [76–78]. As such, non-AMP soils may have had microbial communities better adapted to warmer soil conditions, which in turn could explain why non-AMP soils had greater CO₂ flux at higher soil temperatures. This finding may have implications for the conservation of grassland soil health, including soil organic matter and SOC under future uncertainties associated with variation in growing conditions, including climatic warming.

4.3. Relative Effects of Grazing, Temperature and Moisture on EEA and GHG Fluxes

We used SEM to distinguish between direct effects of grazing on GHG fluxes, and indirect effects that were regulated through the microbial community and the associated levels of EEAs. Overall, the GHGs examined here were more likely to be directly and strongly influenced by soil temperature and moisture, rather than grazing. Contrary to our expectations, grazing had comparatively little impact on soil CO₂ or N₂O fluxes, particularly in comparison to the microclimate conditions. In contrast, soil CH₄ uptake was the lone GHG that demonstrated a strong direct (and positive) response to AMP grazing. This finding suggests that AMP grazing may be altering the soil microbial community in such a way that it leads to increases in the abundance of methanotrophs responsible for CH₄ oxidation. Previous studies have shown that grazing intensity affects the abundance and diversity of active methanotrophs responsible for CH₄ oxidation [23]. Under field conditions, soil CH₄ uptake was found to be influenced by moisture and available substrates [8], and field and incubation experiments showed that soil moisture and labile C and N content are the primary controlling factors for methanotrophy [81,82].

While we hypothesized a linkage would exist between the grazing systems and GHGs via MBC and the measured EEAs, we did not find this association. Instead, CO₂ and CH₄ were independently related to select EEAs, reinforcing the notion that EEAs may be a useful indicator of GHG fluxes [30]. Additionally, the EEAs explored in this study were associated with both MBC, indicating the key role of microbial population size in regulating EEAs [30,83], as well as microclimatic conditions, particularly soil moisture. These results substantiate the conclusion that while the EEAs responsible for C biogeochemical cycling were closely dependent on MBC and soil moisture, soil CO₂ fluxes overall remain more dependent on ambient microclimatic conditions (temperature and moisture) rather than on the prior grazing system.

Finally, we observed divergent effects of the grazing systems over time. Grazing, temperature, and moisture effects were generally more prominent on days 1 and 13 compared to day 102, including in assessing the direct relationship between grazing and CH₄ flux. This may reflect ongoing mineralization of the more labile soil organic matter compounds during the incubation, leading to a general reduction in microbial activity [84], and therefore GHGs. Our results corroborate the resource allocation
theory of microbial enzyme production, which states that soil microbes regulate enzyme production proportionately to the availability of resources such as labile C and N [85].

The lack of a relationship between the grazing systems and either MBC or EEA could be due to our simplified categorical differentiation between grazing systems as either AMP or non-AMP. This separation may be inadequate to test for grazing induced responses, particularly as the non-AMP ranches had more variation in management metrics [86].

As these findings occurred under lab incubation conditions and not in the field where factors such as litter and light levels varied, the most likely reason is a difference in either chemical properties (i.e., SOM composition) or biological properties (i.e., the microbial community). Moreover, this is likely to be the case given that most soil physical and chemical properties did not differ between AMP and non-AMP systems. Under field conditions, many biotic and abiotic factors simultaneously affect C and nutrient cycling, and thus it is imperative to evaluate in situ GHG fluxes from grassland soils under both types of grazing systems to derive results more indicative of fundamental conditions regulating these GHGs. Comparative field studies will provide a better insight into the benefit of AMP grazing in comparison to conventional grazing in terms of GHG dynamics across these northern temperate grasslands.

5. Conclusions

Our results showed that fluxes of different GHGs from grassland soils varied with grazing systems: cumulative CH$_4$ uptake was higher in soils under AMP grazing compared to non-AMP; emissions of N$_2$O were independent of grazing, and grazing interacted with temperature to affect the flux of CO$_2$. The uptake of CH$_4$ and emissions of CO$_2$ and N$_2$O increased with greater moisture levels and soil temperature. Irrespective of grazing systems, MBC had a vital role as an indirect driver on GHG fluxes by influencing the EEA responsible for C and N cycling. Grazing affected CH$_4$ uptake for the first two weeks but, thereafter, the grazing effect became less important and N$_2$O emissions were indirectly influenced by grazing by affecting NAG. We conclude that AMP grazing has the potential to mitigate the effect of a warmer soil on GHG emissions by consuming more CH$_4$ compared to non-AMP-grazed soils. Despite the increased uptake of CH$_4$ in grassland soil under AMP grazing, including soils subject to the higher temperature and moisture conditions, this change was insufficient to offset increases in the other GHGs, particularly of CO$_2$. As a result, environmental conditions favoring high CO$_2$ flux produced the greatest net GHG footprint, and reinforces the importance of maintaining cool soil temperatures within these grasslands, as might occur with the retention of ample litter. This finding has implications for the conservation of grassland soil health, including soil organic matter and SOC, under future uncertainties associated with the variation in growing conditions, including climatic warming in northern temperate grasslands.

Supplementary Materials: The following are available online at http://www.mdpi.com/2073-4395/10/11/1781/s1, Figure S1. Conceptual SEM models for determining effects of grazing systems, soil temperature, and moisture on microbes and resultant fluxes of (a) CO$_2$, (b) CH$_4$, and (c) N$_2$O during the incubation experiment. Table S1: Summary of soil physical and chemical properties for studied grasslands. Table S2. Summary of ANOVA, including the degree of freedom (df), F- and p-values, for the enzyme activities studied during a 102-day incubation. Enzyme activities were analyzed for days 1, 13, and 102 of the incubation period. p-values in bold are ≤0.05. Table S3. Summary of ANOVA, including degrees of freedom (df), F- and p-values, for the enzyme activities studied during a 102-day incubation. Enzyme activities were analyzed for days 1, 13, and 102 of the incubation period. p-values in bold are ≤0.05. Table S4. Summary of ANOVA, including the degree of freedom (df), F- and p-values, for the studied soil microbial biomass carbon (MBC), microbial biomass nitrogen (MBN), and available nitrogen (AN) during a 102-day incubation. Enzyme activities were analyzed for days 1, 13, and 102 of the incubation period. p-values in bold are ≤0.05. Table S5: Fluxes of greenhouse gases (mean ± SE) from incubated soils on selected days of measurement (1, 13, and 102) coinciding with measurements of various extracellular enzyme activities. Table S6: Microbial biomass carbon (MBC), microbial biomass nitrogen (MBN), and available N (mean ± SE) within incubated soils on selected days of measurement (1, 13, and 102) coinciding with measurements of various extracellular enzyme activities. Table S7: Extracellular enzyme activities (mean ± SE μmol g$^{-1}$ h$^{-1}$) within incubated soils on select days of a 102-day incubation.
Author Contributions: Conceptualization and methodology: B.M.S., S.X.C. and D.K.; validation and analyses: B.M.S., E.W.B., C.N.C. and Z.M.; investigation: B.M.S., and D.K.; data curation: B.M.S., D.K. and T.F.D.; writing and editing: B.M.S., E.W.B., C.N.C., S.X.C.; visualization: T.F.D., and Z.M.; resources and supervision, S.X.C., and M.S.B.; funding acquisition and project administration; M.S.B. All authors have read and agreed to the published version of the manuscript.

Funding: Funding for this study was provided by Agriculture and Agri-Food Canada through its Agricultural Greenhouse Gases Program (Project No RES0032548).

Acknowledgments: We acknowledge all participating ranchers for their permission to access their ranches for this project. Richard Teague, Ry Thompson, Steve Apfelbaum, and Jessica Grenke identified the ranches used in this study. We thank Miles Dyck for providing pressure plate extractors for soil moisture determination, and Gleb Kravchensky for assisting with the collection of soil samples and GHG samples during the incubation experiment. Soil texture data were provided by Upama K.C. Laio Sobrinho helped in data analysis in R environment. Finally, we thank Peter Blenis for providing advice on the statistical analysis. We acknowledge anonymous reviewers whose comments helped to improve the earlier version of the manuscript.

Conflicts of Interest: The authors declare no conflict of interest.

Ethical Statement: Researchers conducted the study at the University of Alberta. All pasture management data were collected at the University of Alberta following Human Ethics approval (Human Ethics # Pro00078581).

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