Nitrogen and carbon isotopic dynamics of subarctic soils and plants in southern Yukon Territory and its implications for paleoecological and paleodietary studies

Farnoush Tahmasebi1 *, Fred J. Longstaffe1 *, Grant Zazula2, Bruce Bennett3

1 Department of Earth Sciences, The University of Western Ontario, London, Ontario, Canada, 2 Yukon Palaeontology Program, Department of Tourism and Culture, Government of Yukon, Whitehorse, Yukon Territory, Canada, 3 Yukon Conservation Data Centre, Environment Yukon, Government of Yukon, Whitehorse, Yukon Territory, Canada

* Tahmasebi.farnoush@gmail.com (FT); flongsta@uwo.ca (FL)

Abstract

We examine here the carbon and nitrogen isotopic compositions of bulk soils (8 topsoil and 7 subsoils, including two soil profiles) and five different plant parts of 79 C3 plants from two main functional groups: herbs and shrubs/subshrubs, from 18 different locations in grasslands of southern Yukon Territory, Canada (eastern shoreline of Kluane Lake and Whitehorse area). The Kluane Lake region in particular has been identified previously as an analogue for Late Pleistocene eastern Beringia. All topsoils have higher average total nitrogen δ15N and organic carbon δ13C than plants from the same sites with a positive shift occurring with depth in two soil profiles analyzed. All plants analyzed have an average whole plant δ13C of −27.5 ± 1.2 ‰ and foliar δ13C of −28.0 ± 1.3 ‰, and average whole plant δ15N of −0.3 ± 2.2 ‰ and foliar δ15N of −0.6 ± 2.7 ‰. Plants analyzed here showed relatively smaller variability in δ13C than δ15N. Their average δ13C after suitable corrections for the Suess effect should be suitable as baseline for interpreting diets of Late Pleistocene herbivores that lived in eastern Beringia. Water availability, nitrogen availability, spacial differences and intra-plant variability are important controls on δ15N of herbaceous plants in the study area. The wider range of δ15N, the more numerous factors that affect nitrogen isotopic composition and their likely differences in the past, however, limit use of the modern N isotopic baseline for vegetation in paleodietary models for such ecosystems. That said, the positive correlation between foliar δ15N and N content shown for the modern plants could support use of plant δ15N as an index for plant N content and therefore forage quality. The modern N isotopic baseline cannot be applied directly to the past, but it is prerequisite to future efforts to detect shifts in N cycling and forage quality since the Late Pleistocene through comparison with fossil plants from the same region.
1 Introduction and background information

Stable isotopes of carbon and nitrogen are valuable for studying food webs and tracking transfer of energy and materials through trophic levels [1]. Primary producers in different ecosystems provide varied food sources with distinct carbon ($\delta^{13}C$) and nitrogen ($\delta^{15}N$) isotopic compositions at the base of the food web. The $\delta^{13}C$ and $\delta^{15}N$ of an animal reflect both the isotopic composition of its diet and fractionations during the building of organic tissues. Hence, an isotopic baseline for each ecosystem should be established prior to comparing animals from different regions and times [1]. This is particularly important for paleoecological and paleodietary studies of Late Pleistocene subarctic ecosystems. Such investigations have a special value in the study of whole ecosystem (structure, composition and function) responses to climate change.

Stable isotope data have been widely used in the study of Late Pleistocene subarctic mammals e.g. [2–8]. Determinations of appropriate modern and past local and regional, foodweb isotopic baselines, however, are still underrepresented in the literature. Subarctic ecosystems are not homogeneous. Local and regional ecological mosaics likely existed in the past because of variation in topography, soil moisture, loess deposition, altitude and animal disturbance [9] and are present currently owing to the bioclimatic subzones, patterned ground created by soilfrost processes, different plant communities, and the changes in elevation [10]. The possibility of such microhabitat diversity should be considered when establishing isotopic baselines for these ecosystems.

Soils and plants, which can show an extremely wide range of $\delta^{13}C$ and $\delta^{15}N$, are two main components of all terrestrial ecosystems [11, 12]. A number of studies have addressed plant $\delta^{13}C$ [13, 14] and $\delta^{15}N$ [13, 15–18] variations in Arctic and subarctic ecosystems in North America, Eurasia and Iceland. Wooller et al. [14] conducted the most comprehensive study, analyzing the $\delta^{13}C$ of herbarium modern plants (around 200 taxa) as well as fossil plants from Alaska and Yukon Territory. Schulze et al. [17] studied N isotopic and elemental compositions of different plants in northern Alaska with the aim of investigating their difference in nutrient acquisition. Similar studies have reported $\delta^{15}N$ for various plants from other parts of this ecosystem [15, 16]. The analysis of both $\delta^{13}C$ and $\delta^{15}N$ of plants has also been conducted for Carex from 15 Eurasian Coastal Arctic sites [18] and for lichens and plants from Iceland [13]. While these studies have added greatly to our general isotopic knowledge in these regions, additional local and regional isotopic studies are needed to evaluate the heterogeneity that can exist in these ecosystems.

In the present study, we measured the $\delta^{13}C$ and $\delta^{15}N$ of soils and plants from the south of Yukon Territory (eastern shoreline of Kluane Lake and areas around Whitehorse) (Fig 1). The study’s purpose is to establish a local isotopic baseline for modern foodwebs and to gain a better understanding of main environmental factors affecting C and N isotopic dynamics in this region. The results should also enable future work in which modern and Late Pleistocene isotopic baselines for eastern Beringia are compared, in order to identify environmental factors that affected the baseline. Beringia was a largely ice-free land extending from northwest Canada to northeast Siberia (Fig 1) [19] and was located within the Mammoth Steppe Ecosystem, which was the most extensive biome on Earth during the Last Glacial Maximum [20]. Previous reconstructions of Beringia [9, 19] have shown similarities in main ecosystem contexts (soil and grassland compositions) to the Kluane Lake area [21].

1.1 Study areas: Kluane Lake and Whitehorse, Yukon Territory

The two main regions investigated in this study are located in the southern Yukon Territory: (i) the eastern shoreline of Kluane Lake, and (ii) the Whitehorse area (Fig 1). The first site
consists of grasslands located next to the southeast shore of Kluane Lake, which receives windblown loess from the Slims River delta. The second site consists of grasslands in the Whitehorse valley. Table 1 lists mean air temperature and total precipitation over the last 29 years at these sites as well as these data for 2012 and 2013, when the plant samples were
collected. A few plant samples were also obtained from three sites (S13-13, -14, -15) in the Faro area (Fig 1).

The Kluane Ranges (2000–2800 masl), which are located in the southwest of Yukon Territory, effectively block penetration of Pacific air masses to the Kluane Plateau, resulting in a semiarid continental climate for the Kluane area with cold winters and warm summers [21]. The combination of ice fields at the core of the Kluane Ranges, from which glaciogenic silt and sand are delivered to the Slims River delta, strong winds off of these glaciers, and arid conditions, which amplify evapotranspiration, facilitate continuous transportation and accumulation of loess on the eastern side of Kluane Lake. Such conditions are similar to those reconstructed by Guthrie [22] for eastern Beringia loess formation during the Late Pleistocene.

Loess accumulation in this area occurred during two time periods: Late Pleistocene/early Holocene and Neoglacial [23]. The region still experiences frequent dust storms, particularly during summer months. In most soils from this area, the two loess phases are separated by a reddish-brown paleosol (Fig 2), which is named the “Slims soil” [24].

The vegetation of the Kluane region comprises a mixture of grassland and boreal forest ranging from valley-bottom elevations (781 m) to ~1160 m [25]. The grasslands are composed mainly of *Artemisia-Festuca* communities, which are predominant in drier parts of the area on southwest-facing aspects. The forests consist mainly of white spruce [21].

The Whitehorse valley, like most of Yukon Territory, has a dry, subarctic climate characterized by long and cold winters and short and cool summers. The long-term records (Table 1) show a west to east difference in mean total precipitation, with Whitehorse receiving significantly more precipitation than Kluane Lake. The location of the City of Whitehorse in this valley makes its climate milder than other areas of the Yukon (Table 1). The vegetation is more or less similar to the Kluane Lake area, consisting mainly of boreal forest and grasslands. Shrub communities are also present near the tree line, under a canopy of trees. Grasslands are limited to dry and south-facing slopes, while forests cover many plateaus and valleys [26].

### 1.2 Carbon isotopic composition of soil and terrestrial plants

#### 1.2.1 $\delta^{13}C$ of soil organic carbon

Vegetation following different photosynthesis pathways ($C_3$, $C_4$ and CAM) imparts different $\delta^{13}C$ signatures to soil organic carbon (SOC). This difference is the basis for many studies of past vegetation and climate change [27–30]. Other
processes also affect the δ\(^{13}\)C of SOC. A negative relationship has been observed between soil δ\(^{13}\)C and mean annual precipitation (MAP) in C\(_3\)-dominated ecosystems [28]. This relationship is explained by plant C isotopic response to MAP, which is transmitted to soils [31]. Discrimination against \(^{13}\)C during microbial decomposition also affects soil δ\(^{13}\)COC. This effect depends mostly on the extent of decomposition, which is controlled by climate [32]. Increases in δ\(^{13}\)COC with depth are typically attributed to microbial degradation when <4 ‰ [33, 34], and to C\(_3\)/C\(_4\) vegetation change when >4 ‰ [35].

**1.2.2 Controls on plant δ\(^{13}\)C.** Among vascular plants, C\(_3\) plants are characterized by the lowest δ\(^{13}\)C (–38 to –22 ‰), C\(_4\) plants have higher δ\(^{13}\)C (–21 to –9 ‰) and CAM (Crassulacean Acid Metabolism) plants lie between (–30 to –13 ‰) [36–38]. C\(_3\) plants are overwhelmingly dominant in high latitudes [39, 14], including our study sites (above 60\(^{\circ}\)N).

Other factors affecting plant δ\(^{13}\)C include the δ\(^{13}\)C of source CO\(_2\), relative humidity/water availability, temperature, light intensity and partial pressure of CO\(_2\) (pCO\(_2\)) [31, 36, 37, 40, 41]. Water availability is most important. A negative correlation between MAP and C\(_3\) plant δ\(^{13}\)C [41–44] is attributed to stomatal closure in response to aridity, producing reduced C\(_i\)/C\(_a\), higher water use efficiency and less negative δ\(^{13}\)C [36–38]. Understory growth under closed tree canopies—the “canopy effect”–can lead to more negative plant δ\(^{13}\)C due to fixing of \(^{13}\)C-depleted CO\(_2\) from soil and canopy respiration [45], lower light intensity and higher pCO\(_2\) [45]. No samples in the present study, however, grew under such conditions.

In C\(_3\) plants, photosynthesizing tissues (e.g. leaf) have more negative δ\(^{13}\)C than heterotrophic tissues (e.g. stem, root, inflorescence) [46]. Causes include different macromolecular tissue compositions, growing stage variations in photosynthetic discrimination against \(^{13}\)C, and different contributions of day versus night sucrose with different δ\(^{13}\)C to different tissues [46].

---

**Fig 2. Soil profiles at Kluane Lake and their corresponding depth profiles for δ\(^{13}\)COC, OC and δ\(^{15}\)N.**
(a) profile S13-6, and (b) profile S13-8. In both, data points provide average values for bulk analysis of each soil interval (indicated by the vertical lines). The reddish-brown layer is called the “Slims soil”, and separates Neoglacial loess deposits from underlying Late Pleistocene/early Holocene deposits (photographic credit: Tessa Plint).

https://doi.org/10.1371/journal.pone.0183016.g002
1.3 Nitrogen isotopic composition of soil and terrestrial plants

1.3.1 δ15N of source N. Nitrate (NO$\textsubscript{3}^-$), ammonium (NH$_4^+$) and dissolved organic compounds (e.g. simple proteins, amino acids and amino sugars) [47–49] are the most common forms of soil N taken up by plants. The δ$^{15}$N varies among these sources because of different biochemical reactions in soil (N mineralization, nitrification, denitrification and volatilization) [50, 51]. Bulk soil δ$^{15}$N, however, is not always a good representation of bioavailable N [50]. Adding to the complexity is the δ$^{15}$N gradient with soil depth [49, 52]. Increasing bulk soil δ$^{15}$N with soil depth [53–55] has been attributed to fresh litter input to topsoil, coupled with accumulation of $^{15}$N-enriched, decomposing organic matter with depth [54]. Plants have diverse abilities to acquire N depending on their rooting depth and phenology [55], life form (trees, shrubs and herbs) [17], and preference for different forms of N [16, 56] at different times of year [57].

Mycorrhizal fungi associations also affect the δ$^{15}$N of source N for plants. Plants are likely more reliant on mycorrhizal fungi for N acquisition under conditions of low N availability [11], which commonly is the case in Arctic and subarctic ecosystems [58]. Arbuscular mycorrhizal-, ectomycorrhizal- and ericoid mycorrhizal-associated plants are depleted of $^15$N by 2 ‰, 3.2 ‰ and 5.9 ‰, respectively, relative to non-mycorrhizal plants [11, 59–61]. Herbivory (grazing, trampling, excretory products, soil disturbance) also can change soil and plant δ$^{15}$N. Herbivory affects N dynamics by changing N availability [62], altering the rate of soil N processes [62], and modifying litter quality and plant composition [63, 64]. No systematic pattern of N-isotope effects, however, has emerged from herbivory [65–71].

1.3.2 Controls on plant δ$^{15}$N. While nitrogen in plants mainly originates from soil, plant δ$^{15}$N varies from total soil N [12]. This reflects the range of bioavailable versus non-bioavailable N compounds in bulk soil N, isotopic fractionation during N uptake by plants, and biological processes during N assimilation [12, 50].

1.3.2.1 Nitrogen Isotopic Fractionation during N Uptake and Assimilation. The size of nitrogen isotope fractionation (ε) during plant uptake of NO$_3^-$ and NH$_4^+$ is controlled by two factors: (i) external N concentration [72, 73], and (ii) efflux of $^{15}$N-enriched inorganic N and/or $^{15}$N-depleted organic N from roots after N uptake [72, 74]. Values of ε are small during plant uptake under low concentrations of NO$_3^-$ and/or NH$_4^+$ (~0.5 mol m$^{-3}$); ε increases at higher concentrations [75, 76]. Discrimination against $^{15}$N during plant uptake is negligible under most natural conditions [72, 77]. Enzymatic reactions during N assimilation typically produce large fractionations; ε of +15 ‰ and +17 ‰, respectively, have been reported for nitrate reductase and glutamine synthetase, two main enzymes involved in N assimilation [72].

1.3.2.2 Intra-plant Variation in δ$^{15}$N. Intra-plant variation in δ$^{15}$N can arise from: (i) variation in plant nitrogen sources as different organs form and expand, (ii) different patterns of N assimilation with either NO$_3^-$ or NH$_4^+$ as the primary N source, (iii) reallocation and transportation of N macromolecules between sink and source organs, and (iv) organ-specific efflux of N [72]. Leaves normally have higher δ$^{15}$N than other organs, particularly roots [72, 78], although there are some exceptions [79]. When NO$_3^-$ is the sole N source, significant intra-plant variation occurs, with leaves having much higher δ$^{15}$N than roots. This likely reflects different patterns of NO$_3^-$ vs. NH$_4^+$ assimilation. Whereas NH$_4^+$ is assimilated immediately after root uptake, some NO$_3^-$ is assimilated in roots while the remaining, $^{15}$N-enriched NO$_3^-$ is transported to shoots for N assimilation in leaves [72].

Enzymatic reactions involved in reallocation of N also can produce molecules with lower δ$^{15}$N than the original source and thus cause intra-plant δ$^{15}$N variation [72]. Likewise, loss of NH$_3$ through plant leaves and efflux of organic N from roots can enrich these organs in $^{15}$N [50, 72].
1.3.2.3 Environmental Factors and Plant $\delta^{15}N$. A decrease in soil and plant $\delta^{15}N$ generally accompanies increasing MAP and decreasing mean annual temperature (MAT) [11, 80] for MAT $\geq -0.5^\circ C$. This pattern may be related to changes in the rate and nature of soil and plant N cycling and/or dependence on mycorrhizal association [11]. Changes in amount of rainfall and soil water availability can affect the openness of the N cycle [81]. A more open N cycle in drier sites probably reflects greater N availability because of lower plant N demand; this can stimulate NH$_4^+$ volatilization, leading to higher soil and plant $\delta^{15}N$ [49, 81]. Lower N availability [82] and greater plant reliance on mycorrhizal association for N acquisition [11] can contribute to lower soil and plant $\delta^{15}N$ in wetter ecosystems. In short, N cycling in ecosystems is highly responsive to climatic factors, and the associated changes in plant $\delta^{15}N$ can be traced from primary producers to consumers [6, 83].

2 Materials and methods

2.1 Sample collection and preparation

A total of 79 terrestrial plant samples and 15 soil samples (8 topsoil and 7 subsoil including two soil profiles) were collected during September and August 2012 and 2013 (Fig 1) with permission of the Government of Yukon and agreement of Yukon First Nations (Licenses No. 13-52S&E and No. 14-46S&E). Sample collection and field work did not involve any endangered or protected species. Plant and soil samples were placed in woven poly bags and plastic bags, respectively. At sites S13-8 and S13-10, several topsoil samples were collected in response to observed differences in soil texture and topography (shallow vs steep slopes).

All plant samples were air-dried and separated into different plant parts. These plant tissues were washed with distilled water (DW) and dried at 90°C overnight. The dried plant materials were then ground to a very fine powder using a Crescent Wig-L-Bug and stored in small, sealed glass vials for N and C elemental and isotopic measurements.

All soil samples were air-dried, sieved (<2 mm), ground gently using a metal mortar and pestle, and then stored in plastic containers. Analysis of soil samples for physical and chemical properties (mineral fraction, OM fraction, pH, mineralogy) followed standard methods (see Supporting Information: Section A in S1 Text).

Two methods, (i) acid fumigation [84], and (ii) acid rinsing [85], were used to remove carbonates from soil samples prior to elemental and isotopic analyses of OC. Untreated soil samples were used to determine total nitrogen (TN), total carbon (TC), and $\delta^{15}N$ [84].

2.2 Elemental analysis

Procedures used for elemental analysis are described in Section B of S1 Text.

2.3 Stable isotope analyses

All C and N isotopic results are presented using $\delta$-notation [86]:

$$\delta^{13}C \text{ or } \delta^{15}N (\text{‰}) = \left( \frac{R_{sa}}{R_{std}} - 1 \right) \times 1000$$

where $R_{sa}$ and $R_{std}$ denote $^{13}C/^{12}C$ or $^{15}N/^{14}N$ of the sample and standard for $\delta^{13}C$ and $\delta^{15}N$, respectively. The $\delta$-values of all samples were calibrated to VPDB (carbon) and AIR (nitrogen) using USGS40 and USGS41 [87, 88].

The $\delta^{13}C$ and $\delta^{15}N$ of plant samples, soil OC (after carbonate removal), and soil TN were measured by dry combustion using an EA (Costech Analytical Technologies, Valencia, CA, USA) coupled in continuous flow mode to either a Thermo Scientific Delta$^{+}$PLUS XL or a Thermo Scientific Delta V$^{+}$PLUS IRMS (Thermo Scientific Bremen, Germany). Because of the
low N content of the plant samples, nitrogen isotopic analysis was performed in a separate analytical session from that of carbon; CO₂ generated in the latter sessions was scrubbed from samples using a Carbo-Sorb trap on the EA. Separate analytical sessions were also used to obtain δ¹⁵N for soil TN, using un-acidified samples.

Accuracy and precision of the isotopic analyses were monitored using the laboratory keratin and IAEA-CH-6 (sucrose) standards. The average δ¹³C obtained for keratin was –24.05 ± 0.09 ‰ (n = 96), which compares well with its accepted value of –24.04 ‰. The average δ¹³C obtained for IAEA-CH-6 was –10.46 ± 0.10 ‰ (n = 33), which compares well with its accepted value of –10.45 ± 0.03 ‰ [87]. Reproducibility for sample duplicates was ± 0.10 ‰ for δ¹³C (n = 39). The average δ¹⁵N of keratin was +6.38 ± 0.21 ‰ (n = 108), which compares well with its accepted value of +6.36 ‰. Reproducibility for sample duplicates was ± 0.12 ‰ for δ¹⁵N (n = 27).

2.4 Statistical analysis

All plant samples were categorized into two main functional groups: herbs (including annual and perennial grasses, forbs and sedges), and shrub/subshrubs. An independent-sample t-test was used to test for differences in the C and N isotopic and elemental compositions between: (i) different plant functional groups, and (ii) samples from two sampling years (2012 and 2013). Comparisons of C and N isotopic and elemental compositions for (i) different plant parts, and (ii) plants from different sampling sites were performed using one-way ANOVA followed by means comparison using either Tukey’s HSD test, if variance was homogeneous, or Dunnett’sT3 test, if variance was not homogeneous. Assessments of correlation between (i) plant δ¹³C and δ¹⁵N, (ii) plant N content and δ¹⁵N, and (iii) soil δ¹³C and δ¹⁵N in soil profiles were performed using Pearson correlation coefficient. All statistical analyses were performed in SPSS 20.

3 Results

3.1 Soils

General information for each site sampled is presented in Table 2.

Seven topsoils, two soil profiles and one topsoil sample of the loess source area (Slims River delta) were analyzed for basic physical and chemical properties (see Supporting Information: Section C and Tables A and B in S1 Text), and OC and TN isotopic compositions (Table 3).

All topsoils, except for S13-9, are dominated by silt (avg. 48.9 ± 16.5 wt. %, all ± errors reported hereafter are one standard deviation (SD)). The mineralogy of most soil samples is similar to that of Slims River deltaic sediment, which is representative of the sources of wind-blown deposits in the area (see Supporting Information: Table B in S1 Text).

The total nitrogen isotopic compositions (δ¹⁵Nₜ opinión) of the topsoils range from +2.1 to +5.5 ‰. The δ¹³Cₑₑₑₜ意见 results derived from both types of pretreatment to remove carbonate are very similar except for samples S13-9 and Slims River for which acid fumigation showed more efficacy (Table 3). Accordingly, only results produced using acid fumigation are considered further. The range of δ¹³Cₑₑₑₜ意見 (–25.2 to –24.5 ‰) obtained for topsoils is much smaller than measured for δ¹⁵Nₜ意見, and is characteristic of C₃ vegetation. Slims River sediment, by comparison, has δ¹⁵Nₜ意見 and δ¹³Cₑₑₑₜ意見 of +1.6 ‰ and –20.6 ‰, respectively.

The δ¹³Cₑₑₑₜ意见 of profile S13-6 increases with depth from –24.5 ‰ for topsoil to –22.5 ‰ in the subsoil (Fig 2a). The change in pH with depth in profile S13-6 strongly correlates with inorganic carbon (IC) content, which was calculated by subtracting OC from TC (r = 0.960, p <0.01). The δ¹⁵Nₜ意見 in profile S13-6 also show a positive shift from +4.3 to +6.7 ‰ with increasing depth from topsoil to 60 cm, but then decreases to +4.1 ‰ between 60–70 cm
Table 2. Environmental data for sampling sites and number of soils and plants sampled.

| Site ID | Site Name                        | Latitude | Longitude | Altitude (masl) | Topography                     | Plant Samples | Soil Samples |
|---------|----------------------------------|----------|-----------|-----------------|--------------------------------|---------------|--------------|
| S12-1   | Chinook Lane, Whitehorse         | 60.5938  | -134.8949 | 728             | -                              | 2             | -            |
| S12-2   | Riverdale, Whitehorse            | 60.7056  | -135.0343 | 646             | Top of a steep slope           | 7             | -            |
| S12-3   | Carcross Road                    | 60.6213  | -135.0169 | 782             | Steep slope                    | 5             | -            |
| S12-4   | Schwatka Lake, Yukon River       | 60.6724  | -135.0250 | 678             | -                              | 3             | -            |
| S12-5   | Miles River Canyon, Yukon River  | 60.6614  | -135.0281 | 684             | Flat, adjust to a highway      | 7             | -            |
| S12-6   | Destruction Bay                  | 61.4836  | -138.3666 | 844             | Flat                           | 7             | -            |
| S13-2   | Cultus Bay, Kluane Lake          | 61.3850  | -138.5220 | 783             | South facing slope (~35–40˚)   | 8             | -            |
| S13-3   | Kluane Lake                      | 61.2581  | -138.6161 | 839             | South facing slope (~30–40˚)   | 11            | -            |
| S13-4   | Kluane Lake                      | 61.2461  | -138.5650 | 831             | Flat                           | 4             | -            |
| S13-5   | Research Centre, Kluane Lake     | 61.1836  | -138.4008 | 781             | Flat                           | 1             | -            |
| S13-6   | Kluane Lake                      | 61.3530  | -138.6178 | 838             | Road cut with SW facing slope (~25˚) | 8             | SPa (1)      |
| S13-7   | Kluane Lake                      | 61.2530  | -138.6019 | 829             | Road cut with NE facing slope (~20˚) | 6             | TSb (1)      |
| S13-8   | Kluane Lake                      | 61.3092  | -138.4458 | 821             | Flat                           | 2             | SP (1), TS (2)|
| S13-9   | Kluane Lake                      | 61.3413  | -138.6419 | 813             | Slope (~10˚)                   | -             | TS (1)       |
| S13-10  | Kluane Lake                      | 61.3039  | -138.6611 | 805             | Top & bottom of a steep slope  | 1             | TS (2)       |
| S13-11  | Mount Decoli, Kluane Lake        | 61.0514  | -137.9867 | 1138            | Steep slope                    | 4             | -            |
| S13-13  | Robert Campbell Highway, Faro    | 62.1811  | -133.7603 | 748             | -                              | 1             | -            |
| S13-14  | North Canol Road, Faro           | 61.9972  | -132.3792 | 683             | -                              | 1             | -            |
| S13-15  | Faro                             | 62.3799  | -133.3907 | 1651            | -                              | 1             | -            |
|         | Slims River                      | 61.0016  | -138.5104 | -                | Flat                           | -             | TS (1)       |

a SP: soil profile  
b TS: topsoil

https://doi.org/10.1371/journal.pone.0183016.t002

Table 3. Isotopic compositions of soil TN and OC.

| Sample ID | δ¹⁵N_{TN} (‰, AIR) | δ¹³C_{OC} (‰, VPDB) | Acid-fumigated | Acid-rinsed |
|-----------|--------------------|---------------------|----------------|-------------|
| S13-6 (0–20) | +4.3               | -24.5               | -24.7          |
| S13-6 (20–30) | +6.9               | -24.3               | -24.1          |
| S13-6 (30–40) | +7.2               | -23.8               | -23.7          |
| S13-6 (40–45) | +7.0               | -23.7               | -23.8          |
| S13-6 (45–60) | +6.7               | -23.8               | -23.6          |
| S13-6 (60–70) | +4.1               | -22.5               | -23.0          |
| S13-7 (0–10)  | +4.0               | -24.9               | -24.8          |
| S13-8-1 (0–10) | +4.6               | -25.0               | -24.4          |
| S13-8-1 (10–20) | +4.2               | -24.9               | -25.0          |
| S13-8-1 (20–60) | +5.3               | -24.4               | -24.5          |
| S13-8-2 (0–10) | +4.8               | -24.5               | -24.4          |
| S13-9 (0–10)  | +2.1               | -25.1               | -23.4          |
| S13-10-1 (0–10) | +5.5               | -24.5               | -24.5          |
| S13-10-2 (0–10) | +2.5               | -25.2               | -25.1          |
| Slims River   | +1.6               | -20.6               | -7.5           |

https://doi.org/10.1371/journal.pone.0183016.t003
(Fig 2a). A positive, albeit smaller, shift in both δ¹³C₀₂ (−25.0 to −24.4 ‰) and δ¹⁵Nᵣₑₐₓ (−4.6 to +5.3 ‰) with increasing depth is also observed for the second soil profile (S13-8) (Fig 2b).

3.2 Plants

3.2.1 Isotopic compositions. A total of 31 (September 2012) and 48 (August 2013) C₄ plants representing 15 species (Artemisia frigida, Betula glandulosa, Bromus pumpeilannus, Carex filifolia, Calamagrostis purpurascens, Elymus spicatus, Elymus trachycaulus, Festuca altaica, Linum lewisii, Lepidium ramosissimum, Plantago canescens, Poa glauca, Penstemon gormanii, Rubus idaeus, Salix arctica) were sampled from 18 sites distributed for the most part just east of Kluane Lake, but also including some locations near the City of Whitehorse and in the Faro area, Yukon Territory (Fig 1, Table 2). The samples represent two main functional groups: herbs (including both annual and perennial grasses, forbs and sedges) (n = 66) and shrub/subshrubs (n = 13) [89]. Subshrubs are smaller than shrubs, but still have a woody base and bushy shape; their soft stems die back during cold seasons.

The δ¹³C and δ¹⁵N of leaves (L) and stems (S) were measured for all samples, with additional plant parts analyzed as available (70 root crowns (RC), 68 fine roots (FR), 73 inflorescences (I)) (see Supporting Information: S1 Table). In this study, the root crown is considered as the top part of the root system in herbs and subshrubs. Subshrubs and perennial grasses regrow each spring from buds produced by the root crown. In the herbs, the major vascular changes required for formation of new stems in spring occur in the root crown [90].

Mean δ¹³C₀₂ for all sampled plants ranges from −30.9 to −25.5 ‰, with a mean of −28.0 ± 1.3 ‰ (n = 79). Foliar δ¹⁵N ranges from −8.3 to +7.0 ‰, with a mean of −0.6 ± 2.7 ‰ (n = 77). The foliar δ¹⁵N of two samples (A. frigida, Site: S13-2, year: 2013), which are extremely high (+23.4 and +18.7 ‰), have been excluded from this range and subsequently reported averages. Intense volatilization (ε = 40–60 ‰) at the soil surface resulting from urine/dung fertilization by herbivores at this spot could have caused such extreme ¹⁵N-enrichment [91].

No statistically significant differences in foliar δ¹³C and δ¹⁵N are observed between the two functional groups (δ¹³C: t(14.08) = 1.30, p > 0.05; δ¹⁵N: t(75) = 0.81, p > 0.05), although shrubs and subshrubs have slightly higher mean foliar δ¹⁵N than herbs (+0.1 ‰, n = 11) vs. (−0.6 ‰, n = 66). Within a given plant, there is a clear pattern of lower foliar δ¹³C relative to all other plant parts, as expressed by the difference in plant part vs. leaf δ¹³C (Δ¹³Cplant_part-leaf) (see Supporting Information: S1 Fig); Average values of all individual samples are as follow: Δ¹³Cᵣᵣ₋ₗ = 0.5 ± 0.8 ‰, Δ¹³Cᵢ₋ₗ = 0.6 ± 0.8 ‰, Δ¹³Cₛ₋ₗ = 0.9 ± 1.1 ‰, Δ¹³Cᵢ₋ₛ = 0.6 ± 1.4 ‰. In contrast, no clear pattern of intra-plant variation is observed for δ¹⁵N (see Supporting Information: S2 Fig); Average values of all individual samples are as follow: Δ¹⁵Nᵣᵣ₋ₗ = 0.2 ± 2.5 ‰, Δ¹⁵Nᵢ₋ₗ = −0.1 ± 2.9 ‰, Δ¹⁵Nₛ₋ₗ = 0.1 ± 1.6 ‰, Δ¹⁵Nᵢ₋ₛ = 0.3 ± 1.8 ‰. The range of intra-plant variation in δ¹³C is much smaller than measured for δ¹⁵N (~4 vs. 12 ‰).

Mean δ¹³C and δ¹⁵N for different plant tissues for all plants are illustrated in Fig 3. A one-way ANOVA comparing different plant parts of all plants (F(4,364) = 5.57, p < 0.01) confirms a significant intra-plant variations in δ¹³C with foliar δ¹³C significantly lower than that of fine roots, stems and inflorescences (Table 4). Below-ground plant parts (RCs and FRs) have higher mean δ¹⁵N than above-ground parts (L, S and I), but the difference is not significant (F(4,355) = 0.73, p > 0.05). There is a weak, but statistically significant, positive correlation between foliar δ¹⁵N and δ¹³C for all plant samples from all sites (Fig 4). An independent-samples t-test comparing mean 2012 and 2013 foliar δ¹³C and δ¹⁵N shows no significant difference between the two years (δ¹³C: t(77) = 0.52, p > 0.05 and δ¹⁵N: t(75) = 0.49, p > 0.05). A one-way ANOVA comparison of foliar δ¹³C and δ¹⁵N among sites with ≥5 samples shows a significant difference in δ¹³C₂₀¹₂ and δ¹⁵N₂₀¹₃ between different sites (Δ¹³C₂₀¹₂: F(3, 22) = 5.55, p < 0.01
and δ^{15}N_{2013}: F(3, 27) = 7.79, p < 0.01). The results of ANOVA post-hoc test (Tukey’s HSD/Dunnett’s T3) for these comparisons are presented in Table 5. There are statistically significant differences in foliar δ^{13}C between sites S12-5 and S12-2, and sites S12-5 and S12-6 in 2012, with site S12-5 having a lower foliar mean value (–28.7 ‰) than sites S12-2 (–26.9 ‰) and

![Graph showing δ^{13}C and δ^{15}N of all plant parts analyzed, including fine root, root crown, leaf, stem and inflorescence. Below-ground plant parts have higher mean δ^{15}N than above-ground parts, and foliar δ^{13}C is lower than that of fine roots, stems and inflorescences, but only the latter is statistically significant (see text for details).](https://doi.org/10.1371/journal.pone.0183016.g003)

Table 4. ANOVA post-hoc tests (Dunnett’s T3) results for δ^{13}C differences among plant tissues.

| Plant Part | FR^a | I^b | L^c | RC^d |
|------------|------|-----|-----|------|
| S^e        | 0.951| 0.970| 0.001| 0.654|
| RC         | 0.997| 1.000| 0.048| -    |
| L          | 0.002| 0.024| -    | -    |
| I          | 1.000| -    | -    | -    |
| FR         | -    | -    | -    | -    |

^a Fine root, ^b Inflorescence, ^c Leaf, ^d Root crown, ^e Stem.

Values in boldface font are statistically significant (p ≤ 0.05).

https://doi.org/10.1371/journal.pone.0183016.t004
In 2013, there are statistically significant differences in foliar δ^{15}N between sites S13-6 and S13-7, sites S13-6 and S13-3 and sites S13-6 and S13-2, with site S13-6 having a higher foliar mean values (+1.4 ‰) than sites S13-7 (−2.0 ‰), S13-3 (−1.3 ‰) and S13-2 (−0.74 ‰).

3.2.2 Carbon and nitrogen contents. Foliar N contents range from 0.3 to 4.0 wt. %, with an average of 1.1 ± 0.8 wt. % (n = 79). Foliar C contents range from 34.7 to 48.0 wt. %, with an average of 41.3 ± 2.4 wt. % (n = 79) (Supporting Information: S2 Table). Foliar atomic C/N ranges from 13.3 to 156.7, and averages 59.8 ± 30.9 (Supporting Information: S2 Table). In both 2012 and 2013, root crowns have the highest N content (avg. 1.3 ± 0.3 wt. %, n = 70), and stems the lowest (avg. 0.5 ± 0.5 wt. %, n = 79) (Fig 5). A one-way ANOVA comparison of C and N contents between different plant parts shows a significant difference (C: F(4, 364) = 16.28, p < 0.01 and N: F(4, 364) = 33.77, p < 0.01); Table 6 summarizes the results of ANOVA post-hoc tests (Dunnett’s T3) between different plants parts.

Table 5. ANOVA post-hoc tests (Tukey’s HSD) for foliar isotopic differences between sites (plants sampled > 5).

| Sites (2012) | δ^{13}C (‰, VPDB) | Sites (2013) | δ^{15}N (‰, AIR) |
|--------------|------------------|--------------|------------------|
| S12-6        | 0.855            | S13-7        | 0.471            |
| S12-2        | 0.004            | S13-6        | 0.050            |
| S12-3        | 0.892            | S13-3        | 0.854            |
| S12-5        | 0.027            | S13-2        | 0.821            |
|              | -                | S13-3        | 0.002            |
|              | -                | S13-6        | 0.001            |

Values in boldface are statistically significant (p < 0.05).
On average, foliar C and N contents of shrubs and subshrubs (avg. C (wt. %) = 44.5, avg. N (wt. %) = 2.1) are significantly higher than herbs (avg. C (wt. %) = 40.7, avg. N (wt. %) = 0.9) (Independent-samples t-test: C: t(77) = 6.46, p < 0.01; N: t(13.47) = 4.13, p < 0.01). A statistically significant positive correlation is observed between foliar $\delta^{15}$N and N (wt. %) contents among all plants analyzed ($r = 0.389$, p < 0.01).

Fig 5. Average N and C contents of plant parts according to sampling year. In both 2012 and 2013, root crowns have the highest N content whereas stems have the lowest (see text); likewise, during both years, carbon contents are highest in stems, and lowest in leaves.

https://doi.org/10.1371/journal.pone.0183016.g005

Table 6. ANOVA post-hoc tests (Dunnett’s T3) for differences in C and N contents between different plant parts.

| Plant Part | C (wt. %) | N (wt. %) |
|------------|-----------|-----------|
|            | FR$^a$    | I$^b$     | L$^c$     | RC$^d$    | FR | I | L | RC |
| S$^a$      | 0.000     | 0.000     | 0.000     | 0.000     | 0.000 | 0.002 | 0.000 | 0.000 |
| RC         | 1.000     | 0.943     | 0.551     | -         | 0.000 | 0.000 | 0.209 | - |
| L          | 0.111     | 0.001     | -         | -         | 0.263 | 0.004 | -   | -   |
| I          | 1.000     | -         | -         | -         | 0.043 | -   | -   | -   |

$^a$ Fine root,
$^b$ Inflorescence,
$^c$ Leaf,
$^d$ Root crown,
$^e$ Stem.

Values in boldface font are statistically significant (p < 0.05).

https://doi.org/10.1371/journal.pone.0183016.t006

On average, foliar C and N contents of shrubs and subshrubs (avg. C (wt. %) = 44.5, avg. N (wt. %) = 2.1) are significantly higher than herbs (avg. C (wt. %) = 40.7, avg. N (wt. %) = 0.9) (Independent-samples t-test: C: t(77) = 6.46, p < 0.01; N: t(13.47) = 4.13, p < 0.01). A statistically significant positive correlation is observed between foliar $\delta^{15}$N and N (wt. %) contents among all plants analyzed ($r = 0.389$, p < 0.01).
4 Discussion

4.1 Kluane Lake Soils

All topsoil samples examined along the eastern shoreline of Kluane Lake are rich in silty eolian sediment that likely originated from the Slims River delta [21, 24] except for S13-9, which contains a larger abundance of sand (Supporting Information: Table A in S1 Text). Its higher sand content likely reflects a larger contribution of underlying sandy glaciofluvial deposits resulting from bioturbation or post-fire redistribution [24]. The soil mineralogy is similar to the Slims River sediment, consistent with an eolian source (Supporting Information: Table B in S1 Text).

The average $\delta^{13}C_{OC}$ (−24.8 ± 0.3 ‰) of all topsoils is typical of soil organic matter generated by C$_3$ vegetation, except for the higher $\delta^{13}C_{OC}$ (−20.6 ‰) of the Slims River deltaic sediment. This site (Fig 6), which is one of the two main outlets of the Kaskawulsh glacier [92], is subject to flooding and seasonal and diurnal fluctuations in the water flow, and supports algae and macrophytes growth at low flow stages. Aquatic plants in Yukon Territory have a wide range of foliar $\delta^{13}C_{OC}$ (−41 to −15 ‰), particularly for submerged macrophytes [93]. Less negative $\delta^{13}C_{OC}$ (−16 to −13 ‰) for aquatic plants and algae have also been reported for Arctic continental shelf sediments [94]. Contributions of CAM or C$_4$ plants at modern, northern high-latitude sites, such as this one, are unlikely [39]. Incomplete removal of carbonate can also be ruled out based on the acid fumigation results (Table 3).

In general, the topsoil samples have higher average $\delta^{15}N_{TN}$ and $\delta^{13}C_{OC}$ than plants from the same sites (Fig 7), which is typical of many other terrestrial ecosystems [16, 71]. This can be explained by the general enrichment in $^{13}C$ and $^{15}N$ of plant tissues in soil during organic degradation [12, 32]. The larger isotopic variation among the foliar isotopic compositions relative to topsoil samples likely reflects differences in plant sample size, number of plant species sampled, and the age of plants at different sites. In particular, enzymatic variability among different plant species and different patterns of N acquisition among co-occurring species [72] serves to increase the range of foliar $\delta^{15}N$ among sites and between plants within sites and their topsoil. In contrast, the continuous activities of soil decomposers (mainly fungi and bacteria), the strong role of soil minerals in stabilizing soil chemistry [95] and the open N dynamics of soil input and output reactions reduce N isotopic variations in topsoil from site to site within the same general region.

The increase in $\delta^{13}C_{OC}$ with depth in two soil profiles (Fig 2a and 2b) is in agreement with earlier results for arctic ecosystems [27, 96]. This pattern likely reflects the input of fresh litter with lower $\delta^{13}C$ at the soil surface and accumulation of decomposed and hence more $^{13}C$-rich OM at greater depths [27, 33]. This interpretation is consistent with the strong negative correlation between $\delta^{13}C_{OC}$ and OC content ($r = −0.950, p < 0.05$) and the decrease in OC content with depth (Fig 2a and 2b).

The change in $\delta^{15}N_{TN}$ with depth observed in both soil profiles (Fig 2a and 2b) has been reported previously for such soils from Siberia [27]. The negative shift in $\delta^{15}N_{TN}$ below 60 cm in profile S13-6 may arise from the very small amount of OM (2.9 wt. %) and TN (0.1 wt. %) at this depth, in which case the contribution of inorganic nitrogen to the $\delta^{15}N_{TN}$ signal may be more important at this depth. In soils, organic N typically has higher $\delta^{15}N$ (+5 to +7 ‰) than inorganic forms (NO$_3$- and NH$_4^+$) (−2 to +5 ‰) [12, 56].

Values of $\delta^{15}N_{TN}$ are positive in all topsoil samples, which is typical of alpine and tundra ecosystems [27, 53]. Soil $\delta^{15}N_{TN} > 0$ ‰ point to inputs with compositions higher than those produced by fixation of nitrogen from air and/or N-loss processes that leave the soil N pool enriched in $^{15}N$ [97]. Two samples (S13-9, S13-10-2) sit at the lowest end of the N isotopic range (+2.1 to +5.5 ‰). S13-9 has the highest sand and lowest clay and silt contents of the soils
Fig 6. (a) Areal view of the Slims River delta; (b) Slims River vegetation in August. (photographic credits: Fred Longstaffe and Tessa Plint).

https://doi.org/10.1371/journal.pone.0183016.g006
examined in this study. The content, structure and function of OM associated with mineral particles generally vary with grain size. Sand-sized particles are typically associated with less humified and lower abundances of OM than silt and clay [95]. Several studies of soils underlying grasslands or forest have reported higher $\delta^{15}N_{\text{TN}}$ for clay-sized fractions (~+9 to +12 ‰), which generally have a higher content of stable, humified OM than silt (+5 to +9 ‰) and sand (+2 to +7 ‰) particles [98–101].

Topography may explain the low $\delta^{15}N_{\text{TN}}$ (+2.5 ‰) of sample S13-10-2 at site S13-10. Sample S13-10-2 was collected from top of a steep slope, which made it more susceptible to erosional disturbance, while sample S13-10-1 ($\delta^{15}N_{\text{TN}} = +5.5$ ‰) (from the same site) was collected at the bottom of the slope from a flat and more stable location. Steeply sloping soils can have $\delta^{15}N$ close to atmospheric inputs because of continuous soil removal and soil organic matter rejuvenation, which maintains the soil’s N status far from steady state [80].

4.2 Plant C and N isotopic and elemental compositions

The range of $\delta^{13}C$ (~−32.5 to −23.5 ‰) measured for all plant parts in this study (Fig 8) is typical of C3 vegetation, which dominates high latitude ecosystems [39]. Variation in environmental factors (slope aspect, light, water availability and topography) even within small microhabitats may cause this large spread in $\delta^{13}C$. Wooller et al. [14] reported a very similar range of foliar $\delta^{13}C$ for sedges and grasses from Alaska and Yukon Territory.

The plants analyzed here show great variation in $\delta^{15}N$, spanning ~ 40 ‰ (Fig 8). Variation in $\delta^{15}N$ of vegetation tends to be more pronounced in N-limited ecosystems, which is typical
of Arctic and subarctic regions, and points to utilization of different soil N resources by plants depending on their life forms, type of mycorrhizal association and rooting depth and morphology [16]. Coexisting plant species are known to partition N resources with different δ¹⁵N in these ecosystems [16, 17, 56, 102]. For example, the grass *Calamagrostis canadensis* (δ¹⁵N = +0.9 ‰) in Alaska acquires N from deeper soil horizons, while the evergreens *Picea glauca* and *Picea mariana* (−7.7 ‰) likely utilize ammonium or organic N from fresh litter [17].

There is no statistically significant difference in foliar δ¹³C and δ¹⁵N between herbs and shrubs/subshrubs in this study. Considering that all plants studied here utilize the C₃ photosynthetic pathway, the lack of any clear distinction in foliar δ¹³C is not surprising. The absence of systematic differences in δ¹⁵N between these two groupings is less expected, given potential differences in N conservation and resorption efficiency [103] and patterns of root distribution and rooting depth, which affect resource acquisition [104]. The absence of a statistically meaningful difference in δ¹³N might arise from the unequal sample size of herbs vs. shrubs/subshrubs or confounding environmental factors such as plant growth stage, the range of available N sources, and the varied environmental conditions in the study area.

The shrubs and subshrubs have statistically significantly higher C and N contents than the herbs at both foliar and whole plant levels, as has been observed previously, especially for N
[104]. The difference in N content measured here might be related to different N conservation strategies during the late growing season. Arid and semi-arid perennial grasses have lower N contents in senesced leaves and overall higher N conservation efficiency than shrubs, which is an important adaptive trait for plants from nutrient-limited ecosystems [103]. Plants in this study were sampled in very late growing season (mid-late September in 2012 and late August in 2013), with perennial grasses comprising most herb samples. Hence, the difference in N contents may indicate higher N resorption efficiency from senescing tissues in grasses than in shrubs.

There are likely four main environmental factors that control the isotopic compositions of the herbaceous plants, shrubs and subshrubs sampled in this study: (i) water availability, (ii) N availability, (iii) spacial differences, and (iv) intra-plant variation. Each factor is discussed next.

4.2.1 Water availability. The very weak, but statistically significant positive correlation between foliar $\delta^{15}N$ and $\delta^{13}C$ of all samples analyzed (Fig 4) may point to water availability as at least a minor factor affecting both $\delta^{15}N$ and $\delta^{13}C$ of plants in this ecosystem. Plants capture atmospheric CO$_2$ through leaf stomata and fix it using enzymatic reactions, while N is mainly obtained through roots from soil or symbiotic associations and then assimilated. Given that the sources and pathways determining the $\delta^{13}C$ and $\delta^{15}N$ of plants are different, the observed, albeit weak, correlation may indicate a common environmental factor affecting both isotopic signals. As noted earlier, a change in MAP can affect both $\delta^{13}C$ and $\delta^{15}N$ of plants in the same direction (N: [11, 54, 80]; C: [42, 105, 106]). Ma et al. [43], for example, have reported an aridity-associated positive correlation between $\delta^{13}C$ and $\delta^{15}N$ in plants from northern China and noted that plant isotopic sensitivity to water availability can vary among ecosystems and plant species. Such variations warrant further consideration in paleoecological and paleodietary reconstructions of Arctic and subarctic regions.

4.2.2 N availability. The positive correlation between foliar $\delta^{15}N$ and N content suggests a key role for N availability and N cycling in determining the N isotopic signal acquired by plants in this ecosystem. Such a correlation has been reported previously on local [107], regional [54, 108] and global [11] scales. Higher plant $\delta^{15}N$ reflects higher N availability and a more open N cycle in terrestrial ecosystems [107, 109–111]. The globally observed positive correlation between soil and foliar $\delta^{15}N$ suggest foliar $\delta^{15}N$ as an index for N availability in soils and therefore ecosystems [11]. Nonetheless, the best way to describe N availability for plants in different ecosystems remains unclear [11]. It can be defined in several ways including: (i) an increase in N inputs into the soil from different sources (animal dung, plant materials, microbial N fixation), (ii) increased OM decomposition and N mineralization, (iii) increased NO$_3^-$ production through more nitrification [111], and (iv) less N demand by plants, particularly in drier localities [81]. In any of these scenarios, higher inorganic N availability in soils means that extra N is available to fuel N loss processes (e.g. denitrification and volatilization), which leave the bioavailable N in soils (NO$_3^-$, NH$_4^+$) enriched in $^{15}N$ [49, 50] (Fig 9a). In such ecosystems, plants acquire both higher $\delta^{15}N$ and N content in their leaves, which is characteristic of a more open N cycle. In N-limited ecosystems, by comparison, there is less N-bearing material available for N loss, and hence less opportunity for $^{15}N$ enrichment of the system through such processes. Plants also rely more heavily on mycorrhizal fungi for N acquisition, which is a more $^{15}N$-depleted source [11, 58] (Fig 9b).

4.2.3 Spacial differences. The statistically significant differences observed in foliar $\delta^{15}N$ among sites S13-2, S13-3, S13-7 and S13-6 points to heterogeneity in this ecosystem and the presence of different microhabitats even at small scales. These differences may be related to local variations in soil properties, slope aspects, topography, water availability, animal disturbance and grazing, which in turn can affect local N cycling. While we might predict higher foliar $\delta^{15}N$ at S13-7 than S13-6 given the former’s higher topsoil OM content (9.2 vs. 7.5 wt. %) and
TN (0.4 vs. 0.2 wt. %), the opposite result is obtained (S13-7-avg. δ¹⁵N = −2.0 ‰, n = 6) vs. (S13-6-avg. δ¹⁵N = +1.4 ‰, n = 8). Other factors such as slope aspect and topography may have over-shadowed the influence of soil properties. In S13-6, plant samples were collected from north side of the road on a SW-facing shallow slope, while in S13-7, plants were sampled on south side.
of the road on a NE-facing, shallow slope. This difference influenced the microclimate and vegetation pattern [24], as is indicated by the dominance of subshrubs at S13-7 and herbs at S13-6.

Leaves from S13-6 also have higher $\delta^{15}$N (avg. +1.4‰, n = 8) than those from site S13-3 (avg. −1.3‰, n = 11) and site S13-2 (avg. −0.7‰, n = 6). Soil at site S13-6 may provide plants with N pools having higher $\delta^{15}$N because of its shallower slope than site S13-3, and different topography than site S13-2.

Plants at S12-5 (avg. −28.7‰, n = 7) have significantly lower average foliar $\delta^{13}$C than at S12-2 (avg. −26.9‰, n = 7) and S12-6 (avg. −27.3‰, n = 7). The adjacency of site S12-5 to a main highway may contribute to lower plant $\delta^{13}$C arising from carbon dioxide additions from vehicle fuel combustion (typically −29.3 to −27.6‰; [112]). While Site S12-2 is also located adjacent to a main road, sampling occurred on a steep slope that is substantially elevated from the road.

4.2.4 Intra-plant variations. The average $\Delta^{13}$C_{other plant part-leaf} measured in this study are positive, consistent with previous studies [46, 113]. The differences measured here for most individual specimens, however, are small (<1‰) (Supporting Information, S1 Fig), and should not affect the use of $\delta^{13}$C to infer diet for animals that may prefer one plant part over another as forage.

No clear pattern in $\Delta^{15}$N_{other plant part-leaf} was observed, likely because of confounding factors such as differences in root morphology, depth and distribution, microhabitat, mycorrhizal associations and growing stage. The root crowns examined here have higher N contents (Fig 5) and $\delta^{15}$N (Fig 3) than other plant parts; this difference is statistically significant for N content, but not for $\delta^{15}$N. As discussed earlier, nitrogen contents of different plant parts commonly show variations between active growing and senescent stages in herbaceous plants [114, 115]. The higher N content of the root crowns in this study is best explained by N re-allocation from leaves to below ground parts late in the growing season.

5 Summary and implications

All plants analyzed from this ecosystem follow the C₃ photosynthetic pathway, and have an average whole plant $\delta^{13}$C of −27.5 ± 1.2‰ and foliar $\delta^{13}$C of −28.0 ± 1.3‰. The plants analyzed here showed very little variability in $\delta^{13}$C among different plant parts and sampling sites. Their isotopic composition, with suitable correction for the Suess effect arising from fossil fuel combustion since the Late Pleistocene, provides a suitable baseline for interpreting the diet of ancient herbivores that lived in these grasslands.

The nitrogen isotopic data for these modern plants (average whole plant $\delta^{15}$N = −0.3 ± 2.2‰ and foliar $\delta^{15}$N = −0.6 ± 2.7‰) provide a good baseline for the region’s vegetation under present conditions of nitrogen cycling. The wide range of intra-plant and inter-plant $\delta^{15}$N variations likely arises from multiple factors, including water availability, N availability and topography. Such variation is likely typical of vegetation from different microhabitats within Arctic and subarctic ecosystems. Given this complexity, modern vegetation $\delta^{15}$N is unlikely to adequately approximate Late Pleistocene vegetation in the same region.

A growing body of studies utilizes the isotopic composition of bone collagen and other tissues from Late Pleistocene megaaherbivores in high latitudes to reconstruct their diet. These works are hampered by insufficient isotopic data for plants at the base of food web in these now vanished ecosystems. More data for both modern and well-dated fossil plants from these ecosystems are needed to fully document and understand the variations in plant isotopic baselines both through time and space. The present study provides a starting point for modern vegetation in south and central Yukon Territory, Canada. While the plant N isotopic compositions presented here cannot be applied directly to the past, it provides a baseline for
comparison with its ancient equivalent, which can be obtained by analyzing fossil plants from the same region. We note that the significant positive correlation between foliar $\delta^{15}$N and N content observed for this region has potential as an index for forage quality. Comparison of the modern N isotopic vegetation baseline for this region with its Late Pleistocene equivalent could serve as a proxy for tracing changes in forage quality, which is tightly connected to ecosystem productivity and potentially, the sustainability of megaherbivore populations.

**Supporting information**

**S1 Text.** Section A. Methods for analyzing physical and chemical properties of soil. Section B. Analysis of carbon and nitrogen content. Section C. Results of soil analysis. Table A: Soil characteristics. Table B: Soil mineralogy. (DOCX)

**S1 Table.** $\delta^{13}$C and $\delta^{15}$N of all plant parts. (DOCX)

**S2 Table.** Carbon and nitrogen contents and foliar atomic C/N of all plants. (DOCX)

**S1 Fig.** Differences in $\delta^{13}$C between other plant tissues and leaf ($\Delta^{13}$C). Dashed lines represent means. (TIFF)

**S2 Fig.** Differences in $\delta^{15}$N between other plant tissues and leaf ($\Delta^{15}$N). Dashed lines represent means. (TIFF)

**Acknowledgments**

From the University of Western Ontario, we thank Kim Law and Li Huang of Laboratory for Stable Isotope Science (LSIS) for technical assistance and the late Dr. Jane Bowles (Department of Biology) for assistance with plant identification. We also thank the Yukon Government and Yukon First Nations for permission to collect samples (License No. 13-52S&E and No. 14-46S&E). We are grateful for funding from the Natural Sciences and Engineering Research Council of Canada (FJL), Canada Foundation for Innovation (FJL), Ontario Research Fund (FJL) and Canada Research Chairs Program (FJL), which supported this work. This manuscript has also benefitted greatly from the suggestions of two anonymous reviewers and the PLoS ONE editorial team. This is Laboratory for Stable Isotope Science Contribution #342.

**Author Contributions**

**Conceptualization:** Fred J. Longstaffe.

**Data curation:** Fred J. Longstaffe.

**Formal analysis:** Farnoush Tahmasebi.

**Funding acquisition:** Fred J. Longstaffe.

**Investigation:** Farnoush Tahmasebi.

**Methodology:** Farnoush Tahmasebi.

**Project administration:** Farnoush Tahmasebi, Fred J. Longstaffe.
Resources: Fred J. Longstaffe, Grant Zazula, Bruce Bennett.
Supervision: Fred J. Longstaffe.
Writing – original draft: Farnoush Tahmasebi.
Writing – review & editing: Farnoush Tahmasebi, Fred J. Longstaffe.

References

1. Casey MM, Post DM. The problem of isotopic baseline: Reconstructing the diet and trophic position of fossil animals. Earth-Sci Rev. 2011; 106:131–48.
2. Raghavan M, Themudo GE, Smith CI, Zazula G, Campos PF. Musk ox (Ovibos moschatus) of the mammoth steppe: Tracing palaeodietary and palaeoenvironmental changes over the last 50,000 years using carbon and nitrogen isotopic analysis. Quat Sci Rev. 2014; 102:192–201.
3. Mann DH, Groves P, Kunz ML, Reanier RE, Gaglioti BV. Ice-age megafauna in Arctic Alaska: Extinction, invasion, survival. Quat Sci Rev. 2013; 70:91–108.
4. Bocherens H. Isotopic biogeochemistry and the paleoecology of the mammoth steppe fauna. Deinsea. 2003; 9:57–76.
5. Fox-Dobbs K, Leonard JA, Koch PL. Pleistocene megafauna from eastern Beringia: Paleocological and palaeoenvironmental interpretations of stable carbon and nitrogen isotope and radiocarbon records. Palaeogeogr, Palaeoclimatol, Palaeoecol. 2008; 261:30–46.
6. Szpak P, Gröcke DR, Debruyne R, MacPhee RDE, Guthrie RD, Froese D, et al. Regional differences in bone collagen δ13C and δ15N of Pleistocene mammoths: Implications for paleoecology of the mammoth steppe. Palaeogeogr, Palaeoclimatol, Palaeoecol. 2010; 296:88–96.
7. Bocherens H. Isotopic tracking of large carnivore palaeoecology in the mammoth steppe. Quat Sci Rev. 2015; 117:42–71.
8. Iacumin P, Nikolaev V, Ramigni M. C and N stable isotope measurements on Eurasian fossil mammoths, 40 000 to 10 000 years BP: Herbivore physiologies and palaeoenvironmental reconstruction. Palaeogeogr, Palaeoclimatol, Palaeoecol. 2000; 163:33–47.
9. Zazula GD, Froese Duane G., Schweger CE, Mathewes RW, Beaudoin AB, Telka AM, et al. Ice-age steppe vegetation in east Beringia. Nature. 2003; 423:603. https://doi.org/10.1038/423603a PMID: 12789326
10. Breen AL, Murray DF, Raynolds MK, Timling I, Walker DA. Ecology and evolution of plants in Arctic and Alpine Environments. In: Rajakaruna N, Boyd RS, Harris TB, editors. Plant ecology and evolution in harsh environments. Hauppauge, New York: Nova Science Publishers; 2014. p. 149–77.
11. Craine JM, Elmore AJ, Aidar MPM, Bustamante M, Dawson TE, Hobbie EA, et al. Global patterns of foliar nitrogen isotopes and their relationships with climate, mycorrhizal fungi, foliar nutrient concentrations, and nitrogen availability. New Phytol. 2009; 183:980–92. https://doi.org/10.1111/j.1469-8137.2009.02917.x PMID: 19563444
12. Makarov MI. The nitrogen isotopic composition in soils and plants: Its use in environmental studies (A Review). Eurasian Soil Sci. 2009; 42:1335–47.
13. Wang Y, Wooller MJ. The stable isotopic (C and N) composition of modern plants and lichens from northern Iceland: With ecological and palaeoenvironmental implications. Jökull. 2006; 56:27–37.
14. Wooller MJ, Zazulat GD, Edwards M, Froese DG, Boone RD, Parker C, et al. Stable carbon isotope compositions of eastern Beringian grasses and sedges: Investigating their potential as palaeoenvironmental indicators. AAAR. 2007; 39:318–31.
15. Michelsen A, Schmidt IK, Jonasson S, Quarmby C, Sleep D. Leaf 15N abundance of subarctic plants provides field evidence that ericoid, ectomycorrhizal and non- and arbuscular mycorrhizal species access different sources of soil nitrogen. Oecologia. 1996; 105:53–63. https://doi.org/10.1007/BF00328791 PMID: 28307122
16. Nadelhoffer K, Shaver G, Fry B, Giblin A, Johnson L, McKane R. 15N natural abundances and N use by tundra plants. Oecologia. 1996; 107:386–94. https://doi.org/10.1007/BF00328456 PMID: 28307268
17. Schulze ED, Chapin FS, Gebauer G. Nitrogen nutrition and isotope differences among life forms at the northern treeline of Alaska. Oecologia. 1994; 100:406–12. https://doi.org/10.1007/BF00317862 PMID: 28306929
18. Welker JM, Jonsdottir IS, Fahnestock JT. Leaf isotopic (δ13C and δ15N) and nitrogen contents of Carex plants along the Eurasian Coastal Arctic: Results from the Northeast Passage expedition. Polar Biol. 2003; 27:29–37.
19. Höflé C, Edwards ME, Hopkins DM, Mann DH, Ping C-L. The full-glacial environment of the Northern Seward Peninsula, Alaska, reconstructed from the 21,500-Year-Old Kitluk Paleosol. Quatern Res. 2000; 53:143–53.

20. Zimov SA, Zimov NS, Tikhonov AN, Chapin FS. Mammoth steppe: A high-productivity phenomenon. Quat Sci Rev. 2012; 57:26–45.

21. Laxton NF, Burn CR, Smith CAS. Productivity of loessal grasslands in the Kluane Lake region, Yukon Territory, and the Beringian “Production Paradox”. Arctic. 1996; 49:129–40.

22. Guthrie RD. Fozen Fauna of the Mammoth Steppe, The Story of Blue Babe: The University of Chicago Press Chicago and London; 1990.

23. Denton GH, Stuiver M. Neoglacial chronology, northeastern Saint Elias Mountains, Canada. Am J Sci. 1966; 264:577–99.

24. Sanborn PT, Jull AJT. Loess, bioturbation, fire, and pedogenesis in a boreal forest-grassland mosaic, Yukon Territory, Canada. 19th World Congress of Soil Science, Soil Solutions for a Changing World. 2010.

25. Hoefs M, Cowan I, Krajina V. Phytosociological analysis and synthesis of Sheep Mountain, southwest Yukon territory, Canada. Syesis. 1975; 8:125–228.

26. Cody WJ. Flora of the Yukon Territory. 2 ed: NRC Research Press; 2000.

27. Andreeva DB, Zech M, Glaser B, Erbajeva MA, Chimitdorgieva GD, Ermakova OD, et al. Stable isotope (δ¹³C, δ¹⁵N, δ¹⁸O) record of soils in Buryatia, southern Siberia: Implications for biogeochemical and paleoclimatic interpretations. Quat Int. 2013; 290–291:82–94.

28. Stevenson BA, Kelly EF, McDonald EV, Busacca AJ. The stable carbon isotope composition of soil organic carbon and pedogenic carbonates along a bioclimatic gradient in the Palouse region, Washington State, USA. Geoderma. 2005; 124:37–47.

29. Boutton TW. Stable carbon isotope ratios of soil organic matter and their use as indicators of vegetation and climate change. In: Boutton TW, Yamasaki SI, editors. Mass Spectrometry of Soils 1996. p. 47–82.

30. Pessenda LCR, Valencia EPE, Aravena R, Telles ECC, Boulet R. Palaeoclimatic studies in Brazil using carbon isotopes in soils. In: Waterman JC, Silva-Filho EV, Villas-Boas R, editors. Environmental Geochemistry in the Tropics. Lecture Notes in Earth Sciences. 72: Springer-Verlag, Berlin/New York.; 1998. p. 7–16.

31. Tieszen LL. Natural variations in the carbon isotope values of plants: Implications for archeology, ecology, and paleoecology. J Archaeol Sci. 1991; 18:227–48.

32. Krull ES, Bestland EA, Gates WP. Soil organic matter decomposition and turnover in a tropical Ultisol: Evidence from δ¹³C, δ¹⁵N and Geochmistry. Radiocarbon. 2002; 44:93–112.

33. Bol RA, Harkness D.D, Huang Y, Howard DM. The influence of soil processes on carbon isotope distribution and turnover in the British uplands. Eur J Soil Sci. 1999; 50:41–51.

34. Boström B, Comstedt D, Ekblad A. Isotope fractionation and¹³C enrichment in soil profiles during the decomposition of soil organic matter. Oecologia. 2007; 153:89–98. https://doi.org/10.1007/s00442-007-0700-8 PMID: 17401582

35. Boutton TW, Archer SR, Midwood AJ, Zitzer SF, Bol R. δ¹³C values of soil organic carbon and their use in documenting vegetation change in a subtropical savanna ecosystem. Geoderma. 1998; 82:5–41.

36. Farquhar GD, Sharkey TD. Stomatal conductance and photosynthesis. Annual Reviews of Plant Physiology. 1982; 33:317–45.

37. Tieszen LL, Boutton TW. Stable carbon isotopes in terrestrial ecosystem research. In: Rundel PW, Ehleringer JR, Nagy KA, editors. Stable Isotopes in Ecological Research. New York: Springer; 1989. p. 167–95.

38. Farquhar GD, Ehleringer JR, Hubick KT. Carbon isotope discrimination and photosynthesis. Annu Rev Plant Physiol Plant Mol Biol. 1989; 40:503–37.

39. Welsh SL. Anderson’s flora of Alaska and adjacent parts of Canada. Provo: Brigham Young University Press xvi, 724p. Illustrations, portrait, map. 1974.

40. Diefendorf AF, Mueller KE, Wing SL, Koch PL, Freeman KH. Global patterns in leaf ¹³C discrimination and implications for studies of past and future climate. Proceedings of the National Academy of Sciences. 2010; 107:5738–43.

41. Kohn MJ. Carbon isotope compositions of terrestrial C₃ plants as indicators of (paleo)ecology and (paleo)climate. Proceedings of the National Academy of Sciences. 2010; 107:19691–5.

42. Weiguo L, Xiaohong F, Youfeng N, Qingle Z, Yunning C, Zhisheng AN. δ¹³C variation of C₃ and C₄ plants across an Asian monsoon rainfall gradient in arid northwestern China. Global Change Biol. 2005; 11:1094–100.
43. Ma JY, Sun W, Liu XN, Chen F-H. Variation in the stable carbon and nitrogen isotope composition of plants and soil along a precipitation gradient in Northern China. PLoS ONE. 2012; 7:e51894. https://doi.org/10.1371/journal.pone.0051894 PMID: 23272186

44. Wang G, Feng X, Han J, Zhou L, Tan W, Su F. Paleovegetation reconstruction using δ¹³C of soil organic matter. Biogeosciences. 2008; 5:1325–37.

45. Medina E, Montes G, Guevas E, Rokzandic Z. Profiles of CO₂ concentration and δ¹³C values in tropical rain forests of the Upper RioNegro Basin, Venezuela. J Trop Ecol. 1986; 2:207–17.

46. Cernusak LA, Tcherkez G, Keitel C, Cornwell WK, Santiago LS, Knoll A, et al. Why are non-photosynthetic tissues generally ¹³C enriched compared with leaves in C₃ plants? Review and synthesis of current hypotheses. Funct Plant Biol. 2009; 36:199–213.

47. Näsholm T, Kielland K, Ganeteg U. Uptake of organic nitrogen by plants. New Phytol. 2009; 182:31–48. https://doi.org/10.1111/j.1469-8137.2008.02751.x PMID: 19210725

48. Wei L, Chen C, Xu Z, Näsholm T. Direct uptake and rapid decrease of organic nitrogen by *Wollemia nobilis*. Biol Fertility Soils. 2013; 49:1247–52.

49. Höegberg P. ¹⁵N natural abundance in soil-plant systems. New Phytol. 1997; 137:179–203.

50. Robinson D. δ¹⁵N as an integrator of the nitrogen cycle. TRENDS in Ecology and Evolution. 2001; 16:153–62. PMID: 1179580

51. Houlton BZ, Bai E. Imprint of denitrifying bacteria on the global terrestrial biosphere. Proceedings of the National Academy of Sciences. 2009; 106:21713–6.

52. Handley L, Raven JA. The use of natural abundance of nitrogen isotopes in plant physiology and ecology. Plant, Cell Environ. 1992; 15:965–85.

53. Makarov MI, Malyshева TI, Cornelissen JHC, van Logtestijn RSP, Glasser B. Consistent patterns of ¹⁵N distribution through soil profiles in diverse alpine and tundra ecosystems. Soil Biol Biochem. 2008; 40:1082–9.

54. Martinelli LA, Piccolo MC, Townsend AR, Vitousek PM, Cuevas E, McDowell W, et al. Nitrogen stable isotopic composition of leaves and soil: Tropical versus temperate forests. Biogeochemistry. 1999; 46:45–65.

55. Robinson D, Rorison IH. Relationships between root morphology and nitrogen availability in a recent theoretical model describing nitrogen uptake from soil. Plant, Cell and Environment. 1983; 6:641–7.

56. Miller A, Bowman W. Variation in nitrogen-15 natural abundance and nitrogen uptake traits among co-occurring alpine species: Do species partition by nitrogen form? Oecologia. 2002; 130:609–16. https://doi.org/10.1007/s00442-001-0838-8 PMID: 28547264

57. Shaver GR, Kummerow J. Phenology, resource allocation, and growth of arctic vascular plants. Arctic ecosystems in a changing climate: An ecophysiological perspective. 1992:193–211.

58. Hobbie EA, Hobbie JE. Natural abundance of ¹⁵N in nitrogen-limited forests and tundra can estimate nitrogen cycling through mycorrhizal fungi: A review. Ecosystems. 2008; 11:815–30.

59. Michelsen A, Quarmby C, Sleep D, Jonasson S. Vascular plant ¹⁵N natural abundance in heath and forest tundra ecosystems is closely correlated with presence and type of mycorrhizal fungi in roots. Oecologia. 1998; 115:406–16. https://doi.org/10.1007/s004420050535 PMID: 28308434

60. Hobbie JE, Hobbie EA. ¹⁵N in symbiotic fungi and plants estimates nitrogen and carbon flux rates in Arctic tundra. Ecology. 2006; 87:816–22. PMID: 16676524

61. Hobbie EA, Macko SA, Shugart HH. Insights into nitrogen and carbon dynamics of ectomycorrhizal and saprotrophic fungi from isotopic evidence. Oecologia. 1999; 118:353–60. https://doi.org/10.1007/s004420050736 PMID: 28307279

62. Frank DA, Groffman PM, Evans RD, Tracy BF. Ungulate stimulation of nitrogen cycling and retention in Yellowstone Park grasslands. Oecologia. 2000; 123:116–21. https://doi.org/10.1007/s004420050996 PMID: 28308736

63. Augustine DJ, Frank DA. Effects of migratory grazers on spatial heterogeneity of soil nitrogen properties in a grassland ecosystem. Ecology. 2001; 82:3149–62.

64. Semmartin M, Aguair MR, Distel RA, Moretto AS, Ghersa CM. Litter quality and nutrient cycling affected by grazing-induced species replacements along a precipitation gradient. Oikos. 2004; 107:148–60.

65. Aranibar JN, Anderson IC, Epstein HE, Feral CJW, Swap RJ, Ramontsho J, et al. Nitrogen isotope composition of soils, C₂ and C₄ plants along land use gradients in southern Africa. J Arid Environ. 2008; 72:326–37.

66. Coetsee C, Stock WD, Craine JM. Do grazers alter nitrogen dynamics on grazing lawns in a South African savannah? Afr J Ecol. 2010; 49:62–9.
N and C isotopic dynamics of subarctic soils and plants in southern Yukon Territory

67. Li C, Hao X, Willms WD, Zhao M, Han G. Effect of long-term cattle grazing on seasonal nitrogen and phosphorus concentrations in range forage species in the fescue grassland of southwestern Alberta. J Plant Nutr Soil Sci. 2010; 173:946–51.

68. Golluscio RA, Austin AT, García Martínez GC, Gonzalez-Polo M, Sala OE, Jackson RB. Sheep grazing decreases organic carbon and nitrogen pools in the Patagonian Steppe: Combination of direct and indirect effects. Ecosystems. 2009; 12:686–97.

69. Cook GD. Effects of frequent fires and grazing on stable nitrogen isotope ratios of vegetation in northern Australia. Australian Ecologv. 2001; 26:630–6.

70. Wittmer MHOM, Auerswald K, Schönbach P, Bai Y, Schnyder H. Natural 15N abundance in soils and plants in relation to N cycling in a rangeland in Inner Mongolia. Journal of Plant Ecology. 2010; 3:201–7.

71. Xu Y, He J, Cheng W, Xing X, Li L. Natural 15N abundance in soils and plants in relation to N cycling in a rangeland in Inner Mongolia. Journal of Plant Ecology. 2010; 3:201–7.

72. Evans RD. Physiological mechanisms influencing plant nitrogen isotope composition. Trends Plant Sci. 2001; 6:121–6. PMID: 11239611

73. Kolb KJ, Evans RD. Influence of nitrogen source and concentration on nitrogen isotopic discrimination in two barley genotypes (Hordeum vulgare L.). Plant, Cell and Environment. 2003; 26:1431–40.

74. Craine JM, Brookshire ENJ, Cramer MD, Hasselquist NJ, Koba K, Marin-Spiotta E, et al. Ecological interpretations of nitrogen isotope ratios of terrestrial plants and soils. Plant Soil. 2015; 1–26.

75. Mariotti A, Mariotti F, Champigny ML, Aamarger N, Moyse A. Nitrogen isotope fractionation associated with nitrate reductase activity and uptake of NO3- by pearl millet. Plant Physiol. 1982; 69:880–4. PMID: 16662313

76. Yoneyama T, Matsumaru T, Usui K, Engelaar WMHG. Discrimination of nitrogen isotopes during absorption of ammonium and nitrate at different nitrogen concentrations by rice (Oryza sativa L.) plants. Plant, Cell and Environment. 2001; 24:133–9.

77. Lebauer DS, Tredser KK. Nitrogen limitation of net primary productivity in terrestrial ecosystems is globally distributed. Ecology. 2008; 89:371–9. PMID: 18409427

78. Yoneyama T, Kaneko A. Variations in the natural abundance of 15N in nitrogenous fractions of komatsuna plants supplied with nitrate. Plant Cell Physiol. 1989; 30:957–62.

79. Kolb KJ, Evans RD. Implications of leaf nitrogen recycling on the nitrogen isotope composition of deciduous plant tissues. New Phytol. 2002; 156:57–64.

80. Amundson R. Global patterns of the isotopic composition of soil and plant nitrogen. Global Biogeochem Cycles. 2003; 17:1031.

81. Austin AT, Vitousek PM. Nutrient dynamics on a precipitation gradient in Hawai‘i. Oecologia. 1998; 113:519–29. https://doi.org/10.1007/s004420050405 PMID: 28308032

82. Schuur EA, Matson PA. Net primary productivity and nutrient cycling across a mesic to wet precipitation gradient in Hawaiian montane forest. Oecologia. 2001; 128:431–42. https://doi.org/10.1007/s004420100671 PMID: 24549913

83. DeNiro M, Epstein S. Influence of diet on the distribution of nitrogen isotopes in animals. Geochim Cosmochim Acta. 1981; 45:341–51.

84. Harris D, Horwath WR, Kessel CV. Acid fumigation of soils to remove carbonates prior to total organic carbon or carbon-13 isotopic analysis. Soil Sci Soc Am J. 2001; 65:1853–6.

85. Webb EA, Schwarz HP, Healy PF. Detection of ancient maize in lowland Maya soils using stable carbon isotopes: Evidence from Caracol, Belize. J Archaecol Sci. 2004; 31:1039–52.

86. Coplen TB. Guidelines and recommended terms for expression of stableisotope- ratio and gas-ratio measurement results. Rapid Commun Mass Spectrom. 2011; 25:2538–60. PMID: 21910288

87. Coplen TB, Brand WA, Gehre M, Groning M, Meijer HAJ, Toman B, et al. New guidelines for δ13C measurements. Anal Chem. 2006; 78:2439–41. https://doi.org/10.1021/ac052027c PMID: 16579631

88. Qi H, Coplen TB, Geilmann H, Brand WA, Bohlke JK. Two new organic reference materials for δ13C and δ15N measurements and a new value for the δ13C of NBS 22 oil. Rapid Commun Mass Spectrom. 2003; 17:2483–7. https://doi.org/10.1021/rcm.1219 PMID: 14608617

89. Walker D, Raynolds MK, Danieł FJA, Einarsson E, Elvebakk A, Gould WA, et al. The Circumpolar Arctic vegetation map. Journal of Vegetation Science. 2005; 16:267–82.

90. Chapman GP. The Biology of Grasses: CAB International; 1996.

91. Frank DA, Evans RD, Tracy BF. The role of ammonia volatilization in controlling the natural 15N abundance of a grazed grassland. Biogeochemistry. 2004; 68:169–78.

92. Harris SA. Dynamics and origin of saline soils on the Slims River Delta, Kluane National Park, Yukon Territory. Arctic. 1990; 43:159–75.
93. Plint T, Longstaffe FJ, Hodgetts L, Hodgins G, Rybczynski N, Southon J, et al., editors. Stable isotopic investigations of giant beaver diet and econiche—Not another dam beaver. Geological Associations of Canada Annual Meeting, 2016 June 1–3; Whitehorse, Yukon, Canada. [Abstract]

94. Naidu AS, Cooper LW, Finney BP, Macdonald RW, Alexander C, Semiletov IP. Organic carbon isotope ratios (δ^{13}C) of Arctic Amerasian continental shelf sediments. Int J Earth Sci. 2000; 89:522–32.

95. Christensen BT. Physical fractionation of soil and structural and functional complexity in organic matter turnover. Eur J Soil Sci. 2001; 52:345–53.

96. Menyailo OV, Hungate BA. Carbon and nitrogen stable isotopes in forest soils of Siberia. Dokl Earth Sci. 2006; 409:747–9.

97. Wang C, Wang X, Liu D, Wu H, Lu X, Fang Y, et al. Aridity threshold in controlling ecosystem nitrogen cycling in arid and semi-arid grasslands. Nature Communications. 2014; 5:1–5.

98. Ledgard SF, Freney JR, Simpson JR. Variations in natural enrichment of δ^{15}N in the profiles of some Australian pasture soils. Aust J Soil Res. 1984; 22:155–64.

99. Nacro HB, Larre-Larrouy MC, Mariotti A, Feller C, Abbadi L. Natural nitrogen-15 abundance and carbohydrate content and composition of organic matter particle-size fractions from a sandy soil. Biol Fertility Soils. 2004; 40:171–7.

100. Tiessen H, Karmanos R, Stewart J, Selles F. Natural nitrogen-15 abundance as an indicator of soil organic matter transformations in native and cultivated soils. Soil Sci Soc Am J. 1984; 48:312–5.

101. Guideau SA, Graham RC, Feng X, Chadwick OA. Natural isotopic distribution in soil surface horizons differentiated by vegetation. Soil Sci Soc Am J. 2003; 67:1544–50.

102. McKane RB, Johnson LC, Shaver GR, Nadelhoffe KJ, Rastetter EB, Fry B, et al. Resource-based niches provide a basis for plant species diversity and dominance in arctic tundra. Nature. 2002; 415:68–71. https://doi.org/10.1038/415068a PMID: 11780117

103. Bertiller MB, Sain CL, Carrera AL, Vargas DN. Patterns of nitrogen and phosphorus conservation in dominant perennial grasses and shrubs across an aridity gradient in Patagonia, Argentina. J Arid Environ. 2005; 62:209–23.

104. Sala OE, Golluscio RA, Lauenroth WK, Soriano A. Patterns of nitrogen and phosphorus conservation in dominant perennial grasses and shrubs across an aridity gradient in Patagonia, Argentina. J Arid Environ. 2005; 62:209–23.

105. Murphy BP, Bowman DMJS. The carbon and nitrogen isotope composition of Australian grasses in relation to climate. Funct Ecol. 2009; 23:1040–9.

106. Swap RJ, Araniba JN, Dowty PR, Gilhooly WP, Macko SA. Natural abundances of δ^{13}C and δ^{15}N in C_3 and C_4 vegetation of southern Africa: Patterns and implications. Global Change Biol. 2004; 10:350–8.

107. McLauchlan KK, Ferguson CJ, Wilson IE, Ocheltree TW, Craine JM. Thirteen decades of foliar isotopes indicate declining nitrogen availability in central North American grasslands. New Phytol. 2010; 187:1135–45. https://doi.org/10.1111/j.1469-8137.2010.03322.x PMID: 20553396

108. Craine JM, Lee WG, Bond WJ, Williams RJ, Johnson LC. Environmental constraints on a global relationship among leaf and root traits of grasses. Ecology. 2005; 86:12–9.

109. Hietz P, Turner BL, Wanek W, Richter A, Nock CA, Wright SJ. Long-term change in the nitrogen cycle of tropical forests. Science. 2011; 334:664–6. https://doi.org/10.1126/science.1211979 PMID: 22053047

110. McLauchlan KK, Williams JJ, Craine JM, Jeffers ES. Changes in global nitrogen cycling during the Holocene epoch. Nature. 2013; 495:352–5. https://doi.org/10.1038/nature11916 PMID: 23518563

111. Pardo LH, Templer PH, Goodale CL, Duke S, Groffman PM, Adams MB, et al. Regional assessment of N saturation using foliar and root δ^{15}N. Biogeochemistry. 2006; 80:143–71.

112. Widory D, Javoy M. The carbon isotope composition of atmospheric CO₂ in Paris. Earth Planet Sci Lett. 2003; 215:289–98.

113. Hobbie EA, Werner RA. Intramolecular, compound-specific, and bulk carbon isotope patterns in C_3 and C_4 plants: A review and synthesis. New Phytol. 2004; 161:371–85.

114. Choi WJ, Huang SX, Ro HM. Seasonal changes of shoot nitrogen concentrations and δ^{15}N/δ^{14}N ratios in common reed in a constructed wetland. Commun Soil Sci Plant Anal. 2005; 36:2719–31.

115. Bausenwein U, Millard P, Thornton B, Raven JA. Seasonal nitrogen storage and remobilization in the forb Rumex acetosa. Funct Ecol. 2001; 15:370–7.