An Assessment of Plant Species Differences on Cellulose Oxygen Isotopes From Two Kenai Peninsula, Alaska Peatlands: Implications for Hydroclimatic Reconstructions

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Peat cores are valuable archives of past environmental change because they accumulate plant organic matter over millennia. While studies have primarily focused on physical, ecological, and some biogeochemical proxies, cores from peatlands have increasingly been used to interpret hydroclimatic change using stable isotope analyses of cellulose preserved in plant remains. Previous studies indicate that the stable oxygen isotope compositions ($\delta^{18}O$) preserved in alpha cellulose extracted from specific plant macrofossils reflect the $\delta^{18}O$ values of past peatland water and thereby provide information on long-term changes in hydrology in response to climate. Oxygen isotope analyses of peat cellulose ($\delta^{18}O_{\text{cellulose}}$) have been successfully developed from peat cores that accumulate the same species for millennia. However, to fully exploit the potential of this proxy in species-diverse fens, studies are needed that account for the isotopic variations caused by changes in dominant species composition. This study assesses variation in $\delta^{18}O$ values among peatland plant species and how they relate to environmental waters in two fens informally named Horse Trail and Goldfin, located on the leeward (dry) and windward (wet) side, respectively, of the climatic gradient across the Kenai Peninsula, Alaska. Environmental water $\delta^{18}O$ values at both fens reflect unmodified $\delta^{18}O$ values of mean annual precipitation, although at Goldfin standing pools were slightly influenced by evaporation. Modern plant [mosses and Carex spp. (sedges)] $\delta^{18}O_{\text{cellulose}}$ values indicate that all Carex spp. are higher ($\sim 2.5\%$) than those of mosses, likely driven by their vascular structure and ecophysiological difference from non-vascular mosses. Moss $\delta^{18}O_{\text{cellulose}}$ values within each peatland are similar among the species, and differences appear related to evaporation effects on environmental waters within hummocks and hollows. The plant taxa-environmental water $\delta^{18}O$ differences are applied to the previously determined Horse Trail Fen untreated bulk $\delta^{18}O$ record. Results include significant changes to inferred millennial-to-centennial scale hydroclimatic trends where dominant taxa shift from moss to Carex spp., indicating that modern calibration datasets
are necessary for interpreting stable isotopes from fens, containing a mix of vascular and nonvascular plants. Accounting for isotopic offsets through macrofossil analysis and modern plant-water isotope measurements opens new opportunities for hydroclimatic reconstructions from fen peatlands.

Keywords: hydroclimate, peatland archives, oxygen isotopes (δ18O), cellulose, calibration dataset

INTRODUCTION

Peat core records have long served as geologic archives of paleoenvironmental change, using a range of biological, physical, and biogeochemical proxies. Because peat accumulates under waterlogged conditions, previous studies have related the oxygen isotopic signature preserved in alpha cellulose (δ18O_{cellulose}) extracted from peatland plant macrofossils to the isotopic composition of environmental water, i.e., the source from which the plants are absorbing their water (Vardy, 1997; Wissel et al., 2008; Moschen et al., 2009; Loader et al., 2016). The use of δ18O_{cellulose} values from peatlands was first explored in Sphagnum-dominated bogs using analyses of single species (Brenninkmeijer et al., 1982; Ménot-Combes et al., 2002; Zanazzi and Mora, 2005; Daley et al., 2010, 2016; Kühl and Moschen, 2012; Loader et al., 2016). Single-species cellulose in a core limits variability that could arise from isotopic species effects (Ménot-Combes et al., 2002; Daley et al., 2010; Nichols et al., 2010). Bogs, hydraulically sourced only by precipitation, are generally thought to provide the best record of past precipitation change from peatlands, however, relying only on bogs that accumulate monospecific peat for millennia can considerably limit the utility of this proxy to a small number of sites. For example, many Alaska peatlands are primarily groundwater-sourced fens with high species diversity through space and time. The utility of this proxy was tested in one Alaska fen core by analyzing δ18O changes relative to the plant macrofossils comprising the peat (Jones et al., 2014), and while isotopic shifts occurred that were consistent with other regional records (Fisher et al., 2004; Anderson et al., 2005), questions remain about how closely tied the shifts were to hydroclimatic changes vs. other factors, and most specifically, plant species shifts in response to changes in peatland hydrology.

The primary isotopic fractionation involved in biochemically synthesizing cellulose in environmental water has been relatively well constrained to −27‰ (DeNiro and Epstein, 1979, 1981), and while this process was thought to be insensitive to changes in temperature, Sternberg and Ellsworth (2011) quantified a latitudinal influence associated with differences in mean annual air temperatures (MAT) that is −33‰ for high latitudes such as Alaska. In vascular plants, δ18O_{cellulose} is determined by kinetic and equilibrium fractionation from its environmental water (Amesbury et al., 2015), because values are influenced by both the δ18O values of environmental water and water within the leaf, which are generally enriched in heavy isotopes due to transpiration and plant vascular mediation (Nichols et al., 2010; Loader et al., 2016). Therefore, in Alaskan peats, which are largely composed of a variety of bryophytes and sedges, contrasting physiological processes for water uptake between bryophytes (mosses; non-vascular) and graminoids (seeds; vascular) likely result in significant differences in their cellulose isotope signatures. Bryophytes, including all mosses and liverworts, lack vascular structure and absorb water through cell walls (Proctor, 2000). In contrast, sedges have roots that can obtain water from deeper horizons and regulate moisture loss via stomata, leading to evaporative enrichment from transpiration through their stomata (Yakir et al., 1990; Amesbury et al., 2015). This may potentially lead to differences not only in environmental-water isotope compositions compared to mosses, but also in additional fractionations of water within the plant related to stomatal conductance (Ménot and Burns, 2001).

To further examine oxygen isotope variations of plant cellulose within Alaskan fens, the goals of this study are to (a) determine the δ18O values of modern peatland environmental waters and evaluate their relationship to precipitation and groundwater by comparison with local surface water δ18O values, including Kenai Peninsula lakes, rivers and streams, and the global meteoric water line (GMWL) (Rozanski et al., 1993; Anderson et al., 2016); (b) compare δ18O_{cellulose} values of plant species with the δ18O values of their environmental waters to determine whether species δ18O values significantly differ; and (c) evaluate the range of δ18O_{cellulose} values exhibited by different peatland-plant species spatially. Lastly, the results are used to evaluate the influence of varying species dominance on bulk peat δ18O values determined from a ~14,000-year old core obtained in Horse Trail Fen (HTF) by Jones et al. (2014) that, in turn, influences the hydroclimatic interpretation.

Study Area

The Kenai Peninsula is located in south-central Alaska on the northern coast of the Gulf of Alaska (Figure 1). It is bisected along its eastern edge by the Kenai Mountains, composed of Mesozoic bedrock (Rymer and Sims, 1982), which rise to ~1,025 meters above sea level (m.a.s.l.). The Harding Icefield spans upper elevations of the range, with glaciers terminating at or near sea level to the east and west. The western side of the peninsula is a low-lying landscape shaped by glacial events (Rymer and Sims, 1982; Reger et al., 2008), and is characterized by peatlands, kettle-hole lakes, and boreal forest overlying moraines and glacial outwash. The eastern side of the Kenai Mountains is the wetter, windward side, and precipitation totals are four times higher than on the leeward, western lowlands, spanning a relatively small (<70 km) area. Consequently, the vegetation on the eastern side comprises the western-most edge of the temperate rainforest, whereas
the western lowlands form an ecotone between the boreal and coastal forests.

The climate of the eastern Kenai Mountains is considered maritime, with MAT of 6.5°C with a mean annual precipitation (MAP) of 186.3 cm (Seward, AK), while on the western lowlands, the MAT at Soldotna airport is 6°C and MAP is 46.3 cm (Figure 1D; 1981–2010 climate average; Alaska Climate Research Center, http://climate.gi.alaska.edu). On the western
lowlands, the majority of present-day annual precipitation falls during autumn (August, September) and remains relatively high in winter (October-January) and in the eastern Kenai Peninsula, precipitation is highest from September to January (Alaska Climate Research Center, http://climate.gi.alaska.edu), in response to intensification of the Aleutian Low, a semi-permanent low-pressure system that strengthens over the North Pacific and Gulf of Alaska region in late autumn and subsequently wanes in intensity during spring and summer (Overland et al., 1999).

Horse Trail fen (HTF; informal name, 60.264, −149.356, 110 m a.s.l.), is a large fen complex in the western lowlands near Soldotna, AK. Although located downgradient from the Harding Icefield, watershed analysis using high resolution Digital Elevation Models indicates that the headwater area is isolated from glacial runoff (white outline in Figure 1C). Goldfin (GF, informal; 60.264, −149.356, 2018 m a.s.l.), is a small kettle hole peatland located in a narrow north-to-south trending valley ~30 m above sea level north of Seward, AK. Plant and water isotope samples presented here also include samples obtained from St. Matthew Island (Figure 1A), located in the east-central Bering Sea (60.4°N, −172.7°W, at sea level elevation).

METHODS

Field collections for this study included modern plants [mosses and sedges (Carex spp.)] and associated environmental water. The Horse Trail fen (HTF) site was sampled for water and plants along a 40-m transect from the stream flowing through the peatland to the forest edge at every 10 m in July of 2014. At each point along the transect, surface water associated with plant collections was obtained by submerging 10-mL high-density polyethylene (HDPE) bottles and under the surface of the water table until it filled without bubbles or headspace before sealing under the water. No water was squeezed out of peatland plants. Surface water samples were similarly obtained from near-surface waters of nearby lakes, rivers, and streams (Figure 1C).

At each HTF transect position, the dominant plant species were collected, noting whether the plant was submerged, at the water table, or on a hummock. A similar transect was repeated at GF bog in September 2017, starting at the stream flowing at the edge of the bog to 40 m away from the stream. In this bog, an elevation gradient perpendicular to the stream was less apparent and small pools intersected the transect. At the 20-m sampling location, samples were collected from a small pool at the water table and 10 cm above the water table. At the 40-m location, samples were collected from a small pool and an adjacent hummock 10–20 cm high.

Plant species were bagged and labeled with their transect position and subsequently cleaned with deionized water, identified, and separated by species in the laboratory. If enough material was available, species were separated by their stems and leaves. In some cases, whole plants were analyzed, either in addition to separated stems and leaves or in some instances by themselves. Plants and environmental-water samples from St. Matthew Island, Alaska, were collected June 2018, repeating laboratory methods outlined above.

Plant cellulose was extracted by the Cuprammonium (CUAM) method of Wissel et al. (2008) and Moschen et al. (2009). Samples were first bleached using solution of sodium hypochlorite and acetic acid at 70°C to separate the lignin fraction before they were neutralized and freeze dried. The bleached sample was then placed in 50-ml centrifuge tubes using a cuprammonium solution (Cu(NH)3(4OH)2; “Schweizer solution”) for cellulose dissolution, stirred for 6 h, and left to sit at room temperature for an additional 10 h. Once fully dissolved, the copper complex cellulose solution was centrifuged for 25 min at 2,500 rpm and the supernatant was decanted into a clean 50-ml centrifuge tube, leaving the non-cellulosic material behind. This step was repeated to avoid contamination from other plant material. The supernatant was then loaded with ~3 mL H2SO4 (20%) and cold deionized water to induce cellulose precipitation. The tube was shaken and left to sit for 20 min, before it was centrifuged at 2,500 rpm for 25 min. Additional drops of H2SO4 (20%) were added until the solution turned from blue to clear, to ensure that cellulose precipitation was complete. The cellulose precipitate was subsequently rinsed with deionized water until it reached a neutral pH by centrifuging at 2,500 rpm for 25 min, before it was freeze dried. A mass of 0.2–0.3 mg of dried cellulose sample was packed in silver capsules and stored in a vacuum drier prior to stable oxygen isotopic analysis.

All oxygen and water hydrogen isotope results for cellulose and water are reported in per mil (‰) as the relative difference of isotope ratios (δ) from the international measurement standard Vienna Standard Mean Ocean Water (VSMOW) defined by

\[ \delta^{18}O_{H_2O} = \left[\frac{^{18}O/^{16}O}_{H_2O}/^{18}O/^{16}O_{VSMOW}\right] - 1 \]

\[ \delta^2H_{H_2O} = \left[\frac{^2H/^{1}H}_{H_2O}/^{2}H/^{1}H_{VSMOW}\right] - 1. \]

Water samples for isotopic measurements at the University of Illinois at Chicago (UIC) were analyzed on a Picarro I2130-i analyzer by injecting 5 μl of water into a vaporizer at 110°C where they were evaporated and diluted with nitrogen. The vapor stream was then carried into the cavity of the laser absorption spectrometer. All samples were measured six times and the first three injections were rejected due to the influence of between-sample memory. Additional memory corrections were applied to the last three injections, which were then averaged to generate the reported value. Uncertainty is reported as the absolute range of the last three injections and is <0.2‰ and <1.0‰ for oxygen and hydrogen, respectively. Secondary water isotope standards were run intermittently to assess drift and three in-house span standards were analyzed at the end of each run to correct samples to the Vienna Standard Mean Ocean Water (VSMOW) scale (δ18O: −29.9, −15.9, and −0.9‰, and δ2H: −240, −122, and −0.2‰). The methods closely follow those previously described by Gupta et al. (2009), Brand et al. (2009), and Noone et al. (2013). At Idaho State University, water samples for isotope analysis were filtered through 0.45 μm filter and analyzed using a Thermo Scientific, High Temperature Conversion Elemental Analyzer (TC-EA) interfaced to a Delta V Advantage mass spectrometer through the ConFlo IV system. Isotope values of
$\delta^{2}$H and $\delta^{18}$O are reported as ‰ values relative to the VSMOW scale. Four in-house standards, which are directly calibrated against VSMOW2, SLAP2, and GISP, were used to create a two-point calibration curve to correct the raw data and to monitor the accuracy of the data. Precisions for both $\delta^{2}$H and $\delta^{18}$O values are better than ±2.0‰ and ±0.2‰, respectively.

Plant cellulose samples were measured at the University of Wyoming Stable Isotope Facility and values of $\delta^{18}$O$_{\text{cellulose}}$ were determined by pyrolysis at 1420°C and temperature conversion elemental analysis (TC-EA). Following CO and H$_2$ gas separation using a gas chromatographic column (GS) at 85°C, isotope ratios were measured by a coupled Thermo Scientific DeltaV Plus isotope ratio mass spectrometer (IRMS). If regularly spaced standard uncertainty was larger than 0.3‰, the samples were all re-analyzed until the 2-sigma expanded standard uncertainty of the result was <1‰. The isotopic composition is reported on the ‰ VSMOW scale such that standard reference material IAEA 601 (benzoic acid) and IAEA 602 (benzoic acid), respectively, are +23.3‰ and +71.4‰. At Iowa State University, plant cellulose samples were analyzed on a Finnigan MAT Delta Plus XL mass spectrometer in continuous flow mode connected to a Thermo-Chemical Elemental Analyzer. Reference standards [Sigma-Aldrich alpha cellulose [SAC], benzoic acid [IAEA-601], and sucrose [IAEA-CH-6]] were used for isotopic corrections, and to assign the data to the appropriate isotopic scale. For each isotope value given, two replicates were run for each sample. Analytical precision for the $\delta^{18}$O$_{\text{cellulose}}$ samples are ± <0.3‰. The analytical precision associated with the stable oxygen isotope analyses for the HT Fen core by Jones et al. (2014) was <0.6‰ and is expressed as one standard deviation from the mean based on the results from multiple ($n = 10$) analyses of a laboratory standard (benzoic acid, Fisher Scientific, Lot No 947459) conducted during the run of samples. Statistical relationships were determined using a t-test in SigmaPlot 13.0.

### Horse Trail Fen Core Reanalysis

The Horse Trail fen core (HTF) was previously analyzed on bulk peat ($\delta^{18}$O$_{\text{bulk}}$; Jones et al., 2014), and a select subset of samples, spanning a range of periods of differing species abundances, were analyzed for $\delta^{18}$O$_{\text{cellulose}}$ based on the cellulose extraction method of Wolfe et al. (2001) to determine the relationship between the $\delta^{18}$O$_{\text{bulk}}$ and $\delta^{18}$O$_{\text{cellulose}}$. A statistical regression of the data ($y = 1.0148x-0.0927$ (R$^2 = 0.68856$)) was then used to convert $\delta^{18}$O$_{\text{bulk}}$ to $\delta^{18}$O$_{\text{cellulose}}$ (Supplemental Tables 1, 2). This study utilized the plant macrofossil abundances of mosses and sedges identified by Jones et al. (2014; Figure 6) to scale respective offsets from their environmental water in the modern environment. Analytical errors were propagated by taking the square root of the squared standard deviation of the bulk peat-cellulose regression plus the squared standard deviation of the modern cellulose isotope to water isotope relationship. In each case, one water value at each location was related to the average of multiple moss or sedge values (Tables 2–4).

### RESULTS

#### Water

The average modern GF water $\delta^{18}$O value is $-14.1 \pm 0.1$‰ ($n = 5$) and $\sim 2$‰ statistically higher than the HTF value of $-16.2 \pm 0.1$‰ ($n = 4$; $P = <0.001$) (Tables 2, 3). HTF values from all sampling locations were nearly identical and plot near the intersection between the Local Evaporation Line (LEL), defined by the Kenai lakes measured for this study (Figure 1, Table 1), which are mostly located in the western lowlands (Figure 1), and the GMWL (Tables 1–3; Figure 2). The LEL-GMWL intersection approximates regional mean annual precipitation values (Anderson et al., 2016). The HTF environmental water $\delta^{18}$O values, which lie on the GMWL, closely approximate the $\delta^{18}$O values for mean annual precipitation in the western lowlands (Figure 2).

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**TABLE 1 | Kenai lake water isotope data.**

| Site name            | Date      | $\delta^{18}$O‰ (vs. VSMOW) | $\delta^{2}$H‰ (vs. VSMOW) | $d_\text{ex}$ | Lat (°N) | Long (°W) | Elev. (m a.s.l.) | Comments   |
|----------------------|-----------|-----------------------------|-----------------------------|--------------|----------|-----------|-----------------|------------|
| Horse Trail clearing | 7/21/2014 | -6.0                        | -80                         | -32.14       | 60.43066 | 150.91699 | 122             | fen pool    |
| Arc lake             | 7/24/2014 | -7.4                        | -83                         | -23.30       | 60.44990 | 151.10650 | 60              |            |
| Browns Lake          | 7/23/2014 | -7.1                        | -80                         | -22.88       | 60.48771 | 150.72458 | 88              |            |
| Headquarters Lake    | 7/24/2014 | -7.8                        | -82                         | -19.47       | 60.46341 | 151.07106 | 63              |            |
| Lower Ohmer Lake     | 7/20/2014 | -12.7                       | -108                        | -6.06        | 60.45223 | 150.31454 | 122             |            |
| Bear Mountain Lake   | 7/20/2014 | -13.3                       | -111                        | -4.06        | 60.45512 | 150.25237 | 247             |            |
| Upper Ohmer Lake     | 7/20/2014 | -13.8                       | -112                        | -1.93        | 60.45599 | 150.29001 | 148             |            |
| Bear Lake            | 7/22/2014 | -14.4                       | -109                        | 6.67         | 60.19139 | 149.35844 | 95              |            |
| Summit Lake          | 7/24/2014 | -17.8                       | -134                        | 7.84         | 60.63585 | 149.50703 | 401             |            |
| Skilak Lake          | 7/20/2014 | -16.7                       | -125                        | 8.54         | 60.43827 | 150.32146 | 63              | Glacial lake |
| Tern Lake            | 7/23/2014 | -17.1                       | -128                        | 8.82         | 60.53403 | 149.54771 | 206             |            |
| Kenai Lake           | 7/23/2014 | -17.0                       | -127                        | 9.33         | 60.41246 | 149.38496 | 140             | Glacial lake |
| Portage Lake         | 7/24/2014 | -15.1                       | -109                        | 11.59        | 60.78429 | 148.84002 | 27              |            |
| Browse Lake          | 9/5/16    | -13.9                       | -116                        | -4.46        | 60.56278 | 150.30847 | 131             |            |

1 Deuterium excess, $d_\text{ex} = 8 \times \delta^{18}$O - $\delta^{2}$H.
TABLE 2 | Goldfin (GF) plant cellulose and environmental water isotopes (sampled 9/1/2017).

| Type/location               | Species                        | $\delta^{18}$O$_{cellulose}$ (vs. VSMOW) | $\delta^{18}$O$_{water-cellulose}$* (vs. VSMOW) | $\delta^{2}$H$_{water}$ (vs. VSMOW) |
|-----------------------------|--------------------------------|----------------------------------------|-------------------------------------------------|-------------------------------------|
| **STREAM**                  |                                |                                        |                                                 |                                     |
| Plant–submerged             | Sphagnum subtintens leaves     | 20.1                                   | −34.1                                           |                                     |
| Plant–submerged             | Sphagnum subtintens stems      | 19.8                                   | −33.8                                           |                                     |
| Plant–submerged             | Carex spp.                     | 22.7                                   | −36.7                                           |                                     |
| Stream water                | –                              | −14.0                                  | −109.5                                          |                                     |
| Stream W of site            | –                              | −14.2                                  | −107.5                                          |                                     |
| **1m**                      |                                |                                        |                                                 |                                     |
| Water table                 | Campyllum stelatum (whole plant)| 19.8                                   | −32.4                                           |                                     |
| Water table                 | Sphagnum teres (whole plant)   | 19.2                                   | −31.9                                           |                                     |
| Water table                 | Carex spp.                     | 22.0                                   | −34.6                                           |                                     |
| Environmental water         | –                              | −12.6                                  | −104.9                                          |                                     |
| **20m**                     |                                |                                        |                                                 |                                     |
| Plant–submerged             | Calliergon stramineum          | 21.0                                   | −35.3                                           |                                     |
| Plant–submerged             | Calypogella spagnicola         | 20.9                                   | −35.2                                           |                                     |
| Plant–submerged             | Sphagnum russowii stems        | 20.2                                   | −34.5                                           |                                     |
| Plant–submerged             | Sphagnum russowii leaves       | 20.7                                   | −35.0                                           |                                     |
| Plant–submerged             | Carex spp.                     | 22.4                                   | −36.7                                           |                                     |
| Plant–hummock– 30 cm        | Sphagnum russowii stems        | 20.3                                   | −34.6                                           |                                     |
| Plant–hummock– 30 cm        | Sphagnum russowii leaves       | 20.3                                   | −34.6                                           |                                     |
| Plant–hummock– 30 cm        | Carex spp.                     | 24.0                                   | −38.3                                           |                                     |
| Environmental water         | –                              | −14.3                                  | −109.9                                          |                                     |
| **40m**                     |                                |                                        |                                                 |                                     |
| Plant–hummock– 10–20 cm     | Aulacornum palustre (whole plant)| 21.1                                   | −35.2                                           |                                     |
| Plant–hummock– 10–20 cm     | Sphagnum capillifolium leaves  | 21.3                                   | −35.4                                           |                                     |
| Plant–hummock– 10–20 cm     | Sphagnum capillifolium stems   | 21.2                                   | −35.2                                           |                                     |
| Plant–hummock– 10–20 cm     | Sphagnum russowii stems        | 19.9                                   | −34.0                                           |                                     |
| Plant–hummock– 10–20 cm     | Carex spp.                     | 22.2                                   | −36.3                                           |                                     |
| Plant–submerged             | Sphagnum teres leaves          | 19.6                                   | −33.6                                           |                                     |
| Plant–submerged             | Sphagnum teres stems           | 20.1                                   | −34.1                                           |                                     |
| Plant–submerged             | Carex spp.                     | 22.9                                   | −37.0                                           |                                     |
| Environmental water         | –                              | −14.1                                  | −109.5                                          |                                     |

*The difference between water isotope values and cellulose isotope values.

Plant $\delta^{18}$O$_{cellulose}$ Comparisons by Species and Plant Parts

Within-plant $\delta^{18}$O$_{cellulose}$ differences were determined from measurements of separated stems and leaves of the same species. On average, HTF Sphagnum leaf $\delta^{18}$O$_{cellulose}$ values were higher than stems by $1.3 \pm 0.6\%$ ($n = 6$ pairs). GF leaf $\delta^{18}$O$_{cellulose}$ average was $1.5 \pm 0.2\%$ higher than stems ($n = 5$ pairs; analytical error $0.1 \pm 0.06\%$; Figure 3). However, in neither location was the difference between leaf and stem $\delta^{18}$O$_{cellulose}$ values statistically significant ($P = 0.197$, $P = 0.450$, respectively). The only species for which Sphagnum stem values were higher than leaves was Sphagnum russowii at 40 m ($-0.4\%$) and Sphagnum teres at 20 m ($-0.3\%$). For samples where stems, leaves, and the corresponding whole plant $\delta^{18}$O$_{cellulose}$ values were measured ($n = 7$), whole plant values were both higher or lower by $<2\%$ than corresponding stems or leaves (Table 2), likely driven by the relative proportion of leaves to stems measured and the potential for debris to have been caught in Sphagnum leaves. At GF, brown moss $\delta^{18}$O$_{cellulose}$ values, which includes all non-Sphagnum peat mosses (Tables 2, 3), were consistently higher than Sphagnum by $0.5 \pm 0.1\%$ ($n = 3$ transect positions, incorporating 4 brown mosses and 12 Sphagnum mosses), but the relationship was not statistically significant ($P = 0.454$). At HTF, the relationship was opposite to that of GF ($-0.8 \pm 0.1\%$, $n = 3$ transect positions, incorporating 18 brown moss and 19 Sphagnum samples) and also not statistically significant ($P = 0.908$). In samples where stem and leaf $\delta^{18}$O$_{cellulose}$ were measured from the same sample (HTF only), brown moss leaves were slightly higher by a mean of $0.4 \pm 0.2\%$ than stems ($n = 5$ pairs), but the relationship was not statistically significant ($P = 0.373$). Carex spp. leaf $\delta^{18}$O$_{cellulose}$ values at all HTF and GF sampling locations were higher than their bryophytic (Sphagnum and brown moss) counterparts, but differences varied. However, the average difference between moss and sedge (Carex spp.) values at GF ($n = 16$ mosses, $n = 4$
| Type/location | Species | \( \delta^{18}O \)‰ (vs. VSMOW) | \( \delta^{18}O \)‰ water-cellulose | \( \delta^{2}H \)‰ (vs. VSMOW) |
|---------------|---------|------------------------------|--------------------------------|-----------------|
| 1m            | Plant   | Sphagnum teres (whole plant) | 16.2                          | −32.5           | −         |
|               | Plant   | Sphagnum teres stems         | 16.0                          | −32.3           | −         |
|               | Plant   | Calliergon stramineum (whole plant) | 15.7                          | −32.0           | −         |
|               | Plant   | Calliergon stramineum leaves | 16.6                          | −32.9           | −         |
|               | Plant   | Calliergon stramineum stems  | 16.4                          | −32.7           | −         |
|               | Plant   | Carex spp.                   | 26.3                          | −42.6           | −         |
|               | Plant   | Carex spp.                   | 26.3                          | −42.5           | −         |
|               | Plant   | Carex spp.                   | 25.9                          | −42.2           | −         |
|               | Plant   | Carex spp.                   | 26.1                          | −42.4           | −         |
|               | Environmental water | –                   | −16.3                         | −124            | −         |
| 10m           | Plant   | Calliergon giganteum leaves  | 18.4                          | −34.8           | −         |
|               | Plant   | Calliergon giganteum stems   | 18.2                          | −34.6           | −         |
|               | Plant   | Sphagnum teres (whole plant) | 13.9                          | −30.3           | −         |
|               | Plant   | Calliergon spp. (whole plant) | 16.3                          | −32.7           | −         |
|               | Plant   | Calliergon stramineum stems  | 16.9                          | −33.3           | −         |
|               | Plant   | Sphagnum teres stems         | 16.6                          | −33.0           | −         |
|               | Plant   | Sphagnum teres leaves        | 17.9                          | −34.3           | −         |
|               | Environmental water | –                   | −16.4                         | −125            | −         |
| 20m           | Plant   | Sphagnum teres leaves        | 16.2                          | −32.4           | −         |
|               | Plant   | Sphagnum teres stems         | 16.5                          | −32.7           | −         |
|               | Plant   | Sphagnum teres (whole plant) | 18.0                          | −34.1           | −         |
|               | Plant   | Bulk peat                    | 17.4                          | −33.5           | −         |
|               | Environmental water | –                   | −16.1                         | −123            | −         |
| 30m           | Plant   | Sphagnum spp. leaves         | 19.5                          | −35.6           | −         |
|               | Plant   | Aulacomnium palustre (whole plant) | 20.3                          | −36.4           | −         |
|               | Plant   | Paludella squarrosa (whole plant) | 16.3                          | −32.4           | −         |
|               | Plant   | Drepanocladus spp. (whole plant) | 16.6                          | −32.7           | −         |
|               | Plant   | Sphagnum spp. Stems          | 18.2                          | −34.3           | −         |
|               | Plant   | Paludella squarrosa (leaves) | 16.8                          | −32.9           | −         |
|               | Plant   | Paludella squarrosa (stems)  | 16.3                          | −32.4           | −         |
|               | Plant   | Sphagnum teres leaves        | 20.1                          | −36.2           | −         |
|               | Plant   | Sphagnum teres stems         | 18.3                          | −34.4           | −         |
|               | Plant   | Sphagnum russowi/subfulvum leaves | 19.2                          | −35.3           | −         |
|               | Plant   | Sphagnum russowi/subfulvum stems | 18.5                          | −34.6           | −         |
|               | Plant   | Calliergon stramineum leaves | 17.4                          | −33.5           | −         |
|               | Plant   | Calliergon stramineum stems  | 16.8                          | −32.9           | −         |
|               | Plant   | Calliergon stramineum (whole plant) | 16.6                          | −32.7           | −         |
|               | Plant   | Sphagnum teres (whole plant) | 18.5                          | −34.6           | −         |
|               | Plant   | Drepanocladus revolvens (leaves) | 17.3                          | −33.4           | −         |
|               | Plant   | Drepanocladus revolvens (stems) | 16.6                          | −32.7           | −         |
|               | Plant   | Sphagnum spp. Leaves         | 19.4                          | −35.5           | −         |
|               | Plant   | Paludella squarrosa (whole plant) | 16.9                          | −33.0           | −         |
|               | Plant   | Carex spp.                   | 23.4                          | −39.5           | −         |
|               | Plant   | Carex spp.                   | 23.5                          | −39.6           | −         |
|               | Plant   | Carex spp.                   | 22.7                          | −38.8           | −         |
|               | Plant   | Bulk peat                    | 18.7                          | −34.8           | −         |
|               | Environmental water | –                   | −16.1                         | −123            | −         |

(Continued)
TABLE 3 | Continued

| Type/location | Species                         | \( \delta^{18}O_{\text{cellulose}} \) (vs. VSMOW) | \( \delta^{18}O_{\text{water-cellulose}} \) (vs. VSMOW) |
|---------------|---------------------------------|-----------------------------------------------|--------------------------------------------------|
| 40m*          | Plant: Sphagnum rusowii stems   | 20.2                                          | –                                               |
|               | Plant: Sphagnum rusowii leaves  | 19.8                                          | –                                               |
|               | Plant: Sphagnum rusowii (whole plant) | 18.9                                       | –                                               |
|               | Plant: Bulk peat                | 20.8                                          | –                                               |

*Water-cell calculated with 30 m environmental water value.

The HTF transect was hypothesized to reflect a hydrologic gradient in terms of water table position that was assumed to be at stream level and progressively deeper below the fen surface with distance from the stream to 40 m, which lies at the upland forest edge. Although water isotope values were nearly identical at all transect locations, the bryophyte samples taken at greater distances from the stream showed progressively higher \( \delta^{18}O_{\text{cellulose}} \) values, ranging from \(-16\) to \(19\%)\) with progressively higher values at each location farther from the stream (Table 2). The difference in mean values between the lowest transect position (1 m; \(16.165 \pm 0.360\%\)) and 30 and 40 m (\(19.63 \pm 0.630\%)\) transect are statistically significant (\(P = 0.005\) and \(P < 0.001\), respectively). Of the more limited Carex spp. samples taken at HTF, the sample at 1-m (\(n = 4\)) distance was \(3\% \pm 0.6\) higher than the sample taken at 30-m distance (\(n = 3\)), both of which were higher by variable amounts (Figure 4, Table 3) than the moss samples from the respective transect positions, but additional samples are needed to verify this trend.

The relatively level surface of GF transect was not thought to provide a hydrologic gradient in terms of water table position, but the two sampling locations (referred to here as 20, 40 m) provided a comparison between submerged and sub-aerially exposed hummock plants (Table 2). The mean bryophyte \( \delta^{18}O_{\text{cellulose}} \) value differences between the 20-m site that was submerged (\(n = 4\)): \(19.8 \pm 0.378\%)\) and exposed (\(n = 2\)): \(20.3 \pm 0.044\%)\) were small (\(<0.5\%)\), and the relationship was not statistically significant (\(P = 0.204\)) (Figure 4; Table 2). The mean bryophyte \( \delta^{18}O_{\text{cellulose}} \) value differences between the 40-m site submerged (\(n = 2\)): \(19.8 \pm 0.38\%)\) and hummock (\(n = 4\)): \(20.87 \pm 0.65\%)\) was \(\sim0.5\%)\), and the difference was not statistically significant (\(P = 0.111\)). The differences among sedge \( \delta^{18}O_{\text{cellulose}} \) value values across all transect positions at GF was low (average = \(22.7 \pm 0.7\%)\), although they were on average higher than the difference amongst the bryophytes by \(2.5\%)\) (\(P < 0.001\)).

Mean \( \delta^{18}O_{\text{cellulose}} \) values were calculated for dominant plant types to evaluate the range of variability within a peatland and to better understand how to interpret changes in a peat core. The mean bryophyte \( \delta^{18}O_{\text{cellulose}} \) value at all HTF transect sites was \(17.6 \pm 1.2\%)\) (\(n = 35\); median \(16.9\)\%), with a range of \(15.7–20.3\%)\) (Table 3). The mean bryophyte \( \delta^{18}O_{\text{cellulose}} \) values at the GF sites was \(20.2 \pm 0.6\%)\) (median \(20.3\)\%), with a range of \(19.2–21.2\%)\). The difference in means between HTF and GF \( \delta^{18}O_{\text{cellulose}} \) values is statistically significant (\(P < 0.001\)). In contrast, the mean sedge \( \delta^{18}O_{\text{cellulose}} \) value at HTF is \(24.7 \pm 2.1\%)\) (\(n = 7\); median: \(25.9\)\%) is higher than the GF values of \(22.7 \pm 0.7\%)\) (median: \(22.6\)\%), a difference that is not great enough to reject the null hypothesis (\(P = 0.066\)).

**Plant-Environmental Water \( \delta^{18}O \) Comparisons (\( \Delta \delta^{18}O_{\text{cellulose-water}} \))**

At HTF, the difference between \( \delta^{18}O_{\text{cellulose}} \) values of individual moss species and the \( \delta^{18}O \) values of environmental waters (\( \Delta \delta^{18}O_{\text{cellulose-water}} \)) ranged from \(-30 \) to \(-37\%)\) with an average value of \(-33 \pm 3.2\%)\) (Tables 2, 4). \( \Delta \delta^{18}O_{\text{cellulose-water}} \) values increased from \(-31.2 \) to \(-35.8\%)\) with increasing distance from the stream because although environmental water \( \delta^{18}O \) values

![Figure 2](image-url)
were invariant, plant $\delta^{18}O_{\text{cellulose}}$ values increased. The two sedge samples had a larger $\Delta\delta^{18}O_{\text{cellulose}}-\text{water}$ value, with an average of $-41.1\%o$. The 1-m site had a larger $\Delta\delta^{18}O_{\text{cellulose}}-\text{water}$ value ($-42.4\%o$) than the 40-m site ($-39.4\%o$) (Table 5).

At GF, $\Delta\delta^{18}O_{\text{cellulose}}-\text{water}$ ranged from $-33$ to $-35\%o$ with an average value of $-33.9\%o \pm 1.2$ with larger values for sedge of $-36.4\%o \pm 1.2$. $\Delta\delta^{18}O_{\text{cellulose}}-\text{water}$ for submerged and hummock samples were similar at the 20-m site (0.1$\%o$), but at the 40-m site, the hummock samples were 1.3$\%o$ larger. In general, the range of $\Delta\delta^{18}O_{\text{cellulose}}-\text{water}$ values at GF was lower than at HTF.

To evaluate $\Delta\delta^{18}O_{\text{cellulose}}-\text{water}$ across as wide a range of water values as possible, data from St. Matthew Island is included (Table 4; Figure 5). Considering data from HTF, Goldfin, and St. Matthew Island provided a larger range of water $\delta^{18}O$ values from $-16.2$ to $-9.4\%o$ and $\delta^{18}O_{\text{cellulose}}$ values of 17.5 to $23.9\%o$, indicating a mean $\Delta\delta^{18}O_{\text{cellulose}}-\text{water}$ value for all sites of $-33.4\%o$. The linear regression using data from the three locations provides a slope of 0.79 with an $R^2$ of 0.811 (Tables 2–5; Figure 5).

### Species Effects in the HTF Bulk Peat $\delta^{18}O$ Core Record

While Jones et al. (2014) found no apparent relationship between plant macrofossil assemblage and $\delta^{18}O_{\text{bulk}}$ (Supplemental Figure 1), the results of the modern plant analysis showed a significant difference between $\delta^{18}O_{\text{cellulose}}$ moss and Carex spp. (Figure 4), which at HTF was $7.06 \pm 2.38\%o \ (P < 0.001)$. To illustrate how these adjustments translate to inferred environmental waters, samples that were primarily bryophytic were adjusted by $-33.9 \pm 1.88\%o$ according to the modern calibration, an adjustment supported by the cellulose fractionation factor determined by Sternberg and Ellsworth, (2011). In contrast, samples dominated by Carex spp. were adjusted by $-41.1 \pm 2.4\%o$, based on the HTF sedge-environmental water differences measured on the modern (Table 5; Figure 6). Samples whose bryophyte and Carex spp. macrofossil sum did not equal 100% were omitted so as to not introduce additional sources of error (Supplemental Table 2).

Species-weighted $\delta^{18}O$ calculations were made as follows:

Inferred water $\delta^{18}O = \delta^{18}O_{\text{moss-water}} \times (\%\text{moss abundance}) + \delta^{18}O_{\text{carex-water}} \times (\%\text{Carex abundance}), \ (2)$

Where $\delta^{18}O_{\text{moss-water}} = -33.9 \pm 1.88\%o$ and $\delta^{18}O_{\text{carex-water}} = -41.1 \pm 2.5\%o$ (Figure 6D). Errors were propagated by taking the square root of the sum of the squared sources of error (cellulose errors for mosses and sedges, error of water $\delta^{18}O$ values).

Although the plant species-adjusted $\delta^{18}O_{\text{cellulose}}$ values resulted in a shift toward higher values in the sedge-dominated intervals (Figure 6C), the inferred environmental water $\delta^{18}O$ value adjustment resulted in a downward shift to lower values.
DISCUSSION

Peatland Isotope Hydrology

The water isotope results indicate that the peatland surface water in both HTF and GF plot on the GMWL, suggesting that the water in these systems reflects the local precipitation signature with evaporative effects limited to pools of standing water. (Figure 2). Higher δ18O and δ2H values at GF (−14.1, −109.5‰, respectively), on the windward side of the Kenai Mountains compared to HTF (−16.2, −124‰, respectively), on the leeward side, is consistent with Rayleigh distillation effects (Dansgaard, 1964) across the Kenai Mountain barrier. As moisture is lifted over the Kenai Mountains, heavier isotopes of precipitation are preferentially rained out on the windward side, whereas on the leeward side precipitation values are relatively-isotopically enriched in light isotopes. Consequently, GF waters are 2‰ higher than HTF, and this relationship held for the average range of δ18O_cellulose values for the two peatland sites despite variability (Figure 2).

Table 4: St Matthew Island plant cellulose and environmental water isotopes (sampled 6/2016).

| Type/location | Species            | δ18O‰ (vs. VSMOW) | δ18O‰ water-cell | δ2H‰ (vs. VSMOW) |
|---------------|--------------------|-------------------|------------------|------------------|
| Plant         | Calliergon (whole plant) | 22.0              | −32.9            | −                |
| Environmental water | Stream | −12.6            | −                | −76.1            |
| Plant         | Carex spp.         | 22.8              | −33.7            | −                |
| Environmental water | Stream | −10.8            | −                | −72.5            |
| Plant         | Carex spp.         | 23.5              | −32.9            | −                |
| Environmental water | Standing peatland water | −9.4            | −                | −65.3            |
| Plant         | Drepanocladus unicus | 23.6              | −33.3            | −                |
| Plant         | Carex spp. (living) | 22.5              | −36.6            | −                |
| Water         | Standing peatland water | −9.7            | −                | −72.1            |
| Water         | Standing peatland water | −9.8            | −                | −72.4            |

*Environmental water value of −9.7‰.

Table 5: Plant and water isotope summary and statistics.

| Location               | n = | Mean δ18O‰ (vs. VSMOW) | SD  | Mean δ18O‰ water-plant (vs. VSMOW) | SD  |
|------------------------|-----|-------------------------|-----|------------------------------------|-----|
| HORSE TRAIL FEN        |     |                         |     |                                    |     |
| Water                  | 4   | −16.2                   | 1.0 |                                    |     |
| Moss                   | 37  | 17.6                    | 1.5 | −33.8                              | 1.8 |
| Carex spp.             | 7   | 24.9                    | 1.6 | −41.1                              | 1.9 |
| GOLDFIN BOG            |     |                         |     |                                    |     |
| Water                  | 5   | −13.9                   | 0.7 |                                    |     |
| Moss                   | 16  | 20.4                    | 0.6 | −34.2                              | 0.9 |
| Carex spp.             | 6   | 22.7                    | 0.7 | −36.6                              | 1.0 |
| ST. MATTHEW ISLAND     |     |                         |     |                                    |     |
| Water                  | 5   | −10.5                   | 1.3 |                                    |     |
| Moss                   | 2   | 22.8                    | 1.4 | −33.3                              | 1.9 |
| Carex spp.             | 3   | 23                      | 1.4 | −33.5                              | 1.4 |

(Figure 6D), amplifying periods of the record that were already indicated decreased δ18Obulk values in the unadjusted plots. This was particularly the case at ∼4–3 ka and 11.7–10.8 ka (Figures 6C,D), resulting in inferred environmental water values of −30 to −35‰ during the most extreme lows.
in species values (Tables 2, 3; Figures 4). GF water $^{18}$O values have a wider range; one sample was collected from a standing pool of water at the GF 1 m location had isotopic values that plot on the LEL. This indicates the isotopic effects of evaporation that are also reflected by relatively low $d_{18}O$ value of $-3.67$. In contrast, higher GF isotopic values for the stream and peatland surface waters at GF that plot near the GMWL (compared to HTF) are not affected by evaporation and more probably reflect the combined effects of the site’s location on the windward side of the Kenai Mountains, in the rain-out zone, at an elevation near sea level.

Although previous studies of $^{18}$O_cellulose peat records have focused on bogs, which are peatlands fed by precipitation and therefore presumptively more reflective of isotopically unmodified precipitation (Daley et al., 2010), the HTF water isotope values indicate that in Alaska fens can also host unmodified precipitation as a source, a conclusion that previous hydrologic studies support (Ford and Bedford, 1987; Reeve and Gracz, 2008). Water uptake in plants may also be more susceptible to evaporation in bogs during dry spells, which can lead to the preferential evaporation of light isotopes in water, complicating their interpretations. Similar to studies in lakes (Anderson et al., 2016), understanding of the hydrology of a system and its relation to isotopes of precipitation serves to constrain the interpretation of the sedimentary record. Both of the HTF and GF peatlands are fens that receive some fraction of their water from groundwater. The proximity of the fen water relative to the GMWL suggests the groundwater residence times (seasonal to annual) do not lead to significant evaporative evolution and thereby supports an interpretation of $^{18}$O_cellulose as a proxy for hydroclimate on decadal, centennial, and millennial timescales.

**Intra-Plant Part and Species Effects**

Differences in $^{18}$O_cellulose values between *Sphagnum* stems and branches have been reported (Moschen et al., 2009; Kaislahti Tillman et al., 2010). However, because of the difficulty in separating individual stems and branches while omitting leaves, we only evaluated collections of stems and leaves and found that the differences in $^{18}$O_cellulose values were negligible ($<1\%$) relative to the 5–10% range of observed $^{18}$O_cellulose values in peat core records, similar to results reported by Moschen et al. (2009). Also similar to previous findings (Moschen et al., 2009), the $^{18}$O_cellulose values of *Sphagnum* leaves and all brown moss samples were higher than those in stems in most instances, suggesting a difference in the biochemical synthesis of cellulose in stems vs. leaves and branches. In the few cases where the relationship was reversed, the reversal could be explained by detrital organic matter caught in *Sphagnum* leaves, but this is untested. In general, the intra-plant $^{18}$O_cellulose value differences have little effect on interpretation of larger isotopic shifts in a peat core record.

Different moss-species $^{18}$O_cellulose values across the HTF transect varied by $\sim5\%$ (Figure 4) and distinct patterns emerged among species that reflect local hydrologic conditions (drier vs. wetter). Generally, submerged moss species had lower $^{18}$O_cellulose values relative to those species on hummocks, and values increased from wet to the dry locations with increased distance from streams. Other modern calibration studies have found a strong relationship between relative humidity near the moss growth position and the respective isotopic composition of the cellulose (Loader et al., 2016). The Loader et al. (2016) study recorded relative humidity values between 50 and 70% some evaporative enrichment was observed in subaerially exposed peatland mosses. However, a strong relationship between the environmental water and moss-$^{18}$O_cellulose values indicated that the factors influencing isotopic alteration during the growing season remain stable influences on the $^{18}$O_cellulose values of *Sphagnum*, consistent with findings from this study.

Sedge (Carex spp.) $^{18}$O_cellulose values measured here were always higher ($\sim2.5\%$) than mosses suggesting that while mosses passively uptake water and are governed by the same biochemical synthesis of oxygen from water into cellulose, the $^{18}$O_cellulose values of vascular plants are enriched in heavy isotopes by stomatal regulation of water loss (Figure 4; Ménot-Combes et al., 2002; Zanazzi and Mora, 2005; Amesbury et al., 2015). Further, Amesbury et al. (2015) found a consistent offset of 3% between root-associated water and environmental water in a species of rush, indicating that fractionation during uptake of water in vascular plants also can occur, further contributing to a larger offset between sedges and mosses. We observed no clear trend with respect to sedge and water table position (Figure 4), although the sample size was small ($n = 6$ transect positions). The lack of a trend could be the result of both stomatal regulation of water loss and to the depth at which sedges obtain their water, which could have a different isotopic signature from surface water.

![Figure 5](image-url)
A linear regression between the $\delta^{18}O_{\text{cellulose}}$ values of aquatic moss and its environmental water with a slope of 1:1 suggests $\delta^{18}O_{\text{cellulose}}$ values of aquatic moss are governed by the $\delta^{18}O$ values of environmental water (Zhu et al., 2014). In laboratory experiments where aquatic mosses were not subject to evapotranspiration, the environmental water and cellulose $\delta^{18}O$ slopes range between 0.78 and 0.97 with $R^2$ values between 0.88 and 0.998 (Sauer et al., 2001; Mayr et al., 2013; Zhu et al., 2014). By comparison, the $\delta^{18}O$ regression between our Alaska terrestrial moss and peatland environmental waters in this study...
has a slope of 0.79 and $R^2$ of 0.81 (Figure 5). Although the terrestrial moss regression does not overlap with the aquatic moss regression, likely because of different relationships between peat water and plants compared with lake water and aquatic moss (Loader et al., 2016). The relationship between Alaska peatland moss and environmental water documented here supports inferred environmental-water interpretations of peat core records. The offset of moss $\delta^{18}O_{\text{cellulose}}$ to the $\delta^{18}O_{\text{water}}$ is on average $-33\%o$, which is higher than the previously assumed constant offset of $-27\%o$ identified by DeNiro and Epstein (1981) but is in agreement with a latitudinal-temperature effect identified by Sternberg and Ellsworth (2011), where lower MAAT corresponds to lower $\delta^{18}O_{\text{cellulose}}$.

The results of this study suggest that at a given site, mosses do not differ significantly from one another but that significant difference exists between bryophytic (moss) samples and graminoid (sedge: Carex spp.) samples, likely owing to the ecophysiological differences between vascular and nonvascular plants. This suggests that $\delta^{18}O$ interpretations of peat cores should have modern site-level information on species differences and that these differences need to be accounted for when peat composition transitions from vascular to nonvascular.

Reanalysis of the HTF $\delta^{18}O$ Record
The results of this study show significant species difference between mosses and Carex spp. (Figure 4). Although comparisons between specific peat macrofossil abundances and unaltered $\delta^{18}O_{\text{bulk}}$ revealed no substantial relationship (Supplemental Figure 1), the $+2.5\%o$ difference between sedge and moss $\delta^{18}O_{\text{cellulose}}$ values and $-8\%o$ between sedge and moss inferred-water $\delta^{18}O$ values suggests variations in their respective abundance could lead to substantial deviations from the $\delta^{18}O_{\text{bulk}}$ record (Figure 6). This could have implications for the interpretation of the $\delta^{18}O_{\text{bulk}}$ record that assumed no differences in the way in which vascular and non-vascular plants fractionate water (Jones et al., 2014). Adjusting the $\delta^{18}O_{\text{cellulose}}$ values for abundance of Carex spp. and moss, as outlined above, slightly reduced the overall $\delta^{18}O_{\text{cellulose}}$ range of variability of the record compared to the single-source adjusted values (Figure 6C), and the change in the absolute value of the isotopic signature falls within the errors of each respective curve. However, a larger opposite effect occurs when the species-adjusted values are further adjusted to infer environmental water $\delta^{18}O$ values. Inferred-water $\delta^{18}O$ values are substantially lower during periods of sedge dominance (Figure 6D).

There are additional alternative explanations for the shifts in the HTF $\delta^{18}O_{\text{bulk}}$ values. The unaltered $\delta^{18}O_{\text{bulk}}$ analysis on the entire peat core and subsequent cellulose extraction applied to selected peat samples (Jones et al., 2014) did not remove clay minerals and chitin impurities (Wolfe et al., 2007), which have been shown to be a source of significant contamination in sediment-cellulose $\delta^{18}O$ measurements (Wissel et al., 2008) and cannot be ruled out as contributing to the unusually low bulk peat and inferred water values. The CUAM method has been shown to produce the purest cellulose and higher $\delta^{18}O_{\text{cellulose}}$ values would be expected because they are higher than that of other organic compounds (Zhu et al., 2014). Although Jones et al. (2014) found a compelling relationship between the $\delta^{18}O_{\text{bulk}}$ and $\delta^{18}O_{\text{cellulose}}$, standard deviations are as large as 2.5%o and values declines correspond with changes in %LOI and mineral content (Supplemental Figure 2), which could suggest influence of the mineral component on the $\delta^{18}O$, but the change in %LOI also implies a changing depositional environment and/or climate change. Additional analyses are required to more precisely determine the effect of cellulose extraction methods on the HTF $\delta^{18}O_{\text{bulk}}$ values (Figures 6C,D).

A prominent period when shifts to greater Carex spp. abundances substantially alter the interpretation of the unaltered $\delta^{18}O_{\text{bulk}}$ record occurs between $\sim 11.7$ and $10.8$ ka (Figures 6B,D). The unadjusted record shows values similar to the rest of the early Holocene (Figure 6C), but the species-adjusted inferred $\delta^{18}O$ water values are substantially lower at the beginning of the Holocene ($\sim -27\%o$; 11.7–10.8 ka). The resulting lower values, combined with the species assemblage change, suggest that a previously unrecognized hydroclimatic shift occurred between the early Preboreal (11.7–10.8 ka) and the subsequent early Holocene period (10.8–9 ka). Lower environmental water $\delta^{18}O$ values during this period could reflect the influence of rising sea levels and more extensive sea ice conditions (e.g., Gaglioti et al., 2017). Other mechanisms could include altered atmospheric circulation dynamics, including changing seasonality and longer transport over the continent: lower values are consistent with greater winter precipitation influence on the groundwater signature, as winter precipitation $\delta^{18}O$ values are lower than summer values (IAEA/WMO, 2001; Bailey et al., 2015).

Another period of Carex spp. dominance occurs between $\sim 4$ and $3$ ka, when rising species-adjusted $\delta^{18}O_{\text{cellulose}}$ are higher than unadjusted bulk $\delta^{18}O_{\text{cellulose}}$ values (Figure 6C), leading to inferred water values (Figure 6D) that are the lowest of the entire record ($-35$ to $-32\%o$). In this case, the recorded declines are enhanced by correcting for species, but the overall trend is unchanged. This interval at HTF coincides with a notable decline in organic matter (Supplemental Figure 2), due to an increase in silt percentages that was previously interpreted to reflect a combination of climate-induced peat decomposition and silt deposition, but may reflect the influence of more negative clay mineral water isotope values; bulk peat values may to some degree represent changes in clay mineral abundance.
−30‰. They are unrealistic when compared to the closest (~700 km) modern global network of isotopes in precipitation (GNIP) data from Bethel, Alaska, and Whitehorse, Yukon Territory (IAEA/WMO, 2001), and reported values of last glacial maximum (LGM) water in interior Alaska (−28‰; Lachniet et al., 2012). Notwithstanding the unrealistic absolute values of the species-adjusted inferred water values, the declining trends could suggest declining temperatures, a shift toward winter-dominant precipitation, or more northerly derived moisture during this time interval. It is also possible that the large decrease between ~4 and 3 ka could reflect water from a non-precipitation source, such as an outburst flooding event driven by glacial meltwater, intruded into the peatland, although this hypothesis remains untested. Interestingly, after accounting for age model uncertainty, the timing roughly corresponds (~3.3–2.5 ka) to a similar isotope excursion (Wooller et al., 2012) and flood deposits along the Tanana River in Interior Alaska (Sattler and Jordan, 1987; Mason and Begét, 1991) and from the central Brooks Range (Hamilton, 1981), suggesting widespread flooding. Assuming that at least some of the decrease is related to the plant's environmental water and a non-cellulose source, another hypothesis for the earlier anomalously low δ18O values is that lower Pleistocene-aged meltwater could explain the low δ18O values. Given the proximity of the HTF site to the margin of the ice sheet during the early Holocene, it is conceivable that the low inferred water δ18O values from 11.7 to 10.8 ka were derived from glacial meltwater, as the ice sheet was rapidly melting under a warming climate (Kaufman et al., 2004).

HTF Paleo-Hydroclimate Relative to Modern Results
The HTF δ18Ocellulose values have a mean of 11.9‰, which falls well below the mean δ18Ocellulose estimates (17.5‰ ± 1.4) at HTF today. Furthermore, the mean δ18O water value at HTF today (−16‰) falls well above the Holocene and late-glacial average values (−23.6‰, adjusted; −20.4‰, unadjusted; Figure 6D). Unless the 6–7‰ discrepancies are caused by differences in extraction methods and sample purity, the disparity between the modern and paleo δ18Ocellulose values suggests that (1) the sampling has not fully encapsulate the range of variables affecting the δ18Ocellulose values through time, such as large changes in temperature (Sternberg and Ellsworth, 2011), and (2) that modern hydrologic conditions on the Kenai are largely unlike the conditions of the past 14 ky, with notable exceptions for two periods at ~9.6 ka and ~ 1–1.5 k (Figure 6D). Higher water isotope values generally suggest some combination of warmer temperatures, more proximal moisture sources or those originating from primary moisture sources to the south (i.e., the Gulf of Alaska and North Pacific), or a shift in predominance to summer precipitation. Evidence from other studies has shown intensification of the Aleutian Low over the last 150 years (Anderson et al., 2016; Osterberg et al., 2017), which is supported by the increasing values toward present in the HTF record and the higher modern values reported here.

CONCLUSIONS
In this study, we have determined (a) that Kenai fen water reflects modern precipitation, (b) the relationship between δ18O values of moss and Carex spp. cellulose with environmental water, (c) the range of δ18O values among species of moss and sedge commonly found within Alaskan peatlands, and (d) moss and sedge species separation is necessary for more accurate inferences of variability in environmental water δ18O values from Alaska peat records.

The use of δ18Ocellulose from peat cores has the potential to dramatically improve the spatio-temporal resolution in paleoclimate studies in regions where peatlands are abundant, such as vast areas of the boreal and Arctic. However, the results of this study show that, similar to lake studies, efforts to better characterized peatland hydrology with modern water isotope sampling serves to improve interpretations of paleoclimate studies. Furthermore, prominent shifts in vegetation assemblages, particularly shifts between moss (nonvascular)- and Carex spp. (vascular)-plant dominated intervals common in fens, necessitate modern studies of the relationships between the δ18O values of plants and environmental waters. Fens are the dominant wetland type in Alaska and circumpolar boreal peatlands, and this study has served to highlight their potential as hydroclimatic archives, particularly spanning periods of substantial deglacial sea level and climate change.

AUTHOR CONTRIBUTIONS
MJ conceived of the study, designed the research, analyzed data, and wrote the manuscript. LA helped design the research, analyze data, and helped write the manuscript. KK, BN, and VI processed samples and created protocol to extract cellulose. MW helped write the manuscript and analyze data. CJ separated moss species and plant parts.

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SUPPLEMENTARY MATERIAL
The Supplementary Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/feart.2019.00025/full#supplementary-material
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Conflict of Interest Statement: The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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