Modern calibration of *Poa flabellata* (Tussac grass) as a new paleoclimate proxy in the South Atlantic

Dulcinea V. Groff\(^1,2\), David G. Williams\(^3\), Jacquelyn L. Gill\(^1,2\)

\(^1\)Climate Change Institute, University of Maine, Orono, ME 04469, USA
\(^2\)School of Biology and Ecology, University of Maine, Orono, Me 04469, USA
\(^3\)Department of Botany, University of Wyoming, Laramie, WY 82071, USA

Correspondence to: Dulcinea V. Groff (dulcineavgroff@gmail.com)

Abstract. Terrestrial paleoclimate records are rare in the South Atlantic, limiting opportunities to provide a prehistoric context for current global changes. The tussock grass, *Poa flabellata*, grows abundantly along the coasts of the Falkland Islands and other sub-Antarctic islands. It forms extensive peat records, providing a promising opportunity to reconstruct high-resolution regional climate records. The isotopic composition of leaf and root tissues deposited in these peats has the potential to record variation in precipitation, temperature, and relative humidity over time, but these relationships are unknown for *P. flabellata*. Here, we investigate the isotopic composition of *P. flabellata* plants and precipitation and explore seasonal relationships with temperature and humidity across 4 study locations in the Falkland Islands. We reveal that inter-seasonal differences in carbon and oxygen stable isotopes of leaf \(\alpha\)-cellulose of living *P. flabellata* significantly correlated with monthly mean temperature and relative humidity. The carbon isotope composition of leaf \(\alpha\)-cellulose (\(\delta^{13}C_{\text{leaf}}\)) records the balance of CO\(_2\) supply through stomata and the demand by photosynthesis. The positive correlation between \(\delta^{13}C_{\text{leaf}}\) and temperature and negative correlation between \(\delta^{13}C_{\text{leaf}}\) and relative humidity suggest that photosynthetic demand for CO\(_2\) relative to stomatal supply is enhanced when conditions are warm and dry. Further, the positive correlation between \(\delta^{13}C_{\text{leaf}}\) and \(\delta^{18}O_{\text{leaf}}\) (\(r = 0.88, p < 0.001, n = 24\)) indicates that stomatal closure during warm dry periods explain seasonal variation in \(\delta^{13}C_{\text{leaf}}\). We observed significant differences between winter and summer seasons for both \(\delta^{18}O_{\text{leaf}}\) and \(\delta^{13}C_{\text{leaf}}\), and among study locations for \(\delta^{18}O_{\text{leaf}}\), but not \(\delta^{13}C_{\text{leaf}}\). \(\delta^{18}O\) values of monthly composite precipitation did not differ between seasons or among study locations, yet is characteristic of the latitudinal origin of storm tracks and seasonal winds. The weak correlation between \(\delta^{18}O\) in monthly composite precipitation and \(\delta^{18}O_{\text{leaf}}\) further suggests that relative humidity is the main driver of the \(\delta^{18}O_{\text{leaf}}\). The oxygen isotopes in root \(\alpha\)-cellulose did not reflect, or only partially reflected (at one study location), the \(\delta^{18}O\) in precipitation. Overall, this study supports the use of peat records formed by *P. flabellata* to fill in a significant gap in our knowledge of the long-term trends in Southern Hemisphere climate dynamics.

1 Introduction

The high latitude environments of the South Atlantic are changing rapidly. Over the last century, mean annual temperature in the Falkland Islands (Fig. 1A) has increased by 0.5 °C (Lister and Jones, 2015). This warming has corresponded with an intensification and poleward shift of the southern westerly winds and aridification (Gillett et al., 2008; Thompson and Solomon, 2002; Villalba et al., 2012). These changes are already altering the distribution of
marine animals in the Southern Ocean (Weimerskirch et al., 2012), and warming of the western South Atlantic is projected to alter the distribution of island plants as well (Jones et al., 2013; Upson et al., 2016). The inconsistency of meteorological measurements from the Falkland Islands dating back to 1874 (Lister and Jones, 2015) means we lack critical information on the long-term patterns and whether these are novel conditions. Paleoclimatological archives, such as high-resolution lake sediments and tree rings, can provide useful long-term records documenting and quantifying climate change (Dietl et al., 2015; Dietl and Flessa, 2011; Willis et al., 2010), but such records are lacking for the South Atlantic. The absence of trees and deep lakes across many sub-Antarctic islands especially limits high-resolution, independent paleoclimate reconstructions, which are essential for detecting past abrupt climate change. However, many sub-Antarctic islands support widespread communities of peat-forming C3 tussock grasses (*Poa flabellata*), which provide important habitat and shelter for breeding marine animals such as seals and seabirds. Peat records formed by *P. flabellata* present a promising avenue for paleoclimate reconstructions; peatland vegetation has been used to reconstruct hydrological change and temperature in mid- to high latitudes (Amesbury et al., 2015; Chambers et al., 2012; Pendall et al., 2001).

*P. flabellata* grasslands in the South Atlantic accumulate substantial amounts of peat (Smith and Clymo, 1984), and have the highest carbon accumulation rates of any peatland globally (Payne et al., 2019). Endemic to the South Atlantic, *P. flabellata* only occurs on Tierra del Fuego, the Falkland Islands, Gough Island, and South Georgia. *P. flabellata* grasslands were once widespread throughout the Falklands, but are now greatly reduced because of land-use change and introduced grazers (Strange et al., 1988; Wilson et al., 1993). The term “tussock” is used to describe the clumping growth form of *P. flabellata*, while the species itself is commonly known as “tussac.”

Several factors support the utility of *P. flabellata* peats as a paleoclimate proxy. *P. flabellata* peatlands are formed by a nearly single-species community of *P. flabellata*, which allow very little light or space for other plants to co-occur in the absence of disturbance. Tillers of *P. flabellata* grow on top of a pedestal of decaying roots and leaves (called a “bog”) that can reach 4 meters high (Fig. 2A; (Smith and Clymo, 1984); mature plants thus likely primarily use water from precipitation, as they are not rooted in the soil directly. (Smith and Prince, 1985) established radiocarbon (14C) dates for a *P. flabellata* pedestal and estimated an age of 250 to 330 years. *P. flabellata* grass forms extensive peat deposits of up to 13.3-m deep, with carbon accumulation rates of 139 g C m⁻² yr⁻¹ (Payne et al., 2019; Smith and Clymo, 1984), far greater than peatlands of similar latitude in the Northern Hemisphere (18.6 g C m⁻² yr⁻¹), the tropics (12.8 g C m⁻² yr⁻¹) or Patagonia (22 g C m⁻² yr⁻¹) (Yu et al., 2010). Subfossil *P. flabellata* leaves are abundant in these peats (Fig. 2E), and readily can be separated from root subfossils. Having the highest accumulation rate of any global peatlands, *P. flabellata* peat is ideal for high-resolution climate reconstructions. Basal 14C radiocarbon dates indicate most *P. flabellata* peatlands initiated between ~ 12,500 and 5,500 14C years (Groff, 2018; Payne et al., 2019; Smith and Clymo, 1984).

Grasses exhibiting the tussock growth form often have evergreen leaves and exhibit a profligate/opportunistic water use strategy, due to the high evaporative conditions and pulses of water availability in semi-arid habitats (Moreno-Gutiérrez et al., 2012; Sala et al., 1989; Schwinning and Ehleringer, 2001). The growth phenology of *P. flabellata* is such that it mainly increases in height in summer (39 cm per year) and in winter an increase in basal area occurs with...
the production of new tillers are produced at the base of a "bog" (Stanworth and Upson, 2013). The climate signal recorded in the cellulose of plant tissues (roots, shoots, and leaves) is deciphered using stable isotopes δ¹⁸O, δD (Araguás-Araguás et al., 2000) and δ¹³C. Carbon and oxygen stable isotope ratios record species’ water-use strategies in water-limited environments because of physiological responses such as stomatal conductance and assimilation rates (Farquhar and Sharkey, 1982; Moreno-Gutiérrez et al., 2012). Tussock grasses typically occur in water-limited environments where low water-use efficiency and high stomatal conductance are common functional traits that allow tussock grasses to take advantage of pulses of water (Moreno-Gutiérrez et al., 2012). Correlations between δ¹⁸O of plant cellulose and air temperature and humidity provide information about environmental conditions in the season the cellulose tissue is formed. The δ¹⁸O of leaf water is a primary driver of δ¹⁸O in leaf cellulose, and is influenced by the δ¹⁸O value of plant source water, temperature and humidity (Helliker and Ehleringer, 2002; Roden and Ehleringer, 1999). δ¹⁸O of source water often is correlated with temperature of the environment (Libby et al., 1976). Apart from source water, δ¹⁸O of cellulose also can be influenced by internal exchange among organic molecules and other plant waters (Sternberg et al., 1986). The δ¹³C value of leaf biomass in C3 plants records δ¹³C of source CO₂ and the expression of fractionation effects associated with CO₂ diffusion into and through leaf tissue and carboxylation (Farquhar et al., 1982). The net discrimination against ¹³C during photosynthesis is driven by changes in the supply of CO₂ through stomatal pores and demand for CO₂ by photosynthetic biochemistry (Cernusak et al., 2009; Ferrio and Voltas, 2005); Fig. 2B). The δ¹³C value of roots tends to be 1-3 per mil higher than that of leaves due to a number of post-photosynthetic biochemical fractionations and C allocation pathways (Cernusak et al., 2009).

Plant species vary in the way they isotopically record precipitation and temperature; therefore, peat comprised of a single species is more desirable over a mixture of species (van Geel and Middeldorp, 1988). To test the potential of *P. flabellata* peatlands as a paleoclimate proxy, we conducted a modern calibration study. We measured oxygen (δ¹⁸O) and carbon (δ¹³C) stable isotopes from living *P. flabellata* leaf tissues (α-cellulose) collected monthly at four sites across the Falklands (51° S, 59° W; Fig. 1). We aim to improve our understanding of Southern Hemisphere westerly wind dynamics with a new paleoclimate proxy that leverages the unique properties of *P. flabellata*.

2 Materials and Methods

2.1 Study location description

The Falklands (Fig. 1A) are located approximately 500 km east of southern South America, between 51°0.5' S to 52°28.0' S and 61°22.0' W to 57°40.5’ W. The cool-temperate (mean temperature: January 9.4 °C and July 2.2 °C) climate of the Falklands is driven by the cold Antarctic Circumpolar Current, the waters surrounding the Antarctic Peninsula, the Falklands Current, and the Andes of southern Patagonia to the west (Turner and Pendelbury, 2000). The persistent winds of the southwesterly wind belt average 8.5 m s⁻¹, with gale force winds averaging 70 days per year and annual precipitation generally ranges between 400 to 600 mm (Jones et al., 2013; Lister and Jones, 2015). Study sites were selected to reflect 1) climatic diversity, and 2) the availability of volunteers to collect monthly samples.
for one year. We ultimately selected four sites (Fig. 1B-C): Bleaker Island, Cape Dolphin, Surf Bay, and West Point Island.

2.2 Precipitation, temperature and relative humidity

Precipitation was collected at each site using a Palmex monthly composite collection sampler (Palmex d.o.o., Zagreb, Croatia). The Palmex collector is designed to prevent evaporation and evaporative enrichment of $^{18}$O in precipitation samples without the use of paraffin oil (Gröning et al., 2012), and has been recommended by the Global Network of Isotopes in Precipitation (GNIP). Samples were shipped to the University of Maine prior to analysis. Oxygen ($\delta^{18}$O) and hydrogen ($\delta^D$) stable isotope ratios of water samples were measured at the University of Wyoming Stable Isotope Facility (UWSIF) using a high-temperature conversion elemental analyzer (TCEA) connected to a Thermo Scientific Delta V Plus that is run in continuous flow mode via a ConFlo IV. The technique used injections of 1 µl of water into the TCEA column filled with glassy carbon heated to 1420 °C. Precipitation samples were extracted using cryogenic vacuum distillation (Ehleringer and Osmond, 1989) prior to TCEA analysis to remove aeolian debris, including marine salts. Internal QA/QC working standards calibrated against IAEA international standards Vienna Standard Mean Ocean Water (VSMOW) and Standard Light Antarctic Precipitation (SLAP) and spanning the range of measured values in our study were analyzed with each batch of samples with analytical precision typically better than 0.3 and 2.5 ‰ for $\delta^{18}$O and $\delta^D$, respectively. Isotope values are reported with respect to VSMOW in parts per thousand (per mil, ‰). Explanations of methods for daily average temperature and relative humidity measurements, as well as seasonal wind speed, wind direction, and back trajectory models to determine origins of air masses are found in the Supplemental Text 1.

2.3 Poa flabellata field collection and cellulose extraction, and isotope analyses

Poa flabellata plants were collected at the start of each month at each site from October 2015 through September 2016, from relatively uniform habitats that were undisturbed by grazing or tilling. Up to six P. flabellata plant tillers (leaves, stem, and roots) were collected near each of the four stations each month. Whole plants were stored in paper envelopes stored in a cool, dark, dry location until frozen. Samples collected between September 2015 to February 2016 were frozen in February 2016 and samples collected in March 2016 to August/September 2016 were frozen in August/September 2016. Samples were frozen for eight days at the Falkland Islands Department of Agriculture. For leaf material, the inner developing (youngest) leaves were collected. There was no indication that leaves were morphologically different between summer and winter. Only coarse roots were used and fine roots were excluded. Variation in environmental conditions during the growth of a leaf blade can lead to isotopic variations along the gradient of a single leaf as has been shown with $\delta^{18}$O of cellulose (Helliker and Ehleringer, 2000, 2002); therefore, whole-leaf plant samples were homogenized by drying at 50 °C and pulverizing using a Retsch ball mill at the University of Maine. For each sample, we used 20 mg of pulverized and homogenized leaf or root material for extraction and purification of $\alpha$-cellulose, following an adapted procedure of (Brendel et al., 2000). Samples were vortexed throughout extraction and purification for homogenization and were visually inspected for purity. Further indicators of purity include undetectable amounts of % nitrogen, and analysis of % carbon in cellulose. As an internal
quality control, one leaf sample was selected for extraction and purification of α-cellulose throughout the sample processing in batches of 10 to 12 samples. The δ¹³C and δ¹⁸O of leaf cellulose for the internal quality control samples varied by ± < 0.1 ‰ and ± 0.3 ‰, respectively.

Oxygen (δ¹⁸O) and carbon (δ¹³C) stable isotope ratios of α-cellulose samples were measured at the University of Wyoming Stable Isotope Facility (UWSIF). Oxygen was analyzed using a TCEA coupled to a Thermo Delta V IRMS; δ¹⁸O values are expressed with regard to VSMOW in parts per thousand (per mil) (Craig, 1961; Gonfiantini, 1978). Values were normalized to the VSMOW scale using USGS-42 and IAEA-601 cellulose quality control standard reference materials for oxygen isotopic composition. Analytical precision was ± 0.3 ‰ for δ¹⁸O based on repeated analysis of internal standards, and samples loaded into silver capsules had weights ranging from 0.206 to 0.289 mg. Carbon isotope composition of the cellulose samples were determined using a Costech 4010 Elemental Analyzer coupled to a Thermo Delta Plus XP-IRMS; units are expressed with regard to VPDB in parts per thousand (per mil). Analytical precision was ± 0.1 ‰ for δ¹³C based on repeated internal standards. Quality control standard reference material for carbon isotopic composition included USGS-40 glutamic acid, USGS-41 glutamic acid, and internal UWSIF α-cellulose. Carbon sample weights ranged from 1.937 to 2.194 mg and were loaded into tin capsules.

2.4 Statistical analysis

For both δ¹⁸O and δ¹³C, we analyzed the average of three to four plant leaf samples per month for summer (DJF) and winter (JJA) season at each of the four sites, and the average of up to eight plant root samples (Supplementary Data 1 and 2). We used Pearson’s correlation coefficient, r, to detect associations between δ¹⁸O of cellulose and precipitation samples to test whether δ¹⁸O samples reflects the isotopic value of precipitation. Using Pearson’s correlation coefficient, we tested for a relationship between δ¹³C and δ¹⁸O values of cellulose, temperature, and relative humidity. We tested for a significant difference between summer and winter δ¹⁸O and δD in precipitation using a t-test (n = 24). A separate one-way analysis of variance (ANOVA) to test for significant differences among sites included δ¹⁸O and δD in precipitation samples from the entire year (n = 47).

A two-way ANOVA compared the main effects of season (summer vs. winter) and the four study locations on the carbon and oxygen stable isotopes of α-cellulose of P. flabellata leaves and roots grown in the summer versus winter, followed by a post hoc test (Tukey’s multiple comparison of means). P-values < 0.05 are considered significant. Descriptive and multivariate analyses were conducted with SigmaPlot 12.5.

3 Results

3.1 Environmental measurements

Across all sites, summer (DJF) daily average temperatures ranged from 3.5 °C to 15.6 °C (mean = 10.0 °C) and relative humidity ranged from 64.2 % to 98.1 % (mean = 81.1 %). Winter (JJA) daily average temperatures ranged from -1.8 °C to 7.6 °C (mean = 3.7 °C), and relative humidity ranged from 73.6 % to 100 % (mean 94.3 %). Seasonal temperature
(°C) and relative humidity (%) minimum and maximum ranges for individual study locations are found in Table S1. Between study locations, the daily average temperatures over the year ($F_{(3, 44)} = 0.316, p = 0.813$, Fig. 2C) and relative humidity were not significantly different ($F_{(3, 44)} = 0.674, p = 0.573$, Fig. 2D).

3.2 Wind

The wind rose (Fig. S2) shows that winter winds at Bleaker Island primarily blew from the west and northwest. In winter (JJA), two spokes in the west and NNW direction comprise >30% of the total recorded 15-minute wind directions. In summer (DJF), three spokes in the west, WSW, and SW directions comprise >45% of all 15-minute wind directions. The wind rarely blew from the east, SE, or north. Examining winds from the west in winter, >10% of wind speeds recorded were between 5 and 10 m s$^{-1}$, and the frequency of strongest winds came from the NNE. In summer, >20% winds from the SW were between 5 and 10 m s$^{-1}$, and there was a higher frequency of 10-15 m s$^{-1}$ wind speeds than in winter. Seasonal wind variation deviated from the long-term average (1979-2015). Reanalysis data (ERA Interim; Fig. S3) indicated that the wind speeds during summer (DJF 2015 to 2016) were stronger over the Falklands (5 to 6 m/s) and weaker during winter (JJA 2016).

3.3 Seasonal HYSPLIT air mass trajectory analyses

The daily back trajectory HYSPLIT analysis indicated that during the summer, 89% of the air masses originated ($n = 344$) west of the Falklands. Approximately 11% of summer air masses originated south of the Falklands near the Antarctic Peninsula. In winter, the air mass back trajectories ($n = 332$) were from the west, NW, and SW, while 21% of air masses had backward trajectories south of the Falklands near the Antarctic Peninsula (Fig. S4).

3.4 Monthly composite precipitation, $\delta^{18}O$ and $\delta D$

Each study location had $n = 12$ samples over the year, except for Surf Bay ($n = 11$), which is missing the September 2015 sample. Monthly composite $\delta^{18}O$ and $\delta D$ isotopes in precipitation throughout the year ranged from -12.3‰ to -4.8‰, and from -86‰ to -23‰, respectively. Monthly composite precipitation at each location was used to construct a local meteoric water line using $\delta^{18}O$ and $\delta D$ isotopes ($y = 7.571x + 5.527$; Fig. 3A) from monthly composite precipitation ($n = 47$). The range for winter $\delta^{18}O$ and $\delta D$ was from -8.6‰ to -6.6‰ and -61‰ to -40‰, respectively. Summer values of $\delta^{18}O$ and $\delta D$ in precipitation ranged from -12.3‰ to -5.3‰ and -86‰ to -38‰, respectively, and fit within the range of historical isotopes in precipitation from the Falklands (GNIP; Fig. S5). Summer and winter $\delta^{18}O$ and $\delta D$ isotopes in precipitation ($n = 24$) passed tests for normality (Shapiro-Wilk, $p = 0.297$ and $p = 0.614$, respectively) and failed tests for equal variance (Fisher’s F test, $p < 0.05$). A Mann-Whitney Rank Sum test indicated that the $\delta^{18}O$ isotopes in precipitation were not different for summer (median = -8.3‰) and winter (median = -7.4‰, $U = 39$, $p = 0.061$). For $\delta D$, the summer had a significantly lower median value (median = -64.3‰) than winter (median = -46.5‰, $U = 22$, $p = 0.004$). A one-way ANOVA found no significant difference among sites in $\delta^{18}O$ (F
(3,43) = 0.323, p = 0.809) or δD isotopes (F (3,43) = 0.361, p = 0.785) in precipitation when samples from all months and sites were included (n = 47).

### 3.5 δ¹³C and δ¹⁸O of α-cellulose – temperature, humidity, precipitation

Across all sites, measurements of monthly average leaf oxygen and carbon stable isotope values for α-cellulose extracted from leaf tissues (hereafter δ¹⁸Oleaf and δ¹³Cleaf) had a strong positive correlation (Pearson’s r = 0.877, p < 0.001, n = 24; Fig. 3B) and segregation between winter and summer values. Measurements of δ¹⁸O in precipitation had no significant correlation with δ¹⁸Oleaf or δ¹⁸Oroot across all sites (Table 1).

δ¹⁸Oleaf and δ¹³Cleaf values passed tests for normality (Shapiro-Wilk, p = 0.173 and p = 0.385, respectively) and equal variance (Fisher’s F test, p = 0.865 and p = 0.196, respectively). Thus, a two-way analysis of variance was conducted to detect the influence of independent variables (season and study location) on both δ¹⁸Oleaf and δ¹³Cleaf. Season included two levels (summer and winter) and study location consisted of four levels (Bleaker Island, Cape Dolphin, Surf Bay, and West Point Island). Analysis of combined winter and summer δ¹⁸Oleaf had a mean of 28.9 ‰ ± 1.3 SD, and ranged from 26.3 ‰ to 31.8 ‰ (range of 5.4 ‰; Table S2). The effect of season was significant, with an F ratio of F (1, 16) = 183.2, p < 0.001, and a 2.6 ‰ difference between summer (mean = 30.1 ± 0.8 SD) and winter (mean = 27.5 ± 0.6 SD; Fig. 4; Table S2). The effect of study location yielded an F ratio of F (3, 16) = 4.8, p = 0.014, indicating a significant difference in δ¹⁸Oleaf among study locations. Pairwise multiple comparison (Tukey’s post hoc test) of study locations indicated that Surf Bay is significantly more depleted in δ¹⁸Oleaf than Cape Dolphin (p = 0.016) and Bleaker Island (p = 0.029; Fig. 4). The interaction effect was not significant (p = 0.552). The mean of combined winter and summer δ¹³Cleaf value was -25.4 ‰ ± 1.31 SD, ranging from -30.4 ‰ to -21.9 ‰ (range = 8.4 ‰; Table S2). For δ¹³Cleaf, there was a significant difference between seasonal values (F ratio of F (1, 16) = 40.8, p < 0.001) in summer (mean = -24.2 ‰ ± 1.05 SD) and winter (mean = -26.8 ‰ ± 1.3 SD, Fig. 4; Table S2). Study location (p = 0.861; Figs. 4B and 4D) and the interaction effect (p = 0.638) were not significant. The mean δ¹³C in root α-cellulose (hereafter δ¹³Croot; n = 14) for summer was -25.3 ‰ ± 1.27 SD, and -26.6 ‰ ± 1.38 SD in winter (Table S2). After δ¹³Croot data passed tests for normality (Shapiro-Wilk test; p = 0.085), but not equal variance (p < 0.05), the two-way ANOVA indicated that for δ¹³Croot, the effects for season (p = 0.201) and study location (p = 0.521) were not statistically significant. The interaction effect was not significant (p = 0.886).

The mean δ¹⁸O in root α-cellulose (hereafter δ¹⁸Oroot; n = 14) for summer was 28.8 ‰ ± 1.04 SD, and 28.3 ‰ ± 0.5 SD for winter (Table S2). The δ¹⁸Oroot data passed tests for normality (Shapiro-Wilk test; p = 0.483) and equal variance (Fisher’s F test; p = 0.897); the two-way ANOVA indicated that for δ¹⁸Oroot, the location effect was statistically significant, while season was not. The difference in mean values among seasons (F (1, 8) = 5.4, p = 0.049) and study location (F (2, 8) = 8.7, p = 0.010) were statistically significant. Pairwise multiple comparison (Tukey’s post hoc test)
of study locations indicated that Cape Dolphin was significantly greater than Bleaker Island (p = 0.012) and West Point Island (p = 0.049). The interaction effect was not significant (p = 0.397).

4 Discussion

Significant inter-seasonal differences in $\delta^{13}C_{\text{leaf}}$ and $\delta^{18}O_{\text{leaf}}$ indicate that $P. \text{flabellata}$ tissues record high-resolution patterns of environmental change, supporting the use of $P. \text{flabellata}$ peat records as a paleoenvironmental proxy. The negative correlation between $\delta^{13}C_{\text{leaf}}$ and vapor pressure deficit suggests stomatal conductance is sensitive to atmospheric moisture conditions (Ferrio and Voltas, 2005). The observed positive correlation between $\delta^{13}C_{\text{leaf}}$ and temperature suggests higher temperatures led to an increased assimilation rate and reduced discrimination against $\delta^{13}C$ as shown in other vascular plant studies (Ferrio and Voltas, 2005; Ménot and Burns, 2001). $\delta^{13}C_{\text{leaf}}$ is driven by changes in the ratio of internal leaf partial pressure of CO$_2$ to that of ambient air, and can be explained by a greater influence of either stomatal conductance or increased photosynthetic capacity (Scheidegger et al., 2000).

The $\delta^{18}O_{\text{leaf}}$ is influenced by soil water, leaf water enrichment of $^{18}$O from transpiration, and biochemical fractionations. Leaf water enrichment of $^{18}$O due to transpiration, which is reflected in $\delta^{18}O_{\text{leaf}}$ (Deniro and Epstein, 1979; Roden and Ehleringer, 1999; Sternberg et al., 1986, p.198; Yakir, 1992), depends on relative humidity (Helliker and Ehleringer, 2002). However, the relationship between relative humidity and $\delta^{18}O_{\text{leaf}}$ deteriorated at relative humidity $> 90 \%$ in one C3 species (Helliker and Ehleringer, 2002). Diffusion limitation by stomatal resistance is primarily driven by relative humidity (White et al., 1994). The $\delta^{18}O_{\text{leaf}}$ and relative humidity ($> 60 \%$) were negatively correlated, which is consistent with other studies showing that $\delta^{18}O_{\text{leaf}}$ increases as relative humidity decreases (Barbraud et al., 2012; Helliker and Ehleringer, 2002). At high relative humidity the leaf will more strongly record variation in the isotopic composition of atmospheric vapor, however we have no direct measurements of $\delta^{18}O$ in water vapor.

Previous work on leaf water and cellulose isotopes in grasses demonstrated that the atmospheric-leaf vapor conditions are a strong predictor of $\delta^{18}O_{\text{leaf}}$ (Helliker and Ehleringer, 2002; Lehmann et al., 2018). The $\delta^{18}O$ of the leaf water is captured in the cellulose isotopes and can reflect the effect of changing environmental conditions during the growth of the leaf (Helliker and Ehleringer, 2002; Lehmann et al., 2017). The $\delta^{18}O_{\text{leaf}}$ can also depend on physiological effects, the type of plant anatomical feature used, and stage of development (Lehmann et al., 2017; Liu et al., 2017). Because cellulose records environmental variation along a gradient during leaf growth, we collected and homogenized whole leaves to avoid the complications of $\delta^{18}O$ enrichment (Helliker and Ehleringer, 2002). Our work supports the finding that atmospheric-leaf vapor conditions are reflected in $\delta^{18}O_{\text{leaf}}$, and expands the use of such paleoclimate proxies to peat-forming tussock grasses, which opens up new possibilities for reconstructing paleoclimates across the South...
Atlantic and beyond. At the higher relative humidity range, the $\delta^{18}O_{\text{leaf}}$ is more of a reflection of source water, while the $\delta^{18}O_{\text{root}}$ at low humidity differed greatly from source waters because of evaporative enrichment of $^{18}O$.

The positive correlation between $\delta^{13}C_{\text{leaf}}$ and $\delta^{18}O_{\text{leaf}}$ (Pearson’s $r = 0.88, p < 0.001$, $n = 24$; Table 1; Fig. 3B) suggests that stomatal conductance is the driving force acting on these two proxies, which is a likely scenario when water is not limiting (Saurer et al., 1997; Scheidegger et al., 2000). According to the Scheidegger et al. (2000) model, the decline in stomatal conductance was much more strongly expressed than photosynthetic capacity (maximum net photosynthesis). When air humidity increases, stomatal conductance is assumed to increase. In our study, stomatal conductance is likely driving both $\delta^{13}C_{\text{leaf}}$ and $\delta^{18}O_{\text{leaf}}$ due to relative humidity (Fig. 5). This pattern fits well with the Barbour and Farquhar (2000) model. Measurements of relative humidity allowed us to determine that stomatal conductance was more influential as a possible cause of change in partial pressure of CO$_2$ within the leaf.

In the Falklands, precipitation amount is not highly seasonal, but tends to vary the most in the summer (Fig. S6; data from Jones et al., 2013). Over the year of our study, summer $\delta^{18}O$ in precipitation varied more than winter $\delta^{18}O$ and tended to be more depleted. This is supported by the wind rose from Bleaker Island, which indicated prevailing winds from the SW (Fig. S2). In winter, precipitation tended to be more enriched in $^{18}O$, and most prevailing winds came from the west and NW where $^{18}O$ in equatorward precipitation would be more enriched than $^{16}O$ in high latitude meteoric sources (Fig. S5). The significant negative correlation between $\delta^{18}O$ in precipitation and $\delta^{18}O_{\text{root}}$ at Cape Dolphin (Pearson’s $r = 0.868$, $p = 0.025$, $n = 6$; Table 1) is consistent with the latitudinal origin of storm tracks and seasonal wind data (Fig. S4; Fig. S2). The pattern found at Bleaker Island is less clear, and warrants further investigation.

The observed lack of correlations between $\delta^{18}O_{\text{leaf}}$ and $\delta^{18}O$ values of precipitation (Table 1) demonstrate the overriding influence of humidity on patterns of leaf water $\delta^{18}O$. An alternative explanation for a lack of correlation is that our precipitation sampling density was not sufficient to establish a relationship between $\delta^{18}O_{\text{leaf}}$ and precipitation $\delta^{18}O$. Although $P$. flabellata produces new leaves throughout the year, the growth rate of leaves may not be in sync with shorter precipitation sampling intervals (less than monthly composite precipitation). Examination of leaf waters post-precipitation events would improve our understanding of $\delta^{18}O_{\text{leaf}}$.

The observed relationship between $\delta^{18}O$ in precipitation and $\delta^{18}O_{\text{root}}$ appeared to be less clear in part due to low sample number from only three study locations (Bleaker Island, Cape Dolphin, and West Point Island). At Bleaker Island, there was no correlation between $\delta^{18}O$ in precipitation and $\delta^{18}O_{\text{root}}$, while Cape Dolphin had a strong negative correlation (Table 1). At Cape Dolphin, greatest enrichment of $^{18}O_{\text{root}}$ occurred in summer when $\delta^{18}O$ precipitation was relatively low. In contrast, at Cape Dolphin $\delta^{18}O_{\text{root}}$ was lowest during the winter months when $\delta^{18}O$ of precipitation was high. Despite the strong relationship between $\delta^{18}O$ precipitation and $\delta^{18}O_{\text{root}}$ at Cape Dolphin, we propose that $P$. flabellata ‘bogs’ may not be ombrotrophic, and may potentially source water from fog, sea-spray, groundwater, or a mix. However, the relationship found at Cape Dolphin warrants further analysis of $\delta^{18}O_{\text{root}}$ and source waters. We also consider that the $\delta^{18}O_{\text{root}}$ records a signal of leaf humidity, and is influenced by source water and humidity effects on leaves; sugars produced in leaves are transported to roots to form cellulose, and about half of
the oxygen atoms in root cellulose originate from the leaf water signal. Roots of *P. flabellata* may have represented greater temporal integration of δ18O from precipitation into cellulose due to difficulty in distinguishing new growth in roots, like *Empodisma* in New Zealand (Amesbury et al., 2015).

Further work is needed to understand the relationship between δ18O_{root}, δ18O of precipitation, and that of root and leaf waters. Identifying sources of potential water would also add value, especially considering anecdotes of local differences in fog in the Falklands, which may be an unappreciated source of water for *P. flabellata*. Across the geographic range in the South Atlantic, *P. flabellata* may record a larger latitudinal gradients of isotopes in precipitation, as well as temperature and humidity, than recorded in the Falklands. Thus, establishing the seasonal patterns recorded by *P. flabellata* cellulose in the Falklands enables critical testing of paleoclimate hypotheses regarding the dynamics of Southern Hemisphere westerly wind behavior from broader latitudinal and longitudinal locations where *P. flabellata* occurs. Although *P. flabellata* forms high resolution peat records and is sensitive to inter-seasonal differences, reconstructions using *P. flabellata* peat would represent an integrated signal of broader climate trends, and not inter-annual or lower frequency differences.

5 Conclusion

The scarcity of terrestrial paleoclimate records in the South Atlantic has limited our understanding of past and future climate change and its impacts on ecosystems and people. We found that carbon and oxygen stable isotope values in *P. flabellata* tissues are correlated with seasonal differences in temperature and moisture, providing a promising new avenue for paleoclimate reconstructions in the South Atlantic. *P. flabellata* peats have high accumulation rates, contain abundant leaves, and date back to at least 12,500 14C years, with the potential to provide decadal-scale records of temperature, precipitation, and moisture source. Future work is needed to determine whether δ18O and δ13C of cellulose from *P. flabellata* macrofossils complement other regional proxies for changes in atmospheric temperature and relative humidity during the Holocene. Troublingly, these coastal peatlands are currently threatened by sea level rise and over-grazing, and their reductions means we are losing vital information about past environments in a time when paleoclimate records are needed to provide context for modern climate change in the South Atlantic.

6 Data availability

Datasets for monthly stable isotopes in precipitation, average temperature, and average relative humidity have been submitted to the Global Network of Isotopes in Precipitation (https://nucleus.iaea.org/wiser) and will be publicly
available upon acceptance for publication. Datasets for leaf and root stable isotopes of cellulose can be found at http://dx.doi.org/10.5281/zenodo.3104573, hosted at Zenodo upon acceptance for publication.

7 Author contribution

DG, DW, and JG designed the experiments and DG carried them out. DG performed laboratory analyses. DG prepared the manuscript with contributions from all co-authors.

8 Competing interests

The authors declare that they have no conflict of interest.

9 Acknowledgements

We gratefully thank the citizen scientists who collected samples: Ben Bernsten, Nikki and Mike Summers, Mike and Phyl Rendell, Robert and Elaine Short, Kicki Ericson, Thies Matzen. Paul Mayewski at the University of Maine Climate Change Institute provided precipitation collectors. Logistical, field, and laboratory support was provided by Paul Brickle and Megan Tierney at the South Atlantic Environmental Research Institute, Craig Cook at the University of Wyoming Stable Isotope Facility, Kayla Greenawalt, and Jiemin Guo. Funding: This research was supported by the US National Science Foundation [grant numbers DGE-1144423 and EF-1137336].

10 References

Amesbury, M. J., Charman, D. J., Newnham, R. M., Loader, N. J., Goodrich, J. P., Royles, J., Campbell, D. I., Roland, T. P. and Gallego-Sala, A.: Carbon stable isotopes as a palaeoclimate proxy in vascular plant dominated peatlands, Geochim. Cosmochim. Acta, 164, 161–174, doi:10.1016/j.gca.2015.05.011, 2015.

Araguás-Araguás, L., Froehlich, K. and Rozanski, K.: Deuterium and oxygen-18 isotope composition of precipitation and atmospheric moisture, Hydrol. Process., 14(8), 1341–1355, doi:10.1002/1099-1085(20000615)14:8<1341::AID-HYP983>3.0.CO;2-Z, 2000.

Barbraud, C., Rolland, V., Jenouvrier, S., Nevoux, M., Delord, K. and Weimerskirch, H.: Effects of climate change and fisheries bycatch on Southern Ocean seabirds: a review, Mar. Ecol. Prog. Ser., 454, 285–307, doi:10.3354/meps09616, 2012.

Brendel, O., Iamnetta, P. P. M. and Stewart, D.: A rapid and simple method to isolate pure alpha-<\texttt{tcapcellulose}, Phytochem. Anal., 11(1), 7–10, doi:10.1002/(SICI)1099-1565(20000102)11:1<7::AID-PCA488>3.0.CO;2-U, 2000.

Cernusak, L. A., Tcherkez, G., Keitel, C., Cornwell, W. K., Santiago, I. S., Knohl, A., Barbour, M. M., Williams, D. G., Reich, P. B., Ellsworth, D. S., Dawson, T. E., Griffiths, H. G., Farquhar, G. D. and Wright, I. J.: Why are non-photosynthetic tissues generally 13C enriched compared with leaves in C3 plants? Review and synthesis of current hypotheses, Funct. Plant Biol., 36(3), 199, doi:10.1071/FP08216, 2009.
Cernusak, L. A., Ubierna, N., Winter, K., Holtum, J. A. M., Marshall, J. D. and Farquhar, G. D.: Environmental and physiological determinants of carbon isotope discrimination in terrestrial plants, New Phytol., 200(4), 950–965, doi:10.1111/nph.12423, 2013.

Chambers, F. M., Booth, R. K., De Vleeschouwer, F., Lamentowicz, M., Le Roux, G., Mauquoy, D., Nichols, J. E. and van Geel, B.: Development and refinement of proxy-climate indicators from peats, Quat. Int., 268, 21–33, doi:10.1016/j.quaint.2011.04.039, 2012.

Craig, H.: Standard for Reporting Concentrations of Deuterium and Oxygen-18 in Natural Waters, Science, 133(3467), 1833–1834, doi:10.1126/science.133.3467.1833, 1961.

Deniro, M. J. and Epstein, S.: Relationship Between the Oxygen Isotope Ratios of Terrestrial Plant Cellulose, Carbon Dioxide, and Water, Science, 204(4388), 51–53, doi:10.1126/science.204.4388.51, 1979.

Dietl, G. P. and Flessa, K. W.: Conservation paleobiology: putting the dead to work, Trends Ecol. Evol., 26(1), 30–37, doi:10.1016/j.tree.2010.09.010, 2011.

Dietl, G. P., Kidwell, S. M., Brenner, M., Burney, D. A., Flessa, K. W., Jackson, S. T. and Koch, P. L.: Conservation Paleobiology: Leveraging Knowledge of the Past to Inform Conservation and Restoration, Annu. Rev. Earth Planet. Sci., 43(1), 79–103, doi:10.1146/annurev-earth-040610-133349, 2015.

Ehleringer, J. R. and Osmond, C. B.: Stable isotopes, in Plant Physiological Ecology: Field methods and instrumentation, edited by R. W. Pearcy, J. R. Ehleringer, H. A. Mooney, and P. W. Rundel, pp. 281–300, Springer Netherlands, Dordrecht, 1989.

Farquhar, G., O’Leary, M. and Berry, J.: On the Relationship Between Carbon Isotope Discrimination and the Intercellular Carbon Dioxide Concentration in Leaves, Funct. Plant Biol., 9(2), 121, doi:10.1071/PP9820121, 1982.

Farquhar, G. D. and Sharkey, T. D.: Stomatal Conductance and Photosynthesis, Annu. Rev. Plant Physiol., 33(1), 317–345, doi:10.1146/annurev.pp.33.060182.001533, 1982.

Ferrio, J. P. and Voltas, J.: Carbon and oxygen isotope ratios in wood constituents of Pinus halepensis as indicators of precipitation, temperature and vapour pressure deficit, Tellus B, 57(2), 164–173, doi:10.1111/j.1600-0889.2005.0137.x, 2005.

van Geel, B. and Middeldorp, A. A.: Vegetational history of Carbury Bog (Co. Kildare, Ireland) during the last 850 years and a test of the temperature indicator value of 2H/1H measurements of peat samples in relation to historical sources and meteorological data, New Phytol., 109(3), 377–392, doi:10.1111/j.1469-8137.1988.tb04208.x, 1988.

Gillett, N. P., Stone, D. A., Stott, P. A., Nozawa, T., Karpechko, A. Yu., Hegerl, G. C., Wehner, M. F. and Jones, P. D.: Attribution of polar warming to human influence, Nat. Geosci., 1(11), 750–754, doi:10.1038/ngeo338, 2008.

Gonfiantini, R.: Standards for stable isotope measurements in natural compounds, Nature, 271(5645), 534–536, doi:10.1038/271534a0, 1978.

Groff, D. V.: Modern and Paleoeccological Perspectives on a Terrestrial-Marine Linkage in the Falkland Islands, PhD Thesis, University of Maine, USA 196 pp., 2018.

Gröning, M., Lutz, H. O., Roller-Lutz, Z., Krakl, M., Gourcy, L. and Pöltenstein, L.: A simple rain collector preventing water re-evaporation dedicated for δ18O and δ2H analysis of cumulative precipitation samples, J. Hydrol., 448–449, 195–200, doi:10.1016/j.jhydrol.2012.04.041, 2012.

Helliker, B. R. and Ehleringer, J. R.: Establishing a grassland signature in veins: 18O in the leaf water of C3 and C4 grasses, Proc. Natl. Acad. Sci., 97(14), 7894–7898, doi:10.1073/pnas.97.14.7894, 2000.
Helliker, B. R. and Ehleringer, J. R.: Grass blades as tree rings: environmentally induced changes in the oxygen isotope ratio of cellulose along the length of grass blades, New Phytol., 155(3), 417–424, doi:10.1046/j.1469-8137.2002.00480.x, 2002.

Jones, P. D., Harpham, C. and Lister, D. H.: Construction of high spatial resolution climate scenarios for the Falkland Islands and southern Patagonia, U. K. Falkl. Isl. Trust, 40, 2013.

Lehmann, M. M., Gamarra, B., Kahmen, A., Siegwolf, R. T. W. and Saurer, M.: Oxygen isotope fractionations across individual leaf carbohydrates in grass and tree species, Plant Cell Environ., 40(8), 1658–1670, doi:10.1111/pce.12974, 2017.

Lehmann, M. M., Goldsmith, G. R., Schmid, L., Gessler, A., Saurer, M. and Siegwolf, R. T. W.: The effect of 18O-labelled water vapour on the oxygen isotope ratio of water and assimilates in plants at high humidity, New Phytol., 217(1), 105–116, doi:10.1111/nph.14788, 2018.

Libby, L. M., Pandolfi, L. J., Payton, P. H., Marshall, J., Becker, B. and Giertz-Sienbenlist, V.: Isotopic tree thermometers, Nature, 261(5558), 284–288, doi:10.1038/261284a0, 1976.

Lister, D. H. and Jones, P. D.: Long-term temperature and precipitation records from the Falkland Islands, Int. J. Climatol., 35(7), 1224–1231, doi:10.1002/joc.4049, 2015.

Liu, H. T., Schäufele, R., Gong, X. Y. and Schnyder, H.: The δ18O and δ2H of water in the leaf growth-and-differentiation zone of grasses is close to source water in both humid and dry atmospheres, New Phytol., 214(4), 1423–1431, doi:10.1111/nph.14549, 2017.

Ménot, G. and Burns, S. J.: Carbon isotopes in ombrogenic peat bog plants as climatic indicators: calibration from an altitudinal transect in Switzerland, Org. Geochem., 32(2), 233–245, doi:10.1016/S0146-6380(00)00170-4, 2001.

Moreno-Gutiérrez, C., Dawson, T. E., Nicolás, E. and Querejeta, J. I.: Isotopes reveal contrasting water use strategies among coexisting plant species in a Mediterranean ecosystem, New Phytol., 196(2), 489–496, doi:10.1111/j.1469-8137.2012.04276.x, 2012.

Payne, R. J., Ring-Hrubesh, F., Rush, G., Sloan, T. J., Evans, C. D. and Mauquoy, D.: Peatland initiation and carbon accumulation in the Falkland Islands, Quat. Sci. Rev., 212, 213–218, doi:10.1016/j.quascirev.2019.03.022, 2019.

Pendall, E., Markgraf, V., White, J. W. C., Dreier, M. and Kenny, R.: Multiproxy Record of Late Pleistocene–Holocene Climate and Vegetation Changes from a Peat Bog in Patagonia, Quat. Res., 55(2), 168–178, doi:10.1006/qres.2000.2206, 2001.

Roden, J. S. and Ehleringer, J. R.: Observations of Hydrogen and Oxygen Isotopes in Leaf Water Confirm the Craig-Gordon Model under Wide-Ranging Environmental Conditions, Plant Physiol., 120(4), 1165–1174, doi:10.1104/pp.120.4.1165, 1999.

Saia, O. E., Golluscio, R. A., Lauenroth, W. K. and Soriano, A.: Resource partitioning between shrubs and grasses in the Patagonian steppe, Oecologia, 81(4), 501–505, doi:10.1007/BF00378959, 1989.

Saurer, M., Aellen, K. and Siegwolf, R.: Correlating δ13C and δ18O in cellulose of trees, Plant Cell Environ., 20(12), 1543–1550, doi:10.1046/j.1365-3040.1997.d01-53.x, 1997.

Scheidegger, Y., Saurer, M., Bahn, M. and Siegwolf, R.: Linking stable oxygen and carbon isotopes with stomatal conductance and photosynthetic capacity: a conceptual model, Oecologia, 125(3), 350–357, doi:10.1007/s0044200040466, 2000.

Schwimmen, S. and Ehleringer, J. R.: Water use trade-offs and optimal adaptations to pulse-driven arid ecosystems, J. Ecol., 89(3), 464–480, doi:10.1046/j.1365-2745.2001.00576.x, 2001.
Smith, R. I. L. and Clymo, R. S.: An extraordinary peat-forming community on the Falkland Islands, Nature, 309(5969), 617–620, doi:10.1038/309617a0, 1984.

Smith, R. I. L. and Prince, P. A.: The natural history of Beauchène Island, Biol. J. Linn. Soc., 24(3), 233–283, doi:10.1111/j.1095-8312.1985.tb00374.x, 1985.

Stanworth, A. and Upson, R.: Minefield Restoration Trials Final Report., 2013.

Sternberg, L. D. S. L., Deniro, M. J. and Savidge, R. A.: Oxygen Isotope Exchange between Metabolites and Water during Biochemical Reactions Leading to Cellulose Synthesis, Plant Physiol., 82(2), 423–427, doi:10.1104/pp.82.2.423, 1986.

Strange, I. J., Parry, C. J., Parry, M. C. and Woods, R. W.: Tussac grass in the Falkland Islands, 1988.

Thompson, D. W. J. and Solomon, S.: Interpretation of Recent Southern Hemisphere Climate Change, Science, 296(5569), 895–899, doi:10.1126/science.1069270, 2002.

Turner, J. and Pendelbury, S. (eds.): The International Antarctic Weather Forecasting Handbook, in Sixth Conference on Polar Meteorology and Oceanography, American Meteorological Society, Boston, MA, San Diego, CA., 2000.

Upson, R., Williams, J. J., Wilkinson, T. P., Clubbe, C. P., Maclean, I. M. D., McAdam, J. H. and Moat, J. F.: Potential Impacts of Climate Change on Native Plant Distributions in the Falkland Islands, PLOS ONE, 11(11), e0167026, doi:10.1371/journal.pone.0167026, 2016.

Villalba, R., Lara, A., Masiokas, M. H., Urrutia, R., Luckman, B. H., Marshall, G. J., Mundo, I. A., Christie, D. A., Cook, E. R., Neukom, R., Allen, K., Fenwick, P., Boninsegna, J. A., Snr, A. M., Morales, M. S., Araneo, D., Palmer, J. G., Cuq, E., Aravena, J. C., Holz, A. and LeQuesne, C.: Unusual Southern Hemisphere tree growth patterns induced by changes in the Southern Annular Mode, Nat. Geosci., 5(11), 793–798, doi:10.1038/ngeo1613, 2012.

Weimerskirch, H., Louzao, M., de Grissac, S. and Delord, K.: Changes in Wind Pattern Alter Albatross Distribution and Life-History Traits, Science, 335(6065), 211–214, doi:10.1126/science.1210270, 2012.

White, J. W. C., Ciais, P., Figge, R. A., Kenny, R. and Markgraf, V.: A high-resolution record of atmospheric CO$_2$ content from carbon isotopes in pet, Nature, 367(6459), 153–156, doi:10.1038/367153a0, 1994.

Willis, K. J., Bailey, R. M., Bhagwat, S. A. and Birks, H. J. B.: Biodiversity baselines, thresholds and resilience: testing predictions and assumptions using palaeoecological data, Trends Ecol. Evol., 25(10), 583–591, doi:10.1016/j.tree.2010.07.006, 2010.

Wilson, P., Clark, R., McAdam, J. H. and Cooper, E. A.: Soil erosion in the Falkland Islands: an assessment, Appl. Geogr., 13(4), 329–352, doi:10.1016/0143-6228(93)90036-Z, 1993.

Yakir, D.: Variations in the natural abundance of oxygen-18 and deuterium in plant carbohydrates, Plant Cell Environ., 15(9), 1005–1020, doi:10.1111/j.1365-3040.1992.tb01652.x, 1992.
Table 1. Correlation coefficients (Pearson’s r) of $\delta^{18}$O and $\delta^{13}$C in leaf and root cellulose between $\delta^{18}$O in monthly composite precipitation, monthly average temperature, and humidity by site. Bold values indicate significant correlations >0.600 Pearson’s r. Significance level is p <0.05.

| Site         | n  | $\delta^{13}$C$_{\text{leaf}}$ | $\delta^{18}$O$_{\text{leaf}}$ | $\delta^{18}$O precipitation | Temp | Relative Humidity |
|--------------|----|---------------------------------|---------------------------------|--------------------------------|------|------------------|
| All sites    | 24 | 0.877 (<0.001)                  | -0.201 (0.346)                  | 0.889 (<0.001)                  | -0.877 (<0.001) |
| Bleaker Is.  | 6  | 0.864 (0.026)                   | -0.058 (0.913)                  | 0.947 (0.004)                   | -0.939 (0.005)  |
| Cape Dolphin | 6  | 0.990 (<0.001)                  | -0.357 (0.487)                  | 0.877 (0.021)                   | -0.979 (<0.001) |
| Surf Bay     | 6  | 0.769 (0.074)                   | -0.378 (0.460)                  | 0.952 (0.003)                   | -0.977 (<0.001) |
| West Point Is.| 6 | 0.971 (0.001)                   | -0.330 (0.523)                  | 0.977 (<0.001)                  | -0.900 (0.014)  |
| All sites    | 14 | --                              | -0.302 (0.294)                  | 0.311 (0.279)                   | -0.217 (0.457)  |
| Bleaker Is.  | 6  | --                              | 0.243 (0.642)                   | 0.222 (0.672)                   | 0.219 (0.677)   | -0.116 (0.827)  |
| Cape Dolphin | 6  | --                              | 0.623 (0.186)                   | -0.868 (0.025)                  | 0.701 (0.120)   | -0.694 (0.126)  |
| Surf Bay     | -- | --                              | --                              | --                             | --              | --               |
| West Point Is.| 2 | --                              | --                              | --                             | --              | --               |
| All sites    | 24 | --                              | 0.877 (<0.001)                  | 0.817 (<0.001)                  | -0.759 (<0.001) |
| Bleaker Is.  | 6  | --                              | 0.864 (0.026)                   | 0.843 (0.035)                   | -0.688 (0.131)  |
| Cape Dolphin | 6  | --                              | 0.990 (<0.001)                  | 0.849 (0.032)                   | -0.952 (0.003)  |
| Surf Bay     | 6  | --                              | 0.769 (0.074)                   | 0.780 (0.067)                   | -0.819 (0.046)  |
| West Point Is.| 6 | --                              | 0.971 (0.001)                   | 0.977 (<0.001)                  | -0.816 (0.047)  |
| All sites    | 14 | 0.724 (0.003)                   | --                              | 0.492 (0.074)                   | -0.299 (0.300)  |
| Bleaker Is.  | 6  | 0.832 (0.039)                   | --                              | 0.561 (0.247)                   | -0.273 (0.601)  |
| Cape Dolphin | 6  | 0.570 (0.237)                   | --                              | 0.778 (0.068)                   | -0.718 (0.108)  |
| Surf Bay     | -- | --                              | --                              | --                             | --              | --               |
| West Point Is.| 2 | --                              | --                              | --                             | --              | --               |
Figure 1. Study region. a) Map of the Falkland Islands and western South Atlantic Ocean with ocean currents (black arrows) and frontal zones (dashed lines). Study sites are shown in: b) West Point Island and c) Bleaker Island, Cape Dolphin, and Surf Bay.
Figure 2. a) A single large *Poa flabellata* bog made up of dead and living grass tillers growing on top of a decomposing pedestal at Cape Meredith, Falkland Islands. b) Conceptual figure of the relationship between $\delta^{13}C$ and $\delta^{18}O$ of leaf cellulose and air temperature, and humidity and/or soil moisture. c) Daily average temperature (°C) and d) relative humidity (%) from September 2015-August 2016 at the four study locations calculated from 2 hour measurements. Seven day running averages of daily average temperature and relative humidity are indicated by the lines for each study location. e) Image of a *P. flabellata* peat core section (Photo: D. Groff).
Figure 3. a) $\delta^{18}$O and $\delta$D (‰, VSMOW) isotopes in precipitation for each location (symbol shape) during four seasons (symbol color). The constructed local meteoric water line (LMWL; $y = 7.571x + 5.527$) is shown as a dashed line and global meteoric water line (GMWL: $\delta$D = 8.0 $\delta^{18}$O + 10) is a solid line. b) The relationship between average $\delta^{13}$C$_{\text{leaf}}$ and $\delta^{18}$O$_{\text{leaf}}$ (Pearson’s correlation coefficient, $r = 0.877$, $p < 0.001$, $n = 24$). Open circles are average values for samples collected in winter, solid circles in summer.
Figure 4. Oxygen and carbon stable isotopes of *Poa flabellata*. a) $\delta^{18}O_{\text{leaf}}$ (‰) comparison (mean ± 1 SD) between summer (DJF) and winter (JJA) and, b) at four study sites over one year; c) $\delta^{13}C_{\text{leaf}}$ (‰) comparison (mean ± 1 SD) between summer and winter, and d) at four study locations over one year.
Figure 5. Relationship between carbon and oxygen stable isotopes in cellulose and temperature and relative humidity during winter and summer at four sites. Relationship between $\delta^{13}$C$_{\text{leaf}}$, a) temperature and b) humidity; Relationship between $\delta^{18}$O$_{\text{leaf}}$, c) temperature, and d) humidity. Use Table 1 for Pearson’s r and p-values corresponding to correlations for each site.