Baffin Bay sea ice extent and synoptic moisture transport drive water vapor isotope ($\delta^{18}$O, $\delta^{2}$H, and deuterium excess) variability in coastal northwest Greenland

Pete D. Akers\textsuperscript{1}, Ben G. Kopec\textsuperscript{2}, Kyle S. Mattingly\textsuperscript{3}, Eric S. Klein\textsuperscript{4}, Douglas Causey\textsuperscript{2}, and Jeffrey M. Welker\textsuperscript{2,5,6}

\textsuperscript{1}Institut des Géosciences et l’Environnement, CNRS, 38400 Saint Martin d’Hères, France
\textsuperscript{2}Department of Biological Sciences, University of Alaska Anchorage, 99508 Anchorage, AK, USA
\textsuperscript{3}Institute of Earth, Ocean, and Atmospheric Sciences, Rutgers University, 08854 Piscataway, NJ, USA
\textsuperscript{4}Department of Geological Sciences, University of Alaska Anchorage, 99508 Anchorage, AK, USA
\textsuperscript{5}Ecology and Genetics Research Unit, University of Oulu, 90014 Oulu, Finland
\textsuperscript{6}University of the Arctic (UArctic), c/o University of Lapland, 96101 Rovaniemi, Finland

Correspondence: Pete D. Akers (pete.d.akers@gmail.com)

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Abstract. At Thule Air Base on the coast of Baffin Bay (76.51° N, 68.74° W), we continuously measured water vapor isotopes ($\delta^{18}$O, $\delta^{2}$H) at a high frequency (1 s$^{-1}$) from August 2017 through August 2019. Our resulting record, including derived deuterium excess (dxs) values, allows an analysis of isotopic–meteorological relationships at an unprecedented level of detail and duration for high Arctic Greenland. We examine isotopic variability across multiple temporal scales from daily to interannual, revealing that isotopic values at Thule are predominantly controlled by the sea ice extent in northern Baffin Bay and the synoptic flow pattern. This relationship can be identified through its expression in the following five interacting factors: (a) local air temperature, (b) local marine moisture availability, (c) the North Atlantic Oscillation (NAO), (d) surface wind regime, and (e) land-based evaporation and sublimation. Each factor’s relative importance changes based on the temporal scale and in response to seasonal shifts in Thule’s environment. Winter sea ice coverage forces distant sourcing of vapor that is isotopically light from fractionation during transport, while preventing isotopic exchange with local waters. Sea ice breakup in late spring triggers a rapid isotopic change at Thule as the newly open ocean supplies warmth and moisture that has $\sim$ 10\% and $\sim$ 70\% higher $\delta^{18}$O and $\delta^{2}$H values, respectively, and $\sim$ 10\% lower dxs values. Sea ice retreat also leads to other environmental changes, such as sea breeze development, that radically alter the nature of relationships between isotopes and many meteorological variables in summer. On synoptic timescales, enhanced southerly flow promoted by negative NAO conditions produces higher $\delta^{18}$O and $\delta^{2}$H values and lower dxs values. Diel isotopic cycles are generally very small as a result of a moderated coastal climate and the counteracting isotopic effects of the sea breeze, local evaporation, and convection. Future losses in Baffin Bay’s sea ice extent will likely shift mean annual isotopic compositions toward more summer-like values, and local glacial ice could potentially preserve isotopic evidence of past reductions. These findings highlight the influence that the local environment can have on isotope dynamics and the need for dedicated, multiseason monitoring to fully understand the controls on water vapor isotope variability.

1 Introduction

The Arctic environment is rapidly entering a new state dominated by warmer air temperatures in all seasons, accompanied by dramatic sea ice loss, ecological changes, and ice sheet mass loss. Improving our knowledge of how the Arctic water cycle responds to regional warming and develops feedbacks within the changing climate is key for planning for a more resilient Arctic environment and economy (Meier
et al., 2006; Vihma et al., 2016). While many modeling and satellite-based studies are examining this issue (NOAA, 2020), ground-based observations are still critical for understanding the present-day hydrological cycle and tracking ongoing changes in local environments (e.g., Steen-Larsen et al., 2014; Bonne et al., 2015; Klein et al., 2015). Despite this need, the Arctic is data-sparse in terms of spatial coverage and temporal span of quality surface observations. More environmental monitoring efforts are needed across the region, and particularly at the highest latitudes, to better capture the hydroclimate processes and moisture transport across the Arctic domain.

The stable isotopes of water are well-established environmental tracers of the water cycle, reflecting both local weather conditions and moisture history and synoptic patterns (Craig, 1961; Dansgaard, 1964; Rozanski et al., 1993; Gat, 1996). Although less commonly studied than lower latitude regions, analyses across the Arctic have also shown that local observations of water isotopes can reveal important connections to wider atmospheric parameters such as teleconnections and storm patterns (e.g., Moorman et al., 1996; Welker et al., 2005; Theakstone, 2011; H. L. Bailey et al., 2015; Puntag et al., 2016; Putman et al., 2017). While our knowledge of isotope dynamics in the hydro-sphere is largely based on studies of precipitation and surface waters (e.g., Dansgaard, 1964; Rozanski et al., 1993; Welker, 2000; Gurney and Lawrence, 2004), the development of field-deployable infrared laser spectrometers has fostered a number of recent studies focused on water vapor. With their ability to analyze water isotopes with a high frequency (> 1 min\(^{-1}\)) in a continuous vapor flow, these spectrometers are well suited to long-term monitoring studies (e.g., Sturm and Knohl, 2010; Aemisegger et al., 2012; A. Bailey et al., 2015; Wei et al., 2019). Several studies at land-based sites in the high latitudes have reported continuous water vapor isotope observations for periods ranging from a single season to multiple years (Table S1 in the Supplement), and their data sets are proving highly useful in understanding polar hydroclimate dynamics.

These observational studies are critical for tracking and understanding the ongoing climate changes in high-latitude regions, especially as the impacts of amplified polar warming on global weather patterns are hotly debated (Francis and Vavrus, 2012; Francis et al., 2018; Pithan et al., 2018; Nusbaumer et al., 2019; Cohen et al., 2020). Already, the published results from these polar water vapor isotope sites have highlighted the potential for tracking shifting moisture sources during extreme weather events (Bonne et al., 2015; Klein et al., 2015) and the importance of local geography in short-term isotopic variability (e.g., Kopec et al., 2014; Bréant et al., 2019). However, these sites reflect only a small portion of the vast and diverse polar environment, and fewer than half report data covering multiple years. More spatial coverage and longer periods of records are needed to fully harness water vapor isotope monitoring to resolve unanswered questions on polar hydroclimate processes and to accurately predict and detect future changes.

Greenland is particularly important in our efforts to better understand past, current, and future climate as its immense ice sheet has archived millennia of past climate changes (e.g., Steffensen et al., 2008), and environmental feedbacks from the increasing Greenland ice sheet loss reverberate globally (e.g., Box et al., 2012; Nghiem et al., 2012; Castro de la Guardia et al., 2015). The declining extent and duration of sea ice in its surrounding oceans are altering atmospheric moisture fluxes and transport across the Arctic (Gimeno et al., 2019; Nusbaumer et al., 2019), restructuring marine and terrestrial ecology (Bhatt et al., 2017; Laidre et al., 2020) and harming the health and welfare of indigenous communities (Meier et al., 2006). Northwest Greenland in particular is one of the fastest warming regions on Earth, with massive ice loss observed from glacial retreat and surface ablation (van As, 2011; Carr et al., 2013; Noël et al., 2019). Despite Greenland’s importance, water vapor isotope monitoring has only been reported from four locations on the island (Steen-Larsen et al., 2013; Bonne et al., 2014; Kopec et al., 2014; A. Bailey et al., 2015), and one record (Summit) has not had its data published.

We present here a new multiyear data set from Thule Air Base in northwest Greenland that has recorded stable oxygen and hydrogen ratios (\(\delta^{18}O\) and \(\delta^2H\), respectively) of ambient water vapor nearly continuously from 4 August 2017 through 31 August 2019. As a result, this record is the first reported for Greenland that continuously spans over 2 years, permitting comparative multiyear analysis of isotopic patterns and anomalies. Thule Air Base has been a focus of high Arctic research for many years (e.g., Schytt, 1955; Mastenbrook, 1968; Sullivan et al., 2008; Rogers et al., 2011; Leffler and Welker, 2013; Schaeffer et al., 2013), and our observing station on northern Baffin Bay allows a focus on how changing seasonal and interannual sea ice coverage affect the local climate and water vapor isotopes.

Our research addresses how interactions between the hydroclimate, cryosphere, and ocean are manifested in water vapor isotopes, and we focus here on identifying how these broad interactions are expressed and detected in local weather observations. The long period of record and very high frequency of isotopic (one observation per second, aggregated to 10 min) and meteorological observations (one observation per 10 min) enable us to study these water vapor isotope dynamics across varying temporal scales, from daily to interannual. This Thule data set and our continuing observations are part of the Multidisciplinary drifting Observatory for the Study of Arctic Climate (MOSAiC) project’s Arctic Water Isotope Network (Welker et al., 2019; MOSAiC, 2020), a series of nine pan-Arctic sites simultaneously observing water vapor isotopes. To that end, our research at Thule provides a focused examination of the many local and regional environmental controls on water vapor isotopes at a coastal high Arctic site in northwest Greenland.
Figure 1. Map of the local geography around Thule Air Base, the site of the water vapor and meteorological measurements for this study. The inset map at lower left shows the location of Thule Air Base in Greenland and other important Greenland sites. Elevation data was taken from ArcticDEM, the Polar Geospatial Center (Porter et al., 2018), and ice sheet, land, and ocean extent data were taken from Greenland Ice Sheet Mapping Project (GIMP) and the National Snow and Ice Data Center (NSIDC; Howat et al., 2014; Howat, 2017). Wind rose was created from the Thule airport data covering August 2017–August 2019 (USAF, 2019).

2 Field description

2.1 Local landscape

Our water vapor isotope observation station is located at Thule Air Base on the Pituffik peninsula in far northwest Greenland (Fig. 1). Landmarks are referred to here by their locally known English names together with the Greenlandic names, where known. The main base and airfield, established in 1951 by the United States and Denmark, occupy a low-lying (< 100 m a.s.l.) region along the North River (Pituffiup Kuussua), facing North Star Bay and Bylot Sound, with the small arms of Baffin Bay that are semiprotected by the Saunders and Wolstenholme islands (Appat and Qeqertarsuaq). The main base is bracketed to the south and north by two broad ridges (summits at 300 and 240 m above sea level (a.s.l.), respectively) locally called South Mountain (Akinnarsuaq) and North Mountain. The land-terminating margin of the Greenland ice sheet is approximately 14 km southeast of the main base and extends as low as 300 m a.s.l. This local section of the ice sheet, known as the Tuto Ice Dome, has a summit greater than 800 m a.s.l. and a mass balance that is semi-independent from the main Greenland ice sheet (Schytt, 1955; Hooke, 1970; Reeh et al., 1990).

Other military installations, both abandoned and presently occupied, are scattered throughout the wider Thule Defense Zone that covers the northern half of the Pituffik peninsula. Near the southern edge of the Thule Defense Zone lies higher terrain including Pingorsuit, the highest point of the penin-
sula at 815 m a.s.l. In addition, rugged terrain and mountains over 600 m a.s.l. run along the southern coastline of the peninsula toward Cape York (Sermamiut Nuuat). Wolstenholme Fjord (Uummannaq Kangerlua) forms the northern coast of the Pitufik peninsula, separating it from Steensby Land and the North Ice Cap (summit = > 1200 m a.s.l.). The three marine-terminating glaciers that feed into the fjord have severely retreated and thinned in the past century, including a retreat of over 8 km for the largest glacier (known as Harald Mølke or Ullip Sermia) since the 1940s (Mock, 1966; Hill et al., 2018).

2.2 Local climate

Thule has a polar desert to semidesert climate (Gold and Bliss, 1995; Sullivan et al., 2008), with a mean annual temperature of −10.0 ℃ and mean monthly temperatures ranging from −23.7 ℃ in February to +6.5 ℃ in July (2000–2018 observations; USAF, 2019). The high latitude (76° N) produces long periods of polar night (November–February) and midnight sun (May–August), and the day length changes by 15–30 min each day during transitional months. Extended periods of extremely cold temperatures below −25 ℃ are frequent in winter, and frosts and snowfall are possible in all months. Mean monthly temperatures are above freezing for only the three summer months, but summer can be surprisingly mild under continuous insolation, with inland temperatures sometimes rising above 10 ℃. Coastal sea ice and the seasonal snowpack develop by October and last through May to early June (Barber et al., 2001; Fetterer et al., 2017; Stroeve and Meier, 2018; USAF, 2019). At early spring maximum, sea ice covers Baffin Bay to an extent over 1000 km south of Thule, except for the biologically important North Water Polynya located to the northwest (Barber et al., 2001; Tang et al., 2004; Heide-Jørgensen et al., 2016).

Annual precipitation is 130 mm water equivalent, with half of this precipitation falling mostly as rain in June–August (USAF, 2019). Moisture in western Greenland is primarily sourced from the North Atlantic, but local sources of moisture, such as Baffin Bay, increase in importance during sea ice retreat in summer (Atkinson, 1981), and it brings cooler marine surface air, often associated with fog, several kilometers inland. This sea breeze generally strengthens in the afternoon and weakens at night, although continuous summer insolation often prevents the full shift to nightly katabatic flow that has been observed at other coastal polar sites (e.g., Kopec et al., 2014). These two wind patterns at Thule produce an overall bimodal wind distribution with azimuth peaks at 100° (katabatic) and 270° (sea breeze).

3 Methods

3.1 Equipment setting

A Picarro L2130-i with a standards delivery module (SDM) setup was installed in a temperature-controlled building on the crest of South Mountain (76.514° N, 68.744° W; 229 m a.s.l.) in October 2016, with quality data observations beginning in August 2017 after a system reset. Local air flow on top of South Mountain is unimpeded by topographic barriers from any direction within 20 km. Potential anthropogenic impacts to observations at the monitoring site are limited as the main base lies 2 km to the north and 200 m lower in elevation. Typically, less than 20 vehicles a day travel the access road that is 200 m from the building housing the L2130-i, and visitors are uncommon.

The L2130-i uses cavity ring-down spectroscopy to measure δ18O and δ2H in ambient water vapor and is particularly suited to long-term monitoring with limited human interaction. The vapor collection point is 5 m above the ground surface on the building’s roof edge, away from parking zones and building exhaust, and is outfitted with a plastic cap to prevent precipitation entering the inlet. A total of 5 m of Bev-A-Line IV EVA tubing (0.75 in. or 19 mm diameter) connects the inlet to the L2130-i with an air residence time of less than 3 min. The tubing is not independently heated, but it is routed immediately into the heated building interior which is always warmer than ambient outside air, even during extreme warm events in summer. This limits potential condensation within the tubing, and data quality control included a cull on observations with abnormal isotopic values that suggested possible condensation or precipitation contamination.
Certain tubing types, such as Synflex tubing, are known to adversely affect water vapor isotopic values by a slower vapor response time, increased fractionation of $\delta^2$H, and general smoothing of $\delta^2$H values at timescales less than 30 min (Sturm and Knohl 2010; Tremoy et al., 2011). We see no evidence of substantial tubing effects on our data, as the $\delta^2$H has the same response time and degree of variability as $\delta^{18}$O in our 10 min resolution data, and the strongest $\delta^{18}$O–$\delta^2$H correlations are with zero time lag. Additionally, we do not observe any significant differences in the $\delta^{18}$O–$\delta^2$H relationship between 10 min, hourly, and daily aggregations, which supports the assertion that both isotopic values are varying at a similar short-term frequency.

### 3.2 Isotopic observations and calibrations

Our L2130-i at Thule took an observation of the oxygen and hydrogen isotopic ratios in ambient water vapor approximately once per second, and these ratios are expressed in $\delta$ notation relative to the Vienna Standard Mean Ocean Water (VSMOW) standard as follows:

$$\delta = \frac{R}{R_{\text{VSMOW}}} - 1,$$

where $R$ is the measured ratio of rare to abundant isotopologue ($^{18}$O/$^{16}$O or $^2$H/$^1$H), and $R_{\text{VSMOW}}$ is the matching isotopic ratio of the reference standard water VSMOW. Deuterium excess values ($\Delta x$s) were calculated from each isotope observation by $\Delta x = \delta^2$H − $8 \cdot \delta^{18}$O (Dansgaard, 1964). Observations were continuous from August 2017 through August 2019, except for one period when the analyzer suffered a power failure (5–13 September 2018). Some additional gaps in the data were introduced as a result of quality checking (Sect. S1) and standard calibrations, but these gaps were typically less than 24 h in duration. We acknowledge that technical difficulties and some initial system design choices have prevented our data from being calibrated to the maximum quality level recommended by peer publications, and we attempt to be fully forthright with these issues here.

Every 25 h, two water standards, namely United States Geological Survey (USGS) no. 45 ($\delta^{18}$O = −2.24‰, $\delta^2$H = −10.3‰) and USGS no. 46 ($\delta^{18}$O = −29.80‰, $\delta^2$H = −235.8‰), were injected sequentially by the SDM into a stream of air supplied by a dry air canister in order to normalize to the VSMOW and Standard Light Antarctic Precipitation (SLAP) scales. Each standard was injected for 10 min, and the mean of the last 200 s of each standard injection was kept as the final isotopic value for each standard calibration. Preliminary tests at Thule suggested that acceptably stable isotopic value readings were reached in less than 10 min, and in 51 calibrations that were run in August and September 2019 (Table S2), the final 200 observations (i.e., the data used to calculate calibration values) have flat trends, suggesting acceptable isotopic stability (Table S3; mean slope ±1 standard deviation ($\sigma$) for USGS 45 – $\delta^{18}$O = −0.003 ± 0.0016‰ s$^{-1}$ and $\delta^2$H = 0.0026 ± 0.0062‰ s$^{-1}$; USGS 46 – $\delta^{18}$O = 0.0014 ± 0.0016‰ s$^{-1}$ and $\delta^2$H = 0.0026 ± 0.0064‰ s$^{-1}$). Our calibration interval and duration is longer and shorter, respectively, than in other similar studies (e.g., Bastrikov et al., 2014; A. Bailey et al., 2015; Bréant et al., 2019), but it is deemed a necessary compromise to extend the life of the dry air canister and standard waters at the remote location while still allowing an estimate of the machine’s accuracy drift.

The water standards used in calibration were originally chosen to cover the summer water vapor isotopic values at Thule before a project extension, but unfortunately most nonsummer isotopic values are below the standard waters’ value coverage. As a result, our reported values for nonsummer observations may suffer from undetermined accuracy biases relative to the VSMOW–SLAP scale, particularly at the lowest isotopic values. The potential impact on our analyses is likely limited, as long as any bias is internally consistent and small in magnitude, and a comparison of our reported isotopic values to other polar sites did not reveal any clear sign of major inaccuracies. However, anyone performing a future comparison between our Thule data and the data from another site should certainly take into account the potential for unquantified accuracy bias below the isotopic values of USGS 46. A wider range of standard waters is planned for continued future operation of the Thule observation station.

At lower water vapor mixing ratios (< 1500 parts per million by volume – ppmv), the L2130-i loses accuracy and precision, and a humidity response curve must be developed in order to correct for any analytical bias in isotopic values (Steen-Larsen et al., 2013; Bastrikov et al., 2014; A. Bailey et al., 2015). In July 2019, the two standard waters were each injected for 10 min into dry air at 10 different flow rates to produce a sequence of standard observations between 500 and 7000 ppmv. Humidity response curves were created using nonlinear regression to relate isotopic value offsets due to sensor biases at low humidity with water vapor mixing ratios, and ambient isotopic data were then corrected for sensor bias with these curves based on the ambient mixing ratio at the time of observation (Sect. S2; Fig. S1; Table S4). Ambient observations with a mixing ratio less than 500 ppmv (fewer than 500 individual 1 s$^{-1}$ observations) fell outside our humidity response observations and were thus excluded from further analysis.

The analyzer has a consistent accuracy offset of +1.9‰ and +1.1‰ in all $\delta^{18}$O and $\delta^2$H observations, respectively, at mixing ratios higher than 1500 ppmv. The accuracy bias changes below 1500 ppmv, and the offset for $\delta^{18}$O and $\delta^2$H is −4.5‰ and −20‰, respectively, at 500 ppmv. Ambient data were corrected for these offsets, and 95% confidence intervals for humidity response corrections for $\delta^{18}$O ranged from ±0.24‰ (500 ppmv) to ±0.09‰ (all observations > 1500 ppmv) and for $\delta^2$H ranged from ±1.6‰ (500 ppmv) to ±0.6‰ (all observations > 1500 ppmv). Precision linearly decreases with the re-
ciprocal of the water vapor mixing ratio (Sect. S2; Fig. S1; Table S5). Standard errors of the mean δ¹⁸O values for each flow rate in the humidity response curve ranged from 0.57 ‰ (500 ppmv) to < 0.05 ‰ (> 5000 ppmv), while the standard errors for δ²H ranged from 3.7 ‰ (500 ppmv) to < 0.4 ‰ (> 5000 ppmv). In comparison, each ambient wa-
ter vapor data point from the highest resolution used in our
analysis is the mean value of 600 observations (i.e., 10 min)
compared to the 200 observations in the humidity response
curve, and thus, the precision of our ambient data should as
precise as, or greater than, observed in the humidity response
curves. After the humidity response corrections, data were
quality checked again and offending data (e.g., extreme out-
liers, residual calibration vapor impacts, and extreme mois-
ture spikes unrelated to weather) removed. The final quality-
checked isotope data were aggregated into 10 min, hourly,
and daily means.

The calibration system described as above was installed
in August 2019. Before this installation, standards were in-
jected into an air stream dried by a Drierite column rather
than supplied by a dry air canister. However, the Drierite
column did not fully remove all ambient moisture and in-
creasingly lost effectiveness over time at Thule. The residual
ambient moisture had a significant effect on observed iso-
topic values during calibrations, and very few of the calib-
ration runs prior to the dry air canister installation accurately
recorded the standard values needed to correct for machine
analytical drift. As a result, we do not have accurate daily

calibrations for most of our observation period and cannot
apply a correction to account for possible analytical drift.

We believe that the database can still provide robust an-
alytical results despite the lack of drift correction. Although
we cannot be certain that an estimate of recent stability is rep-
resentative of the stability over the entire observation record
at Thule, the calibrations completed after the dry air installa-
tion show consistently high precision and limited day-to-day
time (Table S2). The standard deviation of the 51 cal-

3.3 Meteorological observations

An automated weather station (SMT – South Mountain), lo-
cated on the roof of the building that houses the L2130-i,
took a reading of air temperature, relative humidity (with re-
spect to the saturation vapor pressure over ice), and station
barometric pressure every 10 min throughout the duration of
isotopic observations (Muscaria, 2018). For analysis at differ-
ent temporal resolutions, these data were also aggregated into
hourly and daily means. The L2130-i takes a reading of the
water vapor mixing ratio with every isotopic observation, and
observations of this variable were added to the SMT data sets
as an independent meteorological observation. The mixing
ratios recorded by the L2130-i have a very strong linear re-
lationship (r² = 0.99) with mixing ratios calculated from the
SMT observations, but the L2130-i mixing ratios are greater
by a consistent factor of 1.23. For analytical simplicity, all
mixing ratio values used in analyses and reported here are
the unaltered observations from the L2130-i.

Hourly mean near-surface wind speeds and azimuths
(USA1F, 2019) were taken at the Thule Airport (THU), ag-
ggregated to daily values, and joined to the hourly and daily
SMT databases. Wind azimuths were converted into υ and
υ components for accurate aggregation and then recon-
verted back into azimuths. While the THU recording station
is 170 m lower than SMT, the two stations are less than
1 km apart, and their meteorological data agree well
(ρtemperature = 0.98, ρv_ew_point = 0.98, ρstation_pressure = 0.96).
Daily climate teleconnection indices for the North Atlantic
Oscillation (NAO) and Arctic Oscillation (AO) were down-
loaded from the US National Weather Service Climate Pre-
diction Center (NOAA, 2019) and joined with the daily SMT
database. Because the AO did not produce significant corre-
lations or results in our analyses, its discussion is limited here
and largely subsumed under discussion of the NAO. Daily
sea ice extent for Baffin Bay was obtained through the Na-
tional Snow and Ice Data Center (NSIDC) Multisensor Ana-
alyzed Sea Ice Extent (MASIE) product (Fetterer et al., 2010).

To reduce issues with the radial nature of azimuth data
during correlation analysis, wind azimuths were converted
to katabatic deviations, where an azimuth of 100° (i.e.,
the mean katabatic wind azimuth) was defined as zero, and other
observations were redefined as their minimum absolute de-
gree distance from 100°. The katabatic deviations therefore
range in value between 0 and 180°. While this does not
fully solve the radial issue (e.g., a katabatic deviation of 80°
can represent both a northerly 20° wind or a southerly 180°
wind), it fares well in distinguishing between the bimodal
“easterly katabatic” and “westerly sea breeze” regimes.

3.4 Statistical analyses

Spearman correlations between isotopic and meteorological
variables were calculated across the full data sets at all three
temporal resolutions. Strong correlations that arise between
variables in this analysis, though, may be spurious due to common responses to seasonal change. Likewise, variables that do not have a seasonal cycle, such as relative humidity, may have stronger correlations once the interference from seasonality is removed. To examine this, we removed the strong seasonal cycles of the isotopic values, air temperature, and mixing ratio by sinusoidal curves fitted to the annual cycle of each full data time series. With these seasonally adjusted (SA) data, we calculated a second set of correlations.

Data at each temporal resolution were also binned by month, and correlations were then calculated for each month to examine how isotope–climate relationships change over the course of 1 year. To assess temporal covariance and possible lead–lag relationships between teleconnection indices (NAO and AO) and isotopic variables, we performed cross-correlation analysis (Addison et al., 2020) with a first-differencing methodology (Peterson et al., 1998) to remove autocorrelations and avoid cross-correlation bias (Olden and Neff, 2001; Runge et al., 2014).

3.5 Back-trajectory analysis

We compiled a Thule-specific moisture-sourcing and transportation quasi-climatological air parcel back-trajectory analysis, using the HYSPLIT tool and based on 3D MERRA-2 wind field data that have a horizontal grid resolution of 0.5° latitude/0.625° longitude and 72 hybrid-eta vertical levels from the surface to 0.01 hPa (Stein et al., 2015; Gelaro et al., 2017). To build this quasi-climatological, we randomly selected 10 d from each month during 1980–2018, as we judged 10 d to provide an acceptable representation of the true climatology while remaining within reasonable computation and data storage limits. We then initiated 10 d back trajectories for each randomly selected day from the MERRA-2 grid point nearest Thule at the four synoptic hours (00:00, 06:00, 12:00, and 18:00 coordinated universal time – UTC) on six vertical levels (10, 100, 200, 500, 1000, and 1500 m a.g.l. – above ground level), resulting in 2880 total trajectories. Following previous studies (e.g., Sodemann et al., 2008; Molina and Allen, 2019), the moisture uptake algorithm implemented in the PySPLIT library (Warner, 2018) infers moisture uptake into the parcel when specific humidity change ($\Delta q$) is greater than +0.2 g kg$^{-1}$ and infers precipitation when $\Delta q$ is less than −0.2 g kg$^{-1}$. Later moisture uptakes are weighted more heavily than uptakes occurring prior to precipitation (Sodemann et al., 2008). A domain covering Baffin Bay from the Davis Strait to Nares Strait was defined, and the percentage of moisture uptake occurring within and outside this domain was calculated with raw values at each grid cell multiplied by the cosine of latitude to compensate for smaller grid cell area with northward extent.

4 Environmental controls on water vapor isotopes at Thule

The mean annual water vapor $\delta^{18}$O, $\delta^{2}$H, and dxs values at Thule are $-33.2 \pm 6.1$‰, $-249 \pm 43$ ‰, and $+16.2 \pm 8.1$ ‰, respectively (10 min data; mean ±1σ). The isotopic records reveal strong annual cycles, with the highest values for $\delta^{18}$O and $\delta^{2}$H in summer and lowest values in winter and early spring, while dxs values show the opposite pattern with minimum values in summer and maximum values in winter (Fig. 2; Table 1; Sect. S3). The isotopic values also have very wide overall ranges, with $\delta^{18}$O, $\delta^{2}$H, and dxs spanning over 30 ‰ ($-49.5$ ‰ to $-17.5$ ‰), 230 ‰ ($-377$ ‰ to $-142$ ‰), and 55 ‰ ($-7.6$ ‰ to 47.5 ‰), respectively, in our 10 min database. Temporal aggregation reduces the overall range of and variability in isotopic data by reducing extreme values, but this does not greatly affect the distribution or means. Conclusions drawn from one level of aggregation are consistent across all levels of aggregation.

The water vapor isotopes also have substantial subseasonal variability, and the magnitude of irregular hourly to weekly variations caused by synoptic weather events can approach 30 ‰–50 ‰ of the entire annual isotopic ranges. These large isotopic changes are typically tied to the different sourcing and transport of warm and cold sector moisture during cyclone passage, as well as changes in air temperature, surface winds and cloud cover (Gedzelman and Lawrence, 1990; Coplen et al., 2008; Dütsch et al., 2016). Similarly, strong anticyclones and blocking events can also dominate local weather and alter atmospheric flow throughout the polar region (Davini et al., 2012; McLeod and Mote, 2016; Wernli and Papritz, 2018). Diel cycles in isotopes are very small (generally $<1$ ‰ for $\delta^{18}$O and dxs), but they help reveal which environmental factors are driving isotopic changes at the local scale.

Based on our statistical analyses (Table 2) and field observations, the variability in water vapor isotope ratios at Thule can be largely explained by the following five interacting factors: (a) local air temperature, (b) local marine moisture availability, (c) the NAO, (d) surface wind regimes, and (e) the evaporation and sublimation of local surface waters and snow. These factors interact and compete with each other, with the dominance of particular factors varying over the course of 1 year with the changing seasons. Together, these factors can be considered as specific expressions of the broad environmental responses to changes in Baffin Bay sea ice extent and polar synoptic flow. To clarify this complex situation, this discussion will first detail the effects of these factors specifically in the Thule environment. The impact of these factors on isotopic systematics will then be examined across seasonal, synoptic, and daily scales to reveal how observed isotopic values are produced.

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Figure 2. Observed data from Thule, Greenland, of (a) water vapor $\delta^{18}$O, (b) water vapor dvs, (c) mean air temperature, (d) water vapor mixing ratio, (e) relative humidity, (f) wind speed, (g) wind azimuth, (h) station pressure, (i) NAO index (NOAA, 2019), and (j) Baffin Bay sea ice extent (Fetterer et al., 2010). The time series of $\delta^{2}$H is very similar to $\delta^{18}$O and not shown. Data shown are daily mean values with hourly values as the lighter backdrop line for higher resolution variables (a–h). All observations were taken at the SMT site, except for wind speed and azimuth which were observed at the Thule Airport (THU). Yellow dashed vertical lines indicate moisture pulse events, and orange dotted vertical lines indicate the timing of sea ice breakup near Thule.
Table 1. Annual and seasonal mean ±1 standard deviation (SD) values of selected water vapor isotopic and meteorological variables for Thule, Greenland, over the analytical period of this study. Values are calculated using 10 min resolution data.

|                  | δ18O (%ε) | δ2H (%ε) | dxx | Air temperature (°C) | H2O mixing ratio (ppmv) | Relative humidity (%) |
|------------------|------------|----------|-----|----------------------|------------------------|----------------------|
| Annual           | −33.2 ± 6.1 | −249 ± 43 | +16.2 ± 8.1 | −8.1 ± 11.1 | 4180 ± 3000 | 78 ± 16 |
| Spring (MAM)     | −35.9 ± 5.8 | −270 ± 40 | +17.7 ± 8.5 | −12.1 ± 9.2 | 2850 ± 1940 | 79 ± 16 |
| Summer (JJA)     | −27.6 ± 3.5 | −212 ± 24 | +8.6 ± 4.9  | +5.4 ± 4.1  | 8310 ± 1290 | 73 ± 18 |
| Autumn (SON)     | −31.3 ± 4.4 | −233 ± 31 | +17.7 ± 6.0 | −8.5 ± 6.4  | 3600 ± 1740 | 78 ± 13 |
| Winter (DJF)     | −38.5 ± 3.6 | −287 ± 26 | +21.6 ± 6.4 | −18.9 ± 5.2 | 1550 ± 900  | 83 ± 16 |

Table 2. Spearman correlation values between isotopic and meteorological variables at Thule. Two sets of correlations are reported here, namely analyses from the original data and analyses from the seasonally adjusted data in which the seasonal cycle was removed for isotopic, air temperature, mixing ratio, and sea ice data. Seasonally adjusting the data weakens nearly all correlations by removing the common response to seasonal change, but strengthens isotopic correlations with relative humidity. Correlations are given for all available levels of temporal aggregation, but some variables do not have data at finer resolutions. Temporal aggregation slightly strengthens correlations but does not generally change the order of variables when ranking by correlation strength. Due to the large sample sizes, all correlations are significant at p < 0.05, except for a few with very weak correlations (italicized).

|                  | ρ (10 min) | ρ (hourly) | ρ (daily) |
|------------------|------------|------------|-----------|
|                  | δ18O       | δ2H        | dxx       | δ18O       | δ2H        | dxx       | δ18O       | δ2H        | dxx       |
| Original         |            |            |           |            |            |           |            |            |           |
| Air temperature  | +0.76      | +0.73      | −0.74     | +0.76      | +0.73      | −0.75     | +0.81      | +0.77      | −0.79     |
| H2O mixing ratio | +0.83      | +0.80      | −0.79     | +0.83      | +0.80      | −0.79     | +0.86      | +0.83      | −0.82     |
| Relative humidity| +0.06      | +0.06      | −0.07     | +0.06      | +0.06      | −0.07     | 0.00       | −0.01      | −0.01     |
| Station pressure | +0.09      | +0.08      | −0.12     | +0.09      | +0.08      | −0.12     | +0.10      | +0.09      | −0.13     |
| Wind speed       | −          | −          | −         | −0.08      | −0.07      | +0.11     | −0.14      | −0.13      | +0.16     |
| Katabatic deviation| −         | −         | −        | −0.50      | +0.48      | −0.50     | +0.48      | +0.47      | −0.41     |
| NAO              | −          | −          | −        | −          | −          | −         | −0.25      | −0.23      | +0.29     |
| AO               | −          | −          | −        | −          | −          | −         | −0.06      | −0.04      | +0.17     |
| Baffin Bay ice extent | −      | −          | −        | −          | −          | −         | −0.69      | −0.71      | +0.47     |

Seasonally adjusted

|                  | ρ (10 min) | ρ (hourly) | ρ (daily) |
|------------------|------------|------------|-----------|
|                  | δ18O       | δ2H        | dxx       | δ18O       | δ2H        | dxx       | δ18O       | δ2H        | dxx       |
| Air temperature  | +0.24      | +0.20      | −0.31     | +0.25      | +0.20      | −0.32     | +0.32      | +0.27      | −0.38     |
| H2O mixing ratio | +0.55      | +0.49      | −0.58     | +0.55      | +0.49      | −0.58     | +0.57      | +0.51      | −0.60     |
| Relative humidity| +0.46      | +0.43      | −0.44     | +0.38      | +0.36      | −0.33     | +0.32      | +0.30      | −0.29     |
| Station pressure | +0.04      | +0.03      | −0.07     | +0.04      | +0.03      | −0.07     | +0.03      | +0.02      | −0.07     |
| Wind speed       | −          | −          | −        | −0.03      | −0.02      | +0.06     | −0.03      | −0.03      | +0.06     |
| Katabatic deviation| −          | −          | −        | +0.19      | +0.17      | −0.18     | +0.10      | +0.09      | −0.03     |
| NAO              | −          | −          | −        | −          | −          | −         | −0.14      | −0.12      | +0.20     |
| AO               | −          | −          | −        | −          | −          | −         | −0.04      | −0.02      | +0.17     |
| Baffin Bay ice extent | −      | −          | −        | −          | −          | −         | −0.02      | −0.03      | +0.05     |

4.1 Local air temperature

Our back-trajectory analysis shows that most water vapor is supplied to Thule from evaporation in more southerly locations and then advected north along the western coast, with a substantial fraction of this water vapor having previously passed over the southern Greenland ice sheet (Fig. 3a). As the air mass cools in this northerly transport, water vapor will condense into clouds and precipitation that selectively remove isotopically heavier water molecules (Dansgaard, 1964; Rozanski et al., 1993). This Rayleigh fractionation results in an air mass with water vapor that is increasingly isotopically lighter (i.e., depleted in 18O and 2H) as it travels to Thule, and colder conditions at Thule create a steeper temperature gradient versus the more southerly moisture source that enhances the degree of fractionation (e.g., Rozanski et al., 1993; Bonne et al., 2014; Kopec et al., 2019). This connection is supported by the strong correlations of δ18O and δ2H with air temperature (ρδ18O,T=10min = +0.76 and ρδ2H,T=10min = +0.73), which are surpassed in strength only by
correlations with the closely linked water vapor mixing ratio (Table 2).

However, strong annual correlations between water isotopes and temperature often result from common responses between air temperature and other variables that also follow seasonal patterns (e.g., Akers et al., 2017). This also appears true for Thule, as the correlations are much weaker when seasonal cycles are removed ($\rho_{\text{SA}-\delta^{18}\text{O} \cdot 10\text{ min}} = +0.24$ and $\rho_{\text{SA}-\delta^2\text{H} \cdot 10\text{ min}} = +0.20$). Most likely, much of the strength of the nonseasonally adjusted correlations is due to the common response with broader environmental changes linked to seasonally covarying sea ice extent. Despite this, the weaker seasonally adjusted correlations show that a temperature effect is still expressed to a weaker degree in shorter term, intraseasonal isotopic variations (Table 2).

Local air temperature is not typically seen as a primary driver of dxx variability; rather, variables related to environmental conditions at the initial moisture source, such as sea surface temperature and relative humidity, are the principal controls (e.g., Merlivat and Jouzel, 1979; Vimeux et al., 1999; Pfahl and Sodemann, 2014). However, dxx values at Thule have a strong correlation with air temperature ($\rho_{\text{dxx} \cdot 10\text{ min}} = -0.74$), and the correlation using seasonally adjusted data is actually stronger ($\rho_{\text{SA}-\text{dxx} \cdot 10\text{ min}} = -0.31$) than the correlations observed in $\delta^{18}\text{O}$ and $\delta^2\text{H}$. Temperature-driven changes on dxx have been reported (Jouzel and Merlivat, 1984; Dütsch et al., 2017; Kopec et al., 2019), particularly for very cold conditions, and it is possible that this is contributing to the short-term correlation between dxx and air temperature as well as the strong negative correlations between dxx and both $\delta^{18}\text{O}$ and $\delta^2\text{H}$. However, it is also possible that the correlation of dxx with temperature is, like with $\delta^{18}\text{O}$ and $\delta^2\text{H}$, actually responding to another environmental factor that covaries with temperature.

4.2 Local marine moisture availability

The isotopic composition of water vapor in an air mass is initially determined by the isotopic composition of the source water and by the environmental conditions during vapor uptake. At synoptic scales, the relative humidity near the sea surface strongly determines the dxx values for the result-

Figure 3. Back-trajectory analysis results for air parcels arriving at Thule for each meteorological season, including (a) trajectory density, (b) regions of water vapor uptake from the boundary layer (BL), and (c) relative humidity for these regions of vapor uptake. Analysis is based on a quasi-climatological that sampled 10 random days from each month during 1980–2018. Trajectory points are binned onto a 1° latitude/longitude grid as (a) probability density functions (PDF) or (b, c) mean values within each grid cell. White and magenta lines show the extent of 95 % and 5 % sea ice concentration (SIC), respectively, for the given season.
ing moisture parcel, with higher humidity producing lower dxs values and vice versa (Craig and Gordon, 1965; Pfahl and Sodemann, 2014), while, at longer timescales, sea surface temperature has traditionally been cited as a primary control on dxs values (e.g., Merlivat and Jouzel, 1979; Vimeux et al., 1999). If these isotopic signatures are well preserved through later moisture transport, isotopic analysis of water vapor or precipitation can be used to remotely infer climate changes at the moisture source and/or shifts to different moisture sources (e.g., Merlivat and Jouzel, 1979; Feng et al., 2009; Bonne et al., 2015, 2019; Dütsch et al., 2017). This has been applied most notably to ice cores in order to reconstruct moisture source changes extending back deep through time (e.g., Vimeux et al., 1999; Steffensen et al., 2008; Steen-Larsen et al., 2013; Osterberg et al., 2015).

Local polar waters have high surface relative humidity when open in summer (Fig. 3c) and supply Thule with water vapor that has high $\delta^{18}O$ and $\delta^2H$ values and low dxs values. At Thule, changes in the seasonal sea ice extent permit or restrict the delivery of local moisture from the nearby ocean, and this availability is a primary control on mean isotopic values. Our back-trajectory analysis shows that the regions of predominant water vapor uptake for air masses arriving at Thule vary substantially over the year (Fig. 3b), supporting the role of sea ice in determining moisture sourcing. Baffin Bay is the dominant evaporative moisture source from the late spring through early winter, contributing $\sim 50 \%$ of water vapor transported to Thule (Table S6). Substantial contributions from the Labrador Sea, Denmark Strait, Hudson Bay, and Canadian Arctic Archipelago regions are also observed during this time.

From January until May, however, local seas are extensively ice covered, and the majority of water vapor present at Thule originates from more distant sources in the Labrador Sea and North Atlantic (Fig. 3b). Large evaporation events in the North Atlantic tend to occur in the cold sector of extratropical cycles (Aemisegger, 2018; Aemisegger and Papritz, 2018) where the dry air (Fig. 3c) produces water vapor with high dxs values. As this moisture travels north to Thule, fractionation from rain out during transport results in isotopically light water vapor, and sea ice coverage in Baffin Bay and other local Thule waters prevents isotopic exchange that would mitigate some of this fractionation. Interestingly, our analysis reveals that a large amount ($>30 \%$) of Thule’s winter water vapor is supplied by sections of southern Baffin Bay and the Labrador Sea, with $5\%$–$95\%$ climatological mean sea ice cover. Moisture originating from such an environment often has very high dxs values from the frequent invasions of very dry Arctic air originating over expanses of continuous sea ice (Kurita, 2011) and/or vapor sourcing from snow accumulated on top of the sea ice (Bonne et al., 2019).

![Figure 4](https://example.com/figure4.png)

**Figure 4.** Cross-correlation analysis results between the NAO index and selected isotopic and meteorological variables. Results shown are all statistically significant at $p < 0.05$ ($n = 737$), and nonsignificant results are not illustrated.

### 4.3 The North Atlantic Oscillation

Phase shifts in the NAO can serve to enhance or limit moisture transport from the south to Thule (Fig. 3a) as atmospheric mass is redistributed between the Arctic and North Atlantic, and changes in water isotopes resulting from these atmospheric shifts have been detected in Arctic snow and ice (e.g., Vinther et al., 2003, 2010; Zheng et al., 2018) and plants (Welker et al., 2005). In western Greenland, the negative phase of the NAO (NAO$-$) is associated with enhanced southerly flow that brings warmer temperatures and greater regional snow, while the positive phase (NAO$+$) has stronger westerlies that limit the northward penetration of southerly air masses (Sodemann et al., 2008; Bjørk et al., 2018). These effects are manifested in our observations where more negative NAO indices are correlated with a stronger Greenland anticyclone (i.e., higher Thule station pressure; $\rho_{\text{NAO-pres}} = -0.60$).

When southerly advection along the western edge of the anticyclone is promoted during NAO$-$ phases, Thule tends to be warmer ($\rho_{\text{NAO-SA-temp}} = -0.31$) and have a higher water vapor mixing ratio ($\rho_{\text{NAO-SA-MR}} = -0.27$). Cross-correlation analysis (Figs. 4 and S2) reveals that shifts to more negative NAO indices produce isotopically heavier water vapor observations with lower dxs values 2–3 d later at Thule, presumably as air advected north over Baffin Bay picks up local moisture, and/or the warmer conditions reduce the degree of fractionation during transport. We did not find noteworthy isotopic correlations with the AO at Thule, likely because changes in AO phase relate to pan-Arctic conditions, while the NAO changes are more focused on Greenland and the North Atlantic.

The relationship between water vapor isotopes and NAO phase across interannual timeframes at Thule is uncertain from our data’s limited period of record, but climatology and ice core studies have argued for clear impacts on Greenland’s precipitation, glacial mass balance, and sea ice
at this scale (Stern and Heide-Jørgensen, 2003; Vinther et al., 2003; Sodemann et al., 2008; Bjørk et al., 2018). NAO indices in summer 2018 were almost continuously positive, coinciding with relatively cool conditions at Thule and below average melt of the northwestern Greenland ice sheet. In contrast, summer 2019 was dominated by NAO− conditions and was nearly 4°C warmer than summer 2018 at Thule (USAF, 2019). Greenland as a whole recorded near-record ice sheet melt over the 2019 melt season, linked in part to extensive warm and sunny conditions over the ice sheet (Maslankik and Stroeve, 1999). Despite these markedly different conditions, warm and sunny conditions over the ice sheet (Maslanik and Stroeve, 1999). Despite these markedly different conditions, mean δ18O and δ2H values are practically identical between the two summers (<0.01% and <1% different, respectively), and mean dxs values differ only by 1%. This suggests that changes in synoptic flow patterns have little impact when local seas are free of sea ice, as isotopic exchange with local marine moisture can buffer any potential isotopic changes from longer distance synoptic transport shifts.

However, spring 2018 was also characterized by NAO+ indices and colder conditions, while spring 2019 had lower NAO indices and warmer weather. In contrast to the summer observations, δ18O and δ2H values are 4.4% and 34% higher, respectively, in spring 2019 than spring 2018, and dxs values are 1.7% lower. As sea ice is still largely intact through spring near Thule, the limited potential for local water input and exchange at this time appears to allow an NAO signal to be expressed in the mean isotopic values. Additional seasons and years of observation with different NAO conditions will help clarify the magnitude and seasonal extent of interannual effects from the NAO on Thule water vapor isotopes.

4.4 Surface wind regimes

Between April and September, the surface winds at Thule alternate between east-southeasterly katabatic winds off the nearby ice sheet and a west-northwesterly sea breeze from Bylot Sound, although not necessarily on a regular daily cycle. Such wind shifts at other coastal sites are isotopically identifiable because katabatic winds supply isotopically light vapor from ice sheets, while sea breezes supply isotopically heavy vapor from the nearby ocean (e.g., Kopec et al., 2014; Bréant et al., 2019). After adjusting for the seasonal biases at Thule, where katabatic winds dominate in colder months and the sea breeze dominates in warmer months, a weak correlation between katabatic deviation (i.e., sea breeze occurrence) and the water vapor isotopes (ρSA,δ18O,hr = +0.19 and ρSA,dxs,hr = −0.18) indicates that on a given day the sea breeze will bring isotopically heavier water vapor with lower dxs values than katabatic flow. Although the difference in mean isotopic values is statistically significant between the two wind regimes for all isotopic species (p < 0.001; Welch’s t test), there is substantial overlap in the overall range (May–September data, mean ±1σ: δ18Osea-breeze = −26.8 ± 3.5‰ vs. δ18O katabatic = −28.5 ± 3.1‰; δ2H sea-breeze = −207±25‰ vs. δ2H katabatic = −219±22‰; dxs sea-breeze = 7.6±4.8‰ vs. dxs katabatic = 9.3±4.8‰; Fig. S3). The isotopic difference between the two wind regimes may be smaller at Thule than other studied sites because of Thule’s location on the west-pointing Pituffik peninsula. Much of the southeasterly katabatic flow here is not sourcing air directly off the main Greenland ice sheet, but rather air that is traveling over Baffin Bay along the western coast and potentially over nearby De Dødes Fjord (Fig. 1). Thus, both the katabatic winds and the sea breeze are bringing water vapor from locations where it has been able to isotopically exchange with local ocean waters.

The slightly lower water vapor isotope values during katabatic flow are then likely due to the vapor’s brief passage over topographic highs of Cape York and the Tuto Ice Dome, while the sea breeze carries moisture onshore, unimpeded, from Bylot Sound. Additionally, we note that as our wind data come from a lower elevation site than our isotopic sampling, it is possible that particularly shallow sea breezes could produce wind observations identified as sea breeze, while the air at the isotopic sampling site on South Mountain ridgetop is still katabatically sourced. However, no consistently clear signs of different air masses affecting the two stations in summer were found when examining periods with anomalous observations (e.g., sea breeze azimuth with high air temperature).

4.5 Evaposublimation of local snow and surface waters

Evaporation and sublimation of local surface waters and snowpack (referred to here as evaposublimation) can also alter local water vapor isotopic values by supplying vapor that is isotopically lighter with higher dxs values relative to the source snow/ice/water’s isotopic composition (Casado et al., 2018; Kopec et al., 2019). Most of the snow at Thule falls in autumn and early winter (USAF, 2019), and surface waters in Thule generally match the isotopic composition of the winter snowpack (Csank et al., 2019). Thus, evaposublimation during warmer months will generally serve to lower δ18O and δ2H values and raise dxs values of water vapor in the local environment, although these two processes may fractionate the water isotopes differently, and the exact isotopic composition of supplied vapor will depend on the relative balance of evaporation to sublimation (Christner et al., 2017). Additionally, very warm events in summer induce surface melt and sublimation across the Tuto Ice Dome and Greenland ice sheet (Box and Steffen, 2001; Nghiem et al., 2012; van As et al., 2012; Neff et al., 2014), and any of this moisture reaching Thule would be depleted in 18O and 2H and have a higher dxs relative to typical summer ocean-sourced moisture as it originates in a higher and colder location on the ice sheet. Isotopic effects from plant transpiration (Gat and Matsui, 1991; Farquhar et al., 2007; Aemisegger et al.,
Figure 5. Summary of effects of environmental factors affecting water vapor isotope composition at Thule. (a) Illustration of temporal scales when each factor is most impactful. Question marks indicate periods of an uncertain degree of impact. (b) Interactions between environmental factors during NAO conditions, with factors that produce higher δ⁸O and δ²H values and lower dxs values in blue, and factors that produce lower δ¹⁸O and δ²H values and higher dxs values in italicized red. The dashed line refers to the possibility that a low sea ice extent reinforces the NAO phase (e.g., Petrie et al., 2015). (c) Same as (b) but for NAO+ conditions.

2014) are assumed to be very weak and inconsequential due to the sparse plant cover at Thule (Gold and Bliss, 1995).

4.6 Summary of environmental drivers of water vapor isotopes at Thule

To summarize, observations at Thule when the water vapor δ¹⁸O and δ²H values are higher and dxs values are lower are linked to (a) warmer local air temperature, (b) more local marine moisture source, (c) synoptic pattern that favors more southerly flow (i.e., NAO−), (d) sea breeze surface winds, and/or (e) lower evaposublimation. Likewise, lower δ¹⁸O and δ²H values and higher dxs values are linked to (a) colder local air temperature, (b) more distant marine moisture sources, (c) synoptic pattern that restricts southerly flow (i.e., NAO+), (d) katabatic surface winds, and/or (e) higher evaposublimation. These different factors operate on different temporal scales (Fig. 5a). For example, shifts in local versus distant moisture sourcing affect isotopic composition at all timescales, but evaposublimation is mostly important only on shorter timescales like diel cycles.

These five factors should not be seen as independent drivers of isotopic change. Rather, these factors are detectable facets expressed in our observational data of the diverse environmental changes controlled by variations in Baffin Bay sea ice extent and synoptic moisture transport to the Arctic. As such, these factors greatly interact with each other, and isotopic variability cannot and should not be reduced down to a single predominant driver. However, these factors largely interact constructively to enhance relationships between environmental conditions and water vapor isotopic values (Fig. 5b and c). For example, the NAO− enhances southerly advection to Thule, which results in warmer local air temperature and potentially reduced sea ice extent that exposes local waters. Similarly, the sea breeze is enhanced during warmer periods of the year when sea ice extent is low, and it directly supplies local moisture to Thule from Bylot Sound. All these factors support isotopically heavier water vapor isotopes with lower dxs values when sea ice extent is low and synoptic conditions favor southerly flow.

The main exception to these constructive interactions is evaposublimation, which supplies isotopically light vapor with high dxs during warmer periods which would otherwise favor higher δ¹⁸O and δ²H values and lower dxs values as previously described. Due to the ample supply of marine moisture in the coastal Thule setting, evaposublimation is not expected to be a predominant moisture source or primary driver of isotopic variability. However, periods where water vapor isotopes are lighter and dxs values higher than expected for a given temperature may be a good identifier of an evaposublimation effect, and such analysis could be used to better quantify ice sheet vapor flux during large surface melt events.

5 Seasonal changes in the drivers of isotopic variability

The strong annual cycle in water vapor isotopes at Thule (Table 1) is directly tied to seasonal changes in weather and the regional environment driven by the presence or absence of
Figure 6. Spearman correlations of meteorological variables (a–f) with δ¹⁸O (blue solid line) and dxs (red dashed line), with data binned by month. The yellow shading highlights the summer season (JJA) and horizontal dotted lines show ρ values of 0.00 and ±0.50 for reference. Correlation patterns for δ²H (not shown) are very similar to those of δ¹⁸O. Correlation values were calculated using the highest available data resolution, namely 10 min for (a–c), hourly for (d), and daily for (e, f).

local sea ice. While the annual sea ice breakup directly influences vapor isotopic composition by allowing local marine moisture supply, it also produces a cascade of other environmental changes that isotopically alter water vapor at Thule (Fig. 5). Beyond simply affecting the mean values of water vapor isotopes, the dramatic environmental transformations that occur at Thule after sea ice breakup also modify correlative relationships between isotopic and meteorological variables (Fig. 6). As a result of all these factors, the seasonal growth and breakup of the Baffin Bay sea ice is readily identifiable in our water vapor isotope record, particularly in the dxs data (Fig. 2).

5.1 Spring and sea ice breakup

In both spring seasons covered by our record, the isotopic and meteorological variables had a very abrupt shift from typical winter values to typical summer values (Fig. 2). During these shifts that began on 14 May 2018 and 29 April 2019, temperatures rose more than 15 °C in a few hours, followed by a ∼10‰ increase in δ¹⁸O, and a ∼10‰–15‰ drop in dxs over the next few days. Sea ice concentrations and satellite imagery from NSIDC (Fetterer et al., 2017) and MODIS (Hall and Riggs, 2015) show that these abrupt spring shifts were associated with the breakup of sea ice near and to the northwest of Thule and a general reduction in sea ice concentration throughout Baffin Bay (Fig. S4). Similar isotopic responses to sea ice breakup have been previously reported in the Arctic Ocean as well (Klein and Welker, 2016). Before the breakups in 2018 and 2019, the NAO index dropped 2 and 4 points, respectively, and the resulting extreme isotopic shifts create very strong correlations in May between isotopic variables and both Baffin Bay sea ice extent (Fig. 6e) and the NAO index (Fig. 6f).

Sea ice breakup and environmental warming is followed in short order by the first sustained sea breeze developments of the year that also aid the delivery of newly available local vapor to Thule. At this time in late spring, radiative heat loss at night drives a semiconsistent diel wind cycle between afternoon sea breezes and nightly katabatic flow, and synoptic storm systems may bring multiple days when winds predominantly come from either the ocean or over the ice sheet, depending on the relative position of the storm system to Thule. The environmental contrast between the relatively warm open ocean and still frigid Greenland ice sheet and tundra maximize isotopic differences in the two wind regimes, and, as a result, the strongest correlations between isotopic variables and katabatic deviation are observed in April through June (Fig. 6d).

5.2 Summer

The height of summer brings striking changes in isotope–climate relationships. Most notably, the δ¹⁸O and δ²H correlations with air temperature switch from positive to negative in summer months (mean ρnon-summer = +0.38 vs. mean ρsummer = −0.32 for δ¹⁸O) with dxs showing a similar, though opposite, pattern (mean ρnon-summer =
−0.53 vs. mean $\rho_{\text{summer}} = +0.40$) (Fig. 6a). As a result, the highest $\delta^{18}O$ and $\delta^2H$ values and lowest dxs values are not observed when air temperatures are greatest but rather near 0°C (Fig. S3a and d). During the same summer period, isotopic correlations with mixing ratios greatly weaken (Fig. 6b), while correlations with relative humidity strengthen (Fig. 6c). The relationship between air temperature and mixing ratio, strongly correlated with a positive slope through most of the year, decouples in summer, and temperatures warmer than 5°C largely result in no higher water vapor content (Fig. S5).

Summer in Thule is typically cool and humid under shallow marine air from local open seas helped onto land by the sea breeze. As the local ocean never warms much above freezing, most of the warmest periods at Thule (i.e., > 5–8°C) require a high-pressure ridge over Greenland, often associated with southerly airflow up along the western Greenland coast and aided by NAO− conditions. In many cases, the position of this high-pressure ridge favors large-scale downsloping and subsidence off the northwestern ice sheet (Fig. 7). This pattern was exceptionally persistent in summer 2019 (Tedesco and Fettweis, 2020), and this period supplied most of the warmest observations in our database. Moisture arriving at Thule through southerly advection and/or downsloping in these warm events is isotopically lighter and less humid than local marine air as it is more distinctly sourced and also must cross topographic highs. When contrasted against cool local marine air that has isotopically heavier vapor with low dxs values, this produces the observed flip in summer correlations between isotopic species and temperature and the strengthened correlations with relative humidity.

Correlations between isotopes and water vapor mixing ratio greatly weaken in summer because mixing ratios do not clearly differentiate between local marine and southerly advected air. Although the shallow marine air has a higher relative humidity, it is also colder than the southerly advected air and, thus, has a lower maximum mixing ratio. In contrast, the relative humidity of southerly advected air is lower, but its warmer temperature means that the actual water content of the air can be very similar to that of local marine air. As a result, the relationship between temperature and mixing ratio decouples (Fig. S5), and isotopic correlations with mixing ratio approach zero (Fig. 6b).
Two additional processes that emerge in summer may also lead to these relationship changes. First, the very warm conditions that occur during strong ridging over Greenland and enhanced southerly advection to Thule are also associated with intense surface melt and sublimation on the Tuto Ice Dome and greater Greenland ice sheet (Nghiem et al., 2012; van As et al., 2012; Neff et al., 2014; McLeod and Mote, 2016; Ballinger et al., 2019). As the ice sheet surfaces are higher in elevation and colder than Thule, water vapor coming from their snow and glacial ice would be particularly isotopically light with high dxs values (Steen-Larsen et al., 2013; Kopec et al., 2014; Bréant et al., 2019), and this would enhance the isotopic signature of southerly advected air as it passes over the ice sheet on its way to Thule.

Second, the high pressure of these warm events often brings sunny weather to Thule that promotes intense heating of the local tundra and exposed rocky surfaces. Outside of polar night, the water vapor mixing ratio typically follows the daily temperature cycle and peaks in the early afternoon. However, on days when the mean temperature is greater than 10°C, the mixing ratio drops as temperature rises and hits its minimum daily value around local noon. This loss of water vapor suggests that daytime heating on the warmest days induces convection (Duynerke and van den Broeke, 1994) that vertically mixes drier air aloft down to the shallow marine surface layer. This results in a net upward water vapor transport (Sherwood et al., 2010; Kiemle et al., 2013; Homeyer et al., 2014), and the drier air that mixes downward from the free troposphere brings vapor with lower δ18O and δ2H and higher dxs values (Bailey et al., 2013). The surface heating and drying during midday could also increase local evaporation from the land and surface waters that, while not enough to overcome the net upward vapor transport from convection, would be an additional source of isotopically light water vapor to near surface moisture.

5.3 Autumn and sea ice growth

The transition to winter-type isotopic and meteorological conditions is much more gradual than the spring transition at sea ice breakup (Fig. 2), in accordance with the comparatively steady growth of sea ice in autumn. The δ18O and δ2H values slowly decrease while dxs values slowly increase from October through November, and values through winter do not generally overlap summer values. While isotopic correlations with sea ice extent are near zero throughout summer (likely because most changes to sea ice extent at this time are occurring too far north or west to directly impact Thule), the autumnal growth of sea ice results in moderately strong correlations in October (Fig. 6e) as the region of sea ice formation again affects Thule. Although small in magnitude, correlations with the NAO index also strengthen in autumn (Fig. 6f), and this is likely due to the close relationship between NAO phase, sea ice extent, and local marine moisture availability. The return of true day–night cycles in September creates a surface wind regime similar to that of spring, and correlations between isotopes and katabatic deviation strengthen in the autumn months until the sea breeze ceases due to polar night in November (Fig. 6d).

5.4 Winter

During the polar night of winter, the water vapor isotopic system at Thule is relatively simple because northern Baffin Bay and all surface waters are frozen over. Mean values for δ18O and δ2H in winter are lower and dxs values higher than other seasons in the absence of a local marine moisture supply (Fig. 3b), and isotopic variability largely reflects extratropical cyclone impacts and broad synoptic flow changes. Strong winter correlations between isotopic species and both air temperature and mixing ratio (Fig. 6) reflect this as warmer and moister air transported to Thule produces higher water vapor δ18O and δ2H values and lower dxs values.

The dxs values in winter are also notably more variable than during summer (Fig. 2). Since nearly all moisture arriving at Thule in summer has to pass over the open water of Baffin Bay, vapor exchange and uptake probably helps stabilize the summer dxs variability. In contrast, when Baffin Bay is frozen, the dxs values from different moisture sources may be preserved better and contribute to the higher winter variability in dxs values. More focused back-trajectory analysis on winter dxs variability in the future may help quantify the variability and effects of different moisture sources.

Interestingly, δ18O and δ2H have very strong positive correlations with the late winter sea ice extent in February and March, while dxs has very strong negative correlations with sea ice extent in February (Fig. 6e). These relationships are opposite to those observed in spring and autumn, when the presence or absence of sea ice plays a clear isotopic role through local water availability. It is not entirely clear what is driving the strong correlations in late winter, as sea ice during these months is near its maximum extent and largely complete in coverage near Thule. These unusual correlations may be due to coincidental extreme events. Shortly after the maximal extent of sea ice was reached in 2018, a period of enhanced southerly moisture advection supplied abnormally heavy water vapor isotopes with very low dxs values. In 2019, the isotopes also became unusually heavy with low dxs for 2 weeks after the peak in sea ice extent, but no associated increase in southerly moisture was identified, and the root cause is unclear. With only 2 years of records, it is difficult to conclude whether the observed cumulative strengths in late winter accurately represent a true change in isotopic character after a certain sea ice threshold is exceeded, or if it is simply due to coincidental occurrence with two unusual late winter weather events. Additional years of observation may help clarify this uncertainty.
5.5 Cold season moisture pulse events

During winter months, rapid shifts to extremely low NAO indices often coincide with intense poleward transport of southerly heat and moisture to Thule that lasts 1–5 d. We refer to these distinct episodes as moisture pulse events as they appear very clearly in the Thule mixing ratio time series (Fig. 2). During these events, air temperature rises 10–15 °C and peaks near or above freezing, while water vapor concentrations can reach 4 times greater than mean winter levels. These pulses appear to notably reduce or slow the growth of sea ice across Baffin Bay (Fig. 2), although the sea ice stays largely intact in the northern reaches near Thule. Although not all these events meet the defined criteria of an atmospheric river (Mattingly et al., 2018), they all have similar impacts to an atmospheric river event due to their anomalously high moisture and heat transport.

Unsurprisingly, such extreme weather changes are reflected by impressive water vapor isotopic responses. During these moisture pulse events, the $\delta^{18}O$ and $\delta^2H$ rise 6‰–10‰ and 50‰–100‰, respectively, to reach values more typical of late spring and early summer. Concurrently, the dxs drops 15‰–25‰, and the minimum dxs values observed in these moisture pulse events match or exceed the minimum values observed at the height of summer. As sea ice coverage prevents the uptake of isotopically heavy moisture with low dxs values from local waters during these moisture pulse events, the anomalous isotopic values must signify the presence of deep southerly moisture transport (Bonne et al., 2015). A focused analysis of these events is currently underway.

The rapid isotopic and environmental shifts coinciding with sea ice breakup in 2018 and 2019 also fit the general pattern of a moisture pulse event, except the isotopic and meteorological variables do not revert back to their preceding values after 1–5 d. Indeed, the shift in 2019 is associated with an atmospheric river event impacting western Greenland and values after 1–5 d. Indeed, the shift in 2019 is associated with an atmospheric river event impacting western Greenland and peaks near or above freezing, while water vapor concentrations can reach 4 times greater than mean winter levels. These pulses appear to notably reduce or slow the growth of sea ice across Baffin Bay (Fig. 2), although the sea ice stays largely intact in the northern reaches near Thule. Although not all these events meet the defined criteria of an atmospheric river (Mattingly et al., 2018), they all have similar impacts to an atmospheric river event due to their anomalously high moisture and heat transport.

6 Diel cycles

6.1 Diel cycles overview

Most polar sites report diel cycling of water vapor isotopes in summer (Steen-Larsen et al., 2013; Bastrikov et al., 2014; Bonne et al., 2014; Kopec et al., 2014; Casado et al., 2016; Ritter et al., 2016; Bréant et al., 2019), but this cycling is attributed to different causes including katabatic wind cycling (Kopec et al., 2014; Bréant et al., 2019), vapor exchange between snow and air (Steen-Larsen et al., 2013; Casado et al., 2016), and dew formation (Bastrikov et al., 2014). In contrast to these sites, Thule exhibits relatively limited isolated isotopic diel cycling in all months except March, with total cycle magnitudes of less than 1‰ for both $\delta^{18}O$ and dxs, even when observations with possible cyclonic system impacts (i.e., lower than average station pressure) are excluded. Additionally, the existence of these small diel cycles is only clearly evident when the larger synoptic variability is removed by averaging multiple days together. This subdued diel cycling is probably due to Thule’s moderate coastal climate (Bréant et al., 2019; Bonne et al., 2020), while lengthy periods of midnight sun and polar night also reduce the day–night contrasts that power diel cycles at lower latitude sites.

Looking broadly at the entire Thule isotopic record, the magnitude of isotopic diel cycles is not a significant driver of isotopic variability. However, we believe that analyzing these cycles gives better insight into the broader environmental controls on water vapor isotopes, including those at synoptic and annual scales. Based on observed isotopic and meteorological patterns, we divide diel cycles at Thule into four regimes, namely polar night, March, summer, and transition (Fig. 8). These regimes are superimposed on the broader seasonal changes in isotopic variability and control previously discussed. Out of the major drivers of isotopic change at Thule, air temperature, surface wind regime, and evaporation-sublimation vary enough on hourly timescales to contribute to diel isotopic cycles, while the NAO and local marine moisture availability are expressed at longer timeframes and/or not on a consistent daily cycle. We note that the magnitude of daily change in the isotopic variables is typically within the confidence intervals of the mean with the exception of the March regime. We believe the daily patterns of variable change are still informative, but a discussion of these trends should be viewed as being more speculative than other conclusions made in this study.

6.2 Polar night regime

The polar night regime includes the months from October through February, when daylight is very short (October and February) or nonexistent (November–January). With the lack of a strong insolation cycle, meteorological variables change little over 1 d, and winds are strong and consistently katabatic. Without any diel changes in the factors that drive iso-
Figure 8. Diel patterns in isotopic (a, b) and meteorological (c–g) variables in Thule, Greenland, showing the mean hourly value (solid thick line) and 95% confidence intervals of mean estimates (colored shading) for each diel regime. Diel patterns for $\delta^{18}$O (a) and are not shown. Variables other than wind speed and katabatic deviation are expressed as deviations from the mean value in each regime, with zero deviation indicated as a dashed horizontal line. Note that because the winds in Thule are binary (katabatic or sea breeze), the diel means of katabatic deviations should be viewed as a probability of being katabatic or sea breeze at a given time and not that the winds are smoothly transitioning from east to west or vice versa. Additionally, because katabatic deviations increase in value in both directions away from 100° (e.g., winds from 80 and 120° both have a katabatic deviation of +20°), the natural variance in purely katabatic winds produces an average value of $\sim$ 25° rather than 0°.

6.3 March regime

The observed isotopic cycles in March (maximum diel range $\delta^{18}$O: 2.4‰ and $\delta^{2}$H: 16‰; dxs – 5.1‰) are by far the largest magnitude of any month of the year at Thule (Fig. 8), although similar cycles with smaller amplitudes are also observed at the end of February. This heightened isotopic response appears due to several coinciding factors, namely a maximized daily insolation cycle near the equinox, an ex-
tensive snowpack to supply water vapor, and the enhanced impact of snow-supplied vapor in the very cold and dry environment (aided by the maximal seasonal extent of sea ice that limits external moisture input). This environment approximates the summer ice sheet setting of two other water vapor isotope observation sites at North Greenland Eemian Ice Drilling (NEEM), Greenland (Steen-Larsen et al., 2013), and Dome C, Antarctica (Casado et al., 2016), and Thule’s March isotopic diel regime appears similar to the cycles reported from those sites.

Moisture at Thule in March predominantly arrives from distant transport through katabatic flow, with resulting low $\delta^{18}O$ and $\delta^2H$ values and high dxs values. However, daytime heating of the snow surface across the landscape promotes the release of water vapor held between snow grains that have equilibrated with the local snow isotopes (Steen-Larsen et al., 2013; Casado et al., 2016). Because most snow in Thule falls in late autumn to early winter and is sourced from Baffin Bay, the equilibrated vapor is isotopically heavier and has a lower dxs than the katabatic-supplied moisture, producing the observed $\delta^{18}O$ and $\delta^2H$ peak and dxs minimum at around local noon. As insolation and air temperature drop in the evening, this vapor release ceases and potentially reverses, allowing the vapor isotopic composition to revert to one of katabatic origin with lower $\delta^{18}O$ and $\delta^2H$ and higher dxs values. The lack of a large drop in relative humidity coinciding with the midday temperature rise suggests that additional vapor is being supplied, and the local snowpack is the only likely source with all local waters frozen. A large isotopic diel cycle predominating in spring has also been reported in the Lena River delta and similarly attributed to the release of water vapor from the preceding winter snowpack (Bonne et al., 2020).

A parallel isotopic regime does not reappear around the autumnal equinox at Thule or at the Lena Delta site (Bonne et al., 2020). This is likely due to little to no extensive snowpack and a much higher water vapor content from warmer temperatures and open nearby seas that buffer any minor potential input from a snowpack vapor exchange. One exception to this lack of autumnal diel cycles, unique in the Thule record, occurred from 20–25 September 2017 when an isotopic diel cycle, very similar to the March regime, appeared (Fig. S6). This cycling occurred shortly after a snowfall and coincided with a diel surface wind cycle where the katabatic winds calmed each afternoon but did not switch to a sea breeze. This appears to have allowed vapor released from the recent snow to raise isotopic values in the afternoon before the returning katabatic winds mixed the snow-derived moisture out and dropped isotopic values in the evening. In other September periods with a similar wind regime but without a recent snow cover, no isotopic cycling was recorded.

6.4 Summer regime

In the summer months (June–August), the midnight sun reduces the diel insolation cycle. Combined with the moderating influence of fully ice-free local seas, most meteorological variables are more stationary over the course of a day. With this reduced environmental variability, diel cycles in isotopes are subdued. The subtle daily patterns in $\delta^{18}O$ and dxs that do exist appear largely attributable to evaposublimation cycles, i.e., daily warming supplies low $\delta^{18}O$ and high dxs water vapor through increased evaposublimation, with the resulting $\delta^{18}O$ minimum and dxs maximum around midday (Fig. 8). Increased boundary layer mixing at midday may also supply moisture from the free troposphere with low $\delta^{18}O$ and high dxs values to produce a similar isotopic effect to evaposublimation.

At Kangerlussuaq (Kopec et al., 2014) and Dumont d’Urville (Bréant et al., 2019), the local surface winds have a clear diel cycle between daily sea breezes and nightly katabatic winds in summer. At Thule, the sea breeze has a diel cycle where it strengthens in the afternoon and weakens at night (Fig. 8g), but there is not a full switch back to katabatic flow at night in summer. This dominance of the sea breeze at Thule is likely due to 24 h summer insolation that can fuel a sea breeze even at “night”, while Kangerlussuaq and Dumont d’Urville still have true night at their lower latitudes. As a result, sea-breeze-supplied local water is likely present throughout a typical day and night at the height of summer in Thule, and the nightly weakening of the sea breeze has a limited isotopic effect without a full shift to katabatic flow.

6.5 Transition regime

The seasonal transition months of April, May, and September have a true day–night cycle that gives meteorological variables a greater diel amplitude than the summer or polar night regimes (Fig. 8). Yet, unlike March, these transitional months are much warmer and more humid with moisture supplied from an open or opening Baffin Bay. Winds exhibit a strong diel cycle as surface heating leads to afternoon sea breeze development (Fig. 8g), and radiative heat loss over the ice sheet at night strengthens katabatic flow. We might expect a clear diel response in water vapor isotopes due to these heightened diel meteorological cycles, but the isotopes have little to no daily cycle (Fig. 8a and b).

This limited isotopic cycling is likely due to the competing effects of diel surface wind regime and evaposublimation cycles. While the sea-breeze-sourced vapor coming directly off Bylot Sound in the afternoon is isotopically heavy, midday heating also supplies isotopically light moisture from evaposublimation and boundary level mixing. Cooling at night brings a wind shift to isotopically lighter katabatic flow but also suppresses evaposublimation and boundary level mixing, while promoting dew condensation. As a result, no clear diel isotopic cycle emerges, although the subtle late evening
peak in $\delta^{18}O$ may arise because evaporosublimation rates are reduced while the sea breeze is still ongoing.

7 Implications

The improved understanding of water vapor isotopic variability granted by our Thule observations can aid the interpretation of regional ice cores. The primary drivers of isotopic changes in water vapor are likely to also affect local precipitation, although there are many additional processes involved in the transfer of an isotopic signature from vapor to precipitation and eventually to ice core that must be considered (Steen-Larsen et al., 2014; Casado et al., 2018; Madsen et al., 2019). Changes in $\delta^{18}O$ and $\delta^2H$ have long been used to reconstruct climate change from deep ice cores, which are typically interpreted as local temperature variability in Greenland (e.g., Dansgaard et al., 1969; Grootes and Stuiver, 1997; Johnsen et al., 2001). While air temperature is strongly correlated with isotopic change at Thule, much of this strength appears to arise as a common response to seasonal change between air temperature and sea ice extent. Air temperature might still be robustly reconstructed from isotopic archives based on our observed strong correlations, but the lack of a causative relationship risks misinterpretation if applied back, far through time, when the local environment may have significantly changed. Of particular warning is our observation that the basic relationship between air temperature and water vapor isotopes inverts in summer, leading to very different isotopic interpretations, depending on the season. While this may be a local and/or coastal effect that is not expressed on ice sheets (Ballinger et al., 2019), we still advise caution.

More recently, additional consideration has been given in ice core analysis, to secondary isotopic variables like $\Delta$Xs and other environmental drivers such as moisture source, sea ice extent, and atmospheric circulation (e.g., Grumet et al., 2001; Vinther et al., 2003; Steffensen et al., 2008; Landais et al., 2018; Kopec et al., 2019). While our 2 year Thule record is too short to statistically determine the strongest drivers of interannual isotopic variability, changes in the duration of sea ice coverage and mean NAO phase appear most likely to control year-to-year differences in mean isotopic composition. In line with recent interpretations in regional ice cores (Osterberg et al., 2015), our results suggest that past periods with decreased sea ice extent in Baffin Bay will have increased local marine moisture sourcing that produces higher $\delta^{18}O$ and $\delta^2H$ and lower $\Delta$Xs values.

In recent years, Baffin Bay has had later sea ice freezes, earlier breakups, and a decrease in overall sea ice extent that accounts for 22% of all recently observed losses in March Arctic sea ice (Onarheim et al., 2018). We expect that the resulting changes in the regional hydroclimate will be isotopically preserved in future glacial ice in Greenland. However, predicting the isotopic responses to these changes is challenging. The future of the Nares Strait ice bridge between Greenland and Ellesmere Island is one potential complication; should it consistently fail to form in a warmer world, the North Water Polynya may see increased sea ice cover that slows the overall sea ice loss impact on Thule (Barber et al., 2001; Puntag et al., 2016; Vincent, 2019). Reductions in Arctic sea ice coverage and duration may also shift precipitation seasonality (Kopec et al., 2016) and atmospheric circulation (McLeod and Mote, 2016; Ballinger et al., 2018; Francis et al., 2018), changing the annual isotopic balance of precipitation. More open winter waters in Baffin Bay could decrease annual $\delta^{18}O$ and $\delta^2H$ values as the relative fraction of isotopically lighter winter precipitation events increases, but this may be counterbalanced in part by the greater sourcing from isotopically heavy local waters.

Enhanced southerly moisture advection aided by NAO−conditions would also favor higher $\delta^{18}O$ and $\delta^2H$ and lower $\Delta$Xs values in northwestern Greenland, though some caution might be warranted for isotopic complications from greater ice sheet surface melt and vertical atmospheric mixing similar to those we have observed during summer NAO−phases. Additionally, the positive relationship between the NAO phase and Baffin Bay sea ice extent (Mysak et al., 1996; Grumet et al., 2001; Stern and Heide-Jørgensen, 2003) makes it difficult to quantifiably split their influences on regional water isotopes. The extreme isotopic values we observe during moisture pulse events suggest that any changes in their frequency and magnitude (and more generally of related atmospheric river events) will have an outsized effect due to the sheer volume of moisture and precipitation they can bring to Greenland. For highly resolved ice cores, it may be possible to identify particularly strong moisture pulse events or years with a high frequency of these events as extreme minima in $\Delta$Xs values. Taken as a whole, it is clear that, in the absence of additional clarifying evidence from other ice core proxies or nearby records, isotopic interpretations should be cautious in assigning the cause solely to one individual environmental factor without further analysis to tease the many highly integrated potential factors apart.

8 Conclusions

Our 2 years of water vapor isotope monitoring in northwestern Greenland have produced a record of unprecedented extent and resolution for high Arctic Greenland and one of the longest records of any polar site reported to date. The variability in water vapor isotopes at Thule is explained by five interrelated environmental mechanisms, namely local air temperature, local marine moisture availability, the NAO, surface wind regime, and evaporosublimation. The relative importance of each mechanism changes over the course of the year and, overall, produces a clear annual isotopic cycle that is closely linked to sea ice extent in Baffin Bay and synoptic polar moisture transport. On top of these seasonal trends, local environmental drivers such as sea breeze development,
vapor supply from surface water and snow, and convection can subtly modify near surface water vapor isotopes through diel cycles.

This analysis highlights the importance of local geography and climate in isotope systematics. Compared to other high-latitude water vapor isotope studies, Thule is more substantially impacted by sea ice fluctuations, both seasonally and interannually. This results in a clearer identification of the isotopic effects of sea ice extent, which is particularly valuable for the interpretation of deep ice cores. Additionally, some aspects of isotopic variability, such as the changes in correlative relationship strength and direction during spring and summer, are so far uniquely reported for Thule, and it is unclear at this time if these seasonal relationship patterns are present at other polar sites. As a result, conclusions based on data at one high-latitude site should not be applied indiscriminately to other sites without extensive validation that such a comparison is warranted.

Caution should also be exercised when making strong conclusions about isotopic controls based on observations of a short duration (i.e., a single year or less). If data are only taken during a single season, any observed relationships may be specific only to that season and should not be assumed to be applicable to the entire year. For variables with relatively long-term variability, such as teleconnection indices, data from a single year are likely not long enough to reveal fully robust correlations. In our Thule data, we were fortunate that the synoptic patterns were different enough between 2018 and 2019 to allow useful analytical comparisons of the NAO at monthly and seasonal resolutions. However, even 2 years of data are not enough to fully clarify the relationships between teleconnections and water vapor isotopes for all months, and additional years of isotopic data from continued observation are needed for an improved understanding.

Moving forward, this isotope and meteorological database offers many opportunities for more in-depth and focused analyses of specific weather events and atmospheric patterns. As previously stated, our identified moisture pulse events are a focus of additional research, and we encourage the use of our data in other focused case studies. An expansion of our quasi-climatology approach could be used to model moisture source environments for direct comparative analysis with the water vapor isotope data, which can help resolve existing questions relating to the nature of the dxs–sea surface temperature relationship in polar regions (Vimeux et al., 1999; Aemisegger and Sjolte, 2018).

Water vapor isotope observations at Thule continue, and additional months and years of data will help refine and verify our conclusions made here. The long temporal coverage of our database makes it an excellent option for validating high-resolution isotope-enabled simulations of air–ice and air–sea interactions in the Arctic. These results from Thule will help greatly in interpreting isotopic variability at other sites in the Arctic Water Isotope Network (Welker et al., 2019), and future collaborative analysis across this network will allow tracking of the effects of synoptic weather patterns on water vapor isotopes in real time across the circumpolar region.

Data availability. Isotopic and meteorological data and R analytical scripts are available at https://doi.org/10.18739/A20000151 (Akers et al., 2020). Meteorological data for SMT are also available at https://doi.org/10.1594/PANGAEA.895059 (Muscari, 2018) and in associated follow-up data sets by Giovanni Muscari, which are also available from PANGAEA. Meteorological data for THU are available upon request from the 821st Weather Squadron of the US Air Force.

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Author contributions. JMW and ESK installed the Picarro L-2130i. PDA and BGK installed the dry air calibration system and performed maintenance on the L-2130i. PDA wrote the code in R to organize and calibrate the data and also for statistical analyses. KSM performed the back-trajectory and water vapor transport analyses. DC performed the cross-correlation analyses. PDA prepared the paper, with contributions from all coauthors.

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