



UiT The Arctic University of Norway

Faculty of Science and Technology, Department of Geosciences

Holocene precipitation seasonality on Svalbard and in Northern Fennoscandia

Reconstructed using organic geochemical and stable isotope proxies

Sofia Elisabeth Kjellman

A dissertation for the degree of Philosophiae Doctor

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Front page image: Ringhorndalen on northern Spitsbergen, Svalbard, seen from helicopter in July 2018. Jodavannet, one of the studied lakes, is seen in the left of the photo.
(Photo: Sofia Elisabeth Kjellman)

Abstract

The high latitudes are experiencing faster climate change than the global average. Feedback mechanisms between the atmosphere, cryosphere and the oceans amplify the warming, and intensify the water cycle. Arctic precipitation is projected to increase because of enhanced winter evaporation due to diminishing sea ice, increased poleward moisture transport in summer, and increased atmospheric moisture content in summer and early fall. Large model uncertainties and different seasonal expressions of these processes emphasize the need to improve our understanding of long-term spatial and temporal variability in the water cycle.

Stable isotopes of oxygen and hydrogen ($\delta^{18}\text{O}$ and $\delta^2\text{H}$) in precipitation are sensitive to the conditions at the moisture source, atmospheric temperature, and the moisture travel history, and are therefore widely used tracers of past and current changes in the water cycle. This study focuses on two aspects of water isotope proxies: 1) reconstructing Holocene precipitation seasonality on Svalbard using $\delta^2\text{H}$ of leaf wax *n*-alkanoic acids from lake sediments (Papers I and III), and 2) exploring the variable influence of inflow (precipitation) seasonality and evaporation on the modern isotopic variability of lake waters in northern Fennoscandia (Paper II). Seasonal distribution of precipitation is important for example because winter precipitation is one of the main controls on glacier mass balance. Understanding the seasonal distribution of precipitation in the past might, therefore, help decipher the drivers of glacier variations.

On Svalbard, we targeted four lakes along a climatic gradient from the relatively warm and humid west coast to the relatively cold and arid Nordaustlandet. We focused on the C_{22} and C_{28} *n*-alkanoic acids, interpreted to reflect lake and soil water, respectively. Our results suggest that Early and Middle Holocene summers were warm with high evapotranspiration, and/or received more proximal moisture. On northern Spitsbergen, the proportion of winter relative to summer precipitation increased in the Early Holocene, as the oceans warmed, and the sea ice retreated. After *c.* 6 cal. kyr BP, summers became cooler with less evapotranspiration, and/or more distally derived moisture. The proportion of winter precipitation decreased as the ocean surface cooled, and the sea-ice cover increased, but periods of increased winter precipitation still occurred. Varying lake hydrology impacted the seasonality reflected in the aquatic leaf waxes; therefore, a good understanding of the individual lake systems is crucial for robust proxy interpretations.

In northern Fennoscandia, surface water was sampled from 135 lakes, from the Norwegian Sea to the Bothnian Bay. We find that both coastal and inland lakes are sensitive to distillation during moisture transport, and that lakes farther from the Norwegian Sea additionally are affected by evaporation. We estimated the inflow isotopic composition for evapo-concentrated lakes using a Bayesian method and compared the values for lakes sampled in both 2019 and 2020. It is suggested that more ^2H -depleted values in 2020 reflect that the lakes received relatively more winter precipitation in 2020, related to strong westerlies bringing greater-than-average amounts of North Atlantic moisture. This implies that evapo-concentrated lakes experience isotopic variability due to both evaporative enrichment and variable inflow isotopic composition, and that coastal lakes are better targets for precipitation reconstructions.

We find that summer precipitation variability across Svalbard follows changes in summer insolation and temperature, whereas the seasonal distribution of precipitation on northern Svalbard is controlled by regional ocean surface conditions, with more winter precipitation during periods of strong Atlantic water advection and reduced sea-ice cover. The inflow isotopic composition to Fennoscandian lakes is influenced by variable strength of the westerlies, by affecting the relative proportion of winter and summer precipitation entering the lake. These results highlight the impact of North Atlantic conditions on the seasonal precipitation patterns on Svalbard and in northern Fennoscandia.

Plain language summary

As the Arctic warms, it is also getting wetter. Exactly why, and by how much is difficult to predict due to lack of long-term observations and complex interactions between the atmosphere, cryosphere (e.g., sea ice and glaciers), and the oceans. The main reason for Arctic precipitation increases is retreating sea ice. The sea ice acts as a lid that prevents evaporation from the ocean surface. In a warmer climate, more winter sea ice will disappear, resulting in more available moisture, and more winter precipitation. Furthermore, the Arctic receives moisture from the mid-latitudes, and this transport is projected to increase, mostly in summer. Additionally, the moisture content in the atmosphere will increase, as warm air can hold more water vapor.

To better understand the climate system, we can use modern and geological data. Stable oxygen and hydrogen isotopes ($\delta^{18}\text{O}$ and $\delta^2\text{H}$) in precipitation are sensitive to changes in the conditions at the moisture source and in atmospheric temperature and circulation patterns and can therefore provide information about various processes in the water cycle. Lakes are excellent climate archives since sediments (e.g., organic mud, silt, sand) accumulate on the lake floor over time, and hold information about the environmental conditions at the time of deposition. Plants living in and around the lake use lake and soil water when they produce leaf wax – a fatty coating to protect against water loss – which incorporates the hydrogen isotopic signal of the source water.

In this thesis, lake sediments are combined with modern lake water and precipitation measurements to study water cycle changes on Svalbard and in northern Fennoscandia (i.e., northern Norway, Finland, and Sweden), with a focus on precipitation seasonality. This is an important aspect for the mass balance of glaciers since the amount of winter precipitation together with the summer temperature decide if a glacier grows or shrinks.

Our results suggest that Holocene variability in summer precipitation on Svalbard follows changes in summer insolation and temperature. Northern Svalbard received more winter precipitation during times of increased inflow of Atlantic water, reduced sea-ice extent, and more available moisture. Modern lake water isotope data from northern Fennoscandia suggest that the lake water isotopic composition changes in response to varying proportion of summer and winter precipitation, which partly depends on the atmospheric circulation patterns over the North Atlantic, and due to evaporative enrichment. This thesis highlights the potential of using organic geochemical and stable isotope proxies to study seasonal changes in precipitation.

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I would like to thank Lena Håkansson for being part of the supervision team and UNIS for having me as a guest PhD student in the beginning of the project. I would also like to acknowledge the co-authors on the scientific papers presented in this thesis: Sandrine Duboscq, Allison Cluett, and Owen Cowling for sample and data processing; Nick McKay for LiPD support and troubleshooting; Wes Farnsworth, Lis Allaart, Skafti Brynjólfsson, and Ólafur Ingólfsson for the many hours spent in the field and for commenting on Paper I and III. I am also grateful to Mike Retelle and others who participating in fieldwork during lake sediment coring, and to John Arne Opheim, Marie Bulínová, and the rest who helped me collect precipitation and lake water samples. Thank you to Owen for training and help in the lab, and for answering questions about lab procedures and data. Thanks to Katie Lovell and Kayla Hollister for sample processing, and the rest of the UB Stable Isotope and Biogeochemistry Lab (Allison, Meg, Devon, and others) for welcoming me into the group during my stay at UB, for sharing code, and for discussing methodology and data.

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part of my bubble during the pandemic, and for your company and encouragement during the last months of writing.

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Preface

This thesis is the result of a PhD project carried out at UiT The Arctic University of Norway between February 2018 and July 2022. The four-year doctoral position was funded by the Faculty of Science and Technology at UiT. Additional funding for fieldwork, laboratory analyses, conference attendance, and mobility was granted by the Svalbard Environmental Protection Fund (17/101 to Schomacker and 20/36 to Schomacker and Kjellman), Olle Engkvists Stiftelse (204-0129 to Schomacker and Kjellman), Helge Ax:son Johnsons Stiftelse (to Kjellman), Nansenfondet (to Schomacker), and the National Science Foundation (EAR-IF 1652274 to Thomas).

The primary tools used in this project are organic geochemical and stable isotope proxies. Sediment cores were collected within other projects between 2015 and 2019, and lake water and precipitation sampling were initiated in 2018 and 2019 as part of this study. In 2018, I spent two months in the University at Buffalo (UB) Organic and Stable Isotope Biogeochemistry Laboratory (OSIBL), learning techniques for extracting and analyzing lipid biomarkers from lake sediments. A second, four-months long, stay at UB was planned in 2020 but cancelled due to the Covid-19 pandemic. The remaining organic geochemical analyses were carried out by technical staff and students in Dr. Thomas's lab at UB. Water samples were analyzed at OSIBL and at the Facility for advanced isotopic research and monitoring of weather, climate, and biogeochemical cycling (FARLAB), University of Bergen.

To fulfil the 25% department and teaching duties, I taught exercises, instructed during field excursions, and provided feedback on student reports on bachelor course GEO-2003 Quaternary Geology (2019-2021) and graduate course GEO-3136/8136 Practical geochronological methods in marine, terrestrial Quaternary Geology and geohazards (2019-2022). I also co-supervised two master students in 2019 and 2020. Part of the duty work included organizing the department's Friday Seminar series (2019-2020), and science communication through establishment of the department's Instagram account (2018-2020), giving presentations to high school students and organizing Geology Day in Tromsø (2020).

To complete the instruction component equivalent to 30 ECTS, I attended GEO-8136 Practical geochronological methods in marine, terrestrial Quaternary Geology and geohazards at UiT, INTPART Arctic Field Summer School 2018: Arctic Coastal Environments in Rapid Transition, in Utqiagvik, Alaska (organized by University of Alaska Fairbanks in collaboration

with UiT and University of Calgary, funded by the Research Council of Norway), and AG-848 Arctic Late Quaternary Glacial and Marine Environmental History at UNIS in 2018. In 2019, I took the mandatory course SVF-8600 Philosophy of Science and Ethics at UiT. Based on the essay I wrote for the ethics course, I later that year published a career column in the world-leading journal *Nature* (Kjellman, 2019).

Over the course of my PhD education, I participated and presented results related to the thesis work at the following international conferences: Joint International Paleolimnology Association and the International Association of Limnogeology (IPA-IAL) Meeting 2018 in Stockholm; 49th International Arctic Workshop in Stockholm, 2019; American Geophysical Union (AGU) Fall Meeting 2020, online; 2nd International Processes and Palaeo-Environmental changes in the Arctic: From Past to Present (PalaeoArc) Conference 2021, online; 35th Nordic Geological Winter Meeting in Reykjavík, 2022. Additionally, I have presented my work at the UiT Geoscience Research Academy of Tromsø (GReAT) Annual Meetings in 2019, 2020, and 2021, and at the Norwegian Research school on changing climates in the coupled earth system (CHESS) Annual Meeting in 2020.

Beyond the thesis work, I have over the course of my PhD program published one additional research paper as a first author (based on my master's thesis; Kjellman et al., 2018) and contributed to seven publications as a co-author (Schomacker et al., 2019; Allaart et al., 2020; Voldstad et al., 2020; Allaart et al., 2021a; Allaart et al., 2021b; Alsos et al., 2021; Vick et al., 2022), whereof two were associated with master's thesis projects that I co-supervised.

The PhD project resulted in three first-authored scientific articles, contributing new knowledge about Arctic precipitation. In this work, we generated seasonal Holocene precipitation isotope records from four lake sediment sequences from Svalbard and investigated modern lake water isotope systematics in northern Fennoscandia.

This thesis consists of an introduction and a summary of the three following research articles:

- I. **Kjellman, S. E.**, Schomacker, A., Thomas, E. K., Håkansson, L., Duboscq, S., Cluett, A. A., Farnsworth, W. R., Allaart, L., Cowling, O. C., McKay, N. P., Brynjólfsson, S., & Ingólfsson, Ó., 2020. Holocene precipitation seasonality in northern Svalbard: influence of sea ice and regional ocean surface conditions. *Quaternary Science Reviews*, 240, 1063882.
<https://doi.org/10.1016/j.quascirev.2020.106388>.

Data related to **Paper I** are freely available at the NOAA/World Data Service for Paleoclimatology website: <https://www.ncdc.noaa.gov/paleo/study/29852>.

- II. **Kjellman, S. E.**, Thomas, E. K., & Schomacker, A., 2022. Arctic and sub-Arctic lake water $\delta^2\text{H}$ and $\delta^{18}\text{O}$ along a coastal-inland transect: implications for interpreting water isotope proxy records. *Journal of Hydrology*, 67, 127556.
<https://doi.org/10.1016/j.jhydrol.2022.127556>.

Data related to **Paper II** are freely available in the [Supplementary Information](#) and in the Water Isotopes Database at www.waterisotopes.org (Project IDs 00292 (surface waters) and 00293 (precipitation)).

- III. **Kjellman, S. E.**, Thomas, E. K., Schomacker, A., Farnsworth, W.R., Cowling, O.C., Allaart, L., & Brynjólfsson, S. Holocene precipitation seasonality along a climatic gradient from western Spitsbergen to Nordaustlandet, Svalbard. In review for *Palaeogeography, Palaeoclimatology, Palaeoecology*.

Data related to **Paper III** will be freely available at the NOAA/World Data Service for Paleoclimatology website and in the Water Isotopes Database.

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Abbreviations

ACL	average chain length
AMS	accelerator mass spectrometry
AO	Arctic Oscillation
cal. kyr BP	calibrated thousand years before present (BP = 1950)
CRediT	Contributor Roles Taxonomy
DCM	dichloromethane
DEM	digital elevation model
d-excess or dxs	deuterium excess
ELA	equilibrium line altitude
ECTS	European Credit Transfer and Accumulation System
FAME	fatty acid methyl ester
FARLAB	Facility for advanced isotopic research and monitoring of weather, climate, and biogeochemical cycling
FID	flame ionization detector
GC	gas chromatograph
GMWL	Global Meteoric Water Line
GNIP	Global Network of Isotopes in Precipitation
IRMS	isotope ratio mass spectrometer
LEL	local evaporation line
LIA	Little Ice Age
LiPD	Linked PaleoData
LMWL	Local Meteoric Water Line
LOI	loss on ignition
MAAT	mean annual air temperature
MAP	mean annual precipitation
m a.s.l.	meters above sea level
MeOH	methanol
NAO	North Atlantic Oscillation
OSIBL	The University at Buffalo Organic and Stable Isotope Biogeochemistry Laboratory
OIPC	Online Isotopes in Precipitation Calculator

RCP8.5	Representative Concentration Pathway 8.5 W/m ² radiative forcing in 2100 = emission scenario “business as usual”
<i>seda</i> DNA	sedimentary ancient DNA
SEM	standard error of the mean
TLE	total lipid extract
UB	University at Buffalo
UiT	UiT The Arctic University of Norway
UNIS	The University Centre in Svalbard
VSMOW	Vienna Standard Mean Ocean Water
WS-CRDS	Wavelength Scanned Cavity Ring Down Spectrometer
XRF	X-ray fluorescence
δ ² H	stable hydrogen isotope ratio (subscript letters indicate the source: aq = aquatic plant waxes; I = lake water inflow; lake = lake water; precip = precipitation; terr = terrestrial plant waxes)
δ ¹⁸ O	stable oxygen isotope ratio
¹⁴ C	radiocarbon

1. Introduction

1.1. Arctic precipitation changes

The Arctic region is experiencing rapid and vast climate changes. The near-surface temperature change over the Arctic is enhanced compared to the global average – a phenomenon called Arctic amplification. This concept was hypothesized already in 1896 by the Swedish scientist Svante Arrhenius, arguing that the effect of changes in the atmospheric carbon dioxide concentration on Earth's surface temperature will be larger by the poles (Arrhenius, 1896). A key driver of Arctic amplification is the albedo feedback. When snow and ice (high albedo) melts, darker surfaces (low albedo) are exposed, and more solar energy is absorbed. Diminishing sea ice lowers the albedo as well as enhances the heat flux to the atmosphere (Serreze et al., 2009; Screen & Simmonds, 2010).

The amplified Arctic warming has huge impacts on the hydrological cycle. Increased evaporation from ice-free seas results in increased precipitation amounts (Bintanja & Selten, 2014; Kopec et al., 2016). A wetter Arctic also results from increased poleward moisture transport (Dufour et al., 2016; Singh et al., 2017; Bintanja et al., 2020), and increased atmospheric moisture content (Screen & Simmonds, 2010; Swann et al., 2010). These processes have different seasonal expression, with meridional moisture transport being more important in summer, atmospheric water vapor content increasing most in summer and early fall, and enhanced evaporation being most prominent in fall and winter. Furthermore, the interannual variability in precipitation is projected to increase (Bintanja et al., 2020), and a greater fraction of the precipitation is expected to fall as rain (Bintanja & Andry, 2017; Førland et al., 2020; McCrystall et al., 2021). Future precipitation increases are projected to be heterogenous, with the greatest annual precipitation increases over the Barents Sea, western Scandinavia, eastern Eurasia, and western North America (Linderholm et al., 2018).

Modeled projections of changes in Arctic precipitation seasonality and humidity are associated with large uncertainties, due to inadequate knowledge of natural climate variations, future anthropogenic greenhouse gas emissions, feedback mechanisms, and model uncertainties (Rawlins et al., 2010; Hanssen-Bauer et al., 2019). To improve our understanding of spatial and temporal variations in precipitation amounts and seasonality over longer timescales, we need paleoclimatic proxy records. Many proxies provide information about temperature, ocean

circulation, and sea ice (e.g., Risebrobakken et al., 2003; de Vernal et al., 2013; Sundqvist et al., 2014; Kaufman et al., 2020). However, our understanding of hydrological changes is still limited, partly because individual proxies often are sensitive to multiple aspects of the water cycle (Konecky et al., 2020). Annual precipitation amounts have been reconstructed using Greenland ice cores (e.g., Dahl-Jensen et al., 1993) and pollen (e.g., Seppä & Birks, 2001; Bjune et al., 2004), but few proxies reflect precipitation seasonality.

In this study, we aim to improve our knowledge about precipitation seasonality on Svalbard and in northern Fennoscandia, two regions projected to experience great and spatially variable precipitation increases (Eklund et al., 2015; Ruosteenoja et al., 2016; Hanssen-Bauer et al., 2017; Hanssen-Bauer et al., 2019). We evaluate precipitation changes across two different timescales: Holocene variability on Svalbard and modern variability in northern Fennoscandia.

One of the reasons we focus on precipitation seasonality is its importance for glacier mass balance. A key question in Holocene climate reconstructions on Svalbard that remains to be answered is why some glaciers re-advanced during the Early Holocene, which was a period characterized by warming and unfavorable conditions for glacier growth (Farnsworth et al., 2020). It has been suggested that warmer oceans and decreasing sea ice – similar mechanisms as are projected to affect precipitation in the future – caused increased winter precipitation amounts in the Late Holocene (D'Andrea et al., 2012; Müller et al., 2012; Røthe et al., 2015). If similar processes acted in the Early Holocene, winter precipitation might have counterbalanced summer temperature-induced glacier ablation.

Stable isotope ratios of oxygen and hydrogen ($\delta^{18}\text{O}$ and $\delta^2\text{H}$) in precipitation are powerful tools for understanding modern-day and past hydrological processes due to their sensitivity to moisture source conditions, atmospheric temperature, and circulation (Dansgaard, 1964; Rozanski et al., 1993; Gat, 1996). In the following sections, I provide an overview of stable water isotopes and the processes influencing the precipitation isotopic composition (section 1.2), and an introduction to the use of leaf wax hydrogen isotopes as a paleo-precipitation proxy (section 1.3).

1.2. Stable water isotopes: basic concepts

Stable isotopes are atoms of the same element with the same number of protons and electrons, but different numbers of neutrons. Stable hydrogen isotopes have 0 or 1 neutrons (^1H and ^2H , the latter also called deuterium), and stable oxygen isotopes have 8, 9, or 10 neutrons (^{16}O , ^{17}O , and ^{18}O). The various isotopes have different mass: the more neutrons, the heavier isotopes, and therefore different physical and chemical properties. Heavy isotopes have higher binding energies and are less volatile than light isotopes, and thus react more slowly. During phase changes (e.g., evaporation and condensation), stable isotopes become enriched in one phase and depleted in the other. This separation is called isotopic fractionation.

Due to the considerably lower abundance of the heavy isotopes, stable isotopes are expressed as the ratio (R) between the rare (heavy) and common (light) isotopes ($^2\text{H}/^1\text{H}$ or $^{18}\text{O}/^{16}\text{O}$), compared to the same ratio in an international standard. Isotope ratios are reported using delta notation (δ) in per mil (‰) relative to Vienna Standard Mean Ocean Water (VSMOW):

$$\delta = 1000 \times \left(\frac{R_{\text{sample}}}{R_{\text{VSMOW}}} - 1 \right) \text{‰} \quad (\text{Equation 1})$$

Stable isotopes are separated from each other through two types of fractionation processes: equilibrium fractionation and kinetic fractionation. Equilibrium fractionation is highly temperature-dependent and occurs during changes between phases in equilibrium. For example, when standard mean ocean water (by definition $\delta^2\text{H} = 0\text{‰}$ and $\delta^{18}\text{O} = 0\text{‰}$) evaporates, isotopically lighter isotopes are preferentially evaporated, and form depleted atmospheric vapor. The ocean surface water, on the other hand, gets isotopically enriched. When the atmospheric water vapor condenses, the resulting precipitation will be enriched relative to the vapor, leaving the remaining vapor depleted (Fig. 1). This process, which results in partitioning of isotopes between two reservoirs as one of the reservoirs decreases in size, is called Rayleigh distillation (Fig. 1). In short, the isotope ratios in the cloud vapor and the condensate are a function of the fraction of remaining water vapor in the cloud (which is a function of temperature), meaning that the precipitation gets more and more depleted as the fraction of water vapor in the cloud gets smaller and smaller.

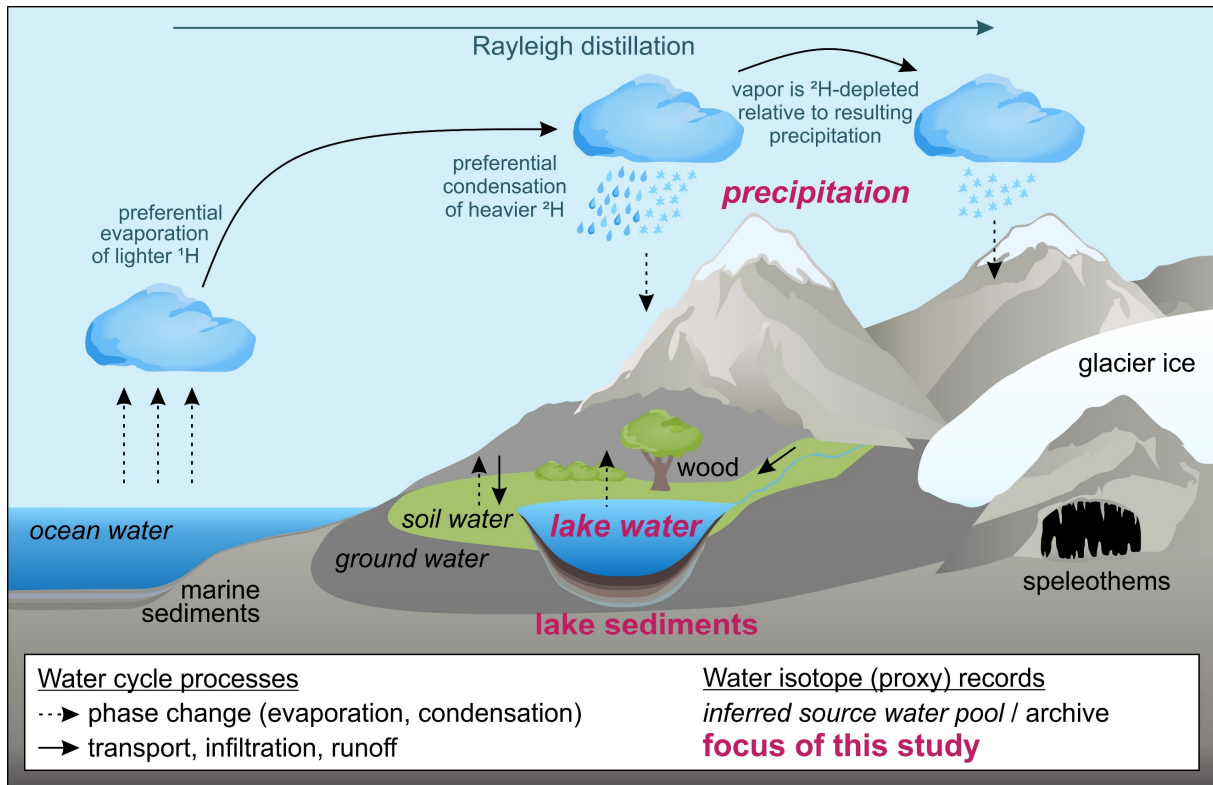


Fig. 1. Conceptual model of the global water cycle, with environmental water pools and archives that can be used to study water isotope proxies. The water pools/proxy records focused on in this study are displayed in red. Arrows show water cycle processes (phase changes and transport). Evaporation of sea water leads to cloud formation and progressive rainout and isotopic fractionation as temperature decreases (Rayleigh distillation). Modified after Konecky et al. (2020).

The spatial and temporal patterns of precipitation $\delta^2\text{H}$ and $\delta^{18}\text{O}$ (hereafter also referred to as precipitation isotopes) are controlled by several mechanisms, including the temperature dependence of equilibrium fractionation, progressive rainout during transport, and fraction of solid precipitation. Due to the mass-dependent fractionation of hydrogen and oxygen isotopes, there is a linear correlation between $\delta^2\text{H}$ and $\delta^{18}\text{O}$. The global average relationship between $\delta^2\text{H}$ and $\delta^{18}\text{O}$ in meteoric waters is described by the Global Meteoric Water Line (GMWL; Craig, 1961):

$$\delta^2H = 8 \times \delta^{18}O + 10 \quad (\text{Equation 2})$$

When a phase reaction cannot reach equilibrium, kinetic fractionation processes can occur. These processes are unidirectional, caused by slower diffusion speed of the heavy isotopes. When water molecules diffuse during evaporation, there is an “excess” of deuterium relative to ^{18}O in vapor (Dansgaard, 1964), causing a deviation from the expected 1:8 ratio. This non-

equilibrium fractionation can be quantified by the so-called deuterium excess (also referred to as d-excess or dxs), which can be calculated as:

$$d - excess = \delta^2H - 8 \times \delta^{18}O \quad (\text{Equation 3})$$

The extent of non-equilibrium fractionation during evaporation (and hence the d-excess value) depends on the air and water temperature, relative humidity above the water surface, and evaporative surface cooling (Merlivat & Jouzel, 1979; Cappa et al., 2003). Furthermore, kinetic fractionation effects can occur during snow formation in mixed-phase clouds (Jouzel & Merlivat, 1984).

Since precipitation isotopes are affected by transport and fractionation processes in the global hydrological cycle, they are widely employed hydrological tracers. As direct measurements of precipitation δ^2H and $\delta^{18}O$ are not available over longer timescales, information about past precipitation isotope variability must be obtained indirectly using paleoclimate proxies. Stable isotopes of hydrogen and oxygen get preserved in various geological records and are therefore commonly applied to understand the paleoenvironment. Some of the most common applications are temperature reconstructions from ice cores (e.g., Dansgaard et al., 1969; Johnsen et al., 2001; Divine et al., 2011), and from foraminifera tests preserved in marine sediments (e.g., Emiliani, 1955; Rasmussen & Thomsen, 2004; Lisiecki & Raymo, 2005). In lake sediments, $\delta^{18}O$ from authigenic carbonates, chironomids (non-biting midges), and diatoms reflects the isotopic composition of the lake water when they formed. As lakes are mainly recharged by meteoric water, these proxies can indirectly archive the isotopic composition of past precipitation, and be used to reconstruct atmospheric circulation, humidity, precipitation seasonality etc. (e.g., Anderson et al., 2001; Hammarlund et al., 2002; Rosqvist et al., 2013; Arppe et al., 2017).

An increasingly popular proxy for high-latitude paleo-precipitation isotopes is δ^2H of sedimentary leaf waxes (e.g., Thomas et al., 2012; Balascio et al., 2013; Thomas et al., 2016; Keisling et al., 2017; Balascio et al., 2018b; Thomas et al., 2018; Thienemann et al., 2019; Thomas et al., 2020; Cowling et al., 2021; Gorbey et al., 2021; Katrantsiotis et al., 2021). This proxy is introduced in the following section and the focus of **Papers I & III**. The key to accurate climate interpretations using water isotope proxies from lacustrine sediments is to understand the processes affecting the lake water isotopic composition (Leng & Marshall, 2004; Jonsson

et al., 2009; Cluett & Thomas, 2020). Therefore, **Paper II** explores the modern water isotopic composition of Arctic and sub-Arctic lakes to evaluate the impact of variable precipitation isotopic composition and seasonality, as well as evaporative enrichment, on the lake water $\delta^2\text{H}$ and $\delta^{18}\text{O}$.

1.3. The leaf wax $\delta^2\text{H}$ paleo-precipitation proxy

Biomarkers are organic molecules that record past environmental and climate conditions. Compound-specific lipid biomarkers are widely used in paleoenvironmental studies due to their persistence to degradation over long geological timescales and their abundance in sedimentary settings (Eglinton & Eglinton, 2008). A variety of lipid biomarkers can be extracted from lake sediments and used to reconstruct different paleoenvironmental conditions within and around the lake basin. Plants produce superficial waxes to keep their moisture balance and protect the leaves (Eglinton & Hamilton, 1967). Here we focus on the leaf wax hydrogen isotopic composition ($\delta^2\text{H}_{\text{wax}}$) of *n*-alkanoic acids, which are straight-chain fatty acids produced by bacteria, algae, and higher plants. $\delta^2\text{H}_{\text{wax}}$ is a useful tool for paleoclimatologists, as it is highly correlated to the $\delta^2\text{H}$ of the source water used by the organism (Sachse et al., 2004; Sachse et al., 2012; McFarlin et al., 2019). There is an offset between $\delta^2\text{H}_{\text{wax}}$ and the $\delta^2\text{H}$ of soil water or lake water (apparent fractionation, ϵ_{app}), e.g., due to biosynthetic fractionation (ϵ_{bio}) occurring during leaf wax generation (Fig. 2; Sessions et al., 1999; Sachse et al., 2012). Different plant types can have different ϵ_{bio} (Daniels et al., 2017; Berke et al., 2019), but it is relatively constant for specific compounds (Sachse et al., 2012; McFarlin et al., 2019).

The environmental water pool used as the hydrogen source differs between organisms, with aquatic plants using lake water and terrestrial plants using soil water during the growing season (Ficken et al., 2000; Aichner et al., 2010b; Shanahan et al., 2013; Wilkie et al., 2013; Thomas et al., 2016; Daniels et al., 2017). Since the hydrogen isotopic composition of aquatic leaf waxes ($\delta^2\text{H}_{\text{aq}}$) reflects the $\delta^2\text{H}$ of lake water ($\delta^2\text{H}_{\text{lake}}$), it will indirectly reflect precipitation $\delta^2\text{H}$ ($\delta^2\text{H}_{\text{precip}}$) in precipitation-fed systems. Additionally, in arid regions where evaporation exceeds precipitation, the lake water $\delta^2\text{H}$ may be overprinted by evaporative $\delta^2\text{H}$ enrichment (Fig. 2; Gibson et al., 2002; Tondu et al., 2013; Cluett & Thomas, 2020). Precipitation is also the ultimate source of soil water used by terrestrial plants. Soil water can be evaporatively enriched,

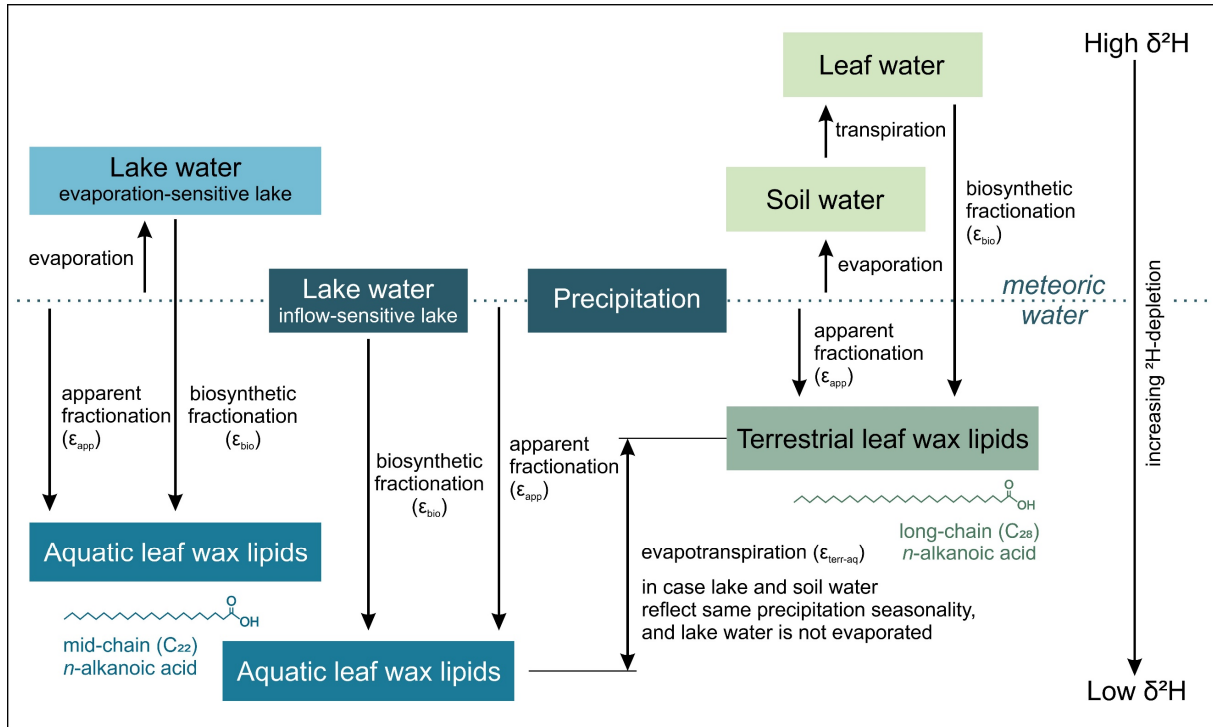


Fig. 2. Isotopic relationships and processes between source water and aquatic and terrestrial leaf waxes (not to scale). Adapted and modified from Sachse et al. (2006) and Aichner (2009).

which means that $\delta^2\text{H}$ of terrestrial plants ($\delta^2\text{H}_{\text{terr}}$) reflects growing season soil water $\delta^2\text{H}$, which is possibly influenced by evapotranspiration (Fig. 2; Kahmen et al., 2013).

The growing season lake and soil water $\delta^2\text{H}$ can reflect different precipitation seasonality. $\delta^2\text{H}_{\text{lake}}$ can reflect summer $\delta^2\text{H}_{\text{precip}}$ if the lake has a short residence time and the water is exchanged throughout summer, mean annual $\delta^2\text{H}_{\text{precip}}$ if the lake has a long residence time and the lake water consists of a mix of summer and winter precipitation, or winter $\delta^2\text{H}_{\text{precip}}$ if the lake receives a higher proportion of the annual precipitation as snow and/or late snowmelt (e.g., Katrantsiotis et al., 2021). Growing season soil water is often assumed to mainly be recharged by summer precipitation (Throckmorton et al., 2016).

In addition to $\delta^2\text{H}_{\text{wax}}$, the following two parameters are useful for leaf wax interpretations. To quantify the isotopic difference between long-chain (C_{28}) and mid-chain (C_{22}) waxes, assuming that the compounds have similar ϵ_{app} (Thomas et al., 2020), we can calculate ϵ_{28-22} , using the equation:

$$\epsilon_{28-22} = 1000 \times \left(\frac{\delta^2\text{H}_{\text{C}_{28}} + 1000}{\delta^2\text{H}_{\text{C}_{22}} + 1000} - 1 \right) \quad (\text{Equation 4})$$

This index can be interpreted as the isotopic difference between the source water pools, i.e., soil and lake water (Fig. 2). Changes in ε_{28-22} can reflect changes in relative humidity (Rach et al., 2017) or precipitation seasonality (Thomas et al., 2020), depending on the hydrology of the lake. The average chain length distribution (ACL) for even-chain C₂₀ to C₃₀ *n*-alkanoic acids is calculated as:

$$ACL_{22-30} = \frac{\sum(n \times C_n)}{\sum(C_n)} \quad (\text{Equation 5})$$

where C_n is the concentration ($\mu\text{g/g}$ sediment) of each *n*-alkanoic acid with *n* carbon atoms. The ACL can give indications of likely source attributions since different plant types preferentially produce different leaf wax homologues.

1.4. Regional setting

The work in this thesis is focused to two research areas: the high-Arctic Svalbard archipelago (**Papers I & III**) and northern Fennoscandia (i.e., northern Norway, Finland, and Sweden; **Paper II**; Fig. 3). Both areas are located in the dynamic Nordic Seas region, where water and air masses originating from low and high latitudes meet, and atmospheric-ocean energy transfer results in varying seasonal climate (Drange et al., 2005). Temperatures in the Nordic Seas (the Norwegian Sea, Greenland Sea, and Iceland Sea) and surrounding areas are higher than at other locations at similar latitudes, mainly due to two mechanisms: 1) meridional heat transport by the North Atlantic Current, and 2) prevailing westerly and southwesterly maritime winds (Årthun et al., 2017). The North Atlantic Current brings warm waters northward along the Norwegian coast, before splitting into the Barents Sea Current and the West Spitsbergen Current, continuing this transport up along the west coast of Spitsbergen (Fig. 3; Skagseth et al., 2008).

In winter, temperature and precipitation variability in the region are strongly associated with the North Atlantic Oscillation (NAO) and intense cyclone activity. The NAO index is defined as the normalized air-pressure difference between the Icelandic Low and the Azores High in winter (Rogers, 1984). The NAO is closely related to the Arctic Oscillation (AO), which is associated with the polar vortex, and causes similar local temperature and precipitation signals (Thompson & Wallace, 1998; Dickson et al., 2000). During positive NAO (or AO) conditions, the pressure gradient between Iceland and the Azores (or between the Arctic and the northern Atlantic and Pacific Oceans) is greater than normal, causing stronger-than-normal westerlies,

(Hurrell, 1995; Uvo, 2003; Lawrence et al., 2020). The NAO also influences the glacier mass balance in northern Fennoscandia and to a lesser extent Svalbard, by regulating the amount of winter accumulation (Pohjola & Rogers, 1997; Bonan et al., 2019). Strong winter cyclones are common over the Norwegian Sea and between Svalbard and Fennoscandia and associated with strong winds and high precipitation amounts (Serreze et al., 1993; Rogers et al., 2005; Wickström et al., 2020).

Summer precipitation is not as strongly related to atmospheric circulation. Cyclones are common also in summer, but they are weaker and more spatially variable (Serreze et al., 1993; Zhang et al., 2004; Wickström et al., 2020). Summer rainfall over Scandinavia is often caused by local convection due to surface heating (Jaagus, 2009; Dyrddal et al., 2015). On Svalbard, low summer temperatures make local convective precipitation less likely to occur. Precipitation amounts are low due to stable, stratified air masses containing low amounts of water vapor (Førland et al., 2009).

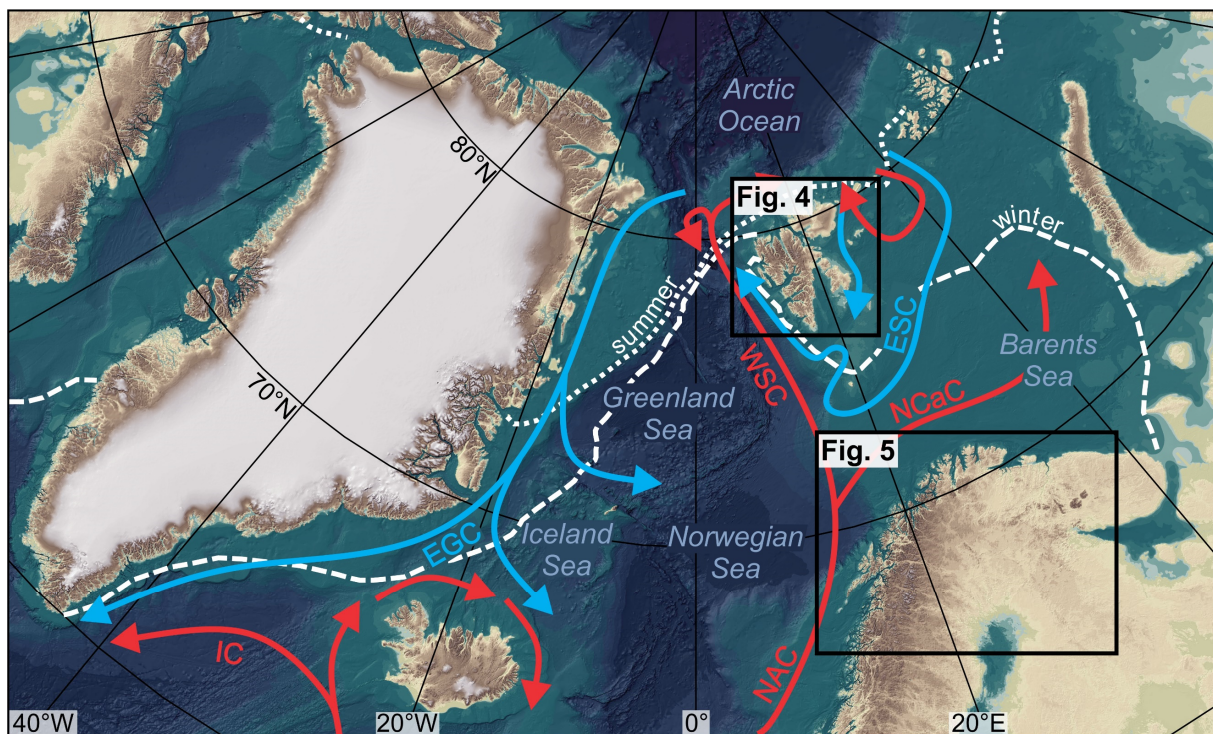


Fig. 3. Overview map of the North Atlantic region. Major ocean surface currents are indicated in red (warm: NAC = North Atlantic Current; NCaC = North Cape Current; WSC = West Spitsbergen Current; IC = Irminger Current) and blue (cold: ESC = East Spitsbergen Current; EGC = East Greenland Current), and median winter and summer sea-ice extent 1981–2010 are shown with white dashed and dotted lines (National Snow and Ice Data Center, 2019). Black boxes mark the two study areas. Background map from IBCAO (Jakobsson et al., 2012).

1.4.1. Svalbard

Svalbard (74–81°N and 10–35°E) is located at the end of the North Atlantic cyclone track and experiences warmer and wetter conditions than other areas at similar latitude (Eckerstorfer & Christiansen, 2011; Serreze & Barry, 2014). The archipelago spans large climate gradients, due to the influence of Arctic and Atlantic air masses and variable ocean heat flux, further influencing the distribution of sea ice (Fig. 4; Hisdal, 1998; Walczowski & Piechura, 2011). The Fram Strait is the world’s northernmost ocean area with essentially ice-free conditions all year round (Fig. 4; Haugan, 1999). The West Spitsbergen Current transports warm, saline water in the east of the strait, along the west coast of Spitsbergen (Skagseth et al., 2008). Svalbard is dominated by southwesterly winds bringing mild Atlantic air, and northeasterly winds bringing dry polar air (Hagen et al., 1993). Easterly moisture from the Barents Sea cause high precipitation amounts over the east coast glaciers (Hagen et al., 1993), whereas extreme precipitation events are favored by south-southwesterly flow (Dobler et al., 2019). Precipitation amounts are highest over the mountains on the west coast and the southeastern islands, whereas the central and northeastern parts are relatively arid (Østby et al., 2017). In addition to these large-scale patterns, Svalbard experiences large local variability (Humlum, 2002; Førland & Hanssen-Bauer, 2003).

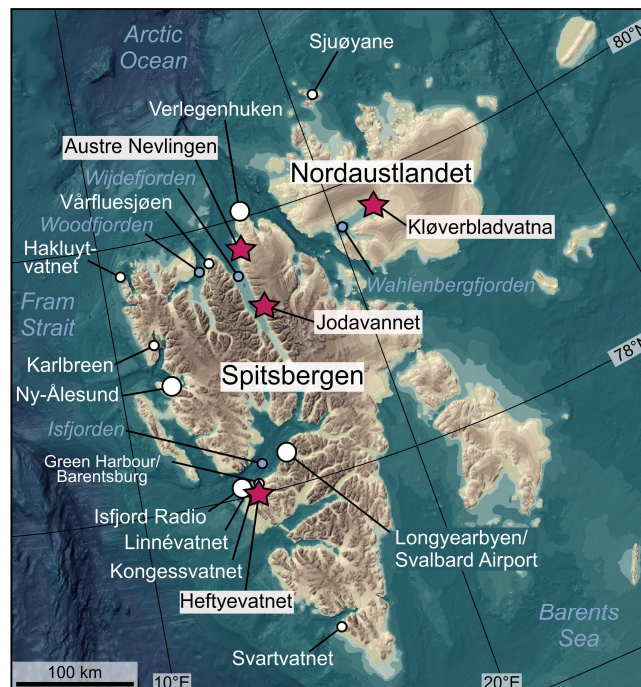


Fig. 4. Overview map of Svalbard. Studied lakes are marked with red stars, meteorological stations with large white circles, and other place names mentioned in the text with small white (terrestrial) and blue (marine) circles. Background map from IBCAO (Jakobsson et al., 2012).

A homogenized, composite monthly mean air temperature time series for Svalbard Airport shows a warming rate of 0.32 °C per decade from 1898 to 2018 (Nordli et al., 2020). After 1991, the rate is 1.7 °C per decade, which is twice the Arctic average and approximately seven times the global average (0.8 and 0.2 °C per decade, respectively; Nordli et al., 2020). From 1971–2000 to 2071–2100, Svalbard is projected to see a temperature increase of almost 10 °C (for emission scenario RCP8.5 = “business as usual”; Hanssen-Bauer et al., 2019). The amplified warming also has an immense impact on the hydrological cycle, through shorter ice-cover seasons in the fjords on the west coast, reduced glacier net mass balance, permafrost warming, and increased moisture transport (Hanssen-Bauer et al., 2019).

In an Arctic context, Svalbard has a unique precipitation record, comprising a number of 50–100-year-long time series of precipitation measurements (Førland et al., 1997; Hanssen-Bauer, 2002; Førland et al., 2011; Hanssen-Bauer et al., 2019). At the four long-term stations (Svalbard Airport, Ny-Ålesund, Bjørnøya, and Hopen), the precipitation amounts have increased by 20–35% over the last 50 years (Førland et al., 2020). There are no long-term precipitation measurements from northeastern Svalbard, but snow accumulation measurements suggest that southern Spitsbergen receives twice as much snow (in water equivalent) compared to central and northern Spitsbergen, and that the east coast receives 40% more snow than the west coast (Sand et al., 2003). From 1971–2000 to 2071–2100, precipitation amounts are projected to increase by 65% (for RCP8.5; Hanssen-Bauer et al., 2019). The largest relative increases are predicted to occur in winter and in the northeast. The heterogenous pattern of snow accumulation has impact on the equilibrium line altitude (ELA) of glaciers, being lower in coastal, and higher in inland settings (Hagen et al., 1993; van Pelt et al., 2016). Furthermore, heavy precipitation events have become more frequent and intense (Hanssen-Bauer et al., 2019), and are projected to continue to do so towards the end of the 21st century (Isaksen et al., 2017).

Our understanding of precipitation isotope variability on Svalbard is limited. Monthly precipitation isotopes have been measured at Isfjord Radio (78.07°N, 13.63°E; 6 m a.s.l.) between 1961–1965 and 1972–1975, and Ny-Ålesund (78.92°N, 11.93°E; 7 m a.s.l.) in 1990–2016, as part of the Global Network of Isotopes in Precipitation (GNIP; IAEA/WMO, 2019). Additionally, precipitation isotope values were monitored at the Hornsund Polish Polar Station in 2013–2014 (Arppe et al., 2017) and in Barentsburg in 1975–1981 and 2016–2017 (Skakun et al., 2020). All these sites are located on the west coast of Spitsbergen, and none of the data

series show a significant seasonal trend in $\delta^2\text{H}$ or $\delta^{18}\text{O}$. The isotopic seasonality is expected to be higher in central and northeastern Svalbard, because of larger contrast in climate between winter and summer.

To reconstruct Holocene climate on Svalbard beyond the instrumental record, we can use paleoclimate records. Many studies have focused on temperature and sea-ice history, but little is known about precipitation. The Early Holocene was characterized by peak summer insolation (Laskar et al., 2004), and high ocean surface and subsurface temperatures due to increased Atlantic heat flux (e.g., Hald et al., 2007; Werner et al., 2016; Mangerud & Svendsen, 2018). This led to reduced sea-ice cover in the eastern Fram Strait (Werner et al., 2016) and in many Svalbard fjords (e.g., Forwick & Vorren, 2009; Bartels et al., 2018; Allaart et al., 2020). Warmer-than-present temperatures have also been suggested by alkenones (van der Bilt et al., 2019) and sedimentary ancient DNA (*sedaDNA*) of vascular plants (Voldstad et al., 2020) from lacustrine sediments. Furthermore, the glacier cover on Svalbard was substantially reduced (Fjeldskaar et al., 2018; Farnsworth et al., 2020). However, glacier re-advances still occurred, and it has been speculated that increased winter precipitation might have been one of the drivers (Farnsworth et al., 2020). Glaciers were absent or much smaller during the Middle Holocene (e.g., Svendsen & Mangerud, 1997; Allaart et al., 2021b), and warm and dry conditions are suggested from lake sediment records (Balascio et al., 2018b; Voldstad et al., 2020). Sea surface temperatures remained high (e.g., Werner et al., 2016; Mangerud & Svendsen, 2018), and sea-ice concentrations low (Werner et al., 2016) until *c.* 6–5 cal. kyr BP. The Late Holocene was characterized by Neoglacial cooling, with declining summer insolation (Laskar et al., 2004), and lower temperatures inferred from alkenones (van der Bilt et al., 2018), plant macrofossils (Birks, 1991), and *sedaDNA* (Voldstad et al., 2020). The sea-ice cover in the Fram Strait and Svalbard fjords increased with weakened Atlantic water advection (Forwick & Vorren, 2009; Werner et al., 2016; Bartels et al., 2018), and many glaciers re-advanced between 4–0.5 cal. kyr BP (Farnsworth et al., 2020). Some of the Neoglacial and Little Ice Age (LIA) glacier advances have been suggested to be driven by increased winter precipitation (D'Andrea et al., 2012; Røthe et al., 2015), possibly due to occasional northward shifts of the sea-ice edge, and increased evaporation (Müller et al., 2012).

Precipitation reconstruction on Svalbard is hampered by the lack of available climate archives reflecting precipitation changes. The dating of ice cores is challenging due to surface melting in summer altering the original ice core records, and the oldest ice core records to date go back

only *c.* 1,200 years (Isaksson et al., 2003; Isaksson et al., 2005; Divine et al., 2011). Furthermore, local pollen production is low, and the long-distance component is high (van der Knaap, 1987; Birks, 1991), limiting the suitability of pollen studies. To better constrain the seasonal variability in precipitation throughout the Holocene on Svalbard, we analyze sedimentary leaf waxes, presented in **Papers I & III**.

1.4.2. Northern Fennoscandia

The second region of interest and focus of **Paper II** is northern Fennoscandia, where we investigate surface water samples from Norway, Finland, and Sweden. The study area spans from the Norwegian Sea in the northwest to Bothnian Bay in the southeast, between 65.74°N to 71.11°N and 18.39°E to 30.06°E (Fig. 5). The northwestern side of the Scandinavian Mountains is characterized by alpine topography shaped by Quaternary glaciations, with steep mountains, U-shaped valleys, and fjords. The highest mountains are the *c.* 1800-meter-high Lyngen Alps (Fig. 5). Towards southeast the mountains become gradually lower, forming a more gently undulating landscape, and reaching sea level by the Bothnian Bay.

The topography has a great impact on the climate in the area. The western side of the mountains has a maritime climate (cool summers and mild winters), whereas the interior parts are more continental (warm summers and cold winters; Førland et al., 2009). Most of the moisture is brought by dynamic low-pressure systems from the North Atlantic and prevailing westerly winds. When the warm humid air reaches land, the steep topography causes orographic lift, and precipitation falls on the windward side of the mountains (Hanssen-Bauer & Førland, 2000). The leeward side is in rain shadow and receives much lower precipitation amounts (Uvo, 2003). The NAO has a large impact on winter precipitation by regulating the strength of the westerlies (Hurrell, 1995; Uvo, 2003; Irannezhad et al., 2014). During NAO+ conditions, Scandinavia receives more precipitation due to a northward shift of the North Atlantic storm track. Also AO+ conditions can result in higher precipitation amounts (Lawrence et al., 2020). Furthermore, winters with mild Arctic weather, low sea-ice concentrations and strongly negative NAO and AO conditions can result in extreme snowfall over northern Europe (Cohen et al., 2013; Bailey et al., 2021). During cold air outbreaks, dry polar air exposed to warm ocean water surfaces creates high heat and moisture flux (Shapiro et al., 1987; Papritz & Sodemann, 2018).

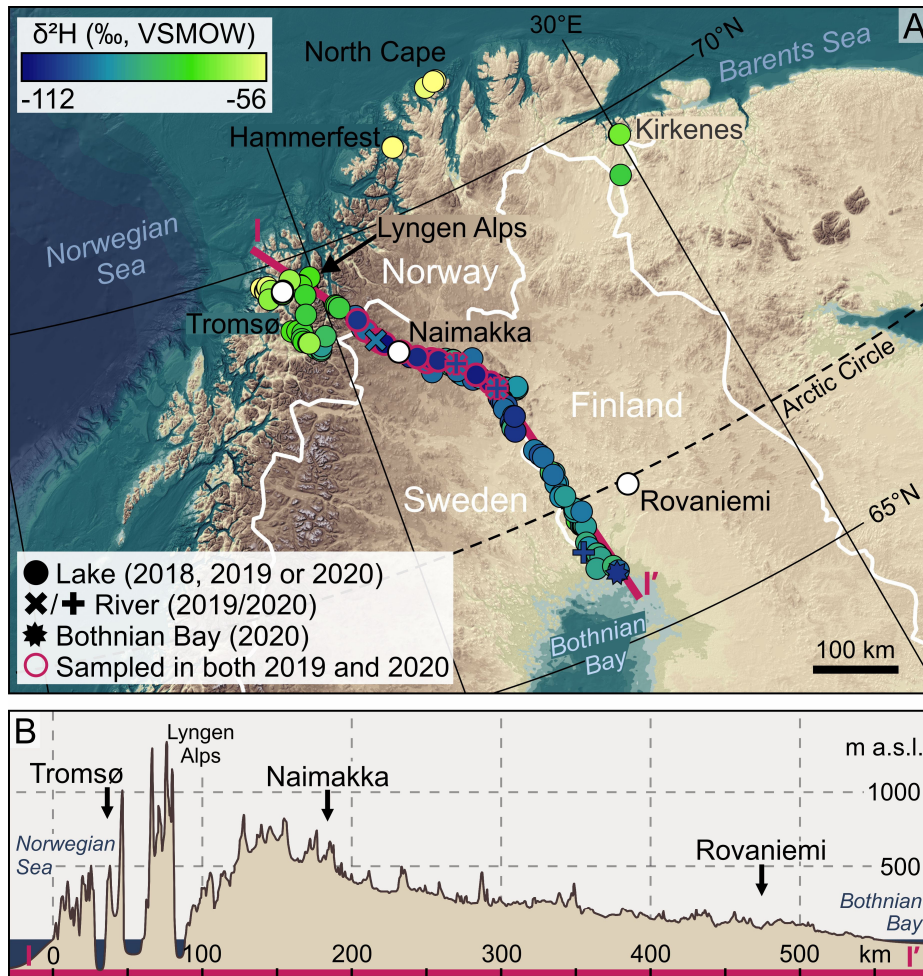


Fig. 5. (a) Map of northern Fennoscandia, showing surface water $\delta^2\text{H}$ from 135 lakes, two rivers and the Bothnian Bay (colored by $\delta^2\text{H}$ value), locations of meteorological stations (white circles), and the extent of the elevation profile in (b) (red line). Background map from IBCAO (Jakobsson et al., 2012). (b) Elevation profile from the Norwegian Sea to the Bothnian Bay.

In northern Norway, the mean annual air temperature (MAAT) has increased by 0.08–0.11 °C per decade from 1900 to 2014 (Hanssen-Bauer et al., 2017). The increase is greatest (0.14–0.15 °C per decade) in spring, and lowest (and not statistically significant) in winter. For emission scenario RCP8.5, MAAT is projected to increase by 5.0–6.1 °C from 1971–2000 to 2071–2100, and the increase is predicted to be largest in winter (Hanssen-Bauer et al., 2017). Finland has experienced a long-term (1847–2013) mean warming of 0.14 °C per decade (Mikkonen et al., 2015), and in northern Finland, the MAAT has increased by 0.3–0.4 °C per decade between 1961–2010 (Aalto et al., 2016). For emission scenario RCP8.5, Ruosteenoja et al. (2016) projected an increase of mean winter air temperature by 6–8 °C and mean summer air temperature by 4–5 °C from 1981–2010 to 2070–2099 in northern Finland and northern

Sweden. This corroborates with data from Sweden (Eklund et al., 2015), predicting similar values for RCP8.5, and the largest increase in winter.

Mean annual precipitation (MAP) amounts have increased by 0.2–1.7% per decade from 1900 to 2014 in northern Norway (Hanssen-Bauer et al., 2017). Also northern Sweden has experienced a wetting trend since 1900, in particular after 1950, and especially in winter (Chen et al., 2021). From 1971–2000 to 2071–2100, MAP amounts over northern Norway are projected to increase by 17–22 % (RCP8.5; Hanssen-Bauer et al., 2017). The increase is projected to occur during all seasons, but there is no consistency between models and scenarios in terms of seasonal trends. For northern Finland and northern Sweden, the precipitation amounts are projected to increase by 20–30% in winter and *c.* 10% in summer from 1981–2010 to 2070–2099 (RCP8.5; Ruosteenoja et al., 2016).

Monthly precipitation $\delta^2\text{H}$ and $\delta^{18}\text{O}$ have been measured at two GNIP stations in our study area: Naimakka (68.68°N, 21.53°E; 403 m a.s.l.) in 1990–1995 and Rovaniemi (66.50°N, 25.75°E; 107 m a.s.l.) in 2003–2014 (Fig. 5; IAEA/WMO, 2019). To monitor the precipitation isotope seasonality at a more coastal site, we installed a precipitation gauge in Tromsø (69.68°N, 18.96°E; 73 m a.s.l.; Fig. 5) in 2019 (see section 2.2.1). Naimakka is the most continental site, located on the eastern side of the Scandinavian Mountains, and displays the most depleted values and the strongest seasonality. Tromsø precipitation $\delta^2\text{H}$ and $\delta^{18}\text{O}$ are relatively enriched and do not show a strong seasonality. Rovaniemi falls in between, having a stronger seasonal variability than Tromsø, but less depleted values than Naimakka.

1.5. Aims and research objectives

Arctic precipitation changes in response to warming during earlier periods in Earth's history can serve as an analogue for the future. However, the Holocene history of Arctic precipitation changes is largely unknown, due to the absence of long instrumental records and lack of robust proxies. In the light of future projections (i.e., variable precipitation increases between different seasons and different locations), we need to increase the spatial distribution of long-term paleo-precipitation records, which requires thoughtful consideration of how lake water proxies record variable seasonal distribution of precipitation. The overall aim of this PhD project is to increase our understanding of the Arctic water cycle, and how changes in time and space are reflected in stable water isotope proxies.

More specifically, the research objectives are to:

- Generate Holocene precipitation records across the understudied Svalbard based on terrestrial and aquatic leaf wax $\delta^2\text{H}$, with a focus on precipitation seasonality (**Papers I & III**).
- Identify Holocene variations in winter precipitation on Svalbard, one of the main controls on glacier mass balance (**Papers I & III**).
- Explore modern controls on lake water $\delta^2\text{H}$ by evaluating variable impact of precipitation $\delta^2\text{H}$ and evaporation on lake water $\delta^2\text{H}$ across northern Fennoscandia (**Paper II**).

2. Scientific approach

In this thesis, I use lipid biomarkers from lacustrine sediment cores from Svalbard to investigate Holocene variability in Arctic precipitation seasonality (**Papers I & III**), and lake surface water samples from northern Fennoscandia to investigate modern controls on lake water $\delta^2\text{H}$ (**Paper II**). The motivation behind the use of two different study areas was mainly practicality. From Svalbard, we had access to a large collection of lake sediment cores that we could use for the Holocene precipitation reconstruction, but few modern data. Getting good spatial coverage of lakes from Svalbard for a lake water study as presented in **Paper II** would require massive logistical support because of the remote location. By targeting northern Fennoscandia, an area with a well-developed road system and easy access to lakes, we could cover a large area and sample many lakes within a few days. It was also feasible to repeat the sampling over several years. Additionally, we could compare the isotopic composition of the lake water to precipitation isotope data from stations along our targeted transect.

Although our southernmost lake on Svalbard is located *c.* 800 km north of the Norwegian mainland and influenced by a range of different processes (e.g., variable sea-ice extent, Arctic air and water masses), there are parallels between the two study regions. Both regions experience climate gradients related to the dominance of North Atlantic moisture and alpine topography, with humid maritime climate on the west coast and drier more continental climate towards the (north/south)east. Furthermore, the west coast (i.e., Isfjord Radio and Ny-Ålesund on Svalbard; Tromsø on the mainland) experiences similar amplitude of seasonal temperature variability (although Svalbard is generally 5–10 °C colder) and receives most precipitation in

fall and winter. In the following sections, I present the scientific approach used in this study, starting with the Holocene sedimentary leaf waxes from lakes on Svalbard, followed by the modern lake water isotopes from northern Fennoscandia.

2.1. Svalbard lakes

For the Holocene precipitation reconstructions, we targeted four lakes on Svalbard to explore spatial and temporal differences in precipitation seasonality (Figs 4 and 6). The lakes were chosen based on their location and estimated residence time (i.e., the average time a given molecule of water remains in the lake). By utilizing sediment cores analyzed in other studies (Schomacker et al., 2019; Voldstad et al., 2020; Farnsworth et al., 2022), we could target sedimentary records that have already been scanned and logged, and which have a good chronological control. A pilot study of one core from Austre Nevlingen was performed at UB in 2018, which together with analyses of a second core generated data for **Paper I**. Heftyevatnet, Jodavannet and Kløverbladvatna are introduced in **Paper III**. Details on the locations of the lakes, and lake and catchment morphometry are given in Table 1.

Table 1. Lake data for Austre Nevlingen, Heftyevatnet, Jodavannet and Kløverbladvatna, Svalbard.

Lake	Latitude (°N)	Longitude (°E)	Lake elevation (m a.s.l.)	Maximum water depth (m)	Lake surface area (km ²)	Catchment area (km ²)	Spring Melt Residence Time (months)	Ice-free Season Residence Time (months)
Austre Nevlingen	79.783	15.786	41	18	0.12	0.48	28.3	156.7
Heftyevatnet	77.998	14.149	43	6.5	0.015	0.15	2.1	15.4
Jodavannet	79.338	16.017	140	6.4	0.020	1.31	0.8	4.2
Kløverbladvatna	79.765	21.715	8	17.5	0.23	1.64	19.0	105.4

2.1.1. Sediment cores

The lacustrine sediment cores presented in **Papers I & III** were collected between 2015 and 2019. The coring, core splitting, logging, and a range of standard bulk sediment analyses are described for Kløverbladvatna, Jodavannet, and Heftyevatnet by Schomacker et al. (2019), Voldstad et al. (2020), and Farnsworth et al. (2022). The Austre Nevlingen cores are presented for the first time in **Paper I**.

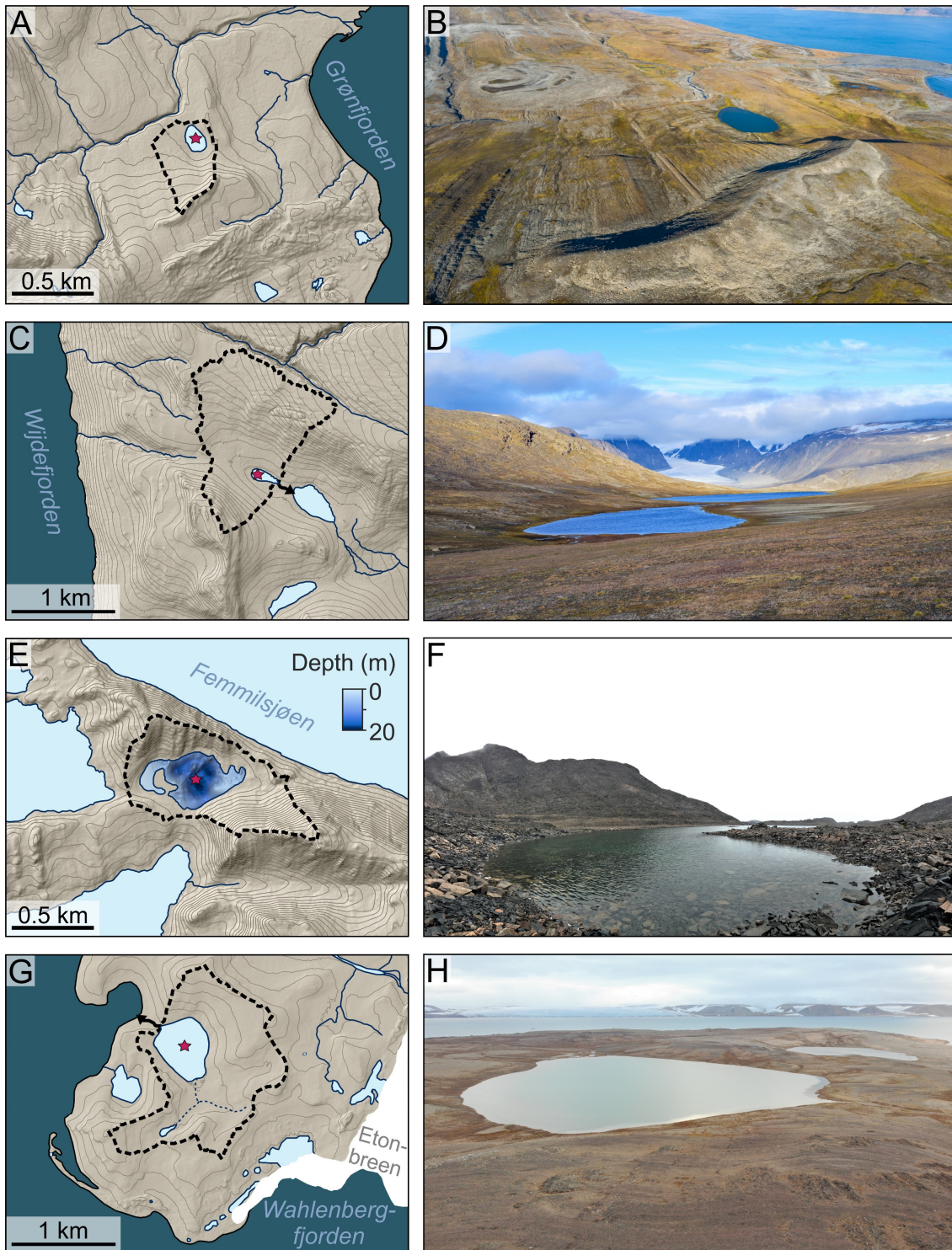


Fig. 6. Catchment overviews of Heftyevatnet, western Spitsbergen (a-b), Jodavannet, northern Spitsbergen (c-d), Austre Nevlingen, northern Spitsbergen (e-f), and Kløverbladvatna, Nordaustlandet (g-h). Background maps in (a), (c), (e), and (g) based on 5x5 m DEM from Norwegian Polar Institute (2014) with 10-m contour lines. The photographs were taken in August 2017 (Erik S. Mannerfelt; Farnsworth et al., 2022), August 2016 (Anders Schomacker), August 2018 (Sofia E. Kjellman), and August 2021 (Sara M. Cohen), respectively.

From all lakes, overlapping cores were collected using a Universal surface corer (120 cm long and Ø68 mm coring tubes) and a hand-held lightweight piston corer (200 cm long and Ø60 mm coring tubes). Coring was performed through a hole in the floor of a small Zodiac in summer, or from the spring lake ice. To identify the deepest basin in each lake, the bathymetry of all lakes was surveyed using a Hondex PS-7 Transducer LCD Digital Sounder for point measurements, and for Austre Nevlingen with a Garmin ECHOMAP™ Plus 73SV with a CV52HW-TM transducer and a 5 Hz receiver, using the Quick Draw contour function.

All lake sediment cores were split open, logged and analyzed in an ITRAX core scanner (Croudace et al., 2006) to record variations in the element-geochemical properties. The ITRAX provides high-resolution X-ray fluorescence (XRF) data, magnetic susceptibility (MS), and optical and radiographic images. We used the titanium (Ti) signal and the calcium-to-iron (Ca/Fe) ratio to identify variations in the minerogenic input, as these can be used to detect lake basin isolation (Larsen et al., 2017) and glacial meltwater input (Kylander et al., 2011). Ti was normalized against the incoherent (inc) and coherent (coh) Rh scatter ($Ti/(inc + coh)$) to remove instrumental effects (Kylander et al., 2011). The XRF data were also used for core correlation (section 2.1.2). To determine the organic matter content, we performed loss on ignition (LOI; Heiri et al., 2001) on 1–2 cm³ samples, that were dried at 110 °C for 24 h and ignited at 550 °C for 4 h. Core descriptions and lithostratigraphy are presented in full in by Schomacker et al. (2019), Voldstad et al. (2020), Farnsworth et al. (2022), and in **Paper I**. An overview of all sediment cores, including core photographs, X-ray images, lithology, and depths for leaf wax and radiocarbon samples is given in **Paper III**.

2.1.2. Core chronologies and composite core construction

To determine the timing of changes in the sedimentary records and interpret variability over time, it is crucial to have a good age control. Radiocarbon (¹⁴C) dating is a widely used method to determine the age of late Quaternary sediments, but can be problematic on Svalbard due to lack of organic material, and the presence of coal or carbonate rocks. To avoid potential lake reservoir effects and minimize the risk of getting too old ages, we targeted lakes with little or no coal and carbonates in the catchment, and preferentially dated terrestrial plant remains, since terrestrial plants uptake carbon directly from the atmosphere (Hajdas, 2008). In regions with crystalline bedrock, autochthonous aquatic macrofossils can be used for dating since the ¹⁴C in lake water and atmosphere is often in equilibrium (Wolfe et al., 2004). The radiocarbon ages

presented in **Papers I & III** are mainly measured on *Salix polaris* leaves or aquatic bryophytes. Another advancing dating method is tephrochronology, using the chemical fingerprint of discrete layers of volcanic ash (tephra) to identify volcanic eruptions of known age. Cryptotephra (non-visible volcanic ash) can travel far from the volcanic source before being deposited and has been found in Svalbard lake sediments (D'Andrea et al., 2012; van der Bilt et al., 2017; Farnsworth et al., 2022). In **Paper III**, the Vedde Ash, identified by Farnsworth et al. (2022), is used as an age constraint in the Heftyevatnet age-depth model.

All core chronologies in this study are based on accelerator mass spectrometry (AMS) radiocarbon dating on handpicked plant macrofossils, identified and isolated through sieving. The samples were dated at the Ångström Laboratory, Uppsala University, Sweden, and Lund University Radiocarbon Dating Laboratory, Sweden. Age-depth models for the sediment cores are presented in **Paper I** (Austre Nevlingen), and by Schomacker et al. (2019; Kløverbladvatna), Voldstad et al. (2020; Jodavannet), and Farnsworth et al. (2022; Heftyevatnet). Three new age-depth models were generated in **Paper III**, to include seven additional ages for Heftyevatnet, three new ages for Jodavannet, and to establish a composite age-depth model for Kløverbladvatna. Since we used multiple cores from all lakes, a composite sequence was constructed for each of the lakes. The overlapping cores were aligned in AnalySeries (v. 2.0.8; Paillard et al., 1996) and correlated using tie points in the XRF data, looking at several different elemental ratios (e.g., Ca/Fe, Fe/Ti, and Ti/(inc+coh)).

To allow us to present proxy data with age model uncertainty, all proxy and chronological data were entered into Linked PaleoData (LiPD) files (McKay & Emile-Geay, 2016). Age-depth models were generated in R (v. 4.1.3; R Core Team, 2021) using Bacon (v. 2.5.7; Blaauw & Christen, 2011) and the IntCal13 (**Paper I**; Reimer et al., 2013) or IntCal20 dataset (**Paper III**; Reimer et al., 2020) within the geoChronR package (v. 1.1.5; McKay et al., 2021). The lower depth (d.max) for each record was set to the lowermost leaf wax or radiocarbon sample depth, and the upper depth (d.min) to the uppermost leaf wax sample. To account for changes in sedimentation rate (as indicated by a major change in the lithology), boundaries were added and the prior mean accumulation rate (bacon.acc.mean) adjusted. All radiocarbon ages are presented in calibrated kiloyears before present (cal. kyr BP; BP = 1950).

2.1.3. Lipid biomarker extraction and analysis

Biogeochemical and isotopic analyses were performed in the Organic and Stable Isotope Biogeochemistry Laboratory at the University at Buffalo, using previously published methods (Thomas et al., 2012). In short, the *n*-alkanoic acids were 1) extracted, 2) purified, and 3) analyzed (Fig. 7). A simplified flowchart summarizing the methodological approach and analytical procedures used for the lacustrine sediment records and modern plant samples, focusing on the *n*-alkanoic acid extraction and analysis, is shown in Figure 8.

In the first step, the samples were freeze-dried, homogenized, and mixed with diatomaceous earth. The total lipid extract (TLE) was extracted with a Dionex 200 Accelerated Solvent Extractor (ASE) using dichloromethane (DCM):methanol 9:1 (v:v), flushed three times for 10 minutes each. The ASE uses high temperature (120 °C) and pressure (1200 psi) to extract the free lipids from the sediments. An internal standard ($C_{20:1}$ *n*-alkanoic acid) was added to allow biomarker quantification. Subsequently, the acid fraction was separated from the neutral fraction using flash columns with an aminopropyl silica gel solid phase. The aminopropyl group acts as a base and forms a strong bond to the acid functional group of the fatty acids. Therefore, when the neutral fraction was eluted in DCM:isopropanol 2:1, the fatty acids stuck to the solid phase. To collect the acid fraction, we used 4% acetic acid in DCM. This stronger acid bonds stronger to the solid phase and the fatty acids are therefore released. After extraction, the fatty

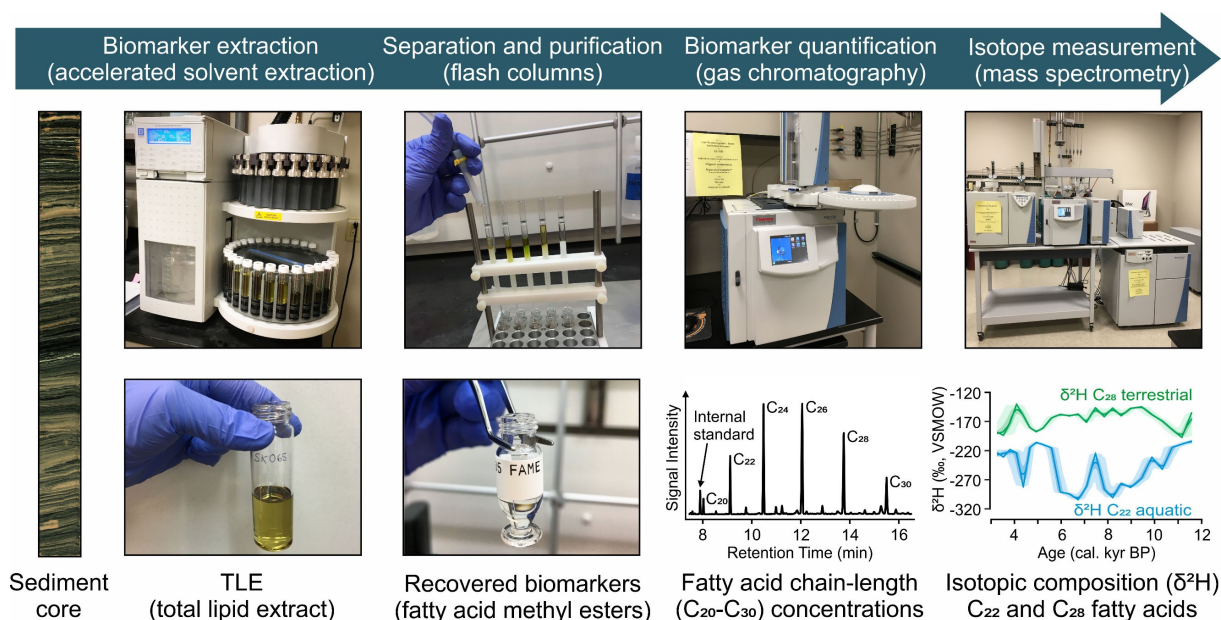


Fig. 7. Simplified methodology for generating leaf wax *n*-alkanoic acid data, from lipid extraction to δ^2H analysis.

acids need to be derivatized to be sufficiently volatile to be analyzed on the gas chromatograph. The acid fraction was methylated in acidified methanol with a known isotopic composition at 60 °C overnight, converting the fatty acids into less polar fatty acid methyl esters (FAMES). After extraction, the FAMES were purified before quantification and $\delta^2\text{H}$ measurement. The FAMES were extracted in hexane three times from a hexane-salt water mixture. A final clean-up was done using flash columns with a silica gel solid phase, eluting FAMES in DCM, after first eluting apolar compounds (e.g., alkanes and hydroxy fatty acids) in hexane.

We quantified the relative abundance of *n*-alkanoic acids on a Thermo Trace 1310 Gas Chromatograph (GC) with two flame ionization detectors operated in parallel and AI1310 autosamplers and two split/splitless injectors. Details on the GC conditions and programs are given in Table 2. Compound peaks were identified using the retention time of the added internal standard as a reference. To convert peak areas into concentrations, we used external calibration curves determined for a C₂₈ FAME standard. The FAME masses were normalized to the dry mass of extracted sample and the recovery of the standard.

FAME $\delta^2\text{H}$ was measured on a Thermo Scientific Delta V Plus isotope ratio mass spectrometer (IRMS) coupled to a Trace 1310 GC via an Isolink II and Conflo IV. The FAME concentrations were used to dilute and inject target amounts of compounds. We used the same GC conditions as during FAME quantification, except for the final flow rate and the carrier gas (Table 2). The H₃⁺ factor was determined at the beginning of every sequence, and samples were run in triplicates (or duplicates or singles for small samples) along with FAME standards of known composition to account for drift, linearity and normalize to the VSMOW scale (Table 2). The FAME values were also corrected for hydrogens added during methylation. We injected C₂₀ and C₂₈ FAME standards at a range of masses to constrain the peak-size effects on the measured isotopic composition. Peaks that were below a threshold (*c.* 2-4 V·s for each sequence) were excluded. For sequences without clear linearity and small peaks (C₂₀ and/or C₃₀), all peaks for that chain length were discarded. For details, see **Paper III**. The total uncertainty of measured $\delta^2\text{H}$ values is given as the Standard Error of the Mean (SEM), i.e., the square root of the sum of the squares of total measurement uncertainty for each sample, divided by the square root of the number of measurements.

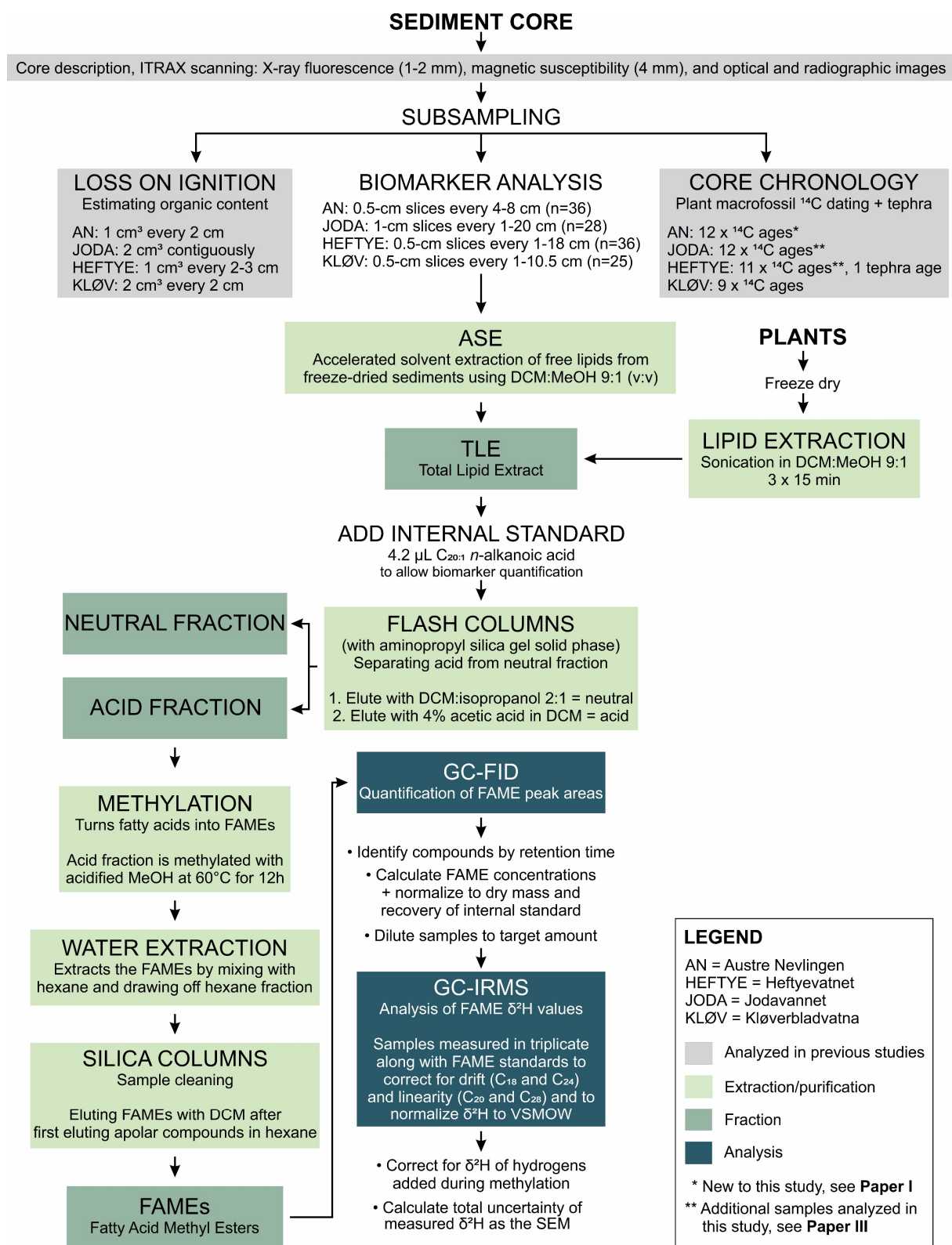


Fig. 8. Flowchart summarizing the methodological approach and geochemical analyses performed in this study, producing data for **Papers I & III**. Abbreviations are listed on pages x–xi.

Table 2. Instrumentation and settings for FAME quantification using gas chromatography with flame ionization detection (GC-FID) and $\delta^2\text{H}$ analysis using gas chromatography-isotope ratio mass spectrometry (GC-IRMS).

	GC-FID	GC-IRMS
Analyzer	Thermo Trace 1310 GC with two FIDs operated in parallel with AI1310 autosamplers and two split/splitless injectors	Thermo Scientific Delta V Plus IRMS coupled via Isolink II and Conflo IV to a Trace 1310 GC
Inlet and flow conditions	Inlets held at 250 °C and operated in splitless mode for 45 s, after which split flow was turned on at 14 mL min ⁻¹ , followed by constant column flow of 3.6 mL min ⁻¹ using hydrogen carrier gas	Inlets held at 250 °C and operated in splitless mode for 45 s, after which split flow was turned on at 14 mL min ⁻¹ , followed by constant column flow of 1.5 mL min ⁻¹ using helium carrier gas
Oven program	Initial temperature of 70 °C for 1 min, then ramped to 230 °C at 27 °C min ⁻¹ , followed by a final ramp to 315 °C at 6 °C min ⁻¹ , and held for 10 min	Initial temperature of 70 °C for 1 min, then ramped to 230 °C at 27 °C min ⁻¹ , followed by a final ramp to 315 °C at 6 °C min ⁻¹ , and held for 10 min
H ₃ ⁺ factor	–	Between 2.19 ± 0.02 and 5.00 ± 0.04 ppm/nA
FAME standards	C ₂₈ (Fisher Scientific)	C ₁₈ and C ₂₄ (drift), C ₂₀ and C ₂₈ (linearity) (A. Schimmelman, University of Indiana)

2.1.4. Chain-length distribution of leaf wax *n*-alkanoic acids

To interpret sedimentary leaf wax $\delta^2\text{H}$, we need knowledge about the plants producing the waxes, and their dominating chain lengths. The classical framework for interpreting leaf wax $\delta^2\text{H}$ is based on studies from low- and mid-latitudes, finding that terrestrial plants produce high concentrations of long-chain waxes and aquatic plants predominantly produce mid-chain waxes (e.g., Ficken et al., 2000; Aichner et al., 2010a; Bush & McInerney, 2013; Diefendorf & Freimuth, 2017). Constraints on modern Arctic vegetation are sparse, but there is an increasing number of studies investigating modern leaf wax *n*-alkanoic acids and *n*-alkanes (e.g., Daniels et al., 2017; Berke et al., 2019; Dion-Kirschner et al., 2020).

In our sedimentary $\delta^2\text{H}$ records, there is a clear separation between mid- and long-chain waxes (except after 3.6 cal. kyr BP in Kløverbladvatna). The isotope values and trends are similar for homologues within each group but differ between the groups. This suggests that mid- and long-chain waxes come from two different sources. In July 2018, we collected a selection of the most common plants growing in the northern Wijdefjorden area (close to Austre Nevlingen). The relative *n*-alkanoic chain-length distribution of plants from four plant groups (dwarf shrubs: *Salix polaris*; rushes: *Luzula confusa*; terrestrial mosses: *Polytrichum* sp. and aquatic mosses:

unidentified) are presented in **Paper I**. The total lipid extract was extracted from freeze-dried plant material using sonication in DCM:methanol 9:1, after which the samples were treated and processed the same way as the sedimentary lipid samples (Fig. 8). *Luzula confusa* and *Salix polaris* had the highest relative C₂₈ concentrations and the lowest concentrations of mid-chain waxes, whereas the mosses contained relatively higher C₂₀ and C₂₂ waxes. Thus, we interpret the mid-chain waxes to mainly be produced by aquatic bryophytes and long-chain waxes by terrestrial plants, in agreement with previous studies (e.g., Thomas et al., 2016; Daniels et al., 2017; Berke et al., 2019).

2.1.5. Precipitation isotope seasonality reflected in the lake water

When interpreting proxy data from a sedimentary record, it is important to know the drivers of modern isotope variability. In this study, we focus on seasonal changes in precipitation, which means that it is crucial to understand what precipitation isotope seasonality is reflected in $\delta^2\text{H}_{\text{terr}}$ and $\delta^2\text{H}_{\text{aq}}$. To help guide our leaf wax $\delta^2\text{H}$ interpretations, we calculated seasonal residence times and collected surface water samples from the lakes. The residence time was calculated based on the lake volume and catchment size, together with the seasonal precipitation amount from the most representative meteorological station (for details, see **Paper III**). Water samples were collected from the lakes in August 2018 (Austre Nevlingen), August 2019 (Jodavannet), and August 2021 (Heftyevatnet and Kløverbladvatna) (Fig. 9). In addition, 21 lakes were sampled on northern Spitsbergen in July-August 2018 (Fig. 9b), three lakes on Sjuøyane in August 2018, three lakes on western Spitsbergen in April 2018, two lakes on northern Spitsbergen in August 2019, three lakes on western Spitsbergen and one lake on Nordaustlandet in August 2021 (Fig. 9a). These samples were collected using an opportunistic approach, i.e., collected during various field trips by me, co-authors, or others, and give a rough idea of the spatial lake water isotopic variability on Svalbard. The lake water isotopes were measured on a Picarro L2130-i WS-CRDS at OSIBL (for instrumentation and procedures, see Table 3). To get indications of the precipitation seasonality reflected by the lake water, we compared the lake isotopes to measured precipitation isotopes on Svalbard and to precipitation isotope values modeled using the Online Isotopes in Precipitation Calculator (OIPC; Bowen et al., 2005; IAEA/WMO, 2019; Bowen, 2021).

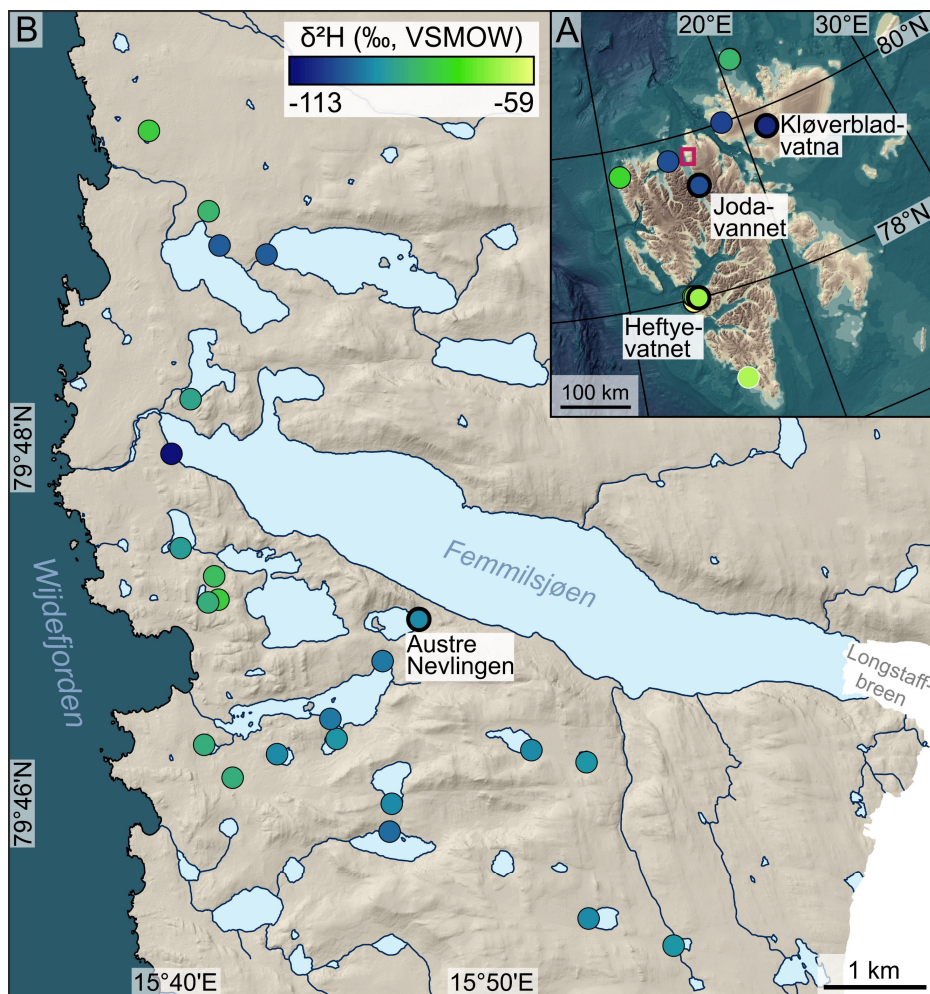


Fig. 9. Spatial summer (July–August) lake water $\delta^2\text{H}$ variability on Svalbard (a) and zoomed in to an area on northern Spitsbergen (b). Lakes analyzed for sedimentary leaf wax $\delta^2\text{H}$ in this study are labelled and outlined with a bold black line. The red box in (a) shows the extent of (b). The southernmost sample in (a) is from Arppe et al. (2017), outlined in white. Background maps in (a) from IBCAO (Jakobsson et al., 2012) and in (b) based on 5x5 m DEM from the Norwegian Polar Institute (2014).

Heftyevatnet is located on the western flank of Grønfjorden, on the humid west coast of Spitsbergen (Figs 4 and 6a-b). The lake is fed by catchment runoff and has a large catchment-to-lake ratio suggesting a short residence time. This was supported by $\delta^2\text{H}_{\text{lake}}$ close to summer $\delta^2\text{H}_{\text{precip}}$ at Isfjord Radio (14 km southwest of the lake). We therefore interpret both $\delta^2\text{H}_{\text{terr}}$ and $\delta^2\text{H}_{\text{aq}}$ from Heftyevatnet to reflect summer $\delta^2\text{H}_{\text{precip}}$ and use the difference between the two to reconstruct evapotranspiration.

Jodavannet is a small lake on the eastern flank of Wijdefjorden, northern Spitsbergen (Figs 4 and 6c-d). The lake receives surface runoff from the surrounding slopes and drains into a larger lake in the southeast. The lake has a large catchment-to-lake ratio, indicating a short residence

time. We targeted Jodavannet because we wanted a summer-biased $\delta^2\text{H}_{\text{aq}}$ record from northern Spitsbergen. However, ^2H -depleted modern $\delta^2\text{H}_{\text{lake}}$ suggested that the lake water is reflecting mean annual to winter precipitation. This could be a result of the exceptionally dry summers in this area (Elvebakk & Nilsen, 2002; Østby et al., 2017), and summer precipitation contributing a relatively small portion of the annual precipitation. Thus, we use Jodavannet $\delta^2\text{H}_{\text{aq}}$ to reconstruct changes in the seasonal distribution of precipitation (for further discussion, see **Paper III**). $\delta^2\text{H}_{\text{terr}}$ is interpreted in terms of summer precipitation $\delta^2\text{H}$ but might also incorporate some winter precipitation due to more winter-biased soil water.

Austre Nevlingen is located *c.* 50 km north of Jodavannet, close to the mouth of Wijdefjorden (Figs 4 and 6e-f). This area is slightly less arid (Østby et al., 2017). The lake has a small catchment-to-lake ratio and ^2H -depleted $\delta^2\text{H}_{\text{lake}}$, indicating a long residence time. $\delta^2\text{H}_{\text{terr}}$ and $\delta^2\text{H}_{\text{aq}}$ are interpreted to reflect the same precipitation $\delta^2\text{H}$ seasonality as in Jodavannet, i.e., seasonal distribution of precipitation and summer precipitation $\delta^2\text{H}$.

Kløverbladvatna is a threshold lake on Oxfordhalvøya, Nordaustlandet (Figs 4 and 6g-h). The peninsula sits between three ice caps, resulting in a relatively dry climate. A dry channel on the southeastern side indicates that the lake has previously received meltwater from Etonbreen, but the lake has no active inlet today. Kløverbladvatna has a small catchment-to-lake ratio and ^2H -depleted $\delta^2\text{H}_{\text{lake}}$ indicating a long residence time. We interpret $\delta^2\text{H}_{\text{terr}}$ to reflect summer $\delta^2\text{H}_{\text{precip}}$ (possibly incorporating some winter $\delta^2\text{H}_{\text{precip}}$), and $\delta^2\text{H}_{\text{aq}}$ to reflect mean annual $\delta^2\text{H}_{\text{precip}}$.

Ideally, we would have precipitation $\delta^2\text{H}$ data from the lake locations to quantify how strong the precipitation isotope seasonality is at present. The only available $\delta^2\text{H}_{\text{precip}}$ are from the humid west coast of Spitsbergen. In August 2019, we installed a 4-inch diameter RG202 Stratus All-Weather Rain Gauge in Adventdalen (*c.* 9 km southeast of Longyearbyen; Fig. 4) to record $\delta^2\text{H}_{\text{precip}}$ at a more arid location. For logistical reasons, the gauge was moved to Longyearbyen and later to Nybyen (2.5 km farther inside Longyeardalen), where UNIS students could help us to collect the samples. There were several issues with the sampling: 1) long periods without precipitation, 2) low precipitation amounts and strong winds resulted in undercatch and very small samples, 3) polar bear presence prevented collection of the samples some days, and 4) the gauge was unattended in periods when we did not have anyone that could collect the samples. Additionally, varying local conditions at the different gauge locations and sampling

by different persons add uncertainty to the measurements. Ultimately, we decided to stop the sample collection in June 2020, since we were not able to generate a coherent dataset. Thus, the Adventdalen/Longyearbyen samples ($n = 45$) have not been analyzed and are not presented in this thesis.

2.2. Northern Fennoscandian lakes

2.2.1. Water sample collection

For **Paper II**, we collected surface water samples from 135 lakes in the summers of 2018, 2019, and 2020 (Figs 5 and 10). The lakes varied in size from $<0.01 \text{ km}^2$ to $>10 \text{ km}^2$, and lake elevation from sea level to *c.* 700 m a.s.l. The water sampling was mainly focused to the first week in July each year to facilitate interannual comparison. In July 2018, the sampling was concentrated to a 5 km^2 area around Tromsø, coastal Norway ($n = 30$). In the end of June 2019, we collected samples along the north Norwegian coast ($n = 9$), and the July 2019 and 2020 campaigns targeted lakes along a coastal-to-inland transect (Fig. 5). The transect comprises easy-accessible lakes along the European route E8, from the Lyngen Alps (Norway) close to the Norwegian Sea in northwest, to Kemi (Finland) by the Bothnian Bay in the southeast, i.e., between 69.4°N , 20.3°E and 65.7°N , 24.6°E . In 2019, the transect reached 230 km inland ($n = 35$), and in 2020 the transect was extended farther southeast, to a total of 460 km ($n = 68$). The transect spans elevations from close to sea level up to *c.* 560 m a.s.l., with the highest elevation close to the tripoint between Norway, Finland, and Sweden (Fig. 5). Eleven of the transect lakes were sampled in 2013 by Balascio et al. (2018a), and twelve of them were sampled in both 2019 and 2020. We also sampled two locations on the Muonio River in both years, as well as resampled some of the coastal lakes ($n = 2$ and $n = 4$ in 2019 and 2020, respectively). The samples were collected from the shoreline, *c.* 10 cm below the surface to avoid evaporatively enriched surface water. The samples were stored in 4 mL glass vials with no headspace, sealed with Parafilm, and kept cold until analysis.

To infer the seasonality of the lake water, we compared the $\delta^2\text{H}_{\text{lake}}$ to $\delta^2\text{H}_{\text{precip}}$ from Naimakka and Rovaniemi (Fig. 5; IAEA/WMO, 2019). Naimakka is located on the eastern side of the Scandinavian Mountains, and Rovaniemi farther southeast, just south of the Arctic Circle. To record the variability in precipitation isotopes closer to the Norwegian Sea, we installed a 4-inch diameter RG202 Stratus All-Weather Rain Gauge on the roof of the UiT Geoscience

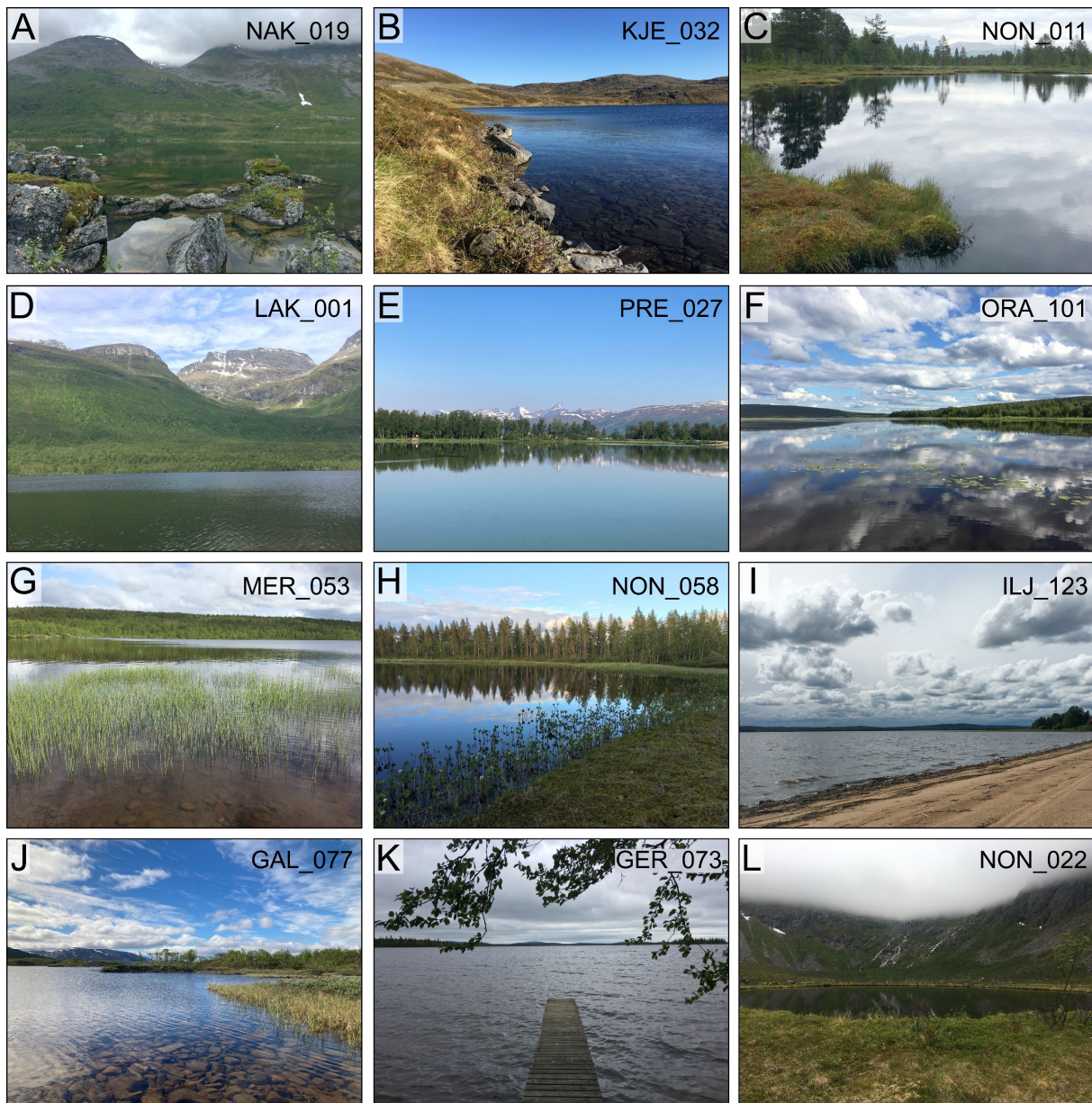


Fig. 10. Photographs of a selection of sampled lakes in northern Norway, Finland, and Sweden. Codes refer to Site IDs in the Supplementary data for **Paper II**.

building (69.68°N, 18.96°E; 73 m a.s.l.) in September 2019. Precipitation samples used to calculate amount-weighted mean monthly values presented in **Paper II** were collected every morning between September 6, 2019, and September 5, 2021 (n = 407). Solid precipitation was collected directly from the 4-inch cylinder, whereas liquid samples were collected using a funnel and an inner 1-inch cylinder. Frozen samples were melted in a sealed container at room temperature. Temperature and precipitation amount (or snow water equivalent) was recorded at every sample collection. All samples were transferred to 4 mL glass vials, sealed with Parafilm, and kept cold until analysis.

2.2.2. Water isotope analysis and inflow $\delta^2\text{H}$ inference

All lake water and precipitation samples were filtered through a 0.2- μm PTFE filter before being analyzed for their $\delta^2\text{H}$ and $\delta^{18}\text{O}$ isotopic composition. Lake water samples collected in 2018–2019 were analyzed at UB, whereas the 2020 lake water samples and 2019–2021 precipitation samples were measured at FARLAB. Information on instrumentation, number of injections per sample, standards, corrections, calibrations, and uncertainties are presented in **Paper II** and in Table 3.

In **Paper II**, we present lake water and precipitation $\delta^2\text{H}$ and $\delta^{18}\text{O}$, as well as d-excess calculated using Equation 3 (section 1.2). To facilitate comparison between our new

Table 3. Instrumentation and procedures used for water $\delta^2\text{H}$ and $\delta^{18}\text{O}$ measurements at the University at Buffalo Organic and Stable Isotope Biogeochemistry Laboratory (OSIBL) and the Facility for advanced isotopic research and monitoring of weather, climate, and biogeochemical cycling (FARLAB), University of Bergen.

	OSIBL	FARLAB
	2018–2019 lake water	2020 lake water 2019–2021 precipitation
Analyzer	Picarro L2130-i WS-CRDS	Picarro L2140-i WS-CRDS
Vaporization module	V1102-i	A0211
Autosampler	CTC PAL	A0325
Total no. of injections per sample	4	12
No. of injections per sample used to get reported value (average)	3	4–6
Range of in-house standards (calibrated against primary standards GISP, VSMOW2, and SLAP2)	–265.86 to +21.43‰ ($\delta^2\text{H}$) –33.62 to +12.41‰ ($\delta^{18}\text{O}$)	–
Range of drift standard and two laboratory standards (calibrated against VSMOW2 and SLAP2)	–	–308.14 to +9.2‰ ($\delta^2\text{H}$) –40.06 to +1.77‰ ($\delta^{18}\text{O}$)
Memory correction	van Geldern and Barth (2012)	Gröning (2011)
Average standard deviation of replicate measurements	0.08‰ ($\delta^2\text{H}$) 0.02‰ ($\delta^{18}\text{O}$)	–
Average standard uncertainties for calibrated measurement values (provided according to Gröning (2018))	–	0.43‰ ($\delta^2\text{H}$) 0.06‰ ($\delta^{18}\text{O}$)

precipitation dataset and published precipitation $\delta^2\text{H}$ data, we converted the daily precipitation data into monthly amount weighted (aw) mean values using the equation:

$$\delta_{aw} = \frac{1}{P} \times \sum_{i=1}^n (p_i \times \delta_i) \quad (\text{Equation 6})$$

where P and p_i are the monthly and daily precipitation amounts in mm, δ_i is $\delta^2\text{H}$ or $\delta^{18}\text{O}$, and n is the number of sampling days.

When using water isotope proxies to infer precipitation $\delta^2\text{H}$, it is important to quantify the possible influence of evaporation. Based on the assumption that isotope values of partially evaporated water will evolve along an evaporation line (EL), a widely used approach is to estimate the isotopic composition of the source water by back-correction to the EL-MWL intersection (Clark & Fritz, 1997). This assumes that all $\delta^2\text{H}_{\text{lake}}$ variation results from evaporation, and that all lakes in a region have the same source water (i.e., precipitation) isotopic composition. At high latitudes and in systems with strong seasonality, the source isotopic composition can vary between years due to changing relative proportion of isotopically depleted snowfall and isotopically enriched rainfall (Tondou et al., 2013). To estimate the source water $\delta^2\text{H}$ for evapo-concentrated lakes, we applied the Bayesian ‘MWL source implementation’ method by Bowen et al. (2018). This approach to estimate source water values includes uncertainty in the MWL and the LEL, thus providing a more rigid estimate of source water $\delta^2\text{H}$ uncertainty. To run the model, the user supplies a MWL equation and a hypothesized LEL slope (both with confidence intervals), lake water $\delta^2\text{H}$ and $\delta^{18}\text{O}$ with uncertainties and covariance (0.95), and the number of iterations (10,000). We used a re-calibrated GMWL ($\delta^2\text{H} = 8.01 \times \delta^{18}\text{O} + 9.57$; $n = 80,672$; Bowen et al., 2018) for the MWL equation, having much less uncertainty than the LMWLs, and applied different LEL slopes for samples collected in 2019 and 2020 (4.82 and 5.18, respectively). These LEL slopes were based on linear regression for the isotopic composition of lakes sampled in both years ($n = 12$), providing a reasonable starting estimate. The modeled mean source water isotope values, referred to as inferred inflow $\delta^2\text{H}$ ($\delta^2\text{H}_i$) were used to compare the seasonality of inflow along the transect and to non-evaporated coastal lakes.

3. Summary of research papers

In this thesis, I present two studies of sedimentary leaf wax $\delta^2\text{H}$ from lakes on Svalbard, and one study of lake water $\delta^2\text{H}$ and $\delta^{18}\text{O}$ from northern Fennoscandia. **Paper I** presents a case study from Austre Nevlingen, reconstructing Holocene precipitation seasonality on northern Spitsbergen. In **Paper III**, Austre Nevlingen is complemented by records from three additional lakes: Heftyevatnet, Jodavannet and Kløverbladvatna. The four lakes are located along a climatic gradient across Svalbard and reflect different precipitation seasonality, allowing us to investigate Holocene seasonal precipitation changes in both time and space. **Paper II** focuses on the modern isotopic composition of 135 lakes in northern Fennoscandia, sampled from 2018 to 2020 and spanning a coastal-to-inland transect from the Norwegian Sea to the Bothnian Bay.

3.1. Paper I

Kjellman, S. E., Schomacker, A., Thomas, E. K., Håkansson, L., Duboscq, S., Cluett, A. A., Farnsworth, W. R., Allaart, L., Cowling, O. C., McKay, N. P., Brynjólfsson, S., & Ingólfsson, Ó., 2020. **Holocene precipitation seasonality in northern Svalbard: influence of sea ice and regional ocean surface conditions.** *Quaternary Science Reviews*, 240, 1063882. <https://doi.org/10.1016/j.quascirev.2020.106388>.

A critical knowledge gap in the Svalbard climate reconstructions is the lack of robust records of Holocene precipitation changes. In **Paper I**, we present sedimentary leaf wax $\delta^2\text{H}$ data from Austre Nevlingen, northern Spitsbergen. We targeted a lake with a long residence time (i.e., with $\delta^2\text{H}_{\text{lake}}$ reflecting mean annual $\delta^2\text{H}_{\text{precip}}$) to allow us to extract information about seasonal variability. The hydrogen isotopic composition of C_{22} and C_{28} *n*-alkanoic acids (mainly derived from aquatic and terrestrial plants, respectively) were used to infer past lake and soil water $\delta^2\text{H}$, interpreted to reflect mean annual and summer precipitation. Our results suggest that northern Spitsbergen experienced large variability in precipitation seasonality throughout the Holocene. ^2H -enriched $\delta^2\text{H}_{\text{terr}}$ in the Early and Middle Holocene (until *c.* 5.6 cal. kyr BP) suggests enhanced summer evaporation and/or ^2H -enriched summer precipitation, indicating warm and stable summer conditions. Decreasing $\delta^2\text{H}_{\text{aq}}$ in the Early Holocene and ^2H -depleted values until *c.* 6 cal. kyr BP likely reflect prominent increase in winter precipitation amounts. We propose that Early Holocene warming had a strong effect on the moisture availability. High summer insolation and increased Atlantic heat flux warmed the waters around Svalbard and caused

reduction in sea-ice cover. More open water promoted evaporation and greater winter precipitation amounts. After *c.* 5.6 cal. kyr BP, decreasing $\delta^2\text{H}_{\text{terr}}$ suggests gradual summer cooling, with less evaporative enrichment and/or more ^2H -depleted summer precipitation. The $\delta^2\text{H}_{\text{aq}}$ trend is not as clear after 6 cal. kyr BP, which we attribute to changes in lake hydrology resulting in large shifts in $\delta^2\text{H}_{\text{aq}}$. We propose that the duration of lake-ice cover might have varied with variable climate, affecting the amount of snowmelt bypass and thus the relative proportion of summer and winter precipitation entering the lake. The Austre Nevlingen record suggests that the Holocene precipitation seasonality on northern Spitsbergen is strongly linked to regional ocean surface conditions and sea-ice cover, which is in agreement with modern observations and future projections.

3.2. Paper II

Kjellman, S. E., Thomas, E. K., & Schomacker, A., 2022. **Arctic and sub-Arctic lake water $\delta^2\text{H}$ and $\delta^{18}\text{O}$ along a coastal-inland transect: implications for interpreting water isotope proxy records.** *Journal of Hydrology*, 67, 127556.
<https://doi.org/10.1016/j.jhydrol.2022.127556>.

Stable isotope proxies that indirectly archive the isotopic composition of hydrogen and oxygen in past precipitation are widely used to reconstruct various aspects of the water cycle. A common approach is to assume a single control on the lake water isotope variability for a given record. One of the outcomes of **Paper I** is that the precipitation seasonality reflected in $\delta^2\text{H}_{\text{lake}}$ can vary through time because of changes in the lake dynamics. Another crucial part of interpreting water isotope proxy records is to separate the precipitation isotope signal from evaporation. In **Paper II**, we measured $\delta^2\text{H}$ and $\delta^{18}\text{O}$ of surface water from 135 lakes in northern Fennoscandia over three years, to study how their isotopic composition varies in time and space. We targeted easy-accessible lakes along a 460-km-long transect from the Norwegian Sea in northwest, to the Bothnian Bay in the southeast. This is a region with strong contrasts in climate, with prevailing westerly winds bringing Atlantic moisture ashore, and the Scandinavian Mountains shielding the leeward side from precipitation. To estimate the source isotopic composition (i.e., the lake water inflow values; $\delta^2\text{H}_i$) for evapo-concentrated transect lakes, we used a Bayesian method. Our results show that there is spatial and temporal variation in $\delta^2\text{H}_{\text{lake}}$ in this region. Both coastal and inland lakes are sensitive to distillation during atmospheric moisture transport, controlling the inflow isotopic composition. In addition, lakes

farther from the Norwegian Sea are more sensitive to evaporation. We suggest that more ^2H -depleted $\delta^2\text{H}_I$ in 2020 than in 2019 is a result of a snow-rich winter 2020, attributed to extreme AO+ conditions and increased moisture supply from the North Atlantic. The large snow amounts resulted in more inflow of ^2H -depleted winter precipitation to the lakes. We conclude that the lake water isotopic variability in northern Fennoscandia reflects seasonal precipitation changes associated with atmospheric circulation changes, as well as catchment-integrated evaporation. Through-flowing coastal lakes are most suited for precipitation reconstructions, whereas $\delta^2\text{H}_{\text{lake}}$ variability in more inland lake is likely to incorporate both changes in inflow variability and evaporation.

3.3. Paper III

Kjellman, S. E., Thomas, E. K., Schomacker, A., Farnsworth, W. R., Cowling, O. C., Allaart, L., & Brynjólfsson, S. **Holocene precipitation seasonality along a climatic gradient from western Spitsbergen to Nordaustlandet, Svalbard.** In review for *Palaeogeography, Palaeoclimatology, Palaeoecology*.

The Svalbard archipelago spans large climate gradients due to complex feedback mechanisms in the atmosphere-ocean-ice system. Climate models project that Svalbard will experience heterogeneous precipitation changes in the future, with largest increase in northeast and during the winter months. In **Paper I**, we suggested that northern Spitsbergen experienced increased winter precipitation during the warm Early Holocene. In **Paper III**, we apply the same method to three additional lakes across Svalbard to explore regional differences in Holocene water cycle changes. From Heftyevatnet, on the relatively warm and humid west coast of Spitsbergen, we interpret $\delta^2\text{H}_{\text{terr}}$ and $\delta^2\text{H}_{\text{aq}}$ to reflect summer precipitation $\delta^2\text{H}$. We use the difference between the two (ϵ_{28-22}) to infer changes in evapotranspiration. The other three lake records are from the relatively cold and arid northeastern Svalbard: Jodavannet and Austre Nevlingen on northern Spitsbergen, and Kløverbladvatna on Nordaustlandet. We interpret $\delta^2\text{H}_{\text{terr}}$ from these three lakes to mainly reflect summer precipitation, but with some incorporation of winter precipitation due to more winter-biased soil water. $\delta^2\text{H}_{\text{aq}}$ reflects variable seasonal distribution of precipitation due to varying lake hydrology. In the Early and Middle Holocene, relatively ^2H -enriched $\delta^2\text{H}_{\text{terr}}$ suggests dry summer conditions and/or more proximal summer moisture across Svalbard, and high ϵ_{28-22} suggests high evapotranspiration. We interpret an Early Holocene decrease in $\delta^2\text{H}_{\text{aq}}$ in both Austre Nevlingen and Jodavannet to reflect an increase in

winter relative to summer precipitation, associated with increased moisture supply as the sea-ice cover decreased with regional warming. Winter precipitation likely stayed high until *c.* 6 cal. kyr BP. After 6–5.5 cal. kyr BP, ^2H -depleted $\delta^2\text{H}_{\text{terr}}$ and lower ε_{28-22} indicate cooler summer, less evapotranspiration, and/or more distal moisture. Intermittent ^2H -depleted $\delta^2\text{H}_{\text{aq}}$ suggests periodic increases in winter precipitation or decreases in summer precipitation inflow to the lakes on northern Spitsbergen. Our results show the potential of the leaf wax $\delta^2\text{H}$ proxy to reconstruct seasonal distribution of precipitation, summer precipitation $\delta^2\text{H}$, and/or evapotranspiration, depending on the hydrology of the lake.

3.4. Author contributions

The papers and manuscript in this thesis are the result of collaboration between researchers from different institutions, contributing to various parts of each study. All co-authors and individual contributions are listed in Table 4.

Table 4. Overview of author contributions to each of the research articles (**Papers I–III**). The contributor roles have been defined according to the CRediT (Contributor Roles Taxonomy) criteria (<https://casrai.org/credit/>). AAC = Allison A. Cluett; AS = Anders Schomacker; EKT = Elizabeth K. Thomas; LA = Lis Allaart; LH = Lena Håkansson; NPM = Nicholas P. McKay; OCC = Owen C. Cowling; ÓI = Ólafur Ingólfsson; SB = Skafti Brynjólfsson; SD = Sandrine Dubosq; SEK = Sofia E. Kjellman; WRF = Wesley R. Farnsworth.

Contribution	Paper I	Paper II	Paper III
Conceptualization	SEK, AS, EKT, LH	SEK, AS, EKT	SEK, AS, EKT
Data curation	SEK	SEK	SEK
Formal analysis	SEK, EKT, OCC	SEK, AS, EKT	SEK, EKT, OCC
Funding acquisition	AS, LA, ÓI, EKT, SD	AS, SEK, EKT	AS, SEK, EKT, WRF
Investigation	SEK ^{1,2,3} , AS ^{1,2,3} , EKT ^{2,3} , LH ² , SD ^{2,3} , AAC ^{2,3} , WRF ^{1,2} , LA ¹ , OCC ² , SB ¹ , ÓI ¹	SEK ^{1,3} , EKT ³ , AS ^{1,3}	SEK ^{1,2,3} , EKT ^{2,3} , AS ^{1,2,3} , WRF ^{1,2} , OCC ² , LA ¹ , SB ¹
Project administration	SEK, AS	SEK, AS	SEK, AS
Software	SEK, EKT, NPM	SEK	SEK
Supervision	AS, EKT, LH	AS, EKT	AS, EKT
Visualization	SEK	SEK	SEK
Writing – original draft	SEK	SEK	SEK
Writing – review & editing	SEK, AS, EKT, LH, SD, AAC, WRF, LA, OCC, NPM, SB, ÓI	SEK, EKT, AS	SEK, EKT, AS, WRF, OCC, LA, SB

4. Discussion

4.1. Source and seasonality of *n*-alkanoic acids

Robust proxy interpretations rely on a well-grounded understanding of the environmental factors driving the isotopic variation. Leaf wax $\delta^2\text{H}$ is a useful tool to infer past hydrological changes because of the strong linear relationship with precipitation $\delta^2\text{H}$ (Sachse et al., 2004; Hou et al., 2008; McFarlin et al., 2019). However, the translation from source water to leaf wax $\delta^2\text{H}$ involves a series of isotopic fractionation processes related to both plant physiology and environmental factors (Sachse et al., 2012). Furthermore, to interpret leaf wax $\delta^2\text{H}$ in terms of seasonal precipitation variability, we need to consider the seasonality of the source (soil or lake water) pools. In this section, I discuss some of the challenges of making accurate leaf wax source attribution and inferring the source water seasonality.

The first step in leaf wax $\delta^2\text{H}$ interpretation is to consider the source of the leaf wax homologues. In **Papers I & III**, the $\delta^2\text{H}$ of all even-chain length *n*-alkanoic acids in our four lakes, except for part of the Kløverbladvatna record, show different trends for the mid- and long-chain homologues. This indicates that they come from two different plant groups. Globally, studies have shown that terrestrial plants primarily produce long-chain waxes (Sachse et al., 2012; Diefendorf & Freimuth, 2017), whereas aquatic plants produce relatively higher concentrations of mid-chain waxes (Ficken et al., 2000; Aichner et al., 2010b). Also in the Arctic, terrestrial plants produce high relative concentrations of long-chain waxes (e.g., Wilkie et al., 2013; Daniels et al., 2017; Berke et al., 2019). The few Arctic studies that examine aquatic plants suggest that they produce higher concentrations of mid-chain than long-chain homologues (Thomas et al., 2016; Berke et al., 2019; Gorbey et al., 2021). On Svalbard, there are almost no aquatic vascular plants, but submerged bryophytes are abundant in some lakes (Balascio et al., 2018b; Voldstad et al., 2020; **Papers I & III**). Aquatic bryophytes from western Greenland ($n = 1$; Thomas et al., 2016) and Baffin Island ($n = 4$; Gorbey et al., 2021) are dominated by C_{24} *n*-alkanoic acids, followed by C_{22} and C_{26} . Balascio et al. (2018b) measured the concentrations of C_{19} – C_{33} *n*-alkanes in one aquatic bryophyte sample from Amsterdamøya, northwestern Svalbard, with C_{27} , C_{25} and C_{21} being the most abundant homologues.

In **Paper I**, we present the average chain-length distribution of aquatic bryophytes ($n = 2$) from northern Spitsbergen, displaying similar concentration of C_{24} , C_{26} and C_{28} *n*-alkanoic acids,

which are the most abundant waxes in the bryophytes (Fig. 11). *Salix polaris* has a similar distribution as the aquatic mosses, whereas *Luzula confusa* is dominated by C₂₈, and *Polytrichum* sp. is dominated by C₂₀ and C₂₂ (Fig. 11). In absolute concentrations (μg *n*-alkanoic acids per g dry biomass), *Salix polaris* and *Luzula confusa* produce the greatest amount of the long-chain waxes per g biomass and *Polytrichum* the greatest amounts of C₂₀ and C₂₂. *Salix polaris* produces higher amounts of all chain lengths compared to the aquatic bryophytes. Furthermore, the average distribution of *n*-alkanoic acids in the sediments vary between the lakes (Fig. 11), reflecting the different plant communities in the catchments. These chain-length observations highlight the importance of the $\delta^2\text{H}$ patterns for source attribution. From the chain-length distribution alone, it is difficult to separate terrestrial and aquatic plants, but the clear separation between mid- and long-chain $\delta^2\text{H}$ trends supports the canonical two-source interpretation. In Kløverbladvatna, all homologues show similar trends after 3.6 cal. kyr BP, which could indicate a source change (i.e., different sources before and same source after 3.6 cal. kyr BP), as the chain-length distribution in the sediments changes (Fig. 11). Another possibility is that the dominant wax producers changed, but more details on the leaf wax sources at this site are needed to confirm the reason for the shift in $\delta^2\text{H}$.

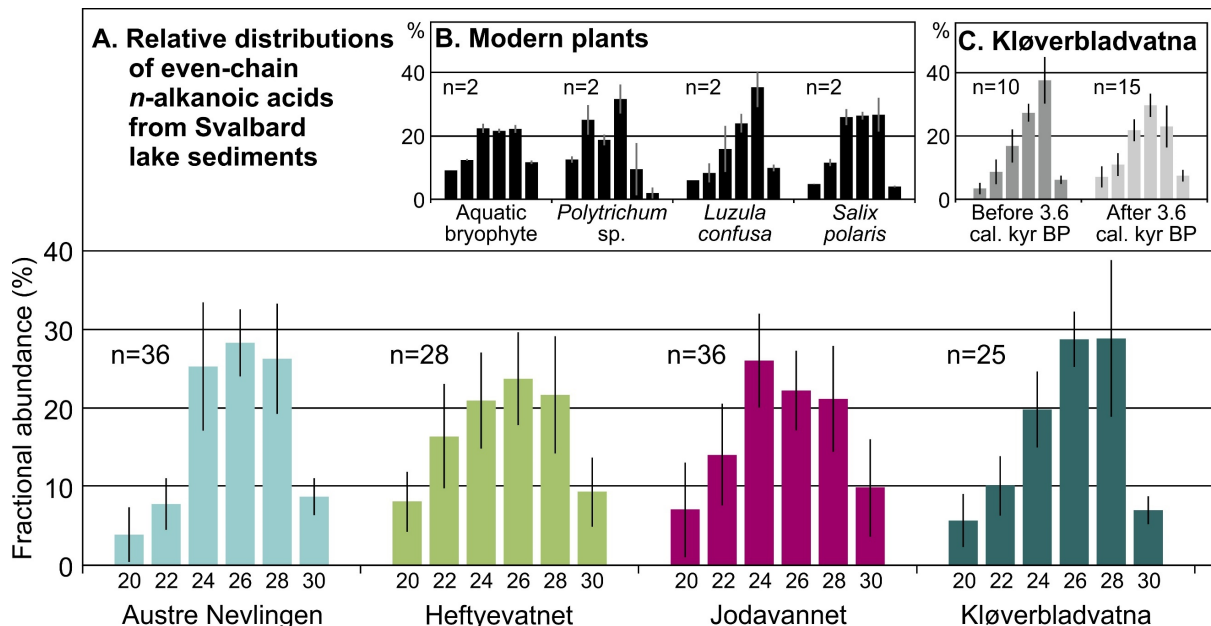


Fig. 11. Distribution of even-chain *n*-alkanoic acids between C₂₀ and C₃₀ from (a) Svalbard lake sediments (**Papers I & III**), (b) modern plants from northern Spitsbergen (**Paper I**), and (c) in Kløverbladvatna sediments before and after 3.6 cal. kyr BP (**Paper III**). Error bars show one standard deviation.

To extract information of seasonal precipitation changes from leaf wax $\delta^2\text{H}$, we need to assess the seasonality reflected in the source water pools used by the plants. In previous Arctic studies, terrestrial leaf wax $\delta^2\text{H}$ is interpreted to reflect summer precipitation $\delta^2\text{H}$ and summer evaporative enrichment, assuming that the soil water during the growing season is mainly recharged by summer precipitation (Cooper et al., 1991; Thomas et al., 2016; Throckmorton et al., 2016; Thomas et al., 2020). In **Paper III**, we suggest that the soil water on northern Svalbard might reflect a mix of summer and winter precipitation. We base this argument on a few lines of evidence. Soil moisture measurements close to the Jodavannet catchment suggest that most of the summer soil moisture comes from snowmelt early in the season (Eidesen et al., 2018). Additionally, summer precipitation amounts on northeastern Svalbard are very low, and might not saturate the active layer by the end of summer, which therefore subsequently gets recharged by winter precipitation during the following snowmelt season. Furthermore, $\delta^2\text{H}_{\text{terr}}$ in the three northeastern lakes are ^2H -depleted compared to Heftyevatnet on the west coast. This could reflect a greater influence of ^2H -depleted winter precipitation on the soil water, and/or more Rayleigh distillation during moisture transport from the Atlantic Ocean.

Aquatic leaf wax $\delta^2\text{H}$ can reflect different precipitation seasonality depending on the residence time of the lake (e.g., Thomas et al., 2020). The residence time can be calculated based on the lake volume, catchment size, and runoff estimated from the precipitation amount (Jonsson et al., 2009). We lack good constraints on precipitation amounts falling over central and northeastern Svalbard, meaning that the calculated residence times are rough estimates. In **Paper III**, Jodavannet had unexpectedly ^2H -depleted $\delta^2\text{H}_{\text{lake}}$ considering the presumably short residence time (Table 1). We suggest three possible explanations for why Jodavannet may not reflect summer precipitation, including low summer precipitation amounts that do not flush the lake in summer, evaporation of summer precipitation before it reaches the lake, and inflow of winter-biased groundwater. Furthermore, the residence time might change through time. In **Papers I & III**, we observe large shifts in $\delta^2\text{H}_{\text{aq}}$ in Austre Nevlingen and Jodavannet, which we suggest reflect variable sensitivity of the lake basins to the seasonal distribution of precipitation. Processes influencing the relative proportion of summer and winter precipitation in the lake during the growing season and/or the residence time include the duration of lake-ice cover, degree of snowmelt bypass, and lake-level changes. In **Paper II**, we show that lake water $\delta^2\text{H}$ is sensitive to variable winter precipitation amounts from one year to the next. In

conclusion, it is important to evaluate temporal changes in lake and catchment processes influencing the precipitation seasonality reflected in lake and soil water isotope proxies.

4.2. Regional variability in paleo-precipitation seasonality

One of the main focuses of all three papers is seasonal changes in precipitation $\delta^2\text{H}$. In **Paper II**, modern lake water $\delta^2\text{H}$ registered variable input of winter (^2H -depleted) relative to summer (^2H -enriched) precipitation from one year to the next, implying that lake water isotope proxies from carefully selected lakes can be interpreted in terms of seasonal distribution of precipitation. Paleoclimate records can provide information about how the hydrological cycle responded to past climate change, but depending on the hydrology of the lake, they may register different precipitation seasonality.

In modern studies it is easier to constrain the drivers behind $\delta^2\text{H}_{\text{lake}}$ (and therefore indirectly $\delta^2\text{H}_{\text{precip}}$) changes because of available meteorological and hydrological data. We can monitor seasonal changes in real time and attribute variations in $\delta^2\text{H}_{\text{lake}}$ to observed processes. For instance, we observed more ^2H -depleted lake water isotopes in summer 2020 than in summer 2019. We interpret the ^2H -depleted values to reflect relatively higher winter input in 2020, supported by large snow amounts falling in winter and spring 2020. In paleo-precipitation studies, we cannot make direct observation of environmental changes during the time of proxy synthesis. This makes inference of the drivers of seasonal precipitation changes from proxy records more challenging. In **Paper I**, we targeted a lake which, based on the modern $\delta^2\text{H}_{\text{lake}}$ and residence time calculations, likely has $\delta^2\text{H}_{\text{aq}}$ reflecting mean annual precipitation $\delta^2\text{H}$. Large shifts in the $\delta^2\text{H}_{\text{aq}}$ during the Middle and Late Holocene suggested that was not always the case, but that hydrological changes could alter the seasonality reflected by the lake.

Papers I & III suggested that northern Svalbard likely received a higher proportion of winter precipitation relative to the annual total in the Early and Middle Holocene, until *c.* 6 cal. kyr BP, and during shorter periods also in the Late Holocene. The Early Holocene increase coincided with high summer insolation and increased advection of warm Atlantic water to Svalbard, and we interpret the increased winter precipitation to be a response to enhanced evaporation from ice-free seas. Reconstructions of seasonal distribution of precipitation can also help assess the importance of lower summer temperature versus higher winter precipitation amounts on glacier activity on Svalbard. There are multiple lines of evidence for greater

precipitation amounts on northern Svalbard in the Early Holocene. ^2H -depleted $\delta^2\text{H}_{\text{aq}}$ in our sedimentary leaf wax records suggests an increasingly greater portion of winter precipitation from *c.* 11.7 to 6 cal. kyr BP (**Paper I & III**), and higher frequency of “snowmelt layers” in lake sediments from Vårfluesjøen (Fig. 4) from the beginning of the record at 10.2 cal. kyr BP to 7 cal. kyr BP (Røthe et al., 2018). The snowmelt layers are fining-upwards silt layers interpreted to reflect short-lived events of increased runoff from snowmelt in the catchment. Winter precipitation has also been suggested to be one of the drivers of Neoglacial (Müller et al., 2012; Røthe et al., 2015) and LIA glacier expansion (D'Andrea et al., 2012; Arppe et al., 2017; Røthe et al., 2018) on Svalbard.

To place our seasonal precipitation trends into a wider context, we can consider paleoprecipitation and glacier records from other Arctic and sub-Arctic regions. In northern Greenland and on northwestern Iceland, increased winter precipitation is suggested to have helped ice caps to survive the HTM (Schomacker et al., 2016; Larsen et al., 2019). Furthermore, asynchronous Middle to Late Holocene glacier expansion in the Arctic indicates that the Neoglacial is not only forced by cooler temperatures associated with declining insolation (McKay et al., 2018), but driven by multiple mechanisms (Solomina et al., 2015; Kjær et al., 2022; Larocca & Axford, 2022). In this context, the effect of precipitation amount and seasonality is poorly understood. Temperature appears to be the dominant long-term control of the size of the Greenland Ice Sheet, but precipitation may drive shorter regional and local events (Alley et al., 2010).

Badgeley et al. (2020) generated independent reconstructions of mean annual temperature and precipitation for the last 20,000 years on Greenland using paleoclimate data assimilation, combining ice-core $\delta^{18}\text{O}$ and accumulation rates with climate-model simulations. The results suggested that annual precipitation closely followed temperature, with low precipitation amounts in the Younger Dryas, followed by an increasing trend in the Early Holocene, and relatively stable precipitation after *c.* 8 cal. kyr BP (Badgeley et al., 2020). On coastal western Greenland, Thomas et al. (2016) interpreted 100‰ more negative sedimentary leaf wax $\delta^2\text{H}_{\text{aq}}$ from 6 to 4 cal. kyr BP than during the Early and Late Holocene to reflect increased winter snowfall, coinciding with high summer temperature, increased heat advection from the West Greenland Current, and decreased winter sea-ice cover in Baffin Bay and the Labrador Sea. Coastal areas were likely more sensitive to local sea-ice changes than the central parts of the

Greenland Ice Sheet. Holocene sea-ice reconstructions suggest different sea-ice dynamics between different Arctic regions (de Vernal et al., 2013), which could partly explain the different timing of ^2H -depleted $\delta^2\text{H}_{\text{aq}}$ in lakes on Svalbard and in Greenland. Periods of enhanced evaporation from seas with reduced ice cover occur at different times on northern Svalbard (Early and Middle Holocene until *c.* 6 cal. kyr BP; **Papers I & III**) and western Greenland (6 to 4 cal. kyr BP; Thomas et al., 2016), yet highlighting that reduced regional sea ice and warmer sea surface temperatures are important controls of the moisture source for Arctic winter precipitation.

4.3. Improvement of the leaf wax $\delta^2\text{H}$ proxy and other water isotope proxies

As discussed in section 4.1, the hydrogen isotopic composition of leaf waxes depends on several biological and environmental factors. Paleo-hydrological interpretations based on $\delta^2\text{H}_{\text{wax}}$ differ between studies due to differences in lake hydrology (e.g., influencing the seasonality reflected in the lake water $\delta^2\text{H}$), but also highlights our incomplete understanding of the factors influencing lipid $\delta^2\text{H}$ variability (Linderholm et al., 2018). There are many possible directions for how to improve water isotope proxy interpretations. In this section, I present some advances made, as well as approaches that might prove fruitful.

4.3.1. Modern water isotope monitoring

A requirement for making accurate proxy interpretation is a good understanding of modern water cycle processes. Monitoring of precipitation $\delta^2\text{H}$ and $\delta^{18}\text{O}$ (e.g., through GNIP; IAEA/WMO, 2019) provides a baseline for interpreting seasonal precipitation variability, but the spatial distribution of high-Arctic measurements remains sparse, and time series often span short and different periods. Furthermore, short-term variation in observational records spanning only one or two years might bias long-term estimates based on those observations (Bowen & Revenaugh, 2003). On Svalbard, all precipitation measurements are from the west coast of Spitsbergen (Arppe et al., 2017; IAEA/WMO, 2019; Skakun et al., 2020). As part of this study, we tried to monitor precipitation isotopes in the more arid Longyearbyen area, but it proved challenging (see section 2.1.5). Future monitoring programs could help constrain the precipitation seasonality across Svalbard, if logistically feasible. In 2018, the Pan-Arctic Precipitation Isotope Network (PAPIN) was established to coordinate precipitation isotope

sampling across key climate types in the Arctic (Mellat et al., 2021). Our precipitation gauge installed in Tromsø to generate data for **Paper II** is still in operation and included as the first coastal Fennoscandian station in the PAPIN network.

We also need better constraints on the seasonal moisture sources. One way to investigate how moisture source changes impact the precipitation at a specific location is to use backtracking modeling. A study from western Greenland demonstrates that moisture source changes can dampen the precipitation isotope seasonality (Cluett et al., 2021). Furthermore, Lagrangian moisture diagnostic and isotope-enabled general circulation models have identified seasonal and interannual variability in moisture sources to Greenland with changing atmospheric circulation patterns and sea-ice cover, subsequently controlling precipitation $\delta^2\text{H}$ and $\delta^{18}\text{O}$ (Werner et al., 2001; Sodemann et al., 2008a; Sodemann et al., 2008b; Nusbaumer et al., 2019). Seasonal moisture source changes have also been suggested to determine precipitation $\delta^2\text{H}$ and $\delta^{18}\text{O}$ in western Siberia, based on in situ water vapor isotope measurements and semi-Lagrangian backward trajectories (Bonne et al., 2020). Climate on Svalbard is tightly linked to atmospheric and oceanic circulation and sea ice and is therefore likely also experiencing moisture source variability on different timescales.

In **Paper II**, we explore lake water isotopic variability over three years. The data show that lake water $\delta^2\text{H}$ can display variable sensitivity to changes in precipitation $\delta^2\text{H}$ and evaporation through time. Proxy records of hydrological variation provide information about a combination of processes but are, for a given record, often interpreted to reflect changes in only one variable such as moisture balance, moisture source change, or precipitation seasonality. By monitoring $\delta^2\text{H}_{\text{lake}}$ changes from year to year and comparing to regional conditions, we can assess the main drivers of modern isotopic variability and target lakes that are sensitive to changes in the process we want to reconstruct. In addition to the samples from 2018–2020 included in **Paper II**, we have repeated the lake water sampling in the first week of July in 2021 and 2022, to continue monitoring the isotopic range and environmental factors controlling lake water isotopes in northern Fennoscandia. Similar efforts on Svalbard would be beneficial to find suitable lakes for future paleo-precipitation studies.

4.3.2. Modern leaf wax isotope monitoring

In addition to the processes determining $\delta^2\text{H}_{\text{precip}}$ and $\delta^2\text{H}_{\text{lake}}$, we need to consider the transformation from source water $\delta^2\text{H}$ to wax $\delta^2\text{H}$. In **Papers I & III**, we focus on qualitative assessment of past hydrological changes, by inferring relative changes in precipitation seasonality and changes in summer $\delta^2\text{H}_{\text{precip}}$. If we want to isolate changes in precipitation $\delta^2\text{H}$, we need to translate $\delta^2\text{H}_{\text{wax}}$ into $\delta^2\text{H}_{\text{precip}}$. As presented in section 1.3, $\delta^2\text{H}_{\text{wax}}$ is depleted by a fractionation factor (ϵ_{app}) relative to $\delta^2\text{H}_{\text{precip}}$ due to climatic and/or plant physiological drivers (Fig. 2; Sessions et al., 1999; Kahmen et al., 2013). To convert $\delta^2\text{H}_{\text{wax}}$ into source water $\delta^2\text{H}$, we need good constraints on ϵ_{app} . Few Arctic studies have investigated terrestrial plant ϵ_{app} (Wilkie et al., 2013; Daniels et al., 2017; Berke et al., 2019), and to our knowledge there are no quantifications of ϵ_{app} for aquatic bryophytes, which are abundant at some of our study sites. Furthermore, a global database of *n*-alkanoic acid ϵ_{app} measured in lake surface sediments, compiled by McFarlin et al. (2019), shows that the median *n*-alkanoic acid ϵ_{app} values are consistently around -100‰ to -110‰ , independent of chain length and source (lake or precipitation) water (Thomas et al., 2020). Given that ϵ_{app} is not well quantified in the Arctic, the uncertainties are large, and ϵ_{app} is relatively constant (at least considering the catchment average for a given chain length), we do not convert $\delta^2\text{H}_{\text{wax}}$ to $\delta^2\text{H}_{\text{precip}}$. More modern studies are needed to better constraints ϵ_{app} in the Arctic, especially for aquatic plants.

Modern studies monitoring wax transport and deposition could help us distinguish aquatic from terrestrial sources. By monitoring transport into and within the lake and separating the two sources, it may be possible to quantify their relative contribution to the sedimentary leaf waxes. Furthermore, their chain-length distribution and isotopic composition could be compared to the chain-length distribution and isotopic composition of modern lake and catchment vegetation to distinguish wax sources to the lake sediments, using a range of recently developed tools, e.g., the Stable Isotope Mixing Model (SIMM; Parnell & Inger, 2016). Additionally, these interpretations would benefit from detailed vegetation mapping to determine the most abundant modern plants.

4.3.3. Spatial and temporal resolution of paleo-precipitation records

A key to solid proxy interpretations is having sufficiently high temporal resolution. Ideally, we want to have enough data points to decipher the rate of change, in addition to the amplitude.

The leaf wax $\delta^2\text{H}$ records presented in **Papers I & III** have an average temporal resolution ranging from 220 to 450 years between subsamples for each lake record. This resolution allows interpretations of general long-term trends but reveals less about short-term variability and complicates comparison between records. Furthermore, each subsample integrates sediments deposited over multiple years. The sample resolution from Arctic lake sediments is commonly limited by low sedimentation rates and low organic content.

In an on-going project in collaboration with researchers from the University of Massachusetts Amherst and Bates College, we analyze leaf wax $\delta^2\text{H}$ from a laminated surface sediment core from Linnévatnet, western Spitsbergen. By high-resolution sampling of the sedimentary sequence covering the instrumental period, we aim to compare the leaf wax $\delta^2\text{H}$ record to precipitation measurements from Isfjord Radio, 3.5 km west of the lake. This new study will hopefully allow us to 1) generate a high-resolution Arctic leaf wax $\delta^2\text{H}$ record for the last *c.* 120 years, and 2) evaluate how well $\delta^2\text{H}_{\text{wax}}$ correlate to seasonal precipitation variability. A similar reconstruction was successfully performed using $\delta^2\text{H}_{\text{wax}}$ from varved proglacial lake sediments on Baffin Island (Thomas et al., 2012).

Since precipitation amounts and seasonality are highly variable in time and space, we need to increase the spatial distribution of long-term paleo-precipitation records. Ice-core records from ice sheets and ice caps, such as the Greenland Ice Sheet, provide information about annual precipitation amount (annual accumulation) (e.g., Dahl-Jensen et al., 1993). Annual precipitation amounts have also been reconstructed from pollen, with available Holocene records from northern Fennoscandia (e.g., Hammarlund et al., 2002; Bjune et al., 2004; Jones et al., 2004), northeast Russia (Melles et al., 2012), and Arctic Canada and Alaska (e.g., Viau & Gajewski, 2009), but few studies differentiate between summer and winter precipitation (Allen et al., 2007). Another popular proxy for investigating various aspects of the water cycle is $\delta^{18}\text{O}$ from diatom biogenic silica, lacustrine carbonates, lake sediment cellulose, and chironomids, studied in lakes in Canada (e.g., Edwards et al., 1996; Wolfe et al., 1996), Alaska (Anderson et al., 2001), Greenland (e.g., Anderson et al., 2004; Lasher et al., 2017), northern Fennoscandia (e.g., Hammarlund et al., 2002; Rosqvist et al., 2013), and Russia (Wolfe et al., 2000). These records span different time scales, and with different temporal resolution, in addition to coming from lakes influenced by different environmental conditions. Some studies interpret $\delta^{18}\text{O}$ variability in terms of moisture balance, whereas others focus on moisture source shifts caused by atmospheric circulation changes and/or seasonality. Precipitation seasonality

has been reconstructed from leaf wax $\delta^2\text{H}$ from lake sediments on Greenland (Thomas et al., 2016; Thomas et al., 2020) and Svalbard (**Papers I and III**), and from a north Norwegian peatland (Nichols et al., 2009). Although there is an increasing number of studies on precipitation seasonality, we still know little about the spatial variability, and need more and higher-resolution records.

4.3.4. Other tools to improve the leaf wax $\delta^2\text{H}$ interpretations

Another way to advance proxy-based paleoclimate interpretations is to use proxy system modeling (e.g., Evans et al., 2013; Dee et al., 2018; Morrill et al., 2019). Since water isotopes are impacted by a combination of factors, this approach could be an important key to constraining the sensitivity of a lake to e.g., changes in the seasonal distribution of precipitation. Proxy system models allow us to use our understanding of modern physical and biological processes affecting the proxy and use sensitivity testing to evaluate which factor, or combination of factors, that most likely caused the isotope variability in our record (Dee et al., 2018). We could for example assess the influence of climate parameters such as winter precipitation amount or relative humidity on aquatic plant leaf wax $\delta^2\text{H}$. To set up lake-specific models and optimize the model performance, we need detailed field measurements of modern lake and catchment parameters, as outlined in sections 4.3.1 and 4.3.2.

Leaf wax $\delta^2\text{H}$ interpretations benefit from being combined with other proxies from the same lake. Plant *sed*aDNA can constrain plant community changes and their possible impact on $\delta^2\text{H}_{\text{wax}}$, and bulk elemental and stable isotope measurements of carbon and nitrogen ($\delta^{13}\text{C}$, $\delta^{15}\text{N}$, and C:N) can be used to examine terrestrial and aquatic contributions to the sediments (Gorbey et al., 2021). Diatom and pollen data can aid $\delta^2\text{H}_{\text{wax}}$ interpretations by providing information about lake conditions and atmospheric temperature (Katrantsiotis et al., 2021). Useful proxies for lake water and air temperatures are alkenones (e.g., D'Andrea et al., 2012; Longo et al., 2020), branched glycerol dialkyl glycerol tetraethers (brGDGTs; e.g., Keisling; Thomas et al., 2018), and chironomids (e.g., Wooller et al., 2004; Arppe et al., 2017). A recent study combined leaf wax $\delta^2\text{H}$ with $\delta^{18}\text{O}$ from chironomids in the same sediment archive to reconstruct precipitation seasonality on western Greenland (Corcoran et al., 2021). However, direct comparison between $\delta^2\text{H}$ and $\delta^{18}\text{O}$ from different proxies requires good constraints on the apparent fractionation factor for each proxy to estimate lake water $\delta^2\text{H}$. Ultimately, multi-proxy

investigations allow a thorough examination of various parameters. The more we know about the lake system, the better our understanding of the leaf wax $\delta^2\text{H}$ proxy becomes.

5. Concluding remarks

The main goal of this PhD project was to contribute to a better understanding of Arctic water cycle changes using stable isotope and organic geochemical proxies to investigate seasonal precipitation variability in time and space. More specifically, the aims were to provide Holocene records of precipitation seasonality from Svalbard, and to examine spatial and temporal variability in modern lake water isotopes and their drivers in northern Fennoscandia.

We present Holocene precipitation reconstructions from four lakes across Svalbard. Our results demonstrate that Svalbard experienced higher proportion of winter precipitation relative to the annual total during times of decreased sea-ice extent and emphasize the role of variable regional ocean surface condition on the seasonal distribution of precipitation on Svalbard. The specific outcomes from the Svalbard leaf wax studies are summaries of **Papers I & III**. The modern lake water study from northern Fennoscandia focused on spatial and interannual variability in lake water $\delta^2\text{H}$, inferred inflow (precipitation) $\delta^2\text{H}$, and evaporative enrichment. The results showed that the interannual variability is mainly controlled by the inflow isotopic composition, and that lakes farther from the Atlantic Ocean are more susceptible to evaporation. Furthermore, variable atmospheric circulation patterns over the North Atlantic influence the relative proportion of winter and summer precipitation entering the lakes. Details on these findings are given in **Paper II**.

Overall, the results presented in this dissertation highlight the potential of organic geochemical and stable isotope proxies to decipher past changes in precipitation seasonality, as well as emphasize the need for careful consideration of lake- and catchment-specific factors influencing lake and soil water $\delta^2\text{H}$. Finally, this dissertation outlines potential future directions for how to advance the understanding and interpretations of organic geochemical and precipitation isotope proxies.

6. Future perspectives

Stable water isotope proxies are powerful tools to infer water cycle changes on different timescales. This study has demonstrated that interpretations require careful consideration, as proxy records provide indirect evidence of climate change and can additionally be affected by environmental processes. By improving our understanding of the processes influencing the isotopic composition of precipitation, lake water, soil water, and plants, we can quantify the water cycle changes more confidently. Preferably, this is achieved through collaborative efforts across disciplines. To forward our interpretations of past water cycle changes on Svalbard, future studies should strive to:

- Improve our understanding of seasonal precipitation variability across Svalbard in modern times, by monitoring precipitation amounts and isotopes in the central, northern, and eastern parts of the archipelago. This requires a massive logistical effort in remote locations but would contribute valuable constraints on spatial variability in Arctic precipitation seasonality. Precipitation isotopes are a bellwether of water cycle change, with larger signal-to-noise ratios than precipitation amount, and greater ability to distinguish changes in moisture source and atmospheric circulation. Thus, having long-term and spatially detailed information about precipitation isotope change will benefit understanding of Arctic system change.
- Investigate modern moisture source variability and how it affects the isotopic composition of precipitation falling on Svalbard. This could be achieved by precipitation backtracking modeling and water-tagging experiments, determining where precipitation originally evaporated.
- Establish regional calibration of the leaf wax $\delta^2\text{H}$ proxy for Svalbard by measuring leaf wax homologue distributions and $\delta^2\text{H}$ values of modern plants, leaf water, soil water, and surface sediments to constrain the apparent fractionation.
- Develop techniques to monitor wax transport to and within the lake, to separate and quantify terrestrial and aquatic wax contribution to the lake sediments. Furthermore, detailed mapping of modern catchment vegetation can help evaluating the contribution of different plant types to the sedimentary leaf wax record.

- Increase the temporal resolution in future lipid biomarker studies. This could help assess whether changes occurred momentarily (possibly reflecting catchment processes altering the seasonality reflected in the lake) or gradually. Furthermore, it would enable more accurate correlations between records.
- Enhance the spatial resolution of Arctic paleo-precipitation records. Precipitation varies substantially on a regional and local scale, and the spatial coverage of proxy data, especially concerning precipitation seasonality, is inadequate to make meaningful comparison between regions.
- Explore proxy system modeling. This might prove to be a key tool to assess precipitation seasonality and to constrain the importance of a combination of processes explaining changes recorded in lake water isotope proxies. This modeling approach uses modern climate variables to simulate lake hydrologic and isotopic balance. Thus, proxy system modeling can help us excel the interpretation from lake water isotope proxies beyond describing relative trends.
- Combine leaf wax $\delta^2\text{H}$ with other proxies. A multi-proxy approach enables reconstruct of multiple climate aspects from the same sedimentary record and provide additional support for interpretations. For example, *sedaDNA* provides evidence of vegetation shifts in response to changes in climate and give indications about the plant wax sources. Moreover, $\delta^{13}\text{C}$ can help constrain the provenance of the waxes, and chironomids, alkenones, and brGDGTs can be used to infer past temperature anomalies.

7. References

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Paper I



Holocene precipitation seasonality in northern Svalbard: Influence of sea ice and regional ocean surface conditions

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ABSTRACT

Arctic precipitation is predicted to increase in the coming century, due to a combination of enhanced northward atmospheric moisture transport and local surface evaporation from ice-free seas. However, large model uncertainties, limited long-term observations, and high spatiotemporal variability limit our understanding of these mechanisms, emphasizing the need for paleoclimate records of precipitation changes. Here we use lipid biomarkers in lake sediments to reconstruct precipitation seasonality in northern Spitsbergen, Svalbard. We measured the hydrogen isotopic ratios ($\delta^2\text{H}$) of *n*-alkanoic acids (C_{20} – C_{30}) from sedimentary leaf waxes in lake Austre Nevlingen, Spitsbergen. We interpret $\delta^2\text{H}$ values of mid-chain (C_{22}) and long-chain (C_{28}) *n*-alkanoic acids to represent $\delta^2\text{H}$ of lake and soil water, respectively. Austre Nevlingen lake water $\delta^2\text{H}$ reflects amount-weighted mean annual precipitation $\delta^2\text{H}$. In contrast, soil water is mostly recharged by summer rainfall, and therefore reflects $\delta^2\text{H}$ values of summer precipitation. Austre Nevlingen leaf wax $\delta^2\text{H}$ values are ^2H -depleted in the Early Holocene, suggesting high winter precipitation amounts. This coincides with high summer insolation, strong Atlantic water advection and reduced spring sea-ice cover in surrounding waters. Winter precipitation continued to dominate until c. 6 cal. kyr BP. After 6 cal. kyr BP, the trend in the biomarker record is not as clear. This could be related to colder conditions causing longer duration of seasonal lake-ice cover, thereby influencing the precipitation seasonality registered by the lake water. The Austre Nevlingen record suggests a close relationship between precipitation seasonality and regional ocean surface conditions, consistent with simulations suggesting that Arctic winter sea-ice loss will lead to increased local evaporation.

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1. Introduction

By the end of the 21st century, warming in the Arctic is predicted to exceed the global average by a factor of 2.2–2.4 (Collins et al., 2013). This amplified warming will affect the amount and seasonality of precipitation, through hydrological intensification (Rawlins et al., 2010; Collins et al., 2013; Bintanja and Selten, 2014). Increased

high-latitude precipitation can be caused by 1) atmospheric circulation changes and enhanced poleward moisture transport, mechanisms mainly influencing summer precipitation (Dickson et al., 2000; Zhang et al., 2013), and 2) increased local surface evaporation, due to warmer Arctic seas and reduced winter sea-ice cover, mainly influencing fall and winter precipitation (Bintanja and Selten, 2014; Kopec et al., 2016). A warmer atmosphere also causes a larger fraction of Arctic precipitation to fall as rain (Førland et al., 2020). These hydrological changes will impact Arctic ecosystems, glacier mass balance, and infrastructure (Bintanja and Andry, 2017; Adakudlu et al., 2019). In order to better assess

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future Arctic precipitation changes, we need improved understanding of the mechanisms causing precipitation variation in the past (Sundqvist et al., 2014; Linderholm et al., 2018). In the absence of instrumental data, paleoclimate (proxy) data are necessary to fill this critical knowledge gap.

Previous reconstructions of the Holocene climate history of Svalbard have mainly focused on temperature. During the beginning of the Holocene (11.7–9 cal. kyr BP), Svalbard experienced warmer-than-present conditions, recorded by alkenones (van der Bilt et al., 2019) and thermophilous mollusks (Mangerud and Svendsen, 2018). Furthermore, glaciers were smaller (Mangerud and Svendsen, 1990; Svendsen and Mangerud, 1997; Røthe et al., 2018) and the sea-ice cover in surrounding waters reduced (Werner et al., 2016; Allaart et al., 2020). Other studies have described Early Holocene re-advances for glaciers on Svalbard (Lønne, 2005; Henriksen et al., 2014; van der Bilt et al., 2015; Farnsworth et al., 2017, 2018; Larsen et al., 2018). There are speculations if enhanced Early Holocene precipitation drove these glacier oscillations (Farnsworth, 2018). The Early Holocene warming was abrupt and could be explained by influx of warm Atlantic water (AW; Risebrobakken et al., 2011; Werner et al., 2016), peak summer insolation (Laskar et al., 2004) or a combination of both. According to Werner et al. (2016), eastern Fram Strait subsurface temperatures remained warm during the Mid Holocene until c. 5 cal. kyr BP, with increasing sea-ice cover from c. 7 cal. kyr BP. In central Spitsbergen, a stepwise cooling began already around 9 cal. kyr BP (Rasmussen et al., 2012). The Late Holocene was characterized by Neoglacial cooling from c. 4–3 cal. kyr BP, inferred from increased glacier activity (Svendsen and Mangerud, 1997; Røthe et al., 2015, 2018; Miller et al., 2017; Bartels et al., 2018; Lovell et al., 2018). A later cooling step, associated with the Little Ice Age, started around 0.7 cal. kyr BP (Werner, 1993; van der Bilt et al., 2015; Miller et al., 2017).

Robust records of Holocene precipitation amount and seasonality are lacking. Leaf wax hydrogen isotope ratios ($\delta^2\text{H}$) provide means to evaluate hydrological changes over long time scales. The potential of the method for high-latitude lacustrine records has been demonstrated in Arctic Canada (Thomas et al., 2012), Greenland (Balascio et al., 2013; Thomas et al., 2016, 2018), Svalbard (Balascio et al., 2018) and northeastern Russia (Wilkie, 2012).

Leaf waxes are straight-chain hydrocarbon compounds produced by both terrestrial and aquatic plants. The *n*-alkanoic acid and *n*-alkane components of leaf waxes are well-preserved in sedimentary records due to their resistance to degradation (Eglinton and Calvin, 1967). Terrestrial plants predominantly produce long-chain wax compounds, whereas aquatic plants produce mid-chain wax compounds (Ficken et al., 2000; Meyers, 2003; Nichols et al., 2009; Gao et al., 2011). The $\delta^2\text{H}$ values of leaf waxes reflect the $\delta^2\text{H}$ values of the plant source water, with a biosynthetic fractionation that is largely constant for specific compounds (Sachse et al., 2012; McFarlin et al., 2019). The source water for terrestrial plants is soil water, which is mostly recharged by summer rainfall (Cooper et al., 1991; Throckmorton et al., 2016). The hydrogen isotope ratios of terrestrial leaf waxes ($\delta^2\text{H}_{\text{terr}}$) therefore reflects summer precipitation isotope values, influenced by some evaporative enrichment of soil and leaf water (Kahmen et al., 2013). In contrast, the source water for aquatic plants is lake water, which may reflect summer or mean annual precipitation $\delta^2\text{H}$ values, depending on the residence time and source of lake water (Cluett and Thomas, 2020; Thomas et al., 2020). Hence, leaf wax $\delta^2\text{H}$ can be used to reconstruct source water $\delta^2\text{H}$ and ultimately $\delta^2\text{H}$ of precipitation and other aspects of climate, including evaporation (Rach et al., 2017). The $\delta^2\text{H}$ of precipitation is influenced by changes in local and source temperature, moisture source location and transport history (Frankenberg et al., 2009), as well as evaporation

for terrestrial waxes (e.g., Sachse et al., 2012; Thomas et al., 2016, 2018).

The hydrogen isotope ratios of aquatic plant leaf waxes ($\delta^2\text{H}_{\text{aq}}$) reflect the isotopic composition of the lake water during the growing season. Therefore, the precipitation isotope seasonality recorded by $\delta^2\text{H}_{\text{aq}}$ (annual mean or summer) depends on the local conditions determining the residence time of the lake water (Thomas et al., 2020). Lakes with short residence times have growing season lake water $\delta^2\text{H}$ values biased towards summer precipitation $\delta^2\text{H}$, whereas water isotopes from lakes with long residence times reflect amount-weighted mean annual precipitation $\delta^2\text{H}$ (Jonsson et al., 2009; Cluett and Thomas, 2020). Balascio et al. (2018) inferred that $\delta^2\text{H}_{\text{aq}}$ changes in lake Hakluytvatnet (Fig. 1), which has a short residence time (i.e., the lake water during the growing season reflects summer precipitation $\delta^2\text{H}$), reflect air temperature and the relative influence of polar and sub-polar air masses. Here, we apply the same concept for another lake in northern Svalbard. By choosing a lake with a longer residence time (i.e., lake water isotopes representing mean annual precipitation $\delta^2\text{H}$), we can extract information about seasonal variation in Arctic precipitation. We use $\delta^2\text{H}$ of C_{22} and C_{28} *n*-alkanoic acids to infer past lake water and leaf water $\delta^2\text{H}$ values, and reconstruct Holocene precipitation seasonality in northern Spitsbergen. By better constraining precipitation seasonality during earlier Holocene warm periods, we can improve our understanding of the mechanisms causing precipitation change in the past, present and in the future.

2. Setting

2.1. Regional setting

The Svalbard archipelago is influenced by strong climate gradients, with air masses transporting heat and moisture from the Atlantic regions meeting cold polar air from the Arctic Basin (Fig. 1a; Førland et al., 2009, 2011; Vikhamar-Schuler et al., 2019). South of Svalbard, the North Atlantic Current splits into the West Spitsbergen Current (WSC) and the North Cape Current (NCaC), transporting warm saline water along the west coast of Svalbard and eastward into the Barents Sea (Fig. 1a). The Barents Sea also receives colder, fresher water masses from the Arctic Ocean transported by the East Spitsbergen Current (ESC). During extreme cold events over the Barents Sea in winter, the strong temperature gradient between cold Arctic air and relatively warm Barents Sea surface water can cause convection and development of low-pressure systems (Rasmussen, 1985). Easterly winds caused by these systems bring moisture to eastern Svalbard, resulting in high precipitation (Hagen et al., 1993).

Since the establishment of the first permanent meteorological stations in Svalbard in the 1910s, temperatures have increased during all seasons, most notably in winter and spring (Førland et al., 2011). The trend in precipitation amount for the same period is not as clear. The robustness of instrumental precipitation data on Svalbard is hindered by 1) a lack of stations - especially in the central and eastern parts of the archipelago, and 2) measurement errors - notably due to snowdrift and undercatch (Førland and Hanssen-Bauer, 2000; Hanssen-Bauer, 2002; Humlum, 2002). However, instrumental records generally display positive trends in autumn and winter precipitation, and negative trends in summer and spring (Adakudlu et al., 2019). On an annual basis, all long precipitation amount time series from Svalbard (i.e., >40 yr) show positive trends, with precipitation increases between 2–8% per decade (Førland et al., 2011). Climate model projections for Svalbard agree that precipitation amount will increase in the future as temperature rises and sea ice retreats. Førland et al. (2011) projected an annual precipitation increase from 1961–1990 to

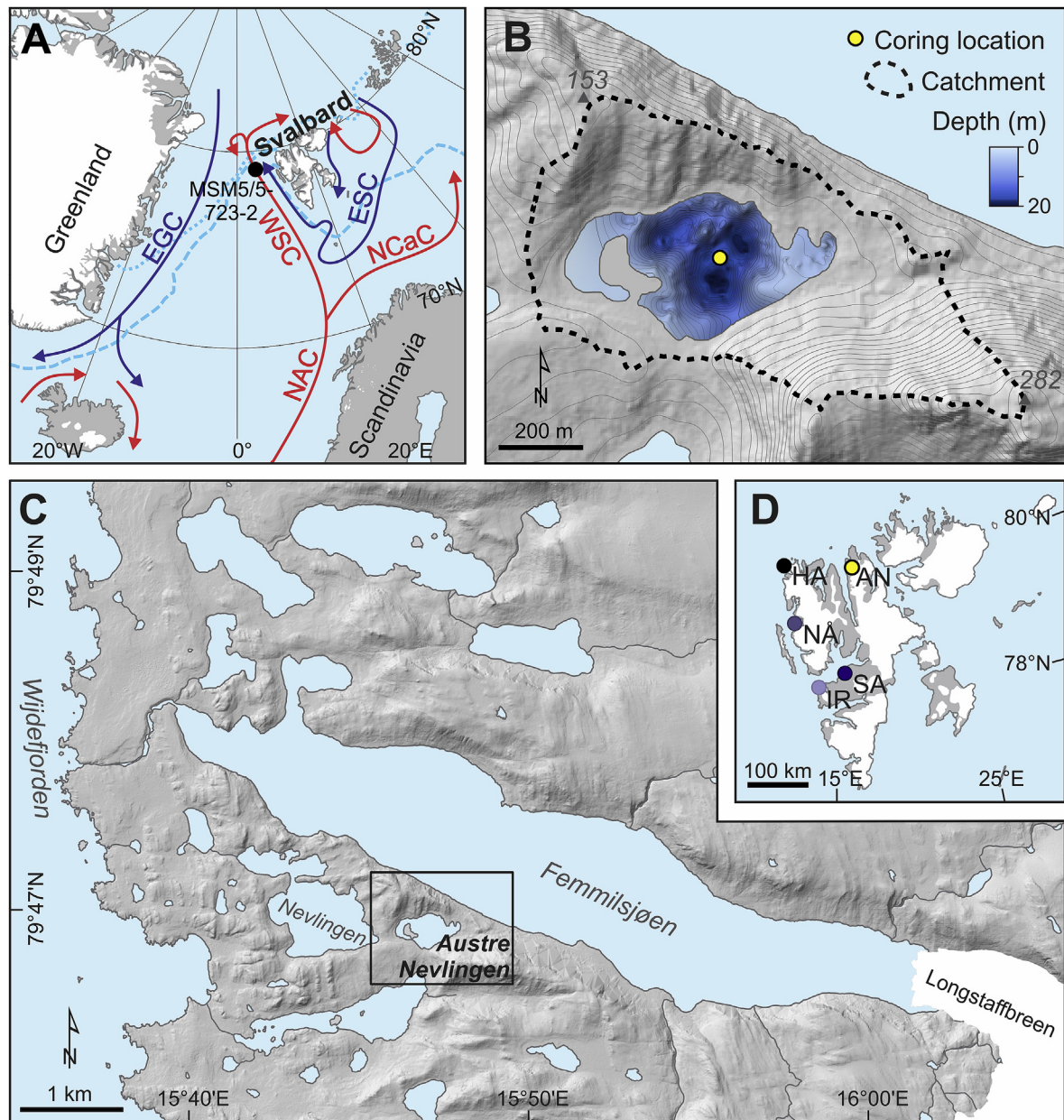


Fig. 1. (A) Map showing the location of Svalbard in the North Atlantic, with major ocean surface currents (warm currents in red, cold in dark blue; NAC = North Atlantic Current; NCaC = North Cape Current; WSC = West Spitsbergen Current; ESC = East Spitsbergen Current; EGC = East Greenland Current), median winter (light blue dotted line) and summer (light blue dashed line) sea-ice extent AD 1981–2010 (National Snow and Ice Data Center, 2019) and location of marine core MSM5/5-723-2 (Werner et al., 2016). (B) Lake Austre Nevlingen catchment, coring location and bathymetry. (C) Map of the study area in northern Spitsbergen. Black box indicates the Austre Nevlingen catchment, shown in (B). (D) Map of Svalbard, showing the locations of Austre Nevlingen (AN), Hakluytvatnet (HA) and meteorological stations (IR = Isfjord Radio; NÅ = Ny-Ålesund; SA = Svalbard Airport). (For interpretation of the references to color in this figure legend, the reader is referred to the Web version of this article.)

2071–2100 of less than 10% in southwestern Spitsbergen, but more than 40% in northeastern Svalbard. This regional climate model simulation was based on the MPIB2 scenario (Max Planck Institute global ECHAM4 model with SRES B2 emission scenario). Based on the RCP8.5 scenario (“business as usual”; Stocker et al., 2013), Adakudlu et al. (2019) projected a 65% annual precipitation increase in Spitsbergen from 1971–2000 to 2071–2100. In the same study, RCP4.5 (reductions after 2040, “medium emissions”; Stocker et al., 2013) gives a median annual precipitation increase of 45%. Both model projections indicate largest precipitation increases in northeastern Svalbard, with greatest change in winter (+54–90%) and least in summer (+27–48%; Adakudlu et al., 2019). The fact that

all models agree that the greatest precipitation increases will occur in the northeast, highlights the precipitation sensitivity of our study area. Model uncertainties and high spatial and temporal variability in precipitation emphasize the need for paleoclimate records to elucidate the mechanisms causing precipitation variability on Svalbard.

2.2. Study area

Lake Austre Nevlingen is located 2 km east of the 108-km-long Wijdefjorden, close to the mouth of the fjord (Fig. 1). Wijdefjorden is associated with the Billefjorden Fault Zone, separating Devonian

sandstones and clastic sedimentary rocks in the west from Precambrian crystalline bedrock in the east (Dallmann, 2015). The landscape around Austre Nevlingen is characterized by undulating terrain with lakes in depressions (Fig. 1b and c). The timing of deglaciation of the study area remains poorly constrained. Cosmogenic exposure ages from Tyrkampen (450 m above sea level (a.s.l.), 25 km to the south), suggest that this highland area was ice free by 18.3 ± 1.2 to 17.9 ± 1.2 kyr ago (Hormes et al., 2013).

The closest meteorological stations to our study site are located in Ny-Ålesund (8 m a.s.l., 125 km SW of Austre Nevlingen), at Svalbard Airport (28 m a.s.l., 172 km S of Austre Nevlingen) and Isfjord Radio (7 m a.s.l., 198 km SSW of Austre Nevlingen; Fig. 1d). Ny-Ålesund mean annual air temperature (1990–2019, $n = 30$) is -4.2 °C and mean annual precipitation (1990–2019, $n = 30$) 477 mm, whereas corresponding values are -3.9 °C ($n = 30$) and 203 mm ($n = 30$) at Svalbard Airport, and -2.6 °C ($n = 11$) and 573 mm ($n = 3$) at Isfjord Radio (Fig. 2; MET Norway, 2020). Hagen et al. (1993) estimated annual precipitation in the Wijdefjorden

area to be as low as 200 mm, based on available measurements from meteorological stations and glacier mass balance and equilibrium line altitude observations. Precipitation $\delta^2\text{H}$ in Ny-Ålesund and Isfjord Radio exhibits no significant seasonal trend (Fig. 2b; IAEA/WMO, 2019), with values close to the Global Meteoric Water Line (GMWL, $\delta^2\text{H} = 8 \times \delta^{18}\text{O} + 10$; Craig, 1961, Fig. 3). There are no measurements of precipitation isotopes in central Spitsbergen, but we would expect greater precipitation isotope seasonality at our study site. Ny-Ålesund and Isfjord Radio are close to open sea conditions all year round, whereas Wijdefjorden is less connected to the main moisture source pathway along the west coast, especially in winter (Fig. 1a).

The inner (southern) and middle parts of Wijdefjorden are classified as high-Arctic steppe, and have unique and high botanically diverse flora for Svalbard (Elvebakk and Nilsen, 2002; Elvebakk, 2005; Eidesen et al., 2018; Voldstad et al., 2020). The outer (northern) part is less vegetated and belongs to the Northern Arctic tundra zone (Norwegian Polar Institute, 2019). The vegetation in the north is dominated by graminoids (e.g., *Luzula confusa*, *Poa arctica*, *P. alpina*). Forbs are abundant (e.g., *Saxifraga oppositifolia*, *Cerastium arcticum*, *Oxyria digyna*, *Dryas octopetala*), mosses (e.g., *Tomentypnum nitens*, *Polytrichum*, *Racomitrium*) and lichens (e.g., *Flavocetraria*, *Stereocaulon*) are common, and dwarf shrubs (e.g., *Salix polaris*, *Cassiope tetragona*) are present (Walker et al., 2005). There are almost no aquatic vascular plants on Svalbard, but submerged bryophytes are common in lakes (Balascio et al., 2018; Voldstad et al., 2020).

2.3. Lake catchment

Austre Nevlingen (79°47'N, 15°47'E; Fig. 1) has a surface area of 0.13 km², a maximum depth of 18 m, and sits at an elevation of 41 m a.s.l. This is below the local marine limit, which can be estimated to c. 60 m a.s.l. based on reconstructed isobases of uplift (Forman, 1990). The Austre Nevlingen catchment is c. 0.48 km², sitting between 153 and 282 m a.s.l. summits (Fig. 1b). The catchment is dominated by frost-shattered bedrock and boulders with almost no soil cover, and dry non-vegetated to sparsely vegetated slopes. During the August 2015 and August 2018 field campaigns, we observed terrestrial vegetation dominated by graminoids (e.g., *Luzula confusa*) and mosses (e.g., *Polytrichum*). Submerged bryophytes, but no other aquatic plants, were observed growing in the

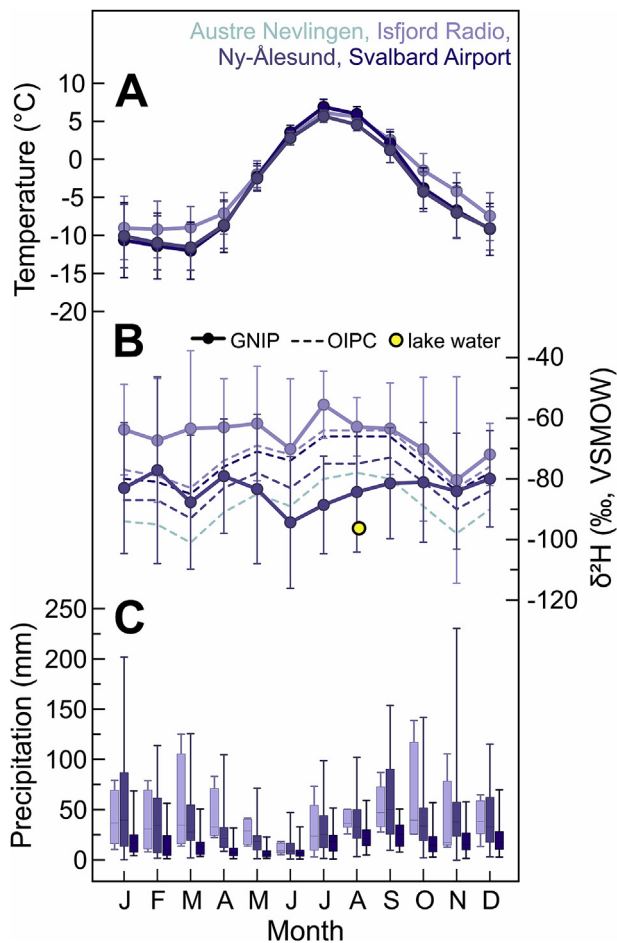


Fig. 2. Climate data from Svalbard Airport, Ny-Ålesund, Isfjord Radio and Austre Nevlingen. For locations, see Fig. 1. (A) Monthly average temperature calculated for all available data from the meteorological stations at Svalbard Airport, in Ny-Ålesund and Isfjord Radio for the period AD 1990–2019 (MET Norway, 2020), with 1σ standard deviations (vertical bars). (B) Global Network of Isotopes in Precipitation (GNIP; IAEA/WMO, 2019) $\delta^2\text{H}$ values from Isfjord Radio (AD 1960–1976) and Ny-Ålesund (AD 1990–2016), precipitation $\delta^2\text{H}$ for all sites (calculated using the Online Isotopes in Precipitation Calculator (OIPC); Bowen et al., 2005; Bowen, 2019; IAEA/WMO, 2019) and lake water $\delta^2\text{H}$ from Austre Nevlingen (2018). (C) Monthly average precipitation calculated for all available data from the meteorological stations Svalbard Airport, Ny-Ålesund and Isfjord Radio, for the period AD 1990–2019 (MET Norway, 2020). For each box plot, the middle line displays the median precipitation, the box represents the 25% to 75% quartile range and whiskers the maximum and minimum values.

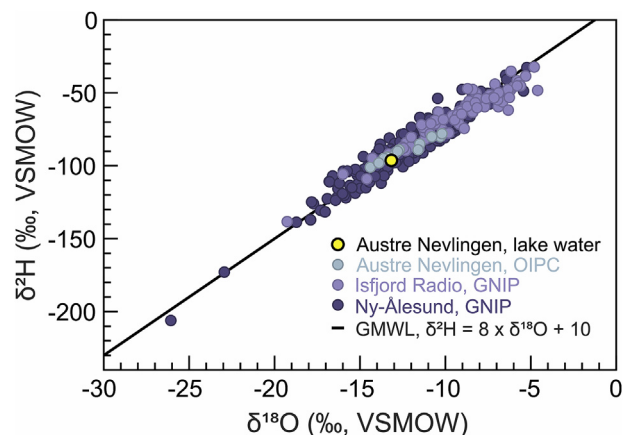


Fig. 3. Lake water isotope data from lake Austre Nevlingen (August 2018), precipitation isotope data from the GNIP stations (IAEA/WMO, 2019) at Isfjord Radio and in Ny-Ålesund, and calculated monthly precipitation $\delta^2\text{H}$ for Austre Nevlingen (OIPC; Bowen et al., 2005; Bowen, 2019; IAEA/WMO, 2019). Black line indicates the Global Meteoric Water Line (GMWL). For locations, see Fig. 1.

lake. The growing season is from mid-June to early September (ORNL DAAC, 2018) and the lake is estimated to be ice-covered from October to the beginning of July (Holm et al., 2012).

The lake is fed by surface runoff, but has no distinct inflow stream. There is also no distinct outflow stream on the surface. To estimate the proportion of lake water replaced annually and seasonally, we multiply the precipitation amount at Svalbard Airport during the different seasons (spring melt season = October to June, ice-free season = July to September) with the catchment area, and divide the product by the lake volume (Jonsson et al., 2009; Thomas et al., 2020). Svalbard Airport (Fig. 1d) has the most similar precipitation amount and seasonality to our study site. On average, approximately 11% of the lake volume is replaced each year: 7% during the spring melt season (i.e., by runoff representing precipitation from months with lake-ice cover) and 4% during the ice-free season (i.e., by runoff during the ice-free months). These values indicate a relatively long residence time, and that the lake water isotopes could be influenced by surface evaporation. Yet, isotope values of lake surface water collected in August 2018 indicate minimal evaporative enrichment. This surface water has a $\delta^2\text{H}$ value of $-96.3 \pm 0.2\text{‰}$ and $\delta^{18}\text{O}$ of $-13.2 \pm 0.02\text{‰}$ (Figs 2b and 3). The deuterium excess ($d\text{-excess} = \delta^2\text{H} - 8 \times \delta^{18}\text{O}$; Dansgaard, 1964) is 9.0‰. These values are well within the range of precipitation $d\text{-excess}$ at Ny-Ålesund and Isfjord Radio (IAEA/WMO, 2019) and close to the GMWL ($d\text{-excess} = 10\text{‰}$). The lack of isotopic evidence for evaporative enrichment suggests that the lake water may be completely flushed each year (Cluett and Thomas, 2020), perhaps via subsurface runoff into the lake and subsurface drainage out of the lake through the frost-shattered bedrock. Lichen-free boulders up to 0.5 m above the current lake level indicate that the lake may have experienced higher water levels in the past, which could indicate subsurface drainage during periods of thawed active layer.

3. Methods

3.1. Bathymetry and sediment core collection

The bathymetry of Austre Nevlingen was surveyed in August 2018, with a Garmin ECHOMAP™ Plus 73SV with a CV52HW-TM transducer and a 5 Hz receiver, using the Quick Draw contour function. Coring in 2015 was guided in real-time with depth sounding data from a Hondex PS-7 Transducer. To minimize disturbance, sediment cores were obtained from the central, deepest part of the basin (Fig. 1b). Coring was performed through a hole in the floor of a small zodiac, which was anchored in a stable position on the lake surface. A 70-cm-long surface core (ANS1) was collected with a Universal surface corer (120 cm long and 68 mm diameter coring tubes). A 129-cm-long piston core (ANP3) was retrieved using a hand-held lightweight piston corer (200 cm long and 60 mm diameter coring tubes). To record modern conditions in the catchment, lake water and leaves from a selection of the most common plants growing in the area were collected in August 2018. Leaves were sampled from multiple plants of the same species to get representative samples, and chosen to represent different plant groups, including rushes (*Luzula confusa*), dwarf shrubs (*Salix polaris*), terrestrial mosses (*Polytrichum* sp.) and aquatic mosses (unidentified).

3.2. Lithology and stratigraphy

The lake sediment cores were split open, logged and analyzed in the sediment lab and ITRAX core facility at the University of Copenhagen. ITRAX scanning was performed to record visual and radiographic imagery and variations in the element-geochemical properties throughout the cores. A rhodium (Rh) tube with a

1 mm resolution and 30 s exposure time was used for X-ray fluorescence (XRF; e.g., Kylander et al., 2011) and 4 mm resolution for magnetic susceptibility (MS; e.g., Sandgren and Snowball, 2002). Information on the ITRAX core scanner is given by Croudace et al. (2006). We present the Ti signal and the Ca/Fe ratio to show variation in minerogenic input, as these can be used to detect glacial meltwater input (Kylander et al., 2011) and lake basin isolation (Larsen et al., 2017), respectively. The Ti peak area was normalized against the incoherent (inc) and coherent (coh) Rh scatter (Ti/(inc + coh)) to remove instrumental effects, as suggested by Kylander et al. (2011).

After scanning, the lithology and stratigraphy of the cores were visually inspected and logged. In order to determine whether the lowermost minerogenic unit was deposited in a marine or lacustrine environment, we searched for marine microfossils (e.g., foraminifera) using a stereo microscope. The total organic content was determined through loss-on-ignition (LOI; Heiri et al., 2001). For this purpose, samples (1 cm³) were collected every 2 cm, dried at 110 °C for 24 h and ignited at 550 °C for 4 h.

3.3. Lipid biomarker extraction and analysis

Biogeochemical and isotopic analyses were performed in the Organic and Stable Isotope Biogeochemistry Laboratory at the University at Buffalo, NY, USA. Samples (3–4 cm³) for lipid biomarker analysis were collected every 5–8 cm from ANS1 and every 4 cm from the organic part (i.e., the uppermost 97 cm) of ANP3, resulting in thirty-six samples for the composite record. The samples were freeze-dried and homogenized, and free lipids were extracted with an Accelerated Solvent Extractor (ASE) 200 (Dionex) using dichloromethane (DCM):methanol 9:1 (v:v). An internal standard (C_{20:1} *n*-alkanoic acid, Fisher Scientific, 4.2 µg) was added to the total lipid extract (TLE) after ASE extraction, allowing biomarker quantification. Preparation for *n*-alkanoic acid analysis followed previously published methods (Thomas et al., 2012). Hydrocarbons were separated by flash-column chromatography using NH₂-functionalized silica gel, DCM:isopropanol 2:1 (v:v) as the neutral eluent and 4% acetic acid in DCM as the acid eluent. The acid fraction was methylated at 60 °C overnight using 5% anhydrous HCl in methanol of known isotopic composition, and purified using silica gel columns, removing hydroxyl-carboxylic acid esters with hexane and recovering the fatty acid methyl esters (FAMES) in DCM.

The FAMES were quantified using a Thermo Scientific Trace 1310 gas chromatograph (GC) equipped with two AI1310 autosamplers, two split/splitless injectors, and two flame ionization detectors (FIDs) operated in parallel for higher throughput. For all analyses, the inlets were held at 250 °C and operated in splitless mode for the first 45 s, after which split flow was turned on at 14 mL/min. Hydrogen carrier gas was used with a constant flow rate of 3.6 mL/min. The oven program was as follows: initial temperature of 70 °C held for 1 min, then a fast ramp of 27 °C/min to 230 °C, followed by a slower ramp of 6 °C/min to 315 °C, with a final hold at 315 °C for 10 min. Compound masses were calculated via external calibration curves determined separately for each detector using a C₂₈ FAME standard (Fisher Scientific), and those masses were normalized to the mass of extracted sediment as well as the recovery of the internal standard to determine the concentration of waxes in sediment.

The $\delta^2\text{H}$ values of the FAMES were measured using a Thermo Scientific Delta V Plus isotope ratio mass spectrometer (IRMS) coupled to a Trace 1310 GC via an Isolink II and Conflo IV. GC inlet flow settings and oven temperature program were identical to those used in GC-FID analysis except for the carrier gas, which was helium at a constant flow rate of 1.5 mL/min. The HTC reactor was held at 1420 °C for all analyses. The H₃ factor was monitored

regularly at the beginning of every sequence. The ANP3 samples were analyzed in November 2017, with an average H_3 factor of $2.8 \pm 0.01\%$, and the ANS1 samples in January 2019, with an H_3 factor of 3.6% . FAME standards of known isotopic composition (C_{18} , C_{20} , C_{24} , Arndt Schimmelman, University of Indiana) were run along with every sequence of samples to correct for drift and the dependence of measured δ^2H on peak size, and to normalize sample results to the VSMOW (Vienna Standard Mean Ocean Water) scale. FAME δ^2H values were corrected for the isotopic composition of the hydrogens added during methylation. All isotope values are expressed in per mil (‰) relative to VSMOW, using standard δ notation:

$$\delta^2H(\text{‰}) = \left(\frac{R_{\text{sample}}}{R_{\text{VSMOW}}} - 1 \right) \times 1000$$

where R is the ratio between deuterium and hydrogen, $^2H/^1H$. Total uncertainty of measured δ^2H values, calculated as the Standard Error of the Mean (SEM; which equals the root sum of squares of uncertainty for the drift and peak size corrections, replicate uncertainty of sample measurements, and uncertainty in the δ^2H value of the methylation hydrogens, divided by the root of the number of measurements), was on average better than $\pm 2.7\%$. Average chain length (ACL) was determined for C_{22} to C_{30} , using the equation:

$$ACL_{22-30} = \frac{\sum (n \times C_n)}{\sum (C_n)}$$

where C_n is $\mu\text{g/g}$ sediment of each n -alkanoic acid with n carbon atoms.

3.4. Chronology and core correlation

The chronology of the sediment cores was established based on accelerator mass spectrometry (AMS) ^{14}C measurements on four terrestrial and eight aquatic plant macrofossils. The macrofossils were identified and isolated from 0.5 mm sieving residues, and measured at the Ångström Laboratory, Uppsala University and Lund University Radiocarbon Dating Laboratory, Sweden. All radiocarbon ages were calibrated using the online OxCal software (v. 4.3; Bronk Ramsey, 2009) and the IntCal13 dataset (Reimer et al., 2013). Calibrated ages are presented in calibrated year before present (cal. yr BP; BP = 1950).

For stratigraphic correlation between the cores, we used visual trends in the XRF data together with the radiocarbon age constraints. The cores were aligned in AnalySeries (v. 2.0.8; Paillard et al., 1996) using tie points in the elemental data. The Ca/Fe ratio was the main signal used (Fig. A, Appendix A), but multiple elemental ratios were compared to construct a common stratigraphic depth scale.

To establish an age-depth relationship for the composite core and incorporate age uncertainty into our biomarker records, we entered all proxy and chronology data into Linked PaleoData (LiPD) files (McKay and Emile-Geay, 2016). The data were analyzed in R (v. 3.6.3; R Core Team, 2020) using the GeoChronR package (McKay et al., 2018) and the IntCal13 dataset (Reimer et al., 2013). An age-depth model was constructed using a prior mean accumulation rate (acc. mean) of 100 years/cm and upper (d.min) and lower (d.max) depths set to the upper- and lowermost depths for the biomarker data (Fig. 4). The age ensemble was mapped to the paleo data, allowing us to display the biomarker data with age uncertainty (Figs 6 and 8) and plot the δ^2H records for both cores together (Fig. 8b). Fine lines in Figs 6 and 8 show the raw data plotted on the median of each age point, bold lines represent median values of the

age model ensembles, and the light and dark shading show the 1σ and 2σ age uncertainty, respectively.

4. Results

4.1. Chronology and core correlation

Radiocarbon ages, calibrated median ages and 2σ age intervals for ANS1 and ANP3 are presented in Table 1 and Figs 4 and 5. The uppermost age in ANP3 (LuS 12222) appeared to be an outlier compared to the rest of the age-depth model, which was otherwise fairly linear, despite different terrestrial and aquatic sources of the samples. We therefore excluded this sample when running the final age-depth model (Fig. 4). Our biomarker record spans 11.5–2.2 cal. kyr BP for ANP3 and 9.2–1.9 cal. kyr BP for ANS1.

4.2. Lithology and stratigraphy

Figure 5 shows optical and X-ray imagery, sedimentological logs, LOI and selected XRF data for the two cores. The cores contain three lithological units.

Unit L1 is only present in ANP3 (129–97 cm; Fig. 5), and consists of light grey, clayey-silty diamict with interlaminated organic material. The LOI is below 2%, and $\text{Ti}/(\text{inc} + \text{coh})$ is higher (0.20–0.60) than in the other units, where values are generally below 0.04. Also, Ca/Fe is higher than in the overlying unit, especially in a coarser grained interval between c. 112–103 cm depth. L1 could be interpreted to represent minerogenic-rich sedimentation, driven by inflow of glacial meltwater, or deposited in a shallow glaciomarine setting. No foraminifera were observed in this unit. The transition from unit L1 to L2 at 97 cm depth, with decreasing minerogenic fines (silt and clay) in the lower 15 cm, indicates either an abrupt termination of glacial meltwater inflow or isolation from the fjord. Both these interpretations are supported by a sharp decrease in $\text{Ti}/(\text{inc} + \text{coh})$, and gradually decreasing Ca/Fe ratio.

Unit L2 (c. 97–34 cm in ANP3, 70–33 cm in ANS1; Fig. 5) consists of light brown to light olive grey stratified gyttja, with sporadic interbedded aquatic bryophytes and high organic content, representing accumulation of organic material with minimal minerogenic input. The organic content is higher in ANP3, increasing gradually from 13% at the lower boundary to 30% near 89 cm depth, and remaining between 30–40% for the rest of the unit. In ANS1, the same unit has LOI values between 17–27%. In both cores, $\text{Ti}/(\text{inc} + \text{coh})$ is generally below 0.03, and Ca/Fe ratio low and stable around 0.015–0.030.

Unit L3 (34–0 cm in ANP3, 33–0 cm in ANS1; Fig. 5) consists of dark brown gyttja, with a gradational lower boundary, and organic-rich beds interbedded with abundant aquatic bryophytes. LOI continues to gradually increase to 37% in ANS1 and just below 50% in ANP3, in the uppermost part of the cores. Ca/Fe ratios are higher and more variable (0.025–0.155), whereas $\text{Ti}/(\text{inc} + \text{coh})$ ratios show values similar to L2, although decreasing in ANS1.

4.3. *n*-alkanoic acid sources and productivity

Large isotopic differences between mid-chain (C_{20} , C_{22} and C_{24}) and long-chain (C_{26} , C_{28} and C_{30}) *n*-alkanoic acids, but similar values within these groups, suggests different sources of mid- and long-chain *n*-alkanoic acids (Fig. 6). This supports the hypothesis that Austre Nevlingen records both summer (terrestrial) and mean annual (aquatic) precipitation isotopes, and that hydrogen isotope ratios of C_{22} and C_{28} leaf waxes can be used as indicators of different plant sources (Gao et al., 2011; Thomas et al., 2016, in press) and to reflect different seasonality.

The analysis of modern plants suggests that different leaf wax

Table 1Radiocarbon ages from sediment cores ANS1 and ANP3 from lake Austre Nevlingen, northern Spitsbergen, Svalbard. Calibrated ages are median ages within the 2 σ age ranges.

Core	Depth (cm)	Lab ID ^a	¹⁴ C-age (yr BP)	Calibrated age (cal. yr BP)	Calibrated 2 σ age ranges (cal. yr BP)	$\delta^{13}\text{C}$ (‰ VPDB)	Dated material
ANS1	4–5	Ua-60808	2200 \pm 33	2232	2318–2131	–25.8	Aquatic moss
ANS1	14–15	Ua-60809	3350 \pm 46	3589	3695–3469	–26.6	Aquatic moss
ANS1	24–25	Ua-60810	4818 \pm 62	5532	5661–5447, 5385–5327	–	Aquatic moss
ANS1	41–42	Ua-60811	6209 \pm 56	7104	7256–6973	–26.0	Aquatic moss
ANS1	55–56	Ua-60812	7410 \pm 52	8250	8370–8156, 8116–8113, 8090–8056	–29.0	<i>Salix polaris</i>
ANP3	9.5–10.5	LuS 12222	4135 \pm 45	4677	4824–4530	–	Aquatic moss
ANP3	25.5–26.5	LuS 12223	4215 \pm 40	4740	4856–4785, 4765–4620	–	Aquatic moss
ANP3	32.5–33.5	LuS 12224	4545 \pm 45	5166	5435–5423, 5320–5046	–	Wood
ANP3	50.5–51.5	LuS 12225	6085 \pm 50	6953	7157–7039, 7033–6831, 6819–6798	–	<i>Salix polaris</i>
ANP3	69.5–70.5	LuS 12226	7850 \pm 45	8636	8952–8919, 8862–8832, 8780–8541	–	Terrestrial plant macrofossil
ANP3	81.5–82.5	LuS 12227	8985 \pm 70	10121	10256–9898	–	Aquatic moss
ANP3	95.5–96.5	LuS 12221	10070 \pm 75	11622	11973–11315	–	Aquatic moss

^a Ua = Ångström Laboratory, Uppsala University, Sweden; LuS = Lund University Radiocarbon Laboratory, Sweden.

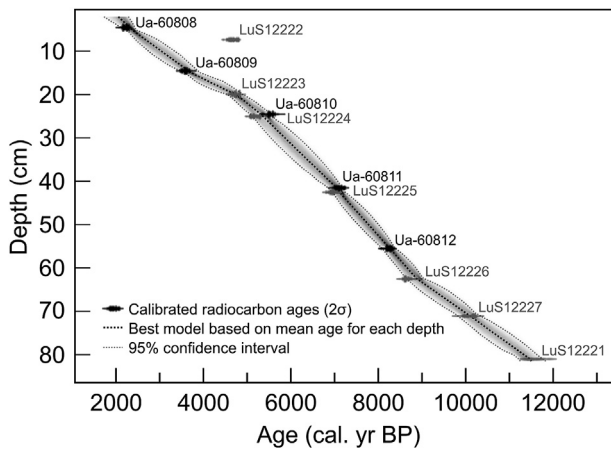


Fig. 4. Composite age depth model for surface core ANS1 (black symbols) and piston core ANP3 (grey symbols), established using GeoChronR (McKay et al., 2018), and the IntCal13 calibration curve (Reimer et al., 2013). Details of each radiocarbon age are given in Table 1.

sources (i.e., aquatic or terrestrial plants) have different chain length distribution. *Luzula confusa* contains the highest relative C₂₈ concentration, followed by *Salix polaris* (Fig. 7). These two terrestrial plant species also show the lowest concentrations of mid-chain waxes. The two mosses, *Polytrichum* sp. and an aquatic bryophyte, record the highest C₂₀ and C₂₂ concentrations, and relatively less C₂₈.

The leaf wax concentrations were highest in the Early and Late Holocene (Fig. 6), suggesting highest productivity during these periods. Long-chain leaf wax concentrations are higher than mid-chain concentrations throughout most of the record, reflecting dominance of terrestrial vegetation or higher leaf wax productivity for terrestrial plants. Generally, the sediment record is dominated by C₂₈, C₂₆ and C₂₄ *n*-alkanoic acids, with relatively higher C₂₀ and C₂₂ concentrations in ANP3 than in ANS1 (Fig. 7).

The average chain length follows a similar trend as the leaf wax concentrations, with high values (>26) during periods of high C₂₈ concentration (Fig. 6). This correlation is weaker for ANS1, where the concentration of mid-chain waxes (especially C₂₄) generally is higher than in ANP3.

4.4. $\delta^2\text{H}$ of sedimentary *n*-alkanoic acids

Based on the evidence presented above, we interpret $\delta^2\text{H}_{\text{C}_{28}}$ to reflect terrestrial plant wax $\delta^2\text{H}$ ($\delta^2\text{H}_{\text{terr}}$) and $\delta^2\text{H}_{\text{C}_{22}}$ to reflect aquatic plant wax $\delta^2\text{H}$ ($\delta^2\text{H}_{\text{aq}}$). $\delta^2\text{H}_{\text{terr}}$ is more stable throughout the

Holocene than $\delta^2\text{H}_{\text{aq}}$ (Fig. 8). The difference in $\delta^2\text{H}_{\text{terr}}$ between the cores is generally less than 10‰ for most of the record, whereas $\delta^2\text{H}_{\text{aq}}$ differs as much as 80‰ at times. When interpreting the $\delta^2\text{H}$ records for the two sediment cores separately, they suggest different stories.

ANP3 covers a longer period and extends further back in time than ANS1 (Fig. 8a). $\delta^2\text{H}_{\text{aq}}$ is relatively ²H-enriched prior to 9.9 cal. kyr BP (values generally above –230‰), for a short interval between 8.2–7.3 cal. kyr BP (–225‰) and after c. 6 cal. kyr BP (–230 to –205‰). For most of the time between 9.9–6 cal. kyr BP, $\delta^2\text{H}_{\text{aq}}$ is nearly 100‰ ²H-depleted (–305 to –270‰) compared to the beginning and end of the record. In contrast, $\delta^2\text{H}_{\text{terr}}$ varies by around 40‰ throughout the Holocene and is relatively ²H-depleted until c. 9.5 cal. kyr BP. After that, $\delta^2\text{H}_{\text{terr}}$ fluctuates between –170 and –145‰ for most of the record, with a slightly decreasing trend.

In ANS1, $\delta^2\text{H}_{\text{aq}}$ is relatively ²H-enriched prior to c. 7.9 cal. kyr BP. On the contrary, $\delta^2\text{H}_{\text{aq}}$ in ANP3 becomes ²H-depleted during this period. $\delta^2\text{H}_{\text{aq}}$ varies by around 40‰ (–300 to –260‰) for the rest of the ANS1 record, with values comparable to the ²H-depleted intervals in ANP3. $\delta^2\text{H}_{\text{terr}}$ is relatively ²H-enriched prior to c. 6.2 cal. kyr BP, with values between –160 and –150‰, compared to –180 to –165‰ for the later part of the record.

5. Discussion

The isotopic composition of *n*-alkanoic acids in the Austre Nevlingen record reveals environmental and climatic changes throughout the Holocene. Even though we are able to correlate ANS1 and ANP3 based on the XRF data (Fig. A, Appendix A), the $\delta^2\text{H}$ records of each core differ significantly, especially for the mid-chain compounds. Despite the large $\delta^2\text{H}_{\text{aq}}$ variability in the upper portion of ANP3, there are several data points with $\delta^2\text{H}$ values similar to ANS1, suggesting that this variability is real. The data do not indicate that one record should be favored over the other. Therefore, we discuss the $\delta^2\text{H}$ records for each core separately (Fig. 8a) and for the composite record (Fig. 8b). The composite record includes both the oldest sediments from ANP3 and the youngest sediments from ANS1, allowing us to discuss our results in full.

5.1. Influence of lake system dynamics on $\delta^2\text{H}$

In Austre Nevlingen, shifts between relatively ²H-enriched and ²H-depleted $\delta^2\text{H}_{\text{aq}}$ values during the Mid and Late Holocene (Fig. 8a and b) suggest that the lake is close to an isotopic threshold. This means that the lake water isotopes could shift between recording summer and mean annual precipitation isotope $\delta^2\text{H}$ depending on processes acting in the catchment (Cluett and Thomas, 2020; Thomas et al., 2020).

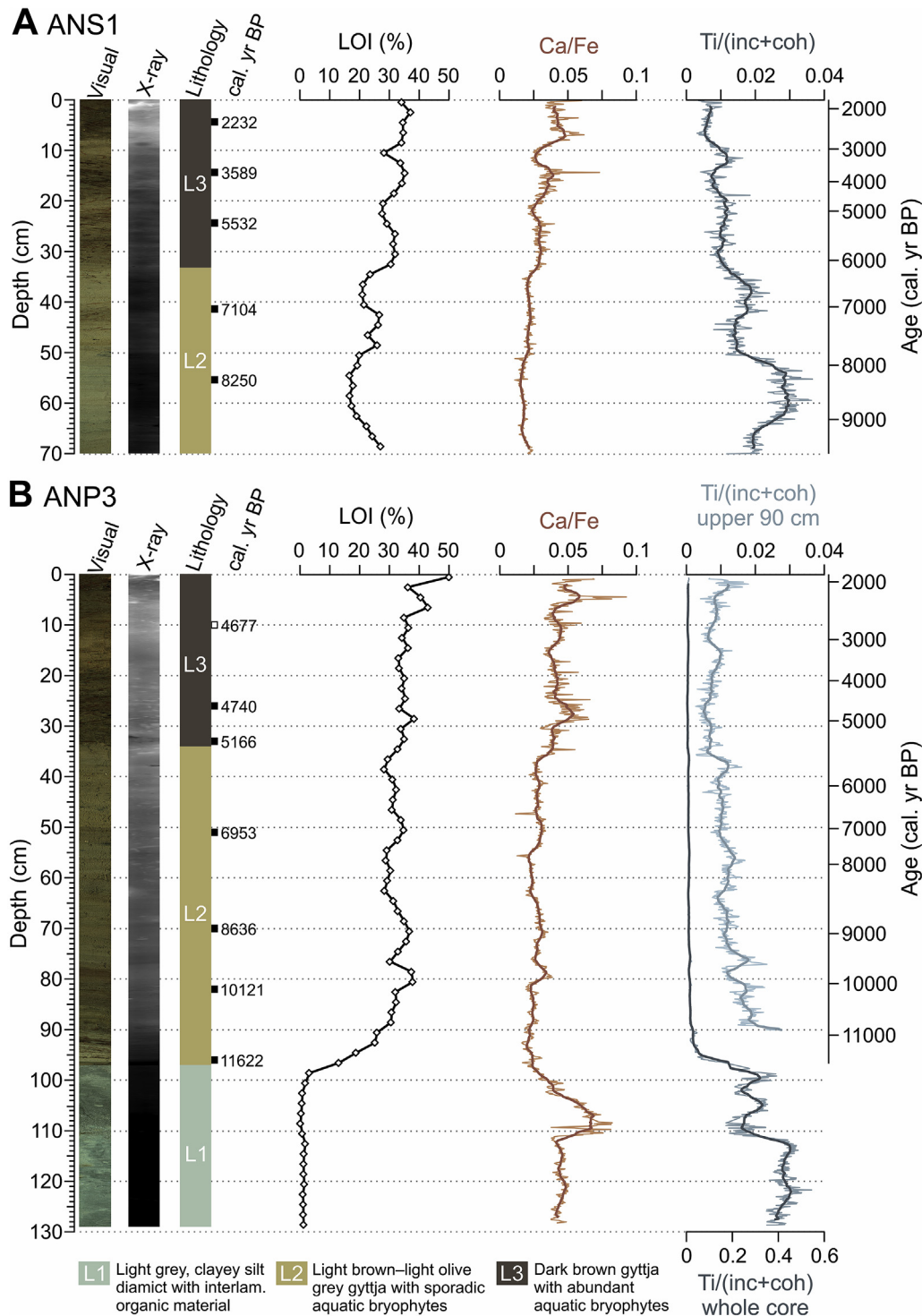


Fig. 5. Selected sediment proxies for (A) Surface core ANS1 and (B) Piston core ANP3. Core photograph, radiographic image, lithology, calibrated ^{14}C ages (median ages; see also Table 1), LOI (%), Ca/Fe and Ti normalized by the incoherent and coherent signal (Ti/(inc + coh)). The XRF data are plotted as raw data and with a 25-point running average.

We suggest that one of the main mechanisms causing the high amplitude (>80‰) variability in Austre Nevlingen $\delta^2\text{H}_{\text{aq}}$ could be the duration of ice on the lake. If lake-ice cover persists during the main spring snowmelt, most of the ^2H -depleted water could bypass the lake in these years (MacDonald et al., 2016). Furthermore, late lake-ice melt could delay the peak in primary production of the lake, and the ^2H -depleted winter precipitation isotope signal may not be incorporated into the leaf waxes. On the contrary, years

without summer lake-ice cover would cause the aquatic plants to receive and incorporate more ^2H -depleted water.

Another possible factor affecting the seasonal signal preserved by the leaf waxes is fluctuating lake level, affecting the residence time of water in the lake. Previously higher lake level (and therefore larger lake volume) is suggested by the light-colored 'bathtub' rim around the lake, implying a previously longer residence time. On the other hand, a lower lake level (and therefore smaller lake

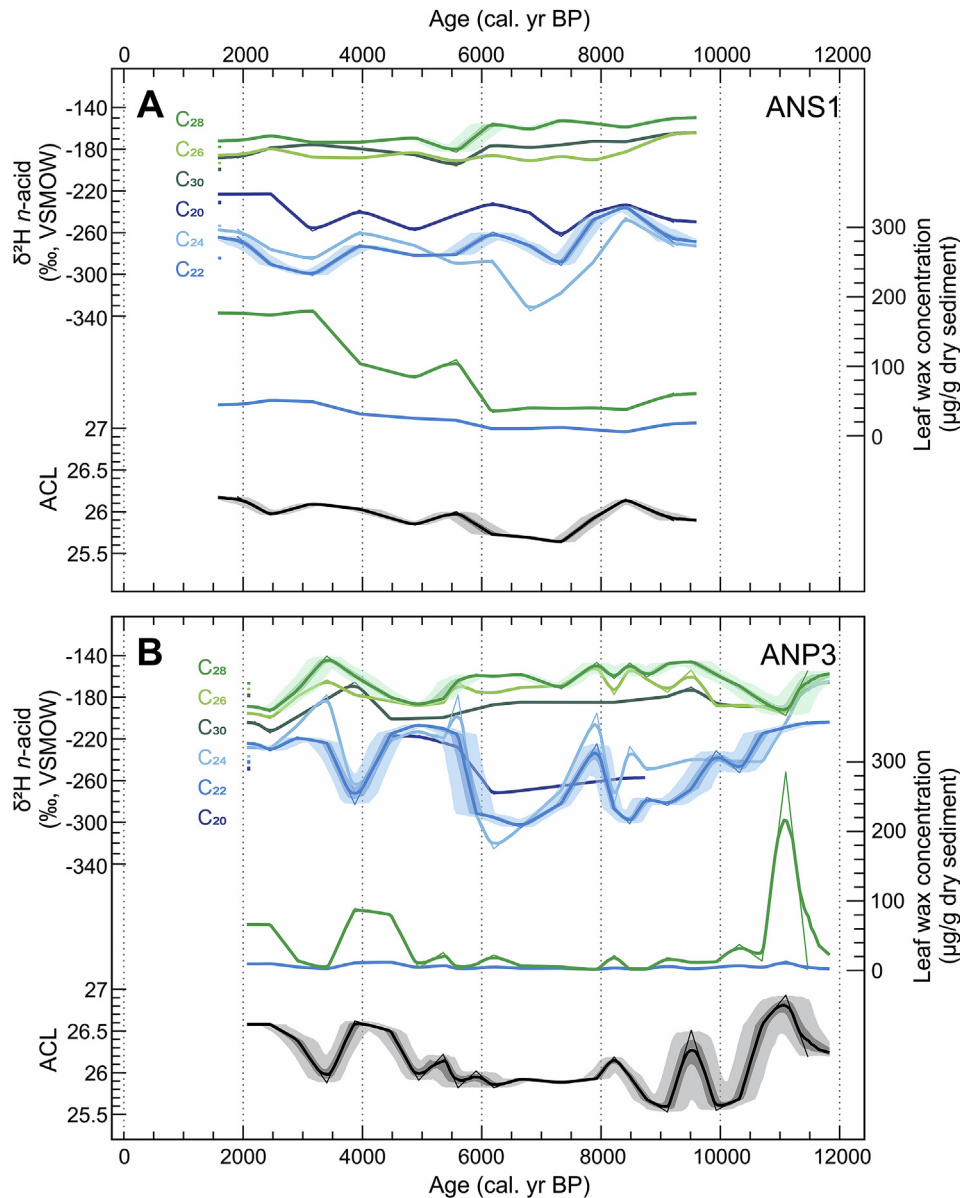


Fig. 6. Austre Nevlingen leaf wax data for (A) Surface core ANS1 and (B) Piston core ANP3. Leaf wax $\delta^2\text{H}$ for C_{20} – C_{24} (blue; aquatic) and C_{26} – C_{30} (green; terrestrial) *n*-alkanoic acids, absolute C_{22} and C_{28} leaf wax concentration and average chain length (ACL) distribution. Fine lines show the raw data plotted on the median of each age point and bold lines represent median values of the age model ensembles. The light and dark shading added to C_{22} , C_{28} and ACL show 1σ and 2σ age uncertainty, respectively (McKay et al., 2018). The vertical colored bars represent the average SEM proxy uncertainty. (For interpretation of the references to color in this figure legend, the reader is referred to the Web version of this article.)

volume) could result in a shorter residence time. If the lake was fully flushed by spring melt in some years but not in others (i.e., because of changes in lake volume), lake water, and therefore $\delta^2\text{H}_{\text{aq}}$, could have shifted between reflecting summer and mean annual precipitation $\delta^2\text{H}$ values. Such lake level shifts, although at a much larger scale, have been described from Hakluytvatnet (Fig. 1). Balascio et al. (2018) interpret a mid-Holocene (7.5–5 cal. kyr BP) hiatus in their sedimentary record to represent desiccation of the lake as a response to warm and/or dry conditions at the time. Today, Austre Nevlingen lake water is not affected by evaporative enrichment, as the *d*-excess measured in August 2018 (9.0‰) is within the range of precipitation *d*-excess. It is possible that evaporation had a significant impact on lake water $\delta^2\text{H}$ during earlier Holocene warm periods as suggested by Balascio et al. (2018), if the ice-free periods were longer and the summers were warmer. Another possible way

to switch the lake water from a mean annual to a summer signal would be by subterranean drainage during periods of deepened active layer. This process could theoretically decrease the lake volume enough to flush the lake completely in summer.

Another explanation for the up to 80‰ difference in $\delta^2\text{H}_{\text{aq}}$ between the Austre Nevlingen cores and between adjacent samples in a single core could be the relative abundance of aquatic bryophytes in the samples, since moss mats occur at different depths in the cores. If some samples mainly contain interbedded bryophytes rather than bulk sediments, $\delta^2\text{H}_{\text{aq}}$ in these samples could be dominated by one source rather than a mix of different sources. However, we could not confirm any correlation between samples dominated by aquatic mosses and significantly ^2H -enriched or ^2H -depleted $\delta^2\text{H}_{\text{aq}}$ values, when revisiting the residues after lipid extraction. Furthermore, there are no distinct shifts in the relative

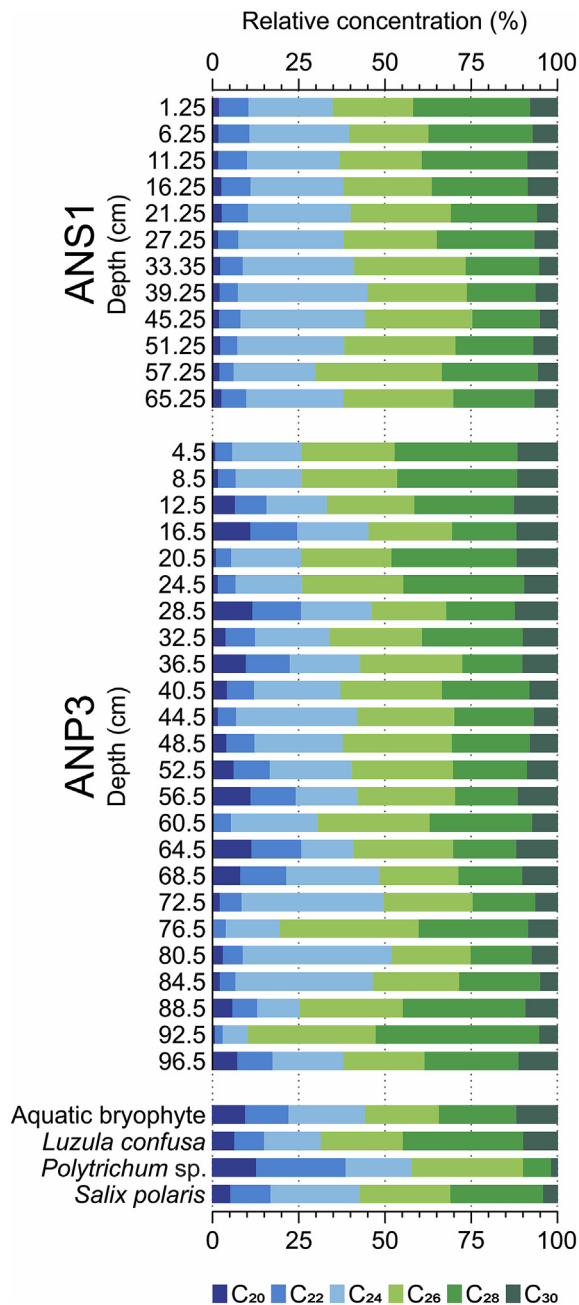


Fig. 7. Relative concentration distribution of even-chained C_{20} – C_{30} n -alkanoic acids in sediment cores ANS1 and ANP3, as well as selection of modern plants growing in the catchment.

chain length concentration associated with the shifts in δ^2H_{aq} values that would suggest different plant sources (Fig. 7). Therefore, the presence of aquatic plant macrofossils does not explain the observed isotopic differences.

More variable δ^2H_{aq} than δ^2H_{terr} (Fig. 8a and b) could reflect that the most prominent changes in precipitation have occurred during winter, since δ^2H_{aq} in Austre Nevlingen is interpreted to represent a mix of summer and winter precipitation, whereas δ^2H_{terr} represents summer rainfall. Additionally, δ^2H_{aq} is relatively more sensitive to local variation within the lake, whereas δ^2H_{terr} represents conditions integrated throughout the catchment.

We compare our isotope values from Austre Nevlingen (Fig. 8a and b) to those from Hakluytvatnet (Fig. 8c; Balascio et al., 2018),

and find some significant differences between the two records. The precipitation isotope seasonality recorded by the two lakes differ because of the large difference in residence time between the lakes. Hakluytvatnet is much smaller than its catchment (Balascio et al., 2018), and therefore has a short residence time (i.e., it is likely fully flushed by spring melt and again by summer precipitation each year). The aquatic leaf wax δ^2H record from Hakluytvatnet should therefore be interpreted in terms of summer precipitation isotopes. This could explain the similar δ^2H patterns for C_{25} and C_{29} n -alkanes in that record (Fig. 8c). Austre Nevlingen has a longer residence time, and is not completely flushed by summer precipitation each year, so likely reflects the mean annual precipitation δ^2H . Because δ^2H for C_{22} and C_{28} in Austre Nevlingen represent different seasons, they show different patterns (Fig. 8a and b). Larger isotopic variation at Austre Nevlingen compared to Hakluytvatnet could also reflect greater seasonality at our site, as it is situated farther from the main moisture source pathway along the west coast of Spitsbergen. Furthermore, the difference in analyzed leaf wax compounds (i.e., n -alkanes for Hakluytvatnet and n -alkanoic acids for Austre Nevlingen) could explain the different trends (Curtin et al., 2019).

The discussed factors highlight the need to interpret lake δ^2H records carefully, depending on the local conditions. For this specific system, we interpret δ^2H_{terr} to reflect summer precipitation and evaporative enrichment and δ^2H_{aq} to reflect mean annual precipitation, sometimes modified by changing lake dynamics. Mean annual precipitation isotopes mainly change as a function of moisture transport (Frankenberg et al., 2009). Even though the main signal preserved can be assumed to reflect climatic changes, the influence of various internal lake dynamics must also be taken into account.

5.2. Early Holocene: 11.7–8.2 cal. kyr BP

Decreasing δ^2H_{aq} and increasing δ^2H_{terr} from the base of our δ^2H record at c. 11.5 to 9.5 cal. kyr BP (Fig. 8a and b) coincide with a June solstice insolation maximum at 80°N (Fig. 8f; Laskar et al., 2004) and rapidly increasing eastern Fram Strait surface and subsurface temperatures (Fig. 8e; Hald et al., 2007; Werner et al., 2016). Similar subsurface temperature trends are reported from the Barents Sea Margin (Risebrobakken et al., 2011). Hald et al. (2007), Werner et al. (2016), and Risebrobakken et al. (2011) interpret this warming as a response to increased influx of warm Atlantic water (AW) to the waters around Svalbard. This led to reduced spring sea-ice cover, evident by strongly decreasing PIP_{25} indices in the eastern Fram Strait (Figs 1a and 8d; $P_{BIP_{25}}$ and $P_{DIP_{25}}$, based on brassicasterol and dinosterol, respectively; Werner et al., 2016). Additionally, Allaart et al. (2020) suggested an Early Holocene decline in sea-ice cover in the nearby Wijdefjorden (Fig. 1c), based on IP_{25} in marine sediments. Increased radiative forcing and greater AW heat transfer has also been suggested to cause warmer-than-present conditions in shallow waters around Svalbard between 11–9 cal. kyr BP (Mangerud and Svendsen, 2018) and peak warmth in three northern Svalbard lakes around 10 cal. kyr BP (van der Bilt et al., 2019).

Increasing Austre Nevlingen δ^2H_{terr} between 11–9.5 cal. kyr BP supports Early Holocene summer warming, suggesting enhanced evaporative enrichment during summer and/or a more proximal moisture source for growing season precipitation. A more proximal moisture source could be explained by reduced spring sea-ice cover and increased local evaporation. Balascio et al. (2018) also interpret increasing δ^2H (for C_{25} to C_{29} n -alkanes) to reflect the high summer insolation and greater influence of mild air masses from the south. These processes could also explain the strong decreasing δ^2H_{aq} during the same period. Mean annual temperatures would have to

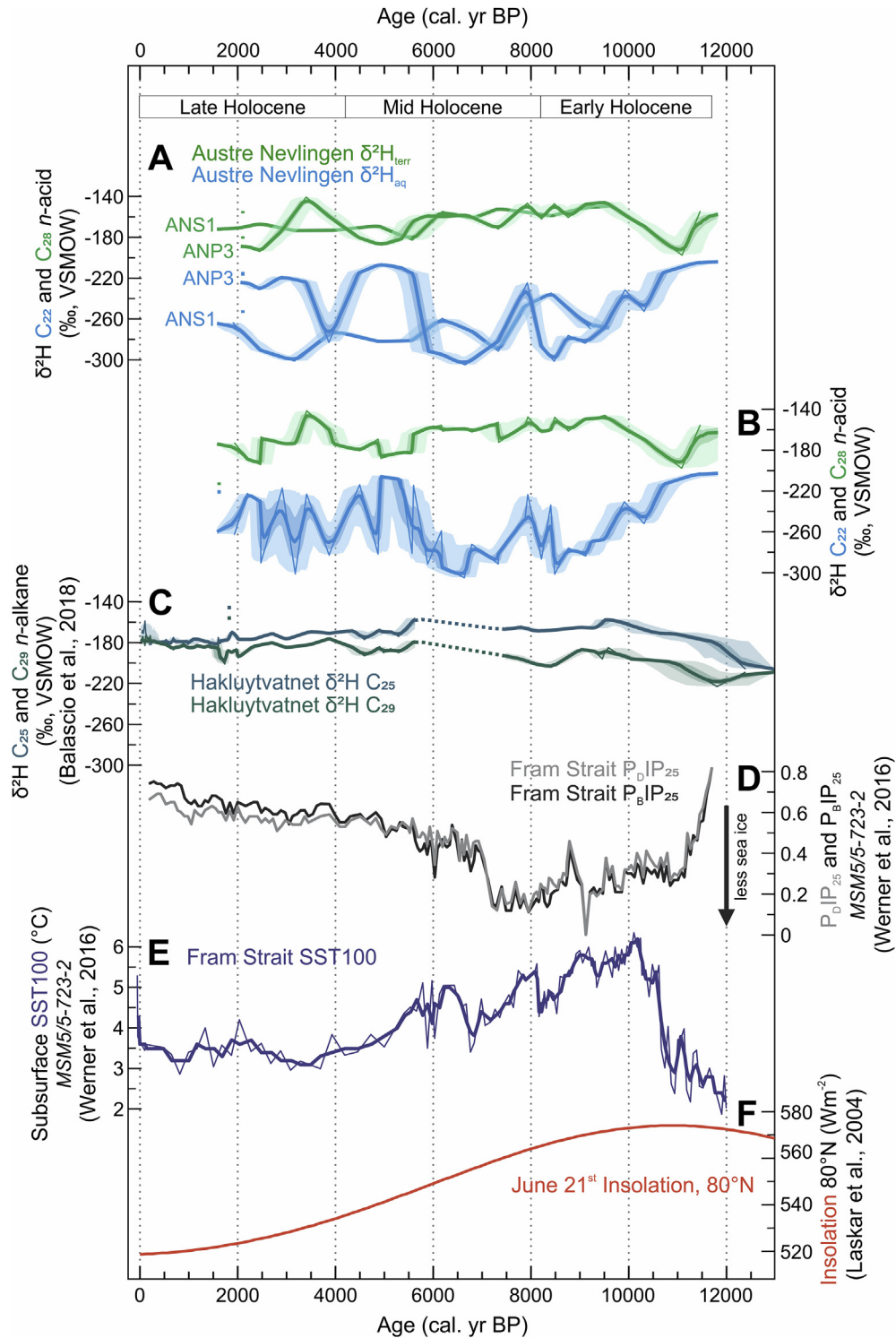


Fig. 8. Austre Nevlingen leaf wax hydrogen isotope data compared to selected regional Holocene climate records. For locations, see Fig. 1. Lines, shading and vertical colored bars shown as in Fig. 6. (A) Austre Nevlingen leaf wax $\delta^2\text{H}$, C_{22} (blue; aquatic) and C_{28} (green; terrestrial) n -alkanoic acids for sediment cores ANS1 and ANP3. (B) Composite leaf wax $\delta^2\text{H}$ record for Austre Nevlingen. (C) Lake Hakluytvatnet leaf wax $\delta^2\text{H}$, C_{25} (blue) and C_{29} (green) n -alkanes (Balascio et al., 2018). Dashed lines in the mid-Holocene denote a possible hiatus, suggested by Balascio et al. (2018). (D) Eastern Fram Strait sea-ice proxies $\text{P}_{\text{a}}\text{IP}_{25}$ (black) and $\text{P}_{\text{b}}\text{IP}_{25}$ (grey) (Werner et al., 2016). (E) Eastern Fram Strait subsurface temperature based on planktic foraminiferal fauna assemblages (fine line; Werner et al., 2016), including 3-point running means (bold line; Husum and Hald, 2012). (F) June 21st insolation at 80°N (Laskar et al., 2004). (For interpretation of the references to color in this figure legend, the reader is referred to the Web version of this article.)

be more than 30 °C cooler to explain the 100‰ depletion, considering a temperature- $\delta^{18}\text{O}$ relationship of 0.34–0.40‰/°C (Kotlyakov et al., 2004), translated to $\delta^2\text{H}$ by multiplying by 8 (Dansgaard, 1964). More likely, the ^2H -depletion reflects a strong winter signal as a response to greater winter ocean evaporation due to the reduced sea-ice cover or more distal moisture transport during this interval.

These interpretations support an early Holocene Climate Optimum, which is in agreement with other recent studies in Svalbard (Werner et al., 2016; Mangerud and Svendsen, 2018; Røthe et al., 2018; de Wet et al., 2018; van der Bilt et al., 2018, 2019; Voldstad et al., 2020), and elsewhere in the Arctic (Lecavalier et al., 2017; McFarlin et al., 2018). Our $\delta^2\text{H}$ record suggests that the regional Early Holocene warming had a strong effect on precipitation at the time, with local moisture from open seas leading to increased winter snowfall. This could also explain why some glaciers advanced in this otherwise warm period (e.g., Farnsworth et al., 2017, 2018). After c. 9.5 cal. kyr BP, $\delta^2\text{H}_{\text{terr}}$ stabilizes around -160‰ , indicating stable summer conditions during this period. $\delta^2\text{H}_{\text{aq}}$ continues to decrease until c. 8.5 cal. kyr BP, reflecting continued sea-ice decrease (Werner et al., 2016; Allaart et al., 2020) and enhanced local evaporation.

5.3. Mid Holocene: 8.2–4.2 cal. kyr BP

In the Mid Holocene, $\delta^2\text{H}_{\text{terr}}$ remains stable around -160‰ until c. 5.6 cal. kyr BP, indicating stable summer conditions. Relatively ^2H -depleted values after 5.6 cal. kyr BP suggest gradual cooling, with less evaporative enrichment and/or more ^2H -depleted summer precipitation during this period. Mangerud and Svendsen (2018) suggest Svalbard ocean water temperatures c. 4 °C warmer than present between 8.2–6 cal. kyr BP, based on the presence of the marine bivalve mollusk *Mytilus edulis*, many of them along the northern coast. East Fram Strait subsurface temperatures stayed warm (up to 6 °C) until c. 5 cal. kyr BP, with a slightly decreasing long-term trend (Fig. 8e; Werner et al., 2016). Both Mangerud and Svendsen (2018) and Werner et al. (2016) interpret this warming to be attributed to intrusion of AW to the Svalbard shelf areas. High Mid Holocene temperatures are also inferred from lake sediment records, with minimal minerogenic accumulation, suggesting greatly reduced glacial activity or ice-free catchments (Svendsen and Mangerud, 1997; Røthe et al., 2015, 2018).

In Austre Nevlingen, ^2H -depleted $\delta^2\text{H}_{\text{aq}}$ values suggest increased winter precipitation starting c. 9.5 cal. kyr BP and persisting into the Mid Holocene until c. 6 cal. kyr BP. An increase in winter precipitation could be explained by continuously low sea-ice extent. Low $\text{P}_{\text{DIP}_{25}}$ and $\text{P}_{\text{IP}_{25}}$ values until c. 7 cal. kyr BP support this interpretation (Fig. 8d; Werner et al., 2016). Higher winter precipitation during the Early-Mid-Holocene is also suggested by Røthe et al. (2018). They propose snowmelt dominated runoff to lake Vårfluesjøen between 10.2–7 cal. kyr BP, inferred from high frequency of ‘snowmelt layers’ in the sediment core. Vårfluesjøen is located on the western side of Wijdefjorden, c. 27 km WSW of Austre Nevlingen. After 7 cal. kyr BP, a lower frequency of snowmelt layers and the occurrence of aeolian sand in Vårfluesjøen, suggest drier Mid Holocene conditions (Røthe et al., 2018). Drier Mid Holocene climate is also proposed by Balascio et al. (2018), inferred from the hiatus (7.5–5 cal. kyr BP) in their sedimentary record. Our Austre Nevlingen record suggests that winter-dominated precipitation sustained until c. 6 cal. kyr BP, without any indications of desiccation of the lake.

5.4. Late Holocene: 4.2–0 cal. kyr BP

From c. 6 cal. kyr BP and into the Late Holocene, both $\delta^2\text{H}_{\text{aq}}$ and

$\delta^2\text{H}_{\text{terr}}$ express larger amplitude variability than in the Early Holocene and first part of the Mid Holocene (Fig. 8a and b). We observe disagreement between the isotopic values in the two cores, but cannot explain exactly why. Some isotopic overlap between the cores (e.g., the ^2H -depleted $\delta^2\text{H}_{\text{aq}}$ value in ANP3 c. 3.9 cal. kyr BP) supports that this large-amplitude variability is real. One potential interpretation is that this variability reflects changes in the precipitation seasonality recorded by the lake. As discussed in Section 5.1, Austre Nevlingen might be close to an isotopic threshold, meaning that the lake water incorporates relatively ^2H -depleted winter precipitation isotopes in some years, but not in others. This process could be explained by changing regional conditions and/or variable conditions in the catchment, amplifying the climatic signal. These processes could affect the seasonality of the isotopes registered by the lake in the Late Holocene $\delta^2\text{H}_{\text{aq}}$ record. Hence, it is difficult to decipher the Late Holocene climate signal, except that the high-amplitude $\delta^2\text{H}_{\text{aq}}$ variability suggests greater climate variability.

The Late Holocene is generally characterized by Neoglacial cooling from c. 5 cal. kyr BP, with decreasing summer insolation (Fig. 8f; Laskar et al., 2004) and ocean cooling around Svalbard (Fig. 8e; Hald et al., 2004; Rasmussen et al., 2014; Werner et al., 2016). Weaker AW advection also led to increased sea-ice cover in eastern Fram Strait (Fig. 8d; Werner et al., 2016), and cooler conditions on land, registered by alkenones (van der Bilt et al., 2018) and *n*-alkanes (Fig. 8c; Balascio et al., 2018) in lakes. Glaciers started to re-advance c. 4–3 cal. kyr BP (Svendsen and Mangerud, 1997; Røthe et al., 2015, 2018; Miller et al., 2017; Bartels et al., 2018; Lovell et al., 2018).

In our Austre Nevlingen record, slightly ^2H -depleted $\delta^2\text{H}_{\text{terr}}$ values after 5.6 cal. kyr BP may be due to this regional cooling trend, as cooler conditions would cause less evaporative enrichment and more ^2H -depleted summer precipitation. Neoglacial cooling could also explain the more ^2H -enriched parts of the $\delta^2\text{H}_{\text{aq}}$ record after 6 cal. kyr BP. Regional cooling and concomitant increases in sea-ice cover would have caused less winter precipitation, in turn causing the lake water to be biased to ^2H -enriched summer precipitation.

6. Conclusions

- Leaf wax hydrogen isotopes from Austre Nevlingen suggest large variability in precipitation seasonality on northern Spitsbergen throughout the Holocene. One of the strengths of our reconstruction is that we can extract both the summer and mean annual precipitation signal. This is possible because terrestrial and aquatic plants use different source water (soil water and lake water, respectively). Furthermore, the lake is not completely flushed by spring melt, with the long residence time allowing winter precipitation isotopes to be incorporated into the aquatic leaf waxes. Contrasting isotopic composition of mid-chain and long-chain *n*-alkanoic acids suggests different leaf wax sources for these two sets of compounds.
- Most prominent Holocene precipitation changes occurred in winter, reflected in greater variability of the aquatic (mean annual) than terrestrial (summer) leaf wax hydrogen isotope ratios.
- Early Holocene regional warming had a strong effect on moisture availability and precipitation seasonality in northern Svalbard. High summer insolation and strong Atlantic water influx contributed to reduced sea ice, which we suggest favored greater local winter evaporation leading to increasing winter precipitation.
- Between 9.5–6 cal. kyr BP, the record is characterized by ^2H -enriched terrestrial and ^2H -depleted aquatic hydrogen

isotope ratios, which we interpret to reflect enhanced summer evaporation and/or ^2H -enriched summer precipitation, reduced sea-ice cover and local moisture contributing to high winter precipitation.

- After c. 6 cal. kyr BP, the $\delta^2\text{H}$ records show greater amplitude variability. We interpret these shifts to be influenced by lake system dynamics, such as the duration of ice cover on the lake, that amplify the climate signal. Hence, we cannot clearly distinguish the amplitude of climate variability in this portion of the record, although the lake variability is most likely in phase with climate variability.
- Our results suggest that the precipitation seasonality in northern Spitsbergen is strongly linked to regional ocean surface conditions. As a result, the positive trend in winter precipitation observed in Svalbard today, may amplify because of warming ocean surface waters and reduction in sea ice.

Data availability

All chronological and geochemical data presented in this paper are freely available at the NOAA/World Data Service for Paleoclimatology website: <https://www.ncdc.noaa.gov/paleo/study/29852>.

Author contributions

AS, EKT, LH and SEK developed the idea; SEK, AS, WRF, LA, SB and ÓI conducted fieldwork; SEK, WRF, AS and LH described and interpreted the lithology; AS and LH identified plant macrofossils for radiocarbon dating; LA processed the bathymetry data; SEK, OCC and SD conducted organic and stable isotope biogeochemical work; SEK, EKT and NPM did the age modelling and leaf wax data analysis and visualization; all authors contributed with interpretation and discussion of results; SEK wrote the paper with contributions from all co-authors.

AS = Anders Schomacker, EKT = Elizabeth K. Thomas, LA = Lis Allaart, LH = Lena Håkansson, NPM = Nicholas P. McKay, OCC = Owen C. Cowling, ÓI = Ólafur Ingólfsson, SB = Skafti Brynjólfsson, SD = Sandrine Dubosq, SEK = Sofia E. Kjellman, WRF = Wesley R. Farnsworth.

Declaration of competing interest

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

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Appendix A

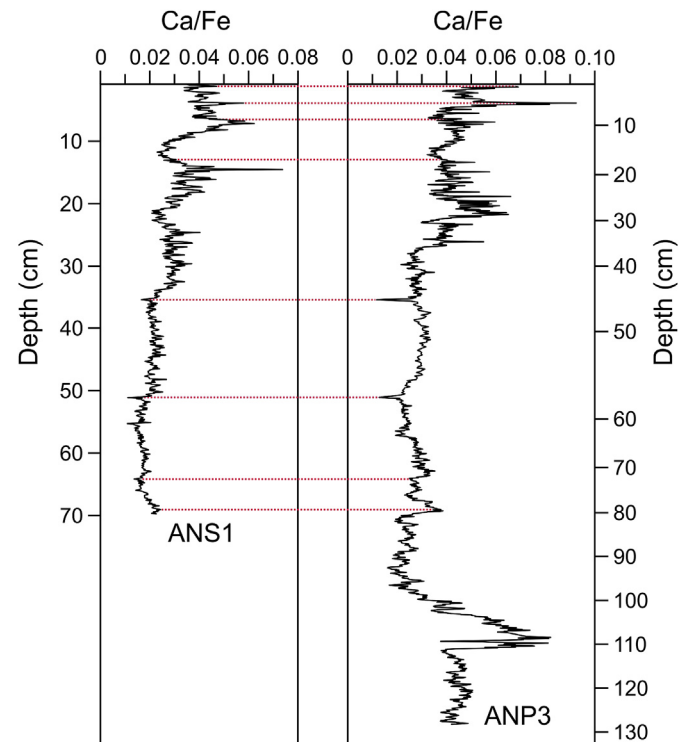


Fig. A. Correlation between surface core ANS1 and piston core ANP3 from lake Austrveiv, Svalbard. The records were aligned in AnalySeries (v. 2.0.8; Paillard et al., 1996), based on tie points in the Ca/Fe data.

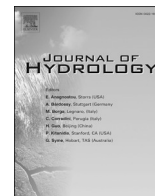
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Paper II



Research papers

Arctic and sub-Arctic lake water $\delta^2\text{H}$ and $\delta^{18}\text{O}$ along a coastal-inland transect: Implications for interpreting water isotope proxy recordsSofia E. Kjellman^{a,*}, Elizabeth K. Thomas^b, Anders Schomacker^a^a Department of Geosciences, UiT The Arctic University of Norway, P.O. Box 6050 Langnes, NO-9037 Tromsø, Norway^b Department of Geology, University at Buffalo, State University of New York, 126 Cooke Hall, Buffalo, NY 14260, USA

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ABSTRACT

High-latitude lakes are sensitive to climate change and store information about large-scale circulation changes and catchment-integrated processes. Lakes are mainly recharged by meteoric water, meaning that some lake sediment proxies may indirectly archive the stable isotopic composition of hydrogen ($\delta^2\text{H}$) and oxygen ($\delta^{18}\text{O}$) of past precipitation. Yet, despite similar precipitation input, lakes within a region may exhibit a wide range of isotopic values due to the varying influence of inflow seasonality and evaporation. Moreover, the relative sensitivity of each lake to these controls may vary through time, something that is difficult to account for. Here, we evaluate the impact of variable inflow $\delta^2\text{H}$ and evaporation on the lake water isotopic composition across northern Fennoscandia (Norway, Finland, and Sweden). We measured lake water $\delta^2\text{H}$ and $\delta^{18}\text{O}$ of 135 lakes spanning from the north Norwegian coast along a 460 km transect to the Bothnian Bay, sampled from 2018 to 2020. Our data show that both coastal and inland lakes are sensitive to distillation during moisture transport, and that lakes farther from the Atlantic Ocean are additionally impacted by evaporation. We estimated the isotopic composition of lake water inflow values for evapo-concentrated transect lakes ($\delta^2\text{H}_i$) using a Bayesian method. Resampled transect lakes had more depleted $\delta^2\text{H}_i$ in 2020 than in 2019, indicating either that precipitation was ^2H -depleted or that more winter precipitation contributed inflow to the lakes in 2020 compared to in 2019. We suggest that the more ^2H -depleted values in 2020 were a response to a snow-rich winter, associated with extremely positive Arctic Oscillation (AO+) conditions and increased moisture supply from the North Atlantic. We find evidence that lake water isotopic variability in this region reflects a combination of seasonal precipitation changes associated with atmospheric circulation changes, and catchment-integrated evaporation. Careful consideration of the variable sensitivity to these processes is essential when making inferences about past climate based on lake water isotope proxies.

1. Introduction

Amplified warming in the northern high latitudes is predicted to cause an intensification of the hydrological cycle (Rawlins et al., 2010; Collins et al., 2013; Bintanja & Selten, 2014; Bintanja et al., 2020). These changes affect both terrestrial and ocean freshwater budgets and have great cryospheric and ecological impact (Bring et al., 2016; Vihma et al., 2016; Wrona et al., 2016). Arctic precipitation change is caused by two main mechanisms with different seasonal imprints. Increased local evaporation mainly occurs in fall and winter, due to accelerated sea-ice loss (Bintanja & Selten, 2014; Bailey et al., 2021). In contrast, poleward moisture transport is greatest in summer, when the meridional temperature and moisture gradients increase the most and continental

moisture sources play a significant role (Vázquez et al., 2016; Singh et al., 2017). Furthermore, greater interannual variability in the summer moisture influx from extrapolar regions is projected to contribute to increasing interannual variability in Arctic precipitation (Bintanja et al., 2020). The projected trends remain uncertain, relying on sparse observational data to describe complex spatiotemporal patterns. To improve the constraints on long-term variability in precipitation and atmospheric circulation, we can use lake-water-derived proxy records (Sundqvist et al., 2014; Linderholm et al., 2018).

Stable isotope ratios of oxygen ($\delta^{18}\text{O}$) and hydrogen ($\delta^2\text{H}$) in precipitation are frequently used to infer processes in the hydrological cycle due to their sensitivity to changes in atmospheric temperature, moisture, and circulation (Dansgaard, 1964; Rozanski et al., 1993; Gat, 1996).

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Much of the variation in precipitation isotopic composition results from equilibrium fractionation during phase changes (e.g., evaporation, condensation, and freezing) during transport from the moisture source to the precipitation site (Gat, 1996). Equilibrium fractionation is highly temperature dependent, and therefore varies both spatially and temporally. The global average relationship between $\delta^2\text{H}$ and $\delta^{18}\text{O}$ in meteoric waters is described by the global meteoric water line (GMWL), defined as $\delta^2\text{H} = 8 \times \delta^{18}\text{O} + 10$ (Craig, 1961). Local meteoric water lines (LMWLs) account for site-specific variation in the isotopic composition due to factors including moisture sources, transport conditions and seasonality. For example, in seasonally snowy regions, slopes of the LMWLs tend to be similar to the GMWL, but the intercepts lower than in other regions due to snow formation processes (Putman et al., 2019).

The isotopic signal of precipitation can be preserved in climate proxies, allowing us to extend our records back beyond the instrumental period. In northern Fennoscandia, the oldest temperature and precipitation amount observations date back to the 1860's (Alexandersson, 2002), whereas precipitation isotope records are limited to a handful of sites, most of them only measuring $\delta^{18}\text{O}$, covering timespans of less than ten years each, and only after 1975 (IAEA/WMO, 2019).

Lakes are excellent archives of past climate change, as their isotopic composition often at least partially reflects precipitation isotopes, and they contain a wide range of proxies that preserve the lake water isotopic composition. In northern Fennoscandia, reconstructions of Holocene precipitation, humidity and atmospheric circulation have mainly been based on $\delta^{18}\text{O}$ from lacustrine carbonates (Hammarlund et al., 2002; Rosqvist et al., 2007), diatom biogenic silica (Shemesh et al., 2001; Rosqvist et al., 2004, 2013; Jonsson et al., 2010), and lake sediment cellulose (St. Amour et al., 2010). Other studies use $\delta^2\text{H}$ of lipid biomarkers to reconstruct different aspects of the water cycle (Nichols et al., 2009; Thienemann et al., 2019; Balascio et al., 2020).

Recent dual biomarker approaches have focused on comparing leaf waxes produced by aquatic and terrestrial plants to reconstruct Holocene and Last Interglacial atmospheric circulation changes, precipitation seasonality and summer moisture balance (Rach et al., 2017; Thomas et al., 2018, 2020; Curtin et al., 2019; Kjellman et al., 2020; Katrantsiotis et al., 2021). These approaches rely on the assumption that the aquatic plant biomarkers are minimally impacted by evaporative enrichment. This might be true for lakes in regions with positive moisture balance (precipitation > evaporation), but not for lakes in more arid areas (Cluett & Thomas, 2020).

When lakes undergo evapo-concentration, they will deviate from their starting isotopic compositions along the MWL due to kinetic fractionation, and evolve along a local evaporation line (LEL; Gat, 1996). The slope of the LEL is controlled by atmospheric conditions (e.g., temperature, relative humidity, and the isotopic composition of atmospheric moisture) and is usually between 5 and 6 at the mid-latitudes and sometimes steeper at high-latitudes (Gibson et al., 2008). If the slope of the LEL is known, it can be used, along with the lake water $\delta^2\text{H}$ and $\delta^{18}\text{O}$ values, to calculate the intersection between the LEL and the MWL, which represents lake water inflow values. One common approach to estimate regional LEL slopes is to adopt a linear regression fit to a group of lakes within the region (Clark & Fritz, 1997). This approach assumes that all the isotopic variation is a result of evaporation, and that all lakes in a region start with the same source isotopic composition (i.e., the lake inflow value; Bowen et al., 2018). This is seldom true, especially for systems with a strong seasonality. At high latitudes, the relative proportion of isotopically depleted snowfall and isotopically enriched rainfall can change between years, resulting in different inflow values from year to year (Tondou et al., 2013). Another approach is to apply theoretical modeling, based on the Craig-Gordon model, which requires knowledge of the isotopic composition of both precipitation and atmospheric moisture (Craig & Gordon, 1965), to infer inflow values for each lake individually. Regardless the technique used to calculate inflow values from evapo-concentrated lakes, most lake water $\delta^2\text{H}$ and $\delta^{18}\text{O}$ values contain information about both the degree of

evaporative enrichment and the inflow isotopic composition. Thus, lake water $\delta^2\text{H}$ and $\delta^{18}\text{O}$ values can be used to infer past change in precipitation $\delta^2\text{H}$ and $\delta^{18}\text{O}$ values and evaporation, but the influence of both inflow and evaporation also confounds isotopic interpretation. Examining spatial and temporal variations in lake water $\delta^2\text{H}$ and $\delta^{18}\text{O}$ values along modern climate gradients can guide interpretations of lake water isotope proxy records.

In northern Fennoscandia, Holocene changes in lake-water-derived $\delta^{18}\text{O}$ and $\delta^2\text{H}$ are often assumed to reflect changes in North Atlantic atmospheric and oceanic circulation (e.g., Hammarlund et al., 2002; Rosqvist et al., 2007; Thienemann et al., 2019). The North Atlantic is the predominant moisture source, but during periods with a more southerly position of the polar front, precipitation can originate from Arctic air masses (Rosqvist et al., 2007). This is further complicated by the fact that Arctic and sub-Arctic lakes can display a wide range of water isotopic compositions, despite similar precipitation input (Gibson & Edwards, 2002; Jonsson et al., 2009; Cluett & Thomas, 2020). This is because meteoric water inputs are modulated by catchment characteristics and additionally modified by lake water evaporation (Leng & Marshall, 2004; Cluett & Thomas, 2020). Lake and catchment morphometry and precipitation amount control the throughflow regime of the lake, since these factors govern the lake water residence time (Jonsson et al., 2009). Furthermore, the travel time of water through surface and groundwater systems and into lakes varies in time and space depending on factors such as catchment size, slope, soil type and seasonality (Botter et al., 2010; Heidebüchel et al., 2012; Jutebring Sterte et al., 2021). Lakes with wetland-dominated catchments generally have shorter residence times and lower evaporation-to-inflow (E/I) ratios (Gibson et al., 2002). To make inferences about past climate based on changes in lake-water-derived $\delta^{18}\text{O}$ and $\delta^2\text{H}$, we therefore need to understand both the regional climate and local hydrology (Jonsson et al., 2009; Jones et al., 2016; Cluett & Thomas, 2020).

Jonsson et al. (2009) measured the water isotopic composition of 11 lakes close to Abisko in northwestern Sweden and found that lakes in this region are mainly recharged by precipitation and shallow groundwater, with little influence of evaporation. They also demonstrated that the lake water isotopic composition varies on a seasonal basis, and that lakes with short residence time (<6 months) are more sensitive to seasonal changes, whereas lakes with longer residence times (>6 months) have lake water isotopic compositions closer to mean annual precipitation. Balascio et al. (2018) measured modern lake water isotopic composition of 15 lakes along a transect from near Tromsø in Norway to the Finnish-Swedish border. The lakes displayed decreasing isotopic values moving inland from the coast, and most of them showed little influence of evaporation. On western Greenland, a recent study of 140 lakes along an aridity gradient between the Labrador Sea and the Greenland Ice Sheet margin demonstrated that lake water $\delta^{18}\text{O}$ and $\delta^2\text{H}$ are impacted by inflow isotopic composition and catchment-integrated evaporation in all lakes, but that the relative importance of those two variables in a given lake depends on climate and lake and catchment morphometry (Cluett & Thomas, 2020).

Here, we investigate the water isotopic composition of lakes across Arctic and sub-Arctic Fennoscandia (Norway, Finland, and Sweden) to better constrain the influence of inflow isotopic composition and seasonality on the lake water isotopic composition throughout this region. We target lakes across a 460-km-long climatic gradient to explore the relative importance of inflow values and evaporative enrichment, and consider the interannual variability by comparing the isotopic composition from lakes sampled in several years. By identifying modern controls on the lake water isotopic composition, we aim to evaluate the sensitivity of lake water to changes in these hydrological and climate processes. This information is crucial to identify lakes that have the potential to accurately record the aspect of the water cycle we are interested in, and to make solid proxy interpretations.

2. Regional setting

The study area spans from the North Atlantic Ocean in the northwest to the Bothnian Bay in the southeast, between 65.74°N to 71.11°N and 18.39°E to 30.06°E (Fig. 1). The Late Weichselian Scandinavian Ice Sheet retreated from the outer coast c. 18–16 cal. kyr BP, whereas the interior parts were not ice-free until 11–10 cal. kyr BP (Hughes et al., 2016; Stroeven et al., 2016). The coast is characterized by alpine topography, with steep mountains intersected by subglacially eroded fjords and U-shaped valleys. The highest mountains are the Lyngen Alps, reaching up to c. 1800 m a.s.l. (Fig. 1b-c). The bedrock is often exposed, especially at high elevation, whereas areas closer to sea level are partly covered by Quaternary deposits (mainly till; NGU, 2021). To the southeast, the Scandinavian Mountains gradually become lower, transitioning into more gently undulating hilly terrain, reaching sea level by the Bothnian Bay (Fig. 1c). East of the Scandinavian Mountains, the till cover becomes thicker (Hirvas et al., 1988) and glaciofluvial deposits (e.g., eskers) are common (GTK, 2021).

Many of the lakes in the region owe their existence to the advances and retreats of the ice sheet and glaciers during the last glacial/interglacial cycle (Brosius et al., 2021). Lakes formed in depressions carved by the ice itself, some were dammed by glacial deposits, and still others are kettles formed due to melting of dead-ice. These processes combined resulted in a landscape with numerous lakes with diverse morphometrics. Many lakes and ponds are also associated with the minerotrophic aapa mires, characteristic for the area (Ruuhijärvi, 1983). These types of mire complexes form in flat areas with high precipitation and low evaporation (Seppä, 2002).

The water in the lakes can reflect different precipitation seasonality depending on duration of lake-ice cover, governing the amount of snowmelt bypass (i.e., how much of the snow that melts off the landscape before the lake becomes ice free; MacDonald et al., 2017). Along the Sweden-Finland border, the average duration of ice cover on the lakes (1961–2000) varies between 200 and 225 days (Laasanen, 1982; Eklund, 1999; Korhonen, 2006). Average lake freeze-up is around October 15 in northwestern Finland and November 5 in the southernmost part of the study area, whereas ice break-up occurs earlier in the

south (May 18) than in the north (June 15) (Laasanen, 1982; Korhonen, 2006). In Norway, north of the Arctic Circle, the duration of ice cover on 18 lakes was monitored for periods of 7–61 years (with an average observation period of 27 years) between 1912 and 1987. Ice-cover duration varies between 175 and 220 days with a mean of 198 days. The average freeze-up date for these lakes is November 10, and the average ice break-up date is May 29 (L'Abée-Lund et al., 2021). Air temperature is the most important factor for the timing of freeze-up and break-up (Palecki & Barry, 1986; Weyhenmeyer et al., 2004), modulated by regional atmospheric circulation (Blenckner et al., 2004). Small and shallow lakes generally freeze over earlier than large and deep lakes, but lake size has less influence on the break-up dates (Korhonen, 2006).

Most of the study area lies in the middle and northern boreal zones, with areas above the tree line classified as alpine (Moen, 1998). The middle boreal zone is characterized by coniferous forest and mires, dominating the southern part of the study area and found in the lowlands along parts of the Norwegian coast. Deciduous trees are also common. The central part of the study area belongs to the northern boreal zone, characterized by birch forests, low-growing coniferous forest, and mires. In mountainous areas, the northern boreal zone transitions into tundra. The low-alpine areas are characterized by continuous cover of dwarf-shrubs and mires, whereas grasses, sedges, mosses, lichens, and forbs dominate the mid-alpine zone. At higher elevations (the high-alpine zone), the vegetation cover is sparse. Plant cover can affect runoff into lakes, and therefore the seasonal lake water inflow values (Gibson & Edwards, 2002).

The topography contributes to a strong hydroclimate gradient, with the Scandinavian Mountains acting as a barrier for air masses from the North Atlantic. The study region is located in the polar front zone, where dynamic low-pressure systems bringing warm moist air masses from the Atlantic meet dry cool air from the Arctic. The prevailing westerly winds bring the humid Atlantic air onshore, and precipitation amounts are highest along the southwestern coast due to orographic lift where the westerlies first meet the mountains (Hanssen-Bauer and Førland, 2000). Most of the moisture falls on the windward side of the mountains, sheltering the leeward side of the mountains from precipitation (Uvo, 2003). The west coast experiences mild winters and cool summers,

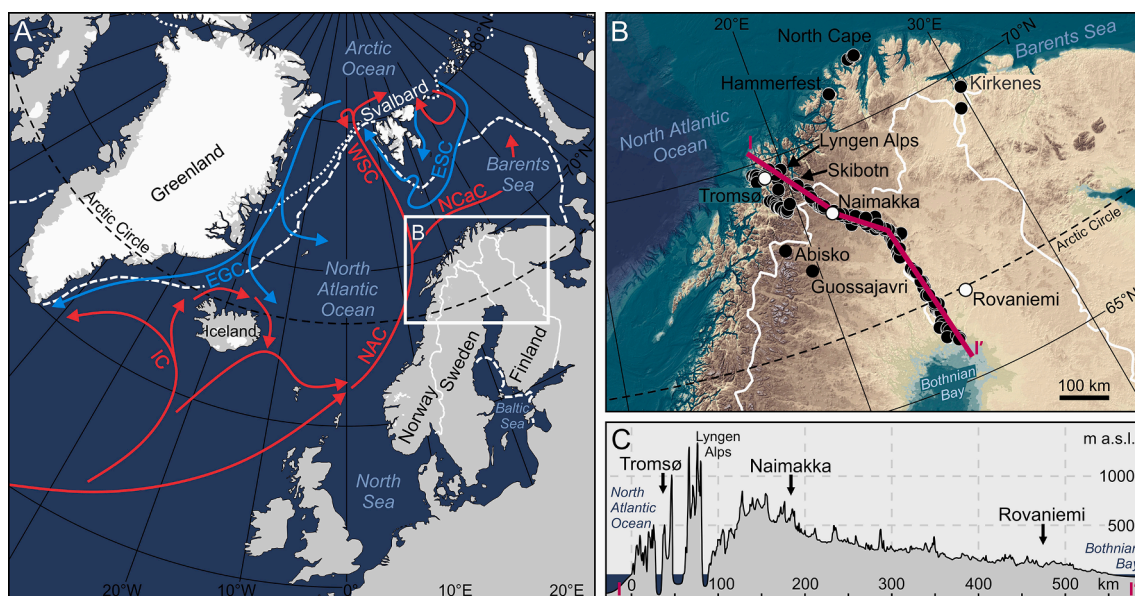


Fig. 1. (a) Overview map of the North Atlantic region, with major ocean surface currents (warm currents in red, cold in blue; NAC = North Atlantic Current; NCaC = North Cape Current; WSC = West Spitsbergen Current; IC = Irminger Current; ESC = East Spitsbergen Current; EGC = East Greenland Current), and median winter (white dashed line) and summer (white dotted line) sea-ice extent 1981–2010 (National Snow and Ice Data Center, 2019). (b) Map of the study area, showing locations of meteorological stations (white circles), sampled lakes and place names mentioned in the text (black circles) and extent of elevation profile in C (red line). Background map from IBCAO (Jakobsson et al., 2012). (c) Elevation profile from the North Atlantic Ocean in the northwest to the Bothnian Bay in the southeast. (For interpretation of the references to color in this figure legend, the reader is referred to the web version of this article.)

whereas the climate in the interior is more continental, with colder winters and warmer summers (Fig. 2a).

Seasonal variation in moisture source and moisture availability can be affected by changes in sea-ice cover. During winter (January–March), the Arctic sea ice reaches its maximum southward extent, covering the northern and eastern parts of the Barents Sea (National Snow and Ice Data Center, 2019; Fig. 1a). In winters with low sea-ice extent in the Barents Sea, more Arctic moisture can reach northern Fennoscandia (Bailey et al., 2021). Sea ice has not formed along the Norwegian coast during the Holocene (Belt et al., 2015) except in inner fjords, where it still forms today (O'Sadnick et al., 2020). Seasonal ice cover in the Bothnian Bay lasts 130–200 days, with sea ice forming in the shallow

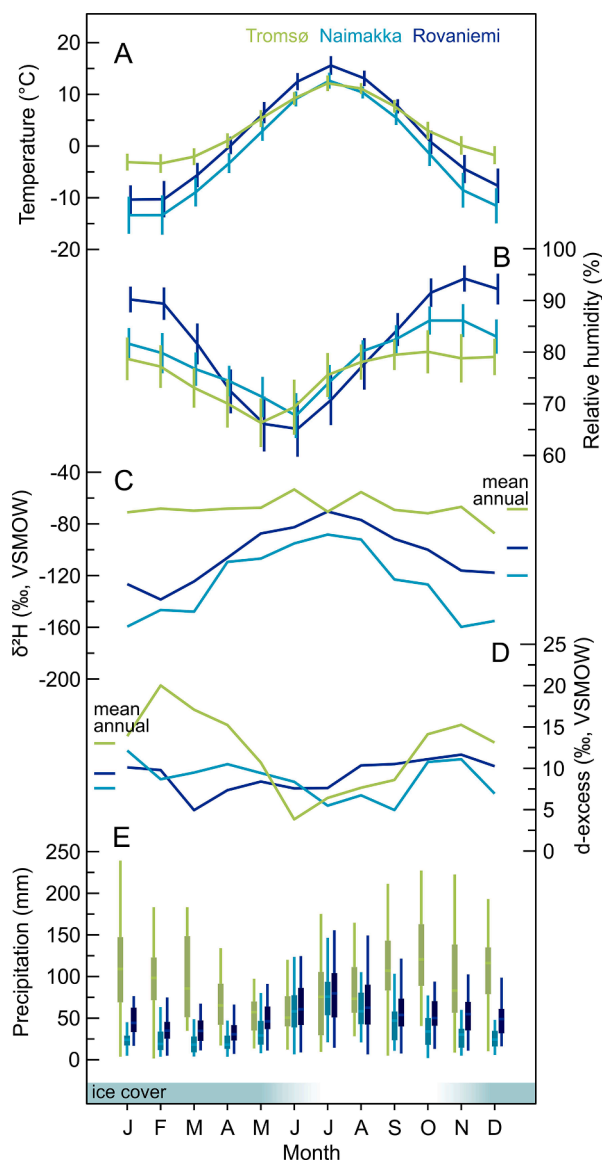


Fig. 2. Climate data from Tromsø, Naimakka and Rovaniemi (for locations, see Fig. 1). Mean monthly air temperatures (a) and relative humidity (b) between 1991–2020 (for Naimakka calculated using all months with >80% data coverage, 1995–2020). Vertical bars represent one standard deviation. (c–d) Amount-weighted monthly $\delta^2\text{H}$ and d-excess in precipitation from Tromsø (2019–2021; this study), and Naimakka (1990–1995) and Rovaniemi (2003–2014), both from IAEA/WMO (2019). (e) Monthly accumulated precipitation, 1991–2020. For each box plot, the middle line displays the median precipitation, the box represents the 25% to 75% quartile range and whiskers the maximum and minimum values. Temperature and precipitation data are retrieved from FMI (2021), MET Norway (2021) and SMHI (2021).

coastal areas in October–November and reaching a maximum ice extent in March (Haapala et al., 2015).

Winter precipitation is also strongly influenced by the North Atlantic Oscillation (NAO), controlling the strength of the westerlies (Hurrell, 1995; Uvo, 2003; Irannezhad et al., 2014). During times of positive NAO index (large atmospheric pressure difference between the Iceland Low and the Azores High), the North Atlantic storm track shifts northwards, increasing the moisture flux to Scandinavia (Hurrell, 1995; Marshall et al., 2001; Uvo, 2003). The NAO is closely related to the Arctic Oscillation (AO), which is linked to the polar low-pressure system, also known as the polar vortex (Thompson & Wallace, 1998). In the winter of 2019/2020 (January–March), which preceded one of our sampling campaigns, an exceptionally strong stratospheric polar vortex related to extreme AO+ conditions (lower than average air pressure over the Arctic and higher than average pressure over the northern Atlantic and Pacific Oceans) caused strong positive temperature (warmer) and precipitation (wetter) anomalies. The highest precipitation anomalies were observed between 60 and 70°N, greatly affecting the Norwegian coast (Lawrence et al., 2020). In contrast, some extreme snowfall events can be attributed to cold air outbreaks during strongly negative NAO and AO, when dry polar air from sea-ice covered areas moves in over a relatively warm ocean, creating large sea-air heat and moisture exchange (Shapiro et al., 1987; Papritz & Sodemann, 2018). One example of an event like this occurred in February 2018, when anomalously high temperatures and low sea-ice extent in the Barents Sea caused a high net moisture flux to the atmosphere and extreme snowfall over northern Europe (Bailey et al., 2021).

In summer, the atmospheric circulation patterns are generally not as strong, and more precipitation forms due to local convection (Jaagus, 2009). Notably, a recent Pan-Arctic study of the isotopic composition of summer 2018 rainfall identified precipitation air masses in Finland originating from two different transport regimes (Mellat et al., 2021). During NAO+ conditions, prevailing westerlies supplied Atlantic moisture, whereas summer cyclone activity related to a deepening low-pressure system over the Kara Sea caused northerly airflow and moisture transport from the Barents Sea (Mellat et al., 2021). These seasonal and synoptic changes in source impact the precipitation isotopic composition in this region (see section 2.1).

At the meteorological station in Tromsø (69.65°N, 18.94°E; 100 m a. s.l.; Fig. 1), the mean annual air temperature (MAAT, 1991–2020) is 3.3 °C and mean annual precipitation (MAP, 1991–2020) is 1091 mm (MET Norway, 2021). At the two more inland sites Naimakka (68.68°N, 21.53°E; 403 m a.s.l.) and Rovaniemi (66.50°N, 25.75°E; 107 m a.s.l.), MAAT is –1.6 °C and 1.5 °C, and MAP 466 mm and 633 mm, respectively (FMI, 2021; SMHI, 2021). Naimakka is the most continental (coldest and driest) of these sites, located on the eastern side of the Scandinavian Mountains (Fig. 1). Rovaniemi is farther from the North Atlantic Ocean, but experiences a warmer and wetter climate than Naimakka, perhaps due to its proximity to the Bothnian Bay. Significant differences in precipitation amount between the coastal and inland sites occur in fall and winter (Fig. 2e). This is the season when the relative temperature difference between (warm) ocean and (cold) land is greatest, causing high precipitation amounts at coastal locations such as Tromsø. In Rovaniemi and Naimakka, the relative humidity is high during the fall and winter months (Fig. 2b), and most precipitation falls during the warm months (Fig. 2e).

2.1. Isotopic setting

Northern Fennoscandia receives moisture from sources with different isotopic composition, where Arctic and Baltic sea surface waters are relatively depleted compared to the predominant North Atlantic source region, and the western Barents Sea (LeGrande & Schmidt, 2006; Bonne et al., 2019). In our study area, precipitation $\delta^2\text{H}$ and $\delta^{18}\text{O}$ have been monitored in Naimakka (1990–1995) and Rovaniemi (2003–2014) as part of the Global Network of Isotopes in Precipitation (GNIP; IAEA/

WMO, 2019; Fig. 1b). These data yield LMWLs with the equations $\delta^2\text{H} = 7.61 \times \delta^{18}\text{O} + 2.59$ (Naimakka) and $\delta^2\text{H} = 7.70 \times \delta^{18}\text{O} + 5.01$ (Rovaniemi; Fig. 3), having slightly lower slopes than the GMWL and lower intercepts. Precipitation is most depleted at Naimakka, where amount-weighted mean annual precipitation $\delta^2\text{H}$ and $\delta^{18}\text{O}$ are -119.94‰ and -16.04‰ , respectively (Figs. 2 and 3). The mean isotopic composition of precipitation $\delta^2\text{H}$ and $\delta^{18}\text{O}$ during the ice-free season (June–October) are -102.38‰ and -13.70‰ , and during the ice-cover season (November–May) -144.14‰ and -19.40‰ . In Rovaniemi, amount-weighted mean annual precipitation $\delta^2\text{H}$ and $\delta^{18}\text{O}$ are -98.56‰ and -13.52‰ , whereas ice-free season mean values are -83.31‰ and -11.59‰ and ice-cover season mean values -113.43‰ and -15.35‰ , respectively. The amount-weighted mean annual values in Rovaniemi are close to Naimakka mean summer (ice-free season) values, and 21.38‰ ($\delta^2\text{H}$) and 2.52‰ ($\delta^{18}\text{O}$) enriched compared to Naimakka amount-weighted annual values.

For regions with no GNIP stations nearby, interpolated precipitation isotope values can be generated using the Online Isotopes in Precipitation Calculator (OIPC; Bowen, 2021). However, these values can be biased towards the closest GNIP station values, not capturing local variability. Balascio et al. (2018) noted that water from lakes close to Tromsø was significantly enriched compared to both annual and summer OIPC estimations, indicating that the OIPC generates too depleted values for Tromsø, located at lower elevation and more coastal than the nearest GNIP stations. To evaluate the modern isotopic range of precipitation by the coast, we analyzed precipitation samples collected in Tromsø (69.68°N , 18.96°E ; 73 m a.s.l.) between September 6, 2019, and September 5, 2021 ($n = 407$). These data yield a LMWL with the equation $\delta^2\text{H} = 6.68 \times \delta^{18}\text{O} - 0.85$, having a lower slope and intercept than Rovaniemi and Naimakka (Fig. 3). The amount-weighted mean annual precipitation $\delta^2\text{H}$ and $\delta^{18}\text{O}$ values in Tromsø are -68.61‰ and -10.20‰ , respectively (Figs. 2 and 3). This is 42.39‰ and 4.70‰ enriched compared to the OIPC estimated mean annual values (-111‰ and -14.9‰) for $\delta^2\text{H}$ and $\delta^{18}\text{O}$ at this site, respectively (Bowen, 2021). The mean annual precipitation values in Tromsø are 29.95‰ ($\delta^2\text{H}$) and 51.33‰ ($\delta^{18}\text{O}$) enriched compared to

Rovaniemi and Naimakka, respectively. Mean ice-free season precipitation $\delta^2\text{H}$ and $\delta^{18}\text{O}$ values in Tromsø are -65.23‰ and -9.19‰ , and ice-cover season mean values are -70.63‰ and -10.81‰ (Fig. 3).

3. Materials and methods

3.1. Water sample collection and isotope analysis

We collected lake water samples from a total of 135 lakes in northern Norway, Finland, and Sweden (Fig. 1, Supplementary Information) between 2018 and 2020. Most of the samples were collected during the first week of July, in 2018, 2019 and 2020. This is after the snowmelt season, although some high elevation lakes are fed by snowmelt throughout summer. In July 2018, we sampled lakes in the coastal Tromsø area ($n = 30$), and at the end of June 2019 along the north Norwegian coast ($n = 9$). The July 2019 and 2020 field campaigns targeted lakes ($n = 35$ and $n = 68$, respectively) along a NW–SE transect from southeast of the Lyngen Alps to the Bothnian Bay (hereafter referred to as the ‘transect lakes’), between 69.4°N , 20.3°E and 65.7°N , 24.6°E (Fig. 1b–c). This transect spans c. 460 km , with lakes spanning elevations from close to sea level in NW and SE, up to c. 560 m a.s.l. close to the tripoint between Norway, Finland, and Sweden (Fig. 1c). We also collected samples from the Muonio and Torne Rivers (two rivers forming the border between Sweden and Finland) and the Bothnian Bay. To evaluate the interannual variability, we sampled 12 of the lakes along the transect and two locations on the Muonio River for two consecutive years. Furthermore, 11 of the transect lakes were sampled in July 2013 by Balascio et al. (2018), and five have been sampled all three years (2013, 2019 and 2020). Additionally, a couple of the coastal lakes were resampled in July 2019 and 2020 ($n = 2$ and $n = 4$, respectively) and/or during other times of the year ($n = 5$; Supplementary Information). Two coastal lakes were sampled for the first time in 2020. We collected the samples from the shoreline, approximately 10 cm below the lake surface to avoid evaporatively enriched water and stored them in 4 mL glass vials with no headspace. The vials were sealed with Parafilm to prevent evaporation during transportation and storage.

Precipitation samples were collected in Tromsø (69.68°N , 18.96°E ; 73 m a.s.l.) from September 6, 2019, to September 5, 2021 ($n = 407$). Samples were collected each morning, using a 4-inch diameter RG202 Stratus All-Weather Rain Gauge. Liquid precipitation was collected using a funnel and an inner 1-inch cylinder, and solid precipitation from the 4-inch cylinder. The samples were transferred to 4 mL glass vials, sealed with Parafilm, and kept at 4°C until analysis. Frozen samples were brought inside and melted in a sealed container at room temperature before transfer. Prior to analysis, all samples (lake water and precipitation) were filtered through a $0.2\text{-}\mu\text{m}$ PTFE filter.

The lake water samples collected in 2018 and 2019 were measured on a Picarro L2130-i WS-CRDS analyzer with vaporization module V1102-i coupled to a CTC PAL autosampler, at the Organic and Stable Isotope Biogeochemistry Laboratory at the University at Buffalo, USA. Each sample was measured four times, and the first injection discarded. Calibration to the VSMOW scale was done using three in-house standards, spanning a range of -265.86 to $+21.43\text{‰}$ for $\delta^2\text{H}$ and -33.62 to $+12.41\text{‰}$ for $\delta^{18}\text{O}$, calibrated against primary standards (GISP, VSMOW2, and SLAP2) from the International Atomic Energy Agency (IAEA). A memory correction was applied, following van Geldern & Barth (2012). Average standard deviations of replicate measurements were 0.08‰ and 0.02‰ for $\delta^2\text{H}$ and $\delta^{18}\text{O}$, respectively. For lake water samples collected in 2020, and for all precipitation samples, $\delta^2\text{H}$ and $\delta^{18}\text{O}$ were measured on a Picarro L2140-i WS-CRDS with a A0211 vaporizer and an A0325 autosampler at the Facility for advanced isotopic research and monitoring of weather, climate, and biogeochemical cycling (FARLAB), University of Bergen, Norway. Each sample was measured 12 times, and the reported values are averaged over the last four to six injections. Calibration was done using a drift standard and two laboratory

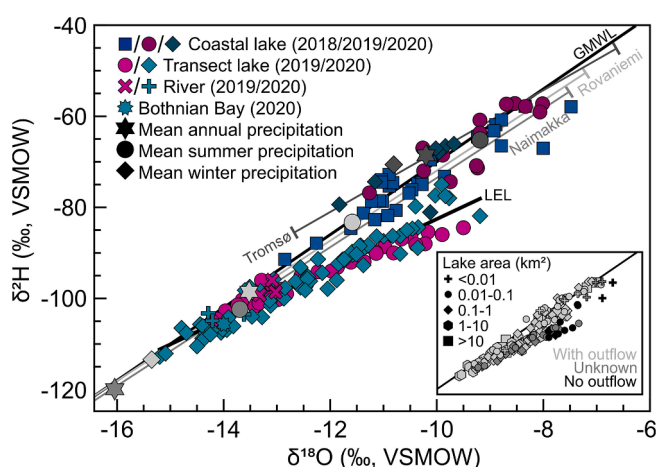


Fig. 3. $\delta^{18}\text{O}$ versus $\delta^2\text{H}$ values for all lakes and rivers sampled from 2018 to 2020, plotted against the global meteoric water line (GMWL, $\delta^2\text{H} = 8 \times \delta^{18}\text{O} + 10$) and local meteoric water lines (LMWLs) from Tromsø ($\delta^2\text{H} = 6.68 \times \delta^{18}\text{O} - 0.85$), Naimakka ($\delta^2\text{H} = 7.61 \times \delta^{18}\text{O} + 2.59$) and Rovaniemi ($\delta^2\text{H} = 7.70 \times \delta^{18}\text{O} + 5.01$). Amount-weighted mean annual, mean ice-cover (November–May = winter) and mean ice-free season (June–October = summer) precipitation are displayed for the three stations in grey. Mean ice-cover season precipitation $\delta^{18}\text{O}$ and $\delta^2\text{H}$ for Naimakka are too depleted (-19.40‰ and -144.14‰ , respectively) to be displayed in the plot. Local evaporation line (LEL) is included for the transect lakes ($\delta^2\text{H} = 5.50 \times \delta^{18}\text{O} - 27.55$). Inset panel has the same axes as the main figure and shows values by lake size (symbol) and throughflow regime (color).

standards, spanning a range of -308.14 to $+9.2‰$ for $\delta^2\text{H}$ and -40.06 to $+1.77‰$ for $\delta^{18}\text{O}$, calibrated against VSMOW2 and SLAP2. To reduce memory effects, a 2-component memory correction was applied, following Gröning (2011). Average standard uncertainties for calibrated measurement values (provided according to Gröning, 2018) were $0.43‰$ and $0.06‰$ for $\delta^2\text{H}$ and $\delta^{18}\text{O}$, respectively.

Isotope ratios are reported in per mil (‰) relative to Vienna Standard Mean Ocean Water (VSMOW), where $\delta = 1000 \times ((R_{\text{sample}}/R_{\text{VSMOW}}) - 1)$ and R is the ratio between $^2\text{H}/^1\text{H}$ or $^{18}\text{O}/^{16}\text{O}$. Deuterium excess (d-excess) was calculated as $d\text{-excess} = \delta^2\text{H} - 8 \times \delta^{18}\text{O}$ (Dansgaard, 1964). The daily precipitation data were converted into monthly amount-weighted mean values using the equation $\delta_{\text{aw}} = \frac{1}{P} \times \sum_{i=1}^n (P_i \times \delta_i)$, where P and P_i are the monthly and daily precipitation amounts (in mm), δ_i is $\delta^2\text{H}$ or $\delta^{18}\text{O}$ and n is the number of sampling days.

3.2. Inferring inflow $\delta^2\text{H}$ values

To estimate lake inflow $\delta^2\text{H}$ and $\delta^{18}\text{O}$ values for the evapo-concentrated transect lakes, we applied the Bayesian ‘MWL source implementation’ method by Bowen et al. (2018). This method allows the user to supply a MWL equation and a hypothesized LEL slope, each with confidence intervals. Since the LMWLs in Naimakka and Rovaniemi are parallel to and have similar intercepts as the GMWL (Fig. 3), we used the default MWL equation provided with the model, which is a re-calibrated GMWL ($\delta^2\text{H} = 8.01 \times \delta^{18}\text{O} + 9.57$; $n = 80,672$; Bowen et al., 2018). The GMWL has much less uncertainty than the LMWLs, which are constrained by only hundreds of data points. To allow for interannual

comparison, we based the LEL estimations on linear regression using the isotopic composition of lakes sampled in both 2019 and 2020 ($n = 12$). To run the model, we supplied the estimated LEL slope for each subset (4.82 and 5.18 for 2019 and 2020, respectively) with a standard deviation of 0.5 (as suggested by Bowen et al., 2018), lake water $\delta^2\text{H}$ and $\delta^{18}\text{O}$ with uncertainties and covariance (0.95), and the number of iterations ($10,000$). We acknowledge the assumptions involved in a regression-based approach for LEL slope estimation, but consider the supplied LEL slopes to be a good starting estimate, because they agree with theoretical LEL slope values (~ 5) modeled by Gibson et al. (2008). Given uncertainties in the LEL regression approach, we do not interpret interannual or spatial differences in LELs, and instead use them as a guide to infer whether lakes have experienced evaporative enrichment. We present modeled mean source water values, hereafter referred to as inferred inflow $\delta^2\text{H}$ ($\delta^2\text{H}_i$). For the discussion where we compare the seasonality of inflow between coastal and transect lakes, we include only the coastal lakes that are through-flowing, as lake water $\delta^2\text{H}$ ($\delta^2\text{H}_i$) in these lakes is not significantly affected by evaporative enrichment, and therefore equal to $\delta^2\text{H}_i$. We exclude small isolated coastal basins, since we do not have a well-constrained estimate of the LEL by the coast.

4. Results

4.1. Lake water isotopic composition

The lake water surface samples from 2018 to 2020 span a broad range of isotopic compositions (Fig. 3). Generally, lakes close to the

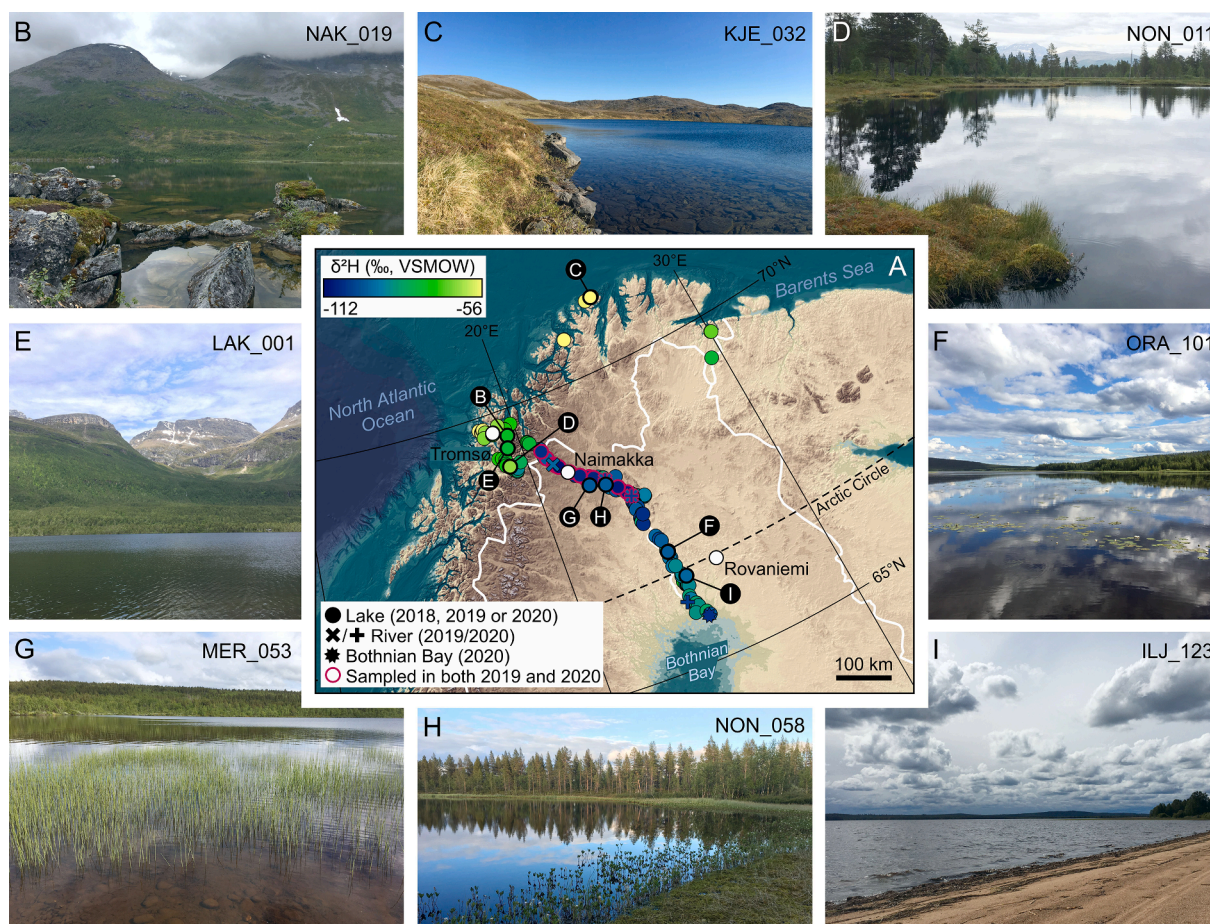


Fig. 4. (a) Geographical distribution of surface water $\delta^2\text{H}$ for 135 lakes, two rivers and the Bothnian Bay, in Arctic and sub-Arctic Norway, Finland, and Sweden. Twelve lakes and two locations along the Muonio River sampled in both 2019 and 2020 are outlined in red. Background map from IBCAO (Jakobsson et al., 2012). (b-i) Photographs of a selection of the sampled lakes. Codes refer to lake IDs (Supplementary Information). (For interpretation of the references to color in this figure legend, the reader is referred to the web version of this article.)

Norwegian (Atlantic) coast lie on or close to the GMWL, with a trend toward more ²H-depleted values moving inland (Figs. 4–7). This trend is likely controlled by the increasing distance from the coast, which is likely a proxy for the degree of distillation resulting from air masses advecting over the mountain peaks (Fig. 5a and 7). The lake elevation, which is often below the highest mountain peaks in the region (Fig. 5b), is less important than the elevation of the mountains over which the moisture has to pass along its transport path (Fig. 7). In the following paragraphs we present the measured lake water δ²H and δ¹⁸O values,

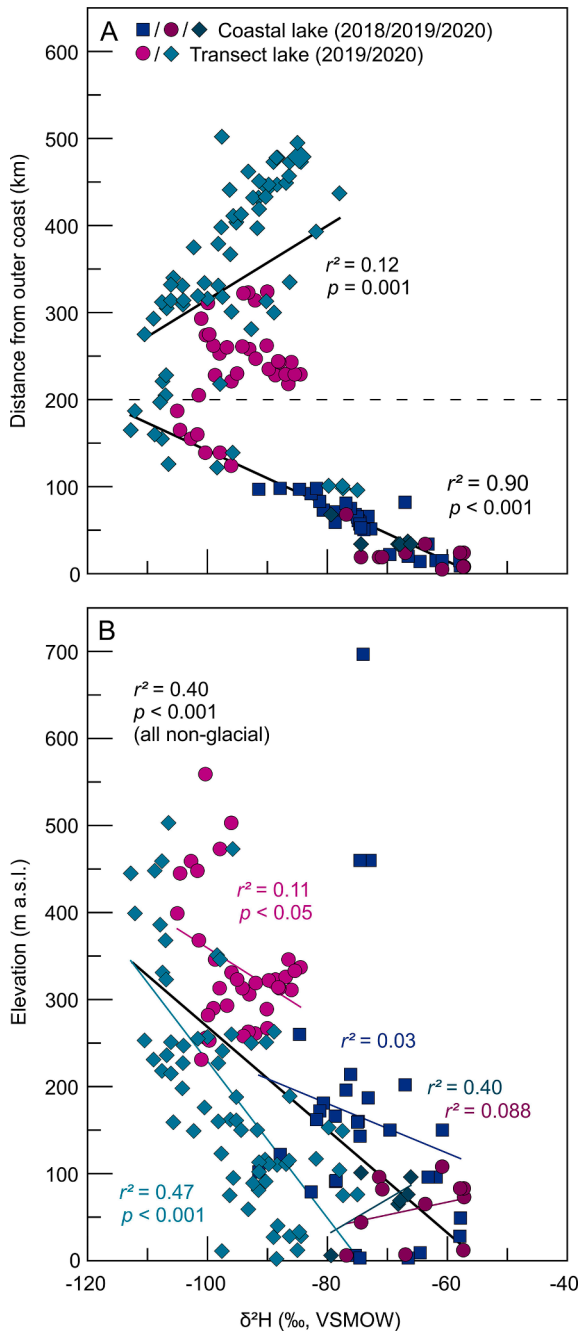


Fig. 5. Spatial trends in lake water δ²H. (a) δ²H versus distance from the outer coast (North Atlantic Ocean or Barents Sea). Linear regression fits are included for all lakes (2018, 2019 and 2020 combined) located less and >200 km from the outer coast (distance = $-3.18 \times \delta^2\text{H} - 176.19$ and distance = $4.25 \times \delta^2\text{H} + 739.39$, respectively), separated by dashed line. (b) δ²H versus elevation, with linear regressions for each subset of coastal and transect lakes (2018, 2019 and 2020), as well as for all non-glacial lakes (excluding three high-elevation lakes (>460 m a.s.l.) and one lake close to sea level receiving glacial meltwater).

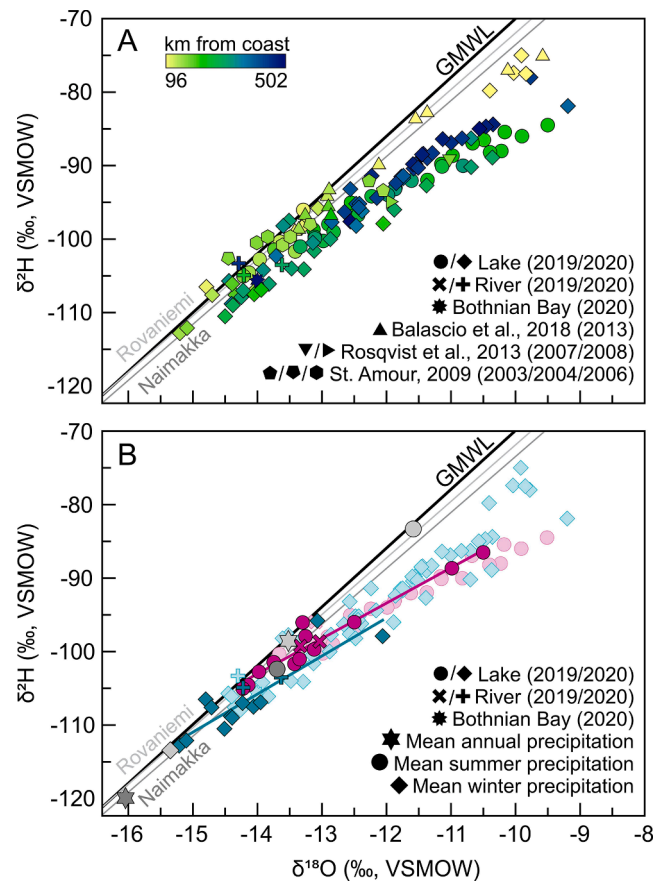


Fig. 6. δ¹⁸O versus δ²H values for transect lakes and rivers sampled in 2019 and 2020, colored by distance to the North Atlantic coast (a), and by sampling year (b). (a) includes lakes sampled in the study area by St. Amour (2009) and Balascio et al. (2018), and lake surface water (depth ≤ 1 m) from Guossajavri (Rosqvist et al., 2013; Fig. 1). Dark symbols in (b) show lakes ($n = 12$) and locations along the Muonio River ($n = 2$) sampled both year, and light symbols are all samples. Local evaporation lines (LELs) for 2019 ($\delta^2\text{H} = 4.82 \times \delta^{18}\text{O} - 35.58$) and 2020 ($\delta^2\text{H} = 5.18 \times \delta^{18}\text{O} - 33.27$) are included for the 12 resampled lakes. Local meteoric water lines and amount-weighted mean annual and seasonal precipitation values (winter = November–May; summer = June–October) for Naimakka and Rovaniemi are displayed in grey.

starting with the coastal lakes, followed by the transect lakes.

All lakes along the outermost coast plot in the upper right corner of the co-isotope plot (Fig. 3). In 2018, δ²H and δ¹⁸O of 30 coastal lakes ranged from -91.43 to -57.82‰ (median = $-74.72 \pm 8.36\text{‰}$ ($\pm 1\sigma$)) and -12.85 to -7.48‰ (median = $-10.64 \pm 1.23\text{‰}$), respectively. Deuterium excess ranged from -3.00 to $+14.37\text{‰}$ (median = $9.10 \pm 3.59\text{‰}$). In 2019, δ²H and δ¹⁸O of 11 coastal lakes ranged from -76.84 to -57.22‰ (median = $-63.69 \pm 7.11\text{‰}$) and -11.27 to -8.02‰ (median = $-9.19 \pm 0.88\text{‰}$), respectively. For these lakes, d-excess ranged from 2.52 to 15.12‰ (median = $9.85 \pm 4.20\text{‰}$). δ²H and δ¹⁸O of six coastal lakes sampled in 2020 ranged from -79.40 to -66.10‰ (median = $-68.05 \pm 4.85\text{‰}$) and -11.83 to -9.68‰ (median = $-10.07 \pm 0.78\text{‰}$), respectively. Deuterium excess ranged from 11.30 to 15.20‰ (median = $12.65 \pm 1.49\text{‰}$). For lakes sampled along the NW–SE transect in 2019 ($n = 35$), δ²H and δ¹⁸O ranged from -105.04 to -84.49‰ (median = $-95.05 \pm 5.82\text{‰}$) and -14.22 to -9.50‰ (median = $-12.44 \pm 1.32\text{‰}$), respectively (Figs. 3–7). Deuterium excess of these lakes ranged from -8.47 to $+10.28\text{‰}$ (median = $3.55 \pm 4.97\text{‰}$). For transect lakes sampled in 2020 ($n = 68$), δ²H and δ¹⁸O ranged from -112.80 to -75.00‰ (median = $-95.75 \pm 9.35\text{‰}$) and -15.20 to -9.19‰ (median = $-12.43 \pm 1.51\text{‰}$), respectively. Deuterium excess ranged from -8.40 to 11.80‰ (median = $3.25 \pm 3.84\text{‰}$). All transect lakes plot along a LEL ($\delta^2\text{H} = 5.50 \times \delta^{18}\text{O} -$

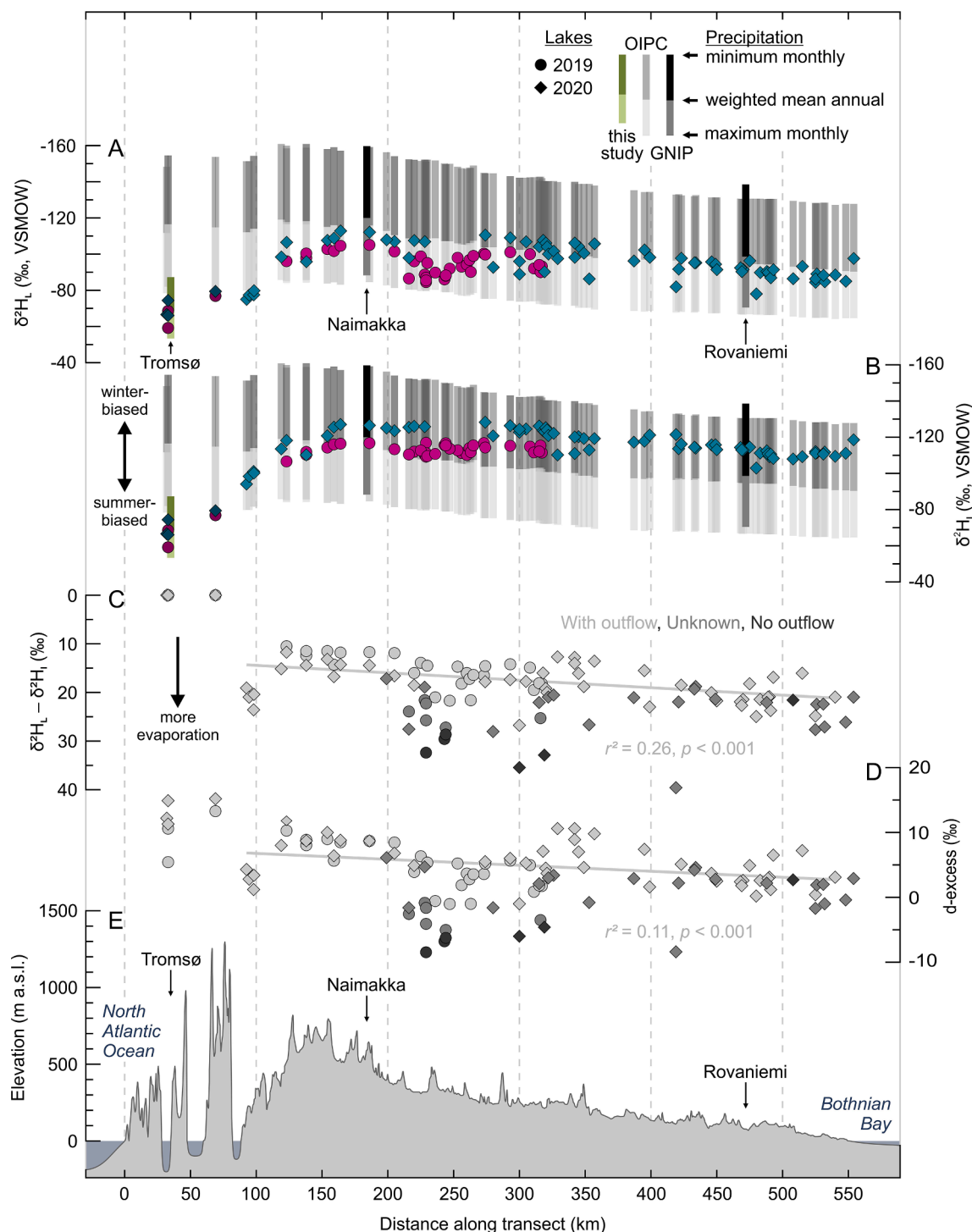


Fig. 7. Lake water $\delta^2\text{H}_L$ (a), inferred inflow $\delta^2\text{H}_I$ (b), offset between $\delta^2\text{H}_L$ and $\delta^2\text{H}_I$ (c), d-excess (d), and elevation (e) for all lakes along transect (colors and shapes as in Fig. 3). Four coastal lakes along an extension of the transect (NW of Skibotn; Fig. 1) are plotted for comparison. Vertical bars in (a-b) represent the ranges between weighted mean annual and maximum and minimum monthly precipitation $\delta^2\text{H}$ in Tromsø (this study), Naimakka and Rovaniemi (GNIP; IAEA/WMO, 2019), and for the location of each lake, modeled using the Online Isotopes in Precipitation Calculator (OIPC; Bowen, 2021). Symbols in (c-d) are colored according to throughflow regime (with outflow, without outflow, or unknown), and linear regression fits are included for all through-flowing lakes ($\delta^2\text{H}_L - \delta^2\text{H}_I = 0.015 \times \text{distance} + 12.89$; d-excess = $-0.0092 \times \text{distance} + 7.69$). Note that the y-axes in (a-c) are inverted.

27.55; Fig. 3), a further indication that some of the lakes undergo evaporation.

Transect lakes sampled in both 2019 and 2020 ($n = 12$) generally showed similar interannual variability (Fig. 6b and 8). Samples collected in 2020 were ^2H -depleted relative to samples collected from the same lakes in 2019 (Fig. 6b and 8) but had similar d-excess both years. The

average offset (2020 minus 2019) was -8.43‰ for $\delta^2\text{H}$ and $+0.90\text{‰}$ for d-excess. One large lake (Kilpisjärvi; 69.05°N , 20.79°E , 37 km^2) was slightly ^2H -enriched in 2020 compared to 2019, with an offset of $+2.12\text{‰}$ for $\delta^2\text{H}$ and $+0.66\text{‰}$ for d-excess (Fig. 8a). Applying linear regression fits to these 12 lakes resulted in LELs with the equations $\delta^2\text{H} = 4.82 \times \delta^{18}\text{O} - 35.58$ and $\delta^2\text{H} = 5.18 \times \delta^{18}\text{O} - 33.27$, for 2019 and 2020

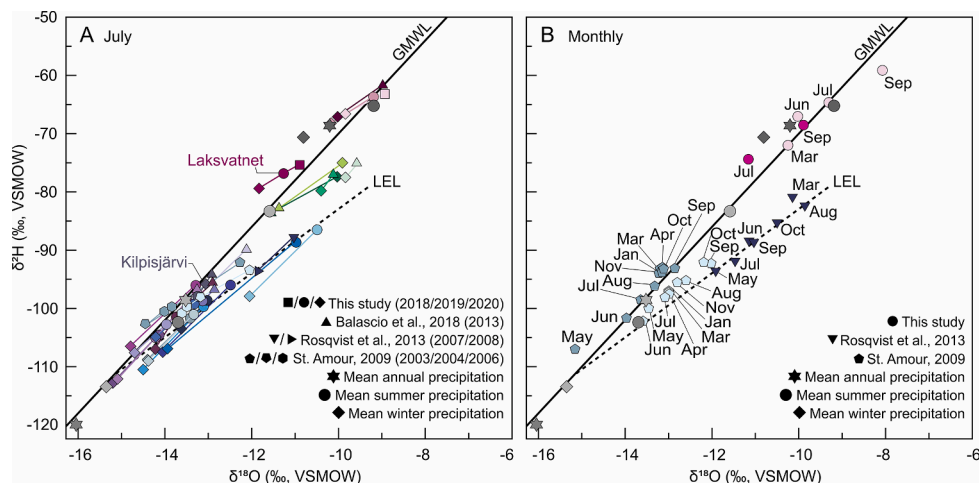


Fig. 8. (a) Interannual variability in summer (July) lake water $\delta^2\text{H}$ and $\delta^{18}\text{O}$ from this study (2018–2020), lakes sampled in the same area (St. Amour, 2009; Balascio et al., 2018) and lake Guossajavri (Rosqvist et al., 2013; 125 km SW of our transect). Lakes water samples collected from the same lakes in July different years (from 2013 to 2020) are illustrated in the same color and connected with lines. Coastal lakes are colored pink, lakes in the Skibotn valley are green, and transect lakes are purple and blue. (b) Monthly average isotopic composition of lakes sampled 1–4 years from 2002 to 2020. All samples were taken at the surface (this study: c. 10 cm below the lake surface) from the shoreline, or from in the middle of the lake (depth ≤ 1 m). Samples from inlets were excluded. Amount-weighted mean annual and seasonal precipitation values (winter = November–May; summer = June–October) are displayed in grey (dark grey = Tromsø; middle grey = Naimakka; light

grey = Rovaniemi), and black dashed line represents the local evaporation line (LEL; $\delta^2\text{H} = 5.50 \times \delta^{18}\text{O} - 27.55$) for transect lakes in this study. For locations, see Fig. 1. (For interpretation of the references to color in this figure legend, the reader is referred to the web version of this article.)

respectively (Fig. 6b). Two locations along the Muonio River exhibited similar interannual variation as the lakes (Fig. 6b), indicating different inflow values in 2019 and 2020. In 2019, the river water had $\delta^2\text{H}$ of -98.62 and -99.24 ‰, respectively. In 2020, the same two sites along the river had $\delta^2\text{H}$ of -103.50 and -104.90 ‰, resulting in an average offset of -5.27 ‰. The average offset for d-excess was $+0.69$ ‰. One sample collected from the Bothnian Bay in 2020 had $\delta^2\text{H}$ and $\delta^{18}\text{O}$ of -105.60 and -14.00 ‰, respectively, reflecting the high river-runoff to this northernmost part of the Baltic Sea (Figs. 3, 4 and 6).

The four northernmost lakes in the transect (located in Skibotn; Fig. 1b) had relatively enriched $\delta^2\text{H}$ and $\delta^{18}\text{O}$ values compared to the rest of the transect lakes (Figs. 4, 8 and S1). $\delta^2\text{H}$ and $\delta^{18}\text{O}$ ranged from -79.80 to -75.00 ‰ (median = -77.45 ± 1.70 ‰) and from -10.40 to -9.84 ‰ (median = -9.97 ± 0.22 ‰), respectively. Deuterium excess ranged from 1.20 to 4.30 ‰ (median = 3.10 ± 1.13 ‰). These four lakes are located within 9 km from the Lyngen fjord, and could therefore be considered coastal (Figs. 1 and 4). However, there is a strong rain-shadow effect in Skibotn, where the mean annual precipitation is 465 mm (MET Norway, 2021), compared to 1091 mm in Tromsø, located on the other side of the Lyngen Alps (Fig. 1c), c. 60 km northwest of Skibotn.

For lakes by the coast and along the beginning of the transect (<c. 200 km from the North Atlantic) there is a significant ($p < 0.001$) trend toward more ^2H -depleted lakes moving away from the coast, whereas farther inland, this trend is still significant ($p = 0.001$), though reversed (Fig. 5a). $\delta^2\text{H}$ also decreases with elevation (Fig. 5b). This trend is significant for the transect lakes ($p < 0.05$ and $p < 0.001$ in 2019 and 2020, respectively), but not for the coastal lakes. Considering the strong correlation between $\delta^2\text{H}$ and $\delta^{18}\text{O}$ for both coastal ($r^2 = 0.84$) and transect ($r^2 = 0.92$) lake samples, we assume that trends in $\delta^{18}\text{O}$ follow $\delta^2\text{H}$ and hereafter primarily focus on $\delta^2\text{H}$ and d-excess.

4.2. Inferred inflow $\delta^2\text{H}$ values

The conventional LEL-MWL intersection approach yielded $\delta^2\text{H}_i$ values for the transect lakes of -104.78 and -112.90 ‰ in 2019 and 2020, respectively. Bayesian inflow modeling, which we did for all 91 transect lakes, yielded $\delta^2\text{H}_i$ ranging from -116.87 to -106.57 ‰ (median = -112.72 ± 2.57 ‰) in 2019, and -128.27 to -94.08 ‰ (median = -115.75 ± 7.53 ‰) in 2020. This is on average 18.27‰ (2019) and 20.55‰ (2020) more ^2H -depleted than measured lake water $\delta^2\text{H}$ (Fig. 7). The 90% confidence intervals for $\delta^2\text{H}_i$ ranged from -148.44 to

-111.52 ‰ for the lower bound and from -118.11 to -83.00 ‰ for the upper bound, with an average of -132.30 and -104.26 ‰, respectively. For comparison, $\delta^2\text{H}_i$ for the coastal lakes ranged from -91.43 to -57.82 ‰ (median = -74.72 ± 8.36 ‰) in 2018, -76.84 to -57.22 ‰ (median = -63.69 ± 7.11 ‰) in 2019, and from -79.40 to -66.10 ‰ (median = -68.05 ± 4.85 ‰) in 2020.

In the area with lakes sampled in both 2019 and 2020 (<320 km from the North Atlantic), $\delta^2\text{H}_i$ values were ^2H -depleted in 2020 compared to in 2019. For the 12 resampled lakes, $\delta^2\text{H}_i$ ranged from -116.72 to -106.57 ‰ (median = -113.79 ± 2.97 ‰) in 2019, and -128.27 to -110.21 ‰ (median = -125.67 ± 4.87 ‰) in 2020, resulting in an average offset of -10.68 ‰ (Fig. 7). Kilpisjärvi had a similar, but slightly more negative inferred $\delta^2\text{H}_i$ value in 2019 (-110.30 ‰) compared to 2020 (-110.21 ‰), with an offset of $+0.09$ ‰. In 2020, the four lakes in Skibotn had $\delta^2\text{H}_i$ values from -101.06 to -94.08 ‰ (median = -99.27 ± 2.69 ‰), comparable to coastal $\delta^2\text{H}_i$, whereas the rest of the transect lakes had $\delta^2\text{H}_i$ ranging from -128.27 to -102.94 ‰ (median = -116.72 ± 6.23 ‰).

Moving inland, $\delta^2\text{H}_i$ decreased rapidly in the first c. 160 km along the transect, after which $\delta^2\text{H}_i$ stayed relatively stable, with a slightly increasing trend towards the Bothnian Bay. Generally, lakes closer to the North Atlantic had $\delta^2\text{H}_i$ closer to inferred $\delta^2\text{H}_i$, for through-flowing lakes $\delta^2\text{H}_i$ - $\delta^2\text{H}$ increased moving southeast along the transect ($+1.53$ ‰ per 100 km), and closed basins had the largest offsets between $\delta^2\text{H}_i$ and $\delta^2\text{H}$ (Fig. 7c).

5. Discussion

Lake water isotopes in our study area display mainly two kinds of variability, caused by differences in 1) inflow value (moving along the MWL) and 2) evaporation (moving below the MWL; Fig. 3). In most cases, lake water inflow values reflect a mixture of sources with different isotopic composition (e.g., summer and winter runoff, groundwater). The seasonality of lake water is primarily governed by the lake water residence time, since this determines the proportion of summer and winter precipitation in the lake (Jonsson et al., 2009). The position of a lake along the MWL and its relation to local precipitation isotopic composition can thus give information about lake water biases toward summer or winter season precipitation. In $\delta^2\text{H}$ - $\delta^{18}\text{O}$ space, lakes experiencing evaporative enrichment (decreasing d-excess) plot below the MWL, and along LELs. In the following sections, we discuss the sensitivity of modern lake water $\delta^2\text{H}$ along the climate gradient to differences

in inflow isotopic composition and evaporation, how these two controls vary in time and space, and why this variability needs to be considered when inferring lake-water-isotope proxy records.

5.1. Regional variability in lake water $\delta^2\text{H}$ and inferred inflow $\delta^2\text{H}$

5.1.1. Coastal lakes

The strong correlation between increasing distance from the coast and decreasing $\delta^2\text{H}_\text{L}$ ($r^2 = 0.90$, $p < 0.001$; Fig. 5a) for the lakes closest to the coast (<c. 200 km; both coastal and the northernmost part of the transect), suggests that distillation during atmospheric moisture transport, prior to precipitation, is the most important control on $\delta^2\text{H}_\text{L}$ for coastal lakes in this area. $\delta^2\text{H}$ and $\delta^{18}\text{O}$ decrease by c. 57‰ and 6.8‰ over 200 km, or 28‰ and 3.4‰ per 100 km, respectively (Fig. 5a). In the Pacific Northwest, USA (mountainous but drier than coastal Norway), the $\delta^{18}\text{O}$ isotopic gradient is 1.2‰ per 100 km (Welker, 2000) and in the Sierra Nevada, USA (wet year-round, most precipitation in winter, some mountains) 5‰ per 100 km (Ingraham & Taylor, 1991). Across the European Plain (from mild and wet coastal Ireland to the continental foothills of the Ural Mountains), the gradient is 0.2‰ per 100 km (Rozanski et al., 1993). This means that our gradient is within ranges found elsewhere and most similar to gradients in similar environments (wet, mountainous, coastal). The least ^2H -depleted lakes are the ones at the northern coast and the outermost coast to the west of Tromsø (Fig. 4), since the transport from the source is shorter and there is not much rain-out prior to precipitation. Three lakes in Kirkenes are slightly ^2H -depleted compared to the other lakes at the outermost coast (Fig. 1b and 4). None of these lakes have distinct outflows, and relatively low d-excess (Supplementary Fig. S1) suggests they are likely experiencing evaporative enrichment. They may also receive source water with a slightly different isotopic signal, being located near the Barents Sea. Lakes near Tromsø are relatively ^2H -enriched compared to lakes slightly farther from the coast (Fig. 4).

In the coastal region, most of the large through-flowing lakes plot close to the MWL, whereas some smaller lakes with low or unknown throughflow and long residence times fall below the MWL, being more affected by evaporation (Fig. 3). A couple of large through-flowing lakes with large mountainous catchments plot above the MWL and close to Tromsø amount-weighted winter precipitation (e.g., Laksvatnet; Fig. 4a and e, 8), suggesting a longer response time, and that the depleted meltwater from snow has not yet been flushed through in early July.

Some of the local variability in coastal lake water $\delta^2\text{H}$ can also be explained by differences in elevation (Fig. 5b), likely reflecting heterogeneous airflow patterns, with fjords funneling onshore winds and mountains causing orographic precipitation. The mountains cause the air masses to rise as they move inland from Tromsø, causing rainout and more negative $\delta^2\text{H}$ values (Figs. 4, 5 and 7). Lakes at a given distance from the coast have similar isotope values, despite differences in elevation (Fig. 5a-b). This is likely because precipitation at a given location forms at the same elevation in the atmosphere, and that elevation rises over the mountains moving inland. Three high elevation coastal lakes (>460 m a.s.l.) have $\delta^2\text{H}_\text{L}$ values comparable to other coastal lakes, despite their high elevation (Fig. 5b). They have high d-excess (average = 14.2 ± 0.3 ‰) and plot above the MWL, suggesting that their isotopic signals are biased toward winter precipitation. In Tromsø, precipitation $\delta^2\text{H}$ is relatively enriched compared to the more inland stations and does not show a strong seasonality. In contrast, d-excess is significantly higher in fall and winter than in summer (Figs. 2 and 3), due to kinetic fractionation effects when snow forms in mixed-phase clouds (vapor supersaturated over ice; Jouzel & Merlivat, 1984). It is common for mountain lakes to receive most precipitation as snowfall and for their catchments to experience late snowmelt, explaining the high d-excess lake water values. Furthermore, two of the lakes have small niche glaciers in their catchments, contributing glacial meltwater. Thus, our dataset suggests that coastal lakes are primarily influenced by varying degrees of distillation during atmospheric

moisture transport, precipitation amount and lake and catchment morphometry, the latter two determining the lake water residence time.

5.1.2. Transect lakes

In contrast to the coastal lakes, most of the transect lakes experience more evaporative enrichment, plotting below the MWL and along a LEL (Figs. 3 and 6). Lake surface water from Oikojärvi (located close to Naimakka; St. Amour, 2009) and Guossajavri (125 km SW of our transect; Rosqvist et al., 2013; Fig. 1b) sampled between 2002 and 2008 fall along similar LELs, suggesting similar evaporation patterns on a regional scale and between years (Figs. 6 and 8). The North Atlantic is still the main moisture source, so precipitation $\delta^2\text{H}$, and as a result lake water $\delta^2\text{H}$, becomes more and more depleted the first c. 160 km along the transect, reflecting Rayleigh distillation during moisture transport (Figs. 4–7).

On the southeastern side of the mountains, where the relief changes from highlands with incised valleys to a gently sloping plain with few hills, there is a larger spread in $\delta^2\text{H}_\text{L}$ (Fig. 7a) and larger and more variable offsets between $\delta^2\text{H}_\text{L}$ and $\delta^2\text{H}_\text{I}$ (Fig. 7c). This coincides with the northern boundary of the northern aapa mire zone, where large, branched mire systems are common (Ruuhijärvi, 1983). Although lakes in wetland-dominated catchments tend to have low E/I (Gibson et al., 2002), many of the lakes in this area are evaporatively enriched, plotting on a LEL and having low d-excess (Figs. 3, 6 and 7d). Hydrologically closed basins, lack of clear surface flow, and/or only seasonal connection by small streams or wetland areas may make these lakes susceptible to evaporation. Furthermore, in continental regions at high latitudes, seasonally arid conditions can increase the evaporative enrichment during the thaw season (Gibson et al., 2005). Smaller inter-lake variation in $\delta^2\text{H}_\text{I}$ than in $\delta^2\text{H}_\text{L}$ (Fig. 7a-c), suggests that most lakes have similar inflow seasonality and sources, whereas evaporation, indicated by decreasing d-excess (-0.92 ‰ per 100 km; Fig. 7d), increases moving southeast. Moreover, some lakes possibly have $\delta^2\text{H}_\text{L}$ values influenced by factors in addition to evaporative enrichment.

Some of the lakes might be groundwater fed, since till and glacio-fluvial aquifers are common in this region (Kortelainen & Karhu, 2004). However, there are no significant differences between the isotopic composition of precipitation and shallow groundwater (till aquifers) in northern Finland and Sweden (Kortelainen & Karhu, 2004). Both follow the same seasonal cycle with most enriched values in late fall, and most depleted values after snowmelt. We do therefore not expect shallow groundwater recharge to the lakes to significantly alter $\delta^2\text{H}_\text{L}$.

Lakes on the southeastern side of the Scandinavian mountains most likely also receive precipitation from other moisture sources, such as the Baltic region and Eurasia. These sources are more depleted (Schmidt et al., 1999; Bailey et al., 2021), but the travel distance may be shorter, potentially resulting in less distillation and precipitation that is relatively enriched compared to Atlantic moisture, a phenomenon also observed in precipitation events in Arctic Alaska (Putman et al., 2017). In summer, precipitation can be induced by local convection, and less influenced by atmospheric circulation (Jaagus, 2009). Local moisture could in this way at least partly explain the slightly enriched $\delta^2\text{H}_\text{L}$ of lakes closest to the Bothnian Bay (Fig. 7a). Furthermore, transpiration, another process most important in summer, can partly compensate for isotopic depletion caused by distillation during moisture transport, by recycling soil moisture without fractionation (Rozanski et al., 1982). The relative contribution of moisture from different source regions and the resulting isotopic compositions of precipitation can be assessed on an event basis at a given station (Bailey et al., 2021; Mellat et al., 2021), but we do not have any data on this over longer time scales (i.e., covering our study period), or know how it varies within our study region. Most likely, the isotopic composition of precipitation along our transect, and therefore lake water inflow, is controlled by a combination of different moisture source regions, local convection, and evapotranspiration.

5.1.3. Lake location and inflow seasonality

Comparing $\delta^2\text{H}$ to precipitation $\delta^2\text{H}$ data from Tromsø, Naimakka and Rovaniemi, as well as OIPC-modeled precipitation $\delta^2\text{H}$ (Bowen, 2021), suggests that winter precipitation is more important for inland lakes, whereas coastal lakes tend to be summer-biased or have an isotopic composition close to mean annual precipitation $\delta^2\text{H}$ (Fig. 7b). A

similar trend is seen on western Greenland, likely reflecting lake water inflow seasonality along an aridity gradient (Cluett & Thomas, 2020). In our study area, the inland region normally receives a greater fraction of the annual precipitation during the summer months (Fig. 2). This lake water bias towards winter precipitation for inland lakes is likely due to lower relative humidity and higher evapotranspiration at inland sites in

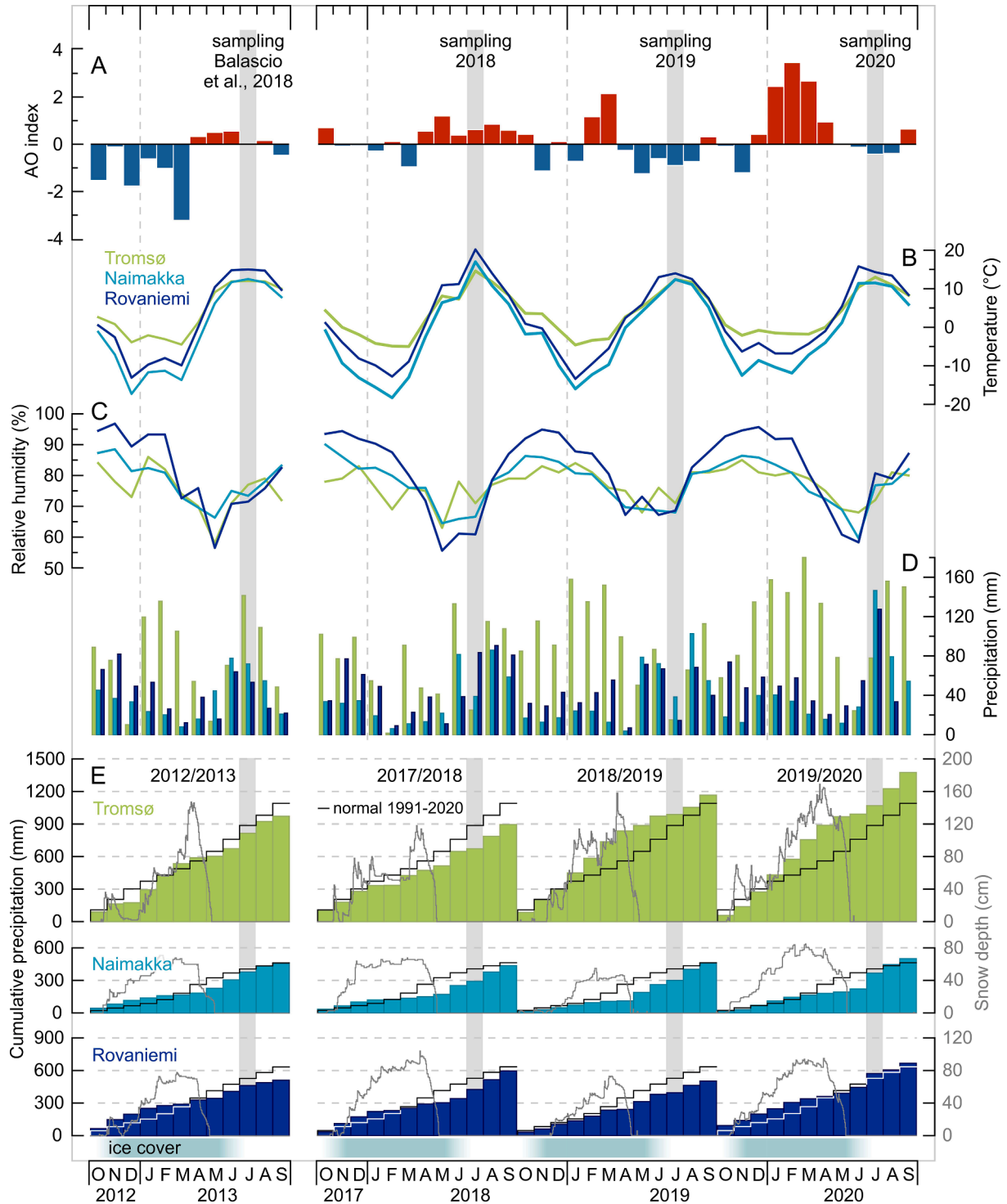


Fig. 9. (a) Monthly Arctic Oscillation (AO) index (National Weather Service Climate Prediction Center, 2021). (b) Mean monthly air temperature, (c) relative humidity, (d) accumulated precipitation and (e) cumulative precipitation and snow depths for the hydrological years (October-September) 2012/2013, 2017/2018, 2018/2019 and 2019/2020 for Tromsø, Naimakka and Rovaniemi (FMI, 2021; MET Norway, 2021; SMHI, 2021). Black stepped lines in (e) illustrate mean cumulative precipitation values for 1991–2020. Snow depths for Naimakka were obtained from Saarikoski (68.81°N, 21.25°E, 450 m a.s.l.), 18 km NW of Naimakka (SMHI, 2021).

summer (Fig. 2), even if a greater fraction of total precipitation falls in summer (Bowen et al., 2018).

That coastal lakes appear to be summer-biased is somewhat unexpected, considering that Tromsø receives more winter precipitation than Naimakka and Rovaniemi (Figs. 2 and 9d-e). OIPC-modeled precipitation $\delta^2\text{H}$ values are based on interpolation using GNIP values. They are therefore biased towards the nearest GNIP stations (i.e., Naimakka and Rovaniemi), which are more continental and experience a larger seasonality than Tromsø (Figs. 2 and 3). We compare the OIPC values with precipitation samples collected in Tromsø in 2019–2021 and can confirm that the modeled coastal values are, indeed, too depleted compared to our new precipitation isotope dataset (Fig. 7a-b). However, it cannot be excluded that the ^2H -depleted winter isotopes might have been efficiently flushed from some coastal lakes with short residence times, especially considering the high precipitation amounts in June 2018 and 2019 (Fig. 9). This suggests that lakes in this region likely reflect a range of summer to mean annual to winter-biased precipitation $\delta^2\text{H}$ (Figs. 3 and 7).

5.2. Lake water $\delta^2\text{H}$ and inferred inflow $\delta^2\text{H}$ changes over time

5.2.1. Interannual variability

Sampling campaigns concentrated in the first week of July in 2018, 2019 and 2020 allowed us to get an insight into the interannual variability in $\delta^2\text{H}_\text{L}$ and $\delta^2\text{H}_\text{I}$. We focus this comparison on the transect lakes, since we sampled coastal lakes in different locations each year (i.e., in the Tromsø area vs. along the northern coast). Only three coastal lakes were sampled two or three of these years, and they follow similar interannual patterns as the transect lakes (Fig. 8a). We sampled the transect lakes in 2019 and 2020, and extended the transect farther southeast in 2020 (Fig. 7). Twelve transect lakes sampled in both years had more ^2H -depleted $\delta^2\text{H}_\text{I}$ in 2020, and a LEL intersecting the MWL at a more depleted value compared to 2019 (Fig. 6b and 7a), suggesting more depleted inflow water in 2020 than in 2019. This is supported by more depleted modeled $\delta^2\text{H}_\text{I}$ in 2020 compared to in 2019 (Fig. 7b). We suggest that the more ^2H -depleted values in 2020 were a response to 1) more precipitation in winter 2019/2020 than in 2018/2019 (Fig. 9d-e), 2) later spring melt in 2020 than 2019 (Fig. 9e) resulting in less time prior to our July sampling for ^2H -enriched summer precipitation to flush the ^2H -depleted winter melt out of the lakes, and/or 3) more May-June rainfall in 2019 than in 2020 (Fig. 9d) contributing to higher snowmelt bypass in 2019 (i.e., more snow melting off the high-elevation portions landscape before the lakes become ice free in the first half of June), which would remove ^2H -depleted winter precipitation from the catchment without entering the lake. The high precipitation amounts in January-March 2020 were associated with extreme AO+ conditions and an exceptionally strong stratospheric polar vortex, causing strong winter westerlies that carried greater-than-average amounts of moisture from the North Atlantic over Fennoscandia (Lawrence et al., 2020; Fig. 9a, d-e).

An additional explanation for ^2H -depleted lake inflow values in 2020 could be a change in moisture source. Surface waters in the Baltic Sea are relatively ^{18}O -depleted compared to the North Atlantic (Schmidt et al., 1999), and a greater influence of Baltic moisture to our study area could therefore contribute to more depleted precipitation. In winter 2018/2019 most of the Bothnian Bay was sea-ice covered, whereas winter 2019/2020 Baltic Sea sea-ice cover set a record for minimum winter extent in the 120-year-long measuring period (Sjöfartsverket & SMHI, 2020a, 2020b). However, predominantly southwesterly winds and pressure systems moving eastward over the Baltic Sea in both years imply a low impact of Baltic Sea moisture to our study area during this period.

Some lakes in the area have been sampled in previous studies, allowing us to extend our interannual comparison further back in time. In 2013, nine of the transect lakes were relatively enriched compared to the same lakes sampled in 2019 and/or 2020 (Balascio et al., 2018;

Fig. 6a and 8). The interannual changes in lake water isotopic composition were parallel to the MWL, suggesting that the lakes had more ^2H -enriched inflow water in 2013 than in 2019 and 2020, but that all years had a similar influence of evaporation. The hydrological year 2012/2013 was relatively dry compared to 2018/2019 and 2019/2020 (especially in Tromsø; Fig. 9d-e). However, June and July 2013 were wet (Fig. 9d), and the lakes were sampled several weeks later than in 2019 and 2020 (July 26, 2013; N. Balascio, personal communication), suggesting that the relatively enriched lake water values reflect greater summer precipitation inflow to the lakes in 2013. Two lakes close to Naimakka (Oikojärvi and Keitjoru) that were sampled by St. Amour (2009) in July 2003, 2004 and 2006 displayed a similar magnitude of interannual variability to the lakes in our dataset (Fig. 8a). The Oikojärvi samples from July 2004 and 2006 were close to the MWL, whereas the 2003 sample was evaporatively enriched along a LEL, perhaps due to exceptional dryness that year, as suggested by St. Amour (2009).

5.2.2. Seasonal variability

The lake water isotopic composition also follows a seasonal cycle. St. Amour (2009) collected monthly samples (excluding February and December) from the two lakes in Naimakka from 2002 to 2005, generally recording lowest values in the beginning of the ice-free season due to isotopically depleted snowmelt, increasing values over summer due to more enriched inflow isotopic composition and/or evaporative enrichment, and high and relatively stable composition during the ice-cover season (Fig. 8b). Guossajavri, northern Sweden, showed a similar trend in 2007 and 2008, with the lake moving up along the LEL during the summer months, and being recharged in winter (Rosqvist et al., 2013; Fig. 8b). This is in agreement with other lakes in northern Sweden (Jonsson et al., 2009) and elsewhere in the Arctic (Leng & Anderson, 2003; Tondur et al., 2013).

We sampled two lakes in Tromsø in March, June, July and/or September in 2018–2020. Except for a low March value, the seasonal isotope patterns in these lakes are similar to interannual patterns in all the lakes: isotope values tend to move parallel to the MWL, implying changes in inflow isotopic composition, rather than changes in evaporation. Although we do not have samples to examine seasonal variability in the transect lakes, given the similarities in interannual variability between the coast and transect, we hypothesize the transect lakes experience similar seasonal changes in inflow isotopic composition, with constant influence of evaporative enrichment.

5.3. Implications for precipitation proxy reconstructions

Interpreting isotope proxy records often assumes that, for a given record, the control on the lake water isotopic variability is constant through time. We have demonstrated that lakes in this region display variable sensitivity to changes in inflow (precipitation) isotopic composition and to evaporative enrichment, and that these controls vary in both time and space. Resolving these controls on modern lake water isotopes is important for making quantitative paleoclimate inferences. Depending on what aspect of the hydrological cycle we aim to reconstruct (e.g., atmospheric circulation, precipitation amount or seasonality, moisture balance), we can target different lakes.

In parts of our study area, modern precipitation isotopes have strong seasonal variability (Figs. 2 and 7), and if this signal is preserved in lake sediment proxy records, we can use it to reconstruct changes in precipitation seasonality (Cluett & Thomas, 2020; Thomas et al., 2020). The seasonality preserved in a proxy record depends on the $\delta^2\text{H}$ or $\delta^{18}\text{O}$ of source (lake or soil) water at the time of proxy synthesis, which can vary between proxies (Corcoran et al., 2021). Lake water isotopic composition and seasonal variability depends on the lake water residence time (Jonsson et al., 2010; Tondur et al., 2013; Ala-Aho et al., 2018), which may also change through time.

Previous paleoclimate work in northern Fennoscandia suggests that changes in Holocene precipitation isotopes are strongly linked to

changes in air temperature and air mass sources. Lacustrine carbonates, biogenic silica, and cellulose from non-evaporated lakes in northern Sweden have been used to reconstruct past changes in lake water $\delta^{18}\text{O}$ and precipitation (Shemesh et al., 2001; Hammarlund et al., 2002; Rosqvist et al., 2004, 2007, 2013; St. Amour, 2009; Jonsson et al., 2010). In most of these studies, depleted summer lake water $\delta^{18}\text{O}$ was inferred to reflect increased influence of Arctic air masses. Furthermore, Rosqvist et al. (2013) suggested that periods of relatively high $\delta^{18}\text{O}_{\text{diatom}}$ reflect a dominance of summer precipitation, related to a positive AO index, whereas low $\delta^{18}\text{O}_{\text{diatom}}$ reflects dominant winter precipitation due to positive NAO.

Studying modern conditions can aid proxy interpretations since it allows us to directly link isotopic changes to observations of atmospheric circulation change. Arctic Ocean and Baltic Sea surface waters are relatively depleted compared to North Atlantic moisture (LeGrande & Schmidt, 2006; Bonne et al., 2019), suggesting that depleted precipitation isotopes, and thereby lake water isotopes, may reflect a change to dominance of Arctic or Baltic air masses. However, this will depend on the location of the lake and the moisture travel distance. Atlantic moisture (which is originally less depleted) that has been transported a long distance may have experienced more distillation than more locally derived moisture with a more depleted starting isotopic composition, and therefore also be relatively depleted (Putman et al., 2017; Cluett et al., 2021). Our ^2H -depleted lake water values in 2020 were likely not caused by a shift in moisture source region, but by high winter precipitation amounts associated with AO+ conditions and an increase in North Atlantic moisture (Lawrence et al. 2020). Greater-than-average winter precipitation amounts over Fennoscandia can result from both extremely positive and negative winter AO conditions, although from different moisture sources (North Atlantic vs. Barents Sea; Lawrence et al., 2020; Bailey et al., 2021). Complicating things further, modern moisture from the Barents Sea has an isotopic composition comparable to North Atlantic moisture, due to increased strength of the North Atlantic Current (Bailey et al., 2021). If conditions were similar in the past, the Atlantic and Barents Sea moisture sources would be difficult to distinguish from one another in proxy records. Our lake water isotope observations imply that seasonal changes in precipitation amounts need to be taken into account when inferring atmospheric circulation from a proxy record in this region, and that changes in precipitation isotope values can be a result of a change in precipitation seasonality, moisture source, or a combination of both.

For reconstructions of atmospheric circulation and precipitation, we should target lakes that are minimally impacted by evaporative enrichment, since evaporation overprints the precipitation signal. In our dataset, most lakes >100 km from the coast have low d-excess values, suggesting that evaporative enrichment affects them to some degree, regardless of their throughflow regime (Figs. 3, 6a and 7). All transect lakes appear to fall on similar LELs, but with slope and intercept varying from year to year, depending on seasonality (Fig. 6). On interannual and seasonal timescales, changes in inflow appear to have a larger impact than evaporation, with isotope values moving along the MWL (Fig. 8). On longer timescales, evaporation may be more important, making it difficult to disentangle how much of the variability derives from each process (i.e., inflow vs. evaporation). Hence, we suggest that through-flowing coastal lakes are better targets for precipitation reconstructions, since changes in their lake water isotopic composition are more likely to primarily reflect changes in the inflow.

For evapo-concentrated lakes there are a few possible approaches to separate the precipitation from the evaporation signal, including 1) comparing two lakes with contrasting residence times (Anderson et al., 2001), 2) comparing terrestrial and aquatic leaf waxes in the same lake (Rach et al., 2017), or 3) measuring both $\delta^2\text{H}$ and $\delta^{18}\text{O}$ in the same records (Gibson et al., 2005). For the first approach, one needs to identify a non-evaporated lake with short residence time (low $\delta^2\text{H}_\text{L}$ - $\delta^2\text{H}_\text{I}$) and one evaporated lake with longer residence time (high $\delta^2\text{H}_\text{L}$ - $\delta^2\text{H}_\text{I}$; Anderson et al., 2001). Moreover, the lakes must receive source water reflecting

the same precipitation seasonality, and evaporation should be the only factor that differs between the two. The offset between $\delta^2\text{H}_\text{L}$ and $\delta^2\text{H}_\text{I}$ for our transect lakes is >10‰ independent of throughflow, suggesting that it will be difficult to identify an unevaporated lake in this region, which would be needed for this approach. Using the second approach, one compares terrestrial and aquatic leaf wax $\delta^2\text{H}$ in the same record, which have source water representing the same seasonality. In areas where terrestrial plant source water is evaporatively enriched, but the aquatic plant source water is not, the isotopic difference between the two reflects leaf water evaporative enrichment (Rach et al., 2017; Thomas et al., 2018; Curtin et al., 2019). In contrast, lake water in arid areas has been suggested to be more evaporatively enriched than terrestrial leaf water, meaning that the difference primarily reflects the evaporative enrichment of lake water (Mügler et al., 2008; Balascio et al., 2013). Along our transect, both aquatic and terrestrial leaf waxes likely incorporate a signal of evaporative ^2H -enrichment due to low relative humidity and warm conditions in summer (Figs. 2 and 9b-c). Thus, a dual leaf wax approach to separate precipitation and evaporation signals might not be applicable along the transect but may be possible at coastal lakes. For the third approach, one measures $\delta^2\text{H}$ and $\delta^{18}\text{O}$ in the same lakes, which may allow estimation of past changes in the slope of the LEL (Gibson et al., 2005) that could subsequently be used to infer past inflow values. Theoretically, this is possible, however, it involves many uncertainties. First, one needs to analyze the same proxies at the same time slice in several lakes. Second, those two proxies (one with $\delta^2\text{H}$ and one with $\delta^{18}\text{O}$) need to reflect the same pool of water for the same window of time in a season. Moreover, one needs to have good constraints on biosynthetic and other fractionation factors (Brandriss et al., 1998; Sachse et al., 2012; Lombino et al., 2021) to be able to estimate lake water $\delta^2\text{H}$ and $\delta^{18}\text{O}$ and directly compare $\delta^2\text{H}$ and $\delta^{18}\text{O}$ from different proxies (Corcoran et al., 2021).

A more recent approach to guide paleoclimate interpretations is to use proxy system modeling, which uses modern climate variables as inputs to mathematical descriptions of lake water isotope and energy balance (e.g., PRYSM2.0; Dee et al., 2018). By conducting sensitivity tests in which we iteratively change values of one or more climate parameters (e.g., lower relative humidity, more winter precipitation), we can quantify the parameter, or combination of parameters, that could explain changes recorded in lake water isotope proxies. For lakes in our transect, we could for example focus on variables that affect the lake water inflow isotopic signal and amount of evaporative enrichment, to assess the sensitivity of the lake water isotopic composition to these variables. If we have an estimate of how a change in moisture source would affect the precipitation isotopes, we can modify the inflow isotope value in the model to see how this would impact the lake water isotope value. One way to simulate how a shift in moisture source would influence the lake water inflow value is to use a one-dimensional Rayleigh-type distillation model along a theoretical trajectory (Sodemann et al., 2008; Steen-Larsen et al., 2011; Cluett et al., 2021). These approaches can provide us with quantitative constraints on past climate changes.

When interpreting lake-water-derived isotope proxy data from northern Fennoscandia, we need to consider the magnitude of variability in both inflow isotopic composition and evaporation. On the coast, inflow causes more variability than evaporation. Considering the variability among the transect lakes, the ranges for $\delta^2\text{H}_\text{L}$ and $\delta^2\text{H}_\text{I}$ are similar, suggesting that inflow and evaporation might be equally important. Interannual variability is largely due to changes in inflow isotopic composition, however, suggesting the importance of inflow isotopic composition even in lakes that are strongly influenced by evaporation. Therefore, we recommend targeting through-flowing lakes by the coast for precipitation reconstructions. If working with more continental records, we need to address the significance of both inflow and evaporation processes. Importantly, our data show that we cannot interpret all isotopic variability in evapo-concentrated basins in terms of evaporation, as there might also be a substantial variability in inflow isotopic composition through time.

Data availability

Water isotope data are freely available in the Water Isotopes Database at www.waterisotopes.org (Project IDs 00292 (surface waters) and 00293 (precipitation)), and in the [Supplementary Information](#).

CRedit authorship contribution statement

Sofia E. Kjellman: Conceptualization, Investigation, Visualization, Writing – original draft, Project administration, Funding acquisition. **Elizabeth K. Thomas:** Conceptualization, Investigation, Writing – review & editing, Supervision, Funding acquisition. **Anders Schomacker:** Conceptualization, Investigation, Writing – review & editing, Supervision, Funding acquisition.

Declaration of Competing Interest

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

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Appendix A. Supplementary data

Supplementary data to this article can be found online at <https://doi.org/10.1016/j.jhydrol.2022.127556>.

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Paper III

