

A modern snapshot of the isotopic composition of lacustrine biogenic carbonates – records of seasonal water temperature variability

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Received: 10 September 2021 – Discussion started: 1 November 2021 Revised: 26 April 2022 – Accepted: 4 May 2022 – Published: 2 June 2022

Abstract. Carbonate shells and encrustations from lacustrine organisms provide proxy records of past environmental and climatic changes. The carbon isotopic composition (δ^{13} C) of such carbonates depends on the δ^{13} C of dissolved inorganic carbon (DIC). Their oxygen isotopic composition (δ^{18} O) is controlled by the δ^{18} O of the lake water and by water temperature during carbonate precipitation. Lake water δ^{18} O, in turn, reflects the δ^{18} O of atmospheric precipitation in the catchment area, water residence time and mixing, and evaporation. A paleoclimatic interpretation of carbonate isotope records requires a site-specific calibration based on an understanding of these local conditions.

For this study, samples of different biogenic carbonate components and water were collected in the littoral zone of Lake Locknesjön, central Sweden (62.99° N, 14.85° E, 328 m a.s.l.) along a water depth gradient from 1 to 8 m. Carbonate samples of living organisms and subfossil remains in surface sediments were taken from the calcifying alga *Chara hispida*, from bivalve mollusks of the genus *Pisidium*, and from adult and juvenile instars of two ostracod species, *Candona candida* and *Candona neglecta*.

Our results show that neither the isotopic composition of carbonates nor the δ^{18} O of water vary significantly with water depth, indicating a well-mixed epilimnion. The mean δ^{13} C of *Chara hispida* encrustations is 4% higher than the other carbonates. This is due to fractionation related to photosynthesis, which preferentially incorporates ¹²C into the or-

ganic matter and increases the δ^{13} C of the encrustations. A small effect of photosynthetic ¹³C enrichment in DIC is seen in contemporaneously formed valves of juvenile ostracods. The largest differences in the mean carbonate δ^{18} O between species are caused by vital offsets, i.e., the species-specific deviations from the δ^{18} O of inorganic carbonate which would have been precipitated in isotopic equilibrium with the water. After subtraction of these offsets, the remaining differences in the mean carbonate δ^{18} O between species can mainly be attributed to seasonal water temperature changes. The lowest δ^{18} O values are observed in *Chara hispida* encrustations, which form during the summer months when photosynthesis is most intense. Adult ostracods, which calcify their valves during the cold season, display the highest δ^{18} O values. The seasonal and interannual variability in lake water δ^{18} O is small ($\sim 0.5\%$) due to the long water residence time in the lake. Seasonal changes in the temperature-dependent fractionation are therefore the dominant cause of carbonate δ^{18} O differences between species when vital offsets are corrected.

Temperature reconstructions based on paleotemperature equations for equilibrium carbonate precipitation using the mean δ^{18} O of each species and the mean δ^{18} O of lake water are well in agreement with the observed seasonal water temperature range. The high carbonate δ^{18} O variability of samples within a species, on the other hand, leads to a large scatter in the reconstructed temperatures based on individual

samples. This implies that care must be taken to obtain a representative sample size for paleotemperature reconstructions.

1 Introduction

The oxygen and carbon isotopic compositions of biogenic carbonates from lake sediments are powerful tools for reconstructing past climatic and environmental changes. Despite the complex influences of climate, hydrology, catchment and lake processes, as well as biological effects, they may be interpreted, depending on the local context, as proxies of temperature, evaporation, the isotopic composition of atmospheric precipitation, or atmospheric circulation patterns (e.g., Holmes et al., 2020; Andersson et al., 2010; Jonsson et al., 2010; Hammarlund et al., 2003; Schwalb, 2003; Hammarlund et al., 2002; von Grafenstein et al., 1999a).

Calcium carbonate (CaCO₃) remains of lacustrine organisms can be identified based on the characteristic morphology of shells, valves or encrustations. For paleoclimate reconstructions, it is important to assess the isotopic composition of specific biogenic carbonates to avoid the difficulties involved in the interpretation of isotopic records obtained on bulk carbonate matter in lake sediments. Changes in the bulk carbonate composition can be caused by variations in the assemblage of calcifying organisms, in clastic carbonate input, or in the amount of inorganic calcite precipitated in the lake (e.g., Bright et al., 2006; Hammarlund and Buchardt, 1996). Furthermore, carbonate minerals such as calcite, aragonite or dolomite differ in their isotopic compositions, so a potentially variable carbonate mineralogy both related to coeval sedimentation and to early diagenesis must be considered (McCormack and Kwiecien, 2021; McCormack et al., 2019).

The δ^{18} O and δ^{13} C of lacustrine biogenic carbonates depend on the isotopic composition of the oxygen and carbon sources, water and dissolved inorganic carbon (DIC), respectively. The isotopic composition of these sources can be variable over space and time, and is modified during carbonate precipitation, mainly through temperature-dependent fractionation and fractionation during physiological processes.

The δ^{18} O of biogenic carbonates carries the isotopic signature of the lake water, which, in turn, depends on the δ^{18} O of atmospheric precipitation in the catchment area, and is increased by evaporation from the lake (Gat, 1996). An increase in air temperature leads to a higher δ^{18} O of lake water through its effect on the δ^{18} O of atmospheric precipitation and on the evaporative enrichment of lake water, and consequently to an increase in carbonate δ^{18} O (see Fig. 1). Biogenic carbonates forming at different water depths may differ in their δ^{18} O, e.g., only those formed above the thermocline are affected by higher temperatures and evaporation during times of thermal stratification of the lake.

The δ^{13} C of biogenic carbonates carries the isotopic signature of DIC (McConnaughey, 1989; Ito, 2001). It depends

on the δ^{13} C signal of DIC in recharging groundwater that is transferred into the lake. The δ^{13} C of groundwater DIC is controlled by exchanges between atmospheric CO₂, vegetation and soil air CO₂, and carbonate bedrock under open or closed system conditions (Clark and Fritz, 1997). In lake water, the δ^{13} C of DIC can vary seasonally, depending on photosynthetic activity in the photic zone, the oxidation of organic matter, and thermal stratification.

The fractionation of oxygen isotopes between water and carbonate is temperature-dependent (Grossman and Ku, 1986; Kim and O'Neil, 1997; Beck et al., 2005; Kim et al., 2007). Biogenic carbonates formed during different times of the year can therefore differ in δ^{18} O due to seasonal water temperature changes, which are especially strong in shallow water. Fractionation factors for carbonate precipitated in isotopic equilibrium with the water are known from empirical studies. However, many earth surface carbonates precipitate out of isotopic equilibrium (Daëron et al., 2019). An increase in water temperature leads to a decrease in the fractionation of oxygen isotopes between water and carbonate (see Fig. 1), and consequently to a decrease in the δ^{18} O of equilibrium carbonate by -0.24% °C⁻¹ (Craig and Gordon, 1965; Kim and O'Neil, 1997). Note that the relationship between atmospheric precipitation δ^{18} O and air temperature is opposite, with a δ^{18} O increase of ~ 0.6 % °C⁻¹ in mid-latitudes (Rozanski et al., 1992). Kinetic effects on fractionation occur in the case of fast carbonate precipitation, observed, e.g., in Chara due to intense photosynthesis (Apolinarska et al., 2016; Andrews et al., 2004), which lowers the carbonate δ^{18} O. Furthermore, variations in the pH have an influence on the isotopic composition of carbonates, with an increasing pH lowering the δ -values (Herbst et al., 2018; Teranes et al., 1999; McConnaughey, 1989).

Fractionation due to physiological processes in a calcifying organism leads to a different isotopic composition of the biogenic calcite compared to inorganic calcite precipitated in isotopic equilibrium. This species-specific vital offset can be quantified through laboratory studies under controlled conditions (Li and Liu, 2010; Chivas et al., 2002; Xia et al., 1997b) or through monitoring in the field (Börner et al., 2017; Decrouy et al., 2011b; Keatings et al., 2002; von Grafenstein et al., 1999b). In addition to the vital offsets, the conditions in the micro-habitat of the carbonate-producing species can play a role as well (Decrouy et al., 2011a; Xia et al., 1997a). If the vital offsets for a given species are known and appear to be constant, they can be corrected for and do not hamper proxy interpretations. However, the causes of vital effects are not yet fully understood, and it has been suggested that varying water chemistry has an influence (Devriendt et al., 2017; Decrouy et al., 2011b).

Recent studies have highlighted the seasonal bias in proxies of past sea surface temperatures and the importance of distinguishing between annual and seasonal temperature reconstructions (Bova et al., 2021; Hertzberg, 2021). Lacustrine biogenic carbonates formed during different times of



Figure 1. A simplified schematic representation of the influences on the oxygen isotopic composition (δ^{18} O) of lacustrine biogenic carbonates. The isotopic composition of precipitation is correlated with air temperature in middle and high latitudes (Rozanski et al., 1992). A higher air temperature leads to a higher lake water δ^{18} O, as indicated in ①, which is reflected in the δ^{18} O of the carbonates. There are additional influences on lake water δ^{18} O, such as catchment and lake hydrology, relative humidity, and the seasonal distribution of atmospheric precipitation, which are not represented in the figure. The fractionation of oxygen isotopes between water and carbonate is also temperature-dependent, as indicated in ②. Note the opposite directions of the temperature influence on ① lake water δ^{18} O and on ② fractionation. As a consequence, temperature has two opposing effects on the δ^{18} O of carbonates. Lastly, "vital effects" which are dependent on the specific physiology of each species, lead to a constant offset between the δ^{18} O of the biogenic carbonate and inorganic carbonate which would precipitate in isotopic equilibrium with the water.

the year can provide insights into seasonal temperature variations. In this context, seasonal influences under modern conditions must be understood. Potential past water depth changes must be considered, because these could impact the location of carbonate formation with respect to a thermocline, changing the relationship between carbonates from organisms living at times of lake water stratification vs. at times of a mixed water column.

For the use of δ^{18} O and δ^{13} C in lacustrine biogenic carbonates as paleoenvironmental proxies, the different influencing factors must be known or constrained. For example, benthic ostracods in deep lakes living below the permanent thermocline form their calcitic valves at a nearly constant temperature around 4 °C, close to the density maximum of freshwater. Variations in the δ^{18} O therefore primarily reflect variations in the δ^{18} O of atmospheric precipitation if evaporative enrichment in ¹⁸O of the lake water is negligible (von Grafenstein and Labuhn, 2021). Under such optimal conditions, the δ^{18} O of ostracod valves provides the basis for reconstructing precipitation δ^{18} O (e.g., von Grafenstein et al.,

1999a). However, ostracod population densities are usually low and carbonates are not always preserved in profundal sediments. Carbonate-producing organisms (including algae that need light for photosynthesis) are often more abundant at littoral sites, where the interpretation of isotope proxies is complicated by a larger seasonal water temperature amplitude and a stronger influence of evaporation.

Disentangling the influences on the isotopic composition of specific biogenic carbonates is possible based on monitoring of the relevant parameters (such as temperature, δ^{18} O and pH of lake water, and the δ^{13} C of DIC), for a sufficiently long time to represent the entire time period of calcification of each organism and to capture seasonal and interannual variability. Monitoring can provide detailed understanding for interpretation of the proxies in the local context of each study (Börner et al., 2017; Meyer et al., 2017; Mayr et al., 2015), but long-term investigations (e.g., von Grafenstein et al., 1999b; Decrouy et al., 2011b) are rare and subject to logistic and financial constraints. In this study, we therefore rely on a single sampling occasion to investigate the oxygen and carbon isotopic composition (δ^{18} O and δ^{13} C) of different carbonate-producing species from Lake Locknesjön in central Sweden. At shallow water sites, the following types of samples were taken from living organisms and their subfossil remains in surface sediments: the calcite encrustations of the algae *Chara hispida*, the aragonitic valves of the bivalve mollusk *Pisidium* sp., and the calcitic valves of two species of ostracods, *Candona candida* and *Candona neglecta*. In addition, two species of charophytes, *Chara aspera* and *Chara hispida* were sampled in the nearby Lake Blektjärnen.

Our goals are (1) to test whether this approach of a "snapshot" study with short-term sampling can narrow down the primary environmental drivers of the isotopic composition of specific biogenic carbonates when long-term monitoring studies are not feasible. We focus on interspecies and intraspecies variations and compare them against the δ^{18} O of local atmospheric precipitation, streams and lake water, considering vital offsets and the timing of calcification for each species. We further (2) assess the uncertainty in seasonal water temperature reconstructions based on isotope measurements of lake water and of carbonates from multiple species. Lastly (3), this snapshot breaks down the details of what could be represented in a single measurement when building long isotope time series from sediment records. We discuss the sensitivity of such a measurement to the choice of the sampled material, and the implications for sub-sampling of sediment cores for paleoenvironmental studies. These investigations will also aid the site-specific paleoenvironmental interpretation of equivalent measurements from Holocene sediments in Lake Locknesjön.

2 Material and methods

2.1 Study site

Lake Locknesjön is located in central Sweden (62.99° N, 14.85° E, 328 ma.s.l., Fig. 2). It is one of the few carbonate lakes in northern Scandinavia, and abundant biogenic carbonate deposits are found in its littoral sediments, with charophyte encrustations as the dominant sediment component. The lake has an area of 27 km^2 , a maximum depth of 57.4 m and a mean depth of 17.6 m (SMHI, 2020a). The catchment area is 134 km², of which about 20% is the lake itself, 65% is coniferous forest, 10% is agricultural land, and the rest is heathland, peat bogs and rural settlements (SMHI, 2020a). There are no other lakes upstream in the catchment area.

The bedrock in the catchment area is dominated by Cambrian to Silurian carbonate-rich sedimentary rock, with some Precambrian crystalline bedrock in the southern part. The lake's central part lies within the Lockne Crater, which was formed by a meteorite impact 458 Ma ago (Ormö et al., 2010). The impact structure consists of impact breccia and resurge deposits (Sturkell et al., 2013). The fracture frequency of the breccia is estimated to be 25 times higher than in the surrounding crystalline bedrock, and the porosity is 2–10 times higher (Bäckström, 2005). These characteristics, together with the karstified carbonate bedrock, facilitate groundwater flow in the catchment area (Dahlqvist et al., 2018). The Quaternary deposits consist mostly of glacial till with a typical thickness of 3–5 m (Dahlqvist et al., 2018). During the last deglaciation, the ice margin retreated northwestward from the study area around 10 000 cal. yr BP (Stroeven et al., 2015; Labuhn et al., 2018).

The hydrological model of the Swedish Meteorological and Hydrological Institute (SMHI) gives an average discharge at the outflow of Lake Locknesjön of $1.28 \text{ m}^3 \text{ s}^{-1}$ (with a mean high water discharge of $3.10 \text{ m}^3 \text{ s}^{-1}$ and a mean low water discharge of $0.55 \text{ m}^3 \text{ s}^{-1}$), and a residence time of 11.5 years. Inflow of surface water to the lake is limited to a few small streams. The modeled average discharge of the largest inflows combined is $0.37 \text{ m}^3 \text{ s}^{-1}$ (SMHI, 2020a). Based on this modeled river discharge data, groundwater can be estimated to make up about 70% of the inflow to the lake. During our field work, groundwater inflows were identified based on a bubbling movement of fine sediment on the lake floor, which was observed on several locations along the sampled transect (see https://doi.org/10.5446/56815, Labuhn et al., 2021). Dahlqvist et al. (2018) described traces of underwater erosion as further indications of groundwater inflow, with water temperatures at such inflows 6–8 °C lower than in the surrounding lake water.

Lake Locknesjön is ice-covered from early December to mid-May (1961–1990 average; SMHI, 2020a). It is a dimictic lake, which is stratified during January and from July to September, and mixed between these periods (Fig. 3a; Eklund, 1998). Autumn mixing occurred at temperatures of about 9 °C in 1957 (the only year for which monthly water temperature measurements are available). The average pH of the lake water is 8.16, with values up to 8.33 above the thermocline during summer (Fig. 3b; data from the Triple Lakes Project, http://www.triplelakes.se/, last access: 16 February 2021).

The average annual air temperature at the meteorological station of Mörsil, 70 km northwest of the lake, is 2.3 °C, and the average annual precipitation is 663 mm (1975–2005 averages; SMHI, 2020b). In the year of sampling (2018), the average May–July temperature was 2 °C higher and the May–July precipitation sum was 40 mm lower than in the reference period (Fig. 4a). The modeled discharge at the lake's inflow and outflow in 2018 shows no significant difference from the average conditions (SMHI, 2020a). The Global Network of Isotopes in Precipitation (GNIP) station closest to our study area, Bredkälen, is located 100 km north of Lake Locknesjön at 400 m a.s.l. and provides monthly measurements of δ^{18} O for the period of 1975–1980, but no measurements of δ D (Fig. 4b; IAEA/WMO, 2019).



Figure 2. (a) Map of Sweden showing the location of the study area (red rectangle), of the GNIP station Bredkälen and of the meteorological station Mörsil. (b) Lake Locknesjön with its catchment (hatched) and Lake Blektjärnen. (c) Locations of the sampling sites on a transect from the shore towards the center of the lake. The area shown in (b) is marked by the red rectangle in (a) and the area shown in (c) is marked in (b) (country outline from USGS, lake and catchment outline from SMHI, 2020a; orthophoto © Lantmäteriet 2020). (d) Photograph of a *Chara hispida* in Lake Locknesjön. An internode (between two nodes) and a branchlet are indicated on the image. The part of the *Chara hispida* visible on the photograph measures about 2 cm in height. (e) Images of different lacustrine biogenic carbonates analyzed in this study: (left) *Pisidium* valve, (middle) adult *Candona candida* valve, (right) *Chara hispida* encrustations.

2.2 Field sampling

In July 2018, a diving team collected samples of water, surface sediments and living *Chara* from the lake. Samples were taken at 13 locations along a transect from the western shoreline towards the center of Lake Locknesjön (Fig. 2c). Water depths at the sampling locations range from 1 to 8 m. Surface sediment samples of about $8 \times 8 \text{ cm}^2$ and less than 1 cm depth were taken with a small shovel and filled into sealable plastic bags. Water samples were taken directly into small glass bottles. Living *Chara hispida* were only found down to 7 m water depth. Up to six individuals were sampled at each location cutting off the whole algae by hand. For a comparison of different species of *Chara*, samples of *Chara hispida* and *Chara aspera* were collected at a depth of 0.3 m in the nearby Lake Blektjärnen. In Locknesjön only *Chara hispida* were found. At 5.6 m water depth in Locknesjön, water from a groundwater inflow was sampled with a syringe. Water samples were also collected at the outflow of Lake Locknesjön, and from surface water of Lake Blektjärnen (Fig. 2b). Further water samples were taken in 2013 and 2014 at depths of 0–20 m at Lake Locknesjön (11 samples), at one of its main tributaries (Musån river, 13 samples), and at its outflow (Forsaån river, 6 samples). Water samples down to 20 m depth were taken with a UWITEC gravity corer, the tributary and outflow samples were taken from the shore into small glass bottles.



Figure 3. (a) Water temperature profiles from Lake Locknesjön. Dark blue dots represent measurements taken between 1951 and 1962, two times per year approximately at the time when the water was coldest during winter (February–March) and warmest during summer (August). Profiles extend from the surface to the bottom at the lake's deepest point. The dashed lines show measurements taken at the same point throughout the year 1957 in the indicated months (data from Eklund, 1998). Other dots represent measurements taken between 2014 and 2016 from surface and bottom water at two sites of shallow (< 11 m, light blue dots) and intermediate (< 20 m, light gray dots) water depth (data from the Triple Lakes Project, http://www.triplelakes.se/, last access: 16 February 2021). Dotted and solid lines connect measurements taken from different depths at the same location and on the same day. The shaded area marks the water depths from which carbonate samples were taken. (b) pH measured in surface and bottom water at two sites of shallow and intermediate depth between 2014 and 2016 (data from the Triple Lakes Project).

2.3 Carbonate sample preparation

The following biogenic carbonate components were obtained from the samples and prepared for stable isotope measurements: encrustations around the internodes and branchlets of *Chara*, valves of *Pisidium*, and valves of juvenile and adult ostracods (see examples in Fig. 2).

Chara species were determined based on the number of cells of the main stem and the differences between primary and secondary cells (Caisova and Gabka, 2009). A further indicator was the relation between the number of cells in the main stem and the number of branchlets in a whorl. Since the whorls of branchlets are not preserved in the sediments (only branchlet fragments), this was only useful for the determination of living algae. Internode and branchlet samples were cut off from different parts of the living *Chara* and dried. The organic matter was removed from the calcite encrustations by putting the samples in 10 % hydrogen peroxide (H₂O₂) for at least 48 h. They were then cleaned with a thin paintbrush to remove fine-grained calcite sticking to the encrustations. The removed fine calcite was also kept for isotope measurements

in order to test if it is composed of inorganically precipitated calcite, or if it originated from the encrustations themselves.

Sediment samples were wet sieved to obtain residual size fractions containing specific biogenic carbonates (> 400 and $> 200 \,\mu\text{m}$). Valves of ostracods and *Pisidium* and encrustations of Chara were picked out under a binocular microscope. Based on their size, the latter were assumed to be fragments of internodes. Ostracod species and Pisidium were identified based on the characteristics of their valves, following Meisch (2000) for ostracod identification. The dominant ostracod species were Candona candida and Candona neglecta. Other species (Cytherissa lacustris, Candona caudata, Cypridopsis vidua, and Limnocythere inopinata) were present in insufficient numbers for isotope measurements. Adult (A) and juvenile instars (A-1 to A-4) of ostracods were separated. A number of living specimens of ostracods and Pisidium were found in the sediment samples. These were briefly boiled so that organic matter could be easily removed from the valves. All valves were rinsed with ethanol and air dried (von Grafenstein et al., 1992; Mischke et al., 2007). In the following, the carbonates obtained from living organisms



Figure 4. (a) Monthly climatology of air temperature and precipitation in the study area from the meteorological station Mörsil for the period of 1975–2005 (data from SMHI, 2020b). The values of the year 2018 until the time of sampling (July) are indicated by the triangles. The average annual air temperature and the average annual precipitation are given at the top of the plot. (b) δ^{18} O in precipitation at the GNIP station Bredkälen for the period of 1975–1980 (data from IAEA/WMO, 2019). See Fig. 2 for the locations of the stations.

are referred to as living samples, the carbonates deposited in surface sediments are referred to as subfossil samples.

In order to obtain sufficient sample weights for isotope measurements, several ostracod valves from the same sampling location, species and instar were combined. Valves of living and subfossil specimens were kept separate. The required numbers of valves in a sample were 2 for adults, 5 for A-1, 8 for A-2, and 15–25 for A-3 and A-4. If the sample weights were not sufficient due to few available valves, successive instars were combined for some locations, leading to combinations of A-2 and A-3, or A-3 and A-4 depending on the sample. *Pisidium* samples were composed of individual valves. *Chara* samples were composed of single stalks from an internode or branchlet.

2.4 Stable isotope measurements

For this study, 49 water samples and 271 carbonate samples were measured. Measurements of δ^{18} O and δ D of water samples were conducted at the Laboratoire des Sciences du Climat et de l'Environnement (LSCE), Gif-sur-Yvette, France. The δ^{18} O was measured with a Finnigan MAT252 isotope ratio mass spectrometer with a precision of $\pm 0.05 \%_0$ (2 standard deviations). The δ D was measured with a Picarro Analyzer (Cavity Ring-Down Spectroscopy) with a precision of $\pm 0.7\%_0$ (1 standard deviation). The results are given relative to the V-SMOW standard.

Carbonate δ^{18} O and δ^{13} C measurements were carried out at MARUM, Bremen, Germany, using a ThermoFisher Scientific 253plus gas isotope ratio mass spectrometer with a Kiel IV automated carbonate preparation device. The standard deviation of the reference standards over the measurement period was 0.06% for δ^{18} O and 0.03% for δ^{13} C. The results are given relative to the V-PDB standard. Some sample weights were $< 20 \,\mu$ g, resulting in low signal intensities (1800–3000 mV for mass 44). A control standard (ground limestone) showed no deviation over this signal range from the expected values within the above errors. Of the samples 29 had a signal intensity $< 1800 \,\text{mV}$ and were excluded from further analyses. One major outlier in δ -values was identified in a sample of living *Candona neglecta*, instar A-2, taken at 5 m water depth (sampling location 5) and also excluded from further analysis. Its δ^{13} C value of -9.63% was below the outer limit of the interquartile range of all ostracod δ^{13} C values.

2.5 Data analysis

Statistical analyses were performed using R v. 3.6.1 (R Core Team, 2018) and the statistical package PAST v. 3.25 (Hammer et al., 2001). All data sets of individual species were tested for normal distribution using the Shapiro-Wilk test (Shapiro and Wilk, 1965). For normally distributed data, correlations were calculated using the Pearson method (Pearson, 1895), and t-tests assuming unequal variances were calculated to compare different sample sets. For samples lacking normal distribution, this comparison was done using the Mann-Whitney test (Mann and Whitney, 1947) and the Spearman method was used to calculate correlations (Spearman, 1904).

Assuming calcite precipitation in isotopic equilibrium with the water, theoretical water temperatures were calcu-

lated from the mean, minimum and maximum δ^{18} O values of water and calcite (corrected for species-specific vital offsets). We used the empirical equation by Kim and O'Neil (1997), with $\delta^{18}O_{\text{calcite}(\text{SMOW})} - \delta^{18}O_{\text{water}(\text{SMOW})}$ as an approximation of the fractionation factor, and the conversion of calcite δ^{18} O values from the PDB to the SMOW scale according to Coplen et al. (1983). For the aragonitic *Pisidium* valves, we assume a temperature dependence of the δ^{18} O like for inorganic calcite (Grossman and Ku, 1986; Kim et al., 2007), and the species-specific offset correction accounts for the offset between the valve carbonate and inorganic calcite.

3 Results

This section presents the results of our stable isotope measurements of water and biogenic carbonates, and explains the influences on their isotopic composition. The subsequent Discussion (Sect. 4) then evaluates how far these influences can be constrained through our "snapshot" approach by comparing different species, and which paleoclimate information can potentially be obtained from the carbonates.

3.1 Isotopic composition of water

The δ^{18} O of monthly precipitation at the Bredkälen GNIP station averaged over the period of 1975–1980 closely follows the annual cycle of air temperature. Values of individual months range between -4.9% and -19.9% (Fig. 4b). Although the precipitation isotope time series and the local water isotope data were collected during different periods, some relationships can be inferred from the distributions of the isotope values.

The δD and $\delta^{18}O$ of water from the tributary of Lake Locknesjön plot close to the Global Meteoric Water Line (GMWL). The seasonal range of $\delta^{18}O$ in the tributary (-13.11% to -11.57%) is reduced compared to the range in precipitation (Fig. 5b), which indicates a substantial groundwater contribution to the stream flow. Still, a clear shift to higher $\delta^{18}O$ and δD values in the tributary shows a marked influence of precipitation on river supply to the lake during the warm season. The isotopic values of the groundwater inflow are more negative than all values of the tributary, and the $\delta^{18}O$ is close to the amount-weighted mean of precipitation for 1976–1980 (-13.5%), which as expected indicates a negligible influence of evaporation.

Water samples from the lake and its outflow, on the other hand, show effects of evaporative enrichment in the heavy isotopes compared to groundwater and the tributary by about 3% for δ^{18} O and 17% for δ D. This enrichment is seen in samples from all investigated years and in all seasons. The inferred a local evaporation line with a slope of 4.9 and an intercept of 29% is typical for lakes with evaporative water loss (Craig and Gordon, 1965; Craig, 1961).

The difference between lake water samples taken in winter and summer is small, with values ranging from -9.66% to -9.24% for δ^{18} O and from -75.4% to 73.6% for δ D. The exceptional meteorological conditions during the summer of 2018 did not lead to a significant effect on the δ^{18} O of lake water, e.g., through increased evaporation. The δ^{18} O of water sampled in 2018 is only about 0.1% higher than in other warm season water samples (Fig. 5c). A potential effect of the dry and warm summer conditions 2018 is either masked through the mixing of lake water, or could arrive in the lake with a delay. The small seasonal and interannual changes of the isotopic composition can be explained by the relatively long water residence time of over 10 years in Lake Locknesjön, caused by its large volume in relation to its limited catchment size (Fig. 2b). Interannual variations in the isotopic composition of precipitation are only reflected in lake water δ^{18} O if they last longer than the lake water residence time (von Grafenstein and Labuhn, 2021).

Variations in δ^{18} O and δ D between 0 and 20 m water depth in April 2014 do not exceed analytical uncertainty, with a range of 0.02% for δ^{18} O and 0.3% for δ D. The δ^{18} O and δ D vary only minimally between 1 and 8 m water depth in July 2018 and do not show any trends with depth. The ranges of 0.17% for δ^{18} O and 0.8% for δ D are only slightly larger than the measurement error, indicating that the lake water is well mixed above the thermocline. Lake Locknesjön displays a distinct summer stratification with a thermocline generally below 10 m (Fig. 3; Eklund, 1998). Although only one summer temperature profile per year is available, it can be concluded that the rapid transition from ice-covered conditions in May to maximum air temperature in July, together with the effects of strong winds, leads to a rather deep and well-mixed summer epilimnion.

Lake Blektjärnen shows a 0.66% lower δ^{18} O and a 13.1% lower δ D compared to Lake Locknesjön in the summer of 2018. These measurements are in agreement with previously published isotope data from Lake Blektjärnen (Andersson et al., 2010).

3.2 Isotopic composition of biogenic carbonates

3.2.1 Pisidium

The δ^{18} O values of the aragonite valves of *Pisidium* samples range between -8.83% and -7.38% and the δ^{13} C values lie between -7.20% and -4.00% (Table 1, Fig. 6a). The relatively narrow ranges can be explained by the accumulation of shell carbonate throughout their lifetime of up to 3 years (Holopainen and Jónasson, 1989). This averages out seasonal and interannual variations in the water isotopic composition and water temperature. There is no significant difference in isotope values between the valves of living specimens sampled in 2018 and the subfossil valves (i.e., which were formed and deposited during previous years), possibly because their time spans of carbonate accumulation



Figure 5. Isotopic composition of water in the study area. The symbols represent different locations and years of sampling. The color scale indicates the month of sampling, with blue values representing the cold period and red values representing the warm period of the year. (a) Average monthly δ^{18} O of atmospheric precipitation for the period of 1975–1980 at the GNIP station Bredkälen (data from IAEA/WMO, 2019, note that δ D is not available for this station). The large black diamond shows the mean δ^{18} O for 1976–1980 weighted by the amount of precipitation. (b) δ D and δ^{18} O of lake water (sampled in 2013–2014 and 2018), the groundwater inflow (2018), and water in the tributary (2013–2014) and outflow (2013–2014) of Lake Locknesjön, and lake water from Lake Blektjärnen (2018). The Global Meteoric Water Line (GMWL), is plotted in black, and a local evaporation line (LEL; $\delta D = 4.9 \cdot \delta^{18}O + 29$) estimated from the isotopic composition of Locknesjön lake water in blue. (c) Zoom on the lake water isotope data from Lake Locknesjön.

overlap. The mean δ^{18} O is the same, and the mean δ^{13} C differs by 0.19%.

The previously reported vital offset on the oxygen isotopic composition of *Pisidium* valves of +0.9% (von Grafenstein et al., 1999b) is confirmed by this study. After correction for this offset, the *Pisidium* δ^{18} O falls close to expected values in equilibrium with the lake water, as observed in other studies of mollusk shells (Apolinarska et al., 2016).

3.2.2 Chara

Samples of calcite encrustations from *Chara hispida* in Lake Locknesjön show the largest range in isotope values compared to other species (Table 1). *Chara* can be annual or perennial (Martin et al., 2003; Pełechaty et al., 2013). It is not known whether overwintering occurs in Locknesjön, but different parts of the algae are formed during different times of the growing season, or possibly during different years. The growth of *Chara* is apical, i.e., calcification at a given

point in time occurs only around these apices (Coletta et al., 2001). Each analyzed sample consisted of only one internode or branchlet, which means that environmental variations within the growing season or between different years are not averaged in individual samples. The samples represent only the short period of time when the respective part of the *Chara* grew. There is no difference between the isotopic compositions of encrustations from living *Chara hispida* and subfossil encrustations collected from surface sediments (Fig. 7a), indicating that the environmental influences on the δ^{18} O and δ^{13} C of the encrustations were similar over the past few years. The possible perennial growth also means that there could be an overlap in the years of growth represented in living and subfossil samples.

Chara growth is restricted to the time of photosynthetic activity, i.e., the summer months in our study area (Hammarlund et al., 2002). The δ^{13} C is driven by fractionation during photosynthesis, which preferentially incorporates

Table 1. Isotopic composition of specific biogenic carbonates from living organisms and from subfossil samples in surface sediments of Lake Locknesjön. Mean, standard deviation (SD), minimum, maximum and range of δ^{18} O and δ^{13} C values, as well as the number of samples are presented for *Chara hispida*, *Candona candida*, *Candona neglecta* and *Pisidium* sp. All values are reported in %₀ relative to V-PDB. Note that these are the measured values without a correction of vital offsets. The results of individual sample measurements are available in the Pangaea database, see Sect. "Data availability".

Species	Instar	No.			$\delta^{18}O$					$\delta^{13}C$		
		samples	Mean	SD	Min	Max	Range	Mean	SD	Min	Max	Range
Chara hispida		100	-9.77	0.94	-12.59	-8.31	4.28	-1.48	0.83	-3.76	0.77	4.53
Candona candida	Adult	13	-5.38	0.77	-7.55	-4.43	3.12	-4.75	0.94	-6.42	-3.46	2.97
	A-1	14	-6.47	0.48	-7.29	-5.76	1.54	-3.88	0.98	-5.3	-2.08	3.23
	A-2, 3, 4	8	-6.97	0.27	-7.42	-6.65	0.77	-4.02	0.75	-5.01	-3.09	1.92
Candona neglecta	Adult	17	-5.13	0.58	-6.16	-4.13	2.03	-5.30	0.59	-6.26	-4.42	1.83
	A-1	13	-5.92	0.44	-6.76	-5.27	1.50	-4.70	0.67	-5.56	-3.47	2.09
	A-2, 3, 4	16	-6.73	0.38	-7.31	-6.08	1.23	-4.66	1.07	-6.9	-3.35	3.55
Pisidium sp.		37	-7.94	0.34	-8.83	-7.38	1.45	-5.29	0.72	-7.20	-4.00	3.20



Figure 6. Isotopic composition of specific biogenic carbonate samples collected from living organisms and from subfossil remains in surface sediments of Lake Locknesjön. (a) δ^{18} O and δ^{13} C of calcite encrustations from *Chara hispida* and of valves from *Candona candida*, *Candona neglecta* and *Pisidium*. (b) The same data with published vital offsets corrected for: δ^{13} C *Chara hispida*: -4% (value from Hammarlund and Keen, 1994), δ^{18} O ostracods: -2.2%, δ^{18} O *Pisidium*: -0.9% (values from von Grafenstein et al., 1999b). The data point marked in brackets was identified as a major outlier and left out from further analysis and figures.

¹²C into the organic matter. Consequently, DIC in the surrounding water and, in turn, the calcitic encrustations are enriched in ¹³C (McConnaughey and Falk, 1991). This process is reinforced by proton pumping, leading to additional ¹³C enrichment in the encrustations (Hammarlund et al., 1997). After correcting *Chara hispida* δ^{13} C for the vital offset of 4%*c*, the values are largely similar to the δ^{13} C values of other biogenic carbonate samples from Lake Locknesjön (Fig. 6b). The magnitude of carbon isotope fractionation in *Chara* encrustations observed here is therefore the same as in previous studies (Hammarlund and Keen, 1994; von Grafenstein et al., 2000).



Figure 7. Isotopic composition of calcite encrustations from different types of *Chara* samples. Internodes, branchlets and fine calcite were collected from living *Chara*, "subfossil fragment" refers to fragments of *Chara* encrustations taken from surface sediments. (a) δ^{18} O and δ^{13} C of *Chara hispida* samples from Lake Locknesjön. (b) δ^{18} O and δ^{13} C of *Chara hispida* and *Chara aspera* samples from Lake Blektjärnen.

The mean δ^{18} O of the *Chara hispida* encrustations in Lake Locknesjön (-9.77%) is close to that of inorganic calcite that would be precipitated in isotopic equilibrium with lake water, which has a mean δ^{18} O of -9.53%. Other studies, on the contrary, have found significant offsets from equilibrium due to kinetic effects during intense photosynthesis associated with fast calcite precipitation (Apolinarska et al., 2016; Andrews et al., 2004).

Fine calcite δ^{18} O values fall within the range of δ^{18} O of the encrustations, but their mean is higher than the mean of the encrustations. The observed δ^{13} C values are similar for fine calcite and encrustations. This indicates that the fine calcite, which also makes up the matrix of the carbonate-rich sediments of Lake Locknesjön, is composed of disaggregated *Chara hispida* encrustations. Allochthonous carbon-

ates would be expected to have a higher δ^{13} C than the fine calcite, close to the δ^{13} C of bedrock in the catchment area (around -0.5%, see Sect. 4.3). Other autochthonous carbonates which are not influenced by fractionation due to photosynthesis have a lower δ^{13} C.

The two *Chara* species sampled in Lake Blektjärnen differ in the isotopic composition of their encrustations (Fig. 7b). The average δ^{13} C in *Chara hispida* is significantly higher (1.04%) than in *Chara aspera*. Such differences can be caused by habitat conditions, such as stand density and by metabolic effects (Pronin et al., 2018; Coletta et al., 2001). The δ^{13} C is driven by photosynthesis. The *Chara hispida* in Blektjärnen grew in dense swards, which limits the water exchange and therefore the removal of DIC enriched in ¹³C due to photosynthesis, leading to a corresponding enrichment of ¹³C of the encrustations (Coletta et al., 2001). The average difference in δ^{18} O (0.46%) is not significant. The δ^{18} O of *Chara* is driven by water δ^{18} O, which leads to a corresponding species-independent δ^{18} O of encrustations formed in the same water.

Internodes and branchlets show small differences in their mean isotope values, but these differences are not consistent between species and lakes. In living *Chara hispida* from Lake Locknesjön, the mean δ^{18} O of branchlets is 0.56% higher and the mean δ^{13} C is not significantly different from internodes. In samples from Lake Blektjärnen, internodes and branchlets are not significantly different in the δ^{18} O and δ^{13} C of *Chara aspera*, nor in the δ^{18} O of *Chara hispida*, whereas the δ^{13} C of *Chara hispida* is higher on average in the internodes. Consequently, it does not seem to be necessary to distinguish between these parts of the alga in isotopic analyses of the encrustations.

3.2.3 Ostracods

The δ^{18} O values of ostracod valves range between -7.55%and -4.13%, and the δ^{13} C values range between -6.90%and -2.08% (Table 1). There is no significant difference between the δ^{18} O values of *Candona candida* and *C. neglecta*. The difference in δ^{13} C is statistically significant, but the mean δ^{13} C is only 0.67\% lower for *C. neglecta*. This general similarity in the isotopic composition of the two species is in agreement with previous studies (von Grafenstein et al., 1999b). With known vital offsets subtracted (2.2\% for both species; von Grafenstein et al., 1999b), the δ^{18} O values of both species are close to the values of other biogenic carbonates from Lake Locknesjön (Fig. 6b).

Comparing different developmental stages of the ostracods, there is a clear decrease in the average δ^{18} O values from adult valves to A-1 to A-2, A-3 and A-4 instars, although the values of individual samples partly overlap (Fig. 8). The timing of the molting of ostracods is controlled by temperature. In southern Germany, it has been observed that co-existing *Candona* instars A-4 to A-2 form their valves during early to mid summer, A-1 at the onset of autumn cool-



Figure 8. Isotopic composition (δ^{18} O and δ^{13} C) of the valves from two species of ostracods, *Candona candida* (blue) and *Candona neglecta* (red), taken from living specimens (encircled symbols) and subfossil samples in the surface sediments of Lake Locknesjön. The symbols represent the instars. See Sect. 2.3 for explanation of the combinations of different instars.

ing and adults between late autumn and early spring (von Grafenstein et al., 1999b). Consequently, seasonal changes in water temperature and water δ^{18} O have an influence on the δ^{18} O of the valves formed during different times of the year. Higher water temperatures lead to reduced fractionation and a decrease in the δ^{18} O of the precipitated carbonate.

Juvenile ostracods form their valves when the lake is thermally stratified during the warm season. As the instars A-2 to A-4 form at the time of maximum water temperature, the mean δ^{18} O of their values is significantly lower than in A-1 and adult samples. There is no significant difference between A-2, A-3 and A-4 instars. The average δ^{18} O of A-1 samples is higher than in younger instars because A-1 instars develop when the water temperature starts to decrease in autumn. Moreover, by the end of the summer the water in the epilimnion is more enriched in ¹⁸O by evaporation compared to the time when the younger instars calcify their valves, leading to a further increase in the δ^{18} O of A-1 valves. The valves of adult ostracods start forming in late autumn. When overturning occurs in the lake, stratification ceases and the isotopically enriched surface water is mixed with deep water with a lower δ^{18} O. Nevertheless, adult ostracods have significantly higher δ^{18} O values on average compared to juvenile instars, because their valves are formed at the lowest water temperature. However, the lowest δ^{18} O value of all ostracod samples was recorded in an adult specimen from 1 m water depth.

The patterns of average δ^{13} C values of ostracod valves from different instars are less clear compared to δ^{18} O. The average δ^{13} C of all juvenile instars are generally similar, whereas the average δ^{13} C of adult ostracods is slightly lower. The highest values, however, were observed in samples of A-1 instars. The δ^{13} C of ostracod valves depends on the carbon isotopic composition of DIC in the surrounding water. The higher δ^{13} C of juvenile ostracods suggests that DIC is slightly enriched in ¹³C as a result of *Chara* photosynthesis and other primary productivity during the summer when juveniles calcify their valves. When adult valves are formed, the lake is no longer stratified and the overturning brings DIC with a lower δ^{13} C from depths below the summer thermocline to the surface.

4 Discussion

4.1 Climate signal in the δ^{18} O of lake water

In order to interpret the isotopic oxygen composition of lacustrine biogenic carbonates, it is important to understand the causes of variations in the δ^{18} O of the oxygen source, i.e., the lake water. Generally, lake water δ^{18} O depends on the water balance as the δ^{18} O of the water input (atmospheric precipitation, potentially modified though processes in the catchment) can be increased through evaporation in the lake (Mayr et al., 2007; von Grafenstein and Labuhn, 2021). The measured values of river and groundwater inflow into Lake Locknesjön show that seasonal variations in the isotopic composition of atmospheric precipitation are smoothed through groundwater supply and long lake water residence time. The difference between tributary and lake water indicates evaporative enrichment in ¹⁸O. In the absence of long-term monitoring, it is not possible to disentangle the relative importance of precipitation and evaporative enrichment for the δ^{18} O of the lake water. However, since both factors increase with increasing temperature, lake water δ^{18} O should contain a signal of multiannual temperature variability. As shown by our lake water δ^{18} O and water temperature profiles (Figs. 3a, 9), the lake water is well mixed down to a depth of about 8 m, i.e., water depth is not a decisive factor across the range of sampled depths.

4.2 Influence of water depth on the isotopic composition of carbonates

None of the investigated species show any significant trends in average δ^{18} O or δ^{13} C, or in the range of isotope values, along the transect from 1 to 8 m water depth (Fig. 9). The largest temperature changes can be expected in shallow water, which would lead to larger ranges of δ^{18} O values at the shallowest sampling depths. However, since no systematic changes were observed in the carbonate isotopic compositions or in the δ^{18} O of the lake water, we conclude that the calcification conditions and the isotopic compositions of the carbon and oxygen sources did not significantly vary within the sampled range of water depths. The effects of water temperature changes and evaporative enrichment in ¹⁸O of lake water are the same, and the water at 8 m depth appears to be well mixed and clearly above the summer thermocline for the time span represented in the living and subfossil samples.



Figure 9. Interspecies and intraspecies variations in the isotopic composition (δ^{18} O and δ^{13} C) of biogenic carbonates sampled along a transect in Lake Locknesjön at water depths from 1 to 8 m. The colors indicate the species, and the shaded areas represent the ranges of their δ -values. The symbols represent different sample types (as in Fig. 7 for *Chara hispida* and as in Fig. 8 for ostracods). The blue asterisks represent δ^{18} O of water samples taken at the same locations. The values of lake water are reported in $\%_0$ relative to V-SMOW.

4.3 Causes of interspecies differences in the isotopic composition of carbonates

Vital offsets are the major cause of differences in the oxygen isotopic composition between species (Table 1, Fig. 6a). With these known and constant offsets corrected for, the mean δ^{18} O values of all species analyzed in our study plot closer to each other and their ranges show larger overlaps (Fig. 6b). The remaining differences can be explained by environmental influences, which vary with the timing and length of the respective calcification periods. The mean values of each species depend on the time of the year when the carbonate is precipitated, involving potentially large differences in water temperature. The range of values of a species depends on the length of the calcification period of each sample, a longer period smoothing seasonal and interannual variability.

The mean δ^{18} O value of *Chara hispida* encrustations lies clearly below the mean δ^{18} O values of all other species. The mean δ^{18} O values of *Pisidium* and juvenile ostracods are higher, and adult ostracods show the highest mean. The timing of calcification in *Chara hispida* is limited to periods of maximum photosynthetic activity in summer, which can last only a few weeks. These periods are likely also the periods of highest temperatures, which would explain the low

 δ^{18} O of *Chara hispida* encrustations compared to other carbonates. Seasonal temperature variations also cause the differences observed between adult and juvenile ostracods (see Sect. 3.2.3). As juvenile ostracods precipitate calcite mainly during the summer months, their values overlap with values of Chara hispida encrustations. The calcification period of Pisidium is the longest of all studied organisms and lasts throughout the ice-free period from May to November (Andersson et al., 2010). Most calcification occurs during summer, whereas their growth is substantially suppressed during winter (von Grafenstein et al., 1999b). The mean δ^{18} O value of *Pisidium* is slightly lower than that observed for juvenile ostracods A-2, A-3 and A-4, although the former are exposed to lower average water temperatures during their calcification period and thus a higher carbonate δ^{18} O would be expected due to temperature-dependent fractionation. The lower Pisidium δ^{18} O values may reflect the influence of relatively ¹⁸O-depleted snow-melt supply to the lake in spring and from mixing of surface water with ¹⁸O-depleted bottom water in autumn, processes that juvenile ostracods are not exposed to.

The range of δ^{18} O is smallest for *Pisidium* (1.45%) and largest for *Chara hispida* (4.28%). *Chara* can be perennial, but the samples measured in this study, encrustations from single internodes or branchlets, were precipitated at different times as the *Chara* grew, i.e., individual samples may represent different times within a growing season and possibly different years, hence explaining their larger isotopic range. The average lifetime of *Pisidium* of 3 years, together with their long annual calcification period probably explain their relatively narrow range of δ^{18} O.

Overall, seasonal water temperature changes seem to be the dominant cause of differences in the δ^{18} O of the investigated biogenic carbonates (after vital offset correction). The δ^{18} O of the lake water, on the contrary, does not have a strong influence due to the small seasonal and interannual variability around the mean of -9.53% during the years of observation.

The carbon isotopic composition of lacustrine biogenic carbonates depends primarily on the $\delta^{13}C$ of DIC in lake water. Observations of δ^{13} C of DIC in groundwater or lake water are not available for Lake Locknesjön, but some general assumptions can be made. The δ^{13} C of groundwater DIC is dependent on the open or closed system conditions, which determine the relative contributions from bedrock carbonates and soil air CO₂ (Clark and Fritz, 1997). The δ^{13} C of carbonate bedrock in the catchment area, i.e., resurge deposits in the Lockne Crater and post-impact sediments, is close to -0.5%(Ormö et al., 2010). The δ^{13} C of microbially respired soil air CO₂ reflects that of the local vegetation, which can be estimated to be around -25% based on tree ring cellulose data from the study area (unpublished data). It is increased by $\sim 4\%$ through fractionation by diffusion as CO₂ outgasses along the pCO_2 gradient between the soil and the atmosphere (Cerling et al., 1991). As a result of the well-mixed character of groundwater recharging the lake as evidenced by water isotope monitoring data (Fig. 5), likely reflecting a relatively long residence time, seasonal variations in δ^{13} C of ground-water DIC are probably small or absent.

On the contrary, DIC in lake water usually shows substantial seasonal variations, increasing in the epilimnion during the stratified warm period and decreasing to the lake-wide mean δ^{13} C during the holomixis (Decrouy et al., 2011a; von Grafenstein et al., 1999b). In Lake Locknesjön, warm season increases in δ^{13} C are most likely mainly induced by the photosynthetic activity of *Chara* (cf. Sect. 3.2.2), which can be assumed to also raise the mean δ^{13} C of lake water DIC compared to groundwater.

Chara hispida encrustations exhibit a significantly higher δ^{13} C than ostracod and *Pisidium* valves (Fig. 6a). While the δ^{13} C of *Pisidium* and ostracod calcite reflects the δ^{13} C of DIC, the δ^{13} C of *Chara hispida* encrustations is influenced by fractionation between DIC and the organic matter produced during photosynthesis. Juvenile *Candona* show a slightly higher δ^{13} C than adult *Candona* and *Pisidium*, which suggests that DIC is slightly enriched in ¹³C by *Chara* photosynthesis in summer when juvenile ostracods calcify their valves (Fig. 8). However, this effect on juvenile ostracods is much weaker than on the carbonate precipitated at the site of photosynthesis, i.e., in the *Chara hispida* encrustations. No other effects related to seasonal changes in δ^{13} C of DIC or lake water temperature on the δ^{13} C of biogenic carbonates can be inferred from our data.

Although probably minor in the carbonate-rich and therefore well-buffered system of Lake Locknesjön, seasonal variations in lake water pH may contribute to the observed interspecies differences in δ^{13} C and δ^{18} O. The pH measured at depths above 8 m ranges from 8.05 to 8.33, the maximum occurring in August. In cultured foraminifera, a slight lowering of δ^{18} O and δ^{13} C has been observed in response to increasing pH (Spero et al., 1997). Field observations of sediment trap calcite and lake water pH confirm this relationship (Teranes et al., 1999). However, further measurements of lake water pH and contemporaneously precipitated calcite would be needed to investigate the influence of pH variations on biogenic carbonates in Lake Locknesjön.

An additional factor that could play a role for interspecies differences is the microhabitat of the respective species. The most prominent spatial change in microhabitat at our sampling sites is the disappearance of *Chara* below 7 m depth. No influence of this change is evident on the carbonate isotopic compositions of the other species (Fig. 9). Furthermore, groundwater inflow may lead to local decreases in temperature and δ^{18} O of the lake water (cf. Sect. 2.1), but such inflows were not identified in the direct vicinity of the sampling sites, and no local decreases in carbonate δ^{18} O related to groundwater inflow were observed. On the contrary, the example of Lake Blektjärnen shows that the δ^{13} C of different *Chara* species can be influenced by their microhabitat, and that consequently a distinction at the species level is necessary for the interpretation of isotope records.

4.4 Uncertainties in water temperature reconstruction

Water temperature measurements contemporaneous with the formation of the analyzed biogenic carbonate samples are not available. Nevertheless, the range of observed seasonal water temperature changes in previous years (Fig. 3) is consistent with the reconstructed water temperature ranges based on mean lake water δ^{18} O and mean carbonates δ^{18} O for the respective species (Fig. 10). The highest reconstructed temperatures were obtained from *Chara hispida* calcifying in summer and the lowest temperatures from adult *Candona* calcifying in winter. This observation confirms that the differences in carbonate δ^{18} O between species (after correcting for vital offsets) are driven by varying water temperatures during their respective periods of calcification.

For a given carbonate δ^{18} O value, reconstructed temperatures calculated based on minimum and maximum observed lake water δ^{18} O values yield a range of 1.9 °C (Fig. 10). The variability in δ^{18} O between individual carbonate samples, on the contrary, has a larger influence on the reconstructed temperature than the variability in lake water δ^{18} O. Temperatures calculated based on the mean δ^{18} O of lake water samples and the minimum and maximum values of carbonates from the respective species yield temperature ranges of 3.6 °C for juvenile Candona candida, > 5 °C for all other ostracod instars, and 21.0 °C for Chara hispida (Fig. 10). The lowest reconstructed water temperatures, near 0 °C for adult Candona, match the lowest observed lake water temperatures at Lake Locknesjön. The maximum reconstructed water temperature of 29°C for Chara hispida, however, is rarely reached in maximum daily air temperatures in the study area, and therefore impossible to occur in the open water column.

A part of these large ranges in reconstructed temperatures is probably related to temporary variations in shallow water temperature that greatly exceed those indicated by the observed annual minimum and maximum values (Fig. 3). In particular, calcification of *Chara* encrustations may take place in secluded microhabitats at anomalously high water temperatures in summer, potentially also involving oxygenisotope disequilibrium effects (Pentecost et al., 2006). In addition, deviations from normal seasonal calcification patterns may occur. For example, summer generations of adult ostracods have been observed, likely as a consequence of thunderstorms, leading to temporary cooling of surface water (von Grafenstein et al., 1999b).

4.5 Implications for paleoclimate studies

Our data exhibit relatively large isotopic variability within one species (Fig. 6a), based on samples collected either from living organisms or from their subfossil remains in surface sediments, i.e., representing calcification during a time interval of a few years in the recent past. Analytical techniques today allow the analysis of small quantities of carbonate corresponding to, e.g., a few ostracod valves or a single stalk of a *Chara* encrustation. However, our results demonstrate that the range of reconstructed temperatures from such individual samples is large (Fig. 10), whereas the temperature reconstructed from an average of all samples per species/instar is in good agreement with observations. Consequently, it must be assured that isotopic records obtained from lacustrine biogenic carbonates for the study of past climate changes yield representative values for the time span covered by each analyzed sediment interval, through either (i) sufficiently large numbers of individual samples, or (ii) homogenized samples incorporating sufficiently large numbers of specimens.

The observed differences between our investigated species highlight the importance of performing isotopic measurements on well-identified remains of specific biogenic carbonate components in lake sediments rather than on bulk carbonate. Isotope records obtained on bulk sediment may reflect changes in the relative contributions of carbonate from different species and variations in detrital carbonate deposition that are not directly related to the environmental variables under study, thus adding complexity to interpretations (Hammarlund and Buchardt, 1996). For example, Chara encrustations are clearly the dominant component of the coarse fraction of the surface sediments of Lake Locknesjön at the sampled locations at water depths of up to 8 m. Changes in lake level at a coring site could lead to disappearance of Chara if wave and ice action (in shallow water) or lack of light (in deep water) hinder their growth, which would shift bulk carbonate δ^{18} O to higher values. If bulk carbonate isotope records are complemented with contemporary monitoring data and proper characterization of the sediment composition, their cautious interpretation may still yield important information. However, the isotopic analysis of specific carbonate components, preferably involving multiple taxa, can provide more detailed insight into past changes in climate because of their differences in the timing of calcification, and consequently water temperature and water δ^{18} O.

Past changes in water depth, caused by sediment accumulation and compaction or hydrological changes, may also influence isotopic records by affecting bottom water temperatures independently of other changes in climate. However, at Lake Locknesjön, potential changes in water depth in the range of 1 to 8 m do not seem to exert any significant influence on the isotopic composition of biogenic carbonates under modern conditions. We can expect that isotope proxy records from sediment cores at this site remain unaffected by possible water depth changes in this range in the past, as long as the sediment surface remains above the summer thermocline.

5 Conclusions

This study presents a modern snapshot of the variability of the carbon and oxygen isotopic compositions of lacustrine biogenic carbonates from living organism and from their sub-



Figure 10. Theoretical temperature calculated from water and carbonate δ^{18} O, assuming the carbonate was precipitated in isotopic equilibrium with the water (according to Kim and O'Neil, 1997). Temperatures are calculated using the mean water δ^{18} O and the range of measured carbonate δ^{18} O values (minimum, mean and maximum) for each species and instar (orange diamonds), as well as using the mean carbonate δ^{18} O of each species and instar and the range of observed water δ^{18} O values (dark blue circles). The measured carbonate δ^{18} O values have been corrected for known vital offsets.

fossil remains in the surface sediments of Lake Locknesjön, which allows us to draw the following conclusions.

- 1. We constrained the controls on the isotopic composition of biogenic carbonates in Lake Locknesjön. The large differences between species are mainly caused by species-specific vital offsets. Additional differences are caused by the timing and length of the calcification periods of the respective species, being exposed to seasonally changing water temperatures. In contrast, seasonal and interannual variations in lake water δ^{18} O are small, at least over the time span of observation (2013–2014 and 2018). The influence of changing lake water δ^{18} O on carbonate δ^{18} O is outweighed by the opposing effect of changing water temperature (see Fig. 1 for different influences of temperature on carbonate δ^{18} O).
- 2. Based on the understanding of these environmental controls, it is possible to estimate seasonal water temperature changes from the δ^{18} O of lake water and of specific biogenic carbonates, provided that components formed during different seasons but sampled from the same sediment layer are analyzed individually. This approach is particularly applicable for groundwater-fed lakes with long residence times like Lake Locknesjön, where lake water δ^{18} O varies within a relatively narrow range. Under such conditions, seasonal changes in lake water temperature are clearly reflected in the δ^{18} O of multiple biogenic carbonates through the temperature effect on isotope fractionation between water and carbonate. Water temperature changes could also exert the dominant control on carbonate δ^{18} O on longer (decadal to millennial) time scales, where significant changes in air temperature can be expected. However, in that case additional factors must be considered, such as the effects of changes in δ^{18} O of atmospheric precipitation brought about by related or independent climate dynamics, or potential variations in evaporative enrichment in the δ^{18} O of the lake water caused by a changing ratio of evaporation to inflow.
- 3. Our results have implications for the design of subsampling strategies aiming at reconstructions of past changes in environment and climate based on isotopic analyses of sediment cores. The intraspecies variability in δ^{18} O and δ^{13} C of biogenic carbonates highlights that care must be taken to obtain representative subsamples of a species for each time interval, especially if working with shallow water environments where water temperature can change rapidly over a short time. The interspecies variability observed in Lake Locknesjön adds to the growing body of evidence demonstrating the importance of performing measurements on specific carbonate components of lake sediments rather than on bulk carbonate. Sampling strategies involving multiple specific biogenic carbonate components are likely to produce more detailed and comprehensive environmental reconstructions, including records of seasonal water temperature variability.

Data availability. Water isotope data obtained in this study are available at https://doi.org/10.1594/PANGAEA.942628 (Labuhn et al., 2022a). Carbonate isotope data are available at https://doi.org/10.1594/PANGAEA.942627 (Labuhn et al., 2022b).

Video supplement. A video showing the movement of sediment on the lake floor at a groundwater inflow is available at https://doi.org/10.5446/56815 (Labuhn et al., 2021).

Author contributions. IL and UvG designed the study and coordinated the field work. Samples were prepared by FT and UvG. Water isotope measurements were performed by BM, carbonate isotope measurements by HK. FT and IL created the figures. IL and FT wrote the paper, with contributions from DH, UvG and HK. All co-authors contributed to the interpretation and discussion of the results. DH and UvG were principal investigators of the project within which this study was conducted. *Competing interests.* The contact author has declared that neither they nor their co-authors have any competing interests.

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Acknowledgements. This study would not have been possible without the scientific diving team of the Limnological Research Station Iffeldorf, Technical University of Munich, Germany: Benedikt Beck, Matthias Galm, Jan Leidholdt and Stefan Zimmermann. The authors are grateful to Östersund Diving School for arranging the diving equipment, to Malin Bernhardsson at the County Administration Board of Jämtland for providing a boat as well as water temperature and pH data from the Triple Lakes project, to Anna Eklund at the Swedish Meteorological and Hydrological Institute for providing archived water temperature data and to Sten-Åke Jonsson for collecting water samples. Wolfgang Bevern, Birgit Meyer and Maike Steinkamp are thanked for helping with the carbonate isotope measurements.

Financial support. This study was funded by the Swedish Research Council as part of the French-Swedish Common Research and Training Programme on Climate and the Environment (grant no. 349-2012-6291) and by the Central Research Development Fund of the University of Bremen (grant no. 2018/ZF08/FB8/Labuhn).

The article processing charges for this open-access publication were covered by the University of Bremen.

Review statement. This paper was edited by Ny Riavo G. Voarintsoa and reviewed by Ola Kwiecien and Karina Apolinarska.

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