Lithium and Sr isotopic composition of salar deposits in the Central Andes across space and time: the Salar de Pozuelos, Argentina

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Abstract
The Central Andes of South America host the largest known lithium resources in a confined area, but the primary lithium sources of the salar deposits and the mobilisation process of lithium are still a matter of speculation. Chemical weathering at or near the surface and leaching in hydrothermal systems of the active magmatic arc are considered the two main mechanisms of Li extraction from the source rock. The lithium and strontium isotope composition of typical salar deposits offer insights into the processes on how Li brine deposits in Andean evaporites are formed. Data from the Salar de Pozuelos indicate near-surface chemical weathering in a cold and dry climate as the dominant mobilisation process of Li, with evaporation being responsible for the enrichment. The Cenozoic ignimbrites are the favoured source rock for the Li, with subordinate additions from the Palaeozoic basement. The identification of the source rocks is supported by radiogenic Nd and Pb and stable B isotope data from salar deposits. A comparison with other Li brine and salt deposits in the Altiplano-Puna Plateau and its western foothills places the Salar de Pozuelos as an endmember of Li solubilisation by chemical weathering with only minor hydrothermal mobilisation of Li.

Keywords Central Andes · Salar deposits · Lithium isotopes · Lithium deposits

Introduction
The arid climate and active tectonics formed and sustain numerous Cenozoic endorheic basins on the Altiplano-Puna Plateau of the Central Andes in NW Argentina, in southern Bolivia, and in the western slope of the Plateau in northern Chile between 20 and 27°S (e.g. Hartley and Chong 2002; Strecke et al. 2007; Quade et al. 2015). An endorheic basin containing a salt lake or a salt pan is called salar in the Andes. The continental sedimentary successions host commonly variably thick evaporite deposits dominated by halite, widespread occurrence of gypsum, and travertine deposits at active or fossil hydrothermal springs (e.g. Stoertz and Ericksen 1974; Alonso et al. 1991; Vandervoort et al. 1995; Risacher and Fritz 2009). The formation of the principal minerals from solutes and the hydro-chemical systematics of evaporite deposits are well understood. Lithium (Li) and boron (B) in brine and borate minerals are of economic interest, but play no or no major role in the formation of the thick evaporite deposits (e.g. seminal work summarised in Hardie and Eugster 1970; Eugster 1980; Warren 2010).

Some of the Andean salars host economic enrichments of boron minerals (Alonso 1999) and world-class brine pool deposits of lithium, presently sourcing approximately 30% of the world’s lithium production and holding more than 50% of the world’s Li resources (Grosjean et al. 2012; Houston et al. 2011; Munk et al. 2016; U.S. Geological Survey 2020). The occurrence of lithium enrichment in the Central Andes is geographically confined to the ‘Lithium Triangle’, an informal term used by the mining industry, which refers to the area between southern Bolivia, NW Argentina and NE Chile.

Boron forms a number of borate minerals within the Andean evaporite deposits and received considerable interest of the mining industry. Its concentration, distribution...
To improve our understanding on how Li brine deposits in Andean evaporites are formed, we analysed the Li and Sr isotope composition of brines, salts and intercalated siliciclastic rocks from the surface and drill cores of the Salar de Pozuelos on the Puna Plateau, NW Argentina. The Salar de Pozuelos is known as a Li brine deposit and has been subject to extensive exploration since 2008, allowing access to surface and drill core sampling. The salar basin and its catchment area is well mapped (Fig. 1; ESM1 Fig. A1), and there is no known complex hydrology or permanent water supply. The catchment area is dominated by siliciclastic marine and continental sedimentary rocks of Ordovician and Palaeogene age, with only minor outcrops of volcanic rocks. This is in contrast to other studied salars in the Central Andes, such as Salar de Hombre Muerto, Salar Olaroz, Salar de Ratones and Salar Centenario (Godfrey et al. 2013; Orberger et al. 2015; Garcia et al. 2020) with widespread outcrops of Cenozoic volcanic rocks (e.g. Schnurr et al. 2006). In order to extend the existing dataset on salar deposits and springs in the Central Andes and the Li-Triangle, we also performed a survey study on Li and Sr isotopes in surface brines and salts from salars from the Li-Triangle in northern Chile and south Bolivia (ESM2 Fig. A1). The Li and Sr isotope signatures from the Salar de Pozuelos deposit are compared with the isotope signatures of potential source rocks of the region, i.e. Cenozoic volcanic rocks, Palaeozoic (Pz) metamorphic, magmatic and sedimentary basement, and their younger sedimentary equivalents in the catchment area of the Salar de Pozuelos (this work; Meixner et al. 2020) and are discussed in the frame of the new and published data from salars of the Li-Triangle (Godfrey et al. 2013, 2019; Godfrey and Álvarez-Amado 2020; Orberger et al. 2015; Munk et al. 2018; García et al. 2020). We also analysed the B isotope composition of the surface brines in order to place the Salar de Pozuelos into the framework of salars from the Puna Plateau and their source rock contribution (Kasemann et al. 2004).

Geological setting of the Salar de Pozuelos

The (semi)arid climate and the formation of endorheic basins with evaporite deposits started around 15 Ma in the Puna Plateau and is ongoing since then (e.g. Alonso et al. 1991; Vandervoort et al. 1995; Strecker et al. 2007; Quade et al. 2015). The vegetation is sparse, and bare rock is the dominant surface in this high altitude cold desert (precipitation <200 mm; Haselton et al. 2002; Garreaud et al. 2009). While physical weathering predominates, in the summer season when water is available, chemical weathering can be a major component of the weathering regime (e.g. Hall et al. 2002). The dominant near-surface rock units of the region are Cenozoic volcanic rocks,
mainly andesite lavas from stratovolcanoes and monoge-meric eruption centres and felsic, large-volume ignimbrites, Palaeozoic (Pz) metamorphic, magmatic and sedimentary basement and their younger sedimentary equivalents (e.g. Reutter and Munier 2006; Schnurr et al. 2006). The main sources of siliciclastic material in these basins relate to the geology of the catchment areas, confined by the surrounding topographic highs. Input of potential air-fall from the active Cenozoic magmatic arc can also originate from outside the catchment areas.
The salars in the Puna Plateau are classified as mature halite and/or immature clastic salars. A decreasing clastic sedimentation and finer grain sizes towards the centre of the salar, followed by a thick evaporite sequence in the sedimentary column is a characteristic of a mature salar. Alternating sequences of fine-grained sediments and evaporitic beds are typical for immature salars (Houston et al. 2011; Vinante and Alonso 2006).

### Lithology of the catchment area

The Salar de Pozuelos is located in the Puna region of northwest Argentina, in the western Salta province, approximately 230 km west of the city of Salta and 150 km east of the Chilean border. The Salar de Pozuelos basin has a reported salar surface area of 82 km$^2$ with the salt flat at approx. 3800 m elevation and a catchment area of 350 km$^2$ (most recent estimates: Hains Engineering Company Limited 2018, investor report). The drainage network of the Salar de Pozuelos basin includes no permanent tributaries, but major surface water flows are expected from the southern and northern sub-basins during pluvial events that accumulated recent fluvial deposits (Fig. 1; ESM1 Fig. A1). The description of the catchment’s lithology follows the geological map and accompanying explanations (Fig. 1; Blasco et al. 1996; Martínez et al. 2018, 2020). The catchment area is dominated by (1) siliciclastic Ordovician marine sedimentary rocks (Copalay Formation) that comprise shales, silt- and sandstones with no or very low-grade metamorphic overprint, and (2) much younger continental sedimentary rocks (mostly red beds of sand, silt and clays) of Eocene (Geste Fm.) and Oligocene (Vizcacheras Fm.) age that were deposited in local fluvial systems and comprise local debris and detritus and intercalated volcanic rocks (tuff layers; Blasco et al. 1996). The Miocene Pozuelos Formation also includes evaporites that formed in a playa depositional environment of an endorheic basin and (3) Quaternary sediments reflect local rock variability (Blasco et al. 1996) and are deposited as alluvial and colluvial fans in the catchment area, and—with shallower morphological gradient—as sheet sands and playasilts and clays at the margin of the salar basin (Martínez et al. 2018). Subordinate Miocene igneous rocks (composed of ignimbrites and volcanic tuffs) occur in the south of the catchment area of the Salar de Pozuelos as isolated intrusions and in the north as intercalations in Miocene sedimentary rocks (Blasco et al. 1996). Apart from the presence of volcanic ash layers in the young sediments, other volcanic detritus, produced from aeolian erosion, transport and deposition can be assumed from the persistent volcanic activity and distribution of ignimbrite in the wider surroundings (e.g. Trumbull et al. 1999; Schnurr et al. 2007; Kay et al. 2010; Brandmeier and Wörner 2016; Wörner et al. 2018), but there is no volcanic edifice in the catchment area. The sedimentary and volcanic cover is underlain by metamorphic, magmatic and sedimentary rocks of the Early Palaeozoic basement that reworked Proterozoic rocks, sourced Palaeozoic sedimentary rocks and magmas, and is the principal crustal component in Cenozoic magmatic rocks of the Andean continental arc (e.g. Trumbull et al. 1999; Lucassen et al. 2001; Kay et al. 2010).

### The salar basin deposits

Salar de Pozuelos is classified as a mature salar with some immature (clastic) characteristics in the northeast of the salar. The surface of the salar is characterised by different types of salt crusts (Fig. 1; ESM1 Fig. A2; Martínez et al. 2020) that developed in different areas depending on the influence of surface runoff from the surrounding catchment area. A hard and rough salt crust is predominant and covers most of the central part of the salar that is distant from surface runoff. A smooth salt crust is developed at the margin comprising patches of a hard salt crust with polygons and a substantial amount of siliciclastic silt and clay component indicating temporary flooding by surface water (Igarzábal 1991; Martínez et al. 2018). Borates (as ulexite) are interbedded as thin beds in the upper two meters of clastic sediments in the north-eastern and western sector of the Salar de Pozuelos basin with immature (clastic) characteristics (e.g. Alonso 1999).

Hot spring activity has ceased in the Salar de Pozuelos, but a travertine field of former Pleistocene hot springs (<100 ka) can be found along the faulted contact between the Ordovician and Paleogene rocks in the northeast sector (Martínez et al. 2018). Currently, only minor spring activity (20 °C) is present at the border of the salar basin (Hains Engineering Company Limited 2018).

The basin infill of the Salar de Pozuelos is known from prospection drill holes. A detailed facies description for the salt pan evaporites and saline mudflat sediments down to ca 183 m depth in the centre of the salar is provided by Martínez et al. (2018) for the cores L01 and DDH1 that we used in this study. It starts from the surface with approximately 95 m of halite comprising scarce siliciclastic components. The halite is coarse crystalline and highly fractured at the top and trends into more competent to massive halite at the lower part. The basal unit is essentially a mixed red clay (clay to fine sandstone) and halite (subordinate) zone that has been traced in a diamond drill hole to 183 m depth (Martínez et al. 2018) and continues to the bottoms of the deepest drill holes at ca 320 to 383 m without reaching the Ordovician sedimentary rocks. Lithium enriched brines are found within the halite matrix and are expected to be present...
down to the base of the salar (Hains Engineering Company Limited 2018; LSC Lithium Cooperation 2019).

**Survey sampling in the Li-Triangle between 18 to 26°S (NW Argentina, southern Bolivia, northern Chile)**

Additional samples of natural brines and spring water from a variety of salars mainly in northern Chile, but also from Salar Uyuni, southern Bolivia and Salar de Antofalla, NW Argentina were collected (for locations, see ESM2 Fig. A1). This was done to broaden the existing database on Li and Sr isotope composition from published detailed studies on single salar systems. All sample locations are within the active magmatic arc with catchment areas that are strongly influenced by the volcanism. Only the Salar de Atacama basin is off the arc and located at the base of the western slope of the Puna Plateau, but it also collects mainly the runoff of the latter, i.e. arc material with extensive ignimbrites of Neogene age (Ramírez and Gardeweg 1982; Reutter et al. 1994).

**Materials and methods**

**Surface samples from the Salar de Pozuelos and the catchment area**

A total of 19 locations of surface outcrops were sampled in February 2011 (Table 1). Brine or water samples \( (n = 13) \) were taken from natural ponds or mechanically dug exploration test pits around the margin and the centre of the salar. Sampling of brines around the salar near the border between the evaporite and outcropping rocks aims at recognising potential compositional variations in the brines related to the influx from local and variable sources (for sample points (sp), refer to Fig. 1 and Table 1; regional overview, ESM2 Fig. A1). In brine ponds close to the southern border \( (n = 1\), sample point sp 10 \) and from the centre \( (n = 2\), sp 1 and 3 \), coexisting brine and salt precipitates were collected in order to constrain potential fractionation of Li during evaporation and salt precipitation. Additional samples of salt \( (n = 2\), sp 2 \) were collected from evaporated brine in an artificial pool at the centre of the salar basin. Two borate (ulexite) samples \( (sp \ 13 \text{ and } 18) \), were sampled from the eastern and western border of the salar, respectively. Unconsolidated siliciclastic sediments were collected from the natural ponds or exploration test pits \( (n = 7\), sp 4, 5, 7 to 9, 11, 13, 16 and 18 \). Furthermore, we sampled fluid from an active spring \( (sp \ 14) \) at the salar’s north-eastern border located within the faulted contact between the Ordovician and Paleogene rocks. The outflow of the spring is presently low, but past higher activity formed a travertine field since the Pleistocene. The travertine (sample point sp 14; sample SP-11–24) was split into 8 subsamples according to the discernible layering to test for compositional changes with time of deposition. Photographs of various sample locations are provided in the supplement (ESM1 Figs. A2 and A3). Salts (halite), brines and sediments were stored in acid-cleaned PE bottles. Silicate rocks from the catchment area of Pozuelos comprise Oligocene (sp 6) and Eocene (sp 15) sedimentary rocks, one Cenozoic ignimbrite (sp 19) and one Ordovician sedimentary rock (sp 17). The latter two samples were already analysed (Meixner et al. 2020). One unconsolidated siliciclastic sediment (sp 12) was collected from a dry riverbed close to Oligocene and Ordovician outcrops.

**Drill cores**

Sampling of two adjacent (11 m distance) commercial drill cores (mining company Ekeko S.A. and Lithea Inc. in 2015) in the centre of the salar (Fig. 1) aims at compositional variations in time. All salt (halite, \( n = 11 \)) samples have been selected from core L01 (SPZ RC001 in the company repository) in approximately 5 to 15 m steps down to 95.5 m. Core DDH1 (SPZ DDH001 in the company repository) has a lower recovery in the halite zone, but comprises in its lower section from 95 m down to 183.5 m a siliciclastic sedimentary sequence with an evaporite (salt) component. Mixed evaporite-siliciclastic samples \( (n = 6) \) were retrieved from DDH1 in approximately 10- to 20-m steps and further separated into three fractions: clay rich, sand rich and water soluble. The respective position of the samples in the cores below surface is given in Table 1. Surface sediment (sample SP-11–07) sampled as salt clay from the centre (sp 3) was also analysed and is treated together with the core.

**Samples from the lithium Triangle**

Additional thermal water, brine and salt samples were collected from thermal springs, salars and lagoons/brine pools in the Lithium Triangle (ESM2 Fig. A1). Sampling includes in northern Chile the Salar Grande, Salar de las Parinas, Salar de la Isla, Salar de Pedernales, Salar de Atacama and Salar de Piedra Parada, the Lagunas de Huasco and Chaxa, and thermal springs from Quebrada Chaco del Norte, Quebrada del Rio Negro and Termas Chiriguaya. Furthermore, two salt samples were collected from the southern end of the Salar de Antofalla and from a salt pond in an ignimbrite field at Campo Negro, both from NW Argentina. Two samples are from the northern and eastern part of the Salar de Uyuni in southern Bolivia.
Sample preparation

All brine and thermal water samples were centrifuged and aliquoted for isotope analyses, and additionally filtered through an unreactive 0.2 μm Minisart® syringe filter for concentration analyses. Salt samples (1 g) from the surface and the drill core (L01) were dissolved in 50 mL, 18 MΩ water and subsequently prepared in the same way as brine samples for isotope and concentration analyses. The borate samples were dissolved in 1 N HCl. The travertine

| Sample point | Coordinates | Brine/water | Sample type | Travertine | Catchment | Description |
|-------------|-------------|------------|-------------|------------|-----------|-------------|
| 1           | 24°43′02.1″S 66°49′30.7″W | SP-11–01 | SP-11–02 | Centre of salar brine and salt sample from the same pond |
| 2           | 24°43′00.4″S 66°49′49.1″W | SP-11–03 | SP-11–04 | Evaporated brines in artificial pool, with pure salt remaining, 03—cubic, 04—flake/dendritic crystal structure |
| 3           | 24°43′00.8″S 66°50′25.0″W | SP-11–05 | SP-11–06 | Brine, salt and clay sample from the same pond |
| 4           | 24°42′28.1″S 66°51′17.4″W | SP-11–08 | SP-11–07 | Western part of salar, brine sample from a pond with minor salt |
| 5           | 24°42′11.8″S 66°51′53.0″W | SP-11–10 | SP-11–11-A | Western part of salar |
| 6           | 24°42′00.3″S 66°52′08.1″W | SP-11–12 | | Oligocene sediment samples; mica rich red sandstones |
| 7           | 24°42′12.2″S 66°50′59.4″W | SP-11–14 | SP-11–11-B | Western part of salar, brine sample and mixture of organics, borates and clay from a test pit |
| 8           | 24°43′01.9″S 66°47′08.7″W | SP-11–15 | SP-11–11-C | Eastern part of salar, brine sample from a pure clay pond |
| 9           | 24°45′58.9″S 66°50′26.7″W | SP-11–16 | SP-11–17 | Southern part of salar |
| 10          | 24°45′11.0″S 66°50′42.9″W | SP-11–18 | SP-11–19 | Southern part of salar, brine and sediment sample from the same pond |
| 11          | 24°44′38.7″S 66°52′11.2″W | SP-11–20 | SP-11–20 | Southern part of salar, sediment sample from a dry river bed |
| 12          | 24°43′49.2″S 66°52′47.2″W | SP-11–21 | SP-11–22* | Western part of salar, sediment sample from a dry river bed |
| 13          | 24°40′49.5″S 66°46′45.6″W | SP-11–23 | SP-11–24 | North-Eastern part of salar, brine sample, no associated salt but ulexite* |
| 14          | 24°40′15.7″S 66°46′33.4″W | SP-11–25 | | North-Eastern part of salar, water sample from a former hydrothermal spring and associated travertine |
| 15          | 24°40′34.2″S 66°44′50.3″W | SP-11–26 | | Eocene sandstone |
| 16          | 24°40′09.5″S 66°49′13.2″W | SP-11–27 | SP-11–28 | Western part of salar, brine samples with no associated salt, sediments are sulfur rich |
| 17          | 24°41′19.6″S 66°50′29.7″W | SP-11–29 | | Ordovician sedimentary rock |
| 18          | 24°41′23.4″S 66°50′23.8″W | SP-11–30 | SP-11–32* | SP-11–31-A | Western part of salar, brine samples with no associated salt but ulexite*, sediments are organic rich |
| 19          | 24°37′50.2″S 66°46′44.2″W | SP-11–31-B | SP-11–34-A | SP-11–34-B | Cenozoic volcanic rock (CVZ), ignimbrite A-fresh, B-altered |
| 20          | 24°43′00.5″S 66°49′49.5″W | L01 | SP-11–34-C | | Drill core (SPZ RC001), pure salt fraction |
| 21          | 24°43′00.5″S 66°49′49.2″W | DDH1 | | Drill core (SPZ DDH001), salt sand and clay fraction |
samples were crushed, powdered, repeatedly washed in 18 MΩ water and dried at 110 °C, and about 100 mg of each sub-sample were dissolved in 1 N HCl. The sample solution was centrifuged and aliquots were taken for Li, B and Sr analyses. Sediment samples were disintegrated with a hammer on a steel plate. Approximately 50 g of clean fragments were collected, washed with 18 MΩ water, dried at 110 °C and powdered in an agate ball-mill. The powders were repeatedly washed in 18 MΩ water, dried at 110 °C and reweighed. The water-soluble fraction (leachate) was saved, dried, dissolved in HCl and aliquoted for Li and Sr isotope analyses. Soft sediments from the salar can contain up to 32% mass fraction of water-soluble salts. Evaporite—siliciclastic mixtures from the core (DDH1) were separated by suspending and centrifuging the sediment repeatedly in 18 MΩ water. The siliciclastic fraction was dried at 110 °C and reweighed. The water-soluble mass fraction of the drill core samples ranges between 4 and 17%. The leachate was dried, redissolved in 18 MΩ water and treated as the brines described above. The elatic fraction was again suspended in 18 MΩ water, and carefully centrifuged to separate a coarse-grained, sand-rich and a fine-grained, more clay-rich fraction (hereafter sand-rich and clay-rich). The clay-rich fraction is the upper most layer in the centrifuge tube, i.e. the bulk of the samples is the sand-rich fraction. After drying, the clay-rich and sand-rich fractions were collected and separately analysed. All washed, leached and separated silicate samples were homogenised in an agate mortar before further digestion. For Li and radiogenic isotopes and elemental analyses, all silicate samples were digested in a 5:1 mixture of triple distilled HF and double distilled HNO₃ at 140 °C during at least 48 h. The sample solution was dried at 80 °C, completely re-dissolved with 2.5 M HNO₃, cooked for 8 h at 140 °C and dried at 85 °C and finally dissolved in 0.15 M HCl (Meixner et al. 2020).

**Major and trace element analyses**

Concentrations of cations were determined by inductively coupled plasma optical emission spectroscopy (ICP-OES) using an Agilent Technologies 700 Series ICP-OES at the University of Bremen (Sediment Geochemistry Group at MARUM). The analytical uncertainty for Li and B determination was found to be <0.1%, <2% for Ca, K, Mg and Sr and <4% for Na⁺ (2 RSD). Apart from the Salar de Uyuni, all Li and B concentrations for fluid and salt surface samples were analysed by ICP-OES. Lithium and boron concentrations of the catchment area, drill core and travertine samples were determined during the course of isotope ratio measurements, using a ThermoScientific Neptune Plus multicollector-inductively coupled plasma-mass spectrometer (MC-ICP-MS) at the University of Bremen, with an uncertainty of <6% (2RSD) based on long term analyses of reference material ZGI-TB (for lithium) and IAEA-B5 (for boron).

**Isotope analyses**

Analyses of hydrogen and oxygen isotopes (δD and δ¹⁸O) were carried out at the Stable Isotope Laboratory at MARUM-Centre for Marine Environmental Sciences, University of Bremen by Isotope Ratio Mass Spectrometry using a ThermoFisher Scientific TC/EA (modified after Gehre et al. 2004) coupled via a ConFlow IV to a ThermoFisher Scientific MAT253. The results are reported relative to four international standards (IAEA-OH 13–16) of the International Atomic Energy Agency with a precision of ±3.0‰ for δD and ±2.7‰ for δ¹⁸O (1sd).

Stable isotopes of Li and B and radiogenic isotopes of Sr, Nd and Pb were analysed in the laboratories of the Isotope Geochemistry Group at MARUM-Centre for Marine Environmental Sciences, University of Bremen. Lithium in aliquots of fluid, salt and silicate rock samples was quantitatively separated from its matrix in a two-step procedure using BioRad AG 50 W-X8 resin (details in Meixner et al. 2020). Li loss during the separation procedure can result in isotopic offsets, and therefore the tails of each separation step were analysed for Li concentrations. Li loss was less than 0.5% of the aliquot in all samples and thus without influence on the Li isotope composition. Reference materials (RM), standard material and the procedural blank were processed together with each sample series for quality control. Li concentrations of sample and RM solutions in 2% HNO₃ were closely adjusted to 25 µg/L Li (±5%). Li isotope ratios were measured in the standard-sample bracketing mode on a MC-ICP MS Neptune plus (ThermoScientific) using the stable introduction system and a high efficiency X-cone. The solvent 2% HNO₃ was measured before and after each standard or sample and the average of both analyses was used as the analytical baseline for correction of the intensities on masses ⁷Li and ⁶Li. The baseline-corrected isotope ratios are reported in the conventional δ⁷Li notation relative to the RM NIST 8545 (LSVEC Li carbonate, Flesch et al. 1973). The Li blank-input during chemical treatment was always less than 0.1% of the sample aliquot and had no effect on the isotope composition of the sample. Processed RMs were analysed in the same batch with the samples to verify the accuracy and precision of the analytical procedure. RM NIST 8545 shows a δ⁷Li value of −0.03 ± 0.18‰ (2sd, n = 18), and internal laboratory seawater standard BSW Susu Knolls has a δ⁷Li value of +3.0 ± 0.1‰ (2sd, n = 11), indicating no isotopic offsets during the whole analytical procedure. The analysed RM ZGI-GM (granite) (−0.7 ± 0.2‰; 2sd, n = 3), ZGI-BM (basalt) (−3.3 ± 0.2‰; 2sd, n = 2), ZGI-TS (black shale) (+4.9 ± 0.1‰; 2sd) and ZGI-TB-2 (shale) (−3.4 ± 0.1‰; 2sd, n = 2) are in line with published values (Meixner et al.
RM USGS-G-3 (granite) gives a value of $+0.1 \pm 0.5\%\,$ (2sd, $n = 2$). The external reproducibility is generally $\leq 0.5\%\,$ (2sd); the repeatability of the individual $\delta^{11}B$ values is reported as two standard deviations based on multiple mass-spectrometer analyses.

Boron was separated from the sample material using microsublimation as described in Hüpers et al. (2016). Boron isotope ratios for all samples were analysed on a MC-ICP MS using the same stable introduction setting and measurement procedure as for lithium and detailed in Hüpers et al. (2016). The analyses of sample material and RM were performed repeatedly in the standard-sample bracketing mode using unprocessed SRM NIST 951 with 100 pg/L boron concentration as reference standard. The isotope data were expressed in the delta notation relative to unprocessed SRM NIST 951 (standard boric acid). Accuracy and repeatability for the separation procedure and measurement was checked through NIST SRM 951 and the internal bottom seawater standard Susu Knolls. SRM NIST 951 shows $\delta^{11}B$ values of $+0.01 \pm 0.13\%\,$ (2sd, $n = 7$), indicating no isotope fractionation during sublimation. The $\delta^{11}B$ values of seawater IAEA-B1 ($+39.9 \pm 0.2\%\,$; 2sd, $n = 5$) and of our laboratory internal seawater standard BSW Susu Knolls (Eastern Manus basin) ($+39.8 \pm 0.5\%\,$; 2sd, $n = 5$) overlap within uncertainty with literature values on modern seawater ($+39.61 \pm 0.2\%\,$, 2sd$_{\text{mean}}$. Foster et al. 2010). Boron isotope composition of groundwater RM IAEA-B3 ($-21.96 \pm 0.02\%\,$; 2sd, $n = 1$) falls near the published values (e.g. Gonfiantini et al. 2003 compiled ($-21.33 \pm 0.9\%\,$, 1sd) and Mahlknecht et al. 2017 ($-21.2 \pm 0.4\%\,$, 2sd)). The repeatability of the individual $\delta^{11}B$ values is given as two standard deviations based on multiple mass-spectrometer analyses.

After general sample preparation, digestion (detailed above), and re-dissolution in 2 N HNO$_3$, chemical separation of Sr, Nd and Pb followed the methods presented in Höppner et al. (2018). Total procedural blanks are < 140 pg for Sr, < 80 pg for Nd and < 60 pg for Pb. Radiogenic isotope ratios of all sample materials were analysed on a thermal ionisation mass spectrometer Triton (ThermoScientific). Nd and Pb isotope ratios were analysed in the static multicollection mode and Sr isotope ratios in the dynamic multicollection mode. Instrumental mass fractionation during Sr isotope analysis was corrected using the stable isotope ratio of $^{86}\text{Sr}/^{88}\text{Sr}$ ($= 0.1194$). The analytical accuracy and external long-term reproducibility for $^{87}\text{Sr}/^{86}\text{Sr}$ of reference material NIST SRM 987 in this study was 0.710246 ± 0.000011 (2SD, $n = 24$; period: May 2015 to May 2017). This is within the range of published values analysed by TIMS of 0.710250 ± 0.000034 (2SD, $n = 1245$, data < 0.7102 and > 0.7103 are discarded) calculated from GeoRem database (query September 2017; http://georem.mpch-mainz.gwdg.de). Instrumental mass fractionation during Nd isotope analyses was corrected to $^{146}\text{Nd}/^{144}\text{Nd}$ ($= 0.7219$). The analytical accuracy and external long-term reproducibility for $^{143}\text{Nd}/^{144}\text{Nd}$ for reference material JNd-1 was 0.512093 ± 0.000009 (2SD, $n = 27$; period: May 2015 to May 2017) and matches the published values of 0.51206 ± 0.000027 (2SD, $n = 211$, data < 0.51204 and > 0.51217 are discarded) analysed by TIMS (GeoRem database; query September 2017; http://georem.mpch-mainz.gwdg.de). The $\varepsilon_{\text{Nd}}$ notation is reported relative to the Chondritic Uniform Reservoir (CHUR) with a value of $^{143}\text{Nd}/^{144}\text{Nd}$ = 0.512638 (Wasserburg and DePaolo 1979).

Instrumental mass fractionation for Pb isotope analysis was corrected using a factor of 1.001 (0.1%) per atomic mass unit based on the repeated analyses of NIST SRM 981. The reproducibility including the correction for mass fractionation is better than 0.1% of the respective ratio, which is the assumed uncertainty on the Pb isotope ratios. The uncertainty of the radiogenic isotope analyses is reported as two standard deviation of the mean (2sd$_{\text{mean}}$).

Results

All new data referred to in the results section are in the Tables listed below, in the supplement, and are also available through the World Data Center Pangaea (https://doi.org/10.1594/PANGAEA.928849, Meixner et al. 2021). Catchment area, salar surface basin, and drill core samples from Salar de Pozuelos were analysed for Li and Sr isotope composition and Li contents (Tables 2, 3, and 4). Surface brines and salts, and salts from core L01 were also analysed for their B isotope composition and elemental contents (Tables 3 and 4), plus O and H isotopic compositions in surface brines (ESM3 Table A1). In addition, Nd and Pb isotope compositions in silicate sediment samples from the catchment area, the salar and the drill core DDH1 were analysed (ESM3 Tables A2 and A3). We also included Li, Sr and B isotope composition and elemental contents of brine samples from salars and thermal springs in a survey study of locations within the Lithium Triangle (Table 5). For locations with solely salt samples, only the Sr isotope composition is available. Travertine Li, B and Sr isotope composition are in ESM3 Table A4. Elemental compositions of surface brine and salt (ESM3 Table A5), salt of core L01 (ESM3 Table A6), and brines of the Li-Triangle (ESM3 Table A7) are also documented.

Elemental composition of the surface brines and salts, and salts from core L01

In the brine, sodium comprises 76 to 96% (molar; most samples > 89%) of the analysed cations (Na, K, Li, B, Ca, Mg, Sr). The total load of dissolved cations as chloride ranges between 0.14 and 5.1 mol/l (ESM1 Fig. A4). The Li contents
Table 2 Isootope and concentration data for the catchment and the salar sediments

| Sample number | Washed rocks | Water-soluble fraction | Washed rocks | Water-soluble fraction |
|---------------|--------------|-------------------------|--------------|------------------------|
| SP-11–11-A    | Oligocene    | 4.3 (1) 29 0.714014 (8) | 14.7 (2) 394 | 0.710634 (8)          |
| SP-11–11-B    | Oligocene    | 4.9 (2) 34 0.718071 (7) | 10.0 (2) 1271 | 0.710555 (6)          |
| SP-11–11-C    | Oligocene    | 9.3 (4) 26 0.71394 (5)  | 22.9 (1) 39   | 0.711295 (7)          |
| SP-11–25      | Eocene       | 12.8 (1) 25 0.717522 (5)| 12.2 (2) 4.3  | 0.715592 (7)          |
| SP-11–29      | Ordovician   | −1.3 (2) 31 0.73130 (1) | 14.3 (2) 8.3  | 0.726735 (6)          |
| SP-11–34-A    | Cenozoic     | −2.5 (2) 24 0.71113 (6) | 9.8 (1) 4.8  | 0.710295 (6)          |
| SP-11–34-B    | Cenozoic     | −4.7 (1) 28 −            | 12.8 (1) 3.1  | 0.710246 (6)          |

Table 3 Isootope and concentration data for fluid and salt surface samples

| Sample number | Brine and water | Salt/Halite* | Borates |
|---------------|-----------------|--------------|---------|
| SP 11–01      | 17.2 (4) 960    | −19.3 (1) 470 | 0.712475 (3) |
| SP 11–05      | 17.5 (1) 1200   | −16.5 (1) 400 | 0.71191 (5) |
| SP 11–08      | 16.8 (3) 220    | −10.4 (1) 210 | 0.712471 (4) |
| SP 11–10      | 18.5 (3) 500    | −8.5 (1) 490  | 0.710801 (4) |
| SP 11–12      | 19.3 (2) 390    | −16.7 (1) 480 | 0.712488 (4) |
| SP 11–14      | 16.0 (3) 230    | −3.4 (1) 340  | 0.715212 (5) |
| SP 11–15      | 15.4 (2) 19     | −2.2 (1) 15   | 0.711478 (4) |
| SP 11–16      | 16.7 (1) 620    | −8.2 (1) 420  | 0.712171 (2) |
| SP 11–18      | 15.9 (1) 180    | −1.8 (2) 120  | 0.711207 (4) |
| SP 11–21      | 16.0 (2) 33     | −11.3 (1) 260 | 0.714932 (4) |
| SP 11–23      | 11.4 (1) 9      | −9.1 (1) 28   | 0.719703 (5) |
| SP 11–23f     | 11.3 (2)        |              |          |
| SP 11–27      | 16.6 (3) 140    | −6.4 (1) 120  | 0.711359 (3) |
| SP 11–30      | 16.9 (2) 360    | −11.8 (1) 340 | 0.712932 (3) |
| SP 11–02      | 17.0 (2) 55     | −19.9 (1) 32  | 0.712451 (5) |
| SP 11–03      | 16.5 (2) 340    | −17.3 (1) 170 | 0.712545 (5) |
| SP 11–03f     | 16.3 (3)        |              |          |
| SP 11–04      | 16.9 (4) 1040   | −16.9(1) 540  | 0.712535(2) |
| SP 11–04f     | 17.0 (2)        |              |          |
| SP 11–06      | 17.3 (2) 74     | −16.5 (1) 31  | 0.711918 (5) |
| SP 11–17      | 16.6 (2) 140    | −8.5 (1) 110  | 0.7121765 (5) |

Uncertainties are valid for the last digits
### Table 4  Isotope and concentration data for the Diamond Core Drill Holes of Salar de Pozuelos

| Hole coordinates | Sample number | Depth mbs | Material     | δ²⁷Li (‰) | Li (µg/g) | δ¹¹B (‰) | B (µg/g) | ⁸⁷Sr/⁸⁶Sr |
|------------------|---------------|-----------|--------------|-----------|-----------|-----------|-----------|------------|
|                  |               |           |              | Salt      | Clastic   |            |           |            |
| L01 (SPZ RC001)  | PZ1-0–1       | −0.5      | Salt/Halite  | 16.9 (0)  | 63        | −16.6 (1) | 237       | 0.714556 (6) |
|                  | PZ1-5–6       | −5.5      | Salt/Halite  | 16.5 (2)  | 26        | −16.4 (1) | 111       | 0.714116 (8) |
| 24°43′00.5"     | PZ1-10–11     | −10.5     | Salt/Halite  | 16.9 (1)  | 36        | −16.3 (1) | 38        | –          |
| 66°49′49.5"     | PZ1-20–21     | −25.5     | Salt/Halite  | 17.0 (2)  | 33        | −18.1 (1) | 11        | 0.713333 (9) |
|                  | PZ1-35–36     | −35.5     | Salt/Halite  | 17.0 (1)  | 14        | −17.3 (1) | 95        | 0.71508 (2)  |
|                  | PZ1-45–46     | −45.5     | Salt/Halite  | 16.0 (3)  | 15        | −14.9 (1) | 423       | 0.715006 (6) |
|                  | PZ1-50–51     | −50.5     | Salt/Halite  | 16.3 (2)  | 49        | −15.4 (1) | 375       | 0.715008 (5) |
|                  | PZ1-65–66     | −65.5     | Salt/Halite  | 16.6 (2)  | 28        | −14.1 (1) | 154       | 0.714439 (7) |
|                  | PZ1-75–76     | −75.5     | Salt/Halite  | 16.8 (3)  | 27        | −14.7 (1) | 146       | 0.714256 (6) |
|                  | PZ1-85–86     | −85.5     | Salt/Halite  | 16.5 (3)  | 28        | −15.0 (1) | 49        | 0.713931 (6) |
|                  | PZ1-95–96     | −95.5     | Salt/Halite  | 17.0 (5)  | 6         | −15.1 (1) | 181       | 0.714839 (8) |
| DDH1 (SPZ DDH001)| Po1-C35       | −96.5     | Sand         | −1.1 (1)  | 125       | –          |           | 0.715725 (6) |
|                  | Po1-C35       | −96.5     | Salt*        | 16.0 (2)  | 935       | –          |           | 0.711918 (9) |
| 24°43′00.5"     | Po1-C43       | −118.5    | Sand         | −1.5 (0)  | 130       | –          |           | 0.715017 (6) |
| 66°49′49.2"     | Po1-C43       | −118.5    | Clay         | −1.1 (0)  | 193       | –          |           | 0.715465 (9) |
|                  | Po1-C43       | −118.5    | Salt*        | 16.9 (3)  | 329       | –          |           | 0.712514 (7) |
|                  | Po1-C47       | −130.5    | Sand         | −2.2 (1)  | 322       | –          |           | 0.711923 (6) |
|                  | Po1-C47       | −130.5    | Clay         | −3.7 (2)  | 370       | –          |           | 0.712729 (6) |
|                  | Po1-C47       | −130.5    | Salt*        | 16.6 (4)  | 413       | –          |           | 0.711916 (6) |
|                  | Po1-C52       | −145.5    | Sand         | −1.9 (2)  | 144       | –          |           | 0.712293 (6) |
|                  | Po1-C52       | −145.5    | Clay         | −3.5 (2)  | 166       | –          |           | 0.718049 (6) |
|                  | Po1-C52       | −145.5    | Salt*        | 17.4 (1)  | 405       | –          |           | 0.711916 (4) |
|                  | Po1-C58       | −165      | Sand         | −2.3 (1)  | 148       | –          |           | 0.711349 (6) |
|                  | Po1-C58       | −165      | Salt*        | 17.4 (2)  | 325       | –          |           | 0.712417 (7) |
|                  | Po1-C62       | −183      | Silt         | −1.1 (1)  | 89        | –          |           | 0.715747 (5) |
|                  | Po1-C62       | −183      | Clay         | −1.8 (0)  | 288       | –          |           | 0.72375 (2)  |
|                  | Po1-C62       | −183      | Salt*        | 17.6 (1)  | 248       | –          |           | 0.712205 (5) |
| Surface sp 3     | SP11-07       | 0         | Sand         | −0.1 (1)  | 63        | –          |           | 0.711379 (4) |
| 24°43′00.8"     | SP11-07       | 0         | Clay         | −0.6 (0)  | 311       | –          |           | 0.715641 (4) |
| 66°50′25.0"     | SP11-07       | 0         | Salt*        | 17.3 (2)  | 74        | –          |           | 0.711917 (4) |

*Water-soluble fraction

Uncertainties are given as 2σd (B, Li) and 2σd mean (Sr) and valid for the last digits
of the brines are related to evaporation and increases with
the total cation load. Saturation appears to be reached at
loads around 5 mol/L, but Li (and also K) further enriches
in the residual brine, because it prefers the brine over the
salt (ESM1 Fig. A4). The Li content in the salts is variable
and distribution coefficients for Li between brine-salt pairs

### Table 5 Isotope and concentration data for samples from the Lithium Triangle

| Sample point | Locality | Coordinates              | Brine/water | Salt δ^7Li (%) | Li (mg/L) | δ^11B (%) | B (mg/L) | ^87Sr/^86Sr | 2sdmean |
|--------------|----------|--------------------------|-------------|----------------|-----------|-----------|-----------|-------------|---------|
| CA1          | Salar Grande | 25°56′45.6″S 68°42′24.3″W | W1          | S1             | 6.8 (1)   | 600       | 16.2 (1)  | 1390        | 0.708649 (6) |
|              |           |                          | W2          | S2             | 6.3 (3)   | 21        | 15.6 (1)  | 50          | 0.708502 (5) |
| CA2          | Salar de las Parinas | 25°48′42.0″S 68°28′39.0″W | W4          | S4             | 6.2 (2)   | 453       | 0.0 (1)   | 505         | 0.707999 (6) |
| CA3          | Salar de la Isla (1) | 25°40′15.0″S 68°34′30.0″W | W6          | S6             | 10.9 (1)  | 70        | −2.7 (1)  | 25.4        | 0.709181 (6) |
|              |           |                          | W8          | S8             | 11.2 (2)  | 66        | −6.5 (1)  | 17.3        | 0.707435 (7) |
| CA4          | Salar de la Isla (2) Volcan ‘Dos Crateros’ | 25°39′10.3″S 68°38′08.4″W | W9          |                | 11.0 (2)  | 37        | −3.7 (1)  | 7.8         | 0.707368 (9) |
| CA5          | Quebrada Chaco del Norte | 25°25′59.9″S 69°16′00.1″W | W11         |                | 8.4 (3)   | 1.5       | 7.8 (1)   | 0.49        | 0.707936 (6) |
| CA6          | Salar de Penedales (1) | 26°21′44.8″S 69°11′22.6″W | W13         |                | 10.7 (1)  | 50        | 16.7 (1)  | 22.6        | 0.708715 (6) |
| CA7          | Quebrada del Rio Negro Banos Rio Negro | 26°30′41.0″S 68°49′10.9″W | W14         |                | 7.3 (2)   | 34        | 3.6 (1)   | 24.4        | 0.707419 (6) |
|              |           |                          | W16         |                | 7.3 (0)   | 28        | 4.6 (1)   | 22.2        | 0.707354 (6) |
| CA8          | Laguna de Huasco | 20°16′38.4″S 68°53′00.0″W | W18         | S18            | 7.0 (2)   | 26        | 6.8 (1)   | 2.1         | 0.706435 (5) |
| CA9          | Terma Chiriguaya | 18°20′27.6″S 69°10′28.8″W | W19         |                | 6.0 (2)   | 1.2       | 5.2 (1)   | 34.2        | 0.707445 (6) |
| CA10         | Salar de Atacama (1) Laguna Chaxa | 23°19′48.9″S 68°10′02.4″W | W20         | S20            | 9.7 (2)   | 279       | −8.3 (1)  | 221         | 0.709773 (6) |
|              |           |                          | W21         | S21            | 9.7 (2)   | 371       | −8.0 (1)  | 290         | 0.709782 (7) |
| CA11         | Salar de Atacama (2) | 23°16′18.8″S 68°10′37.0″W |              | S22            |            |           |           | 0.710434 (6) |
| CA12         | Salar de Atacama (3) | 23°33′58.7″S 68°07′16.8″W |              | S23            |            |           |           | 0.708267 (6) |
| CA13         | Salar de Piedra Parada | 26°22′24.6″S 68°43′48.0″W |              | S24            |            |           |           | 0.707082 (5) |
| CA16         | Salar de Antofalla | 26°20′15.6″S 68°00′55.0″W |              | S25            |            |           |           | 0.707125 (6) |
| CA17         | Salt pond ignimbrite field | 25°47′42.2″S 68°22′40.2″W |              | S30            |            |           |           | 0.708851 (5) |
| CA18         | Salar de Uyuni (1) Ojos del Salar | 20°18′43.0″S 66°59′50.0″W | BU2          |                | 13.3 (1)  | 439       | 1.0 (0)   | 517         | 0.710604 (8) |
| CA19         | Salar de Uyuni (2) Cerro Tunupa | 19°54′26.4″S 67°41′03.3″W | BU4          |                | 13.9 (1)  | 446       | −2.0 (1)  | 323         | 0.708116 (7) |

Uncertainties are valid for the last digits
Locations for Google Earth in ESM2
are 17.5 and 16.2 at locations (sp) 1 and 3 in the centre and 4.4 at location 10 near the rim of the salar. The precipitates (solid salt) in the brine ponds and pools, are mainly halite (four of five samples have NaCl contents of 98 to 99%, one sample 96%). Li contents are variable, ranging between 57 and 140 µg/g in the salt from natural brine ponds, and reaching up to 1040 µg/g in a dendritic salt sample from an artificial brine pool. A salt sample from a completely evaporated artificial pool shows a Li concentration of 340 µg/g. The salt of core L01 contains 99 to 100% NaCl, i.e. is supposed to be nearly pure halite. Li concentrations are generally lower than in the surface salts, ranging between 6 and 63 µg/g and clustering between 30 and 50 µg/g.

Oxygen and hydrogen isotope composition of the surface brines

Oxygen and hydrogen compositions of the brines vary in δ¹⁸O values between −6 and +11‰ and in δ²H values between −68 and −17‰; the spring fluid is somewhat different with δ¹⁸O values of −10‰ and δ²H of −95‰ (Fig. 2). The values of the brines deviate in a linear array from the Global Meteoric Water Line (GMWL; δ²H = 8*δ¹⁸O + 10; Craig 1961) and the Local Meteoric Water Line (LMWL; δ²H = 7.8*δ¹⁸O + 9.7; Aravena et al. 1999) for Northern Chile (Fig. 2). Such arrays of data with slopes that are less steep than the meteoric water line are typical of evaporation (e.g. Fiorella et al. 2015 for the Central Andes) and follows the trend observed for other high-altitude basins in the Central Andes (e.g. Godfrey et al. 2013; Orberger et al. 2015). There is no simple covariation with the progress of evaporation seen in the oxygen isotopes and Li contents (ESM1 Fig. A4).

Lithium and Sr isotope compositions

Salar deposits: surface brines and evaporites

Brines collected at the surface show ranges of δ⁷Li values between +15.4 and +19.3‰ and ⁸⁷Sr/⁸⁶Sr between 0.7108 and 0.7149; salts show ranges of +16.5 to +17.3‰ and 0.7119 to 0.7125. Brine-salt pairs from the same ponds show no differences between their δ⁷Li values within the overlapping uncertainties, but the average δ⁷Li of the brines are slightly higher by 0.1 to 0.2‰ than the average of the coexisting salts. The ⁸⁷Sr/⁸⁶Sr of the brine-salt pairs is indistinguishable within the external reproducibility of the NIST 987 reference material of 14 ppm (2sd_mean). In the hydrothermal spring location, the spring fluid shows a δ⁷Li value of +11.4‰, with a Li concentration of 9 µg/g and Sr isotope signature of 0.7197. The travertine ranges from −0.07 to +1.3‰ (average 0.7 ± 1 (2sd), n = 8) in δ⁷Li, with Li concentrations between 3 and 17 µg/g, and ⁸⁷Sr/⁸⁶Sr between 0.7168 and 0.7247. The δ⁷Li and ⁸⁷Sr/⁸⁶Sr of the travertine are positively correlated (r² 0.05), whereas covariation of Sr and Li isotope composition is absent in all other sample groups. The offset in the δ⁷Li value between the travertine and the spring fluid is −10.7‰. Salts of the upper 95 m of the core L01 show small ranges in δ⁷Li values between +16.0 and +17.0‰ and in ⁸⁷Sr/⁸⁶Sr of 0.7133 and 0.7151 (Fig. 3). Two borate (ulexite) samples have δ⁷Li values of +8.2 and +12.4‰ and ⁸⁷Sr/⁸⁶Sr of 0.7152 and 0.7154, respectively. There is no covariation between Li contents and δ⁷Li values in any of the sample groups (ESM1 Fig. A5).

Salar deposits: siliciclastic sediments and their water-soluble fractions

The δ⁷Li values, Li contents and ⁸⁷Sr/⁸⁶Sr of siliciclastic sediments from the border region of the salar (SP-11–18, 28, 31A and B) vary between −3.6 and +0.8‰, 47 and 590 µg/g and 0.7110 and 0.7151. Compositions of core DDH1 and SP-11–7 range in the sand-rich fraction from −2.3 to −0.1‰, 63 to 322 µg/g, 0.7113 to 0.7157 and in the clay-rich fraction from −3.7 to −0.6‰, 193 to 370 µg/g, 0.7127 to 0.7237. The water-soluble fractions of the core DDH1 show δ⁷Li values between +16.0 and +17.5‰, Li concentrations of 248 to 935 µg/g, and ⁸⁷Sr/⁸⁶Sr of 0.7119 to 0.7125. The water-soluble fraction of surface sediment SP-11–7 from the centre of the salar shows values of +17.3‰, 74 µg/g and 0.7119, and the isotopic composition is indistinguishable from DDH1 (Fig. 3).
Salar catchment area: siliciclastic sediments and igneous rocks and their water-soluble fractions

The Li and Sr isotope composition of the sedimentary rocks (washed) is quite variable, including three Oligocene (δ7Li = +4.3, +4.9, +9.3‰; Li = 29, 34, 26 µg/g; 87Sr/86Sr = 0.7149, 0.7180, 0.7139) and one Eocene (+12.8‰; 25 µg/g; 0.7175) sample, and a recent sediment from a dry drainage channel SP-11–20 (−0.3‰; 25 µg/g; 0.7114). The water-soluble fractions show also considerable variations (δ7Li = +14.7, +10.0, +22.9‰; Li = 394, 1271, 39 µg/g; 87Sr/86Sr = 0.7106, 0.7105, 0.7113) in the Oligocene samples, (+12.2‰; 4.3 µg/g; 0.7156) in the Eocene and in the unconsolidated sediment (−1.9‰; 1.1 µg/g; 0.7109). The fresh and altered ignimbrite samples show δ7Li values of −4.7 and −25.0‰, Li concentrations of 24 and 28 µg/g; and 87Sr/86Sr of 0.7111, respectively. The associated water-soluble fraction has δ7Li values of +9.8 and +12.8‰, Li concentrations of 4.8 and 3.1 µg/g, and 87Sr/86Sr of 0.7103 and 0.7103. The local sample of the Ordovician sedimentary rock in the group of ‘sedimentary rocks’ in the catchment area. The Li isotope composition is uniform with depth and the main compositional difference occurs between the group of siliciclastic fractions and the groups of brine, water-soluble fraction, and salt in the Salar. 87Sr/86Sr of salts from L01 are slightly higher than in the water-soluble fraction DDH1. The variability of 87Sr/86Sr is generally higher in the clay- and sand-rich fraction and the clay-rich fraction scatters towards higher values. Influence of radiogenic Sr isotope compositions of the Palaeozoic basement (local ‘Ord’ Ordovician sedimentary rock in the catchment area) on the young salar deposits is generally small and highest in the travertine. 87Sr/86Sr of the survey samples (Li-Triangle) reflects their position within the active magmatic arc of the CVZ.

Salar catchment area: siliciclastic sediments and igneous rocks and their water-soluble fractions

The Li and Sr isotope composition of the sedimentary rocks (washed) is quite variable, including three Oligocene (δ7Li = +4.3, +4.9, +9.3‰; Li = 29, 34, 26 µg/g; 87Sr/86Sr = 0.7149, 0.7180, 0.7139) and one Eocene (+12.8‰; 25 µg/g; 0.7175) sample, and a recent sediment from a dry drainage channel SP-11–20 (−0.3‰; 25 µg/g; 0.7114). The water-soluble fractions show also considerable variations (δ7Li = +14.7, +10.0, +22.9‰; Li = 394, 1271, 39 µg/g; 87Sr/86Sr = 0.7106, 0.7105, 0.7113) in the Oligocene samples, (+12.2‰; 4.3 µg/g; 0.7156) in the Eocene and in the unconsolidated sediment (−1.9‰; 1.1 µg/g; 0.7109). The fresh and altered ignimbrite samples show δ7Li values of −4.7 and −25.0‰, Li concentrations of 24 and 28 µg/g; and 87Sr/86Sr of 0.7111, respectively. The associated water-soluble fraction has δ7Li values of +9.8 and +12.8‰, Li concentrations of 4.8 and 3.1 µg/g, and 87Sr/86Sr of 0.7103 and 0.7103. The local sample of the Ordovician sedimentary rock in the group of ‘sedimentary rocks’ in the catchment area. The Li isotope composition is uniform with depth and the main compositional difference occurs between the group of siliciclastic fractions and the groups of brine, water-soluble fraction, and salt in the Salar. 87Sr/86Sr of salts from L01 are slightly higher than in the water-soluble fraction DDH1. The variability of 87Sr/86Sr is generally higher in the clay- and sand-rich fraction and the clay-rich fraction scatters towards higher values. Influence of radiogenic Sr isotope compositions of the Palaeozoic basement (local ‘Ord’ Ordovician sedimentary rock in the catchment area) on the young salar deposits is generally small and highest in the travertine. 87Sr/86Sr of the survey samples (Li-Triangle) reflects their position within the active magmatic arc of the CVZ.

Radiogenic (Nd, Pb, Sr) isotope composition of siliciclastic sediments of the salar basin and catchment area

Strontium, neodymium and lead isotope compositions of siliciclastic sediments of the salar basin and sedimentary rocks of the catchment area are within the range of typical large-volume ignimbrite of the Central Andes and show little influence in their Sr isotope composition by signatures of the local (salar catchment area) and regional (Altiplano-Puna Plateau) Palaeozoic basement (ESM1 Fig. A6).
Elemental content and Li and Sr isotopic composition of samples from the Li-Triangle

Sodium in the brine samples comprises 66 to 98% of the analysed (Na, K, Li, B, Ca, Mg, Sr) dissolved cations in the brines with only one sample being as low as 28% (spring water from Quebrada Chaco del Norte in Chile). The Li contents of the fluids ranges from 1.2 to 600 mg/L, showing a positive correlation ($p = 0.05$) between Li and Na, i.e. an evaporation trend. The $\delta_{7}^{11}$B values are between +6.0 and +13.9‰ and are hence, significantly lower than the Pozuelos brines, but in the range of fluids observed from other salars in the Central Andes, such as Hombre Muerto, Ratones and Centenario, and Olaroz (Godfrey et al. 2013; Orberger et al. 2015; Garcia et al. 2020). The $^{87}$Sr/$^{86}$Sr of the fluid samples vary between 0.7064 and 0.7098, nearly perfectly correctly ($p = 0.05$ with the values (0.7062 to 0.7097) from the associated salt samples. Since brine-salt pairs show no differences between their $\delta_{7}^{11}$Li values within uncertainties, we only analysed the salt’s Sr isotope composition. Salt samples without paired fluids show a similar range in $^{87}$Sr/$^{86}$Sr from 0.7071 to 0.7104.

Boron concentration and isotope composition of the surface brines and salts from the surface, core L01 and samples from the Li-Triangle

The boron concentrations of the surface brines range from 15 to 490 µg/g with a total range in the B isotopic compositions from −19.3 to −1.8‰ (Fig. 4). The boron concentration for the salt (31 to 540 µg/g) is similar to the surface brines, but the range in $\delta^{11}$B is more restricted to values between −19.9 and −8.5‰, (average −15.8 ± 8.6, 2sd). The salt from L01 show the most homogeneous $\delta^{11}$B values with an average similar to the surface salts of −15.8 ± 2.4 (2sd, $n = 11$). The B concentration ranges from 11 to 423 µg/g. The travertine shows the most negative $\delta^{11}$B value of about −23.5‰ and two borate minerals show $\delta^{11}$B values of −17.5‰ and −6.7‰. The B concentration for the fluid samples of the survey samples from the Li-Triangle ranges from 0.5 to 1390 mg/L, with the brine samples showing the highest values and spring samples reaching maximum values of 50 mg/L. The total range in the B isotopic compositions is again quite substantial (−6.5 to +16.7‰) but more positive compared to the Salar de Pozuelos samples.

Discussion

Sources of the Li in the Salar de Pozuelos–Sr isotope composition

Aqueous solutions entering the salar reflect the strontium isotope signatures of their sources and the isotopic signature of brine and salt deposits is governed by mixing of different $^{87}$Sr/$^{86}$Sr source-compositions during transport and post depositional processes in the salar. Incongruent, mineral-specific weathering of low and high Rb/Sr minerals from young, Andean magmatic rocks has minor influence on the Sr isotope signature of aqueous solutions, but the influence may be large from selective weathering of such minerals from Ordovician basement rocks. The latter have a given lower limit of Sr isotope signatures for incongruent weathering of a hypothetical very low Rb mineral by their average initial $^{87}$Sr/$^{86}$Sr of 0.715; the average modern $^{87}$Sr/$^{86}$Sr for the case of congruent weathering is 0.759 (Egenhoff and Lucassen 2003). However, the $^{87}$Sr/$^{86}$Sr for our brine and salt samples are not higher than 0.715 but generally lower in most samples, the exclusive origin of Sr from a very low Rb mineral of the Ordovician basement rocks is highly unlikely, and the agreement with the isotopic values of the young siliciclastic salar sediments and the outcropping Andean rock associations present in the catchment area and in the Altiplano-Puna Plateau is very good (Fig. 3; ESM1 Fig. A6, ESM4). Hence, we preclude the Ordovician as the major source of Sr in the Salar.

The Sr isotope signatures of the rock inventory in the Central Andes are well known and distinguish sharply between the major rock sources. The combined Sr and Nd isotopic signatures of the rocks provide an even better tool to identify the sources (ESM1 Fig. A6). The surface geology in the catchment area of Pozuelos is dominated by Ordovician sedimentary rocks, partly covered by much younger Eocene to Miocene sedimentary rocks, partly covered by much younger and Quaternary sedimentary rocks (Fig. 1). Apart from documented tuff layers in these young rocks (Blasco et al. 1996), we assume a frequent widespread supply of volcanic materials from air-falls during eruptions of andesitic stratovolcanoes and ignimbrite flare-ups or windblown volcanic detritus from the still active magmatic arc that is stored and hidden in the young colluvial/alluvial deposits. Besides isolated intrusions of Miocene igneous...
rocks in the catchment area of Pozuelos (Blasco et al. 1996),
relicts of massive ignimbrite exists in the vicinity around
the Salar according to mapping of the surface distribution of
such rocks in the Puna Plateau (Brandmeier and Wörner
2016) and own observations. The surface geology is under-
lain by the Palaeozoic basement.

We use the $^{87}\text{Sr}/^{86}\text{Sr}$ compositions of the local (Salar de
Pozuelos catchment area) and regional (Altiplano-Puna Pla-
teau) rock inventory to characterise the siliciclastic source
rocks of Li deposited in evaporites of the Pozuelos. The
regional averages refer to Palaeozoic basement rocks and
Cenozoic volcanic rocks from the Central Andean arc with
already analysed Li isotope compositions (Fig. 3a; Meixner
et al. 2020). In the catchment area of Pozuelos, the $^{87}\text{Sr}/^{86}\text{Sr}$
of the local Ordovician sedimentary rock (0.731) and the
ignimbrite (0.711) falls within the regional average of the
Palaeozoic sedimentary-metamorphic-magmatic basement
(0.754 ± 0.04 (2sd), n = 27) and the regional averages of
Cenozoic andesite (0.707 ± 0.001 (2sd), n = 21) and ign-
imbrite (0.710 ± 0.001 (2sd), n = 11), respectively. The
$^{87}\text{Sr}/^{86}\text{Sr}$ composition of the local Cenozoic sediments of the
catchment area is 0.711 to 0.718 (0.715 ± 0.002 (2sd), n = 5)
and not dominated by the abundant Ordovician sedimentary
rock, but similar to the composition of Cenozoic ignimbrite.

The siliciclastic sediments from the salar deposits,
including clay- and sand-rich fractions of DDH1 (average
$^{87}\text{Sr}/^{86}\text{Sr}$ 0.715, n = 13), resemble the composition of Ceno-
zoic sedimentary rocks from the catchment area (Fig. 3;
ESM1 Fig. A6). The small range of Nd isotope signatures
of catchment area (0.5122 ± 0.0001 2sd, n = 7) and salar
siliciclastic rocks (0.5122 ± 0.0001 2sd, n = 15) is not
correlated with the more variable $^{87}\text{Sr}/^{86}\text{Sr}$ and both resemble
the variation in regional large volume ignimbrite (ESM1
Fig. A6). The sand-rich fraction of DHH1 shows a small
range of $^{87}\text{Sr}/^{86}\text{Sr}$ (0.711 to 0.716; average 0.713, n = 7) and
the clay-rich fraction is slightly more radiogenic (0.713 to
0.724; average 0.717, n = 5). We speculate that detrital micas
of the Ordovician sedimentary rocks with high $^{87}\text{Sr}/^{86}\text{Sr}$ are
variable in occurrence, but in small amounts enriched in the
clay-rich fraction. This could also explain the uniform Nd
and variable Sr isotope composition of DHH1, due to the
generally low rare earth element contents of micas.

Quantification of distinct components remains ambiguous
considering the potential addition of high $^{87}\text{Sr}/^{86}\text{Sr}$ micas
from the Palaeozoic basement, e.g. the Ordovician sediment-
ary rocks. Mixing models have to consider the uniformity
of Nd isotope signatures in the siliciclastic fractions of DHH1,
other salar sediments, and in the Cenozoic cover (ESM1
Fig. A6). Binary mixing between an ignimbrite endmember
($^{87}\text{Sr}/^{86}\text{Sr}$ of 0.710, Sr of 250 µg/g; ignimbrite database from
Brandmeier and Wörner 2016) and Ordovician sedimentary
rocks ($^{87}\text{Sr}/^{86}\text{Sr}$ of 0.75, Sr of 70 µg/g; Egenhoff and Lucas-
sen 2003) requires 0 to 20% Sr of bulk Ordovician (one
clay-rich sample of DDH1 requires about 30% bulk Ordovi-
cian) to explain the observed $^{87}\text{Sr}/^{86}\text{Sr}$ compositional range
of the Cenozoic siliciclastic sediments.

The Sr isotope compositions of salts and brines from the
Salar de Pozuelos reflect the signatures of the hydrous influx
into the salar and potential processes of homogenisation of
the solutions. All brines and salts from the surface, core L01,
and water-soluble fraction of DHH1 (0.713 ± 0.003 (2sd),
n = 34) plot in general on the local and regional ‘ignimbrite
averages’ of Sr isotope composition and within the composi-
tional range of the Cenozoic sedimentary rocks and their
water-soluble fractions (Fig. 3; ESM1 Fig. A6). Signatures
of the water-soluble fractions of DHH1 (i.e. salt, indicated
as salt in the table) are more homogeneous, generally less
radiogenic (average of 0.7121 ± 0.0005, 2sd) and hence, not
related to the more radiogenic signatures (0.715 ± 0.006, 2sd)
of their silicate hosts. The isotopic signature of the surface
salts (0.7123 ± 0.0005, 2sd) is similar to the water-sol-
uble fraction. The small variability of actual surface brines
from the salar’s border (0.713 ± 0.005, 2sd) is likely caused
by variations in their source contributions. In contrast, the
smaller compositional spread in the salt and water-soluble
fractions of the cores points to a time-integrated homog-
enisation of the run-off from different silicate sources dur-
ing leaching, transport and precipitation of the evaporite
deposit. The $^{87}\text{Sr}/^{86}\text{Sr}$ in surface brines (except SP-11–14
and SP-11–21), the water-soluble fractions of core DDH1,
and surface sediment (SP-11–7) are slightly less radiogenic
(lower) than in the salts of the core L01 (Fig. 3b). Sample
sites (sp 8, 13) of brine SP-11–14 and SP-11–21 border to
an extended outcrop of Ordovician rocks and seem to reflect
the radiogenic $^{87}\text{Sr}/^{86}\text{Sr}$ of this proximal source (Fig. 1). The
$^{87}\text{Sr}/^{86}\text{Sr}$ of SP-11–14 is lower than of SP-11–21, which
locates closer to the spring and travertine deposit. The spring
and the related travertine is hosted in the faulted contact
between the Ordovician and Eocene rocks and samples
slightly variable amounts of radiogenic Sr over time as seen
in the travertine (Fig. 3b; ESM1 Fig. A6). The slight but
systematic offset to more radiogenic $^{87}\text{Sr}/^{86}\text{Sr}$ in the halite
from core L01 could be caused by a higher contribution of
runoff from the Ordovician potentially including enhanced
hydrothermal activity in the past with radiogenic $^{87}\text{Sr}/^{86}\text{Sr}$
as seen in the travertine and related spring.

On the base of the similar Sr isotope signatures in
evaporite and Cenozoic siliciclastic sediments within the
salar and in the catchment area, we favour a dominantly
ignimbrite source rock (binary mixing of about 70 to
100% ignimbrite component and 0 to 30% Ordovician
sedimentary rocks, as discussed above), even if at pre-
sent no significant amount of ignimbrites are exposed in
the catchment area. However, the ignimbrite deposits of
the Central Andes are among the largest on Earth, with
volumes up to several thousands of cubic kilometres and
covering areas of over 10,000 km² (e.g., Brandmeier and Wörner 2016). In addition, aeolian erosion and transport of e.g. poorly indurated ignimbrite are well described (Mandt et al. 2008). Moreover, the Altiplano and the Puna are areas with important dust activity year round (Prospero et al. 2002), potentially since the Late Pliocene age (Goudie and Wells 1995). Thus, a possible scenario includes volcanic detritus deposition either from air-fall or windblown. The volcanic detritus at the surface is also easily transported and potentially accumulated in the young, poorly solidified sedimentary cover, and in the salar basin as evidenced by the composition of siliciclastic salar sediments (Fig. 3b; ESM1 Fig. A6). Such volcanic material is much more sensitive to chemical weathering, due to the large surface and high glass contents (Ruggieria et al. 2010), than the well-solidified massive Ordovician sedimentary rocks.

The Sr isotope composition of the basin sedimentary material and fluids could also be produced by mixing approximately 35 to 60% andesite (Meixner et al. 2020 with references herein) with 40 to 65% Pz-basement. We, however, consider this scenario as less likely. Andesite deposits are not known in the catchment area, and while aeolian transport and deposition of eroded material is possible, the andesites are mostly deposited as lavas produced from stratovolcanoes and monogenetic centres and less prone to erosion than the poorly indurated ignimbrite. In addition, a relative high input of Pz-basement material, the dominant rock lithology (Ordovician sedimentary rock) in the catchment area is necessary for this scenario not only for the addition of Sr, but also to deliver a considerable amount of Li. The host of Li in the Ordovician sedimentary rocks are most likely micas and K-feldspars with a radiogenic Sr isotope composition. An increased input of these minerals would drive the Sr isotope composition of the siliciclastic sediments in the basin to more radiogenic values, a situation that is not observed.

Sources of the Li in the Salar de Pozuelos—Li isotope composition

Silicate minerals are the principal primary host of Li (Teng et al. 2004) and weathering or hydrothermal leaching are the principal processes of mobilisation of Li from the silicate. The extent of fractionation of the Li isotope composition depends on the weathering process. Chemical weathering with bulk solution transfers the δ7Li signature of host minerals to the solute. In contrast, higher δ7Li in solutes depends on the extent of secondary mineral formation in the source rock, during transport, as well as in the deposit area by sequestration of 6Li in secondary clay minerals (e.g. Pistiner and Henderson 2003; Kisakurek et al. 2004; Vigier et al. 2008; Wimpenny et al. 2010; Hindshaw et al. 2019; recent summaries provided by Tomascak et al. 2016; Pogge von Strandmann et al. 2016; Weynell et al. 2017). The fractionation of Li isotopes in hydrothermal systems depends, e.g. on the temperature of the fluid/rock reaction and related formation and stability of secondary minerals. The fractionation, i.e. δ7Li in the fluid from natural and experimental hydrothermal systems, is generally inversely correlated with T and approaches the composition of the host rocks at high T (e.g. Millot and Négre 2007; Millot et al. 2010; Henchiri et al. 2014; Pogge von Strandmann et al. 2016; Cullen et al. 2019: recent summary Tomascak et al. 2016). A hydrothermal spring compositional field was denoted with δ7Li > 0 and < 11‰ (Pogge von Strandmann et al. 2016 and references therein), a range which encompasses many of the δ7Li data in hydrothermal fluids and salar deposits in the Central Andes (see below).

The siliciclastic Ordovician sedimentary rock sample from the Salar de Pozuelos catchment shows a δ7Li value of −1.3‰, which is within the regional (Altiplano-Puna Plateau) average of −1.1 ± 2‰ (2sd, n = 15; Meixner et al. 2020) for Ordovician sedimentary rocks and similar to the regional average of +0.9 ± 3‰ (2sd, n = 15; Meixner et al. 2020) for the Palaeozoic magmatic-metamorphic basement. The Ordovician sedimentary rocks are the dominant surface rocks in the Salar de Pozuelos catchment area and partly covered by Cenozoic sediments with a range in δ7Li value from −0.3 to + 12.8‰ (δ7Li average of +6.2 ± 10‰ (2sd), n = 5). Outcrops of volcanic rocks in the catchment area (local) are restricted to small occurrences of ignimbrite, but influx of detritus from the active arc as ash-fall or windblown material is potentially important. The Li isotope composition of two local ignimbrite samples are −2.5 and −4.7‰; the regional average of silicic (large-volume) ignimbrites is −1.4 ± 7‰ (2sd, n = 10) and +2.3 ± 4‰ (2sd, n = 23) for Central Volcanic Zone (CVZ) andesites (Meixner et al. 2020).

The δ7Li values of brines and salts are in general uniform, pointing to negligible isotope fractionation during precipitation of salt from brine (Godfrey et al. 2013; this work), and higher compared to the values of potential source rocks, i.e. rocks from the catchment area and regional volcanic- and Pz-basement rocks (Fig. 3a;
The δ⁷Li values in surface brines and salts (average +17.5 ± 2‰) are similar to those of the salt and water-soluble fractions in the drill holes (average +16.8 ± 0.9‰) but slightly more variable, which points to homogenisation of the more heterogeneous compositions from proximal sources within the salar (Fig. 3a). Thus, salts and brines of the salar deposits, including the water-soluble fraction of DHH1, show a uniform δ⁷Li composition in three dimensions. There is no covariation, i.e. a mixing relation between δ⁷Li and strongly variable Li concentrations (ESM1 Fig. A5). The concentrations are dominated by evaporation as corroborated by the oxygen isotope composition of the surface brines.

Fig. 5). The δ⁷Li values in surface brines and salts (average +17 ± 2‰) are similar to those of the salt and water-soluble fractions in the drill holes (average +16.8 ± 0.9‰) but slightly more variable, which points to homogenisation of the more heterogeneous compositions from proximal sources within the salar (Fig. 3a). Thus, salts and brines of the salar deposits, including the water-soluble fraction of DHH1, show a uniform δ⁷Li composition in three dimensions. There is no covariation, i.e. a mixing relation between δ⁷Li and strongly variable Li concentrations (ESM1 Fig. A5). The concentrations are dominated by evaporation as corroborated by the oxygen isotope composition of the surface brines.

The spring fluid has a lower δ⁷Li value (+11.4‰) than the brines and salts. The offset in the δ⁷Li value of −10.7‰ between the travertine and the spring fluid is similar to what is observed for the isotopic fractionation between inorganic aragonite and high-salinity fluids (Δδ⁷Li aragonite-artificial seawater of −11‰; Marriott et al. 2004), pointing to original aragonite precipitation. To note is the low Li concentration of approximately 9 μg/g in both the fluid and the travertine. While the actual spring fluid might not represent the original concentration of the Li inflow via the former hot spring, the travertine documents a low Li concentration inflow in the basin and the necessary Li enrichment via evaporation.

A Pleistocene travertine structure from the Salar de Hombre Muerto contains about 42 μg/g of lithium (Godfrey et al. 2013). The siliciclastic sediments associated with the evaporite deposit at surface and in the depth of the Salar de Pozuelos show consistently low δ⁷Li values between +0.8 and −3.7‰ with an average of −1.5 ± 2.8‰ (2sd; n = 16). This average value is lower than the average of the continental crust with ~0‰ (Teng et al. 2004; Sauzéat et al. 2015) and regional andesite, but resembles closely the average of local and regional Ordovician sedimentary rocks and regional ignimbrite. The Li concentrations in the siliciclastic sediments are variable, but higher (>47 μg/g <590 μg/g; average 230 ± 165 μg/g, n = 16) than in average continental crust (30 μg/g; Sauzéat et al. 2015; 35 μg/g; Teng et al. 2004) or in the potential regional source rocks, i.e. andesite, ignimbrite and Ordovician sedimentary rocks (average: 21, 27, and 42 μg/g; Meixner et al. 2020). The comparable high Li concentrations in the salar sediments and lower δ⁷Li values in four out of five clay-rich fractions point to secondary (clay) minerals (ESM1 Fig. A7).

If we tentatively consider the difference between the average δ⁷Li value of +16.8‰ in the evaporite (i.e. the solution) and the salar siliciclastic sediments, assuming the −1.5‰ average of the salar sediments as ‘residue’, the Δδ⁷Li residue-solution is −18.3 ± 1.4‰. This value is close to the difference of −17.3‰ between solids and small river solute from a global compilation (Hindshaw et al. 2019 and references therein), the experimental Δδ⁷Li between clay minerals and solution of −16.6‰ (20 °C; Hindshaw et al. 2019), and extrapolations of experimental data to temperatures down to 5 °C (~19‰; Vigier et al. 2008). Such signatures were also observed in other catchment area—solution systems, e.g. with Δδ⁷Li of −17.8‰ in a similar weathering regime of the Himalayan (and elsewhere; Weynell et al. 2017 and references therein) and modelled (upper limit −16‰ from reactive transport modelling for
high-altitude catchment areas in the Himalayan, Bohlin and Bickle 2019). Thus, the observed δ7Li value of the evaporite deposits in Pozuelos indicates the dominance of near-surface chemical weathering with the formation of secondary clay minerals at low temperature. Bulk solution of Li-bearing silicate host does not fractionate Li and should reflect the source rock’s δ7Li.

In contrast, hydrothermal mobilisation of Li at some depth would cause variable but generally lower δ7Li values, due to the inhibited formation of secondary minerals at higher temperatures (range 1 to 11‰; Pogge von Strandmann et al. 2016 and references therein), than observed in the evaporites of Pozuelos (average of +16.8‰). The actual hydrothermal activity in the catchment area has ceased, and traces of former activity are restricted to one spring on the eastern side of the salar. If the Li isotope signatures of the spring fluid (+11.4‰) and the travertine (~0 to +1.3‰) precipitated from the well are typical for the local thermal waters, then they remained near constant over the evolution of the well, and hydrothermal influx likely played a minor role in the Li (and Sr) isotope budget of the evaporite in Pozuelos. Even if the range of δ7Li in travertine is small (−0.1 to +1.3‰), the values increase with variable radiogenic Sr from the Ordovician basement (Fig. 5), i.e. point to a weak influence of changing source contribution on the δ7Li values.

The most intriguing compositional feature of the Salar de Pozuelos basin is the small variation in 87Sr/86Sr signatures and the uniform high δ7Li values of salt precipitated from solutes in three dimensions of the deposit (Figs. 3 and 6). The age range represented by the drilled strata is unknown. Age models for cores in the Salar de Hombre Muerto (south of Pozuelos, catchment area ~4000 km², salar ~600 km²) indicate ages of approximately 20 ka at 8 m and up to 86 ka at 40 m depth (estimate from the diagrams in Godfrey et al. 2003). Sedimentation in the large Salar de Atacama (surface ~3000 km²; Alonso and Risacher 1996) is faster, with an estimated average rate of 0.9 m/kyr and an age of 106 ka at 100 m depth (Bobst et al. 2001) but represents a different depositional system compared with the conditions of the salars in the Puna Plateau. Even if we consider this fast deposition rate for the Salar de Pozuelos basin, the minimum age there at a depth of ca 100 m should be around 100 ka, and much older than 100 ka at 183 m, the lower end of the sampled core. The climate of the Puna Plateau and surroundings over the last 100,000 years included, compared with the present climate, various prolonged humid periods with glaciations around 13, 22 and 42 ka on the now largely deglaciated plateau (e.g. Flores et al. 1997; Bobst et al. 2001; Placzek et al. 2009; Schildgen et al. 2016; D’Arcy et al. 2019). The δ7Li values of Pozuelos drill cores do not reflect any potential effects of changes in the weathering regime over time (Fig. 3a).

While the isotope composition of salts in the centre of the salar is uniform, the brines and the potential runoff from the catchment area are more variable in composition. Because availability of pluvial water is seasonal and restricted, we were not able to sample the runoff from the catchment area. We focussed instead on the water-soluble fraction of silicate rocks of the catchment area that shows a considerable scatter in δ7Li from +10 to +23‰ with a range in isotope fractionation (Δ7Liwatersoluble-rock) from 0

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Fig. 6 Lithium and Sr isotope composition of source rocks, salar deposits, and (thermal) waters of the Central Andes. (a) δ7Li in the catchment area and salar deposits of Pozuelos, averages of potential regional source rocks of Li in the evaporite deposits (averages are taken from Meixner et al. 2020, and are given in the text and in Fig. 3), range of global hydrothermal waters (Pogge von Strandmann et al. 2016) and various regional evaporite deposits, potential feeders, thermal springs, and groundwater in the active arc of NE Chile around 22°S. (b) 87Sr/86Sr composition. Numbers indicate the respective data sources. Data sources: average host rocks Meixner et al. (2020); Salar de Hombre Muerto (1) Godfrey et al. (2013); Salar de Ratones and Centenario (2) Orberger et al. (2015); Salar Olaroz (3,4) Garcia et al. (2020) ((3) water surroundings of the salar; (4) brine); Salar de Atacama (5,6) Munk et al. (2018) ((5) water from feeders; (6) brine); (7,8) Godfrey and Álvarez-Amado (2020) ((7) groundwater from the surroundings of the salar; (8) brines from the adjacent border of the salar); ground water (9) Godfrey et al. (2019). For locations, see ESM2 Fig. A1
to + 17.5‰, assuming that the water-soluble Li is exclusively derived from the leached rock sample. The variability decreases in the surface brines (+15.4 to +19.3‰), salts (+16.3 to +17.3‰), and water-soluble fraction (+12.7 to +17.2‰) of the siliciclastic material from the salar. The increasing uniformity in the δ7Li values, from the potential catchment runoff to the border and to the centre of the salar, indicates an increasing mixing and homogenisation of the Li signatures, a process that is also observed for the Sr isotope composition (Figs. 3 and 6). Mixing to the uniform composition is likely bound to occasional pluvial events that result in flooding of the salar’s surface. Post depositional mixing of the entire volume of the salar seems unlikely because this would require dissolution of large volumes of halite, also in the siliciclastic sediments. Furthermore, slight systematic differences in the Sr isotope composition between the upper halite and salt (water-soluble fraction) from the lower siliciclastic section are preserved.

Other salar deposits and thermal waters of the Central Andes

Other salar deposits from the wider region, including our Li-Triangle data (mainly the salares of Atacama, Grande, Parina, Isla and Uyuni) show significantly lower δ7Li values than Pozuelos. The ranges of δ7Li values are large, but always higher than in the average regional source rocks, and most samples plot within the range of global hydrothermal waters (Fig. 6a). Only in a few samples, δ7Li values exceed +11‰ and reach up to +14‰.

In detail, most brines from Salar de Atacama show rather uniform δ7Li values from margin to core that range between +9.6 and +12.6‰ (Munk et al. 2018; this study) with generally lower values in feeders starting at +4.4‰. This dataset is resembled by the range of δ7Li values between +3.7 and +12.6‰ from feeder waters and marginal brines at the eastern edge of this salar (Godfrey and Alvarez-Amado 2020). Brines from Salar de Centenario, close to Pozuelos, show δ7Li values of +10.5 and +13.3‰ and from Salar Ratones between 8.9 and +9.8‰ (Orberger et al. 2015). Brines from Salar Olaroz show δ7Li values from +5.9 to +7.1‰ (surface) and +8.1 to +10.2‰ (depth of 150 to 450 m below surface from drill holes; Garcia et al. 2020). The study of the Salar de Hombre Muerto focussed on δ7Li values of rocks and water from feeders (range from 2.6 to +9.4‰) and assumes a δ7Li value of +8.3‰ for the brine in the centre (Godfrey et al. 2013). These salars have large volcanic centres and/or large volume ignimbrite (Salar Olaroz) in their catchment areas as observable in satellite views (ESM2 Fig. A1 Google Earth link) or formed, as the deposits of the Salar de Atacama, by massive inflow of (ground)water from such catchments (Corenthal et al. 2016). Thermal water and groundwater reported from the Argentine Puna (Godfrey et al. 2013), from the Western Cordillera, and adjacent basins in the west (north of Salar de Atacama; Godfrey et al. 2019) show variable δ7Li values between –1 and +13‰, with most samples plotting within the global hydrothermal field (Fig. 6a).

Our data labelled Li-Triangle, the Salar de Atacama, and ground- and thermal waters from the Western Cordillera show 87Sr/86Sr typical of regional andesite and comparable low δ7Li, whereas brines from Salar Olaroz and feeders of Salar de Hombre Muerto show—at similar δ7Li—a range of 87Sr/86Sr typical of large-volume ignimbrite and influence of radiogenic Pz-basement (Fig. 6b).

Active volcanism, arid climate, and internal draining since the Miocene are the background to weathering with generally restricted transport of solutes and solids in the Puna plateau (Alonso et al. 1991; reviews by Strecker et al. 2007; Vandervoort et al. 1995). Catchment areas of the endorheic basins can be highly varied with respect to the prevailing surface geology caused by the rough morphology of the Puna. However, weathering conditions at the surface, humidity and temperature, should be similar to all salar basins (precipitation rates ranging from 25 mm/year for the Salar de Atacama to 150 mm/year for Salar de Uyuni, Houston et al. 2011 with no apparent correlation between annual precipitation rate and the Li isotope composition or assumed source rocks for lithium. The reason for the—compared with Pozuelos—low δ7Li is likely the dominance of bulk dissolution (variable contribution from thermal waters) with a more restricted formation of secondary minerals in hydrothermal systems over weathering at the surface during Li release from the host rocks. This was recently demonstrated for ground waters that were (partially) released with high Li contents from geothermal fields in the Western Cordillera of the active arc and exported to foothill basins interacting with various rock reservoirs in the water table (Fig. 6; Godfrey et al. 2019). The variable 87Sr/86Sr, i.e. the different host rocks of the Li, shows no correlation with the δ7Li.

Insights from boron isotope compositions

It has been shown, that the B isotope composition of fluids, borate minerals and salts from the Puna Plateau can be used to distinguish between the source rocks sampled
by the mineralising fluids (Kasemann et al. 2004; Schmitt et al. 2002). The Palaeozoic basement was identified as the most significant boron source in the eastern Puna and Eastern Cordillera. Another important boron contribution comes from Cenozoic ignimbrites with dominance in the western Altiplano–Puna; Cenozoic andesites and Mesozoic limestones provide a third source. In contrast to lithium, the B isotope composition of the fluids appears to be not affected by secondary processes, i.e. secondary mineral precipitation, but rather directly reflects variations in the isotope composition of the source rocks (Kasemann et al. 2004; Schmitt et al. 2002). The B isotope composition of the Salar de Pozuelos surface brines (−19.3 to −1.8‰) encompasses nearly the whole spread of data measured for brines and thermal spring fluids across the eastern and western Puna (Fig. 4; −18.3 to + 5.7‰; Kasemann et al. 2004). The boron isotope composition for the salt on the surface (−19.9 to −8.5‰) and in the core (−18.1 to −14.1‰) is similar to the surface brines but more homogeneous, duplicating the values reported for halite in the Puna (Kasemann et al. 2004). The two borate (ulexite) samples show significantly different δ11B values. The sample close to the Ordovician outcrop has a δ11B value (−17.5‰) that corresponds to values observed in other salars and thermal springs of the eastern Puna (e.g. Salar de Pocitos −16.7‰). In contrast, the ulexite value of −6.7‰ is much more positive and corresponds to values observed so far only for the Neogene Loma Blanca borate deposit (Kasemann et al. 2004). The spread in the boron isotope data in the fluids and the salts, and the two contrasting values in the borates, suggests mixing of boron leached from the same two sources the lithium was leached from, Palaeozoic basement and the Cenozoic ignimbrites, and supports the results obtained from the strontium data. In contrast to the lithium, the main source for the boron are the basement rocks as seen for other salars in the eastern Puna. The offset between the present day B isotope composition of the spring water (−9.1‰) and the travertine (−23.5‰) points to a pH value of about 8.5 (assuming 20 °C fluid temperature and the δ11B–ocean pH relationship, Klochko et al. 2006); a value that falls well between the pH range of 7.9 and 8.7 reported for thermal springs in the eastern Puna (Kasemann et al. 2004).

The survey samples from the Li-Triangle cover a considerable range in their B isotope composition (−6.5 to +16.7‰). In contrast to Pozuelos, most of the samples cluster approximately in the reported isotope range for thermal springs in the western Puna (−6.5 to +7.8‰) indicating B contributions mainly from the Cenozoic ignimbrites and andesites. Three samples from Salar Grande and Salar de Pedernales in Chile show δ11B values that are significantly more positive (+15.6 to +16.6‰) than previously measured fluids in the Central Andes (e.g. Schmitt et al. 2002; Kasemann et al. 2004) suggesting the presence of marine limestones of Mesozoic age, spatially correlated with the salars.

Conclusions

The Li contents of the Salar de Pozuelos in the Puna Plateau is generated by chemical weathering in a cold and dry climate, and the formation of secondary minerals with the accompanied large Li isotope fractionation dominates the uniform and high δ7Li values of the deposit. The isotope composition does not record changes in the weathering regime over at least 100 ka (or more) of deposition, despite various and well-documented pluvial—glacial periods in the Puna region. Sr and Nd isotope signatures of siliciclastic sediments from core DHH1 and Sr isotope signatures of all other salar basin and catchment area materials resemble those of large-volume ignimbrites as the dominant source for the Li with only minor additions from weathering of the prominent Ordovician rocks of the catchment area and hydrothermal leaching of the underlying Pz basement. The absence of volcanic edifices or extended volcanic deposits in the catchment area indicates the importance of air-fall or windblown material in the Cenozoic sediments of the catchment area and the salar basin. Compared with δ7Li signatures of other Li deposits and Li-bearing waters in the Puna and its western foothills, the Salar de Pozuelos represents the endmember of Li solubilisation from the host rock by near-surface chemical weathering. Sr and Li isotopes indicate that the degree of Li isotope fractionation is independent of the host rock, i.e. andesite, large-volume ignimbrite, and Palaeozoic basement as the principal contributors. Furthermore, δ7Li signatures of the evaporites give a clue on the mobilisation process of the Li between chemical weathering at the surface and processes including a higher proportion of bulk solution of the host rocks (bulk dissolution does not fractionate Δ7Liresidue-solution), e.g. in geothermal waters.

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