Alpine permafrost thawing during the Medieval Warm Period identified from cryogenic cave carbonates

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Abstract

Coarse crystalline cryogenic cave carbonates (CCC\textsubscript{coarse}) dated to the last glacial period are common in central European caves and provide convincing evidence of palaeo-permafrost during this time. Little is known, however, about the exact nature of the environment in which CCC\textsubscript{coarse} formed as no modern analogue setting is known. Here, we report the first findings of sub-recent, albeit inactive, CCC\textsubscript{coarse} from a cave of the Western Alps which is located in the present-day permafrost zone. The globular shape and the presence of ubiquitous euhedral crystal terminations are comparable to previously described aggregates from the last glacial period and strongly suggest that these aggregates formed subaqueously in pools lacking agitation. Furthermore, stable isotope values of mm-sized spheroids point to calcite precipitation in a closed system with respect to CO\textsubscript{2} strongly supporting the hypothesis of a cryogenic origin associated with the freezing of water ponds. U-series analyses revealed three clusters of late Holocene calcite precipitation intervals between 2129 and 751 a b2k. These ages correlate with known periods of elevated summer temperatures, suggesting that warming and thawing of the permafrozen catchment above the cave allowed water infiltration into the karst system. The growth of CCC\textsubscript{coarse} resulted from the re-freezing of this water in the still cold karst cavities.

1 Introduction

The distribution of alpine permafrost and its evolution in a changing climate is being extensively studied to identify potential hazards associated with instable debris slopes and rock-wall activity (e.g. Huggel et al., 2010; Fischer et al., 2012). Little is known, however, about the past evolution of permafrozen areas (French, 2011; Stoffel and Huggel, 2012) and a better identification of freeze and thaw cycles could contribute to the interpretation of geomorphic and ecological responses to specific climatic events (e.g. Gutiérrez, 2005).
Caves represent unique environments to identify present and past cryogenic activity because they are connected to atmospheric processes but are well protected from surface erosion. Field evidence of palaeoglacial activity in the subsurface include frost shattering of cave ceilings, speleothem damage, ice attachments, cryoturbatic movements and remobilization of cave sediments (e.g. Kempe et al., 2009; Luetscher, 2013). While several of these features could be caused by processes other than ice, a new class of carbonate deposits, cryogenic cave carbonates (CCC), has recently emerged as the most reliable indicator of (palaeo)glacial processes which can also be dated by U-series methods (Zak et al., 2004, 2008, 2012).

Cryogenic carbonates form by the segregation of solutes during freezing of water (e.g. Shumskii, 1964; Killawee et al., 1999). Depending on the conditions during calcite precipitation a large range of shapes and sizes of CCC can be observed (Lacelle, 2007; Lacelle et al., 2009; Richter and Riechelmann, 2008; Zak et al., 2008). Fine crystalline carbonate powder (CCC$_{fine}$), whose carbon isotopic composition exhibits large kinetic fractionation effects (Lacelle et al., 2006; Spötl, 2008), is typically associated with rapid (seasonal) freezing under open system conditions (i.e. continuous exchange of CO$_2$ with the cave atmosphere). In contrast, a negative correlation between $\delta^{18}$O and $\delta^{13}$C values indicates that precipitation of coarse crystalline cave carbonates (CCC$_{coarse}$) occurs essentially in a closed system (Zak et al., 2004). Whilst site-specific $\delta^{13}$C offsets have been attributed to cave ventilation regimes before the water started to freeze (Richter et al., 2010), CCC$_{coarse}$$\delta^{18}$O values typically depart from the parent solution (Zak et al., 2004). Richter et al. (2010) concluded that the formation of CCC$_{coarse}$ most likely relates to the progressive freezing of water pools implying a steady heat exchange between the water and its surrounding environment preferentially achieved in the homothermic zone of a permafrozen karst system (cf. Luetscher and Jeannin, 2004).

In contrast to CCC$_{fine}$, which has been reported from alpine and arctic ice caves (Luetscher et al., 2007; Spötl, 2008; Richter et al., 2009; Lacelle et al., 2009), CCC$_{coarse}$ has been exclusively documented from presently ice-free Central European caves, located in former permafrost regions beyond the Pleistocene ice sheets (Zak et al., 2009,
Applying $^{230}$Th/$^{234}$U methods to determine the age of these CCC$_{coarse}$ has recently opened a door to constrain the timing of permafrost occurrences in the Pleistocene (Zak et al., 2012). However, the lack of modern analogues has severely limited a profound understanding of the processes leading to CCC$_{coarse}$ formation. This paper extends for the first time the record of CCC$_{coarse}$ into the Holocene using samples from a partly deglaciated alpine cave located in the present-day permafrost zone. We explore the palaeoclimate significance of these peculiar carbonate formations and demonstrate their potential to reconstruct the evolution of alpine permafrost during the late Holocene.

2 Study site

Leclanché cave is a 130 m-long cave system located at 2620 m a.s.l. (46° 20’ 42” N, 7° 15’ 47” E) in the Sanetsch area, western Swiss Alps (Borreguero et al., 2009). The cave opens with four individual entrances in a south-east facing rock cliff, at the base of the Schrattenkalk Formation, a Cretaceous platform limestone of the Helvetic realm (Wildhorn nappe, Mont-Gond unit; Badoux et al., 1959). The main cave passage, ca. 3 m wide and 1 m high, formed along a regional discontinuity (200/35) and is partly filled with massive congelation ice. In 2004, excavation of sediments obstructing the main conduit at 35 m from the cave entrance allowed speleologists to explore a 15 × 3 m wide chamber in the rearmost part of the cave (Fig. 1). This chamber comprises abundant cryoclasts (i.e. angular rock fragments produced by frost shattering) covered by aggregates of CCC$_{coarse}$ as well as fragments of flowstone. A ca. 10 m$^3$ perennial ice body was still present along the northern wall of the cave chamber in 2012, similar to what was observed during the first cave exploration in 2008.

Cave air temperature recorded since 1998 at the nearby Grotte des Pingouins (46° 21’ 9” N, 7° 16’ 36” E, 2333 m a.s.l.) locate the 0°C isotherm at 2260 m a.s.l., for a measured cave temperature gradient of 0.8°C/100 m (Borreguero and Pahud, 2004). This gradient is consistent with conspicuously dry cave conditions in Leclanché cave.
reflecting the scarcity of water infiltrating the permafrozen karst rock. Still, well-developed conduits may drain substantial amounts of melt water in summer and thus locally transfer heat to the host rock. However, this water is subject to rapid refreezing when intersecting larger cave passages due to an increased rock-water exchange surface.

The surface above Leclanché cave is characterized by a denuded karst comprising widespread karrenfields and numerous cave entrances (Borreguero et al., 2009). The area receives ca. 1900 mm of annual precipitation, a large part of which falls as snow. Morphological observations suggest that small glaciers extended down to an altitude of ca. 2350 m a.s.l. on the Lapi di Bou during the Little Ice Age, but we found no evidence for the presence of glaciers at the cave entrance during the Holocene (Borreguero et al., 2009). Frost shattering is the dominant geomorphological process forming large talus slopes at the base of the rock cliffs. Pioneer vegetation develops sporadically but is otherwise largely absent in the immediate cave surroundings. Alpine tundra vegetation is present up to ca. 2500 m a.s.l. and the local treeline, formed by *Larix decidua* and *Pinus cembra* (supra-subalpine belt) is currently located at ca. 2060 m a.s.l. (Berthel et al., 2012).

### 3 Methods

The CCC\textsubscript{coarse} aggregates were examined using a binocular stereomicroscope as well as in thin sections using transmitted-light (Leica M2 16A) and epifluorescence microscopy (Nikon Eclipse). Gold-coated samples were further examined by field-emission scanning electron microscopy (SEM, DSM 982 Gemini, Zeiss). Raman spectrometry was carried out using a Horiba Jobin-Yvon Labram-HR800 spectrometer. Individual CCC\textsubscript{coarse} aggregates were dissolved in 65 % suprapure HNO\textsubscript{3} and fluorescence properties were analysed using a Perkin-Elmer LS-55 spectrofluorometer. Results were interpreted from excitation-emission matrices obtained by collecting a series of 81 emission scans at 5 nm excitation wavelength intervals between $\lambda_{\text{ex}}$ 200 and 600 nm.
Seven whole CCC\textsubscript{coarse} aggregates were prepared for $^{230}\text{Th}/^{234}\text{U}$ age determination. Chemical separation and multi-collector inductively coupled mass spectrometric (MC-ICPMS) measurement of U and Th isotopic ratios were undertaken at the University of Minnesota using procedures similar to those described in Shen et al. (2012). Samples were pre-treated prior to chemical preparation in order to remove surface impurities from whole CCC\textsubscript{coarse} aggregates. Pre-treatment included either leaching in a weak 2 % HCl solution for 2–3 min (LEC-b and -c) or cleaning ultrasonically in 15 MΩ water for 10 min (LEC-e to -g). Two samples were not pre-treated (LEC-a and -d). Resulting sub-samples for chemical separation and purification were between 17 and 104 mg.

The extent of detrital $^{230}\text{Th}$ contamination was estimated and corrected for by measurement of the long lived chemically equivalent $^{232}\text{Th}$ and assuming a silicate bulk earth initial $^{230}\text{Th}/^{232}\text{Th}$ atomic ratio of $4.4 \pm 2.2 \times 10^{-6}$ (Wedepohl, 1995). Final ages are given as years before 2000 AD (a b2k).

Water samples were collected from active drips at Leclanché cave on 30 July 2012. The O isotope composition of water was determined by equilibration with CO\textsubscript{2} using an on-line continuous flow system (Gasbench II) linked to a Delta\textsuperscript{Plus} XL isotope ratio mass spectrometer. Calibration of the mass spectrometers was accomplished using VSMOW, GISP, and SLAP standards. The 1-sigma analytical errors on the $\delta^{18}\text{O}$ is 0.09 ‰. CCC samples were analysed for $\delta^{18}\text{O}$ and $\delta^{13}\text{C}$ on the Delta\textsuperscript{Plus} XL with an analytical precision (1σ) of 0.08 ‰ for $\delta^{18}\text{O}$ and 0.06 ‰ for $\delta^{13}\text{C}$ (Spötl and Vennemann, 2003), reported on the VPDB scale and calibrated against NBS19.

4 Results

4.1 Petrography

CCC\textsubscript{coarse} occurs as loose aggregates on top of cryoclasts in the rearmost part of Leclanché cave, spread over an area of between 2 and 4 m\textsuperscript{2} (Fig. 2a). No evidence of speleothem deposition postdating the formation of these aggregates was found. The
amber-coloured samples, 1–4 mm in size (2.9 ± 0.9 mm; n = 62), form spheroids which sometimes collate into chains, up to 15 mm long (Fig. 2b–e). A rough estimate identified ca. 400 spheroids per 50 cm², representing a total mass of ca. 15 g of secondary carbonate. Raman spectroscopy confirmed that calcite is the only mineral present, although an earlier calcite generation is commonly observed in the core of the spheroids. No evidence of detrital nuclei, however, was found in thin sections. The CCCcoarse show strong epifluorescence under the microscope and dissolved samples further exhibit fluorescence centres (λex : λem) at 250–280 : 400–460; 260–300 : 325–375, and 320–380 : 400–425 nm.

The globular shape and the presence of ubiquitous euhedral crystal terminations (Fig. 3a, b) strongly suggest that these aggregates formed subaqueously in pools lacking agitation. The internal structure consists of elongated calcite crystals which are themselves composed of crystallites, giving rise to sweeping extinction patterns under cross-polarized light. These rays of crystals form knob-like features on the surface of the spheroids (Fig. 3c, d). Some spheroids, however, show a dent where crystal growth was apparently blocked giving rise to a conspicuously smoother surface (Fig. 3e, f). These concave parts (typically no more than one or two per spheroid) most likely represent the former interface to adjacent spheroids.

4.2 Isotopic composition

δ18O values of the remnant cave ice and active drip waters range between −10.9 and −12.0 ‰, consistent with summer precipitation data at similar altitude in this region (Schürch et al., 2003). Stable isotope values of fossil stalagmites and flowstone from the Sanetsch area vary between −6.8 and −10.2 ‰ for δ18O. Elevated δ13C values in these speleothems, which range between −0.5 and +4.5 ‰, reflect the lack of soil cover in the hydrological catchment which may additionally be affected by kinetic effects. Non-equilibrium fractionation effects are also typical for CCCfine from this cave, with δ13C values reaching up to +15.5 ‰.
In contrast, bulk $\delta^{18}O$ values of $\text{CCC}_{\text{coarse}}$ range between $-15.2$ and $-17.5 \%$ and from $-0.6$ to $+2.2 \%$ for $\delta^{13}C$ (Fig. 4). The two isotopes are strongly anti-correlated ($r^2 = 0.93; n = 22$). Transects milled at 250 $\mu$m increments reveal a continuous enrichment in $^{13}C$ from the core to the spheroid rim along with a decrease in $\delta^{18}O$ by nearly 2 $\%$ (Fig. 5). Both trends follow a third-order power function consistent with a Rayleigh fractionation process and are attributed to progressive freezing of ponded water.

4.3 U/Th dating

Leclanché $\text{CCC}_{\text{coarse}}$ are rich in U, displaying $\text{^{238}U}$ concentrations between 1.9 and 2.3 $\mu$g$^{-1}$ (Table 1). Measured $\text{^{230}Th/^{232}Th}$ atomic ratios are less than $50 \times 10^{-6}$ indicating significant detrital $\text{^{230}Th}$ contamination. Coupled with low concentrations of authigenic $\text{^{230}Th}$, the precision of the final ages is significantly compromised, ranging between 7 and 12 %. Differences in pre-treatment methods appear to have had negligible effect on the final ages, with both leached and ultrasonically cleaned samples yielding coeval ages.

Results obtained from seven $\text{CCC}_{\text{coarse}}$ samples yielded consistent clusters of ages between 751 and 2129 a b2k (Table 1). Two coeval samples were deposited between 751 $\pm$ 55 and 823 $\pm$ 58 a b2k, whereas another four samples provide an average age of 1073$\pm$72 a b2k (1$\sigma$). One sample suggests a significantly older age of 2129$\pm$235 a b2k.

5 Discussion

For the first time, sub-recent $\text{CCC}_{\text{coarse}}$ were observed in a cave located in the alpine permafrost zone. The present-day cave temperature inferred from monitoring data in caves of the surrounding karst system suggests a mean annual air temperature of $-2.5 ^\circ C$ at Leclanché cave. Despite the presence of substantial cave ice deposits leading to the occasional formation of fine crystalline cryogenic powders, no active $\text{CCC}_{\text{coarse}}$ growth was observed in Leclanché cave. The latter supports the hypothesis
that the formation of CCC_{coarse} is a site-specific, possibly short-lived process associated with a particular cave environment.

A variety of morphological types of CCC_{coarse} have been described from former ice caves in Central Europe. Zak et al. (2012) grouped them into three broad categories: (i) individual crystals and random or organized aggregates, (ii) raft-like crystal aggregates, and (iii) fine to coarsely crystalline spherical/globular forms. Members of the first two types are conspicuously absent in Leclanché cave where globular forms are the only type present. Hemispheric spheroids and in particular cupola-shaped forms with a hollow interior as described from the Malachitdom Cave in Germany (Erlemeyer et al., 1992; Schmidt, 1992; Richter and Riechelmann, 2008) are absent as well. Chain-like linked spheroids were likely more abundant in Leclanché cave judging from the dents seen in several of the spheroids but apparently disintegrated when the ice melted. Such aggregates have been reported from several Pleistocene ice caves in Central Europe (e.g. Richter and Riechelmann, 2008; Richter et al., 2010).

The exclusive occurrence of rather uniform spheroids (and short chains thereof) contrasts with the typically more mixed appearance of CCC_{coarse} types in previously described Pleistocene permafrost caves. The latter have been ascribed to several cycles of freezing and complete thawing of these subsurface ice bodies (e.g. Richter et al., 2010). In contrast, the samples from Leclanché cave point to a rather common origin of these spheroids. Preliminary investigations of CCC_{coarse} fluorescence properties further suggest that microbial activity was possibly present during calcite precipitation (Birdwell and Summers Engel, 2010).

Rough estimates of carbonate mass balances confirm that the mass of CCC_{coarse} found in the cave is consistent with the freezing of individual water ponds, typically in the order 0.1 m³. At a permafrost temperature of −2.5°C, which is equivalent to the modern cave temperature, freezing of such water ponds likely takes place within a few hours to a maximum of a few days. This inferred freezing rate is in apparent conflict with the compact, elongated columnar calcite fabric of the CCC_{coarse} spheroids which points to slow crystallization rates. We therefore associate the formation of CCC_{coarse}
with the sporadic infiltration of water due to snow melting above the cave. $^{230}$Th/$^{234}$U dating indicates that all spheroids formed in the Late Holocene. Moreover, six of the samples revealed ages of $978 \pm 160$ a BP (1$\sigma$), i.e. coeval with the Medieval Warm Period (MWP) characterized by elevated summer temperatures (Mangini et al., 2005; Büntgen et al., 2011; Fig. 6). Interestingly, one sample dated at $2129 \pm 235$ a BP, falls within the Roman Warm Period which was also characterized by a succession of warm climate episodes (Büntgen et al., 2011) and reduced glacier extents (Holzhauser et al., 2005).

The formation of CCC$_{coarse}$ during warm climate episodes affecting permafrozen karst environments is consistent with observations of cave ice formation in the Sanetsch area (Borreguero et al., 2009). The rising temperature in the epikarst (i.e. surficial zone of a karst system) favours the infiltration of water which eventually refreezes within the cold cave environment. This process remains active until the conduit becomes completely obstructed by cave ice, preventing further infiltration. In contrast, large water inlets may advect sufficient energy to maintain a local thermal anomaly inhibiting the formation of cave ice. In the presence of irregular (concave) ice surfaces, some of this water may be temporarily confined in pools followed by progressive freezing, ultimately leading to the precipitation of CCC$_{coarse}$.

The progressive freezing process is mirrored by the stable isotope composition across individual CCC$_{coarse}$ spheroids (Fig. 5). $\delta^{13}$C values evolve along a fractionation path consistent with a closed system with respect to carbon, thus supporting the hypothesis of a shallow pool isolated from the cave atmosphere by a surficial ice layer. CO$_2$ released during calcite precipitation escaped the system, either by entrapment in the ice as gas inclusions or by slow diffusion through the ice lid. Laboratory experiments showed that CO$_2$ concentrations of occluded gas bubbles may reach up to 63 vol. % (Killawee et al., 1998) but the resulting effect on carbon isotope fractionation is not fully understood yet. Richter et al. (2010) associated distinct $\delta^{13}$C values of CCC$_{coarse}$ to the cave ventilation regime at the onset of freezing, but seasonal soil activity in the active layer above the cave could also play a role. Lacelle (2007), however,
stated that the $\delta^{13}$C value of calcite during equilibrium freezing is not simply controlled by the initial composition of the water, but also by changing physical and geochemical conditions as freezing progresses. Regardless of the precise process, fractionation of carbon isotopes will proceed concurrently with the precipitation of CCC$_{\text{coarse}}$ (Zak et al., 2004).

The slow freezing of water under closed-system equilibrium conditions gives rise to progressive $^{18}$O depletion in the residual water due to the preferential incorporation of the heavier isotope into the growing ice (O'Neil, 1968; Jouzel and Souchez, 1982; Souchez and Jouzel, 1984). The highest $\delta^{18}$O values of calcite, therefore, likely reflect the composition of the initial water at the onset of calcite precipitation (Zak et al., 2012). The difference of nearly 7 ‰ between inferred $\delta^{18}$O values and the measured seepage water (ca. −11‰), however, strongly suggests that the freezing of the water pond started some time before the precipitation of calcite.

6 Conclusions

Permafrost typically prevents the infiltration of water below the active layer. Although air and water flow in high-permeable karst systems may locally transfer sufficient heat to preserve a rudimentary drainage network, seepage and fracture flow is largely absent from frozen cave passages. In high alpine karst systems, thawing of ice-filled cavities and the subsequent flow of water through ice-free conduits is therefore a perceivable impact of a warming climate.

In contrast, the temporary obstruction of cave passages by ice and the slow freezing of water ponds in the homothermic zone of a karst system represent salient features of a still permafrozen environment. Associated CCC$_{\text{coarse}}$ therefore not only provide a clear indication of permafrost but also provide an archive for dating periods of melting in the hydrological catchment area.

This study demonstrates that CCC$_{\text{coarse}}$ can be successfully used to identify and date Holocene permafrost thawing events. If verified in other caves, CCC$_{\text{coarse}}$ has
the potential to provide precise chronologies of past warm episodes in areas where palaeoenvironmental proxy data are scarce. Cryogenic calcite could therefore contribute to the timing of geomorphic events associated with permafrost degradation, including debris slopes and rock fall activity.

Acknowledgements. This study would not have been possible without the systematic explorations conducted by several generations of speleologists in the Sanetsch area. We are particularly indebted to G. Heiss and P. Tacchini who contributed to the discovery and careful documentation of Leclanché cave. R. Shone is acknowledged for the photographs of the cryogenic cave calcites and K. Pfellant helped with SEM work. M. Luetscher acknowledges support by the Austrian Academy of Sciences (APART; project no. 715010).

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Table 1. U and Th concentrations, isotopic activity ratios and ages of coarse-crystalline cryogenic cave carbonates from Leclanché cave, Switzerland.

<table>
<thead>
<tr>
<th>Sample</th>
<th>Lab Number</th>
<th>238$^{\text{U}}$ Number</th>
<th>238$^{\text{U}}$</th>
<th>232$^{\text{Th}}$</th>
<th>230$^{\text{Th}}$/232$^{\text{Th}}$</th>
<th>δ234$^{\text{U}}$</th>
<th>230$^{\text{Th}}$/238$^{\text{U}}$ (activity) (uncorrected)</th>
<th>230$^{\text{Th}}$ Age (a)</th>
<th>234$^{\text{U}}$ Initial (corrected)</th>
<th>230$^{\text{Th}}$ Age (a b2k)</th>
<th>230$^{\text{Th}}$ Age (a corrected)</th>
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<tbody>
<tr>
<td>LEC-a</td>
<td>GM48</td>
<td>2251 ± 8</td>
<td>7639 ± 155</td>
<td>49.1 ± 1.0</td>
<td>207.3 ± 2.7</td>
<td>0.0101 ± 0.0001</td>
<td>917 ± 6</td>
<td>207.8 ± 2.7</td>
<td>823 ± 58</td>
<td></td>
<td></td>
</tr>
<tr>
<td>LEC-b</td>
<td>GM82</td>
<td>2042 ± 2</td>
<td>11415 ± 229</td>
<td>40.2 ± 0.9</td>
<td>208.4 ± 1.7</td>
<td>0.0136 ± 0.0002</td>
<td>1237 ± 14</td>
<td>209.0 ± 1.7</td>
<td>1090 ± 96</td>
<td></td>
<td></td>
</tr>
<tr>
<td>LEC-c</td>
<td>GM83</td>
<td>1921 ± 2</td>
<td>8288 ± 166</td>
<td>47.0 ± 1.2</td>
<td>206.9 ± 1.9</td>
<td>0.0123 ± 0.0002</td>
<td>1117 ± 16</td>
<td>207.5 ± 1.9</td>
<td>1001 ± 75</td>
<td></td>
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<tr>
<td>LEC-d</td>
<td>GM143</td>
<td>2132 ± 2</td>
<td>29304 ± 587</td>
<td>32.5 ± 0.7</td>
<td>207.8 ± 1.6</td>
<td>0.0271 ± 0.0002</td>
<td>2472 ± 22</td>
<td>209.1 ± 1.6</td>
<td>2129 ± 235</td>
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<tr>
<td>LEC-e</td>
<td>GM174</td>
<td>2321 ± 2</td>
<td>16417 ± 329</td>
<td>31.3 ± 0.7</td>
<td>209.3 ± 1.7</td>
<td>0.0134 ± 0.0001</td>
<td>1218 ± 7</td>
<td>209.9 ± 1.7</td>
<td>1036 ± 121</td>
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<tr>
<td>LEC-f</td>
<td>GM175</td>
<td>2186 ± 2</td>
<td>15476 ± 310</td>
<td>34.6 ± 0.7</td>
<td>208.6 ± 1.6</td>
<td>0.0149 ± 0.0001</td>
<td>1349 ± 8</td>
<td>209.3 ± 1.6</td>
<td>1166 ± 121</td>
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<tr>
<td>LEC-g</td>
<td>GM176</td>
<td>2327 ± 2</td>
<td>7352 ± 147</td>
<td>48.3 ± 1.2</td>
<td>209.3 ± 1.7</td>
<td>0.0093 ± 0.0001</td>
<td>839 ± 12</td>
<td>209.7 ± 1.7</td>
<td>781 ± 55</td>
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</table>

Analytical errors are 2σ of the mean.

a $\delta^{234}\text{U} = ([\text{234}\text{U}/\text{238}\text{U}]_{\text{activity}} - 1) \times 1000$.

b $\delta^{234}\text{U}_{\text{initial}}$ was calculated based on $^{230}\text{Th}$ age (T), i.e. $\delta^{234}\text{U}_{\text{initial}} = \delta^{234}\text{U}_{\text{measured}} \times e^{\lambda_{234}T}$

Following decay constants were used: $\lambda_{230} = 9.158 \times 10^{-6} \text{a}^{-1}$ (Cheng et al., 2000); $\lambda_{234} = 2.826 \times 10^{-6} \text{a}^{-1}$ (Cheng et al., 2000); $\lambda_{238} = 1.551 \times 10^{-10} \text{a}^{-1}$ (Jaffey et al., 1971).

Corrected $^{230}\text{Th}$ ages assume an initial $^{230}\text{Th}/^{232}\text{Th}$ atomic ratio of $4.4 \pm 2.2 \times 10^{-6}$; i.e. values in secular equilibrium with a bulk earth $^{232}\text{Th}/^{238}\text{U}$ value of 3.8. The errors are arbitrarily assumed to be 50%.

c a b2k stands for before 2000 AD.
Fig. 1. (A) Site location (circle) plotted on the Alpine Permafrost Index Map (Boeckli et al., 2012); (B) Leclanché cave entrance (arrow) in the Lé rock cliff; (C) vertical cross section of the Leclanché cave system and location of the coarse crystalline cryogenic cave calcite (CCC\textsubscript{coarse}) findings.
Fig. 2. Coarse-crystalline cryogenic cave carbonate (CCC$_{\text{coarse}}$) observed in Leclanché cave. The loose calcite crystal aggregates occur on the surface of cryoclasts suggesting recent deposition. Calcite crusts (flowstone) coat several rock fragments suggesting an earlier phase of vadose speleothem deposition under non-freezing conditions. Individual CCC aggregates, depicted on the right side, have spherical shapes and are amber-coloured. Photographs courtesy of R. Shone.
Fig. 3. SEM photomicrographs of coarse crystalline cryogenic cave carbonates (CCC\textsubscript{coarse}) aggregates found in Leclanché cave. Note the ubiquitous presence of euhedral (rhombohedral) crystal terminations except for local dents (F) which represent the interface to an adjacent (now detached) spheroid.
Fig. 4. Stable isotope data of cryogenic cave calcite (CCC) found in Leclanché cave (red dots). Values from Central European caves (blue areas; Zak et al., 2012 and references therein) are shown together with the typical range of Alpine speleothems (unpubl. data).
Fig. 5. Stable isotope composition across three individual coarse crystalline cryogenic cave carbonate (CCC$_{coarse}$) spheroids from Leclanché cave. The marked negative correlation ($r^2 = 0.95$) between $\delta^{13}$C and $\delta^{18}$O argues against kinetic fractionation effects and, accordingly, calcite precipitation is believed to have occurred close to isotopic equilibrium with the parent water.
Fig. 6. (A) U-series ages of coarse crystalline cryogenic cave carbonates (CCC\textsubscript{coarse}) from Leclanché cave plotted against (B) June-July-August temperature anomalies and (C) April-May-June precipitation totals over the last 2500 yr reconstructed from Central European tree-rings (Büntgen et al., 2011).