Lost cold Antarctic deserts inferred from unusual sulfate formation and isotope signatures

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The Antarctic ice cap significantly affects global ocean circulation and climate. Continental glaciogenic sedimentary deposits provide direct physical evidence of the glacial history of the Antarctic interior, but these data are sparse. Here we investigate a new indicator of ice sheet evolution: sulfates within the glaciogenic deposits from the Lewis Cliff Ice Tongue of the central Transantarctic Mountains. The sulfates exhibit unique isotope signatures, including $\delta^{34}S$ up to +50‰ for mirabilite evaporites, $\Delta^{17}O$ up to +2.3‰ for dissolved sulfate within contemporary melt-water ponds, and extremely negative $\delta^{18}O$ as low as −22.2‰. The isotopic data imply that the sulfates formed under environmental conditions similar to today’s McMurdo Dry Valleys, suggesting that ice-free cold deserts may have existed between the South Pole and the Transantarctic Mountains since the Miocene during periods when the ice sheet size was smaller than today, but with an overall similar to modern global hydrological cycle.
The ability to understand how Antarctic glaciers and terrestrial ecosystems will change in the future is predicated on knowing how past events impacted on the development of Antarctica's ice mass. One of the contentious factors of the East Antarctic Ice Sheet (E AIS) evolution is whether a massive contraction to 1/3 of its present volume occurred in response to Pliocene warmth, a climate event that may be on par with what is currently occurring as a result of global warming. Present knowledge of the history of the Antarctic continent relies heavily on analyses of ice cores, marginal Antarctica (off-shore sediment cores, the McMurdo Dry Valleys (MDVs)) and a few outcrops along the Transantarctic Mountains (TAM)3-6. In contrast, terrigenous geochemical proxies from the Antarctic interior provide direct evidence for assessing ice volume fluctuations but are scarce due to ice sheet coverage. The terminal moraine environments on the polar side of the TAM are known to receive basal materials from the proximal East Antarctic Polar Plateau and are promising sites for hosting and preserving materials that reflect palaeoclimatological changes in the Antarctic interior7,8.

The Lewis Cliff Ice Tongue (LCIT, 84°14' S, 161°39' E, ~2,200 m a.b.s.l and > 500 km to the coast) moraine is one such site, located along the polar side of the central TAM (Fig. 1), immediately adjacent to the E AIS. Ice flow here is generally northward from the East Antarctic Polar Plateau and is diverted from the Beardmore Glacier and overrides the sub-ice high barrier along the southern edge of Queen Alexandra Range9,10. The glacial ice then enters the Walcott Névé and is eventually stranded by Mount Achernar at the LCIT. In association with ice from the Law Glacier, which produces enormous ice-cored moraine ridges, the moraine materials at the LCIT are forced northward and upward pushing basal glacier materials vertically up to the morainal surface. Support for the occurrence of this process includes the presence of a large population of meteorites at the LCIT9 and Sirius group sedimentary deposits in nearby outcrops (for example, Mount Sirius6). Moraine materials consist of dolerite, basalt, sandstone, siltstone, shale and limestone11,12, which are a collection of bedrock types from the neighbouring regions, namely the Colbert Hills12, the glacier path beside the Queen Alexandra Range and the Beardmore Glacier13. The annual average temperature at the LCIT is ~30°C or colder14. Even at the height of summer, air temperatures rarely exceed ~10°C. Strong Katabatic winds blowing from the ice sheet interior add to the ablation effects on surface ice and also act as a mechanism to limit transport of material from the Ross Sea to the South.

The LCIT moraine consists of a series of alternating ridges where evaporites are found residing on the slope of concavities, with occasional melt-water ponds in surrounding depressions that contain dissolved salts9,15. In the 2005–2006 field season, our team discovered a variety of sulfates in the form of massive clear crystals covered by whitish and powdery materials16,17. Time-resolved X-ray diffraction measurements showed that the evaporites are composed mostly of mirabilite (Na₂SO₄·10H₂O) in the interior, and minor components of thenardite (Na₂SO₄), nacolite (NaHCO₃), trona (Na₂ (CO₃(HCO₃))·2H₂O) and borax (Na₂B₄O₅·(OH)₄·8H₂O) on the exterior surface that appear to be the result of dehydration and weathering processes18. The water ponds are slightly alkaline, with pH ranging from 8.75 to 9.95. The salinity of melt water in ponds ranged from fresh to brackish, with major ions being Na⁺ > Cl⁻ > SO₄²⁻ > , NO₃⁻ > F⁻ (Supplementary Table 1).

Stable isotopes of water. δ¹⁸O and δD values of precipitation (snow), glacial ice, secondary glacial ice (ice lenses) and lake water ranged from −59.2 to −29.7‰ and −456.0 to −231.7‰%, respectively (Supplementary Table 2, Fig. 3). The δ¹⁸O and δD of various water types show that the pond water derives from glacier ice and local snow precipitation and the pond has experienced a small degree of evaporation (Supplementary Discussion and Supplementary Fig. 3).

Stable isotopes of sulfates. The LCIT mirabilite sulfates have uniform isotopic compositions: highly positive δ³⁴S ( +49.1 ± 0.2‰), highly negative δ¹⁸O (−16.9 ± 0.3‰) and negative Δ¹⁷O values (−0.56 ± 0.05‰, Fig. 2). However, compared with mirabilite sulfate, the pond sulfate has lower δ³⁴S values (ranging from +12 to +20‰) and a wider range of
negative $\delta^{18}O$ values (−12.8 to −22.2‰). Also, pond sulfate has the most negative $\delta^{18}O$ value among all known sulfates (−22.2‰; ref. 20). The most important feature of the pond sulfate is its non-mass-dependent $\Delta^{17}O$ enrichment ($D^{17}O = +1.36$ to +2.35‰). This is the only known dissolved sulfate from a natural body of water that possesses positive $\Delta^{17}O$ values (ref. 20 and references therein) and is in sharp contrast to the mirabilite sulfate which does not have positive $\Delta^{17}O$ values. We analysed dissolved sulfates from meromictic Ace Lake (Vestfold Hills, 68.473°S, 78.189°E) for comparison with LCIT sulfates. Isotope compositions from water depths of 5.5 m (aerobic mixolimnion) and 15.5 m (anaerobic monimolimnion) were: water, $\delta^{18}O$ of −16.6 and −16.7‰, respectively; dissolved sulfates, $\delta^{34}S$ of +40.0, +67.1‰; and $\delta^{18}O$ of +3.3, +8.5‰, respectively. These data are consistent with those recorded for samples from Ace Lake taken in 1970 (ref. 23).

Discussion
The stable isotope compositions of LCIT sulfates offer the most diagnostic clues for the formation conditions of sulfates. The unusually high $\delta^{34}S$ (~ +49‰) of mirabilite likely arises from the dissimilatory microbial sulfate reduction (DSMR) reaction when sulfate is converted to reduced sulfur species, leaving a large enrichment of $^{34}S$ in the residual dissolved sulfate. The high solubility of sodium sulfate requires a high salinity for its precipitation. Therefore, formation of mirabilite with a high $\delta^{34}S$ would require DSMR to elevate the $\delta^{34}S_{SO_4}$ throughout the water body before precipitation. Melt-water ponds at the terminus of glaciers along the TAM are typically small, shallow and well aerated (Fig. 1), and they are therefore not suitable for DSMR. Reducing microenvironments may exist if a pond is covered by algal mats or if sufficient sediment depth and stability exists for the development of benthic communities. However, no algae or benthic mats were observed in the ponds at the LCIT. Even with such bio-reduction, the turnover of the sulfur reservoir would be very rapid, resulting in virtually no net sulfur isotope enrichment. On the basis of sulfate isotope signatures and direct assessment of microbial communities, DSMR in Antarctica has been inferred to occur mainly in lakes. Ace Lake is a coastal, marine-derived, meromictic system in the Vestfold Hills where DSMR occurs and it is the only water body measured to date in Antarctica with overall highly enriched $^{34}S$ (average $\delta^{34}S$ of +42‰ (ref. 23)); it therefore appears to be a modern-day
analogue for the production of mirabilite in the LCIT. DMSR is also reported for subglacial environment, but sulfate redox cycling therein produces isotope signatures distinctly different from those of the LCIT. Alternatively, thermochemical sulfate reduction may also introduce high \( \delta^{34}S \), but this process requires high reaction temperature (\( > 100 \, ^\circ\text{C}; \text{ref. 36} \)) and is not known to occur in Antarctica.

The negative \( \delta^{18}O \) values (\( -16.9 \pm 0.3\% \)) of the mirabilites is indicative of the incorporation of light glacier water oxygen. As direct oxygen isotope exchange between sulfate and water is extremely slow\(^{37,38} \), the low \( \delta^{18}O \) values must have been acquired through incorporation of glacial melt-water oxygen during sulfate formation and/or through microbial sulfate reduction and sulfide oxidation cycles\(^{39,40} \). The \( \delta^{34}S \) values in Ace Lake show an apparent steady state for sulfur redox cycling\(^2 \) (Supplementary Table 2), whereas the apparent \( \Delta^{18}O_{\text{SO}_4-\text{H}_2\text{O}} \) at a range of +19.4 to +25.2% (Supplementary Table 2) approaches the predicted sulfate–water oxygen isotopic composition\(^{41} \). Therefore, assuming the LCIT mirabilite sulfate was also in oxygen isotopic equilibrium with water, our data imply that the \( \delta^{18}O \) of water was \(< -45\% \) at the time the LCIT mirabilite precipitated; a value that is close to that of the present-day EAIS\(^3,42 \). The uniform sulfur and oxygen isotope composition of mirabilite throughout the sampling sites also suggests they probably originated from the same water body.

The pond sulfate possesses extreme \( \delta^{18}O \) depletion (\(-12.1 \to -22.2\% \)) and a distinct non-mass-dependent \( \delta^{17}O \) enrichment (\( \Delta^{17}O \) at \(+1.36 \to +2.35\% \)) which is the only case known for modern natural water bodies. The positive \( \Delta^{17}O \) values are a clear indication that a portion of the pond sulfate derives from SAS, as this unique signature came ultimately from ozone and derivatives whose \( \delta^{18}O \) values can also be highly positive and variable in the troposphere and stratosphere\(^{15,44} \). However, despite having positive \( \Delta^{17}O \) values similar to the sulfates in soils of the MDVs and snow/ice cores from the EAIS\(^{45,46} \), the LCIT pond sulfates have much lower \( \delta^{18}O \) values (Fig. 2). Therefore, a source of sulfate in addition to SAS and with extremely low \( \delta^{18}O \) must have contributed to the pond sulfates.

It is noteworthy that the LCIT pond sulfates have \( \delta^{18}O \) values closer to sulfates in the MDVs than in ice cores. The most likely
Some of the materials were ‘dumped’ at the LCIT site. From the TAMs to the Pensacola Basin, marked by the rectangle. The interior cold desert was likely situated in an plateau. I: Transantarctic Mountains, II: Wilkes Basin, III: Aurora Basin and IV: Pensacola Basin. The interior cold desert was likely situated in an area from the TAMs to the Pensacola Basin, marked by the rectangle. (a) Transport model for modern-day LCIT sulfate formations. Arrows represent glacial ice movement from the EAIS to the LCIT. After renewed ice expansion, mirabilite precipitated in lakes and atmospheric sulfate-bearing regolith from the interior cold deserts was exhumed and carried by glacial ice towards the Transantarctic Mountains and the Ross Sea until some of the materials were ‘dumped’ at the LCIT site.

A possible explanation is that both LCIT and DMVs sulfates have a component derived from Antarctic local weathering. Oxidation of sulfide minerals by glacial water can introduce sulfate $\delta^{18}$O as low as $-19\%$ (ref. 47). Assuming the formation pathways of SAS did not change drastically during the past few million years, the $\delta^{18}$O of sulfate from oxidative weathering can be calculated by treating LCIT pond sulfate as a simple mixing of SAS, oxidative weathering and minor sea-salt sulfate (SSS).

$$x\delta^{18}O_{SAS} + y\delta^{18}O_{SSS} + (1-x-y)\delta^{18}O_{Oxidative weathering} = \delta^{18}O_{Pond sulfate}$$  \hspace{1cm} (1)

$$x\Delta^{17}O_{SAS} + y\Delta^{17}O_{SSS} + (1-x-y)\Delta^{17}O_{Oxidative weathering} = \Delta^{17}O_{Pond sulfate}$$  \hspace{1cm} (2)

where $x$ and $y$ are the mole fractions of the SAS and SSS contributions, respectively. Conservatively, $\delta^{18}O_{SAS}$ and $\Delta^{17}O_{SAS}$ were set at $-3\%$ and $+3.5\%$, respectively.\textsuperscript{17,27,45,46,48} (National Snow and Ice database, Fig. 2a) and $\Delta^{17}O_{Oxidative weathering}$, $\delta^{18}O_{SSS}$, $\Delta^{17}O_{SSS}$ and $y$ were set at $-0.50\%$, $+10.3\%$, $-0.1\%$ and $5\%$, respectively. Mass conservation calculations for isotope values show that $\sim 50$ to $70\%$ of the LCIT pond sulfate is SAS, and the $\delta^{18}O_{Oxidative weathering}$ ranges from $-43.5$ to $-58.3\%$, which are much more negative than any previously recorded values. The low values put the glacier water $\delta^{18}O$ at $\sim 60 \pm 10\%$ (ref. 26), which is also in the $\delta^{18}O$ range of cold-based glaciers from higher latitudes of the EAIS. The accumulation of sulfate with positive $\Delta^{17}O$ is characteristic of desert environments exposed to long periods of hyperaridity.\textsuperscript{16,21} The $\Delta^{17}O$ and $\delta^{18}O$ of pond sulfate thus point to the existence of one or more ice-free and arid surfaces (that is, cold deserts) at high latitudes.

As the mirabilite and pond water sulfates have distinct isotopic compositions, the mirabilite evaporites could not have come from the pond water, or vice versa. Instead, the isotopic composition is consistent with an Ace Lake-type reservoir for the formation of the mirabilite sulfates and a hyperarid desert with oxidative weathering for the pond sulfates. These conditions closely resemble the key geochronological and climatic characteristics of present-day MDV, and to an extent, the Vestfold Hills. The cold deserts could have existed in the vicinity of the LCIT and/or a region further towards the interior of the EAIS along the glacier flow path. As the LCIT geological settings cannot generate mirabilite sulfate with high $\delta^{18}S$ (+49.8\%) or accumulate a significant amount of SAS with high $\delta^{18}O$ (+2.1\%), it is very likely that the mirabilite and pond sulfates are terrigenous, having been originally transported from deeper within the Antarctic interior.

Therefore, we argue that the LCIT sulfates originated from a time when the ice cap covered a smaller area than present, along the flow paths of the Beardmore Glacier\textsuperscript{2,9} that includes some of the topographic lowlands along the Queen Alexandra Range and the Pensacola Basin located between the TAM and the East Antarctic Polar Plateau. We suggest that the distribution of cold deserts may be patchy among relatively large ice-free lands, but it is unlikely that a few ice-free outcrops existed within thick ice caps. This once MDV-like environment enabled SAS to accumulate to high levels, glacial melt remnant water-fed lakes allowed for extended periods of DMSR and a relatively warm ambient temperature allowed for oxidative weathering of sulfur-bearing bedrocks. As the climate cooled and glaciers advanced, mirabilite beds may have been formed as a result of salt concentration by freezing of lakes. The mirabilite beds, together with regolith containing SAS and weathering-produced sulfates, would have been disrupted and carried by glaciers and eventually stranded at the LCIT moraine (Fig. 3). The exact location and the size of the cold deserts are yet to be determined. The provenance or geological terrains of these sediments may be traced using radiogenic isotopes and more data from other localities along the TAM\textsuperscript{4}.

The initial age of the mirabilite sulfate salt formation would help determine its origin. Unfortunately, evaporite formation is difficult to date and inferences about age must be gleaned from elsewhere. The ice sheet producing the Beardmore Glacier was suggested to have been overriding the TAM in the Quaternary\textsuperscript{2,9} indicating that the cold deserts would be older than the Quaternary. Glacier palaeotemperature can be estimated based on a robust relationship between $\delta^{18}O$ of precipitation and the local temperature\textsuperscript{29,50}. The $\delta^{18}O$ of the glacier water responsible for the LCIT sulfate $\delta^{18}O$ is comparable, or lower, than those of the Dome C and Vostok ice cores that represent the past hundreds of thousands of years, suggesting that the temperature before emergence of the hypothesized interior cold deserts was as low as it is today\textsuperscript{49}. However, the Miocene ice cap would have had a much higher $\delta^{18}O$ than the present-day glacial $\delta^{18}O$ because: first, the temperature of the Antarctic interior was not expected to be as low today in the Miocene\textsuperscript{51}; and second, the Southern

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**Figure 3** Conceptual model for cold desert formation and LCIT sulfate transport. (a) Illustration of Antarctic landscape during Pliocene warmth with assumption that near or lower than sea level terrains were ice free and the ice sheet remained on Transantarctic Mountains and east Antarctic plateau. I: Transantarctic Mountains, II: Wilkes Basin, III: Aurora Basin and IV: Pensacola Basin. The interior cold desert was likely situated in an area from the TAMs to the Pensacola Basin, marked by the rectangle. (b) Transport model for modern-day LCIT sulfate formations. Arrows represent glacial ice movement from the EAIS to the LCIT. After renewed ice expansion, mirabilite precipitated in lakes and atmospheric sulfate-bearing regolith from the interior cold deserts was exhumed and carried by glacial ice towards the Transantarctic Mountains and the Ross Sea until some of the materials were ‘dumped’ at the LCIT site.
Ocean provided relatively more moisture than it presently does, thus reducing the latitudinal effect for the vapour movement. The δ18O of the EAIS was estimated to be higher than ~35‰ in the Oligocene and Miocene23,32. This is also supported by a recent leaf-wax study that showed that the δD of precipitation along the Antarctic coast was as high as ~50‰ during 20 to 15.5 Myr (ref. 54), which indicates a much higher coeval δ18O of precipitation than today. Our sulfate oxygen isotope data therefore point to formation of LCIT sulfates with coeval waters similar to modern-day glacier waters. This suggests that the LCIT sulfates did not form during the early stages of Antarctic ice cap formation. Instead, the hypothesized interior cold deserts must have formed during a time of relative warmth but a period long after the global hydrological cycle was established similar to that of today. The Pliocene epoch is therefore the most feasible geological time for this occurrence. If we assume the interior cold deserts had 10% of the sulfate concentration as present-day upland soils in the MDVs, the interior cold deserts would have persisted for at least 1 Myr (ref. 16).

Our data support the existence of an ice-free, warmer but dry environment underneath the interior of the high EAIS in the past, likely during the Pliocene warmth. The implied massive ice contraction associated with the emergence of interior cold deserts is supported by recent evidence that during the Pliocene warmth, ~500 km of ice retreated in the Wilkes Basin, and the ice-sheet basal thermal regime transitioned from polythermal to cold4,5. The geological processes leading to the formation of the interior cold deserts following glacier melt are not clear. However, the environment is topographically favourable, with lowlands (for example, Pensacola basin) surrounded by a high-altitude barrier to prevent glacier ice flow and katabatic winds to prevent moisture entry from the sea; conditions that are similar to today’s MDVs.

Methods

Sample collection. Samples from the terminal moraine of the LCIT were collected in an area of ~5 km2 during the 2005–2006 field season (Fig. 1). Samples consist of solid material from the evaporite mounds (mineralogically almost entirely mirabilite), frozen precipitate (ice and snow) and liquids (pond water; Fig.1, Supplementary Fig. 1 and Supplementary Table 2).

Pond water chemical and isotopic analysis. Pond melt-water chemistry was analysed in the Department of Geology and Geophysics at Louisiana State University on an Ion Chromatographer (Dionex ICS-3000) for both cations and anions. Water and ice (δD and δ18O) samples (snow, ice, pond melt water) were analysed using the CO2–H2O equilibrium method50 and the H2O–H2S equilibrium method51. Water isotope measurements were conducted on a Thermo Finnigan MAT 253 via a gas bench in a stable isotope lab at the NASA Johnson Space Center. δD and δ18O values were normalised52. The analytical precisions (1σ) are 0.05‰ and 0.1‰ for δD and δ18O, respectively.

Sulfate isotope analysis. All sulfate samples were converted to BaSO4 and then treated with diethyliamine triacetate and 0.1 M HCl repeatedly to avoid any possible occluded contaminants before the isotope analysis27. δ34S sulfate measurements were conducted in a high-temperature conversion elemental analyser system, through a confo-III interface on a Finnigan MAT 253. The δ17O compositions of sulfate were measured using a Finnigan MAT 253 dual-inlet mass spectrometer, using O2 generated from BaSO4 via a CO2-laser fluorination technique28. The analytical precisions (1σ) are 0.5‰ and 0.08‰ for δ34S, δ17O, respectively. Sulfate oxygen isotope analysis was conducted at Louisiana State University. δ17O sulfate values were measured on a Finnigan MAT 253 mass spectrometer in the Stable Isotope Research Facility at Indiana University. The analytical precision (1σ) is 0.3‰.

Isotope notations and standardization. All δ18O and δD values are reported as standard δ notation with respect to V-SMOW. δ34S sulfate is reported as δ17O–δ18O = −0.52 × δ18O + 25. The value 0.52 was chosen for consistency with results from previous studies27,28. We measured UWWG-2 garnet as the laboratory standard, assuming δ17O_UWWG-2 = 0.52 × δ18O_UWWG-2. δ34S values are reported with respect to V-CDT.

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Author contributions
R.A.S., R.P.H., D.L.B., E.T. and R.C. performed the field works and sampling; T.S., R.A.S. and P.N. conducted chemical and isotopic analysis; D.L.B. conducted XRD analysis; T.S., H.B. and R.A.S. contributed to the data interpretation; T.S., R.C., H.B. and D.L.B. wrote the paper. All the authors contributed to the discussions of this work.

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