Data Repository for "The lead-up to the Sturtian Snowball Earth: Neoproterozoic chemostratigraphy time-calibrated by the Tambien Group of Ethiopia"

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- This document accompanies the discussion contained in the main text. All the Python code
- used for this study, as well as the associated data tables not included in this document, can be
- ³ found at: https://github.com/Swanson-Hysell-Group/2018_Tambien_Group.

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4 Construction of the Chemostratigraphic Composite

- 5 The main text contains a composite δ^{13} C and 87 Sr/ 86 Sr curves for the Tonian and Cryogenian
- 6 that are time-calibrated by the record from Ethiopia and incorporate data from the literature
- 7 from numerous sources. Additional details associated with the data sets within this composite are
- provided below. The Python code used to develop the composite as well as the associated data
- ⁹ table can be found at: https://github.com/Swanson-Hysell-Group/2018_Tambien_Group.

10 Ethiopia

- 11 δ^{13} C and 87 Sr/ 86 Sr data from Ethiopia comes from the Tambien Group and are developed in
- Miller et al. (2009), Swanson-Hysell et al. (2015b), and this study. Combined with U-Pb ID-TIMS
- dates on zircons from Swanson-Hysell et al. (2015b) (815.29 \pm 0.32, 788.72 \pm 0.24, and
- 14 787.38±0.14 Ma) and new U-Pb ID-TIMS dates on zircons from MacLennan et al. (2018)
- $(735.25\pm0.25, 719.58\pm0.56, \text{ and } 719.68\pm0.46 \text{ Ma})$, the Tambien Group is now the source of the
- most temporally well-constrained pre-Sturtian chemostratigraphic dataset to date. We therefore
- use the Tambien Group δ^{13} C curve as the backbone for making correlations with other datasets.
- In the chemostratigraphic composite, the age of the initiation of the Sturtian Glaciation is set to
- ¹⁹ 717 Ma (discussed further in the 'Onset of the Sturtian Snowball' section).

20 Svalbard

- δ^{13} C and δ^{13} C and
- 22 in Halverson et al. (2007a) and Halverson et al. (2007b). However, a slightly stricter threshold for
- 23 87 Sr/ 86 Sr diagenesis is applied to the data included in our composite than in the original
- publication ([Sr] < 500 ppm). Additional $^{87}Sr/^{86}Sr$ data were published in Cox et al. (2016). The
- ²⁵ Polarisbreen Group, which unconformably overlies the Akademikerbreen Group, contains two
- 26 separate diamictite units which have been interpreted to represent the Sturtian and Marinoan

- 27 Glaciations respectively (Halverson et al., 2004). This correlation constrains the Akademikerbreen
- 28 Group to have been deposited prior to the Sturtian Glaciation. No direct geochronological
- 29 constraints exist for the Akademikerbreen Group although thermal subsidence models have been
- used to suggest a ca. 800 Ma age for the Bitter Springs Stage (Maloof et al., 2006). Therefore,
- the δ^{13} C curve from the group is correlated to that of the Tambien Group by aligning the start
- 32 and end of the Bitter Springs Stage and the nadir of the Islay Anomaly. This correlation results
- in a near constant sedimentation rate between these constraints, which is used to estimate the age
- of data that precedes the Bitter Springs Stage.

35 Greenland

- δ^{13} C and δ^{87} Sr/ δ^{86} Sr data from the Eleanore Bay Supergroup are developed in Cox et al. (2016).
- Halverson et al. (2004) approximated the age of the this succession to be ca. 800 Ma based on the
- correlation of lithological and δ^{13} C data to the Akademikerbreen Group of Svalbard, but no direct
- age constraints exist. Therefore, the age model is estimated based on aligning the δ^{13} C data with
- the δ^{13} C curve of the Akademikerbreen Group.

41 Australia

- δ^{13} C data from the Bitter Springs Formation are developed in Swanson-Hysell et al. (2010).
- Further ⁸⁷Sr/⁸⁶Sr data are developed in Cox et al. (2016). Similar to the Akademikerbreen
- 44 Group, the Bitter Springs Formation is unconformably overlain by Sturtian diamictite of the
- 45 Areyonga Formation (Swanson-Hysell et al., 2010), and thus constrains the Bitter Springs
- 46 Formation to have been deposited prior to the Sturtian Glaciation. However, no direct
- 47 geochronological constraints exist for the Bitter Springs Formation. Therefore, the δ^{13} C curve
- 48 from this group is correlated to that of the Tambien Group by aligning the start and end of the
- ⁴⁹ Bitter Springs Stage. Again, this correlation results in a near constant sedimentation rate
- 50 between these constraints, which is used to estimate the age of data that post-dates the Bitter

- Springs Stage. However, if the constant sedimentation rate is applied to δ^{13} C data from the
- 52 Bitter Springs Formation preceding the Bitter Springs Stage, there is a significant mismatch
- between these data and that of the Akademikerbreen Group. Therefore, these data were assigned
- slightly older ages than would be predicted by the constant sedimentation rate assumption in
- order to better match the δ^{13} C curves between these two sections.

56 Canada

71

 δ^{13} C and δ^{87} Sr/ δ^{86} Sr data from Canada come from multiple studies and localities.

 δ^{13} C data and geochronology from the Fifteenmile Group are developed in Macdonald et al. 58 (2010). Additional ⁸⁷Sr/⁸⁶Sr data are developed in Cox et al. (2016). The Fifteenmile Group is 59 unconformably overlain by temporally well-constrained (see the 'Onset of the Sturtian Snowball' section) Sturtian diamictite of the Upper Mount Harper Group (Macdonald et al., 2010). A U-Pb 61 ID-TIMS date on zircons within a tuff of 811.51±0.25 Ma can be tied directly to this curve, and, combined with dates from the Tambien Group $(787.38\pm0.14, 788.72\pm0.24, \text{ and } 815.29\pm0.32 \text{ Ma})$, suggests global synchroneity of the Bitter Springs Stage (Swanson-Hysell et al., 2015b). The nadir of the Islay Anomaly can also be easily identified and correlated. Furthermore, δ^{13} C data that 65 precede the Bitter Springs Stage correlate well with data from the Akademikerbeen Group, and thus were correlated based on similar δ^{13} C values. However, unlike other sections in which the Bitter Springs Stage is observed, the recovery from the interval of low δ^{13} C values appears to be much more gradual. Nevertheless, the end of the minimum δ^{13} C values is assumed to be

interpreted to exist between PF1 and PF3 of the Fifteenmile Group (Macdonald et al., 2010). δ^{13} C and 87 Sr/ 86 Sr data from the Little Dal Group are developed in Halverson (2006) and

Halverson et al. (2007a). A slightly stricter threshold for 87 Sr/ 86 Sr diagenesis is applied to the data included in our composite ([Sr]<250 ppm and Mn/Sr>0.15) than in the original work. A

correlative to the end of the Bitter Springs Stage, and a roughly constant sedimentation rate was

applied to the data between this age and the Islay Anomaly nadir, excluding an unconformity

basalt has been interpreted to conformably overlie the Little Dal Group (Aitken, 1981) and inferred to have erupted ca. 780 Ma based on geochemical similarity to mafic dikes and sills that intrude the Mackenzie Mountain Supergroup (Harlan et al., 2003). Given that the basalt has not been directly dated, there is some uncertainty associated with this interpretation. Nevertheless, correlating the δ^{13} C curve from the Little Dal Group to that of the Tambien Group by aligning the start and end of the Bitter Springs Stage and assuming constant sedimentation rate throughout the rest of the section suggests that the top of the Little Dal Group is ca. 780 Ma, 82 consistent with the inference of Harlan et al. (2003). $\delta^{13}\mathrm{C}$ and $^{87}\mathrm{Sr}/^{86}\mathrm{Sr}$ data and geochronology from the Coates Lake Group are developed in 84 Halverson (2006), Halverson et al. (2007a), and Rooney et al. (2014). A Re-Os isochron date on black shales of 732.2±3.9 Ma can be tied directly to this curve as it comes from strata recording the recovery from the nadir of the Islay δ^{13} C Anomaly. This date provides constraints on the temporal alignment of the curve. Given the uncertainty associated with the date, the correlation is further refined by aligning the nadir and recovery of the excursion with the Tambien Group data. 89 δ^{13} C and 87 Sr/ 86 Sr data from the Shaler Supergroup are developed in Asmerom et al. (1991), although $^{87}\mathrm{Sr}/^{86}\mathrm{Sr}$ data with Mn/Sr>3 and $\delta^{18}\mathrm{O}$ <-10\% are considered to be altered. Further $\delta^{13}\mathrm{C}$ data are developed in Jones et al. (2010). Age constraints on these strata are poor. However, the onset of the Bitter Springs Anomaly and the Islay Anomaly as well as other minor inflexions in the δ^{13} C curve are identifiable in the data. Furthermore, lithostratigraphic correlations between the Shaler Supergroup and the Mackenzie Mountains Supergroup can be made. Therefore, the age model for these data is developed based on the correlation of the δ^{13} C curve as well as the lithostratigraphy between these two supergroups, as in Jones et al. (2010).

8 Scotland

- $_{99}$ $\delta^{13}{\rm C}$ and $^{87}{\rm Sr}/^{86}{\rm Sr}$ data from the Dalradian Supergroup are developed in Sawaki et al. (2010).
- The carbonates from which these data are sourced unconformably underlie a glacial diamictite.

Brasier et al. (2000) argues that ${}^{87}\text{Sr}/{}^{86}\text{Sr}$ values from these carbonates are too low (<0.7065) to 101 be post-Sturtian, and therefore must be pre-Sturtian in age. Besides this argument, no direct 102 geochronological constraints exist for the Dalradian Supergroup. Therefore, the $\delta^{13}\mathrm{C}$ curve from 103 this group is correlated to that of the Tambien Group by aligning the nadir of the Islay Anomaly. 104

Russia 105

122

 δ^{13} C and 87 Sr/ 86 Sr data from Siberia come from multiple sources. 106

 $\delta^{13}\mathrm{C}$ and $^{87}\mathrm{Sr}/^{86}\mathrm{Sr}$ data from the Proterozoic carbonates of the UchurMaya and Turukhansk 107 regions of Siberia are developed in Bartley et al. (2001), with additional ⁸⁷Sr/⁸⁶Sr data from Cox 108 et al. (2016). All available ⁸⁷Sr/⁸⁶Sr data from Bartley et al. (2001) had [Sr]<500 ppm, and as a 109 result Mn/Sr>0.5 is the only threshold applied for diagenesis. Age constraints on these strata are 110 poor. Therefore, the age model applied to these data was based on lithostratigraphic correlation 111 to the Yenisey Ridge and Uchur Maya Region sections, which are temporally constrained to have 112 been deposited ca. 1100-1000 Ma based on geochronological constraints of varying robustness 113 (Gallet et al., 2012). δ^{13} C and 87 Sr/ 86 Sr data from the Karatau Group of the Urals are developed in Kuznetsov et al. 115 (2006), with additional ⁸⁷Sr/⁸⁶Sr data from (Cox et al., 2016). However, a slightly stricter 116 threshold for ⁸⁷Sr/⁸⁶Sr diagenesis was applied to the data included in our composite 117 ([Sr]<350 ppm and Mn/Sr>0.1). Correlation of microbiota across Siberia suggests that the group 118 is younger than ca. 1030 Ma (Kuznetsov et al., 2006). However, no other direct age control is 119 available for this group. Therefore, following Cox et al. (2016), the age model for the Karatau 120 Group data is constructed based on the correlation of one ca. 970 Ma Turukhansk Uplift 121 $^{87}\mathrm{Sr}/^{86}\mathrm{Sr}$ measurement to the start of the Karatau Group data.

123 Cryogenian Successions

Since Tambien Group chemostratigraphy is limited to the Tonian, our Cryogenian δ^{13} C and 87 Sr/ 86 Sr chemostratigraphic composite is a compilation of a number of other Cryogenian datasets. In general, correlations between datasets are made using absolute age constraints where possible - otherwise, characteristic negative δ^{13} C excursions (the ca. 659 Ma Rasthof Excursion, the ca. 655 Ma Taishir Anomaly, and the ca. 643 Ma Trezona Anomaly) are used to align datasets. Unless mentioned otherwise, the same criteria for diagenesis that were used for publication of the original datasets are applied here.

131 Mongolia

 δ^{13} C and 87 Sr/ 86 Sr data from Mongolia come from the Tsagaan-Olam Group and are developed in 132 Bold et al. (2016) and Brasier et al. (1996). Given that data from this group span the entirety of 133 the Cryogenian and into the Ediacaran, we use it as the backbone for our Cryogenian composite. For both datasets we apply a [Sr]<500 ppm and Mn/Sr>0.3 threshold for ⁸⁷Sr/⁸⁶Sr diagenesis. 135 The age model for these data follows that of Bold et al. (2016). A minimum age for the end of 136 the Sturtian Glaciation is constrained by the following: U-Pb ID-TIMS on zircon from a 137 tuffaceous bed overlying Sturtian diamicite in south China yields an age of 662.9 ± 4.3 Ma (Zhou 138 et al., 2004), a Re-Os isochron on black shales overlying Sturtian diamictite in northwest Canada 139 yields an age of 662.4±4.6 Ma (Rooney et al., 2014), and a Re-Os isochron on black shales 140 overlying Sturtian diamictite in Mongolia yields an age of 659.0±4.5 Ma (Rooney et al., 2015). A maximum age for the end of the Sturtian Glaciation is constrained by U-Pb ID-TIMS on zircon 142 from a tuff interbedded with Sturtian diamictite in Australia, which yields an age of 143 663.03±0.11 Ma (Cox et al., 2018). Therefore, for our Cryogenian composite, we set the age of the end of the Sturtian Glaciation (and therefore the age of the base of the Tsagaan-Olam Group) 145 to 660 Ma. 146

A maximum age for the start of the Marinoan Glaciation comes from a U-Pb SHRIMP age on 147 zircon from a tuff underlying Marinoan diamictite in south China of 654.5±3.8 Ma (Zhang et al., 148 2008). However, this tuff is separated from the Marinoan diamictite by a major disconformity, and so the age for the start of the Marinoan Glaciation is likely significantly younger than this 150 U-Pb SHRIMP age. Therefore, following Bold et al. (2016), we set the age for the start of the 151 Marinoan Glaciation in our composite to be 640 Ma. The end of the Marinoan Glaciation is tightly temporally constrained. Zircons from a volcanic 153 ash within and just above Marinoan diamictite in south China yielded U-Pb ID-TIMS dates of 154 635.5±0.6 and 635.2±0.6 Ma respectively (Condon et al., 2005). This constraint is consistent with 155 U-Pb ID-TIMS dates from zircon from tuffs within Marinoan diamictite of 635.5 ± 1.2 Ma in Namibia (Hoffmann et al., 2004), and 636.4±0.5 Ma in Tasmania, Australia (Calver et al., 2013). 157

158 Canada

 δ^{13} C, 87 Sr/ 86 Sr, and geochronological data from the Hay Creek Group are developed in Rooney et al. (2014). A Re-Os isochron on black shales from within this group yielded an age of 662.4 ± 4.6 Ma. The δ^{13} C and 87 Sr/ 86 Sr data correlate well with that from Mongolia.

162 Australia

otal (2010) for the composite. Amadeus Basin and Adelaide Rift Complex are taken from Swanson-Hysell et al. (2010) for the composite. Amadeus Basin data from the Bitter Springs Formation are older than the Sturtian diamictite of the Areyonga Formation which unconformably overlie it. Re-Os isochrons developed for black shales above the Areyonga Formation yielded ages of 643.0±2.4 and 657.2±5.4 Ma (Kendall et al., 2006). Data from the Etina and Trezona Formations of the Adelaide Rift Complex come from between Sturtian and Marinoan glacial deposits. It remains unclear whether or not the Taishir and Trezona Excursions are time equivalent. In this compilation, they are taken to be distinct following Bold et al. (2016) such that the Trezona Anomaly and

subsequent recovery occur temporally close to the initiation of the Marinoan Glaciation. The close temporal connection implied by this age model between the Trezona Anomaly and the initiation of the Marinoan Glaciation needs further work to be substantiated, although dropstones have been documented in the uppermost Trezona Formation (Rose et al., 2012). The data from the Adelaide Rift Complex shows that the δ^{13} C recovers from the nadir of the Trezona Anomaly over \sim 200 m such that recovery from the excursion occurred prior to local ice advance (Rose et al., 2012). While this does not necessarily mean that there is a substantial separation in time between the Marinoan Glaciation and the nadir of the Trezona Anomaly, it does suggest that the δ^{13} C values recovered from the negative anomaly to values near 0% prior to glaciation.

180 Namibia

δ¹³C and ⁸⁷Sr/⁸⁶Sr data from the Otavi Group are developed in Halverson et al. (2005) and
 Halverson et al. (2007a). We applied [Sr]<500 ppm and Mn/Sr>0.1 as the thresholds for
 ⁸⁷Sr/⁸⁶Sr alteration. Apart from the 635.5±1.2 Ma age from Hoffmann et al. (2004) constraining
 the end of the Marinoan Glaciation, no direct geochronological constraints exist for this data.
 Therefore, we align the Trezona Anomaly between the data from Australia and Namibia.

Geochronology

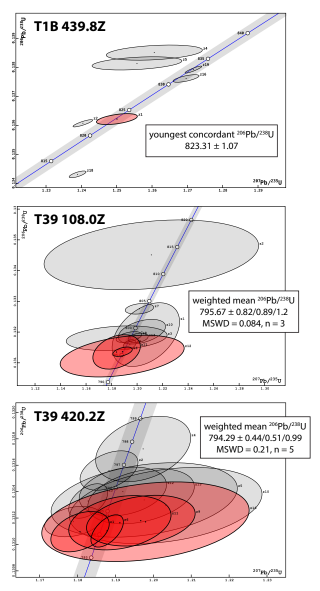


Figure DR1. Concordia diagrams for dates reported in this study. 2σ uncertainties are reported in the format $\pm X/Y/Z$, where X is the internal (analytical) uncertainty in the absence of all external or systematic errors, Y is the uncertainty incorporating the U-Pb tracer calibration error, and Z is the uncertainty including X and Y, as well as the uranium decay constant uncertainty; MSWD = mean square of weighted deviates; n = number of zircon analyses included in the calculated date.

Table DR1. U-Pb data for analyzed zircon from T1B-439.8Z.

	$\overset{\pm}{(2\sigma\%)}$	0.41	0.15	1.03	96.0	0.23	0.12	0.14	0.11
	$a,b,i = \frac{207 \text{Pb}}{206 \text{Pb}} $ (2	0.07	0.07	20.0	0.07	0.07	0.07	0.07	0.07
	$\begin{array}{ccc} & & & \\ + & & & \\ 2\sigma\%) & \overline{2}\overline{\alpha} \end{array}$	0.46	0.22 (1.08	1.01	0.29	0.20	0.19	0.19
	b,i $\frac{207 \text{Pb}}{235 \text{U}}$ (2	1.25 (1.24 (1.26	1.26	1.27	1.32 (1.24 (1.27 (
	$\begin{array}{c} \pm & \frac{26}{2} \\ (2\sigma\%) & \frac{2}{2} \end{array}$	0.14	0.09	0.19	0.16	0.10	0.09	0.09	60.0
ios	a,i $206 Pb$ $238 Pb$ $(238 Pb)$	0.14 (0.14 (0.14 (0.14 (0.14 (0.14 (0.13 (0.14 (
Isotopic Ratios	$ \begin{array}{c} h \\ 206 \text{Pb} \\ 204 \text{Pb} \\ \hline 20 \end{array} $	2348.88	6810.02	908.36	958.46	3610.31	10466.28	7480.36	15958.64 (
I	$\frac{g}{\text{Pb}*}$	37.96	110.86	15.23	16.67	61.23	181.54 1	130.41	266.55 1
	$\begin{array}{c} f \\ \text{Pb}_c \\ \text{(pg)} \end{array}$	0.61	0.62	98.0	0.80	0.22	0.18	0.32	0.18
Composition	e Pb* (pg)	23.13	68.32	13.16	13.33	13.75	32.33	42.34	47.38
Comp	d Th U	0.34	0.35	0.53	29.0	0.65	0.73	0.75	0.57
	c % disc.	0.10	-1.53	-3.65	-3.72	0.47	0.36	2.67	0.12
	corr.	0.51	0.85	0.33	0.37	69.0	0.92	0.73	06.0
	$(2\sigma) \pm$	8.55	3.20	21.58	20.17	4.89	2.59	2.95	2.36
	$^{a,b}_{207\mathrm{Pb}}_{\overline{206\mathrm{Pb}}}$	823.76	809.80	806.54	803.99	834.86	857.86	834.51	833.64
	(2σ)	2.61	1.26	6.10	5.72	1.64	1.17	1.08	1.06
	$\frac{b}{235}$	823.43 2.61	819.07	828.20	825.99	832.22	855.83	818.40	833.16
Ma)	(2σ)	1.07					0.73	0.67	0.69
Dates (Ma)	$a = \frac{206 \text{Pb}}{238 \text{U}}$	823.31 1.07	822.49 0.71	836.29 1.48	834.19 1.29	831.24 0.74	855.05	812.48	832.98
r	fraction	z1	z2	z4	z2	z16	z17	z18	z19

Colored rows indicate fractions included in the calculation of the reported sample age. Isotopic dates calculated using $\lambda 238 = 1.55125 \times 10^{-10}$ and $\lambda 235 = 9.8485 \times 10^{-10}$ (Jaffey et al., 1971).

^a Corrected for initial Th/U disequilibrium using radiogenic ²⁰⁸Pb and Th/U[magma] = 3.50000.

^b Corrected for initial Pa/U disequilibrium using initial fraction activity ratio [231 Pa]/[235 U] = 1.10000.

^c % discordance = $100 - (100 \times (^{206}$ Pb/ 238 U date) / (207 Pb/ 206 Pb date))

^d Th contents calculated from radiogenic 208 Pb and 230 Th-corrected 206 Pb/ 238 U date of the sample, assuming concordance between U-Pb Th-Pb systems.

e Total mass of radiogenic Pb.

f Total mass of common Pb.

 $[^]g$ Ratio of radiogenic Pb (including $^{208}\mathrm{Pb})$ to common Pb.

Measured ratio corrected for fractionation and spike contribution only. Measured ratios corrected for fractionation, tracer and blank.

Table DR2. U-Pb data for analyzed zircon from T39-108.0Z.

	$\pm \frac{(2\sigma\%)}{}$	2.00	6.77	3.28	0.60	0.90	0.79	0.39	1.44	1.49	3.92	0.63
	$^{a,b,i}_{207\mathrm{Pb}}_{\overline{206\mathrm{Pb}}}$	0.07	0.07	0.07	0.07	0.07	0.07	0.07	0.07	0.07	0.07	0.07
	$+ \frac{\pm}{(2\sigma\%)}$	2.19	6.94	3.34	0.65	0.93	0.84	0.42	1.54	1.57	4.09	0.65
	$\frac{b,i}{235}$	1.21	1.21	1.18	1.19	1.20	1.19	1.19	1.20	1.18	1.19	1.20
	$\pm \\ (2\sigma\%)$	0.79	0.84	0.30	0.12	0.13	0.13	0.12	0.30	0.43	0.52	0.12
Ratios	$\begin{array}{c} a,i\\ 206\text{Pb}\\ \overline{238\text{Pb}} \end{array}$	0.13	0.13	0.13	0.13	0.13	0.13	0.13	0.13	0.13	0.13	0.13
Isotopic Ratios	$h = \frac{206 \mathrm{Pb}}{204 \mathrm{Pb}}$	395.48	125.78	233.74	1297.81	813.30	993.36	2223.30	628.68	509.16	238.72	19.34 1218.11
	$\frac{g}{\text{Pb}^*}$	6.29	1.78	3.54	20.81	12.82	16.30	38.31	10.01	7.93	3.54	19.34
u	$f \\ \operatorname{Pb}_c \\ (\operatorname{pg})$	0.25	0.30	1.13	0.27	0.31	0.31	0.33	0.20	0.27	0.39	0.28
Composition	$_{\rm pb}^{e}$	1.59	0.54	4.01	5.66	4.03	5.09	12.47	2.01	2.11	1.37	5.40
Comp	$\frac{d}{d}$	0.57	0.56	0.51	0.47	0.43	0.58	0.73	0.50	0.44	0.42	0.43
	c % disc.	2.29	-3.75	-2.43	-0.16	-1.02	-0.34	0.46	1.40	-1.18	0.18	1.30
	corr.	0.41	0.26	0.25	0.49	0.27	0.46	0.36	0.42	0.33	0.38	0.31
	(2σ)	41.88	142.22	68.88	12.54	18.90	16.56	8.21	30.06	31.21	82.26	13.15
	$^{a,b}_{207\mathrm{Pb}}_{\overline{206\mathrm{Pb}}}$	817.26	783.94	778.96	794.14	795.22	795.41	800.40	810.47	786.05	795.97	808.48
	(2σ)	12.16	38.62	18.38	3.56	5.14	4.66	2.31	8.53	8.65	22.55	3.63
	$\frac{b}{235}$	798.88 5.90 803.74 12.16	805.73	793.16	795.30	801.43	797.63	797.93	802.34	793.15	795.18	
Ma)	(2σ)	5.90	6.45	2.28	0.87	0.98	1.01	0.88	2.28	3.19	3.87	0.88
Dates (Ma)	$ \begin{array}{c} a \\ 206 \text{Pb} \\ 1 \overline{238 \text{U}} (2\sigma) \end{array} $	798.88	813.64 6.45	798.22	795.71 0.87	803.66 0.98	798.42 1.01	797.04	799.42 2.28	795.67 3.19	794.90 3.87	798.28 0.88 800.97
	fraction	z1	z_2	z3	9z	72	82	6z	z10	z11	z12	z13

Colored rows indicate fractions included in the calculation of the reported sample age. Isotopic dates calculated using $\lambda 238 = 1.55125 \times 10^{-10}$ and $\lambda 235 = 9.8485 \times 10^{-10}$ (Jaffey et al., 1971). ^a Corrected for initial Th/U disequilibrium using radiogenic ²⁰⁸Pb and Th/U [magma] = 3.50000. ^b Corrected for initial Pa/U disequilibrium using initial fraction activity ratio [²³¹Pa]/[²³⁵U] = 1.10000.

 $^{^{}c}$ % discordance = 100 - $(100 \times (^{206}\text{Pb}/^{238}\text{U date}) / (^{207}\text{Pb}/^{206}\text{Pb date}))$ discordance = 100 - $(100 \times (^{206}\text{Pb}/^{238}\text{U date}) / (^{207}\text{Pb}/^{206}\text{Pb}/^{238}\text{U date})$ date of the sample, assuming concordance between U-Pb Th-Pb systems.

e Total mass of radiogenic Pb. f Total mass of common Pb.

 $[^]g$ Ratio of radiogenic Pb (including $^{208}{\rm Pb})$ to common Pb. h Measured ratio corrected for fractionation and spike contribution only.

i Measured ratios corrected for fractionation, tracer and blank.

Table DR3. U-Pb data for analyzed zircon from T39-420.2Z.

	$\pm \\ (2\sigma\%)$	0.62	09.0	0.40	1.00	2.00	0.32	1.17	2.06	1.10	0.94	1.13	0.62	2.23
	$a,b,i = \frac{207 \text{Pb}}{206 \text{Pb}}$ (2	0.07	0.07 0	0.07 0	0.07	0.07	0.07	0.07	0.07	0.07	0.07 0	0.07	0.07 0	0.07
	$\begin{array}{ccc} & & a \\ & \pm & 20 \\ (2\sigma\%) & \overline{20} \end{array}$	0.65 0	0.64 0	0.44 0	1.05 0	2.05 0	0.36 0	1.20 0	2.11 0	1.13 0	0.97	1.17 0	0.64 0	2.28 0
	$\frac{b,i}{207 \text{Pb}}$	1.18	1.19	1.18	1.20	1.20	1.19	1.20	1.20	1.19	1.19	1.20	1.18	1.20
	$\overset{\pm}{(2\sigma\%)}$	0.12	0.11	0.10	0.19	0.21	0.09	0.15	0.22	0.16	0.14	0.16	0.09	0.23
Ratios	$\frac{a,i}{206 \mathrm{Pb}}$ $\overline{\frac{238 \mathrm{Pb}}{238 \mathrm{Pb}}}$	0.13	0.13	0.13	0.13	0.13	0.13	0.13	0.13	0.13	0.13	0.13	0.13	0.13
Isotopic Ratios	$h = \frac{206 \text{Pb}}{204 \text{Pb}}$	1391.39	1415.45	2239.52	857.80	427.13	2801.05	731.99	416.84	09.922	903.55	760.42	1363.19	382.41
	$\frac{g}{\text{Pb}_c^*}$	24.28	25.79	40.01	15.02	7.15	50.28	14.19	7.70	13.43	17.46	13.04	27.31	6.40
	$f \\ \mathrm{Pb}_c \\ \mathrm{(pg)}$	0.49	0.59	0.59	0.91	1.71	0.87	0.62	1.38	0.40	0.61	0.40	0.40	1.14
Composition	e Pb* (pg)	11.98	15.22	23.62	13.69	12.24	43.52	8.79	10.65	5.45	10.71	5.25	10.90	7.30
Comp	$\frac{d}{U}$	0.69	0.87	0.76	0.73	0.64	0.77	1.20	1.06	69.0	1.16	99.0	1.30	99.0
	c % disc.	-1.18	-0.73	-1.33	0.82	1.90	0.42	2.12	3.19	86.0	0.51	1.65	-1.80	2.40
	corr.	0.38	0.42	0.48	0.34	0.31	0.55	0.28	0.30	0.28	0.27	0.35	0.29	0.29
	$(2\sigma) \pm$	12.98	12.69	8.35	20.92	41.75	89.9	24.38	42.92	23.04	19.65	23.61	13.14	46.60
	$a,b \\ \frac{207 \mathrm{Pb}}{206 \mathrm{Pb}}$	784.58	790.74	784.69	803.97	810.60	797.18	811.54	821.19	802.10	799.70	808.72	780.29	813.80
	(2σ)	3.59	3.55	2.40	5.80	11.36	1.99	6.63	11.70	6.26	5.34	6.49	3.54	12.64
		791.62	795.17	792.62	799.35	799.47	794.95	90.662	795.25 1.64 802.10 11	796.51	796.87	799.10	790.82	794.55 1.72 799.63 12.64
Ma)	$(2\sigma) +$	0.87	0.85	0.75	1.41	1.61	0.70	1.09	1.64	1.17	1.05	1.19	99.0	1.72
Dates (Ma)	$ \begin{array}{ccc} a & & b \\ \frac{206\text{Pb}}{238\text{U}} & \pm & \frac{207\text{Pb}}{235\text{U}} \end{array} $	794.12 0.87 791.62	796.76 0.85	795.44 0.75	797.69 1.41	795.49 1.61	794.15	794.59 1.09	795.25	794.51	795.85	795.65	794.55 0.66	794.55
'	fraction	z1	z2	z3	z4	z2	9z	6z	z10	z11	z12	z13	z14	z16

Colored rows indicate fractions included in the calculation of the reported sample age. Isotopic dates calculated using $\lambda 238 = 1.55125 \times 10^{-10}$ and $\lambda 235 = 9.8485 \times 10^{-10}$ (Jaffey et al., 1971).

a Corrected for initial Th/U disequilibrium using radiogenic ²⁰⁸Pb and Th/U[magma] = 3.50000.

b Corrected for initial Pa/U disequilibrium using initial fraction activity ratio [²³¹Pa]/[²³⁵U] = 1.10000.

c % discordance = $100 - (100 \times (^{206}\text{Pb})/^{238}\text{U})$ date) / ($^{207}\text{Pb}/^{206}\text{Pb}$ date))

d Th contents calculated from radiogenic ²⁰⁸Pb and $^{230}\text{Th-corrected}$ 206 pb/ ^{238}U date of the sample, assuming concordance between U-Pb Th-Pb systems.

e Total mass of radiogenic Pb.

 $[^]f$ Total mass of common Pb. g Ratio of radiogenic Pb (including 208 Pb) to common Pb. h Measured ratio corrected for fractionation and spike contribution only. i Measured ratios corrected for fractionation, tracer and blank.



Figure DR2. (A) Photograph of the lava flow T1b-439.8Z. (B) Photograph of the ignimbrite T39-108.0Z, with feldspar phenocrysts and fiammed lithic clasts. (C) Photograph of the 30 cm rhyolitic tuff T39-420.2Z, with normally graded lapilli at the base. Hammer points up section in all panels.

Diagenetic Considerations

188 Isotope Conglomerate Test

We compare δ^{13} C and δ^{18} O values of the carbonate clasts from within diamictite of the Negash 180 Formation of the Negash Syncline and Samre Fold-Thrust Belt. In general, the distribution of 190 clast δ^{13} C values is similar to that of the *in situ* Tambien Group carbonates (Fig. DR3A). 191 However, the filtering technique proposes that the stratigraphic distance of a sample to the closest 192 siliciclastic unit is a reasonable predictor for δ^{13} C alteration in in situ Tambien Group 193 carbonates. If such a scenario applied equally to the diamic tite clasts, we might expect the $\delta^{13}{\rm C}$ of the clasts to be pulled to more negative values relative to the in situ carbonates since most of 195 the samples in the *in situ* stratigraphy were extracted from carbonate horizons thicker than the 196 diamictite clasts, but such a distribution is not observed. 197 There are several potential explanations for this apparent inconsistency. First, as discussed in 198 the main text, samples that fall below the threshold d may or may not have had their carbon 199 isotopic composition altered. And so, even though the majority of sampled diamictite clasts have 200 a radius <0.2 m, the δ^{13} C of a significant proportion of these samples need not have been affected 201 by secondary alteration. Second, it is possible that carbon is better buffered in the diamictite 202 relative to the rest of the Tambien Group. Unless 100% of the diamictite's matrix was produced 203 via scouring and redeposition of pre-Snowball Earth siliciclastics with associated organic matter, 204 the matrix likely contains less low δ^{13} C organic carbon relative to the siliciclastic units of the 205 underlying Tambien Group, given that organic productivity was suppressed during the Snowball 206 Earth (Hoffman et al., 2017). The presence of extra-basinal clasts within the diamictite (see main 207 text) suggests that at least some of the protolith was sourced from distal bedrock, and thus the organic component of the diamictite's matrix was likely diluted relative to undisturbed Tambien 200 Group siliciclastics. Third, given that glacial erosion generates a bimodal sediment size 210 distribution (fine grains from scouring and larger clasts from plucking) from the same rock, the

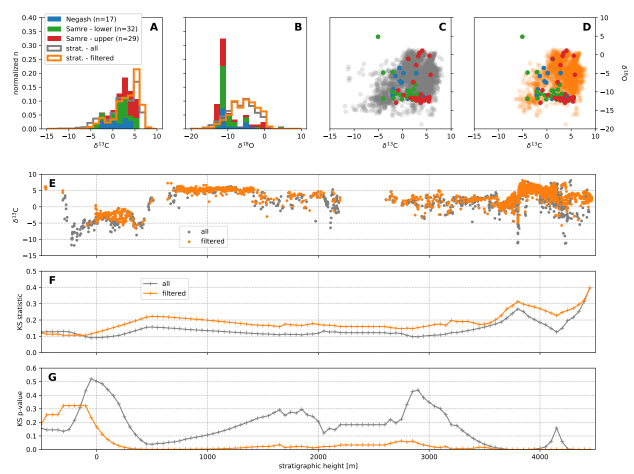


Figure DR3. (A) and (B) Histograms of δ^{13} C and δ^{18} O values of carbonate clasts within the diamictite of the Negash Formation of both the Negash Syncline and Samre Fold-Thrust Belt, compared to all in situ Tambien Group carbonate samples. Both filtered and unfiltered (all) versions of the in situ carbonate data are shown (see main text for a discussion of the filtering method). (C) and (D) Cross plots of δ^{13} C vs δ^{18} O for the clasts and in situ carbonate samples. (E) Filtered and unfiltered versions of the in situ carbonate δ^{13} C data against cumulative stratigraphic height. (F) Degree of correlation (as quantified by the Kolgomorov-Smirnov statistic) between the in situ carbonate δ^{13} C data with the carbonate clasts within the diamictite as samples below a given cumulative stratigraphic height (x-axis) are removed (i.e. the x-axis represents the depth of erosion). Low values suggest that the two datasets are drawn from the same distribution. See accompanying text for further details. (G) Kolgomorov-Smirnov statistic p-value. High values suggest that the two datasets are drawn from the same distribution.

sampled carbonate clasts in the diamictite are likely accompanied by fine carbonate sand from the
same rock. This relatively carbonate-rich diamictite matrix would help to buffer the carbon

isotopic composition of diamictite clasts against changes in δ^{13} C in a way that siliciclastic units 214 within the in situ Tambien Group stratigraphy would not be able to. Finally, it is possible that 215 our sampling of clasts from the diamictite is not representative of the bulk population. The total number of diamictite clasts sampled (n = 78) is substantially smaller than the total number of 217 samples from in situ Tambien Group carbonates (n = 3139). Furthermore, diamictite clasts were 218 only sampled from three discrete stratigraphic horizons, which may have been more carbonate 219 buffered relative to the rest of the diamictite. 220 δ^{18} O values of the diamictite clasts are distinctly different from the spread in values observed in 221 the in situ stratigraphy, and cluster at $\sim 12\%$ (Fig. DR3B). This difference suggests that, unlike 222 the carbon isotopic composition, the oxygen isotope composition of the carbonate clasts was 223 significantly more overprinted following deposition of the diamictite than that of the in situ 224 carbonates. This difference in post-depositional alteration likely arises from the fact that the 225 carbonate clasts in the diamictite are embedded within a predominantly siliciclastic matrix and 226 are therefore less carbonate buffered against altering fluids, whereas most of the samples in the in 227 situ stratigraphy were extracted from carbonate horizons thicker than the diamictite clasts and 228 are therefore more likely to be carbonate buffered. Since carbon is more rock-buffered against 220 altering fluids than oxygen, the δ^{13} C of the clasts are more likely to preserve primary values. Glacial erosion during the Sturtian Glaciation likely preferentially eroded the upper Tambien 231 Group in most places instead of eroding all the way to the base of the Tambien Group. To assess 232 how deep the bulk of glaciers eroded into Tambien Group stratigraphy, we divided the Tambien 233 Group chemostratigraphic composite data collected from the in situ stratigraphy (Fig. DR3E)

into several equal length (50 m) stratigraphic windows, and randomly selected the same number of samples from each of these windows. This Monte Carlo approach is necessary to avoid bias toward relatively heavily sampled intervals of the stratigraphy. We then quantified the similarity in distributions between the δ^{13} C of the Monte Carlo sampled *in situ* stratigraphy with that of the diamictite clasts using the two sample Kolmogorov-Smirnov (KS) statistic, which tests

whether two sets of samples are consistent with being drawn from the same distribution. Low KS 240 statistics and high p-values suggest that the two samples are drawn from the same distribution. 241 We then simulate shallower erosion by iteratively excluding the lowest of these stratigraphic windows and recalculating the KS statistic, moving up the stratigraphy (Fig. DR3F and G). In 243 general, we find that the two distributions are closest when the 'erosion height' is close to the 244 bottom of the Tambien Group (~0 m in Fig. DR3F and G) and near the middle of the Tambien Group (~2900 m). We also observe a distinct trough/peak near the top of the Tambien Group 246 $(\sim 4200 \text{ m})$, although the KS statistic/p-value is not as low/high as at the bottom or near the 247 middle of the Tambien Group. Furthermore, the filtered version of the in situ carbonate δ^{13} C 248 data (see main text) matches the clast data more poorly than all of the in situ carbonate δ^{13} C 249 data, likely as a result of similar post-depositional alteration mechanisms operating throughout 250 the entirety of the Tambien Group. Ultimately, this analysis illustrates the fact that a relatively 251 large proportion of the diamictite clasts have relatively low δ^{13} C (<0\%), and thus the clast δ^{13} C 252 distribution matches the in situ carbonate δ^{13} C distribution when the 'erosion height' is such that 253 it includes a high proportion of samples within the Bitter Springs Stage, the Islay Anomaly, 254 and/or carbonate samples that have likely experienced secondary alteration pulling them to lower δ^{13} C values. Observations of the facies of clasts within the diamictite suggest that they were 256 sourced predominantly from the Matheos and/or Mariam Bohkahko formations (see main text), 257 and thus an 'erosion height' of \sim 2900 m or \sim 4200 m would be consistent with these facies. The 258 KS statistic/p-value at these 'erosion heights' is low/high enough such that we cannot reject the 259 null hypothesis that the diamictite clasts and the in situ carbonate samples from above these 260 heights come from the same distribution. However, we note that erosion into the Mariam 261 Bohkahko/Matheos formations is not observed locally where the diamictite is deposited. This 262 observation requires that the clasts derive from carbonates time-equivalent to these formations 263 deposited elsewhere in the basin, or from carbonates deposited in another basin within an 264 Arabian-Nubian terrane.

Filtering Samples Adjacent to Siliciclastic Units

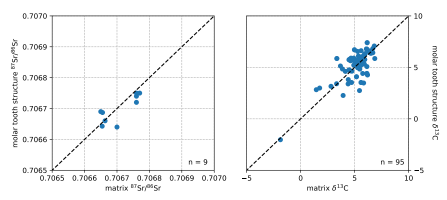


Figure DR4. Cross plots of carbonate matrix vs. adjacent molar tooth structure calcite $^{87}\mathrm{Sr}/^{86}\mathrm{Sr}$ and $\delta^{13}\mathrm{C}$ from Tambien Group samples that meet the filtering thresholds for alteration (see main text). Dashed black lines are the 1:1 lines - samples that fall on these lines have identical isotopic composition between the matrix and molar tooth structure carbonate. The isotopic composition of the matrix is similar to that of adjacent molar tooth structures, and no systematic offsets can be identified.

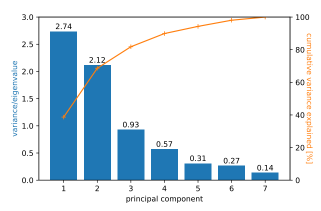


Figure DR5. Eigenvalues and cumulative variance explained for the 7 principal components in the principal components analysis (also known as a scree plot). Notably, the magnitude of the eigenvalues (and the percent variance explained) drops off sharply after the second principal component, indicating that the first two principal components capture the most significant sources of variance in our dataset.

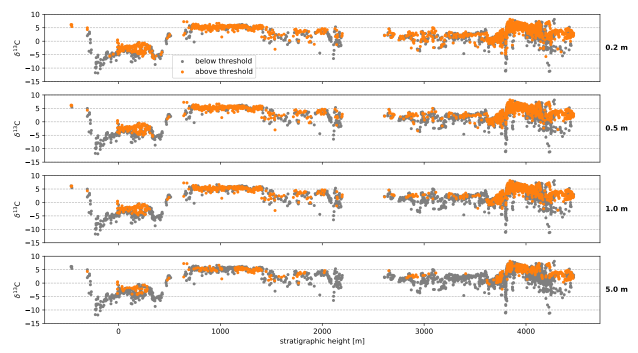


Figure DR6. Resulting composite chemostratigraphy of the Tambien Group as samples below a given d (shown on the right) are filtered out. Note that data that resolve the Islay Anomaly as well as the descent into and recovery out of the Bitter Springs Stage are mostly removed under the d=0.2 m threshold, and completely removed by d=0.5 m.

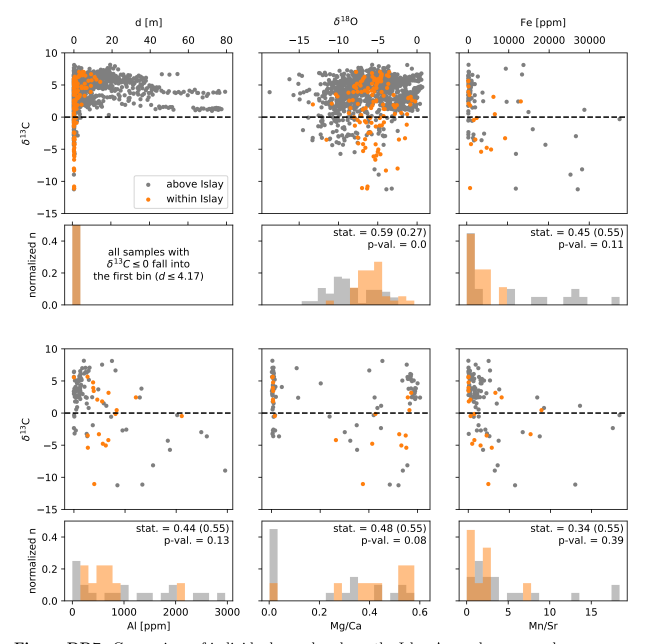


Figure DR7. Comparison of individual samples above the Islay Anomaly vs. samples within/adjacent to the Islay Anomaly. The normalized histograms under each scatter plot compare the distribution of samples with $\delta^{13}\mathrm{C} \leq 0\%$ only. The 'stat.' and 'p-val.' refer to the Kolmogorov-Smirnov statistic and p-value respectively, with the value within the parentheses showing the critical Kolmogorov-Smirnov statistic for rejecting the null hypothesis (see text below).

As per conventions in statistics, the following discussion will use the term 'unit' to refer to an 267 individual carbonate rock, and the term 'sample' to refer to a collection of 'units' from a 268 population. Figure DR7 compares geochemical data of units above the Islay Anomaly to units 269 within/adjacent to the Islay Anomaly. Units within the Islay Anomaly that record $\delta^{13}C \leq 0\%$ 270 appear to exhibit lower Fe, Al, and Mn/Sr than units that record $\delta^{13}C \leq 0\%$ above the Islav 271 Anomaly. This difference in distributions suggests that low δ^{13} C Islay Anomaly units have been less altered by the unbuffered fluids (see main text) than low $\delta^{13}\mathrm{C}$ post-Islav Anomaly units, and 273 thus provides support for the primary nature of the anomaly. To quantify this qualitative 274 interpretation of the data, we compare the distributions of units with low $\delta^{13}\mathrm{C}$ above the Islav 275 Anomaly to units with low δ^{13} C within the Islay Anomaly by computing the two-sample 276 Kolmogorov-Smirnov (KS) statistic. We also compute the critical KS statistic for rejecting the 277 null hypothesis, which is given by $1.36\sqrt{\frac{N_1+N_2}{N_1N_2}}$, where N_1 and N_2 are the number of items in the 278 two samples. If the computed KS statistic is above the critical KS statistic, or the p-value is 279 below 0.05, we can reject the null hypothesis at the 95\% confidence level that the two samples 280 come from the same distribution. We find that the KS test yields ambiguous results for the Fe, 281 Al, and Mn/Sr (Fig. DR7). For all three variables, the KS statistic is below the critical value, and 282 the p-value is above 0.05. These results indicate that we cannot declare at the 95\% confidence 283 level that the two samples come from different distributions - instead, the test indicates that the 284 samples may or may not come from the same distribution. However, the primary reason for this 285 ambiguity is the small number of units used in the test. There are only 20 units with $\delta^{13} C < 0\%$ 286 and element concentration data above the Islav Anomaly, and only 9 units with δ^{13} C <0% and 287 element concentration within the Islay Anomaly, which results in a high critical KS statistic and 288 thus a more 'difficult' test to achieve an unambiguous result in. Therefore, more element 289 concentration data is required in order for the KS test to quantitatively reject the hypothesis at 290 the 95% confidence level that units within the Islay Anomaly that record $\delta^{13}C < 0\%$ exhibit lower 291 Fe, Al, and Mn/Sr than units that record δ^{13} C $\leq 0\%$ 0 above the Islay Anomaly.

293 Sr Isotopes

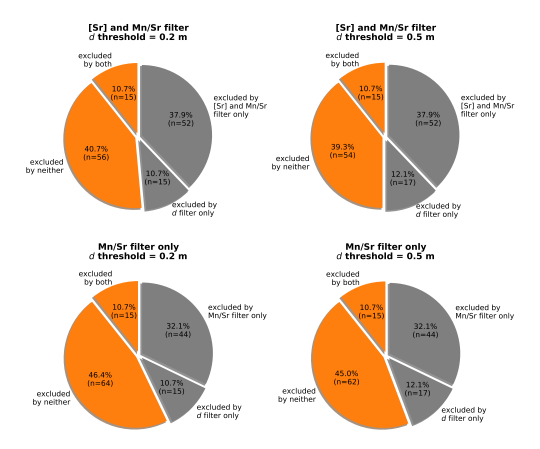


Figure DR8. Comparison of the application of the [Sr] and Mn/Sr filter and the filter based on distance to siliciclastics (d) to the 87 Sr/ 86 Sr data, for both d=0.2 m and d=0.5 m. Orange sectors denote classification agreement between the two filters, and grey sectors denote classification disagreement between the two filters. In the upper row, both the [Sr] and Mn/Sr thresholds are combined to filter samples. In the lower row, only the Mn/Sr threshold is used to filter samples.

Figure DR8 compares the application of the [Sr] and Mn/Sr filter and the filter based on
distance to siliciclastics (d) to the ⁸⁷Sr/⁸⁶Sr data. When both the [Sr] and Mn/Sr thresholds are
combined to filter samples (as in the main text), the [Sr] and Mn/Sr filter and the d filter only
agree on classification for around half of the samples. This lack of agreement results from the fact
that the principal components analysis used for the d filter (see main text) includes Mn/Sr, and
not [Sr], as a variable in the analysis, since [Sr] in carbonates can vary considerably based on

factors other than secondary alteration (e.g. calcite vs. aragonite, Husson et al., 2015). Therefore,
by using the Mn/Sr threshold only to filter samples, agreement between the Mn/Sr filter and the
d filter improves. Still, considerable disagreement between the two filters remain, which highlights
the limitations of the d filter as discussed in the main text. Namely, that in addition to filtering
out samples that have been altered, it is a rather blunt filter and may also filter out samples that
retain relatively pristine geochemistry.

87 Sr/ 86 Sr and the Drivers of Planetary Cooling

307 LIP Analysis

Table DR4 lists the large igneous provinces (LIPs) that were included in the LIP analysis in the
main text. The extent of each LIP was traced in QGIS to generate shapefiles, which were then
added to a paleogeographic model (Swanson-Hysell et al., 2018) to extract the paleolatitude of
the LIPs. We note that the LIP polygons were drawn to include the full areal extent of all dykes,
sills, and volcanics interpreted to be associated with each LIP, which may lead to an overestimate
of the true emplacement extents, since subsurface intrusions could extend over a broader area
than the surface volcanics. Where available, the paleogeographic model honors the paleomagnetic
poles listed in Table DR4.

316 Global Weathering Model

- The Python code used to develop the global weathering model can be found at:
- https://github.com/Swanson-Hysell-Group/2018_Tambien_Group. Table DR5 shows the variables
- and equations used in the global weathering model.

 \mathbf{Table} $\mathbf{DR4}$. Large igneous provinces included in the analysis conducted in the main text.

Name	Craton	Emplacement Age [Ma]	$\begin{array}{c} \textbf{Emplacement} \\ \textbf{Area} \ [\textbf{Mkm}^2] \end{array}$	Polygon Centroid Emplacement Latitude	Age Reference	Polygon Reference	Paleomagnetic Pole Reference
Mackenzie	Laurentia	1267	2.983	11.4	LeCheminant and Heaman (1989)	Ernst and Youbi (2017)	Buchan et al. (2000)
CSDG	Baltica	1255	0.145	-24	Söderlund et al. (2006)	Ernst and Youbi (2017)	Pisarevsky et al. (2014a)
Sudbury	Laurentia	1235	0.056	7.7	Dudas et al. (1994)	Shellnutt and MacRae (2012)	Palmer et al. (1977)
Marnda Moorn	SW. Australia	1210	0.598	65.2	Pisarevsky et al. (2014b)	Ernst and Youbi (2017)	Pisarevsky et al. (2014b)
Abitibi	Laurentia	1141	0.229	48.2	Krogh et al. (1987)	Ernst and Youbi (2017)	Ernst and Buchan (1993)
Umkondo	Kalahari	1109	1.846	8.0	Hanson et al. (2004)	Ernst and Youbi (2017)	Swanson-Hysell et al. (2015a)
Keweenawan	Laurentia	1109	0.414	41.9	Davis and Green (1997)	Ernst and Youbi (2017)	Swanson-Hysell et al. (2014)
SW Laurentia	Laurentia	1090	0.776	28.7	Weil et al. (2003)	Bright et al. (2014)	Weil et al. (2003)
Warakurna - 1	SW. + N. Australia	1070	0.757	38.6	Wingate et al. (2002)	Ernst and Youbi (2017)	Wingate et al. (2002)
Warakurna - 2	SW. + N. Australia	1070	0.444	41.9	Wingate et al. (2002)	Ernst and Youbi (2017)	Wingate et al. (2002)
Dashigou	N. China	925	0.663	3.9	Peng et al. (2011)	Pirajno (2013)	
Gangil-Mayumbia	Congo	920	0.333	-52.3	Tack et al. (2001)	Ernst and Youbi (2017)	
Willouran-Gairdner - 1	S. Australia	827	0.345	24.3	Wingate et al. (1998)	Ernst and Youbi (2017)	
Willouran-Gairdner - 2	S. Australia	827	0.171	29.3	Wingate et al. (1998)	Ernst and Youbi (2017)	
SWCUC - 1	S. China	821	0.914	68.6	Wang et al. (2016)	Ernst and Youbi (2017)	Li et al. (2004)
SWCUC - 2	S. China	821	0.411	65.6	Wang et al. (2016)	Ernst and Youbi (2017)	Li et al. (2004)
Gunbarrel - 1	Laurentia	780	0.21	1-	Harlan et al. (2003)	Ernst and Youbi (2017)	Park et al. (1989)
Gunbarrel - 2	Laurentia	780	0.34	4.2	Harlan et al. (2003)	Ernst and Youbi (2017)	Park et al. (1989)
Mundine Well	N. Australia	755	0.21	25.6	Wingate and Giddings (2000)	Ernst and Youbi (2017)	Wingate and Giddings (2000)
Irkutsk	Siberia	724	0.154	13.6	Ernst et al. (2016)	Ernst et al. (2016)	1
Franklin	Laurentia	720	2.231	-2.3	Denyszyn et al. (2009)	Ernst and Youbi (2017)	Denyszyn et al. (2009)

Notes: LIPs >0.1 Mkm² from the compilation in (Ernst et al., 2008) were included in the LIP analysis in the main text. Some LIPs are comprised of two separate polygons (denoted by 1 and 2).

Table DR5. Variables used in the global weathering model.

Term	Value/Equation	Note
Subaerial		
$[Ca]_{carb}$	302300 ppm	(1)
$[Mg]_{carb}$	47000 ppm	(1)
$[Sr]_{carb}$	610 ppm	(1)
$[Ca]_{rad}$	23750 ppm	(2)
$[Mg]_{rad}$	12800 ppm	(2)
$[Sr]_{rad}$	310 ppm	(2)
$[Ca]_{juv}$	71600 ppm	(3)
$[Mg]_{juv}$	45500 ppm	(3)
$[Sr]_{juv}$	465 ppm	(3)
Hydrothermal		
$H_{Mg-clays}$	$k\cdot [Mg]$	-
k		(4)
$H_{Ca-basalt}$	$lpha_{Mg/Ca} \cdot H_{Mg-clays}$	-
$\alpha_{Mg/Ca}$	1	(5)
$H_{nSr-basalt}$	$lpha_{Sr/Ca} \cdot H_{Ca-basalt}$	-
$\alpha_{Sr/Ca}$	0.0013	(6)
Precipitation		
$P_{Ca-carb}$	$W_{Mg-carb} + W_{Mg-rad} + W_{Mg-juv} - P_{Mg-carb} + W_{Ca-carb} + W_{Ca-rad} + W_{Ca-juv}$	(7)
$P_{Mg-carb}$	$5 \times 10^{10} \text{ mol/yr}$	(8)
$P_{Sr-carb}$	$(Sr/Ca)_{seawater} \cdot K_{Sr} \cdot P_{Ca-carb}$	-
K_{Sr}	0.2	(9)
$^{87}\mathrm{Sr}/^{86}\mathrm{Sr}$		
carbonate	0.70475	(10)
radiogenic	$BABI + (0.2783 \left(\frac{Rb}{Sr}\right)_m (9.3485 + BABI))(1 - e^{-2 \times 10^9 \lambda}) +$	(11)
	$10(0.2783 \left(\frac{Rb}{Sr}\right)_m (9.3485 + BABI))(1 - e^{-\lambda(t-2\times10^9)})$	
juvenile	$BABI + (0.2783 \left(\frac{Rb}{Sr}\right)_m (9.3485 + BABI))(1 - e^{-\lambda t})$	(11)

Notes:

- (1) from Turekian and Wedepohl (1961)
- (2) from Wedepohl (1995)
- (3) taking the mean of Turekian and Wedepohl (1961) and Taylor (1964)
- (4) flux of H₂O in hydrothermal systems, estimated to achieve desired initial steady state, then varied
- (5) assumes 1:1 stoichiometry between Mg and Ca during weathering of the ocean crust
- (6) from Maloof et al. (2010), calculated assuming 200 ppm Sr and 10 wt% CaO
- (7) calculated iteratively assuming carbonate minerals are the only Ca sink
- (8) estimated to achieve desired initial steady state
- (9) homogeneous distribution coefficient for Sr in calcite from Mucci and Morse (1983)
- (10) seawater has roughly constant ${}^{87}\mathrm{Sr}/{}^{86}\mathrm{Sr} \sim 2-1$ Ga (Shields and Veizer, 2002)
- (11) these equations account of ⁸⁷Rb decay, where BABI is the Basaltic Achondrite Best Initial ratio (⁸⁷Sr/⁸⁶Sr =0.69897) from Papanastassiou and Wasserburg (1968), $\left(\frac{Rb}{Sr}\right)_m$ is Rb/Sr of the mantle (0.025), λ is the ⁸⁷Rb decay constant, and t is time since the origin of the Earth.

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