- **1** Genesis of the world's largest rare earth element
- 2 deposit, Bayan Obo, China: protracted mineralization
- 3 evolution over ~1 billion years
- 4 Wenlei Song¹, Cheng Xu^{1*}, Martin P. Smith², Anton R. Chakhmouradian³, Marco
- 5 Brenna⁴, Jindřich Kynický⁵, Wei Chen⁶, Yueheng Yang⁷, Miao Deng¹, and Haiyan
- 6 Tang⁸
- 7 ¹Key Laboratory of Orogenic Belts and Crustal Evolution, School of Earth and Space
- 8 Sciences, Peking University, Beijing 100871, China
- 9 ²School of Environment and Technology, University of Brighton, Brighton BN24GJ, UK
- ³Department of Geological Sciences, University of Manitoba, Manitoba R3T2N2,
- 11 Canada.
- ⁴Department of Geology, University of Otago, Dunedin 9054, New Zealand
- ⁵Department of Geology and Pedology, Mendel University, Brno 361300, Czech Republic
- ⁶State Key Lab of Geological Processes and Mineral Resources, Wuhan 430074, China
- 15 University of Geosciences, China
- ⁷Institute of Geology and Geophysics, Chinese Academy of Sciences, Beijing 100029,
- 17 China
- 18 ⁸College of Earth Science, Jilin University, Changchun 130012, China
- 19 *E-mail: xucheng1999@pku.edu.cn
- 20 ABSTRACT
- 21 The unique, giant, REE deposit at Bayan Obo is the world's largest REE deposit. It
- 22 is geologically complex and its genesis is still debated. Here, we report in situ Th-Pb dating

23	and Nd isotope ratios for monazite and Sr isotope ratios for dolomite and apatite from fresh
24	drill-cores. The measured monazite ages (361-913Ma) and previously reported
25	whole-rock Sm-Nd data show a linear relationship with initial Nd isotope ratio, suggesting
26	a single-stage evolution from a Sm-Nd source that was formed before 913Ma. All
27	monazites show consistent $\epsilon Nd_{(1.3Ga)}$ values (0.3 ± 0.6) close to those of the adjacent 1.3Ga
28	carbonatite and mafic dikes. The primary dolomite and apatite show lower ⁸⁷ Sr/ ⁸⁶ Sr ratios
29	(0.7024–0.7030) than the recrystallized dolomite (0.7038–0.7097). The REE ores at Bayan
30	Obo are interpreted to have originally formed as products of ~1.3Ga carbonatitic
31	magmatism and undergone subsequent thermal perturbations induced by Sr-rich, but
32	REE-poor metamorphic fluids derived from nearby sedimentary rocks.

33 INTRODUCTION

34 The rare earth elements (REE) have become the focus of international attention because of their industrial importance to the development of "low carbon" energy and 35 36 transportation technologies, and because the global REE market is extremely sensitive to 37 geopolitically driven supply limitations (Hatch, 2012). The availability of REE for future 38 markets is a growing concern in the developed world because global demand for these 39 resources is expected to grow significantly (Verplanck and Hitzman, 2016). China, the 40 United States, Russia, Canada, Brazil, Australia, India and Malaysia account for the 41 majority of the world's REE reserves. China presently contains ~40% of the global REE 42 resources (Weng et al., 2015), concentrated primarily in the world's largest REE deposit at 43 Bayan Obo. This deposit has attracted inordinate attention from researchers (over 100 44 papers in peer-reviewed journals just in the past decade) because of its unparalleled 45 endowment in REE (>100Mt REE₂O₃, Weng et al., 2015). The genesis of the Bayan Obo

46	structure and its resources has been the subject of debate for over 50 years. There is no
47	consensus concerning either the age of mineralization (~1300 to ~400Ma; Yang et al.,
48	2017), or the number of mineralization stages (Chao, 1997). Genetic models proposed for
49	Bayan Obo range from sedimentary deposition (Chao, 1997), to metasomatic reworking of
50	metasedimentary marbles by carbonatitic (Smith et al., 1999) or subduction-derived fluids
51	(Yang et al., 2017), to igneous processes related to carbonatite emplacement (Le Bas et al.,
52	2007).

53 Monazite is one of the principal REE hosts in the Bayan Obo deposit. Here we 54 report integrated, in situ, high-precision Th-Pb ages and Nd isotope ratios of monazite 55 samples from an 1776 m long drill core section from the Bayan Obo deposit. The monazite 56 data were combined with in situ apatite and dolomite isotope analyses to show that the 57 Bayan Obo REE mineralization is of Mid-Mesoproterozoic age and of carbonatitic origin, 58 and shows no evidence of any significant REE contribution from external sources. This 59 Mid-Mesoproterozoic mineralization was subsequently modified by younger thermal 60 events.

61 GEOLOGY OF THE DEPOSIT AND SAMPLES

The Bayan Obo deposit is located at the northern margin of the North China Craton (NCC). The basement comprises the Archean Wutai Group (gneisses and migmatites) and Proterozoic Bayan Obo Group. The latter has been subdivided into nine lithological units, conventionally referred to as H1-H9 in ascending chronological order. The Bayan Obo Group is composed predominantly of meta-sandstones and slates, except for the H8 dolomite rock (Fig. DR1 in the GSA Data Repository¹). Volcanic rocks of trachytic, dacitic and rhyolitic composition, as well as mafic dikes, have been found within the H9 group

69 (Zhang et al., 2003; Yang et al., 2011). The metamorphic clastic sequences of the Bayan 70 Obo Group represent sedimentary units deposited within the Bayan Obo pericratonic rift. 71 The REE deposit is hosted by the H8 dolomite rock, which extends for 18km laterally at a 72 width of >1km, and occurs as a spindle-shaped stratiform body. About 100 carbonatite 73 dikes are found adjacent to the deposit, intruding the Bayan Obo Group metasediments 74 (Yang et al., 2011). The REE orebodies consist of disseminated, banded and massive ores, 75 most of which are associated with dolomite, silicates (in particular, alkali clinopyroxene, 76 amphibole and mica), apatite, fluorite and magnetite.

77 The studied drill core was extracted from the Eastern orebody, within the H8 unit, 78 and has a total length of 1776m. Compared to the H8 unit exposed at the surface, which 79 underwent extensive metasomatic alteration and deformation, and contains abundant 80 aegirine, riebeckite, phlogopite and late-stage fluorite-barite veins superposed over the 81 primary mineral assemblage, the drill samples are relatively fresh. The examined rocks are 82 composed predominantly of fine- to coarse-grained dolomite. Most of the dolomite is 83 euhedral to subhedral, and shows evidence of recrystallization with the development of 84 triple grain junctions. Some of the fine-grained, anhedral dolomites occur as a matrix to the 85 porphyritic dolomites (Fig. DR2), defining a primary, igneous texture. The studied drill 86 core shows significant variations in total light REE_2O_3 content, which locally reaches 87 5.8wt.% (Fig. DR3; Table DR1; for methods, see Data Repository). Textural observations 88 show that the early disseminated monazite was usually partially replaced and overgrown 89 by bastnäsite and apatite (Fig. 1a). Late monazite occurs as monominerallic veinlets, or is 90 associated with bastnäsite veinlets (Fig. 1b). Primary apatite was partially corroded and 91 overgrown by a rim of monazite (Fig. 1c). Recrystallized apatite occurs as veinlets and

92 clusters with bastnäsite (Fig. 1d). The textural evolution of REE minerals indicates
93 extensive metamorphic and metasomatic recrystallization (Smith et al., 1999).

94 **RESULTS**

95 The Th-Pb ages were combined with Nd isotopic ratio measurements done 96 independently and in situ on the same monazite grains, to calculate the initial 97 (¹⁴³Nd/¹⁴⁴Nd)_i ratios at the time of formation (Tables DR2,3). These monazites show 98 homogeneous internal textures (Fig. DR4), and have variable ages, ranging from $361 \pm$ 99 $6-913 \pm 15$ Ma. Their corresponding ε Nd(t) values fall between -6.9 and -18. However, all 100 samples show similar T_{CHUR}(Nd) model ages ranging from 1.56 to 1.67Ga, implying 101 derivation from the same source. The inferred ore-forming events at Bayan Obo have been 102 previously constrained chronologically on the basis of whole-rock and mineral 103 assemblages from surface samples, which show a wide range of ages with distinct 104 frequency peaks at ~1.3Ga and ~400Ma (Yang et al., 2017). However, the REE-rich 105 carbonatite dikes adjacent to the orebodies give a consistent Mid-Mesoproterozoic age of 106 ca. 1.3Ga (Fig. 2). The Sm-Nd isochron ages of volcanic rocks and mafic dikes in the 107 Bayan Obo deposit are also close to 1.3Ga. Figure 2 shows the measured monazite ages 108 plotted versus their corresponding $\epsilon Nd(t)$ values, and provides some of the previously 109 reported Sm-Nd isochron ages and $\varepsilon Nd(t)$ values for reference. Notably, the new and 110 published data show a good correlation, indicating a single-stage Nd isotopic evolution 111 from a single source. Late-stage, magma-derived melts or fluids could serve as a source of 112 REE, but this model would require that the later-emplaced magmas had very low Nd 113 isotopic ratios. This is clearly not the case: all reported Neoproterozoic to Carboniferous 114 igneous rocks in the northern NCC plot above the Nd isotopic evolution line for Bayan

Obo (Fig. 2). Therefore, we infer that late-stage changes in the REE mineralization defined by this evolution line were due to remobilization of these elements from the already existing orebody of Mid-Mesoproterozoic age. This interpretation is supported by textural evidence, such as metasomatic replacement of early monazite and apatite accompanied by the development of new REE minerals (Fig. 1). The trace element and isotope data described below further support the interpretation of the timing of primary REE mineralization.

122 The origin of the H8 dolomite rock hosting the deposit has been debated. The two "end-member" hypotheses are igneous crystallization from carbonatitic magma (Le Bas et 123 124 al., 2007) and sedimentary deposition (Chao, 1997). Our Nd isotope evolution line is 125 remarkably different from that characterizing typical sedimentary rocks from units H1-H3, in which the (¹⁴³Nd/¹⁴⁴Nd)_i ratio, calculated from 1.3Ga to 400Ma, is markedly lower than 126 127 in the REE minerals (Fig. 2). Both dolomite and apatite analyzed in this study show high Sr 128 contents (Table DR4), typical of carbonatitic minerals (Hornig-Kjarsgaard, 1998). 129 Different textural types of dolomite and apatite are characterized by distinct REE 130 distribution patterns (Fig. 3). The primary dolomite shows relatively low REE content 131 (La<10ppm) and a flat distribution pattern with $(La/Yb)_{cn} = 1-5$. The recrystallized 132 dolomite is characterized by a much more varied and higher REE content (La = 133 16–109ppm) and stronger enrichment in light REE, with $(La/Yb)_{cn} = 8-32$. The two 134 generations also differ in their key REE ratios, i.e., the primary variety has higher Eu/Eu* 135 and Y/Ho values relative to the recrystallized dolomite. Early disseminated apatite is 136 significantly enriched in REE (La>1400ppm) and shows a higher Eu/Eu* but lower Y/Ho 137 values than the paragenetically later generation confined to the veinlets (Fig. DR5).

138 The textural relations and extreme isotopic variability of the Bayan Obo monazite 139 imply that it is a product of dissolution-reprecipitation processes and that its Th-Pb budget 140 was modified over an extended period of time. The extended history of metasomatism at 141 Bayan Obo is recorded in the monazite ages, spanning from 361 to 913Ma. The primary 142 REE mineralization must have formed earlier than 913Ma and may have occurred ca. 143 1.3Ga, as indicated by the Sm-Nd isochron ages of the ore-bearing H8 unit and spatially 144 associated REE-enriched carbonatites (Fig. 2). These previously studied samples have 145 initial Nd isotopic ratios similar to those of the monazite studied in the present work if 146 calculated for 1.3Ga ($\epsilon Nd_{1.3Ga} = 0.3 \pm 0.6$), implying a common mantle source. In situ Sr isotopic analysis of the primary dolomite and apatite also gave low ⁸⁷Sr/⁸⁶Sr values (Table 147 148 DR5, 0.7024–0.7030), which are far less radiogenic than typical marine carbonates and 149 further support a non-sedimentary origin (Fig. 3). However, the late generations of 150 recrystallized dolomite have variable and high Sr isotopic compositions (0.7038–0.7097). 151 The present-day Sr isotopic ratios measured in the primary dolomite and apatite are considered to approximate the initial ⁸⁷Sr/⁸⁶Sr values because these minerals contain high 152 153 levels of Sr, but negligible Rb and thus, are characterized by very low Rb/Sr ratios. Similar 154 initial Sr isotopes (0.7029–0.7030) have been reported from ~1.3Ga carbonatite dikes 155 without contamination by feldspar from the wall rocks in Bayan Obo (Le Bas et al., 2007).

156

DISCUSSION AND CONCLUSION

157 A newly reported zircon age (1301 \pm 12Ma) on REE-rich carbonatites at Bayan 158 Obo supports the model of Mid-Mesoproterozoic primary mineralization (Zhang et al., 159 2017). The ⁸⁷Sr/⁸⁶Sr ratios of primary dolomite and apatite are close to the Bulk Earth 160 value (0.7029) at 1.3Ga. The ϵ Nd_{1.3Ga} value of monazite is also close to the Chondritic

161 Uniform Reservoir value, implying that the ore-bearing dolomite rock may be derived 162 from a primary, non-depleted mantle source, perhaps residing within the less-accessible 163 parts of the mantle, or in the depleted lithospheric mantle modified by old subducted 164 materials. Experiments show that primary carbonatitic melts derived from carbonated 165 peridotites contain relatively low REE abundances (Foley et al., 2009) and must 166 consequently undergo extensive fractionation in the crust to produce the level of REE 167 enrichment. Initially low REE concentrations in carbonatitic magmas are typically 168 dispersed among such major rock-forming constituents as calcite and apatite 169 (Hornig-Kjarsgaard, 1998), preventing the development of REE mineralization. In 170 contrast, primary carbonatitic magmas can be derived by partial melting of carbonated 171 eclogites (Thomson et al., 2016). In the Trans-North China Orogen of the NCC (i.e., 172 ~300km southeast of Bayan Obo), several occurrences of Paleoproterozoic carbonatite 173 dikes were found to contain high-pressure eclogite xenoliths of recycled crustal origin (Xu 174 et al., 2017a). This discovery provides unambiguous evidence that subducted material is 175 present in the mantle beneath the northern NCC. Seismic imaging of the NCC across the 176 Trans-North China Orogen (Zheng et al., 2009) provides strong support to 177 Paleoproterozoic (1.9–2.1Ga) westward subduction beneath the Western Block of the 178 Craton at the time when it is inferred to have been part of the Columbia supercontinent. 179 Numerous diabase dikes emplaced in the northern NCC (Fig. DR1) are considered to be 180 related to the Mid-Mesoproterozoic breakup of Columbia (see Zhang et al., 2017). The 181 mafic dikes in the northern NCC share geochemical characteristics of both ocean-island 182 basalts and island-arc volcanic rocks, as can be seen in tectonic-setting-based 183 discrimination diagrams (Fig. DR6). Geochemically, these dikes resemble basaltic

184 magmatism whose mantle source was influenced by previous subduction events, and are 185 distinctly different from purely intraplate volcanic rocks in the NCC. In particular, the 186 Bayan Obo mafic dikes and volcanic rocks in unit H9 show consistent negative Nb, Ta and 187 Ti anomalies (normalized to the primary mantle values), and are compositionally similar to 188 arc basalts (Fig. DR7). Crustal contamination as a source of these geochemical deviations 189 can be ruled out because the mafic dikes in the northern NCC show a consistent Nd 190 isotopic signature ($\varepsilon Nd_{1.3Ga} = -0.5 - 1.9$; Yang et al., 2011). Therefore, we consider that 191 subduction modification pre-conditioned the mantle source to generate the Bayan Obo

192

carbonatite REE deposit.

193 Our mineralogical and geochemical results suggest that the primary REE 194 mineralization at Bayan Obo was modified by externally derived fluids, which involved 195 the development of superimposed mineralization and recrystallization of the primary 196 minerals. The metasomatic fluids contain a crustal component, as indicated by a negative 197 shift in Eu/Eu* value and higher ⁸⁷Sr/⁸⁶Sr ratios in the recrystallized dolomite and apatite 198 (Fig. 3, Fig. DR5). The Y/Ho trends exhibited by dolomite and apatite are mutually 199 complementary, indicating structural controls over Y versus Ho partitioning between 200 crystals and the fluids, whereas both minerals show depletion in Eu with recrystallization. 201 The C-O isotope data from the deposit also show a large variation and plot between mantle 202 and sediment fields (Yang et al., 2017). Moreover, the fluids must have been poor in REE, but rich in Sr to explain the positive ⁸⁷Sr/⁸⁶Sr excursion. Caledonian subducted 203 204 slab-derived fluids, as proposed by Yang et al. (2017), are unlikely to be responsible for the 205 observed geochemical trends, because such fluids would be expected to have radiogenic 206 Nd isotopes (Xu et al., 2017b). Their interaction with the H8 unit would inevitably modify

207 its Nd isotopic signature, causing deviation of monazite ε Nd(t) values from the continuous 208 evolutionary line shown in Figure 2. Moreover, the Neoproterozoic to Carboniferous 209 magmatism in the northern NCC could not provide fluids sufficiently depleted in 210 radiogenic Nd to explain the low negative $\varepsilon Nd(t)$ values of young Bayan Obo monazite 211 generations. The sedimentary rocks present in the sequence at Bayan Obo are a viable 212 alternative source of metasomatizing fluids. These rocks show elevated Sr levels (up to 213 radiogenic 580ppm) coupled with a strongly Sr isotopic signature 214 $({}^{87}Sr/{}^{86}Sr_{(985Ma)}=0.7147)$, but are poor in REE (Zhang et al., 2003), and may have 215 contributed this signature to the post-ore metasomatic fluids involved in dolomite and 216 apatite recrystallization. These sedimentary rocks underwent metamorphism to various 217 degrees (from greenschist to low amphibolites facies conditions) and could serve as a 218 persistent fluid source responsible for textural and geochemical changes in the H8 219 dolomite rock.

220 In conclusion, our interpretation of the isotopic and trace element characteristics of 221 monazite, dolomite and apatite support the derivation of primary REE from a 222 Mid-Mesoproterozoic carbonatitic source. The apparent discrepancy in the behavior of Sr 223 and Nd isotopes highlights the importance of multi-systemic approach to geologically 224 complex mineral deposits, and reflects a protracted history of metasomatism induced by 225 Sr-rich, REE-poor fluids. In a similar case, the Nolans Bore REE deposit in Australia has 226 been found to have experienced multiple episodes of recrystallization/internal reworking 227 over a period of at least 1 billion years after primary ore formation (Schoneveld et al., 228 2015). The resetting of the ore system may be common in most REE deposits, and may be 229 critical in the high grade of some deposits. Interpreting geochronological results from the

230	REE orebodies should be undertaken with caution. However, with the utilization of
231	petrographic constrained analyses, recrystallization processes may be related to regional
232	tectonic events, and therefore complex REE orebodies could be used to unravel their
233	tectonic evolution.
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238	REFERENCES CITED
239	Chao, E.C.T., 1997, The sedimentary carbonate-hosted giant Bayan Obo REE-Fe-Nb ore
240	deposit of Inner Mongolia, China: A corner stone example for giant polymetallic ore
241	deposits of hydrothermal origin: U.S. Geological Survey Bulletin, v. 2143, p. 1-65.
242	Foley, S.F., Yaxley, G.M., Rosenthal, A., Buhre, S., Kiseeva, E.S., Rapp, R.P., and Jacob,
243	D.E., 2009, The composition of near-solidus melts of peridotite in the presence of CO_2
244	and H ₂ O between 40 and 60 kbar: Lithos, v. 112, p. 274–283,
245	https://doi.org/10.1016/j.lithos.2009.03.020.
246	Hatch, G.P., 2012, Dynamics in the global market for rare earths: Elements, v. 8,
247	p. 341-346, https://doi.org/10.2113/gselements.8.5.341.
248	Hornig-Kjarsgaard, I., 1998, Rare earth elements in sovitic carbonatites and their mineral
249	phases: Journal of Petrology, v. 39, p. 2105–2121,
250	https://doi.org/10.1093/petrology/39.11.2105.
251	Le Bas, M.J., Yang, X.M., Taylor, R.N., Spiro, B., Milton, J.A., and Zhang, P.S., 2007,
252	New evidence from a calcite-dolomite carbonatite dike for the magmatic origin of the

- 253 massive Bayan Obo ore-bearing dolomite marble, Inner Mongolia, China: Mineralogy
- and Petrology, v. 90, p. 223–248, https://doi.org/10.1007/s00710-006-0177-x.
- 255 Schoneveld, L., Spandler, C., and Hussey, K., 2015, Genesis of the central zone of the
- 256 Nolans Bore rare earth element deposit, Northern Territory, Australia: Contributions
- to Mineralogy and Petrology, v. 170, p. 11,
- 258 https://doi.org/10.1007/s00410-015-1168-x.
- 259 Shao, J.A., Hong, D.W., and Zhang, L.Q., 2002, Genesis of Sr-Nd isotopic characteristics
- 260 of igneous rocks in Inner Mongolia: Geological Bulletin of China, v. 21, p. 817–822.
- 261 Smith, M.P., Henderson, P., and Zhang, P., 1999, Reaction relationships in the Bayan Obo
- 262 Fe-REE-Nb deposit, Inner Mongolia, China: Implications for the relative stability of
- 263 rare-earth element phosphates and fluorocarbonates: Contributions to Mineralogy and
- 264 Petrology, v. 134, p. 294–310, https://doi.org/10.1007/s004100050485.
- 265 Thomson, A.R., Walter, M.J., Kohn, S.C., and Brooker, R.A., 2016, Slab melting as a
- barrier to deep carbon subduction: Nature, v. 529, p. 76–79,
- 267 https://doi.org/10.1038/nature16174.
- 268 Verplanck, P.L., and Hitzman, M.W., 2016, Introduction: Rare earth and critical elements
- 269 in ore deposits: Economic Geology and the Bulletin of the Society of Economic
- 270 Geologists, v. 18, p. 1–4.
- 271 Weng, Z., Jowitt, S.M., Mudd, G.M., and Haque, N., 2015, A detailed assessment of global
- rare earth element resources: Opportunities and challenges: Economic Geology and
- the Bulletin of the Society of Economic Geologists, v. 110, p. 1925–1952,
- 274 https://doi.org/10.2113/econgeo.110.8.1925.

- 275 Xu, C., Kynicky, J., Smith, M.P., Kopriva, A., Brtnicky, M., Urubek, T., Yang, Y.H.,
- 276 Zhao, Z., He, C., and Song, W., 2017b, Origin of heavy rare earth mineralization in
- 277 South China: Nature Communications, v. 8, p. 14598,
- 278 https://doi.org/10.1038/ncomms14598.
- 279 Xu, C., Kynicky, J., Tao, R.B., Liu, X., Zhang, L.F., Pohanka, M., Song, W.L., and Fei,
- 280 Y.W., 2017a, Recovery of an oxidized majorite inclusion from Earth's deep
- asthenosphere: Science Advances, v. 3, p. e1601589,
- 282 https://doi.org/10.1126/sciadv.1601589.
- 283 Yang, K.F., Fan, H.R., Santosh, M., Hu, F.F., and Wang, K.Y., 2011, Mesoproterozoic
- 284 mafic and carbonatitic dikes from the northern margin of the North China Craton:
- 285 Implications for the final breakup of Columbia supercontinent: Tectonophysics,

286 v. 498, p. 1–10, https://doi.org/10.1016/j.tecto.2010.11.015.

- 287 Yang, X., Lai, X., Pirajno, F., Liu, Y., Ling, M., and Sun, W., 2017, Genesis of the Bayan
- 288 Obo Fe-REE-Nb formation in Inner Mongolia, North China Craton: A perspective
- review: Precambrian Research, v. 288, p. 39–71,
- 290 http://doi.org/10.1016/j.precamres.2016.11.008.
- 291 Zhang, S.H., Zhao, Y., and Liu, Y., 2017, A precise zircon Th-Pb age of carbonatite sills
- from the world's largest Bayan Obo deposit: Implications for timing and genesis of
- 293 REE-Nb mineralization: Precambrian Research, v. 291, p. 202–219,
- 294 http://dx.doi.org/10.1016/j.precamres.2017.01.024.
- Zhang, Z.Q., Yuan, Z.X., Tang, S.H., Bai, G., and Wang, J.H., 2003, Age and
 Geochemistry of the Bayan Obo Ore Deposit: Beijing, Geological Publishing House,
- 297 222 p.

- Zheng, T., Zhao, L., and Zhu, R., 2009, New evidence from seismic imaging for
 subduction during assembly of the North China craton: Geology, v. 37, p. 395–398,
 https://doi.org/10.1130/G25600A.1.
- 301 Zhu, X.K., Sun, J., and Pan, C., 2015, Sm-Nd isotopic constraints on rare-earth
- 302 mineralization in the Bayan Obo ore deposit, Inner Mongolia, China: Ore Geology
- 303 Reviews, v. 64, p. 543–553, https://doi.org/10.1016/j.oregeorev.2014.05.015.

304

305 FIGURE CAPTIONS

306

307 Figure 1. Back-scattered-electron images showing the characteristic mineral assemblages

308 and textures observed in the Bayan Obo drill core. Dol, dolomite; Mnz, monazite; Ap,

309 apatite; Bas, bastnäsite; Mag, magnetite.

310

311 Figure 2. Trend of Nd isotopic evolution of Bayan Obo monazite with age compared to 312 other relevant isotopic data. The dashed line is the trend line of the monazite and can be 313 extended to 1.3Ga, where the $\varepsilon Nd(t)$ value is close to zero and similar to the $\varepsilon Nd(t)$ values 314 of the H8 unit (Zhang et al., 2003; Zhu et al., 2015; Yang et al., 2017), mafic dikes (Yang 315 et al., 2011) and volcanic rocks (Zhang et al., 2003) within the H9 unit, and carbonatite 316 dikes adjacent to the deposit (Zhang et al., 2003; Le Bas et al., 2007; Yang et al., 2011). 317 Data for sedimentary rocks (Zhang et al., 2003) from Bayan Obo and igneous rocks (Shao 318 et al., 2002) from the northern NCC are plotted.

- 320 Figure 3. Chondrite-normalized REE patterns and ⁸⁷Sr/⁸⁶Sr ratios of dolomite and apatite
- 321 from the drill core. Average REE abundances were used with error bars of one standard
- 322 deviation.
- 323
- ¹GSA Data Repository item 2018xxx, methods, figures, and tables, is available online at
- 325 http://www.geosociety.org/datarepository/2018/ or on request from
- 326 editing@geosociety.org.



Figure 1





GSA Data Repository 2018XXX 1 W. Song, C. Xu^{*}, M.P. Smith, A.R. Chakhmouradian, M. Brenna, J. Kynický, W. 2 Chen, Y. Yang, M. Deng, and H. Tang, 2018, Genesis of the world's largest 3 rare earth element deposit, Bayan Obo, China: protracted mineralization 4 evolution over ~1 billion years: Geology 5 6 Analytical Methods 7 Whole-rock analysis 8 9 Major and rare earth element abundances in drill core samples were determined by a Spectro Blue Sop inductively coupled plasma optical emission 10 spectrometer (ICP-OES) at the School of Earth and Space Sciences, Peking 11 12 University. The analytical precision is $\pm 5\%$ for all the elements. 13 Element mapping 14 Compositional X-ray maps of monazites were obtained with an Oxford 15 INCA X-MAX50 250+, energy dispersive X-ray spectrometer installed on a FEI 16

Quanta-650FEG scanning electron microscope, at the School of Earth and Space Sciences, Peking University. The backscattered electron and energy-dispersive X-ray data acquired from the samples were combined and processed automatically to generate the most sensitive X-ray mapping. The sample, coated with a conductive Cr layer (10 nm thickness) to prevent sample charging, was analyzed in a high-vacuum mode at standard operating

1 conditions (accelerating voltage of 20 kV, probe current 5 nA).

2

3 Monazite dating

Monazite grains ranging from 50 to 100 µm across were collected from the 4 drill core using conventional heavy liquid separation techniques. 5 Back-scattered electron images show that the crystals are compositionally 6 homogeneous and free of inclusions. The Th-Pb dating of monazite was 7 performed using a Cameca IMS-1280 secondary-ion mass-spectrometer 8 (SIMS) at the Institute of Geology and Geophysics (IGG), Chinese Academy of 9 Sciences (CAS). During the analysis, an O²⁻ primary ion beam was 10 accelerated at 13 kV with an intensity of ca. 2-3 nA. Aperture illumination mode 11 (Kohler illumination) was used with a 200-µm primary beam mass filter 12 aperture to produce even sputtering over the entire analyzed area. The 13 ellipsoidal spot was about 20 × 30 µm in size. Positive secondary ions were 14 extracted with a 10 kV potential. Monazite 44069 was used as a standard. A 15 ²⁰⁷Pb-based common Pb correction method was used. Further instrumental 16 and analytical details can be found in Li et al. (2013). 17

18

19 Trace element analysis

In-situ laser-ablation analyses of dolomite and apatite in thin sections were
 performed by inductively-coupled-plasma mass-spectrometry (ICP-MS) at the
 School of Earth and Space Sciences, Peking University, using a COMPexPro

102 excimer laser and an Agilent7500ce/cs mass-spectrometer. The diameter 1 of an ablation spot was 32 µm. The NIST 610 glass was used as a calibration 2 standard, and the Ca content measured by electron-microprobe analysis, as 3 an internal standard. Signal intensity for indicative trace elements was 4 monitored online during the analysis to ensure that the ablation spot was 5 confined to the area of interest and did not sample other mineral phases or 6 inclusions. The analytical error was estimated to be better than 5% at the ppm 7 level. 8

9

10 Nd-Sr isotopic analysis

The Nd isotopic composition of monazite was measured in situ by 11 12 multi-collector ICP-MS using a Thermo-Finnigan Neptune instrument coupled to a 193-nm ArF excimer laser-ablation system at the IGG, CAS. The diameter 13 of a laser spot and frequency were adjusted to between 10-24 µm and 4-10 Hz, 14 respectively, depending on the Nd concentration in the sample. Each spot 15 analysis consisted of approximately 60 s of signal acquisition. More detailed 16 information on the in-situ Nd isotopic analysis employed in the present work is 17 available in Yang et al. (2008). The Sr isotopic compositions of dolomite and 18 apatite were measured in situ by laser-ablation multicollector ICP-MS 19 (Resonics + Nu instruments) at the State Key Laboratory of Geological 20 Processes and Mineral Resources, China University of Geosciences (Wuhan). 21 The isotopic ratios were quantified in a static multicollector mode at low 22

resolution, using seven Faraday collectors and a mass configuration array
from ⁸²Kr to ⁸⁸Sr to monitor variations in Kr, Rb and Sr signals. The detailed
analytical procedure and data-reduction strategy are described in Tong et al.
(2015).

5

6 Figure captions for Data Repository

Figure DR1. Geological sketch map of the Bayan Obo deposit. a: The
locations of Bayan Obo and ~1.3 Ga mafic dikes in the northern North China
Craton (NCC; Yang et al., 2011; Zhang et al., 2012; Wang et al., 2014); b: The
locations of drill core, carbonatite and mafic dikes in Bayan Obo.

Figure DR2. Drill core samples and their photomicrographs. a, b: Drill cores 11 12 collected from the Eastern Orebody at a depth of 1776 m. c: Photomicrograph of dolomite (Dol) showing re-crystallization texture with the development of 13 elongation and preferred orientation, and triple junctions between crystals. 14 Rare earth minerals (REM) of monazite and REE-fluorocarbonates occur as 15 veinlets. d: Photomicrograph of primary fine-grained dolomite as a matrix to 16 porphyritic dolomite. Disseminated REM is associated with fluorite (FI). 17 Figure DR3. Plot showing the total light REE₂O₃ contents (La-Sm) of the drill 18

core samples with vertical depth.

²⁰ Figure DR4. X-ray compositional maps of representative monazite grains.

Figure DR5. Compositional variation of primary and recrystallized dolomite (Dol) and apatite (Ap) from the Bayan Obo drill cores. a: La/Yb_{cn} (cn -

1 chondrite normalized) vs. total REE; b: Y/Ho vs. Eu/Eu^{*} (Eu anomaly).

Figure DR6. Revised tectonic discrimination diagrams for mafic dikes from the 2 northern NCC. Data of the Wulahada and Wudalianchi volcanic fields in NCC 3 are plotted as reference for cases of basaltic magmatism with the source 4 influenced by previous subduction events (Wulahada at 142 Ma; Zhang et al., 5 2003) and for purely intraplate (OIB-like) volcanism from an enriched source 6 (Wudalianchi 10 Ma to Zhang al., 1995). 7 at recent; et The Mid-Mesoproterozoic mafic dikes (Zhang et al., 2012; Wang et al., 2014) in 8 northern NCC plot in both IAB and OIB, and Bayan Obo data (Wang et al., 9 2003; Yang et al., 2011) mostly in the IAB field, indicating influence of 10 subduction derived fluids in their mantle source. The tectonic discrimination 11 12 diagrams are from Vermeesch (2006). OIB, Ocean Island Basalt; IAB, Island Arc Basalt; MORB, Middle Ocean Ridge Basalt. 13

Figure DR7. Primitive mantle normalized diagram for mafic dikes from the 14 15 northern NCC. Data of OIB is from Sun and McDonough (1989), IAB based on average compositions reported by Jakes and Gill (1970), McCulloch and 16 Gamble, (1991), and with dashed Ta abundance based on the Nb/Ta ratios 17 reported by Stolz et al. (1996). Additional data sources are same as Fig. DR6. 18 Note that the Bayan Obo mafic rocks have Nb, Ta and Ti negative anomalies 19 and Pb and Sr positive anomalies resembling IAB, and have mostly lower 20 21 elemental abundances than OIB, suggesting a subduction influence in their genesis. 22

2 References for Data Repository

3	Jakes, P., and Gill, J., 1970, Rare earth elements and the island arc tholeiitic
4	series: Earth and Planetary Science Letters, v. 9, p. 17-28.
5	Li, Q.L., Li, X.H., Lan, Z.W., Guo, C.L., Yang, Y.N., Liu, Y., and Tang, G.Q.,
6	2013, Monazite and xenotime U-Th-Pb geochronology by ion microprobe:
7	dating highly fractionated granites at Xihuashan tungsten mine, SE
8	China: Contributions to Mineralogy and Petrology, v. 166, p. 65-80.
9	McCulloch, M.T., and Gamble, J., 1991, Geochemical and geodynamical
10	constraints on subduction zone magmatism: Earth and Planetary Science
11	Letters, v. 102, p. 358-374.
12	Stolz, A., Jochum, K., Spettel, B., and Hofmann, A., 1996, Fluid-and
13	melt-related enrichment in the subarc mantle: evidence from Nb/Ta
14	variations in island-arc basalts: Geology, v. 24, p. 587-590.
15	Sun, SS., and McDonough, W.F., 1989, Chemical and isotopic systematics of
16	oceanic basalts: implications for mantle composition and processes:
17	Geological Society, London, Special Publications 42, p. 313-345.
18	Tong, X.R., Liu, Y., Hu, Z., Chen, H., Zhou, L., Hu, Q., Xu, R., Deng, L., Chen,
19	C., and Gao, S., 2015, Accurate Determination of Sr Isotopic
20	Compositions in Clinopyroxene and Silicate Glasses by
21	LA-MC-ICP-MS: Geostandards and Geoanalytical Research, v. 40, p.
22	85-89.

1	Vermeesch, P., 2006, Tectonic discrimination diagrams revisited:
2	Geochemistry Geophysics Geosystems, v. 7, p. Q06017.
3	Wang, Q.H., Yang, H., Yang, D.B., and Xu, W.L., 2014, Mid-Mesoproterozoic
4	(\sim 1.32 Ga) diabase swarms from the western Liaoning region in the
5	northern margin of the North China Craton: Baddeleyite Pb-Pb
6	geochronology, geochemistry and implications for the final breakup of the
7	Columbia supercontinent: Precambrian Research, v. 254, p. 114-128.
8	Wang, Y., Qiu, Y., Gao, J., and Zhang, Q., 2003, Proterozoic anorogenic
9	magmatic rocks and their constraints on mineralizations in the Bayan Obo
10	deposit region, Inner Mongolia: Science in China (D), v. 46, p. 26-40.
11	Yang, K.F., Fan, H.R., Santosh, M., Hu, F.F., and Wang, K.Y., 2011,
12	Mesoproterozoic mafic and carbonatitic dykes from the northern margin of
13	the North China Craton: implications for the final breakup of Columbia
14	supercontinent: Tectonophysics, v. 498, p. 1-10.
15	Yang, Y.H., Sun, J.F., Xie, L.W., Fan, H.R., and Wu, F.Y., 2008, In situ Nd
16	isotopic measurement of natural geological materials by LA-MC-ICPMS:
17	Chinese Science Bulletin, v. 53, p. 1062-1070.
18	Zhang, H.F., Sun, M., Zhou, X.H., Zhou, M.F., Fan, W.M., and Zhang, J.P.,
19	2003, Secular evolution of the lithosphere beneath the eastern North
20	China Craton: evidence from Mesozoic basalts and high-Mg andesites:
21	Geochimica et Cosmochimica Acta, v. 67, p. 4373-4387.
22	Zhang, M., Suddaby, P., Thompson, R.N., Thirlwall, M.F., and Menzies, M.A.,

1	1995, Potassic volcanic rocks in NE China: geochemical constraints on
2	mantle source and magma genesis: Journal of Petrology, v. 36, p.
3	1275-1303.
4	Zhang, S.H., Zhao, Y., and Santosh, M., 2012, Mid-Mesoproterozoic bimodal
5	magmatic rocks in the northern North China Craton: implications for
6	magmatism related to breakup of the Columbia supercontinent:

7 Precambrian Research, v. 222-223, p. 339-367.



Figure DR1



Figure DR2





Figure DR4





Figure DR6



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Depth(m)	TiO ₂	Al ₂ O ₃	FeO	MgO	CaO	MnO	Na ₂ O	K ₂ O	P ₂ O ₅	BaO	SrO	La ₂ O ₃	Ce ₂ O ₃	Pr ₂ O ₃	Nd_2O_3	Sm ₂ O ₃
392	0.03	0.06	17.20	13.53	30.91	4.30	0.08	0.04	0.04	0.07	0.26	0.67	1.67	0.21	0.68	0.06
520	0.06	0.03	6.55	18.65	32.03	1.77	0.07	0.02	0.23	0.44	0.52	0.36	0.69	0.08	0.24	0.02
1048	0.08	0.98	12.23	13.44	24.58	2.35	1.81	1.72	0.31	0.05	0.63	0.93	2.07	0.20	0.47	0.03
1080	0.05	1.59	7.69	13.89	27.33	2.72	0.04	1.89	1.85	0.20	0.23	0.59	1.55	0.22	0.84	0.09
1336	0.04	0.14	7.74	14.62	31.25	2.60	0.08	0.06	1.97	0.99	0.31	0.57	1.05	0.12	0.37	0.03
1336	0.04	0.06	7.74	14.62	31.25	1.76	0.08	0.01	2.54	0.15	0.46	0.56	1.10	0.13	0.46	0.05
1402	0.02	0.07	5.36	14.37	32.16	1.84	0.03	0.03	3.07	0.06	0.19	1.26	1.81	0.17	0.45	0.04
1410	0.02	0.07	8.48	17.69	34.62	2.29	0.03	0.02	1.36	0.04	0.17	0.67	1.25	0.13	0.40	0.04
1461	0.06	0.06	8.38	17.22	33.60	1.74	0.07	0.02	0.63	0.03	0.81	1.79	2.85	0.28	0.74	0.06
1480	0.05	0.10	7.47	12.17	29.23	1.51	0.05	0.02	5.86	0.13	0.49	0.75	1.41	0.16	0.50	0.06
1502	0.06	0.09	7.95	12.92	29.40	1.47	0.03	0.01	2.35	0.15	0.47	1.62	2.61	0.27	0.81	0.08
1532	0.03	0.05	7.65	13.67	27.45	1.47	0.03	0.01	0.36	0.59	0.26	0.33	0.58	0.06	0.19	0.02
1574	0.05	0.04	8.36	16.55	29.48	1.93	0.02	0.01	0.12	1.30	0.35	1.12	1.82	0.19	0.53	0.05
1588	0.36	0.35	13.48	15.07	27.19	2.33	0.06	0.32	1.90	1.86	0.49	0.65	1.41	0.17	0.51	0.05
1612	0.06	0.09	6.52	16.30	38.86	1.32	0.04	0.05	3.34	0.49	0.39	2.00	2.78	0.27	0.69	0.06
1627	0.25	0.26	13.53	14.76	26.04	2.15	0.03	0.29	1.43	0.22	0.30	0.93	1.56	0.16	0.42	0.04
1641	0.02	0.08	11.30	15.65	28.88	2.38	0.05	0.01	0.63	0.18	0.16	0.29	0.55	0.06	0.19	0.02
1649	0.05	0.71	6.69	11.81	36.17	2.08	0.08	0.34	2.14	0.24	0.23	1.08	1.57	0.15	0.38	0.04
1676	0.02	0.07	8.67	14.67	30.31	1.66	0.28	0.10	0.19	0.39	0.57	1.14	1.81	0.17	0.42	0.03
1683	0.06	0.29	6.81	15.30	25.16	1.73	0.30	0.46	2.09	0.53	0.81	1.67	2.46	0.24	0.65	0.05
1692	0.06	0.21	7.40	17.60	27.45	1.48	0.10	0.40	0.75	0.99	0.83	0.91	1.70	0.17	0.44	0.04
1740	0.04	0.12	10.55	15.58	28.96	1.54	0.25	0.25	0.05	1.10	0.50	0.52	0.99	0.11	0.36	0.03
1765	0.13	0.08	9.40	12.52	35.01	1.47	0.08	0.06	2.43	0.06	0.25	1.12	2.41	0.26	0.71	0.06
1776	0.03	0.03	6.71	14.40	34.55	1.30	0.11	0.02	1.64	0.62	0.24	0.91	1.52	0.16	0.46	0.04

Table DR1. Chemical compositions (wt.%) in Bayan Obo drill cores with different depths

²⁰⁸Pb/²³²Th Th/U Th(ppm) Pb/Th age(Ma) sample U(ppm) σ σ BO-1 2308 3.57 646 0.0462 1.7 913 15 BO-2 4197 1.03 4080 0.0438 1.7 866 14 BO-3 2506 3.04 825 0.0415 1.9 822 16 BO-4 0.0410 1841 2.19 839 13 1.6 811 BO-5 2138 2.93 729 0.0401 795 13 1.6 BO-6 2056 2.60 792 0.0400 1.7 792 13 BO-7 1987 2.32 858 0.0377 748 12 1.6 BO-8 2889 2.08 1389 0.0333 1.8 662 12 BO-9 2.19 0.0318 1783 815 1.9 633 12 BO-10 1436 0.97 1488 0.0288 1.9 573 11 BO-11 2294 2.56 897 0.0285 1.6 569 9 BO-12 1551 1.71 907 0.0280 2.5 558 14 BO-13 1302 2.55 511 0.0266 530 9 1.6 BO-14 3134 1.58 1979 0.0259 1.6 517 8 BO-15 4817 1.32 0.0224 2.2 448 10 3643 BO-16 2654 <0.1 >10000 0.0206 1.6 413 7 BO-17 <0.1 0.0206 7 2407 >10000 1.7 413 BO-18 2230 <0.1 >10000 0.0206 2.0 411 8 BO-19 2934 <0.1 >10000 0.0205 1.8 410 7 BO-20 2307 <0.1 >10000 0.0204 1.8 408 8 BO-21 1709 1015 0.0203 406 7 1.68 1.6 7 BO-22 3272 1.42 2312 0.0197 1.8 394 BO-23 1482 1.83 812 370 6 0.0185 1.7 6 BO-24 1530 1.30 1177 0.0180 1.6 361

Table DR2. Monazite dating data from the drill cores in Bayan Obo

Table DR3. In-situ Nd isotope of monazites from Bayan Obo drill cores

sample	¹⁴⁷ Sm/ ¹⁴⁴ Nd	2σ	¹⁴³ Nd/ ¹⁴⁴ Nd	2σ	age (Ma)	$\epsilon_{\text{Nd}}\left(t\right)^{\#}$	Т _{сник} (Ga)
BO-1	0.04239	2	0.511362	19	913	-6.9	1.59
BO-2	0.04607	5	0.511358	32	866	-8.3	1.63
BO-3	0.04606	2	0.511319	22	822	-9.9	1.66
BO-4	0.04487	4	0.511348	23	811	-9.4	1.62
BO-5	0.04645	4	0.511316	27	795	-10.5	1.67
BO-6	0.04651	5	0.511348	22	792	-10.0	1.64
BO-7rim	0.04363	8	0.511309	37	748	-11.3	1.65
BO-7rim	0.04320	4	0.511305	39	748	-11.3	1.65
BO-7core	0.04349	6	0.511318	41	748	-11.1	1.64
BO-7core	0.04633	4	0.511377	25	748	-10.2	1.61
BO-8	0.04561	7	0.511372	22	662	-11.9	1.61
BO-9	0.04394	2	0.511346	20	633	-12.9	1.62
BO-10	0.04507	14	0.511353	25	573	-14.0	1.62
BO-11	0.04652	3	0.511346	21	569	-14.3	1.64
BO-12	0.04497	4	0.511339	19	558	-14.5	1.63
BO-13	0.04375	3	0.511345	24	530	-14.9	1.62
BO-14	0.04410	10	0.511341	34	517	-15.2	1.62
BO-15rim	0.03962	8	0.511283	38	448	-17.5	1.63
BO-15core	0.04118	6	0.511326	14	448	-16.7	1.61
BO-16	0.03988	3	0.511342	26	413	-17.0	1.58
BO-17	0.03938	2	0.511365	25	413	-16.5	1.56
BO-18	0.03588	9	0.511334	27	411	-17.0	1.55
BO-19	0.04002	11	0.511355	27	410	-16.8	1.57
BO-20	0.03950	4	0.511360	23	408	-16.8	1.56
BO-21	0.04533	3	0.511362	33	406	-17.1	1.62
BO-22	0.04763	20	0.511361	24	394	-17.4	1.64
BO-23	0.04527	3	0.511351	27	370	-18.0	1.62
BO-24core	0.04619	7	0.511377	22	361	-17.7	1.61
BO-24rim	0.04616	2	0.511376	23	361	-17.7	1.61

 ${}^{\#}\epsilon_{Nd}(t)$ values are calculated based on present-day (${}^{147}Sm/{}^{143}Nd)_{CHUR} = 0.1967$ and (${}^{143}Nd/{}^{144}Nd)_{CHUR} = 0.512638$.

dolomite					- (1-1- / -			primary								recrysta	allization
Rb	0.05	bdl [#]	bdl	bdl	bdl	bdl	bdl	bdl	0.09	bdl	bdl	0.01	0.01	0.02	0.01	0.03	0.04
Sr	1766	1528	1818	2228	2174	2559	2671	2429	1197	2640	2738	4294	4513	4378	4559	4451	4289
Ва	67.5	9.34	86.1	13.4	133	124	561	90.1	106	69.4	39.5	43.5	61.6	58.3	33.6	56.0	55.5
Y	23.3	28.8	25	27.9	17.3	13.0	19.3	9.31	23.1	31.5	29.6	56.2	58.3	65.3	105.8	78.4	62.0
La	7.18	3.59	9.17	4.92	5.25	5.06	9.63	3.61	15.5	65.7	27.4	36.6	36.6	56.5	109	101	73.8
Ce	25.2	14.0	25.9	17.3	11.2	10.4	19.6	7.69	54.8	162	62.0	128	136	192	362	314	234
Pr	3.83	2.16	3.65	2.58	1.39	1.16	2.36	0.93	7.50	15.1	7.50	17.8	19.6	26.3	51.8	43.1	31.4
Nd	16.9	11.3	16.6	13.4	5.93	5.9	11.4	4.46	32.4	65.3	33.6	77.4	85.2	112	222	184	136
Sm	5.03	3.37	4.10	3.94	1.52	2.67	4.69	1.81	6.54	17.1	8.94	17.9	20.4	24.8	47.3	37.0	28.4
Eu	2.03	1.60	1.99	1.71	0.73	1.15	1.79	0.84	2.47	6.35	3.36	6.52	7.67	8.64	16.3	12.5	9.60
Gd	3.99	3.48	4.12	3.73	1.75	2.46	4.45	2.44	4.79	14.5	7.22	15.3	17.5	19.6	36.8	28.1	21.7
Tb	0.90	0.81	0.76	0.69	0.42	0.41	0.95	0.40	0.85	2.24	1.30	2.84	3.38	3.68	6.45	5.0	3.92
Dy	5.61	6.23	5.31	5.16	2.82	2.86	4.75	2.26	5.17	10.3	6.85	16.2	18.1	20.1	33.7	26.1	20.1
Ho	1.16	1.32	1.23	1.24	0.79	0.53	0.92	0.36	0.90	1.69	1.28	2.8	3.0	3.38	5.5	4.11	3.23
Er	2.89	4.08	3.17	3.37	1.95	1.30	1.89	0.78	2.37	3.07	3.35	5.41	5.72	6.43	10.1	7.40	5.52
Tm	0.35	0.61	0.43	0.38	0.40	0.17	0.23	0.10	0.25	0.32	0.39	0.55	0.53	0.58	0.87	0.64	0.55
Yb	2.01	3.31	2.74	2.03	2.31	0.79	1.17	0.53	1.38	1.39	2.0	2.18	2.18	2.50	3.75	2.90	2.24
Lu	0.20	0.39	0.22	0.17	0.29	0.11	0.14	0.04	0.17	0.12	0.22	0.18	0.20	0.23	0.32	0.21	0.21
Table DI	R4. contii	nued															
apatite						р	rimary									recrystalliz	ation
Rb	0.1	14	1.40	0.26	0.26	3.	01	0.10	0.08	C	0.04	0.12	0.14	0.12	2	0.48	0.18
Sr	51	09	4026	5006	4586	39	82	2834	2663	2	956	2942	2364	325	7	3255	3223

Table DR4. In-situ trace element compositions (ppm) of dolomite and apatite from Bayan Obo drill cores

Ва

Υ

22.9

22.7

62.4

35.3

78.2

64.3

66.7

La	2133	1598	1918	1553	1412	158	157	202	241	564	495	846	339
Ce	7601	5809	6852	5646	6075	585	525	684	779	1505	1387	2372	1114
Pr	1188	937	1035	912	1004	96.7	82	104	109	228	187	335	146
Nd	5830	4406	4954	4553	5151	500	405	516	517	982	827	1394	647
Sm	1227	902	1051	947	1059	106	88.4	103	100	161	144	208	116
Eu	338	251	287	265	278	26.0	21.3	25.5	24.8	37.2	35.9	49.2	27.1
Gd	800	620	714	675	680	61.6	52.8	61.7	60.6	89.5	85.2	116	67.8
Tb	102	76.6	90.0	83.5	78.7	6.75	5.6	6.55	6.05	8.35	9.24	11.5	7.06
Dy	497	376	451	417	377	36.3	29.3	34.2	31.9	40.9	49.4	58.2	33.5
Ho	72.3	54.7	65.9	61.6	53.6	5.79	4.68	5.85	5.38	6.51	8.17	9.48	5.47
Er	132	102	124	118	99.9	13.8	10.7	12.8	12.4	14.7	19.0	22.5	12.1
Tm	11.1	8.50	10.3	9.92	8.02	1.45	1.07	1.37	1.42	1.53	1.94	2.19	1.21
Yb	38.9	30.3	37.1	36.8	28.9	6.76	5.25	6.1	6.95	7.24	8.45	10.3	5.89
Lu	3.07	2.40	2.98	3.03	2.35	0.63	0.50	0.61	0.79	0.70	0.78	0.99	0.58

[#]below determination limits.

	dolomite				apatite		
primary		recrystalliz	ation	primary		recrystalliz	ation
⁸⁷ Sr/ ⁸⁶ Sr	2σ	⁸⁷ Sr/ ⁸⁶ Sr	2σ	⁸⁷ Sr/ ⁸⁶ Sr	2σ	⁸⁷ Sr/ ⁸⁶ Sr	2σ
0.70250	25	0.70606	33	0.70296	19	0.70351	8
0.70241	20	0.70669	67	0.70294	17	0.70323	19
0.70238	22	0.70384	22	0.70293	23	0.70345	13
0.70271	12	0.70946	53	0.70297	19	0.70349	7
0.70293	16	0.70760	81	0.70297	34	0.70357	16
0.70287	9	0.70456	35			0.70357	18
0.70290	14	0.70786	50			0.70367	11
0.70280	9	0.70682	21			0.70347	14
0.70297	10	0.70572	10			0.70347	12
0.70284	14	0.70971	19			0.70364	18
0.70290	9	0.70889	10			0.70341	13
0.70289	4	0.70871	17			0.70354	13
0.70281	9	0.70533	14				
0.70282	5	0.70568	23				
0.70295	8	0.70827	26				
0.70294	3	0.70718	35				
0.70285	7	0.70467	11				
0.70279	4						

Table DR5. In-situ Sr isotope of dolomite and apatite from Bayan Obo drill cores