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models have been proposed for interpreting the formation of the Neoproterozoic BIFs: (1) During the break-up of Rodinia supercontinent!

induced the addition of Fe flux from submarine exhalative (Basta et al., 2011; Cox et al., 2013); (2) The thick ice sheet formed by "Snowball Earth" event isolated the hydrosphere and oxidized atmosphere, which led to the reduction of ocean and the dissolution of a large amount of Fe element. After the ice cover melted, the Fe was oxidized and precipitated into BIF as a result of the contact between ocean and atmosphere (Hoffman et al., 1998; Hoffman and Schrag, 2002; Klein and Ladeira, 2004; Halverson et al., 2011) and (3) during the late Neoproterozoic, there was a reduced S-poor and Fe-rich environment, which was favorable for Fe element existing as divalent ions in seawater, and then to be oxidized and precipitated (Johnston et al., 2010).

In this study, we report a newly identified Neoproterozoic Baijianshan BIF at the southeastern margin of the Tarim Block in NW China. The precipitation mechanism and the oceanic redox condition of the Baijianshan BIF was investigated based on detailed field observations coupled with systematic elemental and isotopic compositions analysis. This study sheds a new light on our understanding of the possible coupling of the Cryogenian snowball Earth event and the formation of Neoproterozoic BIFs.

2 Regional Geology

The Tarim Craton in NW China, covering an area of more than 600,000 km², is one of the main three Precambrian nuclei in China (i.e., North China, South China and Tarim). It is surrounded by the Phanerozoic orogenic belts of Tianshan Mountains to the north, the western Kunlun Mountains to the south, and the Central-Southern Altyn Tagh Mountains to the southeast. Despite most of the Tarim Craton being covered by aeolian sands and difficulties in accessibility, previous work reveals that the craton is characterized by a typical double-layered structure consisting of a Pre-Cryogenian basement and Cryogenian-Cambrian cover sequences (Xinjiang BGMR, 1993; Zhang et al., 2013). Particularly, recent studies demonstrate that the craton docked at the northern fringe of the Rodinia slightly earlier than 760 Ma and then the Cryogenian-Cambrian sedimentary sequences deposited during the breakup of the Rodinia (Zhang et al., 2010, 2016; Xu et al., 2013).

These samples are dominated by magnetite and quartz with banded structures and mainly show mesoband (mm- to cm-scale) and microband (sub-mm- to mm-scale), which are composed of magnetite, quartz, rare hematite, siderite, albite and biotite (Fig. 3e-f). Quartz and other gangue minerals, such as chlorite (Fig. 3g) and sericite (Fig. 3h), occur occasionally as veinlets cutting across the bedding.

The shallow gray tuffs under the four layers of Fe ore bodies have been altered into clays. Nevertheless, the sub-euhedral or euhedral plagioclase and quartz crystal fragments, mostly less than 2 mm, can be seen under magnifier. The crystal fragments account for about 15-20% in the tuffs. One sample was collected from the tuff to constrain the deposition age of the Baijianshan BIF $(C(.\%^*, *)!:. *\%(-.\%:2071)$.

The blackish green basalt layer occurs at the bottom of the BIF layers, with the thickness of 1 meter. The minerals in the basalt were intensively altered and neddle-like albite, chlorite as well as Ti-Fe oxide can be observed in thin sections, three samples were collected from the basalt layer for geochemical analysis (2702H1, 270H2 and 2702H3).

4 Analytical Methods

4.1 In situ zircon U-Pb isotope analysis

Conventional magnetic and density techniques were used to concentrate non-magnetic, heavy fractions and then zircon grains were hand-picked under a binocular microscope. After being mounted in epoxy mount, zircon grains were then polished to section the crystals in half for analysis. All zircons were documented with transmitted and reflected light micrographs as well as cathodoluminescence (CL) images to reveal their internal structures. Zircon U-Pb ages were analyzed using the LA-ICP-MS at the Tianjin Institute of Geology and Mineral Resources (China Geological Survey). Detailed analytical procedures can be found in Li et al. (2009). Zircon standards 91500 and GJ-1 (Jackson et al., 2004) were used to monitor the analysis. The U-Pb concordia plots were processed by ISOPLOT 3.0 and data were presented with 1 errors and 95% confidence limits (Ludwig, 2003). The zircon U-Pb age data are listed in Supplementary Table 1.

4.2 Whole rock geochemistry

Seventeen BIF samples and three basalt samples were carefully selected along the geological location. Whole-rock major compositions were analyzed using standard X-ray fluorescence (XRF) on fused glass beads at the Nanjing Institute of Geology and Mineral Resources (China Geological Survey), following the procedures similar to those described by Li et al., (2006a, b). Analytical uncertainties are between 1 and 5%. In addition, we used wet chemistry technique to measure FeO and Fe_2O_3 , with the procedures described by Andrade et al. (2002). The measured data are listed in supplementary Table 2.

Trace elements were determined using a Perkin-Elmer Sciex ELAN DRC-e ICP-MS at the State Key Laboratory of Ore Deposit Geochemistry, Institute of Geochemistry (Chinese Academy of Sciences), with the analytical accuracy better than 5%. Samples were digested with 1ml of HF and 0.5 ml of $\rm HNO_3$ in screw top PTFE-lined stainless steel bombs at 190 for 12h (Qi et al., 2000). The analytical precision for most elements was better than 1% with concentrations > 200 ppm, and 1-3% when less than 200 ppm. The analytical results are reported in supplementary Table 2.

Sr-Nd isotopes were measured using the Micromass Isoprobe Multi-collector MC-ICP-MS at Tianjin Institute of Geology and Mineral Resources (China Geological Survey), with the analytical procedures similar to those reported by Li et al. (2004). Measured ⁸⁷Sr/⁸⁶Sr and ¹⁴³Nd/¹⁴⁴Nd ratios were corrected for mass-fractionation using ⁸⁶Sr/⁸⁸Sr = 0.1194 and ¹⁴⁶Nd/¹⁴⁴Nd = 0.7219, respectively. The measured values for the NBS SRM 987 standard ⁸⁷Sr/⁸⁶Sr = 0.71025 and the Shin Etsu JNdi-1 standard ¹⁴³Nd/¹⁴⁴Nd = 0.512115. All the initial ¹⁴³Nd/¹⁴⁴Nd values of the BIF are calculated with a probable depositional age (737 Ma) for the Baijianshan BIF.

Iron isotope ratios were measured using MC-ICP-MS at Beijing Createch Testing Technology Co., Ltd. The results of Fe isotope ratio determination as the ten thousandth deviation of the sample from the standard sample is as follows:

$$\delta^{x} Fe/^{54} Fe(\%_{0}) = \left[\frac{(^{x} Fe/^{54} Fe)_{sample}}{(^{x} Fe/^{54} Fe)_{standard}} - 1\right] \times 1000$$

$$(x = 56, 57)$$

The performance of the instrument was assessed by repeated measurements of an in-house standard (CAGS Fe) that yielded deviations relative to the IRMM-014 Fe isotope reference material. The ; $8.6 < H; \qquad ^{56}Fe_{IRMM014} \qquad 2^{56}Fe_{CAGS} \qquad ^{96-}\% \qquad ^{57}Fe_{IRMM014} \qquad 2^{57}Fe_{CAGS} \qquad ^{1}\% \qquad \qquad ribed by Tang et al. (2016).$

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Three basalt. The incompatible Nb/Y ratios (1.05-1 into the alkaline basal ranging from 186 ppm to (Fig. 5a) ($La_N/Yb_N = 9.8-1$ ppm), the basalts are evolved features of the OIB-like basalts

5.2.2 Geochemistry of the BIF

As shown in supplementary Table 1. Fe₂O₃^T concentrations vary between 1. between 26.81% and 55.06% (42.02% oh 3.66% on average), TiO₂, MnO, P₂O₅, CaO, incompatible elements (supplementary Table 2.

Their REE abundances vary from 38.1 ppm to Shale (PAAS; McLennan, 1989), they are characteristic the La_N/Yb_N of 0.23-0.69 (Fig. 5b). The REY pattern of $Ce/Ce^* \sim 1$ and $Pr/Pr^* \sim 1$; Bau and Dulski, 1996, 0.96-1.14; Y/Ho ratios ranging from 23.3 to 31.8 with an

5.2.3 Sr-Nd isotope compositions

Sr-Nd concentrations and isotopic ratios of the Baijian supplementary Table 3. As for the basalts, the basalts exhibit h (87Sr/86Sr)_i ranging from 0.7061 to 0.7084 due to post-eruption

6.9.

Strontium and neodymium concentrations of Baijianshan BIF range from 7.06 ppm to 59.8 ppm, respectively. The nine samples have a large r (0.51231-0.51243) and 87 Sr/ 86 Sr (0.7179- 96) +(I C (t = 7371 to 1.46, with an average of 0.37. However, the BIF samples exhibit a large values between 0.6412 and 0.7175, possibly due to the post-deposition hydroth large range of Rb/Sr ratios varying from 0.02 to 3.47.

5.2.4 Fe isotope compositions

The Fe isotope compositions of the Baijianshan BIF samples are presented in supple The six samples yield ⁵⁶Fe_{IRMM-014} values of 1.24-', ⁵⁷Fe_{IRMM-014} values of Their enrichment in heavy Fe isotopes shares the signatures of the Neoproterozoic BIFs al., 2011; Yan et al., 2011; Shen et al., 2018).

6 Discussions

6.1 Iron sources for the Baijianshan BIFs

Field and microscopic observations revealed that the Neoproterozoic Baijianshan BIF has n

experienced significant metamorphism (Fig. 3b, c and d), indicating the original geochemical characteristics were well retained and can be used to decipher the iron sources and the oceanic environments when it precipitated.

The Baijianshan BIF shows a modern seawater REE signature with significant LREE depletion relative to the HREE (La_N/Yb_N = 0.41) (Fig. 5b, Zhang and Nozaki, 1996; Alibo and Nozaki, 1999). A range of factors can affect the primary REY composition in terms of BIF depositional processes, including post-depositional metamorphism and/or syn-deposition clastic contamination (Alexander et al., 2008; Hu et al., 2020). Previous studies demonstrated that diagenetic and metamorphic events (such as weathering and/or fluid-rock interactions) have negligible effects on the REY of the chemical sediments (Bolhar et al., 2004). However, Rb is a relatively mobile element compared to highly immobile elements such as Th. In Fig. 6a, the well positive correlation between Rb and Th demonstrates insignificant mobility of the LILEs (Large Ion Lithophile Elements) and the absence of significant diagenetic or metamorphic alteration effects, this conclusion is also consistent with thin section observations (Fig. 3c, d, g, h). With respect to the syn-depositional processes, the contents of Al₂O₃ and TiO₂ are consistently low with averages of 3.66% and 0.41%. In addition, despite the positive correlations between Al_2O_3 and TiO_2 (r = 0.93), K_2O (r = 0.79), Rb (r = 0.78), V (r = 0.45), Sc 2 %++! 8 2 %, '! REE (r = 0.62) (Fig. 6b-i) signify the incorporation of 2 %, , ! 8 terrigenous contribution in the deposition processes, most correlation coefficients are much lower than those BIFs sourced from recycled crust (Basta et al., 2011; Cox et al., 2013; Hu et al., 2017, 2020). Besides, the Baijianshan BIF shares similar REE patterns with those BIFs from North China Craton but contains slightly higher contents of Al₂O₃ and TiO₂ (Li et al., 2014). Taken together, we suggest a little incorporation of terrigenous component in the Baijianshan BIF.

Positive Eu anomalies are generally interpreted as the precipitation of high-temperature hydrothermal fluids (Danielson et al., 1992; Bau and Dulski, 1999) while low-temperature hydrothermal fluids usually display weak or no Eu anomalies (Michard et al., 1993; Li et al., 2014). Most samples from the Baijianshan BIF show negligible Eu anomalies (Eu/Eu* = 0.96~1.14, averaging 1.07), which are indicative of low-temperature hydrothermal solutions (Danielson et al., 1992). The C (t) value can be used as a tracer to distinguish different sources and crustal contamination. As shown in Fig. 7, the Baijianshan BIF has slightly average C (t) value (average of C 2 0.37), much lower than those of Baijianshan basalt, indicating that the deposition of the Baijianshan BIF was controlled by a hydrothermal flux with mantle
C (Jacobsen and Pimentel-Klose, 1988). However, the slightly negative correlation between C , (, Ma) and Al₂O₃ contents (Fig. 8a) argues for the low proportion of detrital components inputting.

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Ce/Ce*PAAS values with variable Al₂O₃ contents of the Baijianshan BIF indicate that terrigenous detrital materials played little role in the geochemical budget of Ce. In oxidized seawater, Ce(III) transforms into Ce(IV), and then Ce(IV) is likely to be hydrolyzed and precipitated with Fe-Mn oxyhydroxides, organics and clay which lead to a significant negative Ce anomaly of seawater (Byrne and Sholkovitz, 1996). As shown in the Ce/Ce* vs. Pr/Pr* diagram (Fig. 10), all the Baijianshan BIF samples display no Ce anomalies, similar with most Neoproterozoic BIFs, unambiguously arguing for anoxic environment in the ancient ocean. Although the atmospheric and oceanic oxygen levels during the Neoproterozoic are still under debate, numerous studies revealed that Neoproterozoic oceans underwent a stepwise and protracted oxidation. Anoxic ferruginous deep seawater was a typical feature of the late Neoproterozoic, as inferred from geochemical proxies such as iron geochemistry (e.g., Canfield et al., 2008; Sperling et al., 2015), redox-sensitive elements (e.g., Schröder and Grotzinger, 2007; Rajabi et al., 2015), and framboidal pyrite (e.g., Rajabi et al., 2015). Stern et al. (2013) suggested that this scenario might be attributed to the seawater surfaces being covered by ice sheets, blocking the oxygen from atmosphere dissolving into the widespread anoxic ferruginous ocean. Despite no tillite was identified at the Baijianshan BIF, the coeval tillite sequences were documented in the Quruqtagh area (i.e., ca.740 Ma Beiyixi tillite) of NE Tarim (Xu et al., 2009), which argued for the possibility of ice sheets in the Cryogenian Baijianshan ocean.

In a Fe²⁺-rich marine settings, both abiotic and biotic action were demonstrated as oxidation pathways for the precipitation of iron oxyhydroxides (Fe(OH)₃) from dissolved Fe²⁺ (Konhauser et al., 2011). As an element of variable valency, the fractionation of Fe isotope is affected by the oxidation-reduction condition (Bullen et al., 2001; Zhu et al., 2002; Balci et al., 2006). Experimental studies demonstrated that oxidation of Fe²⁺ to Fe³⁺ in solution causes considerable Fe isotope fraction, generating an enrichment of heavy Fe isotopes in Fe³⁺ (Bullen et al., 2001; Johnson et al., 2002; Balci et al., 2006; Hou et al., 2014). On the other hand, the fractionation caused by pyrite with light-Fe isotopes is considered to play a leading role in the enrichment of heavy Fe isotopes in the residual system relative to the original state (Rouxel et al., 2016), since the pyrite is easier to precipitation from plume. Nevertheless, there is no sulfide observed in the Baijianshan BIF, thus the potential influences of sulfides on the enrichment of heavy Fe isotopes can be excluded. Under oxidized environment, the enrichment of heavy Fe isotopes is shown in oxide or hydroxide of iron trivalent (Fe3+ ppt) and light Fe isotope is displayed in Ferrous solution (Fe²⁺aq) (Johnson et al., 2002). The magnitude of Fe isotope fractionation is controlled by the degree of precipitation from Fe²⁺ to Fe³⁺, which is related to the degree of ocean oxidation (Yan et al, 2010). The Fe in the seawater can be completely precipitated and no fractionation of the Fe isotope occurs when the seawater is completely oxidized, thus the Fe isotope values in iron oxide precipitates (Fe³⁺) can represent the Fe isotope information of seawater. However, when the seawater is partially oxidized, the Fe in the seawater is partially precipitated and the Fe isotope fractionation will occur in iron oxide precipitates (Fe³⁺). Therefore, the Fe isotope value can be used as a proxy to decipher the redox state of seawater (Yan et al., 2010). Compared with some BIFs deposited during the Archean-Paleoproterozoic (Dauphas et al., 2004; Rouxel et al., 2005; Li et al., 2012; Johnson et al., 2003; Hou et al., 2014; Li et al., 2014), the Neoproterozoic Baijianshan Sanjiang (Yan et al., 2010) and Xinyu BIFs (Shen et al., 2008) have higher values ⁵⁷Fe (Fig. 11). Several studies demonstrated that the low 57Fe values of BIFs could be due to the contributions from a continental component and the continental Fe source is best explained by Fe mobilization on the continental margin by microbial dissimilatiry iron reduction (DIR) (Johnson et al., 2008; Li et al., 2015). As for the Fe isotope values of Archean-Paleoproterozoic BIFs we cited, the Fe source of SW Greenland BIF, Anshan-Benxi BIFs, Gongchangling BIFs and Yuanjiacun BIFs are unambiguously demonstrated to be hydrothermal type similar to the scenario of the Baijianshan BIF (Dauphas et al., 2004; Li et al., 2012; Li et al., 2014; Hou et al., 2014), while the Fe of Transvaal, Manjeri, Belingwe Belt and Zimbabwe BIFs are probably from multiple sources (abiologic and biologic processes) (Johnson et al., 2008; Grassineau et al., 2011). The comparison reveals that the environment in which they precipitated was more reduced. Both the iron isotope and elemental geochemistry thus demonstrate that the Neoproterozoic Baijianshan ocean might have the same reduced environment as Archean or Paleoproterozoic, or even more reduced.

6.3 A model of the Baijianshan BIF

During the late Neoproterozoic period (i.e., 750-550 Ma), the Earth experienced long-lived global glaciations, known as the Snowball Earth events (Hoffman et al., 1998). In view of the Neoproterozoic oxygenation event (NOE) at that time, the content of oxygen in atmosphere was much higher than before (Och et al., 2012). Recently, EI-Rahman et al. (2020) argued that some Neoproterozoic BIFs are not glaciogenic due to their significant depleted Nd isotope compositions. Under this scenario, the iron was most likely deriving from hydrothermal alteration of juvenile oceanic crust. The Chondritic-like Nd isotope compositions of the Baijianshan BIF (C 2 %(, do not favor this model. Furthermore, the close temporally-spatially related tillite sequences in the Quruqtagh of NE Tarim (Xu et al., 2009), 765 Ma-735 Ma tillites in Namibia, suggested a local glaciation termed as Kaigas

glaciation (Kay et al., 2001; Cailteux et al., 2005; Hoffmann et al., 2006). Considering the high latitude position of the Tarim in the Rodinia configuration (Li et al., 2008), the Kaigas glaciation (corresponding to the Beiyixi tillite in Tarim) most possibly occurred in Tarim. In combination with previous studies on other BIFs with the data presented in this study, we construct a cartoon model showing the formation mechanism of the Baijianshan BIF (Fig. 12).

Stage 1, the seawater was covered by ice sheet which led to the insulation between atmosphere and And then it induced the emergence of a reductive oceans with the in Н ocean. The iron supplied by low-temperature hydrothermal fluids could exist as the ferrous form, forming a relatively stable anoxic ferruginous reservoir in the ocean.

Stage 2, at interglacial period, the partial melting of the ice sheet caused some oxygen from the atmosphere into the ocean, which emerged a stratified seawater. There were mainly anoxic and Fe²⁺-rich deeper seawater, whereas the Fe²⁺ was gradually oxidized into Fe³⁺ and precipitated into iron oxide in near-shore oxic shallow seawaters. The low-temperature hydrothermal was the dominant source during the precipitation process to interpret the slight positive Eu anomalies and positive values

. Under the effect of the melting ice-sheet, the increasing weathering then contributed to a small amount of terrestrial materials into the shallow seawater, resulting the precipitation of the Baijianshan BIF.

7 Conclusions

The Baijianshan BIF precipitated in glacially-influenced settings during the Neoproterozoic (737 Ma). Low-T hydrothermal fluids, accompanied with limited detritus input, were the dominated iron sources for the formation of Baijianshan BIF.

The Baijianshan ocean was initially covered by ice sheets, giving rise to the extremely reduced environment favored by Fe²⁺. Then it was oxidized into Fe³⁺ and precipitated as magnetite due to the thawing of ice sheets. At the early stage, local reduced ocean enhanced Fe fluxes genetically related to the intensive marine magmatism and hydrothermal activities due to the existence of ice sheets. Thawing of ice sheets induced oxidized oceanic layer, favoring the formation of the Neoproterozoic BIF at the later stage.

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References

- Alexander, B.W., Bau, M., Andersson, P., Dulski, P., 2008. Continentally-derived solutes in shallow Archean seawater: rare earth element and Nd isotope evidence in iron formation from the 2.9 Ga Pongola Supergroup, South Africa. Geochim. Cosmochim. Acta 72, 378-394.
- Alibo, D.S., Nozaki, Y., 1999. Rare earth elements in seawater: particle association, shale normalization and Ce oxidation. Geochim. Cosmochim. Acta 63(3-4), 363-372.
- Andrade, S., Hypolito, R., Ulbrich, H.H.G.J., Silva, M.L., 2002. Iron (II) oxide determination in rocks and minerals. Chem. Geol. 182, 85-89.
- Balci, N., Bullen, T.D., Witte-Lien, K, Shanks, W.C., Motelica, M., Mandernack, K.W., 2006. Iron isotope fractionation during microbially stimulated Fe(II) oxidation and Fe(III) precipitation. Geochim Cosmochim Acta 70, 622-639.
- Basta, F.F., Maurice, A.E., Fontboté, L., Favarger, P.Y., 2011. Petrology and geochemistry of the banded iron formation (BIF) of Wadi Karim and UmAnab, Eastern Desert, Egypt: implications for the origin of Neoproterozoic BIF. Precambrian Res. 187, 277-292.
- Bau, M., Dulski, P., 1996. Distribution of yttrium and rare-earth elements in the Penge and Kuruman iron-formations, Transvaal Super group, South Africa. Precambrian Res. 79, 37-55.
- Bau, M., Dulski, P., 1999. Comparing yttrium and rare earths in hydrothermal fluids from the Mid-Atlantic Ridge: implications for Y and REE behaviour during nearvent mixing and for the Y/Ho ratio of Proterozoic seawater. Chem. Geol. 155, 77-90.
- Bolhar, R., Kamber, BS., Moorbath, S., Fedo, C.M., Whitehouse, M.J., 2004. Characterisation of early Archaean
- chemical sediments by trace element signatures. Earth Planet. Sci. Lett. 222, 43-60.

 Bullen, T.D., White, A.F., Childs, C.W., Vivit, D.V., Schulz, M.S., 2001. Demonstration of significant abiotic iron isotope fractionation. Geology 29, 699-702.
- Byrne, R., Sholkovitz., 1996. Marine chemistry and geochemistry of the lanthanides. In: Gschneider Jr.K.A.,
- Eyring, L. (Eds.), Handbook on the Physics and Chemistry of the Rare Earths. Elsevier, Amsterdam.

 Cailteux, J.L.H., Kampunzu, A.B.H., Batumike, M.J., 2005. Lithostratigraphic position and petrographic G 6 I G 6 -I H ! C @ 7 8 G 6 I G 6 -I Journal of African Earth Sciences 42, 82-94.
- Canfield, D.E., Poulton, S.W., Knoll, A.H., Narbonne, G.M., Ross, G., Goldberg, T., Strauss, H., 2008. Ferruginous Conditions Dominated Later Neoproterozoic Deep-Water Chemistry. Science 321(5891), 949-952. Chen, D.Z., Wang, J.G., Qing, H.R., Yan, D.T., Li, R.W., 2009. Hydrothermal venting activities in the Early

- Cambrian, South China: Petrological, geochronological and stable isotopic constraints. Chemical Geology 258,
- Cox, G.M., Halverson, G.P., Minarik, W.G., Heron, D.P.L., Macdonald, F.A., Bellefroid, E.J., Strauss, J.V., 2013. Neoproterozoic iron formation: an evaluation of its temporal, environment al and tectonic significance. Chem. Geol. 362, 232-249.
- Cox, G.M., Halverson, G.P., Poirier, A., Le Heron, D., Strauss, J.V., Stevenson, R., 2016. A model for Cryogenian
- iron formation. Earth Planet. Sci. Lett. 433, 280-292.

 Danielson, A., Moller, P., Dulski, P., 1992. The europium anomalies in banded iron formations and the thermal history of the oceanic crust. Chem. Geol. 97, 89-100.
- Dauphas, N., van Zuilen, M., Wadhwa, M., Davis, A.M., Marty, B., Janney, P.E., 2004. Clues from Fe isotope variations on the origin of early Archean BIFs from Greenland. Science 306, 2077-2080.
- EI-Rahman, Y.A., Gutzmer, J., Li, X.H., Seifert, T., Li, C.F., Ling, X.X., Li, J., 2020. Not all Neoproterozoic iron formations are glaciogenic: Sturtian-aged non-Rapitan exhalative iron formations from the Arabian-Nubian Shield. Mineralium Deposita 55, 577-596.
- Feng, L.J., Huang, J., Lu, D.B., Zhang, Q.R., 2016. Major and trace element geochemistry of the Neoproterozoic syn-glacial Fulu iron formation, South China. Geological Magazine 154, 1-10.
- Freitas, B.T., Warren, L.V., Boggiani, P.C., Almeida, R.P.D., Piacentini, T., 2011. Tectono-sedimentary evolution of the Neoproterozoic BIF-bearing Jacadigo Group, SW-Brazil. Sedimentary Geology 238, 48-70.
- Grassineau, N.V., Nisbet, E.G., Bickle, M.J., Fowler, C.M.R., Lowry, D., Mattey, D.P., Abell, P., Martin, A., 2000. Antiquity of the biological sulphur cycle: evidence from sulphur and carbon isotopes in 2700 million-year-old rocks of the Belingwe Belt, Zimbabwe. Proc. R. Soc. Lond. B 268, 113-119.
- Gross, G.A., 1980. A classification of iron-formation based on depositional environments. Canadian Mineralogist 18, 215-222.
- Gross, G.A., Mcleod, C.R., 1980. A preliminary assessment of the chemical composition of iron formation in Canada. Canadian Mineralogist 18, 223-229.
- Halverson, G.P., Poitrasson, F., Hoffman, P.F., Nédélec, A., Montel, J.M., Kirby, J., 2011. Fe isotope and trace element geochemistry of the Neoproterozoic syn-glacial Rapitan iron formation. Earth Planet. Sci. Lett. 309(1),
- Hoffman, P.F., Kaufman, A.J., Halverson, G.P., Schrag, D.P., 1998. A Neoproterozoic snowball Earth. Science 281, 1342-1346.
- Hoffman, P.F., Schrag, D.P., 2002. The snowball earth hypothesis: testing the limits of global change. Terra Nova 14(3), 129-155.
- Hoffmann, K.H., Condon, D.J., Bowring, S.A., Prave, A.R., Fallick, A., 2006. Lithostratigraphic, carbon (¹³C) isotope and U-Pb zircon age constraints on early Neoproterozoic (ca. 755 Ma) glaciation in the Gariep Belt, southern Namibia. Snowball Earth Conference, July 16-21, 2006, Montle Namibia, Ticino, Switzerland, 51. Holland, H.D., 2002. Volcanic gases, black smokers, and the great oxidation event. Geochimica et Cosmochimica
- Acta 66(21), 3811-3826.
- Hou, K.J., Li, Y.H., Gao, J.F., Liu, F., Qin, Y., 2014. Geochemistry and Si-O-Fe isotope constraint on the origin of banded iron formations of the Yuanjiacun Formation, Lvliang Group, Shanxi, China. Ore Geology Reviews 57,
- Hu, J., Wang, H., Wang, M., 2017. Geochemistry and origin of the Neoproterozoic Dahongliutan banded iron formation (BIF) in the Western Kunlun orogenic belt, Xinjiang (NW China). Ore Geology Reviews 89,
- Hu, J., Wang, H., Zhang, L.G., 2020. A rare earth element and Nd isotopic investigation into the provenance and deposition of the Dahongliutan banded iron formation and associated carbonates, NW China: Împlications on Neoproterozoic seawater compositions. Precambrian Research 342, 1-29.
- Jackson, S.E., Pearson, N.J., Griffin, W.L., Belousova, E.A., 2004. The application of laser ablation-inductively coupled plasma-mass spectrometry to in situ U Pb zircon geochronology. Chemical Geology 211(1-2), 47-69.
- Jacobsen, S.B., Pimentel-Klose, M.R., 1988. Nd isotopic variations in Precambrian banded iron formations. Geophys Res Lett 15, 393-396.
- James, H.L., 1954. Sedimentary facies of iron-formation. Economic Geology 49(3), 235-293.
- James, H.L., 1966. Chemistry of the iron-rich sedimentary rocks. U.S. Geol. Survey Prof, 440-461.
- James, H.L., 1983. Distribution of banded iron-formation in space and time. Developments in Precambrian Geology 6, 471-490.
- !?B!'%% 6 Johnson, C.M., Beard, B.L., Beukes, N.J., @ ! 8 ! D A
- Earth as inferred from Fe isotope studies of banded iron formations from the Transvaal Craton. Contributions to Mineralogy and Petrology 144(5), 523-547.
- Johnson, C.M., Beard, B.L., Klein, C., Beukes, N.J., Roden, E.E., 2008. Iron isotopes constrain biologic and abiologic processes in banded iron formation genesis. Geochimica Et Cosmochimica Acta 72(1), 151-169.
- Johnson, C.M., Skulan, J.L., Beard, B.L., Sun, H., Nealson, K.H., Braterman, P.S., 2002. Isotopic fractionation between Fe(III) and Fe(II) in aqueous solution. Earth Planet Sci Lett 195, 141-153.
- Johnston, D.T., Poulton, S.W., Dehler, C., Porter, S., Husson, J., Canfield, D.E., Knoll, A.H., 2010. An emerging picture of Neoproterozoic ocean chemistry: insights from the Chuar Group, Grand Canyon, USA. Earth Planet. Sci. Lett. 290, 64-73.
- Kay, R.M., Liyungu, A.K., Njamu, F.M., Somwe, V., Banda, J., Mosley, P.N., Armstrong, R.A., 2001. The western arm of the Lufilian Arc in NW Zambia and its potential for copper mineralization. Journal of African Earth Sciences 33, 503-528.
- Klein, C., 2005. Some Precambrian banded iron-formations (BIFs) from around the world: Their age, geologic setting, mineralogy, metamorphism, geochemistry, and origins. American Mineralogist 90(10), 1473-1499.
- Klein, C., Beukes, N.J., 1992. Time distribution, stratigraphy, sedimentologic setting, and geochemistry of Precambrian iron-formations. In: Schopf, J.W., Klein, C. (Eds.), The Proterozoic Biosphere. Cambridge University Press, Cambridge, 139-146.
- Klein, C., Ladeira, E.A., 2004. Geochemistry and mineralogy of Neoproterozoic banded iron-formations and some selected, siliceous manganese formations from the Urucum district, Mato Grosso do Sul, Brazil. Econ. Geol. 99, 1233-1244
- Konhauser, K.O., Kappler, A.K., Roden, E.E., 2011. The microbial role in iron redox and biomineralization reactions. Elements 7, 89-93.

- change. Precambr. Res. 239, 79-94.
- Change. Precambr. Res. 239, 79-94.
 Sun, S.S., McDonough, W.F., 1989. Chemical and isotopic systematics of oceanic basalts: Implications for mantle composition and processes. Geological Society London Special Publications 42(1), 313-345.
 Tang, S.H., Zhu, X.K., Li, J., Yan, B., Li, S.Z., Li, Z.H., Wang, Y., Sun, J., 2016. New Standard Solutions for Measurement of Iron, Copper and Zine Isotope Compositions by Multi-collector Inductively Coupled Plasma-Mass Spectrometry. ROCK AND MINERAL ANALYSIS 35(2), 127-133 (in Chinese with English
- Wu, Y.B., Zheng, Y.F., 2004. A study on the genetic mineralogy of zircon and its restriction on the interpretation of U-Pb age. CHINESE SCIENCE BULLETIN 49(16), 1589 (in Chinese).
- Xinjiang Bureau of Geology and Mineral Resources (BGMR), 1993. Regional Geology of Xinjiang Uygur
- Autonomous Region. Geological Publishing House, Beijing, 8-33 (in Chinese).

 Xu, B., Xiao, S.H., Zou, H.B., Chen, Y., Li, Z.X., Song, B., Liu, D.Y., Zhou, C.M., Yuan, X.L., 2009. SHRIMP zircon U-Pb age constraints on Neoproterozoic Quruqtagh diamictites in NW China. Precambrian Research 168, 247-258.
- Xu, B., Zou, H.B., Chen, Y., He, J.Y., Wang, Y., 2013. The Sugetbrak basalts from northwestern Tarim Block of northwest China: Geochronology, geochemistry and implications for Rodinia breakup and ice age in the Late Neoproterozoic. Precambrian Research 236, 214-226.

 Xu, D.R., Wang, Z.L., Chen, H.Y., Hollings, Pete, Jansen, Nicholas, H., Zhang, Z.C., Wu, C.J., 2014. Petrography
- and geochemistry of the Shilu Fe Co Cu ore district, South China: Implications for the origin of a
- Neoproterozoic BIF system. Ore Geology Reviews 57, 322-350.

 Xu, Z.Q., He, B.Z., Zhang, C.L., Zhang, J.X., 2013. Tectonic framework and crustal evolution of the Precambrian basement of the Tarim Block in NW China: New geochronological evidence from deep drilling samples. Precambrian Research 235, 150-162.
- Yan, B., Zhu, X.K., Tang, S.H., Zhu, M.Y., 2010. Fe isotopic characteristics of the Neoproterozoic BIF in Guangxi 5.32 841.92 reW*n0 gBT0 Th/fb/incef hold its214hpRc56thif3883/865f AA.66H000008871 0 595.32 841.92 reW*n0 gBT0 Tr/F1 9 Tf1 0 0 1 316 591.7 Tm86.86. A108ETQ

Figure captions

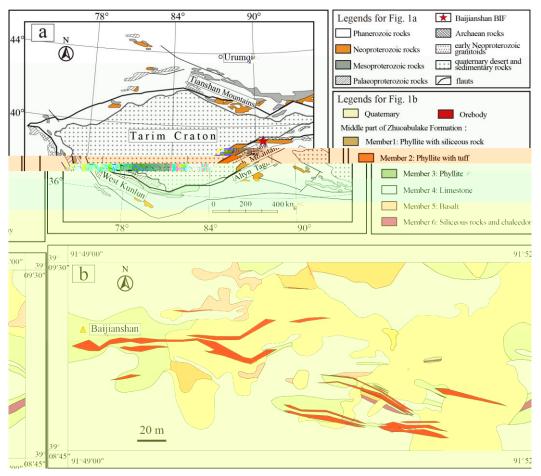


Fig. 1 (a) Sketch Precambrian geological map of Tarim Craton, showing the location of the Baijianshan BIF (modified after Lu et al., 2008). (b) Geological map of the Baijianshan Banded Iron Formation (Baijianshan BIF).

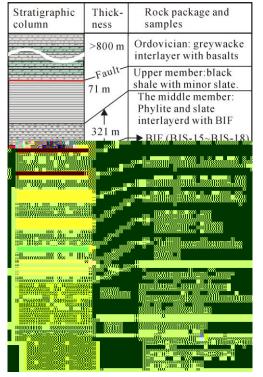


Fig. 2 Stratigraphical column of the Baijianshan BIF.

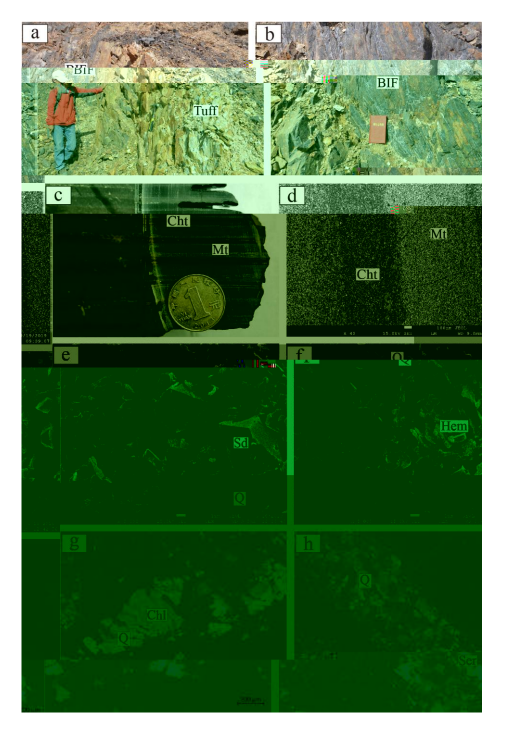


Fig. 3 Field photographs, hand specimen and mineral features of the Baijianshan BIF. (a) The tuff layers in the Baijianshan BIF. (b) Laminated Fe orebodies with local bend. (c) hand specimen. (d) Scanning electron microscopy (SEM) images. Detailed petrographic examination by SEM and transmitted light showing mineral assemblage: (e) Siderite. (f) Hematite. (g) Chlorite. (h) Sericite. Cht = Chert, Mt = Magnetite, Mt = Siderite, Mt = Quartz, Mt = Hematite, Mt = Chlorite, Mt = Chlorite

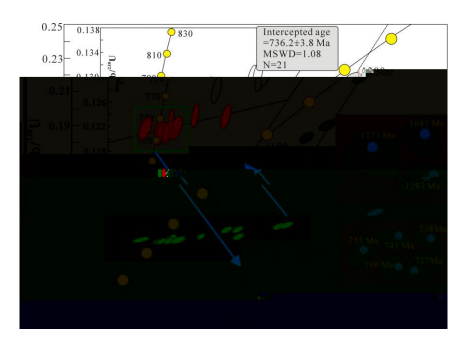


Fig. 4 Concordia diagram of the zircon U-Pb ages of the tuff at the Baijianshan BIF.

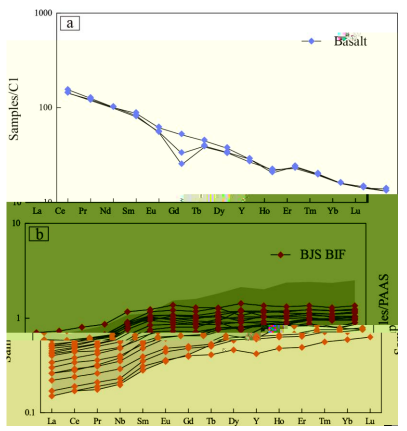


Fig. 5 (a) Chondrite-normalized REE parttern of the basalts. Chondrite-normalized values after Sun and McDonough (1989). (b) PAAS-normalized REE+Y diagrams of BIFs from the Baijianshan BIF. The grayish zone represents the REY data from the Xinyu BIF (Li et al., 2014). PAAS-normalized values after McLennan (1989).



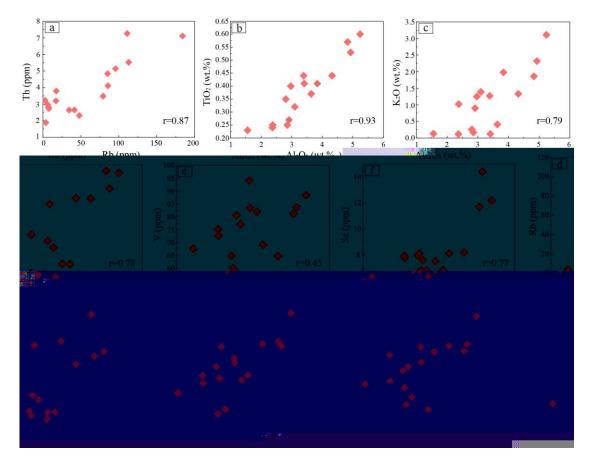


Fig. 6 (a) Bi-variation n between Rb and Th of Baijianshan BIF samples; (b-i) Bi-variation between Al_2O_3 and TiO_2 (b), K_2O (c), Rb (d), V (e), Sc (f), Cs (g), Cr (h) and REE (i) of the Baijianshan BIF. Data for bulk-rock BIF samples are listed in supplementary Table 2.

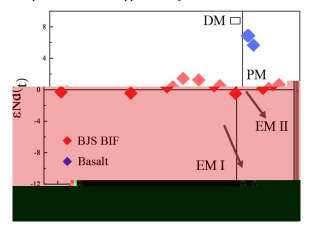


Fig. 7 $\,$ C $\,$ $^{87}Sr/^{86}Sr)_i$ diagram of BIF and basalt samples in Baijianshan.

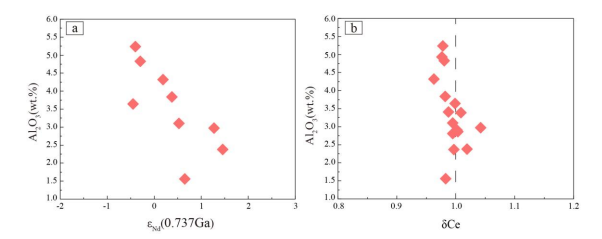


Fig. 8 C (t = 0.737Ga) vs. Al_2O_3 diagram of the Baijianshan BIF; (b) Ce/Ce^*_{PAAS} vs. Al_2O_3 diagram of the Baijianshan BIF.

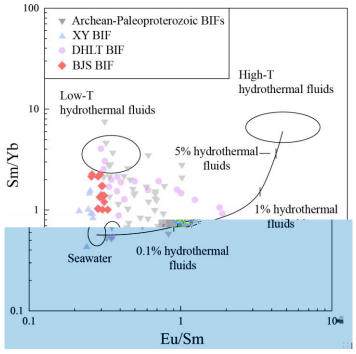


Fig. 9 Sm/Yb vs. Eu/Sm ratios illustrating two component mixing model of the hydrothermal fluid of the Baijianshan BIF (after Alexander et al., 2008). BJS-Baijianshan BIF; XY-Xinyu BIF (Li et al., 2014); DHLT-Dahongliutan BIF (Hu et al., 2017). Data sets of Archean-Paleoproterozoic BIFs are from Planavsky et al. (2010). Average compositions of high-T (> 300 °C) hydrothermal fluids, low-T (< 200 °C) hydrothermal solutions and Pacific seawaters were after Bau and Dulski (1999), Michard et al. (1983) and Alibo and Nozaki (1999), respectively.

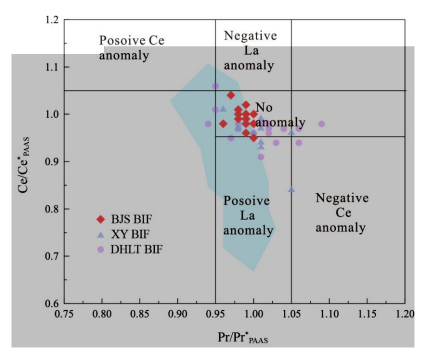


Fig. 10 Pr/Pr*_{PAAS} vs. Ce/Ce*_{PAAS} diagram for the BIF at Baijianshan (after Bau and Dulski, 1996). The light blue area represents the data of Archean-Paleoproterozoic BIFs. Data sets of other BIFs are same as those of Fig. 9.

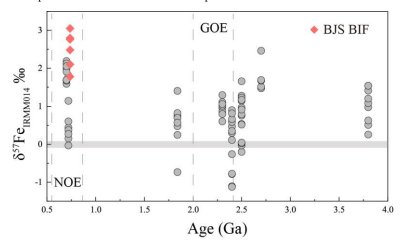


Fig. 11 I ⁵⁷Fe_{IRMM-014} of various ages BIFs in the world (Data of SW Greenland BIFs are from Dauphas et al.(2004); Manjeri Formation, Belingwe Belt, Zimbabwe BIFs are from Rouxel et al. (2005); Anshan-Benxi BIFs are from Li et al. (2012); Transvaal BIFs are from Johnson et al. (2003); Yuanjiacun BIFs are from Hou et al. (2014); Gongchangling BIFs from Li et al. (2014); Sanjiang BIF are from Yan et al. (2010) and Xinyu BIF are from Shen et al. (2008)).

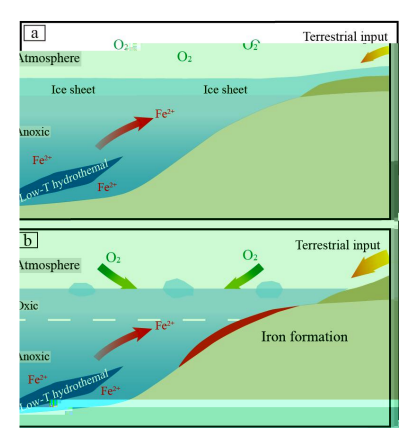


Fig. 12 Cartoon model of the formation of the Baijianshan BIF (see details in the text).

Appendix Table Captions

Appendix-Table 1 LA-ICP-MS U-Pb data for zircon from tuff in the first section of the Lapeiquan Formation

Appendix-Table 2 Geochemical compositions of the Baijianshan BIF

Appendix-Table 3 Sr-Nd isotopic data of the Baijianshan BIF

Appendix-Table 4 Fe isotopic data of the Baijianshan BIF

Supplementary Table 1 Zircon U-Pb age data of the tuff from the Baijianshan BIF

Spot	U ppm	Th ppm	Th/U	²⁰⁶ Pb/ ²³⁸	UAge	²⁰⁷ Pb/ ²³	⁵ UAge	²⁰⁷ Pb / ²⁰⁶ Pb		²⁰⁷ Pb/ ²³⁵ U		$^{206}\text{Pb}/^{23}_{}^{}^{}$	
2071-1	3079	831	0.27	1335	14	1360	21	0.0887	0.0012	2.8156	0.0427	0.2301	0.0025
2071-2	805	427	0.53	898	11	1154	20	0.1027	0.0014	2.1169	0.0369	0.1495	0.0018
2071-3	529	381	0.72	776	8	1009	18	0.0965	0.0016	1.7019	0.0307	0.1279	0.0014
2071-4	155	81	0.52	835	9	1350	22	0.1458	0.0022	2.778	0.0462	0.1382	0.0015
2071-5	184	101	0.55	751	8	1009	18	0.0999	0.0016	1.7022	0.0304	0.1236	0.0013
2071-6	297	252	0.85	1273	14								

Supplementary Table 2 Geochemical compositions of Baijianshan BIF

Sample	BJS-1	BJS-2	BJS-3	BJS-4	BJS-5	BJS-6	BJS-7	BJS-8	BJS-9	BJS-10	_
Major eleme	ents (%)										
Fe_2O_3	53.60	41.52	56.64	52.26	30.01	36.85	41.01	45.22	39.78	48.55	
SiO_2	29.91	38.96	26.81	31.20	40.92	40.79	36.97	41.77	43.66	37.59	
Al_2O_3	1.56										

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TFe ₂ O ₃	21.79	28.86	44.82	42.63	26.98	13.11	40.45	11.61	11.70	11.98
SiO_2	55.06	49.87	41.24	43.65	53.64	59.90	42.80	39.94	40.91	44.55
Al_2O_3	5.24	4.93	2.81	2.86	4.83	7.58	3.84	11.36	11.45	10.76
CaO	0.81	1.02	0.75	0.75	0.76	0.84	0.65	9.77	9.31	7.45
MgO	1.57	1.51	0.74	0.86	1.36	2.32	1.31	8.40	8.54	10.38
TiO_2	0.60	0.53	0.35	0.25	0.59	0.86	0.41	3.66	3.80	3.60
MnO	0.03	0.04	0.05	0.03	0.03	0.02	0.03	0.24	0.25	0.23
P_2O_5	0.20	0.35	0.49	0.40	0.23	0.19	0.54	0.46	0.43	0.38
K_2O	3.11	2.32	0.26	0.17	1.86	3.78	1.98	1.35	1.27	0.36
Na_2O	0.94	1.25	0.14	0.19	0.63	0.77	0.88	0.94	0.98	0.59
LOI	2.06	1.97	4.57	2.89	2.9	3.19	1.92	12.02	11.39	9.60
FeO	7.97	7.26	3.89	5.29	6.06	6.57	5.31	-	-	-
Total	99.38	99.91	100.11	99.97	99.87	99.13	100.12	99.75	100.03	99.88
Trace element	ts (ppm)									
Cr	118	81.0	59.7	10.5	86.2	112	50.9	214	214	306
Ni	15.6	14.5	12.1	11.0	15.6	12.3	12.6	162	166	217
Co	76.1	70.2	33.7	34.7	54.0	27.1	45.3	48.0	62.9	50.7
Sc	12.2	14.4	7.97	8.11	11.7	15.1	8.11	26.4	26.1	25.7
Ba	662	562	111	19.2	359	1219	385	877	864	296
Sr	45.0	53.3	45.8	26.1	32.6	37.0	48.9	205	209	150
Rb	43.0 111	95.6	7.36	6.13	113	184	85.6	34.4	32.2	12.3
		93.4	60.0	87.9	79.9	116		259	251	228
Zr	101						79.7			
Hf	2.63	2.37	1.23	1.50	2.11	2.99	1.92	5.35	5.74	6.00
Ta	0.88	0.75	0.37	0.36	0.64	0.84	0.53	2.24	2.33	2.14
Ga	7.95	7.63	4.39	4.63	7.12	10.3	5.97	18.3	17.2	18.8
Cu	27.6	21.6	8.12	3.56	21.4	36.0	12.4	-	-	-
Zn	62.1	40.0	26.0	23.8	35.3	37.0	29.6	-	-	-
Pb	7.80	8.24	3.54	5.32	7.64	15.4	5.75		-	-
U	0.47	0.69	0.39	0.40	0.72	0.82	0.37	1.45	1.09	0.92
Th	7.26	5.14	2.81	2.97	5.52	7.11	4.11	3.32	3.37	3.18
Nb	8.35	8.21	4.19	4.86	6.80	9.50	6.11	38.0	36.5	33.6
La	26.8	20.3	13.0	9.82	19.1	22.5	17.3	36.9	34.1	33.7
Ce	58.3	44.2	30.2	22.8	42.0	46.1	39.0	77.2	72.9	74.9
Pr	7.03	5.35	3.73	2.77	5.09	5.51	4.81	9.37	9.15	9.28
Nd	29.3	22.4	16.2	11.7	21.3	22.6	20.7	37.9	36.6	40.0
Sm	6.46	4.95	3.67	2.65	4.70	4.96	4.70	8.06	8.20	9.11
Eu	1.33	1.15	0.91	0.66	1.06	1.01	1.12	1.41	1.85	2.91
Gd	6.39	5.03	4.02	3.03	4.64	4.93	5.01	7.69	7.82	8.87
Tb	1.00	0.80	0.67	0.52	0.72	0.79	0.82	1.21	1.22	1.35
Dy	5.98	4.98	4.53	3.71	4.26	4.84	5.36	6.57	7.04	7.04
Y	28.5	24.9	27.9	22.0	20.1	26.1	31.9	34.8	34.0	32.1
Но	1.20	1.04	1.02	0.85	0.86	1.01	1.17	1.25	1.29	1.31
∥ Er	3.23	2.94	3.00	2.59	2.33	2.82	3.38	3.11	3.15	3.20
Tm	0.45	0.43	0.46	0.40	0.34	0.41	0.50	0.39	0.40	0.39
¹ Yb	2.87	2.93	3.06	2.76	2.20	2.69	3.36	2.28	2.36	2.28
Lu	0.43	0.45	0.49	0.45	0.33	0.41	0.53	0.34	0.32	0.32
G: :	150.77	116.95	84.95	64.71	108.93	120.58	107.75	193.68	186.4	194.67
Y/Ho	23.75	23.94	27.35	25.76	23.29	25.84	27.26	27.84	26.36	24.50
La/La*	1.06	1.07	1.09	1.02	1.06	1.09	1.10	1.14	1.06	1.14
Ce/Ce*	0.98	0.98	0.99	1.00	0.98	0.95	0.98	0.98	0.98	1.01
Y/Y^*	0.85	0.88	1.04	0.99	0.84	0.95	1.02	0.89	0.83	0.78
Pr/Pr*	1.00	1.00	0.99	0.99	1.00	1.00	0.99	0.97	0.99	0.95
Eu/Eu*	0.97	1.08	1.10	1.08	1.07	0.96	1.08	0.54	0.69	0.97
La _N /Yb _N	0.69	0.51	0.31	0.26	0.64	0.62	0.38	10.99	9.82	10.04

Sample	87Rb/86Sr	$^{87}Sr/^{86}Sr$	'	$(^{87}Sr/^{86}Sr)_i$	147Sm/144Nd	¹⁴³ Nd/ ¹⁴⁴ Nd	1	T _{DM} (Ga)	С
BJS-1	0.0433	0.717927	0.000004	0.7175	0.1457	0.512425	0.000006	1.63	0.65
BJS-4	4.2608	0.727751	0.000009	0.6829	0.1308	0.512394	0.000003	1.39	1.46
BJS-5	1.4305	0.728893	0.000008	0.7138	0.1378	0.512363	0.000003	1.58	0.19
BJS-7	2.4192	0.729423	0.000009	0.7040	0.1473	0.512376	0.000002	1.78	-0.45
BJS-8	2.8555	0.718681	0.000003	0.6886	0.1238	0.512351	0.000004	1.36	1.27
BJS-10	2.6390	0.723954	0.000005	0.6962	0.1344	0.512364	0.000002	1.51	0.53
BJS-12	7.0891	0.740795	0.000007	0.6662	0.1333	0.512311	0.000003	1.59	-0.40
BJS-16	9.9936	0.746329	0.000008	0.6412	0.1334	0.512317	0.000003	1.58	-0.30
BJS-18	5.0054	0.731954	0.000008	0.6793	0.1373	0.51237	0.000004	1.56	0.38
2702H1	0.4737	0.711034	0.000009	0.7061	0.1286	0.512662	0.000003	0.88	6.90
2702H2	0.4355	0.711156	0.000009	0.7066	0.1354	0.512694	0.000006	0.89	6.88
2702H3	0.2322	0.710822	0.000012	0.7084	0.1377	0.512644	0.000005	1.02	5.69

Suj	pplen	nentary	Table 4 Fe isotopic data of the Baijianshan BIF	
	-	57-	. 56-	

Sample 57Fe_{IRMM-014} 56Fe_{IRMM-014}

