- 1 A review of the potential for rare earth element resources from European red
- 2 muds: examples from Seydişehir, Turkey and Parnassus-Giona, Greece
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12 Abstract

13 Rare earth elements (REE) are viewed as 'critical metals' due to a complex array of 14 production and political issues, most notably a near monopoly in supply from China. 15 Red mud, the waste product of the Bayer process that produces alumina from bauxite, represents a potential secondary resource of REE. Karst-bauxite deposits 16 17 represent the ideal source material for REE-enriched red mud as the conditions 18 during formation of the bauxite allow for the retention of REE. The REE pass 19 through the Bayer Process and are concentrated in the waste material. Millions of 20 tonnes of red mud are currently stockpiled in onshore storage facilities across 21 Europe, representing a potential REE resource. Red mud from two case study sites,

22 one in Greece and the other in Turkey, has been found to contain an average of 23 approximately 1000 ppm total REE, with an enrichment of light over heavy REE. 24 Although this is relatively low grade when compared with typical primary REE 25 deposits (Mountain Pass and Mount Weld up to 80000ppm), it is of interest because 26 of the large volumes available, the cost benefits of reprocessing waste, and the low 27 proportion of contained radioactive elements. This work shows that around 12000 28 tonnes of REE exist in red mud at the two case study areas alone, with much larger 29 resources existing across Europe as a whole.

30 Introduction

31 Processing of the primary aluminium ore, bauxite, to produce alumina (Al₂O₃) using 32 the Bayer process produces a waste material known as red mud. Long term storage 33 of this is expensive and represents an environmental risk, as demonstrated by 34 disasters such as the red mud dam failure at Kolontár, Hungary which caused 35 extensive damage to neighbouring land and waterways and the death of nine 36 people (Gelencsér et al., 2011; Milačič et al., 2012; Anton et al., 2014). To mitigate 37 this risk, as well as to reduce cost, a number of studies have been undertaken to 38 identify possible uses for red mud (e.g. Klauber et al., 2011; Power et al., 2011; 39 Binnemans et al., 2013a).

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Annual global extraction of bauxite in 2013 (the last year for which figures are
available) was approximately 296 million tonnes (Mt), up 16% from 2012 (BGS,
2015). Annually, this results in the production of approximately 150 Mt of red mud

(IAI, 2014). Decades of work on the valorisation of red muds have not yet found a
significant alternative use for this voluminous waste product (Klauber *et al.*, 2011).
Only around 2 Mt (~1.6% of the global total) is currently used each year (Ritter,
2014), mostly in construction materials, as cement additives and in brick making
(Klauber *et al.*, 2011).

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50 The Bayer process produces virtually pure alumina (> 99% Al₂O₃) and therefore 51 almost all non-alumina 'impurities' from the original bauxite end up in the red mud 52 waste. These 'impurities' have been found to include significant amounts of 53 gallium, scandium and REE (Ochsenkühn-Petropoulou et al., 1995, 1996; Smirnov 54 and Molchanova, 1997; Akcil et al., 2013; Rusal, 2014; Borra et al., 2015). Ga and 55 the REE are classed as 'critical metals' (EC, 2014) i.e. those having limited global 56 distribution of production, no known substitutes, highly specialised applications 57 and being of economic significance (Graedel et al., 2014).

58 REE are generally divided into light REE (LREE) from lanthanum to samarium and 59 heavy REE (HREE) from europium to lutetium; Yttrium behaves as a HREE due to its 60 similar physical and chemical properties (Wall, 2014). HREE, followed by LREE, are 61 classified by the European Commission (EC, 2014) and British Geological Survey 62 (BGS, 2012) as the most 'critical' of all raw materials. They are deemed essential to the economies of Canada, Japan and the US among others (CREEN, 2013; 63 Humphries, 2013; Mitsubishi Electric, 2014). Annual global production of REE 64 65 (expressed as rare earth oxides (REO)), was approximately 100 170 t in 2013 (BGS,

66 2015). Of this, over 90% was supplied by China. This geospatial concentration of
67 production has raised concerns over security of supply (Hatch, 2012).

68 The REE are now essential to many green and modern technologies such as smart-69 phone screens, electric cars and wind turbines (Wall, 2014). This has led to 70 increasing global demand for REE with demand growth of 5.6% between 1970 and 71 2010 (Alonso et al., 2012). Present development of electric vehicles and wind 72 turbines relies heavily on dysprosium and neodymium for rare-earth magnets. 73 Future adoption of these technologies may result in large and disproportionate 74 increases in the demand for these two elements (Alonso et al., 2012). Economic 75 exploitation of REE from primary deposits, such as alkaline magmatic rocks, 76 carbonatites and ion adsorption clays, is dependent on the ratio of LREE to HREE in 77 the deposit and metallurgical characteristics of the REE present (Simandl, 2014). 78 Alternative sources of REE are becoming more important due to a range of factors 79 including increasing demand, a poorly developed REE recycling industry (Simand), 80 2014), a more volatile supply chain from China (Hatch, 2012), and the 'balance 81 problem', where demand for specific elements and the natural abundance of those 82 elements in deposits are 'unbalanced', resulting in excess supply of less in-demand 83 REE (Binnemans et al., 2013b; Binnemans and Jones, 2015). In the European Union 84 (EU) this has led to the seventh framework programme (FP7) funding of the EURARE 85 (European Rare Earth Resources, www.eurare.eu) project which aims to research and develop European resources and production of REE in a sustainable, 86 87 economically viable and environmentally friendly way. Europe in this context refers

88 to the EU-28, Liechtenstein, Iceland, Norway and Switzerland, and the candidate 89 and potential candidate countries. Russia and Ukraine are not included in this work. 90 Primary resources of REE have been identified and explored across Europe over the 91 past decades (Goodenough et al., 2016), such as at Norra Kärr, Sweden; Fen, 92 Norway; the Kvanefjeld and Kringlerne deposits in Greenland; and the Aksu Diamas 93 deposit in Turkey. However, none have so far gone into production. Attention has 94 also turned to secondary sources, mainly from the recycling or processing of waste 95 materials. Europe's red muds represent one such possible resource. The work 96 presented here aims to assess the REE potential of European red muds. The study 97 is based on two case examples, Seydisehir, Turkey and Parnassus-Giona, Greece, 98 where the parent karst-bauxites have been identified as REE-bearing (Ochsenkühn-99 Petropoulou et al., 1991; Laskou and Andreou, 2003; Karadağ et al., 2009), and the 100 red muds are stored in accessible 'dry' piles or tailing ponds. These case studies are 101 put in the context of a review of European red mud resources where resource 102 estimates of available REE in European red mud storage facilities are presented.

103 Bauxites and alumina in Europe

Within Europe, bauxite is currently extracted in six countries: Bosnia and
Herzegovina, France, Greece, Hungary, Montenegro and Turkey. Production in
2013 equalled almost 3.5 Mt, approximately 1% of global production (BGS, 2015).
Historically, bauxite has been exploited more widely across Europe, particularly in
the Mediterranean region (Figure 1), representing a significant proportion of global
bauxite production (~15% in 1974 (IGS, 1978)). There are three main categories of

110 bauxite deposit (Bárdossy, 1982): 1. Lateritic bauxite deposits, which derive from in 111 situ weathering of aluminosilicate rocks; 2. Karst bauxite deposits, which occur in 112 the karst topography of limestone and dolomite and comprise aluminosilicate 113 residues that have been transported from either proximal or distal locations; 3. 114 Tikhvin-type deposits, which are laid down on aluminosilicate rocks but have no 115 genetic relationship to them. Karst bauxite deposits can be subdivided further into 116 six main types (Bárdossy, 1982; Bosák et al., 1989), of which the 'Mediterranean 117 type' karst bauxite deposits are of the most interest in Europe due to the amount 118 of REE present. Red mud waste from these Mediterranean bauxite types is the 119 focus of this paper.

120 Formation of karst bauxites begins with muddy sediments being washed into 121 hollows in karst topography. These sediments are subsequently bauxitised through 122 deep lateritic weathering. Bauxitisation is facilitated by moderate temperatures 123 and through-put of mildly alkaline fluids, which allow silica to be incongruently 124 dissolved from minerals such as feldspar and kaolinite leaving an aluminium-rich 125 residue (Gow and Lozej, 1993; Bland and Rolls, 1998). Alongside this, Al may 126 accumulate from a variety of sources including the insoluble residue of limestone, 127 other aluminosilicate material (volcanic ash and clay layers) within the limestone, 128 or the erosion, transportation and weathering of aluminosilicate rocks. The main 129 Al-bearing minerals in bauxite are the monohydrates boehmite ((y-AlO(OH)) and 130 diaspore ((α -AlO(OH)), and the trihydrate gibbsite (Al(OH)₃). The main gangue 131 minerals include hematite (Fe₂O₃), goethite (FeO(OH)), anatase (TiO₂), kaolinite 132 $(Al_2Si_2O_5(OH)_4)$ and minor quartz (SiO_2) .

133 The process of bauxitisation progresses from the top downwards and laterally 134 outward following the direction of drainage (Bárdossy, 1982). Karst-bauxite 135 typically forms under moderate temperatures in subtropical climates usually with 136 an average mean temperature of 26°C (Bárdossy, 1979). Good drainage of slightly 137 alkaline fluids through the developing bauxite is essential for formation. It has been 138 shown that the major element chemical composition of bauxites varies from one 139 region to another and within individual deposits (Akinic and Artir, 2008). The 140 formation conditions also have an important influence on the REE content and 141 distribution within the bauxite. The sources and formation conditions of selected 142 European karst-bauxite deposits, and the implications that these may have for REE 143 content, are discussed below. The size and grade of bauxite deposits is dependent 144 on the duration of the weathering (Bogatyrev et al., 2009), but the effect of the 145 duration of weathering on the REE content is unclear.

146 Alumina production from bauxite has been on-going in Europe since the 147 development of the Bayer process over 125 years ago. To separate alumina from 148 bauxite, the ore is digested in a hot sodium hydroxide solution to convert the 149 alumina to aluminium hydroxide, which dissolves in the hydroxide liquor. The other 150 components of bauxite do not dissolve. The solution is clarified by filtering off the 151 solid impurities, which forms the bauxite residue or red mud (Hind et al., 1999). The 152 composition of red muds depends on the nature of the parent bauxite, the mining location and the parameters of the Bayer process (Hind et al., 1999). 153 154 Mineralogically, red mud is mainly composed of red coloured iron oxides in the 155 form of hematite (Fe_2O_3) and maghemite (γ - Fe_2O_3), aluminium oxides in the form

156 of diaspore (α -AlO(OH)) and gibbsite (Al(OH)₃), and silicon, calcium, sodium and 157 titanium oxides such as calcite (CaCO₃), silicon calcium aluminate hydroxide 158 $(Ca_3Al_2(SiO_4)(OH)_8),$ perovskite (CaTiO₃), rutile (TiO₂) and cancrinite 159 (Na₆Ca₂Al₆Si₆O₂₄(CO₃)₂.2H₂O (Atasoy, 2007). Red muds typically have a density of 160 around 2.75 g/cm³ and a pH of > 12.

161 Red muds are produced at a rate of approximately 150 Mt per year globally. There 162 are now approximately 60 alumina refineries worldwide outside China using the 163 Bayer process, of which 10 are located in Europe (Figure 2). An additional 49 164 refineries were operating in China in 2011 (IAI, 2014). European refineries produce 165 alumina from both European and imported bauxite in order to supply the needs of 166 the European market. The importation of bauxite from a wide range of sources 167 makes it difficult to assess the REE content of the resulting red mud waste. In 168 Europe, the largest alumina refinery is Aughinish in the Republic of Ireland, which 169 has an output of 1.93 Mt of alumina per annum. Each tonne of alumina produced 170 results in the formation of between 0.7 and 2 tonnes of red mud (IAI, 2014). 171 Assuming an average overall ratio of 1:1.5 alumina to red mud (Hamada, 1986; 172 Power et al., 2011), the Aughinish plant is likely to produce almost 3 Mt of red mud 173 per annum.

Worldwide, an estimated 2.7 billion tonnes (Bt) of red muds have been produced
since the development of the Bayer process (Power *et al.*, 2011). These have been
treated in a variety of ways, including storage in onshore lagoons and direct marine
disposal (Power *et al.*, 2011). Historically, large quantities of the waste material

were directly dumped into the sea, however, this practice has been regulated under the 'Convention on the Prevention of Marine Pollution by Dumping of Wastes and Other Matter (IMO, 1972)', the 'London Convention' for short. The practice of disposal at sea is expected to be completely abandoned by 2016 (IAI, 2014). Most commonly, red muds are stored in settling ponds onshore in closed cycle disposal systems; no alumina refineries built after 1970 are known to employ marine disposal of red muds (Power *et al.*, 2011).

185 Statistics on European alumina production since 1972 have been gathered from the 186 British Geological Survey's World mineral statistics database and are shown in 187 Figure 3. Greek production before 2009 is not included, as before then red muds 188 were disposed of at sea. The quantities of red muds likely to have been produced 189 annually in Europe and stored onshore since the introduction of the 'London 190 Convention' have been estimated as at least 200 000 t, using a minimum ratio of 191 1:0.7 tonnes of alumina to residue and a maximum ratio of 1:2 tonnes of alumina 192 to residue, with an average ratio of 1:1.5 tonnes of alumina used (Figure 3).

193 Rare earth elements in bauxite

The characterisation of REE-containing minerals in bauxites was pioneered in the 195 1970s (Bárdossy and Pantó, 1973; Maksimović and Roaldset, 1976; Maksimović and 196 Pantó, 1978; Vukotić, 1983). Of historical note was the first identification of 197 secondary minerals of the bastnäsite group in the San Giovanni Rotondo bauxite in 198 Italy (Bárdossy and Pantó, 1973). Since then, both authigenic and detrital REE-199 bearing minerals have been found in bauxites. The most common authigenic REE-

200 bearing mineral is hydroxylbastnäsite-(Nd) (Nd(CO₃)(OH)), while monazite 201 ((Ce,La,Nd,Th)PO₄)) is the most common detrital REE-bearing mineral (Maksimović 202 and Pantó, 1996). In some occurrences the presence of detrital minerals can result 203 in high REE values, e.g. at Nagyharsány, a Hungarian monazite-bearing bauxite 204 which has total REE (TREE) of up to 10000 ppm (Bárdossy et al., 1976). The leaching 205 of monazite in this deposit led to the precipitation of authigenic bastnäsite-group 206 minerals near the base of the bauxite (Bárdossy et al., 1976; Maksimović and Pantó, 207 1996). In an example from the Southern Apennines of Italy, detrital monazite, 208 xenotime (YPO₄), and supergene LREE-enriched phosphates (high Ce, low La-Nd, 209 Ca minerals and florencite-type ((REE)Al₃(PO₄)₂(OH)₆) minerals) derived from the 210 weathering of monazite were detected in bauxite using SEM-EDS and QEMSCAN® 211 (Mondillo et al., 2011; Boni et al., 2013). In addition, bauxites from the Southern 212 Appenines (Italy) (Boni et al., 2013) and Nurra (Sardinia) (Mameli et al., 2007) 213 contain authigenic high-Ce, Ca-bearing fluorcarbonate minerals (REE₂Ca(CO₃)₃F₂). 214 Parisite $(Ca(Ce,La)_2(CO_3)_3F_2)$ has previously been identified in the basal parts of the 215 Spinazzola bauxite in Italy (Mongelli, 1997; Mongelli and Acquafredda, 1999), 216 however, recent analysis suggests that parisite occurs homogeneously throughout 217 the deposit (Mongelli et al., 2014).

Fractionation of LREE and HREE has been noted in many Mediterranean bauxites
(Maksimović and Pantó, 1991; Mongelli, 1997; Laskou and Economou-Eliopoulos,
2007; Karadağ *et al.*, 2009; Boni *et al.*, 2013), resulting in vertical separation of REE
in bauxite profiles (Boni *et al.*, 2013). Fractionation of Ce, which behaves differently
to the other REE, in particular in the uppermost parts of bauxite, is as a result of the

223 supergene oxidation of Ce⁺³ to Ce⁺⁴. This allows precipitation of cerianite 224 ((Ce⁴⁺,Th)O₂) to occur, with resultant Ce-depleted fluids percolating through the 225 system. Scavenging by iron oxide phases, such as goethite (Boni et al., 2013), of Ce-226 depleted fluids results in further fractionation of REE between the ooids (concentric 227 particles of bauxite with a diameter of between 100 and 1000 µm (Bárdossy, 1982)) 228 and matrix of the bauxite in basal parts of the bauxite (Mongelli, 1997). Subsequent 229 dissolution of cerianite under acidic conditions and per descensum transport of REE 230 can result in the precipitation of LREE-rich minerals, such as parisite, at the base of 231 the deposit (Mongelli, 1997; Mameli et al., 2007). Parisite in particular precipitates 232 under alkaline conditions, such as those found at the base of the bauxite due to the 233 alkaline aquifers in the underlying limestone. Parisite precipitation throughout a 234 bauxite deposits, such as in the Spinazzola bauxite, can be explained by cyclical 235 changes in water table levels and subsequent changes in Eh and pH (Mongelli et al., 236 2014).

237 Preferential uptake of REE by both iron and manganese mineral phases has been 238 identified in other systems such as in marine ferromanganese deposits (Ohta and 239 Kawabe, 2001) and in river estuary systems where iron oxyhydroxide adsorption of 240 LREE in particular occurs (Marmolejo-Rodríguez et al., 2007). The alkaline 'pH 241 barrier' created by the underlying limestone aquifer influences the stability of the 242 REE-carbonate complexes that form at the base of the deposit (Johannesson et al., 243 1995, 1996). As the stability of these complexes increases with atomic number, the 244 HREE are preferentially retained in solution as carbonate complexes. The 245 distribution of the REE along this 'pH barrier' is heterogeneous and they can be

concentrated in lenses and micropores, space fillings and microveins (Maksimovićand Pantó, 1996).

248 Rare earth elements in red muds

249 REE are concentrated into red muds through the Bayer process (Wagh and Pinnock, 250 1987) which results from the association of REE with iron and titanium phases 251 which pass through unchanged (Derevyankin et al., 1981). To illustrate this, red 252 muds from lateritic-type Timan bauxites in Russia have been shown to contain up 253 to 90% of the Nb, Sc and REE present in the original bauxite (Klyucharev et al., 254 2013). Similarly, Ochsenkühn-Petropoulou et al. (1994) and Wagh and Pinnock 255 (1987) have demonstrated up to two-fold increases in REE, including Sc and Y, in 256 red muds when compared to the bulk chemistry of the Greek and Jamaican source 257 bauxites, respectively.

The minerals known to occur in red muds include some that are produced during the Bayer process, e.g. desilicification products such as sodalite (Na₈(Al₆Si₆O₂₄)Cl₂), cancrinite (Na₆Ca₂Al₆Si₆O₂₄(CO₃)₂.2H₂O) and Na-aluminosilicate, and calcite (Atasoy, 2007). However, most minerals are unaltered residual phases from the original bauxite such as monazite, bastnäsite, hematite (Fe₂O₃), diaspore ((α -AlO(OH)), gibbsite (Al(OH)₃) and quartz (SiO₂)(Power *et al.*, 2011).

The mineralogy and petrology of REE-bearing minerals in red muds are relatively poorly constrained (Ochsenkühn-Petropoulou *et al.*, 1996; Borra *et al.*, 2015). Scavenging of REE by iron oxyhydroxides is observed in bauxites (Mongelli, 1997;

267 Mongelli and Acquafredda, 1999; Boni et al., 2013), however, the distribution of 268 REE phases in red muds is less well understood. An association between REE and 269 Sc with iron oxides in red muds has been observed through leaching studies (Borra 270 et al., 2015 and references therein). Sc is heterogenously distributed through the 271 red muds, being mainly contained within detrital mineral phases, such as the Sc-272 enriched zircons found in bauxite (Boni et al., 2013). It may also either be adsorbed 273 onto the surface of iron oxide phases or enriched in the outer layers of iron oxide 274 particles (Borra *et al.*, 2015), which may be the same in the case of REE.

275 The Bayer process involves many steps that affect the characteristics of the red 276 muds and hence potentially the REE distribution within the residue. Milling of the 277 bauxite ore initially changes the particle size and surface area of the minerals, which 278 has an impact on the settling rates within the red mud lagoons, and can also affect 279 chemical properties such as adsorption and dissolution of minerals. It is not possible 280 to apply a general enrichment factor for REE to red muds; it is necessary to 281 investigate the REE enrichment in each red mud locality and compare this, where 282 possible, with the parent bauxite. Further work to define the distribution of REE 283 within red muds is necessary in order to develop a clearer understanding of both 284 the distribution of REE and the phases that host REE within red muds.

Recent advances in the valorisation of red muds for minor metals include the development of Rusal's pilot plant in the Urals, Russia, which is capable of producing 2.5 t/a primary scandium oxide concentrate (RUSAL, 2014). Binnemans *et al.* (2013a) give an overview of current methods of REE extraction from red muds.

289 Bench scale extraction of REE from dehydrated red muds using imidazolium-based 290 ionic liquids is being researched through the EURARE project at National Technical 291 University of Athens (NTUA) (Bourbos et al., 2014; Davris et al., 2014) while 292 leaching experiments are on-going at KU Leuven, Belgium (Borra et al., 2015). The 293 commercial extraction of REE from red muds has been further advanced by the 294 granting of a patent in both Canada and the US to Orbite Aluminae for the 'Orbite 295 Process', (Patent No. 14/371,364, Orbite, 2015), which uses red muds as a 296 feedstock for a new beneficiation stream that aims to extract REE and other minor 297 metals along with alumina, magnesium oxide and titanium dioxide from the 298 residue. It is thus clear that red muds have the potential to be a source for REE for 299 the European economy.

300 European red muds, case studies

301 The Mediterranean region has a complex geological history shaped most recently 302 by the closure of the Lower Palaeozoic to Cenozoic Palaeotethys and Neotethys 303 oceanic system which existed between the continents of Eurasia and Gondwana 304 (Robertson and Mountrakis, 2006). Continental collision resulted in the formation 305 of the Dinarides-Hellenides-Taurides orogenic belt, which extends across the 306 Eastern Mediterranean region, and which includes many carbonate units that 307 provide the karst topography in which karst bauxites have formed. The regional 308 geology is summarised below briefly to set the geological context for the bauxite 309 precursors of the case study localities.

310 Sampling and analytical methodology

311 Four samples of red muds (14/T/15-19) were collected at a depth of approximately 312 20 cm from different locations within the waste ponds at the ETİ Alüminyum S.A. 313 processing plant in Seydisehir, Turkey (inset Figure 4). The samples comprised 300 314 g of wet red muds; each individual sample consisted of three subsamples, which 315 were later recombined and dried. Preparation and analysis of the samples was 316 carried out at the British Geological Survey, Keyworth, Nottingham. Samples were 317 fused with sodium peroxide and 0.2 g of the resulting glass digested with dilute HCl 318 acid and then, HF acid. Dilute sample solutions (in 1% HNO₃+ 0.5% HCL) were then 319 analysed for 56 trace elements in an Agilent 7500CX series ICP-MS. Data for a 320 duplicate analysis of sample 14/T/16, for all elements, were within 0.18%. Data for 321 international standards (AGV-2 (andesite produced by USGS), BCR-2 (Columbia 322 River Basalt produced by USGS) and JR-2 (produced by Geological Survey of Japan)) 323 were always within 6% of published values.

324 Samples of Greek red muds were collected from dewatered stockpiles at the 325 Aluminium S.A. alumina refinery in Agios Nikolaos. Three samples, equating to 326 approximately 5 kg in total, were taken at three different locations (13/Gr/A, 327 13/Gr/B and 13/Gr/Z) across the area of the stockpiled residue. Each of the samples 328 represent red mud residue from different deposits, mined at different times. 329 Preparation of the samples was carried out at the Camborne School of Mines, 330 Penryn, Cornwall. Five grams of each sample were analysed for 31 trace elements 331 (LF100 analysis) by ICP-MS, following lithium borate fusions at Bureau Veritas 332 Minerals Laboratories (BVML), Canada. International standards, BX-N bauxite 333 (reference material produced by ANRT (Association Nationale de la Recherche

Technique)) and STD SO-18 were used by BVML. Values obtained were within 2.7%
of published values. Data from blank samples were checked for contamination and
detection limits. Duplicate analyses (n = 1) of, sample 13/Gr/B were within ± 0.5%.
Results are presented in Table 1, see supplemental data for duplicate, corrections
and certified reference material data.

339 Case studies: red muds from Seydişehir, Turkey and Parnassus-Giona, Greece

340 Case study one: red muds from Seydişehir, Turkey

341 Geology

342 The Anatolide-Tauride terrane is one of three main terranes into which 343 Turkey is subdivided (Sengör and Yılmaz, 1981). The terrane shows Gondwana 344 affinities but was separated from the main mass of Gondwana by the southern 345 branch of Neo-Tethys in the Triassic. Upper Triassic to Upper Cretaceous carbonate 346 deposition formed several thousand metres of thick shallow marine carbonate 347 deposits on a passive margin (Okay, 2008), which have subsequently been 348 overthrust by Upper Cretaceous limestone and ophiolitic material. The karst 349 topography formed on the carbonates in this region hosts a large number of 350 bauxite, Al-laterite and manganese deposits which are mainly Upper Cretaceous to 351 Cenozoic in age. The Mortaş and Doğankuzu Mediterranean-type karst bauxites are 352 located in the central Tauride Mountains, close to Seydisehir, and are hosted in 353 fossiliferous limestones of the Upper Cretaceous Mortas Formation (Figure 4) 354 (Temur et al., 2009).

355 Both the Doğankuzu and Mortas bauxites are mined to provide feedstock for the 356 alumina refinery at Seydişehir. The Mortaş deposit is lens shaped, 950 m in length 357 and 350 m wide, and the adjacent Doğankuzu bauxite elongate, with a strike length 358 of approximately 900 m and a width of 50 m (Karadağ et al., 2009). The deposits 359 occur between 1500 and 2000 m above sea level. The bauxites can be subdivided 360 into four horizons: massive bauxite, oolitic bauxite, breccia-bearing bauxite and 361 earthy bauxite. The main Al-bearing mineral is boehmite, with minor diaspore and 362 anatase, while the main accessory minerals are hematite, kaolinite and goethite. 363 REE are most enriched in the massive and oolitic bauxite horizons relative to the 364 hosting limestone (Karadağ et al., 2009). Bauxite deposition is thought to have 365 resulted from mass flow triggered by tectonic activity with transport of weathered 366 'bauxitic soil' over short distances into depressions and sinkholes in the limestone 367 (Öztürk et al., 2002). The bauxite deposits are between 1 and 40 m thick, and are 368 overlain by thin layers of claystone and argillaceous limestone. These are in turn 369 overlain by c. 50 m of Santonian massive limestone, with an overlying thick cover 370 of bioclastic limestone of Upper Cretaceous age and Eocene limestone and flysch. 371 The bauxites therefore must have formed in a depositional hiatus during the Upper 372 Cretaceous (Öztürk et al., 2002).

373

Bauxite deposits and red muds

The Seydişehir bauxite deposits of Mortaş and Doğankuzu have a total resource of 26.3 Mt high-alumina boehmite bauxite at 55–67% Al_2O_3 , of which approximately 6 Mt had been mined by 2002 (Öztürk *et al.*, 2002; Horkel, 2010).

377 These, the two largest bauxite open pits in Turkey, supply the only aluminium 378 smelter in the country (Horkel, 2010), currently operated by ETi Alüminyum. 379 Bauxite is mined from April to October annually and stockpiled at their plant in 380 Seydişehir; extraction during the winter is suspended due to the inclement weather 381 in the mountains. The extracted bauxite from the two mines is blended to create a 382 plant feed with aluminium content suitable for processing at Seydişehir. Annual 383 production is approximately 800 000 t of bauxite (BGS, 2015), generating 384 approximately 1.5 Mt of red mud. Approximately 10 Mt are stockpiled in the tailing 385 ponds. The red muds collected in the tailing ponds are waste material from both 386 bauxite pits. It is not therefore possible to determine the relative contribution of 387 REE from each of the deposits to the red mud samples.

388 Case study two: red muds from Parnassus-Giona, Greece

389 Geology

390 Greece is geologically subdivided into several tectonic zones, representing 391 an amalgamation of terranes accreted over an extended period from the Lower 392 Palaeozoic to the Mesozoic. The External Hellenides resulted from the Early 393 Tertiary destruction of a Neotethyan oceanic strand known as the Pindos Ocean, 394 which led to the collision between the Apulian and Pelagonian microcontinents 395 (Doutsos et al., 2006 and references therein). The Parnassus-Giona zone is located 396 within the External Hellenides, which is limited to Central Greece and consists 397 almost entirely of limestone and dolomite of Mesozoic age (Doutsos et al., 2006).

398 The Parnassus-Giona tectonic zone is characterised by a nearly continuous 399 sequence of Upper Triassic to Upper Cretaceous epicontinental reef-like 400 carbonates. At the time of formation the Parnassus-Giona zone was a shallow 401 lagoon, with both freshwater and seawater ingress due to changing tectonic and 402 climatic conditions (Valeton et al., 1987). Several phases of uplift, with associated 403 marine transgressions and regressions, permitted the creation of multiple bauxite 404 horizons during the Jurassic and Cretaceous. The materials which formed the 405 bauxites in this area are thought to have been transported by sedimentary 406 processes over a calcareous terrain for a distance of over 30 km (Petrascheck, 407 1989). The bauxite horizons, termed B1-B3, from oldest to youngest, formed during 408 the Upper Jurassic to Middle Cretaceous (Figure 5) (Valeton et al., 1987, Laskou et 409 al., 2010). These bauxite deposits comprise lenses, pockets and irregular masses. 410 The economically most important deposits occur in the B3 horizon which is laterally 411 extensive as a continuous layer with a thickness of between one and ten metres (Petrascheck and Mack, 1978; Valeton et al., 1987; Melfos and Voudouris, 2012; 412 413 Tsirambides and Filippidis, 2012; Laskou and Economou-Eliopoulos, 2013). REE-414 bearing minerals occur in very small concentrations within the bauxites (Valeton et 415 al., 1987). The mineral grains are small (between 1 and 10 μ m) and can be 416 authigenic, detrital, or material from weathered parent lithologies (Bárdossy et al., 417 1976). LREE-bearing minerals include phosphates, such as detrital rhabdophane-418 (Ce) ((Ce,La)PO₄·(H₂O)) and florencite-(Ce) (CeAl₃(PO₄)₂(OH)₆), whereas HREE-419 bearing minerals are Y-phosphates such as detrital churchite (YPO₄.2(H₂O)) and 420 xenotime (Laskou and Andreou, 2003). Monazite and secondary bastnäsite-group

421 minerals, such as hydroxylbastnäsite-(Nd) and hydroxylbastnäsite-(La)
422 ((La)CO₃(OH,F)), and parisite-group minerals occur as micropore and fissure filling
423 aggregates (Lymperopoulou, 1996; Maksimović and Pantó, 1996; Gamaletsos *et al.*,
424 2007).

425 Coal layers, with a maximum thickness of 50 cm, are found in some locations just 426 above the B3 horizon, covered by limestone of Upper Cretaceous age forming the 427 thick layers of the Parnassus-Giona Unit (Kalaitzidis et al., 2010). Oxidation of pyrite 428 inclusions in the coal resulted in the formation of acid fluids that percolated 429 downwards to cause bleaching of the upper parts of the underlying bauxite 430 (Kalaitzidis et al., 2010). This may have impacted on the distribution of REE within 431 the bauxite, as under acidic conditions REE are easily weathered from aluminosilicates (Nesbitt, 1979; Fleet, 1984; Karadağ et al., 2009). 432

433 Bauxite production and red muds

434 The karst bauxite deposits of Greece are among the world's most important 435 sources of bauxite for non-metallurgical products such as for chemicals and fused 436 alumina. The main exploitable deposits are those in the Parnassus-Giona zone, with indicated reserves of 100 Mt (O'Driscoll, 2011). Bauxite production is 437 predominantly from underground mining of the B3 horizon, with less from the 438 439 deeper B2 horizons. The B1 horizon is currently not being exploited due to its 440 greater depth. Alumina is refined at the Aluminium S.A. site using the Bayer process 441 and the resulting waste red muds (700000 tpa (Anagnostou, 2010)) are dewatered

and stockpiled onshore. Deposition of dewatered red mud started in 2009; theapproximate total volume accumulated to end 2015 is 5 Mt.

444 Bauxite REE geochemistry

445 Previous work on the Mortas bauxite from Turkey (Karadağ et al., 2009; Hanilçi, 446 2013) and the Parnassus-Giona bauxites from Greece (Ochsenkühn-Petropoulou et 447 al., 1991, 1994, 1995; Lymperopoulou, 1996; Valeton et al., 1987; Laskou, 1991, 448 2005; Laskou et al., 2010; Laskou and Economou-Eliopoulos, 2007, 2013), shows 449 that they are enriched in the LREE relative to the HREE (Figure 6; Table 1). The 450 Mortaş bauxites show a clear negative europium (Eu) anomaly on a chondrite-451 normalised plot (Figure 6), whilst the Greek bauxites only display a small negative 452 Eu anomaly. These anomalies are likely to be inherited from the bauxite protoliths. 453 The Greek bauxites show a positive Ce anomaly that is not present in the Mortas 454 bauxite. The Ce anomaly could be due to samples being collected from higher, near 455 surface, relatively oxidising conditions where Ce is dominantly present as Ce⁴⁺, rather than Ce³⁺, and therefore fixed in oxides. Laskou and Andreou (2003) also 456 457 identified a positive Ce anomaly in samples from the upper (B3) and middle (B2) 458 bauxite horizons. Overall the Mortas bauxite is more enriched in the REE (614 ppm 459 TREE (Karadağ et al., 2009)) than the Greek bauxite (B3 horizon averages from 417 460 ppm (Valeton et al., 1987; [Economopolou-Kyriakopoulou, 1991]; Laskou, 1991, 2005; Ochsenkühn-Petropoulou et al., 1991; Laskou and Economou-Eliopoulos, 461 2007, 2013) to 527 ppm TREE (mixed bauxite from the B3 and B2 horizons sampled 462 463 at the processing plant) (Ochsenkühn-Petropulu et al., 1994, 1995)); again this

464 could be due to sampling bias as the Greek samples were collected mostly from
465 higher horizons. Published data for karst-bauxite deposits from Italy (Boni *et al.,*466 2013) and Turkey (Karadağ *et al.,* 2009; Hanilçi, 2013) show similar trends in REE
467 content (Table 1), displaying relative enrichments in the LREE, particularly for Nd
468 and Ce.

469

470 Results: rare earth element content of red muds

471 All the sampled red muds show moderate enrichment in TREE (by up to 1.9 times, 472 Figure 6) when compared with the original bauxite (Figure 6; Table 1), with the 473 exception of 14/T/15. Small negative Eu anomalies in all red mud samples are likely 474 to have been inherited from the bauxite protolith. Small positive Gd anomalies are 475 also present in the red muds from both Greece and Turkey, but only occur in one 476 bauxite sample (Greece B3 horizon). The values for REE in the parent bauxites are taken from the literature and will not necessarily correspond exactly to the specific 477 478 red mud sample being analysed.

Turkish red mud samples 14/T/16, 14/T/18 and 14/T/19 have an average content of 1 090 ppm *TREE*, and are all relatively enriched in REE compared with the published values for the parent bauxites (Figure 6; Table 1). Sample 14/T/15, however, shows an overall depletion in REE relative to the Mortaş bauxite (*TREE* 150 ppm). This may be because this sample was taken from the far west end of the pond (inset Figure 4), where the red mud cover is thin and therefore meteoric waters may have leached out the REE, or alternatively, the sample may have come

486 from an unusually low REE bauxite feedstock. Turkish red mud samples 14/T/15-16 487 and 14/T/19 show a positive Ce anomaly, whilst sample 14/T/18 shows a negative 488 Ce anomaly. This could be due to the oxidation state of Ce in the bauxite horizon 489 from which the waste was derived. Sample 14/T/18 also has higher total HREE 490 content, 356 ppm compared with an average of 208 ppm, which could also be 491 explained if it was sourced from deeper within the bauxite. Within the bauxite 492 profile, HREE complexes are more stable in the more alkaline conditions that occur 493 at depth during bauxite formation (Johannesson et al., 1995, 1996; Mameli et al., 494 2007).

495 Greek red mud samples show similar REE patterns to the Turkish samples though 496 they are generally less enriched in REE (Table 1; Figure 6). They also show a more 497 homogeneous spatial distribution of REE, even though samples are from different 498 mines within the B3 and occasionally the B2 layers. TREE values range between 800-499 900 ppm (up to 1000 ppm where Sc is included). All samples are characterised by 500 Ce, La, Nd and Y enrichments. Distinct positive Ce and negative Eu anomalies are 501 visible with minor enrichments in Tb and Gd also present. It should be noted that 502 the samples have been individually analysed by slightly different methods and in 503 different laboratories and as such are not directly comparable. However, these are 504 representative samples from two large red mud storage facilities and so can be 505 used to give an insight into European REE resources from alternative sources.

506

507 Resource potential

Whether red mud in Europe represents a potential resource for REE will largely depend on whether proven bench- and pilot-scale processing and beneficiation techniques for this material can be up-scaled for production. For a detailed overview of these, readers are directed to Binnemans *et al.*, 2015. In addition, a clear understanding of the volume of red muds present in Europe will aid in assessing whether these resources are viable. The remaining discussion will focus on this second aspect.

515 Table 2 utilises the data from Figure 3 and Table 1 to estimate TREE to calculate 516 average total contained REE, Nd and HREE in the red muds of Europe. Taking an 517 average value of 1000 ppm REE in red mud gives a range of resource potentials in 518 Europe. If the REE were to be extracted at an efficiency of 50% (Ochsenkühn-519 Petropoulou et al., 1996), then the minimum quantity of available REE would be 100 520 000 t, while at a maximum there are approximately 270 000 t REE in European red 521 muds. 425 Mt of red mud (see average cumulative figure in Table 2) at 1000 ppm 522 REE would contain 430000 t of REE, which at 50% recovery could produce 210000 t 523 REE. These figures are based on stored red mud stocks produced from alumina 524 refining in Europe since 1972, and as such, are a conservative estimate for the 525 potential resources available from red muds in Europe.

In a global context, around 150 Mt of red muds are currently being produced per year with up to 150000t of contained REE going to waste ponds. This equates to approximately 172 500 t REO, using 1.15 as an average element-to-stoichiometric oxide conversion factor. Annual global production of REE, expressed as tonnes of REO, was approximately 100 170t in 2013 (BGS, 2015). It is thus clear that there is potential for red muds to make a substantial contribution to the global supply of REE.

533 Taking neodymium as an example, the potential Nd resources in red muds across 534 Europe is approximately 20000 - 45000 t. This equates to greater than four years of 535 global production, calculated at approximately 23 000 t per year (2011 data) (EC, 536 2013 and references therein). The figures from Europe as a whole indicate that 537 HREE (Eu to Lu +Y) resources in red muds could be as much as 60 000 t. With the 538 development of selective beneficiation techniques it may be possible to prevent 539 the oversupply of LREE as is the current market situation (in 2015), and 540 preferentially produce the more in-demand elements such as neodymium and 541 dysprosium. An important caveat is that these figures represent red mud waste 542 from all bauxite types processed in European refineries. Annual importation of 543 bauxite into Europe equalled more than 13.5 Mt in 2013, (the last year for which 544 figures are available) (BGS, 2015), with major import partners including Guinea and 545 Brazil (data from UN Comtrade Database, accessed May 2015). Major bauxite 546 deposits in these countries are from lateritic-type bauxites rather than karst-type 547 bauxites. Taking this into consideration, further work on characterising the REE 548 content in lateritic bauxites in Europe and in the known red mud storage facilities 549 across Europe would further refine any initial resource estimates.

550 5 Mt of red muds are currently 'dry' stacked at Aluminium S.A. in Greece while at 551 least 10 Mt of red muds are contained in the Seydişehir tailing ponds in Turkey, with 552 waste material being produced throughout the year. New data for REE in red muds 553 from stored waste facilities in Turkey and Greece demonstrate their potential as 554 REE resource. Total REE in red muds from the two sites is approximately 12 000 t, 555 assuming an average of 1000 ppm REE in the red mud and an 80 % recovery (Borra 556 *et al.*, 2015).

557 Discussion

It is clear that red muds represent a significant potential resource of REE in Europe. Understanding the REE content of red muds is essential for robust resource estimation. Several factors may control the content and distribution of REE within red muds. These include: 1) the REE content and mineralogy of the bauxite precursor; and 2) the mineralogy of the REE within red mud ponds. The development of industrial-scale processes to extract REE from red mud is fundamental in establishing the economic viability of these potential resources.

565

566 *REE in bauxite and red mud*

567 The nature of, and mechanisms causing, variations in REE distribution in bauxites 568 are highly complicated with no one model fitting all deposits. In particular, the 569 effects of fractionation between the LREE and HREE, REE scavenging by Fe-rich 570 mineral phases, and changes in redox are not well understood. Although bauxites 571 are highly variable in their REE contents (e.g. from approximately 100 ppm to 572 greater than 2 300 ppm in bauxite from Parnassus-Giona (Laskou and Andreou, 573 2003)), in general there is often, but not always, an overall passive enrichment 574 downwards towards the limestone footwall. The importance of this is that REE 575 concentrations in red muds will be directly proportional to depth in the original 576 bauxite. If the original bauxite does not contain sufficient levels of REE, then REE 577 extraction from the red mud is unlikely to be viable, even with the recognised 578 enrichment factor of REE, including Sc and Y, from bauxite to red muds of 579 approximately 1:2 (Ochsenkühn-Petropoulou et al., 1994; Wagh and Pinnock, 580 1987). Selection of red muds from processing of a particular bauxite horizon, or a mixture of particular horizons, could potentially be carried out to produce ideal 581 582 blends for REE production.

583 The distribution and nature of the REE-bearing phases and minerals in red mud is

poorly defined. The association of REE with iron and manganese phases is widely acknowledged in the literature, as is the presence of authigenic and detrital REEbearing minerals. However, much more work is necessary in order to fully understand the distribution and mineralogy of REE in red muds. This will, in turn, allow a clear economic quantification of the potential REE resources in European red mud.

590

591 *REE processing*

592 Processing techniques for the two most common REE-containing minerals, 593 bastnäsite and monazite, are quite well understood and have well established 594 beneficiation flow sheets. Processing streams are being developed for other REE-595 bearing minerals such as eudialyte (www.eurare.eu), however, these do not make 596 a significant contribution to current REE production (Jordens et al., 2013). This 597 situation is likely to change with growing commercial demand for REE, with 598 increased opportunities to develop new rare earth element deposits with a wider 599 range and less well-understood mineralogy.

Successful extraction of REE from red muds has been achieved at both bench and pilot scales (Wagh and Pinnock, 1987; Ochsenkühn-Petropoulou *et al.*, 1996, 2002; Smirnov and Molchanova, 1997; Tsakanika *et al.*, 2004; Qu and Lian, 2013; RUSAL, 2014; Borra *et al.*, 2015). However, further development of these techniques is necessary in order to make these resources economically viable. In order for REE to be extracted most economically, ideally an additional processing stream for REE would be developed as an add-on to the Bayer process itself.

Taking an average concentration of 1000 ppm REE for red muds, REE by-product
production from current, karst-bauxite sourced, alumina production,
approximately 1.5 Mt annually (BGS, 2015), could equate to up to 1500 t REE per

610 year. By-product production of REE from European bauxite exploitation of 611 appropriate deposits could result in a steady, secure supply of REE within Europe. 612 In order to benefit from the red muds that are currently stored onshore, it would 613 be advantageous to regard the re-mining of the tailing ponds as a viable additional 614 resource. It has been shown that it is not only the REE that are potential by-products 615 of the red muds, research is also on-going into the extraction of Ga and Sc 616 concentrates from red mud waste at Orbital Aluminae, Canada and Rusal, Russia, 617 respectively. Additionally, base metals and other critical raw materials are 618 frequently found in the waste material, depending on the original composition of 619 the source, and are potential by-products of further extractive processing.

One positive aspect of the onshore stockpiles of red muds in Europe is that these are generally found in politically stable jurisdictions with an obvious history of mining and therefore, possibly without strong local anti-mining sentiment. Additionally, working on stockpiled waste is a more environmentally friendly option as no additional mining, and minimal, if any, grinding, is required. It also could provide an independent source of REE and protect REE-resource poor countries from export quotas and price fluctuations (Binnemans and Jones, 2015).

627 The current lack of plant-scale methods of extracting REE from the red muds is a 628 potential obstacle to the development of these resources. Ongoing research, such 629 as that carried out by the FP7 funded EURARE and Mud2Metals projects, is tackling 630 these challenges. Under the recent patent granted to Orbite Aluminae Inc. (Patent 631 No. 14/371,364, Orbite, 2015) for the 'Orbite Process', a red mud processing plant 632 at Cap-Chat, Québec, Canada, is expected to begin production in the last quarter of 633 2015 and aims to produce a REE concentrate as one of the products. It also aims to reprocess the red muds for alumina, as inefficiencies in the Bayer Process can result 634 635 in losses of up to 25% of the aluminium to the waste. The success of this

beneficiation stream could have a significant impact on the production of REE from red mud waste. However, it has been shown that the extraction of REE from red muds would do little to reduce the volume of red muds produced and so the development of alternative uses for red muds is still a key societal and environmental challenge (Klauber *et al.*, 2011).

641 There is a lack of collated data on red mud production and storage globally, 642 although it has been recognised internationally that the disposal, treatment and 643 maintenance of red mud ponds is of key social and environmental importance. This 644 issue was highlighted in 2010 with the tragic failure of the red mud pond at 645 Kolantár, Hungary. This has been addressed partly by the development of the 646 Bauxite, Residue and Disposal Database (BRaDD) (Gräfe et al, 2011). Maintaining 647 and updating such a large database needs the support of industry but can aid in 648 developing an appropriate strategy for the management of red mud (Power et al., 649 2011), including the possible extraction of valuable metallic commodities.

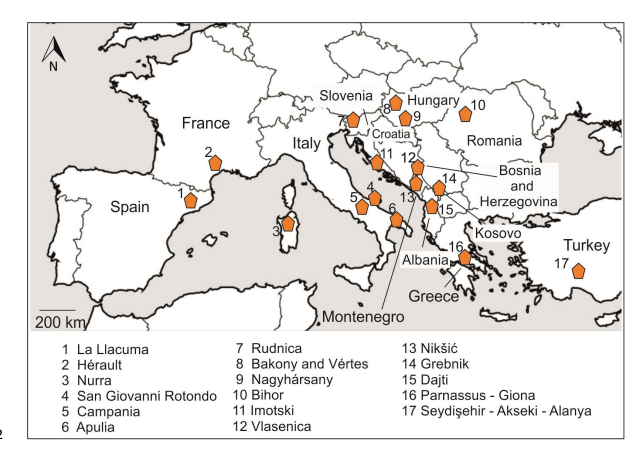
650 Conclusions

651 Potential resources of REE are present in significant quantities in red mud storage 652 facilities globally. Of these red muds, those that are derived from karst-type 653 bauxites are thought to be the most prospective for REE given the alkaline 654 conditions under which they form, where the pH due to the underlying limestone 655 retards mobility of the REE and traps the REE in the bauxite ore. In Europe, karsttype bauxites are most common in the Mesozoic limestone of the Mediterranean. 656 657 Historic and current exploitation of these bauxite deposits has resulted in REE-658 enriched red muds stored onshore across the region. This study has identified 659 combined potential resources of up to a maximum of 15000 tonnes REE contained 660 in red muds in two areas, one in Greece, the second in Turkey. A full exploration

661 campaign would be needed to investigate and evaluate the red mud waste tips. The 662 original heterogeneity of the bauxite could lead to heterogeneous waste tips that 663 would need careful resource estimation. Development of efficient beneficiation 664 and processing techniques and further work on understanding the distribution of 665 REE in red muds, in conjunction with the quantification of resources in European 666 red mud stocks, could lead to the development of sustainable REE production and 667 utilisation of waste material in Europe.

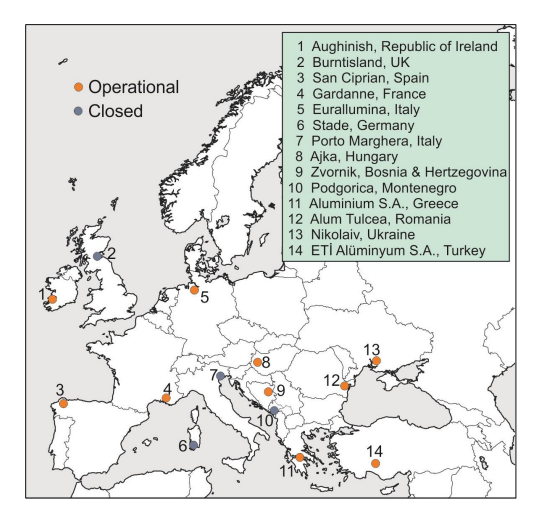
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683 Figure 1: European karst-bauxite deposits and mines in the circum-Mediterranean

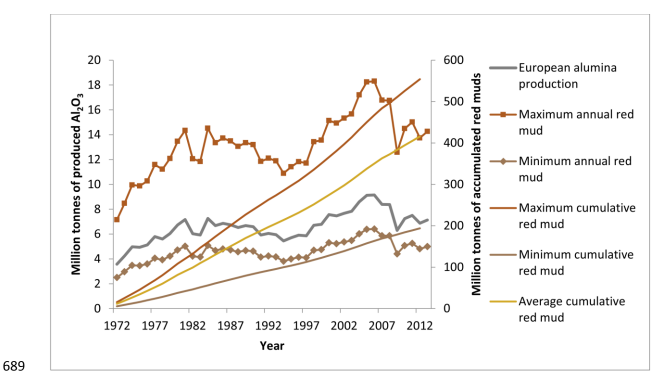
⁶⁸⁴ region (after Patterson, 1967).



685

686 Figure 2: Alumina refineries that utilise the Bayer process, both in use and

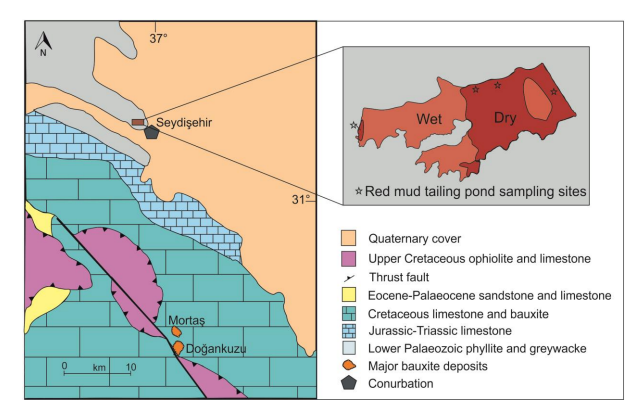
abandoned in Europe (adapted after Power *et al.*, 2011).



690 Figure 3: Time series data of European alumina production since 1972 (data from

691 British Geological Survey World Mineral Statistics database). Greek data are

- 692 excluded before 2009 due to the disposal of red muds at sea. Annual European
- alumina production is on the left-hand vertical axis, and cumulative average,
- 694 minimum and maximum volumes of red muds stored in European onshore tailing
- 695 ponds or 'dry' stacks are on the right-hand vertical axis.

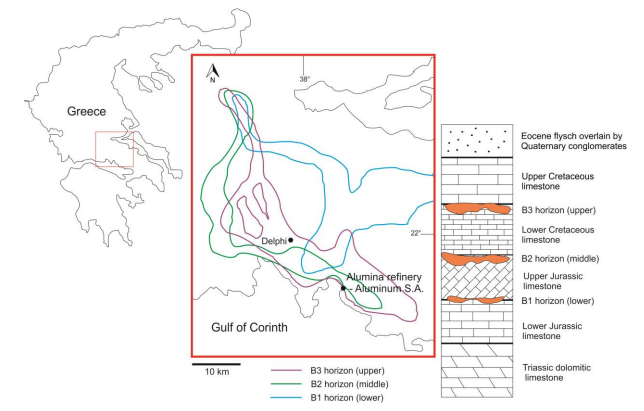




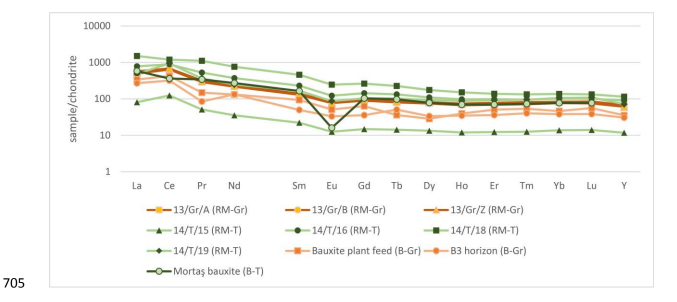
697 Figure 4: Geology of the Seydişehir area, Turkey (adapted with permission from

698 Öztürk et al., 2002 and pers comm.) showing key geological features, the location

699 of the red mud tailing pond and sampling sites.



- 701 Figure 5: Regional distribution and simplified stratigraphic column of the three
- bauxite horizons (Petrascheck and Mack, 1978; Valeton *et al.*, 1987; schematic
- 703 stratigraphic column reproduced with permission after Laskou and Economou-



704 Eliopoulos, 2013).

- 706 Figure 6: Chondrite-normalised (MacDonough and Sun, 1995) REE patterns for red
- 707 mud samples from Greece and Turkey compared with published values for the
- 708 Greek plant feed bauxite and the B3 horizon (Ochsenkühn-Petropoulou et al.,
- 1994, 1995) and the Mortaş bauxite (Karadağ et al., 2009). Red muds: RM;
- 710 bauxites: B; Turkey: T; Greece: Gr.

	Red muds, this study						Bauxite, literature data						
Sample name	13/Gr/A	13/Gr/B	13/Gr/Z	14/T/15	14/T/16	14/T/18	14/T/19	Mortaş bauxite,	Bolkardaği BP bauxite,	Bolkardaği OS bauxite,	Plant feed bauxite, (B3+B2)	Parnassus -Giona B3 bauxite,	Regia Piana bauxite, Italy
	Parnassus-Giona red mud, Greece (this study)			Seydişehir red mud, Turkey (this study)				Turkey*	Turkey*	Turkey*	Greece*	Greece*	(GBRX 8)*
La	134.9	117.2	129.3	19.2	185.9	356.2	114.8	137.6	107.6	35.8	80.2	64	87.7
Ce	416.9	391.3	421	76.1	543.2	728.5	576	221	233.5	159	259.5	195	154.1
Pr	29.6	27	28.3	4.7	49	102.7	33.9	31.7	24.7	7.1	13.8	7.8	19.7
Nd	107.4	99	104	16.2	169	346.7	113.5	124.7	88.1	26	59.8	60.3	74.2
Sm	20.5	19.1	20.5	3.3	34.1	67.9	24.7	24.2	17.2	5.2	13.8	7.3	13.4
Eu	4.5	4.3	4.4	0.7	6.9	13.8	4.9	0.9	3.8	1.2	2.8	1.8	2.8
Gd	19.3	17.8	19.6	0.5	4.8	8.2	3.7	20.6	17.9	6.4	12.5	7.1	12
Tb	3.1	2.8	3	2.9	28.5	52	20.6	3.5	3.2	1.4	1.3	1.8	2.1
Dy	19.7	18.2	19.9	3.3	27	42.9	22.7	19.1	19.6	9.6	7	8.2	11.9
Ho	4.1	3.6	3.9	0.7	5.3	8.2	4.8	3.8	4.2	2.3	2.1	1.9	2.5
Er	12.4	11	11.8	2	15.7	21.9	14.8	11.2	12.5	7.6	8	5.7	7.7
Tm	2	1.7	1.9	0.3	2.4	3.3	2.3	1.8	2	1.3	1.3	1	1.2
Yb	13.3	12.2	13.5	2.2	16.4	21.9	17	12.4	12.2	8.6	7.4	6.1	8.3
Lu	2.1	1.8	2.1	0.3	2.5	3.3	2.7	1.9	1.9	1.3	1.3	0.9	1.2
Total Lanthanoids	790	727.5	783.2	132.5	1090.6	1777.4	956.2	614.4	649.3	674.6	471.3	369.4	497.6
Y	106.2	95.4	95.2	18.4	141.3	180.2	112.7	N/A	78.2	126.2	55.4	48	56.7
TREE	896.2	822.8	878.4	150.9	1231.9	1957.5	1068.9	614.4	727.5	800.8	526.7	417.4	554.3

711

712 Table 1: REE (ppm) contents in red mud samples from Turkey and Greece,

713 (*literature data). Averaged values are for bauxite feed (a blend of horizons B3

and B2) from the Aluminium S.A. alumina refinery, Greece (Ochsenkühn-

Petropulu et al., 1994, 1995); horizon three (B3) of the Parnassus-Giona bauxite

province, Greece (Valeton *et al.*, 1987; [Economopolou-Kyriakopoulou, 1991];

717 Laskou, 1991, 2005; Ochsenkühn-Petropoulou et al., 1991; Laskou and Economou-

Eliopoulos, 2007, 2013); the Mortaş bauxite, Turkey (Karadağ et al., 2009) and, for

719 comparison, the Baharpınarı (BP) and Öşün (OS) bauxite deposits in the Bolkardaği

bauxite province in Turkey (Hanilçi, 2013) and the Regia Piana bauxite (GBRX 8),

721 Italy (Boni *et al.,* 2013).

	Red mud (Mt)	TREE concentration (ppm) ^e	TREE content (Mt) (@ 50% recovery)	Nd concentration (ppm) ^e	Nd content (Mt) (@ 50% recovery)	HREE concentration (ppm) ^e	HREE content (Mt) (@ 50% recovery)
Minimum	200	1000	0.10	150	0.02	200	0.02
cumulative							

Average cumulative 425 1000 0.21 150 0.03 200 0.04	Maximum cumulative	550	1000	0.27	150	0.04	200	0.06
	-	425	1000	0.21	150	0.03	200	0.04

722

Table 2: Cumulative minimum, maximum and average tonnages of red muds

stored onshore in Europe since 1972 and their estimated contents of *TREE*; Nd;

725 and HREE_(Eu-Lu+Y); (^e estimate).

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