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Origin and geochemical evolution from ferrallitized clays to karst bauxite: An example from the Lower Cretaceous of NE Spain

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1	Origin and geochemical	evolution from	ferrallitized clays	to karst bauxite: Ar	example from the

Lower Cretaceous of NE Spain

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8 ABSTRACT

9 Four outcrops of Lower Cretaceous (Barremian) karst bauxites located in Teruel (NE Spain) 10 were analysed. The deposits show a heterogeneous-chaotic lithostructure consisting of 11 pisolitic bauxite blocks embedded in lateritic red clays filling karst cavities. The research has 12 focused on the geochemical study of major, minor, and trace elements (including some critical 13 to industry) of both the bauxites and clays. The objective was to investigate the bauxite 14 precursor material and to characterize the system's geochemical evolution. Geochemical 15 analyses were carried out by inductively-coupled plasma optical emission and mass 16 spectroscopy. An absolute weathering index has been calculated to estimate element mobility, 17 assuming Ti as an immobile element and the Upper Continental Crust (UCC) as parent material. 18 Further, selected samples were observed by field emission scanning electron microscopy. The 19 data indicate that both the bauxites and red clays originated by intense chemical weathering 20 from more mafic argillaceous sediments than the UCC. Ongoing weathering caused the 21 bauxitization of the upper parts of the original profile, preventing the lower parts from being 22 bauxitized, thus producing the ferrallitized clays underlying the pisolitic bauxites. Subsequent 23 karst reactivation gave rise to the current lithostructure. Ferrallitization is related to Fe, Sc, and 24 V enrichment. On the other hand, although bauxites are relatively enriched in some elements 25 compared to clays, the more intense chemical weathering associated with bauxitization led to 26 chemical homogenization and widespread element depletion. During the bauxitization, Al, Ti, 27 Zr, Cr, and probably Hf and the critical element Nb behaved as more immobile elements in the 28 system. Bauxitization also enhanced homogenization and depletion of the REE, which is more 29 pronounced for the LREE. HREE trends seem to be partly related to the concentration of Ti 30 oxides in the bauxites, whereas P-bearing phases, more frequent in the clays, control the LREE. 31 Subsequent to bauxitization, partial kaolinization of the bauxite took place related to the 32 circulation of acid solutions that also caused the karst reactivation. These late processes 33 caused some AI depletion in the bauxites and enhanced Fe loss together with V and, to a lesser 34 extent, Ge.

35 Keywords: karst bauxite; ferrallitization; lower Cretaceous; weathering; geochemistry; REE

36 1. INTRODUCTION

37

38 Bauxitization is a special case of chemical weathering (Bardossy, 1982). Chemical weathering is 39 a major mechanism that partitions elements between crustal rocks and natural water (Taylor 40 and McLennan, 1985); it is completely dependent on the (climate-driven) water cycle (Tardy, 41 1986). During chemical weathering, the differentiation of elements depends on element 42 mobility, and the degree of chemical differentiation depends on the intensity of chemical 43 weathering. Accordingly, major, minor, and trace elements are fractionated during 44 bauxitization, and therefore the distribution of mobile elements can reflect the intensity of 45 weathering and the distribution of immobile elements can be used to identify precursor rocks. 46 Molecular proportions of Al_2O_3 , CaO, Na_2O_3 and K_2O are used to calculate the Chemical Index 47 of Alteration (CIA) (Nesbitt and Young, 1982), which has been utilized in a number of studies to 48 determine the degree of weathering. In bauxite formation, many studies have recognized Al as 49 an immobile element (e.g. Sastri and Sastry, 1982; Valeton et al., 1987; Gow and Lozej, 1993), 50 and high correlations between AI and some major and trace element contents indicate 51 immobile element pairs during the bauxitization process (MacLean, 1990). Most of the 52 geochemical research has also evidenced the relatively immobile behaviour of a group of trace 53 elements (e.g. Sc, Co, Cr, and Zr) in the surficial environment. Furthermore, REE abundance 54 varies in weathering products and is accepted as a marker of their source rocks (e.g. 55 Schellmann, 1986; Taylor and McLennan, 1995; Nyakairu and Koeberl, 2001). What is more, Eu 56 anomalies and (La/Yb)_N ratios depend on fractionation during bauxitization (Cullers and Graf, 57 1983; Maksimovic and Pantó, 1991; Mongelli, 1997; Mameli et al., 2007). In consequence, a 58 number of studies have shown that immobile elements are a powerful tool to decipher the 59 genetic history of karst bauxite deposits (e.g. Maksimovic and Pantó, 1991; MacLean et al., 60 1997; Mongelli, 1997; Mameli et al., 2007; Liu et al., 2013). 61

62 According to Bardossy (1982), bauxite deposits overlying carbonate rocks are called karst 63 bauxite deposits regardless of whether the bedrock surface is karstified or not, or of the 64 degree of karstification (Mameli et al., 2007). Among karst bauxites, a great majority are 65 Mediterranean-type, which frequently shows moderately to strongly karstified bedrock 66 surfaces (Bardossy, 1982). Karst bauxite deposits from NE Spain are of this type and occur in 67 three main zones: the South Pyrenean Zone, Catalonian Coastal Range, and Maestrazgo. Our 68 study focuses on one of the most representative deposits located in the Maestrazgo zone, the 69 Lower Cretaceous (Barremian) Fuentespalda deposit (Teruel, NE Spain) (Molina and Salas, 70 1993). Previous studies based on mineralogical and textural features of the Fuentespalda

71 deposit have proposed a complex sequence of processes to explain the bauxite genesis, mainly

through the transformation of precursor lateritic clays into pisolitic bauxites (Yuste et al., 2015).

73 According to these authors, bauxitization occurred as a result of intense chemical weathering

74 under seasonal subtropical climate conditions that favoured several pulses during the

75 bauxitization process.

76

77 Although the Maestrazgo bauxite deposits no longer have any economic value, understanding 78 the genetic processes and geochemical evolution that led to their formation is a valuable 79 means to decipher the behaviour of chemical elements during bauxitization. This can be of 80 special interest in the case of those elements critical to industry, such as REE, Nb, Ga, and Ge, 81 listed in the 2014 European Union report on critical raw materials and usually found in karst 82 bauxites (Mongelli et al., 2014). In consequence, as Mongelli et al. (2016) summarizes, recent 83 research has dealt with the processes that control the distribution of economic elements in 84 karst bauxite deposits.

85

86 Our research focuses on the geochemical study of major, minor, and trace elements of the 87 Fuentespalda karst bauxite deposit (NE Spain), which is a good example of bauxite closely 88 related to lateritic clay materials. Our goal was to investigate the precursor material of the 89 bauxites and to characterize the geochemical evolution of the bauxitization process.

- 90
- 91 2. GEOLOGICAL FRAMEWORK
- 92

93 The bauxite deposit studied is Early Cretaceous (Barremian) in age and located in the 94 Maestrazgo zone (Fig. 1), which is one of the three most important bauxite regions in NE Spain. 95 During the Mesozoic, the spread of the Tethys westwards and the opening of the North 96 Atlantic Ocean led to the development of a rift system in Iberia. The Iberian rift remained 97 active until the Mid Cretaceous and controlled the late Palaeozoic and the Mesozoic 98 sedimentation in the study area. One of the most active stages of this rifting process took 99 place during the Late Jurassic–Early Creataceous (Salas and Casas, 1993; Van Wees et al., 1998) 100 and was responsible for the creation of several basins with high subsidence rates (Mas et al., 101 1998; Salas et al., 2001; Mas and Salas, 2002). The first of these basins was the Maestrazgo 102 Basin (Guimerà et al., 2004), where a clear predominance of marine carbonates and marls 103 characterized sedimentation during the Late Jurassic (Mas et al., 2004). As the rifting stage 104 progressed, tectonic activity was attenuated and subsidence rates decreased during the Early 105 Cretaceous (Valanginian–Barremian) (Mas et al., 2004). The sedimentary record during this

106 time in the Maestrazgo Basin was controlled by very shallow carbonate platforms subject to 107 significant continental influence, giving rise to large lacustrine-palustrine areas (Mas et al., 108 2004). 109 110 The study area shows considerable development of folds and thrusts affecting the Mesozoic 111 formations, with predominant E-W strike orientations. These structures are conditioned upon 112 basement faults that also controlled the main palaeogeographic characteristics during 113 Mesozoic sedimentation (Molina and Salas, 1993). These features constrain the current 114 location of the study deposits, which crop out in a NW-verging anticlinal thrust affecting 115 Jurassic and Cretaceous rocks surrounded by continental Cenozoic materials (Molina and Salas, 116 1993) (Fig. 1). 117 118 The bauxite and related clay materials constitute bodies filling karst cavities developed in 119 limestones and argillaceous limestones of Upper Oxfordian–Kimmeridgian age, constituting 120 the Polpis Formation (Salas, 1987). Some bodies are lens-shaped and concordant with the 121 stratification of the host rocks. Nevertheless, the original morphology of the cavities, exposed 122 by small quarries and trenches from past mining activity, is frequently modified by brecciation 123 associated with radial acicular calcite cements (Molina and Salas, 1993). Materials filling the 124 cavities have a heterogeneous-chaotic lithostructure (Bardossy, 1982) consisting of up to 125 meter-sized pisolitic bauxite blocks, sometimes fractured or faulted, embedded in red clays 126 (Fig. 2). Pisolitic bauxites are mainly red, but predominantly white zones are also frequently 127 observed in the upper parts of blocks (Fig. 2). The paleokarst is fossilized by the Arenas de 128 Utrillas Formation (Albian) and/or by Cenomanian dolomitized limestones (Molina and Salas, 129 1993). Important outcrops of ferrallitized clays exist in the vicinity of the studied karstic 130 bauxites (Combes, 1969). Lateritic clays from the Barremian Cantaperdius Formation (Salas, 131 1987) crop out in the proximity of the study area (approximately 10–15 km southeast of 132 Fuentespalda) near the localities of Beceite (Teruel) and Fredes (Castellón). These lateritic clays 133 also commonly overlie karstified carbonate strata, show a similar composition to the clay 134 materials related to the studied bauxite and displaying frequent pisolitic levels indicating, as 135 Molina and Salas (1993) pointed out, that they are laterally related to the bauxites and 136 associated clays from Fuentespalda. 137 138 3. METHODS

140 Samples were taken from four outcrops (localities 1-4 in Fig. 1). A set of fourteen samples from 141 the pisolitic bauxite (nine samples) and the red clays (five samples) constituting the infill 142 material of the karst cavities have been analysed. Pisolitic bauxite samples have been divided 143 into red pisolitic and white pisolitic. Further, one host limestone sample and one sample of the 144 observed acicular radial calcite cements have also been analysed. The first number of sample 145 labels in the data tables indicates the locality number. 146 147 Chemical analyses of major and trace elements of bulk samples were performed at Actiabs 148 Laboratories (Canada) using the following techniques (detection limits in brackets): major 149 elements (0.01%, except MnO and TiO₂ with detection limits of 0.001%) and trace elements V 150 (5 ppm), Sr (2), Ba (3), Sc (1), and Be (1) by inductively coupled plasma/optical emission 151 spectroscopy (ICP/OES). The other trace elements, such as Cr (20 ppm), Co (1), Ni (20), Rb (1), 152 Cs (0.1), Th (0.05), U (0.01), Y (0.5), Zr (1), Nb (0.2), Hf (0.1), La (0.05), Ce (0.05), Pr (0.01), Nd 153 (0.05), Sm (0.01), Eu (0.005), Gd (0.01), Tb (0.01), Dy (0.01), Ho (0.01), Er (0.01), Tm (0.005), Yb 154 (0.01), and Lu (0.002) were determined by inductively coupled plasma/mass spectroscopy 155 (ICP/MS). Fused samples (lithium metaborate/tetraborate fusion) were run for major oxides 156 and V, Sr, Ba, Sc, and Be on a combination simultaneous/sequential Thermo Jarrell-Ash ENVIRO 157 II ICP or a Varian Vista 735 ICP. Calibration is performed using seven prepared USGS and 158 CANMET certified reference materials. One of the seven standards is used during the analysis 159 for every group of ten samples. For the analysis of the other trace elements, fused samples are 160 diluted and analysed by Perkin Elmer Sciex ELAN 6000, 6100 or 9000 ICP/MS. Three blanks and 161 five controls are analysed per group of samples. Duplicates are fused and analysed every 15 162 samples. 163

Selected samples (11) were analysed by field emission SEM (FESEM) using backscattered electron (BSE) and energy-dispersive X-ray (EDS) analyses. The observations were performed using a JEOL JSM 6400 (SEM) equipped with an Oxford instrument detector (EDS) and a Carl Zeiss MERLIN FESEM. The accelerating voltage was 4 to 15 kV with a beam current of 1 to 2 nA and a counting time of 50 s. Samples were carbon coated.

169

170 An absolute weathering index (Nesbitt, 1979) has been calculated to estimate element

171 mobility, assuming Ti as an immobile element and the Upper Continental Crust (UCC; Taylor

and McLennan, 1985) as parent material. According to Nesbitt (1979), the percentage increase

173 or decrease (change %) on any element X compared with its concentration in the parent

174 material is given by [(X_{sample}/Ti_{sample})/(X_{UCC}/Ti_{UCC}) - 1]*100. The Chemical Index of Alteration (CIA;

Nesbitt and Young, 1982) has been calculated using molecular proportions of the following
oxides: CIA = (Al2O3/Al2O3+CaO*+Na2O+K2O)*100, where CaO* represents the amount of
CaO associated with the silicate fraction of the rock.

178

179 **4. RESULTS**

180

181 The mineralogical analysis by X-Ray Diffraction was the subject of a previous paper (Yuste et al., 182 2015). Despite this, we include here a brief summary essential to properly discussing and 183 interpreting the geochemical study carried out in the current research. The average 184 mineralogical composition of the samples is shown in Table 1. In whole sample, pisolitic 185 bauxites show kaolinite (KIn), gibbsite (Gbs), diaspore (Dsp), goethite (Gt), and hematite (Hem), 186 accompanied by traces of boehmite (Bhm), anatase (Ant), rutile (Rt), and occasionally quartz 187 (Qtz). The clayey samples are composed of kaolinite, goethite, hematite, diaspore, and illite or 188 muscovite, along with accessory anatase, rutile, guartz, and calcite (Cal). Kaolinite is very 189 abundant in both the pisolitic and the clay samples, whereas gibbsite, always accompanied by 190 boehmite, is only present in some pisolitic samples, both white and red. The other common Al 191 hydroxide in bauxites, diaspore, has been identified in all the samples. Clays and red pisolitic 192 samples always contain hematite and frequent goethite, whereas these minerals are generally 193 absent in the white pisolitic samples. Rutile and, to a greater extent, anatase have been 194 identified in all the samples. In the <2 μ m fraction, kaolinite is the major and almost the only phyllosilicate except in the clays, where illite is always detected and can reach up to 40%. 195 196 Kaolinites from the clay samples are less ordered and less crystalline than those from the 197 pisolitic samples. Among the pisolitic samples, kaolinites from the white ones are the most 198 crystalline. 199

In the Fe₂O₃-Al₂O₃-SiO₂ classification diagram by Aleva (1994) (Fig. 3a), the clays are classified
 as laterites, whereas the red pisolitic samples fall in both the laterite and the bauxite fields and
 the white pisolitic in the kaolinitic bauxite and bauxitic kaolinite fields. The samples falling in
 the bauxite and kaolinitic bauxite fields are those containing gibbsite and boehmite.

204

Si, Al, Fe, and Ti concentrations on average account for 99.04% of the major elements (Table 2) except in the host limestone and the calcite cement sample, where Ca is the most abundant element. In general, major element concentrations reflect the mineralogical composition, with the highest Al contents in samples with gibbsite and boehmite and low Fe contents in white pisolitic samples (scarce or absent Fe oxides). Mg is mainly related to illite, as suggested by its

higher concentrations in the clays. Nevertheless, Mg could be related to some interstratified
smectite layers in illite not detected by XRD due to the low illite contents. Among major
elements, the only significant correlation is that shown by Ti and Al (r = 0.89; r = 0.95 taking
into account the limestone and the calcite sample too) (Fig. 3b), thus constituting an immobile
element pair and indicating that Al- and Ti-bearing minerals concentrate in the same type of
rock. Considering only the pisolitic samples, a negative correlation (r = -0.75) is shown by Al
and Si, as usual in bauxites.

217

218 Major element concentrations (Table 2) reveal that the clay samples are chemically more 219 heterogeneous than the pisolitic bauxites. When normalized to the Upper Continental Crust 220 (UCC) concentrations (Fig. 4a), there is an enrichment in Al and Ti in both the clays and the 221 pisolitic samples. The red pisolitic and the clay samples are also enriched in Fe, but an evident 222 Fe depletion can be observed in the white pisolitic samples. As can be seen in Figure 4a, both 223 the pisolitic and the clay samples are depleted in Si, Mn, Mg, Ca, Na, K, and P although, in 224 general, the clay samples are less depleted in Mn, Mg, Ca, K, and P. On the other hand, the 225 change % values in Figure 4b show that clay samples are more heterogeneous than the 226 pisolitic ones. This figure also shows that Al, P, and to a lesser extent Si, Mn, Ca, and K are in 227 general more depleted relative to Ti in the pisolitic samples. The pisolitic samples, particularly 228 the white ones, tend to be Fe depleted. This is commonplace since the red colour of bauxites 229 comes from ferric iron and therefore the white bauxite samples are indicating at least 230 transient reducing condition invariably leading to mobilization and removal of iron (Bardossy, 231 1982). CIA values (Table 2) are high (>96), with those from the clay samples more 232 heterogeneous and lower (mean value = 98.54) than those from the pisolitic bauxite (mean 233 value = 99.83).

234

235 Figure 5a shows some important trace element concentrations normalized to the UCC. 236 Transition element (Sc, V, Cr, Co, Ni) normalized concentrations show an irregular trend. The 237 only remarkable features are the relatively higher enrichment in Cr and an evident Co 238 depletion in the pisolitic with respect to the clay samples. Also noticeable is the clearly lower V 239 enrichment in the white pisolitic samples. With regard to the critical elements Ga and Ge, only 240 the white pisolitic samples are slightly depleted in Ge. The HFSE (Zr, Nb, Hf, Th, U) are always 241 more abundant in the pisolitic samples than in the clays. Finally, the LILE (Sr, Cs, Ba) are 242 depleted in all the samples, clearly showing that they were all subject to rather intense 243 leaching. The average Ba/Sr ratio is higher in the clays (0.4) than in the red (0.2) and white 244 (0.1) pisolitic samples. With regard to change % values (Fig. 5b), the transition elements are

always depleted in the pisolitic bauxites. The clays are enriched in Sc and V and only slightly
depleted in Cr and Ni. Ga and Ge are also depleted in bauxites and to a lesser extent in clays.
The HFSE, and especially the LILE, are depleted in both the bauxites and the clays and show
similar patterns regardless of the sample type. In general, as for major elements, the change %
values in the clays are more heterogeneous than those of the pisolitic samples.

250

251 Y has been considered along with the REE since there is a high positive correlation between 252 them (r=0.91), indicating similar behaviour. The average REE and Y contents (Table 2) of the 253 clay samples are distinctly higher than those of the pisolitic samples, but they are much lower 254 in the host limestone and the calcite sample. The REE+Y concentrations normalized to the UCC 255 average (Fig. 6a) also show clear differences among the different types of samples considered. 256 In general, the clay samples are clearly enriched in REE+Y, whereas the pisolitic samples are 257 depleted in LREE and enriched in HREE. The limestone and the calcite cement sample are in 258 general depleted in the REE. The highest enrichment in REE+Y in Figure 6a corresponds to the 259 clay sample with the highest illite contents. The REE+Y change % graph (Fig. 6b) also reveals 260 differences: the clay samples are less depleted than the bauxites and are specifically enriched 261 in Y and La and slightly enriched in Lu. The highest enrichment is seen in the clay sample with 262 high illite contents and, especially, in the host limestone and the calcite sample (not in the 263 graph). The chondrite-normalized REE patterns (Fig. 6c) are also different depending on the group of samples considered. In both the pisolitic and the clay samples, LREE fractionation is 264 265 higher than HREE fractionation, but the HREE in the clay samples show an almost horizontal 266 slope, whereas in the pisolitic samples they show a slight upward trend. The most significant 267 difference between the clay and the pisolitic samples (in addition to the higher REE contents in 268 the clay samples mentioned above) is the positive Ce anomaly observed in several pisolitic 269 samples (especially red ones). The host limestone and the calcite cement sample REE patterns 270 are very similar to each other, with LREE and HREE fractionation (the latter higher than that 271 shown by the clay and the pisolitic samples) and a distinct negative Ce anomaly. All the 272 samples show similar Eu anomaly values (Table 1). Plots of (Gd/Yb)c versus (La/Sm)c and 273 Ce/Ce* versus (La/Yb)c in Figure 7 (values in Table 2) allow the three groups of samples to be 274 distinguised. Figure 7a shows higher LREE and HREE fractionation in the clay samples 275 compared to the pisolitic ones. On the other hand, Figure 7b shows lower REE fractionation of 276 the pisolitic samples and higher Ce anomalies of these samples, followed by the clay samples. 277

5. DISCUSSION

280 Several studies on karst bauxite deposits have proposed that bauxite formed from lateritized 281 material precursor (Valeton, 1972; Ling et al., 2015), ferruginous-argillaceous debris 282 (Zarasvandi et al., 2008), or clayey (mainly kaolinitic) material (MacLean et al., 1997; Oggiano 283 and Mameli, 2001). A previous study of the Fuentespalda karst bauxite deposit proposed that 284 pisolitic bauxite originated in situ from precursor lateritic clays through a complex sequence of 285 mineralogical processes (Yuste et al., 2015). According to these authors, the bauxitization 286 process occurred from the top down, preventing the lower part of the profile from being 287 entirely bauxitized and resulting in preservation of the lateritic parent clays underlying the 288 pisolitic bauxite. Karst reactivation subsequent to bauxitization caused the present 289 heterogeneous-chaotic lithostructure of the deposits, following the classic scheme by Combes 290 (1969). 291 292 The high correlation (r=0.94) between Ti and Al (Fig. 3a) is related to the formation of Al-rich 293 minerals and to Ti oxides, indicating an immobile nature for these major elements in the 294 bauxitization process, as traditionally considered (Nesbitt, 1979; Sastri and Sastri, 1982; 295 Valeton et al., 1987; MacLean, 1990; Gow and Lozej, 1993). The trend from clay to pisolitic 296 samples shown in that figure also supports the idea that the clay materials closely associated 297 to the pisolitic bauxite are their precursors. The more heterogeneous chemical character of the 298 clay samples also points in this direction and probably reflects a greater detrital influence 299 related to less and variable weathering compared to the pisolitic samples. This is also 300 supported by the more variable and less crystalline nature of kaolinite from the clay samples, 301 as Yuste et al. (2015) reported. CIA values, higher than those of the UCC (60.11), are more 302 heterogeneous and lower in the clay samples compared to the pisolitic ones, which is in 303 agreement with the above ideas. Regarding the change % data, the widespread higher 304 depletion recorded by the pisolitic bauxite relative to the clays has to be interpreted as a result 305 of more intense weathering that led to the formation of bauxite from the precursor. The 306 mineralogical characteristics of the clay samples and their chemical classification as laterites 307 support the notion that these materials are precursor aluminous clays for the pisolitic bauxite, 308 and are probably related to the lateritic clays from the Cantaperdius Fm. described by Salas 309 (1987) and Molina and Salas (1993). The scenario first included the formation of the clays 310 through ferrallitization and edaphic processes favoured by periodic subaerial exposure in a 311 tropical climate (Molina and Salas, 1993). Palaeogeographic conditions and ongoing 312 weathering permitted bauxitization of this precursor. Subsequently, late kaolinization 313 processes (resilicification) took place (Yuste et al., 2015). The higher Al depletion in the pisolitic

314 samples (Fig. 4b) could be related to these processes subsequent to bauxitization involving

315 partial replacement of gibbsite by kaolinite in the pisolitic bauxite, as Yuste et al. (2015) 316 pointed out. According to these authors, this replacement was more pronounced in the white 317 pisolitic bauxite, which is supported by the slightly higher Si depletion in the red pisolitic 318 samples. Nevertheless, these late processes did not lead to an absolute Si enrichment in the 319 pisolitic samples, as revealed by the change % values. In West Sardinian karst bauxite, for 320 example, SiO₂ addition has been well documented and related to low-temperature solutions 321 linked to Oligocene-Miocene calc-alkaline volcanic activity (Oggiano et al., 1987; Oggiano and 322 Mameli, 2001). In the present case, no post-Jurassic magmatism has been documented, and 323 therefore we must turn to solutions acid enough to induce Al hydroxide dissolution and a 324 decrease in SiO₂ solubility favouring kaolinite recrystallization. The palaeokarst where the 325 bauxites are hosted was fossilized by Albian formations which in the Maestrazgo Basin include 326 materials rich in organic matter and coal. These materials could have played a significant role 327 in the acidification of fluids responsible for karst reactivation and related bauxite kaolinization. 328 329 In general, based on Fe change % values, the more intense weathering related to the 330 bauxitization process led to Fe loss and probably to its accumulation in the underlying clays. 331 Not surprisingly, then, the precursor clays are clearly enriched in this element compared to the 332 pisolitic bauxites. This Fe loss was aggravated by more pronounced late kaolinization processes 333 in the white pisolitic bauxite, as indicated by the higher Fe depletion in these samples. Fe 334 concentrations normalized to the UCC also support this.

335

336 Taking into account concentrations normalized to the UCC, bauxites are relatively more 337 enriched in Cr, Ga, Zr, Nb, Hf, Th, and U and depleted in Co (and in Ge in the case of the white 338 pisolitic samples). The only ones of these elements that show a good positive correlation with 339 each other and with Ti and Al are Zr and Cr (Fig. 8), and thus they can be considered as 340 immobile elements during bauxitization. Nevertheless, in absolute terms, bauxitization has promoted higher trace element depletion (except for Zr, Nb, and Hf, which show similar values 341 342 in both bauxites and clays) than lateritic processes, which is related to the more intense 343 weathering (as deduced from the change % values). This is also supported by the observed 344 Ba/Sr average ratios, since Sr is much more soluble than Ba (Retallack, 1990). The low Cr 345 depletion observed in Figure 5b and the similar values shown by both bauxites and clays 346 support the idea of its less mobile nature along with Zr and probably Nb and Hf. In addition, 347 bauxitization has led to chemical homogenization taking into account the more heterogenous 348 values shown by the clay samples. On the other hand, it is worth mentioning the slight 349 enrichment in Sc and V recorded in the lateritic clays that would be derived from the lateritic

process. The evident V and, to a lesser extent, Sc and Ge depletion observed in the white
samples seems to be linked to the late kaolinization process as it shows a trend similar to Fe.
Most of the REE in karst bauxites are bound to detrital minerals inherited from the source rock
but they also fractionate during the bauxitization process (Maksimovic and Roaldset, 1976;
Maksimovic and Pantó, 1991), and can be adsorbed on the surface of clay minerals (see Wang

356 et al., 2010, and references therein) or Fe phases (Mongeli, 1997; Mameli et al., 2007). In

357 addition, as Maksimovic and Pantó (1978, 1991) have shown, fractionation can take place

- 358 through the formation of authigenic REE minerals.
- 359

360 The higher REE+Y contents in the clay samples could be related to a variety of inherited 361 minerals in agreement with their more detrital nature, supported by the more ubiquitous 362 presence of illite in these samples. In fact, the sample with the highest REE+Y contents is that 363 with the highest illite contents. In addition, the higher REE+Y content in the clay samples would 364 have been enhanced by the downward enrichment of REE in the lowermost part of karst 365 bauxites during the bauxitization process, as D'Argenio and Mindszenty (1995, and references 366 therein) indicated, and Maksimovic and Pantó (1991) showed in the karst bauxite deposits of 367 Yugoslavia and Greece. The transformation of the clays into pisolitic bauxite therefore involved 368 REE+Y loss. High positive correlations have been found between REE and Cs (0.94) and Co (r=0.87), indicating similar behaviour of these elements during bauxitization. REE+Y change % 369 370 values (Fig. 6b) support the above idea since pisolitic samples show a clear depletion in these 371 elements, especially the LREE. From this graph, it can be also deduced that bauxitization tends 372 to homogenize REE+Y concentrations, as observed in trace elements, since clays show more 373 heterogeneous values. This trend is also shown by the REE+Y concentrations normalized to the 374 UCC (Fig. 6a). That is to say, the more intense the chemical weathering (i.e. bauxitization vs 375 ferrallitization), the more LREE depletion can be expected, in agreement with Maksimovic and 376 Pantó (1991).

377

The lower (La/Sm)c, (La/Yb)c, and (Gd/Yb)c average values (lower fractionation) in the pisolitic samples are most likely linked to the concentration of certain immobile phases that act as scavengers for the REE as weathering progresses. In addition, the higher LREE over HREE fractionation indicates that the HREE were less mobile than the LREE. This could be attributed to the concentration of heavy minerals like rutile and anatase, which might have acted as scavengers for HREE, as occurred in the south Turkish bauxite deposits reported by Karadag (2009). On the other hand, some authors (Nyakairu et al., 2001; Kanazawa and Kamitani, 2006;

385 Roy and Smykatz-Kloss, 2007) have reported on the importance of P-bearing phases as LREE 386 carriers. In this regard, SEM observations of the study samples have evidenced a more 387 frequent occurrence of xenotime, apatite, and Al phosphate-sulfate (APS) minerals in the clay 388 samples (Fig. 9), whereas Ti oxides are more common in the pisolitic ones, in agreement with 389 chemical data (Fig. 4). These observations would explain the reported LREE and HREE trends in 390 the pisolitic samples compared to the clay ones. The positive correlation between the LREE 391 and P_2O_5 (r=0.60) would support the above ideas, but the poor negative correlation found 392 between the HREE and TiO₂ (r=-0.21) indicates that minerals other than Ti oxides are related to 393 the REE behaviour during bauxitization. As Bardossy and Panto (1973), Maksimovic and Pantó 394 (1991), Mongelli (1997), and Mameli et al. (2007) have indicated, fluorocarbonate minerals of 395 the bastnäsite group are the most frequent REE minerals in Mediterranean karst bauxite 396 deposits, although these minerals have not yet been identified in the Fuentespalda deposit. 397

398 With regard to the cited frequent positive Ce anomaly in the pisolitic samples, several 399 researchers (Braun et al., 1990; Nyakairu et al., 2001; Compton et al., 2003) have reported that 400 Ce is usually retained in the upper parts of the weathering profiles due to the oxidation of 401 Ce(III) to the less mobile Ce(IV). This supports the idea of in-situ bauxitization occurring from 402 the top down suggested by Yuste et al. (2015). The negative Ce anomaly recorded in some 403 pisolitic samples would be related to variable oxidation conditions favouring Ce(III) over Ce(IV), 404 probably related to the afore-mentioned acidic and reducing solutions, which promoted late 405 kaolinization stages. In fact, white pisolitic samples show a lower average Ce anomaly value. 406

407 Bardossy and Aleva (1990) stated that the textures and composition of most lateritic bauxites 408 can be used to relate them directly to the underlying source rocks, but this is not the case in 409 karst bauxites. In the present study, the pisolitic bauxite is derived from the clays with which 410 they are closely associated in the karst cavities they fill. As Mongelli (1993) (among others) has 411 indicated, the Eu anomaly has proven to be retained during intense weathering and the fact 412 that the observed Eu anomaly values in both the pisolitic and the clay samples are very similar 413 supports their genetic relationship. In addition, as pointed out above, Al, Ti, and Cr have 414 behaved as immobile elements during bauxitization and therefore the TiO₂/Al₂O₃ and Ti/Cr 415 ratios can be considered as sensitive indices of parental affinity. Plots of these ratios against Eu 416 anomaly values have been used in other Mediterranean karst bauxites to decipher the 417 parental affinity of the bauxites, that is, the bauxite deposits from Nurra (Italy) studied by 418 Mameli et al. (2007). As can be observed in Figure 10a, Eu/Eu* is similar to the UCC, but 419 TiO_2/Al_2O_3 departs from the UCC and tends towards more mafic compositions. This may be

- 420 due in part to the AI depletion relative to Ti deduced from the change % values, as commented
- 421 above. Similarly, Ti/Cr values also depart from the UCC with an average value half-way
- 422 between the UCC and the average basalt composition (Fig. 10b). This points to an intermediate
- 423 source with a parental affinity slightly more mafic than the UCC for the argillaceous sediments
- 424 from which first aluminous clays, and subsequently bauxite developed through intense
- 425 chemical weathering.
- 426

427 6. CONCLUSIONS

- 428
- 429 This research indicates that the studied karst bauxites are derived from the Al-rich clays with 430 which they are closely related. Bauxitization of the clays was the result of intense weathering 431 taking place from the top down. This prevented the lower parts of the weathering profile from 432 being bauxitized and resulted in the aluminous clays underlying the pisolitic bauxite. 433 Consequently, the clays geochemistry reflects variable weathering and a greater detrital 434 influence. Compared to ferrallitization, bauxitization led to chemical homogenization and 435 widespread higher element depletion. During the process, Al and Ti behaved as an immobile 436 element pair, whereas Fe was variably leached. Nevertheless, processes subsequent to 437 bauxitization caused some Al depletion in the bauxite, related to the partial replacement of 438 gibbsite by kaolinite. This replacement, which increased the loss of Fe, was favoured by the 439 circulation of acid solutions that triggered karst reactivation, which gave rise to the present 440 heterogeneous-chaotic lithostructure of the deposits. 441

UCC-normalized trace-element concentrations have revealed that bauxites are enriched in some elements compared to clays, including the critical element Ga. Nevertheless, in absolute terms, ferrallitization can be linked to Sc and V enrichment, whereas bauxitization has promoted chemical depletion, except for Zr, Cr, and probably the critical elements Nb and Hf, which can be considered immobile during the process. On the other hand, V and, to a lesser extent, Ge show behaviour similar to Fe, and therefore they were depleted during the late kaolinization process.

449

The higher degree of weathering related to bauxitization also involved the homogenization of REE+Y concentrations and their depletion during the transformation of ferrallitized clays into bauxites. This depletion is more pronounced for the LREE. The mobility and fractionation behaviour of the REE is linked to the concentration of minerals that act as scavengers for these elements. Ti oxides preferably scavenge the HREE, but other minerals, like fluorocarbonates of

455	the bastnäsite group, cannot be ruled out. Further research needs to be carried out on this
456	point since no fluorocarbonates have been detected yet in the studied karst bauxites. On the
457	other hand, the LREE seem to show a preference for P-bearing phases, more frequent in the
458	clay precursor of the bauxites.
459	
460	Intense chemical weathering, responsible for the development of the aluminous clays first and
461	subsequently the bauxites, did not modify the Eu anomaly. This anomaly, together with the Ti,
462	Al, and Cr ratios, suggests an intermediate character slightly more mafic than the UCC for the
463	argillaceous sediments from which the Al-rich clays and bauxites originated.
464	
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466	
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- 600

601 FIGURE CAPTIONS

- 602
- Figure 1. Geological framework of the study area located in the Maestrazgo zone (NE Spain)
- and geological sketch of the location of the study outcrops (stars). I-I' on the map corresponds
- to the cross-section below. M: Middle Jurassic, J1: Oxfordian, J2: Polpís Fm., A: Arenas de
- 606 Utrillas Fm., C: Cenomanian, T: Tertiary.
- 607 Figure 2. Field appearance of the heterogeneous-chaotic lithostructure of the study materials
- in one of the outcrops (locality 2 in Fig. 1) and hand specimens of red (left) and white (right)
- 609 pisolitic bauxites. Scale: coin is 2 cm in diameter.
- 610 Figure 3. a: Plot of Al_2O_3 vs. TiO₂. b: Fe₂O₃-SiO₂-Al₂O₃ diagram after Aleva (1994) of bauxite
- 611 chemical classification.
- 612 Figure 4. a: Average and range values of major element concentrations normalized to the
- 613 Upper Continental Crust (UCC) concentrations (Taylor and McLennan, 1985). b: Average and
- 614 range of major elements change % values.
- 615 Figure 5. a: Average and range values of some important trace elements normalized to the
- 616 UCC. b: Average and range change % values of trace elements.
- 617 Figure 6. a: Average and range REE+Y concentrations normalized to the UCC. b: Average and
- 618 range change % values of the REE+Y. c: Chondrite-normalized REE+Y patterns of the study
- 619 samples.
- 620 Figure 7. Plots of a: (La/Sm)c vs. (Gd/Yb)c, and b: (La/Yb)c vs. Ce/Ce*.
- 621 Figure 8. Plots of Cr and Zr vs. TiO_2 and Al_2O_3 and Zr vs. Cr.
- Figure 9. Backscattered electron (BSE) images of clay samples. a: Y-rich mineral (xenotime)
- 623 showing an irregular growth rim of probable apatite (Ap). b: Al phosphate-sulphate(APS)
- 624 mineral. Xen: xenotime; Ill: illite; Kln: kaolinite; Fe Ox.: iron oxides.
- Figure 10: Plots of Eu/Eu* vs. TiO₂/Al₂O₃ (a) and Ti/Cr (b). Granite and basalt compositions are
- 626 from Condie (1993).

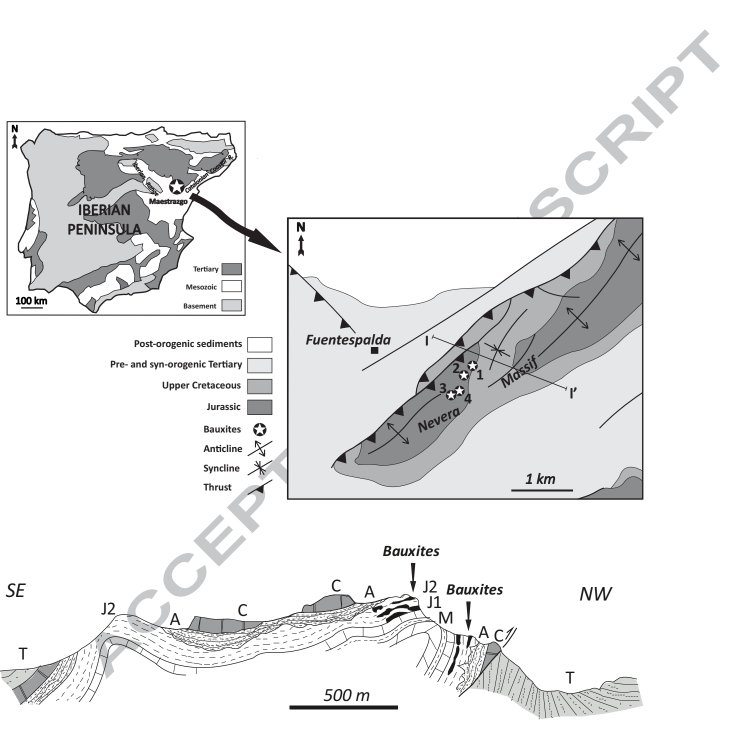


Fig. 1



Fig. 2

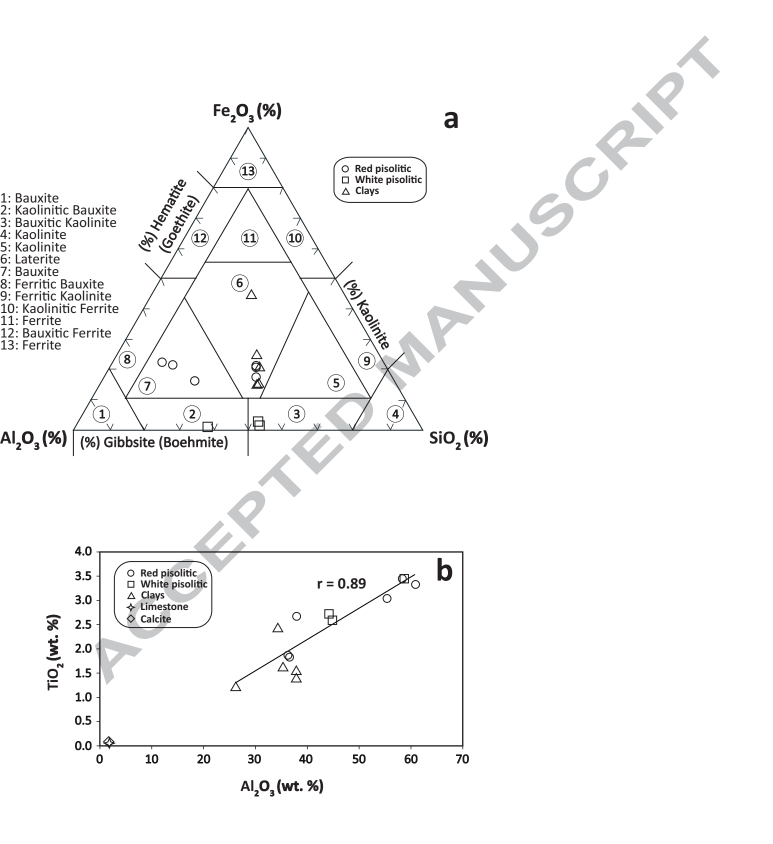
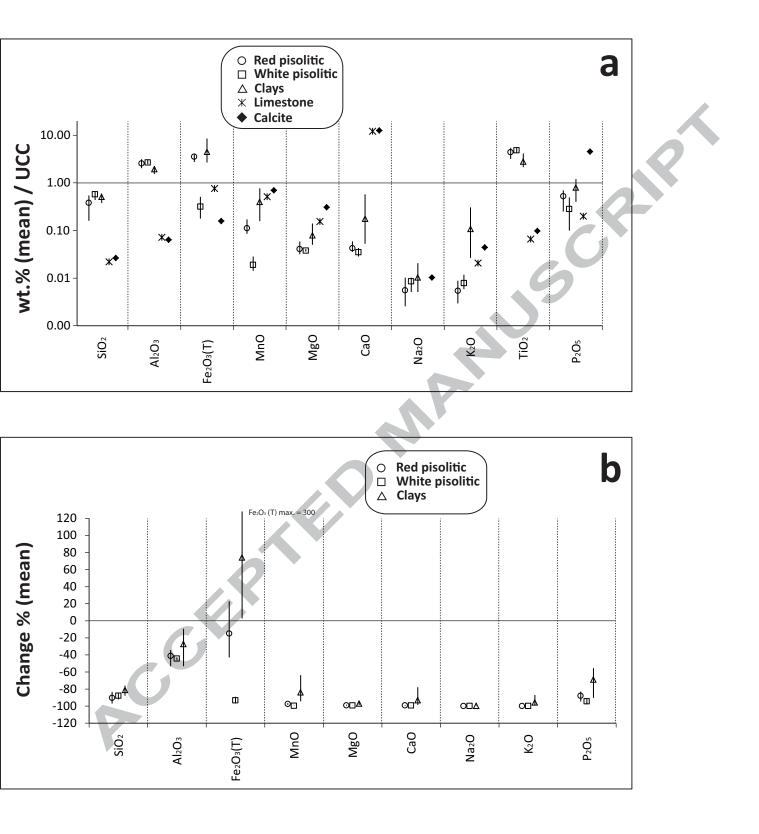
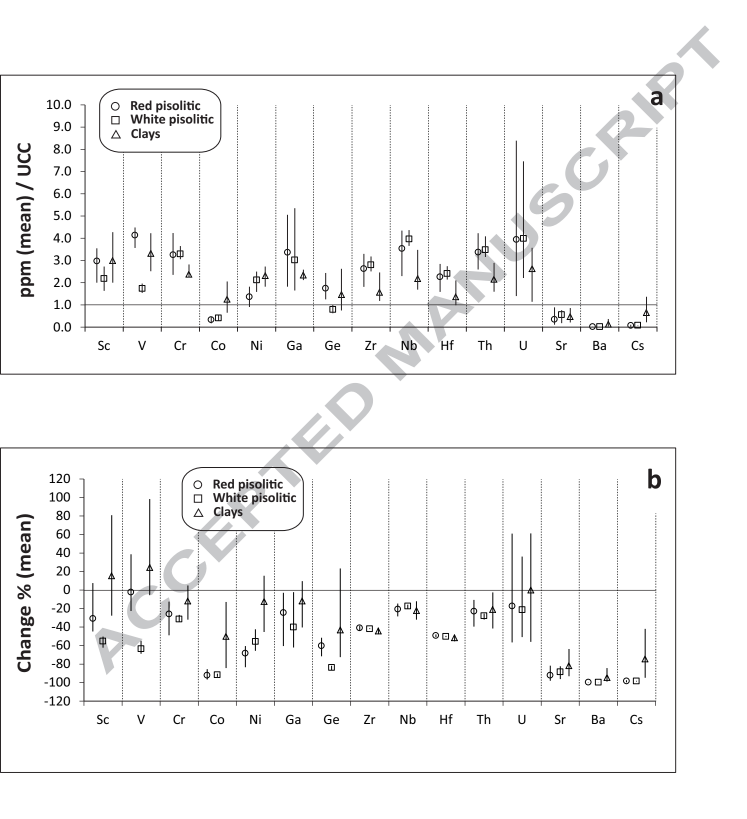


Fig. 3





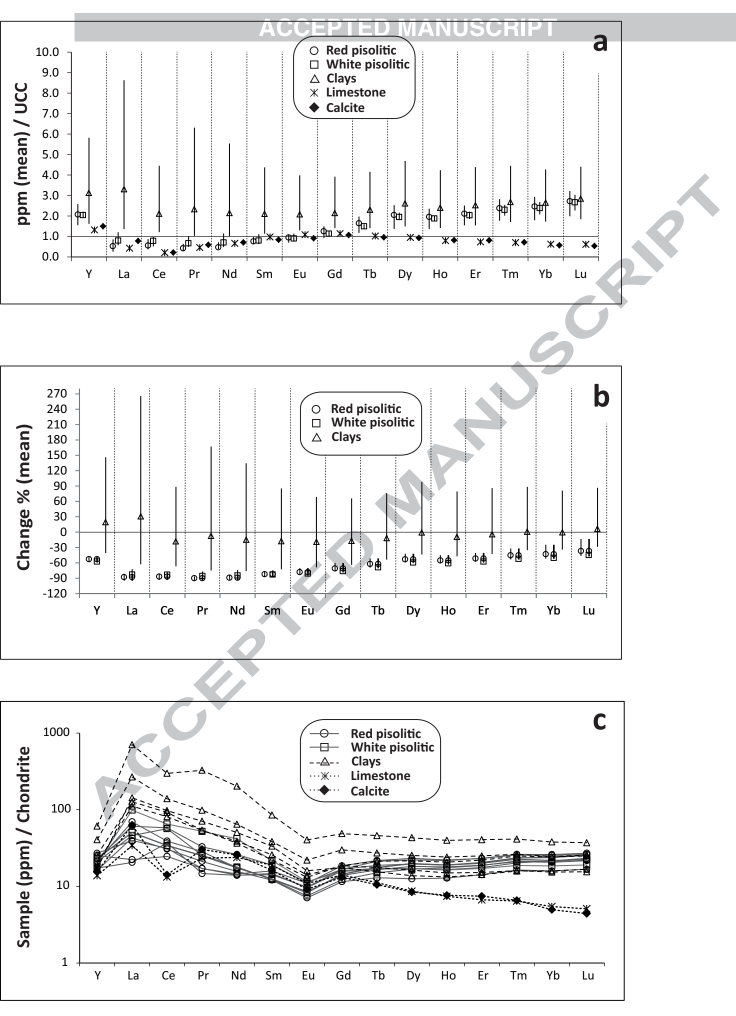


Fig. 6

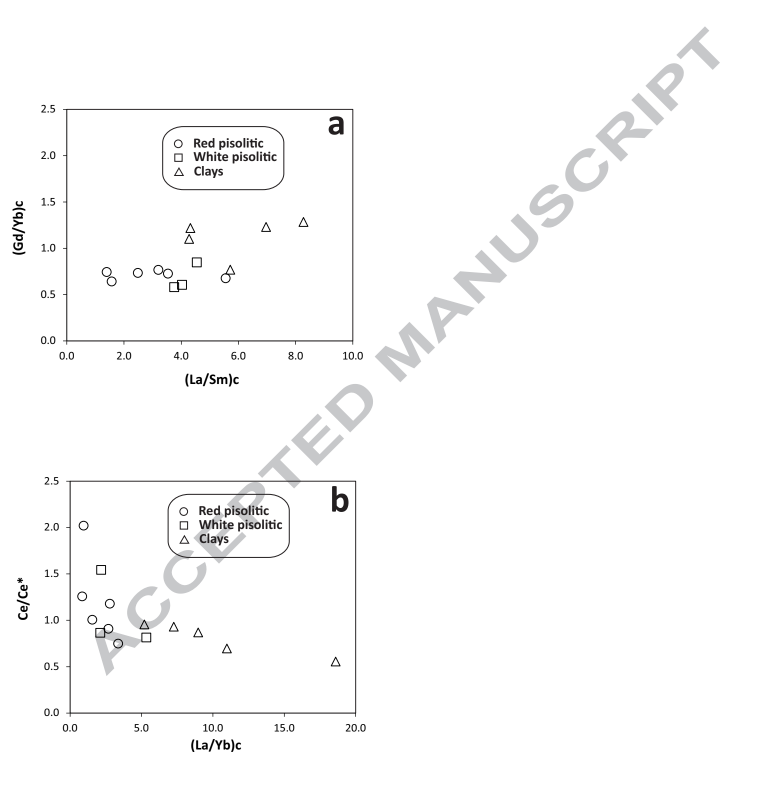


Fig. 7

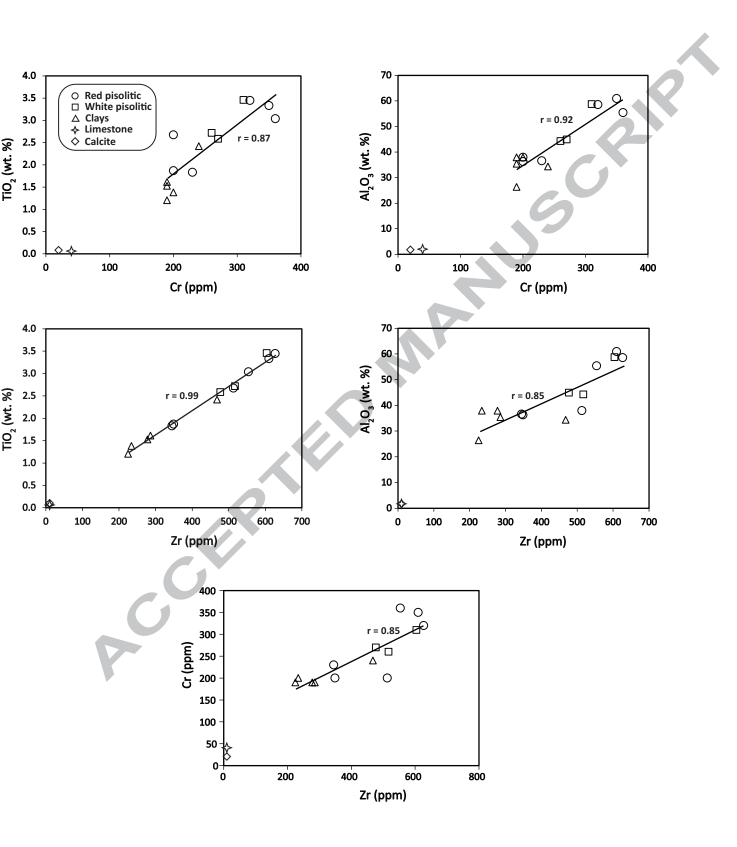
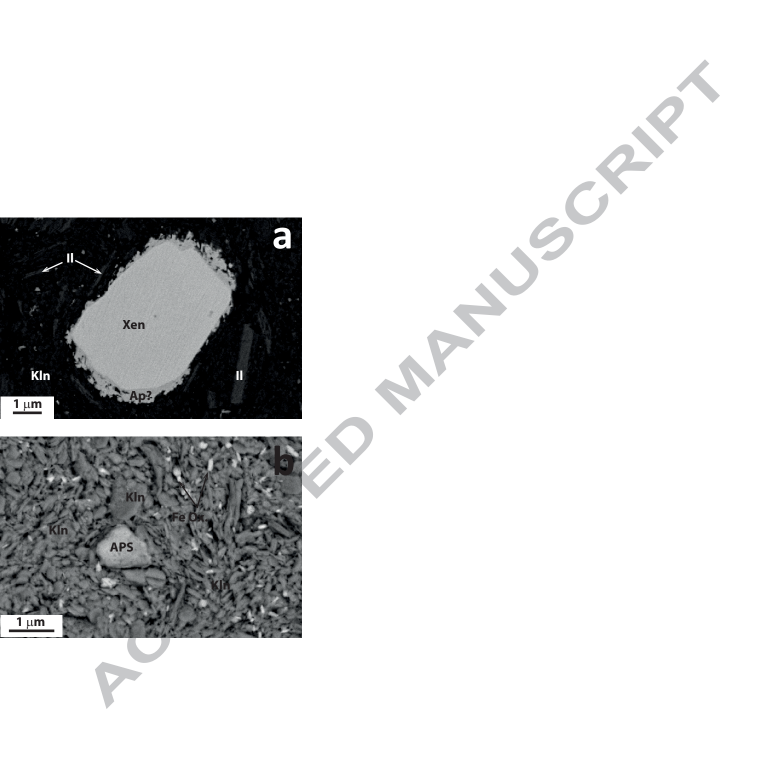


Fig. 8



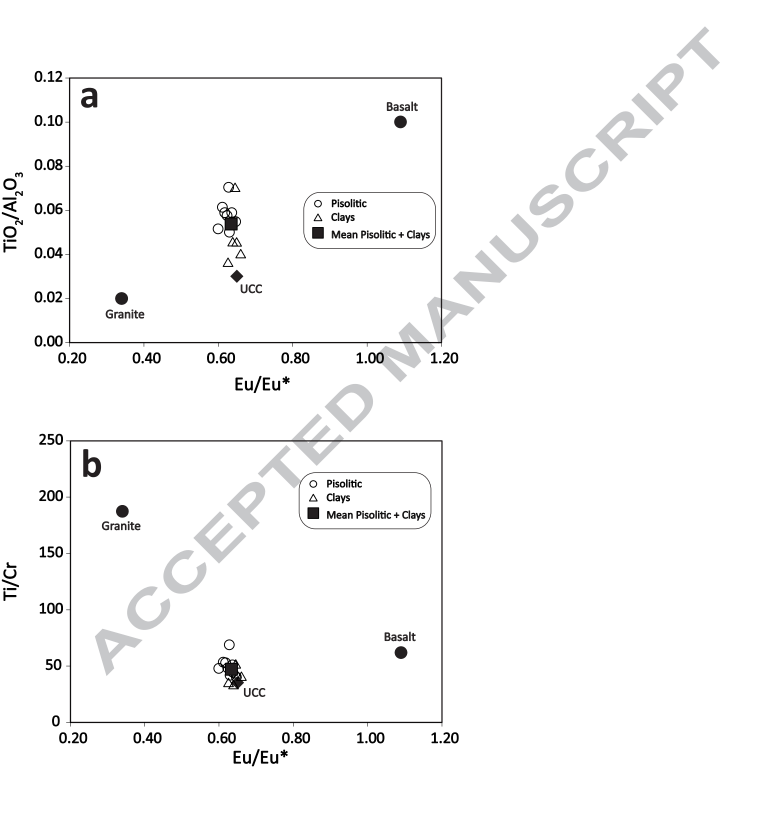


Fig. 10

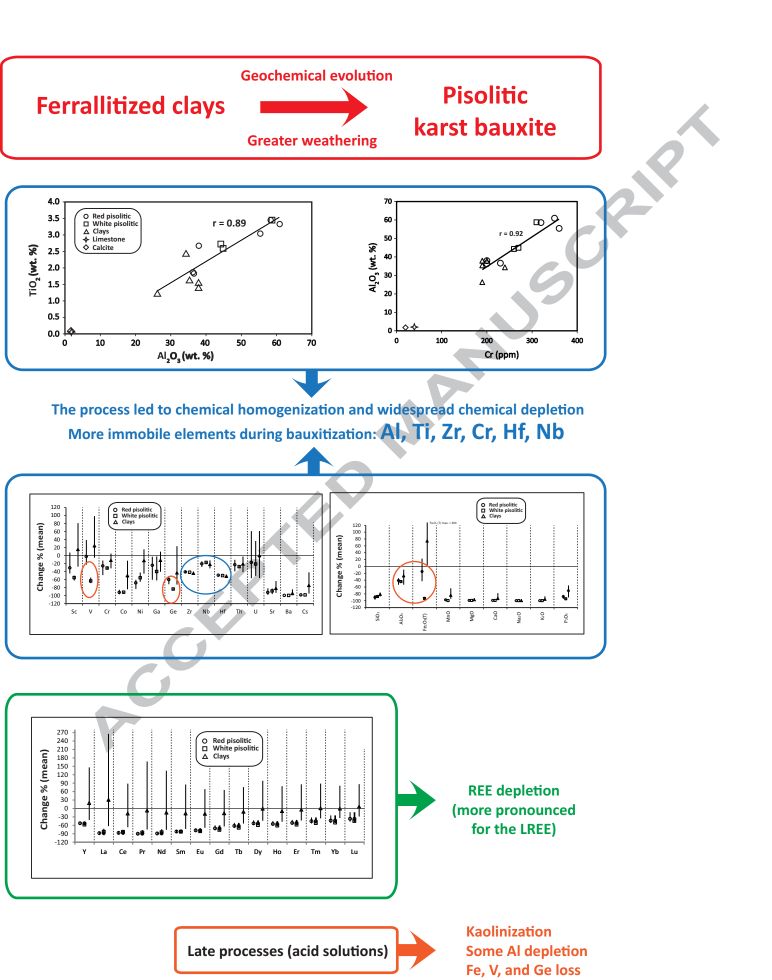
Table 1. Mineralogical composition (%) of the study samples (min.: minimum; max.: maximum; st. dv.: standard deviation). Kln: kaolinite; Gbs: gibbsite; Gt: goethite; Hem: hematite, Dsp: diaspore; Bhm: boehmite; Ant: anatase; Rt: ruitle; Ill: illite; Qtz: quartz; Cal: calcite; tr: traces.

					,	Whole	e samı	ple					< 2µm fraction		
sample description		Kln	Gbs	Gt	Hem	Dsp	Bhm	Ant	Rt	111	Qtz	Cal	III	Kln	
	mean	55	22	3	10	6								100	
Red pisolitic (n=6)	min.	28	0	0	8	5	tr	tr. tr. tr.		tr	-	tr.	100		
neu pisontic (n=0)	max.	83	45	7	12	8	и.		и.	-	и.	-	u.	100	
	st. dv.	22	22	3	1	1								0	
	mean	78	10	2		6								100	
Vhite pisolitic (n=3)	min.	57	0	0	-	5 7	tr. tr.	tr.	-	-	-	tr.	100		
	max.	92	31	5		7								100	
	st. dv.	15	15	2		1								0	
	mean	66		12		4				4		1	11	89	
Clays (n=5)	min.	44	-	0	0	0	0 9	tr.	tr.	0	tr.	0	0	60	
	max.	79		25	18	9				18		5	40	100	
	st. dv.	13		9	7	4				7		2	16	16	
											\$	P			
							Ù								

Table 2. Chemical composition of the analyzed samples. Major elements in weight per cent (wt. %) and trace elements in parts per million. wt. % oxides recalculated to 100 % on a volatile-free basis.

	sample	SiO₂	Al ₂ O ₃	Fe ₂ O ₃ (T)	TiO ₂	MnO	MgO	CaO	Na₂O	K ₂ O	P_2O_5	S	Sc	v	Cr	Со	Ni	Cu	Rb	Sr	Y	Zr	Nb	Cs	Ва	Hf
	F1-2	41.09	36.61	20.03	1.83	0.01	0.08	0.21	0.03	0.01	0.09	0.01	38	403	230	8	50	10	<1	101	36.3	345	27.6	0.4	13	9.2
	F1-3	40.65	36.26	20.66	1.87	0.01	0.15	0.19	0.05	0.04	0.13	0.01	22	475	200	5	50	10	<1	69	34	349	28.8	0.3	13	9.3
Red	F2-3	13.48	60.88	21.56	3.33	0.01	0.13	0.32	0.01	0.04	0.17	0.08	39	476	350	3	60	10	<1	39	56.7	610	52.1	0.4	14	15.6
pisolitic	F2-4	42.12	37.94	16.88	2.67	0.01	0.08	0.20	0.02	0.01	0.06	0.00	28	381	200	5	80	20	<1	134	48.1	513	47.1	0.1	6	13.5
	F3-1R	25.50	55.37	15.57	3.04	0.01	0.10	0.19	0.01	0.03	0.15	0.04	33	443	360	5	80	90	<1	312	44.7	554	49.1	0.4	11	14.6
	F4-2	16.94	58.53	20.53	3.45	0.01	0.11	0.20	0.03	0.01	0.18	0.02	36	480	320	8	40	< 10	<1	79	53	627	50.1	0.5	14	16.5
	F2-2	49.89	44.25	2.70	2.72	0.00	0.11	0.21	0.05	0.05	0.02	0.01	24	165	260	5	70	20	<1	63	43.9	517	46.6	0.4	12	13.6
White pisolitic	F2-6	50.70	44.89	1.42	2.58	0.00	0.10	0.17	0.05	0.02	0.06	0.01	18	210	270	7	110	30	<1	268	42.2	477	43.9	0.4	9	12.4
pisointic	F3-1B	36.38	58.72	1.01	3.46	0.00	0.10	0.15	0.03	0.03	0.13	0.01	30	183	310	9	100	140	<1	258	48.5	604	52.4	0.3	11	15.9
	F1-1	43.02	37.91	14.11	1.53	0.02	0.14	2.83	0.04	0.24	0.15	0.02	24	297	190	12	80	20	12	251	35.6	278	21.2	2.1	48	7.7
	F2-1	28.08	26.34	43.24	1.20	0.06	0.12	0.58	0.03	0.14	0.16	0.04	39	452	190	23	100	50	6	73	41.5	225	20.3	1.1	24	5.7
Clays	F2-5	38.91	34.33	23.66	2.42	0.03	0.14	0.29	0.02	0.10	0.09	0.01	33	420	240	11	100	30	5	100	54.1	468	41.7	1	23	12.2
	F1-4	41.27	35.35	20.18	1.61	0.01	0.36	0.36	0.05	0.44	0.28	0.09	22	337	190	26	110	80	22	99	85.1	286	22.9	4.5	71	7.8
	F3-2	43.80	37.88	14.83	1.38	0.04	0.25	0.26	0.09	1.21	0.25	0.01	47	269	200	35	120	100	52	300	128	234	24.9	6.3	203	6.1
li	mestone	2.52	1.90	5.97	0.06	0.06	0.59	88.68	0.00	0.12	0.07	0.03	12	36	40	<1	< 20	10	4	75	28.9	10	1.1	0.9	11	0.2
calcite	e cement	2.99	1.66	1.22	0.08	0.08	1.17	90.37	0.07	0.26	1.56	0.54	3	24	20	<1	< 20	< 10	7	45	32.7	10	< 0.2	1.3	12	0.3

	sample	Th	U	La	Ce	Pr	Nd	Sm	Eu	Gd	Tb	Dy	Но	Er	Tm	Yb	Lu	CIA	∑REE	∑LREE	ΣHREE	Eu/Eu*	Ce/Ce*	(La/Sm)c	(Gd/Yb)c	(La/Yb)c
	F1-2	30.7	3.92	7.53	35.4	2.01	9.91	3.39	0.837	4.86	1	6.5	1.41	4.38	0.721	5.31	0.888	99.81	84.15	58.24	25.07	0.63	2.02	1.40	0.74	0.96
	F1-3	27.8	3.95	15.7	29.3	2.9	11.1	2.79	0.615	3.52	0.75	4.78	1.09	3.54	0.582	3.94	0.635	99.68	81.24	61.79	18.84	0.60	0.91	3.54	0.72	2.69
Red	F2-3	45.2	23.5	14.5	30	2.34	10.3	3.67	0.951	5.67	1.26	8.83	1.88	5.75	0.919	6.27	0.996	99.90	93.34	60.81	31.58	0.64	1.01	2.49	0.73	1.56
pisolitic	F2-4	29.8	6.92	22.3	55.4	4.5	18.5	4.38	0.972	5.11	1.09	7.44	1.61	4.97	0.804	5.41	0.84	99.87	133.33	105.08	27.27	0.63	1.18	3.20	0.77	2.79
	F3-1R	39.9	13.7	25.5	37.3	3.89	12.4	2.89	0.742	4.25	0.95	6.93	1.52	4.69	0.746	5.11	0.825	99.91	107.74	81.98	25.02	0.65	0.75	5.55	0.67	3.37
	F4-2	43.3	14.2	8.19	23.5	2.35	9.96	3.28	0.852	5.1	1.23	8.4	1.85	5.77	0.931	6.45	1.03	99.91	78.89	47.28	30.76	0.64	1.26	1.57	0.64	0.86
White	F2-2	33.8	6.38	16.9	54.1	3.49	12.7	2.83	0.651	3.74	0.87	6.39	1.49	4.65	0.764	5.24	0.859	99.71	114.67	90.02	24.00	0.61	1.54	3.76	0.58	2.18
pisolitic	F2-6	34.3	6.2	36.2	62.1	7.14	29.7	5.01	0.999	4.78	0.97	6.48	1.39	4.27	0.666	4.58	0.726	99.77	165.01	140.15	23.86	0.62	0.81	4.55	0.85	5.34
	F3-1B	43.7	20.9	18.5	32.6	3.44	12.4	2.89	0.72	4.39	1.02	7.54	1.61	5.09	0.838	5.9	0.971	99.88	97.91	69.83	27.36	0.62	0.86	4.03	0.60	2.12
	F1-1	22.5	3.2	52.5	93.3	9.69	35.9	7.72	1.39	5.36	0.9	5.24	1.13	3.54	0.563	3.95	0.637	99.18	221.82	199.11	21.32	0.66	0.87	4.28	1.10	8.98
	F2-1	20.2	9.61	40.9	77.6	7.35	27.6	5.95	1.22	5.71	1.02	6.2	1.27	3.78	0.572	3.8	0.59	99.24	183.56	159.4	22.94	0.64	0.93	4.33	1.22	7.27
Clays	F2-5	30.9	8.69	46.4	88.2	7.39	26	5.11	1.14	5.7	1.23	8.2	1.76	5.48	0.887	6.02	0.949	99.56	204.47	173.1	30.23	0.65	0.96	5.72	0.77	5.21
	F1-4	17.1	5.29	98.1	133	13.5	45.9	8.86	1.91	9.15	1.58	9.66	2.05	6.22	0.932	6.03	0.973	98.47	337.87	299.36	36.60	0.65	0.70	6.97	1.23	10.99
	F3-2	24.6	10	259	285	44.8	144	19.7	3.51	14.9	2.66	16.4	3.39	10.1	1.47	9.41	1.41	96.27	815.75	752.5	59.74	0.63	0.55	8.28	1.28	18.60
li	mestone	1.08	1.74	12.5	12.7	3.2	16.9	4.35	0.947	4.3	0.65	3.29	0.63	1.66	0.229	1.35	0.194	-	62.90	49.65	12.30	0.67	0.43	1.81	2.58	6.26
calcite	e cement	1.04	6.03	23.1	13.5	4.16	18.2	3.74	0.795	4.02	0.61	3.21	0.65	1.85	0.233	1.23	0.168	-	75.47	62.7	11.97	0.63	0.28	3.89	2.65	12.69



628 HIGHLIGHTS

- 629 This paper focuses on the geochemical evolution from ferrallitized clays to karst
- 630 bauxites.
- 631 The process led to chemical homogenization and widespread element depletion.
- Al, Ti, Zr, Cr, Hf, and Nb behaved as more immobile elements during bauxitization.
- 633 Bauxitization enhanced REE depletion, more pronounced for the LREE.
- Late processes related to acid solutions caused Al depletion and Fe, V, and Ge loss.
- 635 636