



Weathering Products of a Dismantled Variscan Basement. Minero-Chemical Proxies to Insight on Cretaceous Palaeogeography and Late Neogene Palaeoclimate of Sardinia (Italy)

Paola Mameli^{1*}, Giovanni Mongelli², Rosa Sinisi² and Giacomo Oggiano¹

¹ Department of Chemistry and Pharmacy, University of Sassari, Sassari, Italy, ² Department of Sciences, University of Basilicata, Potenza, Italy

OPEN ACCESS

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*Correspondence:

Paola Mameli
mameli@uniss.it

Specialty section:

This article was submitted to
Geochemistry,
a section of the journal
Frontiers in Earth Science

Received: 19 March 2020

Accepted: 22 June 2020

Published: xx June 2020

Citation:

Mameli P, Mongelli G, Sinisi R and
Oggiano G (2020) Weathering
Products of a Dismantled Variscan
Basement. Minero-Chemical Proxies
to Insight on Cretaceous
Palaeogeography and Late Neogene
Palaeoclimate of Sardinia (Italy).
Front. Earth Sci. 8:290.
doi: 10.3389/feart.2020.00290

This study compares, for the first time, the mineralogy and geochemistry of two residual-clay deposits in NW Sardinia (Nurra district) that formed at different times in tropical and sub-tropical climates. Both deposits represent palaeosols with deep-weathered residual profiles and overlie Mesozoic carbonate rocks that were deposited on the south European palaeomargin. The older alterite is Cenomanian–Turonian in age and grades upward into a horizon of karstic bauxite, whereas the younger unit occurs within alluvial deposits of Late Neogene age. The Cretaceous palaeosol represents the precursor of the overlying bauxite and formed from unknown sedimentary parent rocks. In contrast, the Messinian weathering products formed by alluvium that was sourced from the Variscan metamorphic basement. Chemical Index of Alteration values, REE fractionation index values, and the results of R-mode factorial analysis suggest a common initial weathering path and a common precursor for the deposits. However, the latter stages of weathering of the Cretaceous palaeosols resulted in lateritic alteration and bauxite production, whereas weathering of the Late Neogene palaeosols produced 2:1 clay minerals under less extreme conditions. Comparison of these residual products constrains the parental material and weathering trends and allows insight into the relationship between palaeoclimate and regional/local palaeogeography of southern Europe during Upper Cretaceous and during Messinian.

Keywords: clays, chemical index of alteration, variscan basement, Cretaceous, Messinian

INTRODUCTION

Breakdown of rock at the Earth's surface produces a thin porous covering that represents a section of the recently defined "Critical Zone" (National Research Council, 2001; Brantley et al., 2006), and according to Eggleton (2001), referred to as "regolith." Typical regolith displays considerable vertical (distinguishable layers of weathered rocks) and lateral (variations in landscape and associated soils) heterogeneity (Taylor and Eggleton, 2001; Anderson et al., 2007). The rate and intensity of chemical weathering are controlled by various factors (Singer, 1984; Garzanti and Resentini, 2016), including vegetation, drainage of the parent rock, and climate.

115 Weathering products such as clays are an important component
116 of clastic sediments (e.g., siltstone, shale, and greywacke) and can
117 record the environmental conditions of the source area (Zabel
118 et al., 2001; Sinisi et al., 2014).

119 Clay mineral assemblages in residual and transported
120 sediments are controlled primarily by climatic conditions. Soils
121 and sediments at high latitudes and in cold climates generally
122 experience a low rate of chemical weathering that results in
123 the production of illite and chlorite association (Robert and
124 Maillot, 1990; Ehrmann and Mackensen, 1992; Ehrmann et al.,
125 2005). Conversely, soils and terrigenous sediments in tropical
126 wet regions undergo heavy leaching that produces kaolinite
127 and gibbsite assemblages (Macias Vazquez, 1981; Deepthy and
128 Balakrishnan, 2005; Bauluz et al., 2014). Furthermore, the degree
129 of weathering controls the accumulation/leaching of elements
130 from the precursor rocks to the residual material (Net et al.,
131 2002). Different elements show varying behavior during chemical
132 weathering. The low field strength elements (LFSE; e.g., Na, Ca,
133 and Sr) have a high affinity for aqueous phases and are therefore
134 easily removed from exposed rocks in wet regions (Nesbitt et al.,
135 1980; Nesbitt and Markovics, 1997). Other elements, such as
136 K, Rb, Mg, and Ba, are easily removed from primary minerals
137 but are often rapidly incorporated into secondary clay minerals
138 and insoluble hydroxides within weathering profiles and residual
139 deposits. Chemical weathering therefore redistributes major and
140 trace elements in rocks, particularly in sedimentary deposits.
141 During weathering, some elements are depleted in the parent
142 rock and residual products, whereas others are conservative and
143 are retained in moderately weathered rocks or concentrated in
144 residual deposits (Nesbitt et al., 1980; Peuraniemi and Pulkkinen,
145 1993; Condie et al., 1995; Nesbitt and Markovics, 1997).
146 Hence, mineralogical and geochemical investigations of alterites
147 (including palaeosols) can constrain both the palaeoclimate and
148 the nature of the parent rocks.

149 The present study compares the mineralogical and
150 geochemical features of two weathering products that were
151 generated at different times on the south European crust
152 (Sardinia-Corsica microplate).

153 In NW Sardinia (Nurra district), alterites overlie Mesozoic
154 carbonates and occur in two distinct stratigraphic contexts.
155 The older alterite deposit is overlain by Upper Cretaceous
156 karstic bauxites and rests above Upper Jurassic/Lower Cretaceous
157 carbonates. In contrast, the younger alterites occur as palaeosols
158 within a continental sequence of alluvial fan and braided river
159 deposits, late Miocene in age, which also overlie the same
160 Mesozoic carbonate sediments.

161 Between Lower and Upper Cretaceous, clayey alterites and
162 associated bauxite formed in a monsoonal climate on the south
163 European palaeomargin at 30°N palaeolatitude (Dercourt et al.,
164 1985; Mameli et al., 2007). The major questions on this clayey
165 material associated to bauxite are concerned with its precursor.
166 Indeed, the dissolution of high-grade carbonates is unlikely to
167 have generated the observed thick continuous layer of karstic
168 bauxite. Accordingly, it has been proposed that the bauxite
169 was derived from allochthonous debris, deposited on Mesozoic
170 limestone (MacLean et al., 1997), which in turn may have been
171 sourced from Variscan basement (Mameli et al., 2007). This

172 hypothesis, if confirmed, is consistent with an Upper Cretaceous
173 palaeogeographic setting in which portions of the carbonate shelf
174 were uplifted and eroded causing the basement to be exposed. In
175 contrast, the precursor of the Messinian alterites is known to be
176 debris derived from the Variscan metamorphic basement of NW
177 Sardinia (Mongelli et al., 2012), but what is still uncertain is the
178 palaeoclimate during the age of their formation.

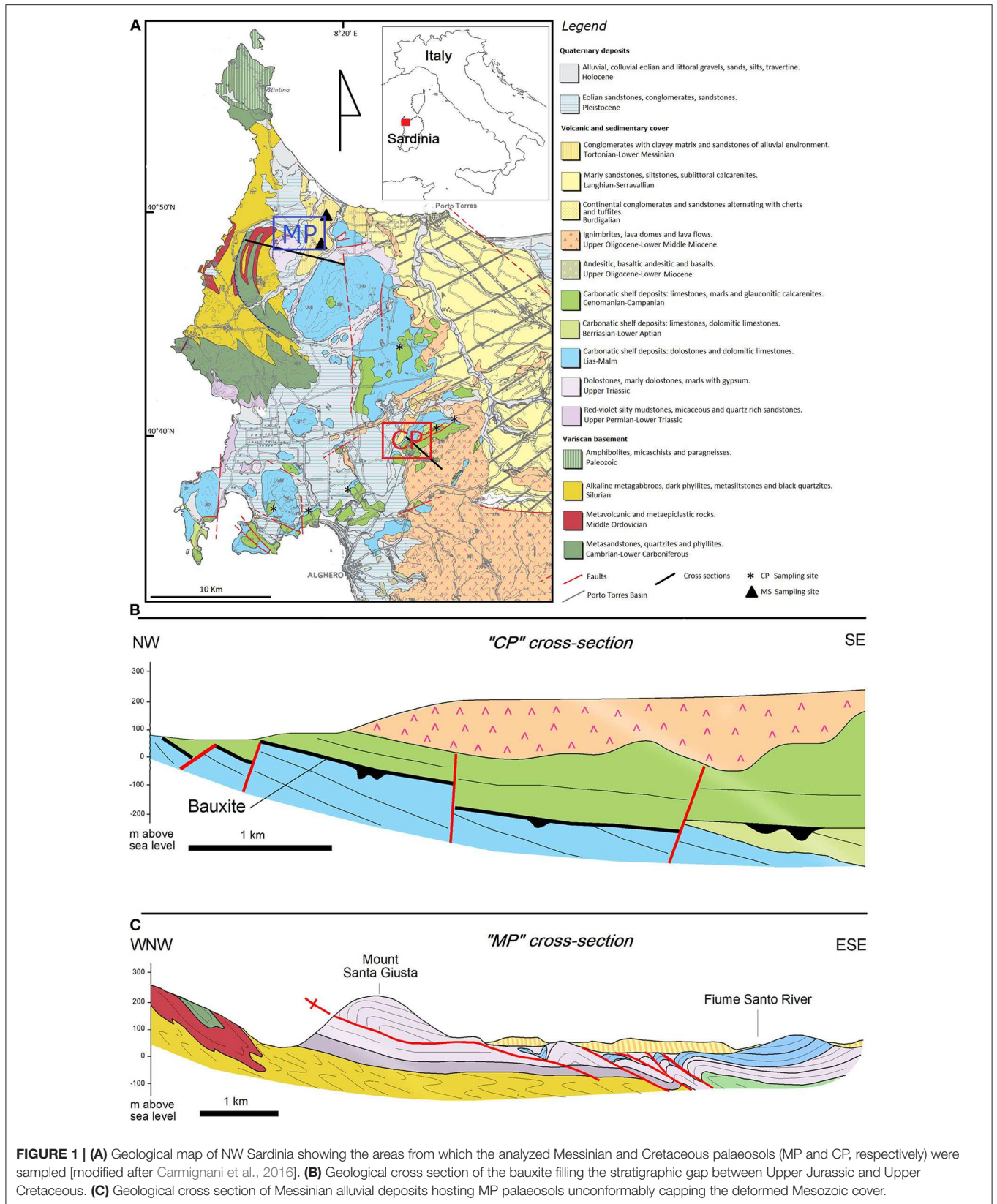
179 Messinian is an age known for the so called Mediterranean
180 salinity crisis (MSC—i.e., its partial desiccation), one of the most
181 appealing and debated palaeoclimatic and palaeogeographic
182 events in the geological history of this sea. It is still controversial
183 whether the desiccation is controlled by climatic or tectonic
184 factors (Fauquette et al., 2006; Achalhi et al., 2016). Most of
185 palaeoclimatic and palaeogeographic reconstructions are based
186 on palynological and isotopic data on continental and marine
187 sediments, respectively (Fauquette et al., 2006; Jiménez-Moreno
188 et al., 2013; Bertini and Menichetti, 2015). The attempts to test the
189 geochemical and mineralogical proxies on weathering product
190 are scanty (Mongelli et al., 2012) and are worthy of further
191 investigations able to give hints of palaeoclimatic relevance.

192 By comparison between the mineralogy and geochemistry of
193 the two alterites, we are expected to disclose new aspects in
194 Cretaceous palaeogeography and contribute in the assessment of
195 Messinian palaeoclimate.

196 In detail we aim to provide further insights into: (1)
197 the provenance of the Cretaceous alterite, which is the
198 parental material of bauxite; (2) the palaeogeography of
199 the south European margin during the Late Cretaceous; (3)
200 the palaeoclimatic conditions in the western Mediterranean
201 during the Messinian. Overall we also aim to contribute to
202 our understanding of the interplay among palaeoclimate and
203 precursor material during the weathering of Al-rich rocks to clay.
204

205 GEOLOGICAL SETTING 206

207 The Sardinian microplate was part of the South Europe margin
208 until the Aquitanian (Carmignani et al., 2004), before its counter-
209 clockwise drift and the opening of the Ligurian-Provencal
210 Basin in Burdigalian time (Thomas and Gennesseaux, 1986;
211 Gattaceca et al., 2007; Oudet et al., 2010). NW Sardinia
212 (**Figure 1A**) is characterized by a structural high caused by
213 the tilting of the Ligure-Provencal rift shoulder, during early
214 Miocene. In present coordinates the tilting results toward
215 the east; progressively older formations are therefore exposed
216 toward the west. The oldest rocks crop out near the coast
217 and comprise mainly phyllite with intercalated metarhyolite and
218 metabasite of the Variscan basement, which northward, due to
219 the prograde metamorphism, pass into gneiss, amphibolite, and
220 anatectic granite (Oggiano and Mameli, 2006; Cuccuru et al.,
221 2018). Late Permian and Mesozoic cover units are exposed
222 along a hilly/flat area to the east of the basement. Farther
223 east, the Mesozoic cover is overlain by Tertiary (Aquitanian–
224 Langhian) volcano-sedimentary successions of the Porto Torres
225 Basin (Funedda et al., 2000). The Mesozoic marine succession
226 is almost continuous from the Triassic (Ladinian) to the
227 uppermost Cretaceous (Campanian) and comprises evaporite,
228



dolostone, and limestone, with few intercalations of marlstone. There exists a stratigraphic gap embracing the Albian–Turonian interval, related to a period of emergence (Mameli et al.,

2007). This gap is represented by an almost continuous horizon of karstic bauxite, and it is constrained, on paleontological and micropaleontological bases (Philip et al., 1978; Philip,

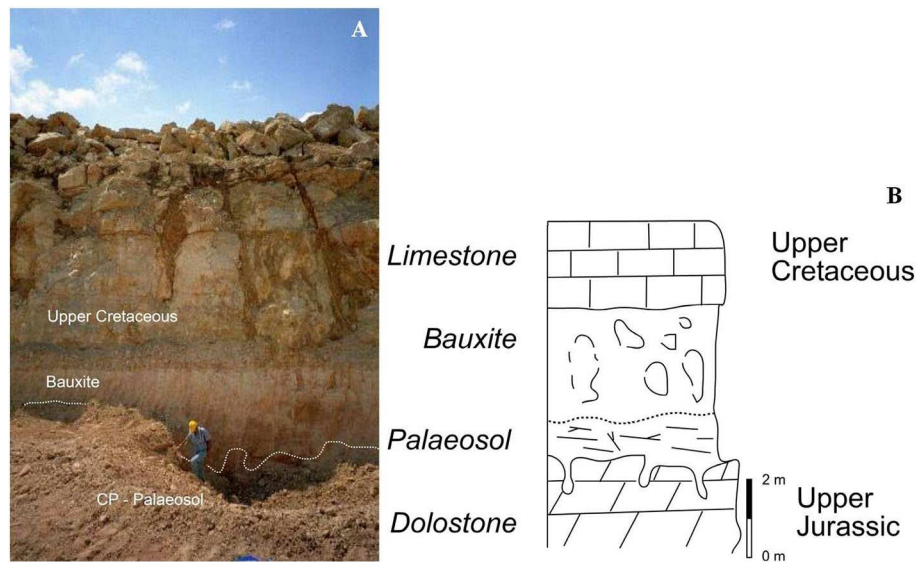


FIGURE 2 | (A) Clay-bearing CP palaeosol below bauxite at Olmedo mine. **(B)** Schematic stratigraphy of the Cretaceous deposits.

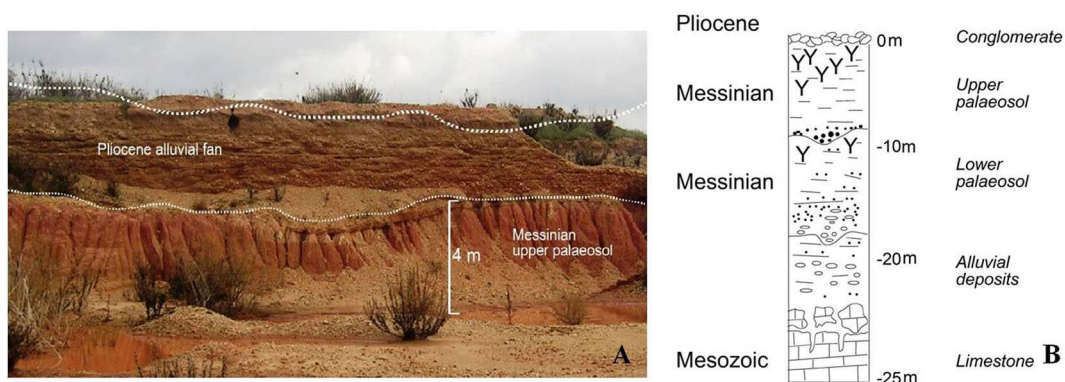


FIGURE 3 | (A) Continental deposit comprising channelized gravel and clay-rich horizons displaying evidence of pedogenesis (modified after Mongelli et al., 2015). **(B)** Sketch of the stratigraphy of the MP deposit.

1983) between the Aptian and the Coniacian. The bauxite unconformably overlies different rocks of different ages of the Mesozoic carbonate shelf that include pure limestone, dolostone, and minor marlstone (Figures 1B, 2A,B). Below the bauxite, green-gray and purple clay horizons locally occur, which display a mottled appearance and contain carbonaceous remnants of fossil plants. These clays are interpreted as palaeosols, where the green-gray portions with scattered carbonized vegetal remnants could represent reworked humic horizons. The passage into the overlying bauxite is gradational, marked by lowering in kaolinite, increasing Al_2O_3 and appearance of pisolites (Mameli et al., 2007).

The Messinian palaeosols are embedded within a wide alluvial system that linked up with upstream basement areas. The alluvial deposits unconformably overlie Mesozoic carbonate rocks that are underlain by Permian clastic sediments, which are the oldest, deposited on the Variscan basement (Figure 1C). Interbedded

channelized debris contains pebbles of metamorphic quartz, schist, and quartzite, consistent with being sourced from the adjacent basement. The alluvial deposit is locally several tens of meters thick (Figure 3A) and, together transported clay, hosts at least two horizons of residual clays, which feature as palaeosols. The primary palaeosol is well exposed along road cuts and quarries, and has a maximum thickness of 10 m. Another palaeosol horizon, which was intersected during coring and in one excavation, is generally separated from the former by a discontinuous gravelly/sandy deposit (Figure 3B). In the excavation the transition between the two palaeosols is a horizon of fine sands that yielded a tusco-sardinian land mammal palaeofauna of Turolian age, leading to consider as Messinian the upper palaeosol (Abbazzi et al., 2008).

The upper palaeosol consists of reddish clay (5/8–2Y; mottled with gray in reduced areas) to ochre (8/8–7.5Y) with rare rhizoliths (Figure 4), and locally is cut by overlying, channelized

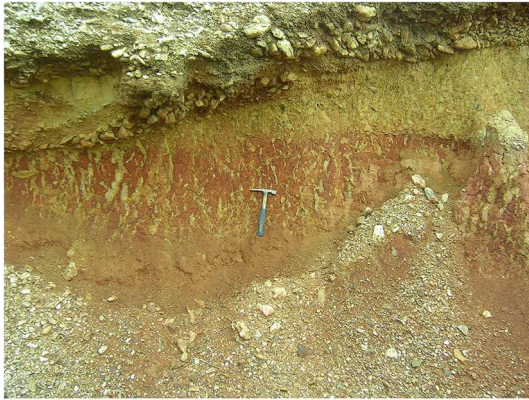


FIGURE 4 | Rhizoliths within pedogenized clay (Mongelli et al., 2015).

quartz-bearing gravel deposit. This deposit consists of mild sorted, quartz-dominated gravel, clast supported, with no matrix or with fine sandy matrix. The gravel clasts are sub-rounded to sub-angular. The particle size ranges from 1.2 cm pebbles to 10–15 cm, cobbles in places cemented by manganese oxyhydroxides. This poorly graded, relatively disorganized sediment can be referred to streamflows-dominated alluvial fan, possibly of Pliocene age.

MATERIALS AND METHODS

We performed mineralogical and geochemical analyses on 34 representative samples: 12 from the Cretaceous palaeosols (CP) and 22 from the Messinian palaeosols (MP). CP samples were collected in two vertical profiles (3 per each profile from bottom up to the boundary with bauxite) in a mined open pit, other 6 come from individual surficial outcrops scattered over a wide area, in order to get samples representative of different Mesozoic bedrocks. The upper Messinian palaeosol was sampled in two quarry cuts (samples MP1–MP12) at different levels, the lower (MP 13–MP 22) in two cores drilled during brick clay prospecting.

Whole-rock samples were dried and reduced to fine powder in a planetary mill (Retsch) equipped with two agate jars and agate milling balls. Mineralogical analysis was performed on randomly oriented whole-rock powders using a Siemens D5000 powder X-ray diffractometer (Cu-K α radiation, 40 kV, and 30 mA) at 2θ angles of 6–70° and a step size of 0.02°. Mineral phases were identified from powder XRD patterns using the ICDD PDF-2 database and the Bruker DIFFRACplus EVA 14.2 software package.

Major, trace, and rare earth element concentrations were measured by inductively coupled plasma–optical emission spectrometry (ICP-OES) and inductively coupled plasma–mass spectrometry (ICP-MS) at Activation Laboratories, Ancaster, Canada. The powder samples were dissolved by fusion with lithium metaborate/tetraborate and the resulting molten bead was rapidly digested in a weak nitric acid solution or a multiacid one (for Cu, Zn, and Ni determination), as expected from 4Litho and 4B1 packages. GXR-1, NIST 694, DNC-1, GXR-4,

SDC-1, GXR-6, LKSD-3, TDB-1, NOD-P-1, W-2a, DTS-2b, SY-4, CTA-AC-1, BIR-1a, NCS DC86312, ZW-C, NCS DC70009 (GBW07241), OREAS 100a (Fusion), OREAS 101a (Fusion), OREAS 134a (Fusion), and JR-1 were the standards used. Analytical uncertainties were <5%, except for elements occurring at concentrations of ≤ 10 ppm, which yield uncertainties of 5–10%. Loss on ignition (LOI) was estimated using gravimetric analysis following combustion at 950°C.

The texture, mineralogy, and microchemistry of polished sections of representative samples were analyzed using an ESEM Zeiss LaB6 scanning electron microscope equipped with an energydispersion spectrometer (EDS).

The samples were then labeled according to their provenance.

The degree of weathering of the palaeosols was estimated through calculation of the chemical index of alteration (CIA = $\text{Al}_2\text{O}_3 / (\text{Al}_2\text{O}_3 + \text{CaO}^* + \text{Na}_2\text{O} + \text{K}_2\text{O}) \times 100$; where elements are in molecular proportions and CaO* represents CaO hosted in silicate minerals only; Nesbitt and Young, 1982). CIA values were not calculated for samples with CaO concentrations higher than that of Post Archaean Australian Shales (PAAS) (1.3 wt.%; Taylor and McLennan, 1985).

Inter-elemental relationships were determined using element variation diagrams and R-mode factor analysis, performed with XLStat software package. In detail we used varimax rotation and subsequent R-mode factor analysis using principal components in the two sets of palaeosol samples. A standardized correlation matrix with equal weighting for all variables was used, enabling conversion of the principal component vectors into factors. Factor weightings were determined separately for the Cretaceous and Messinian samples. For both datasets, we only discuss those variables with absolute weightings of >0.65.

RESULTS

Mineralogy

XRD patterns from whole-rock disoriented powders indicate that all samples are composed of dominantly clay minerals and quartz, with minor Al and/or Fe-oxyhydroxides and Ti-oxides.

The main mineral assemblage in the CP samples comprises clay minerals (kaolinite + illite \pm montmorillonite) + quartz + Al/Fe-oxyhydroxides (boehmite/gibbsite + goethite + minor hematite) + Ti-oxides (anatase and/or rutile) (Figure 5).

The main mineral assemblage in the MP samples comprises clay minerals (illite + kaolinite) + quartz + Al/Fe-oxyhydroxides (gibbsite + goethite) + Ti-oxides (rutile + minor anatase) \pm feldspar. Palygorskite occurs in three core samples from the lower palaeosol and halite was detected in four samples (Figure 6).

The minerals phases are generally well-resolved except for some Cretaceous samples from Olmedo mine and some core samples from the Messinian deposit.

In addition to XRD, SEM EDS observation evidenced: (i) the occurrence of tiny bipyramidal magnetite crystals in the Al rich MP1 and MP2 samples; (ii) the coexistence of detrital, irregularly edged, kaolinite plates with tiny, exagonal, growing crystal stacks; (iii) the survival of detrital muscovite laminae in some quartz rich millimetric layers within the upper Messinian palaeosol; (iv) the occurrence of detrital kaolinite also in the Cretaceous clays.

Chemistry

The major and trace element concentrations of the analyzed sets of samples are listed in **Table 1**. Major elements and their variability are summarized below.

CP samples yield SiO₂ concentrations of 32.09–43.94 wt.% (mean = 39.10 wt.%; *n* = 12), indicating depletion relative to

the Upper Continental Crust (UCC) [66.62 wt.%; (Rudnick and Gao, 2014) and references therein] and post-Archaean Australian shale (PAAS) (62.80 wt.%; Taylor and McLennan, 1985). The samples contain 23.44–31.64 wt.% Al₂O₃ (mean = 26.98 wt.%), more than the UCC (15.40 wt.%) and PAAS (18.90 wt.%). They contain 7.90–17.24 wt.% Fe₂O₃ (mean = 12.77 wt.%), 0.25–4.28

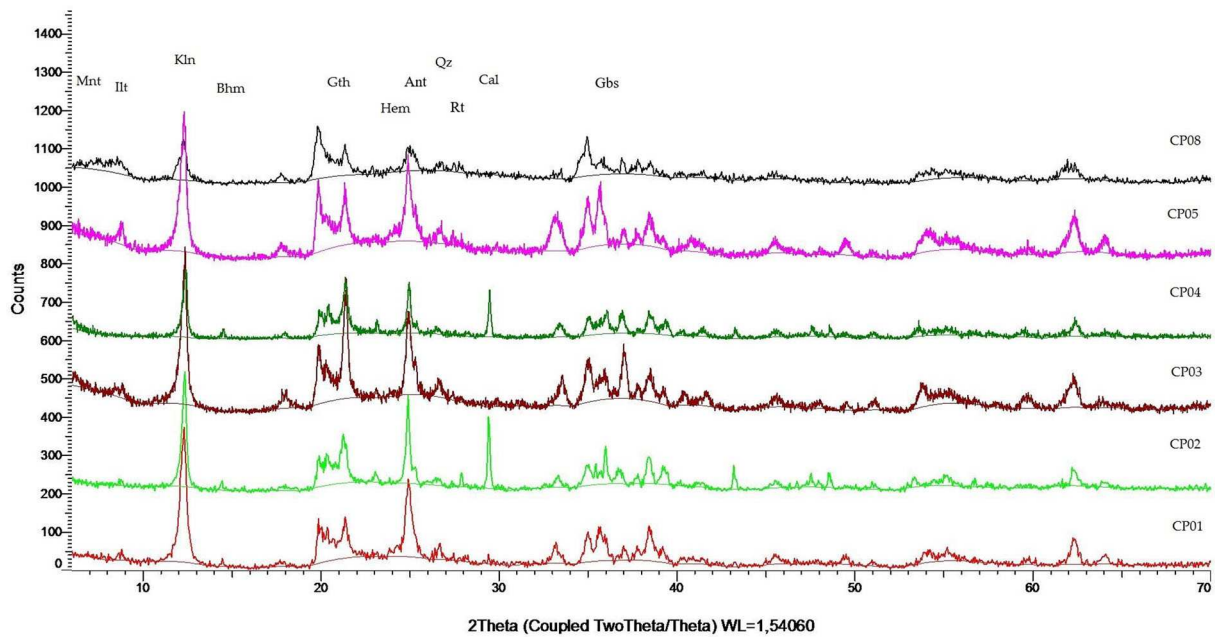


FIGURE 5 | XRPD patterns of representative CP palaeosol samples. Legend. Kln, Kaolinite; Illt, Illite; Mnt, Montmorillonite; Plg, Palygorskite; Qz, Quartz; Fsp, Feldspar; Cal, Calcite; Rt, Rutile; Ant, Anatase; Hem, Hematite; Gth, Goethite; Gbs, Gibbsite; Bhm, Boehmite; Hl, Halite.

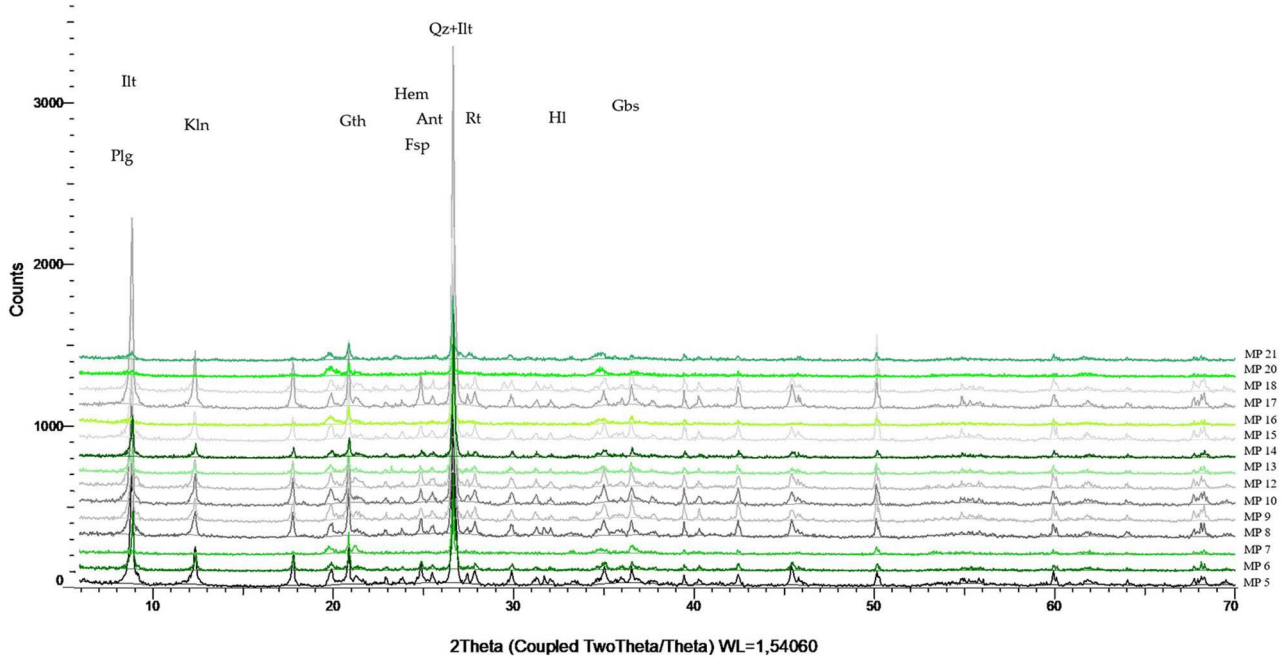


FIGURE 6 | XRPD patterns of representative MP palaeosol samples. Legend. See **Figure 5**.

TABLE 1 | Chemical compositions of the analyzed samples.

Samples	CP01	CP02	CP03	CP04	CP05	CP06	CP07	CP08	CP09	CP10	CP11	CP12	Mean	Median	O	
wt%																
D.L.																
Al ₂ O ₃	0.01	31.64	27.00	29.43	28.07	29.02	26.73	25.59	27.85	25.18	24.32	25.49	23.44	26.98	26.87	2.35
CaO	0.01	0.72	4.51	0.24	4.70	0.25	0.37	0.37	0.50	0.34	0.42	0.41	0.85	1.14	0.42	1.63
Fe ₂ O ₃ (T)	0.01	13.64	17.24	17.18	15.14	15.30	11.56	12.54	9.62	10.51	12.33	10.25	7.90	12.77	12.44	3.00
K ₂ O	0.01	1.01	0.50	1.56	0.25	1.93	3.13	3.72	3.68	3.93	3.73	4.00	4.28	2.64	3.41	1.49
MgO	0.01	0.29	0.42	0.89	0.47	0.84	1.41	1.59	1.54	1.78	1.65	2.02	2.11	1.25	1.48	0.64
MnO	0.001	0.009	0.049	0.018	0.077	0.020	0.017	0.024	0.006	0.022	0.024	0.021	0.015	0.025	0.021	0.019
Na ₂ O	0.01	0.16	0.14	0.15	0.12	0.19	0.25	0.23	0.20	0.24	0.22	0.21	0.38	0.21	0.21	0.07
P ₂ O ₅	0.01	0.04	0.07	0.06	0.04	0.06	0.05	0.11	0.05	0.09	0.12	0.09	0.08	0.07	0.07	0.03
SiO ₂	0.01	36.82	32.09	35.42	34.04	37.52	39.70	39.83	43.94	42.30	40.91	42.93	43.64	39.10	39.77	3.92
TiO ₂	0.001	1.722	1.517	1.625	1.203	1.570	1.532	1.444	1.391	1.371	1.308	1.327	1.298	1.442	1.418	0.153
LOI	0.01	12.15	15.25	14.17	16.48	12.88	14.09	13.18	10.31	13.40	13.92	13.25	14.40	13.62	13.66	1.54
Total		98.21	98.78	100.70	100.60	99.57	98.84	98.63	99.07	99.16	98.95	99.99	98.39	99.24	99.01	0.81
SiO ₂ /Al ₂ O ₃		1.16	1.19	1.20	1.21	1.29	1.49	1.56	1.58	1.68	1.68	1.68	1.86	1.47	1.52	0.24
ppm																
Ba	2	152	66	126	95	190	233	226	198	209	231	203	216	179	201	56
Co	1	9	59	33	23	21	23	19	17	18	21	17	27	24	21	13
Cr	20	190	140	230	300	220	180	180	130	170	170	160	200	189	180	45
Cs	0.5	5.4	4.0	10.0	1.0	12.3	20.5	19.4	22.2	22.7	21.5	21.7	21.3	15.2	20.0	8.1
Cu	1	5	34	4	12	9	6	13	15	10	11	14	95	19	12	25
Ga	1	38	29	30	38	32	42	38	36	38	40	32	35	36	37	4
Hf	0.2	10.7	12.3	13.2	8.5	13.5	11.1	10.2	10.5	10.8	10.7	8.7	7.9	10.7	10.7	1.8
Nb	1	35	35	29	23	30	30	23	35	32	30	22	35	30	30	5
Ni	1	89	265	168	103	167	165	246	98	150	211	122	179	164	166	56
Pb	5	75	30	50	35	111	37	110	87	56	74	42	68	65	62	28
Rb	2	44	37	71	11	94	151	159	191	186	178	184	196	125	155	69
Sc	1	33	27	33	45	36	29	28	26	29	28	27	24	30	29	6
Sr	2	77	33	99	22	118	109	167	84	119	122	111	140	100	110	41
Th	0.1	21.5	23	27.2	26.8	25.6	21	16.6	20.1	19.1	21.1	17.5	19.6	21.6	21.1	3.5
U	0.1	8.1	10.1	8.2	4.1	8.4	6.3	4.8	4	5.6	6.3	4.1	4.9	6.2	6.0	2.0
V	5	342	442	407	310	331	250	272	195	213	298	197	209	289	285	82
Y	1	39	99	105	80	63	91	73	38	142	118	86	54	82	83	31
Zn	1	42	77	76	22	82	120	152	85	142	165	128	181	106	103	50
Zr	2	405	456	398	287	399	438	460	404	433	439	377	333	402	405	51
La	0.1	46.2	227.0	392.0	66.1	118.0	124.0	212.0	60.4	316.0	324.0	184.0	79.3	179.1	154.0	116.7
Ce	0.1	207.0	185.0	463.0	205.0	251.0	670.0	156.0	90.4	208.0	188.0	108.0	108.0	236.6	196.5	167.3
Pr	0.05	9.23	58.60	82.80	23.50	28.40	28.80	59.90	10.70	89.70	91.10	46.70	19.60	45.75	37.75	30.24
Nd	0.1	32.2	210.0	257.0	84.7	102.0	110.0	221.0	34.9	330.0	331.0	173.0	75.8	163.5	141.5	106.3
Sm	0.1	7.2	35.0	46.9	20.5	24.7	25.3	41.2	6.2	62.7	62.1	30.7	15.1	31.5	28.0	19.0
Eu	0.05	1.70	7.47	9.43	4.53	5.36	5.10	7.14	1.34	11.60	11.20	5.52	3.07	6.12	5.44	3.38
Gd	0.1	7.4	31.2	38.7	15.6	19.0	20.4	22.9	5.5	44.6	41.2	21.5	13.4	23.5	21.0	12.9
Tb	0.1	1.4	4.1	4.9	2.5	3.1	3.3	3.2	1.2	5.7	5.4	2.9	2.1	3.3	3.2	1.5
Dy	0.1	8.7	20.6	23.7	15.8	18.2	18.3	17.2	8.0	30.8	28.8	15.3	12.6	18.2	17.7	7.1
Ho	0.1	1.7	3.7	4.5	3.2	3.4	3.4	3.0	1.7	5.5	5.3	2.8	2.4	3.4	3.3	1.2
Er	0.1	5.5	10.4	12.8	9.5	10.1	10.2	9.0	5.9	15.6	15.0	8.1	7.6	10.0	9.8	3.2
Tm	0.05	0.92	1.47	1.80	1.37	1.66	1.62	1.42	1.11	2.11	2.09	1.18	1.09	1.49	1.45	0.39
Yb	0.1	6.4	9.0	10.4	8.3	10.8	10.9	9.5	8.4	12.9	13.0	7.4	7.0	9.5	9.3	2.2
Lu	0.01	0.96	1.31	1.54	1.21	1.68	1.79	1.45	1.31	1.86	1.97	1.10	1.08	1.44	1.38	0.33
ΣREE		336.51	804.85	1349.47	461.81	597.4	1033.11	764.91	237.06	1137.07	1120.16	608.2	348.14	733.22	686.56	362.85
(La/Yb) _{ch}		4.88	17.04	25.47	5.38	7.38	7.69	15.08	4.86	16.55	16.84	16.80	7.66	12.14	11.38	6.66
(La/Sm) _{ch}		4.04	4.08	5.26	2.03	3.01	3.08	3.24	6.13	3.17	3.28	3.77	3.31	3.70	3.29	1.09
(Gd/Yb) _{ch}		0.94	2.81	3.02	1.52	1.43	1.52	1.95	0.53	2.80	2.57	2.35	1.55	1.92	1.75	0.80
Eu/Eu*		0.71	0.69	0.68	0.77	0.76	0.69	0.71	0.70	0.67	0.68	0.66	0.66	0.70	0.69	0.04
Ce/Ce*		2.35	0.38	0.60	1.22	1.02	2.63	0.32	0.83	0.29	0.26	0.27	0.64	0.90	0.62	0.81
CIA		91.95		90.82		88.90	82.31	79.39	80.58	78.38	78.37	78.20	73.68	82.26	79.99	6.17

(Continued)

TABLE 1 | Continued

Samples	MP01	MP02	MP03	MP04	MP05	MP06	MP07	MP08	MP09	MP10	MP11	MP12	Mean	Median	O	
wt%																
D.L.																
Al ₂ O ₃	0.01	32.66	30.45	26.92	25.42	22.95	22.24	19.70	21.71	21.25	21.80	20.99	21.13	23.94	22.02	4.10
CaO	0.01	0.10	0.10	0.13	0.08	0.08	0.27	0.70	0.61	0.17	0.10	0.20	0.20	0.23	0.15	0.21
Fe ₂ O ₃ (T)	0.01	1.62	3.69	9.81	7.34	8.31	7.94	9.17	7.98	7.17	7.06	8.17	7.98	7.19	7.96	2.30
K ₂ O	0.01	5.38	5.06	4.20	4.37	3.79	3.91	3.44	3.68	3.47	3.87	3.65	3.37	4.02	3.83	0.64
MgO	0.01	0.58	0.63	0.98	0.62	0.67	0.72	2.26	0.81	0.54	0.61	0.61	0.64	0.81	0.64	0.47
MnO	0.001	0.006	0.014	0.131	0.014	0.022	0.223	0.258	0.131	0.339	0.095	0.240	0.143	0.135	0.131	0.111
Na ₂ O	0.01	1.00	0.88	0.65	0.73	1.28	0.55	0.15	1.31	0.69	0.64	0.61	0.66	0.76	0.68	0.32
P ₂ O ₅	0.01	0.07	0.05	0.09	0.06	0.07	0.06	0.08	0.08	0.06	0.06	0.13	0.12	0.08	0.07	0.02
SiO ₂	0.01	49.82	50.15	46.38	52.60	53.97	53.79	48.92	54.34	60.47	58.25	56.69	57.16	53.55	53.88	4.19
TiO ₂	0.001	1.074	1.068	1.006	1.200	1.065	1.012	0.785	1.034	1.141	1.145	1.101	1.169	1.067	1.071	0.108
LOI	0.01	7.73	7.55	10.00	7.46	6.76	8.62	15.13	8.89	5.67	5.63	7.12	6.32	8.07	7.51	2.57
Total		100.10	99.63	100.30	99.89	98.97	99.34	100.60	100.60	101.00	99.26	99.50	98.89	99.84	99.76	0.69
ppm																
Ba	2	772	695	570	614	896	683	303	626	677	620	636	521	634	631	142
Co	1	2	7	33	7	11	25	40	21	35	16	29	19	20	20	12
Cr	20	120	130	90	120	80	110	130	80	80	80	100	70	99	95	22
Cs	0.5	20.8	18.3	24.0	15.7	11.7	11.2	11.7	12.7	11.6	12.4	10.9	12.0	14.4	12.2	4.3
Cu	1	10	24	43	36	34	27	29	34	31	30	30	30	30	30	8
Ga	1	40	39	34	34	30	28	51	31	29	29	28	29	34	31	7
Hf	0.2	3.2	3.7	5.1	5.4	6.0	6.0	5.8	6.4	6.6	6.0	8.7	7.7	5.9	6.0	1.5
Nb	1	27	27	27	32	20	19	15	22	22	22	20	24	23	22	5
Ni	1	19	21	70	33	54	49	69	79	101	50	55	55	54	52	24
Pb	5	157	29	36	20	35	32	39	40	34	41	42	42	46	38	36
Rb	2	262	239	208	204	176	199	186	185	171	183	190	176	198	188	27
Sc	1	24	24	23	21	21	20	20	20	19	19	19	19	21	20	2
Sr	2	297	197	169	162	138	133	72	138	119	135	128	127	151	137	55
Th	0.1	11.7	13.5	16.1	18.9	18.0	17.7	12.9	17.5	18.7	15.9	24.3	21.3	17.2	17.6	3.6
U	0.1	2.7	2.9	3.4	3.6	3.3	3.9	3.3	3.4	3.7	3.2	4.4	3.9	3.5	3.4	0.5
V	5	161	169	221	160	151	150	216	148	141	144	170	178	167	161	26
Y	1	14	17	25	29	42	35	40	49	52	41	60	50	38	41	14
Zn	1	23	36	107	49	62	88	74	85	90	74	95	99	74	80	26
Zr	2	114	135	188	192	221	186	198	233	240	228	269	294	208	210	51
La	0.1	157.0	32.0	68.1	43.5	61.8	54.1	79.4	59.2	77.3	54.6	93.3	75.7	71.3	65.0	31.7
Ce	0.1	113.0	52.2	134.0	79.5	121.0	121.0	176.0	123.0	144.0	111.0	170.0	156.0	125.1	122.0	35.4
Pr	0.05	19.10	6.51	13.10	9.78	13.60	13.60	18.50	13.20	20.10	12.40	23.60	17.80	15.11	13.60	4.80
Nd	0.1	46.2	22.7	41.4	35.1	49.6	45.1	61.0	48.4	77.9	45.4	78.4	66.0	51.4	47.3	16.6
Sm	0.1	5.2	4.0	6.6	6.2	9.4	9.0	11.4	9.4	15.0	8.8	14.9	12.4	9.4	9.2	3.6
Eu	0.05	0.96	0.81	1.33	1.27	2.03	1.93	2.33	2.12	3.49	2.00	3.16	2.68	2.01	2.02	0.83
Gd	0.1	3.9	3.9	6.2	6.2	8.0	7.7	8.6	8.9	13.8	8.0	12.7	10.4	8.2	8.0	3.1
Tb	0.1	0.5	0.5	0.9	0.9	1.4	1.2	1.3	1.5	2.1	1.3	1.9	1.6	1.3	1.3	0.5
Dy	0.1	2.8	3.2	5.0	5.5	7.8	6.9	7.1	8.1	11.4	7.1	10.8	9.1	7.1	7.1	2.7
Ho	0.1	0.5	0.6	1.0	1.1	1.5	1.4	1.4	1.6	2.1	1.4	2.2	1.7	1.4	1.4	0.5
Er	0.1	1.7	1.9	2.8	3.2	4.4	3.9	4.1	4.6	6.1	4.0	6.2	5.1	4.0	4.1	1.4
Tm	0.05	0.27	0.31	0.45	0.49	0.66	0.59	0.58	0.68	0.90	0.61	0.89	0.75	0.6	0.60	0.20
Yb	0.1	1.8	2.1	3.0	3.2	4.4	3.7	3.5	4.3	5.7	4.0	5.6	4.7	3.8	3.9	1.2
Lu	0.01	0.29	0.32	0.46	0.47	0.63	0.56	0.50	0.62	0.82	0.58	0.83	0.71	0.57	0.57	0.17
ΣREE		353.22	131.05	284.34	196.41	286.22	270.68	375.71	285.62	380.71	261.19	424.48	364.64	301.19	285.92	83.68
(La/Yb) _{ch}		58.94	10.30	15.34	9.19	9.49	9.88	15.33	9.30	9.16	9.22	11.26	10.88	14.86	10.09	14.06
(La/Sm) _{ch}		19.00	5.04	6.49	4.42	4.14	3.78	4.38	3.96	3.24	3.91	3.94	3.84	5.51	4.05	4.33
(Gd/Yb) _{ch}		1.76	1.51	1.67	1.57	1.47	1.69	1.99	1.68	1.96	1.62	1.84	1.79	1.71	1.68	0.16
Eu/Eu*		0.65	0.63	0.64	0.63	0.72	0.71	0.72	0.71	0.74	0.73	0.70	0.72	0.69	0.71	0.04
Ce/Ce*		0.48	0.85	1.05	0.90	0.98	1.05	1.08	1.03	0.86	1.00	0.85	1.00	0.93	0.99	0.16
CIA		76.64	76.63	77.85	76.13	74.56	75.41	75.09	72.08	76.23	75.46	75.43	76.57	75.67	75.80	1.43

(Continued)

TABLE 1 | Continued

Samples	MP13	MP14	MP15	MP16	MP17	MP18	MP19	MP20	MP21	MP22	Mean	Median	σ	Mean T	Median T	σ T	
wt%																	
D.L.																	
Al ₂ O ₃	0.01	20.51	20.54	21.26	17.92	20.14	19.47	18.94	19.29	16.65	16.41	19.11	19.38	1.65	21.74	21.13	4.00
CaO	0.01	0.22	0.14	0.11	0.46	0.15	0.96	0.12	0.67	0.51	0.49	0.38	0.34	0.29	0.30	0.20	0.25
Fe ₂ O ₃ (T)	0.01	7.69	7.67	7.64	8.68	7.08	7.35	7.62	8.85	7.72	4.96	7.53	7.66	1.05	7.34	7.67	1.81
K ₂ O	0.01	4.02	3.46	3.60	3.44	3.50	3.38	3.29	3.95	5.64	2.91	3.72	3.48	0.74	3.88	3.68	0.69
MgO	0.01	0.99	0.59	0.55	1.94	0.58	0.65	0.69	2.71	2.69	0.51	1.19	0.67	0.90	0.98	0.65	0.71
MnO	0.001	0.027	0.034	0.090	0.152	0.097	0.173	0.171	0.103	0.076	0.048	0.097	0.094	0.054	0.118	0.103	0.090
Na ₂ O	0.01	0.52	0.60	0.57	0.28	0.80	1.13	1.30	0.17	0.29	0.79	0.65	0.59	0.37	0.71	0.66	0.34
P ₂ O ₅	0.01	0.09	0.07	0.07	0.16	0.08	0.08	0.10	0.08	0.18	0.05	0.10	0.08	0.04	0.09	0.08	0.03
SiO ₂	0.01	56.18	57.28	60.81	53.07	59.86	59.13	59.12	48.47	56.00	65.40	57.53	58.20	4.59	55.36	56.00	4.73
TiO ₂	0.001	1.213	1.038	1.064	0.803	1.055	1.058	1.052	0.701	0.853	0.970	0.981	1.045	0.152	1.028	1.055	0.134
LOI	0.01	8.78	7.23	5.10	13.01	6.45	6.35	6.69	14.75	9.16	6.70	8.42	6.97	3.13	8.23	7.46	2.78
Total		100.30	98.66	100.90	99.91	99.79	99.72	99.09	99.74	99.76	99.25	99.71	99.75	0.62	99.78	99.76	0.65
ppm																	
Ba	2	484	488	533	300	611	642	529	251	326	525	469	507	133	559	570	159
Co	1	10	9	14	20	32	32	24	21	16	7	19	18	9	20	20	11
Cr	20	140	120	80	90	70	70	80	120	100	80	95	85	24	97	90	23
Cs	0.5	11.8	10.1	12.5	10.4	12.2	11.8	10.0	12.0	9.2	8.8	10.9	11.1	1.3	12.8	11.8	3.7
Cu	1	27	26	31	30	30	36	29	30	14	24	28	30	6	29	30	7
Ga	1	27	26	29	24	28	28	25	51	36	23	30	28	8	32	29	8
Hf	0.2	7.5	7.6	6.5	6.3	6.9	6.5	7.8	5.5	4.7	10.8	7.0	6.7	1.6	6.4	6.4	1.6
Nb	1	23	19	21	16	21	21	27	13	15	18	19	20	4	21	21	5
Ni	1	43	40	66	55	61	84	54	67	54	39	56	55	14	55	54	19
Pb	5	14	21	30	31	43	63	27	37	14	35	32	31	15	39	35	28
Rb	2	193	171	174	162	165	166	160	195	154	150	169	166	15	185	183	27
Sc	1	19	18	19	17	18	18	16	20	15	14	17	18	2	19	19	3
Sr	2	117	122	128	75	129	125	126	67	91	105	109	120	23	132	128	48
Th	0.1	19.8	22.5	18.5	13.9	19.9	18.0	22.9	13.0	10.3	20.3	17.9	19.2	4.2	17.5	18.0	3.8
U	0.1	3.9	3.5	3.2	3.3	3.6	3.7	3.8	3.2	2.3	4.1	3.5	3.6	0.5	3.5	3.5	0.5
V	5	142	132	138	156	132	134	134	208	133	103	141	134	27	156	150	29
Y	1	39	38	43	35	46	54	46	48	33	35	42	41	7	40	40	11
Zn	1	72	84	68	86	84	92	93	69	68	121	84	84	16	78	84	22
Zr	2	248	272	240	217	254	240	283	191	164	355	246	244	52	226	228	54
La	0.1	61.8	82.2	66.1	61.3	66.9	61.8	80.2	44.2	47.0	70.4	64.2	64.0	12.2	68.1	66.1	24.6
Ce	0.1	123.0	162.0	131.0	107.0	142.0	131.0	153.0	148.0	95.4	144.0	133.6	136.5	20.7	129.0	131.0	29.3
Pr	0.05	13.40	19.10	15.10	14.70	15.40	13.70	19.20	11.60	10.70	15.90	14.88	14.90	2.78	15.00	14.70	3.93
Nd	0.1	45.0	64.6	54.7	50.1	56.2	50.5	71.4	41.5	36.1	51.7	52.2	51.1	10.4	51.8	50.1	13.8
Sm	0.1	8.7	12.2	10.2	9.7	10.7	9.7	12.5	9.8	7.4	9.8	10.1	9.8	1.5	9.7	9.7	2.8
Eu	0.05	1.90	2.45	2.19	2.10	2.35	2.19	2.57	2.21	1.69	2.01	2.17	2.19	0.26	2.08	2.10	0.63
Gd	0.1	7.5	9.0	8.6	8.0	9.3	9.2	12.0	9.5	7.0	7.8	8.8	8.8	1.4	8.5	8.5	2.4
Tb	0.1	1.3	1.3	1.4	1.1	1.5	1.5	1.6	1.4	1.1	1.3	1.4	1.4	0.2	1.3	1.3	0.4
Dy	0.1	7.6	7.7	7.8	6.3	8.5	8.2	9.0	8.4	6.1	7.4	7.7	7.8	0.9	7.4	7.6	2.1
Ho	0.1	1.6	1.6	1.5	1.3	1.7	1.6	1.7	1.7	1.2	1.5	1.5	1.6	0.2	1.5	1.5	0.4
Er	0.1	4.7	4.6	4.3	3.7	4.9	4.8	4.9	4.8	3.5	4.6	4.5	4.7	0.5	4.2	4.4	1.1
Tm	0.05	0.70	0.68	0.65	0.55	0.73	0.70	0.73	0.67	0.49	0.67	0.66	0.68	0.08	0.63	0.66	0.16
Yb	0.1	4.5	4.3	4.2	3.4	4.7	4.4	4.7	4.0	2.9	4.1	4.1	4.3	0.6	4.0	4.1	1.0
Lu	0.01	0.64	0.62	0.62	0.48	0.67	0.65	0.69	0.59	0.45	0.65	0.61	0.63	0.08	0.58	0.62	0.14
ΣREE		282.34	372.35	308.36	269.73	325.55	299.94	374.19	288.37	221.03	321.83	306.37	304.15	46.11	303.54	299.94	67.72
(La/Yb) _{ch}		9.28	12.92	10.64	12.18	9.62	9.49	11.53	7.47	10.95	11.60	10.57	10.79	1.62	12.91	10.64	10.46
(La/Sm) _{ch}		4.47	4.24	4.08	3.98	3.94	4.01	4.04	2.84	4.00	4.52	4.01	4.02	0.46	4.83	4.04	3.24
(Gd/Yb) _{ch}		1.35	1.70	1.66	1.91	1.60	1.69	2.07	1.92	1.96	1.54	1.74	1.70	0.22	1.73	1.69	0.19
Eu/Eu*		0.72	0.71	0.71	0.73	0.72	0.71	0.64	0.70	0.72	0.70	0.71	0.71	0.02	0.70	0.71	0.04
Ce/Ce*		1.00	0.96	0.97	0.84	1.04	1.05	0.91	1.53	1.00	1.01	1.03	1.00	0.19	0.97	1.00	0.18
CIA		73.71	76.12	76.36	73.70	74.76	70.47	72.66	72.49	62.66	72.21	72.51	73.18	3.90	74.24	75.09	3.19

$(La/Yb)_{ch} = (La/La_{ch})/(Yb/Yb_{ch})$; $(La/Sm)_{ch} = (La/La_{ch})/(Sm/Sm_{ch})$; $(Gd/Yb)_{ch} = (Gd/Gd_{ch})/(Yb/Yb_{ch})$; $Eu/Eu^* = Eu/Eu_{ch}/\sqrt{(Sm/Sm_{ch} \times Gd/Gd_{ch})}$; $Ce/Ce^* = Ce/Ce_{ch}/\sqrt{(La/La_{ch} \times Pr/Pr_{ch})}$. D.L., Detection Limit. In red are reported mean, median and standard deviation relative to the whole CP and MP sample sets, respectively; in blue are reported mean, median and standard deviation relative to upper and lower sub-sets of the MP palaeosol.

wt.% K₂O (mean = 2.64 wt.%), 0.12–0.38 wt.% Na₂O (mean = 0.21 wt.%), and 0.29–2.11 wt.% MgO (mean = 1.25 wt.%). Two samples yield relatively high CaO concentrations (>4.50 wt.%) due to the presence of carbonate (calcite) fragments, which were detected in XRD patterns.

MP samples, as a whole, yield SiO₂ concentrations of 46.38–65.40 wt.% (mean = 55.36 wt.%; *n* = 22), lower than UCC and PAAS. The samples contain 16.41–32.66 wt.% Al₂O₃ (mean = 21.74 wt.%), more than the UCC and PAAS. They also contain 1.62–9.81 wt.% Fe₂O₃ (mean = 7.34 wt.%), 2.91–5.64 wt.% K₂O (mean = 3.88 wt.%), 0.08–0.96 wt.% CaO (mean = 0.30 wt.%), and 0.51–2.71 wt.% MgO (mean = 0.98 wt.%). Four samples yield relatively high Na₂O concentrations (>1.10 wt.%) due to the presence of halite, which was detected in XRD patterns.

DISCUSSION AND EVALUATION OF PALAEOWEATHERING

By the comparison of the two alterites the purpose of this work is to obtain information on the parental rock of a palaeosol whose boundary palaeoclimatic conditions are known (CP) and, on the other hand, to get information on the palaeoclimatic conditions from a palaeosol of which, instead, the parent rock is well identified (MP). Beyond the mechanisms of alteration, the cross-comparison of the mineralogical and geochemical features of the two alterites can reveal new insights on the Cretaceous palaeogeographic frame of the south European margin and, on the other end, add new constraints on the climatic scenario during Messinian time.

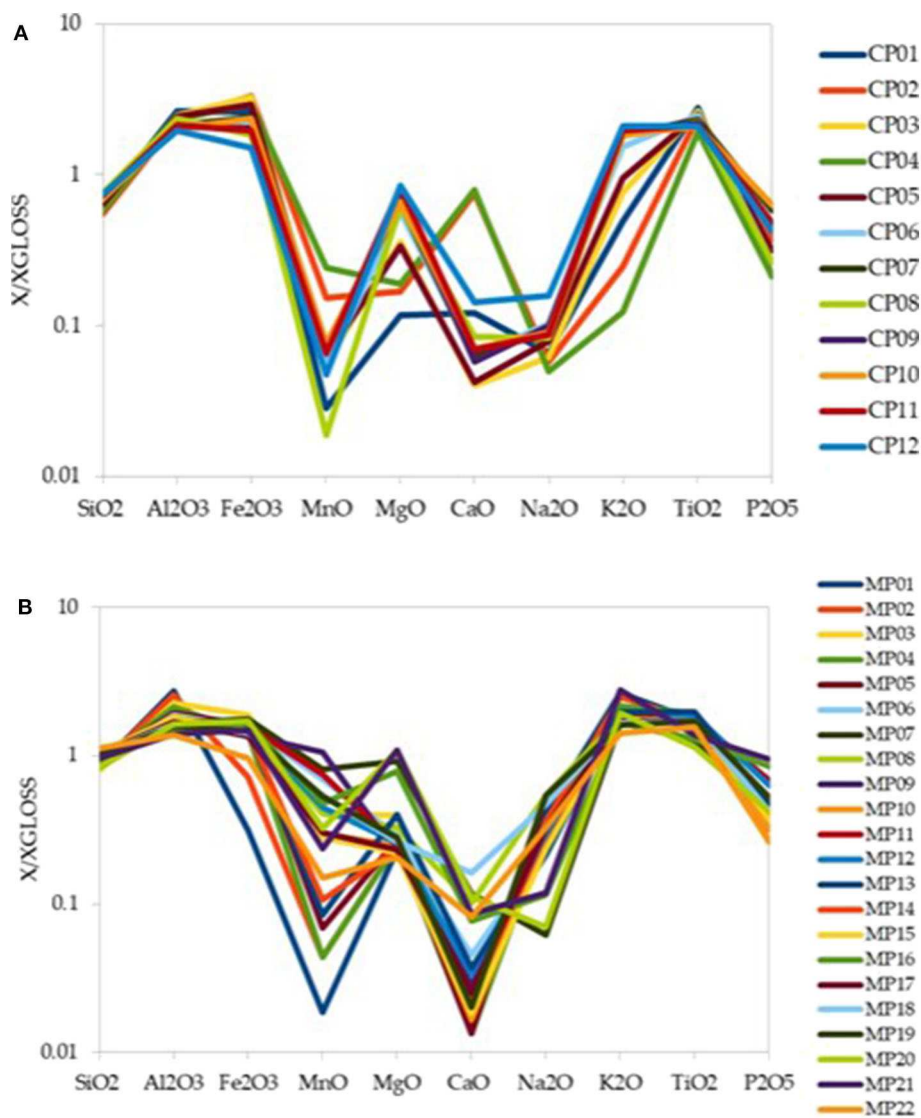


FIGURE 7 | Global Subducting Sediment (GLOSS)-normalized major elements patterns of (A) the CP palaeosols, and (B) the MP palaeosols. GLOSS values are from Plank and Langmuir (1998).

In fact few is known about the mechanism that drove the emergence of the Mesozoic shelf at the beginning of Upper Cretaceous (Albian-Turonian) generating bauxite and associated palaeosols. Eustatism and tectonics could both account for such emergence (Durand et al., 1988; Combes and Peybernes, 1991; Haq and Huber, 2017; Jež and Otoničar, 2018). In the first case the thick carbonate shelf must have experienced weak erosion; whereas in the second case an important uplift could have caused the complete erosion of the carbonate covers leading to the exposition of the basement.

Uncertainties also affect the Messinian scenario dominated by the salinity crisis (Gardner, 1970) for which the contribute of a warm-dry climate was generally invoked (Suc and Bessais, 1990), even if some authors claim that tectonics is the only cause (Fauquette et al., 2006; Achalhi et al., 2016). In any case the shift to a wetter climate is supposed only at the end of this age (Griffin, 2002; Willett et al., 2006).

Below, the results related to the mineral phases, the major, and trace elements are discussed separately, in the light of their contribution in clarifying the relationships between weathering conditions and precursors in the formation of residual rocks as well as in palaeoclimatic and palaeogeographic reconstructions.

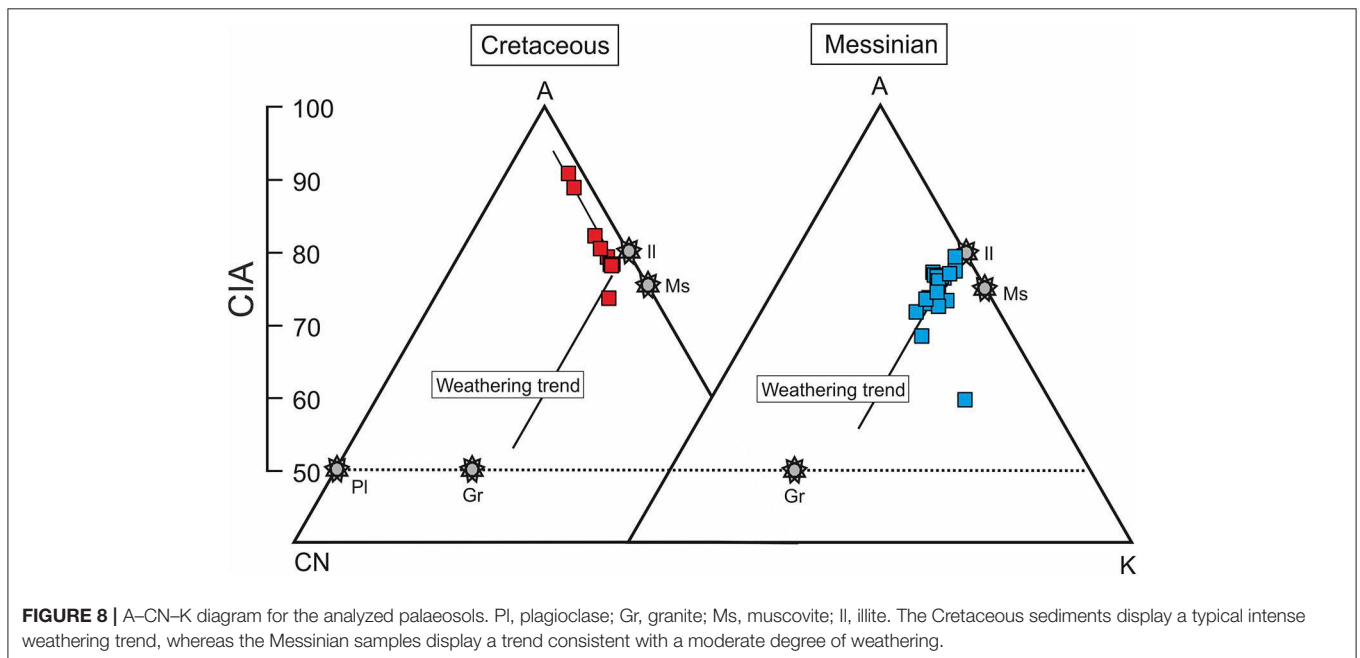
Major Elements Variations and Mineral Phases

We observed several differences between the compositions of the Cretaceous and Messinian palaeosols. The Cretaceous samples yield high TiO_2 , Al_2O_3 , Fe_2O_3 , MgO , and CaO concentrations, whereas the Messinian samples yield higher concentrations of SiO_2 , Na_2O , and K_2O (Figures 7A,B).

These differences, which include variations in both aluminum and alkaline element concentrations, affect the CIA values. The lower Messinian palaeosol yields a mean CIA of 72.51

(median = 73.18; $\sigma = 3.90$), the upper palaeosol yields a slightly higher CIA of 75.67 (median 75.80; $\sigma = 1.43$). These values indicate a moderate/intermediate degree of weathering, whereas Cretaceous samples yield a mean CIA of 82.26 (median = 79.99; $\sigma = 6.17$), suggesting more intense weathering, according to the different mineralogical evolution (illite rich vs. kaolinite dominated). These inferences are supported by the different proportions of kaolinite and illite in the two sets of samples. On an $\text{Al}_2\text{O}_3\text{-CaO} + \text{Na}_2\text{O-K}_2\text{O}$ diagram (Figure 8), the Cretaceous samples plot along a weathering trend parallel to the A-K join moving toward the A apex, indicating the leaching of alkaline elements from primary minerals coupled with the transformation of metamorphic phyllosilicates to kaolinite and in turn to Al-hydroxides (Yuste et al., 2015, 2017). In contrast, most of the Messinian palaeosols samples plot along a weathering trend that is sub-parallel to the A-CN join, indicating the leaching of Ca and Na from primary minerals coupled with the formation of 2:1 phyllosilicates. More in general, the A-CN-K diagram of both the CP and MP subsets clearly show weathering trends moving from a pristine “granitoid,” UCC-like, composition toward a clayey composition devoid of Ca and Na. Trends parallel to the A-K join moving toward the K apex are not observed, thus excluding any K-metasomatism modification. This is consistent with the SEM-EDS observations showing that illite and muscovite are always detrital.

The observed mineral distribution in the Messinian alterites is consistent with the hypothesis that weathering intensity was greatest at the surficial palaeosol and diminishes with depth. These characteristics, together with the occurrence of palygorskite, in the lower palaeosol indicate that the initial stage of weathering occurred under semi-arid conditions (Elidrissi et al., 2018 and references therein).



Noticeably the clay of the stratigraphically higher, Messinian, palaeosol contains non-negligible amount of gibbsite. This indicates a high water/rock ratio and the dissolution of kaolinite rather than the direct formation of gibbsite from feldspars, which requires a pH of ~ 3 (Gardner, 1970), which is inconsistent with the composition of meteoric water in NW Sardinia (present-day pH = 7–9; Mongelli et al., 2013) draining calcareous formations. Kaolinite, instead, could have formed by direct weathering of the basement rocks, before being transported, together other clay minerals, in the form of mud and debris flows, over the Mesozoic carbonate succession, where further weathering generated the kaolinite rich palaeosols. A high water content in the Messinian palaeosol is also testified by the occurrence of magnetite.

Trace Elements Variations

The Cretaceous palaeosols yield higher median concentrations of most trace elements (including Ga), transition metals (V, Cr, Ni, and Zn), high field-strength elements [HFSE; Y, Zr, Nb, total REE

(\sum REE), Hf, Th, and U] Cs, and Pb than the Messinian samples. In contrast, the Messinian palaeosols are decidedly enriched only in Ba and Cu relative to the Cretaceous samples (Table 1).

Both sets of samples yield median concentrations of HFSE, U, large ion lithophile elements (LILE; Rb, Cs, and Pb), and some transition metals (Sc, V, Cr, and, in the Cretaceous set only, Ni) that are higher than those of average global subducting sediment (GLOSS; Plank and Langmuir, 1998). In contrast, Cu, Sr, and Ba are depleted relative to GLOSS (Figures 9A,B).

Chondrite-normalized patterns for both sets of samples (Figures 10A,B) display a similar degree of LREE–HREE (Light Rare Earth Elements and Heavy Rare Earth Elements respectively) fractionation, as indicated by their median (La/Yb)_N values (CP = 11.38; MP = 10.64), which are higher than that of GLOSS (7.05). In contrast, the degrees of LREE–MREE (Medium Rare Earth Elements) [median (La/Sm)_N: CP = 3.29; MP = 4.04] and MREE–HREE fractionation [median (Gd/Yb)_N: CP = 1.75; MP = 1.69] are similar to those of GLOSS (3.14 and 1.67, respectively). The palaeosols yield median Eu

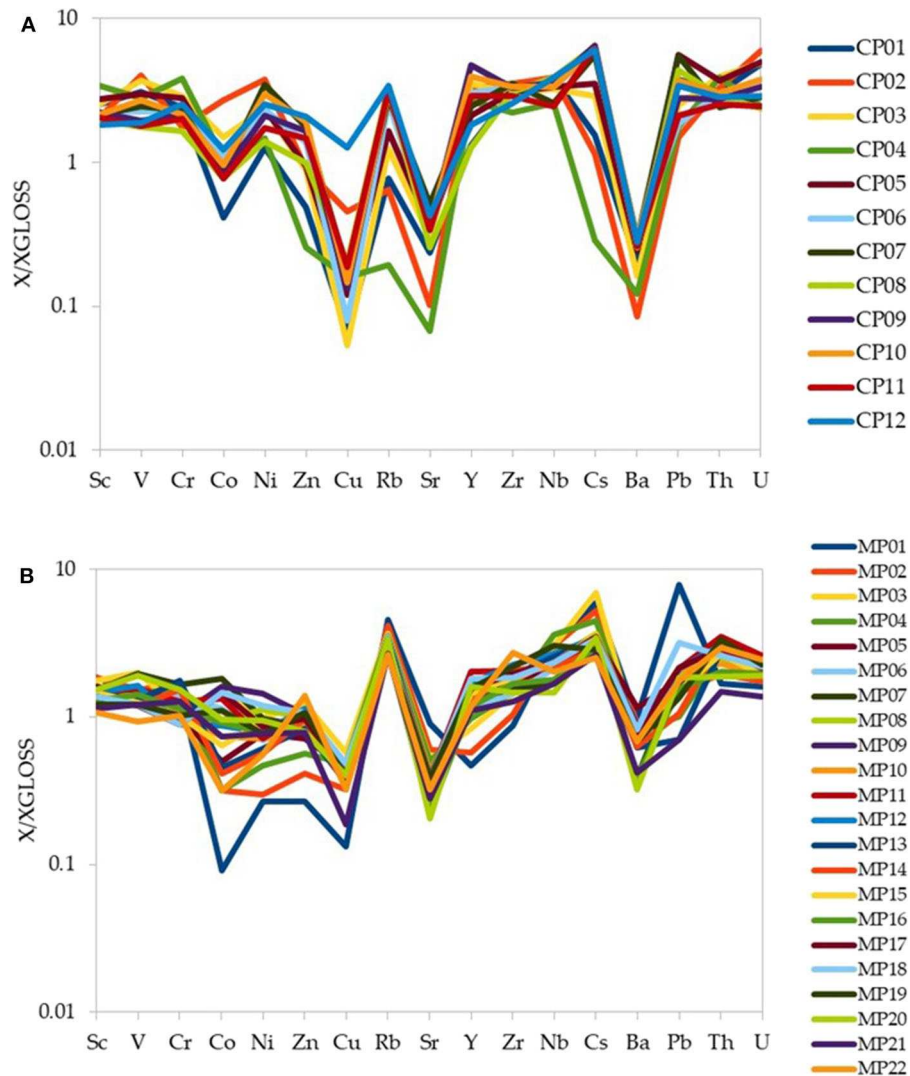


FIGURE 9 | GLOSS-normalized trace elements patterns of (A) the CP palaeosols, and (B) the MP palaeosols. GLOSS values are from Plank and Langmuir (1998).

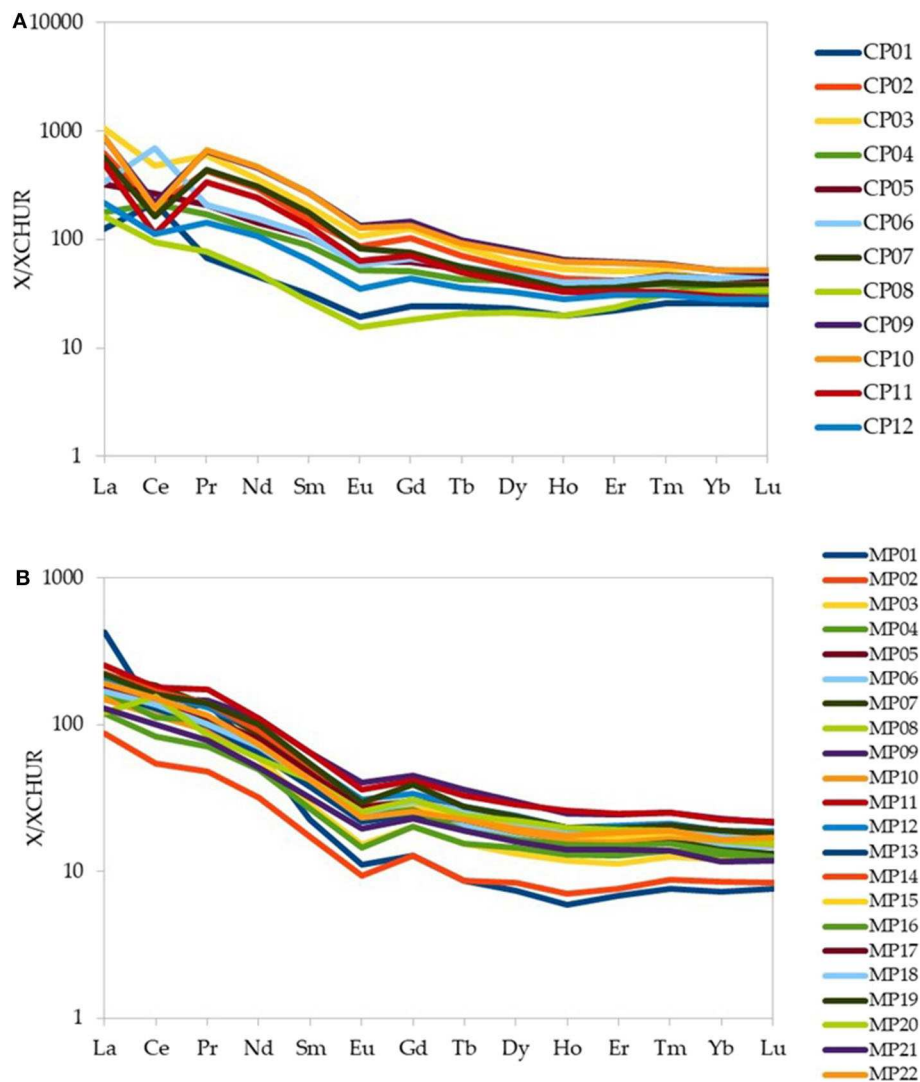


FIGURE 10 | Chondrite-normalized REE patterns of (A) the CP palaeosols, and (B) the MP palaeosols. Chondrite values are from Taylor and McLennan (1981).

anomalies (Eu/Eu^*) (CP = 0.69; MP = 0.71) that are similar to that of GLOSS (0.72). These fractionation indexes, that are those more affected by provenance, especially the Eu/Eu^* ratio, suggest both the sets share a provenance characterized by a composition close to the average Upper Continental Crust one.

Finally, the Messinian palaeosols yield Ce anomalies that are generally close to 1 with the exception of sample MP1 ($\text{Ce}/\text{Ce}^* = 0.48$). The Cretaceous samples instead exhibit a wide range of Ce/Ce^* values (0.26–2.63) likely as a consequence of local Eh fluctuations, as supported by the presence of remnants of a reducing humic horizon and oxyhydroxide phases (Braun et al., 1990; Mongelli, 1997; Mameli et al., 2008).

Inter-elemental Relationships

The relationships between major and trace elements in the Cretaceous and Messinian samples were evaluated using varimax

rotation and subsequent R-mode factor analysis. We selected different variables for the two datasets due to their differences in chemical evolution and mineralogy. Most Cretaceous samples display Ce anomalies; consequently, La (representing the LREE) and Yb (representing the HREE) were included in addition to Ce to enable the evaluation of inter-elemental fractionation among the REE. In contrast, as the Messinian samples lack significant Ce anomalies, we included only total REE. We also included a LILE (Rb) because the most abundant mineral in these samples is a 2:1 phyllosilicate.

Cretaceous Palaeosols

Three factors can explain 77.8% of the total variance in the composition of the Cretaceous palaeosols (Table 2).

The first factor (F1) explains 38.1% of the total variance and has positive weightings for Al_2O_3 , Sc, Cr, and Th. The Al_2O_3 concentrations are higher in soils with a higher weathering rate,

and are indicative of abundant clay minerals and Al-hydroxides. Al-bearing minerals in soils are also generally coupled with resistate phases that control the abundance of low-solubility elements, such as Sc and Th (e.g., Mongelli et al., 2014a,b), which are characterized by low water/rock partition coefficients (Rudnick and Gao, 2014). As for Cr, it occurs as insoluble Cr_2O_3 over much of its Eh–pH space, although in nature most Cr^{3+} is hosted in chromite, and Cr in soils occurs mainly in the residual immobile fraction (Kabata-Pendias, 2010). Thus, F1 likely represents the capacity of some elements to be concentrated during intense weathering.

The second factor (F2) explains 24.2% of the total variance and has significant positive weightings for Fe_2O_3 and V, and a negative weighting for Ga. The positive weightings for Fe_2O_3 and V suggest that V concentrations in the Cretaceous palaeosols were controlled mainly by Fe-oxyhydroxides. As for Ga, in the Ga–O–H system, it is hosted in söhngite $\text{Ga}(\text{OH})_3$, which is a common compound in soils (Kabata-Pendias, 2010), over much of its Eh–pH space ($\text{pH} = 6\text{--}10$ for $a_{\text{Ga}} = 10^{-8}$). Therefore, F2 might record competition between Fe- and Ga-hydroxides.

The third factor (F3) explains 15.1% of the total variance and has significant positive weightings for La and Yb. This factor accounts for the processes governing the distribution of REE in the Cretaceous palaeosols, with the notable exception of Ce. In general, soil solution composition is the main factor that drives the differential transport of REE during weathering and pedogenesis (Laveuf and Cornu, 2009 and references therein). HREE form complexes with organic matter (Henderson, 1984; Byrne and Li, 1995; Sonke, 2006) and bicarbonates (Cantrell and Byrne, 1987; Lee and Byrne, 1993; Pourret et al., 2007) that are more stable than those involving LREE, resulting in the preferential leaching of HREE during weathering. The soil solution composition also affects adsorption onto secondary minerals, and the balance of REE concentrations between soil solutions and mineral surfaces is controlled mainly by pH-driven adsorption/desorption reactions. REE are adsorbed onto mineral surfaces as pH increases and LREE are more efficiently adsorbed than HREE (Pourret et al., 2010 and references therein). The order in which REE are adsorbed onto mineral surfaces with

increasing pH is as follows: LREE > MREE > HREE, and REE are released from mineral surfaces in the same order during a period of decreasing pH (Gammons et al., 2005; Johannesson et al., 2006; Leybourne and Johannesson, 2008; Welch et al., 2009). Therefore, circa-neutral to acidic soil solutions enhance REE fractionation in soils, explaining the higher $(\text{La}/\text{Yb})_{\text{N}}$ and $(\text{La}/\text{Sm})_{\text{N}}$ values in the palaeosols relative to GLOSS.

Due to its redox chemistry, Ce is commonly fractionated relative to the other REE in soils and sediments that form through intense weathering (e.g., Mongelli et al., 2014a and references therein). Samples that yield negative Ce anomalies typically acquire their REE budget through the scavenging of Ce-depleted solutions resulting from Ce oxidation and precipitation in an environment devoid of organic matter (Usman, 2008; Mongelli et al., 2015).

Messinian Palaeosols

Four factors can explain 85.0% of the total variance in the composition of Messinian palaeosols (Table 3).

The first factor (F1) explains 39.4% of the total variance and has strong positive weightings for Al_2O_3 , Sc, and Rb. The 2:1 phyllosilicates formed during weathering; thus, Al_2O_3 concentrations are mainly related to the clay mineral content.

The second factor (F2) explains 22.8% of the total variance and has significant positive weightings for Fe_2O_3 , Co, Ni, and Cu. Most transition elements can be adsorbed onto Fe-oxyhydroxides (Usman, 2008); however, Acosta et al. (2011) showed that also Fe-rich mica-like clay minerals such as biotite, chamosite, and chlorite, which occur in the Variscan basement, may control the transition element budget in soils. Thus, F2 accounts for the capacity of Fe-oxyhydroxides and possibly 2:1 clay minerals to host heavy metals in soils.

The third factor (F3) explains 14.7% of the total variance and has significant positive weightings for TiO_2 and Th, and a negative weighting for Ga. In the Messinian palaeosols, the most abundant Ti-bearing mineral is detrital rutile. The relationship

TABLE 2 | R-mode factor analysis results for the CP composition.

	Factor 1	Factor 2	Factor 3
Al_2O_3	0.68		
$\text{Fe}_2\text{O}_3(\text{T})$		0.74	
Sc	0.93		
V		0.81	
Cr	0.82		
Ni			
Ga		−0.85	
La			0.80
Ce			
Yb			0.90
Th	0.82		

TABLE 3 | R-mode factor analysis results for the MP composition.

	Factor 1	Factor 2	Factor 3	Factor 4
Al_2O_3	0.91			
$\text{Fe}_2\text{O}_3(\text{T})$		0.83		
TiO_2			0.86	
Sc	0.97			
V				
Cr				
Co		0.79		
Ni		0.81		
Cu		0.85		
Ga			−0.81	
Rb	0.91			
Zr				
Th			0.78	
ΣREE				0.94

between Ti and Th may indicate that the abundance of rutile covaries with that of detrital Th-bearing minerals such as monazite. The negative weighting for Ga suggests that it is not present in high concentrations in accessory and detrital phases.

The fourth factor (F4) explains 8.1% of the total variance and has significant positive weightings for REE. Thus, F4 accounts for the processes controlling the REE distribution in the samples.

General Implications About the Palaeoclimate Conditions and Palaeogeography

In siliciclastic and residual sediments is generally retained (e.g., Perri and Ohta, 2014; Perri et al., 2015a,b) that relationships among Al_2O_3 , Sc, and Rb indicate that these elements covary with the abundance of clay minerals, with Rb likely occurring as interlayer cation. Thus, F1 in MP represents the resistance of some trace elements, including LILE such as Rb, to leaching during a weathering less intense with respect to CP. It is interesting to observe that the factors explaining the higher variance in both the sets (F1 Cretaceous vs. F1 Messinian) support what depicted by the mineralogical composition and the A-CN-K diagrams about the difference in palaeoclimate during the formation of the two different residual clay deposits.

The Eu anomaly, recording in sediments the proportion of exposed crust in the source area (e.g., McLennan et al., 1993; Cullers, 2000; Condie et al., 2001; Roser et al., 2002; Perri et al., 2013; Sinisi et al., 2014), has proved to be an

effective and reliable proxy of provenance also during intense weathering (e.g., Mongelli et al., 2014a, 2016). Similarly to the Eu/Eu^* proxy, the Sm/Nd ratio reflects chemical differentiation, since only minor fractionation of Sm and Nd occurs during intense tropical weathering (Viers and Wasserburg, 2004) and, as a consequence, the Sm/Nd ratio has been profitably used to determine parental affinities for evolved alterites (Mongelli et al., 2014a, 2016). In the Eu/Eu^* vs. Sm/Nd binary diagram (Figure 11), both the CP and MP subsets show large similarity and fall close to the average cratonic sediment suggesting that the Variscan basement, which patently was the source of the Messinian alluvial deposits, was also the source of the Cretaceous alterites.

Such evidence implies that during the Cenomanian-Turonian interval a wide region north of Sardinia (in pre-drift coordinates) underwent strong uplift and emergence that caused the erosive elision of the Mesozoic carbonatic shelf (at least, 1.5–2 km thick), leading to the exposition and dismantling of the underlying basement. In Nurra the pre-bauxite sequence was eroded down to the Oxfordian dolostone, causing the elision of about 400 m of carbonatic succession. The chrono-stratigraphic gap matches that of Nerthe in Provence (Masse and Philip, 1976), where more than 600 m of lower Cretaceous deposits were eroded (Guyonnet-Benaize et al., 2010). From this match can be argued that north Sardinia was located on the prolongation of the Durancian isthmus where Turonian bauxite is widespread (Laville, 1981; Combes, 1990). This well known structural high developed between the Vocontian basin to the north and

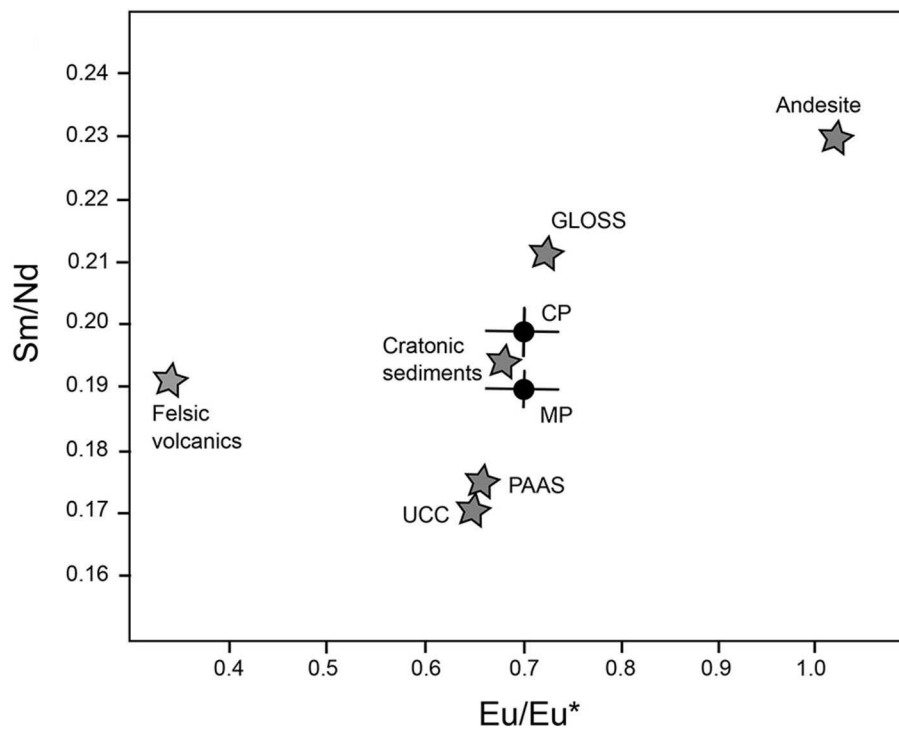


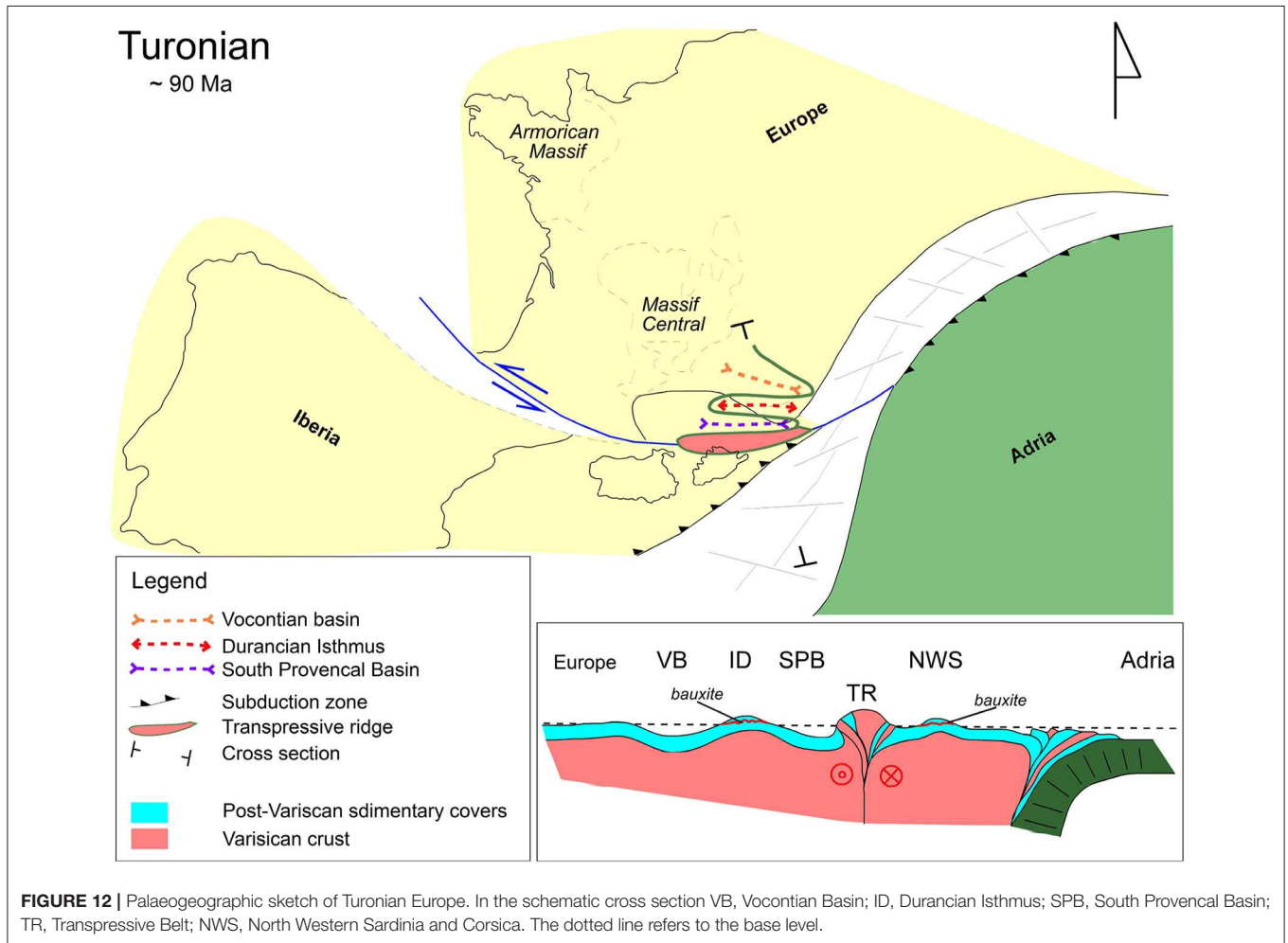
FIGURE 11 | Eu/Eu^* vs. Sm/Nd binary plot showing average standard lithologies. Circles are CP and MP palaeosols. Felsic volcanics and Cratonic sediments in Condie (1993); PAAS and Andesite in Taylor and McLennan (1985); GLOSS in Plank and Langmuir (1998).

1711 the South Provencal basin in the south (Masse and Fenerci-
 1712 Masse, 2013). The trigger for the uplift was recently referred
 1713 to a wide restraining zone, which caused N–S shortening and
 1714 long wave E–W crustal folding during the left lateral motion
 1715 between Iberia and Europe along a huge wrench fault (Schreiber
 1716 et al., 2011). Eastward, across the Ligure-Piemontese Ocean this
 1717 fault acted as a trench-trench transform fault separating two
 1718 oceanic domains with opposite subduction polarity (**Figure 12**;
 1719 Carmignani et al., 1995; Oggiano et al., 2009). In any case,
 1720 the uplift was limited to the carbonate cover, hence no bauxite
 1721 lying directly on basement rocks is known in the Durancian
 1722 isthmus. And yet, between Sardinia and Provence, there must
 1723 have been a ridge, with exposed basement, which fed with Al-
 1724 rich debris the continental plains and the bordering basins.
 1725 The relics of this structural relief (“Massif Meridional;” Hennuy,
 1726 2003) are to be found in the allochthonous slices of basement
 1727 in the Maure Massif and in basement outcrops buried in the
 1728 Ligure-Provencal basin (Fournier et al., 2016). Such transpressive
 1729 ridge faced the beginning subduction of the Ligure-Piemontese
 1730 Ocean beneath the European plate (Carmignani et al., 1995;
 1731 Argnani, 2009; Oggiano et al., 2009) and was more elevated—and
 1732 independent—compared to the Durancian rise (**Figure 12**). The
 1733

1768 occurrence of severely uplifted portions of the south European
 1769 margin are also in agreement with the south Variscan provenance
 1770 of detrital zircons in both proximal and distal Upper Cretaceous
 1771 flysches in “Alpine Corsica” (Lin et al., 2018) and in western
 1772 Liguria (Mueller et al., 2018), respectively. However, the latter
 1773 authors, disregarding the Cretaceous left lateral motion between
 1774 Iberia and Europe, refer the uplift, which should have caused the
 1775 complete elision of the thick Mesozoic cover—and this is not the
 1776 case—to a lithospheric bulge involving the Var-Maures Massif
 1777 (i.e., the Durancian isthmus). This bulge is thought caused by
 1778 the down-bending of the south European margin while it was
 1779 approaching the Alpin2e accretionary wedge.

1780 To define the degree of weathering of the basement exposed
 1781 over this structural high is not easy; in fact mantled alterites
 1782 on the Variscan basement have been hardly spared by the post-
 1783 Cretaceous erosion, where the protection due to transpressive
 1784 covers is lacking.

1785 Some residual alterites mantling the Variscan basement,
 1786 nevertheless, are preserved around the Massif Central; they
 1787 were classically assigned to the Tertiary (e.g., the siderolithic
 1788 formation), but palaeomagnetic and thermochronological dating
 1789 revealed a Cretaceous age (Thiry et al., 2006). Such alterites
 1790



1825 mostly consist of kaolinite and Fe oxy-hydroxides rich paleosols
1826 (Simon-Coinçon et al., 2000). A similar, already weathered,
1827 material partially contributed to feed the alluvial deposits that
1828 evolved into mature palaeosols and then into bauxite over the
1829 Upper Cretaceous calcareous palaeosurface (Combes et al., 1993)
1830 of Sardinia.

1831 Paleoclimate proxies, including the Ba/Sr ratio and the
1832 $\text{Na}_2\text{O}/\text{K}_2\text{O}$ molar ratio (Retallack, 1997; Retallack et al., 2001;
1833 Sheldon, 2006) support a not negligible water/rock ratio also
1834 during the formation of the Messinian deposits. The Ba/Sr
1835 ratio is largely used as leaching proxy during weathering since
1836 Sr is significantly more soluble than Ba, and Ba/Sr values >2
1837 indicate acidic/leached conditions under wet climate, and in
1838 the Messinian upper subset the average value of this proxy
1839 is 4.42. Similarly the $\text{Na}_2\text{O}/\text{K}_2\text{O}$ index, the proxy measuring
1840 the salinization of a palaeosol, is decidedly low (average value
1841 0.17) excluding a dry climate coupled with intense evapo-
1842 transpirative condition.

1843 These evidences do not meet the dry Messinian hypothesis
1844 (Suc and Bessais, 1990) and suggest a long interval of sub-
1845 tropical, wet climate during this age. How long this time span
1846 was, in the absence of a continental fossil record, is difficult to
1847 establish. In any case it is reasonable to argue that the lower
1848 alterite in the alluvial sequence matches one of the dry late
1849 Tortonian-early Messinian intervals predating the MSC in the
1850 Apennines (Bertini and Menichetti, 2015) and in Southern Spain
1851 (Casas-Gallego et al., 2015). The upper palaeosol, indicative of
1852 more wet subtropical climate reflects a time span wide enough to
1853 encompass the entire MSC, considering that so thick, kaolinite-
1854 enriched, paleosols take a few hundred thousand years to be
1855 formed (Thiry et al., 2006; Bronger, 2007).

1856 At the Messinian-Pliocene boundary a more seasonal,
1857 temperate climate with alternating dry and wet periods prevailed
1858 in western Mediterranean (Bertini and Martinetto, 2008). Soil
1859 formation was hindered by prevailing erosional dynamics during
1860 dry intervals (Günster and Skowronek, 2001). One of these, in
1861 north-western Sardinia, could be referred to the gravelly deposit
1862 that covers the Messinian alterite.

1864 CONCLUSIONS

1865
1866 Some interesting consequences rise from the comparison of
1867 the geochemical and mineralogical features of CP and MP
1868 weathering products:

- 1869 i. given that the two different alterites sheared a common source,
1870 any difference in chemical and mineralogical feature are to
1871

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1882 be referred to different weathering paths and, ultimately, to
1883 different climates;

- 1884 ii. during the Cenomanian and Turonian, the Variscan basement
1885 was exposed along a possible transpressive belt located
1886 between the South Provencal basin and Sardinia;
1887 iii. the dismantling of this ridge supplied kaolinite-rich material
1888 to the emerged Sardinian carbonate platform, where it was
1889 converted into a mature palaeosol and, after a long lasting
1890 period of alteration driven by monsoonal climate, into bauxite;
1891 iv. as for the Messinian alterites, the base of the alluvial
1892 sequence—possibly late Tortonian-early-Messinian—shows
1893 pedogenic clays dominated by illite, palygorskite, and
1894 oxidative conditions, which are indicative of dry warm
1895 climate, similar to that of actual savannah; conversely the
1896 Messinian upper palaeosol yields an unexpected weathering
1897 trend toward 2:1 clay minerals and, at less extent, gibbsite that
1898 are suggestive of an evolution toward more severe weathering
1899 in a wet climate;
1900 v. MP and CP palaeosols shared similar alteration in the
1901 first stages of the weathering path but MP, despite strong
1902 Al enrichment, never ended in lateritic soils, and bauxite.
1903 Though unexpectedly wet the climate was not warm enough
1904 to generate ferrallitic alteration.

1906 DATA AVAILABILITY STATEMENT

1907
1908 All datasets presented in this study are included in the
1909 article/**Supplementary Material**.

1911 AUTHOR CONTRIBUTIONS

1912
1913 All authors listed have made a substantial, direct and intellectual
1914 contribution to the work, and approved it for publication. All
1915 authors contributed ideas and assisted with writing and editing
1916 the manuscript.

1917 FUNDING

1918
1919 This work was funded by a research contract on Messinian
1920 clays awarded to the University of Sassari by the Municipality
1921 of Sassari.

1922 SUPPLEMENTARY MATERIAL

1923
1924 The Supplementary Material for this article can be found
1925 online at: [https://www.frontiersin.org/articles/10.3389/feart.
1926 2020.00290/full#supplementary-material](https://www.frontiersin.org/articles/10.3389/feart.2020.00290/full#supplementary-material)

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Conflict of Interest: The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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