

1 CELADONITE AND SMECTITE FORMATION IN THE 2 Úrkút Mn-CARBONATE ORE DEPOSIT (HUNGARY)

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13 ABSTRACT

14 Synsedimentary and early diagenetic oxygen levels are estimated by evaluating celadonite-
15 smectite formation in marine Jurassic black shale-hosted manganese-carbonates. Celadonite
16 formed under suboxic-dysaerobic conditions, Al-rich Fe-smectite formed at suboxic-anaerobic
17 conditions, and nontronite formed at anoxic-anaerobic conditions during sedimentary burial. A
18 genetic pathway by direct precipitation from solution is proposed for the enormous mass of
19 celadonite, based on mineral and textural evidence. Lamination of the manganese ore is
20 independent of clay-mineral composition and was given by a series of mineralized microbial Fe-
21 rich biomats.

22 **Key words:** Jurassic, Mn carbonate ore, celadonite, nontronite, microbial Fe oxidation

23 1. Introduction

24 Clay minerals are useful environmental indicators. For example, the occurrence of kaolinite
25 indicates intense acidic chemical leaching, whereas montmorillonite indicates a slightly alkaline

26 environment, reflecting the redox conditions at the modern ocean floor (Hein et al., 1979) and in
27 pore waters during alteration of volcanic rocks (Bustillo and Martínez-Frías, 2003). However, for
28 black shale-hosted Mn-carbonate ore assemblages, clay minerals have not been used even though
29 the laminated nature and color of the deposits (brown-green-grey) indicate a possible clay-
30 mineral utilization.

31 Investigations on the clay mineralogical composition of the Jurassic (Lias-Toarcian) Úrkút Mn
32 deposit started in the early 1980s (Fig. 1), when smectite and authigenic celadonite, as main
33 components of the fine grained rhodochrosite ore, were identified by XRD (Kaeding et al.,
34 1983), suggesting clay minerals precipitated from porewater and/or seawater influenced by
35 hydrothermal fluids. Varentsov et al. (1988) emphasized that these deposits resemble the
36 nontronite-celadonite metalliferous sediments of marine hydrothermal areas (i.e. the Galapagos
37 Rift Zone, southeast Pacific, Red Sea deeps), products of moderate to low-temperature
38 hydrothermal systems formed in oxygen-deficient environments.

39 Alteration of volcanic material did not result in clay mineral formation in the Úrkút deposit
40 (Polgári et al., 2012a) and Weiszbürg et al. (2004) suggested that celadonite formed by
41 authigenic precipitation from pore fluids. The minute size of the crystallites of the ore deposit
42 made characterization of the clay minerals difficult. For celadonite, the dominant crystallite size
43 is several tens of nm in thickness and some 100 to 1000 nm in length (Cora, 2008). The silicate
44 flakes are chemically heterogeneous and have various Fe-Al-Mg-Si ratios and K contents that are
45 typical of interlayer-deficient micas (0.7-0.8 atoms per formula unit-a.p.f.u.; Weiszbürg et al.,
46 2004; Cora, 2008).

47 Smectite crystallites are several 100 nm long and very thin, with the dominant cations being K-
48 Mg-Fe-Al. This silicate is also chemically heterogeneous and forms two subgroups, Al-rich Fe-

49 smectite and Al-poor Fe-smectite (Cora, 2008). The Mg and Fe contents are higher in the clay
50 minerals in laminae where Fe-rich minerals (pyrite, goethite) are abundant. The smectite is
51 mainly nontronite (Cora, 2008; Tóth et al., 2010).

52 The lamination of the ore is not mineralogically induced (Polgári et al., 2012b).

53 This paper determines the mineralogy, microstructure, and distribution of clay minerals
54 stratigraphically through the ore section; and the characterization of clay mineral genetic
55 pathways, which reflect fluctuations in redox conditions during ore formation.

56 **2. Geological setting and ore description**

57 The Jurassic (Lias, Toarcian) Úrkút Mn deposit is located in the Transdanubian Range (Fig. 1,
58 Fig. 2A). This black shale-hosted Mn-carbonate deposit is among the ten largest in the World. Its
59 reserves are 80 million tonnes of Mn-carbonate ore averaging 20 wt.% Mn and 10 wt.% Fe, with
60 an areal extent of tens of square kilometers. The ore deposit occurs within marine sedimentary
61 rocks composed mainly of bioclastic limestone, radiolarian clay marlstone, and dark-gray to
62 black shale. The Mn-carbonate ore beds conformably overlie middle Lias cherty limestone. The
63 rhodochrosite ore is composed of laminated, alternating gray, green, brown, and black sections
64 composed of mixtures of very fine-grained carbonate minerals and clay. Fine-grained (1–2 μm)
65 rhodochrosite rock lacks coarse and fine detrital clastics (Polgári et al., 2012a). Ore accumulation
66 took place in a structurally-controlled small marine basin in a low-energy, and low temperature
67 depositional environment. The deposit is unmetamorphosed, and was not effected by diagenetic
68 thermal overprint proved by stable O isotope data and interpretation (Polgári et al., 2012a). The
69 ore deposit consists of three ore beds (10-, 3-, and 1 m thick), separated by a 20- and 4 m-thick
70 black shale (Fig. 2B).

71 A genetic model shows, that two cycles of bacterial activity triggered ore formation (Polgári et
72 al., 2012a). Cycle 1 is a near-seabed aerobic chemolithoautotroph cycle which was essential in
73 sequestering metal ions (Mn^{2+} , Fe^{2+}) from solution via enzymatic Mn(II) oxidation. Mn-oxide
74 proto-ore was deposited in the sediment pile, serving as a paleoenvironmental indicator of oxic
75 conditions. Cycle 2 represents an anaerobic/suboxic heterotrophic bacterial cycle in the frame of
76 which early diagenetic bacterially mediated Mn(IV) and Mn(III) reduction processes took place
77 via organic matter oxidation and Mn-carbonate mineralization (Polgári et al., 1991; Polgári et al.,
78 2012a). The ore sequence is laminated in the millimeter scale (Fig. 2C, Fig. 3) reflecting a series
79 of Fe-rich biomats (Polgári et al., 2012b). On a meter scale, the deposit shows color variations,
80 the lower and upper part of the main ore bed is green and the middle part is brown. Towards the
81 middle black shale, the top of the main ore bed is grey.

82 *Fig. 1.*

83 *Fig. 2.*

84

85 **3. Samples and methods**

86 X-ray powder diffraction on 56 samples (252 subsamples) was performed using a Philips
87 diffractometer (PW 1710) with carbon monochromator and Cu $K\alpha$ radiation, accelerating voltage
88 - 45 kV and current - 35 mA Mineral composition was determined on randomly powdered
89 samples by semi-quantitative phase analysis according to the modified direct method
90 of Bárdossy et al. (1980), using previously defined intensity factors.

91 Oriented samples (112) were taken along a 917 cm complete section from the footwall to
92 hanging wall from five sections dissecting the main ore bed and one section dissecting the upper
93 ore beds (Fig. 2B). Bulk and separated lamina sub-samples were studied to determine

94 macroscopic features that cause color and grain-size variations. Ninety thin sections were studied
95 by microscopy.

96 **4. Results**

97 The Mn-carbonate ore beds are composed of Ca-rhodochrosite, celadonite, smectite, goethite,
98 and siderite. Moderate components are quartz, Mn-bearing calcite, apatite, pyrite, and barite; and
99 minor components are manganite, kutnohorite, gypsum, and feldspar (Fig. 4; Polgári et al.,
100 2012ab). Discrete nontronite layers of variable thicknesses occur in the ore deposit. Sub-lamina
101 XRD results show that celadonite and smectite occur separately in some laminae and in other
102 laminae both celadonite and nontronite occur, in variable amounts, and without recognizable
103 stratigraphic or spatial patterns.

104 Thin sections show that the entire ore bed is composed of a millimeter or thinner series of woven
105 structures, interpreted by Polgari et al. (2012b) as biomats (Fig. 3). The Fe-rich biomat structures
106 are composed of goethite. The matrix consists of Ca rhodochrosite, celadonite, and nontronite as
107 main components.

108 *Fig. 3.*

109 *Fig. 4.*

110 **5. Discussion**

111 **5.1. Authigenic mineral formation**

112 Recognition that Fe-rich biomats created the mm-scale lamination in the ore deposit makes it
113 possible to constrain the formation of authigenic clay minerals. The most probable microbial
114 Fe(II) oxidizing microbe was a *Gallionella*-like (*Mariprofundus ferrooxidans*) form living in a
115 neutrophylic, non-photosynthetic suboxic/dysaerobic environment (0.3 V Eh using recent
116 analogs; Konhauser, 1998; Hallbeck and Pedersen, 1990; Emerson et al., 2010). Most of the

117 modern seabed shows neutral pH conditions. Mineralogical and textural investigations show that
118 celadonite, goethite, and manganite formed at the sediment/water interface as synsedimentary
119 authigenic minerals.

120 **5.2. Celadonite formation**

121 The formation of celadonite has particular importance for understanding the genesis of the Úrkút
122 manganese ores. It is generally accepted that Fe-micas form preferentially (in limited quantities)
123 in submarine hydrothermal environments such as amygaloidal infilling and veinlets in basalt
124 (type 1, Pichler et al., 1991, Bustillo and Martínez-Frías, 2003), or during oxidic submarine
125 alteration of volcanic rocks (type 2, basaltic rocks, alteration rinds, often replaced or overgrown
126 by saponite; Butuzova et al., 1979, 1983; Varentsov et al., 1983). In addition, large
127 accumulations of hydrothermal green clays occur in mounds of the Galapagos Rift Zone, in
128 sediments of the East Pacific Rise and Bauer depression, in the spreading zones of the Gulf of
129 California, in the TAG and the Famous areas along the Mid-Atlantic Ridge, in Red Sea deeps,
130 and in the Gulf of Aden. Varentsov et al. (1988) interpreted the celadonitic phases in these green
131 clays as products of post-sedimentary (mainly diagenetic) transformation of Fe-smectite, which
132 in turn were formed in the initial crystallization stages of a siliceous-ferruginous gel under weak
133 reducing conditions (type 3; Butuzova et al., 1979, 1983; Varentsov et al., 1983).

134 These processes are highly unlikely at Úrkút because there are no volcanic rocks or substantial
135 amounts of volcanic debris. Nevertheless, there is an enormous amount of very fine-grained
136 celadonite, with an estimated mass of some tens to a hundred million tonnes based on ore reserve
137 and composition calculations. Detailed mineralogical observations (Weiszburg et al., 2004; Cora,
138 2008) indicate that celadonite directly precipitated from solution. Hence, celadonite is not a
139 diagenetic product from a precursor mineral phase (type 4, Úrkút; marine basin with distal

140 hydrothermal discharge and primary celadonite formation). Regardless of the differences in
141 precursor phases (rock, mineral, and solution), the formation of celadonite is similar. The
142 composition of celadonites from different formation types are similar, but textures (vein fillings,
143 alteration rinds), accompanying mineralogy (volcanic rock components), and chemical
144 composition distinguishes the four types of celadonite.

145 The Úrkút authigenic celadonite formed at low temperatures (17-23° C; Polgári et al., 2012a).
146 Celadonite at Úrkút shows evidence for a slight variation in redox conditions (Fig. 5). It starts
147 from an oxidizing water-dominated environment (microbial Mn(II) enzymatic oxidation, aerobic
148 system with dissolved oxygen (DO) of more than 2 ml/l. That is followed by dysoxic conditions
149 (DO: 0.2-2.0 ml/l of H₂O) where microbial Fe(II) oxidation occurs (Fe-rich biomats, DO: 0.3
150 ml/l of H₂O), and then suboxic conditions (DO: 0-0.2 ml/l of H₂O), where primary celadonite
151 formation (DO: 0.1-0.2) occurs in a neutrophylic environment. DO values are based on Hallbeck
152 and Pedersen (1990), Wignall (1994), Konhauser (1998), Emerson et al. (2010), and the results
153 of nontronite synthesis laboratory experiments (Harder, 1976).

154 *Fig. 5.*

155 Conditions changed to a more reducing, sediment-dominated environment during burial, where
156 first Al-rich Fe-smectite and later Al-poor Fe-smectite (nontronite), formed. During this latter
157 stage, pyritization of goethite took place (Polgári et al., 2012b). Pyrite is always accompanied by
158 smectite, indicating reducing, anoxic conditions. Celadonite did not form from Fe-smectite
159 because the diagenetic process trends to oxygen depletion during increasing burial.

160 **5.3. Smectite formation**

161 The Úrkút smectites could have formed from (a) direct precipitation from hydrothermal fluids, as
162 like as nontronites in the modern ocean basins, characterized by Fe-rich compositions, low Al

163 and Mg, the presence of Ba, and relatively warm temperatures (T) (Bischoff, 1972; Cole and
164 Shaw, 1983; Cole, 1983; Iizasa et al., 1998), or Red Sea brine smectites (Butuzova et al., 1979,
165 1983). However, relatively high temperatures (T) are not supported by O isotope results. A more
166 probable mechanism is (b), low T reaction of Fe oxyhydroxide and silica, which is characterized
167 by high Fe, low Al, and probably low Mg contents if formation took place with FeOOH
168 formation; and moderate to high Fe, moderate Al, and low- to moderate Mg contents if the
169 smectite formed at great distances from where FeOOH formed (Heath and Dymond, 1977;
170 Polgári et al., 2012a).

171 According to Hein et al. (1979), two more general mechanisms can be presumed for the
172 formation of authigenic smectite in the deep sea: (c) precipitation from solutions at low
173 temperature into vesicles and fractures of basalts (seawater or leached from basalt by seawater),
174 which are common at oceanic spreading centres (Seyfried et al., 1978); and (d) alteration of
175 volcanic rock fragments and glass in the marine environment, which is the most commonly
176 reported mechanism (Hein and Scholl, 1978; Bustillo and Martínez-Frías, 2003). Such smectites
177 are characterized by a range of compositions, from Fe- and Mg-rich and moderate Al contents, to
178 moderate to high Fe and Mg and higher Al contents than in types (a) and (b).

179 The composition of Úrkút smectites are Fe-Mg-rich with low Al and occasionally contain K
180 (Polgári et al., 2012a), support formation by mechanisms (a) or (b). The (b) type smectite forms
181 near the sediment-water interface where Mg is abundant and this may be the case for Úrkút,
182 where the bottom waters were oxygenated (Polgári et al. 2012a), and consequently, smectite may
183 not have formed until after some burial, perhaps 10-20 cm where conditions were more reducing
184 (Hein et al., 1979). In this regard, smectite formed (Al-rich Fe-smectite) under more reduced

185 conditions than those under which celadonite formed, and nontronite formed in a fully reducing
186 environment, probably from celadonite or Al-rich Fe-smectite.

187 Another mechanism for nontronite and celadonite formation is through microbially mediated
188 processes (Köhler et al., 1994; Iizasa et al., 1998; Tazaki, 1997; Bustillo and Martínez-Frías,
189 2003). This mechanism cannot be excluded for the Úrkút, where a series of Fe-rich biomats
190 occur; however, microbially mediated diagenetic overprints probably eliminated primary textural
191 evidence for that process.

192 **5.4. Transformations of clay minerals and goethite**

193 On the basis of microscopy, the Fe-biomaat texture is well preserved in most of the ore sections.
194 Only at the top part of the main and second goethite was transformed to pyrite. The textural
195 connection between well-preserved Fe-biomaat goethite and the clay mineral matrix shows an
196 oxygen decrease from suboxic conditions, which probably occurred during early diagenesis after
197 some burial. Clay mineral-rich sediment accumulated prior to or at the same time as Fe-biomaat
198 growth. The source of Fe(II) was most probably low T geothermal fluids ascending and mixing
199 with seawater (Polgári et al., 2012a). This is an aspect that needed clarification because the
200 microbially mediated goethite (originally ferrihydrite) reacting with silica can form clay minerals
201 as well, but textural observations do not support that sequence of events. Nontronite → goethite
202 and celadonite → goethite transformations are not considered likely because of Eh constraints;
203 the sediment pile did not become more oxic during diagenesis.

204 **6. Conclusion**

205 Clay minerals constrain the paleoenvironmental conditions during formation of the Úrkút Mn ore
206 deposit. The clay minerals formed synchronously in different parts of the sediment column under
207 different redox conditions, celadonite at the sediment/water interface or very close to it, and

208 smectite in the deeper, more reducing parts. A pathway of celadonite formation is proposed by
209 direct precipitation from solution en masse. Nontronite formed in the reducing part of the
210 sediment column. These results show that celadonite and nontronite indicate palaeo-oxygen level
211 fluctuations in the environment. Biomats reflect suboxic conditions, which changed to anoxic
212 with increasing burial. Color in the Úrkút deposit does not reflect oxygen conditions as
213 commonly thought; for example, brown can be anoxic (nontronite) and green suboxic
214 (celadonite), so color as a paleoproxy must be used with caution.

215 **Acknowledgments**

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217

218 **References**

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293

294 **Figure captions**

295 Fig. 1. Geological sketch map of the Úrkút manganese deposit (after Szabó and Grasselly, 1980).

296 The locality GPS data: 47°04'55"N; 17°38'40"E

297 Fig. 2. Locality map (A), Geological profile with sample locations of the Úrkút Mn-carbonate
298 deposit (B). (Sampling 2009, Úrkút Mine, Shaft No. III, deep level, +180 m; total No. of
299 samples 112 spanning 917 cm from the base to the top of the three ore layers and the
300 intervening black shale). Key: fragm-fragmented sample; cont.-continuous sampling; not
301 cont.-not continuous sampling; numbers on the stratigraphic columns are sample
302 numbers; * indicates samples for XRD; 0 indicates samples for thin sections (in brackets
303 the number of thin sections, total 90); Profiles 1, 3, 4, 5 are from main ore bed, Profile Za
304 and Zb are from second ore bed. Patterns show only color varieties and not sedimentary
305 structures; (C1) green (C2) brown Mn-carbonate ore.

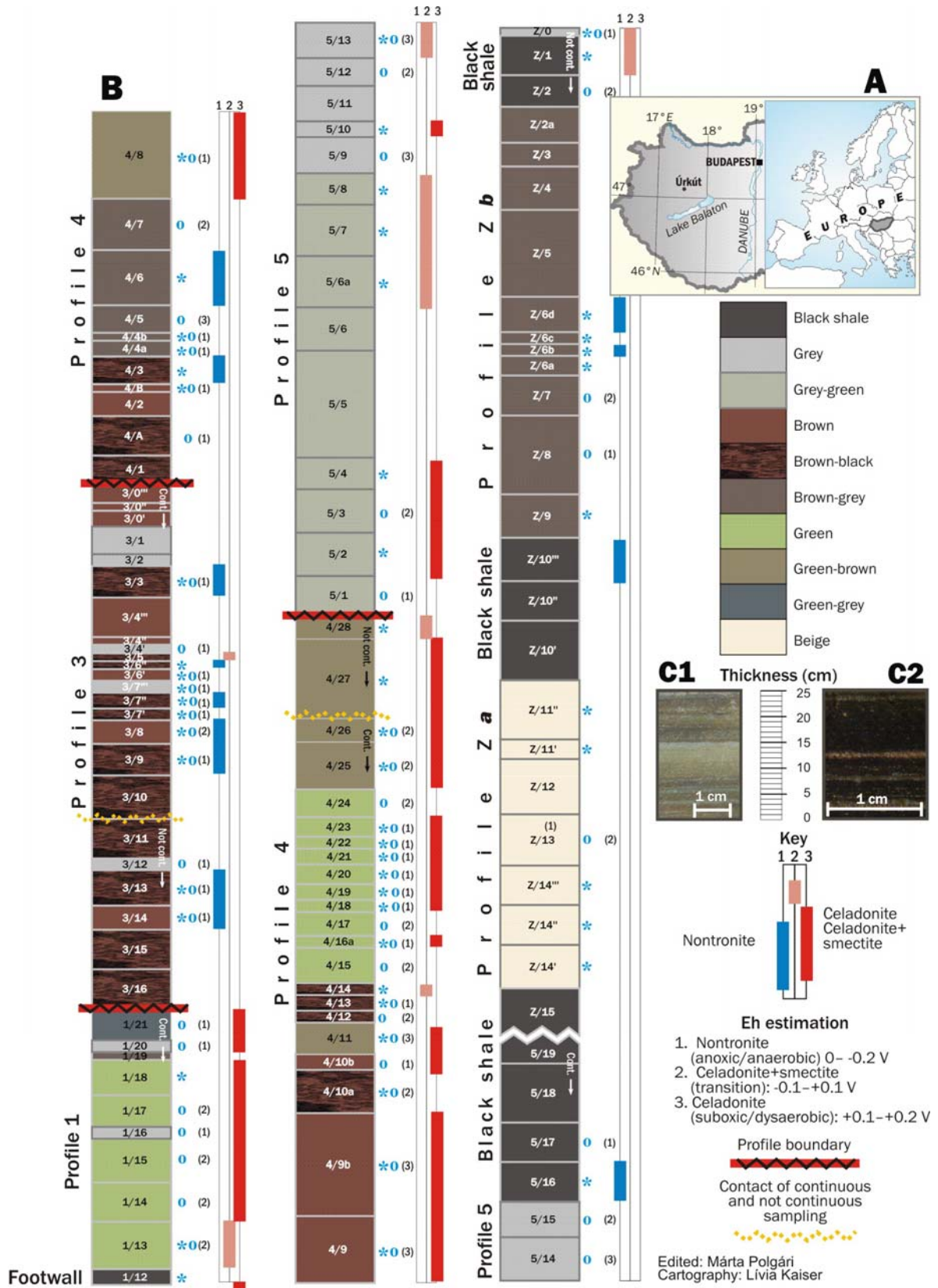
306 Fig. 3. Thin section photos showing Fe-rich biomat structures for different representative
307 samples and magnifications. The thickness of the thin sections and the density of biomats
308 are variable. Arrows show representative parts of mineralized filamentous structures. For
309 sample locations see Fig. 1.

310 Fig. 4. Mineral composition (XRD) of Mn-carbonate ore samples and macroscopically separated
311 subsamples normalized to 100% (for sample locations see Fig. 1). The graph was made
312 by coding the estimated quantity of selected minerals (graphics by Gergely Rózsás,
313 Pázmány Péter University). Key: gry-grey; grn-green; brn-brown; blk-black; avg-average.

314 Fig. 5. Estimated formation conditions of the clay minerals in the black shale-hosted Mn
315 carbonate ore deposit (Úrkút), zone of microbial Fe(II) oxidation (+0.3 V Eh using recent
316 analogs; Konhauser, 1998; Hallbeck and Pedersen, 1990; Emerson et al., 2010), and the
317 results of nontronite synthesis laboratory experiments (by dashed lines, Harder, 1976,
318 1978). Note: in general oxidizing conditions mean high oxygen concentration (high Eh

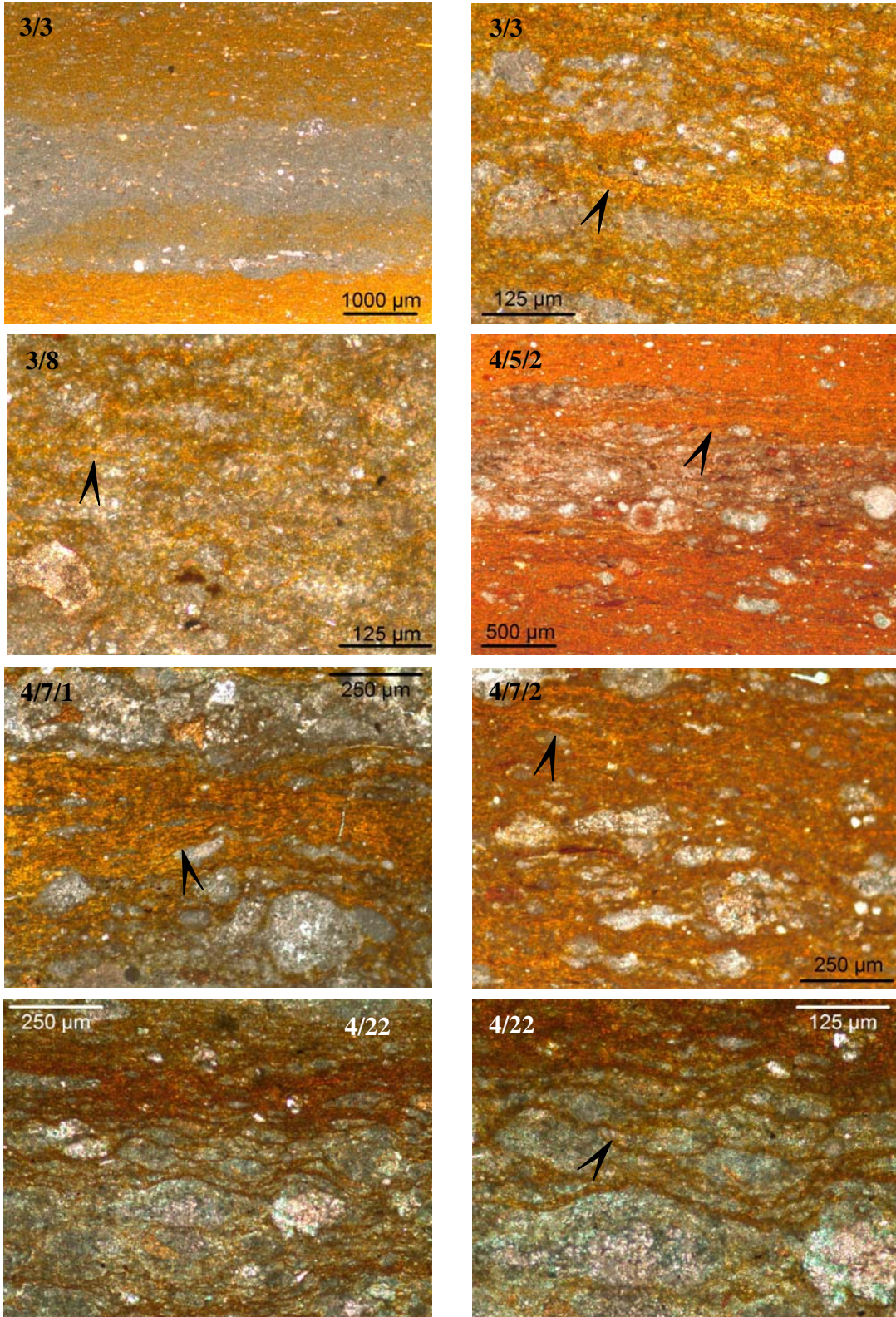
319 values up to +0.4V), while reduced conditions reveal a lack of oxygen (low Eh down to -
320 0.2V. Eh estimation is based on Wignall, 1994).

321 Fig. 1.



322 Fig. 2.

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368 Fig. 3.

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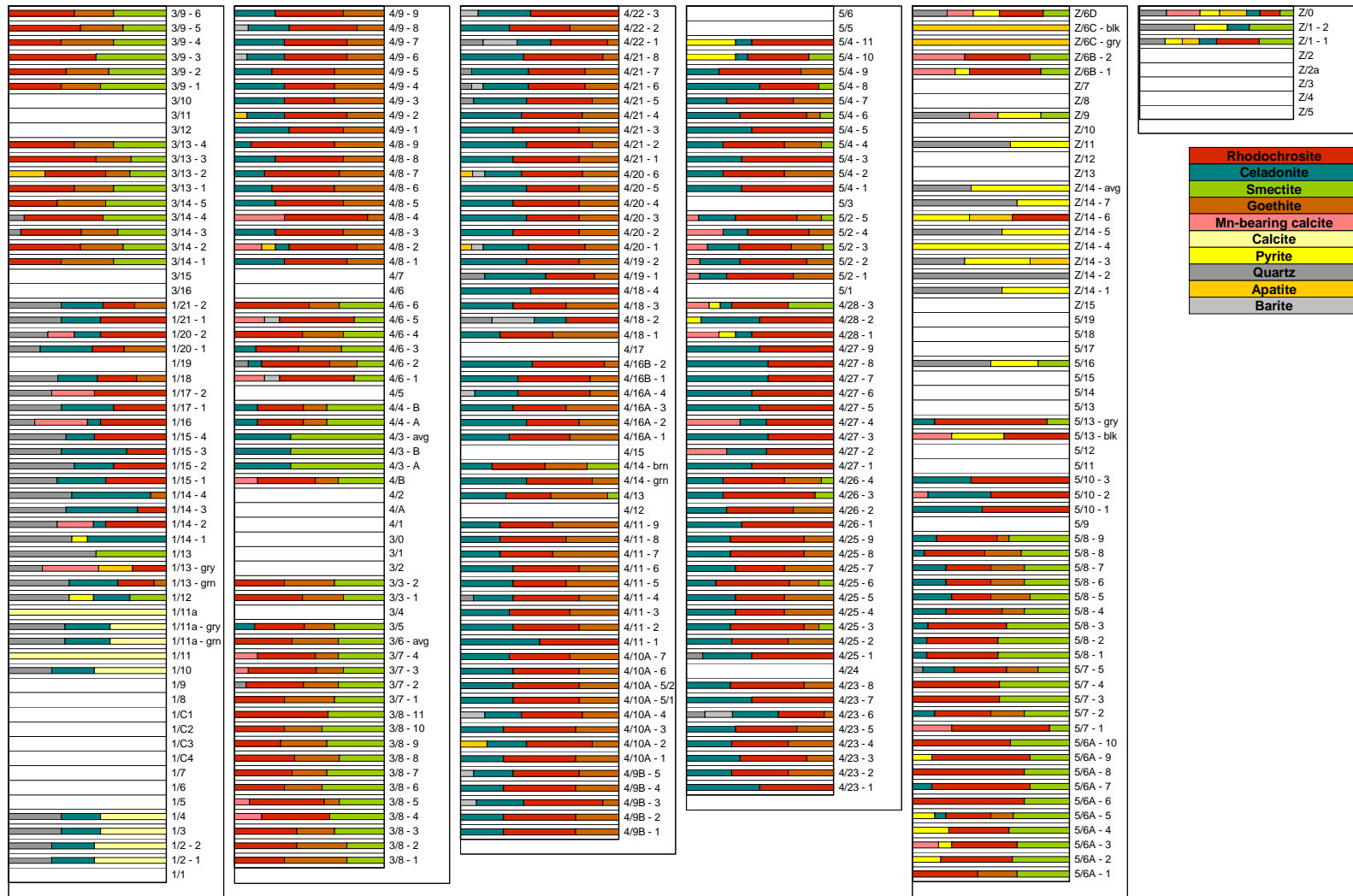
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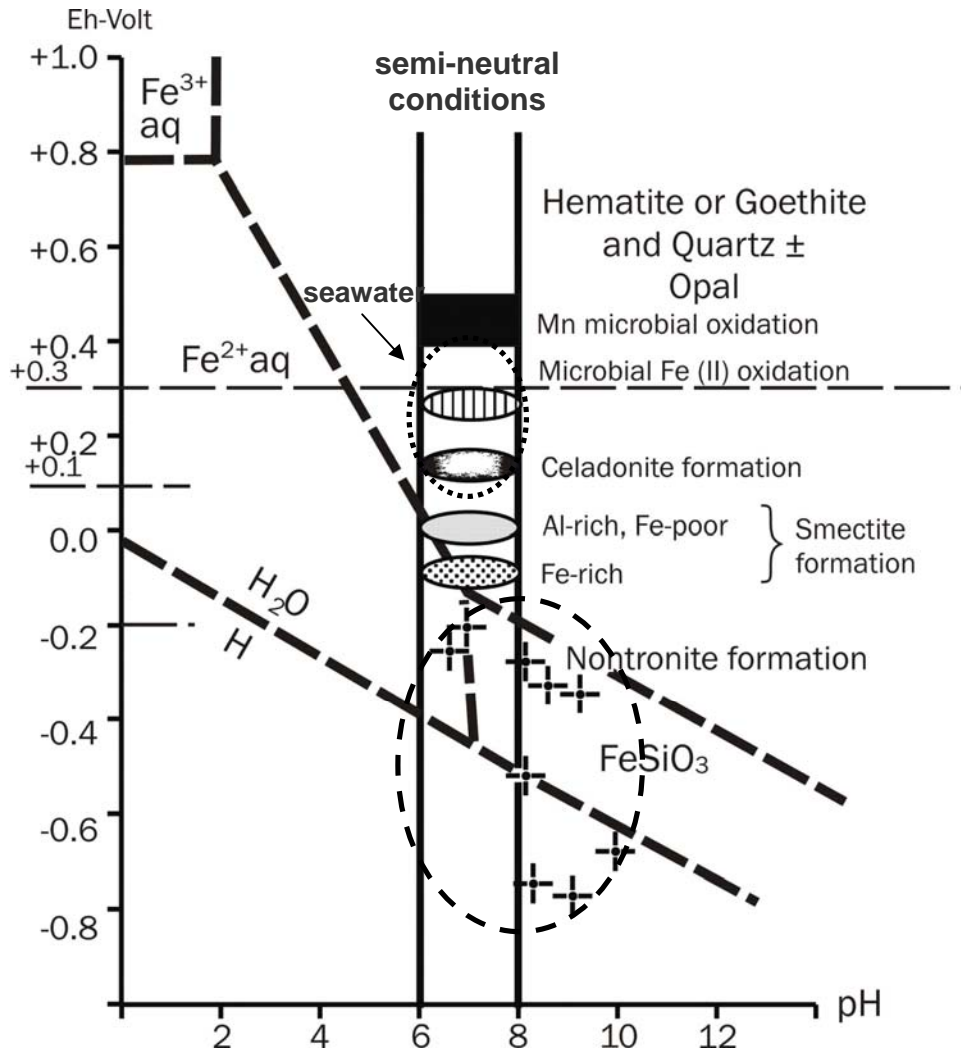
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400 Fig. 4.



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