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**Weathering and mineralogical evolution in a high Alpine soil chronosequence: A combined approach using SEM–EDX, cathodoluminescence and Nomarski DIC microscopy**

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1 **Weathering and mineralogical evolution in a high Alpine soil**  
2 **chronosequence: a combined approach using SEM-EDX,**  
3 **cathodoluminescence and Nomarski DIC microscopy**

4

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17

18 **Abstract**

19 Physical and chemical weathering of primary minerals of granitic till in the proglacial area of  
20 Morteratsch (Switzerland) was investigated using cathodoluminescence (CL), Nomarski  
21 differential interference contrast (DIC) microscopy and scanning electron microscope (SEM-  
22 EDX). The investigated time-span ranges from 0 to 140 yr of sediment exposure. For the  
23 very early stage of weathering or soil formation only little information is available. The main  
24 aim of our investigation was consequently to see whether weathering of primary minerals can  
25 be detected in such a short time-span using for the first time for soils well-established  
26 methods as CL and Nomarski DIC microscopy in geo- and material science such. For that  
27 purpose, the fine earth fraction (< 2 mm) of topsoil samples was investigated. Some physical

28 weathering had taken place within 140 years. The delamination of biotite seems to increase  
29 with time. SEM and CL analyses also demonstrate early weathering of quartz by evidencing  
30 edge pits and structural bonds – such as Si-O-Si in quartz – that start to break and to  
31 transform into free radicals. K-feldspar and plagioclase contain Fe. When using Fe<sup>3+</sup> as  
32 reference point (680-700 nm) to standardise the CL spectra, the Al-O-Al defects of K-  
33 feldspar exhibit a relative decrease with time; this was not the case for plagioclase. The CL  
34 measurements showed that the investigated apatite contained REE (rare earth elements) in  
35 the crystal structure. However, none of the other techniques (DIC, SEM-EDX) were helpful in  
36 detecting any specific weathering features for apatite. In the time span of 140 years, epidote  
37 weathering was evidenced using XRD in a previous investigation and here using DIC  
38 microscopy (morphologic changes). Several mineral changes could be traced within a very  
39 short weathering sequence using the applied techniques. These changes include physical  
40 (e.g. biotite), chemical or crystal structure (K-feldspar, biotite) features. Such an analytical  
41 combination is promising, therefore, for the detection of chemical, physical and mineralogical  
42 characteristics and changes in very young glacial sediments.

43

44 **Keywords:** cathodoluminescence; Nomarski DIC microscopy; SEM-EDX; weathering; soil  
45 formation; proglacial area; feldspar; plagioclase; quartz; apatite

46

47

## 48 **Introduction**

49 Due to global warming, glaciers retreat and new areas are exposed to weathering. Glaciers  
50 and discontinuous permafrost in these ecosystems react sensitively to atmospheric warming  
51 because the year-round temperature of their surrounding is not greatly below their melting  
52 point (Haeberli and Beniston, 1998). The abrasive action of active ice masses produces  
53 sediments (i.e. moraines), which tend to weather as a function of their exposure time. In the  
54 chemical sense, soil development is roughly synonymous with weathering. Elemental and  
55 mineralogical compositions of soils evolve due to complex feedbacks among geochemical

56 and geomorphic processes within and at the boundaries of the soil layer. Geochemical  
57 processes involve dissolution, leaching, precipitation and colloidal translocation (Yoo and  
58 Mudd, 2008), whereas geomorphic processes include the conversion of parent material to  
59 soil materials and the colluvial transport of the soil materials.

60 It is often assumed that weathering mechanisms in cold regions are slow due to the low  
61 temperatures. The proglacial area is, however, a potential zone of a) high geochemical  
62 reactivity due to the availability of freshly-ground reactive material (subglacially derived), b)  
63 high water-to-rock ratios and contact times, c) high permeability, and d) a constant supply of  
64 dilute waters (meltwaters and rain/snowmelt) percolating through the deposits. All these  
65 factors should favour chemical weathering. There is no agreement about the velocity of  
66 reaction in proglacial areas. Anderson et al. (1997, 2000) concluded from their  
67 measurements that silicate weathering reactions in proglacial areas may be important only  
68 after a vegetation cover is established. In contrast, Wadham et al. (2001), Egli et al. (2003)  
69 and Hosein et al. (2004) suggest that glacially derived material is subjected to intense  
70 chemical weathering, starting immediately after deposition in the proglacial zone and  
71 subsequently continuing for thousands of years after glacier retreat.

72 Hosein et al. (2004) and Föllmi et al. (2009a,b) measured high weathering rates in proglacial  
73 areas in the Alps ( $10^{-15}$  to  $10^{-13}$  mol biotite  $m^{-2} s^{-1}$  for a 140-270 yr-old exposed proglacial  
74 area). Biotite weathering in young soils was generally much higher than in old soils. The  
75 calculated weathering rates of biotite were several orders of magnitude higher than known  
76 field weathering rates (e.g., Swoboda-Colberg and Drever, 1993; Murphy et al., 1998). This  
77 seems to be related to the predominance of fine-grained particles ( $< 63 \mu m$ ) in glacial  
78 sediment that are mechanically disaggregated and preferentially leached.

79 In general, the availability of data about weathering rates and alteration of primary rock-  
80 forming minerals in young and cold areas is scarce. The detection of changes is often limited  
81 by the choice of techniques. The combination of transmitted-light cathodoluminescence (CL),  
82 Nomarski DIC (Differential Interference Contrast) microscopy and scanning electron  
83 microscopy (SEM-EDX) can give important insights into the chemical composition and crystal

84 arrangement of minerals (i.e. point defects, cationic changes, etc.). Most materials have  
85 distinct luminescence properties that allow a rapid identification of phase distribution and  
86 transformation. Nomarski DIC microscopy is an optically-based technique first documented  
87 by Nomarski and Weill (1951) and Nomarski (1955). It allows the observation of  
88 micromorphological features in natural materials and/or metal alloys, and has been applied in  
89 reflected light microscopy in metallurgy and petrographic research (e.g. Pearce et al., 1987;  
90 Keevil and Walker, 1992). Despite a relatively straightforward preparation of the samples and  
91 the high amount of information that can be retrieved by the combination of CL and Nomarski  
92 DIC, their coupled application is rare (Götze and Siedel, 2004, 2007). Transmitted-light CL  
93 microscopy is a technique applied to building materials (Michalski et al., 2002; Götze and  
94 Siedel, 2007; Götze, 2009), archaeology (Lapuente et al., 2000; Götze and Siedel, 2004) and  
95 geosciences (e.g. Götze et al., 2000; 2004; Götze and Zimmerle, 2000; Richter et al., 2003).  
96 The aim of this study is to trace mineralogical and crystallographic changes in soils that have  
97 developed across a recently exposed granitic till. For that purpose, the proglacial area of the  
98 Morteratsch glacier was selected. The combination of preferential weathering patterns and  
99 elemental depletion (for the main structure modifier elements) for primary mineral phases  
100 were taken into account.

101 For this purpose, CL and Nomarski DIC microscopy – well-known analytical techniques in  
102 material science but not applied so far for soil studies – analyses were performed in  
103 combination with SEM-EDX. An additional aim was consequently to test the power and  
104 suitability of these techniques for studying short-term and initial weathering processes in  
105 young soils.

106

## 107 **Study area**

108 The proglacial area of Morteratsch is located in Upper Engadine, SE Switzerland (Fig. 1).  
109 The valley runs in N-S direction and the current length of the exposed area is approx. 3 km.  
110 The glacier has been continuously retreating without re-advancements since the 1850s (end  
111 of the Little Ice Age; Burga, 1999). The altitude of the investigated proglacial valley ranges

112 from 1900 m asl to approx. 2150 m asl, with no abrupt stacks and slopes along its main axis.  
113 Geologically, the glacial till consists of granitic material. The morainic material was produced  
114 through glacial transport within a small area of relatively homogeneous parent material  
115 (Mavris et al. 2010, 2011). The proglacial area of Morteratsch is set in the Lower  
116 Austroalpine Bernina Nappe, which is mainly composed of plutonic rocks, such as  
117 granodiorites, diorites, syenites and alkaligranites. Rarerly accessory rocks such as  
118 dolomites, gabbros and serpentines are also reported (Büchi, 1994). In the investigated area,  
119 these rock types were not observed (Mavris et al. 2010, 2011). The rock units underwent  
120 greenschist facies metamorphism during the High Alpine orogenesis (Oligocene-Eocene;  
121 Trommsdorff and Dietrich, 1999). This event caused the saussuritisation of primary rock-  
122 forming plagioclases into albite, epidote and calcite (D'Amico et al., 1998).

123 The vegetation cover of the Morteratsch proglacial area has been studied by Burga (1999).  
124 The first flowering plants colonising the young deglaciated surfaces are scattered individuals  
125 of *Epilobium fleischeri* and *Linaria alpina* that appear after about 7 years. First plants of the  
126 community *Oxyrietum digynae* appear after c. 12 years and disappear after c. 27 years. The  
127 establishment of *Larici-Pinetum cembrae* forests takes place after about 77 years on sites  
128 where the soil is more intensely weathered (Burga, 1999). The soils are weakly developed  
129 and are mostly Lithic Leptosols (IUSS Working Group WRB, 2006; Table 1). Present climatic  
130 conditions for the Morteratsch site are 0.5 °C mean annual temperature and 1000-1300 mm  
131 mean annual precipitation.

132

133

## 134 **Material and methods**

135

### 136 *Soil sampling and preparation*

137 Soil samples were collected from 10 soil pits distributed over the whole proglacial area (Fig.  
138 1). This procedure resulted in the collection of a soil chronosequence of surfaces ranging  
139 from 0 (starting from the glacier front or unweathered parent material) to 140 years old. Soil

140 profiles were excavated down to the C horizon (Table 1). For each horizon, 1–2 kg of  
141 material was collected. The parent material ( $t = 0$  yr) is the unweathered glacial sediment  
142 that can be found either at the glacier front or below the soils (C horizon). The unweathered  
143 parent material used for the optical investigations was collected at the bottom of the  
144 excavated profiles or glacial front ( $n = 7$  C horizon samples were considered). The soil and  
145 parent material samples were oven-dried and sieved to 2 mm (Department of Geography,  
146 University of Zurich, Switzerland). An aliquot of the fine earth sample ( $< 2$  mm) was washed  
147 with de-ionised water and, when present, the suspended organic fraction ( $< 1$  g/cm<sup>3</sup>) was  
148 removed by floating. These samples were used for the preparation of thin sections. Fine  
149 earth samples (material with a diameter  $< 2$  mm) were dispersed with an ultrasonic  
150 disperser, then embedded in polyester resin and cut to a thickness of c. 30  $\mu$ m in order to  
151 ensure a good transparency of the sample. Thin sections were then polished and carbon  
152 coated. The grain size of the investigated minerals was in the range of 10 to 800  $\mu$ m.  
153 In total,  $> 20$  grains were analysed using the Nomarski DIC microscopy or SEM-EDX.

154

#### 155 *SEM-EDX*

156 The fine-earth fraction of  $n = 4$  topsoil samples and  $n = 1$  parent material were analysed  
157 using scanning electron microscopy (SEM) and energy-dispersive spectroscopy (EDS). The  
158 analysis was performed using a Dual Beam Quanta 200 3D FEI coupled with EDX, with Dual  
159 BSD detector and W emitter operating at an accelerating voltage of 20 kV. The EDS detector  
160 is equipped with an ultra-thin window allowing detection of mineral elements and carbon.  
161 EDS provided the elemental composition of the solid phases and helped to identify them  
162 (point analyses and elemental maps). Investigations were performed on both thin sections  
163 (carbon coated) and loose granular samples (uncoated) at the Institute for Building Materials  
164 (ETH Zurich, Switzerland).

165

#### 166 *Cathodoluminescence*

167 Luminescence is a common phenomenon in inorganic and organic compounds, resulting

168 from an emission transition of anions, molecules or a crystal from an excited electronic state  
169 to a ground or other state having less energy (Götze, 2009). In the present study,  $n = 4$   
170 topsoils and  $n = 1$  parent material were investigated. CL measurements were carried out at  
171 the Institute of Mineralogy of the TU Bergakademie Freiberg (Germany) using a hot cathode  
172 CL microscope HC1-LM (Neuser et al., 1995). The system was operated at 14 kV  
173 accelerating voltage and a current density of about  $10 \mu\text{A}/\text{mm}^2$ . Luminescence images were  
174 taken on-line during CL operations using a Peltier-cooled digital video camera (KAPPA 961-  
175 1138 CF 20 DXC). CL spectra were recorded in the wavelength range 320 to 900 nm using  
176 an Acton Research SP-2356 digital triple-grating spectrograph with a Princeton Pixi 256B  
177 Spec 10 CCD detector that was attached to the CL microscope by a silica-glass fibre guide.  
178 CL spectra were measured under standardised conditions (wavelength calibration by a Hg-  
179 halogen lamp, spot width  $30 \mu\text{m}$ , measuring time 1–5 s).

180 Spectral data evaluation was performed considering the relative intensity (counts/s) of the  
181 luminescence emission from each sample. The CL spectra depend on the crystal orientation  
182 (Barbarand and Pagel, 2001; Finch et al., 2003). Therefore, a comparison of the same  
183 mineral type in different samples must be done carefully if the orientation is not the same.  
184 Because we were analysing soil systems, a similar orientation of the investigated minerals (in  
185 a time series) could not be achieved. A quantitative analysis of weathering in the considered  
186 time-span was therefore not possible. To have, however, a semi-quantitative indication for K-  
187 feldspar and plagioclase, peaks in the CL spectra were normalised to the Fe peak of the  
188 minerals in the parent material and in the soil of interest.

189

#### 190 *Nomarski DIC microscopy*

191 Nomarski DIC microscopy is a modern technique applied in material science to visualise  
192 different phases and/or to image the surface relief of samples (Fig. 2). In the present study,  $n$   
193 = 4 topsoils and  $n = 1$  parent material were investigated. The polarisation objectives used  
194 have a magnification/numerical aperture of 20x/0.50, 40x/0.75 and 50x/0.80 with specific DIC  
195 prisms for transmitted and reflected light studies (Götze, 2009). The microscope was coupled

196 with a digital video camera. The investigations and observations were carried out at the  
197 Institute of Mineralogy of the TU Bergakademie Freiberg (Germany).

198

#### 199 *Total element analysis*

200 Element pools in the soil (Ca, Mg, Na, K, Si, Fe, Al, Ti, and Mn) were determined by a total  
201 dissolution method for the parental material (average of n = 7 C horizons) and the topsoils  
202 having an age > 100 yr (n = 4; Table 2). Oven-dried samples were dissolved using a mixture  
203 of HF, HCl, HNO<sub>3</sub>, and H<sub>3</sub>BO<sub>3</sub> (Hossner, 1996) in a microwave oven and at a pressure of c.  
204 25 bar in a closed system. Concentrations were determined by an AAAnalyst 600 Perkin  
205 Elmer Atomic Absorption Spectrometer (AAS) at the Department of Geography, University of  
206 Zurich (Switzerland).

207

#### 208 **Results**

209 The soil mineralogy of the Morteratsch proglacial area has a predominantly silicatic character  
210 due to the granitoid parent rock (Mavris et al. 2010, 2011 and Table 2). In the parent material  
211 the most abundant phases detected were quartz, K-feldspar and Na-rich plagioclase followed  
212 by Fe-rich mica (biotite) and epidote. Apatite was also detected. Furthermore, accessory  
213 phases such as pyrite, titanite and ilmenite were detected, but were not the object of this  
214 study. Secondary phases due to weathering were almost undetectable. The total  
215 concentration of elements remains stable with exposure time (Table 2). The variability is  
216 predominantly due to some inhomogeneities of the glacial sediment.

217 The morphology of the minerals is characterised by smooth surfaces and sharp angles –  
218 typical for a low weathering degree and limited transport (see Figs. 3 and 4). The amount of  
219 rounded grains is low, which is common for morainic material having a minimal chemical  
220 leaching and limited transport.

221

#### 222 *Quartz*

223 SEM-EDX. Quartz grains of the parent material have sharp edges and conchoidal surfaces,

224 typical for short glacial transport. Quartz from the 140 year old soil features a much more  
225 uneven surface. The degree of roundness of the grains is higher, and at high magnifications  
226 weak traces of chemical corrosion could be detected (Fig. 3). Elemental mapping of quartz  
227 grains did not give any evidence of chemical changes following 140 years of weathering,  
228 because, apart Si or O, all other elements were below the detection limit of EDS.

229 CL. The luminescence for the investigated quartz grains was homogeneously dark brown-  
230 blue. Primary internal structures (e.g. growth zoning) were usually not observed in the quartz  
231 crystals (Fig. 4). CL spectra of quartz indicated some structural changes. The observed  
232 broad CL bands of quartz at 450 nm and 650 nm are typical for granitoid-pegmatitic quartz  
233 (Richter et al., 2003). The first band represents a defect due to an oxygen vacancy (Stevens  
234 Kalceff and Phillips, 1995). The second band is caused by 'nonbridging oxygen hole centres'  
235 (NBOHC) (Götze et al., 2001; Fig. 2). A detailed observation of the quartz spectrum of the  
236 fresh sediment revealed a small contribution at 620 nm. This is attributed to Si-OH (silanol  
237 groups) and becomes hindered by the broad 'NBOHC' band with time (Stevens Kalceff and  
238 Phillips, 1995; Fig. 5). In addition, a pronounced band was detected at 710 nm for the quartz  
239 of the parent material. This band most likely indicates a substitutional incorporation of Fe<sup>3+</sup>  
240 into the quartz lattice as observed in other silicate structures. This would be typical for the  
241 alkali-metasomatic alteration (finitisation) of the granite (Götze et al., 2001).

242 Nomarski DIC microscopy. This technique revealed a quite homogeneous morphology for  
243 quartz. The rare fractures detected along the surface of the crystals showed that only – if  
244 ever – a slight corrosive pre-exposure occurred. In the oldest soil, no clear weathering  
245 features were observed.

246

#### 247 *Feldspar*

248 SEM-EDX. K-feldspar and Na-plagioclase display similar chemical stability with respect to  
249 weathering (Allen and Hajek, 1989). Morphologically, we were not able to clearly distinguish  
250 the two phases using SEM. In the parent material, the grains appeared to be coarsely-  
251 shaped, with sharp, regular cleavage edges, indicating short transport and mechanical

252 abrasion. Across the grain surfaces, a diffuse roughness was observed in both parent  
253 material and topsoil samples.

254 CL. The luminescence of K-feldspar showed a blue-violet colour (Fig. 4). Plagioclase showed  
255 green luminescence, slightly fading as a function of time. The heterogeneous distribution of  
256 CL colours in feldspar minerals indicates weathering/alteration already in the initial rock  
257 material. In the CL spectra two emission bands could be observed (Fig. 5). The band at c.  
258 460 nm, whose intensity decreases with soil age, is related to Al-O<sup>-</sup>-Al defects (e.g. Marfunin,  
259 1979). The band at c. 680 nm is generated by Fe<sup>3+</sup> impurities in Al lattice positions (Götze et  
260 al., 1999).

261 Nomarski DIC microscopy. Na-plagioclase and K-feldspar exhibited similar features (Fig. 6).  
262 Using parallel and crossed Nicols polarised light, the grains revealed surfaces having  
263 distinctive, cross-cutting preferential cleavage fissures. Already in the glacial till, two sets of  
264 weathering surfaces were distinguishable (Fig. 6). The first one (red arrow) is observed at the  
265 outer part of the grain from where the fissures started to penetrate and propagate randomly  
266 into the inner part of the mineral grain. The second one (black arrow) developed along  
267 parallel sets of cleavage surfaces.

268

#### 269 *Biotite*

270 SEM-EDX. Biotite occurs as layered aggregates with some typical perfect basal cleavages  
271 (Fig. 7). In both the parent material and topsoil mechanical deformation and delamination of  
272 the layers due to glacial abrasive activity was observed. Furthermore, Ti-containing minerals  
273 (titanite?) and feldspar were detected among the mica layers (Fig. 8). SEM-EDX did not  
274 evidence a development of secondary phases. In the surface layer and along fissures of the  
275 biotite platelets, K appeared depleted compared to Fe due to leaching processes (Fig. 8).

276 CL. Biotite shows a very weak CL intensity because of the quenching by structural iron.  
277 Therefore, neither images nor CL spectra were obtained.

278 Nomarski DIC microscopy. Within the weathering sequence, an increasing delamination of  
279 biotite with time could be observed. Using the Nomarski technique and SEM, larger spaces

280 between the mica layers were detectable in the older samples (Fig. 7). As a consequence of  
281 the delamination process of mica, a much higher surface area became available for chemical  
282 weathering. However, the presence of structurally weaker phases, i.e. newly-formed clay  
283 minerals, could not be detected within the biotite platelets. This confirms physical - rather  
284 than chemical - alteration of biotite.

285

#### 286 *Apatite (accessory phase)*

287 SEM-EDX. In the glacial till, apatite was detected as sporadically occurring, euhedral crystals  
288 having a size of up to 30  $\mu\text{m}$ . Despite a rather low resistance to weathering in soil profiles  
289 having low pH (Allen and Hajek, 1989; Taunton et al., 2000), it was still possible to find  
290 apatite crystallites in the oldest topsoil. The EDX measurements did not evidence any  
291 significant chemical change in the considered time-span.

292 CL. Apatite was observed having a strong yellow luminescence emission but did not feature  
293 any rim (or growing) structures in the crystal, reflecting a homogeneous composition of  
294 apatite (Fig. 4). The CL emission spectra of apatite evidenced sharp emission lines of REE  
295 together with a strong  $\text{Mn}^{2+}$  band at c. 560 nm (Mariano, 1988). A strong enrichment in  $\text{Ce}^{3+}$ ,  
296  $\text{Nd}^{3+}$ ,  $\text{Sm}^{3+}$  and  $\text{Eu}^{2+}$  is typical for apatite originating from alkaline magmatic complexes  
297 (Zhang et al., 1985; Fleischer and Altschuler, 1986; Mariano, 1988; Boudreau and Kruger,  
298 1990; Kempe and Götze, 2002).

299 Nomarski DIC microscopy. The detected apatite particles were too small (i.e.  $< 30 \mu\text{m}$ ) to be  
300 properly analysed from a morphological point of view.

301

#### 302 *Epidote*

303 SEM-EDX. Epidote occurs mainly as elongated crystals having a size of  $> 50 \mu\text{m}$  (Fig. 9). In  
304 the glacial till, no pre-exposure weathering traces were detected on the epidote crystals. EDX  
305 revealed two varieties, a common Fe-rich and a sporadically-occurring REE-bearing epidote  
306 (data not shown; see also Mavris et al., 2010). The occurrence of the REE-bearing phases  
307 correlates with the dismantlement of pegmatite veins (cf. Büchi, 1994). Epidote did not show

308 any distinct chemical or mechanical weathering alterations in the oldest topsoil.

309 CL: Epidote featured a very weak (dark green) luminescence glow. The needle structure of  
310 the small crystals, coupled with the strong interfering luminescence of the hosting feldspar  
311 did not allow spectral CL measurement of epidote.

312 Nomarski DIC microscopy. In contrast to SEM-EDX, a distinct epidote alteration was  
313 traceable (Fig. 9). Epidote appeared to have a physically and chemically affected crystal  
314 structure when compared to the associated (surrounding) phases. The differential  
315 interference contrast evidenced this feature where epidote was associated to tectosilicate-  
316 structured phases. Neither secondary clay minerals nor oxyhydroxides were observed on the  
317 surface of epidote.

318

### 319 **Discussion**

320 The collapse of structure bonds is paramount for the transformation of primary tectosilicatic  
321 structure into secondary phases (e.g. smectite; Aoudjit et al., 1995). Physical weathering  
322 processes (such as freeze-thawing cycles) tend to weaken mineral crystal structures such as  
323 large cation-bond tectosilicatic crystal lattices and thus facilitate a further step of elemental  
324 leaching by circulating waters (e.g. Hall et al., 2002). The chemical and mineralogical  
325 structure of mineral, such as the Fe(II) content, oxygen sharing (the resistance of a primary  
326 mineral to weathering increases the degree of sharing of oxygen between adjacent Si  
327 tetrahedra in the crystal lattice), lattice distortion etc. strongly determine weathering stability  
328 (Curtis, 1976).

329 In the investigated till, it appears that structural bonds – such as Si-O-Si in quartz – start to  
330 break and to transform into free radicals at very early stages of weathering (Figs 3 and 5).

331 This process was also shown in controlled dissolution experiments using quartz (Bennett et  
332 al., 1988). The surface of quartz in aqueous solutions consists entirely of species derived  
333 from the hydrolysis and hydroxylation of broken =Si-O-Si= and the subsequent protonation  
334 and deprotonation of these sites (Furrer and Stumm, 1986; Brady and Walther, 1990). CL  
335 spectroscopy was a helpful tool to detect such nano-scale features (Fig. 5). The increase of

336 both oxygen vacancy and 'NBOHC' bands is the direct consequence of the bond  
337 disintegration and a subsequent leaching of Si (or other accessory impurities present, if any).  
338 In both, the parent material and the oldest soil of the glacier forefield, the presence of apatite  
339 could be verified. Apatite, the main source of inorganic P in the Morteratsch proglacial area,  
340 neither displays significant chemical and structural changes nor specific dissolution patterns,  
341 even after 140 years of weathering. This is in contrast to the findings of Föllmi et al. (2009b).  
342 Their observations of the progressive change in surface morphology of whole apatite grains,  
343 the systematic shift in the composition of P phases from detrital to iron-bound and organic P  
344 and estimations of the weathering rate of detrital P in the moraine samples indicated that  
345 apatite is biogeochemically actively weathered in the proglacial areas of the Rhône and  
346 Oberaar glaciers. That we do not observe such a trend in the proglacier area of Morteratsch  
347 might be due to a different chemical composition of the apatite or the low number of  
348 observations. The CL measurements seem to indicate a weathering of REE in the  
349 investigated apatite; however, due to technical constraints, a definitive conclusion is not  
350 possible.

351 The amount of epidote decreased as a function of time (see Mavris et al., 2010). Apparently,  
352 the sorosilicate structure of epidote undergoes a leaching of major cations (predominantly  
353  $\text{Ca}^{2+}$ ), thus featuring a chemically-changed crystal lattice (Fig. 9). However, an extended X-  
354 ray absorption fine structure (EXAFS) spectroscopy study on Fe in the Damma glacier  
355 forefield showed that the Fe fraction within epidote remained constant over the  
356 chronosequence (of c. 150 years), indicating a rather low Fe weathering rate of epidote  
357 (Kiczka-Cyriac, 2010).

358 Within the 140 years of weathering the Al-O-Al bonds of K-feldspar seemed to decrease  
359 (Fig. 5) which might be due to a weakening of the crystal structure. Compared to Fe, less Al  
360 structural bonds are present in the mineral structure. These structural changes were also  
361 confirmed by the Nomarski DIC microscopy (Fig. 6). This reveals remarkable chemical  
362 variations most often on the grain surface and at a lattice level. The standardised CL spectra  
363 of plagioclase were overwhelmingly overlapping (Fig. 5F) and, consequently, no major

364 changes could be observed. Finch et al. (2003) reported about a (structurally) ordered  
365 plagioclase. The peak ratios after CL excitation/stimulation changed up to 34.3% with the  
366 different crystallographic directions of the plagioclase. One way to detect the effective  
367 change of specific bonds in a time series is the normalization (ratio) of the band of interest to  
368 an 'inert' compound (such as  $\text{Fe}^{3+}$  in oxic environments). If this ratio varies between two  
369 samples more than 34.3%, a real crystallographic change is more likely. If we assume that  
370 such a change as a function of orientation is also valid for the plagioclase in the Morteratsch  
371 area then no changes can be measured after 140 yr because the peak ratio changes remain  
372 below this value. The Al-O<sup>-</sup>-Al/Fe (460/680 nm) ratio is for 0 years (parent material) = 0.51  
373 and for 140 years = 0.48. The change in the Al-O<sup>-</sup>-Al/Fe ratio of K-feldspar, however,  
374 between 0 years (2.43) and 140 years (1.24) is much higher than one would expect from a  
375 spectra-change of the same mineral with varying orientation (according to Finch et al., 2003).  
376 Consequently, a part of the Al-O<sup>-</sup>-Al decrease in the K-feldspar (Fig. 5E, F) must be  
377 attributed to weathering (and not only to a change in orientation of the mineral).

378 Both chemical and physical transformations - namely, elemental depletion of K over Fe, and  
379 increased delamination over time - confirmed biotite weathering in the proglacial sediments  
380 (Figs 7 and 8). As observed by other authors, biotite typically expands with time, delaminates  
381 and contributes to an increase in porosity of the weathered rock (Meunier et al., 2007; Rossi  
382 and Graham, 2010; Graham et al., 2010, Caner, 2011). The opening of fissures in mineral  
383 grains and the increase in porosity finally give rise to accelerated alteration rates (chemical  
384 weathering; Meunier et al., 2007). Fast weathering rates, not only of biotite, can usually be  
385 observed at the beginning of soil formation when fresh and reactive surfaces are available  
386 (Egli et al., 2003; Mavris et al. 2011). With time, these rates often decrease (Kump et al.,  
387 2000; Brantley and Mellott, 2000; Peters, 2009) due to the progressive occlusion of pores by  
388 secondary minerals and a decreased chemical potential in the alteration product (Caner,  
389 2011).

390 The combined CL/Nomarski DIC/ESEM observations of the Morteratsch sediments are  
391 based on only a small number of observations. Consequently, our findings cannot be

392 conclusive and, in particular, more CL measurements on individual minerals are required.

393 The total elemental concentration in the investigated topsoils reveals that the loss of  
394 structure-modifying cations within the investigated time span of 140 y is not yet high enough  
395 to give rise to a chemical trend along the proglacial area (Table 2). However, on the much  
396 smaller atomic and mineral structural scale weathering features can be detected using a  
397 combination of CL, Nomarski and SEM-EDX techniques.

398

399

#### 400 **Conclusions**

401 Active chemical and physical weathering in a high Alpine chronosequence of recently  
402 exposed, granitic sediments could be documented. We have the following main findings:

- 403 • Biotite shows distinct mechanical weathering due to delamination of layers within the  
404 observed time-span of 140 years.
- 405 • Although the bulk chemistry of the soils did not change over time, K-feldspar and Na-  
406 plagioclase showed some chemical transformations with respect to Al (K-feldspar) and Mn  
407 (plagioclase). SEM-EDX analyses indicated a loss of K in biotite.
- 408 • Apatite seemed to be resistant to weathering.
- 409 • Epidote, however, shows clear weathering features (such as physical structure  
410 disintegration) after 140 years.

411 The combined use of chemical, mineralogical and spectroscopic analyses proved to be a  
412 useful approach in deciphering weathering patterns even at the atomic bond scale (crystal  
413 structures). CL and Nomarski DIC have a great potential in detecting and analysing  
414 chemical, physical and mineral structural changes due to weathering also in soils. Such an  
415 approach should in future enable a qualitative and semi-quantitative estimation of lattice  
416 position losses in primary and accessory minerals.

417

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425

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## Figure captions

Fig. 1. Overview of the investigated area with the monitored sites in the proglacial area. The isochrones are according to Burga et al. (1999).

Fig. 2. Schematic sketch (modified after Götze, 2009) showing: a) simplified structure of quartz with some typical features detectable using cathodoluminescence (modified after Götze et al., 2001). b) Nomarski DIC microscopy. Optical diagram with the structure of the DIC microscope; bottom: the phase difference ( $\Phi$ ) is explained by geometric (step height 'd') and physical factors (difference in the material specific phase shift 'a' and 'b').

Fig. 3. SEM micrographs with a) quartz grain from the 140 yr old topsoil and b) surface morphology showing an early chemical corrosion of quartz. The arrows point to distinct surface dissolution pits.

Fig. 4. Cathodoluminescence micrographs of apatite, quartz, plagioclase and alkali feldspar of the parent material (0 yr) and 140 yr old topsoil. *Apatite*: yellow luminescing crystals due to the activation of  $Mn^{2+}$ ; *quartz*: blue-brown CL; *plagioclase*: green luminescing areas with strongly emitting  $Mn^{2+}$  peak; *alkali feldspar*: violet emitting grains, related to both  $Al-O^-Al$  defects and  $Fe^{3+}$  in Al-sites.

Fig. 5. Cathodoluminescence emission spectra of A) quartz, B) alkali feldspar, C) plagioclase and D) apatite. The bold line denotes the 140 yr topsoil and the thin line the parent material. The spectra E (alkali feldspar) and F (plagioclase) are normalised to the Fe-spectrum of the parent material.

Fig. 6. Plagioclase from the parent material (0 yr) and the 140 yr old topsoil. In the parent material, two sets of weathering patterns are underlined: geologically inherited (along cleavage surfaces; red arrows), and pedogenic (mostly on the outer surface of the grains; black arrow). The image shows the grains in thin section in parallel and crossed Nicols transmitted (polarised) light.

Fig. 7. (*above*) Delamination of biotite crystallites in the parent material ( $t = 0$  yr) and in the oldest topsoil ( $t = 140$  yr) observed using SEM; (*below*) biotite delamination as a function of the time documented using Nomarski DIC microscopy.

Fig. 8. Elemental mapping (SEM-EDX) of K and Fe in biotite of the parent material and the oldest soil.

Fig. 9. Micrographs of epidote from the parent material (0 yr) and the oldest topsoil aged (140 yr). The image shows the grains in thin section in parallel and crossed Nicols transmitted (polarised) light.

Figure 1  
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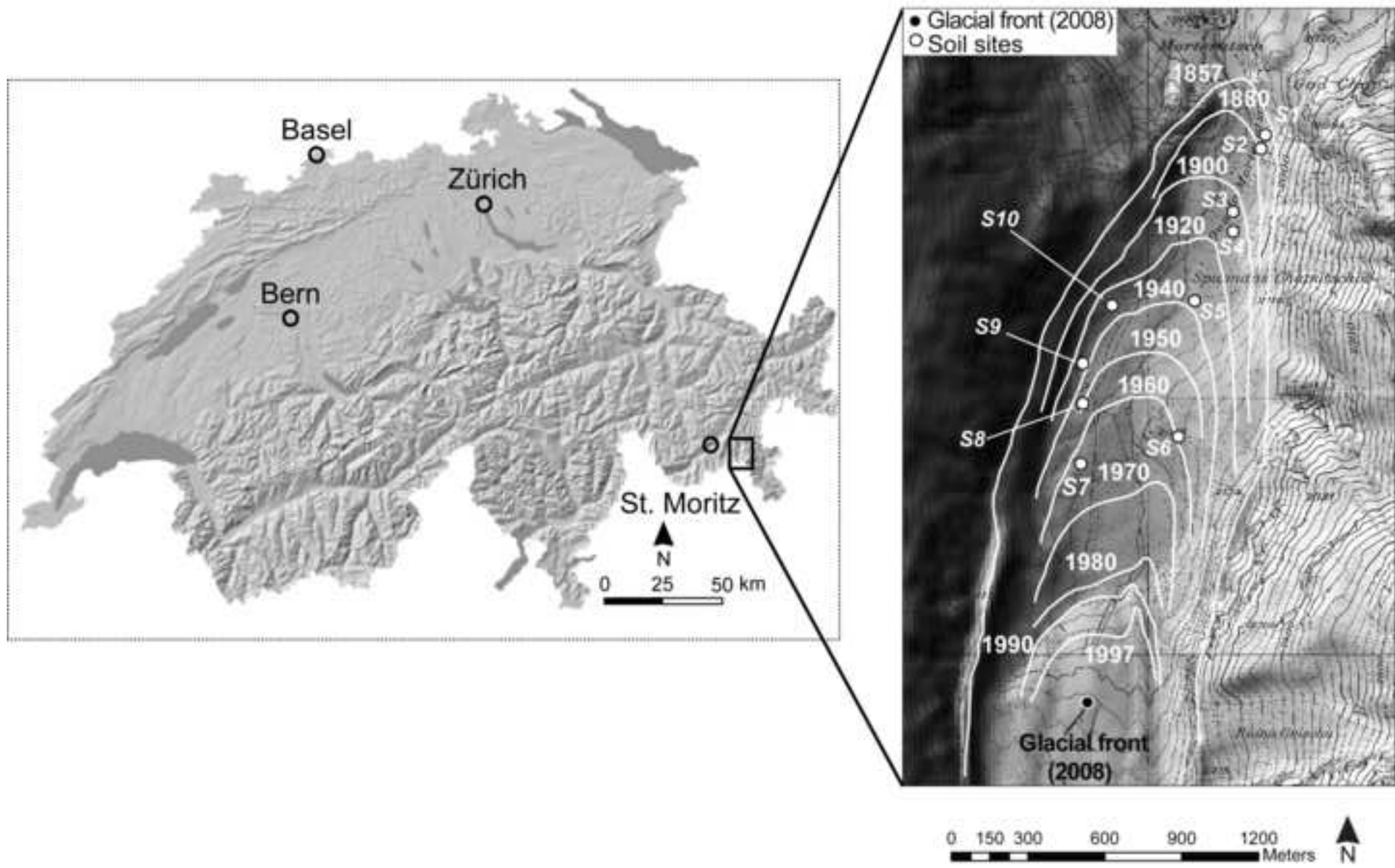
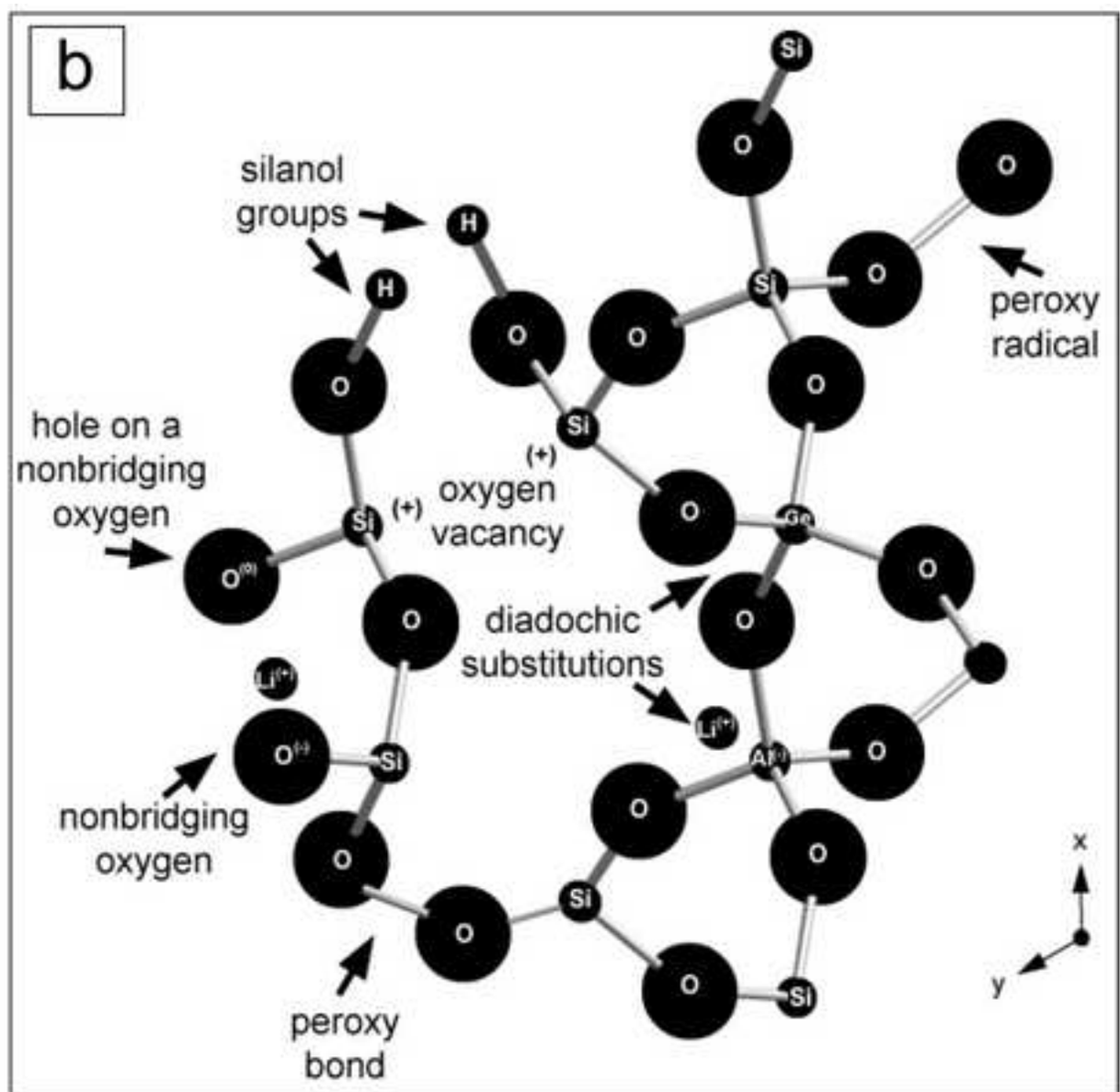
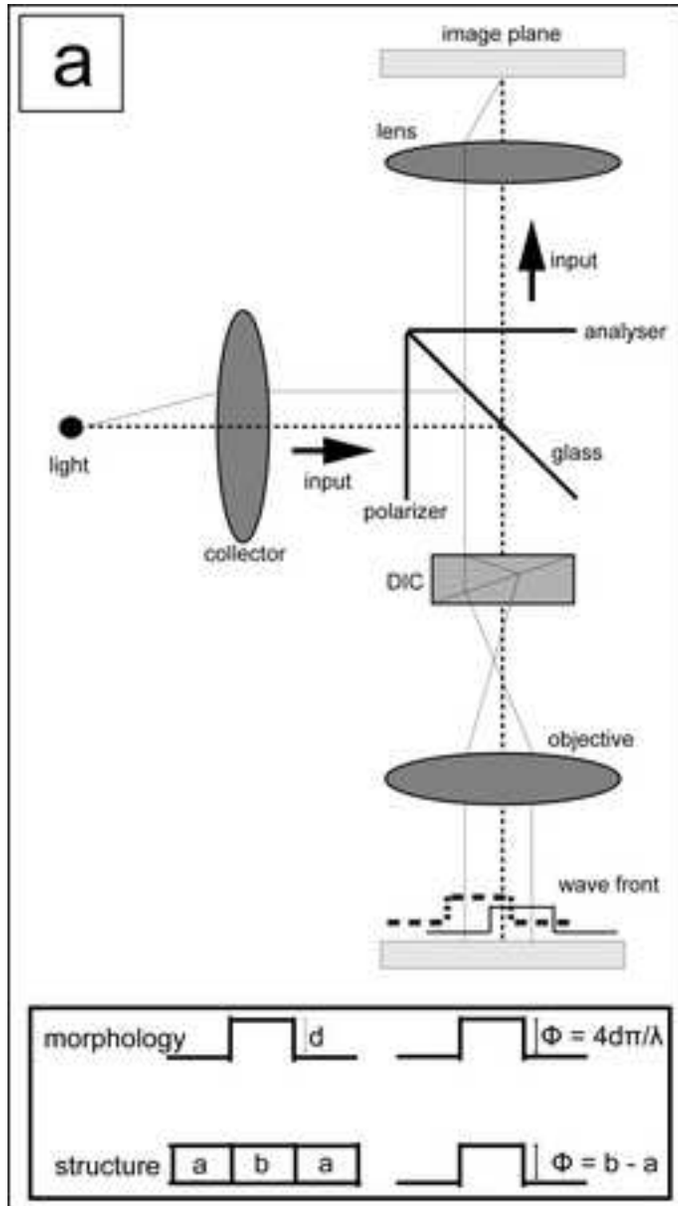


Figure 2  
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**Figure 3**  
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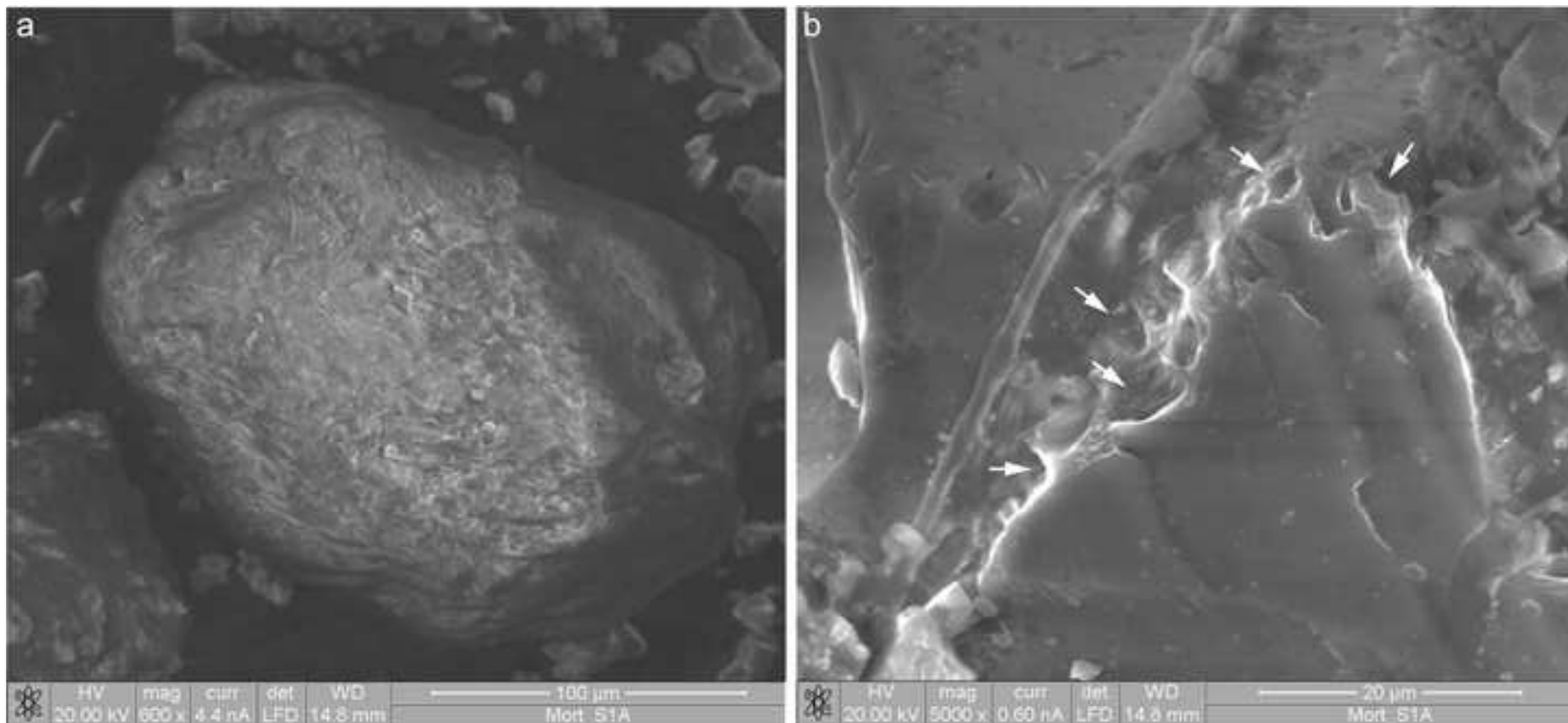


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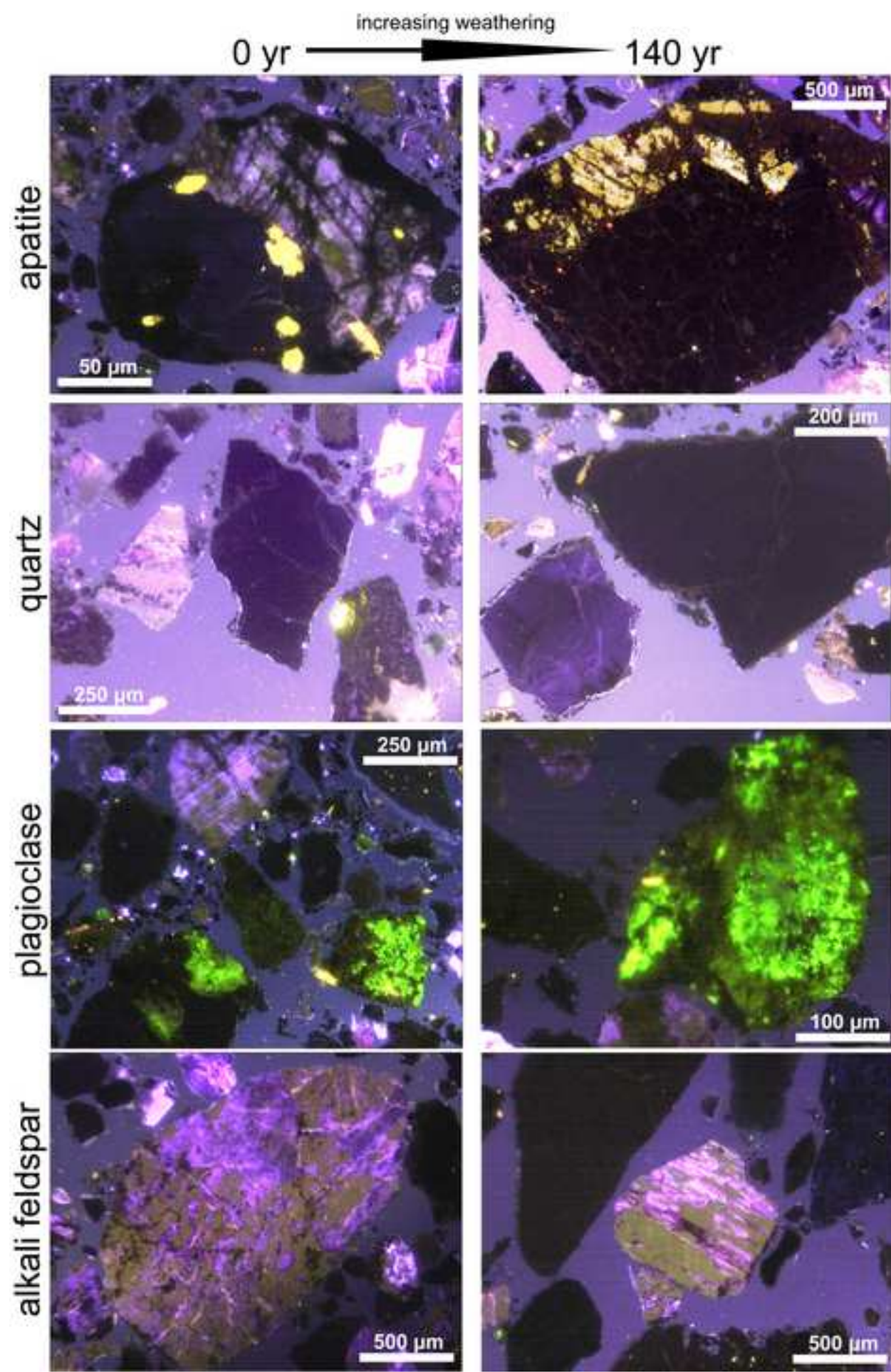


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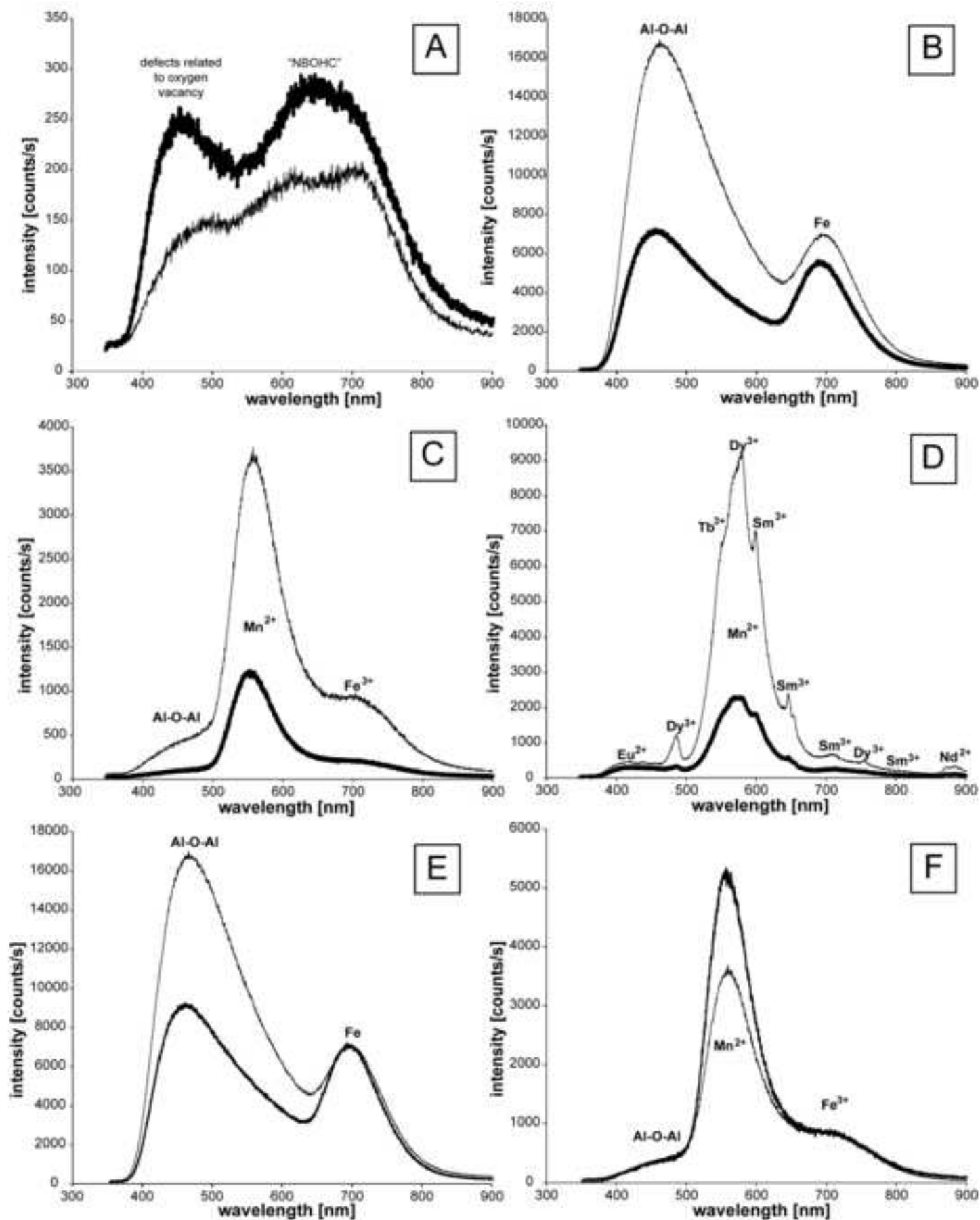


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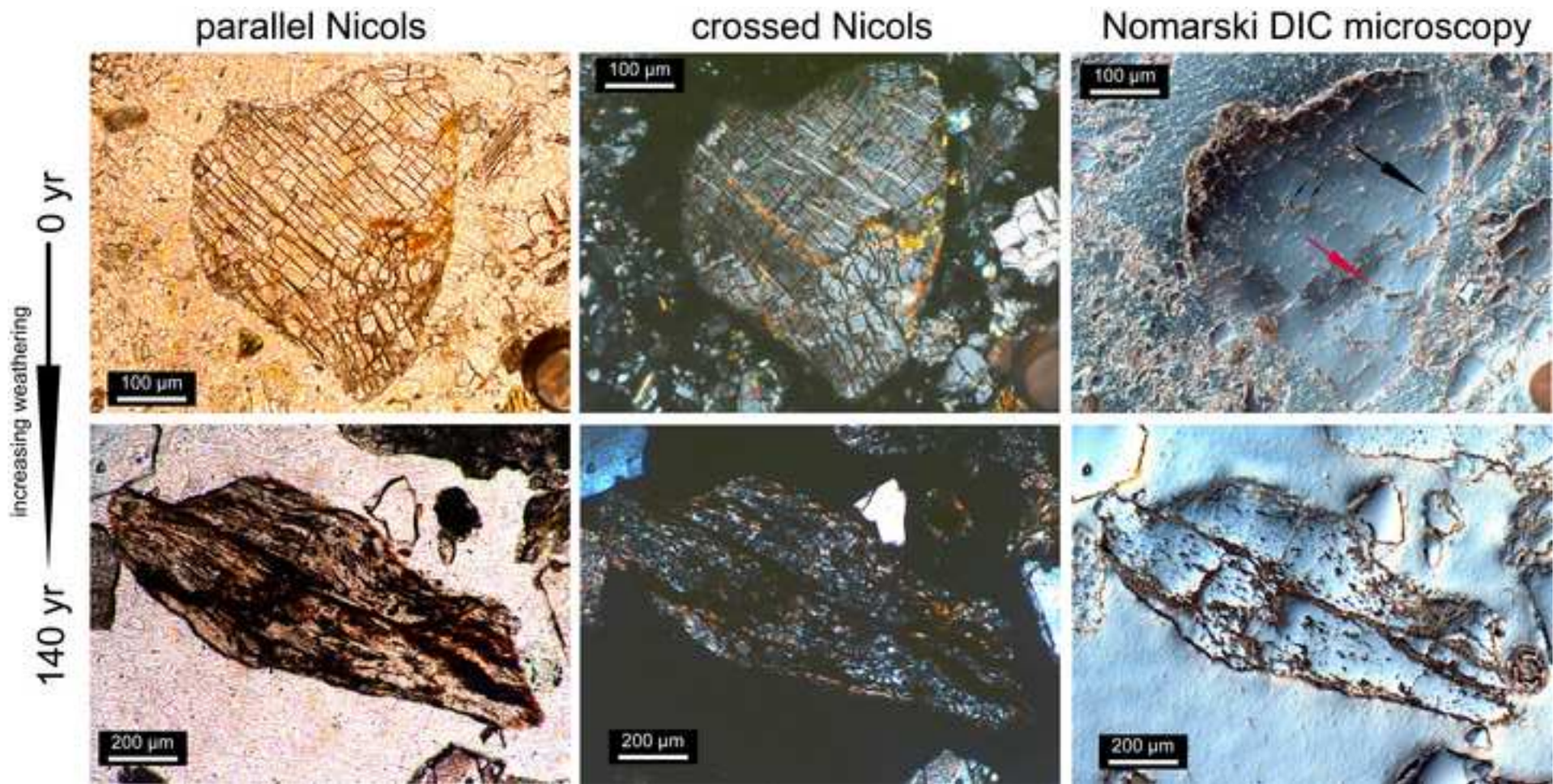
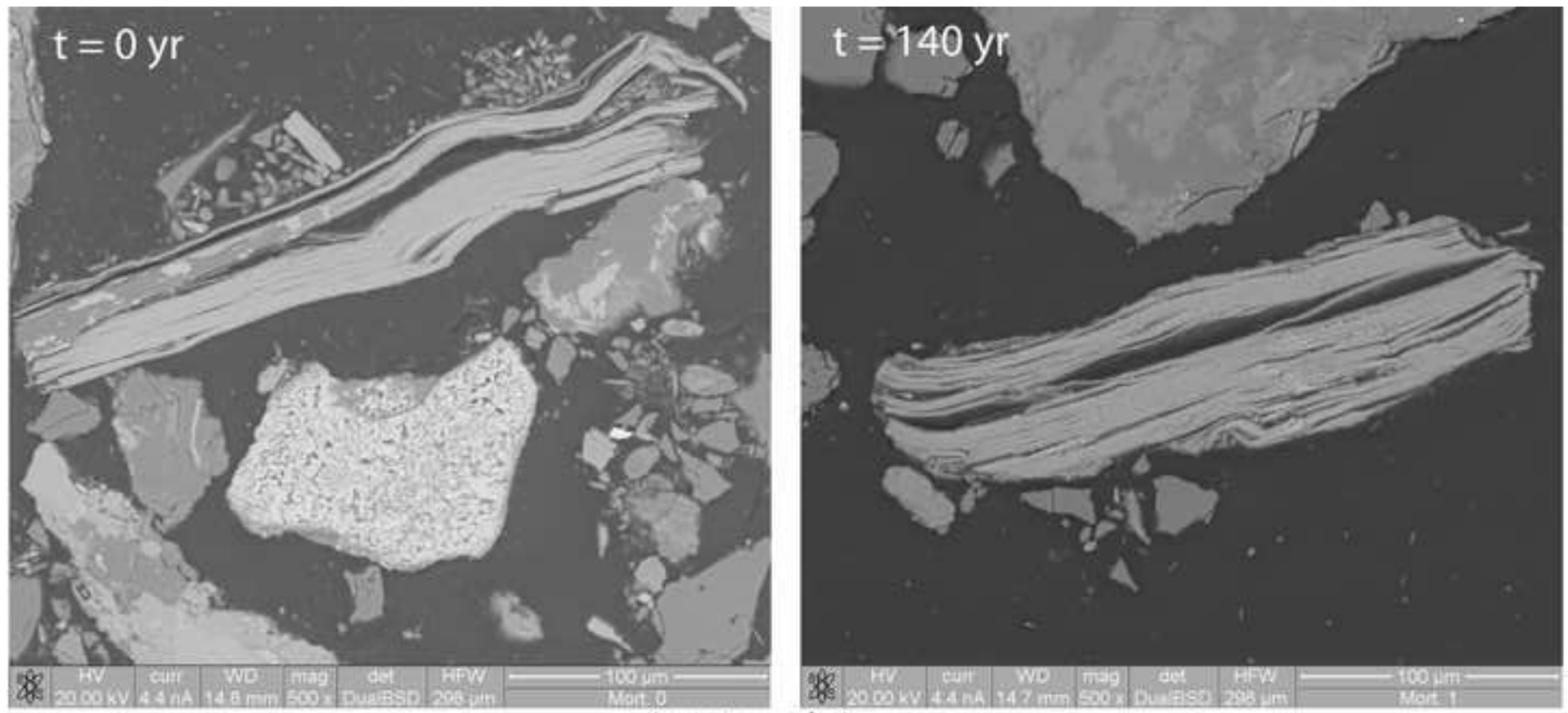


Figure 7  
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increasing weathering

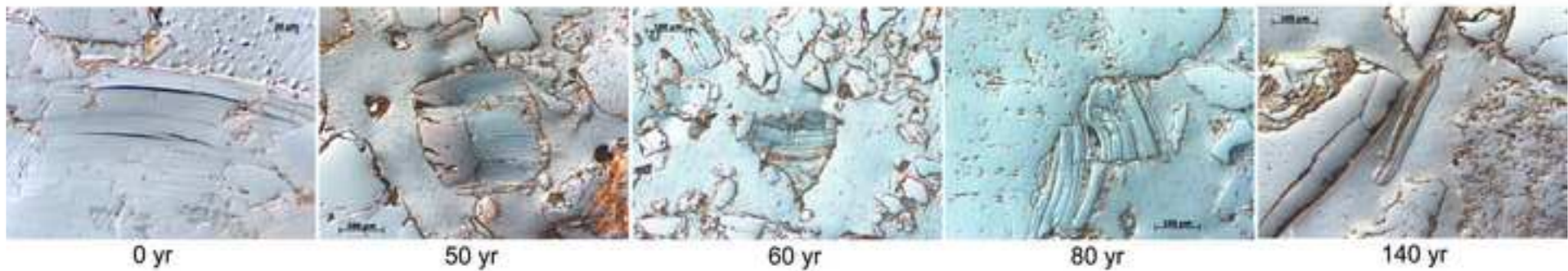


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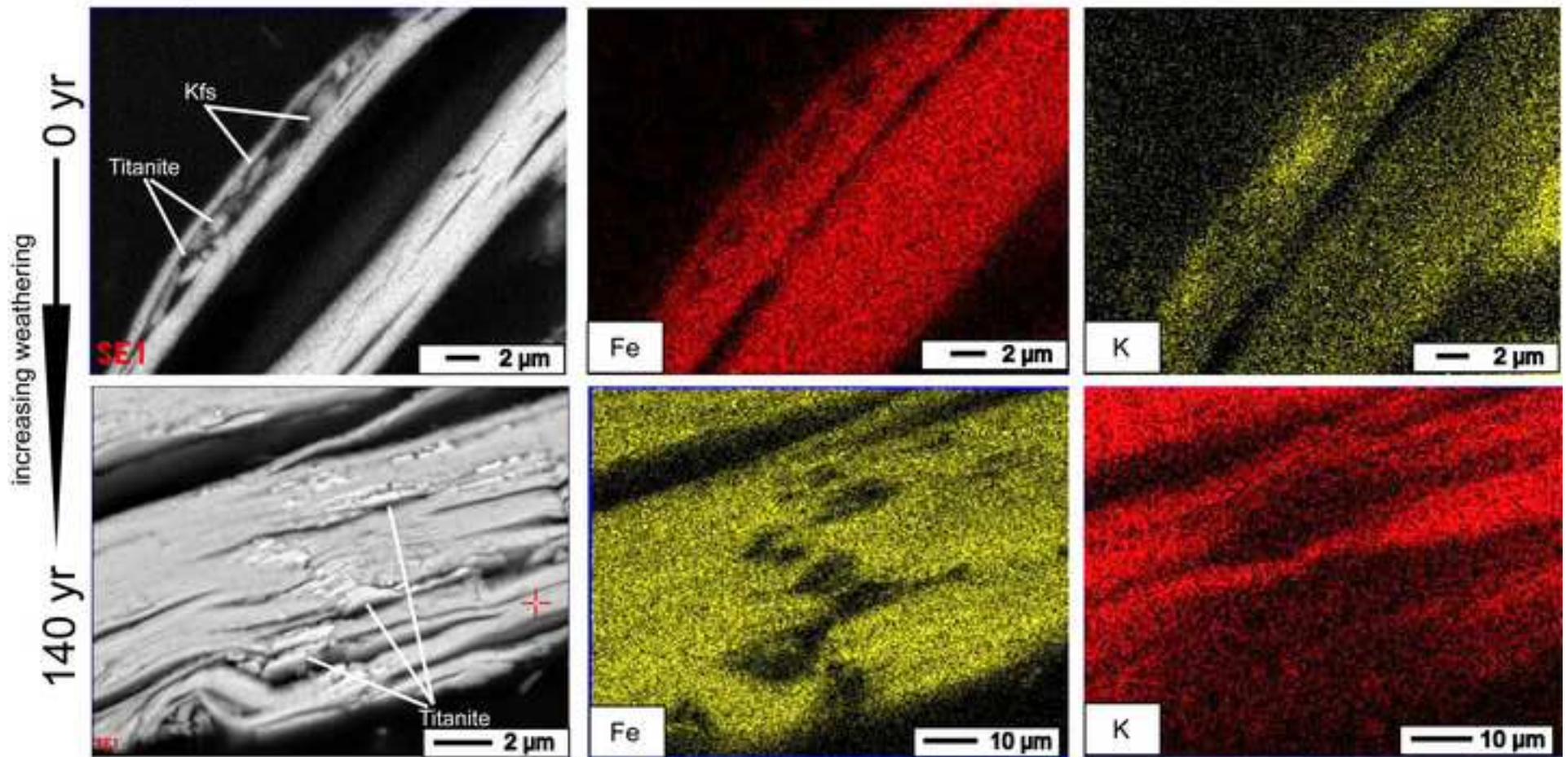
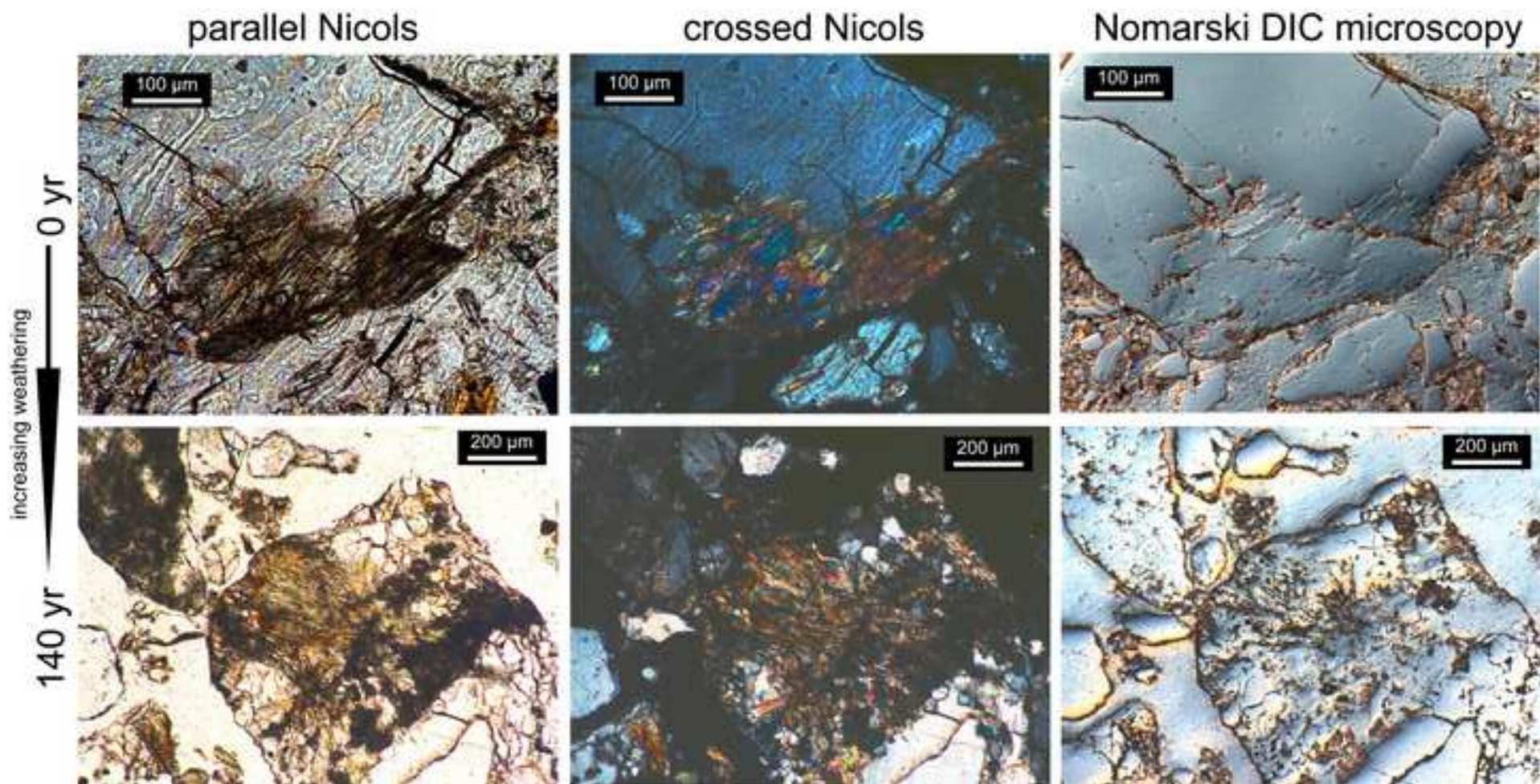


Figure 9  
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**Table 1**[Click here to download Table: Table1.doc](#)Table 1. Properties of the monitored soil sites. Grain sizes: sand (2000-63  $\mu\text{m}$ ), silt (63-2  $\mu\text{m}$ ) and clay (< 2  $\mu\text{m}$ ).

Site/Soil	Soil age (yr)	Horizons	Depth (cm)	Skeleton (wt. - %)	Sand (g/kg)	Silt (g/kg)	Clay (g/kg)	pH (CaCl <sub>2</sub> )
S1/Humi-Skeletal Leptosol	138	O	0-6	41	n.m.	n.m.	n.m.	4.60
		A	6-9	50	777	184	39	4.80
		BC	9-14	53	830	158	12	4.70
		C	14-30	40	757	222	21	4.60
S2/Humi-Skeletal Leptosol	128	A	0-10	64	754	204	42	4.85
		AC	10-40	68	695	272	33	4.90
S3/Humi-Skeletal Leptosol	108	A	0-3	54	667	265	68	5.10
		AC	3-15	70	677	281	42	4.50
S4/Humi-Skeletal Leptosol	98	A	0-1	55	n.m.	n.m.	n.m.	5.30
		AC	1-5	51	939	61	15	5.20
		C	5-30	70	931	57	12	5.20
S5/Humi-Skeletal Leptosol	68	A1	0-1	7	n.m.	n.m.	n.m.	4.85
		A2	1-4	1	530	432	38	4.55
		C1	4-9	36	573	387	40	4.65
		C2	9-20	64	570	372	58	4.60
S6/Skeletal Leptosol	48	A	0-2.5	64	n.m.	n.m.	n.m.	4.80
		C	2.5-25	68	852	129	19	5.00
S7/Skeletal Leptosol	48	A	0-4	26	n.m.	n.m.	n.m.	6.10
		C1	4-11	37	823	146	31	5.20
		C2	11-34	67	747	211	42	5.10
S8/Skeletal Leptosol	58	OA	0-12	63	n.m.	n.m.	n.m.	4.60
		C	12-33	48	712	220	68	4.40
S9/Humi-Skeletal Leptosol	73	O	0-3	44	n.m.	n.m.	n.m.	5.15
		AC	3-10	65	785	175	40	4.40
		C	10-36	58	832	133	35	4.65
S10/Humi-Skeletal Leptosol	78	A1	0-2	49	n.m.	n.m.	n.m.	4.70
		A2	2-10	68	818	143	39	4.50
		AC	10-25	84	733	219	48	4.80

n.m. = not measured

**Table 2**[Click here to download Table: Table 2.doc](#)

Table 2. Total chemical analysis of the &gt; 100 yr topsoils (n = 4) and the parent material (n = 7) with standard deviation (SD).

	Al (g/kg)	Si (g/kg)	Ti (g/kg)	Ca (g/kg)	Mg (g/kg)	K (g/kg)	Na (g/kg)	Fe (g/kg)	Mn (g/kg)
parent material	76.8	310.2	8.00	17.2	9.53	28.7	26.8	22.5	0.58
SD	6.5	27.4	2.38	6.4	3.20	4.1	3.87	6.9	0.16
topsoils (age >100 yr)	65.0	330.4	5.90	27.3	7.32	28.1	25.3	29.1	0.62
SD	5.8	24.0	2.25	22.3	2.77	2.4	2.6	16.1	0.14