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1 **Mercury speciation in a tropical soil association;**
2 **Consequence of gold mining on Hg distribution in French Guiana.**

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23 **Abstract.**

24 Mercury (Hg) speciation was compared in French Guiana pristine soils and in
25 Hg contaminated soils impacted by former (~1950's) gold-mining activities which
26 used Hg for gold amalgamation. Four selective extractions were performed on soil
27 samples to assess the fraction of Hg present as Hg(II) and bond to organic matter
28 (extracted by NH₄OH and KOH), to amorphous iron oxides (ascorbate) and to soil
29 components other than refractory minerals (HCl/HNO₃). In addition, pyrolysis was
30 used to quantify the content of elemental Hg in contaminated soils. X-ray diffraction
31 (XRD) and X-ray fluorescence micro-mapping (μ XRF) were used in combination to
32 selective extractions to assess the nature of targeted components, the possible
33 overlaps between the different extraction procedures and the spatial correlation
34 between Si, K, Fe, Au and Hg.

35 In soil profiles from pristine toposequences, Hg concentrations (0.01-
36 0.49 $\mu\text{g.g}^{-1}$) decreased with increasing depth in soil matrix. Hg concentrations also
37 decreased from ferralsols to acrisols and further to gleysols. In pristine soil matrix, Hg
38 was mainly associated to the clay size fraction (< 2 μm) which was mainly constituted
39 of amorphous and crystalline Fe oxides (Al-substituted goethite and hematite),
40 gibbsite and fine organic matter (OM), whose relative abundances vary along the soil
41 association. Total Hg concentration was positively correlated with sulfurs and organic
42 carbon suggesting the association of Hg with OM sulfur-bearing functional groups.
43 Gleysols were depleted in Hg because of the prevailing reducing conditions that lead
44 to the dissolution of iron oxides. In the same soil profiles, Hg concentrations in
45 ferruginous nodules, which make up most of the soil coarse fraction (> 2 mm), were
46 similar to those reported in the pristine soil matrix. These nodules mainly contained
47 Al-substituted hematite and goethite and were especially abundant upslope in

48 ferralsols and Acrisols. Gold-mined gleysols were strongly disorganized by former
49 activities as neither the original structure nor the texture was preserved. Soil
50 granulometry was dominated by gravels, sands and silts. Hg concentrations (0.09-
51 9.22 $\mu\text{g}\cdot\text{g}^{-1}$) largely exceeded those in pristine soils. μXRF allowed the identification
52 of Au-amalgamated Hg and of elemental Hg droplets. Pyrolysis confirmed Hg to be
53 mainly present in its elemental form in contaminated soils. Selective extractions
54 showed additional minor contributions of Hg(II) associated to OM, and to Al or Fe
55 oxides.

56 The combination of selective extractions with XRD and μXRF data showed
57 that extraction efficiency is strongly dependent on the soil type, and that this
58 efficiency needs to be determined on a soil-by-soil basis for Hg speciation studies.
59 KOH extraction was especially delicate as crystalline and amorphous oxides were
60 extracted together with organic matter.

61 Key-words: Mercury, Tropical soils, French Guiana, gold-mining, XRD, selective
62 extraction, speciation.

63 **1. Introduction**

64 Tropical soils from the Amazonian basin are known to have accumulated
65 atmospheric Hg over millions of years, thus inducing geochemical backgrounds
66 higher than those reported in boreal and temperate climate soils (Carmouze et al.,
67 2001). In addition, past and current gold mining activities, which represent the main
68 contribution to anthropogenic emissions (Lacerda, 1997; Roulet et al., 1998; Guedron
69 et al., 2006), contribute to increase further the total Hg concentrations in these soils.
70 Small-scale mining activities have increased significantly during the worldwide gold
71 rush, created by the gold price increase in the 1980's. UNIDO (United Nations
72 Industrial Development Organization) estimates that in 2004 10 to 15 million people
73 were working in small-scale gold mines releasing 650-1000 tons of Hg annually
74 (Veiga and Baker, 2004). In addition to this major atmospheric emission, gold mining
75 activities release large amounts of Hg-rich particles into hydrosystems where
76 methylmercury (MMHg) production has been clearly identified in anoxic and suboxic
77 areas (Benoit et al., 2003; Coquery et al., 2003). In turn, MMHg accumulation along
78 the aquatic food chains is a threat to Amerindian populations whose diet relies mainly
79 on fish (Lebel et al., 1996; Frery et al., 2001; Barbosa et al., 2003). To assess Hg
80 mobility and its availability to methylating microorganisms, it is thus crucial to
81 determine how Hg is bond to soil particles.

82 The retention capacity of tropical soils varies as a function of soil components
83 which depends on soil types and horizons (Charlet and Sposito, 1987; Charlet and
84 Sposito, 1989; Fontes and Gomes, 2003). In particular, the relative proportions of soil
85 components responsible for Hg retention [clay minerals, Fe-, Al-, and Mn-
86 (oxyhydr)oxides, organic matter] vary with the geochemical conditions which in turn

87 depend on the topographic position along the soil association (Boulet et al., 1993; do
88 Valle et al., 2005; Fritsch et al., 2005; Fritsch et al., 2006).

89 Hg has a high affinity for soil organic matter, especially for reduced sulfur
90 groups such as thiol ligands (Schuster, 1991; Skyllberg et al., 2000; Khwaja et al.,
91 2006; Skyllberg et al., 2006). High Hg levels can also be associated to Fe-
92 (oxyhydr)oxides. In particular, amorphous or poorly crystalline oxides are known as
93 sinks for metal contaminants owing to their large surface area and/or microporous
94 structure (Trivedi and Axe, 2001). Ferruginous nodules, formed during the lateritic
95 weathering process, also contain elevated Hg contents (Roulet and Lucotte, 1995;
96 Roulet et al., 1998; De Oliveira et al., 2001; Brabo et al., 2003).

97 Solid-phase chemical speciation has been widely used on sediment samples
98 to decipher metal distribution in contaminated systems (Wallschlager et al., 1998;
99 Rodriguez Martin-Doimeadios et al., 2000; Beldowski and Pempkowiak, 2003; Bloom
100 et al., 2003; Boszke et al., 2006). Most of the recent studies of Hg speciation in
101 natural or contaminated sediments, soils and tailings used sequential extraction
102 procedure (SEP) or Hg pyrolysis procedure following published recommendations
103 (Eagenhouse et al., 1978; Di Giulio and Ryan, 1987; Biester and Scholz, 1997;
104 Bloom et al., 2003). While some studies strictly follow the described SEP (Sanchez et
105 al., 2005; Beldowski and Pempkowiak, 2007), many authors adapt it to the studied
106 matrix. Each extraction step of the SEPs for soils and sediments is critical, as it is
107 challenging to obtain an adequate recovery by preventing losses, contamination or
108 speciation changes, and limiting the interferences (Leermakers et al., 2005).

109 In the present study, the speciation of Hg among soil components was
110 determined in soil profiles along pristine soil toposequences and in soils
111 contaminated with Hg by former gold mining in a French Guiana watershed.

112 Separate selective extractions were used to quantify Hg distribution between soil
113 carrier phases, in order to avoid cumulative errors intrinsic to SEPs. In addition, the
114 combination of X-ray diffraction (XRD) and chemical analysis allowed the
115 identification of the extracted phases and of possible overlaps between selective
116 extractions (cross recoveries). Intrinsic limitations of the selective extraction
117 procedure are discussed in the light of these results.

118 **2. Materials and methods.**

119 *2.1. Research areas*

120 The former gold mine flat is located in French Guiana (Fig. 1) on the Combat
121 Creek watershed (52°23'W, 4°35'N), a small catchment of ~1 km² covered by tropical
122 rain forest. Except for the two hilltops and for the lowland, the relief is characterized
123 by steep slopes (15-30%). The climate is tropical humid, with an annual average
124 rainfall of ~4000 mm (Barret, 2004).

125 The Combat Creek watershed is located on the 'Amina series' of the Guiana
126 Proterozoic shield consisting primarily of darkschist and thin sandstones (Milési et al.,
127 1995). Large gravel deposits from ancient rivers contain large amounts of gold,
128 resulting from the weathering of auriferous quartz veins from the Proterozoic shield
129 (Milési et al., 1995).

130 Soil distribution within the Combat Creek watershed is typical of French Guiana
131 and is related to soil position along the slopes (Boulet et al., 1979; Grimaldi et al.,
132 2001; Guedron et al., 2006). Ferralsols dominate upslope (Deckers et al., 1998),
133 having typically a high clay (< 2 µm size fraction) content and a micro-aggregated
134 structure extending over 1 m depth, which allows a good vertical water drainage
135 (Guehl, 1984). Ferruginous nodules are present throughout the entire ferralsol

136 profiles. Halfway down the slopes, ferralsols steadily evolves to Acrisols, a massive
137 alteritic horizon with a high content of fine silts at shallow depth (< 1 m). During
138 rainfalls, a perched aquifer forms on the slope above this massive alteritic horizon.
139 Following rainfalls, the aquifer is evacuated laterally, exporting preferentially clay-size
140 particles and dissolution products (Molicova et al., 1997; Grimaldi et al., 2004).
141 Downslope, the soils become hydromorphic, with dominant sands. A permanent
142 aquifer is present and slowly drained by the river, thus imposing reducing conditions.
143 Whatever the soil type, field observations have shown that quartz, kaolinite, and Fe-
144 Al-(oxyhydr)oxides dominate mineralogy, quartz being the sole remnant from the
145 primary rock-forming minerals. In the massive alteritic horizon, a few muscovite
146 crystals were also observed. Soil pH ranged from 4.2 to 4.9. The upper part of the
147 studied watershed (ferralsols and Acrisols) was considered to be pristine based on
148 the soil structure and texture.

149 In the lowland, ancient 'Long Tom' sluices and gold-bearing gravel heaps attest
150 of former gold mining activities dated from the early 1950's. These activities involved
151 the removal of fine particles with water, in order to concentrate heavier gold-rich
152 particles. Soils are thus strongly disorganized, and neither original structure nor
153 texture is preserved. Present day soil granulometry reflects the former activity as
154 gravels, sands and silts are dominant, while the clay-size fraction is almost absent.
155 Hg droplets were also observed in these gold-mined gleysols during a prospective
156 "panning" campaign. These soils can be described as disorganized gleysols
157 according to their present geochemical characteristics.

158 *2.2. Sampling collection and analysis*

159 *2.2.1 Soil collection*

160 Three toposequences were sampled in the pristine part of the watershed, (i)
161 the first in the upper part of the basin, (ii) the second on the west (mountain) side and
162 (iii) the third next to the outlet of the basin (Fig. 1). Toposequences II and III reached
163 the gold-mined flats (profiles II-0 and III-0 in Fig. 1).

164 Thirteen profiles were collected in the former gold-mined flat. SL0 to SL9
165 profiles were mesh-sampled over a restricted 250 m² area in the flat zone (Fig. 1). G5
166 profile was sampled on a gravel heap. Soil profiles were sampled systematically
167 every 10 or 20 cm, down to 1-2 m depth, using an auger and collected in sterile
168 polyethylene bags.

169 *2.2.2 Soil conditioning, granulometry and chemical analysis*

170 Granulometry was determined with standard sieving and sedimentation
171 procedures (Avery and Bascomb, 1974; Rowell, 1994). Identification of the various
172 fractions was performed according to the Udden-Wentworth classification scheme
173 (Wentworth, 1922), and “gravel”, “sand”, “silt” and “clay” thus refer to particle
174 diameters (x) of $x > 2$ mm, 2 mm $< x < 50$ μ m, 50 μ m $< x < 2$ μ m and $x < 2$ μ m,
175 respectively. Total soil organic carbon ([C]), sulfur ([S]), and nitrogen ([N])
176 concentrations were determined from the dry combustion of soil sample aliquots
177 crushed and sieved to 63 μ m, using a Fisons 1500CHNS auto-analyzer.

178 Soil samples aliquots used for Hg analysis, selective extractions and XRD
179 analysis were freeze-dried and sieved to $x < 2$ mm. This size fraction was
180 subsequently crushed to $x < 63$ μ m. Ferruginous nodules were crushed also to x
181 < 63 μ m and subsequently freeze-dried. Total Hg concentrations ([HgT]) were

182 determined by atomic absorption spectrophotometry after dry mineralization and gold
183 amalgamation with an automatic mercury analyzer (Altec, Model AMA 254). All
184 analyses were duplicated. The relative error was routinely $\pm 5\%$ and always
185 under $\pm 10\%$ (Roos-Barraclough et al., 2002). Detection limit (defined as three times
186 the standard deviation (SD) of the blank) is $0.005 \mu\text{g g}^{-1}$. Concentrations obtained for
187 repeated analyses of certified reference materials (CRMs) never exceeded the
188 published range of concentration ($0.090 \pm 0.012 \mu\text{g g}^{-1}$ and $0.091 \pm 0.008 \mu\text{g g}^{-1}$ for
189 CRMs 7002, and MESS-3, respectively). Total dissolved Hg concentration $[(\text{HgT})_{\text{D}}]$
190 were analyzed by cold vapor atomic fluorescence spectrometry (CVAFS) after
191 conversion of all mercury species to Hg^0 (Bloom and Fitzgerald, 1988), using a
192 Tekran[®] instrument (Model 2500). Additional information on the method is available
193 elsewhere (Bloom and Fitzgerald, 1988). $(\text{HgT})_{\text{D}}$ concentrations obtained for
194 repeated analyses of CRMs never exceeded the published range of concentration
195 ($12.6 \pm 1.1 \text{ ng L}^{-1}$ for ORMS-3). The detection limit is 0.05 ng L^{-1} (Bloom and
196 Fitzgerald, 1988).

197 Dissolved Fe and Al were determined by inductively coupled plasma-atomic
198 emission spectrometry (ICP-AES) using a Perkin-Elmer optima 3300DV. Calibration
199 was performed with using standard solutions for Fe and Al (Fluka[®] 44903 for Fe and
200 Fluka[®] 06155 for Al). Relative analytical errors were estimated for each sample from
201 triplicate measurements and were lower than 10%.

202 *2.3. Selective extraction procedure for Hg and major element analysis*

203 *2.3.1 Sample selection*

204 Along soil profiles, samples were selected on the basis of characteristics such
205 as granulometry, C and/or Hg contents, reflecting the textural and structural

206 variations along soil profiles. In pristine soils, the selection criteria were extreme
207 contents of C, Hg, clay-size fraction, and fine-silt fraction which increased in alterites.
208 In contaminated gleysols from the gold-mined flat, the selection relied mainly on the
209 abundance of the clay- and sand-size fractions, as well as on C and Hg
210 concentrations.

211 In addition to selected samples, selective extractions were performed for
212 quality assurance/quality controls (QA/QC) on 2 CRMs, namely a light sandy soil
213 (CRM 7002 - Czech Metrological Institute) and a marine sediment (MESS-3 -National
214 Research Council of Canada - Ottawa) and on two natural samples. The latter two
215 samples [acrisol I-1 (0-10 cm), and contaminated gleysol SL6 (30-50 cm)] were
216 chosen as models for the pristine and contaminated parts, respectively, of the
217 watershed. On all samples [HgT] was measured on both the supernatant and the
218 solid residue to assess extraction recovery. In addition, XRD data was systematically
219 collected on solid residues to identify dissolved mineral species.

220 *2.3.2 Selective extraction materials and procedure*

221 Four chemical extractions were chosen to selectively extract the main Hg
222 carrier phases reported in the literature for tropical soils: organic matter, amorphous
223 Fe oxides, and crystalline Fe/Al oxides.

224 For a given sample, selective extractions were performed on separate aliquots
225 of dry soil (~ 40 mg). All analytical procedures were conducted using ultra clean
226 sample handling to avoid laboratory contamination of low-level sample extracts and a
227 cross-contamination of high-level samples (Cossa and Gobeil, 2000). Specifically, all
228 materials in contact with the samples were previously acid washed (1 week in HNO₃
229 20%, 1 week in HCl 10%). All solutions were prepared from reagent grade chemicals
230 and supra-pure acids. Selective extractions were performed under a laminar hood in

231 8 ml Teflon[®] vessels at a 1:100 solid-to-liquid ratio (Bloom et al., 2003). At the end of
232 the extraction time, samples were centrifuged to 3000 rpm for 20 min. The
233 supernatant was analyzed for major elements. The solid residue was rinsed 3 times
234 with 4 mL MilliQ[®] water to obtain a final volume of ~16 mL, which was subsequently
235 freeze-dried before Hg analysis. The Hg content extracted by a given extraction was
236 calculated as:

$$237 \quad [\text{Hg}]_{\text{extracted}} (\text{ng g}^{-1}) = ([\text{HgT}] \times M_{\text{sample}} - M_{\text{Hg in residue}}) / M_{\text{sample}} \quad (1)$$

238 where M_{sample} is the mass of the aliquot used (in g) and $M_{\text{Hg in residue}}$ the mass of Hg
239 measured in the residue (in ng).

240 To extract Hg bond to fulvic and humic compounds, a sample aliquot was
241 digested in NH_4OH (1M) for 1h (Di Giulio and Ryan, 1987; Rodriguez Martin-
242 Doimeadios et al., 2000; Veiga and Baker, 2004). Another extraction focused on
243 organo-chelated Hg was performed by a 18h KOH digestion (1M) (Bloom et al.,
244 2003). Extraction of Hg bond to amorphous Fe(III) oxide minerals and ferrihydrite
245 was performed by adding ascorbate to a sample aliquot and allowing it to react at
246 room temperature for 24h (Kostka and Luther III, 1994). Finally, extraction of Hg
247 associated to all soil components except refractory minerals (i.e., silicates, primary
248 minerals) was determined after the digestion of soil sample aliquot in a Suprapur
249 HCl/HNO_3 solution (1/9 v/v ratio) for 10h at 70°C (Coquery et al., 1997; Bloom et al.,
250 2003). All species associated to a specific extractant are labeled as follow: $(i)_j$ where i
251 is the element and the subscribe j describes the extractant (i.e., $(i)_{\text{NH}_4\text{OH}}$ for NH_4OH ,
252 $(i)_{\text{KOH}}$ for KOH , $(i)_{\text{Asco}}$ for ascorbate and $(i)_{\text{HCl}/\text{HNO}_3}$ for HCl/HNO_3) and square brackets,
253 $[i]_j$, are used for calculated concentrations (1). The concentration of Hg related to
254 crystalline and refractory minerals (RC) is calculated as the difference between $[\text{HgT}]$
255 and ascorbate plus NH_4OH extractions:

256
$$[\text{Hg}]_{\text{RC}} (\text{ng g}^{-1}) = (([\text{HgT}] \times M_{\text{sample}}) - M_{\text{Hg Asco}} - M_{\text{Hg NH}_4\text{OH}}) / M_{\text{sample}} \quad (2)$$

257 This should be equivalent to a complete HF/HCl/HNO₃ extraction. Similar calculations
258 were made to estimate the fraction of Hg bond to crystalline Fe and Al oxides ([i]_{Cryst}).
259 For this purpose, concentrations obtained after HCl/HNO₃ extraction were considered
260 as related to amorphous and crystalline Fe and Al oxides but not to refractory
261 minerals. Thus, [i]_{Cryst} was calculated as the difference between HCl/HNO₃ and
262 ascorbate extraction :

263
$$[i]_{\text{Cryst}} (\text{mg g}^{-1}) = ([i]_{\text{HCl/HNO}_3} \times M_{\text{HCl/HNO}_3} - [i]_{\text{Asco}} \times M_{\text{Asco}}) / M_{\text{sample}} \quad (3)$$

264 A complementary pyrolysis procedure was used to assess the amount of Hg⁰ present
265 in the soils (Biester, 1994; Biester and Scholz, 1997; Bollen et al., 2008). After
266 heating a ~1g sample aliquot for 48 hours at 180°C in a porcelain crucible, the
267 content of Hg⁰ was determined from the difference in Hg concentrations between raw
268 and treated samples.

269 *2.4. Physical methods*

270 *2.4.1 X-ray diffraction*

271 X-ray diffraction (XRD) was performed on selected samples [Nodules I-2 (10-
272 20 cm), I-1 (30-40 cm) and I-1 (180-200 cm)] for routine mineralogical identification,
273 but also to identify the minerals targeted by the various selective extractions on CRM
274 7002, and MESS-3 standards, as well as on samples I-1 (0-10 cm) and SL6 (30-
275 50 cm). For this purpose, XRD patterns were collected on solid residues of the
276 different extractions, and compared to those obtained on the untreated samples.

277 XRD patterns were recorded on randomly oriented powders using a Bruker
278 D5000 powder diffractometer equipped with a Si(Li) solid-state detector from Baltic
279 Scientific Instruments and CuK α radiation ($\lambda = 1.5418 \text{ \AA}$). Intensities were recorded

280 from 5 to $80^{\circ}2\theta$, using $0.04^{\circ}2\theta$ steps, and 40 s counting time per step.
281 Measurements were duplicated on the CRM 7002 set of samples to ensure sampling
282 homogeneity and selective extraction reproducibility (data not shown).

283 *Mineralogical identification and unit-cell refinement*

284 Basic data processing operations (e.g. background removal, diffractogram
285 normalization) and mineralogical identification, based on peak positions and relative
286 intensities, were performed using the Bruker Difract^{plus} EVA software and the ICDD
287 database.

288 Relative intensities of XRD lines varied between raw and treated samples as
289 the result of specific mineral dissolutions. To overcome the intensity variations due to
290 the amount of measured sample, all XRD patterns were normalized to a selected
291 quartz peak, as quartz was present in all samples and not dissolved by the reagents
292 used. The peak used for normalization was selected so that (i) there is no peak
293 overlap with other minerals and (ii) that its intensity is statistically meaningful. For iron
294 oxides (hematite and goethite) present in Al-rich environments, experimental peak
295 positions may differ significantly from theoretical ones as the result of isomorphic Al-
296 for-Fe substitutions. In such case, theoretical unit-cell parameters of iron oxides were
297 refined using the U-Fit program (Evain, 1992) to assess their Al-contents.

298 *2.4.2 Micro X-ray fluorescence mapping and elemental analysis.*

299 SL6 (30-50 cm) sample was freeze-dried, sieved to 300 μm and used for
300 elemental mapping focused on Hg, Au, Al, Cl, Fe, Cr, Si, Ti, Mn, Ni, Ca and K. X-ray
301 fluorescence yields (XRF) were measured with an Eagle III μ -XRF spectrometer
302 (Röntgenanalytik Messtechnik GmbH) equipped with a Rh anode and a 40 μm poly-
303 capillary. XRF was measured for 300 s per point over a $1.28 \times 1.00 \text{ mm}^2$ surface area.
304 The spectrometer was operated under vacuum at 25 kV and 500 μA so as to

305 maximize the fluorescence yield at the Hg L_α edge (9.99 keV) while keeping a
306 statistically significant signal on light elements (down to Al K_α radiation - 1.49 keV).
307 Further details on the method are given elsewhere (Doering et al., 2004).

308 *2.5 Statistical treatment.*

309 Since geochemical data were not normally distributed, the following
310 parameters are reported: mean, standard error on the mean (SEM), median, 25th and
311 75th percentiles (25th perc. and 75th perc., respectively) and the number of
312 observations (N - Webster, 2001). In addition, non parametric Mann-Whitney rank
313 sum test (*U* test) and Kruskal-Wallis one way analysis of variance on ranks (*H* test)
314 were used to compare two or more than two data sets, respectively. Pearson
315 correlations were applied to compare multiple data set pairs. Correlation coefficient
316 (CC) and P values (P) are reported.

317 **3. Results**

318 *3.1 Blanks and CRMs*

319 *3.1.1 Mineralogy (QA/QC)*

320 XRD patterns obtained on treated CRM samples indicate that halite was
321 dissolved by all reagents (Supplementary data - Figs E1b. to E1e. and E2b to E2e).
322 For CRM 7002, the chlorite component was partly dissolved by both NH₄OH and
323 KOH. Exception made of halite, the effect of ascorbate was negligible. Most of the
324 minerals identified by XRD were thus only affected by HCl/HNO₃: clinocllore, calcite
325 and dolomite were totally dissolved, and an intensity decrease was observed for
326 hornblende and, to a lesser extent, for kaolinite (Supplementary data – Figs E1e. and
327 E2e.).

328 *3.1.2 Extraction recoveries (QA/QC)*

329 For NH₄OH, KOH and HCl/HNO₃ extractions, summation of M_{Hg} in both the
330 supernatant and the solid residue led to total Hg concentration consistent with those
331 of CRM 7002, MESS-3 and I-1 (10-20cm) sample. Relative error was routinely less
332 than ±10% (Tab. E1). Hg recovery was lower with ascorbate extraction, possibly due
333 to the loss of Hg⁰ in the supernatant following Hg^{II} reduction (Battke et al., 2008).
334 Consistent with this hypothesis, additional measurements of [(Hg)_D] in the
335 supernatant confirmed a steady decrease with time at the end of the extraction.

336 *3.2. Pristine soils*

337 *3.2.1 Granulometry, chemistry and mineralogy*

338 In the pristine area, the relative proportion of the clay-size fraction (< 2 µm)
339 decreased from ~90% in ferralsols to less than 30% in acrisols and gleysols (Fig. 2).
340 Ferruginous nodules made up most of the abundant gravel size fractions (x > 2 mm)
341 in ferralsols and acrisols, whereas for gleysols vegetal debris prevailed in this size
342 fraction. In the < 2 mm fraction, total C and S contents in superficial horizons
343 decreased from ferralsols and acrisols to gleysols while the C/N ratios were similar
344 for all soils, thus indicating an homogeneous mineralization of the fine organic matter
345 (OM - Bravard and Righi, 1991).

346 Minerals extracted by HCl/HNO₃ in sample I-1 (0-10cm) were identified as Al-
347 substituted iron oxides (goethite and hematite) and gibbsite (Fig. 3). Quartz, anatase,
348 kaolinite, and muscovite being essentially unaffected by the HCl/HNO₃ extraction
349 (Supplementary data – Fig. E3e.).

350 For HCl/HNO₃ and ascorbate extractions, Fe and Al concentrations were
351 correlated to the content of the clay size fraction (Tab. 1) showing that crystalline and

352 amorphous Fe and Al oxides were central in this fine fraction. Large Al and Fe
353 concentrations were measured in supernatants following KOH, ascorbate and
354 NH₄OH extractions. XRD confirmed that no crystalline Fe or Al oxides were dissolved
355 by NH₄OH, thus supporting the dissolution of amorphous (or poorly crystalline) Fe
356 and/or Al minerals. In addition, goethite was partially dissolved by ascorbate and
357 hematite by KOH, as indicated by the ~10-20% intensity decrease of their
358 characteristic XRD lines after these chemical extractions (Fig. 3). For all pristine soil
359 samples, partial dissolution of these well-crystallized species is supported by the
360 higher Al and Fe concentrations measured in KOH (median = 52.6% and 12.1% of
361 [Al]_{HCl/HNO₃} and [Fe]_{HCl/HNO₃}) and ascorbate (median = 24.7% and 21.7% of [Al]_{HCl/HNO₃}
362 and [Fe]_{HCl/HNO₃}) extracts compared to NH₄OH extracts (median=9.1% and 3.6% of
363 [Al]_{HCl/HNO₃} and [Fe]_{HCl/HNO₃}) (*H* test, *P* < 0.001).

364 [Fe]_{HCl/HNO₃}, [Al]_{HCl/HNO₃}, [Fe]_{Asco} and [Al]_{Asco} decreased steadily along the slopes from
365 ferralsols to gleysols (Fig. 4), [Fe]_{Asco} and [Al]_{Asco} being commonly lower than [Fe]_{Cryst}
366 and [Al]_{Cryst} for all profiles (Fig. 4). In ferralsols and acrisols, amorphous Fe and Al
367 oxides were most abundant in the A horizon and in the top 50 cm of the B horizon,
368 where weathering was most intense and granulometry dominated by clays. In the C
369 horizon, the relative proportion of amorphous oxides decreased while well
370 crystallized oxides and residual primary minerals became predominant, more
371 especially in the fine silt size fraction. Al-for-Fe substitutions were identified in
372 goethite (~33 Al mol%) and hematite (~10 Al mol%) from XRD unit-cell
373 determinations and confirmed by the positive correlations between [Fe]_{Cryst} and
374 [Al]_{Cryst} in B and C horizons (Fig. 4 - CC = 0.76; *P* < 0.01). Similar correlation was
375 observed between [Fe]_{Asco} and [Al]_{Asco} (CC = 0.98; *P* < 0.01) for the whole profiles of
376 pristine soils. The better correlation found between [Fe] and [Al] for ascorbate than

377 for crystalline oxide is suggestively due to an increase in [Al] related to Al extraction
378 from gibbsite and in a lesser extent from kaolinite during HCl/HNO₃ extraction.

379 3.2.2 Hg in pristine soils

380 Total mercury concentrations measured in pristine soils matrix (0.01-
381 0.49 µg.g⁻¹ - Fig. 2) were consistent with those recently reported for French Guiana
382 soils (0.09-0.50 µg.g⁻¹ - Richard et al., 2000; Grimaldi et al., 2001; Guedron et al.,
383 2006), the highest concentrations being measured in ferralsols. [HgT] decreased
384 along the slope in acrisols and in gleysols (Fig. 2).

385 [HgT] were positively correlated with [Hg] extracted by HCl/HNO₃, KOH,
386 NH₄OH and ascorbate (Tab. 1). In turn, [Hg] extracted by the four reagents were
387 positively correlated to the content of the clay-size fraction, [S] and [C]. The clay-size
388 fraction, which is composed of OM (S-groups) and Fe and Al minerals, thus
389 represents the main Hg reservoir (Tab. 1).

390 In general, [Hg] extracted by all four reagents decreased with depth and from
391 ferralsols to gleysols. However, Figure 5 shows that Hg extraction yields can be
392 separated in two sets: (i) ferralsols and superficial acrisols and (ii) deep acrisol
393 alteritic (C) horizons and gleysols. In both sets, NH₄OH extraction yields were
394 significantly larger than ascorbate ones but followed the same trend. (Hg)_{NH₄OH} and
395 (Hg)_{Asco} extraction yields were highest in superficial horizons of both ferralsols and
396 acrisols and steadily decreased with depth. Only in the upstream profiles I-1 and I-2
397 (Hg)_{NH₄OH} and (Hg)_{Asco} extraction yields were constant with depth. In gleysols and
398 deep alteritic horizons of acrisols, both (Hg)_{NH₄OH} and (Hg)_{Asco} decreased rapidly with
399 depth, whereas (Hg)_{RC} extraction yields increased strongly with increasing depth.
400 (Hg)_{RC} decreased with increasing depth in the B horizons of ferralsols and acrisols,
401 then increasing steadily in the ferralsol C horizons. (Hg)_{KOH} extraction yields were

402 highly variable with highest values in ferralsol and Acrisol B horizons. In pristine
403 gleysols (profile II-1), $(\text{Hg})_{\text{KOH}}$ extraction yields increased with increasing depth over
404 the first 50 cm, while decreasing in the C horizon.

405 *3.3 Ferruginous nodules in pristine soils*

406 *3.3.1 Chemistry and mineralogy*

407 Ferruginous nodules (> 2 mm) were abundant (mean \pm RSD = 45.8 \pm 28.6%
408 total weight), especially in superficial horizons of the pristine toposequences, their
409 relative abundance decreasing from ferralsols to gleysols. Nodules from samples I-2
410 (10-20 cm), I-2 (30-40 cm) and I-2 (180-200 cm) contained quartz, Al-substituted
411 hematite and goethite, gibbsite and kaolinite. Figure 6 shows the steady increase of
412 muscovite (Mu), paragonite (Pa), gibbsite (Gb) and rutile (Ru) contents from surface
413 (sample I-2, 10-20 cm) to depth (sample I-2, 180-200 cm - Fig E4). This
414 mineralogical evolution is consistent with chemical data as $[\text{Fe}]_{\text{HCl/HNO}_3}$ and $[\text{Al}]_{\text{HCl/HNO}_3}$
415 steadily decreased with increasing depth.

416 Position of the iron oxide peaks (goethite and hematite) shifted to lower
417 angular values with increasing depth (Fig. 6, inset), thus indicating the lengthening of
418 unit-cell parameters, and thus the probable decrease of Al-for-Fe substitution, since
419 the ionic radius of Al is smaller than Fe (Shannon, 1976). The extent of this
420 substitution was assessed for the two oxides by refinement of the unit-cell
421 parameters and empirical relations between the Al content and the lattice parameters
422 (Schulze, 1984; Stanjek and Schwertmann, 1992). Al substitutions in hematite
423 decreased from 17 mol% (surface) to 10 mol% (deep alteritic horizons) and were
424 almost constant (~20 mol%) over the entire profile for goethite. Uncertainty on lattice
425 parameters (and therefore on Al substitution quantification) was low for hematite

426 owing to the sharp and numerous diffraction peaks, but higher for goethite. Indeed,
427 few isolated diffraction peaks were available for the determination of the unit-cell
428 parameters and peaks were broadened.

429 The intensity of hematite diffraction lines increased with decreasing depth,
430 thus indicative of a higher proportion of hematite towards the surface, consistent with
431 its status of final alteration product of iron-rich phases (Schwertmann and Latham,
432 1986). Hematite and goethite are indeed the most thermodynamic stable iron oxides
433 in surface conditions. Amorphous oxides were present as minor constituents since Al
434 and Fe extracted by ascorbate and NH_4OH never exceeded 7% of their HCl/HNO_3
435 counterparts (Fig. E5). As in pristine soils matrix, KOH extraction likely induced the
436 partial dissolution of gibbsite as $[\text{Al}]_{\text{KOH}}$ was large (median = 51.5% of $[\text{Al}]_{\text{HCl}/\text{HNO}_3}$).

437 *3.3.2 Hg in ferruginous nodules of pristine soils*

438 For surface horizons, $[\text{HgT}]$ were much lower in nodules than in the soil matrix.
439 $[\text{HgT}]$ increased with increasing depth, similar values being obtained for nodules and
440 soil matrix in the deepest sample of acrisol I-1 profile. $[\text{HgT}]$ were even larger in
441 nodules than in soil matrix for ferralsol I-2 samples below 50 cm depth (Fig. E5).

442 $[\text{HgT}]$ was positively correlated with $[\text{Hg}]_{\text{HCl}/\text{HNO}_3}$ and $[\text{Hg}]_{\text{KOH}}$ ($P < 0.05$) and
443 with $[\text{Fe}]_{\text{HCl}/\text{HNO}_3}$ and $[\text{Al}]_{\text{HCl}/\text{HNO}_3}$ ($P < 0.05$). $(\text{Hg})_{\text{HCl}/\text{HNO}_3}$ extraction yields were larger
444 in ferralsol nodules from deep horizons compared to surface ones. Conversely, in
445 acrisols they were larger for surface horizons. For all nodules these yields scattered
446 about 80%. $(\text{Hg})_{\text{Asco}}$ and $(\text{Hg})_{\text{NH}_4\text{OH}}$ extraction yields rarely exceeded 10% and rapidly
447 decreased with increasing depth in both acrisols and ferralsols. $(\text{Hg})_{\text{KOH}}$ (44.8
448 $\pm 30.2\%$) extractions yields were systematically larger than the sum $(\text{Hg})_{\text{Asco}} +$
449 $(\text{Hg})_{\text{NH}_4\text{OH}}$, and progressively decreased downwards in I-1 and I-2 profiles.

450 3.4 Contaminated gold-mined soils

451 3.4.1 Granulometry, mineralogy and chemistry

452 In the gold-mined flat, contaminated gleysols contained a large proportion of
453 gravel size fractions (mainly quartz gravels and stones, mean \pm RSD = 42.7 \pm 24.6%
454 total weight) and the < 2 mm fraction was dominated by sands in contrast with
455 pristine soils (Fig. 2b). Total organic C largely exceeded the concentrations
456 determined in pristine soils and elevated C/N ratios were characteristic of poorly
457 mineralized OM (data not shown). Leaves and wood debris were visible in the litter.

458 Minerals present in sample SL6 after HCl/HNO₃ extraction were quartz,
459 kaolinite, muscovite, diaspore, rutile and microcline. Gibbsite was for the most part
460 dissolved by HCl/HNO₃, which also possibly partially dissolved kaolinite. The
461 absence of Fe oxides is consistent with the low [Fe]_{HCl/HNO₃} values (0.35 mg.g⁻¹).

462 Amorphous and crystalline Fe and Al oxides were scarce in contaminated soils
463 as shown by the low Al and Fe concentrations extracted by any reagent compared to
464 pristine soils. For all soil profiles, [Al]_{HCl/HNO₃} and [Fe]_{HCl/HNO₃} were maximum in
465 surface horizons, and sharply decreased to ~5 mg.g⁻¹ at ~40 cm depth, except for II-
466 0, SL0 and G5 profiles.

467 As for pristine soils, partial gibbsite dissolution by KOH was evidenced by
468 chemistry data as [Al]_{KOH} (median = 76.3% of [Al]_{HCl/HNO₃}) were larger than [Al]_{Asco}
469 (median = 55.7% of [Al]_{HCl/HNO₃}) and [Al]_{NH₄OH} (median = 28.5% of [Al]_{HCl/HNO₃}) (*U* test,
470 *P* < 0.05). Fe oxides were also more attacked by ascorbate ([Fe]_{Asco} median = 31.1%
471 of [Fe]_{HCl/HNO₃}) than by KOH and NH₄OH ([Fe]_{KOH} : median = 17.3% of [Fe]_{HCl/HNO₃}
472 and [Fe]_{NH₄OH} : median = 14.8% of [Fe]_{HCl/HNO₃}).

473 3.4.2 Hg in contaminated gold mined soils

474 [HgT] values measured in gleysols from the gold-mined flat (up to $\sim 10 \mu\text{g}\cdot\text{g}^{-1}$ in
475 SL6 profile) were up to 5 times higher than those determined in gleysols from the
476 pristine area (Fig. 2b). Although extremely variable, even over very short distances,
477 these values were consistent with those determined in Hg-contaminated soils from
478 different gold-mining regions (Lechler et al., 1997; Lechler et al., 2000; Richard et al.,
479 2000; Sladek and Gustin, 2003), and positively correlated with [Hg] extracted by all
480 four reagents (Tab. 2a and 2b). Contrarily to the pristine area, where gleysols were
481 the pole of lowest [HgT], the only significant correlation of [HgT] with other soil
482 parameters was observed with the sand-size fraction ($P < 0.05$).

483 μ -XRF map (Fig. 7) shows that Hg was not homogeneously distributed in
484 sample SL6 (30-50cm). Indeed, Hg is essentially present as “hot-spots” of 30 to
485 100 μm in diameter. In these hot-spots, Hg concentration is not correlated with any
486 element (data not shown), except for a covariation with Au (Fig. 7). These spots are
487 thus likely Hg^0 droplets, possibly amalgamated with Au. Consistently, pyrolysis
488 measurements showed that almost 60% of total Hg was desorbed at 180°C .
489 However, lower percentages of desorbed Hg were determined for SL8 and SL7
490 samples, thus suggesting that Hg^0 was not systematically the main Hg form over the
491 entire former gold-mined flat (Fig. 8). Consistently, selective extractions have shown
492 that Hg was partially associated to OM and amorphous oxides for some samples as
493 shown by the NH_4OH and ascorbate extraction yields. Such associations remained
494 however minor as $(\text{Hg})_{\text{NH}_4\text{OH}}$ (median = 28.4%) and $(\text{Hg})_{\text{Asco}}$ (median = 14.7%)
495 extraction yields only seldom exceeded 50 and 30%, respectively (Fig. 8). Relative
496 proportion of Hg associated to refractory minerals was negligible as $(\text{Hg})_{\text{HCl/HNO}_3}$
497 extraction yields were systematically $\sim 100\%$. In this case, large $(\text{Hg})_{\text{RC}}$ reflect the

498 proportion related to Hg^0 (Fig.8). KOH extraction yields also were high (median =
499 86.5%), being always higher than 80% in the uppermost 40 cm.

500 **4. Discussion**

501 *4.1 Selective extractions*

502 Further investigation was performed to unravel the reactions leading to Hg
503 reduction and loss during ascorbate extraction, although $[\text{Hg}]_{\text{Asco}}$ were not biased,
504 being calculated from the analysis of the solid residue. Results of the pyrolysis
505 procedure were also questionable as the pyrolysis (180°C) of marine sediment
506 MESS-3 has shown no significant desorption of Hg, while almost 70% of total Hg
507 desorbed from light sandy soil CRM 7002 under the same conditions. It is thus likely
508 that in addition to Hg^0 , Hg bond to OM was partially desorbed during pyrolysis.
509 Pyrolysis temperature (180°C) was indeed close to the temperature range reported
510 by Biester and Scholz (1997) for the thermal release of Hg bound to organic
511 compounds (200-350°C). This is especially relevant for the assessment of the
512 different Hg-compartments in organic-rich contaminated soils.

513 Cross recoveries between selective extractions were observed for both CRMs
514 and natural samples, but it was also possible to attribute selective mineral extractions
515 to each reagent. The selective character of a given extraction depends on the
516 considered soil matrix, as reagent selectivity depends both on the nature of
517 components present and on their reactivity with respect to a given reagent.
518 Overlapping extractions further increase the uncertainty on the relative proportion of
519 Hg associated to any carrier phase. Although chemical data may help to assess
520 cross recoveries, the restricted and variable selectivity of the different reagents
521 precludes the use of selective extractions as standardized procedures for

522 environmental studies. Specifically, the use of selective extractions needs preliminary
523 investigation to assess reagent selectivity for a given soil matrix.

524 *4.2 Hg distribution in pristine soils*

525 The study of Hg distribution in a soil has to take into account the original soil
526 cover formation, its current evolution and its present topographic position, that is its
527 position in the soil association (Fritsch et al., 2006). These factors are indeed
528 responsible, at least partially, for the relative abundance and the reactivity of soil
529 organic and mineral components, including Hg carriers, thus providing additional
530 insight in the Hg natural equilibrium and contents.

531 *4.2.1 Hg carrier phases in soil profiles*

532 A positive correlation between the relative amount of the clay-size fraction and
533 C, S, [Fe] and [Al] extracted by HCl/HNO₃, NH₄OH and ascorbate is shown in Table
534 2. Hg is thus likely associated with finely divided Fe and Al organic complexes and/or
535 with organic coatings on Fe/Al oxides. Several authors have already shown that in
536 Amazonian micro-aggregated horizons, micropeds develop from the bonding
537 between clays, organic matter and iron oxides through the formation of carbon-
538 coatings on mineral surfaces (Tandy et al., 1990; Malengreau and Sposito, 1997; Do
539 Nascimento et al., 2004).

540 Specific bonding of Hg with sulfur groups from finely divided organic matter
541 may be inferred from the above described correlations for all pristine soils. This is
542 consistent with previous reports of organics, and more especially of reduced sulfur
543 groups, being the most effective sorbents for Hg in acidic environments (pH < ~4.5-
544 5.0), compared to iron oxides and clay minerals (Andersson, 1979; Schuster, 1991;
545 Roulet and Lucotte, 1995; Skyllberg et al., 2000; Skyllberg et al., 2006; Manceau and

546 Nagy, 2008).

547 In addition to this essential organic contribution, Hg contents in soils from
548 French Guiana are also controlled by the actual nature of Fe and Al (oxyhydr)oxides
549 (Roulet and Lucotte, 1995; Roulet et al., 1998). In pristine soil matrix, large $[Hg]_{Asco}$
550 and $[Hg]_{HCl/HNO_3}$ values were indeed attributed to Al-substituted amorphous and
551 crystalline Fe oxides. As a result of their larger surface areas compared to their
552 crystalline counterparts, amorphous or poorly crystalline species have an increased
553 adsorption capacity, (Schwertmann and Latham, 1986; Schwertmann and Cornell,
554 2000). The positive correlation between $[Hg]_{KOH}$ and $[Al]_{KOH}$ supports also the
555 importance of Al oxides as a reservoir for Hg. This reservoir is accessible through
556 KOH and, to a lesser extent, through ascorbate extractions, thus the large $[Al]_{Asco}$
557 plead for the partial dissolution of (amorphous) Al (oxyhydr)oxides (Schwertmann
558 and Latham, 1986).

559 In nodules, as in soil matrix, Hg is bond to Al and Fe oxides. Ferruginous
560 nodules are abundant in ferralsols and acrisols, where they represent ~50% of total
561 soil materials and thus represent large Hg reservoirs, $[HgT]$ being similar in nodules
562 and soil matrix. In addition to Al-substituted hematite and goethite, gibbsite
563 contributes significantly to Hg retention in soil nodules, especially in surface horizons,
564 as suggested by the large $(Hg)_{KOH}$ extraction yields associated with large $[Al]_{KOH}$
565 values. This observation is consistent with the strong affinity of Hg(II) for gibbsite
566 (Kim et al., 2004a; Kim et al., 2004b; Weerasooriya et al., 2007).

567 Finally, the contribution of refractory minerals (identified as muscovite,
568 anatase, rutile and kaolinite) is non negligible, although minor, in pristine soils as the
569 median $(Hg)_{HCl/HNO_3}$ extraction yield is only 89.8%.

570 4.2.2 Influence of soil structure on Hg distribution in soil profiles

571 From the above description, it appears that the vertical distribution of OM,
572 Al/Fe (oxyhydr)oxides and ferruginous nodules in the soil matrix rules Hg distribution.
573 In the following section, pedological and chemical characteristics responsible for the
574 spatial distribution of these components in soil profiles will be discussed.

575 Upslope, large [HgT] have been measured in the matrix of the ferralsol micro-
576 aggregated horizons (i.e., A and B horizons). The micro-aggregated structure confers
577 a good water drainage capacity to ferralsols, favoring water flow and oxidizing
578 conditions. As a result, Hg is spread throughout these micro-aggregated horizons as
579 the result of both significant leaching and migration of atmospheric inputs (Guedron
580 et al., 2006). Plant litter is rapidly mineralized in surface horizons, and the rapid
581 turnover of humic substances leads to the migration of fulvic acids in deeper organo-
582 mineral horizons where they are complexed to Fe and Al oxides (Roulet and Lucotte,
583 1995). Thus, Hg originating from plant litter decomposition and/or from atmospheric
584 fallouts can migrate into the organo-mineral horizons as organic-Hg complexes and
585 progressively adsorb onto Al/Fe (oxyhydr)oxides. Soil ecosystem engineers (soil
586 invertebrates and roots) also play an important role in the particulate transfer of Hg
587 by enhancing aggregation (Velasquez et al., 2007) and soil hydraulic conductivity
588 (Grimaldi et al., 2008) through biogenic soil organization. In the micro-aggregated B
589 horizon of ferralsols, the distribution of amorphous Fe oxides is positively correlated
590 to [HgT], although their relative abundance is lower than or similar to that of the
591 crystalline varieties. The comparison of $[\text{Fe}]_{\text{Cryst}}$ and $[\text{Fe}]_{\text{Asco}}$ indicates that the relative
592 proportion of crystalline Fe oxides is increased further in the superficial layer and in
593 the deep alteritic horizons compared to the micro-aggregated B horizon of ferralsols.
594 In addition, the positive correlations between $[\text{Hg}]_{\text{RC}}$ and $[\text{Fe}]_{\text{Cryst}}$ or $[\text{Al}]_{\text{Cryst}}$

595 throughout the entire soil profile suggest that $[Hg]_{RC}$ is partially associated to Al-
596 substituted crystalline Fe oxides. In this case, the sum of $[Hg]$ adsorbed onto
597 amorphous and crystalline Fe oxides would be comparable or larger than the part
598 related to organic compounds.

599 The micro-aggregated structure of ferralsols disappears progressively with
600 increasing depth, giving rise to deep mineral horizons with a continuous structure
601 composed of large mineral relics. Parental material relics “dilute” Al/Fe oxides and
602 thus reduce the overall sorption capacity of deep horizons. Another essential factor is
603 the lower permeability of deep horizons which limits the downward migration of Hg
604 (Guedron et al., 2006). Hg content thus mainly results from rock weathering and
605 pedogenesis (Guedron et al., 2006), $[HgT]$ being mainly associated with coarse-
606 grained goethite and hematite.

607 In ferralsol nodules, Hg is mainly bond to Al-substituted hematite and goethite,
608 and to gibbsite. As in ferralsol matrix, highest $[HgT]$ values were measured in the
609 micro-aggregated horizon. Indeed, the downward decrease of $(Hg)_{HCl/HNO_3}$ extraction
610 yields and the mineralogical evolution suggest that A and B horizons nodules are
611 primarily composed of neo-formed minerals, thus indicating an advanced stage of
612 pedogenesis and weathering in superficial horizons. Mineral neof ormation also
613 appears to favor the incorporation of atmospheric Hg inputs.

614 Halfway down the slope, in acrisols, a similar mineralogical and structural
615 contrast is observed between superficial (A and B horizons) and alteritic horizons.
616 This transition is sharp as evidenced by the increase of the silt fraction and by the
617 decrease of $[HgT]$. As for ferralsol, the increase of refractory minerals with increasing
618 depth is most likely responsible for the observed decrease of $[HgT]$.

619 Downslope, in gleysols, [HgT] were lower since fine organic fraction, S and
620 amorphous oxides are less abundant than upslope, even though carrier phases were
621 similar to those identified in ferralsols and Acrisols. Gleysols are depleted in iron
622 oxides and the main minerals are kaolinite, gibbsite, muscovite and quartz. Hg
623 extracted by NH₄OH never exceeds 40% of [HgT] in gleysol surface horizons and
624 rapidly decreases with depth. Hg bond to amorphous oxides never exceeds 20% of
625 [HgT], and almost 50% of [HgT] is likely associated to Al-substituted crystalline Fe
626 oxides, gibbsite and to refractory minerals. Consistently, Hg, Fe, and Al extracted by
627 HCl/HNO₃ are together positively correlated in the gleysol profile. In depth, Hg is less
628 abundant and mainly associated to refractory minerals since (Hg)_{RC} extraction yields
629 increase with increasing depth.

630 *4.2.3 Influence of soil formation on the distribution of Hg over the soil association*

631 The soil association results from the ferralsols current imbalance, which leads
632 to temporal and spatial evolutions (Boulet et al., 1993; Grimaldi et al., 2004).
633 Because of their major role as Hg reservoirs, the following description of Hg
634 distribution along the toposequence will thus focus on OM and Fe oxides behavior in
635 the soil association.

636 In organic horizons, the [HgT] decrease along the slopes correlates with the
637 decrease of their C and S contents. However, organic horizons in gleysols are
638 characterized by an increased proportion of OM in gravel size fractions (> 2 mm) and
639 thus by a lower degree of mineralization of coarse OM because of hydromorphy.
640 Consequently, the overall reactivity of coarse OM towards Hg (including the reactivity
641 of S groups) decreases when all soil fractions are considered even if C and S remain
642 abundant (Andersson, 1979; Kerndorff and Schnitzer, 1980; Schnitzer and Kerndorff,
643 1981; Ochs et al., 1994; Gasper et al., 2007).

644 The decreasing proportion of Fe and Al (oxyhydr)oxides down the slope
645 results from the combined effect of diffuse erosion and leaching due to the lateral
646 drainage of horizons overlying the alteritic horizons of Acrisols and of the progressive
647 occurrence of hydromorphy (Boulet et al., 1979; Fritsch et al., 1986; Fritsch et al.,
648 2006; Grimaldi et al., 2004; Lucas et al., 1996). Drainage induces a texture
649 coarsening from the original clayey texture of Ferralsols to the sandy-loam texture of
650 Gleysols, whereas hydromorphy induces the dissolution of crystalline and amorphous
651 Al-bearing Fe oxides and the export of the reduction products through the
652 hydrographic network (Grimaldi et al., 2008). Both processes result in the relative
653 increase of coarse fractions composed mainly of quartz and neoformed kaolinite
654 (Boulet, 1978; Roulet et al., 1998) which possess a low affinity for Hg (Sarkar et al.,
655 1999; Sarkar et al., 2000).

656 *4.3 Influence of former gold mining activities on anthropogenic Hg pool in soils*

657 In contrast to pristine soils, Hg appears to be associated mainly with the
658 prevailing sandy fraction. In highly contaminated sites Hg has been identified as Hg⁰,
659 insoluble cinnabar and metacinnabar (Lechler et al., 1997; Slowey et al., 2005;
660 Slowey and Brown, 2007). In the present study, mercury sulfides (cinnabar and
661 metacinnabar) are most likely insignificant as no positive correlation was found
662 between Hg and S. (Hg)_{HCl/HNO₃} extraction yields are systematically close to 100%
663 and support further the low amount of mercury sulfides which are essentially
664 insoluble in concentrated HCl/HNO₃ (Mikac et al., 2002; Mikac et al., 2003)..

665 The probable presence of Hg⁰ is supported by the observation of macroscopic
666 Hg⁰ droplets during a prospective “panning” in the sandy horizon of gold-mined
667 Gleysols (50-100 cm depth) next to profile III-0. It was however impossible to assess

668 the distribution of the Hg^0 droplets along vertical profiles. μXRF confirmed the
669 presence of isolated Hg^0 droplets ($\sim 100 \mu\text{m}$ in diameter) in sample SL6 (30-50cm),
670 possibly amalgamated with micrometric gold particles. Consistently, pyrolysis
671 measurements have shown that Hg^0 generally prevail in the former gold-mined flat
672 soils, with some exceptions (e.g., SL7 and SL8 profiles). In the latter profiles, the
673 reduced contribution of Hg^0 can result from its oxidation, known to be fast in tropical
674 environments (Dominique et al., 2007). If geochemical conditions are not
675 permanently reducing, as in the surface horizons, Hg^0 can be oxidized, allowing
676 adsorption onto OM and minerals. The relative proportion of Hg bond to OM is
677 however lower than in pristine soils, $(\text{Hg})_{\text{NH}_4\text{OH}}$ extraction yields rarely exceeding
678 50%. As previously discussed, the overall limited mineralization of OM is most likely
679 responsible for the low reactivity of OM for Hg. Similarly, the relative proportion of Hg
680 bond to amorphous Fe oxides is low (up to 15%), these oxides being scarce in
681 hydromorphic environments. Finally, the part related to refractory minerals (i.e.,
682 quartz, kaolinite and muscovite) appears to be negligible since $(\text{Hg})_{\text{HCl/HNO}_3}$ extraction
683 yields are always close to 100%.

684 **5. Conclusion**

685 The combination of selective extractions with physical techniques (XRD and
686 μXRF) significantly improved the determination of Hg speciation by allowing the
687 identification of Hg carrier phases targeted by the different extractants. However, the
688 quantification of Hg associated to a given carrier phase is affected by intrinsic bias,
689 especially because of probable overlaps between the different extraction procedures.
690 Thus, selective extractions can not be used as a standard procedure for natural

691 samples since the use of selective extractions requires the preliminary assessment of
692 reagent selectivity for a given soil matrix.

693 By coupling XRD and selective extractions, the main Hg carrier phases in
694 pristine soils were identified as sulfur groups from organic matter, amorphous and
695 crystalline Al-substituted Fe oxides and, to a lower extent, Al oxides. The relative
696 contribution of these carriers to the overall Hg concentration was dependent on their
697 relative abundance which is controlled by the pedogenetic evolution along the slopes,
698 and more especially by redox conditions, hydric properties, and ecosystem
699 engineers. In the former gold-mined flat, Hg was found mainly as Hg⁰, possibly
700 amalgamated with Au, and as oxidized Hg^{II} associated with organic matter, Fe and/or
701 Al oxides in surface horizons.

702 *5.1 Concluding Remarks*

703 These findings have major implications on the contamination of hydrosystems
704 in tropical regions, where soil erosion due to deforestation, gold mining and
705 agricultural processes are intense and lead to the release of Hg-rich clay-size
706 particles as shown by the low proportion of clay-size fractions in gold-mined gleysols.
707 When entering suboxic and anoxic environments (e.g. river or dam sediments), Hg
708 bond to clay-size particles indeed becomes accessible to ferri- and sulfato-reducing
709 bacteria that are the main factors of Hg methylation (Fitzgerald and Lamborg, 2003;
710 Fleming et al., 2006). Hg bound to OM and Fe oxides is specifically concerned as
711 these major components of the clay-size fraction are essential nutrients for
712 methylating micro-organisms and final electron acceptors of their metabolism. In
713 addition, former gold mined soils are potential important sources of Hg for
714 methylating bacteria. Indeed, Dominique et al. (2007) have shown that anthropogenic

715 Hg⁰ from former gold-mining activities also promotes the production of MMHg in the
716 Amazonian hydrosystems and its availability to trophic webs.

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Table 1: Pearson correlation matrix (correlation coefficient (CC), P value (P) and number of samples (N)) of pristine soils extracted Hg, Hg extraction yields, Fe, Al, Clay size fraction, Fine silts, C and S for total and (a) HCl/HNO₃ (labeled HCl) and ascorbate (labeled Asco) selective extractions, and (b) KOH and NH₄OH selective extractions.

Table 2: Pearson correlation matrix (correlation coefficient (CC), P value (P) and number of samples (N)) of contaminated soils extracted Hg, Hg extraction yields, Fe, Al, Clay size fraction, Fine silts, C and S for total and (a) HCl/HNO₃ (labeled HCl) and ascorbate (labeled Asco) selective extractions, and (b) KOH and NH₄OH selective extractions.

Figure captions

Fig. 1: Top left: french Guiana map with the studied site location (closed circle). Right and bottom: detailed map of the studied site, including general watershed scheme with soil sampling locations, and associated legend.

Fig. 2: Evolutions with depth of total Hg, C and C/N ratio for (a) pristine and (b) contaminated soils. Evolutions with depth of the clay- and sand-size fractions are reported for pristine and contaminated soils, respectively.

Fig. 3: XRD patterns normalized on quartz peak (refer to the text for details) for raw samples, and following ascorbate, KOH, NH₄OH and HCl/HNO₃ extractions.

Fig. 4: Top : Evolutions with depth of Fe and Al concentrations extracted by ascorbate and corresponding Al/Fe ratio. Bottom: same graphs as top, focused on calculated crystalline + residual fraction in pristine soils.

Fig. 5: Evolutions with depth of (Hg)_{HCl/HNO₃}, (Hg)_{CR}, (Hg)_{Asco}, (Hg)_{NH₄OH}, and (Hg)_{KOH} extraction yield (%) in pristine soils. Triangles, circles and squares refer to toposequence I, II and III, and white, grey and black filled colours refer to ferralsols, acrisols and gleysols, respectively.

Fig. 6: XRD patterns of nodules from pristine soils I-2 (10-20cm), I-2 (30-40cm), and I-2 (180-200cm).

Fig. 7: μ XRF mapping of Si, K, Fe, Au and Hg of SL6 (30-50 cm) sample and counts measured on L α edge fluorescence of Hg and Au versus K α radiation of Si, K and Fe, and comparison on counts measured for Hg (Y axis) versus Au (X axis).

Fig. 8: Evolution with depth of (Hg)_{HCl/HNO₃}, (Hg)_{CR}, (Hg)_{Asco}, (Hg)_{NH₄OH}, (Hg)_{KOH} and (Hg⁰) pyrolysis extraction yields (%) in contaminated soils.

Fig. 1.

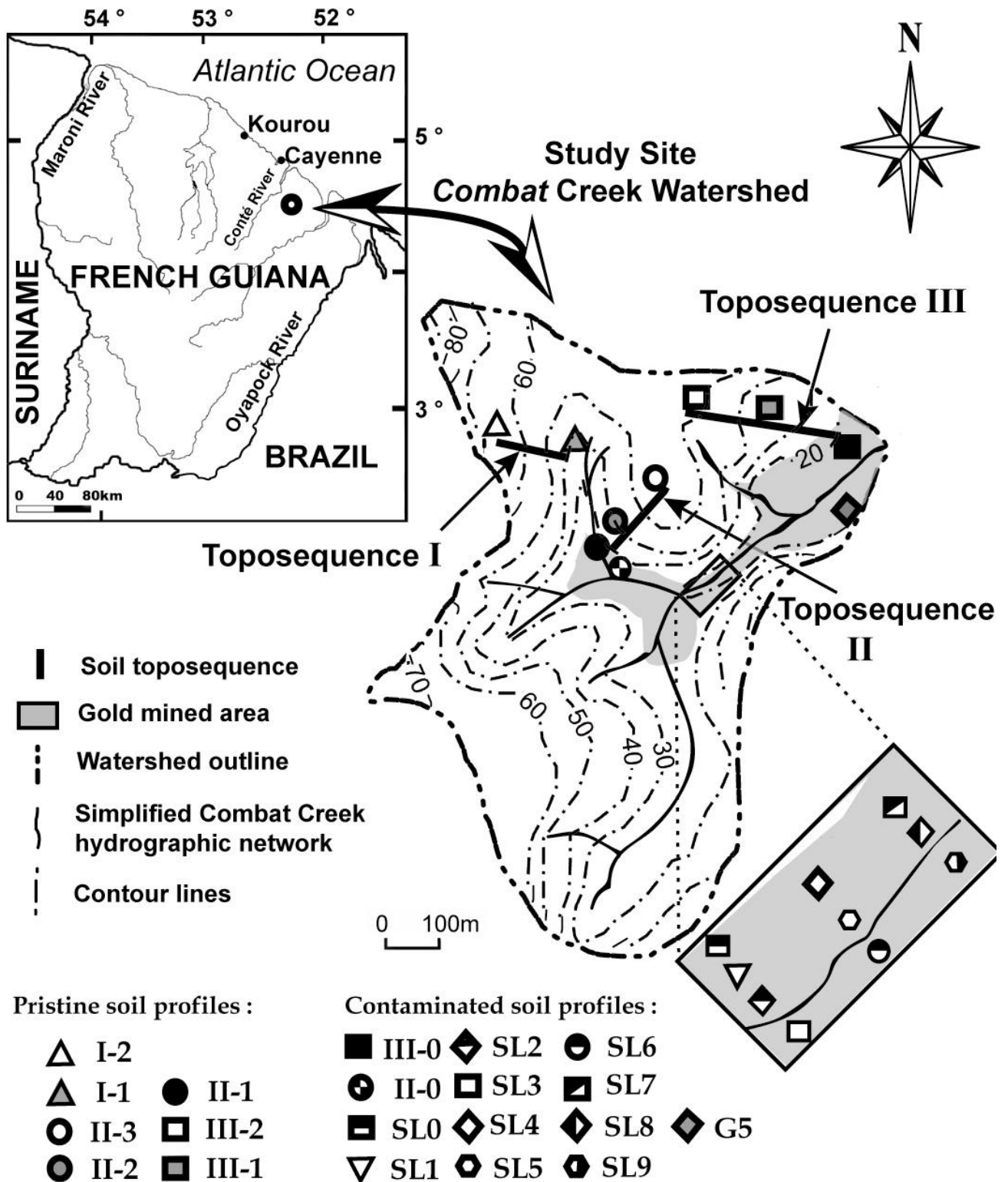


Fig. 2.a.

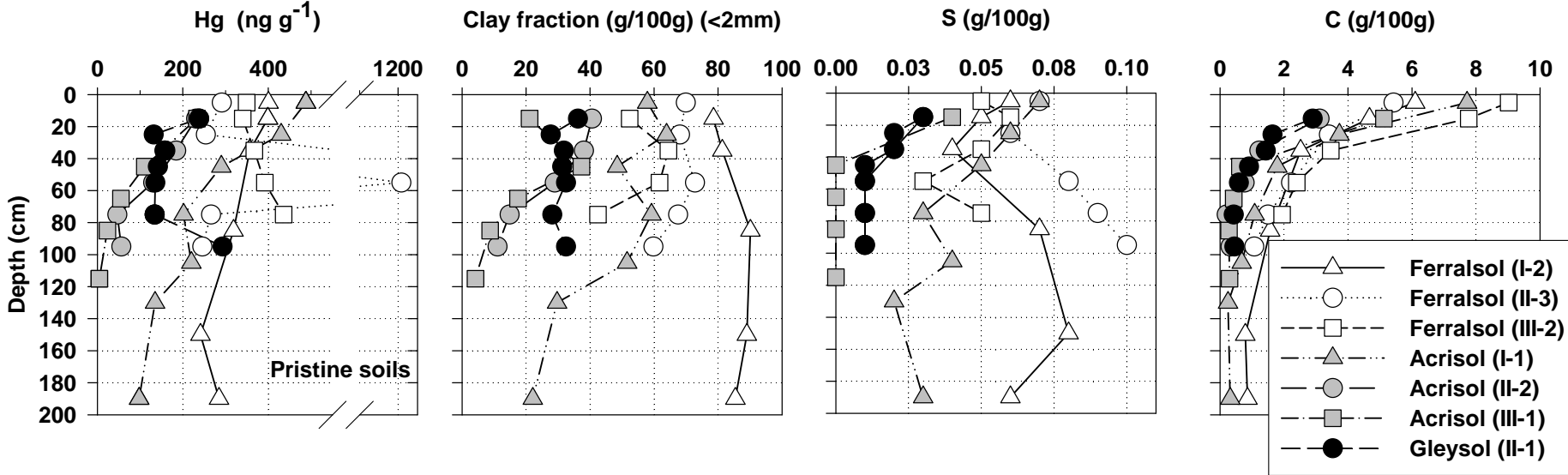


Fig. 2.b.

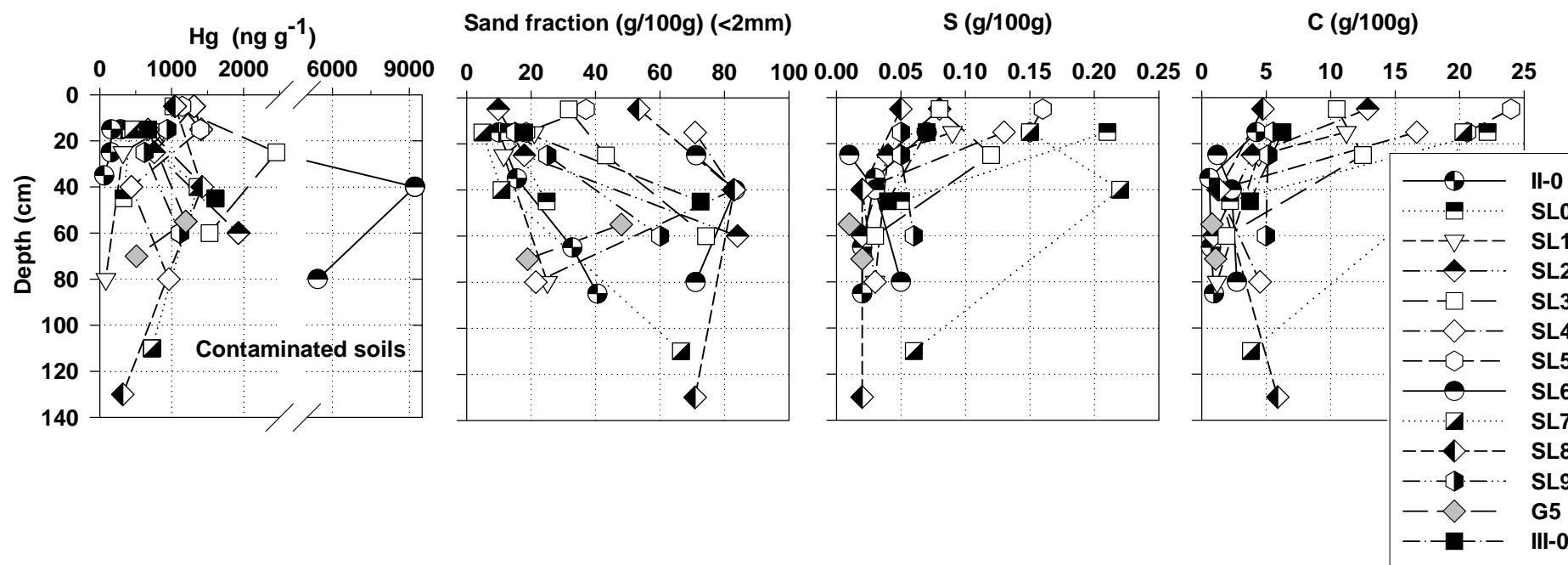


Fig. 3.

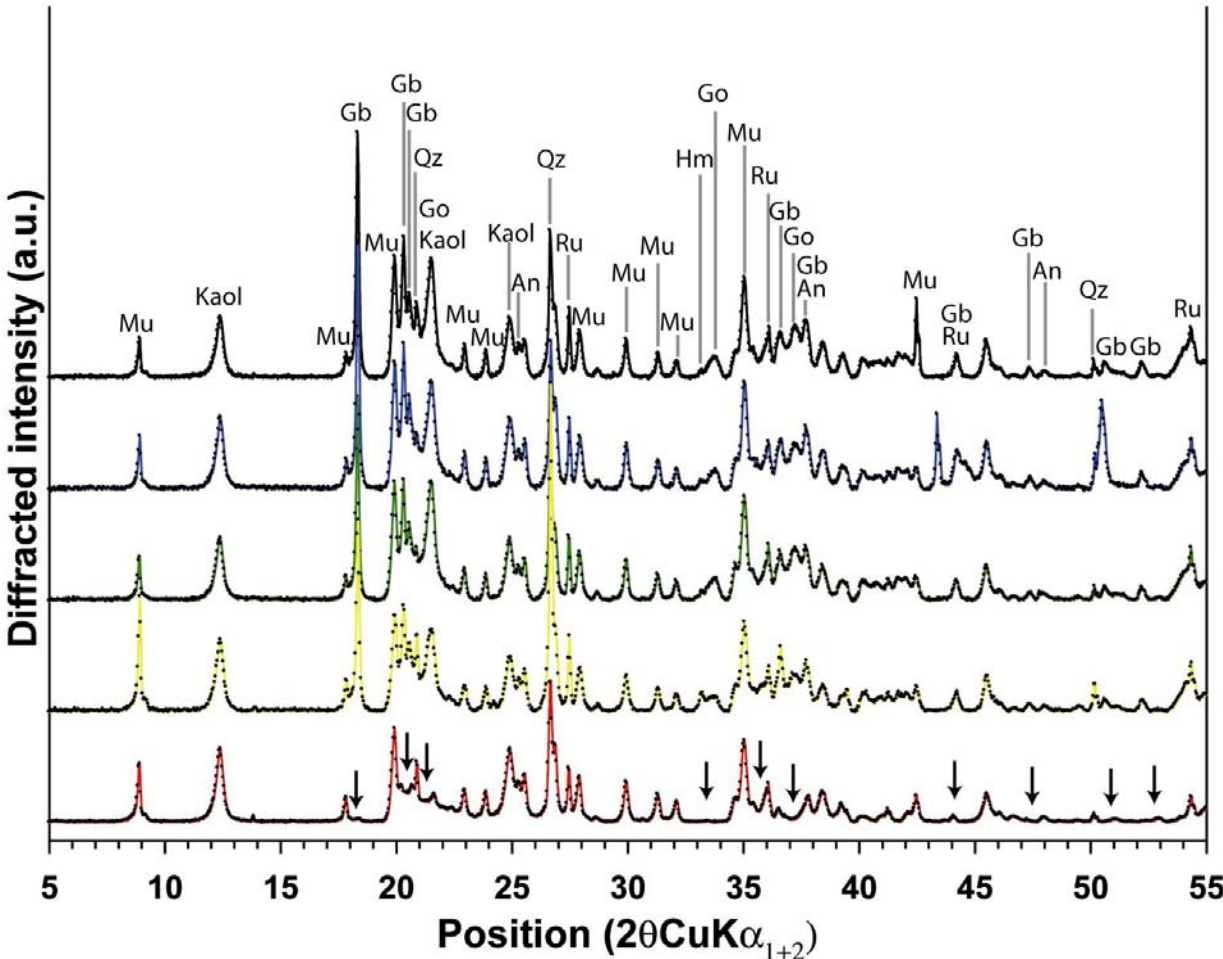


Fig. 4.

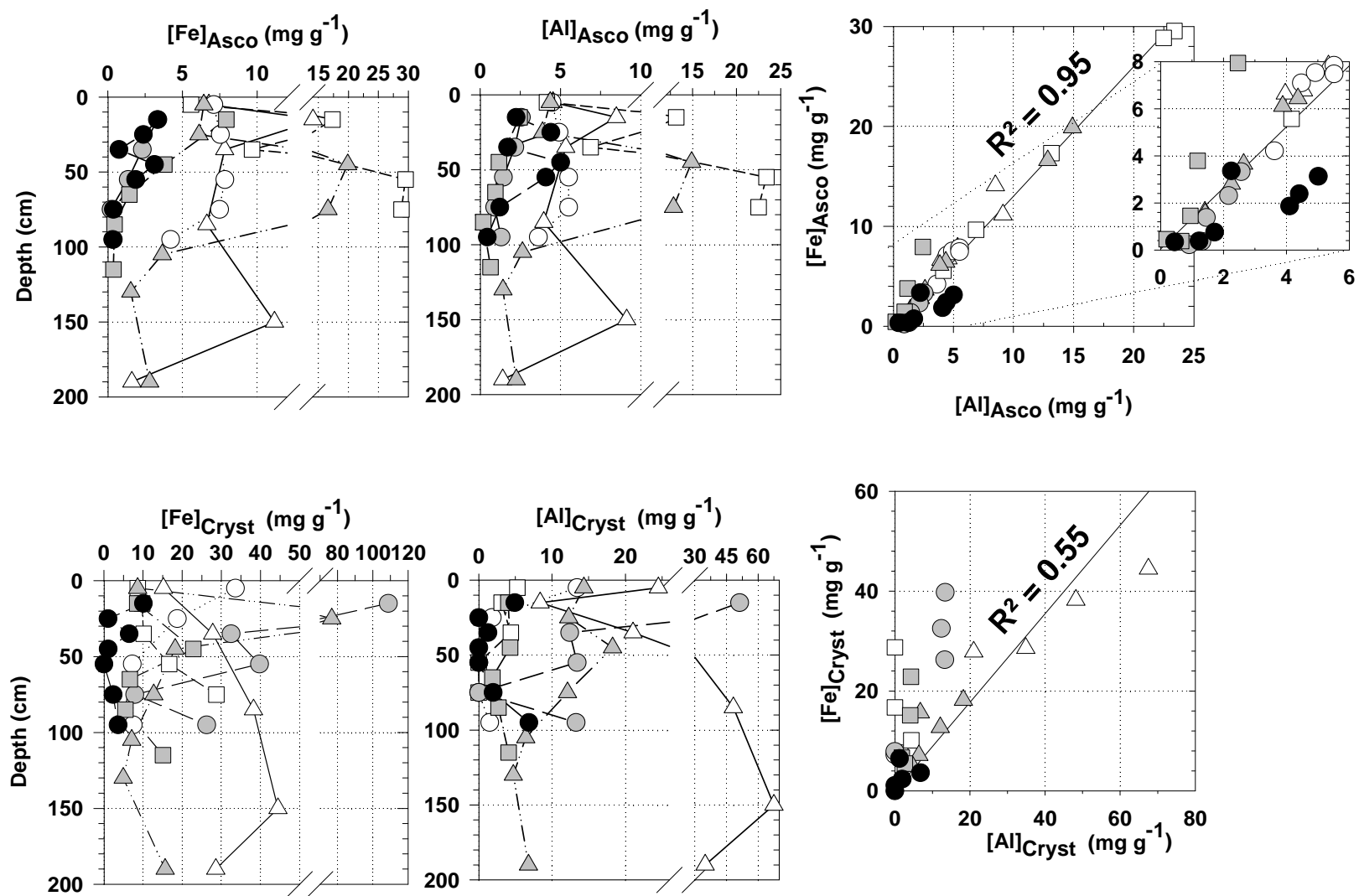


Fig. 5.

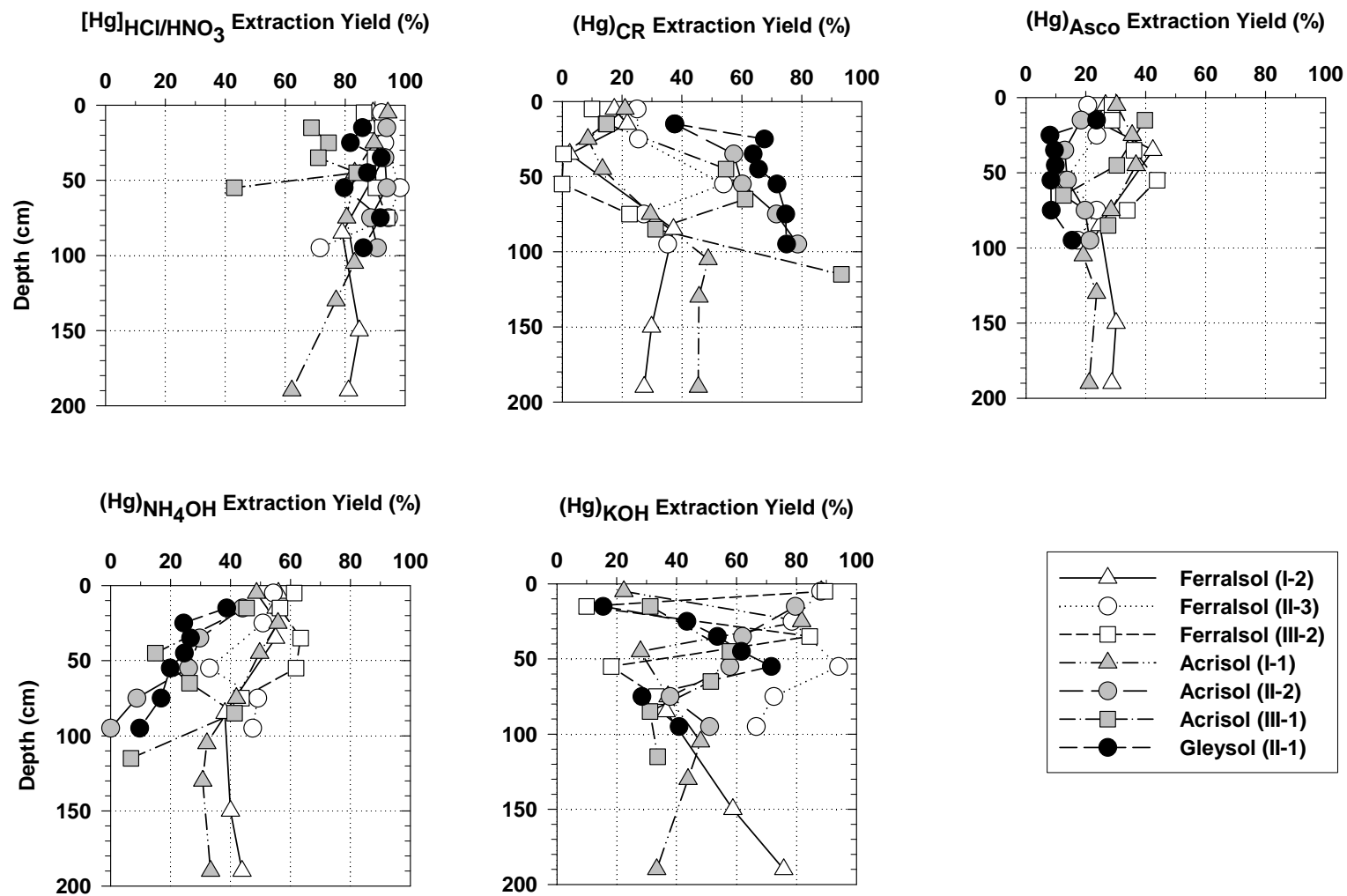


Fig. 6.

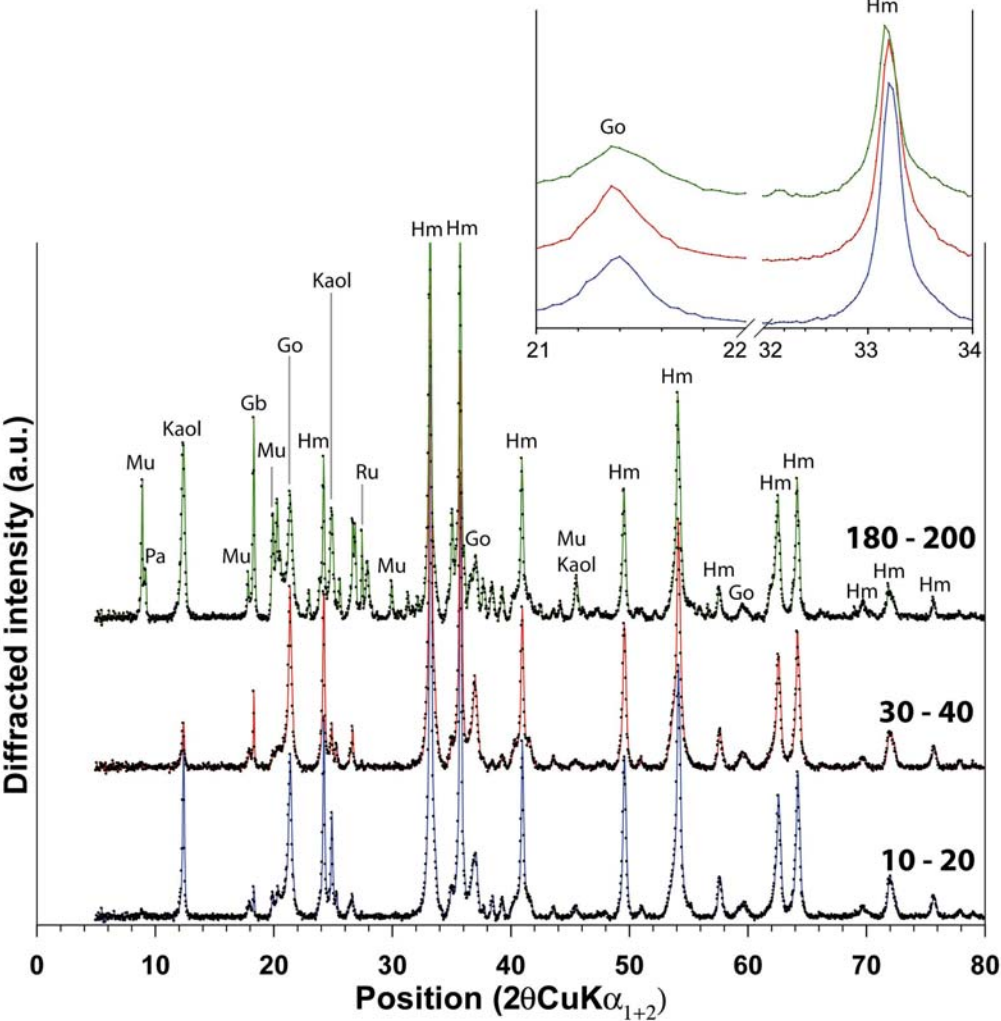


Fig . 7.

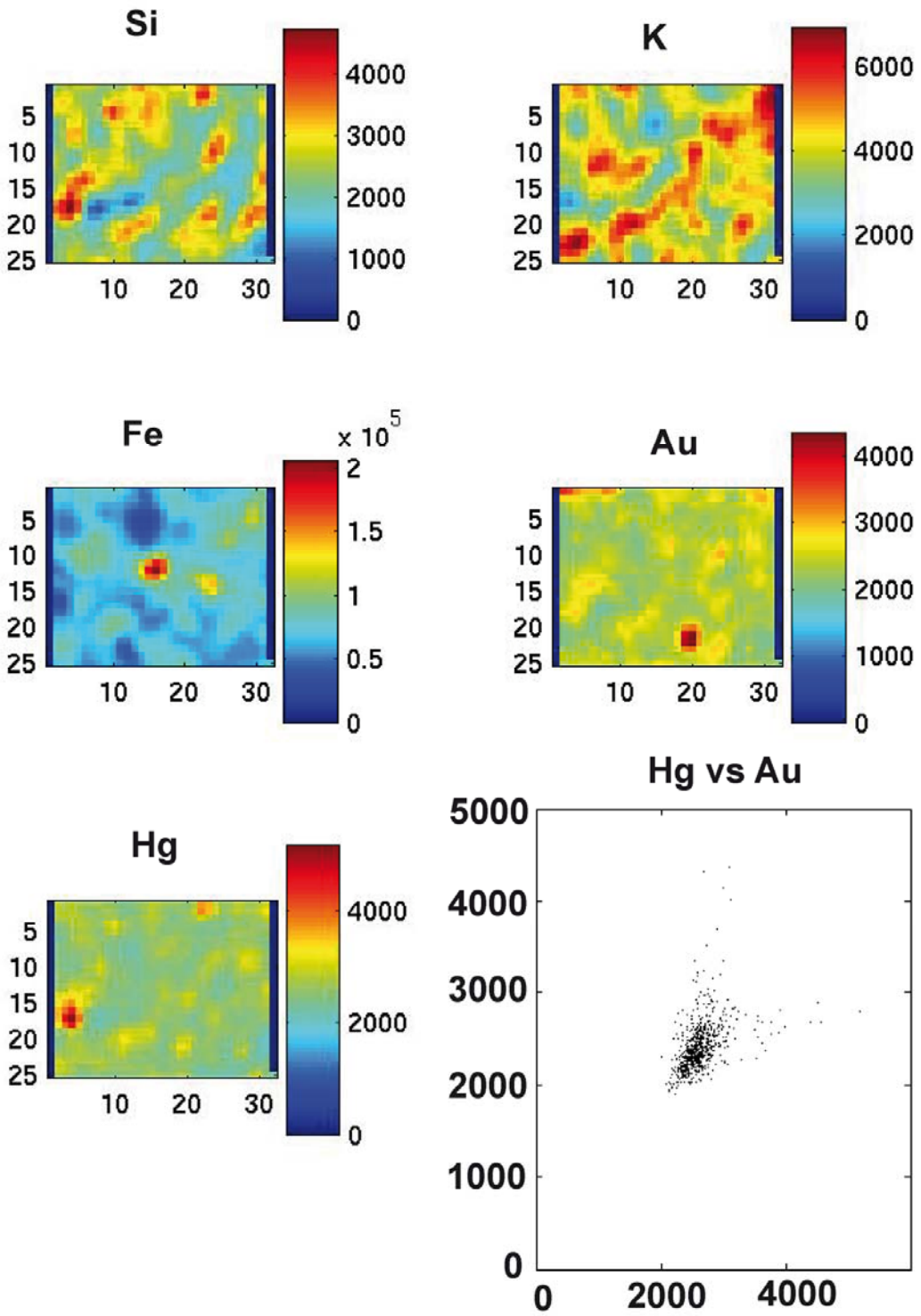
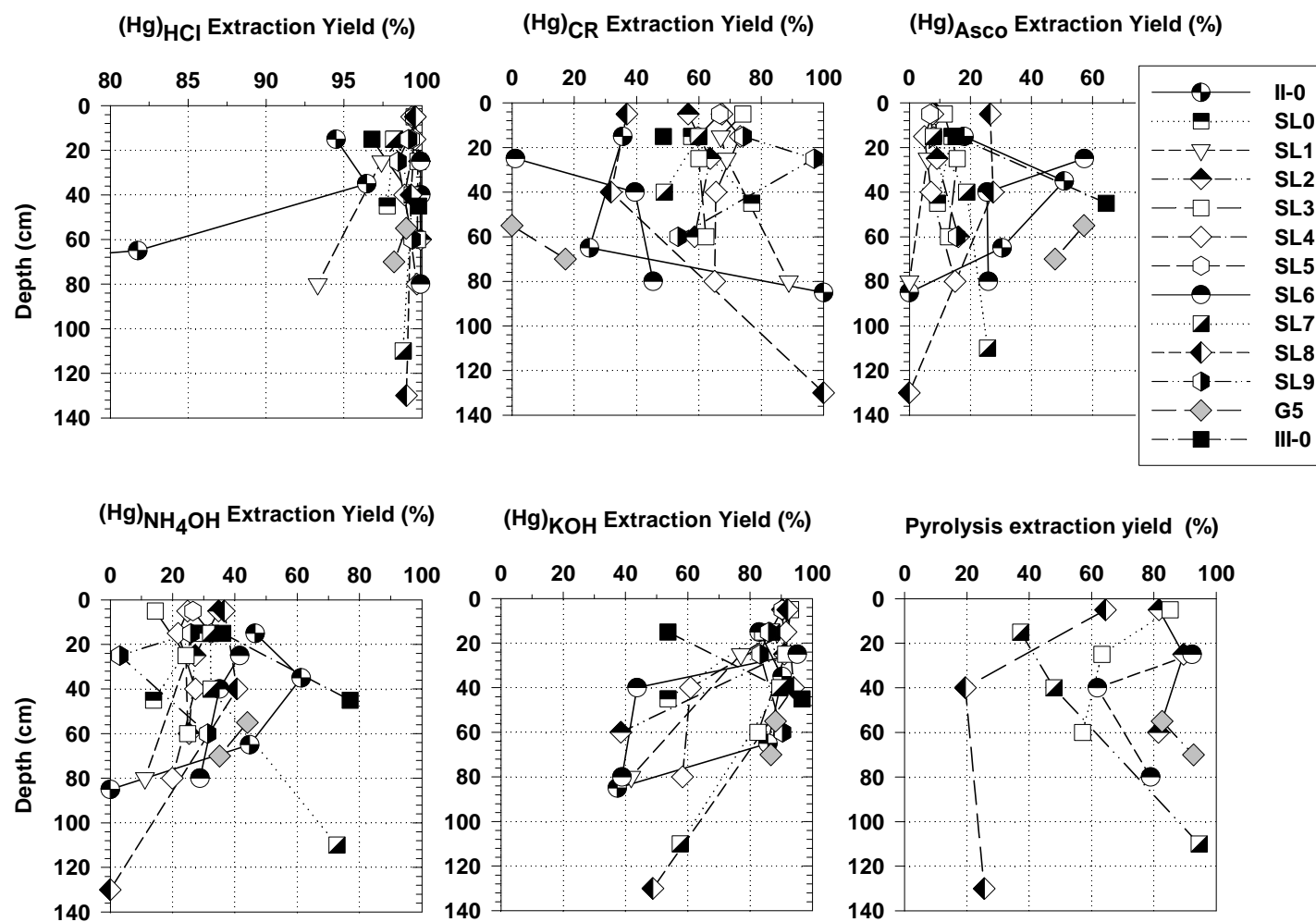


Fig . 8.



Supplementary data 2

(Electronic table)

Table E1: Percent recoveries of Hg in supernatant and solid residue for Certified materials CRM 7002, MESS-3 and sample I-1 (0-10 cm) for ascorbate, NH₄OH , KOH and HCl/HNO₃ selective extractions.

Sample	Ascorbate			NH ₄ OH		KOH			HCl/HNO ₃		
	CRM 7002	MESS- 3	I-1 (0-10 cm)	CRM 7002	MESS- 3	CRM 7002	MESS- 3	I-1 (0-10 cm)	CRM 7002	MESS- 3	I-1 (0-10 cm)
Mean	98.8	68.7	60.4	102.1	93.5	106.5	96.4	87.9	101.0	114.9	109.5
Std. Error	12.3	13.4	5.31	9.57	10.9	13.6	5.45	3.27	6.46	19.1	9.70
Median	89.9	62.9	60.4	99.0	88.2	94.5	96.5	90.4	96.7	95.6	108.5
0.25 Perc.	75.5	55.6	53.5	90.3	80.4	90.5	89.3	83.7	96.7	91.0	95.7
0.75 Perc.	116.2	77.8	67.3	114.8	107.9	117.8	103.5	91.5	110.7	127.6	123.3
Size (N)	6	5	3	3	3	5	3.0	3	7	5	4

Supplementary Figure captions

Fig. E1a: X-ray diffraction pattern of CRM Mess-3 (black trace) showing identified minerals (colored ticks)

Fig. E1b: Normalized X-ray diffraction patterns of CRM Mess-3 before and after NH_4OH extraction (black and red traces, respectively) showing identified minerals (colored ticks)

Fig. E1c: Normalized X-ray diffraction patterns of CRM Mess-3 before and after KOH extraction (black and red traces, respectively) showing identified minerals (colored ticks)

Fig. E1d: Normalized X-ray diffraction patterns of CRM Mess-3 before and after ascorbate extraction (black and red traces, respectively) showing identified minerals (colored ticks)

Fig. E1e: Normalized X-ray diffraction patterns of CRM Mess-3 before and after HCl/HNO_3 extraction (black and red traces, respectively) showing identified minerals (colored ticks)

Fig. E2a: X-ray diffraction pattern of CRM 7002 (black trace) showing identified minerals (colored ticks)

Fig. E2b: Normalized X-ray diffraction patterns of CRM 7002 before and after NH_4OH extraction (black and red traces, respectively) showing identified minerals (colored ticks)

Fig. E2c: Normalized X-ray diffraction patterns of CRM 7002 before and after KOH extraction (black and red traces, respectively) showing identified minerals (colored ticks)

Fig. E2d: Normalized X-ray diffraction patterns of CRM 7002 before and after ascorbate extraction (black and red traces, respectively) showing identified minerals (colored ticks)

Fig. E2e: Normalized X-ray diffraction patterns of CRM 7002 before and after HCl/HNO₃ extraction (black and red traces, respectively) showing identified minerals (colored ticks)

Fig. E3a: X-ray diffraction pattern of sample I-1(0-10m) (black trace) showing identified minerals (colored ticks)

Fig. E3b: Normalized X-ray diffraction patterns of sample I-1(0-10m) before and after NH₄OH extraction (black and red traces, respectively) showing identified minerals (colored ticks)

Fig. E3c: Normalized X-ray diffraction patterns of sample I-1(0-10m) before and after KOH extraction (black and red traces, respectively) showing identified minerals (colored ticks)

Fig. E3d: Normalized X-ray diffraction patterns of sample I-1(0-10m) before and after ascorbate extraction (black and red traces, respectively) showing identified minerals (colored ticks)

Fig. E3e: Normalized X-ray diffraction patterns of sample I-1(0-10m) before and after HCl/HNO₃ extraction (black and red traces, respectively) showing identified minerals (colored ticks)

Fig. E4: X-ray diffraction pattern of nodules from sample I-1(180-200m) (black trace) showing identified minerals (colored ticks)

Fig. E5: Total Hg, (Hg)_{HCl/HNO₃}, (Hg)_{residual}, (Hg)_{Asco}, (Hg)_{NH₄OH}, and (Hg)_{KOH} extractions extraction yield (%) vertical profiles in nodules of pristine soils. Triangles refer to toposequence I, triangles up refers to surface nodules, and

triangle down to deeper horizons III. white and grey filled colours refer to oxisol and ultisol, respectively. Filled triangles refer to soil matrix and hair-crossed symbols refer to nodules.

Fig. E1a. X-ray diffraction pattern of CRM Mess-3 (black trace) showing identified minerals (colored ticks)

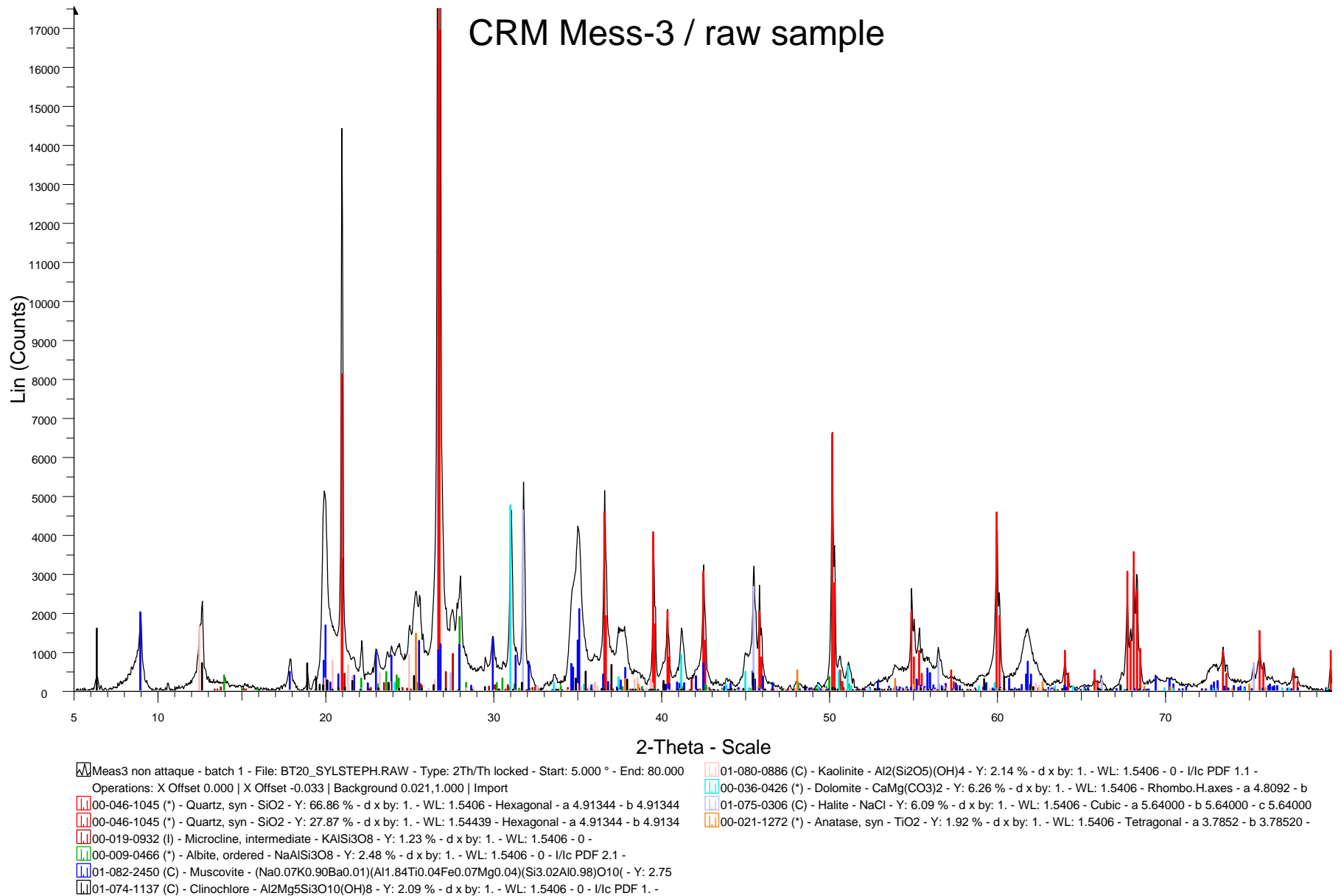
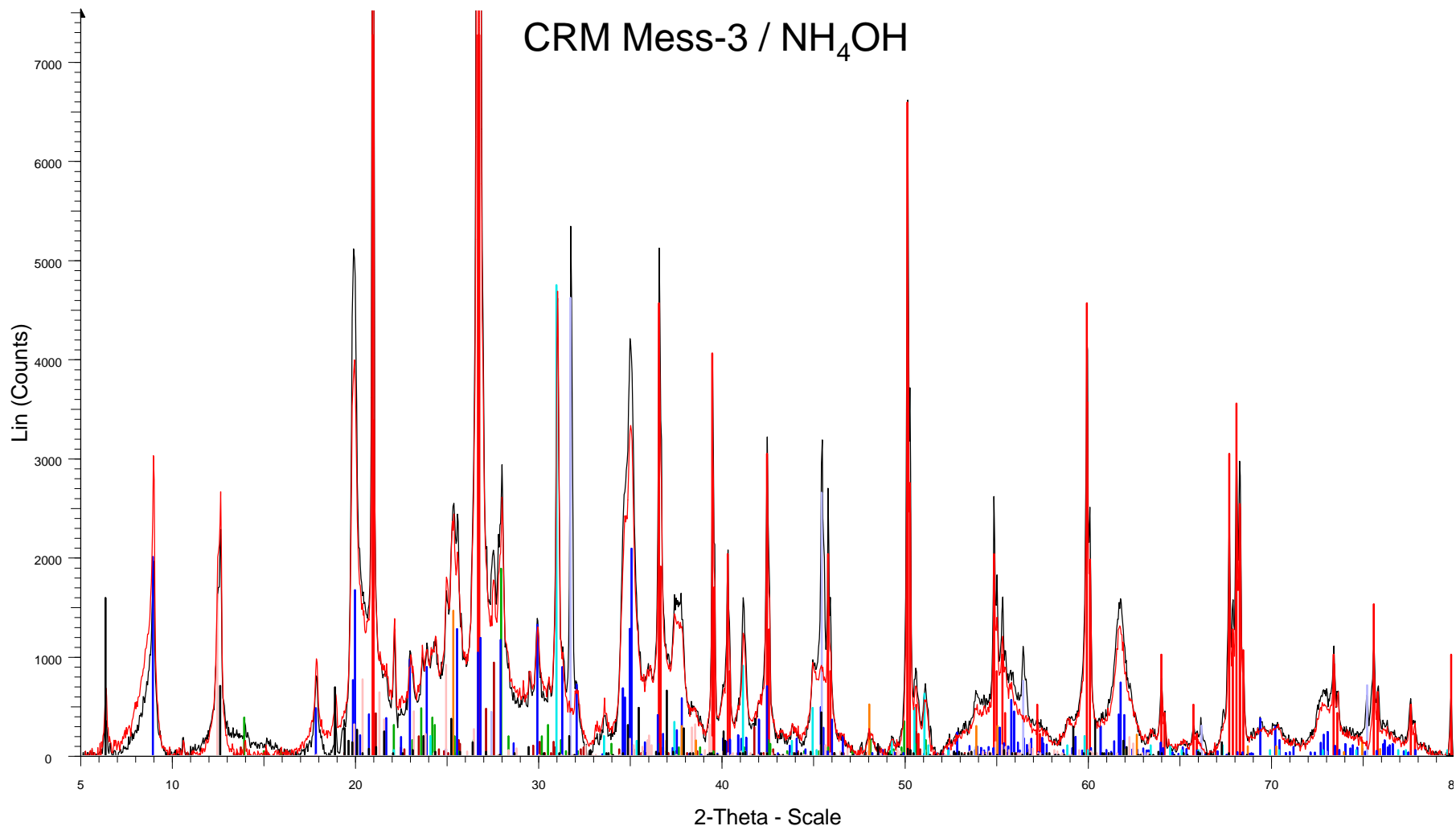
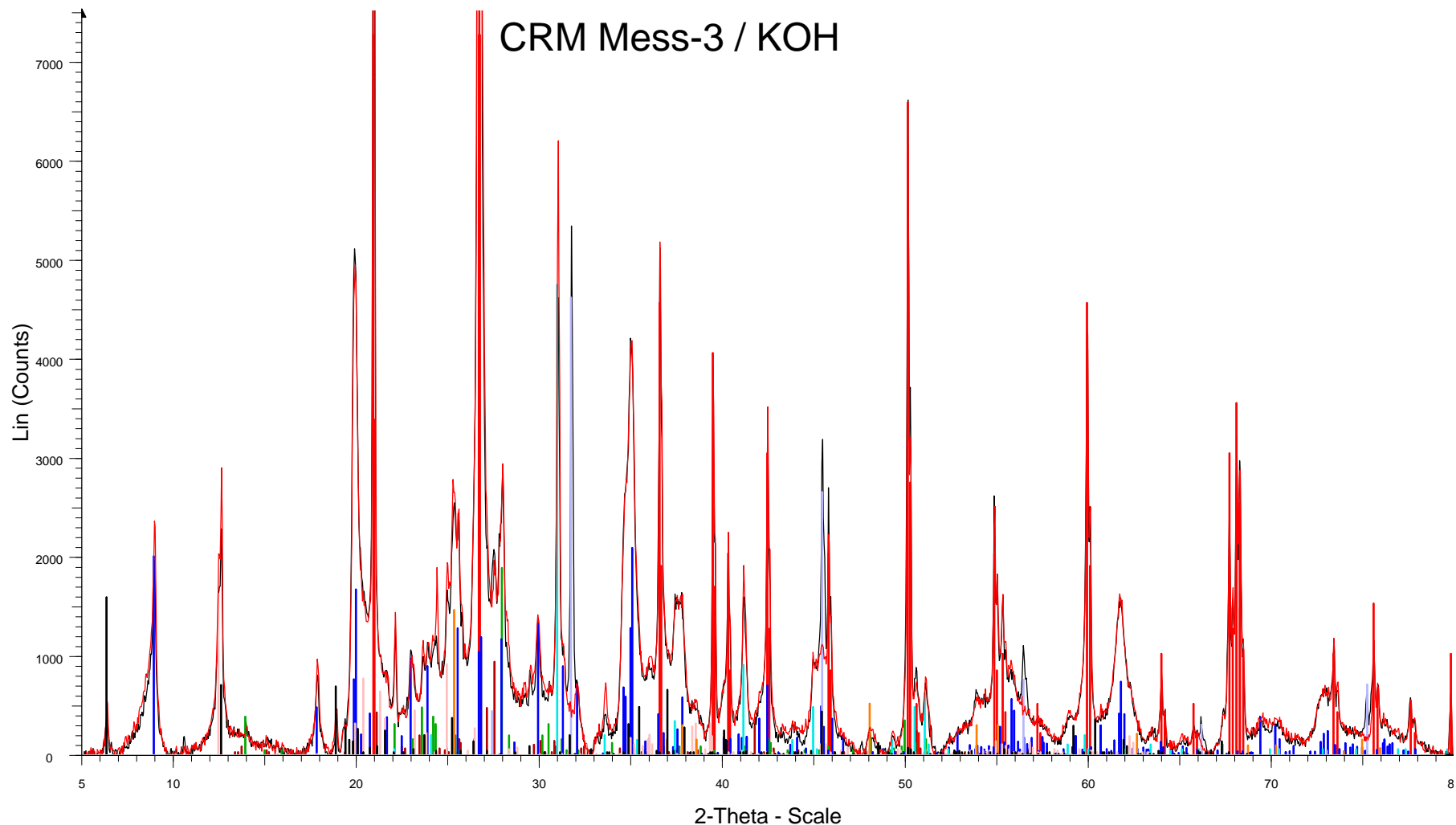


Fig. E1b. Normalized X-ray diffraction patterns of CRM Mess-3 before and after NH_4OH extraction (black and red traces, respectively) showing identified minerals (colored ticks)



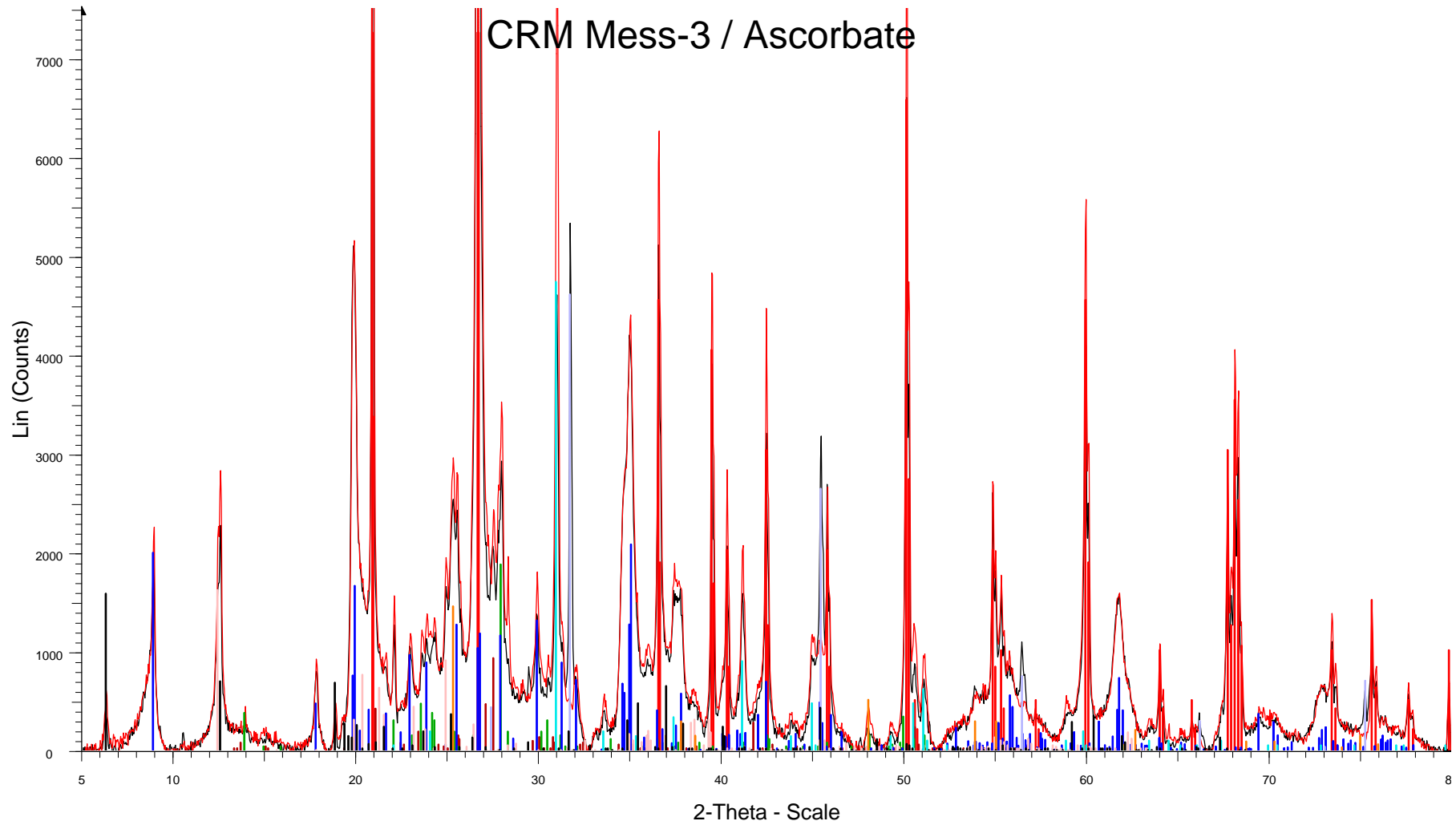
- Meas3 non attaque - batch 1 - File: BT20_SYLSTEPH.RAW - Type: 2Th/Th locked - Start: 5.000 ° - End: 80.000
 Operations: X Offset 0.000 | X Offset -0.033 | Background 0.021,1.000 | Import
- N1 (true) batch1 - File: BT26_SylSteph_1scan.RAW - Type: 2Th/Th locked - Start: 5.000 ° - End: 80.000 ° - Step
 Operations: Background 0.014,1.000 | Import
- 00-046-1045 (*) - Quartz, syn - SiO_2 - Y: 56.62 % - d x by: 1. - WL: 1.5406 - Hexagonal - a 4.91344 - b 4.91344
 00-046-1045 (*) - Quartz, syn - SiO_2 - Y: 23.60 % - d x by: 1. - WL: 1.54439 - Hexagonal - a 4.91344 - b 4.91344
 00-019-0932 (I) - Microcline, intermediate - KAlSi_3O_8 - Y: 1.04 % - d x by: 1. - WL: 1.5406 - 0 -
 00-009-0466 (*) - Albite, ordered - $\text{NaAlSi}_3\text{O}_8$ - Y: 2.10 % - d x by: 1. - WL: 1.5406 - 0 - I/lc PDF 2.1 -
- 01-082-2450 (C) - Muscovite - $(\text{Na}0.07\text{K}0.90\text{Ba}0.01)(\text{Al}1.84\text{Ti}0.04\text{Fe}0.07\text{Mg}0.04)(\text{Si}3.02\text{Al}0.98)\text{O}10$ - Y: 2.33
 01-074-1137 (C) - Clinocllore - $\text{Al}_2\text{Mg}_5\text{Si}_3\text{O}_{10}(\text{OH})_8$ - Y: 1.77 % - d x by: 1. - WL: 1.5406 - 0 - I/lc PDF 1. -
 01-080-0886 (C) - Kaolinite - $\text{Al}_2(\text{Si}_2\text{O}_5)(\text{OH})_4$ - Y: 1.81 % - d x by: 1. - WL: 1.5406 - 0 - I/lc PDF 1.1 -
 00-036-0426 (*) - Dolomite - $\text{CaMg}(\text{CO}_3)_2$ - Y: 5.30 % - d x by: 1. - WL: 1.5406 - Rhombo.H.axes - a 4.8092 - b
 01-075-0306 (C) - Halite - NaCl - Y: 5.16 % - d x by: 1. - WL: 1.5406 - Cubic - a 5.64000 - b 5.64000 - c 5.64000
 00-021-1272 (*) - Anatase, syn - TiO_2 - Y: 1.63 % - d x by: 1. - WL: 1.5406 - Tetragonal - a 3.7852 - b 3.78520 -

Fig. E1c. Normalized X-ray diffraction patterns of CRM Mess-3 before and after KOH extraction (black and red traces, respectively) showing identified minerals (colored ticks)



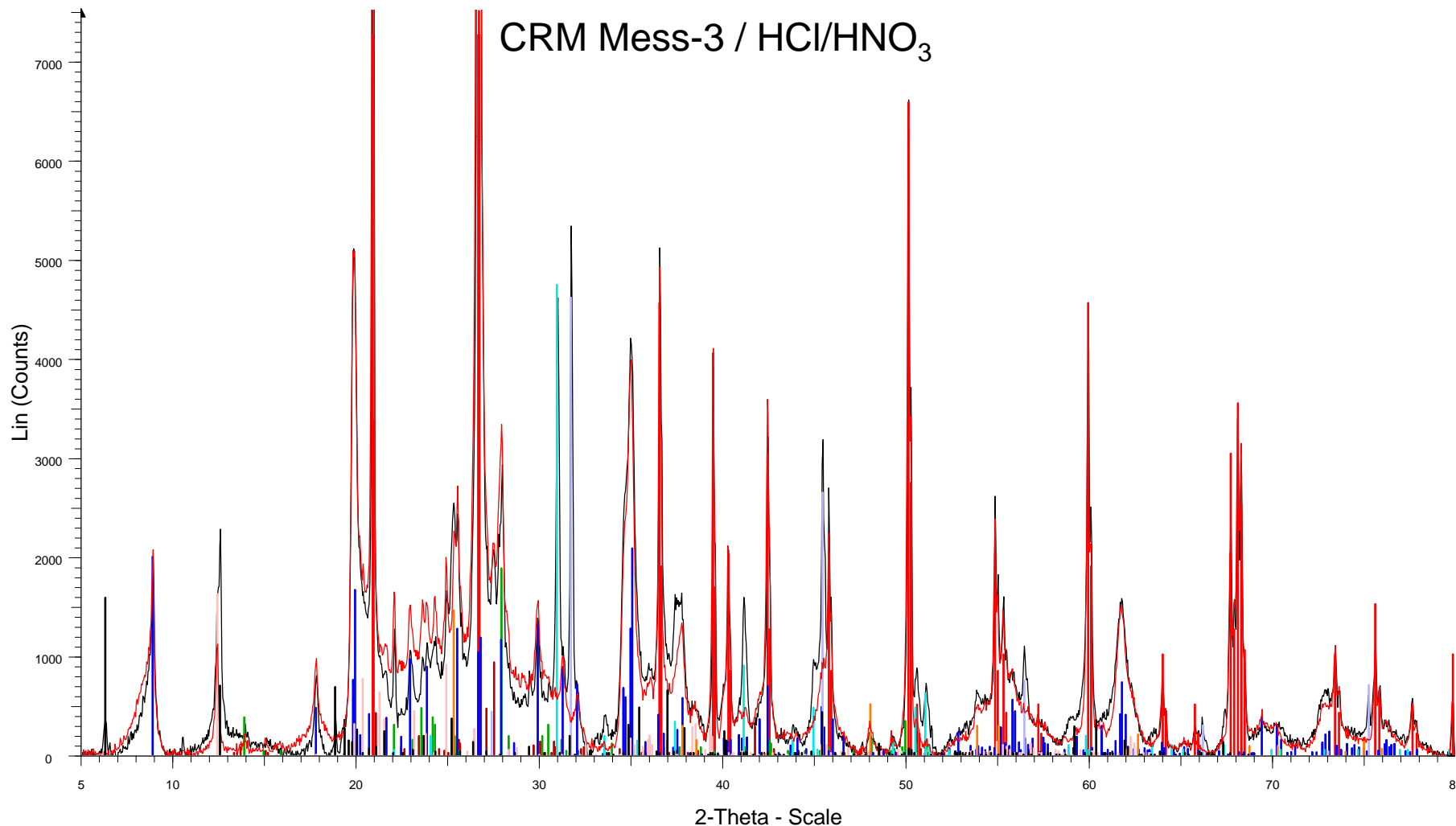
- | | |
|---|--|
| <ul style="list-style-type: none"> ▮ Meas3 non attaque - batch 1 - File: BT20_SYLSTEPH.RAW - Type: 2Th/Th locked - Start: 5.000 ° - End: 80.000
 Operations: X Offset 0.000 X Offset -0.033 Background 0.021,1.000 Import ▮ MESS3 KOH - MESSK2 Batch1 - File: BT22_SylSteph.RAW - Type: 2Th/Th locked - Start: 5.000 ° - End: 80.000
 Operations: Y Scale Mul 1.000 Background 0.014,1.000 Import ▮ 00-046-1045 (*) - Quartz, syn - SiO₂ - Y: 56.62 % - d x by: 1. - WL: 1.5406 - Hexagonal - a 4.91344 - b 4.91344 ▮ 00-046-1045 (*) - Quartz, syn - SiO₂ - Y: 23.60 % - d x by: 1. - WL: 1.54439 - Hexagonal - a 4.91344 - b 4.91344 ▮ 00-019-0932 (I) - Microcline, intermediate - KAlSi₃O₈ - Y: 1.04 % - d x by: 1. - WL: 1.5406 - 0 - ▮ 00-009-0466 (*) - Albite, ordered - NaAlSi₃O₈ - Y: 2.10 % - d x by: 1. - WL: 1.5406 - 0 - I/lc PDF 2.1 - | <ul style="list-style-type: none"> ▮ 01-082-2450 (C) - Muscovite - (Na_{0.07}K_{0.90}Ba_{0.01})(Al_{1.84}Ti_{0.04}Fe_{0.07}Mg_{0.04})(Si_{3.02}Al_{0.98})O₁₀(- Y: 2.33 ▮ 01-074-1137 (C) - Clinocllore - Al₂Mg₅Si₃O₁₀(OH)₈ - Y: 1.77 % - d x by: 1. - WL: 1.5406 - 0 - I/lc PDF 1. - ▮ 01-080-0886 (C) - Kaolinite - Al₂(Si₂O₅)(OH)₄ - Y: 1.81 % - d x by: 1. - WL: 1.5406 - 0 - I/lc PDF 1.1 - ▮ 00-036-0426 (*) - Dolomite - CaMg(CO₃)₂ - Y: 5.30 % - d x by: 1. - WL: 1.5406 - Rhombo.H.axes - a 4.8092 - b ▮ 01-075-0306 (C) - Halite - NaCl - Y: 5.16 % - d x by: 1. - WL: 1.5406 - Cubic - a 5.64000 - b 5.64000 - c 5.64000 ▮ 00-021-1272 (*) - Anatase, syn - TiO₂ - Y: 1.63 % - d x by: 1. - WL: 1.5406 - Tetragonal - a 3.7852 - b 3.78520 - |
|---|--|

Fig. E1d. Normalized X-ray diffraction patterns of CRM Mess-3 before and after ascorbate extraction (black and red traces, respectively) showing identified minerals (colored ticks)



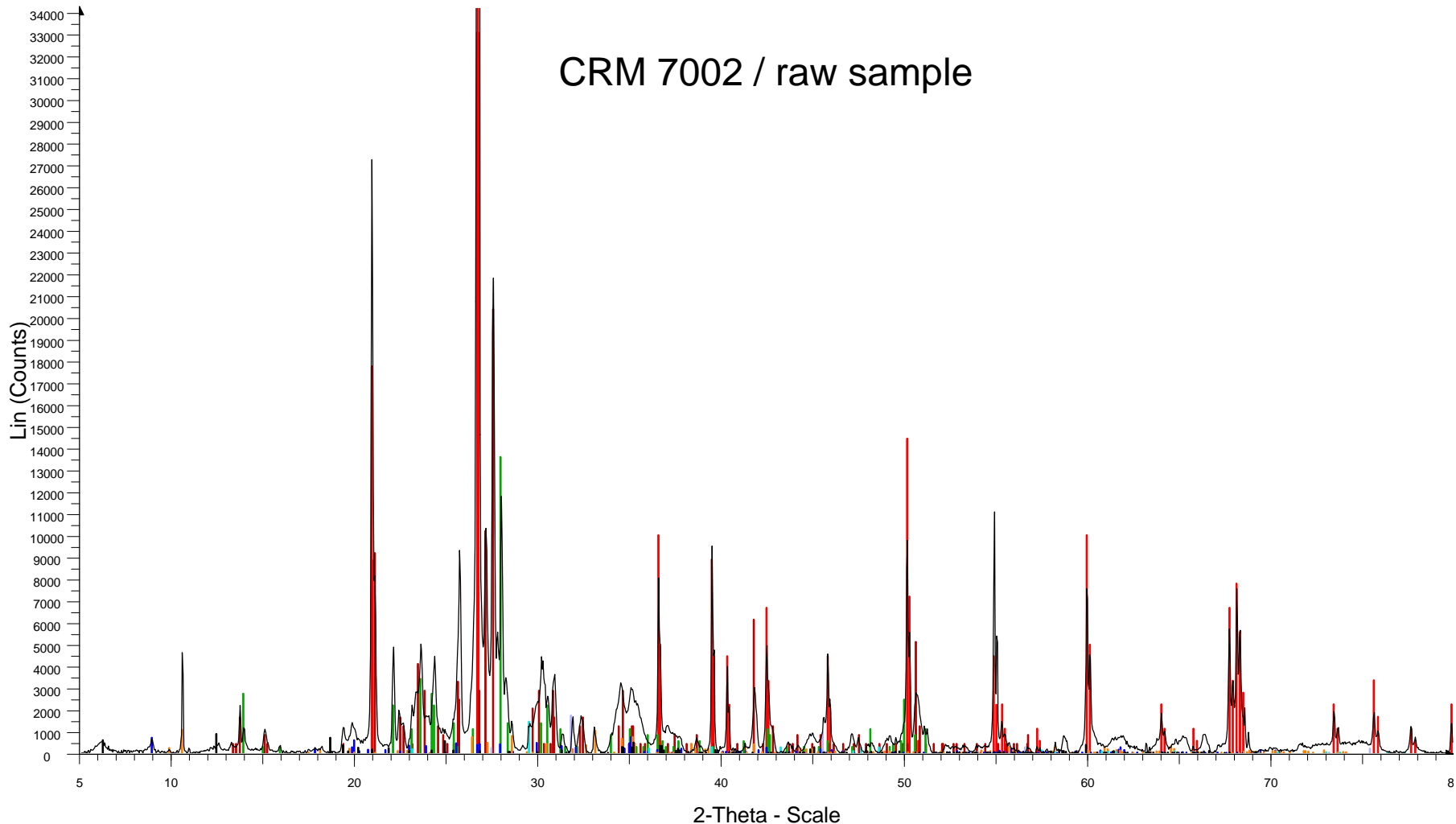
- ▮ Meas3 non attaque - batch 1 - File: BT20_SYLSTEPH.RAW - Type: 2Th/Th locked - Start: 5.000 ° - End: 80.000
 Operations: X Offset 0.000 | X Offset -0.033 | Background 0.021,1.000 | Import
- ▮ Sample A4 - batch1 - File: BT17_SylSteph.RAW - Type: 2Th/Th locked - Start: 5.000 ° - End: 80.000 ° - Step: 0.
 Operations: Background 0.014,1.000 | Import
- ▮ 00-046-1045 (*) - Quartz, syn - SiO₂ - Y: 56.62 % - d x by: 1. - WL: 1.5406 - Hexagonal - a 4.91344 - b 4.91344
▮ 00-046-1045 (*) - Quartz, syn - SiO₂ - Y: 23.60 % - d x by: 1. - WL: 1.54439 - Hexagonal - a 4.91344 - b 4.91344
▮ 00-019-0932 (l) - Microcline, intermediate - KAlSi₃O₈ - Y: 1.04 % - d x by: 1. - WL: 1.5406 - 0 -
▮ 00-009-0466 (*) - Albite, ordered - NaAlSi₃O₈ - Y: 2.10 % - d x by: 1. - WL: 1.5406 - 0 - I/lc PDF 2.1 -
- ▮ 01-082-2450 (C) - Muscovite - (Na_{0.07}K_{0.90}Ba_{0.01})(Al_{1.84}Ti_{0.04}Fe_{0.07}Mg_{0.04})(Si_{3.02}Al_{0.98})O₁₀ - Y: 2.33
▮ 01-074-1137 (C) - Clinocllore - Al₂Mg₅Si₃O₁₀(OH)₈ - Y: 1.77 % - d x by: 1. - WL: 1.5406 - 0 - I/lc PDF 1. -
▮ 01-080-0886 (C) - Kaolinite - Al₂(Si₂O₅)(OH)₄ - Y: 1.81 % - d x by: 1. - WL: 1.5406 - 0 - I/lc PDF 1.1 -
▮ 00-036-0426 (*) - Dolomite - CaMg(CO₃)₂ - Y: 5.30 % - d x by: 1. - WL: 1.5406 - Rhombo.H.axes - a 4.8092 - b
▮ 01-075-0306 (C) - Halite - NaCl - Y: 5.16 % - d x by: 1. - WL: 1.5406 - Cubic - a 5.64000 - b 5.64000 - c 5.64000
▮ 00-021-1272 (*) - Anatase, syn - TiO₂ - Y: 1.63 % - d x by: 1. - WL: 1.5406 - Tetragonal - a 3.7852 - b 3.78520 -

Fig. E1e. Normalized X-ray diffraction patterns of CRM Mess-3 before and after HCl extraction (black and red traces, respectively) showing identified minerals (colored ticks)



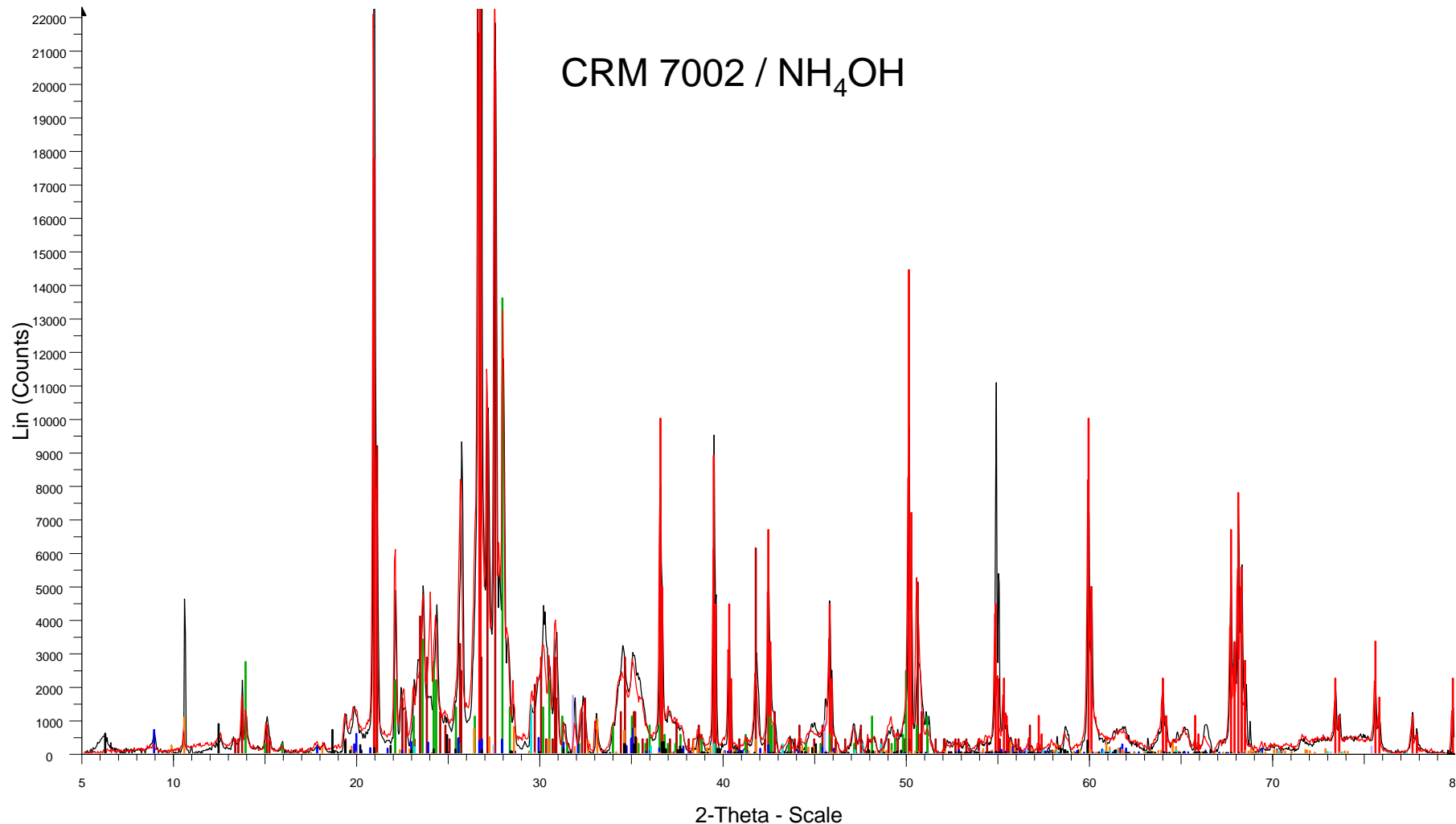
- ▮ Meas3 non attaque - batch 1 - File: BT20_SYLSTEPH.RAW - Type: 2Th/Th locked - Start: 5.000 ° - End: 80.000
 Operations: X Offset 0.000 | X Offset -0.033 | Background 0.021,1.000 | Import
- ▮ MESS3 H1 batch1 - File: BT21_SylSteph.RAW - Type: 2Th/Th locked - Start: 5.000 ° - End: 80.000 ° - Step: 0.0
 Operations: Y Scale Mul 0.750 | Background 0.014,1.000 | Import
- ▮ 00-046-1045 (*) - Quartz, syn - SiO₂ - Y: 56.62 % - d x by: 1. - WL: 1.5406 - Hexagonal - a 4.91344 - b 4.91344
- ▮ 00-046-1045 (*) - Quartz, syn - SiO₂ - Y: 23.60 % - d x by: 1. - WL: 1.54439 - Hexagonal - a 4.91344 - b 4.91344
- ▮ 00-019-0932 (I) - Microcline, intermediate - KAlSi₃O₈ - Y: 1.04 % - d x by: 1. - WL: 1.5406 - 0 -
- ▮ 00-009-0466 (*) - Albite, ordered - NaAlSi₃O₈ - Y: 2.10 % - d x by: 1. - WL: 1.5406 - 0 - I/lc PDF 2.1 -
- ▮ 01-082-2450 (C) - Muscovite - (Na_{0.07}K_{0.90}Ba_{0.01})(Al_{1.84}Ti_{0.04}Fe_{0.07}Mg_{0.04})(Si_{3.02}Al_{0.98})O₁₀(- Y: 2.33
- ▮ 01-074-1137 (C) - Clinoclchlore - Al₂Mg₅Si₃O₁₀(OH)₈ - Y: 1.77 % - d x by: 1. - WL: 1.5406 - 0 - I/lc PDF 1. -
- ▮ 01-080-0886 (C) - Kaolinite - Al₂(Si₂O₅)(OH)₄ - Y: 1.81 % - d x by: 1. - WL: 1.5406 - 0 - I/lc PDF 1.1 -
- ▮ 00-036-0426 (*) - Dolomite - CaMg(CO₃)₂ - Y: 5.30 % - d x by: 1. - WL: 1.5406 - Rhombo.H.axes - a 4.8092 - b
- ▮ 01-075-0306 (C) - Halite - NaCl - Y: 5.16 % - d x by: 1. - WL: 1.5406 - Cubic - a 5.64000 - b 5.64000 - c 5.64000
- ▮ 00-021-1272 (*) - Anatase, syn - TiO₂ - Y: 1.63 % - d x by: 1. - WL: 1.5406 - Tetragonal - a 3.78520 - b 3.78520 -

Fig. E2a. X-ray diffraction pattern of CRM 7002 (black trace) showing identified minerals (colored ticks)



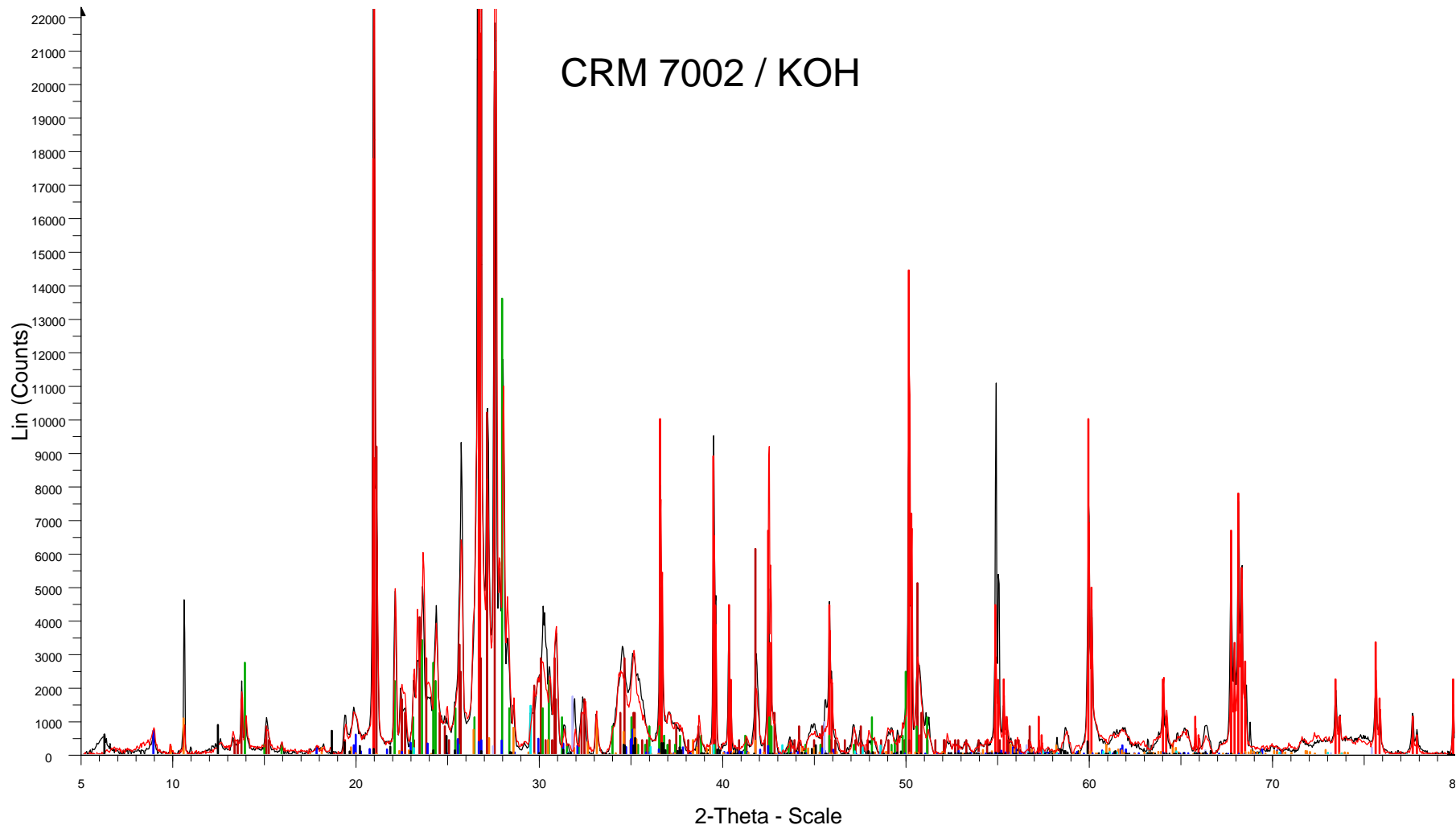
- Batch3 - CRM 7002 non attaqué - File: BT3_SylSteph.RAW - Type: 2Th/Th locked - Start: 5.000 ° - End: 80.000
 Operations: Background 0.120,1.000 | Import
- | | |
|---|---|
| <input checked="" type="checkbox"/> 00-046-1045 (*) - Quartz, syn - SiO ₂ - Y: 68.14 % - d x by: 1. - WL: 1.5406 - 0 - I/c PDF 3.4 - | <input checked="" type="checkbox"/> 01-071-1060 (C) - Hornblende - Na ₉ K ₄ Ca _{1.6} Mg _{2.9} Fe _{1.4} Ti ₅ Al _{2.4} Si ₆ O ₂₄ - Y: 0.65 % - d x by: 1. - WL: 1.5406 |
| <input checked="" type="checkbox"/> 00-046-1045 (*) - Quartz, syn - SiO ₂ - Y: 33.87 % - d x by: 1. - WL: 1.54439 - 0 - I/c PDF 3.4 - | <input checked="" type="checkbox"/> 00-005-0586 (*) - Calcite, syn - CaCO ₃ - Y: 0.88 % - d x by: 1. - WL: 1.5406 - Rhombo.H.axes - a 4.98276 - b 4. |
| <input checked="" type="checkbox"/> 00-019-0932 (I) - Microcline, intermediate - KAlSi ₃ O ₈ - Y: 12.51 % - d x by: 1. - WL: 1.5406 - 0 - | <input checked="" type="checkbox"/> 00-005-0628 (*) - Halite, syn - NaCl - Y: 1.05 % - d x by: 1. - WL: 1.5406 - Cubic - a 5.63080 - b 5.63080 - c 5.63 |
| <input checked="" type="checkbox"/> 00-009-0466 (*) - Albite, ordered - NaAlSi ₃ O ₈ - Y: 8.34 % - d x by: 1. - WL: 1.5406 - 0 - I/c PDF 2.1 - | |
| <input checked="" type="checkbox"/> 01-082-2450 (C) - Muscovite - (Na _{0.07} K _{0.90} Ba _{0.01})(Al _{1.84} Ti _{0.04} Fe _{0.07} Mg _{0.04})(Si _{3.02} Al _{0.98})O ₁₀ (- Y: 0.44 | |
| <input checked="" type="checkbox"/> 01-082-0038 (C) - Clinocllore Ilb-4 (Cr-, Mg-rich) - (Mg _{0.99} Al _{0.01}) ₅ (Al _{0.67} Fe _{0.33})(Si _{3.02} Al _{0.98})O ₁₀ (OH) ₈ - Y: | |

Fig. E2b. Normalized X-ray diffraction patterns of CRM 7002 before and after NH₄OH extraction (black and red traces, respectively) showing identified minerals (colored ticks)



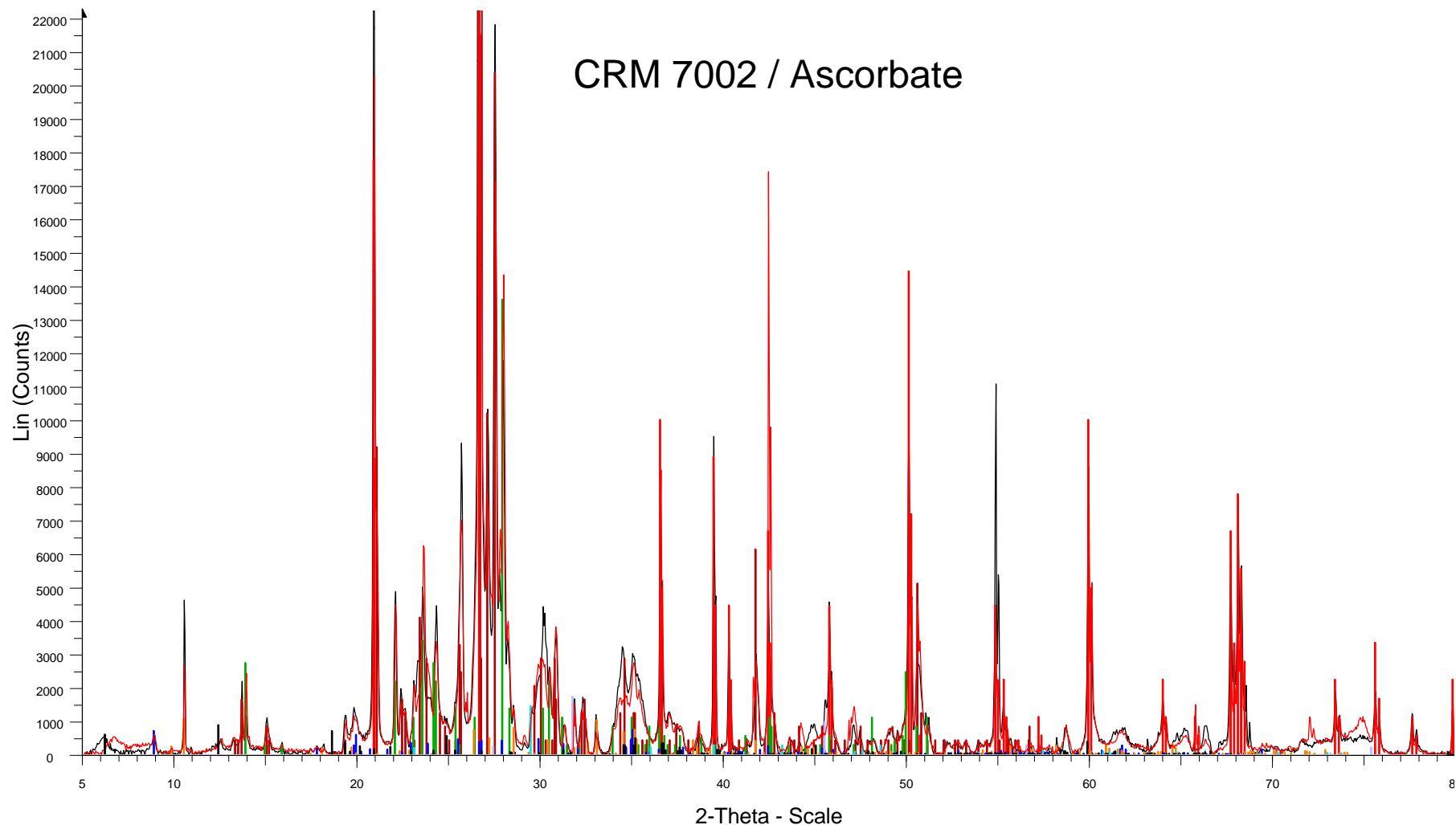
- | | |
|---|---|
| Batch3 - CRM 7002 non attaqué - File: BT3_SylSteph.RAW - Type: 2Th/Th locked - Start: 5.000 ° - End: 80.000
Operations: Background 0.120,1.000 Import | 01-082-2450 (C) - Muscovite - (Na0.07K0.90Ba0.01)(Al1.84Ti0.04Fe0.07Mg0.04)(Si3.02Al0.98)O10(- Y: 0.44 |
| N2 - Batch1 - File: BT23_SylSteph.RAW - Type: 2Th/Th locked - Start: 5.000 ° - End: 80.000 ° - Step: 0.040 ° -
Operations: Y Scale Mul 1.458 Background 0.014,1.000 Import | 01-082-0038 (C) - Clinoclchlore Ilb-4 (Cr-, Mg-rich) - (Mg0.99Al0.01)5(Al0.67Fe0.33)(Si3.02Al0.98)O10(OH)8 - Y: |
| 00-046-1045 (*) - Quartz, syn - SiO2 - Y: 68.14 % - d x by: 1. - WL: 1.5406 - 0 - I/lc PDF 3.4 - | 01-071-1060 (C) - Hornblende - Na.9K.4Ca1.6Mg2.9Fe1.4Ti.5Al2.4Si6O24 - Y: 0.65 % - d x by: 1. - WL: 1.5406 |
| 00-046-1045 (*) - Quartz, syn - SiO2 - Y: 33.87 % - d x by: 1. - WL: 1.54439 - 0 - I/lc PDF 3.4 - | 00-005-0586 (*) - Calcite, syn - CaCO3 - Y: 0.88 % - d x by: 1. - WL: 1.5406 - Rhombo.H.axes - a 4.98276 - b 4. |
| 00-019-0932 (I) - Microcline, intermediate - KAISi3O8 - Y: 12.51 % - d x by: 1. - WL: 1.5406 - 0 - | 00-005-0628 (*) - Halite, syn - NaCl - Y: 1.05 % - d x by: 1. - WL: 1.5406 - Cubic - a 5.63080 - b 5.63080 - c 5.63 |
| 00-009-0466 (*) - Albite, ordered - NaAlSi3O8 - Y: 8.34 % - d x by: 1. - WL: 1.5406 - 0 - I/lc PDF 2.1 - | |

Fig. E2c. Normalized X-ray diffraction patterns of CRM 7002 before and after KOH extraction (black and red traces, respectively) showing identified minerals (colored ticks)



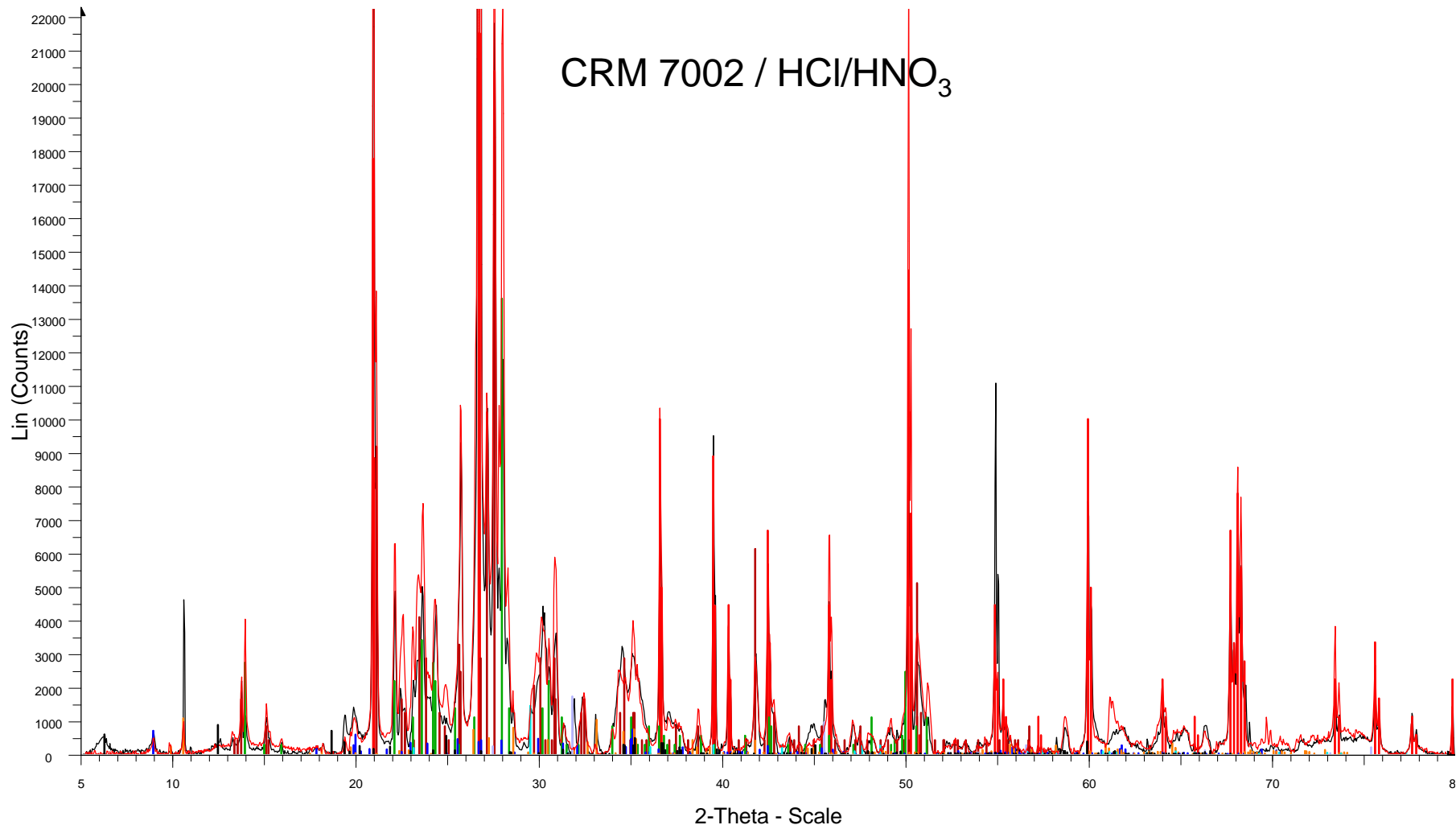
- | | |
|--|---|
| <ul style="list-style-type: none"> Batch3 - CRM 7002 non attaqué - File: BT3_SylSteph.RAW - Type: 2Th/Th locked - Start: 5.000 ° - End: 80.000
Operations: Background 0.120,1.000 Import Sample F batch1 - File: BT12_SylSteph.RAW - Type: 2Th/Th locked - Start: 5.000 ° - End: 80.000 ° - Step: 0.04
Operations: Y Scale Mul 1.125 Background 0.014,1.000 Import 00-046-1045 (*) - Quartz, syn - SiO₂ - Y: 68.14 % - d x by: 1. - WL: 1.5406 - 0 - I/c PDF 3.4 - 00-046-1045 (*) - Quartz, syn - SiO₂ - Y: 33.87 % - d x by: 1. - WL: 1.54439 - 0 - I/c PDF 3.4 - 00-019-0932 (I) - Microcline, intermediate - KAlSi₃O₈ - Y: 12.51 % - d x by: 1. - WL: 1.5406 - 0 - 00-009-0466 (*) - Albite, ordered - NaAlSi₃O₈ - Y: 8.34 % - d x by: 1. - WL: 1.5406 - 0 - I/c PDF 2.1 - | <ul style="list-style-type: none"> 01-082-2450 (C) - Muscovite - (Na_{0.07}K_{0.90}Ba_{0.01})(Al_{1.84}Ti_{0.04}Fe_{0.07}Mg_{0.04})(Si_{3.02}Al_{0.98})O₁₀(- Y: 0.44 01-082-0038 (C) - Clinocllore Ilb-4 (Cr-, Mg-rich) - (Mg_{0.99}Al_{0.01})₅(Al_{0.67}Fe_{0.33})(Si_{3.02}Al_{0.98})O₁₀(OH)₈ - Y: 01-071-1060 (C) - Hornblende - Na₉K₄Ca_{1.6}Mg_{2.9}Fe_{1.4}Ti₅Al_{2.4}Si₆O₂₄ - Y: 0.65 % - d x by: 1. - WL: 1.5406 00-005-0586 (*) - Calcite, syn - CaCO₃ - Y: 0.88 % - d x by: 1. - WL: 1.5406 - Rhombo.H.axes - a 4.98276 - b 4. 00-005-0628 (*) - Halite, syn - NaCl - Y: 1.05 % - d x by: 1. - WL: 1.5406 - Cubic - a 5.63080 - b 5.63080 - c 5.63 |
|--|---|

Fig. E2d. Normalized X-ray diffraction patterns of CRM 7002 before and after ascorbate extraction (black and red traces, respectively) showing identified minerals (colored ticks)



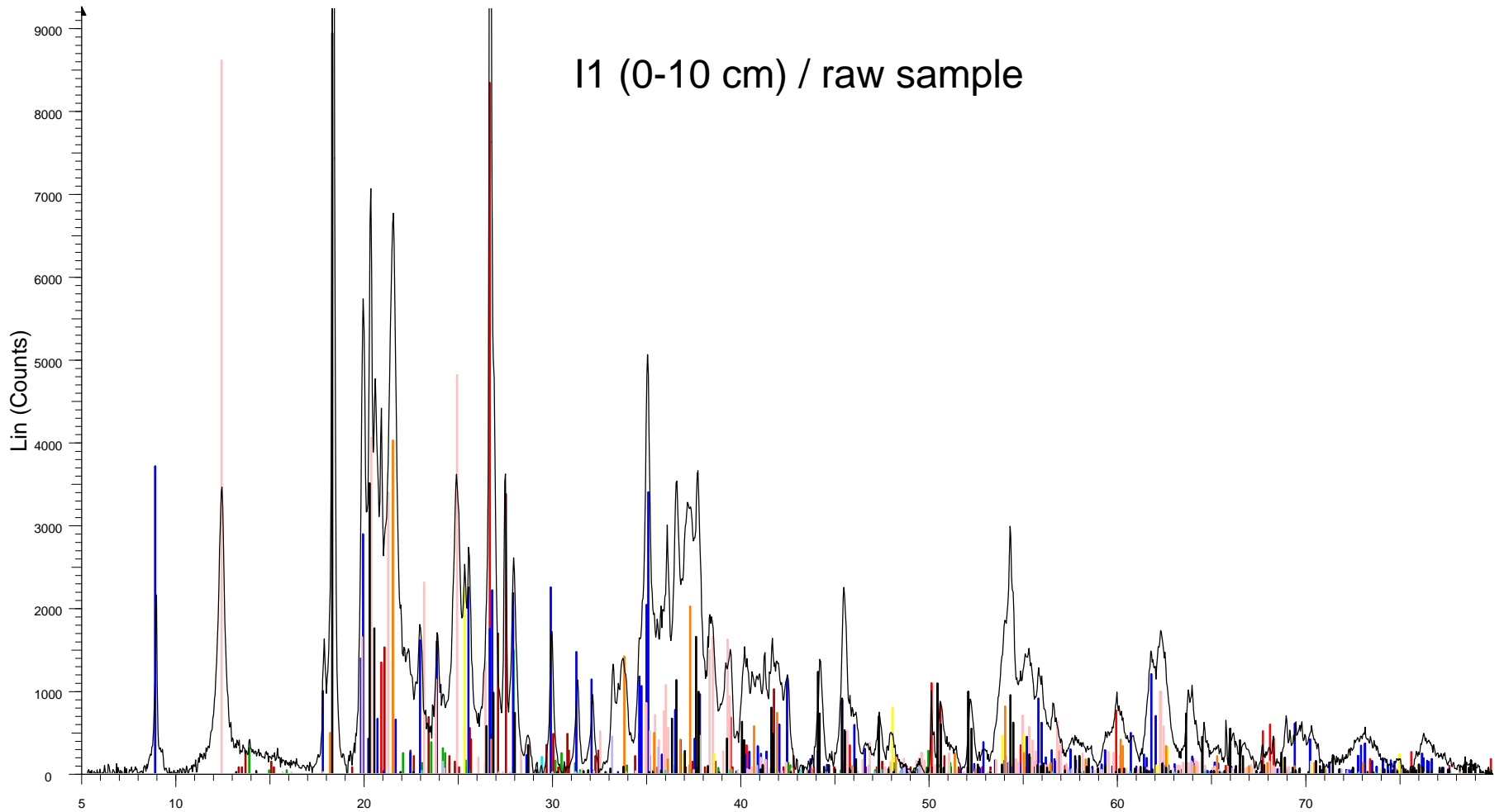
- ▮ Batch3 - CRM 7002 non attaqué - File: BT3_SylSteph.RAW - Type: 2Th/Th locked - Start: 5.000 ° - End: 80.000
 Operations: Background 0.120,1.000 | Import
- ▮ Sample A3 - Batch3 - File: BT16_SylSteph.RAW - Type: 2Th/Th locked - Start: 5.000 ° - End: 80.000 ° - Step: 0.
 Operations: Background 0.014,1.000 | Import
- ▮ 00-046-1045 (*) - Quartz, syn - SiO₂ - Y: 68.14 % - d x by: 1. - WL: 1.5406 - 0 - I/c PDF 3.4 -
▮ 00-046-1045 (*) - Quartz, syn - SiO₂ - Y: 33.87 % - d x by: 1. - WL: 1.54439 - 0 - I/c PDF 3.4 -
▮ 00-019-0932 (I) - Microcline, intermediate - KAlSi₃O₈ - Y: 12.51 % - d x by: 1. - WL: 1.5406 - 0 -
▮ 00-009-0466 (*) - Albite, ordered - NaAlSi₃O₈ - Y: 8.34 % - d x by: 1. - WL: 1.5406 - 0 - I/c PDF 2.1 -
- ▮ 01-082-2450 (C) - Muscovite - (Na_{0.07}K_{0.90}Ba_{0.01})(Al_{1.84}Ti_{0.04}Fe_{0.07}Mg_{0.04})(Si_{3.02}Al_{0.98})O₁₀(- Y: 0.44
▮ 01-082-0038 (C) - Clinocllore Ilb-4 (Cr-, Mg-rich) - (Mg_{0.99}Al_{0.01})₅(Al_{0.67}Fe_{0.33})(Si_{3.02}Al_{0.98})O₁₀(OH)₈ - Y:
▮ 01-071-1060 (C) - Hornblende - Na₉K₄Ca_{1.6}Mg_{2.9}Fe_{1.4}Ti₅Al_{2.4}Si₆O₂₄ - Y: 0.65 % - d x by: 1. - WL: 1.5406
▮ 00-005-0586 (*) - Calcite, syn - CaCO₃ - Y: 0.88 % - d x by: 1. - WL: 1.5406 - Rhombo.H.axes - a 4.98276 - b 4.
▮ 00-005-0628 (*) - Halite, syn - NaCl - Y: 1.05 % - d x by: 1. - WL: 1.5406 - Cubic - a 5.63080 - b 5.63080 - c 5.63

Fig. E2e. Normalized X-ray diffraction patterns of CRM 7002 before and after HCl extraction (black and red traces, respectively) showing identified minerals (colored ticks)



- | | |
|---|---|
| Batch3 - CRM 7002 non attaqué - File: BT3_SylSteph.RAW - Type: 2Th/Th locked - Start: 5.000 ° - End: 80.000
Operations: Background 0.120,1.000 Import | 01-082-2450 (C) - Muscovite - (Na0.07K0.90Ba0.01)(Al1.84Ti0.04Fe0.07Mg0.04)(Si3.02Al0.98)O10(- Y: 0.44 |
| Batch 2 Extraction HG - CRM + HCl - File: BT2_SylSteph.RAW - Type: 2Th/Th locked - Start: 5.000 ° - End: 80.
Operations: Background 0.014,1.000 Import | 01-082-0038 (C) - Clinocllore Ilb-4 (Cr-, Mg-rich) - (Mg0.99Al0.01)5(Al0.67Fe0.33)(Si3.02Al0.98)O10(OH)8 - Y: |
| 00-046-1045 (*) - Quartz, syn - SiO2 - Y: 68.14 % - d x by: 1. - WL: 1.5406 - 0 - I/c PDF 3.4 - | 00-005-0586 (*) - Calcite, syn - CaCO3 - Y: 0.88 % - d x by: 1. - WL: 1.5406 - Rhombo.H.axes - a 4.98276 - b 4. |
| 00-046-1045 (*) - Quartz, syn - SiO2 - Y: 33.87 % - d x by: 1. - WL: 1.54439 - 0 - I/c PDF 3.4 - | 00-005-0628 (*) - Halite, syn - NaCl - Y: 1.05 % - d x by: 1. - WL: 1.5406 - Cubic - a 5.63080 - b 5.63080 - c 5.63 |
| 00-019-0932 (I) - Microcline, intermediate - KAISi3O8 - Y: 12.51 % - d x by: 1. - WL: 1.5406 - 0 - | |
| 00-009-0466 (*) - Albite, ordered - NaAlSi3O8 - Y: 8.34 % - d x by: 1. - WL: 1.5406 - 0 - I/c PDF 2.1 - | |

Fig. E3a. X-ray diffraction pattern of sample I-1(0-10m) (black trace) showing identified minerals (colored ticks)

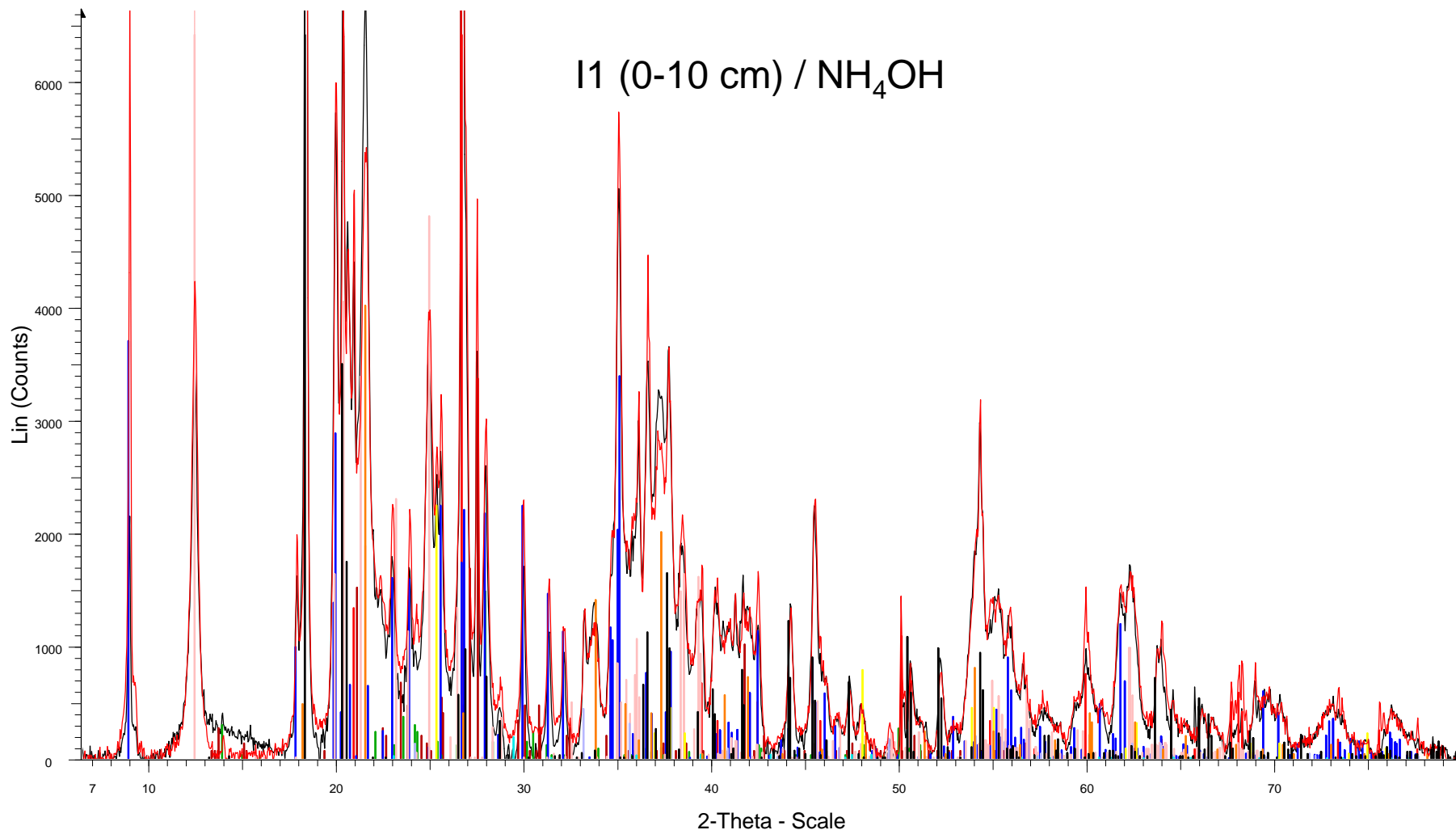


I1 (0-10 cm) / raw sample

2-Theta - Scale

- Batch 7 -- M1 sans attaque - File: BT7_SylSteph_bl.RAW - Type: 2Th/Th locked - Start: 5.000 ° - End: 80.000 °
 Operations: Background 0.014,1.000 | Import
- 00-046-1045 (*) - Quartz, syn - SiO₂ - Y: 57.51 % - d x by: 1. - WL: 1.5406 - Hexagonal - a 4.91344 - b 4.91344
 - 00-019-0932 (I) - Microcline, intermediate - KAlSi₃O₈ - Y: 23.23 % - d x by: 1. - WL: 1.5406 - Triclinic - a 8.560 - b 8.560 - c 5.450
 - 00-009-0466 (*) - Albite, ordered - NaAlSi₃O₈ - Y: 10.22 % - d x by: 1. - WL: 1.5406 - Triclinic - a 8.144 - b 12.7 - c 5.450
 - 01-080-0742 (C) - Muscovite 2M1 - (K_{0.82}Na_{0.18})(Fe_{0.03}Al_{1.97})(AlSi₃)O₁₀(OH)₂ - Y: 25.54 % - d x by: 1. - WL: 1.5406 - Triclinic - a 5.15770 - b 8.560 - c 5.450
 - 01-080-0886 (C) - Kaolinite - Al₂(Si₂O₅)(OH)₄ - Y: 59.38 % - d x by: 1. - WL: 1.5406 - Triclinic - a 5.15770 - b 8.560 - c 5.450
 - 00-005-0586 (*) - Calcite, syn - CaCO₃ - Y: 1.31 % - d x by: 1. - WL: 1.5406 - Rhombo.H.axes - a 4.989 - b 4.989 - c 3.989
 - 00-021-1272 (*) - Anatase, syn - TiO₂ - Y: 15.48 % - d x by: 1. - WL: 1.5406 - Tetragonal - a 3.7852 - b 3.78520 - c 9.514
 - 01-070-2038 (C) - Gibbsite - Al(OH)₃ - Y: 69.49 % - d x by: 1. - WL: 1.5406 - Monoclinic - a 8.68400 - b 5.07800 - c 3.570
 - 00-033-0664 (*) - Hematite, syn - Fe₂O₃ - Y: 3.03 % - d x by: 1. - WL: 1.5406 - Rhombo.H.axes - a 5.0356 - b 5.0356 - c 3.570
 - 00-029-0713 (I) - Goethite - Fe+3O(OH) - Y: 27.70 % - d x by: 1. - WL: 1.5406 - Orthorhombic - a 4.56000 - b 9.514 - c 3.570

Fig. E3b. Normalized X-ray diffraction patterns of sample I-1(0-10m) before and after NH_4OH extraction (black and red traces, respectively) showing identified minerals (colored ticks)



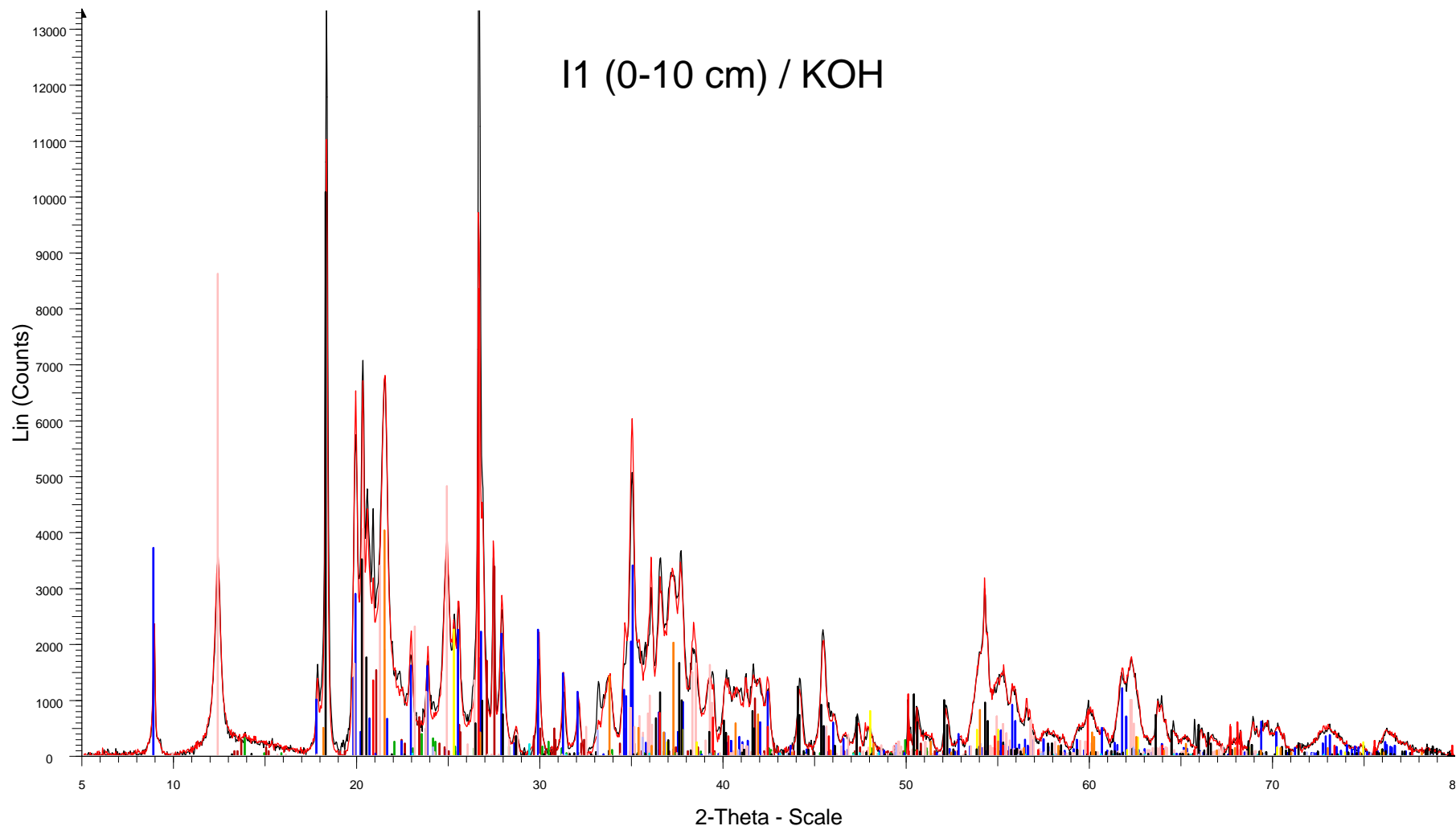
▮ Batch 7 -- M1 sans attaque - File: BT7_SylSteph_bl.RAW - Type: 2Th/Th locked - Start: 5.000 ° - End: 80.000 °
 Operations: Background 0.014,1.000 | Import

▮ N1-batch1 - File: BT25_BL.RAW - Type: 2Th/Th locked - Start: 5.000 ° - End: 80.000 ° - Step: 0.040 ° - Step tim
 Operations: Y Scale Mul 1.000 | Y Scale Mul 1.500 | Background 0.000,1.000 | Import

▮ 00-046-1045 (*) - Quartz, syn - SiO_2 - Y: 42.31 % - d x by: 1. - WL: 1.5406 - Hexagonal - a 4.91344 - b 4.91344
▮ 00-019-0932 (I) - Microcline, intermediate - KAlSi_3O_8 - Y: 17.09 % - d x by: 1. - WL: 1.5406 - Triclinic - a 8.560 - b 8.560
▮ 00-009-0466 (*) - Albite, ordered - $\text{NaAlSi}_3\text{O}_8$ - Y: 7.52 % - d x by: 1. - WL: 1.5406 - Triclinic - a 8.144 - b 12.78
▮ 01-080-0742 (C) - Muscovite 2M1 - $(\text{K}_0.82\text{Na}_0.18)(\text{Fe}_0.03\text{Al}_1.97)(\text{AlSi}_3\text{O}_{10})(\text{OH})_2$ - Y: 18.79 % - d x by: 1. - WL: 1.5406 - Triclinic - a 5.15770 - b 8.560

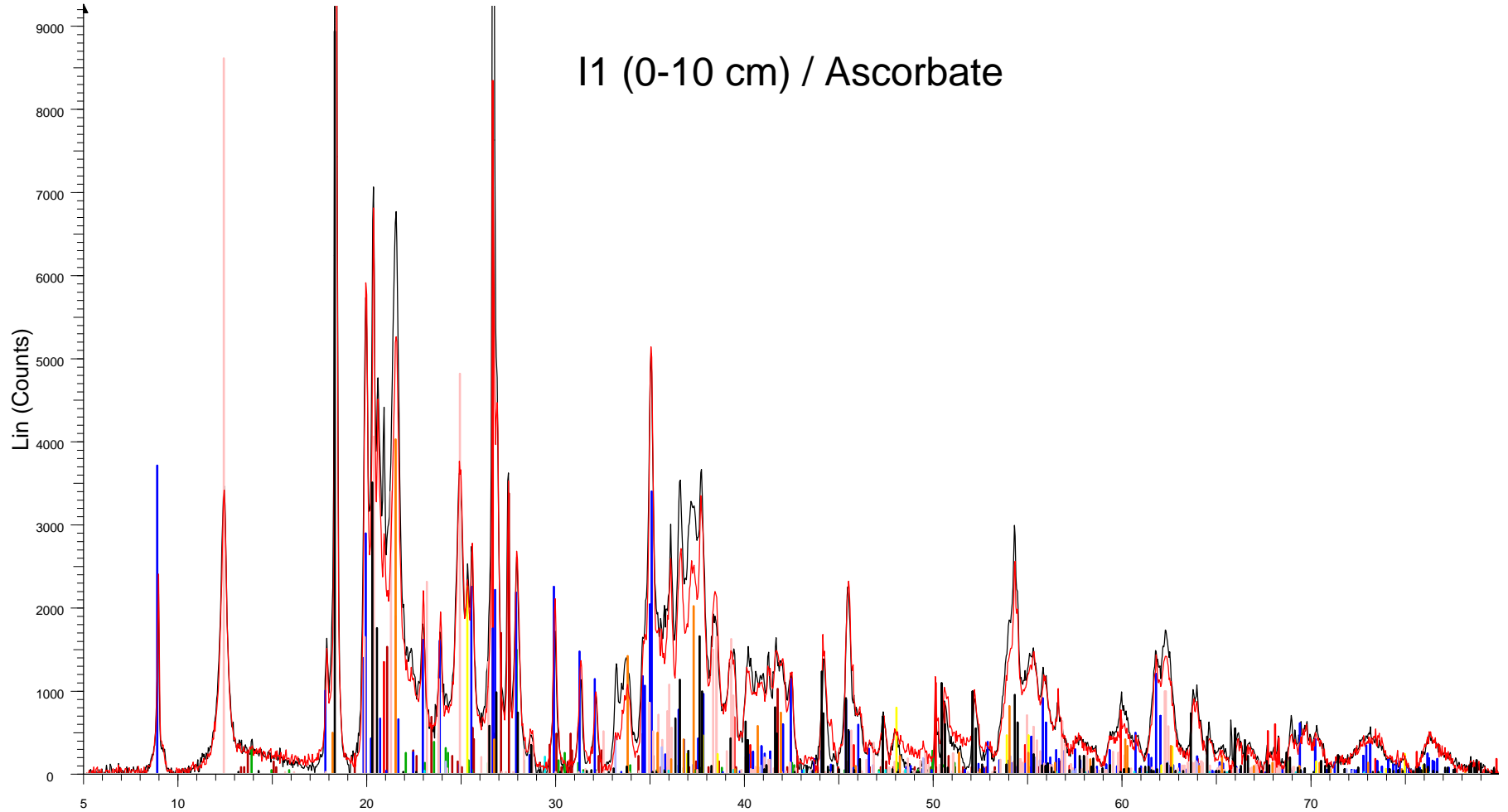
▮ 01-080-0886 (C) - Kaolinite - $\text{Al}_2(\text{Si}_2\text{O}_5)(\text{OH})_4$ - Y: 43.69 % - d x by: 1. - WL: 1.5406 - Triclinic - a 5.15770 - b 8.560
▮ 00-005-0586 (*) - Calcite, syn - CaCO_3 - Y: 0.97 % - d x by: 1. - WL: 1.5406 - Rhombo.H.axes - a 4.989 - b 4.989
▮ 00-021-1272 (*) - Anatase, syn - TiO_2 - Y: 11.39 % - d x by: 1. - WL: 1.5406 - Tetragonal - a 3.7852 - b 3.78520
▮ 01-070-2038 (C) - Gibbsite - $\text{Al}(\text{OH})_3$ - Y: 51.13 % - d x by: 1. - WL: 1.5406 - Monoclinic - a 8.68400 - b 5.07800
▮ 00-033-0664 (*) - Hematite, syn - Fe_2O_3 - Y: 2.23 % - d x by: 1. - WL: 1.5406 - Rhombo.H.axes - a 5.0356 - b 5.0356
▮ 00-029-0713 (I) - Goethite - $\text{Fe}+3\text{O}(\text{OH})$ - Y: 20.38 % - d x by: 1. - WL: 1.5406 - Orthorhombic - a 4.56000 - b 9.00000

Fig. E3c. Normalized X-ray diffraction patterns of sample I-1(0-10m) before and after KOH extraction (black and red traces, respectively) showing identified minerals (colored ticks)



- | | |
|---|--|
| Batch 7 -- M1 sans attaque - File: BT7_SylSteph_bl.RAW - Type: 2Th/Th locked - Start: 5.000 ° - End: 80.000 °
Operations: Background 0.014,1.000 Import | 01-080-0886 (C) - Kaolinite - Al ₂ (Si ₂ O ₅)(OH) ₄ - Y: 59.38 % - d x by: 1. - WL: 1.5406 - Triclinic - a 5.15770 - b 8. |
| Batch 8 -- Ech H - File: BT8_SylSteph.RAW - Type: 2Th/Th locked - Start: 5.000 ° - End: 80.000 ° - Step: 0.040
Operations: Background 0.014,1.000 Import | 00-005-0586 (*) - Calcite, syn - CaCO ₃ - Y: 1.31 % - d x by: 1. - WL: 1.5406 - Rhombo.H.axes - a 4.989 - b 4.98 |
| 00-046-1045 (*) - Quartz, syn - SiO ₂ - Y: 57.51 % - d x by: 1. - WL: 1.5406 - Hexagonal - a 4.91344 - b 4.91344 | 00-021-1272 (*) - Anatase, syn - TiO ₂ - Y: 15.48 % - d x by: 1. - WL: 1.5406 - Tetragonal - a 3.7852 - b 3.78520 |
| 00-019-0932 (I) - Microcline, intermediate - KAlSi ₃ O ₈ - Y: 23.23 % - d x by: 1. - WL: 1.5406 - Triclinic - a 8.560 - b 12.7 | 01-070-2038 (C) - Gibbsite - Al(OH) ₃ - Y: 69.49 % - d x by: 1. - WL: 1.5406 - Monoclinic - a 8.68400 - b 5.07800 |
| 00-009-0466 (*) - Albite, ordered - NaAlSi ₃ O ₈ - Y: 10.22 % - d x by: 1. - WL: 1.5406 - Triclinic - a 8.144 - b 12.7 | 00-033-0664 (*) - Hematite, syn - Fe ₂ O ₃ - Y: 3.03 % - d x by: 1. - WL: 1.5406 - Rhombo.H.axes - a 5.0356 - b 5. |
| 01-080-0742 (C) - Muscovite 2M1 - (K _{0.82} Na _{0.18})(Fe _{0.03} Al _{1.97})(AlSi ₃)O ₁₀ (OH) ₂ - Y: 25.54 % - d x by: 1. - WL | 00-029-0713 (I) - Goethite - Fe+3O(OH) - Y: 27.70 % - d x by: 1. - WL: 1.5406 - Orthorhombic - a 4.56000 - b 9. |

Fig. E3d. Normalized X-ray diffraction patterns of sample I-1(0-10m) before and after ascorbate extraction (black and red traces, respectively) showing identified minerals (colored ticks)

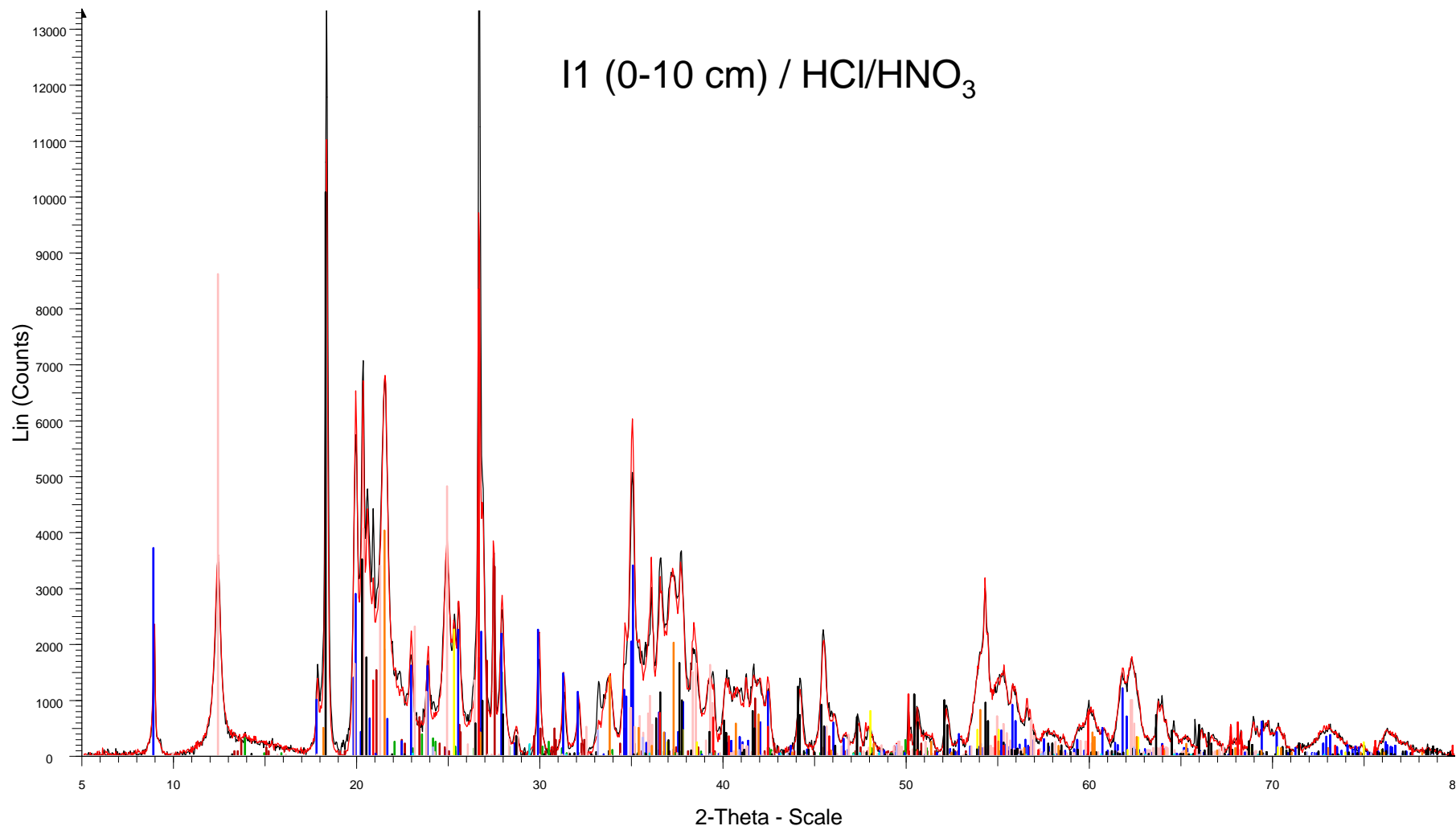


I1 (0-10 cm) / Ascorbate

2-Theta - Scale

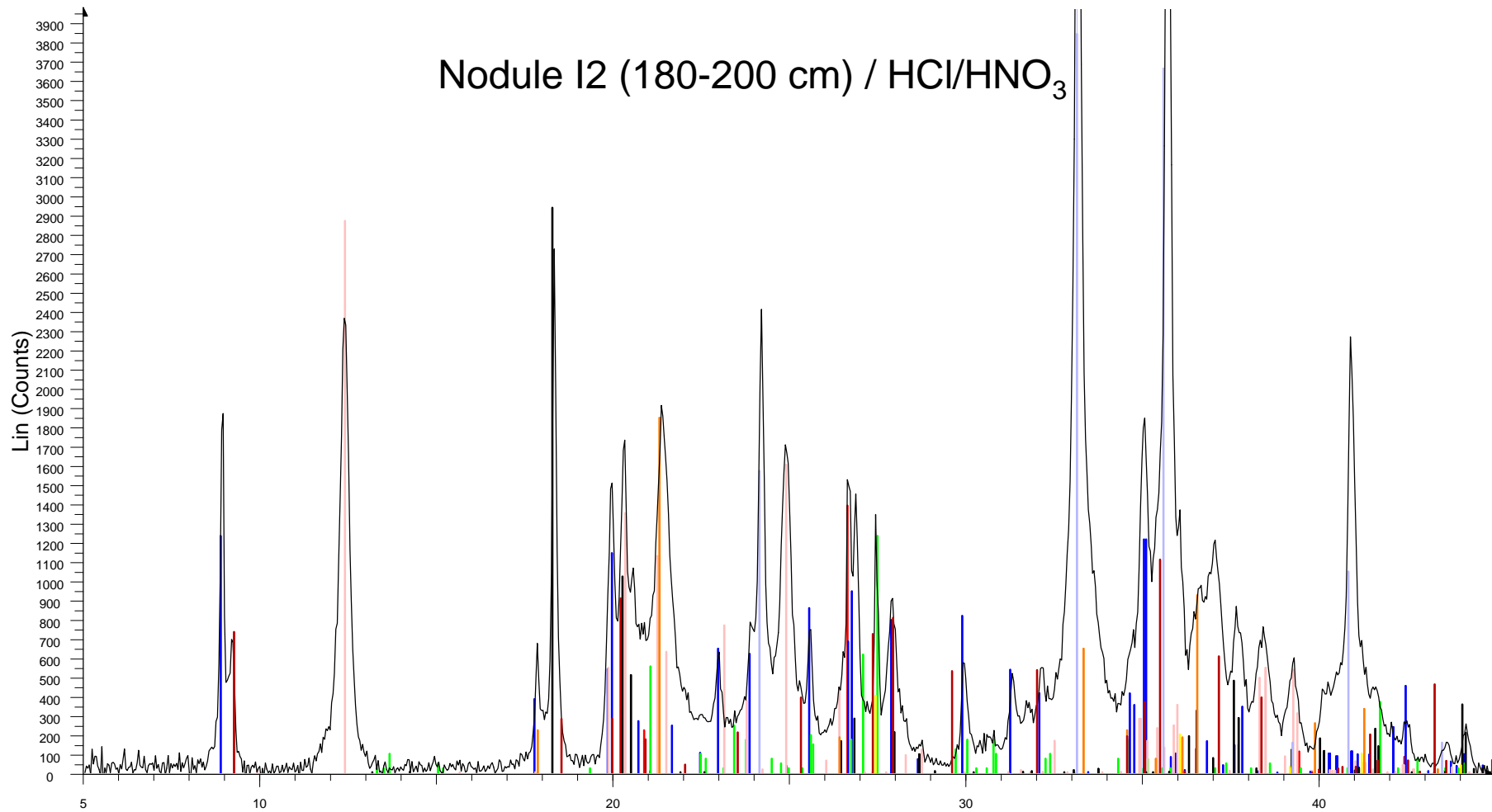
- | | |
|--|--|
| <p>Batch 7 -- M1 sans attaque - File: BT7_SylSteph_bl.RAW - Type: 2Th/Th locked - Start: 5.000 ° - End: 80.000 °
Operations: Background 0.014,1.000 Import</p> <p>Sample A8 - Batch1 - File: Bt18_BL.RAW - Type: 2Th/Th locked - Start: 5.000 ° - End: 80.000 ° - Step: 0.040 ° -
Operations: Background 0.000,1.000 Import</p> <p>00-046-1045 (*) - Quartz, syn - SiO₂ - Y: 57.51 % - d x by: 1. - WL: 1.5406 - Hexagonal - a 4.91344 - b 4.91344</p> <p>00-019-0932 (I) - Microcline, intermediate - KAlSi₃O₈ - Y: 23.23 % - d x by: 1. - WL: 1.5406 - Triclinic - a 8.560 - b 12.7</p> <p>00-009-0466 (*) - Albite, ordered - NaAlSi₃O₈ - Y: 10.22 % - d x by: 1. - WL: 1.5406 - Triclinic - a 8.144 - b 12.7</p> <p>01-080-0742 (C) - Muscovite 2M1 - (K_{0.82}Na_{0.18})(Fe_{0.03}Al_{1.97})(AlSi₃)O₁₀(OH)₂ - Y: 25.54 % - d x by: 1. - WL: 1.5406 - Triclinic - a 5.15770 - b 8.4989</p> | <p>01-080-0886 (C) - Kaolinite - Al₂(Si₂O₅)(OH)₄ - Y: 59.38 % - d x by: 1. - WL: 1.5406 - Triclinic - a 5.15770 - b 8.4989</p> <p>00-005-0586 (*) - Calcite, syn - CaCO₃ - Y: 1.31 % - d x by: 1. - WL: 1.5406 - Rhombo.H.axes - a 4.989 - b 4.989</p> <p>00-021-1272 (*) - Anatase, syn - TiO₂ - Y: 15.48 % - d x by: 1. - WL: 1.5406 - Tetragonal - a 3.7852 - b 3.78520</p> <p>01-070-2038 (C) - Gibbsite - Al(OH)₃ - Y: 69.49 % - d x by: 1. - WL: 1.5406 - Monoclinic - a 8.68400 - b 5.07800</p> <p>00-033-0664 (*) - Hematite, syn - Fe₂O₃ - Y: 3.03 % - d x by: 1. - WL: 1.5406 - Rhombo.H.axes - a 5.0356 - b 5.0356</p> <p>00-029-0713 (I) - Goethite - Fe+3O(OH) - Y: 27.70 % - d x by: 1. - WL: 1.5406 - Orthorhombic - a 4.56000 - b 9.45600</p> |
|--|--|

Fig. E3e. Normalized X-ray diffraction patterns of sample I-1(0-10m) before and after HCl extraction (black and red traces, respectively) showing identified minerals (colored ticks)



- | | |
|---|---|
| <ul style="list-style-type: none"> ▮ Batch 7 -- M1 sans attaque - File: BT7_SylSteph_bl.RAW - Type: 2Th/Th locked - Start: 5.000 ° - End: 80.000 °
 Operations: Background 0.014,1.000 Import ▮ Batch 8 -- Ech H - File: BT8_SylSteph.RAW - Type: 2Th/Th locked - Start: 5.000 ° - End: 80.000 ° - Step: 0.040
 Operations: Background 0.014,1.000 Import ▮ 00-046-1045 (*) - Quartz, syn - SiO₂ - Y: 57.51 % - d x by: 1. - WL: 1.5406 - Hexagonal - a 4.91344 - b 4.91344 ▮ 00-019-0932 (I) - Microcline, intermediate - KAlSi₃O₈ - Y: 23.23 % - d x by: 1. - WL: 1.5406 - Triclinic - a 8.560 - b 12.7 ▮ 00-009-0466 (*) - Albite, ordered - NaAlSi₃O₈ - Y: 10.22 % - d x by: 1. - WL: 1.5406 - Triclinic - a 8.144 - b 12.7 ▮ 01-080-0742 (C) - Muscovite 2M1 - (K_{0.82}Na_{0.18})(Fe_{0.03}Al_{1.97})(AlSi₃)O₁₀(OH)₂ - Y: 25.54 % - d x by: 1. - WL: 1.5406 - Triclinic - a 5.15770 - b 8. | <ul style="list-style-type: none"> ▮ 01-080-0886 (C) - Kaolinite - Al₂(Si₂O₅)(OH)₄ - Y: 59.38 % - d x by: 1. - WL: 1.5406 - Triclinic - a 5.15770 - b 8. ▮ 00-005-0586 (*) - Calcite, syn - CaCO₃ - Y: 1.31 % - d x by: 1. - WL: 1.5406 - Rhombo.H.axes - a 4.989 - b 4.98 ▮ 00-021-1272 (*) - Anatase, syn - TiO₂ - Y: 15.48 % - d x by: 1. - WL: 1.5406 - Tetragonal - a 3.7852 - b 3.78520 ▮ 01-070-2038 (C) - Gibbsite - Al(OH)₃ - Y: 69.49 % - d x by: 1. - WL: 1.5406 - Monoclinic - a 8.68400 - b 5.07800 ▮ 00-033-0664 (*) - Hematite, syn - Fe₂O₃ - Y: 3.03 % - d x by: 1. - WL: 1.5406 - Rhombo.H.axes - a 5.0356 - b 5. ▮ 00-029-0713 (I) - Goethite - Fe₃O(OH) - Y: 27.70 % - d x by: 1. - WL: 1.5406 - Orthorhombic - a 4.56000 - b 9. |
|---|---|

Fig. E4. X-ray diffraction pattern of nodules from sample I-1(180-200m) (black trace) showing identified minerals (colored ticks)



Nodule I2 (180-200 cm) / HCl/HNO₃

2-Theta - Scale

- M1 180-200 Batch1 - File: BT34_SylSteph_uni.RAW - Type: 2Th/Th locked - Start: 5.000 ° - End: 80.000 ° - Ste
 Operations: Background 0.000,0.000 | Import
- 00-046-1045 (*) - Quartz, syn - SiO₂ - Y: 23.52 % - d x by: 1. - WL: 1.5406 - Hexagonal - a 4.91344 - b 4.91344
 - 01-086-1384 (C) - Muscovite 2M1 - K_{0.894}Al_{1.93}(Al_{0.943}Si_{2.829}O₁₀)(OH)1.744F_{0.256} - Y: 20.84 % - d x by:
 - 01-083-2129 (C) - Paragonite 3T - NaAl₂(Si₃Al)O₁₀(OH)₂ - Y: 18.76 % - d x by: 1. - WL: 1.5406 - Hexagonal - a
 - 00-019-0932 (I) - Microcline, intermediate - KAlSi₃O₈ - Y: 20.84 % - d x by: 1. - WL: 1.5406 - Triclinic - a 8.560 -
 - 01-080-0886 (C) - Kaolinite - Al₂(Si₂O₅)(OH)₄ - Y: 48.56 % - d x by: 1. - WL: 1.5406 - Triclinic - a 5.15770 - b 8.
 - 01-070-2038 (C) - Gibbsite - Al(OH)₃ - Y: 49.73 % - d x by: 1. - WL: 1.5406 - Monoclinic - a 8.68400 - b 5.07800
 - 00-033-0664 (*) - Hematite, syn - Fe₂O₃ - Y: 88.55 % - d x by: 1. - WL: 1.5406 - Rhombo.H.axes - a 5.0356 - b
 - 00-029-0713 (I) - Goethite - Fe₃O(OH) - Y: 31.26 % - d x by: 1. - WL: 1.5406 - Orthorhombic - a 4.59072 - b 9.
 - 00-021-1276 (*) - Rutile, syn - TiO₂ - Y: 6.78 % - d x by: 1. - WL: 1.5406 - Tetragonal - a 4.5933 - b 4.59330 - c

Fig. E5.

