

1 **First crystal-structure determination of chromites from an**
2 **acapulcoite and ordinary chondrites**

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23 **ABSTRACT** – We report the first crystal structure determinations of chromites from an
24 acapulcoite and from ordinary chondrites. Cell edges range from 8.3212 (3) to 8.3501
25 (1) Å, while the oxygen positional parameters are in the range 0.2624 (3) to 0.26298 (9).
26 Their compositions show they are very close to the chromite end-member FeCr_2O_4 with
27 limited Al and Mg content. Titanium oxide content exceeds 1%, whereas the amount of
28 Fe^{3+} is negligible. Extraterrestrial chromite is readily distinguished from terrestrial
29 analogues on the basis of cell edge and oxygen positional parameter. These distinctions
30 will facilitate ongoing attempts to reconstruct the paleoflux of meteorites to Earth from
31 resistant extraterrestrial spinel grains recovered from ancient sediments.

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33 Keywords: chromite; acapulcoite; ordinary chondrite; X-ray single crystal diffraction;
34 meteorite flux

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1. INTRODUCTION

36 Chromite is a minor but regular constituent of ordinary chondrites and its chemistry
37 in meteorites has been studied since the 1960's. Bunch *et al.* (1967) were the first to
38 give a survey of the chemical composition of chondritic chromites. They suggested that
39 there is a relationship **between chromite Fe-Mg contents and olivine Fe-Mg contents** in
40 the H, L and LL groups of equilibrated chondrites. Besides the main components FeO
41 and Cr_2O_3 , chromite contains minor amounts of MgO and Al_2O_3 . The increase in
42 $\text{Fe}^{2+}/(\text{Fe}^{2+}+\text{Mg})$ from H to LL chondrites is evident. **Nonetheless**, Wlotzka (2005)
43 argued it was not possible to distinguish between the H and L groups from the chromite
44 composition alone, as has been attempted for fossil chondrites (Bridges *et al.*, 2007),
45 because of the overlapping $\text{Fe}^{2+}/(\text{Fe}^{2+}+\text{Mg})$ ratios of chromites. Besides Al and Mg, Ti
46 is a major element in the chromite. Although there is overlap in the composition of

47 chromites from H, L and LL chondrites, the average element composition of a larger set
48 of chromite grains from fossil meteorite gives an indication of the dominant origin of
49 the grains (Schmitz, 2013). This relation has been confirmed by oxygen isotopic
50 analyses of chromites from fossil Ordovician meteorites showing that these indeed are
51 dominantly pseudomorphosed L chondrites, as indicated by their chromite chemical
52 composition (Heck *et al.*, 2010). Acapulcoites are coarsely crystalline, unshocked
53 objects of chondritic composition and igneous texture (Palme *et al.*, 1981; Folco *et al.*,
54 2006; Rubin, 2007). Equilibration temperatures for acapulcoites are much higher than
55 those estimated for the most equilibrated ordinary chondrites (950-1250 K; Dodd,
56 1969). Oxygen-three isotopes indicate possible aqueous alteration of acapulcoites prior
57 to the onset of thermal metamorphism (Greenwood *et al.*, 2012), and a comparison of
58 chromite from acapulcoites and ordinary chondrites is thus of interest.

59 It is well known that **terrestrial** Cr-bearing spinel can be considered as a petrogenetic
60 indicator because chromite composition is a rich source of information on the origin and
61 evolution of its parent magmas (Irvine, 1967; Dick and Bullen, 1984). The relationships
62 between composition, structural parameters and genetic environment of Cr-bearing
63 spinels from different terrestrial occurrences have been considered by several authors
64 for a better understanding of their genesis and/or oxidation mechanisms (Della Giusta *et*
65 *al.*, 1986; Princivalle *et al.*, 1989; Carbonin *et al.*, 1999; Carraro, 2003; Bosi *et al.*,
66 2004; Uchida *et al.*, 2005; Lenaz *et al.*, 2010, 2011; Derbyshire *et al.*, 2013; Princivalle
67 *et al.*, 2014).

68 This study presents the first structural and chemical data of meteoritic chromite. Data
69 are given for a total of nine chromite **grains, including one from an acapulcoite (labeled**
70 **ACAP), and four each from the Kernouvé H6 ordinary chondrite (H6) and from an ~**
71 **470 Ma old fossil meteorite, Gol 001 (GOL), found in Ordovician limestone at**
72 **Kinneulle in Sweden and originating from the coeval disruption of the L-chondrite**

73 parent body (Schmitz *et al.*, 2001; Heck *et al.*, 2010; Schmitz, 2013). Recently,
74 Alwmark and Schmitz (2009) and Heck *et al.* (2010) distinguished terrestrial and
75 extraterrestrial chromite using relict silicate inclusions and high-precision oxygen three-
76 isotopes SIMS analyses, respectively. As these analyses are time-consuming and
77 necessitate a lot of material, our study aims to determine whether structural parameters,
78 such as cell edge and oxygen positional parameter, can be used to distinguish
79 extraterrestrial from terrestrial chromite. Since structure refinement requires much less
80 material, this will facilitate ongoing attempts to reconstruct the paleoflux of meteorites
81 to Earth from resistant extraterrestrial spinel grains recovered from ancient sediments.

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2. MATERIALS AND BACKGROUND

84 Acapulcoites belong to the class of acapulcoite-lodranite (A-L) meteorites (McCoy *et al.*,
85 1993). A high degree of recrystallization and mineral chemical data indicate
86 formation of acapulcoites under redox conditions intermediate between those of H- and
87 E-chondrites, at about 1100°C, from which it cooled at a rate $> 10^{\circ}\text{C}/\text{Ma}$ (Dodd, 1969).
88 The major element composition is within the range of H-chondrites. Troilite and metal
89 are heterogeneously distributed. Associated trace elements are heterogeneously
90 distributed whereas chromium content is a factor of two higher than in H-chondrites.
91 Acapulcoites consist of an assemblage of orthopyroxene, olivine and feldspar associated
92 with comparatively large amounts of metallic nickel-iron, and minor amounts of
93 diopside, troilite, chromite and phosphate. The chromite is rather constant in
94 composition from grain to grain and appears to be distinct from chromite in ordinary
95 chondrites (Bunch *et al.*, 1967), silicate inclusions of iron meteorites (Bunch *et al.*,
96 1970), pallasites, mesosiderites, and eucrites (Bunch and Keil, 1971). The distribution
97 of divalent cations fits into the correlation observed for equilibrated meteorites. The

98 high concentrations of Cr, Mn and Zn in chromites are also observed in chromite from
99 silicate inclusions of iron meteorites. ZnO concentrations in acapulcoite chromites range
100 from 0.9 to 1.3%; these relatively high contents reflect the unusually high bulk Zn
101 content of acapulcoite (Palme *et al.*, 1981).

102 Kernouvé is an ordinary chondrite of the H group from an observed fall in France in
103 1869. It is texturally metamorphosed and of petrologic type 6. It contains abundant
104 metals, was heated for a significant period of time before the onset of partial melting,
105 and is minimally shocked (Ford *et al.*, 2008). It contains relict chondrules set in a
106 coarse-grained, recrystallized matrix of olivine (Fo₈₀), orthopyroxene (Fs_{15.7-18.1}),
107 plagioclase, Fe-Ni metal, troilite, chromite and phosphate (Ford *et al.*, 2008). Chromite
108 grains from this chondrite are chemically homogeneous and have the typical
109 equilibrated H-chondritic composition with generally somewhat lower TiO₂ (~2.5 wt%)
110 and higher Al₂O₃ (~6.0 wt%) contents than equilibrated L chondrites (typically TiO₂
111 ~3.0 wt% and Al₂O₃ ~ 5.2 wt%) (Bunch *et al.*, 1967; Schmitz *et al.*, 2001).

112 Meteorite Gol 001 (formal name Österplana 029) is a relatively large (6 x 9 x 2 cm)
113 mid-Ordovician fossil meteorite found in the Golvsten bed in the Thorsberg quarry in
114 southern Sweden (Schmitz *et al.*, 2001). The meteorite contains abundant, large and
115 well-preserved chromite grains. Chromite grains are homogeneous in chemical
116 composition throughout the meteorite, and the average composition of several of the
117 grains plots with equilibrated L chondrites (Schmitz *et al.*, 2001). Based on these data
118 and the maximum chromite grain diameter, Bridges *et al.* (2007) assigned it to the L6
119 group and type. Oxygen-three isotopic analyses of the chromite grains confirm an L or
120 LL chondritic origin (Greenwood *et al.*, 2007; Heck *et al.*, 2010). The cosmic-ray
121 exposure age of meteorite Gol 001 is ca. 300 kyr based on noble (neon) gas isotopic
122 measurements of chromite grains from the meteorite (Heck *et al.*, 2004). This age
123 concurs very well with the estimated timing of the break-up of the L chondrite parent-

124 body based on the stratigraphic first appearance of abundant sediment-dispersed
125 chromite grains from L-chondritic micrometeorites (Schmitz, 2013). Altogether the
126 isotopic and chemical data show that the chromite grains from this meteorite are
127 excellently preserved despite having been buried in marine limestone for 470 Ma.

128 3. METHODS

129 X-ray diffraction data for the nine chromite grains analyzed were recorded on an
130 automated KUMA-KM4 (K-geometry) diffractometer at the Department of
131 Mathematics and Geosciences (University of Trieste), using MoK α radiation,
132 monochromatized by a flat graphite crystal. Chromites from the studied meteorites are
133 usually rather small. The longer axis in these chromite grains is up to about 120 μm , but
134 usually below 100 μm , the other axis is between 20 and 50 μm so that in some cases the
135 grain is nearly "two-dimensional" with a thickness of only 20 μm . Given that, for the
136 largest grains, twenty-four equivalent reflections of (12 8 4) peak, at about 80° of 2 θ ,
137 were accurately centered at both sides of 2 θ , and the α_1 peak barycenter was used for
138 cell parameter determination. For the smallest grains, forty-eight reflections of (8 4 0)
139 and (8 4 4) peaks, at about 45-50° of 2 θ , have been used. Data collection was made,
140 according to Della Giusta *et al.* (1996), up to 50° of θ in the ω -2 θ scan mode, scan
141 width 1.8°2 θ , counting time from 20 to 50 seconds depending on the peak standard
142 deviation. Corrections for absorption and background were performed according to
143 North *et al.* (1968). Structural refinement using the SHELX-97 program (Sheldrick,
144 2008) was carried out against $F_o^2_{\text{hkl}}$ in the Fd-3m space group (with origin at -3*m*),
145 since no evidence of different symmetry appeared. Scattering factors were taken from
146 Prince (2004) and Tokonami (1965). Neutral scattering curves, Mg vs. Fe in T site and
147 Cr vs. Al in M site, were assigned, with the constraints of full site occupancy and equal
148 displacement parameters. Oxygen ionization was varied from one grain to another in

149 order to reach the best fit between structural refinement and chemical analyses and to
150 obtain the best value for all conventional agreement factors. Results are in Table 1.

151 After X-ray data collection, the same crystals used for X-ray data collection were
152 mounted on glass slides, polished and carbon coated for electron microprobe analyses
153 on a CAMECA-CAMEBAX microprobe at IGG-CNR, Padua, operating at 15kV and
154 15nA. A 20-s counting time was used for both peak and total background. Synthetic
155 oxide standards (MgO, Fe₂O₃, MnO, ZnO, NiO, Al₂O₃, Cr₂O₃, TiO₂ and SiO₂; purity
156 99.99 %) and synthetic chromite and Mg-chromite (Lenaz *et al.*, 2004b) were used.
157 Raw data were reduced by PAP-type correction software provided by CAMECA.
158 Results are in Table 2.

159 Several different procedures may be adopted to determine cation distribution, and
160 very satisfactory results have recently been obtained by combining data from single-
161 crystal X-ray structural refinements and electron microprobe analyses. This approach
162 simultaneously takes into account both structural and chemical data and reproduces the
163 observed parameters by optimizing cation distributions. Differences between measured
164 and calculated parameters are minimized by a function $F(X)$ taking in consideration
165 different parameters as the observed quantity and their standard deviations, cation
166 fractions in T and M sites, unit cell and oxygen parameter, mean atomic numbers of T
167 and M sites, atomic proportions given by microprobe analyses, and constraints imposed
168 by crystal chemistry (total charges and occupancies of T and M sites) (see Appendix).
169 Several minimization cycles of the equation containing these parameters were
170 performed until convergence according to the procedures described in Carbonin *et al.*
171 (1996) and Lavina *et al.* (2002) (Table 2 and Table 3). **We should note that the program**
172 **operates in order that the number of cations totals 3. As a result, the GOL samples have**
173 **large $F(X)$ values and errors because the number of cations never exceeds 2.985. A**
174 **similar situation arose with the Franzl chromite analyzed by Carbonin *et al.* (1999).**

175 Considering the possible valence state partitioning (Papike *et al.*, 2005) we tested the
176 effects of V₂O₅ instead of V₂O₃ without improvement. In Table 3 it is possible to
177 observe the observed and calculated values of parameters involved in the minimization
178 process.

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4. RESULTS AND DISCUSSION

181 The cell edge of chromites in H6 ranges from 8.3480 (2) to 8.3501 (1) Å, while in
182 GOL it varies from 8.3212 (3) to 8.3463 (3) Å (Table 1). The oxygen positional
183 parameter ranges from 0.2627 (2) to 0.26298 (9) in H6 and from 0.2624 (3) and 0.26267
184 (9) in GOL (Table 1). These values show that there are minor differences in chromite
185 composition and structure from the two types of chondrites. The ACAP chromite shows
186 a cell edge equal to 8.3384 (1) Å and an oxygen positional parameter of 0.26251 (9)
187 (Table 1).

188 The chemical composition of H6 and ACAP samples is rather homogeneous while
189 GOL chromites present different compositions from one point to the other. This feature
190 creates patch-like surfaces with local enrichment/depletion in major oxides. The
191 analyzed chromites point to a nearly pure end-member composition with limited Al↔Cr
192 (Al < 0.26 apfu) and Mg↔Fe²⁺ (Mg ≤ 0.15 apfu, except for the ACAP sample)
193 substitutions. Ferric iron is negligible in all the analyzed chromites (Table 2), a feature
194 characteristic of other phases too, in extraterrestrial materials (Papike *et al.*, 1998). All
195 the GOL chromites here analyzed have been assigned to the L-chondrite group (TiO₂ in
196 the range 2.9 – 3.3 wt.%) (Table 2).

197 The large excess octahedral crystal field stabilization energy of Cr³⁺ (Δ CFSE_(oct-tet) is
198 about 160 kJ mol⁻¹; O'Neill and Navrotsky, 1984) should ensure that Cr-bearing spinels
199 have an almost completely normal cation distribution (Urusov, 1983), meaning that
200 about ¾ of the M site is filled by Cr. Titanium and vanadium can also be assumed to

201 occupy this site, consequently only about 0.3 cation could be filled by other cations.
202 Cation assignment, taking in account structural and chemical parameters, shows an
203 ordered distribution where Al almost completely fills the balance of the M site (Table
204 2).

205 Calculation of the formulae for the chromites in the GOL samples from Kinnekulle
206 gives a total number of cations less than 3 and a total number of electrons per formula
207 unit significantly higher than that derived from structural refinement. In terrestrial
208 samples this fact, coupled with a low oxygen parameter value (< 0.2620), suggested the
209 presence of vacancies (Carbonin *et al.*, 1999; Bosi *et al.*, 2004; Lenaz *et al.*, 2014a,
210 2014c). The low oxygen positional parameter in terrestrial samples (in the range
211 $0.2608-0.2620$) is a product of the $3\text{Fe}^{2+} \rightarrow 2\text{Fe}^{3+} + \text{vacancies}$ transformation under
212 oxidizing conditions. This causes the development of a magnetite component as seen
213 from a combined X-ray single crystal diffraction and Mössbauer (MS) approach (Lenaz
214 *et al.*, 2014a, 2014c) and exemplified by Lenaz *et al.* (2013) by comparing the results of
215 structural refinement (SREF), microprobe analyses, point-MS and powder-MS collected
216 on several chromite samples. They showed that in some cases there was a large
217 discrepancy between SREF and powder-MS because commonly used powder-MS needs
218 a lot of grains where oxidation degree could be very different among the grains, while
219 SREF and point-MS have been performed on the same single crystal yielding a good
220 comparison. These phenomena in the terrestrial environment are considered as the result
221 of oxidation processes that happened after primary chromite formation at temperatures
222 between $600-700^{\circ}\text{C}$. It is unlikely this oxidation processes in the present case took place
223 on the sea floor or during diagenesis, consequently we think that this non -
224 stoichiometry should be primary. Alwmark and Schmitz (2007) noticed in some
225 extraterrestrial chromites from the Lockne crater in central Sweden, Zn enrichment and
226 oxidation that they suggested to be a result of the hydrothermal system induced by the

227 impact. In the present case there is undoubtedly a non - stoichiometry, but the absence
228 of Zn enrichment as well as the oxygen parameters higher than 0.2624 suggest that the
229 non-stoichiometry is not due to oxidation. Moreover, this feature has been already
230 noticed in other extraterrestrial mineral phases (Weill *et al.*, 1971; Herd and Papike,
231 1998; Yurimoto *et al.*, 2001; Abreu *et al.*, 2014). Yurimoto *et al.* (2001) suggested a
232 rapid crystallization to explain the non - stoichiometry of some spinels from a CO
233 chondrite, while Abreu *et al.* (2014) suggested shock metamorphism for non-
234 stoichiometry in CR chondrite **grains of several unknown ferromagnesian silicates**
235 ($\text{Fe}_5\text{MgSi}_2\text{O}_{10}$, $\text{Fe}_4\text{MgSi}_3\text{O}_{11}$ and other similar compounds). We suggest that the spinels
236 studied here present a non - stoichiometry for similar reasons.

237 By comparison with the structural parameters of chromites from terrestrial occurrences,
238 in an oxygen positional parameter vs. cell edge diagram (Fig. 1) the extraterrestrial
239 chromites fall in a field close to that of chromites from komatiites (Lenaz *et al.*, 2004a),
240 kimberlites and included in diamonds (Lenaz *et al.*, 2009), and not far from that of
241 layered intrusions (Lenaz *et al.*, 2007, 2012) and the synthetic MgCr_2O_4 - FeCr_2O_4 series
242 (Lenaz *et al.*, 2004b). When comparing the meteoritic spinels studied by us with the
243 mantle occurrences from a structural point of view we can see that they fall in a field
244 more or less parallel to that of the MgCr_2O_4 - FeCr_2O_4 binary synthetic series (Lenaz *et*
245 *al.*, 2004b). Considering that the MgAl_2O_4 and FeAl_2O_4 spinels have cell edges equal to
246 8.0855 and 8.1646 Å and oxygen positional parameter equal to 0.2613 and 0.2642,
247 respectively, changes along the x-axis roughly suggest a $\text{Cr} \leftrightarrow \text{Al}$ substitution (see also
248 Lenaz and Skogby, 2013, for the hercynite-chromite synthetic series) while changes
249 along the y-axis roughly suggest a $\text{Mg} \leftrightarrow \text{Fe}^{2+}$ (Fig. 2). Our data suggest a limited
250 $\text{Cr} \leftrightarrow \text{Al}$ substitution (Al_2O_3 is close to 6.5 wt. % for H6 and to 5.8 wt. % for GOL) and a
251 larger $\text{Mg} \leftrightarrow \text{Fe}^{2+}$ (MgO ranges from 2 to 7 wt. %). Therefore the chromites in our
252 meteorites (excluding the non-stoichiometric ones), those included in diamonds (Lenaz

253 *et al.*, 2009) and those from komatiites (Lenaz *et al.*, 2004a) are on a trend ($R^2=0.92$).
254 As all of these occurrences are mantle related, it is possible that this is a consequence of
255 the degree of differentiation of the liquid from which they originated.

256 Hazen and Ferry (2010) defined three eras and ten stages of mineral evolution in our
257 solar system. During the first era, the so-called Era of Planetary Accretion (>4.55 Ga),
258 perhaps 60 different mineral species appeared as primary condensates. These planet-
259 forming materials developed as planetesimals. As asteroids grew and began to
260 experience modest heating by the decay of short-lived radionuclides such as ^{26}Al , the
261 melting of ice and the subsequent alteration at low temperatures increased mineral
262 diversity to 250 (McCoy, 2010). In fact, one of the many mechanisms that drove
263 mineral diversification is increase in the ranges of intensive variables such as pressure,
264 temperature and the activities of H_2O , CO_2 and O_2 (Hazen and Eldredge, 2010). **Even**
265 **while this aqueous alteration and thermal metamorphism were occurring in asteroids, an**
266 **era of intense bombardment began. These impacts formed new minerals as chondritic**
267 **asteroids were subjected to intense heat and pressure (McCoy, 2010).** As heating
268 continued, the solidus temperature of about 950°C was exceeded **and melting began.**
269 **The earliest partial melts formed from FeNi metal and the iron sulfide troilite, but they**
270 **also incorporated minerals like chromite and phosphates (McCoy, 2010).** By using the
271 geothermometer proposed by Princivalle *et al.* (1999) an intracrystalline equilibration
272 temperature equal or higher than about 1250°C (Table 2) has been found for H6 and
273 ACAP chromite grains, showing that the temperature reached after heating was very
274 high. Calculated temperatures for GOL chromites are unreliable ($> 1700^\circ\text{C}$). This
275 happened also for some terrestrial occurrences (Derbyshire *et al.* 2013) where zoning is
276 present, **so that it is possible to assume that where well-defined core-rim zoning or**
277 **patches are present, as in the GOL chromites, it is not possible to use the thermometer.**
278

279

5. CONCLUSIONS

280 Because meteoritic material on the Earth surface is rapidly altered or weathered away
281 with the exception of chromite, similarly to what happens for terrestrial mafic –
282 ultramafic material (Lenaz *et al.*, 2009, 2014b), different methodologies have been
283 recently developed to recognize the extraterrestrial detrital chromitic material such as
284 relict silicate inclusions (Alwmark and Schmitz, 2009) and high-precision oxygen three-
285 isotopes SIMS analyses (Heck *et al.*, 2010). This work demonstrates that a structural
286 study can, as well, distinguish extraterrestrial from terrestrial material and also that
287 different chondritic (H and L) origins can be recognized. This will facilitate ongoing
288 attempts to reconstruct the paleoflux of meteorites to Earth from resistant extraterrestrial
289 spinel grains recovered from ancient sediments.

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APPENDIX

300 *I) Cation distribution*

301 The cation distribution for the present samples has been achieved by using the Lavina *et*
302 *al.* (2002) model. This model yields cation distribution by minimizing the weighted
303 differences between observed crystal chemical data and data calculated from site atomic

304 fractions. This cation distribution in the tetrahedral (T) and octahedral (M) sites must be
 305 consistent with the assumptions that:

306 1 – The mean atomic number (m.a.n.) corresponds to:

$$307 \quad \text{m.a.n. T} = \sum_i {}^{\text{IV}}X_i N_i \quad (\text{A1})$$

$$308 \quad \text{m.a.n. M} = \sum_i {}^{\text{VI}}X_i N_i \quad (\text{A2})$$

309 where ${}^{\text{IV}}X_i$ and ${}^{\text{VI}}X_i$ are chemical species in T and M respectively and N is their atomic
 310 number

311 2 – The site bond length arises from a linear contribution of each species to the
 312 tetrahedral (T-O) and octahedral (M-O) coordination distances so that:

$$313 \quad \text{T-O} = \sum_i {}^{\text{IV}}X_i {}^{\text{IV}}D_i \quad (\text{A3})$$

$$314 \quad \text{M-O} = \sum_i {}^{\text{VI}}X_i {}^{\text{VI}}D_i \quad (\text{A4})$$

315 where ${}^{\text{IV}}D_i$ and ${}^{\text{VI}}D_i$ are the cation-to-oxygen bond distances of each cation in tetrahedral
 316 and octahedral coordination, respectively.

317 To summarize, site atomic fractions ${}^{\text{IV}}X_i$ and ${}^{\text{VI}}X_i$ must not only satisfy the above
 318 equations, but also correspond to the atomic proportions from the chemical analyses and
 319 obey three crystal-chemical constrains: occupancies of T and M sites and formal
 320 valence. Consequently, ${}^{\text{IV}}X_i$ and ${}^{\text{VI}}X_i$ may be calculated by minimizing the following
 321 sum of residuals:

$$322 \quad F(X_i) = \frac{1}{n} \sum_{j=1}^n \left(\frac{O_j - C_j(X_i)}{\sigma_j} \right)^2 \quad (\text{A5})$$

323 Where O_j are the observed quantities with their standard deviation σ_j . O_j are the four
 324 observed crystallographic parameters (a , u and m.a.n. of T and M sites) and the
 325 chemical proportions for a total of n . $C_j(X_i)$ are the corresponding quantities calculated
 326 by means of variable cation fractions X_i .

327

328 *II) Temperature calculation*

329 Della Giusta *et al.* (1996) fitted the data of their heating experiments on spinels with the
330 following empirical equation

$$331 \quad T(^{\circ}\text{C}) = C1 - C2 \cdot B + C3 \cdot B^2 \quad (\text{A6})$$

332 where

$$333 \quad B = \text{Al}(\text{T})/\text{Al}_{\text{tot}} + C4 \cdot (1 - \text{Mg}(\text{T}) - \text{Al}(\text{T}) + C5 \cdot (2 - (\text{Al}(\text{M}) - \text{Mg}(\text{M}))) \quad (\text{A7})$$

334 and C1 to C5 are the fitting coefficients. The second term of A7 takes into account the
335 compositional influence of ($\text{Fe}^{2+} + \text{Fe}^{3+} + \text{Si} + \text{Mn} + \text{Zn}$) in the T site, and the third term
336 ($\text{Cr} + \text{Fe}^{2+} + \text{Fe}^{3+} + \text{Ti} + \text{Ni}$) in the M site. Princivalle *et al.* (1999) revised the above
337 equations obtaining the linear equation

$$338 \quad T(^{\circ}\text{C}) = 6640 \cdot B \quad (\text{A8})$$

339 with new coefficients C4 and C5 being 0.101 and 0.041, respectively. Obtained
340 temperatures for the **chromites studied by us** are in Table 2. As happened for severely
341 altered terrestrial chromites (Derbyshire *et al.*, 2013), the temperatures of non-
342 stoichiometric chromites are unreliable and, consequently, not reported.

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530 Figure captions

531 Fig. 1: Oxygen positional parameter, u vs. cell edge, a . Red circles: H6, this study; red
532 squares: GOL, this study; black diamond: ACAP chromite, this study; blue
533 diamonds: chromite in kimberlites and included in diamonds (Lenaz *et al.*, 2009);
534 purple squares: chromites in komatiites (Lenaz *et al.*, 2004a); yellow circles:
535 chromites in layered intrusions (Lenaz *et al.*, 2007; 2012); green circles: MgCr₂O₄-
536 FeCr₂O₄ synthetic spinels (Lenaz *et al.*, 2004b).

537 Fig. 2: Oxygen positional parameter, u vs. cell edge, a for synthetic series and the
538 meteoritic chromites studied here. Blue squares: MgAl₂O₄-FeAl₂O₄ (Andreozzi and
539 Lucchesi, 2002); orange diamonds: FeAl₂O₄-FeCr₂O₄ (Lenaz and Skogby, 2013);
540 green circles: MgCr₂O₄-FeCr₂O₄ spinels (Lenaz *et al.*, 2004b). The **meteorite**
541 **chromites** of this study are represented by red circles.

Table 1: Results of crystal structure refinements of chromite

Sample	H6-1	H6-2	H6-C	H6-D	ACAP	GOL-A	GOL-B	GOL-C	GOL-D
a (Å)	8.3488 (1)	8.3501 (1)	8.3480 (2)	8.3485 (2)	8.3384 (1)	8.3463 (3)	8.3349 (3)	8.3347 (3)	8.3212 (3)
<i>u</i>	0.26286 (7)	0.26291 (7)	0.26298 (9)	0.2627 (2)	0.26251 (9)	0.26267 (9)	0.2625 (1)	0.2626 (1)	0.2624 (3)
T-O	1.994 (1)	1.995 (1)	1.995 (1)	1.992 (3)	1.987 (1)	1.990 (1)	1.985 (2)	1.989 (2)	1.980 (5)
M-O	1.9856 (5)	1.9856 (5)	1.9845 (7)	1.986 (2)	1.9854 (6)	1.9864 (7)	1.9852 (8)	1.986 (2)	1.983 (2)
m.a.n.T	23.7 (4)	24.0 (6)	24.2 (5)	24.5 (6)	21.3 (2)	23.3 (3)	22.0 (4)	24.0 (5)	22.1 (5)
m.a.n.M	22.1 (5)	22.1 (7)	22.5 (6)	22.0 (5)	22.5 (3)	22.4 (3)	21.9 (6)	22.6 (5)	22.3 (5)
U (M)	0.0038 (1)	0.0039 (1)	0.0045 (1)	0.0039 (2)	0.0045 (1)	0.0046 (1)	0.0059 (1)	0.0052 (1)	0.0059 (3)
U (T)	0.0068 (1)	0.0071 (2)	0.0075 (2)	0.0073 (3)	0.0073 (1)	0.0073 (1)	0.0081 (2)	0.0082 (2)	0.0086 (5)
U (O)	0.0059 (2)	0.0060 (3)	0.0060 (3)	0.0070 (5)	0.0063 (2)	0.0068 (2)	0.0084 (3)	0.0072 (4)	0.0085 (9)
N. refl.	155	152	122	117	177	155	127	118	85
R1	2.01	1.86	1.80	3.01	1.67	2.04	2.23	2.43	3.55
wR2	2.77	3.11	2.67	6.65	3.31	3.62	3.34	4.64	5.50
GooF	1.241	1.088	1.119	1.173	1.154	1.258	1.145	1.137	1.296
Diff. peaks	2.91; -0.99	1.63; -0.64	1.12; -0.89	2.26; -1.53	1.74; -0.84	2.33; -1.29	1.62; -0.61	2.28; -0.63	1.75; -1.21

a: cell parameter (Å); *u*: oxygen positional parameter; T-O and M-O: tetrahedral and octahedral bond lengths (Å), respectively; m.a.n.T and M: mean atomic number; U(M), U(T), U(O): displacement parameters for M site, T site and O; N. Refl.: number of unique reflections; R1 all (%), wR2 (%), GooF as defined in Sheldrick (2008). Diff.peaks: maximum and minimum residual electron density ($\pm e/\text{Å}^3$). Space Group: Fd-3m. Origin fixed at $-3m$. Z=8. Reciprocal space range: $-19 \leq h \leq 19$; $0 \leq k \leq 19$; $0 \leq l \leq 19$. Estimated standard deviations in *brackets*.

Table 2. Mean chemical analyses and cation distribution of chromite

Sample	H6-1	H6-2	H6-C	H6-D	ACAP	GOL-A	GOL-B	GOL-C	GOL-D
MgO	3.35 (8)	3.3 (1)	3.2 (1)	3.0 (1)	6.9 (3)	2.3 (2)	2.9 (3)	2.0 (1)	2.67 (9)
Al ₂ O ₃	6.5 (2)	6.4 (1)	6.46 (8)	6.7 (1)	6.1 (1)	5.9 (1)	5.8 (1)	5.8 (3)	5.78 (4)
TiO ₂	2.3 (2)	2.32 (7)	2.29 (4)	2.10 (7)	1.18 (4)	3.08 (4)	3.2 (1)	2.9 (1)	3.32 (3)
V ₂ O ₃	0.74 (6)	0.67 (5)	0.66 (3)	0.71 (6)	0.57 (3)	0.67 (1)	0.73 (5)	0.65 (5)	0.69 (1)
Cr ₂ O ₃	56.9 (4)	56.9 (5)	57.2 (1)	56.4 (5)	61.7 (5)	57.0 (7)	57.2 (9)	57.0 (8)	56.62 (6)
MnO	0.91 (6)	0.87 (6)	0.95 (7)	0.84 (4)	1.6 (5)	0.68 (7)	0.71 (6)	0.57 (6)	0.93 (6)
FeO	28.7 (3)	29.1 (3)	29.0 (2)	29.2 (2)	21.1 (4)	29.5 (1.0)	29.0 (1.0)	28.9 (1.5)	28.6 (3)
NiO*	0.03 (3)	0.02 (3)	0.00 (0)	0.01 (1)	0.01 (1)	0.01 (1)	0.01 (1)	0.01 (1)	0.01 (1)
ZnO*	0.37 (2)	0.36 (5)	0.31 (3)	0.33 (4)	1.0 (1)	0.30 (3)	0.33 (6)		0.7 (1)
Sum	99.74	99.92	100.00	99.19	100.13	99.39	99.23	98.8	99.3
T Site									
Mg	0.152 (4)	0.107 (6)	0.110 (4)	0.101 (6)	0.32 (1)	0.105 (9)	0.16 (1)	0.07 (1)	0.089 (4)
Al	0.009 (1)	0.013 (1)	0.0090 (4)	0.020 (1)	0.017 (1)	0.027 (2)	0.0001 (1)	0.037 (5)	0.0702 (9)
Mn	0.027 (2)	0.026 (2)	0.027 (2)	0.025 (1)	0.05 (1)	0.020 (2)	0.022 (2)	0.017 (2)	0.028 (2)
Fe ²⁺	0.803 (6)	0.843 (8)	0.845 (5)	0.844 (7)	0.59 (1)	0.82 (2)	0.51 (1)	0.86 (3)	0.783 (6)
Zn	0.009 (1)	0.009 (1)	0.009 (1)	0.009 (1)	0.025 (3)	0.0079 (8)	0.010 (2)		0.018 (4)
Vac.						0.0184 (5)	0.13 (1)	0.0114 (8)	0.0104 (8)
M Site									
Mg	0.027 (2)	0.067 (5)	0.058 (3)	0.057 (4)	0.033 (4)	0.022 (4)		0.040 (7)	
Al	0.260 (5)	0.252 (5)	0.260 (2)	0.262 (4)	0.231 (4)	0.225 (6)	0.230 (6)	0.22 (1)	0.174 (1)
Ti	0.060 (3)	0.062 (2)	0.061 (1)	0.056 (2)	0.031 (1)	0.082 (2)	0.082 (3)	0.078 (3)	0.0889 (8)
V	0.021 (2)	0.019 (1)	0.0187 (8)	0.020 (2)	0.0157 (8)	0.0191 (4)	0.020 (1)	0.019 (2)	0.0197 (3)
Cr	1.593 (8)	1.588 (9)	1.592 (5)	1.584 (9)	1.68 (1)	1.61 (2)	1.52 (2)	1.61 (3)	1.609 (5)
Fe ²⁺	0.038 (1)	0.0076 (1)	0.0090 (5)	0.018 (1)	0.012 (1)	0.044 (5)		0.020 (5)	0.049 (1)
Vac.							0.006 (2)	0.0086 (7)	0.0050 (6)
T (°C)	1245	1374	1265	1512	1377				

T (°C) is the intracrystalline temperature calculated by using the thermometer of Princivalle et al. (1999). *Ni and, occasionally, Zn not present in cation distribution because of the deviation higher than 2σ . Estimated standard deviations are in brackets. High F(X) values are caused by the non-stoichiometry of the spinels with a number of cations in the range 2.980-2.984. Temperatures for non-stoichiometric chromites are unreliable and, consequently, not reported (see Appendix for details).

Table 3. Comparison of observed number of electrons, cell edge a and oxygen positional parameter u with corresponding values calculated with the MINUIT program.

Sample	H6-1	H6-2	H6-C	H6-D	ACAP	GOL-A	GOL-B	GOL-C	GOL-D
e^-T_{obs}	23.7 (4)	24.0 (6)	24.2 (5)	24.5 (6)	21.3 (2)	23.3 (3)	21.9 (4)	24.0 (5)	22.1 (5)
e^-T_{calc}	23.7	24.3	24.3	24.3	21.3	23.7	20.5	24.2	23.6
e^-M_{obs}	22.1 (5)	22.1 (7)	22.5 (5)	22.0 (5)	22.5 (3)	22.4 (3)	21.9 (6)	22.6 (6)	22.3 (5)
e^-M_{calc}	22.4	22.1	22.1	22.2	22.5	22.5	22.8	22.4	22.6
a_{obs}	8.3488 (1)	8.3501 (1)	8.3480 (2)	8.3485 (2)	8.3384 (1)	8.3463 (3)	8.3349 (3)	8.3439 (3)	8.3212 (3)
a_{calc}	8.3488	8.3501	8.3480	8.3485	8.3384	8.3463	8.3348	8.3439	8.3215
u_{obs}	0.26286 (7)	0.26291 (7)	0.26298 (9)	0.2627 (2)	0.26251 (9)	0.26267 (9)	0.2625 (1)	0.2626 (1)	0.2624 (2)
u_{calc}	0.26284	0.26287	0.26294	0.2627	0.26248	0.26266	0.2620	0.2626	0.2622
F(X)	0.16	0.14	0.22	0.72	0.09	0.54	4.08	0.51	8.94

F(x): minimisation factor, which takes into account the mean of square differences between calculated and observed parameters, divided by their standard deviations.



