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High-temperature structure of Co_{3}O_{4}: Understanding spinel inversion using in situ and ex situ measurements

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1	The high-temperature structure of Co_3O_4 : Understanding spinel inversion using <i>in</i>					
2	$situ$ and $ex\ situ$ measurements					
2	Taylor D. Sparks*					
3	Department of Materials Science and Engineering					
4 6	University of Utab Solt Lake City Utab 8/112 USA					
5	Chebelisting of Count, Such Dance Court, Otanie, 04112, OBM					
6	Aleksander Gurlo					
7	Institute for Material Science and Technologies,					
8 Technische Universität Berlin, D-10587 Berlin, Germany						
9	Maged F. Bekheet					
10	Fachaebiet Keramische Werkstoffe. Institut für Werkstoffwissenschaften und -technologien.					
11	Technische Universität Berlin, Hardenbergstrae 40, 10623 Berlin, Germany					
12	Michael W. Gaultois					
13	Leverhulme Research Centre for Functional Materials Design,					
14	The Materials Innovation Factory, Department of Chemistry,					
15	University of Liverpool, 51 Oxford Street, Liverpool, L7 3NY, United Kingdom.					
16	Gennady Cherkashinin					
17	Fachbereich Material- und Geowissenschaften Fachgebiet Oberflächenforschung,					
18	Technische Universität Darmstadt, Darmstadt, 64287, Germany					
19	Laetitia Laversenne					
20	Université Grenoble Alpes, CNRS, Grenoble INP, Institut Néel, 38000 Grenoble, France					
21	David R. Clarke					
22	School of Engineering and Applied Sciences,					
23	Harvard University, Cambridge,					
24	Massachusetts, 02138, USA					
25	(Dated: February 1, 2019)					
26	In this paper we investigate the high-temperature structure of Co_3O_4 , a compound that has been					
27	studied extensively over the last 60 years due to its unresolved high-temperature structure. In					
28	situ thermal analysis and X-ray diffraction confirm previously reported high-temperature structural					
29	changes and show that these changes are unrelated to the high-temperature decomposition to CoO.					
30	Raman active peaks are also extinguished over the same temperature range. By considering the					
31	changing lattice parameter, A-O, and B-O bond lengths as well as cation size we are able to cal-					
32	culate degree of inversion which reaches a maximum of 0.6. To further study the structure in this					
33	experimentally inaccessible range we quench samples and perform <i>ex situ</i> measurements including					
34	redox titration, X-ray photoelectron spectroscopy, and neutron diffraction. We do not observe any					
35	evidence of large oxygen vacancy concentrations or octahedral $\operatorname{Co}_B^{\to \pm}$ ions with high spin state. How-					
36	ever, we do show an evolution in the magnetic moment from magnetic structure refinement from $(2,4)$ by $(2,7)$ by the second structure refinement from $(2,4)$ by $(2,7)$ by the second structure refinement from $(2,4)$ by $(2,7)$ by the second structure refinement from $(2,4)$ by $(2,7)$ by the second structure refinement from $(2,4)$ by $(2,7)$ by the second structure refinement from $(2,4)$ by $(2,7)$ by the second structure refinement from $(2,4)$ by $(2,7)$ by the second structure refinement from $(2,4)$ by $(2,7)$ by the second structure refinement from $(2,4)$ by $(2,7)$ by the second structure refinement from $(2,4)$ by $(2,7)$ by the second structure refinement from $(2,4)$ by $(2,7)$ by the second structure refinement from $(2,4)$ by $(2,7)$ by the second structure refinement from $(2,4)$ by $(2,7)$ by the second structure refinement from $(2,4)$ by $(2,7)$ by the second structure refinement from $(2,4)$ by $(2,7)$ by the second structure refinement from $(2,4)$ by $(2,5)$ by the second structure refinement from $(2,4)$ by $(2,5)$					
37	$(2.4 \mu_{\rm B})$ to $(2.7 \mu_{\rm B})$ that coincides exactly with the high-temperature anomaly and suggests partial					
38	inversion (0.40) of the spinel structure in fairly good agreement with the inversion calculated from					
39	bolid lengths.					

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I.

Cobalt oxide spinel, Co₃O₄, is a well-known antiferro-⁵³ 41 magnetic oxide investigated most recently as a promis- 54 42 ing catalyst for oxygen evolution reactions in artifi-55 43 cial photosynthesis.^{1,2} The compound crystallizes in the ⁵⁶ spinel crystal structure featuring twice as many six-57 coordinate (octahedral) cation sites as four-coordinate 58 (tetrahedral) sites. In order to maintain charge neutral-⁵⁹ ity with the oxygen anions, there must be two Co^{3+} ions 60 and one Co^{2+} ion per formula unit. 61

INTRODUCTION

How the two different ions arrange themselves over the tetrahedral cation sites (often called the A site) or octahedral cation sites (often called the B site) dictates many important electrical and magnetic properties in spinels. The low-temperature structure of Co_3O_4 is considered the archetypal *normal* spinel because the Co^{2+} ions completely occupy the tetrahedral A site and Co^{3+} ions completely occupy the octahedral B site. Neutron diffraction and magnetic susceptibility measurements suggest the d⁶ electrons from Co^{3+} pair up to completely fill the octa-hedral t_{2g} triplet, whereas the d⁷ electrons from Co^{2+} on the tetrahedral site leave three unpaired electrons on t_2

⁶² orbitals that account for the observed magnetic moment¹¹⁷ ⁶³ of $2.6 \,\mu_{\rm B}$ to $3.25 \,\mu_{\rm B}$.^{3–5} ¹¹⁸

In contrast, the details of the high-temperature crys-119 64 tal structure of Co_3O_4 are not well understood and have¹²⁰ 65 been an active area of research. A variety of obser-121 66 vations including a sudden non-linear increase in lat-122 67 tice parameter,^{6–8} excess thermal expansion,⁹ electromo-123 68 tive force (EMF) measurements on the oxygen partial¹²⁴ 69 pressure, 7,10 opposite changes in the A–O and B–O bond¹²⁵ 70 lengths with increasing temperature,⁸ electrical conduc-126 71 tivity and thermopower,^{9,11} and a phase transformation from heat capacity measurements,¹² all indicate a high-72 73 temperature structural anomaly. 74 127 The exact origin of the high-temperature structural

75 anomaly is obfuscated by two potential mechanisms as $_{128}$ 76 well as the incipient decomposition to rock-salt type CoO_{129} 77 (1165 K to 1270 K).⁷ One mechanism is cation anti-site¹³⁰₁₃₀ 78 disorder, termed *inversion* of the spinel structure where $\frac{1}{131}$ 79 Co^{3+} ions sit on the tetrahedral A site and Co^{2+} ions on 80 the octahedral B site. The other mechanism is a low-spin 81 (LS) to high-spin (HS) spin state transition of Co^{3+} ions³³ 82 on the octahedral sites between 1000 K and 1200 K that 83 has been suggested by many researchers. 7,8,10,12 There is $^{139}_{136}$ 84 currently no consensus on how much, if any, cation anti-85 site disorder is present. Chen¹³ summarizes that some $_{136}^{137}$ 86 authors suggest a complete inversion of the structure 7^{130}_{139} 87 (*i.e.* Co_A^{3+} and $\operatorname{Co}_B^{2+}\operatorname{Co}_B^{3+}$), while others favor complete ¹³⁹ anti-site disorder,⁸ and still others propose a retention of the normal spinel structure with only 5–10% disorder.¹²¹¹ 88 89 90

High-temperature in situ magnetic susceptibility, X_{-143}^{-142} 91 ray photoelectron spectroscopy (XPS), and neutron 92 diffraction measurements to settle the ambiguity $\operatorname{are}_{_{145}}$ 93 hampered by the low Néel temperature ${\approx}40\,{\rm K}$ of ${\rm Co_3O_4}_{_{146}}$ 94 and lack of high-temperature instrumentation. Never-147 95 the less, there are several in situ techniques remaining to $_{148}$ 96 study the high-temperature structure of Co_3O_4 includ-97 ing Raman spectroscopy and thermogravimetric analysis 98 (TGA) discussed in this work, and thermopower which will be discussed in detail in a subsequent contribution.¹⁴ 99 100 152 Furthermore, in this contribution we attempt to $\operatorname{preserve}_{\scriptscriptstyle 153}$ 101 the high-temperature structure via quenching and then $_{154}$ 102 use ex situ XPS, neutron diffraction, and redox titra-103 tion measurements on the samples to gain additional in- $_{156}$ 104 sight. Finally, in situ phase composition is monitored $_{157}$ 105 via spectroscopy and diffraction to ensure that the high-106 temperature anomaly is unrelated to the transformation₁₅₉ 107 from Co_3O_4 to CoO. 108 160

II. EXPERIMENTAL

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Disc-shaped pellets were fabricated from nanometer-165 sized Co_3O_4 powder (Sigma Aldrich, 99.9 wt.%, 50 nm166 particles). The powders were first cold-pressed with a167 6 ton load in a $\frac{1}{2}$ inch die and then sintered in air.168 Pure phase, dense samples of Co_3O_4 and CoO were ob-169 tained by firing the samples from room temperature up170 to 1148 K over 2 hours and holding for 6 hours, ramping171 to 1373 K over 1 hour and holding for 6 hours, ramping back to 1148 K over 2 hours and holding for 2 hours. From this point, samples were quenched in air to obtain phase-pure Co₃O₄ or slowly cooled at 1 K/min to obtain phase-pure Co₃O₄. Intermediate cooling rates resulted in mixtures of CoO and Co₃O₄ phases. *Ex situ* magnetic structure determination, XPS, and redox titration measurements were performed on samples that were quenched from high-temperature by submersion in liquid nitrogen.

A. Property measurement

X-ray diffraction from room temperature up to 1273 K was performed in air using a PANalaytical XPert Pro diffractometer in continuous-scanning mode at 40 kV 45 mA with a vertical $\theta - \theta$ scanning geometry from 20° to 100° with a step size of 0.017° and a rate of 1.40 s/step. The height of the sample in the furnace was zeroed by breaking the beam and tilt was corrected by rocking the sample. Iron-filtered Co K α radiation was used, except for the room temperature XRD of CoO quenched and intermediate cooled samples, where nickel-filtered Cu K α radiation was used. Measurements at temperature were made with an Anton-Paar HTK1200N furnace to heat the samples for high-temperature data collection. The temperature fluctuation within the furnace was less than 1 K, and the sample was heated slowly at 5 K/min and allowed to equilibrate 30 minutes before each scan was performed. Measurements were performed at 300 K, 800 K, 875 K, 950 K, 1025 K, 1100 K, 1140 K, 1163 K and 1273 K, and an additional measurement was made after cooling back to 300 K.

Raman spectra were obtained as a function of temperature in air using a confocal LabRAM ARAMIS Raman Microscope (HORIBA Scientific) with the sample in a Linkam heating stage (Scientific Instruments Ltd., Waterford Surrey, England). The spectra were excited with a 515 nm laser (Kimmon Electric US, Ltd. IK Series He-Cd LASER Englewood, USA). Spectra were recorded from room temperature (298 K) up to a temperature of 1223 K and back down to room temperature and with a constant heating and cooling rate (10 K/min). Spectral analysis such as baseline adjustment, smoothing and normalization, was performed using GRAMS software package (Galactic Industries Corporation Salem, USA).

Neutron diffraction measurements were performed on the two-axis diffractometer D1B at the Institut Laue Langevin (ILL, Grenoble, France). Powdered sample was loaded into a vanadium can and placed in a cryostat; diffractograms were recorded with a wavelength of $\lambda = 2.52$ Å.

Nuclear structure determination and refinement was performed using the full-pattern Rietveld method based on least-squares refinement in the GSAS and EXPGUI software package.^{15,16} Magnetic structure determination was performed with the full-pattern Rietveld method us-

ing FullProf and BASIREPS software packages.^{17,18} The 172 magnetic structure was refined based on the known struc-173 ture previously determined by Roth.⁴ Heat capacity mea-174 surements performed elsewhere exhibit a sharp, lambda-175 type anomaly at the Neél temperature,^{19,20} suggesting 176 the transition is strongly second-order. Accordingly, the 177 space group above the ordering temperature was used for 178 refinement of the magnetic structure. 179

Thermogravimetry (TG) - Differential Thermal Anal-180 vsis (DTA) characterization was performed on TGA 181 Q5000 (TA Instruments, New Castle, USA) on pow-182 dered samples in dynamic conditions (synthetic air or Ar, 183 $10 \,\mathrm{ml/min}$) with a heating and cooling rate of $10 \,\mathrm{K/min}$. 184 Finely ground powders of Co₃O₄ annealed and 185 quenched from 300 K, 773 K, and 1173 K were pressed 186 into indium foil, mounted on an electrically grounded 187 sample holder, and analyzed with a chamber base pres-188 sure of $\sim 10^{-9}$ mbar. XPS spectra were collected using 189 a PHI 5000 VersaProbe spectrometer and a monochro-190 matic AlK α (1486.7 eV) X-ray source; the diameter of 191 the analysis area was $\sim 100 \,\mu \text{m}$, and the electron escape 192 angle was fixed at 45° . High-resolution photoelectron 193 spectra of the Co₂p, Co₃s, O₁s and C₁s core-levels were 194 collected using a pass energy of 11.75 eV and a step size 195 of $0.05 \,\mathrm{eV}$. At this pass energy, the resolution is less than 196 $0.6 \,\mathrm{eV}$, determined by analysis of the Ag $3d_{5/2}$ FWHM 197 of a clean Ag foil; the precision of the measured bind-198 ing energies (BE) is better than $\pm 0.10 \,\text{eV}$. A dual-beam 199 charge neutralizer was used to prevent surface charging 200 during measurements; the system is equipped with a low 201 voltage electron gun and a floating ion gun generating 202 low energy ion beam. All spectra were calibrated using 203 the C1s core-line arising from adventitious carbon with 204 a fixed value of 284.8 eV, and a Shirley-type function was₂₂₇ 205 applied to remove the background arising from electron₂₂₈ 206 energy loss. 207 229

The mean oxidation state of cobalt was determined by₂₃₀ 208 the redox titration method adapted from Gayduk and₂₃₁ 209 Pantaler.²¹ In a typical analysis, the powdered sample₂₃₂ 210 (9 mg) was dissolved in 100 mL hydrochloric acid solu-233 211 tion (6 N) mixed with $35 \,\mathrm{mL}$ of indigo carmine solution₂₂₄ 212 (0.004 N). The resulted solution was cooled to room tem-₂₃₅ 213 perature and the excess of indigo carmine was titrated₂₃₆ 214 by potassium bromate $(0.001 \,\mathrm{N})$. 215 237

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III. RESULTS AND DISCUSSION

According to the calculated binary Co-O phase²⁴² 217 diagrams, 22,23 Co₃O₄ decomposes at elevated tempera-²⁴³ 218 tures to CoO, and then to Co. The region of Co_3O_4 sta-²⁴⁴ 219 bility depends strongly on oxygen partial pressure as well₂₄₅ 220 as the size of Co₃O₄ crystallites and their surface hydrox-246 221 vlation, shifting to higher temperatures with decreasing₂₄₇ 222 crystallite size and increasing oxygen partial pressure.248 223 For example, bulk Co₃O₄ decomposes to CoO at 1242 K₂₄₉ 224 when $p(O_2) = 1$ bar, while it decomposes at 909 K when₂₅₀ 225 when $p(O_2) = 10^{-6}$ bar.^{22,23} There are two studies that₂₅₁ 226





FIG. 1. X-Ray diffraction patterns for (a) single-phase CoO obtained by fast cooling, (b) mixed phase Co₃O₄ and CoO samples obtained by intermediate cooling rates, and (c) single-phase Co₃O₄ obtained by slow cooling. The poor signal-to-noise ratio for CoO is due to fluorescence from using Cu K α radiation. Asterisks denote peaks arising from the Al₂O₃ crucible used for the reaction.

report significantly lower temperatures. Results of *in situ* heating infrared emission spectroscopy have indicated the disorder of the spinel Co_3O_4 begins at 873 K and conversion to the rock-salt CoO occurs at 923 K.²⁴ There is also a high-temperature X-ray diffraction study that reports the transformation of Co_3O_4 to a disordered spinel occurs at 1150 K.⁸

Our initial attempts to produce dense sample of Co_3O_4 by sintering resulted in a mixture of spinel Co_3O_4 and rock-salt CoO. Accordingly, we varied the sintering conditions and heating/cooling rates to determine the optimum conditions for producing phase-pure Co_3O_4 . Through repeated trials, we found that dense samples of phase-pure spinel Co_3O_4 could be reproducibly prepared by sintering at 1148 K and then slowly cooling at 1 K/min. CoO was produced if the samples were quenched from 1148 K instead of cooling slowly. These results are summarized in Figure 1.

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Additional information concerning the phase stability of spinel Co_3O_4 was obtained by Raman spectroscopy, thermal analysis, and high-temperature XRD. Figure 2 shows the Raman spectroscopy results of samples heated from room temperature to 1200 K and then cooled. At room-temperature five Raman peaks are observed corresponding to the F_{2g} , E_g , and A_{1g} Raman active phonon

🛯 CoO

modes reported by Hadjiev for Co₃O₄.²⁵ With increas-₃₁₀ 252 ing temperature these peaks broaden, shift to lower311 253 wavenumbers and decrease in intensity. Rock-salt CoO₃₁₂ 254 has no Raman active modes and none are observed at₃₁₃ 255 the highest temperatures. However, the Raman modes₃₁₄ 256 for Co₃O₄ are unexpectedly extinguished in the same₃₁₅ 257 temperature regime, 600 K to 1000 K, where other high-316 258 temperature anomalies are observed. Upon cooling, the₃₁₇ 259 CoO to Co_3O_4 reaction is reversible and the same behav-318 260 ior is observed. 319 261

A more probable reason for the Raman mode extinc-320 262 tion is locally changing the temperature at the spot illu-321 263 minated by Raman laser. This suggestion is supported³²² 264 by previous studies on cobalt and iron oxide demonstrat-323 265 ing that even low power laser pulses can cause enough³²⁴ 266 local heating sufficient to meaningfully alter Raman spec-325 267 tra and oxidation states.^{26,27} On other hand other possi- 326 268 ble effects, such photoluminescence, which is superposed³²⁷ 269 with the Raman modes as temperature increases or in-328 270 version of the spinel structure cannot be completely ruled³²⁹ 271 out. One notes the inversion implies that Co^{2+} ions³³⁰ 272 will increasingly occupy the B site in the spinel struc-³³¹ 273 ture yielding an octahedral arrangement similar to rock332 274 salt making them dominant in the vibration properties³³³ 275 of Co₃O₄ and thereby removing Raman active modes.³³⁴ 276 However, the further investigations are needed to specify³³⁵ 277 the dominating effect. 336 278

The thermal analysis data shown in Figure 3 indicates³³⁷ that Co₃O₄ decomposition in air starts at ~1165 K and is³³⁸ completed by ~1270 K. The observed weight loss of $6.4\%^{339}$ corresponds well to the theoretical weight loss of 6.6% for³⁴⁰ the decomposition: Co₃O₄ \rightarrow 3 CoO + $\frac{1}{2}$ O₂ ³⁴¹

²⁸⁴ Upon slow cooling (10 K/min) in air, the increase in the³⁴² ³⁸⁵ specimen weight is observed at T < 1100 K, owing to the³⁴³ ²⁸⁶ reoxidation of CoO to Co₃O₄. The reoxidized specimen³⁴⁴ ²⁸⁷ nearly reaches its initial weight; the small weight differ-³⁴⁵ ²⁸⁸ ence of about 0.5% corresponds to x = 0.08 in Co₃O_{4-x},³⁴⁶ ³⁴⁹ implying a nearly stoichiometric composition with a min-³⁴⁷ ²⁹⁰ imal concentration of oxygen vacancies. ³⁴⁸

In an argon atmosphere, the Co_3O_4 decomposition³⁴⁹ starts at a lower temperature (~1084 K) and is completed³⁵⁰ by ~1200 K with the same weight loss. Upon slow cool-³⁵¹ ing, the increase in the specimen weight is observed at³⁵² T < 990 K, likely owing to incomplete oxidation of CoO_{353}^{353} to Co_3O_4 by residual oxygen in the instrument.

Quenching samples is performed to preserve the high-297 temperature electronic and crystal structure. However, 298 given that inversion is accomplished by the transfer of 299 an electron, and electron dynamics are generally asso-300 ciated with picosecond timescales, quenching in liquid 301 nitrogen is likely insufficiently quick to preserve any pos-302 sible electron transfer that is present only at high tem-303 perature. Nevertheless, neutron diffraction and XPS and 304 redox titration results both provide evidence of a sys-357 305 tematic electronic/magnetic/crystal structure transition₃₅₈ 306 as a function of quenching temperature. This systematic₃₅₉ 307 change in the quenched samples could be due to oxygen₃₆₀ 308 vacancies. However, as with TGA, redox titration (see₃₆₁ 309

Figure 4) indicates a negligible concentration of oxygen vacancies below 1100 K.

The results of the *in situ* XRD characterization agree with the results of the thermal analysis. According to analysis of the X-ray diffraction patterns, the spinel Co_3O_4 remains phase-pure from room temperature up to 1163 K upon heating in air (Figure 5). By 1273 K, Co_3O_4 had completely decomposed to the rock-salt structured CoO (see Table I, Figure 5). This decomposition was reversible: upon cooling in air the CoO oxidized back to phase-pure spinel Co_3O_4 . Specimens quenched to retain the rock-salt CoO phase remain stable in air on heating from room temperature until 950 K to 1188 K, where they are oxidized to Co_3O_4 . As the temperature is increased further, above 1188 K Co_3O_4 again decomposes reversibly to 100 wt% CoO.

The results of the quantitative Rietveld analysis (phase fraction, lattice parameter and oxygen fractional parameter u) are recorded in Table I.

As can be seen in Figure 6, the lattice parameter aof spinel Co_3O_4 exhibits a linear increase with temperature up to 950 K. Above 950 K, the lattice parameter was rapidly increased, while the oxygen fractional parameter u was sharply decreased. These results can be attributed to the cations disorder over tetrahedral and octahedral sites, which are in good agreement with previous report.⁸ In order to get more information about the cations disorder in spinel Co_3O_4 with temperature, the A–O and B-O bond length in tetrahedral and octahedral site, respectively, were calculated by Rietveld refinement (Figure 6). Similar to lattice parameter, both A-O and B-O bond length show a linear increase with temperature up to 950 K, which is due to the thermal expansion of the lattice. Above 950 K, a significant decrease in A–O bond length was observed, which can be explained by the partial substitution of smaller Co^{3+} for larger Co^{2+} cations $[r_{tetr} (\text{Co}^{3+}, \text{high spin}) = 0.45 \text{ Å}, r_{tetr} (\text{Co}^{2+}) = 0.58 \text{ Å};$ both ions are 4-fold coordinated].²⁸ On the other hand, the sharp increase in B-O bond length above 950 K could be attributed to the partial substitution of larger Co^{2+} for smaller Co³⁺ cations $[r_{oct} (\text{Co}^{3+}, \text{low spin}) = 0.53 \text{ Å},$ r_{oct} (Co²⁺) = 0.72 Å; both ions are 6-fold coordinated].²⁸ The cation distributions in Co_3O_4 spinel and the degree of inversion (i), which is equal to the fraction of B-cations on A sites or A-cations on B sites, can be calculated from bond lengths d_{A-O} and d_{B-O} from the relationships.

$$d_{A-O} = ir_{tetr}(Co^{3+}) + (1-i)r_{tetr}(Co^{2+}) + r_{tetr}(O^{2-})$$
(1)

$$d_{B-O} = \frac{i}{2} r_{oct}(Co^{2+}) + \frac{(2-i)}{2} r_{oct}(Co^{3+}) + r_{tetr}(O^{2-})$$
(2)

By using r_{tetr} (O²⁻) = 1.38 Å,²⁹ the A–O and B–O bond lengths for normal Co₃O₄ spinel (i.e. i = 0) are calculated to be 1.96 Å and 1.91 Å, respectively. Since the experimental values of A–O and B–O bond lengths are 1.935(3) and 1.9202(3) Å determined from Rietveld re-



FIG. 2. (a) In situ Raman spectroscopy of samples heated from room-temperature up to 1200 K and then cooled again. (b) Peaks corresponding to Co_3O_4 are observed but are extinguished between 600 K and 1000 K, long before the transformation to CoO that begins at 1165 K (determined by TGA).

TABLE I. Rietveld structure refinement details for slowly cooled Co_3O_4 as a function of temperature (standard deviation in parenthesis).

T (K)	$Wt\%Co_3O_4$	Wt%CoO	$a \operatorname{Co}_3 \operatorname{O}_4$	$u \operatorname{Co}_3 \operatorname{O}_4$	a CoO	$R_{\rm wp}$
298	100	-	8.0838(1)	0.2632(3)	-	2.95%
623	100	-	8.1023(3)	0.2630(8)	-	2.53%
800	100	-	8.1158(1)	0.2632(3)	-	2.54%
875	100	-	8.1225(1)	0.2632(3)	-	2.56%
950	100	-	8.1311(1)	0.2633(4)	-	2.55%
1025	100	-	8.1438(1)	0.2622(5)	-	2.71%
1093	100	-	8.1651(2)	0.2606(5)	-	2.78%
1100	100	-	8.1672(1)	0.2601(6)	-	2.53%
1113	100	-	8.1755(3)	0.2599(8)	-	2.53%
1140	100	-	8.1884(1)	0.2593(7)	-	2.50%
1153	100	-	8.1996(2)	0.2597(4)	-	2.59%
1163	100	-	8.2031(1)	0.2574(7)	-	2.46%
1273	-	100	-	-	4.3207(1)	3.79%
300	100	-	8.0842(3)	0.2628(3)	-	2.88%

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finement, a multiplication factors, $1.935/1.96 = 0.98778_{366}$ and 1.9202/1.91 = 1.00534 for A- and B-site respectively, 367 were introduced in the above equations to be: 368

$$d_{A-O} = [ir_{tetr}(Co^{3+}) + (1-i)r_{tetr}(Co^{2+}) + r_{tetr}(O^{2-}] \times 0.987$$
(3)₃₇₂
(3)₃₇₂

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$$d_{B-O} = \left[\frac{i}{2}r_{oct}(Co^{2+}) + \frac{(2-i)}{2}r_{oct}(Co^{3+}) + r_{tetr}(O^{2-})\right] \times \mathbf{1}_{\mathbf{J}_{\mathbf{J}}}^{373}$$

$$(4)^{375}$$

The increase in A-O and B-O bond lengths due to thermal expansion was determined by the linear fitting of experimental bond lengths below 950 K and extrapolating to higher temperature data. The obtained values were subtracted from experimental values of A-O and $7^{\circ}B$ -O bond lengths before calculating the degree of in-

version. As shown in Figure 5, Co_3O_4 remains ordered spinel below 1000 K, and then a significant increase in $r_4O_5M_4$ cation disordering was observed above that temperature. The highest degree of inversion in Co_3O_4 spinel



FIG. 3. Thermogravimetric Analysis and Differential Thermal Analysis of Co_3O_4 samples in air (a) and Ar (b). The sample transforms to CoO at 1165 K in air and 1084 K in Ar. In both cases, the weight loss corresponds to that expected by the loss of oxygen in Co_3O_4 . Unexpectedly, the sample in Ar regains a portion of the lost weight, likely due to reaction⁴⁰⁵ with residual oxygen in the instrument chamber.

calculated from A–O and B–O bond lengths were 0.54_{407} 376 and 0.60, respectively, at $1163 \,\mathrm{K}$, which is in good agree-377 ment with previous reported values.⁸ The difference be- $_{409}$ 378 tween the values of degree of inversion calculated from $_{410}$ 379 A-O and B-O bond lengths is negligible below $1113 K_{411}$ 380 and becomes remarkable at higher temperatures, which₄₁₂ 381 can be explained by transition of Co^{3+} from low-spin_{412} to high-spin state $[r_{oct} (\text{Co}^{3+}, \text{high spin}) = 0.61 \text{ Å}] \text{ or}_{414}$ the partial reduction of smaller Co^{3+} into larger Co^{2+}_{415} 382 383 384 cations. In fact, thermogravimetric analysis $(Figure 3)_{_{416}}$ 385 reveals a slight weight loss in Co_3O_4 spinel heated in₄₁₇ 386 air above $1100 \,\mathrm{K}$, which suggests the releasing of oxygen_{418} 387 from Co_3O_4 lattice and the partial reduction of Co^{3+} into 388 Co^{2+} as the charge compensation. This finding was $\mathrm{also}_{_{420}}$ 389 confirmed by XPS and redox titration analysis (Figures 4_{421} 390 and 9). 391 422

Rietveld refinement was performed on neutron diffrac-423 392 tion data collected above and below the Neel tempera- $_{424}$ 393 ture in order to determine the magnetic structure and₄₂₅ 394 moment (see Figure 7). At $1.5 \,\mathrm{K}$, the intensity of some₄₂₆ 395 fundamental reflections (e.g. (111), (200), (113), (222),427 396 (331) and (420)) increased markedly but no extra reflec-428 397 tions of magnetic origin were observed. This suggests₄₂₉ 398 that the magnetic unit cell is the same as the nuclear one430 399 and the magnetic structure can be described by a prop-431 400 agation vector k = (0, 0, 0). The magnetic irreducible₄₃₂ 401 representations for (8a) and (16d) sites propagation vec-433 402 tor k = (0, 0, 0) were determined using the BASIREPS₄₃₄ 403 program.¹⁸ The magnetic reducible representation for₄₃₅ 404



FIG. 4. Redox titration analysis of Co^{3+} reveals a small amount of oxygen vacancies, which increases with quenching temperature of Co_3O_4 .

(8a) site comprises two 3-dimensional irreducible representations as:

$$\Gamma_{Co1}^{8a} = 1\Gamma_7^{(3)} + 1\Gamma_{10}^{(3)} \tag{5}$$

The basis vectors of $1\Gamma_7^{(3)}$ and $1\Gamma_{10}^{(3)}$ are [(x, y, z) + (-x, -y, -z)] and [(x, y, z) + (x, y, z)], respectively. The diffraction patterns of all samples can be well fitted with the magnetic structure corresponding to representation Γ_7 (Figure 7). The refinements of the magnetic structure using the representation Γ_{10} for (8a) site as well as all irreducible representations determined for (16d) site yield poor fitting to the observed diffraction data. These results suggests that the magnetic structure of Co_3O_4 is due to antiferromagnetic ordering of spins in the A sites only, which is in good agreement with previous work.⁴ As the quenching temperature of the samples increases, the refined magnetic moment (Figure 8) remains roughly constant $(2.4 \,\mu_{\rm B})$ until 823 K, where the magnetic moment increases sharply until it reaches $2.71 \,\mu_{\rm B}$ in the sample quenched from 1200 K. The magnetic moment of the samples quenched from low temperature (i.e. $T < 823 \,\mathrm{K}$) is in good agreement with Chen and Selloni who calculate the Co²⁺ magnetic moment to be 2.59 $\mu_{\rm B}$ in bulk Co₃O₄.⁵ The increase in the magnetic moments in A-site with increasing the quenching temperature can be explained by cations disorder and the partial substitution of Co^{2+} by high-spin Co^{3+} in that tetrahedral site. On other hand, the diffusion of Co^{2+} into the octahedral *B*-site, which maintains the high spin state, is expected to induce a small magnetic moment in B-site. Nevertheless, no magnetic moment was observed in the B-site for all samples. However, degree of inversion in Co_3O_4 spinel can be also estimated from the overall effective magnetic moment μ_A



FIG. 5. In situ X-ray diffraction of Co_3O_4 on heating from room temperature to 1273 K. The sample converts entirely to CoO between 1163 K and 1273 K.

⁴³⁶ in the *A*-site using the equation:

$$\mu_A^2 = i\mu_{Co^{3+}}^2 + (1-i)i\mu_{Co^{2+}}^2 \tag{6}$$

where $\mu_{Co^{2+}}$ and $\mu_{Co^{3+}}$ are spin only effective moments of Co²⁺ (3.87 $\mu_{\rm B}$) and Co³⁺ (4.89 $\mu_{\rm B}$), respectively. The 437 438 theoretical effective moment in A-site for normal Co_3O_4 439 spinel (*i.e.* i = 0) calculated from the equation given 440 above is $3.87 \,\mu_{\rm B}$. Since the experimental effective mo-441 ment value of unheated Co_3O_4 spinel, which is expected 442 to be a normal spinel, was $2.4 \,\mu_{\rm B}$, a multiplication factor, 443 3.87/2.4 = 1.61, was introduced in the above equations 444 to be: 445

$$(\mu_A \times 1.61)^2 = i\mu_{Co^{3+}}^2 + (1-i)i\mu_{Co^{2+}}^2 \tag{7}$$

As shown in Figure 1, Co_3O_4 quenched from low temperature (i.e. T < 823 K) remains ordered spinel, and a significant increase in the cation disordering was observed in the samples quenched from temperatures above 823 K (See Figure 8). The highest degree of inversion in Co_3O_4 spinel quenched from 1163 K was 0.46, which is very close to the value obtained from *in situ* XRD experiments.

453 X-ray photoelectron spectroscopy (XPS) was per-454 formed to investigate the oxidation and spin states of 455 the Co centers. 3d transition metals (TM) in the oxide 456 compounds exhibit complicated nature of the electronic 457 structure characteristic for strongly correlated systems;



FIG. 6. (top) Lattice parameter, (middle) bond length, and (bottom) degree of inversion calculated from A–O and B–O bond lengths plotted as a function of temperature.



FIG. 7. Rietveld refinement was performed using neutron diffraction patterns of a quenched Co_3O_4 sample measured (a) below (1.5 K) the Neél temperature, and (b) above (300 K). The nuclear and magnetic structures are well-described by the model used to determine the magnetic moment of Co^{3+} .



FIG. 8. The magnetic moment of Co_3O_4 (*left axis*) and degree of inversion calculated from the magnetic moment (*right axis*) increase with the quench temperature. Samples quenched from higher temperature have a greater amount of oxygen vacancies, which increases the amount of Co^{2+} .

the shape of the photoelectron spectra is strongly in-458 fluenced by the final state effects and depends also on 459 the $TM \, 3d - O \, 2p$ hybridization.³⁰ Well defined photo-460 electron peaks at the binding energies of \sim 780 eV and 461 ${\sim}795\,\mathrm{eV}$ are due to spin-orbit splitting of the Co 2p spec- $_{439}$ 462 trum into $\text{Co} 2p_{3/2}$ and $\text{Co} 2p_{1/2}$ photoemissions (Fig-490 463 ure 9b). The spectral feature extended to higher binding₄₉₁ 464 energies from the main Co2p photoelectron line is the₄₉₂ 465 charge transfer satellite. The presence of the satellite₄₉₃ 466 peak, S, at $\sim 10 \,\mathrm{eV}$ is a sign of the contribution of octa-494 467 hedral Co^{3+} ions with the low spin state, while the Co^{2+}_{495} 468 ions contribute to the satellite structure at $\sim 6 \,\mathrm{eV}$ from₄₉₆ 469 the main photoelectron emission (Figure 9b). The diva-497 470 lent Co ions in tetrahedral coordination do not exhibit₄₉₈ 471 the strong intensive satellite structure (denoted as $S^*_{_{499}}$ in Figure 9b) in opposite to the octahedral Co²⁺ ions.³¹ 472 473 500 Samples quenched from lower temperatures (e.g., $300 \,\mathrm{K}_{501}$ 474 and 773 K) have Co 2p spectra consistent with previous₅₀₂ 475 reports of Co_3O_4 .^{31,32} 476 503

As the quenching temperature is increased (up to_{504} 477 1173 K), there is no evolution in the satellite peak S cor-505 478 responding to Co³⁺ ions, while, a slight increase in the 506 479 satellite peak S^{*} corresponding to Co²⁺ was observed.⁵⁰⁷ 480 Thus, ex situ XPS measurements suggest a small amount⁵⁰⁸ 481 of octahedral Co²⁺ ions is present in the sample quenched⁵⁰⁹ 482 from 1173 K,³¹ which confirms the results of *in situ* XRD₅₁₀ 483 and neutron diffraction experiments. Another noticeable511 484 difference: Co $2p_{3/2}$ and Co $2p_{1/2}$ core-line spectra from 512 485 samples quenched at high temperature (1173 K) demon-513 486 strate the main Co 2p photoelectron peaks with more⁵¹⁴ 487 broadening and less intensities at lower binding ener-515 488



FIG. 9. (a) O 1s and (b) Co 2p photoelectron spectra of Co_3O_4 samples quenched at low and high temperatures. The spectral features S and S^{*} are typical for Co_3O_4 with the Co^{3+} ions with LS state (satellite S) and Co^{2+} ions (satellite S^{*}); the shape of the Co 2p spectra is similar for both samples. $Co 2p_{3/2}$, $Co 2p_{1/2}$ and O 1s core-line spectra from samples quenched at high-temperature (1173 K) have peaks with less intensity, highlighted by the difference curve.

gies, e.g. ~779.2 eV and ~780.3 eV, see Figure 9b). Since the characteristic peaks of Co^{3+} ions are expected to be at binding energy lower than those of Co^{2+} ions,^{31,33–37} thus, this result indicates the increase in Co^{2+} ions in the sample quenched from 1173 K in comparison with those quenched from lower temperature. However, the precise determination of the surface $\text{Co}^{3+}/\text{Co}^{2+}$ ratio in samples by fitting XPS spectra is very difficult due to the controversial methods used to fit XPS spectra of Co_3O_4 in the literature.^{31,33–37}

The O 1s spectra of both samples can be decomposed into an intense peak at \sim 529.4 eV and a broader and less intense one at $\sim 530.8 \,\mathrm{eV}$, which can be assigned to lattice oxygen species and the surface adsorbed oxygen or surface oxygen defect species, respectively.^{31,33–37} As shown in Figure 9a, the intensity of the peak at \sim 530.8 eV in the spectrum of Co_3O_4 quenched from 1173 K is higher than that of the sample quenched from 300 K, which suggests the increase in the surface oxygen vacancies with increasing the quenching temperature. These results agree with the results of redox titration (Figure 4), which shows an increase in the oxygen vacancies in Co₃O₄ spinels with increasing quenching temperatures. According to the electroneutrality principle, the larger amount of surface vacancies in the sample quenched from high temperature is due to the partial reduction of Co^{3+} into Co^{2+} at elevated temperature.

The results of all our high-temperature characteriza-561 516 tion methods (XRD, Raman spectroscopy, thermal anal-517 vsis) unambiguously show that the decomposition of_{552} 518 Co_3O_4 to CoO occurs in air at a temperature above₅₅₃ 519 1100 K. This decomposition temperature is in good agree-520 ment with the majority of previous studies including₅₆₅ 521 those on which the phase diagrams for the Co–O sys- $_{566}$ 522 tem have been calculated.^{22,23} This is important because 523 it signifies that the anomalous broadening and weaken- $_{568}$ 524 ing of the Raman peaks reported here (Figure 2), as well₅₆₉ 525 as those reported by others between $600 \,\mathrm{K}$ and $1000 \,\mathrm{K}_{2570}$ 526 must be due to changes in the Co_3O_4 spinel structure₅₇₁ 527 itself rather than as a result of a phase transformation to_{572} 528 CoO. 529 573

575 The structural observations listed above have been at- $\frac{1}{576}$ 530 tributed to two main effects; inversion of the spinel struc-531 ture and spin unpairing of Co^{3+} ions on the octahe-532 dral site. Inversion of the structure implies a degree of_{579} 533 cation mixing where both Co^{2+} and Co^{3+} ions are dis-534 ordered over octahedral and tetrahedral sites. Koumoto₅₁₁ 535 and Yanagida suggested this leads to the rise in elec- $_{\scriptscriptstyle 582}$ 536 trical conductivity, as electron exchange (hopping) can₅₈₃ 537 occur between Co^{2+} and Co^{3+} ions on either site.¹¹ Ef⁵⁸³₅₈₄ 538 occur between Co²⁺ and Co²⁺ ions on either site. E1-584 fectively, this is equivalent to an electron transfer from 585 Co²⁺ to Co³⁺ (hopping): Co²⁺_A ($e_g^4 t_2^3$, S = 3/2) + 586 Co³⁺_B (LS, $t_{2g}^6 e_g^0$, S = 0) \leftrightarrow Co³⁺_A (HS, $e_g^3 t_2^3$, S = 2) 587 + Co²⁺_B ($t_{2g}^6 e_g^1$, S = 1/2). On the other hand, spin 588 unpairing involves the octahedral Co³⁺ electrons going 589 from a LS to a HS state as in: Co³⁺_B (LS, $t_{2g}^6 e_g^0$, S = 0) 590 539 540 541 542 543 544 $\rightarrow \operatorname{Co}_B^{3+}(\operatorname{HS}, t_{2q}^4 e_g^2, S=2)^{38}$ with possible formation of₅₉₁ 545 an intermediate state, IS, Co_B^{3+} , with $t_{2g}^5 e_q^1$, $S = 1.^{39}$ 546

The measurements reported in this work do not pro-547 vide any additional evidence of a spin state transition in⁵⁹³ 548 spinel Co_3O_4 . However, the results of in situ XRD ex-594 549 periments and the evolution of the magnetic moment in₅₉₅ 550 the quenched samples does suggest inversion of the struc-596 551 ture. It is also possible that the extinction of Raman ac-597 552 tive modes in Co_3O_4 could be associated with inversion₅₉₈ 553 of the structure. While these findings suggest that ex situs 554 measurements on quenched samples could potentially of-600 555 fer some insight into high-temperature crystal structures⁶⁰¹ 556 that are otherwise experimentally inaccessible, it is likely₆₀₂ 557 that the electronic transitions associated with inversion603 558 and spin state in Co_3O_4 are too rapid to be observed.⁶⁰⁴ 559 Additional *in situ* measurements are needed. 560 605

IV. CONCLUSIONS

X-ray diffraction, thermal analysis, and Raman scattering of Co_3O_4 spinel are reported from room temperature up to 1273 K. Previous changes in lattice parameter and bond length were confirmed, and we report a newly observed extinction of Raman peaks between 600 K and 1000 K. This anomaly occurs over the same temperature range as many other reported high temperature anomalies in this compound. The phase evolution of Co_3O_4 to CoO was carefully monitored (with an observed onset in air at ~1165 K), and does not contribute to this anomaly.

By considering the different cation sizes of Co^{3+} and Co^{2+} in octahedral and tetrahedral sites along with the changing A–O and B–O bond lengths determined by *in situ* X-ray diffraction, the degree of inversion, *i*, was calculated to reach 0.6. Samples were also quenched from high temperature for *ex situ* characterization by neutron diffraction, XPS, and chemical titration. In situ TGA investigation of the Co₃O₄ to CoO reaction suggests the spinel Co₃O₄ is close to the ideal oxygen stoichiometry, without large fractions of oxygen vacancies. Ex situ redox tirtation reveals quenching samples from above 800 K introduces a small amount of oxygen vacancies, up to ~4% by 1100 K.

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Magnetic structure refinement performed on neutron diffraction patterns reveals the A-site magnetic moment increases from $2.4 \,\mu_{\rm B}$ to $2.7 \,\mu_{\rm B}$ with quenching temperature, suggesting a maximum degree inversion of 0.46, in fairly good agreement with the value based on *in situ* bond length changes. No direct evidence of spin unpairing (spin state transition from LS to HS) was observed.

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^{606 *} email: sparks@eng.utah.edu

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