# On the relative timing of listwaenite formation and chromian spinel equilibration in serpentinites

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#### ABSTRACT

Ultramafic rocks exposed at the Earth's surface generally record multiple stages of evolution that may include melt extraction, serpentinization, carbonatization, and metamorphism. When quantitative thermometry based on mineral chemistry is applied to such rocks, it is often unclear what stage of their evolution is being observed. Here, in peridotites with extensive replacement of silicate minerals by carbonates (listwaenites), we present a case study that addresses the timing of carbonate formation relative to closure of exchange reactions among relict primary minerals.

Massive and schistose serpentinized peridotites of Neoproterozoic age outcrop at Gabal Sirsir, South Eastern Desert, Egypt (northwestern corner of the Arabian-Nubian Shield or ANS). Petrography, bulk composition, and mineral chemistry are all consistent with a strongly depleted mantle harzburgite protolith for the serpentinites. Bulk compositions are low in Al<sub>2</sub>O<sub>3</sub> and CaO and high in Mg# [molar Mg/(Mg+Fe) = 0.89-0.93]. Relict spinel has high Cr# [molar Cr/(Cr+Al)] and low Ti, while relict olivine has high Mg# and NiO contents. Based on compositions of coexisting relict olivine and chromian spinel, the protolith experienced 19 to 21% partial melt extraction. Such high degrees of partial melting indicate a supra-subduction zone environment, possibly a forearc setting.

Along thrust faults and shear zones, serpentinites are highly altered to form talc-carbonate rocks and weathering-resistant listwaenites that can be distinguished petrographically into Types I and II. The listwaenitization process took place through two metasomatic stages, associated first with formation of the oceanic crustal section and near-ridge processes (~750-700 Ma) and subsequently during obduction associated with the collision of East and West Gondwana and escape tectonics (~650-600 Ma). In the first stage, Mg# of chromian spinel in the serpentinites continuously changed due to subsolidus Mg– $Fe^{2+}$  redistribution, while the Mg# of chromian spinel in the Type I listwaenites was frozen due to the absence of coexisting mafic silicates. During the second stage, the Type II listwaenites formed along shear zones accompanied by oxidation of relict chromian spinel to form ferritchromite and Crbearing magnetite in both serpentinites and listwaenites. The high Cr# of chromian spinel relics in both serpentinites and listwaenites preserves primary evidence of protolith melt extraction, but divalent cations are more easily mobilized at low temperature. Hence, relict chromian spinel in listwaenites shows significantly higher Mg# and lower MnO than that in serpentinite, suggesting that nearly complete alteration of ultramafic rocks to form listwaenite took place prior to re-equilibration between chromian spinel and the surrounding mafic minerals in serpentinites. Furthermore, the ferritchromite in the serpentinites has higher Mn content (1.1-2.1 wt%) than that in the listwaenites (0.6-0.9 wt%), indicating its formation after carbonatization since carbonate minerals are a favorable sink for Mn.

Keywords: Arabian-Nubian Shield, neoproterozoic, serpentinite, listwaenite

# **INTRODUCTION**

Listwaenite is a term used for carbonated meta-ultramafic rocks that commonly bear a fuchsite-quartz-carbonate mineral paragenesis (e.g., Halls and Zhao 1995; Gahlan et al. 2015a) and represent the end products of various degrees of carbonatization, potassium alteration, and silicification. Alternate spellings include listvenite and listvanite (Kelemen et al. 2011) and roughly interchangeable names for the same rock type include "Barramiya rocks" and "carbonated meta-ultramafics" (e.g., Rittmann 1958; Azer 2013; Gahlan et al. 2015a; Boskabadi et al. 2017; Sofiya et al. 2017). They are often associated with talc-carbonate rocks. During the listwaenitization process, primary ferromagnesian silicate minerals in ultramafic rocks are replaced by carbonate minerals, and the released silica forms quartz (Uçurum 2000). Fuchsite, the common name for the green Cr-rich variety of muscovite, is the result of potassium metasomatism. Carbonated meta-ultramafic rocks in the Eastern Desert

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of Egypt, especially listwaenites, have drawn attention because of their economic potential, including gold mineralization (e.g., Botros 1993; Osman 1995; Botros 2002, 2004; Ramadan 2002; Ramadan et al. 2005; Abd El-Rahman et al. 2012; Azer 2013; Gahlan et al. 2015a; Boskabadi et al. 2017) that has been mined since Pharaonic times (Harraz 2000; Klemm et al. 2001).

Chromian spinel is a nearly ubiquitous accessory mineral in ultramafic and many mafic rocks. It has been used successfully as a geotectonic and petrogenetic indicator mineral (e.g., Irvine 1965, 1967; Dick and Bullen 1984; Arai 1992, 1994) because its composition is notably sensitive to changes in temperature, pressure, oxygen fugacity, bulk-rock and fluid composition (e.g., Irvine 1965, 1967) and because it resists low-temperature alteration processes that may affect every other phase in an ultramafic sample. The sensitivity of spinel Cr# in particular to the degree of partial melt extraction has been widely applied as a geotectonic indicator for upper-mantle derived peridotites (e.g., Dick and Bullen 1984; Arai 1994).

The obduction of ophiolites in the Arabian-Nubian Shield (ANS) was accompanied by several kinds of alteration (e.g., Stern et al. 2004; Azer and Stern 2007; Azer 2013; Boskabadi et al. 2017). A characteristic feature in the ANS ophiolites is the major abundance of carbonate alteration along shear zones in the ultramafic rocks, but the sources and fluxes of the CO<sub>2</sub>-rich fluids that drove this carbonate alteration and its timing relative to other events in the evolution of the rocks are unknown.

The current research offers a detailed report on the geological, mineralogical, and geochemical characteristics of the serpentinized peridotites and their alteration products at Gabal (G.) Sirsir, Wadi Hodein district, South Eastern Desert, Egypt. No detailed petrological or geochemical study of this occurrence has been published before. Although the serpentinization and carbonatization processes largely obliterated primary petrological characteristics of the G. Sirsir peridotites, primary relics of mantle olivine and chromian spinel have been used as petrogenetic indicators. We focus on the evidence of alteration and achievement of equilibrium between chromian spinel and coexisting minerals, which provides a good opportunity to constrain the relative timing of the epoch in the evolution of the rocks recorded by the spinel chemistry and of the events leading to carbonatization of the ultramafic rocks to form listwaenite. A similar approach to the listwaenite occurrences in Ethiopia was recently published by Sofiya et al. (2017); we compare our results and conclusions to theirs below.

# **GEOLOGICAL OUTLINE**

The Arabian-Nubian Shield (ANS) is a juvenile tract of continental crust that forms the northern part of the East African Orogen (e.g., Stern 1994, 2002; Jarrar et al. 2003; Johnson and Woldehaimanot 2003; Meert 2003; Stoeser and Frost 2006). It formed in the Neoproterozoic through multiple stages of accretion and arc development along several suture zones. Ophiolitic rocks are scattered through most of the ANS. They represent fragments of oceanic lithosphere obducted on the continental margins during collision between East and West Gondwana and closure of the Mozambique Ocean. Such Neoproterozoic ophiolites are common in the central and south Eastern Desert of Egypt (Fig. 1), the northwestern corner of the ANS. Reconstructing the pseudo-lithostratigraphic column of the Egyptian ophiolites shows that



**FIGURE 1.** Distribution of ophiolitic rocks in the Eastern Desert of Egypt (modified after Shackleton 1994; Stern and Hedge 1985); the location of the study region (Fig. 2) is indicated.

they represent remnants of oceanic lithosphere that formed in a supra-subduction zone setting during Neoproterozoic time (e.g., Stern et al. 2004; Azer and Stern 2007; Ali et al. 2010; Azer et al. 2013; Khalil et al. 2014; Gahlan et al. 2015b; Obeid et al. 2016).

Complete ophiolite sections are rare (but present) in the Eastern Desert of Egypt. They generally consist of a lower serpentinized ultramafic unit and an upper unit of layered and isotropic gabbros, sheeted dikes, and massive or rarely pillow basalts (e.g., El Sharkawy and El Bayoumi 1979; El Bayoumi 1983; Gahlan et al. 2015b). However, most of the Egyptian ophiolites are variably dismembered, deformed, altered, and metamorphosed due to serpentinization and interaction with a large flux of CO<sub>2</sub>-bearing fluids (Stern and Gwinn 1990; Gahlan et al. 2015a; Boskabadi et al. 2017). The alteration is focused along shear zones and faults and leads to formation of various types of carbonate-bearing meta-ultramafics.

The G. Sirsir area forms the southern part of the Wadi Hodein district in the South Eastern Desert of Egypt, ~20 km west-northwest of Shalatin City (Fig. 1). It features outcrops of Neoproterozoic basement rocks including an ophiolite sequence, an island arc association, and late- to post-tectonic granites (Fig. 2). Basement rocks in the area were affected by a regional compressional regime with shortening along the west-southwest (WSW) to east-northeast (ENE) direction, expressed by northwest-trending folids and WSW-verging thrust faults, but many earlier phases of structural deformation can be recognized as well (e.g., Abdeen et al. 2008). The G. Sirsir ophiolite fragments are polydeformed allochthonous megashear masses in a tectonic mélange and include serpentinized ultramafics, metagabbros and pillow metabasalts. The island-arc association is represented by variably deformed metasediments, metavolcanics, and a metagabbro-diorite complex. The deformed metagabbro-diorite rocks intrude the ophiolitic mélange matrix and the island arc metavolcaniclastics. The metavolcaniclastics, serpentinites, and carbonatized serpentinites (listwaenites) are subsequently intruded by late- to post-orogenic granites, with

notable granite offshoots.

The variably serpentinized and carbonatized ophiolitic ultramafic rocks form the mantle section of the G. Sirsir ophiolite (Fig. 2). They form lenses and sheets striking northwest (NW) and dipping ~50° to the southwest ~50° SW, tectonically incorporated in a mélange with metasediments and metavolcaniclastics. Serpentinites are generally massive and homogeneous, dominated by harzburgite with less-abundant dunite. Along shear zones and toward the sole thrust, they are transformed into schistose serpentinites, listwaenites, and talc-carbonate rocks. Mesoscopic network veins (2–20 cm thick) of magnesite and quartz cross-cut both serpentinites and talc-carbonates. Also, small masses and irregular pockets of magnesite are observed in the sheared carbonatized serpentinite. Talc-carbonate rocks occur as variably sheared blocks, sheets, and lenses along the NW and NNW striking fault zones.

The talc-carbonate rocks are brownish to greenish gray and can be either massive or highly schistose. On the other hand, listwaenites form rootless reddish cream-colored sheets, ridges, and irregular lenses and masses along shear zones and fault



FIGURE 2. Geologic map of Gabal Sirsir area, modified after Zoheir (2011).

planes (up to 3000 m long and 500 m wide). They express positive geomorphic relief relative to the surrounding rocks due to their resistance to arid-climate weathering. Structural elements in the listwaenite rocks are generally conformable to the main plano-linear fabric of the host country rocks (Fig. 2). Along contacts, the listwaenites show no chilled margins, reaction haloes, or xenoliths from the host. The characteristic reddish cream weathered surface and porous texture are the results of supergene oxidative weathering of Fe-bearing carbonates to form iron oxides. Two types of listwaenite have been identified, Types I and II (Fig. 3). Type I is typical listwaenite. Although Type II might best be called a listwaenite-like rock, we will refer to it herein as Type II listwaenite. Type I is fuchsite-bearing and low in modal quartz; it is found spatially associated with serpentinites in yellowish-green colored NW striking deformed lenses with prominent schistosity (Figs. 3a and 3b). By contrast, Type II is fuchsite free with more abundant quartz and is spatially associated with granites in west-northwest striking, undeformed, reddish-cream or yellowish masses, locally brecciated and fractured (fractures are filled with carbonate veinlets and fine quartz ribbons) (Figs. 3c and 3d).

#### METHODOLOGY

Mineral identification in polished thin sections was accomplished using a Polarizing Nikon Microscope, environmental scanning electron microscope (ESEM), and electron probe micro-analyzer (EPMA), supplemented by powder X-ray diffraction (XRD) analyses. XRD analyses used a BRUKER D8 advanced X-ray diffractometer in the Central Metallurgical and Development Institute, Cairo, Egypt, with Cu radiation and a secondary monochrometer. Scanning speed was  $2\theta = 1^{\circ}/min$  at constant voltage 40 kV and current 40 mA. Mineral identification was referenced to both *d*-spacing and relative intensities of reflections using the American Standard Test Materials (ASTM) cards.

Some polished sections of listwaenite samples were examined with a Philips XL30 ESEM at the Nuclear Materials Authority in Egypt, operating at 25 kV and equipped with energy-dispersive analytical X-ray (EDAX) capability. The spectrometer detects elements with atomic number greater than 4 (e.g., B) and a count rate ~1000–1500 counts per second.

Mineral chemistry of relict primary (olivine and spinel) and metamorphic minerals were determined with an electron microprobe at the Geology and Metallogeny Laboratory, Orléans, France. Operating conditions were 15 kV accelerating voltage, 20 nA beam current and 3  $\mu$ m beam diameter. Suitable synthetic and natural silicate and oxide standards were applied for calibration. Carbonate standards were not used for the carbonate mineral analyses; instead carbonate analyses were checked for quality by computing the weight of CO<sub>2</sub> necessary to stoichiometrically balance the measured FeO+MnO+MgO+CaO. Samples whose total estimated this way did not yield 100  $\pm$  2 wt% were discarded.

Whole-rock chemical analyses (major oxides and trace elements) of powdered



FIGURE 3. (a) Field photo showing sheared serpentinite and listwaenite (type-I). (b) Hand specimen of sheared listwaenite (type-I) with oriented quartz and carbonates. (c) Field photo showing massive listwaenite (type-II), and (d) hand specimen of massive listwaenite (type-II) with quartz veins.

rock samples (12 listwaenites and 8 serpentinites) were carried out at ACME Analytical Laboratories in Vancouver, Canada. The analyzed serpentinite samples were carefully selected to avoid carbonate or quartz veinlets. Samples were analyzed by inductively coupled plasma-mass spectrometry (ICP-MS) following a lithium metaborate/tetraborate fusion and nitric acid digestion of a 0.2 g sample. The detection limits are between 0.01 and 0.04% for major elements, and between 0.01 and 0.5 ppm for most of the trace elements, except Au (0.5 ppb) and Ni (20 ppm). Analytical precision, as calculated from replicate analyses, is 0.5–1.0% for major elements and varies from 5 to 20% for trace elements. Loss on ignition (LOI) is determined by weight difference after ignition at 1000 °C.

# PETROGRAPHY

# Serpentinites

The massive and sheared serpentinites of G. Sirsir are notably similar in composition, but the latter has a schistosity defined by subparallel alignment of serpentine flakes. The massive serpentinite consists mainly of antigorite with lesser amounts of chrysotile, lizardite, magnesite, talc, chlorite, brucite, and magnetite (quantitative mineral identification by XRD). There are trace relics of primary olivine and chromian spinel. Locally, a few serpentinite samples are rich in carbonates (up to 40 vol%). The serpentine minerals replacing olivine exhibit mesh texture (Fig. 4a), while those replacing orthopyroxene exhibit bastite texture (10–15 vol%) (Fig. 4b), suggesting dunite and harzburgite protoliths.

Antigorite forms fibro-lamellar or flame-like crystals and interpenetrating texture. Occasionally, antigorite is intergrown with magnesite (Fig. 4c). The chrysotile long fibers form veinlets cross-cutting the antigorite matrix, indicating protracted serpentinization. Carbonates are stained reddish brown by iron oxides. In general, the carbonates occur as scattered aggregates (Fig. 4d), as cross-cutting cryptocrystalline veinlets and crack-fill, or as replacement of mesh texture. Talc occurs as flakes or fine-grained matrix associated with carbonates. Brucite appears as platy or fibrous crystals intermixed with serpentines and as veinlets.

Chromian spinel morphology appears to be independent of whether the host rock is schistose. It forms subhedral and anhedral crystals in serpentinized dunites and amoeboid or vermicular (Fig. 4e) crystals in serpentinized harzburgites. Rarely, it forms euhedral cubic crystals and cumulate layers. Along grain boundaries and cracks, chromian spinel is partially replaced by an irregular ferritchromite mantle (light-gray colored) and a magnetite rim (gray-white colored) (Fig. 4e). A few altered chromian spinel rims are surrounded by aureoles of faintly pleochroic flakes, from violet to deep violet, of chromian chlorite (the variety commonly called kämmererite). Magnetite forms scattered anhedral to euhedral crystals throughout serpentinites. In addition, magnetite develops as rims around chromian spinel or by replacing mesh centers and rims. Disseminated sulfides-including pyrite, pentlandite, and rare chalcopyrite-are observed scattered throughout the serpentinites and variably altered. Pyrite occurs as euhedral to subhedral disseminated crystals or as inclusions in magnetite. Pentlandite occurs as rounded grains, slightly altered to garnierite and magnetite along margins.

# Listwaenites

Petrographically, two types of listwaenite can be identified: Type I (typical listwaenite with fuchsite; Fig. 4f) and Type II (listwaenite-like rocks without fuchsite). Although listwaenites form along shear zones, most samples do not show any preferred orientation or schistosity. Type I listwaenite is composed essentially of carbonates (50-60 vol%), guartz (35-45%), and minor fuchsite, with accessory Fe-Ti oxides, serpentine, chromian spinel, and chlorite. The carbonates are mainly magnesite and breunnerite, with minor amounts of calcite and dolomite. Breunnerite clearly formed at the expense of primary carbonates and is sometimes in turn replaced by Fe-Mn oxides. Fuchsite has a dark emeraldgreen color. It forms flakes as well as fine disseminated crystals and thin bands with a perfect cleavage in one direction (Fig. 4f), sometimes intergrown with chlorite. A few magnesite veinlets are observed cross-cutting the groundmass. The chromian spinel forms subhedral-fractured grains with ferritchromite developed along grain boundaries and cracks (Fig. 4g). The ferritchromite zone around primary spinel is consistently thinner in Type I listwaenites (Fig. 4g) than that in serpentinites. Polished thin sections revealed minor disseminated sulfides and scarce native gold.

The Type II listwaenite (fuchsite-free listwaenite-like rocks) has typically slightly more modal quartz (40–45%) and lower abundance of carbonate minerals (45-60%) than Type I listwaenite. The accessory minerals include Fe-Ti oxides, chromian spinel, muscovite, chlorite, and sulfides. Reflected-light microscopy shows that Type II listwaenites are richer in opaque minerals (~2.0-2.5% by volume) than Type I listwaenites (<1.5%). The opaque minerals include pyrite, chalcopyrite, galena, sphalerite, covellite, goethite, magnetite, and gold. The carbonate minerals are mainly magnesite with less calcite and dolomite. Chromian spinel is mostly anhedral, fractured, and highly altered to ferritchromite (Fig. 4h). The alteration of chromian spinel in the Type II listwaenite is more extensive than that in the Type I listwaenite. In general, the petrography of our Type II listwaenite appears similar to all the samples described from the Ethiopian occurrence by Sofiya et al. (2017).

# Magnesite veins and masses

The magnesite in veins and masses is nearly monomineralic and cryptocrystalline. Petrographic, XRD, and EDAX analyses indicate that the magnesite veins consist of >98 vol% coarse (0.1–0.4 mm) magnesite with minor amounts of serpentine minerals and iron oxides. The massive magnesite is composed essentially of anhedral fine-grained (0.05–0.1 mm) magnesite (85–90 vol%) with minor serpentine minerals, iron oxides, dolomite, and calcite. Carbonate minerals in the magnesite masses display evidence of shearing in the forms of stretched grain ribbons and recrystallized bulges along crystal boundaries. Dolomite and calcite fill cavities and vugs in the magnesite masses and are somewhat coarser (0.5–0.8 mm) than magnesite.

#### Talc-carbonates

The talc-carbonate rocks occur as massive bodies or sheared mylonitic masses. The massive type consists essentially of carbonate minerals and talc, with minor amounts of amphibole (tremolite, anthophyllite), quartz, chlorite, altered chromian spinel, and magnetite. Talc occurs as microcrystalline fibers and fine shreds, or rarely as coarse- to medium-grained flakes. In a few samples, coarse talc flakes occur as nests and openspace filling veins, indicating different generations of talc.



**FIGURE 4.** Photomicrographs showing petrographic textures under crossed nicols (**a**, **b**, **c**, **d**, and **f**) and reflected light (**e**, **g**, and **h**). Abbreviations: Ol = olivine, Cb = carbonates, Spl = spinel, Fu = Fuschite, Srp = serpentine, Qtz = quartz, Mt = magnetite. (**a**) Fresh relic of olivine within serpentinite; (**b**) bastite texture after orthopyroxene; (**c**) antigorite crystals intergrown with magnesite; (**d**) scattered blocky aggregates of carbonates within serpentinite; (**e**) large chromian spinel crystal with pristine core followed by an irregular light-gray ferritchromite zone and a gray-white magnetite rim; (**f**) fuchsite flakes with a perfect cleavage in one direction; (**g**) chromian spinel in listwaenite overprinted by thin rims of ferritchromite along margins and later cracks; and (**h**) chromian spinel in Type II listwaenite-like rock highly altered to ferritchromite.

Chromian spinel is completely altered to ferritchromite. The mylonitic talc-carbonates consist of carbonate minerals and talc mixed with nodules of serpentinite and other rock types. Chromian spinel and magnetite occur as highly altered cataclastic and brecciated crystals. In a few samples of mylonitic talc-carbonate, quartz veinlets are observed cutting the talc groundmass.

# MINERAL CHEMISTRY

The whole microprobe data set is given in the Electronic Appendix<sup>1</sup> (Supplementary Tables S1–S8). Not all phases could be analyzed; anthophyllite and tremolite were recognized petrographically but not quantitatively analyzed.

#### Olivine

Rare relics of primary olivine can be found in the serpentinites. Microprobe analyses, calculated structural formulas, and end-member components are given in Supplementary<sup>1</sup> Table S1. MgO (48.2–52.4 wt%) and NiO (0.3–0.5 wt%) contents are high, similar to those in primary mantle olivines (e.g., Arai 1980; Takahashi et al. 1987) and other Egyptian ophiolite mantle sequences. Their Fo contents (89.4–93.6; avg. 92.1) are clearly distinct from those in olivine from the non-ophiolitic maficultramafic intrusions in Egypt (Fig. 5a).

#### **Chromian spinel**

Electron microprobe analyses and calculated structural formulas of fresh cores and altered rims of chromian spinel are given in Supplementary<sup>1</sup> Tables S2 and S3. In most grains, Al<sub>2</sub>O<sub>3</sub>, Cr<sub>2</sub>O<sub>3</sub>, and MgO show a systematic decrease from core to rim, whereas  $Fe_2O_3^T$  increases outward. MnO does not show any systematic variation within fresh spinel cores but is concentrated in the ferritchromite alteration zone in serpentinite samples. The Al–Cr–Fe<sup>3+</sup> triangular plot reveals differences in chemical composition between various zones, with a well-developed chemical gap between spinel primary cores and altered rims (Fig. 5b). The fresh cores are rich in Al<sub>2</sub>O<sub>3</sub> and Cr<sub>2</sub>O<sub>3</sub> and lie along the Cr-Al join. The ferritchromite phase lies close to the Cr-Fe<sup>3+</sup> join; Al is lost preferentially to Cr. The Cr-magnetite outer rims are nearly devoid of Al<sub>2</sub>O<sub>3</sub> and lie along the Cr-Fe<sup>3+</sup> join with progressive Cr depletion and ferric iron enrichment.

The chromian spinel cores in type-I listwaenite have distinctly higher Mg# (0.5–0.6) than those in serpentinites (Mg# = 0.3-0.5) and type-II listwaenite (Mg# = 0.3-0.4), with minimal overlap between the populations (Fig. 5c). There are a few large disseminated chromian spinel crystals in the serpentinites with high Mg# (~0.5) cores similar to those recorded in the listwaenite. The interpretation of these Mg# data is discussed below and is a central part of the argument of this work.

Ferritchromite was analyzed in both serpentinites and listwaenites. The ferritchromite in type I listwaenite is richer in MgO (2.3–4.0 wt%), Al<sub>2</sub>O<sub>3</sub> (3.2–4.6 wt%), and Cr<sub>2</sub>O<sub>3</sub> (35.3–40.3 wt%) and lower in FeO<sup>T</sup> (49.5–55.1 wt%) and MnO (0.6–0.9 wt%) than that in serpentinites (0.4–1.9 wt% MgO, 0.5–1.7 wt% Al<sub>2</sub>O<sub>3</sub>, 14.0–33.1 wt% Cr<sub>2</sub>O<sub>3</sub>, 59.0–81.1 wt% FeO<sup>T</sup>, and 1.1–2.1 wt% MnO) and type II listwaenite (1.0–3.4 wt% MgO, 1.0–2.0 wt% Al<sub>2</sub>O<sub>3</sub>, 22.0–34.9 wt% Cr<sub>2</sub>O<sub>3</sub>, 56.1–71.7 wt% FeO<sup>T</sup>, and 1.0–2.2 wt% MnO). The interpretation of MnO distributions in the spinel family minerals is discussed below. Minor but resolvable silica contents are detected in ferritchromite (0.5-1.2 wt%) in serpentinite and 0.4-1.3 wt% in listwaenite); silica in the spinel structure has been attributed elsewhere to effects of alteration and metamorphism (Burkhard 1993).

# Mica

Chemical composition and structural formulas of the analyzed muscovite species are given in Supplementary<sup>1</sup> Table S4. Two types of mica can be recognized: fuchsite (green Cr-muscovite) in Type I listwaenite and muscovite (white mica) in Type II listwaenite. Fuchsite in Type-I listwaenite has 6.1–6.3 apfu Si, which completely overlaps the range of Si in muscovite in the Type-II listwaenite. Fuchsite has higher Cr<sub>2</sub>O<sub>3</sub> (1.8–3.7 wt%), FeO<sup>T</sup> (2.5–3.7 wt%), and MgO (1.3–2.5 wt%) with lower Al<sub>2</sub>O<sub>3</sub> (30.1–32.3 wt%) and K<sub>2</sub>O (7.4–9.4 wt%) than muscovite (0.2–0.8 wt% Cr<sub>2</sub>O<sub>3</sub>, 1.6–2.5 wt% FeO, 1.2–1.6 wt% MgO, 31.8–34 wt% Al<sub>2</sub>O<sub>3</sub>, and 9.4–10.1 wt% K<sub>2</sub>O). Cr<sub>2</sub>O<sub>3</sub> contents in fuchsite are negatively correlated with Al<sub>2</sub>O<sub>3</sub> contents (Fig. 5d).

# Magnetite

Disseminated magnetite crystals and magnetite rims around chromian spinels could only be analyzed in serpentinite samples; chemical composition and structural formulas are given in Supplementary<sup>1</sup> Table S5. The disseminated magnetite has negligible  $Cr_2O_3$  (0.1–1.3 wt%), whereas the magnetite developed around spinel has  $Cr_2O_3$  contents between 2.0 and 7.8 wt%. On the Cr–Al–Fe<sup>3+</sup> plot (Fig. 5b), both magnetites plot along the Cr–Fe<sup>3+</sup> join. Outer Cr-magnetite rims around the ferritchromite in serpentinites are depleted in MnO (0.1–0.9) (see below).

# Chlorite

Chlorite was analyzed in serpentinites and both listwaenite types; chemical composition and structural formulas are given in Supplementary<sup>1</sup> Table S6. In the serpentinites, chlorite occurs in two forms, as an aureole around chromian spinel and as isolated sparse crystals in the groundmass. Likewise, in Type I listwaenites, chlorite occurs both intergrown with fuchsite and as disseminated grains. The abundances of SiO<sub>2</sub>, Al<sub>2</sub>O<sub>3</sub>, FeO, and Cr<sub>2</sub>O<sub>3</sub> in chlorite vary systematically with its mode of occurrence. Chlorite from serpentinites contains higher  $Al_2O_3$  (19.3–21.2) wt% in aureoles and 20.0-21.8 wt% in disseminated chlorite) and lower SiO<sub>2</sub> (26.8-28.2 wt% in aureoles and 27.1-28.8 wt% in disseminated chlorite) than those from listwaenite (16.2-18.8 wt% Al2O3 and 28.5-31.0 wt% SiO2). The chlorite in the aureoles around chromian spinel is Cr-bearing chlorite (kämmererite), with higher  $Cr_2O_3$  (2.5 to 3.9 wt%) and MgO (22.2–23.9 wt%) than the disseminated chlorite (Cr-poor and Fe-rich). Cr-rich chlorite in the serpentinites is classified mainly as ripidolite, whereas the disseminated chlorite includes ripidolite and pycnochlorite (Supplementary<sup>1</sup> Table S5).

The chlorite intergrowth with fuchsite in the Type I listwaenite is rich in MgO (20.4–22.1 wt%) with somewhat elevated  $Cr_2O_3$  (1.9–2.4 wt%), whereas disseminated chlorite in the Type I listwaenite has elevated FeO<sup>T</sup> (18.4–20.3 wt%) and low  $Cr_2O_3$ (1.1–1.6 wt%). According to the classification of Hey (1954), chlorite in Type I listwaenite is classified mainly as pycnochlorite



**FIGURE 5. (a)** Mg# vs. NiO (wt%) of olivine from G. Sirsir serpentinized peridotites. The mantle olivine array is adopted from Takahashi et al. (1987); the field of olivine in Egyptian ophiolites is drawn based on the published data of Khalil and Azer (2007), Khalil et al. (2014), Gahlan et al. (2015), and Obeid et al. (2016); and the field of Egyptian layered intrusions is based on published data of Ghoneim (1989), Helmy and El Mahallawi (2003), Farahat and Helmy (2006), Azer and El-Gharbawy (2011), Helmy and Mahallawi (2003), Abd El-Rahman et al. (2012), and Azer et al. (2016, 2017). (b) Cr-Al-Fe<sup>3+</sup> plot for chromian spinel and its alteration products in the serpentinized peridotites and listwaenite. (c) Cr# vs. Mg# of chromian spinel cores in serpentinite and Type I listwaenite (adopted from Stern et al. 2004). MORB = mid-ocean ridge basalt. (d) Cr<sub>2</sub>O<sub>3</sub> vs. Al<sub>2</sub>O<sub>3</sub> of fuchsite in Type I listwaenite.

with minor diabantite. Chlorite in Type II listwaenite occurs as sparse crystals and veinlets. Only the veinlet chlorite of veinlets could be analyzed; it has  $FeO^T \ge MgO$  and low  $Cr_2O_3$  (<1.2 wt%) and is classified as pycnochlorite and diabantite.

# Talc

A few analyses of talc from serpentinites are given in Supplementary<sup>1</sup> Table S8. Talc has Mg# of 93.3 to 94.8 and analytical totals of 94.3 to 95.2%.

#### Carbonates

The results of chemical analyses of carbonate minerals are listed in Supplementary<sup>1</sup> Table S7. The loss of total in the analyzed carbonates is due to CO<sub>2</sub>, which is assumed to be present in stoichiometric proportions. The analyses divide the carbonate minerals into magnesite, ferroan magnesite (breunnerite), dolomite, and calcite.

Magnesite in serpentinites and Type I listwaenite essentially consists of MgO (45.9–47.5 wt% in serpentinites and 44.7–48.3

wt% in listwaenite), with less than 1.4 wt% CaO. Magnesite in serpentinite contains distinctly higher MnO (1.1–1.4 wt%) than that in Type I listwaenite (0.3–0.7 wt%). Breunnerite is observed in Type I listwaenite; it is defined by significant MgO (39.5–41.3 wt%) and FeO<sup>T</sup> (10.3–11.2 wt%) contents with very low concentrations of CaO (<0.9 wt%) and MnO (<0.2 wt%). MgO and FeO<sup>T</sup> are negatively correlated among the breunnerite analyses.

Dolomite is observed in serpentinite and in both types of listwaenite. The major components in the analyzed dolomite include MgO (18.7–20.2 wt% in serpentinites, 19.1–19.4 wt% in Type I listwaenite, 19.3–21.0 wt% in Type II listwaenite) and CaO (29.3–30.8 wt% serpentinites, 30.3–31.8 wt% in Type I listwaenite and 29.0–34.6 wt% in Type II listwaenite). Calcite is recorded in both types of listwaenite. It consists essentially of CaO (52.4–57.8 wt%) with minor amounts of MgO (0.4–0.9 wt%) and FeO<sup>T</sup> (1.7–1.9 wt%).

# **GEOCHEMISTRY**

Twenty-one samples (9 serpentinites and 12 listwaenitic rocks) were analyzed to evaluate the whole-rock geochemical characteristics of the G. Sirsir serpentinized peridotites and listwaenites (Table 1). Serpentinites were selected carefully after rigorous petrographic examination to avoid the presence of any carbonate or quartz veins.

The serpentinites of G. Sirsir have 37.0-40.4 wt% SiO<sub>2</sub>, 0.3-1.0 wt% Al<sub>2</sub>O<sub>3</sub>, 38.3-40.1 wt% MgO, and 6.0-7.6 wt%

total Fe as Fe<sub>2</sub>O<sub>3</sub> (Fe<sub>2</sub>O<sub>3</sub><sup>T</sup>). Low-CaO (<0.8 wt%) contents in the serpentinized peridotite suggest that carbonatization had no effect on these samples, despite their proximity to extensively altered talc-carbonates and listwaenites along nearby shear zones. Serpentinites have high volatile contents (presumed to be mostly H<sub>2</sub>O, but including CO<sub>2</sub>) based on their LOI values (12.6–15.4 wt%). The high concentrations of Cr (1519-2814 ppm), Ni (1718–2274 ppm), and Co (96–120 ppm) and low concentrations of K<sub>2</sub>O (<0.02 wt%) and Na<sub>2</sub>O (<0.04 wt%) are all consistent with an ultramafic protolith for the serpentinites. Mg# [100 molar Mg/(Mg+Fe<sup>T</sup>)] of serpentinized peridotites ranges from 90.0 to 92.3 with an average of 91.0. This is generally consistent with oceanic peridotites (Mg#>0.89, Bonatti and Michael 1989) and similar to the Mg# of other reported serpentinites in the Eastern Desert of Egypt (e.g., Azer et al. 2013; Khalil et al. 2014; Gahlan et al. 2015; Obeid et al. 2016).

The population of listwaenite samples encompasses significant variability in all major oxides, in particular SiO<sub>2</sub>, Fe<sub>2</sub>O<sub>3</sub><sup>T</sup>, CaO, MgO, K<sub>2</sub>O, and volatile contents (LOI). The Type I listwaenites are generally lower in SiO<sub>2</sub> and CaO but higher in Al<sub>2</sub>O<sub>3</sub>, MgO, Fe<sub>2</sub>O<sub>3</sub><sup>T</sup>, Na<sub>2</sub>O, K<sub>2</sub>O, and LOI than the Type II listwaenites. The Type I listwaenites are distinctly higher in olivine-compatible trace elements Cr (2147–2765 ppm), Ni (1244–2021 ppm), and Co (51–78 ppm) than the Type II listwaenites (598–1256 ppm Cr, 897–1213 ppm Ni, and 28–63 ppm Co).

Several trace elements (V, Ta, Nb, Th, and Cs) show no systematic differences between the population of serpentinite

TABLE 1. Major and trace elements of serpentinites and listwaenitic rocks of G. Sirsir area, south Eastern Desert, Egypt

Rock type	Serpentinites									Listwaenitic rocks											
											Тур	e I listw	aenite				Туре	II listw	aenite		
Sample no.	SR2	SR12	SR15	SR22	SR31	SR37	SR40	SR44	SR49	SRS	SR7	SR10	SR17	SR20	SR23	SR26	SR28	SR31	SR34	SR46	SR52
									Majo	oxide	s in wt	%									
SiO <sub>2</sub>	40.41	39.86	36.97	37.06	38.47	37.82	38.39	38.05	37.81	38.8	9 41.3	5 40.28	39.72	42.02	46.17	47.57	48.48	56.21	49.36	47.76	45.68
TiO <sub>2</sub>	0.05	0.04	< 0.02	0.03	0.03	0.03	0.02	< 0.02	0.04	0.1	0.17	0.24	0.2	0.15	0.06	0.08	0.28	1.14	0.14	0.09	0.06
$AI_2O_3$	0.54	0.29	0.51	1.01	0.41	0.65	0.78	0.28	0.54	1.2	3 1.99	1.52	2.06	1.71	1.22	1.16	0.93	0.68	0.92	0.74	0.86
Fe <sub>2</sub> O <sub>3</sub> <sup>T</sup>	6.88	6.56	7.62	7.21	7.08	5.96	7.57	6.09	7.13	8.2	8.36	9.09	8.17	7.96	6.43	6.04	5.04	4.03	5.73	6.63	6.34
MnO	0.07	0.07	0.07	0.05	0.12	0.08	0.12	0.09	0.07	0.0	0.13	0.08	0.11	0.13	0.08	0.11	0.17	0.08	0.07	0.06	0.13
MgO	39.06	38.83	39.22	40.13	38.26	39.86	38.42	39.34	39.13	24.3	8 22.2	4 21.53	22.11	20.58	20.84	21.3	17.92	13.44	19.99	21.31	21.72
CaO	0.03	0.31	0.04	0.83	0.05	0.09	0.65	0.07	0.15	6.8	9 5.34	7.06	6.61	7.05	12.58	10.23	12.65	10.19	8.81	8.94	9.93
Na <sub>2</sub> O	0.01	0.04	0.02	0.01	< 0.01	< 0.01	0.01	< 0.01	0.01	0.4	0.96	0.62	1.27	0.86	0.13	0.25	0.08	0.11	0.09	0.15	0.14
K <sub>2</sub> O	0.02	0.01	< 0.01	0.02	< 0.01	< 0.01	0.02	< 0.01	0.01	1.2	1.24	1.87	2.51	1.66	0.11	0.13	0.22	0.56	0.21	0.19	0.12
$P_2O_5$	< 0.01	< 0.01	0.01	0.03	< 0.01	< 0.01	< 0.01	0.02	0.02	0.0	9 0.08	0.06	0.07	0.11	0.04	0.02	0.03	0.05	0.01	0.02	0.03
LOI	12.55	13.44	14.94	13.3	14.84	15.06	13.34	15.43	14.83	17.8	9 18.0	9 17.11	16.89	17.51	11.46	12.48	13.27	14.11	14.78	13.62	14.45
Total	99.62	99.45	99.4	99.68	99.26	99.55	99.32	99.37	99.74	99.5	3 99.9	5 99.46	99.72	99.74	99.12	99.37	99.07	100.6	100.11	99.51	99.46
Mg#	91.0	91.3	90.2	90.8	90.6	92.3	90.0	92.0	90.7	84.	) 82.6	80.9	82.8	82.2	85.2	86.3	86.4	85.6	86.1	85.1	85.9
								Trace	elemer	nts in p	arts pe	r millio	n								
Sc	6.5	5.8	7.6	6.1	7.7	8.3	3.2	2.6	6.1	4.2	10.2	13.5	7.8	11.3	3.7	4.4	5.1	3.2	6.9	4.5	3.3
V	32.4	26.1	31.8	32.7	27.5	33.4	41.2	b.d.l.	35	47	39	50	47	36	30	38	41	34	45	29	33
Cr	2436	1789	2814	2237	2090	2562	1802	1519	2088	224	9 214	7 2316	2471	2765	987	1256	598	1026	815	752	867
Со	96.4	117.2	102.6	105.2	120.3	100.1	118.2	104.6	98	69	58	51	62	78	29	28	32	42	58	63	37
Ni	2058	1718	1984	2005	2132	2134	1921	2274	1931	176	4 197	9 1244	1837	2021	897	1051	1121	956	1214	965	941
Rb	0.2	<0.1	0.1	<0.1	0.5	0.3	0.2	<0.1	0.2	2.3	1.7	4.8	2.5	3.6	1.9	1.2	3.7	5.6	3.4	1.8	1.7
Sr	2	3	5	3	7	0.9	1	4	0.6	112	127	164	142	169	110	123	134	128	102	137	164
Y	0.1	<0.1	<0.1	0.2	0.2	<0.1	<0.1	0.1	0.1	0.8	1.6	0.5	1.1	1.5	0.8	1.4	1.1	0.5	0.7	1.3	0.4
Nb	<0.1	<0.1	<0.1	0.1	0.1	<0.1	<0.1	0.1	<0.1	0.1	<0.7	0.2	<0.1	0.2	0.3	0.4	0.4	<0.1	0.2	<0.1	0.3
Cs	<0.1	0.2	<0.1	0.3	<0.1	0.1	<0.1	0.1	0.2	<0.	0.1	<0.1	0.2	0.1	0.4	<0.1	0.1	<0.1	<0.1	0.3	0.2
Pb	0.2	0.1	0.2	0.3	0.5	0.4	0.5	0.2	0.2	4	3	2	1	3	11	6	5	8	4	7	5
Та	0.1	<0.1	<0.1	0.2	<0.1	0.2	<0.1	0.1	<0.1	0.1	<0.	0.1	<0.1	0.2	0.2	<0.1	0.1	0.2	<0.1	0.1	0.3
Ba	4	7	6	5	5	6	4	3	2	72	94	108	137	114	32	21	35	27	18	22	29
Th	<0.2	<0.2	<0.2	0.3	0.2	<0.2	<0.2	<0.2	0.1	<0.	2 0.2	<0.2	0.3	<0.2	0.2	<0.2	0.2	<0.2	0.3	0.4	<0.2
U	<0.1	<0.1	0.1	0.3	<0.1	<0.1	0.1	<0.1	0.2	0.2	0.4	0.2	<0.1	0.3	0.2	0.5	0.4	0.7	0.3	0.4	0.6
Au (ppb)	10.7	6.5	4.7	6.3	8.1	5.8	4.1	5.8	5.7	144	6 573	871	1157	2008	2011	4987	2231	1854	1502	6584	1871
Sb	1.7	0.9	0.8	1.2	1.4	1.3	0.8	0.7	1.1	0.7	1.9	4.3	6.8	7.5	1.41	2.8	5.6	6.9	8.8	8.7	9.2
As	9.4	8.9	6.2	7.3	5.6	7.8	8.2	6.7	7.4	156	183	211	256	282	223	197	301	326	254	207	186
Cu	53	28	33	40	47	53	31	24	35	32	78	44	56	73	114	95	82	109	77	100	93
Ag	<0.1	0.1	0.2	0.2	<0.1	0.1	0.2	<0.1	0.2	0.3	0.4	0.5	0.6	0.3	0.9	0.7	1.1	1	0.8	1.7	1.4
Zn	39	33	28	32	31	29	41	35	36	65	113	58	107	105	80	113	44	87	103	64	57

samples and either group of listwaenites. However, elevated and variable Rb, Ba, and Sr concentrations are only found in listwaenites; these fluid-mobile elements are at or below detection limits in the serpentinites. Au contents are also variable and highly enriched in both Type I (573–2008 ppb) and Type II listwaenites (1502–6584 ppb) compared to serpentinites (4.1–10.7 ppb). Cu, Zn, Pb, Sb, As, and Ag are enriched in both types of listwaenites relative to the serpentinites, but the highest values of these elements generally occur in the Type II listwaenites, presumably hosted in the abundant sulfide minerals observed in polished section.

Broadly, if we assume that the protoliths of the listwaenites resembled the average serpentinite in bulk composition, then we infer that the alteration and carbonatization processes were accompanied by significant modification of the whole-rock compositions, as discussed below.

#### DISCUSSION

# Protolith and geodynamic setting of serpentinites

A variety of tectonic settings for the Egyptian ophiolites has been proposed, including mid-ocean ridge (MOR) and suprasubduction zone (SSZ) environments. Zimmer et al. (1995) assigned a MOR setting based on the geochemistry of the ophiolitic metavolcanic sequences. Other workers have argued that the tectonic setting of Precambrian ophiolites cannot be accurately defined from metavolcanic bulk composition due to the effects of fractional crystallization and subsequent alteration (e.g., Stern et al. 2004; Azer and Stern 2007). More recently, a SSZ tectonic setting has become the consensus view due to detailed work on the mantle sequences, with disagreement focusing on the more detailed question of back-arc (e.g., El Sayed et al. 1999; Ahmed et al. 2001; Farahat et al. 2004; Azer and Khalil 2005) vs. forearc position (e.g., Azer and Stern 2007; Abd El-Rahman et al. 2009; Azer 2014; Khalil et al. 2014; Gahlan et al. 2015b; Obeid et al. 2016; among others).

Relict textures (bastite and mesh) and chromian spinel morphology (Fig. 4) together with high Cr, Ni, and Co in the G. Sirsir serpentinized mantle lithologies (Table 1) indicate a depleted, harzburgite-dunite complex protolith. Their low modal abundance of pyroxene (10–15 vol%), low MgO/SiO<sub>2</sub> ratios (<1.04), and low TiO<sub>2</sub> contents (<0.05 wt%) are all features of supra-subduction zone depleted peridotites (e.g., Deschamps et al. 2013; Salters and Stracke 2004). More specifically, their very low CaO (<0.65 wt%) and Al<sub>2</sub>O<sub>3</sub> (<0.78 wt%) contents resemble typical fore-arc peridotites (e.g., Ishii et al. 1992) and their very low Al<sub>2</sub>O<sub>3</sub>/SiO<sub>2</sub> ratios (<0.02) also suggest a forearc setting for the partial melting of the protolith. Serpentinization is thought to have negligible influence on these indicators, particularly the Al<sub>2</sub>O<sub>3</sub>/SiO<sub>2</sub> ratio (e.g., Paulick et al. 2006; Deschamps et al. 2013), if quartz veins are avoided.

The chemistry of primary mantle mineral relics such as chromian spinel, olivine, and pyroxene can be used as petrogenetic and geodynamic setting indicators to further elucidate the protolith of the serpentinites (e.g., Dick and Bullen 1984; Barnes and Roeder 2001; Ohara et al. 2002; Coish and Gardner 2004; Sobolev and Logvinova 2005; Arif and Jan 2006; Page and Barnes 2009; Moghadam et al. 2015; Robinson et al. 2015). The high Cr# (0.61–0.74) (Fig. 5c) with low TiO<sub>2</sub> content (<0.2 wt%) of relict primary chromian spinel and high Fo content of coexisting olivine (Fo<sub>89–93</sub>) in both serpentinites and listwaenites argue for a depleted mantle protolith that suffered extensive partial melting in a suprasubduction zone setting, most likely in the fore-arc (Fig. 6) (e.g., Dick and Bullen 1984; Pearce et al. 1984; Ishii et al. 1992; Bloomer et al. 1995; Proenza et al. 2004). This conclusion is consistent with proposed tectonic settings for most Egyptian ophiolites, assigned by several recent authors to incorporated fragments of oceanic lithosphere emplaced above a subduction zone in a forearc setting (e.g., Azer and Stern 2007; Abd El-Rahman et al. 2009; Azer et al. 2013; Khalil et al. 2014; Gahlan et al. 2015b).

The Cr# and Al<sub>2</sub>O<sub>3</sub> values of spinels are commonly used to constrain the nature of mantle peridotites and the degrees of partial melt extraction they have experienced (Dick and Bullen 1984; Hellebrand et al. 2001). Most studies have found that only the divalent cation contents of chromian spinel are vulnerable to subsolidus re-equilibration during hydrothermal alteration and low-grade metamorphism, such that indicators such as Mg# can suggest faulty petrogenetic interpretations (Mellini et al. 2005; Saumur and Hattori 2013; Singh and Singh 2013; Colás et al. 2014; Qiu and Zhua 2017). Spinels with high Cr# ( $\geq 0.6$ ) can be found in residues of high-degree melt extraction and can also crystallize from complementary highly magnesian magmas (Mg-rich tholeiitic or boninitic-type), formed by high-degree partial melting in arc-related tectonic settings (Dick and Bullen 1984; Zhou et al. 1996; Beccaluva et al. 2004). Based on the empirical equation  $[F = 10 \ln (Cr\#) + 24]$  of Hellebrand et al. (2001), the G. Sirsir peridotite protoliths experienced degrees of partial melt extraction ranging from 19 to 21%. Such degrees of melting are again consistent with formation in a forearc setting (e.g., Pearce et al. 2000).

The Mg-rich nature of the analyzed olivines (Fo = 89.4-93.4) is similar to mantle olivine in typical fore-arc peridotites that represent residues after extensive melt extraction (e.g., Pearce et al. 2000; Coish and Gardner 2004). The high NiO contents in the analyzed olivines support their primary mantle character (Takahashi et al. 1987). The plot of Cr# of spinel vs. Fo content of coexisting olivine relics (Fig. 6) shows the consistency of these two minerals in indicating high degrees of melting and forearc affinity (Uysal et al. 2012), probably representing an early stage of development of a subduction zone (Bonatti and Michael 1989; Bloomer et al. 1995).

#### Timing and genetic models of listwaenitization

The serpentinized ultramafics from ophiolitic mantle sections in the Eastern Desert of Egypt are in many cases conspicuously altered and the alteration products provide valuable indicators for mineral exploration. Alteration is attributed to the effects of circulation and infiltration by hydrothermal fluids that can, depending on fluid composition and temperature, produce various assemblages including serpentinite, listwaenite, and talc-carbonate rock. However, the timing of this alteration—relative to other events in the formation and obduction of oceanic crust—is controversial. The origin of fluids that drove alteration of the ANS ophiolites is unclear as well (e.g., Newton and Stern 1990; Stern and Gwinn 1990; Azer 2013; Boskabadi et al. 2017; Hamdy and Gamal El Dien 2017), having been attributed



**FIGURE 6.** Cr# of spinel vs. Fo content of coexisting olivine from serpentinized peridotites (Arai 1992, 1994). PM = primitive mantle; OSMA = olivine-spinel mantle array. The degree of melt extraction is based on spinel composition (after Hellebrand et al. 2001 and Uysal et al. 2012).

variously to: (1) mantle-derived CO<sub>2</sub>-bearing fluids during nearridge oceanic crust formation (e.g., Lebda 1995; Boskabadi et al. 2017; Hamdy and Gamal El Dien 2017); and (2) meteoric and metamorphic hydrothermal fluids penetrating along tectonic fractures during or even after exhumation to upper crustal levels (e.g., Shukri and Lotfi 1959; Salem et al. 1997; Ghoneim et al. 1999, 2003; Hamdy and Lebda 2007). In fact, systematic stable isotope (C, O, H) studies are generally needed to distinguish the temperature of alteration and the composition and source of the metasomatic agents (Fallick et al. 1991; Zedef et al. 2000; Mirnejad et al. 2008; Bjerga et al. 2015; Boskabadi et al. 2017).

In the study area, all occurrences of both Type I and Type II listwaenites are structurally controlled by faults and shear zones, indicating a close link between alteration and the tectonic evolution, metamorphism, and regional geology of the area. Although Abdeen et al. (2008) comprehensively discussed the tectonic evolution of the G. Sirsir area, the position of listwaenite formation in this tectonic evolution has not been considered before. Deformation and schistosity in the Type I listwaenites indicate that the listwaenitization was contemporaneous with serpentinization. On the other hand, the absence of deformation in the Type II listwaenites and their spatial association with nearby granites suggests that they postdate serpentinization.

The fluid CO<sub>2</sub> activity and temperature ranges necessary to drive replacement reactions of the silicates in ultramafic rocks (forsteritic olivine and enstatite-rich orthopyroxene) with carbonates (principally magnesite) has been reviewed by several authors, including the experimental study of Saldi et al. (2012) and Falk and Kelemen's (2015) study of listwaenite along the basalt thrust of the Samail Ophiolite, Oman. The reader is referred to those references for detailed thermodynamic and kinetic discussions.

The structurally distinct deformed (Type I) and undeformed (Type II) listwaenites are also compositionally (Table 1) and mineralogically (Fig. 4) distinct. Type I listwaenite has higher MgO values due to the presence of magnesite as the most common carbonate mineral and high values of  $Fe_2O_3^T$  hosted by ferroan magnesite (breunnerite). The clear enrichment of this group in K<sub>2</sub>O, Na<sub>2</sub>O, and Al<sub>2</sub>O<sub>3</sub> (1.5–2.2 wt%) reflects the presence of a potassium-bearing phase, fuchsite.

The correlation of structurally and compositionally distinct types indicates that hydrothermal listwaenitization took place through two metasomatic stages. The first metasomatic stage was coincident with serpentinization of the original residual peridotites and so likely took place during the development of the ophiolite sequence as fore-arc oceanic crust near a spreading center (~750-700 Ma; Stern and Hedge 1985; Ali et al. 2010). In near-ridge settings, penetrative serpentinization of the uppermost mantle section is thought to be driven by magmatic hydrothermal systems fed by seawater-derived hydrous fluids that gain buoyancy as they are drawn inward and downward toward sites of focused upward discharge. However, noting that forearc settings in particular are positioned above the accretionary prism of a subduction zone, in such cases there is an additional source of CO<sub>2</sub>-rich fluids, generated below the oceanic crust by decomposition of subducted, carbonate-bearing sedimentary rocks at high temperature. These buoyant CO<sub>2</sub>-rich fluids infiltrate upward along fractures and fault planes and can develop a magnesite-talc-quartz-fuchsite (listwaenite) mineral paragenesis locally. As far as can be reconstructed from the coincidence of serpentinite and massive magnesite textures, these two fluid infiltrations-penetrative seawater from above and focused CO<sub>2</sub>-rich fluid from below—occurred simultaneously. In the areas transformed to listwaenite at this stage, all the mafic silicates, primary or secondary, that might later exchange with chromian spinel are removed. Hence, despite a protracted history of cooling after this stage, the Mg# of chromian spinel in listwaenite is not affected by any temperature-sensitive changes in Mg-Fe<sup>2+</sup> distribution coefficients between spinel and mafic silicates. On the other hand, the Mg# of chromian spinel in the host serpentinites continuously evolves by sub-solidus Mg-Fe2+ exchange with mafic silicates as the section of oceanic lithosphere cools and ages.

The second metasomatic stage then took place during the East and West Gondwana continental collision (~650-600 Ma) marked in the field area by development of the Najd fault system and granitic intrusions. During this stage, the Type II listwaenites formed along shear zones from serpentinites bearing chromian spinels whose divalent cation contents had already been reset over the ~100 Ma of subsolidus evolution since their emplacement. Fluids introduced at this stage were highly oxidizing and contributed to oxidation of relict chromian spinel to form ferritchromite and/or Cr-magnetite in both serpentinites and listwaenites.

#### **Chromian spinel alteration**

It is rare for disseminated chromian spinels found in Egyptian ophiolites to preserve zoning profiles in their distributions of Al,  $Fe^{3+}$ , and Cr or of Mg,  $Fe^{2+}$ , and Mn (e.g., Ahmed et al. 2001; Farahat 2008; Azer 2014; Gahlan et al. 2015b; Abdel-Karim and El-Shafei 2017). This suggests an unfortunate gap between the timescales of trivalent and divalent ion redistribution in spinel, with timescales of trivalent ion diffusion generally longer than orogenic events and of divalent ion diffusion generally shorter than events. Nearly complete modification of divalent element distributions during alteration and metamorphism complicates the use of spinel as a petrogenetic indicator (e.g., Barnes and Roeder 2001; González-Jiménez et al. 2009; Proenza et al. 2008; Zhou et al. 1996). However, listwaenite presents a special case. The Mg–Fe<sup>2+</sup> exchange that can occur in serpentinites is arrested in listwaenite because spinel does not readily exchange with carbonate minerals. The chromian spinels in Type I listwaenite are homogeneous in the cores. We argue that the common preservation of unaltered chromian spinel cores with high Mg# in Type I listwaenite implies preservation of high-temperature signatures in the chromian spinel of listwaenite due to early carbonation.

However, Mg# in the chromian spinel cores of Type II listwaenites more closely resembles the values in the serpentinites. This is attributed to ongoing equilibration between chromian spinel and surrounding silicate as the serpentinite protoliths of the listwaenite-like rocks followed the same cooling path as the serpentinites until later carbonate alteration.

Low-temperature serpentinization produces ferritchromite zones with, in the Type II listwaenites, magnetite outer zones, as a result of fluid ingress along cracks and around the grain boundaries of Cr-spinel. During the ferritchromite formation Fe and to a lesser extent Mn are introduced into Cr-spinel, while Al, Mg, and Cr diffuse outward. Hence ferritchromite rims have higher Cr/(Cr+Al) and lower Mg/(Mg+Fe<sup>2+</sup>) than unaltered cores. The depletion of MnO in ferritchromite of Type I listwaenite can be attributed to the presence of carbonate minerals, a favorable sink for Mn (Deer et al. 1992). This indicates that the ferritchromite formed after carbonatization of Type I listwaenites but before carbonatization of Type II listwaenites. The absence of Crmagnetite around the ferritchromite zone of chromian spinel in the Type I listwaenites indicates that the ferritchromite did not re-equilibrate with the surrounding silicates, and consequently Cr-magnetite is absent.

Sofiya et al. (2017) studied listwaenite formed by replacement of serpentinite in the Tulu Dimtu tectonic mélange in Ethiopia, about 1500 km further south along the East African Orogen belt than the G. Sirsir study area. Their petrographic description makes no mention of fuchsite and it appears that all the listwaenite sampled by those authors resembles the Type II listwaenite defined here. Indeed, based on the Mg# of chromian spinel in the Ethiopian case, Sofiya et al. (2017) reach a similar conclusion to ours about the timing of spinel equilibration compared to carbonatization. However, the absence of Type I samples in the Ethiopian case means there is no evidence for the first stage of fluid infiltration at that locality.

Kämmererite is not typically an important product of serpentinization. Its presence in the current suite may instead reflect replacement of chromian spinel during later alteration or regional metamorphism. The significant Cr content of kämmererite and its petrographic relationship to primary relics of Cr-spinel suggest this type of chlorite forms after primary chromian spinel, perhaps during its alteration to ferritchromite. During alteration of chromian spinel, most Cr and Fe enter into ferritchromite, whereas Al and Mg are released to the surrounding silicate minerals. The excess Al and Cr react with serpentine minerals to produce kämmererite (Azer and Stern 2007). Cr-chlorite aureoles are not observed in the Type I listwaenites because the carbonates prevent exchange of Cr with mafic silicate minerals.

# Element mobility during serpentinization and listwaenitization: Application

The chemical changes during the different styles of serpentinite alteration into listwaenite and talc-carbonate rocks are dominated by the addition and removal of CO<sub>2</sub> and H<sub>2</sub>O and by the redistribution of SiO<sub>2</sub>, MgO, and CaO accompanying growth of carbonate minerals at the expense of silicates. Although Mg<sup>2+</sup> and Ca<sup>2+</sup> can be leached during hydrous fluid alteration, the high activity of CO<sub>2</sub> in the present case instead stabilizes Mg- and Ca-carbonate phases that reduce the mobility and loss of these cations. Alteration also caused redistribution of trace elements, with some being locally remobilized within the rock, some being added from a fluid phase, and others being leached out of the rock. Serpentinites were enriched in fluid-mobile elements (K, As, Rb, Sr, Sb, Cs, Ba, Pb, and U) relative to primitive (McDonough and Sun 1995) or depleted mantle values (Salters and Stracke 2004), first due to modification in the mantle wedge (e.g., Hattori and Guillot 2003, 2007; Deschamps et al. 2011, 2012) and perhaps also from serpentinizing fluids; in any case they were retained in the serpentine minerals (Deschamps et al. 2013). Both types of listwaenite show further enrichment of these elements, suggesting both efficient retention by the carbonates and additional contribution from the CO<sub>2</sub>-rich alteration fluid.

Listwaenite rocks have long been considered an indicator of Au mineralization, although the relationship between Au enrichment and listwaenite formation is still unclear. Gold shows similar behavior, to some extent, to the fluid-mobile elements. Its concentrations in the G. Sirsir serpentintes are commonly higher than in typical serpentinized mantle peridotite (i.e.,  $\sim 3-5$ ppb; Buisson and Leblanc 1987), average ophiolitic rocks (i.e.,  $\sim 2.8$  ppb; Crocket 1991), or depleted mantle (i.e.,  $\sim 1$  ppb; Salters and Stracke 2004). Compared to serpentinites, the listwaenite rocks of the G. Sirsir then show much higher concentrations of Au (673–2008 ppb in Type I listwaenites and 987–6584 ppb in Type II listwaenites), suggesting that intense alteration of the serpentinites to form listwaenite within shear zones was accompanied by fluid concentration that enriched these rocks in Au (as well as As and Ag).

The current results are comparable to other studies worldwide that have demonstrated hosting of Au mineralization in listwaenite (e.g., Buisson and Leblanc 1985, 1986, 1987; Ash and Arksey 1990; Aydal 1990; Koç and Kadioglu 1996; Uçurum and Larson 1999; Uçurum 2000; Belogub et al. 2017) as well as specifically within the Eastern Desert of Egypt (Botros 1993, 2002; Osman 1995; Oweiss et al. 2001; Ramadan et al. 2005; Zoheir and Lehmann 2011; Azer 2013). Some authors have argued that the relation between gold mineralization, listwaenitization, and shear zones reflects their combination of weak shear strength and high permeability, in strong rheological contrast with more coherent rocks such as the granitoids commonly found in contact with listwaenites in ANS gold deposits (e.g., Buisson and Leblanc 1987; Zoheir and Lehmann 2011; Azer 2013; Zoheir and Moritz 2014).

#### Chlorite geothermometry

Compositions of chlorite are sensitive to prevailing physicochemical conditions during its formation because distribution coefficients between precursor ferromagnesian minerals and chlorite vary, especially as a function of temperature (deCaritat et al. 1993). Therefore, the chemical composition of chlorite has been used to determine the temperature of its formation by replacement reactions (e.g., Kranidiotis and MacLean 1987; Cathelineau and Nieva 1985; Cathelineau 1988; Hillier and Velde 1991; Bourdelle et al. 2013; Bourdelle and Cathelineau 2015; Yavuz et al. 2015). Kranidiotis and MacLean (1987) corrected the linear correlation shown by Cathelineau and Nieva (1985) between tetrahedral Al and temperature in chlorites saturated with an Al-rich phase with as estimate of the effect of Fe/(Fe+Mg) on the temperature. The resulting equation is suitable for chlorite that coexists with aluminous spinel or muscovite. The opposite, Al-undersaturated boundary of the chlorite stability field lies about 50 °C higher; given the presence of Cr in the coexisting spinels and muscovites in most of the analyzed rocks, it is possible that the Kranidiotis and MacLean (1987) thermometer systematically underestimates chlorite temperatures in these rocks by ~10 °C. The calculated temperatures for chlorite formation according to the geothermometer equation of Kranidiotis and MacLean (1987) are listed in Supplementary<sup>1</sup> Table S6; the equation is given as a footnote to the Table. The temperatures obtained are higher in serpentinites (286-311 °C in aureoles and 277-300 °C in disseminated chlorite) than in Type I listwaenite (243-274 in chlorite associated with fuchsite and 229-270 °C in disseminated chlorite) or Type II listwaenite-like rock (233-249 °C). Interestingly, Fe-rich chlorites in Type II listwaenite-like rocks give the lowest temperature range (avg. 240 °C), suggesting they sample a unique hydrothermal stage, likely the last to affect the G. Sirsir bodies.

Zoheir (2011) obtained quite similar temperatures (~270–340 °C) for chlorite in listwaenite in south Eastern Desert of Egypt. We note that the chlorite temperatures from both our study and that of Zoheir (2011) are plainly higher than the inferred maximum temperatures of ~65 °C for listwaenite formation proposed by Wilde et al. (2002) or the magnesite clumped isotope temperatures of 65–114 °C for Oman ophiolite basal listwaenite inferred by Falk and Kelemen (2015), whereas they are much more similar to temperatures estimated by numerous authors for other listwaenites worldwide (e.g., Andrew 1985; Buisson and Leblanc 1987; Weir and Kerrick 1987; Madu et al. 1990; Spiridonov 1991; Schandl and Wicks 1993; Schandl and Gorton 2012; Oskierski et al. 2013).

#### IMPLICATIONS

The G. Sirsir serpentinized ultramafics form tectonic sheets and lenses within a tectonic mélange, elongated in the northwestsoutheast (NW–SE) direction. They are extensively altered along thrust and shear zones into quartz-carbonate rocks (listwaenite assemblages) and talc-carbonates. Chromian spinel relics in both serpentinites and listwaenites apparently preserve pristine ratios of trivalent cations, with high Cr# typical of spinel in residual peridotites. However, relict chromian spinel in the Type I listwaenite has significantly higher Mg# and lower MnO than that of serpentinites, suggesting nearly complete alteration of ultramafic rocks to form listwaenite took place prior to re-equilibration between chromian spinel and the surrounding mafic minerals in serpentinites. The lower Mn-content of ferritchromite rims in the listwaenites compared to the serpentinites indicates this stage of spinel alteration also postdates carbonatization in the listwaenites, since carbonate minerals are favorable sinks of Mn.

Based on field and textural observations, it is very likely that carbonation affected serpentinites through two stages to form the two types of listwaenites. The first stage was apparently contemporaneous with serpentinization and emplacement of the mantle section of the G. Sirsir ophiolite into oceanic lithosphere. The second stage (~650–620 Ma), forming the unsheared Type II listwaenite-like rocks, is younger than the presumed protolith age and associated with the Najd fault system, collisional tectonics, and ophiolite obduction. The implication is that both these stages of deformation were associated with infiltration of CO<sub>2</sub>-rich fluids. Carbonate source rocks likely decomposed to release such fluids both from the slab underlying the original forearc setting of spreading and ophiolite formation and again in the footwall of the structures that accommodated ophiolite obduction onto a continental shelf. At some level, then, both stages of listwaenitization and associated Au mineralization can be traced to the tropical setting of the Mozambique Ocean and the resulting abundance of carbonate sediments in the marine sequences trapped within the Pan-African orogeny.

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#### **Endnote:**

<sup>1</sup>Deposit item AM-18-76473, Supplemental Tables. Deposit items are free to all readers and found on the MSA web site, via the specific issue's Table of Contents (go to http://www.minsocam.org/MSA/AmMin/TOC/2018/Jul2018\_data/Jul2018\_data.html).