Hydrothermal alteration of Ni-rich sulfides in peridotites of Abu Dahr, Eastern Desert, Egypt: Relationships among minerals in the Fe-Ni-Co-O-S system, f_{O_2} and f_{S_2}

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ABSTRACT

The Neoproterozoic peridotites of Abu Dahr, Eastern Desert of Egypt, consist mainly of highly depleted harzburgites that have experienced multiple stages of serpentinization (lizarditization and antigoritization) and carbonation/listvenitization in a forearc environment. The Abu Dahr forearc harzburgites are more oxidized than oceanic mantle, with the oxygen fugacity (f_{02}) values ranging from FMQ+0.41 to FMQ+1.20 (average = +0.60 FMQ), and were equilibrated at temperatures of 910-1217 °C and pressures of 4.1-7.8 kbar. This study has documented for the first time the presence of various Ni-rich Ni-Fe (-Co) sulfide and metal phases along with Fe-oxides/oxyhydroxides in serpentinized-carbonated peridotites of the Abu Dahr forearc. Here I concentrate on the relationship between redox state and Fe-Ni-Co-O-S minerals with emphasis on the role of hydrothermal processes in upgrading magmatic sulfide tenors, desulfurization (sulfur-loss) of magmatic pentlandite and hydrothermal upgrading of the sulfide phases in Abu Dahr forearc environment. The minerals involved are high-Ni pentlandite ($Fe_4Ni_5S_8$), cobaltian pentlandite ($Fe_{3.47}Ni_{4.78}Co_{0.75}S_8$), heazlewoodite ($Fe_{0.07}Ni_{2.93}S_2$), godlevskite (Fe_{0.26}Ni_{8.73}Co_{0.01}S₈), millerite (Fe_{0.01}Ni_{0.98}Cu_{0.01}S), awaruite (Ni₇₅Fe₂₁) and native Ni (Ni₉₃Fe₃), and nickeliferous magnetite and goethite. Chalcopyrite is a rare mineral; other Cu-phases, Fe-sulfides and Ni-arsenides/phosphides are not present. Texturally, Ni-sulfide and alloy minerals occur as interstitial disseminated blebs of either solitary phases or composite intergrowths with characteristic replacement textures, documenting strong variations in oxygen and sulfur fugacities (f_{02} - f_{52}). Sulfide assemblages are divided into three main facies: (1) pentlandite-rich; (2) godlevskite-rich; and (3) millerite-rich. Textural relationships imply the following sequence: (a) primary pentlandite \rightarrow cobaltian pentlandite, with partial replacement of the latter by awaruite and/or heazlewoodite along with magnetite; (b) heazlewoodite is replaced by godlevskite, which in turns is replaced by millerite; (c) Ni-rich awaruite breaks down to millerite; and finally, (d) magnetite is completely replaced by goethite. The sulfide mineralogy reflects the magmatic and post-magmatic evolution of the complex. The primary magmatic processes gave rise to pentlandite, whereas the secondary Ni-sulfides together with the metallic alloys formed in response to changing f_{0} , and f_{s} , conditions associated with post-magmatic serpentinization and carbonation. Serpentinization-related Ni-Fe-Co remobilization from magmatic olivines resulted in; (1) upgrading the Ni-Co tenors of pre-existing primary pentlandite, and desulfidation to form low-sulfur sulfides (mainly heazlewoodite) and awaruite under extremely low f_{o_2} and f_{s_2} conditions; (2) in situ precipitation of secondary Ni-sulfides in the presence of extra sulfur as aqueous H_2S derived from the desulfurization of magmatic pentlandite or native Ni when f_{s_2} approaches 0; (3) transformation from low-sulfur pentlandite- and godlevskite-rich assemblages to the high-sulfur millerite-rich assemblages related to later carbonation with increasing f_{07} ; and (4) partial dehydration of antigorite serpentinites under high-pressure conditions (>1 GPa) generated Ni-rich awaruite in equilibrium with the prograde assemblage antigorite-metamorphic olivine at higher f_{02} and f_{52} within subduction channel. The mineralogical, chemical, and thermal similarities with other serpentinite-related Ni-sulfides worldwide suggest that Ni minerals in the Fe-Ni-Co-O-S system record changing f_{O_2} and f_{S_2} during progressive serpentinization and carbonation.

Keywords: Abu Dahr forearc, Ni-rich sulfides, oxygen and sulfur fugacities, replacement textures, serpentinization and carbonation, desulfurization, subduction channel.

INTRODUCTION

Peridotite serpentinization and carbonation play important roles in upgrading the mineralogy and tenor (i.e., the Ni content of the sulfide fraction) of pre-existing primary Ni-sulfides. Serpentinization is accompanied by the formation of magnetite and release of Ni, Fe, and Co from the primary silicates (e.g., olivine) (Groves et al. 1974; Eckstrand 1975; Donaldson 1981; Kamenetsky et al. 2016). Ni released from olivine during serpentinization may be added to or substituted for Fe in the pre-existing sulfide phases (e.g., pentlandite), producing high-Ni tenor pentlandite and/or precipitation of new hydrothermal sulfides when enough H₂S is available (e.g., Eckstrand 1975; Donaldson 1981; Keays and Jowitt 2013). Sulfide and metal mineralogy during hydrothermal alteration reflect strong variations in fluid redox conditions, i.e., variations in oxygen fugacity (f_{0_2}) and sulfur fugacity (f_{s_2}) (Frost 1985; Alt and Shanks 1998; Marques et al. 2007; Klein and Bach 2009; Schwarzenbach et al. 2012, 2014; 2021; Foustoukos et al. 2015; de Obeso and Kelemen 2020). Initial serpentinization, with relict primary mantle minerals (olivine, orthopyroxene, and clinopyroxene) along with the low-sulfur assemblage pentlandite-heazlewoodite-awaruite

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represents a highly reducing and extremely low f_{02} and f_{52} environment (Eckstrand 1975; Alt and Shanks 1998; Schwarzenbach et al. 2014). Completely serpentinized peridotites preserve high-sulfur assemblages and magnetite (Eckstrand 1975; Alt and Shanks 1998; Schwarzenbach et al. 2014). Talc-carbonate alteration also results in the enrichment of Ni in talc and, at high sulfur activity, millerite grows at the expense of the low-sulfur sulfides (heazlewoodite and godlevskite) (e.g., Donaldson 1981; Marques et al. 2007; Klein and Bach 2009; Schwarzenbach et al. 2014). Therefore, hydrothermal Ni-upgrading might cause the ophiolitic serpentinized peridotites to be viewed as prospective for Ni deposits.

Several serpentinite-hosted Ni deposits such as the Dumont Sill, Quebec (Eckstrand 1975; Sciortino et al. 2015), Bou Azzer, Morocco (Ahmed et al. 2009), Avebury Ni sulfide deposit, Tasmania (Keays and Jowitt 2013; Kamenetsky et al. 2016), Hayachine, Japan (Shiga 1987), Eastern Metals, Canada (Auclair et al. 1993), Epoch deposit, Zimbabwe (Pirajno and González-Álvarez 2013), Doriri Creek deposit, Papua, New Guinea (González-Álvarez et al. 2013), Elizabeth Hill, Western Australia (Hoatson and Sun 2002), Black Swan, Western Australia (Barnes et al. 2009), and Sarah's Find Ni-Cu-(PGE) deposit, Western Australia (Le Vaillant et al. 2016) have been interpreted as having hydrothermal origins.

The oxidation state of the forearc mantle wedge is attributed either to oxidizing components, derived from subducted slab and added to the mantle wedge cause metasomatic changes and/or partial melting (Evans 2006, 2012; Bénard et al. 2018; Kilgore et al. 2018), or to deserpentinization (Evans and Frost 2021). The redox budget (Evans 2006, 2012; Evans and Tomkins 2011; Malaspina et al. 2009, 2017) of subducted slab-derived fluids could change the redox state of the mantle wedge by transporting such redox sensitive elements as iron, carbon and sulfur into it (Rielli et al. 2017; Bénard et al. 2018; Debret et al. 2020). Generally, the f_{02} of the mantle wedge is higher than that of the oceanic mantle (Malaspina et al. 2009; Evans et al. 2012, 2017; this study).

In this study, I contribute new mineralogical and f_{02} information to constrain redox conditions in the Fe-Ni-Co-O-S system and the role of hydrothermal processes in upgrading magmatic sulfide tenors, desulfurization (sulfur-loss) of magmatic pentlandite and hydrothermal upgrading of the sulfide phases in Abu Dahr forearc environment. Then, I correlate these data with those from global Ni-rich serpentinites.

The redox conditions, expressed in terms of f_{0_2} and f_{s_2} , are controlled by the mineralogy and stabilities of Ni-rich phases in serpentinization environments (Frost 1985; Klein and Bach 2009; Schwarzenbach et al. 2014, 2021; Foustoukos et al. 2015; de Obeso and Kelemen 2020). In subduction zones, slab-derived fluids migrate upwards and cause serpentinization of the adjacent mantle wedge and subsequent partial melting (Zheng et al. 2020). The products of serpentinization depend on the primary mineralogy of the protolith, water/rock ratios, temperature, pressure and fluid composition (e.g., Frost 1985; Frost and Beard 2007; Schwarzenbach et al. 2014). Variations in water/rock ratios lead to strong variations in f_{02} and f_{S2} conditions (Frost 1985; Bach et al. 2006; Delacour et al. 2008; Ishimaru et al. 2009; Klein and Bach 2009; Frost et al. 2013; Schwarzenbach et al. 2014; Evans et al. 2017; de Obeso and Kelemen 2020). Serpentinization at low water/rock ratios commonly leads to lower f_{O_2} and f_{S_2} and favors the formation of low-sulfur sulfide assemblage of heazlewoodite, Fe-Ni alloys (e.g., taenite, awaruite), native metals, magnetite and hydrogen (Frost 1985; Alt and Shanks 1998, 2003; Delacour et al. 2008). In contrast, serpentinization at higher water-rock ratios leads to higher f_{02} with an assemblage of millerite, pyrite, hematite, and sulfate (Frost 1985; Alt and Shanks 1998, 2003; Delacour et al. 2008). The sulfide mineral assemblages provide constraints on f_{02} and f_{52} prevailing during hydrothermal alteration (Alt and Shanks 1998, 2003; Delacour et al. 2008 Schwarzenbach et al. 2012, 2014, 2021).

This study has documented for the first time the presence of various Ni-rich Ni-Fe(-Co) sulfide and metal phases along with Fe-oxides/oxyhydroxides in serpentinized-carbonated peridotites of Abu Dahr forearc. The main goals of this study are to: (1) describe the mineralogy, mineral assemblages and textures of the high-Ni sulfide and intermetallic minerals in Abu Dahr as potentially major Ni suppliers for Ni resource exploration, compared with other Ni-rich serpentinites worldwide; (2) identify the paragenetic sequence of the Ni-rich phases based on mineral textures and associations; (3) determine the composition of the Ni-rich phases and their relationship to f_{02} and f_{52} conditions; and (4) discuss the role of hydrothermal processes in upgrading disseminated magmatic sulfide tenors, desulfurization of magmatic sulfides and secondary upgrading of the sulfide phases, and their reflection on the magmatic and post-magmatic evolution of the Abu Dahr complex.

GEOLOGICAL SETTING AND SAMPLES

The Abu Dahr area (23° 30'-23° 40' N, 35° 00'-35° 10' E, Fig. 1) is located in the South Eastern Desert of Egypt, ~70 km NW of Shalatin city on the Red Sea, and covers the largest part of Beitan province. It forms a part of the Arabo-Nubian Shield (890-570 Ma, Stern et al. 2004) and comprises Neoproterozoic Beitan gneisses, dismembered ophiolite, island arc metavolcanic-plutonic assemblage, volcaniclastic metasediments (mélange matrix), and syn- to late-tectonic granite intrusions (Ashmawy 1987; Abdel-Khalek et al. 1992; El Tahlawi et al. 1997; Zoheir et al. 2008; Khedr and Arai 2013, 2016; Gahlan et al. 2015; Abdel-Halim et al. 2020; Khedr et al. 2022). According to field relationships, the following sequence was established by El Tahlawi et al. (1997) starting with the uppermost rock unit: (1) Abu Dahr Nappe, (2) Arayis ophiolitic mélange, (3) Egat metavolcaniclastics, and (4) Beitan gneisses. The Abu Dahr ophiolite consists, from the base upward, of a mantle section (Abu Dahr Peridotite Nappe), ultramafic-mafic cumulates, and crustal metagabbros and metabasalts (Gahlan et al. 2015; Abdel-Halim et al. 2020; Khedr et al. 2022). The crustal rocks are associated with the Arais and Balamhindit ophiolite complexes (Khedr and Arai 2013, 2016).

The Peridotite Nappe of Abu Dahr (~150 km²) is the second largest, well-exposed and preserved Neoproterozoic ultramafic rocks (after El Gerf) in the South Eastern Desert, Egypt. The Abu Dahr Nappe is composed mainly of highly depleted clinopyroxene-poor harzburgites with minor dunites (+ chromitites) and pyroxenites. The nappe, structurally, forms the uppermost lithotectonic unit, thrusted southwestwards onto Beitan gneisses. It is divided into two major domains separated by EW-trending Abu Khayl fracture zone: Abu Dahr North and South (Abdel-Halim et al. 2020). The Abu Dahr North consists of harzburgite with minor dunite (+ chromitite), cut by a network of olivine–orthopyroxenite dikes. The harzburgites are partially serpentinized to form lizardite-bearing serpentinites (Lz-serpentinites), but pristine harzburgites are more abundant at **FIGURE 1. (a)** Distribution of ophiolitic rocks in the Eastern Desert of Egypt (after Shackleton 1994). (b) Geological map of Abu Dahr [(modified from the Geologic Map of Baranis, Geological Survey of Egypt (1992), Scale 1:250 000]. Numbers indicate locations of the observation points sampled. (c) Thematic Mapper of Abu Dahr massif for possible lithological discrimination.



Wadi Abu Mastoura (~4 km long). The chromitites are massive to disseminated high-Cr chromitites composed of magnesio-chromite with high Cr# values (83-93) containing inclusions of platinumgroup minerals (PGM) + base-metal sulfides (BMS) + Au (Abdel-Halim et al. 2020). Massive chromitites form large pods within dunite "envelopes" in harzburgites host. Magnesite veins (up to 1 m wide) are common in the serpentinized harzburgites. Gabbroic and doleritic dikes are also encountered. Antigorite serpentinites (Atg-serpentinites) constituting Abu Dahr South were transformed to carbonate-rich serpentinites (listvenites) along fractured and sheared zones. The Abu Dahr Nappe is intruded by concentric intrusive complexes of Um Eliega and Rahaba from the north and east, respectively (Zoheir et al. 2008; Abdel-Karim et al. 2021; Azer and Asimow 2021). These complexes consisting of a mafic core (gabbro and diorite) surrounded by tonalite in the middle to granodiorite at the margin, formed in an active continental margin setting during the subduction of the oceanic slab (Abdel-Karim et al. 2021).

The samples studied were collected from surface outcrops of northern and southern Abu Dahr. The sample localities are shown in Figure 1 and their coordinates are listed in Table 1. The samples include harzburgites, dunites, chromitites, and orthopyroxenites. The harzburgites considered here, are partially to completely serpentinized.

Analytical Methods and Calculations of P-Tand f_{O_2}

More than 200 polished thin sections (PTS) were investigated at various magnifications (100 and 200×) using dual-purpose optical microscope incorporating both transmitted- and reflected-light options and scanning electron microscope (SEM) hosted in the Department of Geology at Assiut University, Egypt. The modal abundances of sulfides were determined by point counting technique at 500× magnification.

The mineral chemistry of Ni-sulfides, metal alloys, olivines, pyroxenes, amphiboles, Cr-spinels and Fe-oxides/oxyhydroxides was determined by Electron Microprobe (EMP) analysis using a JEOL (JSM-6310) instrument by wavelengthdispersion spectroscopy (WDS) and energy-dispersive spectroscopy (EDX) at the Institute of Earth Sciences, Karl Franz University, Graz, Austria. Operating conditions of the electron beam were 20 kV accelerating potential, 20 nA beam current, 1 μ m beam diameter, and 10 s measurement time (peak) for Cu, Co, Fe, Mg, Ni, O, S and Si. Natural and synthetic silicate and oxide standards were used for calibration. Silicate standards were quartz for Si; titanite for Ca and Ti; synthetic corundum for Al; garnet for Fe; olivine for Mg; jadeite for Na; adularia for K; chromite for Cr; rhodonit for Mn; and nickel oxide for Ni. Standards for base-metal minerals were chalcopyrite for S, Fe, and Cu; cobalt for Co; nickel for Ni; silver for Ag; Cabri-141 (NiSb) for Sb; and Cabri-526 (PtAs2) for As. The relative analytical error is better than 1% (1 σ)

Major- and trace-element contents of bulk-rocks were determined by X-ray fluorescence spectrometry (XRF) and inductively coupled plasma mass spectrometry (ICP-MS) at the Institute of Earth Sciences, Karl Franz University, Graz, Austria. Major elements were determined by XRF using fused glass disks, excited with X-ray radiation, normally generated by an X-ray tube operated at a potential of between 10 and 100 kV, with relative precision of <1%. The analytical error is less than 0.6 wt% for all major elements. Loss on ignition (LOI) was calculated after heating the powdered samples at 1050 °C for 2 h. Trace-element concentrations were measured by ICP-MS using an Agilent 7500 s quadrupole ICP-MS system attached to a New Wave/Merchantek UP213 laser ablation system. The analyses were carried out with a beam diameter of 15 µm and a 5 Hz repetition rate. Helium was used as carrier gas and N2 was added to optimize the detection limits. The energy density was 4 J/cm², and the repetition rate was set at 6 Hz. Quantitative results were obtained through calibration of relative element sensitivities using the NIST-610 standard glass. The analytical error is less than 0.1 ppm by weight for most elements.

Calculations of P-T and f_{02} . Equilibration temperatures for the studied harzburgites were determined using the two-pyroxene thermometers (Wood and Banno 1973; Wells 1977; Brey and Köhler 1990) and Ca-in-orthopyroxene thermometer of Brey and Köhler (1990). The two-pyroxene geobarometer suggested by Putirka (2008) was used to determine the pressure conditions under which the peridotites could have formed. The f_{02} of the Abu Dahr harzburgites was determined using the oxybarometry of Ballhaus et al. (1991) based on spinel-olivine-orthopyroxene oxybarometry equilibrium at a pressure of 1 GPa, which is approximately the center of the spinel stability field (see Fig. 13 of Simon et al. 2008; Dare et al. 2009; Birner et al. 2016), and adopted Tk from the equilibrium temperatures of Brey and Köhler (1990) in f_{02} , calculations.

The f_{o_2} of peridotites can be calculated using phase equilibrium between olivine (Ol), orthopyroxene (Opx), and spinel (Spl) (R1):

$$6Fe_2SiO_4(Ol) + O_2 = 3Fe_2Si_2O_6(Opx) + 2Fe_3O_4(Spl).$$
(1)

Location	OP	Latitudes	Longitudes	No. of
sampled		North	East	samples
Abu Dahr North	300	23° 36′ 36″	35° 04′ 38″	19
	301	23° 36′ 40″	35° 04′ 42″	8
	302	23° 36′ 41″	35° 05′ 53″	12
	303	23° 36′ 42″	35° 05′ 56″	9
	304	23° 36′ 40″	35° 05′ 57″	8
	305	23° 36′ 40″	35° 05′ 57″	17
Wadi Abu Khayl	306	23° 34′ 16″	35° 07′ 02″	24
	307	23° 34′ 29″	35° 04′ 59″	5
	308	23° 33′ 24″	35° 06′ 46″	25
	309	23° 33′ 21″	35° 06′ 49″	23
	310	23° 32′ 53″	35° 07′ 07″	2
	311	23° 33′ 03″	35° 07′ 15″	4
	312	23° 32′ 58″	35° 07′ 26″	4
	313	23° 32′ 57″	35° 07′ 26″	3
	314	23° 32′ 38″	35° 07´ 46″	3
	315	23° 32′ 44″	35° 07′ 54″	1
Wadi Abu Mastoura	316	23° 35′ 19″	35° 05´ 22″	20
	317	23° 35′ 10″	35° 05´ 28″	6
	318	23° 35′ 21″	35° 05′ 12″	16
	319	23° 35′ 32″	35° 04′ 54″	11
	320	23° 35′ 38″	35° 04´ 48″	3
	377	23° 35′ 10″	35° 05′ 48″	8
	378	23° 35′ 12″	35° 05′ 43″	3
	379	23° 35′ 22″	35° 05´ 21″	6
	380	23° 35′ 30″	35° 05′ 14″	4
	381	23° 35′ 31″	35° 05´ 08″	8
	382	23° 35′ 33″	35° 05´ 00″	6
	383	23° 35′ 41″	35° 04′ 39″	7
	384	23° 35′ 52″	35° 04´ 17″	2
Wadi Abu Sayyal	321	23° 34′ 01″	35° 00′ 53″	23
	322	23° 34′ 26″	35° 01´ 05″	15
	323	23° 34′ 22″	35° 01′ 28″	9
	324	23° 34′ 28″	35° 01' 21"	17
	325	23° 34′ 40″	35° 01′ 38″	5
	326	23° 30′ 38″	35° 06′ 47″	5
	327	23° 30′ 38″	35° 06′ 46″	15
	328	23° 30′ 45″	35° 07′ 06″	4
	329	23° 31′ 09″	35° 08′ 00″	18
Gebel Abu Savval	330	23° 35′ 47″	35° 01′ 09″	34
,,u	331	23° 35′ 43″	35° 01′ 13″	22

 TABLE 1. GPS coordinates of the observation points (OP) sampled from

 Abu Dabr peridotites

The calculation of f_{0_2} depends on the pressure and temperature of equilibrium. For this study, we used the two-pyroxene thermometers (Wood and Banno 1973; Wells 1977; Brey and Köhler 1990) and Ca-in-orthopyroxene thermometer of Brey and Köhler (1990). The f_{0_2} values are calculated relative to the fayalite-magnetitequartz (FMQ) buffer in log units using the olivine-spinel oxybarometry of Ballhaus et al. (1991) and the formula of them assuming a pressure of 1 GPa as follows (R2):

$$\begin{aligned} & \Delta \log(f_{02})^{FMQ} = 0.27 + 2505/T - 400P/T - 6 \log (XFe^{01}) - 3200(1 - XFe^{01})^2/T + \\ & 2\log (XFe^{2+Sp1}) + 4 \log (XFe^{3+Sp1}) + 2630 (XAI^{Sp1})^2/T. \end{aligned}$$

The uncertainty in the calculated f_{0_2} originates from analytical errors and errors associated with activity estimates of components in Equation 1. Analytical errors in silicate compositions contribute little to the f_{0_2} uncertainty. For example, olivine composition has an error of less than ±0.003 in XFe, which translates into an uncertainty of ±0.15 log units in f_{0_2} . Orthopyroxene grains are more heterogeneous in composition than olivine in individual samples; however, propagated uncertainties from this heterogeneity are also small, on the order of 0.1 log units in f_{0_2} or less. Inaccurate Fe³⁺ contents in spinel are the major cause of error in calculated f_{0_2} . The oxygen fugacity recorded by mantle minerals is unaffected by nearby low-temperature serpentinization, and is representative of mantle processes (Birner et al. 2016).

RESULTS

Peridotite petrography

The harzburgites show a wide range of serpentinization from 0% to 100%, and exhibit a sequence of hydrothermal alteration from harzburgites \rightarrow Lz-serpentinites \rightarrow Atg-serpentinites \rightarrow Atg-talc-magnesite \rightarrow quartz-carbonate listvenites. Figure 2 shows the paragenetic sequence for Abu Dahr ophiolitic rocks and associated

silicates, carbonates, oxides, and Ni-sulfides and metal alloys. The fresh harzburgites show a typical porphyroclastic texture, characterized by millimeter-sized forsterite-rich olivine (Mg# is 91-93), orthopyroxene with lesser amounts of clinopyroxene lamellae (<1 vol%), and Cr-spinel, as well as Ca-amphiboles (magnesiohornblende and tschermakite). Orthopyroxene porphyroclasts are surrounded by domains of polygonal or irregularly shaped olivine neoblasts (Figs. 3a and 3b). Lz-serpentinites from Abu Dahr North are primarily composed of low temperature serpentine (i.e., lizardite and/or chrysotile) forming mesh and bastite textures after olivine and orthopyroxene, respectively, with disseminated magnetite and interstitial sulfides (Figs. 3c and 3d). Brucite is developed in the centers of narrow serpentine veinlets, and occurs as thin veins associated with serpentine and/or magnetite (Fig. 3d), similar to brucite texture from Santa Elena (see Fig. 8 of Schwarzenbach et al. 2016) and Mariana (see Fig. 2 of Albers et al. 2020). Cr-spinel is virtually unaltered but some grains exhibit variably thick ferritchromite rims. Atg-serpentinites from Abu Dahr South consist of ~100% antigorite with an interlocking texture, minor carbonate (magnesite and dolomite), Fe-oxides/oxyhydroxides, talc, and Ni-sulfides and Ni-rich awaruite (Fig. 3e). Well zoned chromites with ferritchromite and nickeliferous magnetite rims are widely distributed in samples from both areas (Fig. 3f).

Disseminated Ni mineralization of Abu Dahr

Primary sulfides. The major sulfides of mantle rocks, pentlandite, pyrrhotite, and chalcopyrite are present in low abundance and occur interstitially among the silicates (e.g., Harvey et al. 2016; Lorand and Luguet 2016; Kiseeva et al. 2017; Vaughan and Corkhill 2017). At Abu Dahr, pentlandite (Fe,Ni,Co)₉S₈, the dominant primary mantle sulfide mineral, occurs as single mineral grains (pentlandite/cobaltian pentlandite) from 10 to 2000 µm in size (Figs. 4a and 4e), and/or as composite intergrowth (10–50 µm across) of pentlandite and its replacement secondary phases (Figs. 5a, 5b, and 5a). Chalcopyrite is a rare mineral; other Cu-phases, Fe-sulfides, and Ni-arsenides/phosphides are not present in the investigated samples.



FIGURE 2. The paragenetic sequence for the Abu Dahr ophiolitic rocks and associated silicates, carbonates, oxides, and Ni-sulfides and native alloys. Mineral abbreviations: see Table 2.



FIGURE 3. Optical (a and b) and backscattered electrons (c-f) images showing microtextures of Abu Dahr peridotites. (a and b) Harzburgites showing a typical porphyroclastic texture, characterized by millimeter-sized porphyroclasts of olivine and orthopyroxene embedded in a fine-grained neoblastic matrix. Olivine occurs as small inclusions in orthopyroxene. Recrystallized grains of olivine usually rimming olivine and orthopyroxene porphyroclasts (0.5 cm). Unaltered chromite grain with orthopyroxene and amphibole inclusions. (c and d) Partially serpentinized harzburgites showing meshtextures typical of serpentine (lizardite and/or chrysotile) forming mesh textures after olivine. Alongside serpentine, secondary brucite, magnetite, sulfides and native alloys, and are present. Brucite is developed in the centers of narrow veinlets during the initial serpentinization. (e) Atg-serpentinites consist of ~100% antigorite with an interlocking texture; (f) zoned chromite grain up to 2 mm across commonly rimmed/fractured by secondary magnetite. Mineral abbreviations: Ol = olivine, Opx = orthopyroxene, Spl = spinel, Brc = brucite, Mag = magnetite, Atg = antigorite, Tlc = talc, Mgs = magnesite, Srp = serpentine.

Mineral assemblages In Abu Dahr serpentinized peridotites, various high-Ni sulfide, and metal phases occur, including high-Ni pentlandite, cobaltian pentlandite, heazlewoodite, godlevskite, millerite, awaruite, and native Ni along with magnetite and goethite. The mineral abbreviations (after Whitney and Evans 2010) and stoichiometric formulas of the primary and secondary sulfides and metals mentioned in the text are given in Table 2. The Ni-sulfide and metal alloy minerals are disseminated in every polished thin section, and display low abundance (<<1% of the bulk rock; Kiseeva et al. 2017). Based on the dominant Ni-bearing sulfide mineral, sulfide assemblages are divided into three distinct facies:

(1) pentlandite-rich, (2) godlevskite-rich, and (3) millerite-rich. Magnetite is present in almost all assemblages as fine-grained disseminations in serpentinites. Typical assemblages and intergrowths are illustrated in Figures 4–6.

Pentlandite-rich assemblages. Pentlandite-awaruite-magnetite and cobaltian pentlandite-heazlewoodite-magnetite are the dominant assemblages (Figs. 4a–f) in partially serpentinized peridotites (lizarditization stage).

Godlevskite-rich assemblages. Cobaltian pentlandite-heazlewoodite-godlevskite-magnetite and heazlewoodite-godlevskitemagnetite are the main mineral assemblages (Figs. 5a–d) in fully serpentinized rocks (antigorite stage).

Millerite-rich assemblages. Cobaltian pentlandite-heazlewoodite-millerite and millerite-native Ni-goethite assemblages are the major Ni-sulfide/native Ni/Fe-oxyhydroxide minerals of the



FIGURE 4. Backscattered electron (**a**–**e**) and optical (**f**) images showing textural characteristics of the pentlandite-rich assemblage in Abu Dahr serpentinized peridotites. (**a**) Individual grains of cobaltian pentlandite and magnetite in serpentine. (**b**) Euhedral grain of heazlewoodite encapsulating magmatic pentlandite relics. (**c**) Zoned euhedral grain of Ni-rich awaruite (native Ni) core rimmed by stoichiometric awaruite. (**d** and **e**) Large desulfurized pentlandite grains partly replaced and rimmed by awaruite. (**f**) Disseminated blebs, up to 50 µm in diameter, comprise a granular aggregate of godlevskite which is replaced by a reticulate pattern of lamellar millerite (veinlets ~5 µm in width), C.P. Numbers indicate the EMP point analyzed performed in the Tables 3 and 4. Mineral abbreviations: see Table 2.



FIGURE 5. Backscattered electron images showing texturalmineralogical characteristics of the godlevskite-rich assemblage in Abu Dahr serpentinized peridotites. (a) Composite intergrowths of the godlevskite-rich assemblages in serpentine. Enlarged area is shown in **b**. (b) Pentlandite intergrown with heazlewoodite, godlevskite, and magnetite. (c) Heazlewoodite intergrown with godlevskite and mantled by magnetite. (d) Enlarged area of box in **c** showing a BSE image of heazlewoodite replaced by godlevskite. Mineral abbreviations: see Table 2.

carbonation stage (Figs. 6a and 6b) in steatized (talc-carbonated) serpentinites (carbonation stage).

Replacement textures of Ni-rich phases

Based on their textural relations, primary pentlandite occurs as finely disseminated discrete grains of Ni-rich pentlandite/ cobaltian pentlandite (less than 20 μ m) associated with the serpentine matrix (Fig. 4a), or in rare cases as euhedral grains of pentlandite replaced by heazlewoodite (Fig. 4b). Awaruite occurs as finely disseminated euhedral grains (< 10 μ m) of Ni-rich awaruite (native Ni) core rimmed by stoichiometric awaruite (Fig. 4c) and as interstitial disseminations (>2 mm) of desulfurized pentlandite (Figs. 4d–f). Heazlewoodite commonly replaces cobaltian pentlandite forming composite intergrowths of cobaltian pentlanditeheazlewoodite (Fig. 6a). In fully serpentinized rocks, godlevskite replaces heazlewoodite-forming composite intergrowths of cobaltian pentlandite-heazlewoodite-godlevskite (Fig. 5a), cobaltian pentlandite-heazlewoodite-godlevskite (Fig. 5b), and heazlewoodite-godlevskite (Fig. 5c). Magnetite commonly



FIGURE 6. Backscattered electron images showing texturalmineralogical characteristics of the millerite-rich assemblage in Abu Dahr serpentinized peridotites. (a) Two composite grains with characteristic replacement textural patterns as a result of serpentinization. One shows replacement of cobaltian pentlandite by heazlewoodite, and the other showing complex replacement patterns of cobaltian pentlandite by heazlewoodite and millerite in talc-carbonate rocks. (b) Millerite intergrown with native Ni. Millerite rimmed by native Ni and nickeliferous goethite. Euhedral tabular goethite crystals surrounding millerite and scattered in serpentine. Numbers indicate the EMP point analyzed performed in the Tables 3 and 4. Mineral abbreviations: see Table 2.

TABLE 2. Primary and secondary Ni-bearing phases in serpentinized peridotites of Abu Dahr with refering to mineral abbreviation used in this study

	Primary Ni-bearing p	hases	Secondary Ni-bearing phases					
Mineral	Abbreviation	Stoichiometric formula	Mineral	Abbreviation	Stoichiometric formula			
Pentlandite Cobaltian pentlandite	Pn CoPn	(Ni,Fe)₅S ₈ (Ni,Fe, Co)₅S ₈	Heazlewoodite Godlevskite	Hzl Gv	Ni_3S_2 Ni_9S_8			
			Millerite Chalcopyrite Awaruite Native Ni Magnetite Goethite	Mlr Ccp Awr native Ni Mag Gth	NiS CuFeS ₂ Ni ₃ Fe Ni Fe ₃ O ₄ (Fe Ni)OOH			



◄ FIGURE 7. (a) Compositional relationship between Fo content of olivine and Cr#s of coexisting Cr-spinels of mantle harzburgites of Abu Dahr; the olivine-spinel mantle array (OSMA) after Arai (1994). Fields for abyssal and forearc peridotites are from Dick and Bullen (1984), Parkinson and Pearce (1998), and Pearce et al. (2000). FMM-fertile MORB mantle. (b) Plot of Mg# vs. Cr# of Cr-spinels. Fields for abyssal peridotite is from Dick and Bullen (1984), the Mariana forearc field from Ishii et al. (1992). (c) Plot of ∆log (f_{02}) fayalite-magnetite-quartz (FMQ) vs. Cr# in Cr-spinel for the Abu Dahr harzburgites. Discrimination boundaries for MOR dunite and harzburgite, and SSZ dunite and harzburgite are from Dare et al. (2009).

mantles heazlewoodite and godlevskite (Fig. 5d). Heazlewoodite co-occurring with godlevskite may also contain small cobaltian pentlandite inclusions, which are lacking in godlevskite. This indicates that godlevskite did not directly grow at the expense of cobaltian pentlandite; it is more likely that godlevskite replaces heazlewoodite in fully serpentinized rocks. Pentlandite is also replaced by secondary magnetite forming the sulfide-magnetite composite grains (Fig. 5b).

With increasing the degree of steatitization (carbonation), low-sulfur Ni-sulfides are progressively replaced by sulfur-rich Ni-sulfides. Millerite grows at the expense of the low-sulfur heazlewoodite and godlevskite (Figs. 4f and 6a). Godlevskite is replaced by a reticulate pattern of lamellar millerite veinlets of ~5 μ m width (Fig. 4f). Millerite in these veinlets is fibrous in habit and appears to follow structural directions in the replaced godlevskite (Hudson and Travis 1981). The completely steatized rocks contain rounded and irregular grains of millerite (Fig. 6b). The Ni-rich awaruite (native Ni) rims the millerite grain boundaries (Fig. 6b). Euhedral tabular goethite crystals occur in association with millerite and forms rims on millerite (Fig. 6b). These textural patterns occur as a result of sulfide replacement processes (Rottier et al. 2016). Chalcopyrite (CuFeS₂) is not present and rarely occurs as small supergene grains intergrown with the millerite-rich assemblage (Fig. 6a).

Paragenesis

The paragenetic sequence of the Ni-bearing phases of Abu Dahr peridotites accompanying the serpentinization and carbonation processes is summarized in Figure 2. Textural relationships suggest this generalized paragenetic sequence: pentlandite \rightarrow cobaltian

TABLE 3. Representative EMF	³ analyses of pentland	te Pn and cobaltian pentlandite Co	Pn in Abu Dahr serpentinized peridotites
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			· · · ·			El	ement	wt%		Element at%			Mineral Formulas (apfu)				Ratios			
Sample	Figure	Point	Assemblage	Mineral	Fe	Co	Ni	S	Total	Fe	Со	Ni	S	Fe	Со	Ni	S	Ni/ Co	Ni/ Fe	Metal/ S
316-3				Pn*	26.06	2.14	36.69	35.12	100	20.99	1.63	28.11	49.27	3.72	0.29	4.99	8.00	17.25	1.34	1.03
316-5				Pn*	27.95	1.96	34.58	35.51	100	22.44	1.49	26.43	49.66	4.01	0.27	4.72	8.00	17.74	1.18	1.01
322-16				Pn*	27.80	1.59	36.63	33.98	100	22.54	1.22	28.26	47.98	3.90	0.21	4.89	8.00	23.16	1.25	1.08
306-7				Pn*	26.69	2.72	35.25	35.33	100	21.46	2.07	26.97	49.49	3.82	0.37	4.81	8.00	13.03	1.26	1.02
				Average	27.13	2.10	35.79	34.98	100	21.86	1.60	27.44	49.10	3.86	0.29	4.85	8.00	17.79	1.26	1.04
				St.dev.	0.91	0.47	1.04	0.69	0.00	0.76	0.36	0.89	0.76	0.12	0.07	0.12	0.00	4.16	0.07	0.03
306-7	4a	1		Co Pn*	25.57	5.17	36.85	32.41	100	20.96	4.02	28.74	46.28	3.51	0.67	4.82	8.00	7.15	1.37	1.16
306-7	4a	2		Co Pn*	26.23	5.28	35.51	32.98	100	21.59	4.12	27.81	46.48	3.63	0.69	4.68	8.00	6.75	1.29	1.15
306-7	4a	3		Co Pn*	24.91	5.00	35.68	34.41	100	20.16	3.83	27.48	48.53	3.52	0.67	4.81	8.00	7.17	1.36	1.06
309-5	6a	15	Pn-Hzl-Mag	Co Pn	25.54	5.10	34.36	35.00	100	20.59	3.90	26.36	49.16	3.64	0.69	4.67	8.00	6.76	1.28	1.03
309/5			Pn-Hzl-Mag	Co Pn	24.84	5.04	36.30	33.82	100	20.18	3.88	28.06	47.87	3.48	0.67	4.85	8.00	7.23	1.39	1.09
322-7A	5b	7	Pn-Hzl-Gv-Mag	g Co Pn	25.57	5.17	35.12	34.14	100	20.73	3.97	27.09	48.21	3.60	0.69	4.71	8.00	6.82	1.31	1.07
322-7A			Pn-Hzl-Gv-Mag	g Co Pn	23.15	6.22	36.72	33.91	100	18.81	4.79	28.39	48.00	3.25	0.83	4.92	8.00	5.93	1.51	1.08
309-5	ба	17	Pn-Hzl-Mlr	Co Pn	24.66	6.28	35.57	33.49	100	20.08	4.85	27.56	47.51	3.44	0.83	4.73	8.00	5.68	1.37	1.10
309-5	6a	18	Pn-Hzl-Mlr	Co Pn	22.39	7.53	36.17	33.91	100	18.20	5.80	27.98	48.02	3.15	1.00	4.85	8.00	4.82	1.54	1.08
				Average	24.76	5.64	35.81	33.79	100	20.14	4.35	27.72	47.78	3.47	0.75	4.78	8.00	6.48	1.38	1.09
				St.dev.	1.24	0.86	0.79	0.76	0.00	1.05	0.67	0.71	0.92	0.17	0.11	0.09	0.00	0.83	0.09	0.04

Notes: *discrete individual mineral



FIGURE 8. (a) Chemical composition (at%) of the analyzed Nisulfides and awaruite-native Ni of the studied Abu Dahr peridotites on the Fe-Ni-S phase diagram (after Misra and Fleet 1973). (b) Mineral stabilities in the ternary diagram Fe_9S_8 -Ni₉S₈-Co₉S₈ (after Kaneda et al. 1986) with coexisting of the Abu Dahr pentlandites occupy the stability field at 200 °C.

pentlandite \rightarrow awaruite/native Ni (and/or heazlewoodite), and heazlewoodite \rightarrow godlevskite \rightarrow millerite, and awaruite \rightarrow millerite and goethite. This sequence is consistent with the desulfidation during initial serpentinization; progressive serpentinization allows the formation of high-sulfur assemblages. On the basis of predominant minerals, mineral textures and associations, there were four main stages of formation and modification of Ni-bearing minerals: stage 1 involved replacement of orthomagmatic pentlandite/ cobaltian pentlandite by awaruite and/or heazlewoodite; stage 2 replacement of heazlewoodite by godlevskite; stage 3 replacement of godlevskite by millerite; and stage 4 Ni-rich awaruite broke down to millerite and goethite.

Although pyrrhotite occurs in many serpentinized peridotites described in the literature (e.g., Shiga 1987; Abrajano and Pasteris 1989; Lorand 1989; Miller 2007), it is absent in all the investigated samples from Abu Dahr. Even the pyrrhotite was present, and based on other case studies, the formation of magnetite decreases the Fe:Ni ratio of the sulfides, increasing the ratio of Ni-bearing sulfides to Fe-bearing sulfides (pyrrhotite), until pyrrhotite disappears. This also explains that the mineral pyrrhotite may be occurred, but disappears with progressive serpentinization and magnetite formation.

Mineral chemistry

Primary mantle assemblage Primary mineral compositions of Abu Dahr harzburgites are reported in Online Materials¹ Table S1. Olivine in the harzburgites has forsterite (Fo) values of 91–93 (average = 92). Their NiO and MnO contents are in the range of 0.36–0.57 wt% (corresponding to 2828–4478 ppm Ni) and 0.02–0.3 wt%, respectively. Spinel Cr# [= 100 Cr/(Cr+Al)] values range from 55 to 65 (average = 62) and Mg# [= 100 Mg/(Mg+Fe)] from 53 to 57. The compositional plots of spinel Cr# vs. olivine Fo and spinel Cr# vs. Mg# are illustrated on Figures 7a and 7b. Orthopyroxene is enstatite with high Mg#, which ranges from 91.7 to 92.3 (average = 92). Clinopyroxene (exsolution lamellae in the orthopyroxene) is diopside with Mg# ranging from 94.1 to 95.2. Primary amphiboles of the Abu Dahr harzburgite are magnesio-hornblende (Mg# 94–97) and tschermakite (Mg# 78–80) of the calcic-amphibole group.

Ni-rich sulfide and alloy phases

Pentlandite and cobaltian pentlandite. Pentlandite is the dominant sulfide mineral at Abu Dahr and shows stoichiometric compositions ranging from (Fe_{3.86}Ni_{4.85}Co_{0.29})₉S₈ to (Fe_{3.72}Ni_{4.99}Co_{0.29})₉S₈ (Table 3). All of the pentlandite grains analyzed (13 grains) contain between 26.06-27.95 (average = 27.13 ± 0.91) wt% Fe, 34.58-36.69 (average = 35.79 ± 1.04) wt% Ni, and 1.59-2.72 (average = 2.10 ± 0.47) wt% Co (Table 3, Fig. 8a). In this study, pentlandite has a high-Ni tenor (up to 5.00 apfu (atoms per formula unit) Ni), with Ni/Fe ratios ranging from 1.18 to 1.34 (average = 1.26 ± 0.07). The atomic metal/sulfur ratios displayed by the Abu Dahr pentlandites are 1.01-1.08 (average = 1.04 ± 0.03). Pentlandite contains up to 2.72 wt% Co, and the Ni/Co ratio is 13.0-23.2 (average = 1.79 ± 4.16). In the ternary system Fe₉S₈-Ni₉S₈-Co₉S₈ (Kaneda et al. 1986), the Abu Dahr pentlandites plot within the pentlandite stability field at 200 °C (Fig. 8b).

The average cobaltian pentlandite composition is $(Fe_{3.47} Ni_{4.78}Co_{0.75})_9S_8$ Co with average values of 24.76 ± 1.24 wt% Fe, 35.81 ± 0.79 wt% Ni, and 5.64 ± 0.86 wt% Co (Table 3, Fig. 8a). The Ni/Fe ratios are 1.28–1.54 (high-Ni). The atomic metal/sulfur ratios (1.03–1.16) are consistent with those of pentlandite. The cobaltian pentlandite is characterized by, remarkably, high Co contents of 5.00 to 7.53 (average = 5.64) wt% Co with Ni/Co ratios (4.82–7.23, average = 6.48 ± 0.83) typical of mantle sulfides worldwide (Wang et al. 2010). Binary plots (Figs. 9a–d) were generated to compare the Abu Dahr pentlandites with sulfides from forearc



FIGURE 9. Binary plots (wt%) between the metals in Abu Dahr pentlandites compared with pentlandites from different localities worldwide (see text). (**a**) Co vs. Ni, (**b**) Co vs. Fe, (**c**) Co vs. Fe+Ni, and (**d**) Fe vs. Co+Ni. Additional data are from Vourinos, Greece (Tzamos et al. 2016); Massif du Sud Peridotite Nappe, New Caledonia (Augé et al. 1999); Cemetery Ridge, Arizona (Haxel et al. 2018); Santa Elena Nappe, Costa Rica (Schwarzenbach et al. 2014); and Latao, SE Sulawesi (Rafianto et al. 2012).

							Elemei	nt wt%				Ele	ment at ^c	%	
Sample	Figure	Point	Assemblage	Mineral	Fe	Со	Ni	Cu	S	Total	Fe	Co	Ni	Cu	S
306-7	4b	4		Hzl*	0.40	0.00	70.36	0.00	29.24	100	0.34	0.00	56.64	0.00	43.09
306-7		Hzl*		0.39	0.00	71.19	0.00	28.42	100	0.33	0.00	57.62	0.00	42.10	0.02
322-7A	6a	16	Pn-Hzl-Mag	Hzl	0.41	0.00	71.77	0.00	27.58	100	0.35	0.00	58.38	0.00	41.41
322-7A	5a	19	Pn-Hzl-Mlr	Hzl	0.51	0.00	72.40	0.00	26.75	100	0.44	0.00	59.17	0.00	40.52
322-7A	5b	8	Pn-Hzl-Gv-Mag	Hzl	1.87	0.00	71.16	0.00	26.91	100	1.61	0.00	58.12	0.00	40.32
322-7A	5d	13	Hzl-Gv-Mag	Hzl	3.60	0.00	70.15	0.00	26.25	100	3.11	0.00	57.61	0.00	39.46
322-7A	5a	11	Pn-Hzl-Gv-Mag	Hzl	3.78	0.00	68.99	0.00	27.23	100	3.24	0.00	56.28	0.00	40.66
				Average	1.57	0.00	70.86	0.00	27.48	100	1.35	0.00	57.69	0.00	41.08
				St.dev.	1.54	0.00	1.13	0.00	1.04		1.33	0.00	1.00	0.00	1.22
322-7A	5b	9	Hzl-Gv-Mag	Gv	2.33	0.00	67.57	0.00	29.71	100	1.96	0.00	54.10	0.00	44.06
322-7A	5a	10	Pn-Hzl-Gv-Mag	Gv	2.44	0.00	66.32	0.00	29.92	99	2.04	0.00	52.77	0.00	45.31
322-7A	5b	12	Pn-Hzl-Gv-Mag	Gv	1.59	0.16	68.42	0.00	29.34	100	1.33	0.13	54.78	0.00	43.68
322-7A	5d	14	Pn-Hzl-Gv-Mag	Gv	1.31	0.20	67.60	0.00	30.89	100	1.10	0.16	53.82	0.00	45.03
				Average	1.92	0.09	67.48	0.00	29.97	99	1.61	0.07	53.87	0.00	44.52
				St.dev.	0.55	0.11	0.86	0.00	0.66		0.46	0.08	0.84	0.00	0.77
309-5	6b	23	Mlr-native Ni-Gth	Mlr	0.44	0.00	66.00	0.00	33.55	100	0.36	0.00	49.44	0.00	50.25
309-5				Mlr	0.47	0.00	64.08	0.00	35.46	100	0.38	0.00	48.37	0.00	51.20
309-5				Mlr	0.78	0.00	64.21	0.00	35.01	100	0.64	0.00	49.73	0.00	49.63
309-5				Mlr	0.83	0.12	64.16	0.00	34.89	100	0.67	0.00	48.59	0.00	50.54
322-7A	ба	20	Pn-Hzl-Mlr	Mlr	0.53	0.32	64.46	1.79	33.02	100	0.44	0.26	49.83	1.31	48.08
322-7A	бa	21	Pn-Hzl-Mlr	Mlr	0.85	0.35	65.13	0.57	33.10	100	0.70	0.27	51.10	0.41	47.54
				Average	0.65	0.13	64.67	0.39	34.17	100	0.53	0.09	49.51	0.29	49.54
				St.dev.	0.19	0.16	0.75	0.72	1.07		0.15	0.14	0.98	0.53	1.44
322-7A	бa	22	Pn-Hzl-Mlr	Сср	29.60	0.00	1.49	33.67	35.25	100	24.28	0.00	1.16	24.27	50.37
				Average	29.60	0.00	1.49	33.67	35.25	100	24.28	0.00	1.16	24.27	50.37
306-7	4c	5	Awr-native Ni	Awr (rim)	21.76	1.15	76.41	0.11	0.00	99	22.61	1.14	76.10	0.12	0.00
306-7	4c	6	Awr-native Ni	Awr (core)	13.45	0.96	85.03	0.33	0.00	100	14.06	0.95	84.76	0.30	0.00
				Average	17.61	1.06	80.72	0.22	0.00	100	18.34	1.04	80.43	0.21	0.00
309-5	6b	24	Mlr-native Ni-Gth	native Ni	4.96	0.39	92.46	2.13	0.00	100	5.19	0.41	91.16	2.00	0.00
309-5			Mlr-native Ni-Gth	native Ni	6.23	0.46	91.88	1.44	0.00	100	6.55	0.47	91.58	1.35	0.00
309-5			Mlr-native Ni-Gth	native Ni	4.84	0.31	92.75	2.09	0.00	100	4.95	0.30	92.57	1.88	0.00
309-5				native Ni	6.28	0.57	91.70	1.45	0.00	100	6.87	0.59	90.64	1.40	0.00
				Average	5.58	0.43	92.20	1.78	0.00	100	5.89	0.44	91.49	1.66	0.00
				St.dev.	0.78	0.11	0.49	0.38	0.00		0.96	0.12	0.82	0.33	0.00
Notes: *di	screte indiv	vidual m	ineral												

 TABLE 4.
 Representative EMP analyses of heazlewoodite Hzl, godlevskite Gv, millerite Mlr, chalcopyrite Ccp, awaruite Awr, and native Ni in Abu

 Dahr serpentinized-carbonated peridotites

(Table continues on next page)

mantle wedge settings worldwide (e.g., Vourinos, Greece: Tzamos et al. 2016; Massif du Sud Peridotite Nappe, New Caledonia: Augé et al. 1999; Cemetery Ridge, Arizona: Haxel et al. 2018; Santa Elena Nappe, Costa Rica: Schwarzenbach et al. 2014; Latao, SE Sulawesi: Rafianto et al. 2012). The Abu Dahr pentlandites display a positive correlation between Co and Ni (Fig. 9a) and negative correlation between Fe and Fe+Ni (Figs. 9b and 9c). Similarly, Fe shows a strong negative correlation with Co+Ni (Fig. 9d), similar to typical mantle wedge occurrences worldwide.

Heazlewoodite. The average composition of analyzed heazlewoodite is $(Fe_{0.07}Ni_{2.93})_3S_2$ with 1.57 ± 1.54 wt% Fe, 70.86 \pm 1.13 wt% Ni (2.93 apfu Ni) and no detectable Co (Table 4, Fig. 8a). The atomic metal/sulfur ratio of heazlewoodite ranges from 1.32 to 1.54 (average = 1.44 ± 0.07), with lower values where heazlewoodite replaces cobaltian pentlandite (1.32), and slightly elevated values when heazlewoodite is associated with godlevskiterich assemblages. Heazlewoodite contains variable amounts of Fe (0.39–3.78 wt%); heazlewoodite associated with relict cobaltian pentlandite is Fe-poor (0.41–1.87 wt%), and Fe-rich where it is associated with or mantled by magnetite (3.60–3.78 wt%).

Godlevskite. The average composition of godlevskite is $(Fe_{0.26}Ni_{8.73}Co_{0.01})_9S_8$ with average concentrations of 1.92 ± 0.55 wt% Fe and 67.48 ± 0.86 wt% Ni) (8.73 apfu Ni) (Table 4, Fig. 8a). The atomic metal/sulfur ratio of godlevskite ranges from 1.21 to 1.29 (average = 1.25). The average ratio is higher than the theoretical value of 1.125. Metal/sulfur ratios are slightly higher where godlevskite replaces heazlewoodite (1.27–1.29), and lower

where godlevskite coexists with cobaltian pentlandite (1.21-1.22). The Fe and Co contents of godlevskite (2.33, 0.20 wt%, respectively) are elevated when godlevskite is associated with relict cobaltian pentlandite.

Millerite. The average composition of millerite is (Fe_{0.01} Ni_{0.98}Cu_{0.01})S. Average element concentrations are 0.65 ± 0.19 wt% Fe, 64.67 \pm 0.75 wt% Ni (0.98 apfu Ni), 0.13 \pm 0.16 wt% Co, and 0.39 \pm 0.72 wt% Cu (Table 4, Fig. 8a). The atomic metal/sulfur ratio of millerite ranges from 0.95 and 1.10 (average = 1.02 ± 0.06), with slightly elevated values of 1.10 (average = 1.09) where millerite replaces heazlewoodite. Millerite associated with native Ni shows elevated Fe contents (up to 0.83 wt% Fe, with no Co or Cu). The Fe content also increases when millerite is associated with relict cobaltian pentlandite-heazlewoodite-chalcopyrite (0.85 wt%), and the Co and Cu contents of this millerite are up to 0.35 wt% and 1.79 wt%, respectively.

Chalcopyrite. The average composition of chalcopyrite is (Fe_{0.98}Cu_{0.98}Ni_{0.05})₂S₂. The element concentrations are 29.60 wt% Fe, 33.67 wt% Cu, and 1.49 wt% Ni, with no detectable Co (Table 4, Fig. 8a).

Awaruite and native Ni. Awaruite exhibits a wide compositional range from $Ni_{75}Fe_{21}$ (stoichiometric Ni_3Fe) to $Ni_{93}Fe_5$ (native Ni) (Table 4, Fig. 8a); only a few analyses are classified as awaruite sensu stricto (Ni_3Fe). Euhedral awaruite shows Ni zonation, with a stoichiometric awaruite rim of 21.76 wt% Fe and 76.41 wt% Ni and Ni-rich core containing 13.47 wt% Fe and 85.03 wt% Ni. Co is present in minor amounts of less than 1.15 wt%. Native nickel

TABLE 4.—EXTENDED

		Miner	al Formulas	s (apfu)		
Sample	Fe	Co	Ni	Cu	S	Metal/S
306-7	0.02	0.00	2.98	0.00	2.00	1.32
306-7	0.00	2.98	0.00	2.00	1.38	
322-7A	0.02	0.00	2.98	0.00	2.00	1.42
322-7A	0.02	0.00	2.98	0.00	2.00	1.47
322-7A	0.08	0.00	2.92	0.00	2.00	1.48
322-7A	0.15	0.00	2.85	0.00	2.00	1.54
322-7A	0.16	0.00	2.84	0.00	2.00	1.46
	0.07	0.00	2.93	0.00	2.00	1.44
	0.06	0.00	0.06	0.00	0.00	0.07
322-7A	0.31	0.00	8.69	0.00	8.00	1.27
322-7A	0.33	0.00	8.67	0.00	8.00	1.21
322-7A	0.21	0.02	8.77	0.00	8.00	1.29
322-7A	0.18	0.03	8.79	0.00	8.00	1.22
	0.26	0.01	8.73	0.00	8.00	1.25
	0.07	0.02	0.06	0.00	0.00	0.04
309-5	0.01	0.00	0.99	0.00	1.00	0.99
309-5	0.01	0.00	0.99	0.00	1.00	0.95
309-5	0.01	0.00	0.99	0.00	1.00	1.01
309-5	0.01	0.00	0.98	0.00	1.00	0.97
322-7A	0.01	0.00	0.96	0.03	1.00	1.08
322-7A	0.01	0.01	0.97	0.01	1.00	1.10
	0.01	0.00	0.98	0.01	1.00	1.02
	0.00	0.00	0.01	0.01	0.00	0.06
322-7A	0.98	0.00	0.05	0.98	2.00	0.99
	0.98	0.00	0.05	0.98	2.00	0.99
306-7	22.61	1.14	76.10	0.10	0.00	0.00
306-7	14.06	0.95	84.76	0.31	0.00	0.00
	18.34	1.04	80.43	0.21	0.00	0.00
309-5						0.00
309-5						0.00
309-5						0.00
309-5						0.00
						0.00
						0.00

grains have Ni contents of 91.7 to 93.0 wt% (average = 92.20 ± 0.49), with variable Fe contents (4.96–6.28 wt%, average = 5.58 ± 0.78) and minor Co and Cu contents (0.43 ± 0.11 and 1.78 ± 0.38 wt%, respectively).

Fe-oxides/oxyhydroxides

Nickeliferous magnetite and goethite. The Cr-spinel alteration phases (ferritchromite and magnetite) and goethite from Abu Dahr are enriched in Ni (Table 5); however, primary chromites are Ni-free (Online Materials¹ Table S1). Ferritchromite has relatively high Ni contents (0.24-1.19, average = 0.63 wt% NiO). Magnetite rims to ferritchromite have the highest Ni contents (0.40-1.24, average = 0.75 wt% NiO, Table 5), than the serpentinization-related magnetite from the Dumont sill, which contains average 0.07 wt% NiO (Sciortino et al. 2015). Magnetite intergrown with pentlanditeheazlewoodite-godlevskite (Fig. 5b) has not yet been analyzed. The nickeliferous goethite (Fe, Ni)O(OH) has Ni concentrations of 1.91 to 2.12 (average = 2.02) wt% NiO (Table 5).

Equilibrium P-T and oxygen fugacity calculations

The results of calculated *P*-*T* and the f_{O_2} values [expressed as the deviation from FMQ in log units, $\Delta \log(f_{O_2})$ FMQ] for Abu Dahr harzburgites are given in Table 6.

The equilibrium temperatures estimated using the Brey and Köhler (1990) Ca-in-orthopyroxene and two-pyroxene thermometers, suggest that the Abu Dahr harzburgites equilibrated at temperatures of 910–1217 °C. Equilibration pressures using twopyroxene geobarometer of Putirka (2008) gave pressure values between 4.1 and 7.8 kbar (0.4–0.8 GPa). The calculated f_{02} values for Abu Dahr harzburgites range from FMQ+0.41 to FMQ+1.20 and correlated positively with melting (spinel Cr#) (Fig. 7c).

Whole-rock geochemistry

The results of whole rock major and trace element analyses of five samples from Abu Dahr are reported in Table 7. Bulk-rock geochemistry is strongly affected by hydrothermal alteration, consistent with findings from subduction-related and abyssal serpentinites worldwide (e.g., Deschamps et al. 2010; Marchesi et al. 2013; Uno and Kirby 2019), and the high loss on ignition (LOI) values 9.22-12.71 wt% indicate the presence of H₂O and CO₂ (serpentine, brucite, magnesite). Major elements are normalized to 100% anhydrous compositions, with Mg# values and total FeO of most of the harzburgites falling within a restricted range (89-91 and 8.71-10.54 wt%, respectively). The MgO and SiO₂ contents are variable (41.24-46.83 and 42.12-46.92 wt%, respectively), consistent with increased depletion of the harzburgites. These samples have low concentrations of Al₂O₃ (0.41-0.90 wt%) and TiO₂ (0.01-0.06 wt%) indicating a highly depleted nature, which is consistent with a highly refractory mantle residue after high-degree partial melting (Pearce et al. 2000). The CaO contents are generally low (0.20-0.34 wt%) reflecting the scarcity of clinopyroxene in the peridotites, but two samples (322/3 and 324/4) have relatively high 1.83 and 1.49 wt% CaO. The MgO values can be used effectively as an indicator of the degree of depletion. Any increase in the MgO content is consistent with increased depletion and higher olivine contents (Wu et al. 2018). On Harker diagrams (Figs. 10a-d), the Abu Dahr harzburgite samples show relatively linear decreases in SiO2, Al2O3, and CaO contents with increasing MgO, but total FeO increases slightly with increasing MgO. These trends indicate the depleted nature of Abu Dahr peridotites and are similar to those of ultramafic rocks from similar settings (e.g., Mariana forearc: Parkinson and Pearce 1998; Cemetery Ridge, Arizona: Haxel et al. 2018; Santa Elena Nappe, Costa Rica: Schwarzenbach et al. 2014). All samples are significantly depleted relative to the primitive mantle (Figs. 10a-d).

Trace element analyses show that the serpentinized harzburgites of Abu Dahr are enriched in Cr (2461–7849 ppm), Ni (1489–2590 ppm), and Co (119–144 ppm) relative to fresh harzburgites of northern Abu Dahr. In contrast, the analyzed samples are notably depleted in Cu (0.1–10.9 ppm), below the average Cu concentration in spinel harzburgites 35 ± 11 ppm (Ciazela et al. 2015). The Abu Dahr harzburgites exhibit very low concentrations of the incompatible trace elements (Zn, Sc, V and Y), LILE (K, Rb, Ce, Sr, Ba), and HFSE (Zr, Y, Nb, Hf, Ta, U); reflecting the highly residual nature of the peridotites and their low clinopyroxene content (<1.0 vol%). The rare-earth element (REE) contents are also very low.

DISCUSSION

Based on the textural evolution of Ni-bearing phases and their mineralogical and chemical composition data, the important aspects to be discussed here are the relationship between redox state (changes in f_{0_2} and f_{s_2}) and Fe-Ni-Co-O-S minerals. In the following, we will first discuss the petrogenesis, oxidation state, and serpentinization of Abu Dahr harzburgites; and then discuss: (1) remobilization of Ni from magmatic olivines and upgrading of magmatic sulfide tenor; (2) relationship between redox state and Fe-Ni-Co-O-S minerals; (3) high Co contents of pentlandite in Abu Dahr; (4) similarities to Ni-sulfides and alloys from forearc mantle elsewhere; and finally (5) genetic model for Abu Dahr Ni-sulfides and metal alloys.

Petrogenesis of the Abu Dahr protolith

Peridotites of Abu Dahr are mainly highly depleted clinopyroxene-poor harzburgites with forsterite-rich olivine (Fo 91-93) and high Cr# spinel (55–65, average = 62) and are similar to harzburgites from other areas of the ophiolites in the Eastern Desert (e.g., Azer and Stern 2007; Khedr and Arai 2013, 2016). Spinel cores chemistry is extensively used as a petrogenetic (Dick and Bullen 1984; Arai 1992; 1994; Hellebrand et al. 2001) and tectonic setting (Arai 1992; Dare et al. 2009) indicator in ultramafic and mafic rocks. Based on olivine and spinel chemistry, the compositional plots of spinel Cr# vs. coexisting olivine Mg# and spinel Mg# vs. Cr# have shown that the Abu Dahr harzburgites are residues of high degrees of partial melting (~35%), plotted within the field of forearc peridotites and on modern Mariana forearc peridotite field (Figs. 7a and 7b). On the tectonic discrimination diagram of f_{Ω_2} vs. spinel Cr# (Dare et al. 2009), the Abu Dahr harzburgites plotted within the field of SSZ harzburgites (Fig. 7c).

Oxidation state of the Abu Dahr forearc

The spinel-olivine-orthopyroxene oxybarometry of Ballhaus et al. (1991) was used to estimate the oxidation state of Abu Dahr forearc peridotites, assuming a 1 GPa equilibrium pressure. The calculated f_{02} values for Abu Dahr forearc peridotites range from FMQ+0.41 to FMQ+1.20 (average = +0.60 FMQ), fall within the range from FMQ+0.3 to FMQ+2 of forearc mantle wedge (Parkinson and Arculus 1999), and are within the ± 2 FMQ of the upper mantle (Frost and McCammon 2008). The calculated f_{02} is positively correlated with spinel Cr# (Fig. 7c), indicating that higher melting degrees may influence the oxidation state of the mantle wedge, as demonstrated in previous studies (Parkinson and Arculus 1999; Kelley and Cottrell 2009; Kilgore et al. 2018).

The oxidation state of the Abu Dahr forearc harzburgites (+0.41 to +1.20 FMQ) is in agreement with the previous studies, which concluded that the mantle wedge is generally more oxidized than oceanic mantle (e.g., Ballhaus 1993; Parkinson and Arculus 1999; Arai and Ishimaru 2007; Malaspina et al. 2009; Evans et al. 2012 (FMQ 1–4); Bénard et al. 2018 (FMQ+0.5 to +1.5); Kilgore et al. 2018; Wang et al. 2020), and subcontinental lithospheric mantle (Perinelli et al. 2012, FMQ-1.52 to -0.23).

Serpentinization of Abu Dahr peridotites

The Abu Dahr peridotites have experienced multiple stages of serpentinization (lizarditization and antigoritization) and carbonation/listvenitization. Lizardite-brucite-magnetite from olivine and bastites from pyroxene and amphibole characterize the lizarditization stage (T < 300 °C; P < 4 kbar; depth <13 km; lower aCO_2 and f_{O_2} and f_{S_2} (Pirajno 2009; Schwartz et al. 2013; Debret et al. 2019; Boskabadi et al. 2020). Alteration of primary amphiboles is a Ca-releasing reaction and contributes to the Ca budget of the bulk peridotites. Alteration of olivine (R3 and R4) can be described in the following reactions (after Wang et al. 2009):

$$\begin{aligned} & 6(Mg, Fe)_{2}SiO_{4} + 7H_{2}O = 3(Mg, Fe)_{3}Si_{2}O_{3}(OH)_{4} + Fe_{3}O_{4} + \\ & H_{2}(aq). \end{aligned} \tag{3}$$

$$\begin{aligned} & 6(Mg, Fe)_2 SiO_4 + 9H_2 O = 3(Mg, Fe)_3 Si_2 O_5 (OH)_4 + \\ & 3Mg_{0.8} Fe_{0.2} (OH)_2. \end{aligned} \tag{4}$$

The alteration of amphibole releases aqueous Ca and SiO_2 (R5, after Wang et al. 2009):

$$\begin{aligned} Ca_2Mg_{4.8}Fe_{0.2}Si_8O_{22}(OH)_2 + 16/13H_2O + 38/13H^+ + 7/130Fe^{2+} \\ = 24/13Mg_{2.6}Fe_{0.4}Si_2O_5(OH)_4 + 2Ca^{2+}(aq) + 56/13SiO_2(aq). \end{aligned}$$

Magnetite formation is accompanied by the release of iron from early formed phases such as Fe-serpentine R3 or Fe-brucite

 TABLE 5. Representative EMP analyses and Ni contents (ppm) of magnetite Mag, ferritchromite Fe-chromite and goethite Gth from Abu Dahr

 Nappe

		SiO ₂	TiO ₂	Al ₂ O ₃	Cr ₂ O ₃	FeO _{Total}	MgO	MnO	CaO	NiO	Total	Ni
Sample	Mineral	wt%	wt%	wt%	wt%	wt%	wt%	wt%	wt%	wt%	wt%	ppm
318-9	Mag	0.49	0.05	0.06	7.75	83.9	0.93	0.19	0.11	0.40	93.90	3142
318-9	Mag	0.38	0.03	0.00	5.72	83.9	0.88	0.00	0.00	0.45	91.39	3535
319-8	Mag	1.03	0.01	0.00	1.45	82.4	0.90	0.10	0.02	1.24	87.16	9741
319-8	Mag	0.78	0.00	0.34	3.52	83.9	1.19	0.03	0.03	0.67	90.46	5263
319-8	Mag	0.53	0.00	0.09	4.52	84.5	0.87	0.11	0.07	1.05	91.73	8248
319-8	Mag	0.80	0.00	0.15	5.92	81.2	1.53	0.00	0.05	0.77	90.46	6049
331-19	Mag	1.03	0.28	0.00	0.17	88.8	0.44	0.11	0.02	0.70	91.59	5499
331-19	Mag	1.21	0.00	0.01	0.00	86.3	1.79	0.00	0.08	0.60	89.96	4713
331-19	Mag	0.84	0.03	0.00	4.65	85.2	0.91	0.00	0.00	0.88	92.53	6913
	Average	0.79	0.04	0.07	3.74	84.47	1.05	0.06	0.04	0.75	91.02	5900
318-9	Fe-chromite	0.82	0.02	2.72	31.79	57.3	3.03	0.42	0.01	0.78	96.88	6127
318-9	Fe-chromite	0.87	0.06	0.69	28.76	61.8	1.62	0.66	0.03	0.91	95.40	7148
318-9	Fe-chromite	0.85	0.13	1.06	31.00	58.9	2.11	0.54	0.02	0.41	95.03	3221
319-8	Fe-chromite	0.86	0.08	0.16	12.71	76.0	0.99	0.25	0.07	1.19	92.34	9348
330-9	Fe-chromite	0.90	0.01	0.23	19.81	68.9	0.84	0.39	0.07	0.94	92.06	7384
330-9	Fe-chromite	0.91	0.00	0.84	26.09	60.6	1.53	0.82	0.05	0.44	91.28	3456
330-9	Fe-chromite	0.89	0.00	0.81	27.99	56.9	1.40	0.82	0.09	0.24	89.12	1885
330-9	Fe-chromite	0.86	0.00	0.65	30.50	56.9	1.97	0.84	0.09	0.33	92.11	2592
330-9	Fe-chromite	0.86	0.02	0.12	17.68	71.8	1.34	0.15	0.00	0.71	92.64	5577
331-19	Fe-chromite	0.87	0.13	3.21	32.46	53.9	2.89	0.47	0.01	0.38	94.33	2985
	Average	0.87	0.05	1.05	25.88	62.29	1.77	0.54	0.04	0.63	93.12	4973
309-5	Gth					59.3				1.91	61.22	15004
309-5	Gth					62.2				2.12	64.33	16654
	Average					60.76				2.02	62.78	15829
Notes: npm	$Ni = wt\% NiO \times 100$	00/1.273										

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TABLE 6. Calculated equilibrium *P*–*T* and oxygen fugacities for Abu Dahr harzburgites

Lithology	Harzburgite													
Sample	316-3-1	316-3-2	316-3-3	316-3-4	316-3-5	316-5-1	316-5-2	318-9-1	318-9-2	318-9-3				
Spinel														
Mg#	55.7	56.7	54.9	55.1	52.7	55.1	55.4	56.2	57.4	56.0				
Cr#	60.0	61.1	64.1	63.2	56.4	54.7	63.8	63.5	64.6	62.1				
Fe ³⁺ #	5.1	5.0	4.9	4.1	5.9	3.9	4.5	4.5	2.3	4.4				
XFe ²⁺	0.443	0.433	0.451	0.449	0.473	0.449	0.446	0.438	0.426	0.440				
XFe ³⁺	0.051	0.050	0.049	0.041	0.059	0.039	0.045	0.045	0.048	0.045				
XAI	0.380	0.369	0.342	0.352	0.410	0.435	0.345	0.348	0.336	0.362				
Olivine														
Fo	91.6	91.3	91.5	91.8	91.7	91.7	92.2	91.0	91.8	92.5				
XFe	0.084	0.087	0.084	0.081	0.081	0.081	0.077	0.089	0.080	0.074				
Orthopyroxene														
Mg#	91.9	91.8	92.0	91.7	92.3	92.2	92.2	92.0	91.7	92.3				
XFe	0.081	0.082	0.080	0.083	0.077	0.078	0.081	0.081	0.077	0.080				
Clinopyroxene														
Mg#	94.1	94.7	95.1	94.2	94.2	94.9	95.2	94.6	94.2	95.0				
XFe	0.059	0.053	0.049	0.058	0.058	0.051	0.048	0.054	0.058	0.050				
T1 ℃	1766	1764	1763	1758	1766	1776	1775	1762	1768	1774				
T2 °C	1775	1774	1769	1768	1768	1782	1789	1770	1770	1784				
T3 ℃	1207	1204	1140	1162	1196	1219	1166	1184	1182	1191				
ТК	1480	1477	1413	1435	1469	1492	1439	1457	1455	1464				
T4 ℃	922	969	880	880	880	969	946	880	927	919				
P (KPar)	4.7	4.1	4.8	5.3	7.8	6.9	4.1	5.6	6.7	5.8				
$\Delta log(f_{O_2})^{FMQ}$	0.71	0.55	0.60	0.41	1.14	0.41	0.63	0.32	0.65	0.77				

Notes: Spl = spinel; Ol Fo = olivine forsterite; Opx = orthopyroxene; Cpx = clinopyroxene; $XFe^{2+} = Fe^{2+}/(Fe^2+Mg)$; $XFe^{3+} = Fe^{3/}(Fe^2+Fe^3)$; $XAI = AI(AI+Cr+Fe^3)$; $Fe^{3+} = Fe^{3+}/(Fe^{3+}+Cr+AI)$. T1, T2, and T3 are the equilibrium temperatures calculated using two-pyroxene geothermometers of Brey and Köhler (1990). T4 is the equilibrium temperatures calculated using two-pyroxene geothermometers of Brey and Köhler (1990). T4 is the equilibrium temperatures calculated using two-pyroxene geothermometers of Brey and Köhler (1990). T4 is the equilibrium temperatures calculated using two-pyroxene geothermometers of Brey and Köhler (1990). T4 is the equilibrium temperatures calculated using the oxybarometry of Ballhaus et al. (1991). Pressure of 1 GPa is assumed throughout. FMQ = fayalite-magnetite-quartz.

R6 (Bach et al. 2006; Frost and Beard 2007; Wang et al. 2009; Frost et al. 2013; Boskabadi et al. 2020):

$$9Fe(OH)_2 + 4SiO_2(aq) = 2Fe_3Si_2O_5(OH)_4 + Fe_3O_4 + 4H_2O + H_2(aq).$$
(6)

An antigoritization stage is a high-temperature event (T > 320-410 °C; P > 4 kbar; depth >13 km; higher aCO₂, aSiO₂, and f_{O_2} and f_{S_2} (Pirajno 2009; Schwartz et al. 2013; Debret et al. 2019; Boskabadi et al. 2020) and is characterized by antigorite after lizardite in fully serpentinized samples. The onset of the carbonation stage occurred when infiltration of CO₂-rich fluids destabilized lizardite/chrysotile to form antigorite and Fe-magnesite (R7). More infiltration of CO₂-rich fluids destabilizes antigorite to form magnesite (R8) and talc (R9). Formation of antigorite-talc-magnesite rock is based on the following reactions (R7–R9, after Boskabadi et al. 2020):

$$17(Mg,Fe)_{3}Si_{2}O_{5}(OH)_{4} + 3CO_{3} = (Mg,Fe)_{48}Si_{34}O_{85}(OH)_{62} + 3Mg(Fe)CO_{3} + 3H_{2}O$$
(7)

$$Mg_{48}Si_{34}O_{85}(OH)_{62} + 48CO_2(aq) = 48MgCO_3 + 34SiO_{2(aq)} + 31H_2O$$
(8)

$$\begin{split} Mg_{48}Si_{34}O_{85}(OH)_{62} + 30SiO_2(aq) &= 16Mg_3Si_4O_{10}(OH)_2 + \\ 15H_2O. \end{split} \tag{9}$$

The dehydration of Atg-serpentinites. Metamorphic olivine records external fluid infiltration during serpentinite dehydration (Clarke et al. 2020). Metamorphic olivine is formed during breakdown of brucite or antigorite (high-pressure serpentine) at ~400 and ~650 °C, respectively (Scambelluri et al. 2004). Partial dehydration (formation of prograde olivine at ~600 °C under high pressure >1 GPa) of Atg-serpentinites of Abu Dahr South occurs within subduction channel. Antigorite, stable to 620 °C at 1 GPa

(Ulmer and Trommsdorff 1995), breaks down into metamorphic (secondary) olivine through antigorite dehydration during prograde metamorphism under high *P-T* conditions in subduction zones (Padron-Navarta et al. 2011; Debret et al. 2013; Deschamps et al. 2013). Antigorite commonly coexists with secondary olivine and clinopyroxene, suggesting high-temperature serpentinization at \sim 450–550 °C (Murata et al. 2009). At Abu Dahr, the occurrence of secondary olivine and clinopyroxene (Fo 87 and Mg# 98, respectively, Online Materials' Table S1 and talc in serpentinites suggests that the serpentinization and eclogitization took place under similar conditions (600 °C, 20 kbar; Guillot et al. 2000). In areas with high hydrothermal alteration, serpentines have been transformed to fully carbonated peridotites (listvenites).

Remobilization of Ni from magmatic olivines and upgrading of magmatic sulfide tenor

Olivine is the main Ni-bearing mineral in the harzburgite and dunite protoliths and the major source of Ni forming the disseminated Ni mineralization (Marques et al. 2007; Kamenetsky et al. 2016). Partial melting of mantle peridotites increases the Ni contents of residual olivines (Herzberg et al. 2016). The high-Ni contents are consistent with high-Fo olivines (Sobolev et al. 2005, 2007; Nikkola et al. 2019). At Abu Dahr, harzburgite olivines are highly magnesian (Fo 91–93, average = 92) with higher NiO concentrations (0.33–0.57, average = 0.41 wt% NiO) than those reported from mantle wedge harzburgites worldwide (e.g., Mariana: 0.30–0.44 wt% NiO, Wang et al. 2009; Himalaya: 0.38–0.39 wt%, Hattori and Guillot 2007; Tonga: 0.22–0.41 wt%, Birner et al. 2017; Aladag: 0.31–0.51 wt%, Lian et al. 2018; Oman: 0.38–0.39 wt%, Negishi et al. 2013; New Caledonia: 0.30–0.41 wt%, Secchiari et al. 2020).

At Abu Dahr, pentlandite is the dominant primary mantle sulfide mineral and has Ni tenor ranging up to 37 wt% (apfu Ni = 5). Serpentinization-related high-Ni tenor disseminated sulfide deposits hosted within olivine-rich mantle peridotites are consistent with olivine compositions of Fo91–92 and attributed to the release of Ni from magmatic olivine and enrichment of the tenor of pentlandite by Fe/Ni exchange between olivine and magmatic sulfides (R10, after Barnes et al. 2013):

$$FeS + NiO = FeO + NiS.$$
(10)

Remobilization of Ni from magmatic olivines by postmagmatic fluids during serpentinization of Abu Dahr peridotites resulted in upgrading the Ni tenors of pre-existing primary pentlandite, explaining why Abu Dahr pentlandites have high Ni tenors and therefore, referred to here as high-Ni pentlandites (Table 3).

Relationship between redox conditions and Fe-Ni-Co-O-S minerals

In Abu Dahr, the redox conditions (changes in f_{0_2} and f_{s_2}) recorded by Ni-rich phases are varying from highly reducing conditions during partial serpentinization stage to strongly oxidizing conditions during late stages of serpentinization and carbonation. Evidence thereof is that Abu Dahr exhibits three main Ni-rich assemblages: (1) pentlandite-rich assemblages (pentlanditeawaruite-magnetite and pentlandite-heazlewoodite-magnetite) in Lz-serpentinites; (2) godlevskite-rich assemblages (pentlandite-

TABLE 7. Bulk-rock major oxides (wt%) normalized on a volatilefree basis and trace elements (ppm) contents of Abu Dahr serpentinized harzburgites

Sample	379/5	381/3	322/3	324/4	329/11
		Norma	lized wt%		
SiO ₂	42.12	43.91	45.02	44.33	46.92
TiO ₂	0.01	0.01	0.01	0.04	0.06
AI_2O_3	0.65	0.41	0.53	0.90	0.73
FeO _{Total}	10.02	8.74	8.71	8.85	10.54
FeO	9.02	7.86	7.84	7.96	9.48
MnO	0.14	0.13	0.15	0.13	0.15
MgO	46.83	46.52	43.74	44.24	41.24
CaO	0.20	0.27	1.83	1.49	0.34
K ₂ O	0.00	0.00	0.00	0.00	0.01
Na₂O	0.00	0.00	0.00	0.00	0.00
P_2O_5	0.02	0.01	0.01	0.01	0.01
Total	100.00	100.00	100.00	100.00	100.00
LOI	10.7	9.66	11.83	9.22	12.71
Mg#	90	91	91	91	89
-		r	opm		
Sc	0.00	13.70	0.00	0.00	0.00
V	29.00	23.40	23.00	31.70	32.90
Cr	7849.00	2461.30	2607.90	2468.80	3649.10
Co	143.90	127.80	119.10	121.70	122.70
Ni	2589.50	2378.60	2198.40	2246.60	1489.10
Cu	7.20	0.10	4.20	5.20	10.90
Zn	64.10	54.50	54.00	52.10	60.70
Ga	0.00	0.00	0.00	0.00	0.00
Rb	0.00	2.40	3.80	5.00	5.50
Sr	2.26	0.00	14.57	7.62	15.72
Y	3.80	6.10	6.00	6.30	4.90
Zr	0.00	28.90	0.00	29.40	35.60
Nb	0.00	0.00	0.00	0.00	0.00
Cs	18.00	0.00	0.00	0.00	46.60
Ba	0.00	0.00	0.00	0.00	0.00
La	0.00	0.00	0.00	0.00	0.00
Ce	0.00	0.00	0.00	0.00	0.00
Pr	0.00	0.00	0.00	5.70	0.00
Hf	0.00	0.00	0.00	0.00	0.00
Та	0.00	0.00	0.00	0.00	0.00
Pb	0.00	0.00	0.00	0.00	0.00
Th	15.10	14.90	10.80	12.50	13.80
U	0.00	0.00	0.00	0.00	0.00
Notes: Mg	# = (100×Mq0)/40.32)/(MgO	/40.32+FeO/71	.85).	

heazlewoodite-godlevskite-magnetite and heazlewooditegodlevskite-magnetite) in Atg-serpentinites; and (3) millerite-rich assemblages (pentlandite-heazlewoodite-godlevskite-millerite and millerite-native Ni-goethite) in carbonated serpentinites. Similar redox conditions during serpentinization and talc-carbonation are recorded in the Fe-Ni-Co-O-S system (Eckstrand 1975; Frost 1985; Klein and Bach 2009; Foustoukos et al. 2015; Sciortino et al. 2015; Evans et al. 2017), Cu-Fe-S-O-H system (Schwarzenbach et al. 2014; 2021), Fe-Ni-Cu-O-S system (de Obeso and Kelemen 2020), and Ni-Fe-S-As minerals (González-Jiménez et al. 2021).

The phase diagram displayed in Figure 11 (modified after Frost 1985) illustrates the phase stability in the Fe-Ni-Co-O-S minerals as a function of $\log f_{O_2}$ and $\log \Sigma S [aS_2(aq) + aSO_2(aq) + aH_2S(aq)]$ at 300 °C and 2 kbar. In fact, f_{O_2} and f_{S_2} were not measured here (which is actually extremely difficult), but an assessment was made based on the comparison of existing parageneses with available diagrams (e.g., Frost 1985). Therefore, Ni minerals in the Fe-Ni-Co-O-S system have been used here in a qualitative way to trace changes in f_{O_2} and f_{S_2} during serpentinization and carbonation (discussed below). In the following, I focused on (1) desulfurization (sulfur-loss) of magmatic pentlandite and formation of awaruite and magnetite, and (2) hydrothermal upgrading of Ni-rich sulfide phases.

Desulfurization (sulfur-loss) of magmatic pentlandite and awaruite formation. The assemblage pentlandite-awaruite-magnetite is linked to lizarditization of Abu Dahr forearc harzburgites; and is commonly observed in partially serpentinized peridotites as the result of destabilization of pentlandite (Eckstrand 1975; Filippidis 1985; Frost 1985; Klein and Bach 2009; Schwarzenbach et al. 2014, 2021; Evans et al. 2017; Arai et al. 2020; González-Jiménez et al. 2021). High quantities of aqueous H₂ released during serpentinization produced a highly reducing and extremely low f_{02} and f_{S2} environment, which caused desulfurization (sulfur-loss) of pentlandite to form awaruite, magnetite, and aqueous H₂S according to the reaction (R11, Klein and Bach 2009; Schwarzenbach et al. 2014; 2021):

$$Ni_{4.5}Fe_{4.5}S_8 + 4H_2(aq) + 4H_2O = 1.5Ni_3Fe + Fe_3O_4 + 8H_2S(aq).$$
(11)

A temperature of 200 °C is suggested for the equilibration of pentlandite/cobaltian pentlandite from Abu Dahr based on its Co content (Fig. 8b; Kaneda et al. 1986). Previous work by Sciortino et al. (2015) and Olade (2019) has documented awaruite formed after primary sulfides with a narrow compositional range of approximately 70 to 75 wt% Ni, whereas Fe-Ni alloys associated with serpentine, olivine, and magnetite show more compositional variation up to 100 wt% Ni. At Abu Dahr, nearly all Ni-Fe alloy awaruite exhibits a wide compositional range from 75 to 93 wt% Ni. Awaruite shows two forms: euhedral and anhedral. Euhedral awaruite embedded in serpentine (below 20 µm; Fig. 4c) may form by primary crystallization from a Ni-rich melt, upgraded to high-Ni awaruite (native Ni) up to 85 wt% Ni (the remaining Ni released from olivine during serpentinization). The solitary phase, euhedral microstructure, and characteristic compositional zonation with a stoichiometric awaruite rim (75 wt% Ni) and Ni-rich awaruite core (85 wt% Ni) indicate that awaruite was formed at higher temperatures from a Ni-rich melt. The most abundant awaruite grains are anhedral spherical usually formed

FIGURE 10. Variation diagrams of Abu Dahr bulk rock MgO vs. (a) SiO_2 , (b) Al_2O_3 , (c) FeO_7 , and (d) CaO and compared with ultramafic rocks from other localities worldwide (see text). Additional data are from Mariana forearc (Parkinson and Pearce 1998); Cemetery Ridge, Arizona (Haxel et al. 2018); and Santa Elena Nappe, Costa Rica (Schwarzenbach et al. 2014). PM primitive mantle (Palme and O'Neill 2003). The arrows indicate the depletion trend of the samples.



by primary Ni-bearing sulfide pentlandite desulfurization (>2 mm; Fig. 4d and 4e). Considering the *P*-*T* conditions estimated for initial serpentinization of the investigated Abu Dahr forearc (\leq 300 °C and <1 GPa), the association of awaruite and magnetite could be stable and in equilibrium with lizardite at log $f_{02} \sim -41$ to -38 and log Σ S <-2 (Fig. 11).

Partial dehydration of serpentinites and Ni-rich awaruite formation. Ni-rich awaruite or native nickel has been reported at various localities in association with heazlewoodite (Augé et al. 1999) and Ni-rich magnetite (Hudson and Travis 1981). Some of these Ni-rich awaruite (native Ni) grains might have crystallized from Ni-rich hydrothermal fluids if f_{s_2} approaches zero and stabilize at sufficiently high f_{o_2} (Dekov 2006), or formed by desulfurization (sulfur-loss) of magmatic pentlandite under a strongly reducing conditions (Ramdohr 1980; Hudson and Travis 1981; Marques de Sá et al. 2018). In contrast, partial dehydration of the La Cabana serpentinites (South-Central Chile) under high-pressure conditions (>1 GPa) generated Ni-rich awaruite in equilibrium with the prograde assemblage antigorite-metamorphic olivine formed within subduction channel at higher f_{o_2} and f_{s_2} (González-Jiménez et al. 2021).

At Abu Dahr, native Ni grains (rimming millerite-Figure 6b) have a composition almost exclusively of Ni (up to 93 wt% Ni) with minor Fe (up to 6.23 wt%) and traces of Cu (up to 1.66 wt% Cu) and Co (up to 0.44 wt% Co). Partial dehydration of the Atgserpentinites of Abu Dahr South under high-pressure conditions (>1 GPa) generated Ni-rich awaruite in equilibrium with the prograde assemblage antigorite-metamorphic olivine at higher f_{02} and f_{52} . These Ni-rich awaruites (native Ni) could register the uptake of Ni from pre-existing Ni-rich sulfides or serpentine, indicating substantial Ni mobility during high *P-T* metamorphism within



FIGURE 11. Phase stability in the Fe-Ni-S-O-H system as a function of $\log f_{O_2}$ and $\log \Sigma S$ (activity of sulfur species) at 300 °C and 2 kbar. The diagram illustrates stabilities for Ni minerals of interest in Abu Dahr serpentinites at 300 °C and 2 kbar (after Frost 1985). Red solid lines mark boundaries between dominant Ni-sulfides-awaruite-magnetite of Abu Dahr. Blue solid lines mark boundaries between Fe-rich phases. Dashed lines are contours of $\log f_{S_2}$ in the fluid. Mineral abbreviations: see Table 2.

subduction channel.

Magnetite is ubiquitous in serpentinites from various environments, ranging from high-temperature magmatic to low-temperature hydrothermal environments (Dare et al. 2014; Nadoll et al. 2014). At Abu Dahr, Ni-rich magnetite (0.40-1.24 wt% NiO) with low Cr content (up to 7.75 wt% Cr₂O₃) reflects its post-magmatic origin. Magnetite formation is commonly described by the reactions R3 and R6. Hydrothermal oxidation of pentlandite (Fig. 5b) to form Ni-rich magnetite has been observed elsewhere (e.g., Ramdohr 1980; Ballhaus 1993; Prichard et al. 2013).

Hydrothermal upgrading of Ni-rich sulfide phases. The assemblage pentlandite-heazlewoodite-magnetite formed during the lizarditization stage with atomic sulfur/metal ratio <1, is indicative of reducing conditions and records very low f_{O_2} and f_{S_2} (Frost 1985; Frost and Beard 2007; Klein and Bach 2009; Marchesi et al. 2013; Evans et al. 2017; González-Jiménez et al. 2021). Such low f_{O_2} is associated with serpentinization at low water/rock ratio (Frost 1985; Alt and Shanks 1998). The desulfurization of pentlandite to form heazlewoodite (Figs. 5b and 6a) can be described by the reaction (R12, after Klein and Bach 2009):

$$Fe_{4.5}Ni_{4.5}S_8 + 6H_2O(aq) = 1.5Ni_3S_2 + 1.5Fe_3O_4 + H_2 + 5H_2S(aq).$$
 (12)

With increasing f_{S_2} the assemblage pentlandite-awaruitemagnetite becomes unstable and further reaction of awaruite with H₂S from fluids may result in the formation of heazlewoodite according to reaction R13 (after Klein and Bach 2009):

$$3Ni_{3}Fe + 6H_{2}S(aq) + 4H_{2}O = 3Ni_{3}S_{2} + Fe_{3}O_{4} + 10H_{2}(aq).$$
 (13)

In the presence of extra sulfur (released from desulfurization of pentlandite or released from the subducted slab (Li et al. 2020) as H_2S , Ni remaining after pentlandite upgrade forms new discrete euhedral heazlewoodite grains in partially serpentinized peridotites by the following reaction (R14, after González-Jiménez et al. 2021):

$$3NiO_{(in silicate)} + S_{2(aq)} = Ni_3S_2 + 3/2 O_2.$$
 (14)

The assemblage pentlandite-heazlewoodite-godlevskite is linked to antigoritization stage accompanied by increasing f_{o_2} and f_{s_2} . Because low-sulfur godlevskite is found exclusively in fully serpentinized rocks, it is considered that godlevskite replaces heazlewoodite and the formation of godlevskite-rich assemblages (Figs. 5a–d) occur with increasing f_{o_2} and f_{s_2} (log $f_{o_2} \sim -32$ to ~ -30 (Fig. 11) in the final stage of serpentinization. Formation of godlevskite by direct replacement of heazlewoodite can be described by the reaction R15.

$$3Ni_3S_2 + 2H_2S(aq) = Ni_9S_8 + 2H_2(aq).$$
 (15)

The assemblage pentlandite-heazlewoodite-godlevskitemillerite is linked to carbonation of the serpentinized peridotites with increasing f_{02} and f_{82} . During carbonation of serpentinized peridotites millerite grows at the expense of the low-sulfur heazlewoodite and godlevskite (Fig. 6a). Formation of millerite occurred by direct replacement of low-sulfur sulfides heazlewoodite or godlevskite according to reactions R16 and R17 (after Klein and Bach 2009):

$$\begin{aligned} Ni_{3}S_{2} + H_{2}S(aq) &= 3NiS + H_{2}(aq). \end{aligned} \tag{16} \\ Ni_{9}S_{8} + H_{2}S(aq) &= 9NiS + H_{2}(aq). \end{aligned} \tag{17}$$

These reactions indicate increasing f_{o_2} and f_{s_2} to values above the quartz-fayalite-magnetite (QFM) buffer. According to the phase relations constructed here (Fig. 11), the reactions occur at a range

of $\log f_{02} > -30$ and a higher value of $\log \Sigma S \sim -1$. The assemblage millerite-native Ni-goethite is linked to partial dehydration of the Atg-serpentinites of Abu Dahr South under highpressure conditions (>1 GPa). The Ni-rich awaruite is generated in equilibrium with the prograde assemblage antigorite-metamorphic olivine. The increases in $aSiO_{2}, f_{02}$, and f_{S2} suggest the destabilization of Ni-rich awaruite to more oxidizing assemblages (Sciortino et al. 2015). Under these conditions, millerite is formed by direct replacement of Ni-rich awaruite (Fig. 6b) according to reaction R18:

$$2Ni_{3}Fe + 6H_{2}S(aq) + 2O_{2} = 6NiS + 2FeOOH + 5H_{2}(aq).$$
 (18)

With an increasing degree of steatitization (carbonation), magnetite is further oxidized to form goethite (Fig. 6b), indicating higher f_{02} . As a result, Fe²⁺ in magnetite is dissolved and precipitated as Fe³⁺ in goethite incorporating Ni (Fe,Ni)O(OH). The overall reactions of magnetite dissolution and goethite formation are represented by R19 and R20 (after He and Traina 2007):

$$Fe_{3}O_{4} + 2H^{+}(aq) + 2H_{2}O = Fe^{2+} + 2Fe(OH)_{3}.$$
 (19)

$$2Fe(OH)_3 + 0.5O_2 + 2H^+(aq) = 2FeOOH + 3H_2O.$$
 (20)

Chalcopyrite is absent in all investigated samples from Abu Dahr and occurs only within a steatized sample associated with millerite-rich assemblage (Fig. 6a). The addition of Cu might be explained by hydrothermal fluid infiltration from proximal mafic lithologies (Schwarzenbach et al. 2014). At Abu Dahr, primary chalcopyrite is not present and the rare mineral chalcopyrite formed by Cu-bearing hydrothermal fluid infiltration from proximal Um Eleiga gabbro complex intruded Abu Dahr Nappe. Cu-dominant Cu-Ni-PGE sulfides (chalcopyrite, bornite, pyrite, pyrrhotite, pentlandite, and mackinawite) do occur in the Um Eleiga gabbro intrusion (Abdel-Halim 2006).

High Co contents of pentlandite in Abu Dahr

Pentlandite, nominally (Fe, Ni)₉S₈, can accommodate variable amounts of Co and Ni in its structure. At Abu Dahr, the cobalt content of pentlandites varies between 1.60 to 7.53 wt%. Based on the reported compositions of Co-pentlandite from different parts of the world (Harris and Nickel 1972; Brickwood 1986; Hughes et al. 2016), I classify pentlandite as end-member pentlandite (<3 wt% Co) or cobaltian pentlandite (>3 up to 20 wt% Co). Cobalt pentlandite (>20 wt% Co) is not detected at Abu Dahr and generally, is a very rare Co-bearing sulfide mineral worldwide. The host peridotites contain magmatic pentlandites, which gradually become enriched in Co during serpentinization. Whole-rock geochemical data show that Ni and Co concentrations range from 1489 to 2590 ppm and 119 to 144 ppm, respectively. There is no significant difference in Ni and Co concentrations between partially and fully serpentinized rocks, suggesting that the serpentinization process occurred isochemically between Ni and Co. Fe, Ni, and Co released from primary silicates during serpentinization incorporated into newly formed magnetite and upgrading the Ni-Co tenors of preexisting primary pentlandite.

The highest concentration of Co in Abu Dahr pentlandites (7.53 wt% Co) is higher than the Co concentrations in pentlandites of komatiitic dunite at Betheno, Western Australia (up to 0.37 wt% Co, Barnes et al. 2011), Dumont pentlandites (average 3.8 wt% Co, Sciortino et al. 2015), and abyssal peridotite pentlandites (up to 4.79 wt% Co, Marchesi et al. 2013). Pentlandites with higher concentrations of Co are reported from Vourinos, Greece (17.7 wt% Co, Tzamos et al. 2016), Massif du Sud, New Caledonia (13.1 wt% Co, Augé et al. 1999), Cemetery Ridge, Arizona (34.4 wt% Co, Schwarzenbach et al. 2014), and Latao, SE Sulawesi (22.0 wt% Co, Rafianto et al. 2012). High-Co sulfides are also reported from plume magmatism-related mantle peridotite xenoliths (e.g., Wang et al. 2010; Hughes et al. 2016).

Globally, significant Co mineralization is associated with ophiolites (e.g., Bou Azzer, Morocco: Ahmed et al. 2009; and Outokumpu, Finland: Peltonen et al. 2008) or slow-spreading ridges (e.g., Rainbow fields, Mid-Atlantic Ridge; Douville et al. 2002; Mozgova et al. 1996; Borodaev et al. 2007). At Abu Dahr, Co contents in pentlandite are ascribed to the sequestration of Ni into heazlewoodite and awaruite, and, furthermore, from the binary plots (Figs. 9a-d), it can be inferred that Co→Fe substitution is dominant in Abu Dahr pentlandites. In addition, strong negative correlations between Co and Fe+Ni and Fe and Co+Ni imply that these elements substituted each other. Similar substitutions were observed in pentlandites from forearc mantle wedge settings worldwide (e.g., Vourinos, Greece: Tzamos et al. 2016; Massif du Sud Peridotite Nappe, New Caledonia: Augé et al. 1999; Cemetery Ridge, Arizona: Haxel et al. 2018; Santa Elena Nappe, Costa Rica: Schwarzenbach et al. 2014; Latao, SE Sulawesi: Rafianto et al. 2012). The cobalt content of pentlandite varies with the degree of serpentinization (Duke 1986), the mineral assemblage with which it is associated (Mozgova et al. 1996), and/or associated with the subduction of Co-enriched oceanic crust (Hughes et al. 2016). One possible explanation for the widespread occurrence of Co-rich pentlandite in Abu Dahr, is that serpentinization took place by reaction of high salinity chloride-rich fluids with cobalt-bearing Ni-Fe alloys, and Co present in the cobalt-bearing Ni-Fe alloys was then released to the fluid and remobilized into the pentlandite.

Similarities to Ni-sulfides and alloys from forearc mantle elsewhere

Online Materials¹ Table S2 illustrates the relationships of Nisulfides and alloys from Abu Dahr with those reported from mantle wedge serpentinites worldwide (e.g., Vourinos, Greece: Tzamos et al. 2016; Massif du Sud Peridotite Nappe, New Caledonia: Augé et al. 1999; Cemetery Ridge, Arizona: Haxel et al. 2018; Santa Elena Nappe, Costa Rica: Schwarzenbach et al. 2014; Latao, SE Sulawesi: Rafianto et al. 2012; Bou Azzer, Morocco: Ahmed et al. 2009). The minerals show mineralogical and chemical composition similarities (Fig. 12). Some of these occurrences contain beside Ni-sulfides and awaruite: Fe-sulfides, arsenides, and sulfersenides (e.g., Bou Azzer, Morocco; Cemetery Ridge, Arizona). Figure 12 shows the distribution of Ni proportions (Ni molar%) of Abu Dahr Ni-sulfide and alloy minerals compared with those reported from Vourinos, New Caledonia, Cemetery Ridge, Santa Elena, Latao, and Bou Azzer. Besides Ni sulfides \pm alloys, the Cemetery Ridge peridotites contain two Ni-arsenides and, less commonly, pyrrhotite (Haxel et al. 2018). Santa Elena Nappe contains Cupentlandite (Cu <3 wt%) and chalcopyrite (Schwarzenbach et al. 2014), while the Massif du Sud harzburgites of New Caledonia ophiolite contain Pt-rich Fe-Ni alloy awaruite and native copper (Augé et al. 1999). The Bou Azzer (Morocco) is dominated by Co-Ni arsenide ores, but according to Ahmed et al. (2009) sulfides occur as accessory phases. The chemical, mineralogical and redox conditions reported in this paper suggest that all forearc mantle wedge settings were subjected to similar processes (e.g., Augé et al. 1999; Rafianto et al. 2012; Schwarzenbach et al. 2014, 2021; Tzamos et al. 2016; Haxel et al. 2018).

Genetic model for Abu Dahr Ni-sulfides and metal alloys

The disseminated Ni mineralization of Abu Dahr consists of high-Ni tenor sulfides and Ni-rich awaruite (native Ni) without Fe sulfides or Ni arsenides. Abu Dahr is one of the serpentinizationrelated Ni-Co mineralization worldwide (e.g., Dumont serpentinites (Canada): Sciortino et al. 2015; Bou Azzer serpentinites (Morocco): Ahmed et al. 2009; abyssal serpentinized and talcaltered peridotites: Alt and Shanks 1998; Klein and Bach 2009; Marchesi et al. 2013; Latao (SE Sulawesi): Rafianto et al. 2012; Decar (British Columbia): Britten 2017; Cemetery Ridge (Southwest Arizona): Haxel et al. 2018; Dangoma (Nigeria): Olade 2019; among others). In all of these occurrences, Ni mineralization is related to hydrothermal alteration accompanying serpentinization of magmatic olivine that liberates Ni-Fe-Co and subsequently forms Ni-rich phases, documenting variations in f_{O2} and f_{S2} . A similar scenario may be suggested in the case of Abu Dahr Ni mineralization, which involved three distinct genetic processes. These are (1) desulfurization of magmatic pentlandite and associated sulfur loss that led to the formation of heazlewoodite and awaruite during initial serpentinization recording extremely low f_{02} and f_{s_2} conditions; (2) in situ precipitation of secondary Ni-sulfides in the presence of extra sulfur as aqueous H2S derived



FIGURE 12. Pie charts showing the wt% proportions of Fe, Co, Ni, Cu, and S in Ni-sulfides and alloys of Abu Dahr comparable with those reported from mantle wedge serpentinites worldwide. Data sources are from Online Materials¹ Table S2.

from the desulfurization of magmatic pentlandite or, native Ni when f_{S_2} approaches 0; and (3) hydrothermal upgrading of Ni-rich sulfide phases from low-sulfur heazlewoodite to godlevskite in complete serpentinization, and finally to high-sulfur millerite-rich assemblage related to later carbonation recording much higher f_{O_2} .

IMPLICATIONS

The disseminated Ni mineralization in Abu Dahr comprises high nickel concentrates of Ni-rich sulfides and awaruite (native Ni) with neither Fe-sulfides nor Ni-arsenides. Within Abu Dahr forearc environment, high-Ni tenor sulfide and alloy phases such as pentlandite (37 wt% Ni; 7.5 wt% Co), heazlewoodite (71 wt% Ni), godlevskite (67 wt% Ni), millerite (66% Ni wt%) occasionally with awaruite (75-84 wt% Ni) and native Ni (93 wt% Ni) are potentially present. In this serpentinized environment, pyrite and pyrrhotite are not present. This facilitates production of high-grade nickel concentrate after metallurgical process according to IGE Nordic (2010): Ni-rich sulfides + no Fe sulfides + no Ni arsenides constitute a high grade nickel concentrate. The Abu Dahr awaruite contains more Ni content than the compositional range of Dumont awaruite Ni60Fe40-Ni90Fe10 (Sciortino et al. 2015) and those described from the Shetland ophiolite (Prichard et al. 1994) and podiform chromitites (Ahmed and Arai 2003). Compared with other nickel potential localities worldwide, the Abu Dahr could be a prospective for hydrothermally upgraded Ni-sulfides and Ni-rich awaruite (native Ni). Heat and fluids supplied by the nearby granite intrusions may transport the newly formed sulfides and alloys to suitable trap-sites. Further mineralogical and geochemical studies and geophysical exploration will be required to evaluate the economic potential of the Abu Dahr property.

ACKNOWLEDGMENTS AND FUNDING

The insightful comments of Margaux Le Vaillant, B. Ronald Frost, and Mervat A. Elhaddad are highly appreciated. I would like to express my sincere gratitude to Suzanne K. Birner and Xiaolu Niu for providing me with Excel spreadsheet programs for P-T and oxygen fugacity calculations. I thank Hongwu Xu, American Mineralogist Editor, and Kate Kiseeva, Associate Editor, for their help through research submission and their valuable comments and suggestions on my manuscript. The reviewers Katy A. Evans and Michael Zelenski are thanked for constructive reviews and their insightful comments and suggestions that improved the manuscript. This research was supported, in part, by the project "The Economic Assessment of Platinum in the Eastern Desert of Egypt," Assiut University, Egypt.

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MANUSCRIPT RECEIVED JULY 13, 2021

MANUSCRIPT ACCEPTED FEBRUARY 23, 2022

ACCEPTED MANUSCRIPT ONLINE MARCH 3, 2022

MANUSCRIPT HANDLED BY KATE S. KISEEVA

Endnote:

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