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Tissemouminites: A new group of primitive achondrites spanning the transition between acapulcoites and winonaites

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Abstract–The Northwest Africa (NWA) 090 meteorite, initially classified as an acapulcoite, presents petrological, chemical, and isotopic characteristics comparable to a group of seven primitive winonaites: Dhofar 1222, NWA 725, NWA 1052, NWA 1054, NWA 1058, NWA 1463, and NWA 8614. Five of these samples were previously classified as acapulcoites or ungrouped achondrites before being reclassified as winonaites based on their oxygen isotopic compositions. These misclassifications are indicative of the particular compositional nature of these primitive achondrites. All contain relict chondrules and a lower closure temperature of metamorphism of \(820 \pm 20\) °C compared to other typical winonaites, as well as mineral elemental compositions similar to those of acapulcoites. The oxygen isotopic signature of these samples, \(\delta^{17}O\) of \(1.18 \pm 0.17\)‰, \(\delta^{18}O\) of \(3.18 \pm 0.30\)‰, and \(\Delta^{17}O\) of \(-0.47 \pm 0.02\), is in fact resolvable from both acapulcoites and winonaites. We investigate the relationship between these eight primitive achondrites, typical winonaites, and acapulcoites, to redefine petrological, mineralogical, and geochemical criteria of primitive achondrite classification. Distinguishing between winonaites, acapulcoites, and this group of eight primitive achondrites can be unambiguously done using a combination of several mineralogical and chemical criteria. A combination of olivine fayalite content and FeO/MnO ratio, as well as plagioclase potassium content allow us to separate these three groups without the absolute necessity of oxygen isotope analyses. NWA 090 as well as the other seven primitive achondrites, although related to winonaites, are most likely derived from a parent body distinct from winonaites and acapulcoites–lodranites, and define a new group of primitive achondrites that can be referred to as tissemouminites.

INTRODUCTION

Acapulcoites–lodranites and winonaites are primitive achondrites (Prinz et al., 1983), meaning they have crystalline igneous, hence achondritic, textures, but with near-chondritic bulk compositions (Krot et al., 2014; Weisberg et al., 2006). These meteorites have sampled partially melted chondritic parent bodies, formed only a few Ma after the formation of calcium-aluminum inclusions (CAIs; Schulz et al., 2010). As such, they provide a window into the early stages of protoplanetary differentiation. Acapulcoites and lodranites most likely originated from a single chondritic parent body, close in composition to H ordinary chondrites (McCoy, Keil, Clayton, et al., 1997; Mittlefehldt, 2014; Palme et al., 1981). These primitive achondrites represent residues of different degrees of partial melting of their chondritic precursor that occurred in a moderately reducing environment (Keil & McCoy, 2018). Lodranites were heated to higher temperatures (1100–1250 °C), resulting in a course-grained texture (540–700 μm) and in 5%–10% of Fe,Ni–FeS cotectic and basaltic partial melting. Migration of these
melts has been observed through the depletion of troilite and plagioclase in some loderanites relative to acapulcoites. Acapulcoites are finer grained achondrites (150–230 µm), as they were heated to lower temperatures (950–1050 °C), producing lower degrees of Fe,Ni–FeS coticet partial melting (1%–4%) without any melt migration. Hence, these meteorites kept a chondritic mineralogy and bulk composition (Floss, 2000; McCoy et al., 1996; McCoy, Keil, Clayton, et al., 1997; McCoy, Keil, Muenow, et al., 1997; Mittlefehldt et al., 1996; Palme et al., 1981; Patzer et al., 2004). Winonaites are generally more fine-grained than acapulcoites (90–250 µm; Zeng et al., 2019); experienced limited Fe,Ni–FeS partial melting, if any (900–1200 °C); and show evidence for brecciation and high metamorphism (similar to type 6 chondrites) on the chondritic precursor body (Benedix et al., 1998, 2005). While the chondritic mineralogy is similar between winonaites and acapulcoites, the winonaites precursor would have formed in a more reduced environment than the acapulcoite–lodranite parent body, with the major element composition lying between enstatite chondrites and H ordinary chondrites (Benedix et al., 1998; Kimura et al., 1992) and similarities in trace element abundances to CM carbonaceous chondrites (Hunt et al., 2017). However, chondritic precursors of both winonaites and acapulcoites–lodranites are unlike any known chondritic groups and our best witnesses for primitive achondrite parent bodies are relic chondrule-bearing samples (Rubin, 2007; Schrader et al., 2017; Zeng et al., 2019). These relic chondrule-bearing samples have been suggested to be called “chondritic acapulcoites” (Neumann et al., 2018). As such this nomenclature, as well as the term “chondritic winonaites,” will be used in the text for simplicity.

To date, 9 of 69 classified winonaites (as of May 2022) contain relic chondrules: Dhofar 1222; Pontlyfni; Mont Morris (Wisconsin); Northwest Africa (NWA) 725, NWA 1463; NWA 1052 and its pairs NWA 1054; NWA 1058; and NWA 8614 (Benedix et al., 1998, 2003; Moggi-Cecchi & Caporali, 2015; Farley et al., 2015; Néri, 2019; Rubin, 2007; Zeng et al., 2019). One should note that Mont Morris is an ambiguous member of this subgroup; while this meteorite is often cited as containing relic chondrules in the literature (Benedix et al., 1998; Greenwood et al., 2012; Schrader et al., 2017), Benedix et al. (1998) actually identified only one suspected chondrule, while Davis et al. (1977), Graham et al. (1977), and Bevan and Grady (1988) did not report any. Moreover, Bevan and Grady (1988) concluded that Mount Morris (Wisconsin) is a detached silicate inclusion from the IAB iron Pine River. Therefore, Mount Morris will not be considered further in this article. Interestingly, from the eight chondritic winonaites, only Pontlyfni, NWA 1463, and NWA 8614 were originally classified in the Meteoritical Bulletin as winonaites. Dhofar 1222, NWA 1052, and NWA 1054 were initially classified as acapulcoites as their mineralogy and chemistry seemed consistent with these primitive achondrites, while NWA 1058 and NWA 725 were classified as ungrouped achondrites. Recently, these samples have been reclassified as winonaites based on their oxygen isotope compositions (Greenwood et al., 2012; Zeng et al., 2019) or cosmic ray exposure (CRE) ages that do not match with the restricted 4–7 Ma CRE age of the acapulcoite–lodranite clan (Eugster & Lorenzetti, 2005). These somehow regular misclassifications highlight two important points: (i) the need to revise the mineralogical and geochemical distinguishing criteria between winonaites and acapulcoite primitive achondrites, some of which are obviously outdated, and (ii) the peculiar nature of this group of chondritic winonaites.

Modal abundance, mineralogy, and chemistry are typically used to distinguish acapulcoites from winonaites (Benedix et al., 1998; Kimura et al., 1992; Yugami et al., 1998); lodranites, being coarser grained, are readily identified from the two previously mentioned groups. Compositions of mafic minerals are generally used to classify primitive achondrites into either the acapulcoite or winonaites groups. A correlated variation plot between ferrosilite (Fs) content of low-Ca pyroxene and fayalite (Fa) content of olivine in primitive achondrites is often used for this purpose. Criteria to distinguish winonaites from acapulcoites include: (i) highly reduced assemblage, among which typically Mg-rich olivine and Mg-rich pyroxene, as well as the presence of highly reduced accessory minerals such as alabandite, daubréelite, and schreibersite; (ii) modal abundance of metal, troilite, and olivine; (iii) major element composition of pyroxene and chromite; (iv) bulk composition (Floss, 2000); and (v) oxygen isotopes (Clayton & Mayeda, 1996; Greenwood et al., 2012, 2017). Because the latter is not always measured to classify a meteorite (e.g., Keil & McCoy [2018] report that more than 50% of classified acapulcoites are lacking oxygen isotopic data), meteoricists rely mostly on these mineralogy and chemistry criteria. However, acapulcoites and winonaites share many common characteristics making their distinction rather delicate, and the rise of primitive achondrite discoveries over the last 20 years has exposed the limit of mineralogical and geochemical distinguishing criteria. This is the case of some acapulcoites that are highly reduced (Fa < 7 mole%) such as Allan Hills (ALH/ALHA) 81187 and ALH 84190, which is suspected to be its pair, as well as Superior Valley (SuV) 014, NWA 13041, NWA 13839, NWA 4816, and NWA 090. While ALH 81187 and ALH 84190 have acapulcoite oxygen isotope signatures, the other five samples lack this information. Hence, a reclassification might be required.

Most chondritic winonaites exhibit an abnormal oxidized state compared to typical winonaites

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*References*

- Benedix et al., 1998
- Greenwood et al., 2012
- Schrader et al., 2017
- Zeng et al., 2019
- Keil & McCoy, 2018
- Eugster & Lorenzetti, 2005
- Hunt et al., 2017
- Clayton & Mayeda, 1996
- Greenwood et al., 2012, 2017
- Clayton & Mayeda, 1996
- Yugami et al., 1998
- Moggi-Cecchi & Caporali, 2015
- Farley et al., 2015
- Néri, 2019
- Rubin, 2007
- Zeng et al., 2019
- Davis et al., 1977
- Graham et al., 1977

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*Notes*

- 9 of 69 classified winonaites contain relic chondrules.
- Mont Morris is an ambiguous member of this subgroup.
- Dhofar 1222, NWA 1052, and NWA 1054 were initially classified as acapulcoites.
- NWA 1058 and NWA 725 were classified as ungrouped achondrites.
- NWA 1058 and NWA 725 were reclassified as winonaites.
- Acapulcoites and winonaites share many common characteristics.
- Superior Valley (SuV) 014, NWA 13041, NWA 13839, NWA 4816, and NWA 090 also exhibit acapulcoite characteristics.
(Fa > 5.2 mole% and/or Fs > 7.1 mole%): Dhofar 1222, NWA 725, NWA 1052 and its pairs NWA 1054, NWA 1058, NWA 1463, and NWA 8614. Several studies have already noted the primitive nature of these chondritic winonaites (Benedit et al., 2003; Farley et al., 2015; Floss et al., 2008; Irving & Rumble, 2006; Néri, 2019; Zeng et al., 2019). In fact, they are all characterized by a protogranular very fine-grained texture, the presence of relatively abundant relic chondrules, higher oxidation state compared to typical winonaites, and a low temperature of metamorphism. In fact, these criteria are indicative of a lack of igneous fractionation processes, otherwise observed in other typical winonaites. Most likely, they sampled the regolith of the winonaites parent body (Floss et al., 2008; Zeng et al., 2019), direct proxy of the chondritic precursor, being the least heated zone of parental bodies. Interestingly, these samples have been suggested likely to be paired and to be named W chondrites (Irving & Rumble, 2006). In fact, it has been recently highlighted that distinction between highly metamorphosed chondrites and primitive achondrites is vague (Tomkins et al., 2020) and chondritic achondrites are transitional between strongly metamorphosed chondrites and primitive achondrites (Greenwood et al., 2012).

Here, we investigate the petrography, mineralogy, chemistry, and isotopic compositions of NWA 090, classified as an acapulcoite (Bouvier et al., 2017), as well as a group of seven primitive winonaites, namely Dhofar 1222, NWA 725, NWA 1052, NWA 1054, NWA 1058, NWA 1463, and NWA 8614 (five of them previously classified as acapulcoites or ungrouped achondrites). These meteorites lie between acapulcoites and winonaites, highlighting the need to revisit the classification criteria of primitive achondrites. Our data indicate a new group of primitive achondrites, likely derived from a parent body different to those of acapulcoite–lodranite and winonaites, despite sharing similar oxygen isotope characteristics as that of winonaites. We suggest naming this new group tissemouminites after the location where the type specimen NWA 725 was found.

SAMPLE AND ANALYTICAL TECHNIQUES

Samples

Northwest Africa 090 was purchased in Erfoud, Morocco in 2000 (Bouvier et al., 2017). NWA 090 shows medium weathering and low shock and a well-developed fusion crust. Several rounded mineral aggregates were identified as well as a radial pyroxene aggregate. NWA 090 was then classified as an acapulcoite based on its mineral major element composition. We study here the same polished thin section of NWA 090 used for the first classification, prepared at Buseck Center for Meteorite Studies, Arizona State University. Northwest Africa 1052 and its pair NWA 1054 section from the Department of Earth Sciences, Firenze, were also analyzed as some mineral chemical composition about these samples were lacking.

Secondary Electron Microscope and Electron Probe Microanalyses

Polished thin section of NWA 090 was studied using optical and scanning electron microscopy (SEM) methods. Microscopic observations under transmitted and reflected light performed at IAPS sample preparation facility were used to determine the main mineral phases and study the presence of relic chondrules. Backscattered electron (BSE) images of minerals and relic chondrules were obtained with the ZEISS EVO MA15 SEM (MEMA Center, University of Firenze). Elemental X-ray maps of Al, Ca, Cr, Cu, Fe, K, Mg, Mn, Na, Ni, P, S, Si, and Ti were collected to identify pyroxene, olivine, feldspar, phosphate, troilite, chromite, taenite, kamacite, and accessory minerals in NWA 090. Chemical characterization of olivine and pyroxene in NWA 090 was obtained using the Cameca SX-100 electron microprobe at the University of Arizona. Chemical characterization of all silicate minerals (olivine, pyroxene, feldspar) and oxides (troilite, chromite, taenite, and kamacite) was carried out with the JEOL JXA 8230 (Department of Earth Sciences, University of Firenze). Using the Cameca SX-100 electron microprobe at the University of Arizona, quantitative olivine analyses were performed in wavelength-dispersive mode, with an accelerating potential of 15 kV. A focused beam of 20 nA was used to analyze pyroxenes. Quantitative mineral analyses done with the JEOL JXA 8230 were performed in wavelength-dispersive mode, with an accelerating potential of 15 kV. ZAF correction was applied to all measurements. Typical detection limits were 0.02%–0.05% for major element oxide abundances. Mineral modal abundances were calculated using XMapTools (Lanari et al., 2014).

Thermodynamic Modeling

Closure temperatures were determined using geothermometers based on two different mineral pairs. We used both chromite–olivine and two-pyroxene systems to determine temperatures, using the MELTS calculator (Sack & Ghiorsio, 1991). Oxygen fugacity was calculated from both these closure temperatures using quartz–iron–fayalite (QIFa) and quartz–iron–ferrosilite (QIFs) equilibria, as in Gardner-Vandy et al. (2012).
Laser Fluorination—Oxygen Isotope Analyses

Oxygen isotope analysis was performed at the Open University (OU), using an infrared laser-assisted fluorination system (Greenwood et al., 2017) on NWA 090. Oxygen was released from the sample (approximate weight of 2 mg) by heating in the presence of BrF5. The released oxygen gas was purified by passing it through two cryogenic nitrogen traps and over a bed of heated KBr. Oxygen gas was analyzed using a MAT 253 dual inlet mass spectrometer. Recent levels of precision obtained on the OU system, as demonstrated by 38 analyses of an in-house obsidian standard, were as follows: 0.053‰ for δ17O; 0.095‰ for δ18O; 0.018‰ for Δ17O (2 SD) (Starkey et al., 2016). Oxygen isotope analyses are reported in standard delta notation. The δ18O value is calculated as δ18O = ([18O/16O]sample/[18O/16O]ref − 1) × 1000 ‰. The δ17O value is calculated as δ17O = ([17O/16O]sample/[17O/16O]ref − 1) × 1000 ‰. The reference for both is Vienna Standard Mean Ocean Water VSMOW. The deviation from the terrestrial fractionation line has been calculated using the linearized format of Miller et al. (2002): Δ17O = 1000 × ln(1 + δ17O/1000) − k1000 ln(1 + δ18O/1000) where k is 0.5247. The sample was pretreated with ethanolamine thioglycolate (EATG) solution to remove terrestrial weathering products.

RESULTS

Petrography

NWA 090, which retains a well-developed fusion crust (Figs. 1 and 2a), displays a protogranular texture, with typical curvilinear grain boundaries between olivine and other silicates (Mercier & Nicolas, 1975) and a lack of 120° triple junctions. The sample is dominated by olivine, low-Ca pyroxene, FeNi metal, and troilite. The granoblastic texture as well as the high amount of opaque phases in NWA 090 (consisting of metal, troilite, chromite, and oxidation products) is clearly noticeable from the BSE map of the thin section (Fig. 1). Terrestrial weathering products are present, resulting in oxidation of FeNi metallic phases, mostly around grains surrounded by porosity or through small cracks (few micrometers; e.g., Fig. 2c). We estimated the amount of terrestrial weathering products at 8.9 wt%. Hence, a W2 weathering grade, indicating a moderate oxidation of metal, is justified (Bouvier et al., 2017; Wlotzka, 1993). The average silicate grain size was uniformly ~80 μm, smaller than the 200 μm initially reported in the Meteoritical Bulletin (Bouvier et al., 2017). Holly-leaf-shaped Fe-Ni metal grains and troilite (dominantly occurring as single crystal) are largely present in the section, as well as few well-crystallized chromite grains.
Fig. 2. BSE images of NWA 090 highlighting various features: a) relict porphyritic olivine chondrule near the fusion crust of NWA 090, (b) large relict porphyritic olivine chondrule, (c) rounded relict porphyritic olivine chondrule. Identification of relict chondrules was aided by the presence of voids around almost all relict chondrules. Notice also patches and veins of terrestrial alteration products, oxidized metallic phases (TW). d) Large relict radial pyroxene chondrule. Most chondrules also contain small blebs of opaque phases. e) Chromite grain in contact with kamacite and a relict chondrule. f) Kamacite overgrowth on a taenite grain. The taenite–kamacite interface is marked by a large Ni spike (cf. Fig. 8).
However, typical Fe,Ni–FeS veins are absent from the section. Porosity in NWA 090 is estimated to be 8% (using ImageJ program), mostly surrounding relict chondrules and grain boundaries (e.g., Fig. 2e).

Modal abundances of NWA 090 have been calculated with XMapTools software (Lanari et al., 2014) based on X-ray maps (Fig. 3; Table 1) (wt%): 11.4 olivine, 37.6 low-Ca pyroxene, 3.6 high-Ca pyroxene, 11.1 feldspar, 0.4 phosphates, 12.9 troilite, 0.5 chromite, 22.4 FeNi metallic phases. Some accessory mineral phases indicative of highly reducing conditions, such as daubreeelite (FeCr₂S₄), alabandite (MnS), and...
schreibersite ([Fe,Ni]$_3$P), were not found. These are typically found in winonaites and enstatite chondrites, while rarely in acapulcoites (Benedix et al., 1998). Only a few schreibersite grains were found in NWA 090.

Finally, 12 relict chondrules were also identified in the section (Fig. 2a–e), with an average diameter of ~400 μm (from 150 to 700 μm). Most of them are easily identifiable as they have retained their primary textures. We estimate that 83% of relict chondrules are porphyritic, mainly olivine-rich, porphyritic olivine (PO) chondrules (Fig. 2a–c and 2e). One large radial pyroxene chondrule has been identified (Fig. 2d) as well as a probable granular olivine chondrule. Relict chondrules represent about 1.5 vol% of the thin section studied here.

Mineral Major Elemental Compositions

Silicates

Composition summary and detailed electron microprobe analysis data for silicate phases in NWA 090 and other chondritic winonaites are reported in Tables 2 and 3, respectively, and presented in Figs. 4–6. Olivine fayalite (Fa) content is 6.1 ± 0.2 and FeO/MnO 12.9 ± 0.8 ($n = 11$) (Figs. 4 and 5). The composition of low-Ca pyroxene is homogeneous, with ferrosilite (Fs) content of 7.4 ± 0.3, wollastonite (Wo) content of 1.3 ± 0.4, and FeO/MnO of 9.8 ± 1.3 ($n = 18$) (Figs. 4 and 6). The Fs and Fa compositions of NWA 090 lie toward the higher range for winonaites and place NWA 090 in the overlapping zone with acapulcoites values (Fig. 4), leading to some confusion in classifying this meteorite. High-Ca pyroxene composition is Fs 3.2 and Wo 45.4 ± 0.6, with a Cr$_2$O$_3$ content of 1.18 ± 0.04 wt% and an Na$_2$O content of 0.70 ± 0.02wt% ($n = 11$) (Fig. 6). Interestingly, relict chondrules are often enriched in high-Ca pyroxenes compared with the matrix. Plagioclase in NWA 090 is enriched in the orthoclase component compared to typical winonaites, with an average composition of An$_{12.2}$Ab$_{82.2}$Or$_{5.3}$ (Figs. 5 and 6).

Opaques

Detailed electron microprobe analysis data for opaque phases in NWA 090 and other chondritic winonaites are

<table>
<thead>
<tr>
<th>Samples</th>
<th>Opx</th>
<th>Ol</th>
<th>Pl</th>
<th>Cpx</th>
<th>Php</th>
<th>Tro</th>
<th>Chr</th>
<th>Metal</th>
<th>Acc.</th>
<th>Silicate</th>
<th>Opaque</th>
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<td>40.3</td>
<td>11.8</td>
<td>10.1</td>
<td>5</td>
<td>0.7</td>
<td>10.3</td>
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<td>0.4</td>
<td>20.1</td>
<td>—</td>
<td>68.6</td>
<td>27.5</td>
<td>Moggi-Cecchi and Caporali (2015)</td>
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<td>36.8</td>
<td>26.3</td>
<td>15.8</td>
<td>5.3</td>
<td>—</td>
<td>6.2</td>
<td>0.1</td>
<td>9.5</td>
<td>84.2</td>
<td>15.8</td>
<td>Moggi-Cecchi and Caporali (2015)</td>
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<td>16</td>
<td>5.3</td>
<td>0.4</td>
<td>7.5</td>
<td>0.5</td>
<td>8.5</td>
<td>—</td>
<td>85.2</td>
<td>16.1</td>
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<td>46</td>
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<td>10.5</td>
<td>3.8</td>
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<td>10.7</td>
<td>0.7</td>
<td>14.8</td>
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<td>73.7</td>
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<td>37.6</td>
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<td>0.4</td>
<td>12.9</td>
<td>0.5</td>
<td>22.4</td>
<td>tr. 64.1</td>
<td>35.8</td>
<td>This study</td>
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<tr>
<td>NWA 8614</td>
<td>44.3</td>
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<td>20.4</td>
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<td>0.1</td>
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<td>25.1</td>
<td>Hunt et al. (2017)</td>
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</table>

Range of modal abundances for typical winonaite is given for comparison. Tr. for trace as few grains of schreibersite have been identified in NWA 090 section. One should note that modal abundance estimations were done with diverse methods.

Table 2. Summary of silicate chemical compositions of winonaites with relict chondrules.

<table>
<thead>
<tr>
<th>Samples</th>
<th>Opx</th>
<th>Ol</th>
<th>Fa</th>
<th>Opx Fs</th>
<th>Opx Wo</th>
<th>Cpx Fs</th>
<th>Cpx Wo</th>
<th>An</th>
<th>Ab</th>
<th>Or</th>
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<td>Zeng et al. (2019)</td>
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<td>11.7</td>
<td>7.9</td>
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<td>3.3</td>
<td>46.1</td>
<td>13.4</td>
<td>81.2</td>
<td>5.4</td>
<td>Patzer et al. (2004)</td>
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<tr>
<td>NWA 1052</td>
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<td>—</td>
<td>7.6</td>
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<td>6.4</td>
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<td>1.3</td>
<td>3.3</td>
<td>45.4</td>
<td>14.3</td>
<td>79.5</td>
<td>6.3</td>
<td>Metbull</td>
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<tr>
<td>Dhofar 1222</td>
<td>6.6</td>
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<td>7.7</td>
<td>1.3</td>
<td>3.9</td>
<td>45.6</td>
<td>13.3</td>
<td>80.7</td>
<td>6.0</td>
<td>Néri (2019)</td>
<td></td>
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<tr>
<td>NWA 090</td>
<td>6.1</td>
<td>12.9</td>
<td>7.4</td>
<td>1.3</td>
<td>3.2</td>
<td>45.4</td>
<td>12.2</td>
<td>82.2</td>
<td>5.3</td>
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<tr>
<td>ISD</td>
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<td>0.8</td>
<td>0.3</td>
<td>0.4</td>
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<td>0.1</td>
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<td>1.4</td>
<td>3.5</td>
<td>45.5</td>
<td>13.6</td>
<td>79.9</td>
<td>6.4</td>
<td>Farley et al. (2015)</td>
<td></td>
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<td>Pontlyfni</td>
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<td>1.2</td>
<td>1.6</td>
<td>1.1</td>
<td>46.1</td>
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<td>—</td>
<td>—</td>
<td>Benedix et al. (1998)</td>
<td></td>
</tr>
<tr>
<td>NWA 1463</td>
<td>3.2</td>
<td>—</td>
<td>7.4</td>
<td>1.2</td>
<td>—</td>
<td>—</td>
<td>12.6</td>
<td>82.8</td>
<td>4.6</td>
<td>MetBull</td>
<td></td>
</tr>
</tbody>
</table>

Table 1. Modal abundances of winonaites containing relict chondrules.
reported in Table 4. FeNi metal is mainly represented by kamacite. Taenite is also present and a crystal of taenite with an overgrowth of kamacite was found in the section (Fig. 2f). Ni profiles across the taenite–kamacite show an Ni spike of about 5 wt% in the taenite that occurs at the flat taenite/kamacite interface (Fig. 7). Kamacite nucleation and growth around a taenite matrix occurred upon cooling (Wood, 1967) and can be used as metallographic cooling rate of the sample. The NWA 090 cooling rate was determined using the taenite central Ni content method (also known as the Wood method). We used the cooling rate curves calculated in ordinary chondrites by Willis and Goldstein (1981) and extrapolated by Taylor et al. (1987), in a similar manner as done by Scott et al. (2014b). Considering a taenite half-width of ~80 μm with a central Ni concentration of 17.5 wt%, we
estimated a cooling rate for NWA 090 of about 100 °C yr/C0. The only sulfides in NWA 090 are troilite. Cr and Ni were below the detection limit. Well-crystallized chromite is also present (Fig. 2e) and shows an enrichment in V2O3 compared to typical winonaites, with an average of 0.52/C6 0.07 wt%, as well as an enrichment in iron compared to winonaites, with FeO content of 16.92/C6 0.50 wt% (Fig. 8). The presence of taenite (Fig. 2f) as well as well-crystallized oxidized mineral like chromite, and no trace of typical highly reduced accessory minerals like daubréelite and alabandite are indicative of NWA 090 being FeO-rich compared to winonaites (Benedix et al., 1998; Li et al., 2011; McCoy et al., 1996; Patzer et al., 2004; Yugami et al., 1998).

Thermodynamic Modeling

The closure temperatures and corresponding oxygen fugacities for both olivine–chromite and two-pyroxene systems are presented in Table 5 and Fig. 9. Two-pyroxene thermometry for NWA 090 yields a closure temperature of 827 °C. For comparison, we also recalculated the temperature of Dhofar 1222, NWA 1058, and NWA 725, which yield similar temperatures of 818 °C, 805 °C, and 831 °C, respectively. The olivine–chromite thermometry of NWA 090 gives a temperature of 657 °C. Similarly, Dhofar 1222, NWA 1058, and NWA 725 temperatures were estimated at 657 °C, 640 °C, and 649 °C, respectively. The lower temperature recorded by the olivine–chromite system is consistent with Fe-Mg exchange between these minerals closing at lower temperature than Ca-transfer between pyroxenes. The olivine–chromite oxygen fugacity recorded by the four sections ranges from 2.56 to 2.49 log units below the iron–wustite (IW) oxygen buffer, with an average 2.53/C6 0.05 log units. The pyroxene oxygen fugacity ranges from 2.17 to 2.14 log below the IW buffer, with an average 2.16/C6 0.03 log units. Olivine–chromite and two-pyroxene thermometers exhibit a strong correlation among the four samples.

Oxygen Isotopic Composition

The oxygen isotopic composition of NWA 090 and other samples is presented in Fig. 10 and in Table 6, along with other primitive achondrites with relict chondrules. Analysis of NWA 090 by laser fluorination
gave δ17O = +1.12, δ18O = +3.32, and Δ17O = −0.46. δ17O and δ18O lie on the fractionation line of winonaites, although values are lower. Oxygen isotopic composition of NWA 090 is similar to those of Dhofar 1222, NWA 725, NWA 1052, NWA 1054, NWA 1058, NWA 1463, and NWA 8614.

**DISCUSSION**

**Northwest Africa 090 and Other Chondritic Winonaites in the Winonaites–Acapulcoite Dichotomy: A New Group of Primitive Achondrites**

The petrographic, chemical, and isotopic characteristics of NWA 090 conflict with the traditional dichotomy between acapulcoites and winonaites. In fact, most of NWA 090’s characteristics are ambiguous. Petrographically, NWA 090 displays typical curvilinear grain boundaries between olivine and other silicates and a lack of 120° junctions. These two features suggest less extensive solid-state recrystallization in NWA 090 than typical winonaites, which usually exhibit equigranular textures and abundant triple junctions (Benedix et al., 1998). The average 80 μm silicate grain size measured in NWA 090 is also typically lower than in winonaites (Zeng et al., 2019), and within the range of other chondritic winonaites, that is, Dhofar 1222 and NWA 725 (~45 to 55 μm; Néri, 2019). The characteristics of easily identified relict chondrules are similar to those in H chondrites and chondritic acapulcoites (Friedrich et al., 2015; Gooding & Keil, 1981; Scott & Krot, 2014), with a diameter range of 400 μm (OCs: 450 μm;...
A new group of primitive achondrites

(a) Mgo (wt.% low-Ca px) vs. Low-Ca pyroxene FeO (wt.%)

(b) Na₂O (wt.% high-Ca px) vs. Cr₂O₃ (wt.% high-Ca px)

(c) A₂O₃ (wt.% high-Ca px) vs. Mg# high-Ca px

(d) Cr₂O₃ (wt.% high-Ca px) vs. Mg# high-Ca px

(e) Or vs. Ab

Winonaites
Acapulcoites
Dhofar 1222
NWA 090
NWA 725
NWA 1052
NWA 1054
NWA 1058
NWA 1058
NWA 8614
NWA 1463

AlHA B1187
Table 4. Average kamacite, taenite, troilite, and chromite chemical composition of NWA 090 and some winonaites with relic chondrules.

<table>
<thead>
<tr>
<th></th>
<th>NWA 090</th>
<th>NWA 725</th>
<th>Dhofar 1222</th>
<th>NWA 1058</th>
</tr>
</thead>
<tbody>
<tr>
<td>Fe</td>
<td>92.07 ± 0.60</td>
<td>75.62 ± 9.25</td>
<td>63.15 ± 0.06</td>
<td></td>
</tr>
<tr>
<td>Ni</td>
<td>6.93 ± 0.10</td>
<td>23.39 ± 8.39</td>
<td>0.01 ± 0.02</td>
<td></td>
</tr>
<tr>
<td>Mn</td>
<td>—</td>
<td>—</td>
<td>0.01 ± 0.01</td>
<td></td>
</tr>
<tr>
<td>S</td>
<td>—</td>
<td>—</td>
<td>37.36 ± 0.07</td>
<td></td>
</tr>
<tr>
<td>Cu</td>
<td>—</td>
<td>—</td>
<td>0.03 ± 0.03</td>
<td></td>
</tr>
<tr>
<td>Total</td>
<td>99.00 ± 0.60</td>
<td>99.01 ± 0.86</td>
<td>100.56 ± 0.07</td>
<td></td>
</tr>
</tbody>
</table>

Data on NWA 725 and Dhofar 1222 are from Néri (2019); data on NWA 1058 are from Patzer et al. (2004).

acapulcoites: 400–700 μm; Friedrich et al., 2015) and a greater abundance of porphyritic relict chondrules.

As for most of the winonaites, low-Ca pyroxene is the most abundant mafic silicate, consistent with the reduced nature of winonaites compared to ordinary chondrites (Benedix et al., 1998). The modal abundances of NWA 090 and its olivine/opaque ratio (Fig. 3) are evocative of winonaites (Moggi-Cecchi & Caporali, 2015; Hunt et al., 2017; Zeng et al., 2019). The overlapping of Fs and Fa signatures between NWA 090 and acapulcoites (Fig. 4) and the presence of taenite and the absence of other highly reduced accessory minerals attest to NWA 090 being less reduced and more FeO-rich than typical winonaites. Interestingly, chemical elemental compositions of low-Ca and high-Ca pyroxenes set NWA 090 in the overlapping zone between winonaites and acapulcoite (Fig. 6). Low-Ca pyroxene in NWA 090 has an MnO content of 0.54 wt%, closer to the acapulcoite range (0.48–0.84 wt%) than to the winonaites range (0.17–0.53 wt%) (cf. Fig. 6a) (Benedix et al., 1998; Keil & McCoy, 2018; Kimura et al., 1992; Li et al., 2011; Yugami et al., 1998; Zeng et al., 2019). Similarly, high-Ca pyroxene in NWA 090 has an Na₂O content of 0.70 wt %, also within the acapulcoite range (0.63–0.84 wt%) rather than the winonaites range (0.30–0.69 wt%, apart from Tierra Blanca at 1.51 wt%) (cf. Fig. 6b and references therein). However, when plotting Al₂O₃ or Cr₂O₃ of high-Ca pyroxene versus its Mg# (Mg/

[Table 4. Average kamacite, taenite, troilite, and chromite chemical composition of NWA 090 and some winonaites with relic chondrules.]

<table>
<thead>
<tr>
<th></th>
<th>NWA 090</th>
<th>NWA 725</th>
<th>Dhofar 1222</th>
<th>NWA 1058</th>
</tr>
</thead>
<tbody>
<tr>
<td>TiO₂</td>
<td>0.87 ± 0.12</td>
<td>0.86</td>
<td>0.89</td>
<td>0.93</td>
</tr>
<tr>
<td>Al₂O₃</td>
<td>3.10 ± 1.40</td>
<td>5.2</td>
<td>5.39</td>
<td>5.29</td>
</tr>
<tr>
<td>V₂O₃</td>
<td>0.52 ± 0.07</td>
<td>0.53</td>
<td>0.52</td>
<td>—</td>
</tr>
<tr>
<td>Cr₂O₃</td>
<td>65.71 ± 1.71</td>
<td>64.56</td>
<td>63.7</td>
<td>64.89</td>
</tr>
<tr>
<td>FeO</td>
<td>16.92 ± 0.50</td>
<td>16.31</td>
<td>16.24</td>
<td>16.34</td>
</tr>
<tr>
<td>MnO</td>
<td>2.49 ± 0.41</td>
<td>2.47</td>
<td>2.26</td>
<td>2.56</td>
</tr>
<tr>
<td>MgO</td>
<td>8.56 ± 0.68</td>
<td>9.12</td>
<td>9.36</td>
<td>9.34</td>
</tr>
<tr>
<td>Total</td>
<td>98.17 ± 0.58</td>
<td>99.07</td>
<td>98.4</td>
<td>99.35</td>
</tr>
</tbody>
</table>

Data on NWA 725 and Dhofar 1222 are from Néri (2019); data on NWA 1058 are from Patzer et al. (2004).

Interestingly, most of NWA 090’s features match those of other classified chondritic winonaites, that is, Dhofar 1222, NWA 725, NWA 1052, NWA 1054, NWA 1058, NWA 1463, NWA 8614 (Figs. 3–6, 8). It is critical to note the special case of Pontlyfni regarding the other classified winonaites with relict chondrules. Pontlyfni does not fit with the other eight samples for any criteria (e.g., petrography, mineralogy, chemistry, or isotopy). In fact, the relict chondrules in Pontlyfni are not straightforward to identify (Benedix et al., 1998) and were not even identified by Graham et al. (1977), while the relict chondrules in the other samples are numerous (1.5–5 vol%) and barely altered (Moggi-Cecchi & Caporali, 2015; Farley et al., 2015; Rubin, 2007; Zeng et al., 2019). As such, it seems that Pontlyfni does not belong to this particular “chondritic achondrite” group and might be the only real winonaites-bearing relict chondrule. In terms of modal abundances, Dhofar 1222, NWA 1052, NWA 8614, NWA 1463, and NWA 725 have an olivine/opaque ratio below 0.8 (Fig. 3), concordant with NWA 090 and a winonaite affiliation. NWA 1058, NWA 1052, and its pair NWA 1054 do not concur with the other chondritic winonaites. However, the large range of modal abundances among the typical winonaites indicate that the parent body of winonaites must have been heterogeneous (Floss et al., 2008; Hunt et al., 2017). As such, this criterion cannot be used to exclude samples from a primitive achondrite group. Interestingly, NWA 1058 has been suggested to be paired with NWA 725, NWA 1052, and NWA 1463 (Irving & Rumble, 2006). However, regarding the variations in modal abundances among these samples (Table 1), we
argue against such a pairing, favoring a common parent body origin. Nonetheless, these eight samples match extremely well for all the other criteria of composition (Figs. 3–6 and 8; Tables 2 and 3).

The estimated closure temperature of 830 °C for NWA 090 (Fig. 9) as well as the absence of sulfide veins indicate that this primitive achondrite is one of the least thermally metamorphosed samples from primitive achondrites, since most winonaites and acapulcoites record metamorphic temperatures higher than 950 °C (Benedix et al., 1998; Hunt et al., 2017), between the Fe, Ni–FeS cotectic temperature (Kullerud, 1963) and the basaltic partial melting temperature (Morse, 1980). As such, NWA 090 did not experience any silicate partial melting. Dhofar 1222, NWA 725, and NWA 1058 recorded similar closure temperatures (Fig. 9), between 805 and 830 °C, while Farley et al. (2015) estimated a closure temperature at ~820 to 860 °C for NWA 8614. The lack of Fe veins, Ni-metal, and troilite indicates that NWA 1463 did not reach the Fe,Ni–FeS cotectic (Hunt et al., 2017). While two-pyroxene thermometry likely gives a closure temperature lower than the actual peak temperature of metamorphism (Benedix et al., 1998), it seems that none of the chondritic winonaites experienced any partial melting. Given these recorded temperatures, the oxygen fugacity for the grouplet of chondritic winonaites is estimated from two-pyroxene thermometry at 2.16 ± 0.03 log units below the IW oxygen buffer and between winonaites and acapulcoites (Wadhwa, 2008).

Indeed, winonaites and silicate inclusions in IAB irons range from 2.3 to 3.2 log units below the Fe-FeO buffer (Benedix et al., 2005) while acapulcoites range from ~1.85 to ~3.09 log units (Benedix & Lauretta, 2006; Righter et al., 2016). The closure temperatures recorded by these samples are rather similar to the H5/H6 ordinary chondrite range of peak temperatures (Huss et al., 2006; McSween et al., 2002). Criteria of Van Schmus and Wood (1967) to determine the petrologic type of chondrite involve, among others, heterogeneity of silicate minerals based on the PMD (percent mean deviation) of FeO content, the presence of well-crystallized feldspar (Kovach & Jones, 2010), and the presence of taenite and texture of chondrules (Huss et al., 2006). The presence of chondrules that are readily delineated, a homogenous olivine composition (PMD = 2.5) and slightly heterogenous pyroxene composition (PMD = 5.1) argue for a type 5 for NWA 090, similar to that inferred for NWA 1463 (Floss et al., 2008; Hunt et al., 2017), while the presence of taenite and abundant plagioclase >50 μm (modal abundance 11.1%) is indicative of a highly metamorphosed chondrite closer to a type 6, similar to that inferred for NWA 8614 (Farley et al., 2015). As a matter of fact, Irving & Rumble (2006) inferred that NWA 1054 was a chondrite type 5/6 related to the winonaites and suggested to call these samples W chondrites. In their metamorphic facies series for
equilibrated stony meteorites, Tomkins et al. (2020) showed that NWA 725 falls within the plagioclase facies, meaning that it has equilibrated sufficiently to grow plagioclase over 10 μm in size, but has not yet undergone silicate partial melting. They also estimated this facies to be equivalent to petrologic types 5 and 6. The pristine characteristic of these samples has already been suggested: Benedix et al. (2003) and Floss et al. (2008) highlighted the unique features of NWA 1463 compared to common winonaites, in particular bulk and trace element abundances, and inferred that it sampled the winonaite chondritic precursor. Equally, Zeng

Fig. 8. Two different microprobe Ni traverses across the taenite–kamacite intergrowth. An Ni spike occurs at the flat taenite/kamacite interface. (Color figure can be viewed at wileyonlinelibrary.com.)

Table 5. Thermodynamic properties determined for this study using data from Tables 3 and 4 for Dhofar 1222, NWA 090, NWA 1058, NWA 725.

<table>
<thead>
<tr>
<th>Meteorite</th>
<th>Olivine–chromite Temperature (°C)</th>
<th>QIFa log(fO₂)</th>
<th>Δ IW</th>
<th>Two-pyroxene Temperature (°C)</th>
<th>QIFs log(fO₂)</th>
<th>Δ IW</th>
<th>ΔT</th>
</tr>
</thead>
<tbody>
<tr>
<td>Dhofar 1222</td>
<td>657</td>
<td>−25.4</td>
<td>−2.49</td>
<td>818</td>
<td>−20.7</td>
<td>−2.16</td>
<td>161</td>
</tr>
<tr>
<td>NWA 090</td>
<td>657</td>
<td>−25.5</td>
<td>−2.54</td>
<td>827</td>
<td>−20.5</td>
<td>−2.17</td>
<td>170</td>
</tr>
<tr>
<td>NWA 1058</td>
<td>640</td>
<td>−26.1</td>
<td>−2.56</td>
<td>805</td>
<td>−21.0</td>
<td>−2.17</td>
<td>165</td>
</tr>
<tr>
<td>NWA 725</td>
<td>649</td>
<td>−25.7</td>
<td>−2.53</td>
<td>831</td>
<td>−20.3</td>
<td>−2.14</td>
<td>182</td>
</tr>
<tr>
<td>Average</td>
<td>651</td>
<td>−25.7</td>
<td>−2.53</td>
<td>820</td>
<td>−20.6</td>
<td>−2.16</td>
<td>170</td>
</tr>
<tr>
<td>2SD</td>
<td>14</td>
<td>0.5</td>
<td>0.05</td>
<td>0.5</td>
<td>0.03</td>
<td></td>
<td>18</td>
</tr>
</tbody>
</table>

characteristic of these samples has already been suggested: Benedix et al. (2003) and Floss et al. (2008) highlighted the unique features of NWA 1463 compared to common winonaites, in particular bulk and trace element abundances, and inferred that it sampled the winonaite chondritic precursor. Equally, Zeng
et al. (2019) and Farley et al. (2015) proposed that NWA 725 and NWA 8614, respectively, had sampled the chondritic protolith of the winonaite parent body, remained unaltered in the parent body’s regolith. As such, they span the transition between winonaites and acapulcoites, as well as between primitive achondrites and highly metamorphosed chondrites.

 Nonetheless, from a petrological point of view, that is, protogranular texture and abundant relict chondrules, and a mineralogic point of view, that is, olivine, low-Ca pyroxene, high-Ca pyroxene, and chromite composition, NWA 090, Dhofar 1222, NWA 725, NWA 1052 and its pairs NWA 1054, NWA 1058, NWA 1463, and NWA 8614 do not match the composition of typical winonaites and are rather close to acapulcoite characteristics. Oxygen isotopes have been referred as the only tool able to clearly distinguish between winonaites and acapulcoites (Benedix et al., 1998; Greenwood et al., 2012; Li et al., 2011). While NWA 090 entails an oxygen isotope signature that would attribute this primitive achondrite to the winonaite group rather than the acapulcoite–lodranite clan (Fig. 10), one should note that all the relict chondrule-bearing winonaites have δ17O and δ18O values lower than typical winonaites. This has already been highlighted by Greenwood et al. (2012). Indeed, the oxygen isotopic compositions of winonaites, treated by EATG for removing any terrestrial
alteration, which is known to shift the isotopes, are δ^{17}O of 2.01 ± 0.68‰ (1σ), δ^{18}O of 4.81 ± 1.29‰ (1σ), and Δ^{17}O of −0.51 ± 0.04 (1σ) (Greenwood et al., 2012). Considering the average of only EATG-treated samples, that is, NWA 725, NWA 1052, NWA 1463, NWA 8614, and NWA 090 (Table 6 and references therein), the oxygen isotopic signatures for this group of primitive achondrites are narrowed to δ^{17}O of 1.18 ± 0.17‰, δ^{18}O of 3.18 ± 0.30‰, and Δ^{17}O of −0.47 ± 0.02. This subgroup of primitive achondrites is resolvable in terms of their δ^{18}O values from the typical winonaites oxygen isotopic compositions (Fig. 10), although plotting on the same
Table 6. Oxygen isotope signatures of primitive achondrites.

<table>
<thead>
<tr>
<th>Meteorite</th>
<th>Weathering treatment</th>
<th>δ17O/‰</th>
<th>1σ</th>
<th>δ18O/‰</th>
<th>1σ</th>
<th>Δ17O</th>
<th>1σ</th>
<th>References</th>
</tr>
</thead>
<tbody>
<tr>
<td>NWA 725</td>
<td>EATG residue</td>
<td>1.03</td>
<td>0.05</td>
<td>2.92</td>
<td>0.11</td>
<td>−0.5</td>
<td>0.01</td>
<td>Greenwood et al. (2012)</td>
</tr>
<tr>
<td>NWA 1058</td>
<td>Not specified</td>
<td>1.33</td>
<td>0.09</td>
<td>3.57</td>
<td>0.21</td>
<td>−0.53</td>
<td>0.09</td>
<td>Russell et al. (2003)</td>
</tr>
<tr>
<td>NWA 1052</td>
<td>EATG residue</td>
<td>1.01</td>
<td>0.12</td>
<td>3.50</td>
<td>0.21</td>
<td>−0.48</td>
<td>0.01</td>
<td>Greenwood et al. (2012)</td>
</tr>
<tr>
<td>Dhofar 1222</td>
<td>Untreated</td>
<td>1.38</td>
<td>0.17</td>
<td>3.54</td>
<td>0.30</td>
<td>−0.43</td>
<td>0.05</td>
<td>Farley et al. (2015)/MB</td>
</tr>
<tr>
<td>NWA 1054</td>
<td>Not specified</td>
<td>1.44</td>
<td>0.12</td>
<td>3.52</td>
<td>0.21</td>
<td>−0.45</td>
<td>0.01</td>
<td>Greenwood et al. (2012)</td>
</tr>
<tr>
<td>NWA 8614</td>
<td>EATG residue</td>
<td>1.42</td>
<td>0.17</td>
<td>3.54</td>
<td>0.30</td>
<td>−0.46</td>
<td>0.05</td>
<td>This study</td>
</tr>
<tr>
<td>NWA 1463</td>
<td>EATG residue</td>
<td>1.25</td>
<td>0.15</td>
<td>3.3</td>
<td>0.27</td>
<td>−0.48</td>
<td>0.02</td>
<td>Greenwood et al. (2012)</td>
</tr>
<tr>
<td>Average (n = 8)</td>
<td></td>
<td>1.25</td>
<td>0.17</td>
<td>3.32</td>
<td>0.29</td>
<td>−0.47</td>
<td></td>
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</tr>
<tr>
<td>1σ</td>
<td></td>
<td>0.17</td>
<td></td>
<td>0.29</td>
<td></td>
<td></td>
<td>0.03</td>
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</tr>
<tr>
<td>Average treated samples (n = 5)</td>
<td></td>
<td>1.18</td>
<td></td>
<td>3.18</td>
<td></td>
<td>−0.47</td>
<td></td>
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<tr>
<td>1σ</td>
<td></td>
<td>0.17</td>
<td></td>
<td>0.30</td>
<td></td>
<td>−0.29</td>
<td>0.02</td>
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</table>

Classification Scheme of Primitive Achondrites

Since these classified chondritic primitive achondrites, aka tissemouminites, conflict with the traditional dichotomy between acapulcoites and winonaites, we propose a revised criteria traditionally used for primitive achondrite classification (Benedix et al., 1998; Kimura et al., 1992; Li et al., 2011; Yugami et al., 1998). This will allow us to ascertain if, as proposed previously (Li et al., 2011), solely using the silicate mineralogy as a distinguishing criterion between primitive achondrites is risky, requiring the bulk oxygen isotopic composition as the only unambiguous way of distinguishing primitive achondrites. One of the first criteria used is the Fa and Fs content of olivine and low-Ca pyroxene, respectively (Fig. 4). Kimura et al. (1992) defined winonaites with an Fa < Fa6. As of now, acapulcoites and winonaites overlap in the Fa4–7 zone (Fig. 4), making this criterion probably the most confusing one for primitive achondrite classification. In fact, this criterion has led most of tissemouminites to be classified as acapulcoites, then reclassified as winonaites while being in fact affiliated to a distinct primitive achondrite group (The Meteoritical Bulletin database). All winonaites have a Fa < Fa7 (Fig. 4). So far, tissemouminites have an Fa content between Fa3.2 and Fa6.6 (Table 2 and references therein), that is, also less than Fa7. This translates as a primitive achondrite; having an Fa > Fa7 is undeniably an acapulcoite. In fact, only seven classified acapulcoites yield Fa content lower than Fa7, meaning they could be potential winonaites or tissemouminite candidates: Yamato (Y) 983,237 (Fa6.7), NWA 4618 (Fa5.6), SuV 014 (Fa4.6), ALHA 81187 (Fa4.2) and its pairs ALH 84190 (Fa4.4), NWA 13041 (Fa4.1), and NWA 13839 (Fa4.8) (Keil & McCoy, 2018). The bulk oxygen isotopes of ALHA 81187 and ALH 84190 have identified them as acapulcoites while this information is lacking for the other samples. These seven acapulcoites will be discussed below through the criteria. Conversely, an Fa < Fa3.2 would be consistent with a winonaites affiliation. Between Fa3.2 and Fa7, this criterion is not conclusive and should be used with caution.

Modal abundances of silicate minerals as well as phosphates and opaques are generally similar between lodranites, acapulcoites, and winonaites (Fig. 2). In fact, modal abundances of olivine, low-Ca pyroxene, and opaque minerals vary drastically within each of these three groups of primitive achondrites (cf. Zeng et al. [2019] and references therein). Kimura et al. (1992) have noted that the abundance ratios of olivine to orthopyroxene in
Winonaites are generally lower than those of acapulcoites, but since then, discoveries of new winonaites have made this criterion nonconclusive. However, abundance ratios of olivine to opaque phases (i.e., <0.8) can be used to distinguish winonaites and tissuemouminites from the acapulcoite–lodranite clan, that is, a low abundance of olivine (<15 wt%); acapulcoites ranging from 15 to 41 wt%; Keil & McCoy, 2018) together with a high opaque content (>30 wt%); acapulcoites ranging from 1 to 27 wt%; Keil & McCoy, 2018) (Fig. 2a). That is, an olivine to opaque ratio lower than 0.8 will exclude an acapulcoite affiliation (Fig. 2b). It also follows that winonaites and tissuemouminites cannot be distinguished based on modal abundances alone, their modal abundances being extremely similar for all phases.

Kimura et al. (1992) have noted chemical composition differences between low-Ca pyroxenes, as well as high-Ca pyroxenes, in acapulcoites and winonaites. Low-Ca pyroxenes in winonaites tend to be FeO and MnO poorer than acapulcoites ones, while high-Ca pyroxenes are Na2O and Cr₂O₃ poorer in winonaites compared to acapulcoites. As for modal abundances and Fa-Fs contents, the gap initially present between these two primitive meteorite clans has been filled, with more winonaite and acapulcoite analyses (Fig. 6). As such, while the assertions made by Kimura et al. (1992) are in average still correct, some overlap occurred between low-Ca and high-Ca pyroxene chemical composition of acapulcoites and winonaites. Interestingly, the eight samples of interest, aka tissuemouminites, are intermediate in composition between acapulcoites and winonaites. As such, these minerals are not decisive proxies to distinguish between the three groups of primitive achondrites, except for extremely poor or low content of MnO and FeO content of low-Ca pyroxene, or extremely poor or rich content of Na₂O, Cr₂O₃, Al₂O₃, and MgO of high-Ca pyroxene (cf. Fig. 6). However, while plagioclase composition of acapulcoites and winonaites is indistinguishable, with composition of An₃₋₅Αb₈₋₉Or₀₋₂₋₆₋₇ for acapulcoites and winonaites, respectively (Keil & McCoy, 2018), tissuemouminites are typically enriched in K₂O content (0.87–1.06 wt%; Fig. 6e) (Néri, 2019; Patzer et al., 2004; Zeng et al., 2019), yielding a range of composition of An₁₂₋₁₄Αb₈₋₁₀Or₀₋₁₂₋₆ (Table 3 and references therein). Similarly, vanadium composition of chromite can distinguish between winonaites and acapulcoite (Fig. 8) but will not discriminate the new group of primitive achondrites. One should note here that the number of primitive achondrite chromite composition in the literature is limited.

Olivine might be the most useful mineral to classify primitive achondrites. Indeed, FeO/MnO versus Fa content of olivine in acapulcoites and winonaites defines, remarkably, two trends (Fig. 5). As a matter of fact, the classified acapulcoites NWA 13041 and NWA 13839 plot on the winonaites trend, concordant with their relatively low Fa-Fs contents (Fs₄₋₅ Fa₃₋₅; Fs₄₋₅ Fs₄₋₅, respectively). The acapulcoite ALHA 81187, while also being more reduced than typical acapulcoites, plots at the origin of the winonaites trend, also coherent with its acapulcoite oxygen isotopic signature. Surprisingly, winonaites NWA 4937 plots with the tissuemouminites along the acapulcoite trend, which does not match its oxygen isotopic composition (Weisberg et al., 2008). This FeO/MnO versus Fa content allows us to discriminate winonaite from the other two groups of primitive achondrites, apart from a few cases, but not tissuemouminites from acapulcoites. As such, we decided to combine olivine composition, that is, FeO/MnO ratio and Fa content, with plagioclase composition (Or content) in a ternary diagram to enable the classification of primitive achondrites (Fig. 11). Remarkably, the three groups of primitive achondrites can be clearly separated from one another. Classified acapulcoites NWA 13839 and NWA 13041 fit within winonaite. As such, we suggest a reclassification of NWA 13839 and NWA 13041 as winonaite. In fact, low-Ca pyroxene (Fs₄₋₅; Fs₄₋₅, respectively), olivine (Fa₃₋₅, FeOMnO = 22.8; Fa₄₋₅, FeOMnO = 18, respectively), and plagioclase compositions (An₂₋₁₄Ab₈₋₁₀Or₀₋₁₂₋₆) (data from The Meteoritical Bulletin; Figs. 4–6 and 10) are evocative of winonaites rather than acapulcoite. Acapulcoite ALHA 81187, while being a highly reduced sample, lies consistently within the acapulcoite field. On the contrary, the classified winonaites NWA 4937 has oxygen isotopes in the range of winonaites but plots within the acapulcoite area (Fa₇, FeO/MnO = 12.8, Or₂₋₃) (Weisberg et al., 2008) (Figs. 4–6 and 10). As such, a thorough examination of this sample might be necessary since oxygen isotopic signature and chemistry are contradictory on its group affiliation. Few data are available for SuV 014 and NWA 4816, but investigation should be conducted to determine if they are indeed acapulcoites, winonaite, or tissuemouminites. The presence of relict chondrules in NWA 4816 might suggest a link to the tissuemouminite group, as well its FeO/MnO (12) and fayalite content (Fa₅₋₆) of olivine (Weisberg et al., 2008) (Fig. 5). However, no data on plagioclase exists, which would enable the possibility to test its position in our classification scheme. Finally, winonaites NWA 6448 plots along the tissuemouminite newly defined group and examination should also be conducted to ascertain its correct classification. As such, we suggest detailed chemical analysis of primitive achondrite silicates for classification. This could be useful since systematic chemical composition of primitive achondrites permits a classification within winonaite, acapulcoite–lodranite, or tissuemouminite group, without the absolute need of oxygen isotopes.

Another criterion that could be used is the CRE age. As established by Eugster and Lorenzetti (2005), all measured acapulcoites and lodranites have CRE ages...
ranging between 4 and 7 Ma. NWA 1058 yields a CRE age of 38 Ma, which has led these authors to suggest a reclassification of this meteorite from acapulcoite to winonaite, since CRE ages of Winona, Pontlyfni, and Mount Morris range from 20 to 90 Ma (Benedix et al., 1998). However, few literature data are available, and this criterion cannot be used to distinguish tissemuominites from winonaites.

Oxygen isotopes are often considered as the primary tool to distinguish between meteorite groups (Greenwood et al., 2012). Indeed, the three primitive achondrite groups, acapulcoite–lodranites, winonaites, and tissemuominites, are resolvable from one another on a three-oxygen isotope plot (Fig. 10). This criterion has been argued to be the sole criterion that can accurately discriminate winonaites and acapulcoites (Li et al., 2011). As highlighted by Greenwood et al. (2012, 2017), samples need to be pretreated for the removal of terrestrial weathering products. The absence of EATG treatment produces consequent shift in oxygen isotopes. In fact, when considering untreated samples, it is not possible to resolve tissemuominites from winonaites. Moreover, they are on the same fractionation line. In any case, oxygen isotopes are not always measured to classify a meteorite. Here, we have demonstrated that a combination of olivine compositions, that is, FeO/MnO ratio and Fa content, with plagioclase compositions (Or content) enables the classification of meteorites within the three groups of primitive achondrites, and silicate elemental composition might be sufficient to accurately classify them.

Fig. 11. Ternary diagram of mineral compositions in NWA 090, acapulcoites, and winonaites. The three endmembers represent Or content of plagioclase, Fa content of olivine, and FeO/MnO of olivine. Each value has been normalized to 100: for example, for the fayalite endmember, the equation is Fa/(Fa + Or + FeO/MnO) × 100. As such, correlation observed in previous plots is artificially enhanced to observe the diverse groups of primitive achondrite compositions. NWA 090 and the other relict chondrule-bearing winonaites do indeed plot apart from both acapulcoites and winonaites. Data are from references in Figs. 4–6. (Color figure can be viewed at wileyonlinelibrary.com.)
Petrogenesis of the Tissoumouinite Parent Body

Based on the petrology, mineralogy, and chemistry of the eight hereby classified tissoumouinites, they most probably sampled regolith of a primitive achondrite parent body, being related to type 5/6 chondrites (Benedix et al., 2003; Floss et al., 2008; Zeng et al., 2019). Farley et al. (2015) have already proposed that various winonaitic protoliths might have existed, due to the lack of Fe-Mg equilibrium between olivine and pyroxene in NWA 8614 (Fig. 4), arguing against a single metamorphic FeO reduction trend. This assertion is consistent with NWA 8614 being part of a newly defined primitive achondrite group. The lack of equilibrium between olivine and pyroxene Fe-Mg in fact true for all the eight tissoumouinites (Fig. 4). More importantly, a recent study of Mo isotopes of IAB iron-complex meteorites and primitive achondrites has also suggested a minimum of three separate parent bodies (Worsham et al., 2017). The IAB–iron–meteorite complex is characterized by the common presence of silicate inclusions and is classified in various groups (Wasson & Kalllemeyn, 2002): the main group (MG) and five chemical subgroups (sLL, sLM, sLH, sHL, and sHH), defined by abundances of Au and Ni. The IAB-MG has been linked to the winonaitic parent body, as a result of their similar petrography and common oxygen isotopic signature (Benedix et al., 2000; Clayton & Mayeda, 1996; Greenwood et al., 2012). Interestingly, while IAB MG iron meteorites and winonaites share a common Mo isotopic signature, supporting the proposed genetic link, the IAB sHH and IAB sHL have a distinct Mo isotopic composition. As such, they suggested that IAB MG and IAB sHH/sHL should have been derived from distinct parent bodies. Interestingly, NWA 725, now part of the tissoumouinite group, has a similar Mo isotopic composition to the IAB sHH/sHL, which could be seen as evidence for a common origin, although scarcity of information calls for caution in evaluating this potential genetic link (Worsham et al., 2017).

Nonetheless, metal–silicate Hf–W segregation model age estimated by Worsham et al. (2017) is 3.4 ± 0.7 Ma after CAIs for IAB MG while ranging from 0.3 to 1.6 Ma CAIs for IAB sHH and IAB sHL. As such, IAB sHH and IAB sHL protoliths were formed during 26Al lifetime, when protoliths were melted to form metallic cores and silicate mantles (Bizzarro et al., 2005; Kleine et al., 2009). If a genetic link between IAB-sHH or IAB-sHL and tissoumouinites were to be demonstrated, these primitive achondrites would have formed before winonaites. The metallographic cooling rate of 100 °C Ma⁻¹ estimated for NWA 090 and NWA 1463 (Scott et al., 2014a) is in the range of winonaites, although the number of data points is very limited (Benedix et al., 2005). As such, it means that either the thermal history of the tissoumouinite parent body was homogenous throughout or that these meteorites sampled a very narrow region of this parent body, as also suggested by their extremely similar compositions. However, while it has been argued that they could be paired (Irving & Rumble, 2006), their modal abundance variation prevents a pairing. As discussed previously, the oxygen fugacity of tissoumouinites is higher than the range observed in winonaites, although the number of data points is very limited (Benedix et al., 2005). They fit in the range calculated for acapulcoites, meaning that the oxidation state of their parent body is similar to that of the acapulcoite–lodranite parent body. It also means, as suggested previously by Righter and Drake (1996), that all primitive achondrites as well as HED and IAB irons have an fO2 approximately of 2 log units below the IW buffer and that the reservoir from which they formed...
should have been homogenous. That corroborates the similarity of silicate and opaque phase compositions between the primitive achondrite groups and the difficulty of distinguishing and classifying them.

CONCLUSION

NWA 090 and other relict chondrule-bearing winonaites, or chondritic winonaites (Dhofar 1222, NWA 725, NWA 1052, NWA 1054, NWA 1058, NWA 1463, NWA 8614), define a distinct group of primitive achondrites, the tissuemouminites that most likely came from a distinct parent body compared to those of winonaites and acapulcoites, although sampling similar nebular material as winonaites in terms of oxygen isotopic composition. Their modal abundances and chemical compositions lie between acapulcoites and winonaites, confounding with their traditional dichotomy. Nonetheless, by combining several chemical criteria, that is, fayalite content and FeO/MnO in olivines, as well as potassium content of plagioclases, these three groups of primitive achondrites can be clearly separated, highlighting the usefulness of mineral chemistry in primitive achondrite classification. In addition, oxygen isotopes, often cited as the only trustworthy criterion, confirm these eight samples to come from a distinct parent body than both winonaites and acapulcoites, hence also confirming the existence of this new tissuemouminite group. Tissuemouminites are pristine primitive achondrites, close to a petrographic type 5/6, as shown by the presence of relatively abundant relict chondrules and a closure temperature of metamorphism estimated at 820 °C. They probably sampled their parent body’s regolith. Other samples, like NWA 4972, SuY 014, NWA 4816, and Y 983237, might be related to this group, and more information is needed to classify them with certainty. It is important to note that not all chemical information is available for all the samples, which sets some limit to this classification scheme. Classification of primitive achondrites, and in general achondrites, needs to provide detailed mineral chemistry of samples that permit an appropriate association to known or new meteorite groups.

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Data Availability Statement—The data that support the findings of this study are available in the tables of this article.

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REFERENCES


