Olivine zoning and retrograde olivine-orthopyroxene-metal equilibration in H5 and H6 chondrites

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Abstract—Electron microprobe studies of several H5 and H6 chondrites reveal that olivine crystals exhibit systematic Fe-Mg zoning near olivine-metal interfaces. Olivine Fa concentrations decrease by up to 2 mol% toward zoned taenite + kamacite particles (formed after relatively small amounts of taenite undercooling) and increase by up to 2 mol% toward zoneless plessite particles (formed after ~200 °C of taenite undercooling).

The olivine zoning can be understood in terms of localized olivine-orthopyroxene-metal reactions during cooling from the peak metamorphic temperature. The silicate-metal reactions were influenced by solid-state metal phase transformations, and the two types of olivine zoning profiles resulted from variable amounts of taenite undercooling at temperatures <700 °C. The relevant silicate-metal reactions are modeled using chemical thermodynamics.

Systematic olivine Fe-Mg zoning adjacent to metal is an expected consequence of retrograde silicate-metal reactions, and the presence of such zoning provides strong evidence that the silicate and metallic minerals evolved in situ during cooling from the peak metamorphic temperature.

INTRODUCTION

Ordinary chondrites contain 35–50 wt% olivine, ~30 wt% pyroxene, and 5–20 wt% Fe-Ni metal. In 1916, G.T. Prior observed that important relationships exist between the silicate and metallic minerals in chondrites: “the less the amount of nickel-iron in chondritic stones, the richer it is in nickel and the richer in iron are the magnesium silicates” (Prior 1916). Prior’s Rules, which can be understood in terms of a redox-controlled distribution of Fe among olivine, pyroxene, and metal, illustrate that the silicate and metallic minerals interacted in order to approach or attain chemical equilibrium.

Most studies of olivine-orthopyroxene-metal equilibration in chondrites assume that the olivine and pyroxene compositions were established at the peak metamorphic temperature where high diffusion rates promote equilibration (Larimer 1968; McSween and Labotka 1993). According to Keil and Fredrickson (1964), Dodd (1969), Van Schmus and Wood (1967), Olsen and Bunch (1984), and Miyamoto et al. (1986), olivine crystals in types 4–6 ordinary chondrites are thought to be compositionally homogeneous. This result suggests that 1) olivine-orthopyroxene-metal equilibration was complete at the peak metamorphic temperature and 2) retrograde reactions among these minerals were insignificant. However, previous researchers did not investigate the possibility of systematic rim-to-core olivine Fe-Mg zoning that may have formed during slow cooling.

Several workers have attempted to quantify aspects of olivine-orthopyroxene-metal equilibria in chondrites. Mueller (1963) and Mueller and Olsen (1967) derived the mass balance equations that describe how olivine, orthopyroxene, and metal compositions and modes vary during isothermal redox reactions. Other workers used chemical thermodynamics to investigate the temperatures, pressures, and oxygen fugacities that prevailed during chondrite formation. Craig (1964) attempted to calculate chondrite equilibration temperatures on the basis of Fe-Mg partitioning between olivine and orthopyroxene. Mueller (1964), Larimer (1968), and Williams (1971) calculated chondrite equilibration temperatures and oxygen fugacities on the basis of the Fe distribution among olivine, orthopyroxene, and metal. Finally, McSween and Labotka (1993) calculated chondrite oxygen fugacities from olivine, orthopyroxene, and
metal compositions. None of these studies, however, considered the effect of olivine-orthopyroxene-metal equilibration during cooling from the peak metamorphic temperature.

Van Schmus and Koffman (1967) recognized that chondrite orthopyroxene-diopside Fe-Mg exchange temperatures might reflect closure temperatures during cooling rather than peak metamorphic temperatures. More dramatically, chemical zoning in taenite ($\gamma$, face-centered-cubic) and kamacite ($\alpha$, body-centered-cubic) are the result of taenite-kamacite equilibration during cooling (Wood 1967). It seems reasonable to expect, by analogy, that some degree of retrograde olivine-orthopyroxene-metal equilibration occurred.

The present study investigates the importance of olivine-orthopyroxene-metal equilibration in several H5 and H6 chondrites during cooling from the peak metamorphic temperature. These chondrites experienced metamorphic temperatures and time scales that were sufficient to homogenize individual olivine crystals (Miyamoto et al. 1986). Therefore, any systematic olivine zoning observed in these samples must have developed during cooling from the peak temperature. This study focuses on H-group chondrites because they are metal-rich and consequently contain a high density of silicate-metal interfaces where silicate zoning (resulting from silicate-metal reaction) might occur. Olivine and orthopyroxene crystals are analyzed near olivine-orthopyroxene, olivine-metal, and orthopyroxene-metal interfaces in order to search for silicate zoning. Additionally, a chemical thermodynamic model is developed to investigate the effect of taenite transformations (i.e., changing taenite compositions) on the equilibrium olivine and orthopyroxene compositions during cooling. This study represents the first attempt to investigate the importance of retrograde olivine-orthopyroxene-metal equilibration in chondrites.

**BACKGROUND:**

**TAE-NITE TRANSFORMATIONS DURING COOLING**

In this paper, we demonstrate that silicate-metal reactions occurred in ordinary chondrites during cooling from the peak metamorphic temperature, and that these reactions were closely tied to the taenite transformations during cooling. Therefore it is important to review what is known about taenite transformations in ordinary chondrites.

There are two mechanisms by which taenite, which is stable at the peak metamorphic temperature, can transform to a more stable taenite + kamacite mixture upon cooling. In mechanism 1, taenite particles that are polycrystalline at high temperatures exsolve kamacite directly by the taenite $\rightarrow$ taenite + kamacite transformation (Reisener and Goldstein 2003a, 2003b). The two fundamentally different reaction paths for taenite decomposition are summarized below.

**Polycrystalline Taenite: The Equilibrium Taenite Path**

Consider the behavior of a parent taenite particle that is polycrystalline and contains 10 wt% Ni, a typical composition for metal particles in H chondrites (Fig. 1a). The taenite particle will cool below the taenite/(taenite + kamacite) solvus at ~700 °C (Fig. 1a). Kamacite will precipitate heterogeneously along internal taenite grain boundaries after relatively small amounts of undercooling below the solvus (Fig. 1b, left) (Reisener and Goldstein 2003a, 2003b). The grain boundary kamacite precipitate will grow by consuming one of the adjacent taenite grains (Reisener and Goldstein 2003b).

Initially, taenite and kamacite volume diffusion rates are sufficiently high that total taenite-kamacite equilibrium is maintained, i.e., taenite and kamacite remain homogeneous with compositions defined by the Fe-Ni phase diagram equilibrium tie-lines. However, volume diffusion rates decrease exponentially during cooling and mass transfer eventually becomes too sluggish to maintain total equilibrium. The classic M-shaped Ni profile develops in taenite and an Agrell Ni depletion develops in kamacite (Fig. 2b). Local equilibrium is maintained at the taenite/kamacite interface throughout the cooling process, even though taenite and kamacite do not maintain total equilibrium (Fig. 2b). Taenite-kamacite equilibrium is also maintained locally at taenite/silicate and kamacite/silicate interfaces (Fig. 2b) because Fe and Ni atoms diffuse across taenite and kamacite surfaces much faster than they diffuse through the taenite and kamacite lattices (Willis and Goldstein 1983).

The equilibrium taenite composition, $\gamma_{\text{E}}$, at taenite/kamacite and taenite/silicate interfaces follows the “equilibrium taenite path” during cooling (Fig. 1a, line A-E). The final metal microstructure formed from a polycrystalline taenite particle is a “zoned taenite + kamacite particle” such as the one shown on the left in Fig. 2a. At temperatures <500 °C, the taenite interior further decomposes by a variety of complex solid-state transformations (Yang et al. 1997).

Zoned taenite and kamacite frequently appear isolated from each other when viewed on the plane of sectioning. Sequential polishing, however, has shown that zoned taenite and kamacite are always connected to each other in the third dimension (Willis and Goldstein 1983).

**Monocrystalline Taenite: The Metastable Taenite Path**

Many parent taenite particles in type 5 and 6 ordinary chondrites were monocrystalline at the beginning of cooling due to the effect of taenite grain growth at high temperatures.
A taenite particle that is monocrystalline will experience large amounts of undercooling because taenite/taenite boundaries, which serve as kamacite nucleation sites, are not present (Fig. 1b, right) (Reisener and Goldstein 2003a, 2003b). The undercooled taenite particle remains homogeneous and metastable (Fe-supersaturated at 10 wt% Ni) as it cools through the taenite + kamacite field (the metastable taenite composition, $\gamma_M$, is indicated by a vertical broken line in Fig. 1a). At ~500 °C (after ~200 °C of undercooling) the homogeneous taenite particle will cool below the martensite-start temperature (Fig. 1a, dashed line) and will transform to martensite ($\alpha_2$, strained bcc) by the diffusionless taenite $\rightarrow$ martensite transformation (Fig. 1b). The homogeneous martensite, itself metastable, will subsequently decompose by the martensite $\rightarrow$ taenite + kamacite reaction during continued cooling (Fig. 1b). The final microstructure of the undercooled taenite particle is a two-phase, micron-scale mixture of taenite and kamacite known as plessite (Fig. 2a, right). These metal particles are named “zoneless plessite particles” because they lack systematic rim-to-core Ni zoning (Fig. 2b) (Sears and Axon 1975; Reisener and Goldstein 2003b).

In many well-equilibrated (type 5 and 6) ordinary chondrites, zoned taenite + kamacite particles and zoneless plessite particles coexist within microscopic distances of each other (Fig. 2a). Both types of metal particles had identical starting compositions and experienced the same cooling history, but the different parent taenite structures (polycrystalline versus monocrystalline) initiated different types of phase transformations resulting in different final metal microstructures (zoned taenite + kamacite particles versus zoneless plessite particles).

**PROCEDURE**

Polished thin sections of four low-shock (S1–S2), type 5–6 H-group chondrites were selected for detailed studies of olivine and orthopyroxene zoning: Estacado (H6), Guarena (H6), Kernouve (H6), and Nuevo Mercurio (H5). The four chondrites have well-recrystallized textures and coherent metallographic cooling rates indicating that they did not experience brecciation after cooling from the peak metamorphic temperature. Each chondrite contains two types of Fe-Ni metal particles: 1) zoned taenite + kamacite particles that formed after small amounts of taenite undercooling, and 2) zoneless plessite particles that formed after monocrystalline taenite undercooled ~200 °C below the taenite/(taenite + kamacite) solvus, as described by Reisener and Goldstein (2003a, 2003b).

The thin sections were polished with diamond paste from 15 μm down to 0.06 μm, and lightly etched with Nital (2 vol% HNO$_3$, balance ethyl alcohol) in order to reveal the fine-scale
A Cameca SX50 electron microprobe, operated at an accelerating voltage of 15 kV and a beam current of 20 nA, was used to analyze Si, Fe, Mg, Mn, Ca, Ni, Cr, and Al in olivine and orthopyroxene crystals using a 40-second counting time for each element. These conditions gave a one standard deviation uncertainty of <0.1 wt% for SiO₂, FeO, and MgO. The microprobe standards were synthetic fayalite (for Fe and Si), Shallowater enstatite (Mg), Broken Hill rhodonite (Mn), Kiglapait labradorite (Ca and Al), synthetic NiO (Ni), and Stillwater chromite (Cr). Quantitative analyses were obtained using the PAP ϕ(ρz) routine.

Silicate-metal interfaces were studied using both reflected-light and transmitted-light microscopy. The positions of silicate-metal interfaces, as viewed in reflected light, often shift toward the silicate when viewed in transmitted light. This optical effect indicates that an opaque mineral such as metal or troilite underlies silicate below the plane of sectioning. Electron microprobe analyses were collected only from olivine and orthopyroxene crystals that do not overlie opaque minerals, minimizing the possibility that Fe X-rays were generated by subsurface metal or troilite.

Olivine and orthopyroxene crystals were analyzed using line scans (1 μm step intervals) across olivine-orthopyroxene interfaces, olivine-metal interfaces, and orthopyroxene-metal interfaces. Additionally, point analyses were obtained at olivine cores and at olivine rims (including olivine-metal interfaces) in order to rapidly characterize compositional zoning patterns in olivine.

**RESULTS**

**Silicate Crystals Isolated from Fe-Ni Metal**

Individual olivine and orthopyroxene crystals that are isolated from metal have core and rim compositions that are identical within analytical uncertainties. However, silicate compositions vary by up to ~1 mol% Fa or Fs from one crystal to another in each chondrite. Figure 3 shows fayalite measurements in olivine crystals that are isolated from metal in the four H chondrites (the analyses are represented by circles). Electron microprobe line scans collected across olivine-orthopyroxene interfaces in Estacado (H6) and Kernouve (H6) are shown in Fig. 4. No systematic FeO/(FeO + MgO) zoning was observed at the interfaces. In each chondrite, olivine Fa concentrations are higher than orthopyroxene Fs concentrations (Fig. 4). The measured distribution of Fe and Mg between olivine and orthopyroxene is consistent with the work of Keil et al. (1964) and Larimer (1968) and also is consistent with experimental FeO-MgO-SiO₂ system equilibria (Larimer 1968).

In addition to the major elements, olivine contains 0.4–0.6 wt% MnO and 0.01–0.05 wt% Ca; orthopyroxene contains 0.5–0.6 wt% MnO and ~0.6 wt% CaO.
Silicate Crystals Abutting Zoned Taenite + Kamacite Particles

Electron microprobe line scans, traversing olivine crystals toward zoned taenite + kamacite particles, are reported as squares in Fig. 5. Olivine Fa concentrations decrease systematically by ~2 mol% as a zoned taenite + kamacite particle is approached. The zoning extends 30–40 μm from the metal, and is independent of whether the olivine abuts zoned taenite or the coexisting kamacite. An olivine Fa concentration decrease is also shown in Fig. 3, in which analyses collected within 10 μm of a zoned plessite particle are reported as triangles. MnO concentrations are homogeneous along the Fe-Mg zoning profiles. Orthopyroxene Fs concentrations appear to remain constant as zoneless plessite particles are approached.

Silicate Crystals Abutting Zoneless Plessite Particles

Electron microprobe line scans, traversing olivine crystals toward zoneless plessite particles, are reported as triangles in Fig. 5. Olivine Fa concentrations increase systematically by ~2 mol% as a zoneless plessite particle is approached; the zoning extends 30–40 μm from the metal. An olivine Fa concentration increase is also shown in Fig. 3, in which analyses collected within 10 μm of a zoneless plessite particle are reported as triangles. MnO concentrations are homogeneous along the Fe-Mg zoning profiles. Orthopyroxene Fs concentrations appear to remain constant as zoneless plessite particles are approached.

Olivine Stoichiometry

Olivine and orthopyroxene crystals in all three textural settings are stoichiometric with respect to SiO₂, FeO, MgO, and MnO. Cation totals in olivine are 3.00 ± 0.05, even along the Fe-Mg zoning profiles.

DISCUSSION

Is the Olivine Zoning Real?

It is important to evaluate whether the observed olivine Fa variations could be artifacts of the electron probe...
microanalysis. When an olivine crystal is analyzed near an olivine-metal interface, electron-sample interactions and X-ray fluorescence can potentially excite Fe atoms in the metal resulting in apparent Fa concentrations that are erroneously high. It is conceivable that electron-sample interactions and/or X-ray fluorescence could explain the elevated olivine Fa concentrations measured near zoneless plessite particles (neither effect, however, can explain the low olivine Fa concentrations measured near zoned taenite + kamacite particles). The potential effects of electron-sample interactions and X-ray fluorescence are evaluated below.

**Electron-Sample Interactions: Primary X-ray Generation Volume**

The spatial resolution of the electron microprobe is determined by the volume of primary X-ray generation. When the electron beam is situated in olivine near an olivine-metal interface, the volume of X-ray generation may potentially overlap metal resulting in Fa concentrations that are erroneously high. It is conceivable that electron-sample interactions and/or X-ray fluorescence could explain the elevated olivine Fa concentrations measured near zoneless plessite particles (neither effect, however, can explain the low olivine Fa concentrations measured near zoned taenite + kamacite particles). The potential effects of electron-sample interactions and X-ray fluorescence are evaluated below.

**Characteristic X-ray Fluorescence**

Characteristic X-rays traveling though a sample can ionize atoms resulting in the fluorescence of other X-rays. The volume of X-ray fluorescence is much larger than the volume of electron interaction because X-rays penetrate solids more efficiently than electrons. However, the characteristic X-rays generated by the major elements in olivine (Si, Fe, Mg, and Mn) have insufficient energy to excite Fe Kα radiation (7.11 keV) in the metal. Therefore, the
elevated olivine Fa concentrations measured near zoneless plessite particles cannot be explained by characteristic X-ray fluorescence.

Continuum X-ray Fluorescence

The background continuum radiation produced during electron probe microanalysis provides X-rays with energies ranging from 0 eV up to the electron beam energy. For the 15 keV beam used in this study, the continuum radiation is capable of exciting Fe X-rays (K edge = 7.11 keV) and Ni X-rays (K edge = 8.33 keV) in metal. However, less than 0.05 wt% Ni is measured during olivine analyses (even near olivine-metal interfaces). This result suggests, by analogy, that continuum fluorescence of metallic Fe does not make a significant contribution to the measured Fe intensities.

The importance of continuum fluorescence was also investigated experimentally. A piece of San Carlos olivine (Fo90) and a piece of synthetic Fe-Ni alloy (Fe10Ni) were investigated experimentally. A piece of San Carlos olivine (Fo90) and a piece of synthetic Fe-Ni alloy (Fe10Ni) were clamped together along flat surfaces. An electron microprobe line scan (1 μm step interval) was used to measure the composition of olivine as the olivine-metal interface was approached. Olivine Fa concentrations were constant until the electron beam was within ~3 μm from the olivine-metal interface. Within 3 μm of the metal, olivine Fa concentrations increase by several mol% because the electron interaction volume overlaps the metal. This experiment confirms that X-ray fluorescence of metallic Fe cannot explain the elevated olivine Fa concentrations that extend up to 30 μm from zoneless plessite particles.

Olivine Stoichiometry

The measured stoichiometry of the olivine provides additional evidence that the Fe-Mg zoning is real. If the elevated olivine Fa concentrations measured near zoneless plessite particles were caused by electron-sample interactions or X-ray fluorescence, cation totals (Si + Fe + Mg + Mn) would exceed 3.0 due to Fe X-ray contributions from metal. Cation totals along the olivine Fe-Mg zoning profiles are 3.00 ± 0.05, indicating that negligible amounts of Fe X-rays were generated in the adjacent metal.

In summary, neither electron excitation nor X-ray fluorescence can explain the elevated olivine Fa concentrations measured near zoneless plessite particles. Moreover, neither of these effects can explain the low olivine Fa concentrations measured near zoned taenite + kamacite particles. The authors conclude that the systematic olivine Fe-Mg zoning measured near Fe-Ni metal particles is real.

Origin of Olivine Zoning by Solid-State Reaction

Two fundamentally different mechanisms can cause systematic Fe-Mg zoning in olivine: 1) nonequilibrium crystallization of olivine from a melt (igneous zoning) and 2) solid-state equilibration between olivine and other Fe-bearing minerals (metamorphic zoning). Igneous zoning in olivine is characterized by Fa concentrations that increase systematically from crystal core to crystal rim. Olivine crystals in type 3 ordinary chondrites display igneous zoning due to rapid chondrule solidification rates, but in type 5 and 6 chondrites such zoning would have been completely homogenized during metamorphism by solid-state diffusion (Miyamoto et al. 1986).

Olivine crystals in the H5 and H6 chondrites exhibit Fe-Mg zoning only where they abut Fe-Ni metal, suggesting an origin by silicate-metal reaction in the solid state. Moreover, the observation that olivine zoning profiles adjacent to zoned taenite + kamacite particles differ from olivine zoning profiles adjacent to zoneless plessite particles suggests that the zoning is linked to the metal transformation path, i.e., the olivine zoning developed during cooling from the peak metamorphic temperature.

The meteorite literature contains few reports of olivine zoning resulting from solid-state reactions between silicate and metal. In pallasites, consisting mainly of olivine crystals surrounded by metal, olivine exhibits slight Fe-Mg zoning from crystal core to rim (Miyamoto 1997). In Esquel and Yamato 8451 (a pyroxene-bearing pallasite) olivine Fa decreases by ~1 mol% from crystal core to rim. In Imilac, some olivine crystals show Fa-enrichment toward the rim while other olivine crystals show the opposite zoning pattern. Miyamoto (1997) did not discuss the origin of the pallasite olivine zoning profiles in detail, but he did entertain the possibility that the zoning formed during subsolidus cooling.

In another relevant study, Dohmen et al. (1998) addressed the formation of olivine FeO-enrichment halos around Fe-Ni metal inclusions in Allende (CV3). Experimental studies of the system olivine + metal led to the unexpected conclusion that Mg-rich olivine can react with metallic Fe, even in the absence of other Mg-bearing minerals that provide a sink for Mg transported away from the olivine-metal interface. The olivine-metal reaction is possible because Mg is lost from the interface via a gaseous phase.

As we shall see, olivine zoning may develop in chondrites by silicate-metal reaction without loss of Mg from the system. The experimental study by Dohmen et al. (1998) may be relevant, however, in explaining intergranular transport of chemical components within chondrites via a dry vapor phase composed of Fe, Mg, SiO, O2, O, etc.

Silicate-Metal Equilibria: Thermodynamic Analysis

In this section, chemical thermodynamics will be used to investigate the compositional evolution of olivine + orthopyroxene + metal assemblages in ordinary chondrites during cooling from the peak metamorphic temperature. A new analysis of silicate-metal equilibration incorporates two
major effects that were neglected during previous studies: 1) taenite composition changes during cooling and 2) the tendency for many taenite particles to undercool within the taenite + kamacite field (i.e., the occurrence of equilibrium taenite and metastable taenite within the same chondrite).

Reactions

Chondritic olivine and orthopyroxene are essentially binary solutions of the Fe-Mg endmembers, fayalite-forsterite and ferrosilite-enstatite respectively. These silicates coexist with the Fe-Ni metal phases taenite and kamacite. Taenite is the stable metallic phase at temperatures above the taenite/(taenite + kamacite) solvus, while taenite and kamacite are in equilibrium at temperatures below the solvus. In addition, the oxygen fugacity \( (f_O^2) \) plays an important role during silicate-metal equilibration.

Three reactions completely describe the equilibria among olivine, orthopyroxene, taenite, and kamacite:

\[
\begin{align*}
\text{Olivine} + \text{Orthopyroxene} + \text{Gas} & \rightarrow \text{Taenite} + \text{Taenite} + \text{Kamacite} \\
\text{Olivine} & \rightarrow \text{Orthopyroxene} + \text{Ni} + \text{Fe} \\
\text{Taenite} + \text{Kamacite} & \rightarrow \text{Taenite} + \text{Kamacite}
\end{align*}
\]

An increase in oxygen fugacity will drive Reaction 1 to the right producing olivine at the expense of orthopyroxene and metallic Fe (oxidation). Olivine becomes FeO-enriched as iron is transferred from metal to silicate, and the residual metal becomes Fe-depleted (relatively Ni-enriched). Simultaneously, equilibrium between olivine and orthopyroxene (Reaction 2) will result in Fe-enrichment of orthopyroxene. Mass balance is maintained during oxidation via the partitioning of SiO\(_2\) between orthopyroxene and olivine.

A decrease in oxygen fugacity (reduction) will drive Reaction 1 to the left producing orthopyroxene and metallic Fe at the expense of olivine; the olivine and orthopyroxene will become MgO-enriched and the metal will become Fe-enriched (relatively Ni-depleted). Partitioning of SiO\(_2\) between olivine and orthopyroxene allows mass balance to be maintained during the reduction process.

It is important to note that analogous redox reactions can be driven by variations in taenite composition. At a given temperature and oxygen fugacity, an increase in taenite Fe concentration will drive Reaction 1 to the left (oxidation) while a decrease in taenite Fe concentration will drive the reaction to the right (reduction).

It should also be noted that Reaction 1 is written in terms of taenite rather than kamacite. This treatment is clearly valid at temperatures above the taenite/(taenite + kamacite) solvus because taenite is the only stable metallic phase. This approach is also justified at temperatures below the taenite/(taenite + kamacite) solvus because the reaction that describes taenite-kamacite equilibrium (Reaction 3) is independent of the reactions describing olivine-orthopyroxene-taenite equilibria (Reactions 1 and 2). At temperatures <700 °C, taenite-kamacite equilibrium forces the taenite composition to follow the taenite/(taenite + kamacite) solvus of the Fe-Ni phase diagram. Therefore, the equilibrium taenite and kamacite compositions can be obtained directly from the Fe-Ni phase diagram at any temperature.

Formalism

Olivine and orthopyroxene in equilibrated ordinary chondrites can be regarded as binary solutions of the Fe-Mg endmembers. In reality, olivine contains 0.4–0.6 wt% MnO and less than 0.1 wt% CaO and orthopyroxene contains 0.4–0.6 wt% MnO and 0.5–1 wt% CaO. The low concentrations of MnO and CaO, however, should have little influence on Reactions 1 or 2. Indeed, there is close experimental agreement between Fe-Mg partitioning studies in which olivine and orthopyroxene are Ca-free and Fe-Mg partitioning studies in which olivine and orthopyroxene contain up to 1.0 wt% Ca (Larimer 1968).

Using the ionic solution model (e.g., Ganguly and Saxena 1987) the activity, \( a \), of an end-member component in a phase \( m \) with a general formula \( (i, j, \ldots)_mQ \) can be expressed as:

\[
a^m_{iQ} = (\chi_i \cdot \lambda^m_i)^{\alpha} \tag{2}
\]

where \( \chi \) is the component’s molecular concentration, \( \lambda \) is the component’s activity coefficient, and \( \alpha \) represents the number of different crystallographic mixing sites in the mineral. The value of \( \alpha \) for taenite and orthopyroxene is 1; for olivine, the value is 2. The activity coefficients of fayalite, forsterite, ferrosilite, and enstatite are approximated by the symmetrical solution model (Ganguly and Saxena 1987):

\[
\lambda_i^m = \exp \left[ \frac{(1 - \chi_i^m)^2 W_{Fe-Mg}^i}{\alpha R T} \right] \tag{3}
\]

where \( T \) is the temperature, \( R \) is the gas constant, and \( W_{Fe-Mg}^i \) is the Fe-Mg interaction parameter for olivine or orthopyroxene (Berman and Aranovich 1996). The activity coefficient of Fe in taenite is approximated by the subregular asymmetrical solution model:

\[
\lambda_{Fe} = \exp \left[ x_i^2 W_{FeNi} + 2(x_i W_{FeNi} - W_{NiFe})x_i \right] \tag{4}
\]

where \( W_{FeNi} \) and \( W_{NiFe} \) are the Fe-Ni interaction parameters (Hsieh et al. 1987).
The equilibrium constants for Reactions 1 and 2 are:

\[
K_1 = \frac{\left( \frac{O_2}{V} \right)^{1/2} \left( \chi_{\text{Fe}} \cdot \lambda_{\text{Fe}} \right)}{\left( \chi_{\text{Fe}} \cdot \lambda_{\text{Fe}} \right)^2}
\]

and

\[
K_2 = \frac{\left( \chi_{\text{Mg}} \cdot \lambda_{\text{Mg}} \right)}{\left( \chi_{\text{Fe}} \cdot \lambda_{\text{Fe}} \right)^2}
\]

Reactions 1 and 2 are related to their respective equilibrium constants by the law of mass action:

\[
\Delta G_{\text{rxn}} = -RT \ln K
\]

where \(\Delta G\) is the Gibbs free energy change of the reaction.

For each reaction, the Gibbs free energy change can be calculated from the Gibbs free energies of formation of the individual mineral components:

\[
\Delta G_{\text{rxn}, T} = \sum v_i \Delta G_{f, T(\text{products})} - \sum v_i \Delta G_{f, T(\text{reactants})}
\]

where \(v_i\) are the stoichiometry coefficients. The standard Gibbs free energy of formation of each mineral component is given by:

\[
\Delta G_{f, T} = \Delta H_{f, T} - T \cdot S_{T} + \int \frac{C_p dT}{T} - T \cdot \int \frac{C_p dT}{298.15}
\]

where \(\Delta H_{f, T}\) and \(S_{T}\) represent the standard enthalpy of formation from elements and standard entropy (respectively) of a mineral component at 298 K and 1 bar, and \(C_p\) represents the heat capacity of the mineral component. No pressure correction is included in Equation 10 because pressure has little effect on Reactions 1 or 2 at lithostatic pressures below several kilobars (Larimer 1968). The standard thermodynamic properties of the silicate minerals are from Berman and Aranovich (1996), and those of oxygen and metallic Fe are from Chase (1998).

**Calculations**

The compositions of olivine and orthopyroxene in equilibrium with Fe-Ni metal during cooling can be calculated using chemical thermodynamics. Such calculations will be used to trace the silicate compositions from an assumed peak metamorphic temperature of 900 °C to a final temperature of 500 °C (a temperature below typical silicate closure temperatures). Three assumptions are made: 1) olivine, orthopyroxene, and taenite are fully equilibrated (hence homogeneous) at the onset of cooling; 2) all parent taenite particles have bulk compositions of 10 wt% Ni, typical of metal particles in H chondrites (Fig. 1a); and 3) the oxygen fugacity is defined by olivine-orthopyroxene-metal equilibria and is homogeneous throughout the chondrite at all times.

**Olivine and Orthopyroxene Compositions at the Peak Metamorphic Temperature (900 °C)**

Geothermometry shows that type 6 ordinary chondrites reached a peak metamorphic temperature of 750–950 °C (Dodd 1981). The compositions of olivine and orthopyroxene at an assumed peak temperature of 900 °C can be estimated from measured compositions and thermodynamic data. Figure 6 shows the equilibrium Fe-Mg distribution between olivine and orthopyroxene, calculated by simultaneously solving Equations 3, 6, 8, 9, and 10. The closure temperature for olivine-orthopyroxene Fe-Mg exchange in Estacado (H6) and Kernoué (H6) can be estimated from the compositions of coexisting crystals of olivine and orthopyroxene shown in Fig. 4 (these crystals are isolated from metal and hence homogeneous within microprobe counting statistics). In both chondrites, the olivine composition is ~18.5 Fa and the orthopyroxene composition is ~17.0 Fs, suggesting a final equilibration temperature of ~600 °C (Fig. 6). The olivine and orthopyroxene compositions at 900 °C can be found by extrapolation of the measured compositions (Fig. 4) to the 900 °C isotherm in Fig. 6 (mass conservation constrains the olivine and orthopyroxene compositions to a line of slope = −1). The extrapolation yields an olivine composition of ~19.0 Fa and an orthopyroxene composition of ~16.5 Fs at 900 °C.

**The “Chondritic fO₂-T Path”**

Before calculating equilibrium olivine and orthopyroxene compositions as a function of temperature, it is necessary to derive the relationship among chondrite oxygen fugacity, mineral compositions, and temperature. By rearranging Equation 5, the oxygen fugacity of an olivine + orthopyroxene + taenite assemblage at a given temperature can be expressed as:

\[
\log fO_2 = \frac{1}{2} \left( \log K_1 - \log \left( \frac{O_2}{\text{Taenite} \cdot \text{Fe}} \right) \right)
\]

**fO₂-T Relations between 900 and 700 °C:** As olivine + orthopyroxene + taenite assemblages cool from 900 to 700 °C, each taenite particle contains 10 wt% Ni (Fig. 1a) and remains chemically homogeneous throughout cooling. However, the compositions of both olivine and orthopyroxene...


**Fig. 6. Compositions of coexisting olivine and orthopyroxene between 900 and 500 °C due to Fe-Mg exchange (Reaction 2).** The open diamond represents the compositions of coexisting olivine and orthopyroxene measured in this study. The dots trace the compositional evolution of the olivine + orthopyroxene pair between 900 and 500 °C. Mass conservation during Fe-Mg exchange constrains the silicate compositions to a slope of 900 and 500 °C. The dashed line represents the condition that olivine Fa = orthopyroxene Fs.

**fO₂-T Relations between 700 and 500 °C:** At temperatures <700 °C, taenite particles that are polycrystalline will precipitate kamacite at taenite/taenite boundaries while taenite particles that are monocrystalline will undercool without forming kamacite. The two different metal reaction paths produce two populations of taenite compositions: polycrystalline taenite will have the equilibrium taenite composition, γₑ, defined by the Fe-Ni phase diagram (Fig. 1a) whereas monocrystalline taenite will retain the metastable composition (γₑ) of ~10 wt% Ni throughout cooling. Clearly, whole-rock silicate-metal equilibrium is not possible in the presence of two populations of taenite compositions (γₑ and γₑ) (Fig. 1a). Olivine, orthopyroxene, and taenite can establish local equilibria, however, either by adjustments in the local oxygen fugacity or by exchange of Fe between metal and silicates.

Assuming that the oxygen fugacity is defined by olivine-orthopyroxene-taenite equilibria, the two populations of taenite compositions will produce two fO₂-T paths at temperatures <700 °C. The fO₂-T path defined by an ol + opx + γₑ assemblage can be calculated using the olivine and orthopyroxene compositions in Fig. 6 (dots) and taenite compositions along the equilibrium taenite path in Fig. 1a. Similarly, the fO₂-T path defined by an ol + opx + γₘ assemblage can be calculated using the silicate compositions in Fig. 6 (dots) and a constant taenite composition of 10 wt% Ni (equilibrium is assumed to exist between metastable taenite, olivine, and orthopyroxene). Calculations for the 700 to 500 °C temperature interval show that the oxygen fugacity defined by an ol + opx + γₑ assemblage is up to 0.1 log unit higher than the oxygen fugacity defined by an ol + opx + γₘ assemblage. The two fO₂-T paths diverge during cooling because the equilibrium taenite and metastable taenite compositions diverge as temperature decreases (Fig. 1a).

The existence of two different fO₂-T paths within the same chondrite is inherently unstable and some degree of oxygen fugacity homogenization will occur. Oxygen fugacity homogenization will be promoted by the slow cooling rates of ordinary chondrites (typically on the order of 1–100 °C/MY; Wood 1967; Pellis and Storzer 1981; Willis and Goldstein 1983) and high chondrite permeabilities (Sugiura et al. 1986). A “chondritic fO₂-T path,” representing the whole-rock oxygen fugacity after homogenization, can be approximated by averaging the fO₂-T path defined by an ol + opx + γₑ assemblage and the fO₂-T path defined by an ol + opx + γₘ assemblage. The resulting fO₂-T path is shown as a solid line in Fig. 7a. The ferrosilite-iron-fayalite (FIF) buffer and the intrinsic oxygen fugacities measured in H chondrites (Brett and Sato 1984) are plotted in the same figure for reference. The chondritic fO₂-T path calculated in the present study is lower than the intrinsic oxygen fugacity path, but the calculated fO₂-T path is consistent with H chondrite oxygen fugacities calculated by McSween and Labotka (1993) on the basis of mineral compositions. The intrinsic oxygen fugacities measured by Brett and Sato (1984) lie above the FIF buffer at temperatures <750 °C and consequently are incompatible with ordinary chondrite mineralogy. As explained below, we propose that it is the chondritic (homogenized) oxygen fugacity that drives redox reactions at silicate-metal interfaces.

**Olivine and Orthopyroxene Compositions During Cooling**

As an olivine + orthopyroxene + taenite assemblage cools, the olivine and orthopyroxene compositions will vary due to the combined effects of redox and Fe-Mg exchange equilibria (Reactions 1 and 2). The compositions of olivine and orthopyroxene in equilibrium with metal can be calculated by simultaneously solving Equations 3–10 for the 900–500 °C temperature interval. Specifically, the silicate compositions coexisting with equilibrium taenite (γₑ) can be calculated using taenite compositions along the equilibrium taenite path (Fig. 1a) and oxygen fugacities along the chondritic fO₂-T path shown in Fig. 7. Similarly, the silicate compositions coexisting with metastable taenite (γₘ) can be...
calculated using a constant taenite composition of 10 wt% Ni and oxygen fugacities along the chondritic $f_{O_2}$-$T$ path (Fig. 7). The calculated silicate compositions in ol + opx + $\gamma_E$ assemblages are reported as solid lines in Fig. 8. The calculated silicate compositions in ol + opx + $\gamma_M$ assemblages are reported as broken lines in the same figure.

**Cooling from 900 to 700 °C:** Olivine and orthopyroxene will have initial (900 °C) compositions of 19.0 Fa and 16.5 Fs respectively. As olivine + orthopyroxene + taenite assemblages cool to the taenite/(taenite + kamacite) solvus at 700 °C, all taenite particles remain homogeneous at 10 wt% Ni (Fig. 1a). The oxygen fugacity (defined by olivine + orthopyroxene + taenite equilibria) is constant everywhere within the chondrite. During cooling, olivine Fa concentrations decrease slightly and orthopyroxene Fs concentrations increase slightly (Fig. 8) due to Fe-Mg exchange (Reaction 2).

**Cooling from 700 to 500 °C:** At 700 °C, the taenite particles cool into the two-phase taenite + kamacite field (Fig. 1a). If a parent taenite particle is polycrystalline, kamacite will precipitate at the taenite/taenite grain boundaries by the reaction taenite $\rightarrow$ taenite + kamacite (Fig. 1b). The equilibrium Ni concentration in taenite ($\gamma_E$) at the taenite/silicate interface increases along the equilibrium taenite path throughout cooling (Fig. 1a).

If a parent taenite particle is monocrystalline, it will undercool below the taenite/taenite + kamacite solvus without forming kamacite. The composition of the metastable taenite ($\gamma_M$) at the taenite/silicate interface will remain constant at ~10 wt% Ni throughout cooling (Fig. 1b) (Reisener and Goldstein 2003b). The compositions of $\gamma_E$ and $\gamma_M$ at taenite/silicate interfaces will diverge along the $\gamma_E$ and $\gamma_M$ paths throughout cooling (Fig. 1a).

From 700 to 500 °C, olivine and orthopyroxene compositions change dramatically (Fig. 8) due to the effect of redox reactions (Reaction 1) initiated by metal phase transformations. The chondritic oxygen fugacity (after homogenization) is lower than the oxygen fugacity in equilibrium with ol + opx + $\gamma_E$ assemblages, causing these assemblages to undergo local reduction. The retrograde reduction reaction is manifested in ordinary chondrites by olivine Fa concentrations that decrease toward zoned taenite + kamacite particles (formed after relatively small amounts of taenite undercooling). The oxygen fugacities are shown as deviations from the chondritic (averaged) oxygen fugacity shown in Fig. 7a.

Fig. 7. a) The chondritic oxygen fugacity calculated by averaging the oxygen fugacities defined by olivine + orthopyroxene + $\gamma_E$ assemblages and olivine + orthopyroxene + $\gamma_M$ assemblages (metastable $f_{O_2}$-$T$ path) and olivine + orthopyroxene + $\gamma_M$ assemblages (equilibrium $f_{O_2}$-$T$ path). The oxygen fugacities are shown as deviations from the chondritic (averaged) oxygen fugacity shown in Fig. 7a.

The olivine and orthopyroxene compositions shown in Fig. 8 were calculated by assuming that the metastable taenite composition ($\gamma_M$) remains constant at 10 wt% Ni throughout cooling. In reality, $\gamma_M$ may become somewhat Ni-enriched during cooling as Fe becomes oxidized and incorporated into the silicates. Nonetheless, the simplified calculations are justified because the $\gamma_M$ composition will remain Ni-poor relative to $\gamma_E$ throughout cooling.

According to the scenario presented above, the retrograde redox reactions in ordinary chondrites were not caused by external oxygen fugacity changes. Rather, they were caused by variable amounts of taenite undercooling and were facilitated by whole-rock homogenization of the internally controlled oxygen fugacity. The redox reactions would not have occurred if every taenite particle in the chondrite had experienced the same amount of undercooling, as olivine-orthopyroxene-metal equilibria would have defined the oxygen fugacity at a single value everywhere throughout the chondrite and no disequilibrium would exist to drive changes in silicate compositions.
Fig. 8. The compositional evolution of olivine and orthopyroxene in ordinary chondrites during cooling. At temperatures between 900 and 700 °C, olivine Fa concentrations decrease slightly and orthopyroxene Fs concentrations increase slightly due to Fe-Mg partitioning (Reaction 2). At temperatures below 700 °C, olivine and orthopyroxene compositions undergo significant changes due to redox reactions (Reaction 1). The solid lines show the compositions of olivine and orthopyroxene that coexist with γ_E, while the broken lines show the compositions of olivine and orthopyroxene that coexist with γ_M.

**Reaction Kinetics and the Absence of Measured Orthopyroxene Zoning**

Olivine crystals in H5 and H6 chondrites were compositionally homogeneous at the peak metamorphic temperature, and the high temperature compositions are preserved at olivine cores. However, silicate-metal reactions during cooling produced zoning of 3–4 mol% Fa within ~30 μm of olivine-metal interfaces. The zoning developed at temperatures <700 °C, the temperature at which taenite compositions diverge along the γ_E and γ_M paths.

The average distance of diffusion in olivine can be estimated using the Einstein relation, \( x = (2Dt)^{1/2} \), where \( x \) is distance, \( D \) is the olivine Fe-Mg interdiffusion coefficient, and \( t \) is time. Diffusion distances in olivine at 700, 600, and 500 °C, calculated for one million year isothermal heat treatments using interdiffusion coefficients reported by Chakraborty (1997), are 500, 100, and 10 μm, respectively. The diffusion calculations show that 30 μm long olivine Fa gradients, such as those measured in this study (Fig. 5), can indeed form at temperatures <700 °C. The relatively short diffusion profiles of ~30 μm indicate that the zoning developed at temperatures well below 700 °C. This interpretation is consistent with the expectation that the driving force responsible for the development of zoning is zero at 700 °C and becomes greater during cooling as the γ_E and γ_M compositions diverge.

The thermodynamic calculations (Fig. 8) indicate that orthopyroxene Fs concentrations should decrease toward zoned taenite + kamacite particles and increase toward zoneless plessite particles. Electron microprobe line scans across orthopyroxene crystals toward metal, however, did not reveal Fe-Mg zoning. It is possible that Fe-Mg gradients are present in orthopyroxene but are too narrow to measure with an electron microprobe. At a given temperature Fe-Mg interdiffusion in orthopyroxene is 2 orders of magnitude slower than Fe-Mg interdiffusion in olivine (Ganguly and Tazzoli 1994; Chakraborty 1997). Since diffusion distance scales with \( D^{1/2} \), diffusion distances in orthopyroxene will be 1 order of magnitude smaller than those in olivine. The measured Fe-Mg diffusion distances in olivine are <30 μm (Fig. 5), suggesting that orthopyroxene Fe-Mg profiles will be <3 μm long. Such short diffusion profiles in orthopyroxene would be difficult to resolve with an electron microprobe.

Fe-Mg interdiffusion coefficients in olivine and orthopyroxene increase as oxygen fugacity increases. Since ordinary chondrites evolved under more reducing conditions than the experimental conditions used to measure the diffusion coefficients, the diffusion distances calculated in this paper may be regarded as maximum distances.

Finally, it is important to note that Fe-Mg zoning profiles can potentially develop in olivine (near Fe-Ni metal) without the formation of analogous zoning profiles in orthopyroxene. Olivine zoning profiles measured near zoned taenite + kamacite particles (decreasing Fa toward metal) are opposite from olivine zoning profiles measured near zoneless plessite particles (increasing Fa toward metal). The opposite Fe-Mg zoning profiles can be explained by exchange of Fe\(^{2+}\) and Mg\(^{2+}\) between olivine crystals via grain boundary diffusion. In this case, grain boundaries need not migrate and orthopyroxene is neither formed nor consumed. Experimental studies by Dohmen et al. (1998) demonstrate that Fe and Mg transport can occur through a gaseous phase (i.e., along grain boundaries) when the partial pressures of both species are quite low.

**Implications for Ordinary Chondrites**

**Final Equilibration Temperatures**

In H-group chondrites, the redox reactions responsible for the formation of olivine Fe-Mg zoning adjacent to metal were initiated at temperatures <700 °C (the temperature at which the metal cooled below the taenite/(taenite + kamacite) solvus). The temperature at which redox reactions apparently ceased can be estimated by comparing the olivine Fe-Mg zoning pattern measured at olivine-metal interfaces (Fig. 5) to the calculated olivine compositions shown in Fig. 8. In the chondrites we examined, olivine Fa concentrations decrease by ~2 mol% near zoned taenite + kamacite particles and increase by ~2 mol% near zoneless plessite particles (Fig. 5). The olivine composition range of 3–4 mol% Fa suggests that the redox reactions effectively ended at ~600 °C (Fig. 8). This final equilibration temperature (600 °C) is identical to the Fe-
Mg exchange closure temperature obtained for olivine-orthopyroxene pairs that are isolated from metal (compare Figs. 4 and 6).

The final equilibration temperature of 600 °C for both the redox reaction (Reaction 1) and the olivine-orthopyroxene Fe-Mg exchange reaction (Reaction 2) is below the closure temperatures recorded by orthopyroxene-diopside Fe-Mg exchange and orthopyroxene-diopside Ca exchange equilibria (820–830 °C) (Olsen and Bunch 1984). However, it is similar to olivine-spinel Fe-Mg exchange closure temperatures measured in several chondrites. For example, olivine-spinel temperatures in Bjurböle (L/LL4), Aldsworth (LL5), and Appley Bridge (LL6) are <600 °C (Johnson and Prinz 1991) and olivine-spinel temperatures in Carcote (H5) are between 570 and 720 °C (Kleinschrodt and Okrusch 1999).

Metal particles in well-equilibrated H-group chondrites have bulk compositions of ~10 wt% Ni, and consequently cooled into the two-phase taenite + kamacite field at ~700 °C (Fig. 1a). This temperature was sufficiently high to allow detectable progress of silicate-metal reaction during subsequent cooling. In contrast, L and LL chondrites have whole-rock metal compositions of 15–30 wt% Ni and cooled into the taenite + kamacite field at temperatures <600 °C (Fig. 1a). The present study was limited to the examination of H chondrites (owing to the high density of silicate-metal interfaces), but the authors suspect that relatively sluggish diffusion rates at temperatures <600 °C may have limited the importance of retrograde silicate-metal reactions in L and LL chondrites.

Interpretation of “Anomalous” Olivine Compositions

Random collection of olivine analyses using the electron microprobe will occasionally sample Fe-Mg diffusion profiles that developed by retrograde silicate-metal reactions. Even olivine crystals that do not appear contiguous with metal in the plane of sectioning may be compositionally zoned due to the complex geometries of olivine-metal interfaces.

Clearly, any interpretation of olivine compositions in highly quenched H chondrites should recognize the important role of retrograde silicate-metal reactions. In one relevant study, Rubin (1990) reported olivine composition histograms for several H5 and H6 chondrites. In several of the chondrites, a few olivine analyses vary by up to ~2 mol% Fa from the host chondrite’s mean olivine composition (see Fig. 7.7 in Rubin 1990). Although these chondrites lack obvious brecciation textures such as angular clasts or light-dark structures, Kallemeyn et al. (1989) and Rubin (1990) proposed that the anomalous olivine compositions represent aberrant crystals that were incorporated into the host chondrite after cooling from the peak metamorphic temperature. The same authors suggested that these post-metamorphic fragmental breccias were cemented by shock lithification at relatively low ambient temperatures (i.e., low lithification temperatures prevented the aberrant olivine crystals from equilibrating with other crystals). We do not dispute the interpretations of Kallemeyn et al. (1989) and Rubin (1990), but we wish to emphasize that small amounts of olivine Fe-Mg zoning are an expected consequence of retrograde silicate-metal reaction. In fact, the existence of olivine Fe-Mg zoning adjacent to metal provides strong evidence that the silicate and metallic minerals evolved in situ during cooling from the peak metamorphic temperature.

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