

An evolutionary system of mineralogy, Part V: Aqueous and thermal alteration of planetesimals (~4565 to 4550 Ma)

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ABSTRACT

Part V of the evolutionary system of mineralogy explores phases produced by aqueous alteration, metasomatism, and/or thermal metamorphism—relicts of ancient processes that affected virtually all asteroids and that are preserved in the secondary mineralogy of meteorites. We catalog 166 historical natural kinds of minerals that formed by alteration in the parent bodies of chondritic and non-chondritic meteorites within the first 20 Ma of the solar system. Secondary processes saw a dramatic increase in the chemical and structural diversity of minerals. These phases incorporate 41 different mineral-forming elements, including the earliest known appearances of species with essential Co, Ge, As, Nb, Ag, Sn, Te, Au, Hg, Pb, and Bi. Among the varied secondary meteorite minerals are the earliest known examples of halides, arsenides, tellurides, sulfates, carbonates, hydroxides, and a wide range of phyllosilicates.

Keywords: Philosophy of mineralogy, classification, mineral evolution, natural kinds, meteorite mineralogy, thermal metamorphism, aqueous alteration, metasomatism

INTRODUCTION—HISTORICAL NATURAL KINDS

The evolutionary system of mineralogy classifies “historical natural kinds” (Boyd 1991, 1999; Hawley and Bird 2011; Magnus 2012; Khalidi 2013; Ereshefsky 2014; Godman 2018; Cleland et al. 2020) based on paragenetic modes of minerals, as manifest in their distinctive combinations of attributes. In accord with Godman’s (2018) concept of “historical essences,” our approach to mineral classification relies on a closely linked pairing of “individuation” and “causal explanation.” In other words, mineral classification in a historical context must be based equally on diagnostic suites of mineral properties and the inferred processes by which those distinctive properties arose.

We contend that the information-rich natures of different mineral kinds, including their trace and minor elements, isotopic ratios, structural defects, solid and fluid inclusions, morphologies, zoning, twinning, and myriad other physical and chemical characteristics, are direct consequences of their physical, chemical, and/or biological modes of origin and, in many cases, subsequent alteration (Hazen 2019). The evolutionary system thus embraces the intrinsic data-rich characters and varied historical contexts of minerals while building on standard protocols of the Commission on New Minerals, Nomenclature and Classification of the International Mineralogical Association (IMA), which discriminate among mineral “species” based exclusively on idealized major element chemical composition and atomic structure (e.g., Burke 2006; Mills et al. 2009; Schertl et al. 2018; Hatert et al. 2021; Hazen 2021).

The first five parts of the evolutionary system collectively encompass the variety of condensed phases formed in presolar environments and during the first 15 to 20 Ma of the solar system,

most of which are accumulated and preserved in meteorites. Part I (Hazen and Morrison 2020) cataloged stellar minerals that predate our solar nebula, i.e., prior to 4.567 Ga. Subsequently, in Part II, we explored primary interstellar and nebular condensates commencing ~4.567 Ga (Morrison and Hazen 2020), while the primary mineralogy of chondrules from ~4.566 to 4.561 Ga was the focus of Part III (Hazen et al. 2021). Part IV summarized the primary asteroidal mineralogy of non-chondritic meteorites from ~4.566 to 4.560 Ga, as well as high-pressure impact mineralogy preserved in meteorites (Morrison and Hazen 2021). Note that primary and secondary minerals in IDPs (e.g., Rietmeijer 1999; Brownlee 2016) and comets (e.g., Brownlee 2014) will be summarized in a later part of this series.

These first four parts of the evolutionary system of mineralogy were relatively straightforward in their blending of diagnostic attributes with causal explanation, as required for a valid enumeration of historical kinds (Godman 2018). For example, stellar minerals possess characteristic isotopic anomalies that derive directly from nucleosynthetic processes in evolving stars—attributes that set them apart from all other mineral occurrences. Likewise, the primary condensates of calcium-aluminum-rich inclusions (CAIs) and amoeboid olivine aggregates (AOAs), the primary igneous phases of chondrules, and the primary minerals of differentiated asteroids display ranges of physical and chemical characteristics that reveal direct links between their presumed modes of origin and their diagnostic mineral attributes. Similarly, in the case of impact minerals, the appearance of micrometer-scale, dense high-pressure phases in the context of lower-pressure assemblages provides a clear connection between mineral properties and the rapid and violent shock events that formed them. However, in Part V we encounter a more nuanced and potentially problematic situation.

Part V continues our systematic exploration of pre-terrestrial mineralogy with an examination of “secondary” asteroidal

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minerals formed by aqueous alteration, metasomatism, and/or thermal metamorphism in planetesimals. These events occurred primarily during the first ~15 Ma of solar system history (McSween et al. 1988; Krot et al. 2006; Kleine et al. 2018), with significant alteration occurring in the first 5 Ma as a consequence of heating by short-lived radioisotopes (Ghosh et al. 2006; Fujiya et al. 2012; Doyle et al. 2015). [Note that we distinguish between secondary minerals formed in planetesimals (included here) from those formed much more recently through terrestrial weathering (to be reviewed in a later contribution).] The principal difficulty in dealing with these varied secondary phases in meteorites is that they form gradually from “primary” minerals, with sometimes subtle shifts in composition and structure that span thousands to millions of years. Consequently, even though we may be able to recognize suites of secondary mineral properties (e.g., zoning, site order/disorder, defect density, and/or exsolution) that can be directly linked to causal events (thermal and aqueous alteration), in several instances no sharp boundary exists between the primary and secondary forms of minerals. An additional degree of uncertainty arises from the hypothesis that some presumably secondary asteroidal minerals, including halides, Fe²⁺ phases, and clay minerals, also might have formed via nebular processes, for example oxidation, sulfidization, or hydration through pre-accretionary interaction with warm nebular gas (Krot et al. 1995, their Table 2; Bischoff 1998; Ciesla et al. 2003).

A degree of irony exists in this situation, as some philosophers of science (e.g., Ereshefsky 2014; Godman 2018) have argued that historical natural kinds can only be valid in a system with the property of common descent; e.g., “historical essentialism of a Kind such as a particular species demands some reproductive or near reproductive relation” (Godman 2018, p.13). However, in the context of mineral historical kinds, primary phases—minerals formed *de novo* from a vapor or liquid with no prior minerals, and thus with no possible analogy to common descent—provide the least ambiguous examples of valid historical kinds (Cleland et al. 2020). The stellar condensate *AGB spinel* is clearly a distinct historical kind from *CAI spinel* by virtue of different isotopic systematics. *CAI spinel*, in turn, is distinct from primary igneous chondrule spinel (*PC spinel*) on the basis of oxygen isotopes, morphology, and petrologic context. In each of these instances, a diagnostic suite of attributes is directly linked to a specific historical causal context without pre-existing minerals.

By contrast, oxide spinel minerals from primary asteroid assemblages may blend continuously into those from secondary assemblages. This mineralogical analog of “common descent,” by which secondary oxide spinels gradually “evolve” from primary oxide spinels and other phases, leads to ambiguity in classification. How do we recognize a primary/secondary boundary in such minerals? In this example, the philosophical literature on biological speciation has some relevance. For example, Ereshefsky (2014) explores the conundrum of defining a new biological species as a “founder population” begins to branch off from a prior species. He contends that one can recognize the new biological species only in retrospect—only in the historical context of subsequent evolution. Furthermore, there exists a transition regime of individuals that are not strictly of the parent species nor of the daughter species. Such a biological scenario in some ways parallels that of primary minerals transforming

gradually to secondary minerals, especially in the sense that there may exist intermediate stages of transition that do not unambiguously belong to either the primary or the secondary mineral assemblage.

In this contribution, we bypass this issue, at least in part, by focusing exclusively on the appearance of new secondary species in the IMA sense—i.e., phases with a new combination of atomic structure and major element composition compared to their precursors. Thus, for example, the appearances of sulfates, carbonates, clay minerals, and hydroxides are all treated as secondary meteorite minerals in Part V. Similarly, we consider plagioclase formed by thermal devitrification of primary chondrule feldspathic glass as a secondary phase. However, we do not distinguish between primary chondritic olivine, which is characterized by disparate compositions in adjacent chondrules, as opposed to modified olivine in fully equilibrated chondrites, which experienced diffusion that led to more uniform mineral compositions in adjacent chondrules owing to thermal alteration.

SECONDARY PROCESSES IN PLANETESIMALS

All chondritic and non-chondritic meteorites were subjected to alteration processes in their parent bodies (McSween et al. 1988; Sears and Dodd 1988; Zolensky and McSween 1988; Brearley and Jones 1998; Mittlefehldt et al. 1998; Krot et al. 2006, 2014; Mittlefehldt 2014; Rubin and Ma 2017, 2021; Russell et al. 2018). Consequently, the mineralogy of many meteorites has been complicated by repeated episodes of rapid and gradual heating, shock transformation and impact brecciation, and interactions with aqueous fluids—processes that, as noted above, often caused continuous, gradual alterations that blur the boundaries between “primary” and “secondary” minerals. Gradations may also exist among low-temperature aqueous alteration and hydrothermal/metamorphic processes, which span wide ranges of temperature-composition space at pressures <0.2 GPa (McSween et al. 1988; Zolotov 2009). Therefore, rather than invoke arbitrary distinctions among the phases formed by these varied secondary processes, we lump them all into “secondary asteroidal” (“*SA*”) minerals. Nevertheless, we acknowledge that debates remain unresolved regarding the primary vs. secondary origins of several meteorite phases.

Our decision to lump together all secondary minerals of a given species is, admittedly, subjective. For example, we could have differentiated secondary olivine formed by thermal metamorphism and dehydration of phyllosilicates (e.g., Tomeoka et al. 1989c) from secondary olivine formed by metasomatic exchange reactions with a Fe-rich fluid (Varela et al. 2012). Other minerals with likely multiple modes of secondary formation include troilite, calcite, calcic clinopyroxene, and plagioclase. However, until more data are available on the diagnostic chemical and physical attributes imparted by these different secondary modes of mineral formation, we group them into single historical natural kinds.

Secondary alteration of chondrite meteorites

The mineralogical consequences of aqueous alteration, metasomatism, and thermal metamorphism differ among different groups of meteorites. [For nomenclature of the many kinds of chondrite and achondrite meteorites, as well as their components, see Krot et al. (2014).] Alteration of chondrites

has been reviewed by numerous authors (Brearley and Jones 1998; Brearley 2006; Huss et al. 2006; Krot et al. 2006). Van Schmus and Wood (1967) introduced a numerical scale that, in its current guise, defines the least altered (i.e., “unequilibrated”) chondrites as “3.0,” with greater numbers up to 7 representing increasing degrees of thermal alteration to temperatures of ~950 to 1000 °C—the highest temperature at which these meteorites still retain some of their distinctive compositional and/or textural characteristics (Dodd 1981; McSween et al. 1988; McSween and Patchen 1989; Huss et al. 2006). At modest degrees of thermal alteration, higher resolution scales between 3.0 and 3.9, as well as from 3.00 to 3.15, have been devised for ordinary and CO chondrites (e.g., Grossman and Brearley 2005).

Common thermal metamorphic changes in chondrites include gradual equilibration of silicate compositions, especially Fe/Mg ratios in olivines and pyroxenes, through element diffusion, as well as devitrification of silicate glass, most commonly manifest as nucleation of feldspar and possibly augite (Sears et al. 1980; Sears and Hasan 1987; Scott et al. 1994). Thermal metamorphism also resulted in dehydration of phyllosilicates, gradual oxidation of Fe metal to Fe²⁺, exsolution of new phases, and varied solid-state transformations (Rubin 2005; Rubin and Ma 2017, 2021). A chondrite of type 3.9 corresponds to metamorphism at ~600 °C, at which the compositions of olivines in diverse chondrules have largely equilibrated and feldspars may exhibit Al-Si disorder (Sears et al. 1995). Type 6 or 7 chondrites experienced temperatures close to 1000 °C, based on changes in metal compositions (McSween et al. 1988).

In a similar fashion, chondrites affected by aqueous processes display alteration of metal alloys and silicates, with corresponding textural changes (e.g., Van Schmus and Wood 1967; McSween 1979). Increasing extents of aqueous alteration are designating by decreasing numbers below 3.0, from 2.9 to 1.0 (Van Schmus and Wood 1967; Browning et al. 1996; Rubin et al. 2007; Marrocchi et al. 2014). These varying degrees of aqueous alteration and their associated clay mineralogy have also been correlated to reflectance spectra of CM and CI chondrites and their presumed parent bodies (Takir et al. 2013). Estimated temperatures for aqueous alteration range from close to 0 °C to <150 °C (Clayton and Mayeda 1984; Zolensky and McSween 1988).

Note that these numbering schemes to designate degrees of thermal and aqueous alteration are complemented by numerical scales for shock alteration (S1 to S7; Stöffler et al. 1991, 2018) and terrestrial weathering (W0 to W6; Wlotzka 1993; Bland et al. 2006).

SYSTEMATIC MINERALOGY OF SECONDARY ASTEROIDAL MINERALS

In the previous part of this series (Morrison and Hazen 2021), we listed 40 high-pressure minerals that formed through rapid shock alteration. In this contribution, we consider 166 historical natural kinds that arose through the more gradual changes to primary asteroidal minerals caused by the range of low-pressure (<0.2 GPa) processes subsumed under aqueous alteration, metasomatism, and thermal metamorphism (Table 1).

Aqueous alteration occurred when anhydrous minerals became hydrated and/or, in some instances, oxidized, carbonated, or sulfidized (Zolensky and McSween 1988; Zolensky et al.

1993; Krot et al. 1995; Brearley 2006). Such alteration occurred in asteroids and planetesimals as H₂O-dominant fluids were mobilized in warming events that melted precursor ice, though in some instances hydration may have also resulted from interaction with nebular gas (e.g., Bischoff 1998). Aqueous alteration also affected isotopic compositions; three major oxygen reservoirs existed in the protoplanetary disk, each with different initial O isotopes: CO, H₂O, and silicates. Krot (2019) reviewed this rich O-isotope record and described how, for example, aqueous alteration by ¹⁶O-depleted H₂O is revealed in ¹⁶O-depleted minerals.

Thermal metamorphism resulted from heating in asteroids, primarily within the first 10 Ma of the solar system as a consequence of the decay of short-lived radioisotopes such as ²⁶Al and ⁶⁰Fe, as well as electromagnetic induction (McSween et al. 1988; Ghosh et al. 2006) and solar heating for asteroids with orbits that passed close to the Sun (Wittmann et al. 2011; Libourel et al. 2015). In the case of chondrite parent bodies, which were sedimentary accumulations comprised of diverse objects from various sources, a major consequence of thermal metamorphism was increasing degrees of equilibrium among their diverse collections of chondrules. Thermal metamorphism may also be accompanied by reduction, which at times produced Fe alloys at the expense of more Fe-rich silicates (Rambaldi and Wasson 1982; McSween et al. 1988; Wasson et al. 1993; Menzies et al. 2005; Huss et al. 2006; Simon et al. 2016).

Note that in most instances, these two processes appear to have been independent, as thermal metamorphism was usually an anhydrous process (McSween et al. 1988). For example, CM carbonaceous chondrites, epitomized by the Murchison CM2 meteorite, were affected by aqueous alteration but not significantly by thermal metamorphism (McSween 1979). Nevertheless, several meteorites, notably ordinary and carbonaceous chondrites (Krot et al. 2004; Dobrică and Brearley 2014; Fintor et al. 2014; Harries and Zolensky 2016; Vacher et al. 2019), display effects of metasomatism and hydrothermal activity. For example, Dyl et al. (2012) presented evidence for short-lived (1 to 10 years) hydrothermal activity in a metamorphosed ordinary chondrite at an estimated temperature of 800 °C and 1 bar water pressure, based on variations in feldspar and oxygen isotopic compositions.

Even when aqueous and thermal effects are decoupled, some secondary minerals may have formed through a sequence of processes (e.g., Zolensky et al. 1993; Krot et al. 2004). For example, Krot et al. (1997a) proposed that secondary fayalite in cracks and as rims on phenocrysts in CV chondrites arose from the thermal metamorphism of phyllosilicates, which formed by aqueous alteration of forsterite. Similarly, Ikeda and Prinz (1993) and Kimura and Ikeda (1992) described a sequence of mineralization in the Belgica-7904 carbonaceous chondrite, with phyllosilicate formation followed by dehydration. Furthermore, the chondrules in oxidized CV chondrites have mesostases that were metasomatized by Fe-alkali-halogen-bearing fluids. The result was the replacement of mesostasis glass and plagioclase by nepheline and sodalite, plus minor grossular, wollastonite, andradite, kirschsteinite, and hedenbergite (Ikeda and Kimura 1995; Kimura and Ikeda 1995), as well as merrillite (Murakami and Ikeda 1994). In addition, superimposed on these mineralogical changes was the potential for additional heating events, including effects of violent impact processes, which may have

TABLE 1. Secondary asteroidal minerals in chondrite and achondrite meteorites

Group species (formula)	Natural kind	Paragenesis	References
Native elements and alloys			
Iron or "kamacite" (α -Fe,Ni) ^a	SA iron	Thermal metamorphism of silicates in reducing C-rich environments	1,2
Taenite (γ -Fe,Ni) ^a	SA taenite	Thermal metamorphism/oxidation of Fe in kamacite	3
Tetrateaenite (Fe,Ni)	SA tetrateaenite	Thermal metamorphism/annealing and Fe-Ni ordering of taenite	4
Awaruite (Ni ₃ Fe)	SA awaruite	Thermal metamorphism/preferential oxidation of Fe in Fe-Ni alloys	5–7
Wairauite (CoFe)	SA wairauite	Thermal metamorphism/oxidation of Fe-Ni alloys	8–10
Copper (Cu)	SA copper	Thermal metamorphism/oxidation of Fe-Ni alloys	7,11–13
Mercury (Hg)	SA mercury	Thermal metamorphism/sublimation in asteroid interior	14
Platinum-iron Alloy (Pt,Fe)	SA Pt-Fe alloy	Thermal metamorphism/exsolution from Fe-Ni alloys	15,16
Osmium Alloy (Os)	SA Os alloy	Thermal metamorphism/exsolution from Fe-Ni alloys	15,16
Gold Alloy (Au)	SA Au alloy	Thermal metamorphism/exsolution from Fe-Ni alloys	15–17
Niggliite (PtSn)	SA niggliite	Thermal metamorphism/exsolution from Fe-Ni alloys	15,16
Rustenburtite (Pt ₃ Sn)	SA rustenburtite	Thermal metamorphism/exsolution from Fe-Ni alloys	15,16
Graphite (C) ^a	SA graphite	Annealing of carbonaceous material; exsolution from C-rich metal	18–23
Sulfur (S)	SA sulfur	Alteration of pyrrhotite, with which it is usually associated	24,25
Carbides			
Cohenite [(Fe,Ni) ₃ C]	SA cohenite	Thermal metamorphism/exsolution from Fe-Ni alloys	26–28
Haxonite [(Fe,Ni) ₂₃ C ₆]	SA haxonite	Thermal metamorphism/exsolution from Fe-Ni alloys	26
Phosphides			
Schreibersite [(Fe,Ni) ₃ P]	SA schreibersite	Thermal metamorphism/exsolution from Fe-Ni alloys	29,30
Florenskyite (FeTiP)	SA florenskyite	Thermal metamorphism/exsolution from Fe-Ni alloys	31,32
Andreyivanovite (FeCrP)	SA andreyivanovite	Thermal metamorphism/exsolution from Fe-Ni alloys	33
Melliniite [(Fe,Ni) ₄ P]	SA melliniite	Thermal metamorphism/exsolution from Fe-Ni alloys	34
Halides			
Halite (NaCl)	SA halite	Precipitation from an aqueous fluid in chondrite meteorites	35,36
Sylvite (KCl)	SA sylvite	Precipitation from an aqueous fluid in chondrite meteorites	35,36
Chlormayenite [Ca ₁₂ Al ₁₄ O ₃₂ (□ ₄ Cl ₂)]	SA chlormayenite	Aqueous precipitation in the NWA 1934 carbonaceous chondrite	37
Unnamed (BiCl ₃)	SA unnamed BiCl ₃	Aqueous precipitation in the LAP 04840 R chondrite	38
Sulfides			
Troilite (FeS) ^a	SA troilite	Common as sulfidation of Fe-Ni alloys; exsolution	7,39–42
Pyrrhotite (Fe ₇ S ₉) ^a	SA pyrrhotite	Common in CI and CM matrices; oxidation of troilite; hydrothermal	39,42–46
Pyrite (FeS ₂)	SA pyrite	A minor secondary phase; oxidation and/or hydrothermal	13,17,47,48
Greigite (Fe ₃ S ₄)	SA greigite	A minor sulfidation/hydrothermal phase in enstatite chondrites	21
Smythite (Fe ₃ S ₁₁)	SA smythite	A minor sulfidation/hydrothermal phase in enstatite chondrites	21
Millerite (NiS)	SA millerite	A minor sulfidation/hydrothermal phase in CK chondrites	17
Heazlewoodite (Ni ₃ S ₂)	SA heazlewoodite	A minor sulfidation/hydrothermal phase in carbonaceous chondrites	46,49,50
Pentlandite [(Ni,Fe) ₉ S ₈] ^a	SA pentlandite	Common as sulfidation of Fe-Ni alloys; aqueous alteration	39,42,50–52
Shenzhuangite (FeNi ₅ S ₂)	SA shenzhuangite	Sulfidation of taenite in the Suizhou ordinary chondrite	53
Covellite (CuS)	SA covellite	Alteration of djerfisherite in enstatite chondrites	21
Chalcopyrite (CuFeS ₂)	SA chalcopyrite	Rare secondary phase in carbonaceous chondrites	13,17,54–55
Idaite (Cu ₃ FeS ₄)	SA idaite	Alteration of djerfisherite in enstatite chondrites	21
Bornite (Cu ₃ FeS ₄)	SA bornite	Alteration of djerfisherite in enstatite chondrites	21
Cubanite (CuFe ₂ S ₃)	SA cubanite	Aqueous alteration at T < ~200 °C	47,56
Isocubanite (CuFe ₂ S ₃)	SA isocubanite	Rare aqueous alteration phase in matrices of CI chondrites	56,57
Brezinaite (Cr ₂ S ₃)	SA brezinaite	Thermal metamorphism/sulfidation of chromite	58,59
Murchisite (Cr ₂ S ₆)	SA murchisite	Exsolution from Cr-rich Fe-Ni alloy at T ≤ 300 °C	60
Daubréelite (FeCr ₂ S ₄)	SA daubréelite	Common sulfidation product in enstatite chondrites	39,59,61,62
Sphalerite [(Zn,Fe)S]	SA sphalerite	Thermal metamorphism in the Qingzhen enstatite chondrite	63
Rudashevskyite [(Fe,Zn)S]	SA rudashevskyite	Thermal metamorphism in enstatite chondrites	63–65
Unnamed [(V,Fe,Cr) ₂ S ₃]	SA unnamed [(V,Fe,Cr) ₂ S ₃]	A rare secondary phase in the Sierra Gorda CB chondrite	59
Molybdenite (MoS ₂)	SA molybdenite	Sulfidation of refractory metals in CV chondrites	66–69
Wassonite (WS)	SA wassonite	Sulfidation of refractory metals in Yamato 691 enstatite chondrite	70
Cinnabar (HgS)	SA cinnabar	Thermal alteration and sublimation	14
Erlichmanite (OsS ₂) and Laurite (RuS ₂)	SA erlichmanite	Sulfidation of refractory metals in carbonaceous and R chondrites	17,48
Cooperite (PtS ₂)	SA cooperate	Sulfidation of refractory metals in CK chondrites	17
Petrowskaite [(Au,Fe,Ag) ₂ S]	SA petrowskaite	Sulfidation of refractory metals in the LEW 87009 CK chondrite	17
Unnamed [(Fe,Au,Co) ₂ S ₃]	SA unnamed [(Fe,Au,Co) ₂ S ₃]	Sulfidation of metal alloys in CK chondrite	17
Nuwaite (Ni ₆ GeS ₂) and Butianite (Ni ₆ SnS ₂)	SA nuwaite	Sulfidation of metal alloys in Allende CV chondrite	71
Djerfisherite [K ₆ (Fe,Cu,Ni) ₂₅ S ₂₆ Cl]	SA djerfisherite	Aqueous alteration of sulfides	21,63,72
Unnamed hydrated [(Na,Cu ⁺)CrS ₂]	SA unnamed hydrated [(Na,Cu ⁺)CrS ₂]	Aqueous alteration of caswellsilverite	21
Unnamed hydrated Na-Cu-Zn-Cr sulfide	SA unnamed hydrated Na-Cu-Zn-Cr sulfide	Aqueous alteration of caswellsilverite	21
Tochilinite {6(Fe _{0.9} S) ₅ [(Mg,Fe)(OH) ₂]} ^a	SA tochilinite	Aqueous alteration phase in chondrite matrices	30,73–76
Arsenides			
Sperrylite (PtAs ₂)	SA sperrylite	Aqueous alteration in oxidized R chondrites	15,16
Irarsite (IrAsS)	SA irarsite	Aqueous alteration in oxidized R chondrites	15–17
Tellurides			
Moncheite [(Pt,Pd)(Te,Bi) ₂]	SA moncheite	Aqueous alteration in R and CK chondrites	15–17,77,78
Unnamed Au-Pt-Fe telluride	SA unnamed Au-Pt-Fe telluride	Aqueous alteration in CK chondrites	17
Altaite (PbTe)	SA altaite	Possible post-magmatic alteration in iron meteorites	79,80

(Continued on next page)

TABLE 1.—CONTINUED

Group species (formula)	Natural kind	Paragenesis	References
Sulfates			
Anhydrite (CaSO ₄) ^a	SA anhydrite	Aqueous alteration in carbonaceous chondrites	81–84
Gypsum (CaSO ₄ ·2H ₂ O) ^a	SA gypsum	Aqueous alteration in carbonaceous chondrites	24,84–89
Bassanite (CaSO ₄ ·0.5H ₂ O) ^a	SA bassanite	Aqueous alteration in carbonaceous chondrites	81,82
Hexahydrate (MgSO ₄ ·6H ₂ O)	SA hexahydrate	Aqueous alteration in carbonaceous chondrites	24,89
Epsomite (MgSO ₄ ·7H ₂ O)	SA epsomite	Aqueous alteration in carbonaceous chondrites	24,89
Thenardite (Na ₂ SO ₄)	SA thenardite	Aqueous alteration in Murray CM chondrite	90
Unnamed Mg-Al-Fe sulfate	SA unnamed Mg-Al-Fe sulfate	Aqueous alteration in Murray CM chondrite	73
Blödite [Na ₂ (Mg,Ni)(SO ₄) ₂ ·4H ₂ O]	SA blödite	Aqueous alteration in carbonaceous chondrites	24,25,91
Barite (BaSO ₄)	SA barite	Aqueous alteration in CV chondrites	7,92
Carbonates			
Calcite (CaCO ₃) ^a	SA calcite	Aqueous alteration in ordinary and carbonaceous chondrites	26,84,86,91,93–98
Dolomite [CaMg(CO ₃) ₂] ^a	SA dolomite	Aqueous alteration in ordinary and carbonaceous chondrites	39,91,99,100
Magnesite (MgCO ₃) and siderite (FeCO ₃) ^a	SA breunnerite	Aqueous alteration in ordinary and carbonaceous chondrites	39,91,100–102
Rhodochrosite (MnCO ₃)	SA rhodochrosite	Aqueous alteration in Yamato 82162 carbonaceous chondrite	54,102
Aragonite (CaCO ₃)	SA aragonite	Aqueous alteration in ordinary and carbonaceous chondrites	103
Phosphates			
Chlorapatite [Ca ₅ (PO ₄) ₃ Cl] ^a	SA chlorapatite	Aqueous alteration or thermal metamorphism of Fe-Ni alloys	13,40,48,49, 103–107
Hydroxylapatite [Ca ₅ (PO ₄) ₃ OH]	SA hydroxylapatite	Aqueous alteration of carbonaceous chondrites	103,105,108
Fluorapatite [Ca ₅ (PO ₄) ₃ F]	SA fluorapatite	Aqueous alteration of alkaline clasts	109,110
Merrillite [Ca ₉ NaMg(PO ₄) ₆] ^a	SA merrillite	Thermal metamorphism of P-bearing Fe-Ni alloys	104–106,111
Whitlockite [Ca ₉ Mg(PO ₃ OH)(PO ₄) ₆]	SA whitlockite	Low-temperature aqueous alteration	68,112
Sarcopside [Fe ₃ ²⁺ (PO ₄) ₂] and chopinite [Mg ₃ (PO ₄) ₂]	SA sarcopside	Oxidation of P-rich metal	113
Farringtonite [(Fe,Mn) ₂ (PO ₄) ₃]	SA farringtonite	Oxidation of P-rich metal	113
Chladniite [Na ₂ CaMg ₇ (PO ₄) ₆] and johnsomervilleite [Na ₁₀ Ca ₈ Mg ₁₈ Fe ₂₂ (PO ₄) ₃₆]	SA chladniite	Oxidation of P-rich metal	113–115
Brianite [Na ₂ CaMg(PO ₄) ₂]	SA brianite	Oxidation of P-rich metal	116
Oxides			
Magnetite (Fe ₂ O ₃) ^a	SA magnetite	Oxidation of kamacite or troilite	3,30,39,84,117,118
Chromite (Fe ₂ ²⁺ Cr ₂ O ₄) ^a	SA chromite	Oxidation of metal or sulfides	84,107,119–121
Spinel (MgAl ₂ O ₄)	SA spinel	Thermal metamorphism of carbonaceous chondrites	17,35
Hercynite (Fe ²⁺ Al ₂ O ₄)	SA hercynite	Thermal metamorphism of carbonaceous chondrites	35,40
Ulvöspinel (Fe ²⁺ Ti ⁴⁺ O ₄)	SA ulvöspinel	Thermal metamorphism of CO chondrites	122
Coulsonite [(Fe,Mg)V ₂ O ₄]	SA coulsonite	Thermal metamorphism of Allende CV chondrite	40,123
Periclase (MgO) and wüstite (FeO)	SA magnesiowüstite	Oxidation of carbonaceous chondrites	54,124,125
Corundum (Al ₂ O ₃)	SA corundum	Thermal alteration of hibonite in Allende CV chondrite	126,127
Maghemite [(Fe _{0.67} □ _{0.33})Fe ₂ ³⁺ O ₄]	SA maghemite	Oxidation of matrix in Semarkona ordinary chondrite	26
Ilmenite (FeTiO ₃) ^a	SA ilmenite	Thermal metamorphism of carbonaceous chondrites	3,17,41,128–131
Eskolaite (Cr ₂ O ₃)	SA eskolaite	Thermal alteration of kamacite	30,58,124
Rutile (TiO ₂)	SA rutile	Thermal metamorphism of ordinary chondrites	121,132
Pyrophanite (MnTiO ₃)	SA pyrophanite	Thermal metamorphism of the Raguli ordinary chondrite	133
Scheelite (CaWO ₄)	SA scheelite	Oxidation of refractory metal in Allende CV chondrite	40
Powellite (CaMoO ₄)	SA powellite	Oxidation of refractory metal in Allende CV chondrite	112
Unnamed Mg-Fe molybdate	SA unnamed Mg-Fe molybdate	Oxidation of refractory metal in Allende CV chondrite	40
Kamiokite [(Fe,Mg) ₂ Mo ₃ O ₈]	SA kamiokite	Oxidation of monipite in Allende CV chondrite	134
Majindete [(Mg,Fe) ₂ Mo ₃ O ₈]	SA majindete	Oxidation of Mo-rich phase in Allende CV chondrite	135
Tugarinovite (MoO ₂)	SA tugarinovite	Oxidation of monipite in Allende CV chondrite	134
Beckettite (Ca ₂ V ₆ Al ₆ O ₂₀)	SA beckettite	Oxidation of monipite in Allende CV chondrite	134
Unnamed [(Nb,V,Fe)O ₂]	SA unnamed [(Nb,V,Fe)O ₂]	Oxidation of monipite in Allende CV chondrite	134
Chihuahuaite [(Fe,Mg)Al ₁₂ O ₁₉]	SA chihuahuaite	Fe-alkali metasomatism of hibonite in Allende CV chondrite	136
Baddeleyite (ZrO ₂)	SA baddeleyite	Exsolution from ilmenite; thermal alteration	133,137
Hydroxides			
Brucite [Mg(OH) ₂]	SA brucite	Aqueous alteration of Mg silicates in carbonaceous chondrites	25,138
Amakinite [Fe(OH) ₂]	SA amakinite	Alteration by Fe-rich fluids in CM chondrites	76
Ferrihydrite [Fe ₁₀ O ₁₄ (OH) ₂]	SA ferrihydrite	Aqueous alteration of carbonaceous chondrite matrices	139,140
Limonite [FeO(OH)·nH ₂ O]	SA limonite	Aqueous alteration and oxidation by Fe-rich fluids	25
Silicates			
Quartz (SiO ₂)	SA quartz	Thermal metamorphism; exsolution from metal	107,141
Forsterite (Mg ₂ SiO ₄) and fayalite (Fe ₂ SiO ₄) ^a	SA olivine	Dehydration of phyllosilicates; oxidation of Fe metal	3,39,86,105,110, 118,142–148
Larnite (Ca ₂ SiO ₄)	SA larnite	Rare secondary phase from CV chondrites	149,150
Monticellite (CaMgSiO ₄)	SA monticellite	Thermal metamorphism of melilite in Allende CV chondrite	3,39,151
Kirschsteinite (CaFeSiO ₄)	SA kirschsteinite	Thermal alteration of CALs in carbonaceous chondrites	3,41,152
Grossular (Ca ₃ Al ₂ Si ₃ O ₁₂) ^a	SA grossular	Thermal alteration of CALs and matrices in carbonaceous chondrites	3,39,153–157
Andradite (Ca ₃ Fe ₂ ³⁺ Si ₃ O ₁₂) ^a	SA andradite	Thermal alteration of CALs and matrices in carbonaceous chondrites	3,39,155–158
Hutcheonite [Ca ₃ Ti ⁴⁺ (SiAl ₂)O ₁₂]	SA hutcheonite	Thermal alteration of CALs in the Allende CV chondrite	159
Goldmanite [Ca ₃ V ₂ ³⁺ (SiAl ₂)O ₁₂]	SA goldmanite	Thermal alteration of CALs in the Leoville CV chondrite	160
Titanite (CaTiSiO ₅)	SA titanite	Thermal alteration in carbonaceous and ordinary chondrites	131,161
Adrianite [Ca ₁₂ (Al ₆ Mg ₃ Si ₃)O ₃₂ Cl ₆]	SA adrianite	Alkali-halogen metasomatism of Ca-silicates	162
Wadalite [Ca ₆ Al ₅ Si ₂ O ₁₆ Cl ₃]	SA wadalite	Metasomatism of Ca silicates by Cl-rich fluids	162,163

(Continued on next page)

TABLE 1.—CONTINUED

Group species (formula)	Natural kind	Paragenesis	References
Enstatite (MgSiO ₃) and ferrosilite (FeSiO ₃) ^a	SA orthopyroxene	Thermal metamorphism/inversion of clinopyroxene	39,105
Diopside (CaMgSi ₂ O ₆) ^a	SA diopside	Thermal alteration of CAIs and matrices in carbonaceous chondrites	39,95,154,156, 164–166
Hedenbergite (CaFeSi ₂ O ₆) ^a	SA hedenbergite	Metasomatism by Fe-rich fluids	3,39,154,164,167
Donpeacorite or Kanoite (MnMgSi ₂ O ₆)	SA donpeacorite	Metasomatism	168
Wollastonite (CaSiO ₃) ^a	SA wollastonite	Metasomatism of CAIs in the Allende CV chondrite	3154,155,158,169
Magnesio-arfvedsonite [NaNa ₂ (Mg,Fe ³⁺)Si ₆ O ₂₂ (OH) ₂]	SA magnesio-arfvedsonite	Aqueous alteration in the Kaidun polymict breccia	111
Anthophyllite [(Mg,Fe) ₂ Si ₈ O ₂₂ (OH) ₂]	SA anthophyllite	Aqueous alteration in the Allende CV chondrite	170
Magnesio-hornblende [□Ca ₂ (Mg,Fe ²⁺) ₂ (Si,Al) ₂ O ₂₂ (OH) ₂]	SA magnesio-hornblende	Aqueous alteration in the Allende CV chondrite	170
Winchite [□NaCa(Mg,Al)Si ₆ O ₂₂ (OH) ₂] and barroisite [□NaCa(Mg ₃ Al ₂)Si ₆ O ₂₂ (OH) ₂]	SA winchite	Hydrothermal alteration in the Tieschitz ordinary chondrite	171
Jimthompsonite [(Mg,Fe) ₂ Si ₆ O ₁₆ (OH) ₂]	SA jimthompsonite	Aqueous alteration in the Allende CV chondrite	170
Aspidolite [NaMg ₃ AlSi ₂ O ₁₀ (OH) ₂]	SA aspidolite	Aqueous alteration in CV chondrites	3
Phlogopite [KMg ₃ AlSi ₃ O ₁₀ (OH) ₂]	SA phlogopite	Aqueous alteration in CV chondrites	172,173
Paragonite [NaAl ₂ Si ₃ O ₁₀ (OH) ₂]	SA paragonite	Aqueous alteration in the Murray CM chondrite	76
Clintonite [CaAlMg ₂ SiAl ₃ O ₁₀ (OH) ₂]	SA clintonite	Aqueous alteration in the Allende CV chondrite	174
Margarite [CaAl ₂ Al ₂ Si ₂ O ₁₀ (OH) ₂]	SA margarite	Aqueous alteration in the Allende CV chondrite	174
Greenalite [(Fe ²⁺ ,Mg) ₂ Si ₂ O ₇ (OH) ₄] and chrysotile [(Mg,Fe ²⁺) ₃ Si ₂ O ₇ (OH) ₃] ^a	SA greenalite	Aqueous alteration of carbonaceous chondrites	39,141,175–177
Cronstedtite [(Fe ²⁺ ,Fe ³⁺) ₃ (Si,Fe ³⁺) ₂ O ₇ (OH) ₄] ^a	SA cronstedtite	Aqueous alteration of carbonaceous chondrites	30,39,85,105,178
Berthierine [(Fe ²⁺ ,Mg,Fe ³⁺) ₃ (Al,Si) ₂ O ₇ (OH) ₄] and amesite [(Mg,Al) ₃ (Al,Si) ₂ O ₇ (OH) ₄] ^a	SA berthierine	Aqueous alteration of CM and CO chondrites	35,76,179–181
Talc [Mg ₃ Si ₄ O ₁₀ (OH) ₂]	SA talc	Aqueous alteration of carbonaceous chondrites	54,170
Saponite and ferrosaponite [(Ca,Na) _{0.5} (Mg,Fe ²⁺) ₃ (Si,Al) ₄ O ₁₀ (OH) ₂ ·4H ₂ O] ^a	SA saponite	Aqueous alteration of carbonaceous chondrites	110,120,173, 182–185
Montmorillonite [(Na,Ca) _{0.5} (Al,Mg) ₂ Si ₄ O ₁₀ (OH) ₂ ·nH ₂ O]	SA montmorillonite	Aqueous alteration of carbonaceous chondrites	3,180,186
Nontronite [Na _{0.5} Fe ³⁺ (Si,Al) ₄ O ₁₀ (OH) ₂ ·nH ₂ O]	SA nontronite	Aqueous alteration of carbonaceous chondrites	26
Vermiculite [Mg _{0.35} (Mg,Fe ³⁺ ,Al) ₃ (Si,Al) ₄ O ₁₀ (OH) ₂ ·4H ₂ O]	SA vermiculite	Aqueous alteration of carbonaceous chondrites	175,180
Clinochlore [Mg ₂ Al(AlSi ₃ O ₁₀)(OH) ₃]	SA clinochlore	Aqueous alteration of the Nagoya CM chondrite	175
Chamosite [(Fe ²⁺ ,Mg,Al,Fe ³⁺) ₄ (Si,Al) ₄ O ₁₀ (OH) ₃]	SA chamosite	Aqueous alteration of carbonaceous chondrites	76,86,187
Tochilinite-cronstedtite intergrowths (Fe-Ni-Si-S-O) ^a	SA TCI	Aqueous alteration of CM and other carbonaceous chondrites	39,85,87,188,189
Anorthite (CaAl ₂ Si ₂ O ₈) and albite (NaAlSi ₃ O ₈) ^a	SA plagioclase	Thermal metamorphism; glass devitrification	13,39,98,108, 154–156,190,191
Celsian (BaAl ₂ Si ₂ O ₈)	SA celsian	Metasomatically altered CAI in Allende	192
Nepheline [Na ₃ (Al ₃ Si ₄ O ₁₆) ^a]	SA nepheline	Thermal metamorphism, especially of CAIs	3,39,41, 155–157,164,193,194
Sodalite [Na ₄ (Si ₃ Al ₃)O ₁₂ ·Cl] ^a	SA sodalite	Thermal metamorphism of CAIs	3,39,130,148, 155,194,195
Marialite (Na ₄ Al ₃ Si ₃ O ₂₆ Cl)	SA marialite	Thermal metamorphism in an ordinary chondrite	196
Chabazite-Na [(Na,K)Al ₄ Si ₈ O ₂₄ ·11H ₂ O]	SA chabazite-Na	Aqueous alteration and heating in the Kaidun polymict breccia	32
Roedderite [(Na,K) ₂ Mg ₂ (Mg ₃ Si ₁₂)O ₃₀] and merrihueite [(K,Na) ₂ (Fe,Mg) ₂ Si ₂ O ₃₀]	SA roedderite	Thermal metamorphism of enstatite chondrites	197–199
Wilkinsonite [Na ₄ (Fe ²⁺ ,Fe ³⁺) ₄ (Si ₁₂ O ₃₆)]	SA wilkinsonite	Metasomatism in the Kaidun polymict breccia	32
Aenigmatite [Na ₄ (Fe ²⁺ ,Ti) ₂ O ₄ (Si ₁₂ O ₃₆)]	SA aenigmatite	Metasomatism, possibly of melilite	32,200
Gehlenite (Ca ₂ Al ₂ SiO ₇) and äkermanite (Ca ₂ MgSi ₂ O ₇)	SA melilite	Metasomatism with Ca-rich fluids	193
Indialite [Mg ₂ Al ₂ (AlSi ₃ O ₁₀) ₃]	SA indialite	Thermal metamorphism in carbonaceous chondrites	201,202
Dmisteinbergite (CaAl ₂ Si ₂ O ₈)	SA dmisteinbergite	Hydrothermal alteration of CAIs in NWA 2086 CV chondrite	203–205
Rankinite (Ca ₂ Si ₂ O ₇)	SA rankinite	Occurs with larnite in the Bali CV chondrite	192
Tilleyite [Ca ₂ Si ₂ O ₇ (CO ₃) ₂]	SA tilleyite	With secondary phases in a CAI from Allende CV chondrite	149
Organic minerals			
Whewellite [Ca(C ₂ O ₄)·H ₂ O]	SA whewellite	Reaction with carbonaceous matter at T < 480 °C	87
Kerogen (C,H,O,N,S) ^a	SA kerogen	Aqueous and thermal alteration in carbonaceous chondrites	20,22,85, 87,206–208

^a Volumetrically significant secondary asteroidal minerals.

References: 1 = Wlotzka (1972); 2 = Goodrich et al. (1987); 3 = Krot et al. (1995); 4 = Scott and Rajan (1981); 5 = Pederson (1999); 6 = Taylor et al. (1981); 7 = Kurat et al. (1989); 8 = Afiattalab and Wasson (1980); 9 = Rubin (1990); 10 = Hua et al. (1995); 11 = Rubin (1994); 12 = Fuchs (1966); 13 = Rubin and Kallemeyn (1994); 14 = Caillat Komorowski (2012); 15 = Schulze et al. (1994); 16 = Schulze (1998); 17 = Geiger and Bischoff (1995); 18 = Brearley (1996); 19 = Brearley (1990); 20 = Abrea and Brearley (2011); 21 = El Goresy et al. (1988); 22 = Piani et al. (2012); 23 = El Goresy et al. (2005); 24 = DuFresne and Anders (1962); 25 = Boström and Fredriksson (1966); 26 = Hutchison et al. (1987); 27 = Herndon and Rudee (1978); 28 = Rubin (1983); 29 = Endress et al. (1994); 30 = Palmer and Lauretta (2011); 31 = Ivanov et al. (2000); 32 = Zolensky and Ivanov (2003); 33 = Zolensky et al. (2008); 34 = Pratesi et al. (2006); 35 = Barber (1981); 36 = Rubin et al. (2002); 37 = Ma et al. (2011a); 38 = McCanta et al. (2008); 39 = Brearley and Jones (1998); 40 = Armstrong et al. (1985); 41 = Kojima et al. (1995); 42 = Kerridge et al. (1979a); 43 = Herndon et al. (1975); 44 = Harries and Zolensky (2016); 45 = Kallemeyn et al. (1994); 46 = El Goresy et al. (1979); 47 = Gomes and Keil (1980); 48 = Bischoff et al. (1994); 49 = Blum et al. (1989); 50 = Haggerty and McMahon (1979); 51 = Berger et al. (2016); 52 = Brearley and Prinz (1992); 53 = Bindi and Xie (2018); 54 = Ikeda (1992); 55 = McSween (1976); 56 = Kerridge et al. (1979b); 57 = Buchwald (1975); 58 = Prinz et al. (1994); 59 = Ivanova et al. (2019); 60 = Ma et al. (2011b); 61 = Scott (1988); 62 = Ulyanov (1981); 63 = Rambaldi et al. (1986a); 64 = El Goresy and Ehlers (1989); 65 = Britvin et al. (2008); 66 = Fegley and Post (1985); 67 = Fuchs and Blander (1977); 68 = Hutcheon et al. (1987); 69 = Blum et al. (1988); 70 = Nakamura-Messenger et al. (2012); 71 = Ma and Beckett (2018); 72 = Grossman et al. (1985); 73 = Lee and Greenwood (1994); 74 = Vacher et al. (2019); 75 = MacPherson and Davis (1994); 76 = Pignatelli et al. (2016); 77 = Connolly et al. (2006); 78 = Grady et al. (2015); 79 = Karwowski and Muszyński (2008); 80 = Litasov et al. (2018); 81 = Greenwood et al. (1994); 82 = Lee (1993); 83 = Brearley (1993a); 84 = Fuchs et al. (1973); 85 = Bunch and Chang (1980); 86 = Armstrong et al. (1982); 87 = El Goresy et al. (1984); 88 = Nagy and Alexander (1964); 89 = Richardson (1978); 90 = King and King (1981); 91 = Fredriksson and Kerridge (1988); 92 = Wlotzka and Wark (1982); 93 = Sylvester et al. (1993); 94 = Davis et al. (1991); 95 = Mao et al. (1990); 96 = Noguchi (1994); 97 = Weisberg et al. (1993); 98 = Ichikawa and Ikeda (1995); 99 = Riciputi et al. (1994); 100 = Endress and Bischoff (1996); 101 = Endress et al. (1996); 101 = Johnson and Prinz (1993); 102 = Tomeoka et al. (1989b); 103 = Müller et al. (1979); 104 = Rubin and Grossman (1985); 105 = Jones et al. (2014); 106 = Lewis and Jones (2016); 107 = Zanda et al. (1994); 108 = Keller et al. (1994); 109 = Ivanov et al. (2003); 110 = Kimura et al. (1992); 111 = Rubin (1991); 112 = Bischoff and Palme (1987); 113 = Grew et al. (2010); 114 = Olsen and Steele (1993); 115 = McCoy et al. (1994); 116 = Kimura and Ikeda (1995); 117 = Herndon et al. (1975);

(Caption continues on next page)

occurred before, during, and/or after the more gradual aqueous and thermal alterations, and solar heating near perihelion, notably for asteroids that crossed the orbit of Mercury and thus experienced cyclic near-surface temperatures exceeding 700 °C (Wittmann et al. 2011; Libourel et al. 2015).

Here we summarize the secondary mineralogy of chondrite and achondrite meteorites, including 166 historical natural kinds that have been ascribed to these secondary processes (Table 1). These varied secondary minerals encompass 169 IMA-approved species (some of which are lumped together), plus nine as yet unnamed and not fully characterized species and three amorphous or intergrown nanoscale phases. They collectively incorporate 41 different essential chemical elements (i.e., a defining element in one or more phases; Fig. 1), including the earliest known occurrences of minerals with essential Co, Ge, As, Nb, Ag, Sn, Te, Au, Hg, Pb, and Bi. Note, however, that Sc, Y, Rh, and Re, which are found as essential (if minor) elements in primary condensates of ultra-refractory inclusions in chondrite meteorites (Morrison and Hazen 2020), do not to our knowledge occur as essential elements in secondary asteroidal minerals.

Each mineral natural kind is given a binomial designation: here, the first name is “SA” (for “Secondary Asteroidal”) for all examples, whereas the second name in most instances conforms to the name of an approved IMA mineral species. However, in several cases we deviate from IMA nomenclature:

- We designate secondary α -(Fe,Ni) “SA iron” in conformity with IMA nomenclature. However, to avoid confusion, we often employ the common name “kamacite” for this Fe-Ni alloy. Similarly, whereas “enstatite” is the IMA-approved name for Mg-orthopyroxene, we designate this phase as “ortho-enstatite” for clarity.

- In several instances of isostructural mineral pairs with continuous solid solutions, we lump two end-member species. In most instances (except as noted parenthetically) we employ the name of the more prevalent end-member: erlichmanite-laurite, nuwaite-butianite, magnesite-siderite (*SA breunnerite*), sarcopsidite-chopinite, chladniite-johnsomervilleite, periclase-wüstite (*SA magnesiowüstite*), forsterite-fayalite (*SA olivine*), enstatite-ferrosilite (*SA orthopyroxene*), winchite-barrosite, greenalite-chrysolite, berthierine-amesite, saponite-ferrosaponite, albite-anorthite (*SA plagioclase*), roedderite-ferrihueite, and gehlenite-åkermanite (*SA melilite*).

- We recognize three amorphous and/or complexly intergrown phases: (1) *SA limonite*, including nanoscale intergrowths of iron

oxide/hydroxides; (2) “tochilinite-cronstedtite intergrowths” (*SA TCI*), which are common in CM chondrites; and (3) the amorphous carbonaceous phase, *SA kerogen*.

Native elements and metal alloys

Iron and nickel form the most abundant metal alloys in meteorites. In general, the fraction of metal in chondrites decreases with metamorphic grade from type 3 to type 6, including in chondrules and in the matrix, implying that kamacite and taenite are not generally secondary phases (Afiatlab and Wasson 1980). However, these alloys commonly display effects of secondary alteration; for example, equilibrated Fe-Ni alloys incorporate secondary exsolved phases such as graphite, chromite, phosphate, and silica owing to oxidation (Zanda et al. 1994). On the other hand, kamacite is described as a secondary phase in some ureilites, in which ferrous iron-bearing phases are reduced during thermal metamorphism in the presence of carbonaceous material (Wlotzka 1972; Goodrich et al. 1987; Rubin and Ma 2021).

Iron α -(Fe,Ni). Ni-poor *SA iron* (commonly known as “kamacite” in the meteoritics literature) occurs as metallic blebs in 10 to 100 μm thick reduced silicate rims in association with cohenite and troilite (Wlotzka 1972; Goodrich et al. 1987). Kamacite is thought to form when Fe^{2+} -bearing olivine or pyroxene grains are thermally metamorphosed in contact with carbonaceous material, hydrogen gas, or other reducing agent. Note that kamacite is well known to recrystallize during thermal metamorphism, causing rhythmic or concentric growth lines possibly owing to cyclic solar heating at perihelion, for example, in the Social Circle (IVA) and Indian Valley (IIAB) iron meteorites (Buchwald 1977; Wittmann et al. 2011). However, we do not consider this iso-mineral alteration a secondary occurrence.

Taenite γ -(Fe,Ni). Though not often explicitly reported as such, we suggest that *SA taenite* is one of several Ni-enriched alloys that forms when Fe in kamacite is preferentially oxidized to a Fe^{2+} -bearing phase such as magnetite, leaving a secondary alloy more enriched in Ni (e.g., Krot et al. 1995; Rubin and Ma 2021).

Tetrataenite (FeNi). Tetrataenite is a tetragonal (space group *P4/mmm*) variant of taenite with an ordered arrangement of Fe and Ni that forms from taenite during slow cooling or annealing. We consider tetrataenite to be primary when it forms by gradual cooling in the core of a differentiated planetesimal (e.g., Mittlefehldt et al. 1998). However, *SA tetrataenite* occurs as a secondary alloy phase when it occurs in an environment where a precursor metal phase has been oxidized and annealed,

TABLE 1.—CONTINUED

118 = Brearley (1995); 119 = Wlotzka (2005); 120 = Kessel et al. (2007); 121 = Buseck and Keil (1966); 122 = Kojima et al. (1995); 123 = Ma et al. (2016); 124 = Kimura and Ikeda (1992); 125 = Zinner et al. (1991); 126 = Steele (1995); 127 = Simon et al. (2001); 128 = Snetsinger and Keil (1969); 129 = Kornacki and Wood (1985); 130 = McGuire and Hashimoto (1989); 131 = Tomioka et al. (1992); 132 = Brearley (1993b); 133 = Krot et al. (1993); 134 = Ma et al. (2014); 135 = Ma and Beckett (2016); 136 = Ma (2010); 137 = El Goresy et al. (1978); 138 = Mackinnon (1980); 139 = Tomeoka and Buseck (1988); 140 = Keller and Buseck (1990b); 141 = Kimura et al. (2005); 142 = Tomeoka et al. (1989c); 143 = Akai (1990); 144 = Zolensky et al. (1991); 145 = Brigham et al. (1986); 146 = Wasson and Krot (1994); 147 = Hua and Buseck (1995); 148 = Kimura and Ikeda (1997); 149 = Ganino and Libourel (2017); 150 = Krot et al. (2020); 151 = Wark (1987); 152 = Greenwood et al. (1992); 153 = Fuchs (1974); 154 = Chaumard et al. (2014); 155 = MacPherson and Grossman (1984); 156 = Allen et al. (1978); 157 = Hashimoto and Grossman (1985); 158 = Fuchs (1971); 159 = Ma and Krot (2014); 160 = Simon and Grossman (1992); 161 = Delaney et al. (1984); 162 = Ma and Krot (2018); 163 = Ishii et al. (2010); 164 = Clayton et al. (1984); 165 = Zolensky et al. (1996); 166 = MacPherson et al. (1981); 167 = Sheng et al. (1991); 168 = Kimura and El Goresy (1989); 169 = Barber et al. (1984); 170 = Brearley (1997a); 171 = Dobrica and Brearley (2014); 172 = Kimura and Ikeda (1996); 173 = Tomeoka and Buseck (1990); 174 = Keller and Buseck (1991); 175 = Zolensky et al. (1993); 176 = Tomeoka et al. (1989a); 177 = Tomeoka and Buseck (1982a); 178 = Marrocchi et al. (2014); 179 = Ikeda (1983); 180 = Zolensky and McSween (1988); 181 = Richardson and McSween (1978); 182 = Alexander et al. (1989); 183 = Keller and Buseck (1990a); 184 = Lee et al. (1996); 185 = Cohen et al. (1983); 186 = Tomeoka and Buseck (1982b); 187 = Gooding (1985); 188 = Tomeoka and Buseck (1985); 189 = Vacher et al. (2019); 190 = Noguchi (1993); 191 = Dyl et al. (2012); 192 = A. Krot, personal communication; 193 = Wark et al. (1987); 194 = Tomeoka et al. (1992); 195 = Blander and Fuchs (1975); 196 = Alexander et al. (1987); 197 = Fuchs et al. (1966); 198 = Rambaldi et al. (1986b); 199 = Ikeda (1989); 200 = Wark (1986); 201 = Fuchs (1969); 202 = Mikouchi et al. (2016); 203 = Ma et al. (2013); 204 = Park et al. (2013); 205 = Fintor et al. (2014); 206 = Buseck and Hua (1993); 207 = Cody et al. (2008); 208 = Kebukawa et al. (2019).
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ELEMENTS IN SECONDARY ASTEROIDAL MINERALS

Major mineral-forming elements

Minor mineral-forming elements

1 H																	2 He
3 Li	4 Be											5 B	6 C	7 N	8 O	9 F	10 Ne
11 Na	12 Mg											13 Al	14 Si	15 P	16 S	17 Cl	18 Ar
19 K	20 Ca	21 Sc	22 Ti	23 V	24 Cr	25 Mn	26 Fe	27 Co	28 Ni	29 Cu	30 Zn	31 Ga	32 Ge	33 As	34 Se	35 Br	36 Kr
37 Rb	38 Sr	39 Y	40 Zr	41 Nb	42 Mo	43 Tc	44 Ru	45 Rh	46 Pd	47 Ag	48 Cd	49 In	50 Sn	51 Sb	52 Te	53 I	54 Xe
55 Cs	56 Ba	57 *La	72 Hf	73 Ta	74 W	75 Re	76 Os	77 Ir	78 Pt	79 Au	80 Hg	81 Tl	82 Pb	83 Bi	84 Po	85 At	86 Rn
87 Fr	88 Ra	89 #Ac	104 Rf	105 Db	106 Sg	107 Bh	108 Hs	109 Mt	110 Ds	111 Rg	112 Cn	113 Nh	114 Fl	115 Mc	116 Lv	117 Ts	118 Og

FIGURE 1. Secondary minerals from chondritic and nonchondritic meteorite parent bodies formed by aqueous alteration and thermal metamorphism primarily from 23 different essential elements that appear in 5 or more minerals, with important additional contributions from 18 minor elements that appear in fewer than 5 scarce phases. Included among these elements are the earliest known appearances of minerals with essential Co, Ge, As, Nb, Ag, Sn, Te, Au, Hg, Pb, and Bi. (Color online.)

for example, in CV chondrites, where it is found in association with awaruite, Fe-Ni sulfides, magnetite, and various silicates, as well as in CH, CO, and CR carbonaceous chondrites (Scott and Rajan 1981; Rubin and Ma 2021).

Awaruite (Ni₃Fe). *SA awaruite* forms via secondary processes in chondrules and chondrite matrices as Fe in Fe-Ni alloys is preferentially oxidized (Pederson 1999; Rubin and Ma 2021). Awaruite is found in the matrices of various chondrites, including ordinary chondrites (Taylor et al. 1981), as the dominant metal in so-called “dark inclusions” in CV chondrites in association with magnetite and minor pentlandite and merrillite (Kurat et al. 1989; Rubin 1991), and in R chondrites (Rubin and Kallemeyn 1989).

Wairauite (CoFe). Afittalab and Wasson (1980) and Rubin (1990) reported the cobalt-rich alloy *SA wairauite* with up to 33 wt% Co in association with a sulfide and high-Ni alloy in several ordinary chondrites. Hua et al. (1995) analyzed wairauite with 39 wt% Co, associated with troilite and pentlandite, in the matrix of the anomalous, oxidized Ningqing CK chondrite.

Copper (Cu). *SA copper* is a rare mineral in dark inclusions of CV chondrites, where it occurs in association with pentlandite and troilite (Kurat et al. 1989). Copper is a volumetrically minor but widespread phase in equilibrated ordinary chondrites (Rubin 1994), in the Hvittis enstatite chondrite in association with djerfisherite (Fuchs 1966), and in association with oxides and sulfides in the PCA 91002 R chondrite (Rubin and Kallemeyn 1994).

Mercury (Hg). Caillet Komorowski et al. (2012) describe an unusual association of *SA mercury* with cinnabar and copper sulfides in the Tieschitz unequilibrated ordinary chondrite. They suggest that the mercury minerals formed through sublimation in an asteroidal interior.

Platinum-iron Alloy (Pt,Fe). In highly oxidized Rumuruti chondrites, noble metals commonly form discrete micrometer-scale grains of *SA Pt-Fe alloy*, in association with telluride, arsenide, and stannide phases that represent ~0.0001 vol% of these meteorites. Schulze et al. (1994) and Schulze (1998)

investigated noble metal grains in several R chondrites, in which almost all noble metals are concentrated. Confirmed phases include Pt-Fe-dominant alloys (often the most abundant phase), Os-dominant alloys, and Au-dominant alloys, as well as chengbolite (PtTe₂), sperrylite (PtAs₂; at times Sn-rich), irarsite (IrAsS), niggliite (PtSn), rustenbergite (Pt₃Sn), erlichmanite (OsS₂), and laurite (RuS₂). These secondary phases often occur in association with Fe-Ni sulfides. Simon and Grossman (1992, their Table 2) examined PGE element exsolution from Fe-Ni alloys in opaque assemblages from the Leoville CV3 chondrite; they suggest that equilibration occurred in a post-accretionary environment at ~600 °C.

Osmium alloy (Os). *SA osmium alloy* occurs as a minor phase in micrometer-scale grains in highly oxidized Rumuruti chondrites, in association with Pt-Fe alloy, tellurides, and arsenides (Schulze et al. 1994; Schulze 1998).

Gold alloy (Au). Geiger and Bischoff (1995) report Au-dominated alloys in Acfer 217, which we designate *SA gold alloy*. Schulze et al. (1994) describe an Au-dominant alloy with minor Fe, Ni, and Pt from the Rumuruti R chondrite, while Schulze (1998) observed similar occurrences in other R chondrites.

Niggliite (PtSn). *SA niggliite* occurs as a minor phase in micrometer-scale grains in highly oxidized Rumuruti chondrites, in association with Pt-Fe alloy, tellurides, arsenides, and rustenbergite (Schulze et al. 1994; Schulze 1998).

Rustenbergite (Pt₃Sn). *SA rustenbergite* occurs as a minor phase in micrometer-scale grains in highly oxidized Rumuruti chondrites, in association with Pt-Fe alloy, tellurides, arsenides, and niggliite (Schulze et al. 1994; Schulze 1998).

Graphite (C). Graphite is one of many minerals that occur as both a primary and secondary phase in meteorites, at times with a degree of ambiguity regarding its paragenesis. Poorly graphitized carbon occurs in the matrices of CV chondrites (Brearley 1996). *SA graphite* occurs in opaque assemblages in association with kamacite in ordinary chondrites as a conse-

quence of annealing “poorly graphitized carbon” at temperatures above 300 °C (Brearley 1990; Abreu and Brearley 2011). In enstatite chondrites, secondary graphite occurs in association with kamacite, perhaps as exsolution from C-rich metal as a consequence of thermal metamorphism (El Goresy et al. 1988), or by metal-catalyzed graphitization of insoluble organic matter (Piani et al. 2012). Graphite is also a common phase in highly metamorphosed acapulcoites, lodranites, and winonaites (Benedix et al. 1998; El Goresy et al. 2005; McCoy et al. 2006).

Sulfur (S). *SA sulfur* has been reported to occur in the matrices of CI chondrites, most likely as an alteration product of pyrrhotite, with which it is often associated (DuFresne and Anders 1962; Boström and Fredriksson 1966).

Carbides

Cohenite [(Fe,Ni)₃C]. *SA cohenite* was reported by Hutchison et al. (1987) as a minor accessory phase, as veinlets associated with an Fe sulfide in the matrices of the Semarkona ordinary chondrite. Krot et al. (1997b) proposed that cohenite and haxonite in the carbide-magnetite assemblages of ordinary chondrites formed by reaction of Fe-Ni alloys with CO-bearing fluids, possibly through precipitation of Fe-Ni carbonyls. Secondary cohenite also occurs as exsolution lamellae in kamacite in enstatite chondrites as a consequence of thermal metamorphism (Herndon and Rudee 1978; Rubin 1983). In the LEW 88774 ureilite, thermally metamorphosed chromite grains display rims with Cr-rich cohenite, in association with breznaitite (Cr₃S₂) and eskolaite (Cr₂O₃)—reduced phases formed by reaction with carbonaceous material (Rubin and Ma 2021).

Haxonite [(Fe,Ni)₂₃C₆]. *SA haxonite* in association with cohenite was reported by Hutchison et al. (1987) as a likely minor accessory phase in the matrices of the Semarkona ordinary chondrite. Haxonite in the carbide-magnetite assemblages of ordinary chondrites may have formed by the reaction of Fe-Ni alloys with CO-bearing fluids (Krot et al. 1997b).

Phosphides

Schreibersite [(Fe,Ni)₃P]. Schreibersite occurs as both a primary and secondary phase in chondrites. A likely secondary occurrence of *SA schreibersite* is as a rare accessory phase in dark inclusions of CR chondrites (Endress et al. 1994). Palmer and Laurretta (2011) observed schreibersite as a common alteration product of kamacite in CM chondrites, in association with a P-bearing sulfide, tochilinite, and eskolaite.

Florenskyite (FeTiP). *SA florenskyite* was reported from Kaidun polymict breccia meteorite (Ivanov et al. 2000; Zolensky and Ivanov 2003), with micrometer-scale grains of average composition [Fe_{1.01}(Ti_{0.87}Ni_{0.13}Cr_{0.03}V_{0.02}Co_{0.01})(P_{0.97}Si_{0.03})]. These phosphide grains, found encased in serpentine, may have formed through exsolution from cooling metal, though a hydrothermal origin is also possible.

Andreyivanovite (FeCrP). *SA andreyivanovite* is the Cr-dominant isomorph of florenskyite, with which it coexists in the Kaidun polymict breccia. Micrometer-scale grains of average composition [Fe(Cr_{0.587}Fe_{0.15}V_{0.11}Ti_{0.08}Ni_{0.06})P] were described by Zolensky et al. (2008).

Melliinite [(Fe,Ni)₄P]. Pratesi et al. (2006) described *SA melliinite* in association with kamacite and nickel phosphide (the Ni

isomorph of schreibersite) from the NWA 1054 acapulcoite. Crystals to ~100 μm have average composition [(Ni_{2.30}Fe_{1.64}Co_{0.01})P_{1.05}].

Silicides

The origins of iron-nickel silicides in meteorites are enigmatic. We previously included suessite (Fe₃Si), carltonmooreite (Ni₃Si), and perryite [(Ni,Fe)₈(Si,P)₃] as primary minerals in differentiated asteroids (Morrison and Hazen 2021). However, Keil et al. (1982) suggested that suessite might have formed by reduction of Fe metal in the presence of carbonaceous material (a secondary process), though perhaps induced by a shock event. In any event, unless additional evidence is forthcoming, we do not recognize silicides as secondary asteroidal minerals.

Halides

Halides, including halite, sylvite, chlormayenite, and an unnamed bismuth chloride, are scarce aqueous alteration phases in chondrite meteorites. Keil (1968) reported an occurrence of the iron chloride lawrencite (FeCl₂) from enstatite chondrites; however, Rubin (1997) suggests that lawrencite is a product of terrestrial weathering. Droninoite [Ni₆Fe₂³⁺Cl₂(OH)₁₆·4H₂O], described by Chukanov et al. (2009) from the Dronino weathered iron meteorite, is also a halide formed by terrestrial weathering.

Halite (NaCl). *SA halite* was found as an accessory phase in the carbonaceous matrices of several ureilites (Berkley et al. 1978) and as submicrometer grains in the “waxy” organic-rich matrix of CM chondrites (Barber 1981). Striking dark blue to purple halite grains with fluid inclusions occur in euhedral crystals up to 0.5 cm diameter in the matrices of the Zag and Monahans (1998) ordinary chondrite breccias (Zolensky et al. 1999; Rubin et al. 2002). In Monahans, halite with up to ~1 mol% KCl occurs in association with minor sylvite. Of special note are 4.5 billion year old organic-rich brine inclusions in halite from Zag and Monahans (1998) (Zolensky et al. 1999; Chan et al. 2018).

Sylvite (KCl). Berkley et al. (1978) identified *SA sylvite* as an accessory phase in the carbonaceous matrices of several ureilites, and Barber (1981) recorded sylvite as sub-micrometer grains in the “waxy” organic-rich matrix of CM chondrites. Rubin et al. (2002, their Table 3) record sylvite of composition [(K_{0.81}Na_{0.17})Cl] from the Monahans (1998) ordinary chondrite as inclusions in more abundant halite.

Chlormayenite [Ca₁₂Al₁₄O₃₂(□₄Cl₂)]. Ma et al. (2011a) described a new alteration mineral, which they named brearleyite, from the NWA 1934 carbonaceous chondrite. However, this phase was subsequently recognized as equivalent to *SA chlormayenite*, which occurs in association with krotite in a refractory inclusion.

Unnamed (BiCl₃). McCanta et al. (2008) reported an unidentified bismuth chloride (provisionally *SA unnamed BiCl₃*) as a rare accessory phase in sub-micrometer grains in the R chondrite LAP 04840.

Sulfides

With at least 33 different natural kinds (Table 1), secondary asteroidal sulfides formed by oxidation and sulfidation of primary asteroidal phases are among the most diverse meteoritic minerals. Several of these phases, including [(V,Fe,Cr)₄S₅], [(Fe,Au,Co)₂S₃], and a Na-Cr-sulfide, are not yet fully described and are listed here provisionally. In addition, several research-

ers have identified an unknown P-bearing sulfide as inclusions in troilite or tochilinite in CM chondrites (Bunch and Chang 1980; Devouard and Buseck 1997; Nazarov et al. 2009; Palmer and Lauretta 2011). The composition of this phase or possibly mixture of phases is not yet known, though it is rich in Fe and Ni and incorporates minor O and Co. However, until a more complete description is available, we do not include this phase in our tabulation.

In addition, several meteoritic sulfide minerals are likely to have formed as a consequence of much later terrestrial weathering and have not been confirmed as pre-terrestrial secondary phases (Rubin and Ma 2021; A. Rubin, personal communication, 7 June 2020). These minerals include cronosite ($\text{Ca}_{0.2}\text{CrS}_2 \cdot 2\text{H}_2\text{O}$; Britvin et al. 2001), unnamed Cu-Cr-sulfide (Bevan et al. 2019), digenite ($\text{Cu}_{1.8}\text{S}$; Kimura et al. 1992), galena (PbS ; Nyström and Wickman 1991), mackinawite $[(\text{Fe},\text{Ni})_{1+x}\text{S}]$; Gomes and Keil 1980; Buseck 1968), schörlhornite ($\text{Na}_{0.3}\text{CrS}_2 \cdot \text{H}_2\text{O}$; Ivanov et al. 1996), and violarite (FeNi_5S_4 ; Chukanov et al. 2009), as well as two hydrous Na-Cr sulfides, designated A and B by El Goresy et al. (1988).

Troilite (FeS). Troilite, an important primary sulfide in a wide range of chondritic and achondritic meteorites, also occurs as an alteration phase. *SA troilite* is a product of secondary sulfidization in a range of chondritic environments, including the opaque assemblages known as Fremdlinge (Armstrong et al. 1985), with nepheline in CAIs of CO chondrites (Kojima et al. 1995), with exsolved pentlandite and pyrrhotite in the matrices of CM chondrites (Kerridge et al. 1979a; Brearley and Jones 1998, and references therein), and as a common phase in dark inclusions in association with pentlandite in CV chondrites (Kurat et al. 1989).

Pyrrhotite (Fe_7S_8). *SA pyrrhotite* is the most common sulfide in the matrices of CI chondrites, where it occurs in association with pentlandite and cubanite (Kerridge 1970; Kerridge et al. 1979b; Brearley and Prinz 1992). Herndon et al. (1975) suggested that pyrrhotite and coexisting magnetite formed from the oxidation of troilite at $T < 400$ °C, while Berger et al. (2016) demonstrated formation temperatures between 25 and 135 °C. Pyrrhotite, often in association with troilite and pentlandite, is a relatively common constituent of CM chondrite matrices (Kerridge et al. 1979a; Bunch and Chang 1980; Brearley 1995). Harries and Zolensky (2016) reported pyrrhotite from the Kaidun brecciated meteorite, including grains in the monoclinic 4C polytype that formed under extreme hydrothermal conditions at temperatures and pressures possibly as high as 300 °C at 85 bars water pressure, implying alteration deep within the parent body. Pyrrhotite and pentlandite occur in association with metal in the chondrules of CR chondrites (Kallemeyn et al. 1994), and it is a common mineral in the Fremdlinge of CV chondrites (El Goresy et al. 1979; Brearley and Jones 1998, and references therein).

Pyrite (FeS_2). *SA pyrite* is a minor secondary opaque phase in CK chondrites, where it occurs with various Fe-Ni-Cu sulfides (Geiger and Bischoff 1995). Pyrite in association with troilite, pyrrhotite, and pentlandite is also a minor mineral in R chondrites, (Bischoff et al. 1994; Rubin and Kallemeyn 1994). Gomes and Keil (1980) reported a Ni-rich variety of pyrite, known as “bravoite,” as a minor secondary phase that occurs in association with pentlandite, cubanite, and other secondary minerals in equilibrated ordinary chondrites.

Greigite (Fe_3S_4). El Goresy et al. (1988, their Table 16) reported *SA greigite* of near-ideal composition in association with smythite in both Na-Cr-rich clasts and low-temperature “sulfide patches” in altered EH chondrites.

Smythite (Fe_9S_{11}). *SA smythite* occurs in association with greigite as a sulfidation product in altered EH chondrites (El Goresy et al. 1988).

Millerite (NiS). *SA millerite* with ~5 wt% Fe is a rare secondary phase in the opaque assemblages of CK chondrites (Geiger and Bischoff 1995).

Heazlewoodite (Ni_3S_2). *SA heazlewoodite* is a minor component of Fremdlinge from CV chondrites (El Goresy et al. 1979; Blum et al. 1989). It also occurs as an opaque phase in carbonaceous chondrites in association with troilite, pyrrhotite, and pentlandite (McSween 1977; Haggerty and McMahon 1979).

Pentlandite ($[(\text{Ni},\text{Fe})_9\text{S}_8]$). *SA pentlandite* with a wide range of Ni/Fe (Brearley and Jones 1998; Berger et al. 2016, their Table 1 and Fig. 1) is a common secondary meteoritic sulfide in the opaque assemblages of carbonaceous chondrites, where it often is found with pyrrhotite (Haggerty and McMahon 1979). For example, pentlandite coexists with pyrrhotite and troilite in CM chondrites (Kerridge et al. 1979a; Bunch and Chang 1980; Brearley 1995); it occurs in association with pyrrhotite and cubanite in CI chondrite matrices (Kerridge et al. 1979b; Brearley and Prinz 1992)—an alteration that Berger et al. (2016) determined occurs in an aqueous environment at $T < 135$ °C; and it replaces troilite in chondrules in CO3 chondrites (Scott and Jones 1990). In CV chondrites, secondary pentlandite is found in Fremdlinge (El Goresy et al. 1978); it is the dominant sulfide in so-called “dark inclusions” in association with awaruite and troilite (Kurat et al. 1989); and it occurs in opaque assemblages in association with magnetite, awaruite, and minor merrillite (Rubin 1991).

Shenzhuangite (FeNiS_2). Bindi and Xie (2018) discovered the rare Fe-Ni isomorph of chalcopyrite, *SA shenzhuangite*, with the empirical formula $[(\text{Ni}_{0.7}^{2+}\text{Cu}_{0.3}^{+})(\text{Fe}_{0.7}^{2+}\text{Fe}_{0.3}^{3+})\text{S}_2]$ in the Suizhou L6 ordinary chondrite. They suggested that shenzhuangite is an alteration phase by sulfidation of taenite. This phase may be equivalent to the “Fe-Ni monosulfide,” i.e., $[(\text{Fe},\text{Ni})\text{S}]$, reported by El Goresy et al. (1979) from Fremdlinge of CV chondrites.

Covellite (CuS). El Goresy et al. (1988) observed *SA covellite* as an alteration product of djerfisherite in association with troilite, idaite, and bornite in enstatite chondrites.

Chalcopyrite (CuFeS_2). *SA chalcopyrite* was reported as a rare minor phase in the unusual Bench Crater carbonaceous chondrite, which was collected on the Moon’s surface by Apollo 12 (McSween 1976), and in the matrix of the unusual metamorphosed carbonaceous chondrite Yamato 82162 (Ikeda 1992). Chalcopyrite is also a rare secondary phase in the opaque assemblages of CK chondrites (Geiger and Bischoff 1995), as well as in R chondrites (Rubin and Kallemeyn 1994; Schulze et al. 1994).

Idaite (Cu_3FeS_4). *SA idaite* occurs as an alteration product of djerfisherite in association with troilite, bornite, and covellite in enstatite chondrites (El Goresy et al. 1988).

Bornite (Cu_5FeS_4). *SA bornite* is one of several sulfide minerals observed by El Goresy et al. (1988) as an alteration product of djerfisherite in enstatite chondrites.

Cubanite (CuFe₂S₃). Orthorhombic (*Pcnm*) *SA cubanite* and its cubic (*Fm3m*) isomorph *isocubanite* have been reported in association with pyrrhotite and pentlandite in the matrices of CI chondrites (Kerridge et al. 1979b), and as a minor secondary phase in equilibrated ordinary chondrites (Gomes and Keil 1980). Berger et al. (2015) determined that the presence of cubanite indicates low-temperature aqueous alteration at < ~200 °C.

Isocubanite (CuFe₂S₃). *SA isocubanite*, also referred to as “Cu-rich pyrrhotite” by Buchwald (1975), is reported to occur in the matrices of CI chondrites (Kerridge et al. 1979b). However, we cannot confirm that this phase is distinct from cubanite.

Brezinaite (Cr₃S₄). Prinz et al. (1994) reported *SA brezinaite* in the LEW 88774 monomict Cr-rich ureilite, occurring as rims on thermally metamorphosed chromite grains in association with Cr-rich cohenite and eskolaite—reduced phases formed by reaction with carbonaceous material (Rubin and Ma 2021). V-rich brezinaite [(Cr_{2.05}V_{0.62}Fe_{0.33})₃S₄], in association with V-rich daubréelite [Fe(Cr,V)₂S₄] and a new V-rich sulfide [(V,Fe,Cr)₄S₅], was identified in the CBa chondrite Sierra Gorda 013 by Ivanova et al. (2019).

Murchisite (Cr₂S₆). Ma et al. (2011b) found a new sulfide with empirical composition [(Cr_{4.6}V_{0.1}Fe_{0.1})S₆] occurring as subhedral to rounded grains to 4 μm diameter in the Murchison CM chondrite. *SA murchisite* evidently transformed at relatively low-temperature (≤300 °C) from a Cr-S phase that exsolved from an iron alloy at a higher temperature.

Daubréelite (FeCr₂S₄). Daubréelite, which is a common primary phase in enstatite chondrites, aubrites, and iron meteorites (Morrison and Hazen 2021), also occurs as a secondary phase (*SA daubréelite*) in chondrites (Scott 1988; Brearley and Jones 1998). Ivanova et al. (2019) described a V-rich daubréelite in association with V-rich brezinaite [(Cr,V,Fe)₃S₄] and a new V-rich sulfide [(V,Fe,Cr)₄S₅] in the CBa chondrite Sierra Gorda 013. In addition, Ulyanov (1991) reported a Cu-rich variety of secondary daubréelite.

Sphalerite [(Zn,Fe)S]. Rambaldi et al. (1986a) reported the occurrence of sphalerite with composition [(Zn_{0.7}Fe_{0.3})S] from millimeter-scale metal-sulfide nodules in the Qingzhen EH3 enstatite chondrite that appear to have been reheated and remelted. This relatively Zn-rich example coexists with a second population of Ga-bearing rudashevskyite with the more typical meteoritic Fe/Zn ~1.5 to 1.7 (see below).

Rudashevskyite [(Fe,Zn)S]. Most meteoritic “sphalerite,” both primary and secondary, is Fe-rich, typically with Zn-Fe zoning and at times with exsolution lamellae of troilite. Examples have been described from several EH enstatite chondrites (El Goresy and Ehlers 1989; Britvin et al. 2008; Rubin and Ma 2021). Britvin et al. (2008) documented the Fe-dominant analog of sphalerite, *SA rudashevskyite*, as a presumably secondary matrix phase in the Indarch enstatite chondrite, in which they found the average composition of 31 grains (5 to 120 μm maximum dimension) to be [(Fe_{0.61}Zn_{0.35}Mn_{0.04}Cu_{0.01})S], with a range from 54 to 69 mol% FeS end-member. A continuous Zn-Fe solid solution from the ZnS end-member to >60 mol% exists (Barton and Toulmin 1966). However, the majority of secondary meteoritic examples appear to fall in the Fe-rich range from ~40 to ~70 mol% FeS—occurrences that we name *SA rudashevskyite*. Rambaldi et al. (1986a) recorded unusual Ga-rich examples

(56 to 63 mol% FeS; 2.1 to 3.7 wt% Ga) from millimeter-scale reheated and remelted metal-sulfide nodules in the Qingzhen EH3 enstatite chondrite.

Unnamed [(V,Fe,Cr)₄S₅]. *SA unnamed [(V,Fe,Cr)₄S₅]* with empirical formula [(V_{1.55}Fe_{1.52}Cr_{0.92})S₅] was described by Ivanova et al. (2019) as a secondary phase in association with V-rich daubréelite and V-rich brezinaite [(Cr,V,Fe)₃S₄] in the CBa chondrite Sierra Gorda 013.

Molybdenite (MoS₂). *SA molybdenite* was observed in altered CAIs in the Allende CV carbonaceous chondrite (Fegley and Post 1985; Fuchs and Blander 1977), as well as in opaque assemblages known as “Fremdlinge” in CV chondrites (El Goresy et al. 1978; Hutcheon et al. 1987). Molybdenite is thought to have formed by secondary oxidation/sulfidation of refractory metals (Blum et al. 1988).

Wassonite (WS). Nakamura-Messenger et al. (2012) reported wassonite (we assume secondary *SA wassonite*, though a primary origin cannot be ruled out) with empirical formula [(Ti_{0.93}Fe_{0.06}Cr_{0.01})S] in the mesostasis of a barred olivine chondrite in association with forsterite, enstatite, Fe-Ni metal, and other sulfides from the Yamato 691 enstatite chondrite.

Cinnabar (HgS). Caillet Komorowski et al. (2012) describe an unusual association of cinnabar with native mercury and copper sulfides in the Tieschitz unequilibrated ordinary chondrite. They suggest that the mercury minerals formed through sublimation in an asteroidal interior. Cinnabar occurs as ~5 μm diameter grains in troilite-pentlandite rims surrounding the troilite-rich cores of dark inclusions from the Allende CV chondrite (Kurat et al. 1989).

Erlichmanite (OsS₂) and laurite (RuS₂). Both Ru-dominant (to 85 mol%) and Os-dominant (to 75 mol%) disulfides, in some instances with significant Ir (to 20 wt%), as well as Pt, Fe, and Ni, occur as minor phases in opaque assemblages of CK carbonaceous chondrites (Geiger and Bischoff 1995, their Table 6). We lump these cubic (space group *Pa3*) PGE disulfides into *SA erlichmanite*, because Os often appears to be the dominant element. Micrometer-scale grains of Os-Ru-Ir-Pt disulfide also occur in highly oxidized Rumuruti chondrites, in association with Pt-Fe alloy, tellurides, and arsenides (Schulze et al. 1994; Schulze 1998). This phase appears similar to an unidentified Os-Ru-Fe sulfide in the Acfer 217 R chondrite (Bischoff et al. 1994).

Cooperite (PtS₂). Geiger and Bischoff (1995, their Table 7) reported the tetragonal Pt-dominant disulfide, *SA cooperite*, with ~6 wt% Fe and/or Ni as micrometer-scale grains in opaque assemblages of the ALH 82135 and EET 87519 CK carbonaceous chondrites.

Petrowskaite [(Au,Fe,Ag)₂S]. Geiger and Bischoff (1995, their Table 7) described a single occurrence of an Au-Fe-Ag sulfide [(Au_{0.65}Fe_{0.25}Ag_{0.07})₂S] that appears to be related to petrowskaite (ideally AuAgS) in an opaque assemblage of the LEW 87009 CK carbonaceous chondrites. We provisionally name this phase *SA petrowskaite*.

Unnamed [(Fe,Au,Co)₂S₃]. Geiger and Bischoff (1995, their Table 7) reported *SA unnamed [(Fe,Au,Co)₂S₃]* of composition [(Fe_{0.71}Au_{0.25}Co_{0.05})₂S₃] in opaque assemblages of the EET 87514 CK carbonaceous chondrites.

Nuwaite (Ni₆GeS₂) and butianite (Ni₆SnS₂). Ma and Beckett (2018) reported micrometer-scale grains of Ni-Ge-Sn

sulfides associated with grossular, melilite, heazlewoodite, and Fe-Ni alloys, and filling cracks in igneous diopside, as a vapor-deposited alteration product in CAIs of the Allende CV carbonaceous chondrite. They called a grain with empirical formula $[(\text{Ni}_{5.95}\text{Fe}_{0.16})(\text{Ge}_{0.60}\text{Sn}_{0.23})(\text{S}_{1.72}\text{Te}_{0.33})]$ nuwaite, whereas a grain with empirical formula $[(\text{Ni}_{5.93}\text{Fe}_{0.13})(\text{Sn}_{0.52}\text{Ge}_{0.41})(\text{S}_{1.56}\text{Te}_{0.45})]$ (i.e., Sn > Ge) was called butianite. We lump these isostructural phases into *SA nuwaite*, because both minerals have sub-equal amounts of Ge (41 to 60 mol%) and Sn (23 to 52 mol%), evidently in a continuous solid solution, as well as significant Te substitution for S (up to 23 mol%).

Djerfisherite $[\text{K}_6(\text{Fe,Cu,Ni})_{25}\text{S}_{26}\text{Cl}]$. The alkali Cu-Fe sulfide *SA djerfisherite* is a minor phase in enstatite chondrites (Fuchs 1966; Rambaldi et al. 1986a; El Goresy et al. 1988). Unlike most other unusual sulfides in enstatite chondrites, djerfisherite always occurs external to sulfide-rich chondrules (Grossman et al. 1985), and it is always a secondary alteration mineral (A. Rubin, personal communication, 7 June 2020).

Unnamed NaCr-sulfides. El Goresy et al. (1988, their Tables 13 and 14) described at least two new, but as yet not fully characterized, Na-Cr hydrated layer structure sulfides, which are thought to be aqueous alteration products of caswellsilverite (NaCrS_2), from the unequilibrated Yamato 691 EH enstatite chondrite. One of these phases is hydrated $[(\text{Na,Cu}^+)\text{CrS}_2]$, and therefore similar in composition to caswellsilverite, while the other is described as a hydrated “Na-Cu-Zn-Cr-sulfide.” Both are listed below. Two additional hydrated Na-Cr-sulfides, one lower in Na and higher in Cr and S than caswellsilverite, the other with low analytical totals (~71 wt% total) and thus perhaps significantly hydrated, were observed by El Goresy et al. (1988, their Table 15) in the Qingzhan enstatite chondrite but were not sufficiently characterized to list here.

Unnamed hydrated $[(\text{Na,Cu}^+)\text{CrS}_2]$. *SA unnamed hydrated $[(\text{Na,Cu}^+)\text{CrS}_2]$* is an alteration phase of caswellsilverite in the Yamato 691 enstatite chondrite (El Goresy et al. 1988, their Table 13). Six analyzed grains display a range of Na/(Na+Cu) from 0.48 to 0.96, while minor Fe and Zn substitute for Cr.

Unnamed hydrated (Na-Cu-Zn-Cr-sulfide). *SA unnamed hydrated (Na-Cu-Zn-Cr-sulfide)* is an alteration phase of caswellsilverite in the Yamato 691 enstatite chondrite (El Goresy et al. 1988, their Table 14). Four analyses reveal well-constrained Cr (~33 wt%) and S (~43 wt%), but significant variations in Na (2 to 13 wt%), Zn (6 to 10 wt%), Cu (4 to 8 wt%), and Fe (0.7 to 4.7 wt%).

Tochilinite $\{6(\text{Fe}_0.9\text{S})\cdot 5[(\text{Mg,Fe})(\text{OH})_2]\}$. The tochilinite group of minerals are layered phases with alternating ~5 Å brucite-type $[\text{Mg}(\text{OH})_2]$ and ~6 Å mackinawite $[(\text{Fe,Ni})_{1-x}\text{S}]$ layers, including Fe-dominant tochilinite as well as closely related haapalaite (Buseck and Hua 1993) and vallerite (Ackermann and Rasse 1973), which are Ni- and Cu-bearing phases approved as $\{2[(\text{Fe,Ni})\text{S}]\cdot 1.61[(\text{Mg,Fe})(\text{OH})_2]\}$ and $\{2[(\text{Fe,Cu})\text{S}]\cdot 1.53[(\text{Mg,Al})(\text{OH})_2]\}$, respectively. However, we suggest that members of the tochilinite group are better represented as a single natural kind—a complex mixed-layer solid solution: $\{2[(\text{Fe,Mg,Cu,Ni},\square)]\text{S}\cdot 1.57\text{--}1.85[(\text{Mg,Fe,Ni,Al,Ca})(\text{OH})_2]\}$ (Rubin and Ma 2021). We lump all of these occurrences as *SA tochilinite*.

Tochilinite is a relatively common phase in altered CAIs, altered chondrules, and matrices of CM chondrites, where it may

occur in association with sulfides and schreibersite (MacPherson and Davis 1994; Lee and Greenwood 1994; Palmer and Lauretta 2011). Vacher et al. (2019) point to alteration of kamacite to tochilinite, which commences between 120 and 160 °C. Palmer and Lauretta (2011) suggest that small tochilinite grains in the matrices of CM chondrites with comparatively low P, Ni, and Co may have formed by sulfidation of magnetite. Even in the least altered CM chondrites, tochilinite occurs intergrown with cronstedtite as rims replacing kamacite (Pignatelli et al. 2016).

A variety of fine-grained layered phases in altered meteorites, once termed “poorly characterized phases” or PCPs (e.g., Barber et al. 1983), are more properly referred to as tochilinite-cronstedtite intergrowths or “TCI” (e.g., Vacher et al. 2019). TCIs occur in multiple morphotypes, including needle-like enrolled layers and as undulating layers with a dominant 10.8 Å spacing (Barber et al. 1983). Mackinnon and Zolensky (1984) proposed that some examples of TCI are predominantly tochilinite, but the tochilinite story is complicated by its common occurrence as layered intergrowths with the serpentine group mineral cronstedtite (Nakamura and Nakamura 1996), at times in locally ordered 17 or 24 Å repeats, representing 1:1 and 2:1 serpentine:tochilinite sequences, respectively (Zolensky et al. 1993; Mackinnon and Zolensky 1984; Tomeoka and Buseck 1985; see below).

Arsenides

Arsenides are rare alteration phases in chondritic meteorites. For example, El Goresy et al. (1978) reported possible Pt-Os-Ru-Fe arsenides with PGE alloys in CAIs of the Allende and Leoville carbonaceous chondrites; Schulze et al. (1994) identified sperrylite (PtAs_2) and irarsite (IrAsS) with significant Pt, Rh, and Ru substitution in R chondrites; and Geiger and Bischoff (1995, their Table 7) described an arsenide-sulfide phase similar to irarsite from extremely oxidized CK chondrites. These varied phases appear to represent a complex solid solution among compositions corresponding to sperrylite, iridarsenite $[(\text{Ir,Ru})\text{As}_2]$, löllingite (FeAs_2), omeite (OsAs_2), anduoite (RuAs_2), and possibly their sulfide isomorphs. A complication arises because at least three distinct structure types exist: (1) sperrylite and irarsite have the cubic pyrite structure (space group *Pa3*); (2) anduoite, löllingite, and omeite, as well as the related minerals rammelsbergite (NiAs_2), ruarsite (RuAsS), and safflorite (CoAs_2), have the orthorhombic marcasite structure (space group *Pnmm*); and (3) iridarsenite is reported to be monoclinic (space group *P2₁/c*). However, no structural information is available for meteoritic arsenides. Until additional information about compositions, structures, and phase relationships is forthcoming, we assign these minerals to two secondary arsenides for which chemical analyses are available: *SA sperrylite* and *SA irarsite*. In addition to these secondary meteorite minerals, Nyström and Wickman (1991) ascribe several meteoritic arsenides to terrestrial weathering processes, including cobaltite (CoAsS), gersdorffite (NiAsS), maucherite ($\text{Ni}_{11}\text{As}_8$), nickeline (NiAs), orcelite ($\text{Ni}_{5-x}\text{As}_2$), rammelsbergite (NiAs_2), and safflorite (CoAs_2).

Sperrylite (PtAs_2). *SA sperrylite*, at times Sn-rich, occurs as a minor phase in micrometer-scale grains in highly oxidized Rumuruti chondrites, in association with Pt-Fe alloy, tellurides, stannides, and irarsite (Schulze et al. 1994; Schulze 1998).

Irarsite (IrAsS). *SA irarsite*, with significant Pt, Rh, and Ru

contents, occurs as a minor phase in micrometer-scale grains in highly oxidized Rumuruti chondrites, in association with Pt-Fe alloy, tellurides, stannides, and sperrylite (Schulze et al. 1994; Schulze 1998). In addition, Geiger and Bischoff (1995, their Table 7) described a single grain of an Ir-As-S phase, presumably irarsite, of composition $[(\text{Ir}_{0.71}\text{Pt}_{0.19}\text{Fe}_{0.08}\text{Os}_{0.07})\text{As}_{1.10}\text{S}_{0.85}]$ from the EET 87860 CK chondrite.

Tellurides

Moncheite $[(\text{Pt},\text{Pd})(\text{Te},\text{Bi})_2]$. *SA moncheite* occurs as a minor phase in micrometer-scale grains in highly oxidized Rumuruti chondrites, in association with Pt-Fe alloy, tellurides, and arsenides (Schulze et al. 1994; Schulze 1998). Moncheite has also been reported from altered opaque assemblages in CK carbonaceous chondrites (Geiger and Bischoff 1995, their Table 7; Connolly et al. 2006; Grady et al. 2015). Note that the name “chengbolite,” at times used for a Bi-poor variety of moncheite, is not an approved IMA species.

Unnamed Au-Pt-Fe telluride. Geiger and Bischoff (1995) identified several tellurides in the altered opaque assemblages of CK carbonaceous chondrites, including *SA unnamed Au-Pt-Fe telluride*, which may be a solid solution among krennerite (Au_3AgTe_8), calaverite (AuTe_2), chengbolite (PtTe_2), and frobergite (FeTe_2).

Altaite (PbTe) . Karwowski and Muszyński (2008) and Litasov et al. (2018) report rare occurrences of *SA altaite* from the Morasko and Masyanino IAB iron meteorites. Litasov et al. (2018) offer alternative possible modes of origin, including primary crystallization from a highly differentiated melt, post-magmatic alteration, or terrestrial weathering. We provisionally adopt a secondary origin as more consistent with other sulfide, arsenide, and telluride meteoritic minerals.

Sulfates

Sulfates are an important component of the altered matrices of CI chondrites, in which they can exceed 10 vol% as both veins and isolated grains (Boström and Fredriksson 1966; Richardson 1978; Fredriksson and Kerridge 1988; Johnson and Prinz 1993; Endress and Bischoff 1996). Uncertainty exists regarding the timing of sulfate formation: Gounelle and Zolensky (2001) observed the formation of sulfate veins and efflorescences by reaction with the atmosphere during museum storage of CII carbonaceous chondrites and they suggest that all reports of meteoritic sulfates may represent terrestrial weathering. With that caveat, we tentatively tabulate nine alkali, alkaline earth, and Fe sulfates that might be products of secondary aqueous alteration and oxidation on a parent body.

In addition, Gooding (1981) and Gooding et al. (1991) described meteoritic hydrous sulfates produced by terrestrial weathering, including coquimbite $[\text{Fe}_2(\text{SO}_4)_3 \cdot 9\text{H}_2\text{O}]$, kieserite $(\text{MgSO}_4 \cdot \text{H}_2\text{O})$, slavikite $[\text{NaMg}_2\text{Fe}_2(\text{SO}_4)_7(\text{OH})_6 \cdot 33\text{H}_2\text{O}]$, szomolnokite $(\text{FeSO}_4 \cdot \text{H}_2\text{O})$, and voltaite $[\text{K}_2\text{Fe}_3\text{Al}(\text{SO}_4)_{12} \cdot 18\text{H}_2\text{O}]$. Other terrestrial weathering minerals include starkeyite $(\text{MgSO}_4 \cdot 4\text{H}_2\text{O})$; Zolensky and Gooding (1986) and paraotwayite $[\text{Ni}(\text{OH})_{2-x}(\text{SO}_4, \text{CO}_3)_{0.5x}]$; Zubkova et al. 2008]. Uncertain examples that we suspect are terrestrial weathering products include honessite $[(\text{Ni},\text{Fe})_8\text{SO}_4(\text{OH})_{16} \cdot n\text{H}_2\text{O}]$; Buchwald 1977], jarosite $[\text{KFe}_3(\text{SO}_4)_2(\text{OH})_6]$; Buchwald 1977], melanterite

$(\text{FeSO}_4 \cdot 7\text{H}_2\text{O})$; Ulyanov 1991), and schwertmannite $[\text{Fe}_6^{3+}(\text{OH},\text{SO}_4)_{12-13}\text{O}_{16} \cdot 10\text{H}_2\text{O}]$; Pederson 1999].

Anhydrite (CaSO_4) . Greenwood et al. (1994) report *SA anhydrite* intergrown with bassanite and in association with calcite from altered CAIs in the Cold Bokkeveld CM carbonaceous chondrite. Anhydrite also occurs as a minor phase in matrices of CM chondrites (Fuchs et al. 1973; Lee 1993). Brearley (1993a) reports fibrous vein-filling anhydrite (2 μm maximum dimension) in the matrix of the ALH A77307 CO chondrite.

Gypsum $(\text{CaSO}_4 \cdot 2\text{H}_2\text{O})$. *SA gypsum* is an alteration product of melilite in CAIs and refractory inclusions of CM chondrites (Bunch and Chang 1980; Armstrong et al. 1982; MacPherson et al. 1983; El Goresy et al. 1984). Gypsum is a rare phase in the matrices of CI chondrites, in which it occurs in veins and as individual grains (DuFresne and Anders 1962; Nagy and Andersen 1964; Fuchs et al. 1973; Richardson 1978).

Bassanite $(\text{CaSO}_4 \cdot 0.5\text{H}_2\text{O})$. *SA bassanite* (also referred to as “hemihydrate”) intergrown with anhydrite was reported by Greenwood et al. (1994) from altered CAIs in CM Cold Bokkeveld. Bassanite also occurs in veins with anhydrite in association with calcite in the matrices of CM chondrites (Lee 1993).

Hexahydrate $(\text{MgSO}_4 \cdot 6\text{H}_2\text{O})$. DuFresne and Anders (1962) and Richardson (1978) reported *SA hexahydrate* from the matrices of CI chondrites, in which it may be a terrestrial weathering phase (Gounelle and Zolensky 2001).

Epsomite $(\text{MgSO}_4 \cdot 7\text{H}_2\text{O})$. *SA epsomite* occurs as veins and grains in the matrices of CI chondrites, where it may be associated with hexahydrate (DuFresne and Anders 1962; Richardson 1978).

Thenardite (Na_2SO_4) . *SA thenardite* is a rare secondary sulfate in meteorites, tentatively recorded by King and King (1981) from chondrule rims in the Murray CM2 carbonaceous chondrite.

Unnamed Mg-Al-Fe sulfate. Lee and Greenwood (1994, their Table 4) detected a secondary Mg-Al-Fe sulfate with an approximate empirical formula (exclusive of OH/H₂O) of $[(\text{Na}_{0.4}\text{Mg}_{4.0}\text{Al}_{2.4}\text{Fe}_{0.9}\text{S}_{0.9})\text{O}_{12}]$ in altered CAIs of the Murray CM2 carbonaceous chondrite. Lee and Greenwood (1994) tentatively identified this phase as aluminocopiapite $[(\text{Al},\text{Mg})\text{Fe}_3^+(\text{SO}_4)_6(\text{OH},\text{O})_2 \cdot 20\text{H}_2\text{O}]$; however, the element ratios are inconsistent with this identification. Until more information is forthcoming, we designate this phase as *SA unnamed Mg-Al-Fe sulfate*.

Blödite $[\text{Na}_2(\text{Mg},\text{Ni})(\text{SO}_4)_2 \cdot 4\text{H}_2\text{O}]$. The matrices of some CI chondrites contain the hydrous Na-Mg-sulfate *SA blödite* (DuFresne and Anders 1962; Boström and Fredriksson 1966; Fredriksson and Kerridge 1988), including Ni-rich examples in the Ivuna CI chondrite that approach the composition of nickelblödite, though with $\text{Mg}/(\text{Mg}+\text{Ni}) = 0.67$ (Fredriksson and Kerridge 1988, their Table 3).

Barite (BaSO_4) . Wlotzka and Wark (1982) reported *SA barite* in altered CAIs of the Leoville CV chondrite, where it occurs in veins associated with Ba-rich feldspar. Kurat et al. (1989) recorded barite as a rare accessory phase in dark inclusions from the Allende CV carbonaceous chondrite, where it is associated with unusual sulfide-andradite objects that contain native copper, Cu- and Ti-rich magnetite, perovskite, and calcite.

Carbonates

Carbonates are important low-temperature ($0 < T < 130$ °C) aqueous alteration phases in the matrices of carbonaceous

chondrites, where they occur in association with phosphates, sulfates, sulfides, and magnetite, and they are minor secondary constituents of ordinary chondrites (Alexander et al. 2015). They represent on average ~5 vol% of CI chondrites, occurring in matrix clasts up to several millimeters in maximum dimension (Nagy and Andersen 1964; Fredriksson and Kerridge 1988; Johnson and Prinz 1993; Endress and Bischoff 1996; Endress et al. 1996). In matrices of CM chondrites, ubiquitous carbonate minerals are commonly intimately intergrown with clay minerals (Zolensky and McSween 1988). Fujiya et al. (2012) employed ^{53}Mn - ^{53}Cr dating of calcite to determine that carbonates in CM chondrites formed within 7 Ma of the formation of CAIs (4567.3 \pm 0.16 Ma; Connelly et al. 2012).

In addition, several carbonates have been reported as terrestrial alteration products in meteorites, including barringtonite ($\text{MgCO}_3 \cdot 2\text{H}_2\text{O}$; Ulyanov 1991), chukanovite [$\text{FeCO}_3(\text{OH})_2$; Pekov et al. 2007], hydromagnesite [$\text{Mg}_5(\text{CO}_3)_4(\text{OH})_2 \cdot 4\text{H}_2\text{O}$; Zolensky and Gooding 1986; Velbel 1988], nesquehonite [$\text{Mg}(\text{HCO}_3)(\text{OH}) \cdot 2\text{H}_2\text{O}$; Zolensky and Gooding 1986; Velbel 1988], nyerereite [$\text{Na}_2\text{Ca}(\text{CO}_3)_2$; Ulyanov 1991], reevesite [$\text{Ni}_6\text{Fe}_2(\text{CO}_3)(\text{OH})_{14} \cdot 4\text{H}_2\text{O}$; Buchwald 1977], and vaterite (CaCO_3 ; Okada et al. 1981).

Rhombohedral carbonates [(Ca,Mg,Fe,Mn)CO₃]. End-member rhombohedral carbonates (space group $R\bar{3}c$ or $R\bar{3}$) include calcite (Ca), magnesite (Mg), siderite (Fe), and rhodochrosite (Mn), whereas intermediate cation-ordered species include dolomite (CaMg), ankerite (CaFe), and kutnohorite (CaMn). In addition, the intermediate Mg-Fe carbonate, though not an approved species, is often termed “breunnerite” in the meteoritics literature.

Calcite (CaCO₃). *SA calcite* has been observed in numerous carbonaceous and ordinary chondrites. In CV chondrites, calcite occurs with Al-diopside and sodalite (Sylvester et al. 1993), as veins in forsterite (Davis et al. 1991), and as irregular masses (Mao et al. 1990) in the Vigarano CV chondrite; as the dominant secondary phase in the Leoville CV chondrite (Mao et al. 1990); and with Ca-phosphate in CAIs of in the Coolidge CV4 metamorphosed carbonaceous chondrite (Noguchi 1994). Calcite is also a common alteration product in the CAIs of CM chondrites (Armstrong et al. 1982; Lee and Greenwood 1994; MacPherson and Davis 1994), and it is the most common carbonate in the matrices of CM chondrites (Fuchs et al. 1973; Bunch and Chang 1980; Barber 1981; Tomeoka et al. 1989a; Brearley 1995), as well as in CK chondrites (Noguchi 1993). Calcite of near end-member composition is a minor phase in the matrix of CI chondrites (Fredriksson and Kerridge 1988; Johnson and Prinz 1993; Endress and Bischoff 1996), and it occurs in CR chondrites, including as a component of matrices (Weisberg et al. 1993; Ichikawa and Ikeda 1995; Alexander et al. 2015). Near end-member calcite occurs as an important secondary phase in the matrices of the Semarkona ordinary chondrite, where it is found in association with smectite (Hutchison et al. 1987).

Dolomite [CaMg(CO₃)₂]. *SA dolomite* is the most common carbonate in the matrices of CI chondrites, where it coexists with breunnerite and calcite (Fredriksson and Kerridge 1988). Endress and Bischoff (1996) reported a range of compositions [$(\text{Ca}_{0.35-0.53}\text{Mg}_{0.34-0.51}\text{Mn}_{0.00-0.15}\text{Fe}_{0.02-0.13})\text{CO}_3$] (Brearley and Jones 1998, Figs. 130 and 131), which underscores the extensive solid

solution possible in this system. Trace element compositions point to crystallization from brines that are analogous to those of terrestrial deposits (Riciputi et al. 1994). Dolomite is much less common than calcite in the matrices of CM chondrites (Johnson and Prinz 1993). Rubin et al. (2007) suggest that dolomite has replaced calcite in the most altered examples.

Magnesite (MgCO₃) and siderite (FeCO₃). *SA breunnerite*, a solid solution between magnesite and siderite (and thus sometimes referred to as “ferroan magnesite”), is a common secondary phase in the matrix of CI chondrites, where it coexists with dolomite (Fredriksson and Kerridge 1988; Johnson and Prinz 1993; Endress and Bischoff 1996; Endress et al. 1996). Fredriksson and Kerridge (1988) reported a compositional range of [$(\text{Ca}_{0.00-0.05}\text{Mg}_{0.53-0.78}\text{Mn}_{0.02-0.15}\text{Fe}_{0.15-0.39})\text{CO}_3$] (see Brearley and Jones 1998, their Fig. 132). Breunnerite with Mg/(Mg+Fe) from 0.73 to 0.77 dominates the carbonate mineralogy of Yamato 82162, an unusual highly metamorphosed carbonaceous chondrite (Tomeoka et al. 1989b, their Table 5).

Rhodochrosite (MnCO₃). Ikeda (1992) reported a single occurrence of the Mn carbonate, *SA rhodochrosite*, in the highly metamorphosed carbonaceous chondrite Yamato 82162; however, further compositional information was not provided. MnCO_3 is also a common component of Ca-, Mg-, and Fe-rich carbonates and the Yamato 82162 occurrence may be closer to an intermediate variety. For example, Tomeoka et al. (1989b, their Table 5) report an Mn-rich carbonate with empirical formula [$(\text{Mg}_{0.61}\text{Fe}_{0.18}\text{Mn}_{0.21})\text{CO}_3$] from Yamato 82162. In addition, Zolensky and McSween (1988) list the Ca-Mn carbonate, kutnohorite, as a secondary meteorite phase though, again, complete analyses were not provided.

Aragonite (CaCO₃). *SA aragonite* was identified by electron diffraction in the matrices of CM chondrites, in which it coexists with more abundant calcite (Müller et al. 1979; Barber 1981). In most instances, structural investigation of calcium carbonate is not undertaken; therefore, aragonite may be more abundant than reported (M. Zolensky, personal communication, 5 October 2020).

Phosphates

Several chondritic phosphate minerals, notably chlorapatite and merrillite, occur as secondary phases that formed through aqueous alteration and/or thermal metamorphism of prior P-bearing phases (e.g., Rubin and Grossman 1985; Jones et al. 2014). In addition, phosphates that are the likely result of terrestrial weathering include arupite [$\text{Ni}_3(\text{PO}_4)_2 \cdot 8\text{H}_2\text{O}$; Buchwald 1977], collinsite [$\text{Ca}_2(\text{Mg,Fe,Ni})(\text{PO}_4)_2 \cdot 2\text{H}_2\text{O}$; Buchwald 1977], lipscombite [$(\text{Fe,Mn})\text{Fe}_2(\text{PO}_4)_2(\text{OH})_2$; Buchwald 1977], monazite [$(\text{Ce,L,Th})\text{PO}_4$; Yagi et al. 1978], and vivianite [$\text{Fe}_3(\text{PO}_4)_2 \cdot 8\text{H}_2\text{O}$; Buchwald 1977].

Apatite group [Ca₅(PO₄)₃(Cl,F,OH)]. Apatite group minerals, which are commonly encountered as both primary and secondary meteorite phases, commonly display complex solid solution, most notably among Cl, F, and OH. Reports indicate that Cl-dominant examples are abundant, whereas F- and OH-rich apatite occurs only in restricted settings. Therefore, we recognize three different natural kinds.

Chlorapatite [Ca₅(PO₄)₃Cl]. *SA chlorapatite* commonly occurs in association with merrillite as a product of aqueous

alteration or thermal metamorphism of P-bearing Fe-Ni alloys in various meteorites. In ordinary chondrites, chlorapatite is a common minor phase (Jones et al. 2014), including in troilite-chlorapatite and metal-chlorapatite assemblages in the matrices of unequilibrated ordinary chondrites (Ahrens 1970; Rubin and Grossman 1985), as well as in equilibrated ordinary chondrites of types 3.6 to 6, in which phosphates [with Cl/(Cl+F) 0.65 to 0.87] and coexisting plagioclase may have experienced metasomatism by alkali-halogen-bearing fluids during metamorphism (Zanda et al. 1994; Lewis and Jones 2016, their Table 3). Chlorapatite in ordinary chondrites typically incorporates up to ~30 mol% of the F-bearing fluorapatite component (Jones et al. 2014, their Table 2 and Fig. 7). Chlorapatite occurs in R chondrites with merrillite, ilmenite, Cu metal, and phyllosilicates (Bischoff et al. 1994; Rubin and Kallemeyn 1994; Kallemeyn et al. 1996), as well as in altered CAIs of the Allende CV carbonaceous chondrite and in sulfide-phosphate assemblages in the matrices of CV chondrites (Armstrong et al. 1985; Blum et al. 1989).

Hydroxylapatite [Ca₅(PO₄)₃OH]. *SA hydroxylapatite* lacking measurable Cl or F has been reported from the matrix of the Cochabamba CM chondrite (Müller et al. 1979), as well as in the Bali CV3 chondrite (Keller et al. 1994). Note, however, that apatite group minerals in ordinary chondrites rarely exceed 400 ppm H₂O (Jones et al. 2014; their Fig. 8).

Fluorapatite [Ca₅(PO₄)₃F]. Kimura et al. (1992, their Table 4) analyzed *SA fluorapatite* with F/(F+Cl) = 0.93 in association with diopside from the Yamato 75305 winonaite. Ivanov et al. (2003, their Table 3) reported *SA fluorapatite* in association with arfvedsonite, aenigmatite, and wilkinsonite as inclusions in albite crystals from unusual alkaline and subalkaline clasts from the Kaidun polymict breccia; note, however, that F/Cl was not reported.

Merrillite [Ca₉NaMg(PO₄)₇]. *SA merrillite*, commonly in association with chlorapatite, occurs as a product of thermal metamorphism of P-bearing Fe-Ni alloys in ordinary chondrites (Ahrens 1970; Jones et al. 2014; Lewis and Jones 2016), in which merrillite is the primary host of rare earth elements (Crozas and Zinner 1985; Jones et al. 2014). Meteoritic merrillite in ordinary chondrites is typically Mg-dominant, but it incorporates up to 12 mol% of the Fe²⁺ ferromerrillite component (Jones et al. 2014, their Table 2). Abundant, though volumetrically minor, sulfide-merrillite assemblages are found in CV3 carbonaceous chondrites (Rubin and Grossman 1985), who propose that phosphate minerals formed by reaction of schreibersite with Ca, O, and Cl from silicates. Merrillite also occurs as a minor phase in opaque assemblages of CV chondrites in association with magnetite, awaruite, and pentlandite (Rubin 1991). The schreibersite formed previously by exsolution from P-rich Fe-Ni alloys that condensed from nebular gas. A Ca-rich phosphate, either merrillite or whitlockite (see below), occurs in the matrices of CI chondrites (Nagy and Andersen 1964; Boström and Fredriksson 1966).

Whitlockite [Ca₉Mg(PO₃OH)(PO₄)₆]. Some confusion exists in the meteorite literature regarding occurrences of merrillite vs. whitlockite, both of which are valid Ca-Mg phosphate species of the whitlockite group according to the IMA. While we are unable to resolve this issue, we suspect that anhydrous merrillite may form during thermal metamorphism, whereas

SA whitlockite is a plausible product of aqueous/hydrothermal alteration. Hutcheon et al. (1987) and Bischoff and Palme (1987) found whitlockite in a sulfide-rich Fremdlinge of Allende CAIs. A Ca-rich phosphate, either whitlockite or merrillite, occurs in the matrices of CI chondrites (Nagy and Andersen 1964; Boström and Fredriksson 1966).

Sarcopsidite [Fe³⁺(PO₄)₂] and chopinite [Mg₃(PO₄)₂]. Grew et al. (2010) reported secondary Mg-Fe-Mn orthophosphates of the chopinite-sarcopsidite solid solution, i.e., [(Fe,Mg,Mn)₃(PO₄)₂], in association with farringtonite from the altered GRA 95209 acapulcoite. Analyzed grains span the range from the 1 to 89 mol% Mg end-member, with up to 23 mol% zavalaiite component [Mn₃(PO₄)₂] in the more Fe-rich samples. Until more compositional data on these secondary phosphates are available, we lump them into *SA sarcopsidite*, because they are all members of a continuous solid solution and most samples are Fe-dominant. Grew et al. (2010) suggest that these phosphates formed by oxidation of P-rich metal, with subsequent exchange of Mg and Mn for Fe. They also note the possible influence of modest impact pressures, because chopinite typically forms at pressures above 0.4 GPa (Brunet et al. 1998).

Farringtonite [(Mg,Fe)₃(PO₄)₂]. Grew et al. (2010) reported *SA farringtonite*, a polymorph of chopinite, in association with grains of the chopinite-sarcopsidite solid solution in the altered GRA 95209 acapulcoite. They observe 6 to 14 mol% of the graftonite component [Fe₃(PO₄)₂] and suggest that phosphate formed by oxidation of P-rich metal, with subsequent replacement of Fe by Mg from silicates. Note that Grew et al. (2010) did not observe graftonite, the Fe-rich isomorph of farringtonite, from which it is separated by a miscibility gap.

Chladniite [Na₂CaMg₇(PO₄)₆] and johnsomervilleite [Na₁₀Ca₆Mg₁₈Fe₂₅(PO₄)₃₆]. Minerals of the chladniite-johnsomervilleite solid solution are rare phases in meteorites, in which they occur as both primary phosphates in iron meteorites (Olsen and Steele 1993; McCoy et al. 1994) and presumably secondary phases, for example, in the GRA 95209 acapulcoite (Grew et al. 2010). Grew et al. (2010, their Fig. 7) reported a range of compositions, almost all of which have Mg > Fe and thus lie in the chladniite field. Therefore, we designate these occurrences as *SA chladniite*.

Brianite [Na₂CaMg(PO₄)₂]. *SA brianite* is one of several minor secondary phases reported by Kimura and Ikeda (1995) in opaque awaruite-sulfide-magnetite spherules in association with whitlockite and apatite in the Allende CV carbonaceous chondrite. However, no compositional data were provided.

Oxides

Oxidation plays a significant role in the secondary alteration of meteorites. Consequently, we tabulate 23 species of secondary oxides, 10 of which are also known as primary meteorite minerals. It is often difficult to differentiate these secondary minerals from recently formed terrestrial weathering products. Until more information is available, we ascribe the following meteorite phases to terrestrial alteration: Ca-armalcolite (CaTi₂O₅; Lin and Kimura 1996), cuprite (Cu₂O; Ulyanov 1991), hematite (Fe₂O₃; Buchwald 1977), olkhonskite (Cr₂Ti₃O₉; Schmitz et al. 2016), pseudobrookite (Fe₂TiO₅; Ramdohr 1973; Krot et al. 1993), thorianite (ThO₂; MacPherson et al. 1988), and trevorite

($\text{NiFe}_3^+\text{O}_4$; Buchwald 1977).

Oxide spinel group [(Mg,Fe²⁺,Zn)(Al,Fe³⁺,Cr,V,Ti)₂O₄].

Several members of the oxide spinel group have been reported as secondary phases in ordinary and carbonaceous chondrites. Extensive solid solution occurs in these phases (e.g., El Goresy 1976; Brearley and Jones 1998, their Table A3.39; Kessel et al. 2007, their Table 2); nevertheless, the following six secondary minerals appear to represent distinct compositional regimes in meteorites.

Magnetite (Fe₃O₄). *SA magnetite* is a common alteration phase in carbonaceous chondrites, in which it may form by oxidation of kamacite or troilite (e.g., Herndon et al. 1975; Krot et al. 1995; Zolensky et al. 2010; Palmer and Lauretta 2011; Chan et al. 2016). Magnetite is the most abundant oxide in the matrices of CI chondrites, where it occurs as spherules, framboids, and platelets in aggregates to a few tens of micrometers in maximum dimension (Brearley and Jones 1998, and references therein). Zolensky and Ivanov (2003) document secondary magnetite formed by oxidation of pyrrhotite in the highly altered Kaidun polymict breccia. Magnetite is found as a minor phase in the matrices of many CM chondrites (Fuchs et al. 1973; Bunch and Chang 1980; Barber 1981; Brearley 1995), in which it is an alteration product of kamacite from microenvironments with limited S and Si (Palmer and Lauretta 2011). However, magnetite in association with interlayered saponite/serpentine is a major matrix phase in the Bells CM2 chondrite (Brearley 1995). Magnetite is also an alteration product of kamacite in the chondrules of CO3 meteorites (Scott and Jones 1990) and in CK chondrites (Geiger and Bischoff 1995). Secondary magnetite occurs in CV chondrites (Krot et al. 1995), both in the fine-grained matrix, at times with framboidal morphology (Keller et al. 1994), and in opaque assemblages with awaruite and pentlandite (Rubin 1991; Lee et al. 1996). McSween (1977) divided CV3 chondrites into reduced and oxidized subgroups based on their different metal-to-magnetite ratios. Secondary magnetite often deviates from ideal Fe₃O₄ (Brearley and Jones 1998, and references therein); for example, V-rich magnetite with up to 10 mol% coulsonite (FeV₂O₄) component was reported from Fremdlinge in Allende (Armstrong et al. 1985, their Table 2; Hutcheon et al. 1987). Dunn et al. (2016) studied magnetite minor element compositions in oxidized CK and CV chondrites and documented systematic changes in the contents of Mg, Ni, Al, Cr, and Ti with metamorphic grade.

Chromite (Fe²⁺Cr₂O₄). Cr-rich oxide spinels, both chromite and a range of Cr-bearing phases (Wlotzka 2005; Hazen et al. 2021), are common primary phases in the full range of equilibrated and unequilibrated ordinary chondrites (Bunch et al. 1967; Dodd 1969; Brearley and Jones 1998, their Fig. 191; Wlotzka 2005; Kessel et al. 2007). With increasing metamorphic grade, the abundance of chromite increases significantly, revealing that chromite also forms by secondary processes during thermal metamorphism. Chromite in equilibrated ordinary chondrites typically incorporates Cr/(Cr+Al) ~ 0.85 to 0.90 and Fe/(Fe+Mg) ~ 0.73 to 0.92 (Wlotzka 2005, their Table 1; Kessel et al. 2007). Wlotzka (2005) found that chromite compositions become more uniform in Mg-Fe with increasing petrologic type while retaining heterogeneity in Cr/(Cr+Al). Note, however, that a second more heterogeneous population of Cr-bearing oxide spinels in metamorphosed ordinary chondrites displays a wide range of compo-

sitions in the spinel-hercynite-chromite-magnesiocromite solid solution field (Wlotzka 2005, their Table 2).

SA chromite occurs as exsolution lamellae in ilmenite in thermally metamorphosed OC meteorites (Buseck and Keil 1966), as exsolution lamellae in Fe-Ni metal in metamorphosed ordinary and carbonaceous chondrites (Zanda et al. 1994), and in the matrices of CM chondrites (Fuchs et al. 1973; Barber 1981).

Spinel (MgAl₂O₄). *SA spinel* with minor Fe, Cr, V, and Ti occurs in the matrices of CM chondrites (Barber 1981). Spinel with significant hercynite and minor gahnite (ZnAl₂O₄) components exsolves from magnetite in thermally metamorphosed CK carbonaceous chondrites (Geiger and Bischoff 1995, their Table 2). Spinel is also a minor phase that occurs as irregular regions up to 15 μm maximum dimension precipitated from aqueous fluids in grossular-rich veins in CAIs from the Allende CV chondrite (A. Krot, personal communication, 11 October 2020).

Hercynite (FeAl₂O₄). *SA hercynite*, at times with minor Mg, Cr, and Ti, is an important indicator mineral for thermal metamorphism in the matrices of CI and CM chondrites (Barber 1981; Tonui et al. 2014). Armstrong et al. (1985, their Table 7) identified an unusual V-rich spinel in Fremdlinge of the Allende CV chondrite, where it occurs in association with V-rich magnetite and fassaite. They record a typical composition as [(Mg_{0.44}Fe_{0.56})(Al_{1.26}V_{0.56}Cr_{0.14}Ti_{0.02}Si_{0.01})O₄], which is closest to the hercynite end-member (56 mol%), but with significant components of magnesiocoulsonite (MgV₂O₄; 28 mol%), magnesiocromite (MgCr₂O₄; 7 mol%), and spinel (7 mol%). In addition, secondary hercynite that replaces grossite and krotite in a CAI from Yamato 81020 contains up to 10 wt% ZnO (M. Zolensky, personal communication, 5 October 2020).

Ulvöspinel (Fe₂⁺TiO₄). Kojima et al. (1995) report an occurrence of *SA ulvöpsinel* in association with ilmenite replacing perovskite in an altered CAI from CO3 chondrites. However, no compositional information is given.

Coulsonite [(Fe,Mg)V₂O₄]. Vanadium-rich oxide spinels have long been recognized as secondary meteorite minerals. As noted above, Armstrong et al. (1985, their Tables 2 and 7) recorded both V-rich magnetite with up to 10 mol% coulsonite component and V-rich hercynite with 28 mol% magnesiocoulsonite in Fremdlings of the Allende CV carbonaceous chondrite. However, Ma et al. (2016) documented the first confirmed example of *SA coulsonite*, with 60 mol% coulsonite, 27 mol% hercynite, and 12 mol% spinel [(Fe_{0.87}Mg_{0.12})(V_{1.19}Al_{0.81})O₄; C. Ma, personal communication, 28 July 2020]. *SA coulsonite* occurs in a V-rich CAI from the Allende CV chondrite in association with beckettite and other secondary minerals.

Other oxides

Periclase (MgO) and wüstite (FeO). “Magnesiowüstite” is a commonly employed, though unapproved, name for intermediate oxide phases from the periclase-wüstite (MgO-FeO) solid-solution series. *SA magnesiowüstite* with composition (Mg_{0.70}Fe_{0.30}) occurs in Fremdlinge from the Vigarano CV carbonaceous chondrite (Zinner et al. 1991). Magnesiowüstite with Mg/(Mg+Fe) from 0.55 to 0.75 in association with carbonates was identified in the Yamato 82162 (Ikeda 1992) and Yamato 86029 (Tonui et al. 2014) altered CI carbonaceous chondrites. Kimura and Ikeda (1992, their Table 4) analyzed samples with

Mg/(Mg+Fe) from 0.38 to 0.40 in the Belgica 7904 carbonaceous chondrite. In addition, we have added end-member MgO, *CAI periclase*, to our list of primary condensates (see Addendum, forthcoming).

Corundum (Al₂O₃). Corundum occurs uncommonly as a primary phase in the CAIs of carbonaceous chondrites (Morrison and Hazen 2020). In addition, Steele (1995) and Simon et al. (2001) reported *SA corundum* in association with nepheline in the Allende CV carbonaceous chondrite—a consequence of open-system alteration of primary melilite (M. Zolensky, personal communication, 5 October 2020).

Maghemite [(Fe_{0.67}□_{0.33})Fe₂³⁺O₄]. *SA maghemite* has been reported as a minor accessory phase associated with smectite in the altered matrices of the Semarkona unequilibrated ordinary chondrite (Hutchison et al. 1987).

Ilmenite (FeTiO₃). *SA ilmenite* in association with chromite is a common accessory phase in thermally metamorphosed ordinary chondrites (Snetsinger and Keil 1969; Rubin and Ma 2021). Ilmenite also occurs as a product of metasomatism of CAIs in CV carbonaceous chondrites (Krot et al. 1995); replacing perovskite in CAIs from CV (Kornacki and Wood 1985; McGuire and Hashimoto 1989), including Mg-rich varieties (11 to 28 mol% MgTiO₃; Steele 1995); in CAIs of CO chondrites as replacement of perovskite, sometimes as rims on perovskite (Tomeoka et al. 1992; Kojima et al. 1995); and exsolved from Cr-bearing magnetite in thermally metamorphosed CK carbonaceous chondrites (Noguchi 1993; Geiger and Bischoff 1995).

Eskolaite (Ti₂O₃). Palmer and Lauretta (2011) observed *SA eskolaite* as a common alteration product of kamacite in CM chondrites, in association with sulfides, tochilinite, and schreibersite. In the LEW 88774 ureilite, thermally metamorphosed chromite grains display rims with eskolaite in association with Cr-rich cohenite and breznaitite—reduced phases formed by reaction with carbonaceous material (Rubin and Ma 2021). Kimura and Ikeda (1992, their Table 4) detected eskolaite as 3 μm thick secondary rims on “Cr-rich ovoids” in magnesian chondrules of Belgica 7904 carbonaceous chondrite. Prinz et al. (1994) described aluminous eskolaite with average composition [(Cr_{0.74}Al_{0.23}Ti_{0.03})₂O₃] from the LEW 88774 monomict Cr-rich ureilite. They ascribed this occurrence to thermal equilibration deep within the ureilite parent body, followed by excavation by an impact event and re-equilibration.

Rutile (TiO₂). *SA rutile* occurs as a minor secondary phase in thermally metamorphosed ordinary chondrites in association with ilmenite and chromite, at times as exsolution lamellae in ilmenite (Buseck and Keil 1966; Rubin and Ma 2021). Brearley (1993b) recorded TiO₂ (presumably secondary rutile) in association with the highly reduced and enigmatic Magnéli phases, Ti₅O₉, and Ti₈O₁₅, in the altered matrix of the Bells CM2 chondrite.

Pyrophanite (MnTiO₃). Krot et al. (1993) reported an occurrence of near end-member *SA pyrophanite* in the Raguli ordinary chondrite, in association with ilmenite and baddeleyite, possibly formed through metamorphism on the parent body.

Scheelite (CaWO₄). Armstrong et al. (1985, their Table 5) reported the first known occurrences of meteorite phases from the powellite-scheelite solid solution as a minor phase in an unusual Fremdling from the Allende CV chondrite. W-rich

examples of *SA scheelite* with intergrown V-rich magnetite display a range of W/(W+Mo) from 0.87 to 0.96.

Powellite (CaMoO₄). Bischoff and Palme (1987, their Table 6) analyzed samples of the powellite-scheelite solid solution spanning the range (W_{0.56}Mo_{0.44}) to (W_{0.21}Mo_{0.79}) from Fremdlinge of the Allende CV chondrite. Because Mo-rich examples are more commonly encountered, we ascribe all such occurrences to *SA powellite*. Bischoff and Palme (1987) suggest formation by oxidation of refractory metal alloys.

Unnamed Mg-Fe molybdate. Armstrong et al. (1985, their Table 6) reported an as yet undescribed secondary Mg-Fe molybdate phase, which we provisionally call *SA unnamed Mg-Fe molybdate*. It occurs as a minor phase with V-rich fassaite, V-rich magnetite, Ni-Fe metal, Fe-Ni sulfide, and several minor phases in an unusual Fremdling from the Allende CV chondrite.

Kamiokite [(Fe,Mg)₂Mo₃O₈]. *SA kamiokite* with composition [(Fe_{1.56}Mg_{0.32}Ca_{0.07}Ni_{0.07})Mo₃O₈] was reported by Ma et al. (2014) as an alteration phase of primary monipite (MoNiP) in a CAI from the Allende CV meteorite.

Majindeite [(Mg,Fe)₂Mo₃O₈]. Ma and Beckett (2016) described *SA majindeite* [(Mg_{1.57}Fe_{0.43})Mo₃O₈], the Mg-dominant isomorph of kamiokite, from CAIs in the Allende CV chondrite. Majindeite, which occurs as sub-micrometer crystals in association with Fe-Ni and PGE alloys, apatite, and an unnamed Nb-rich oxide (see below), is thought to have formed during subsolidus oxidation of a Mo-rich precursor, possibly secondary kamiokaite. Note that it is also possible that the kamiokite-majindeite solid solution is continuous and represents a single natural kind of secondary Mo oxides.

Tugarinovite (MoO₂). Ma et al. (2014) identified *SA tugarinovite* as one of several alteration phases of monipite (MoNiP) in a CAI from the Allende CV meteorite.

Beckettite (Ca₂V₆Al₆O₂₀). Ma et al. (2016) reported a new alteration phase, *SA beckettite*, in a V-rich CAI from the Allende CV meteorite. A member of the sapphirine group, beckettite was found as 4 to 8 μm diameter crystals with secondary coulsonite, grossular, anorthite, hercynite, and corundum. The empirical formula is [Ca_{2.0}(V_{3.5}Al_{1.4}Ti_{0.6}⁴⁺Mg_{0.25}Sc_{0.1})(Al_{5.7}Si_{0.3})O₂₀].

Unnamed [(Nb,V,Fe)O₂]. An as yet undescribed cubic Nb-rich oxide, *SA unnamed [(Nb,V,Fe)O₂]*, with 46 wt% Nb₂O₅ and empirical formula [(Nb_{0.54}V_{0.27}Fe_{0.15}Mg_{0.05}Al_{0.04})O₂] was suggested by Ma et al. (2014) to be one of several alteration phases of primary monipite (MoNiP) from a CAI in the Allende CV carbonaceous chondrite.

Chihuahuaite [(Fe,Mg)Al₂O₉]. *SA chihuahuaite*, originally “hibonite-(Fe)” but subsequently renamed for the Mexican state in which the host Allende CV meteorite was found, was discovered by Ma (2010) in altered CAIs, in which it is associated with nepheline, ilmenite, hercynite, and “Fe²⁺-rich spinel” (i.e., an Fe-bearing aluminous spinel with Mg > Fe²⁺). Meteoritic chihuahuaite with Fe/(Fe+Mg) ~0.6, which coexists with near end-member hibonite [Fe/(Fe+Mg) ~ 0.06], is thought to have formed by Fe-alkali metasomatism of primary hibonite.

Baddeleyite (ZrO₂). Baddeleyite, which is a rare primary phase in the refractory inclusions of chondrites as well as in eucrites and aubrites, has also been described as a secondary phase. *SA baddeleyite* occurs in the Fremdlinge of Allende and Leoville CV carbonaceous chondrites (El Goresy et al. 1978)

and exsolved from ilmenite in association with pyrophanite in the Raguli (H3.8) ordinary chondrite (Krot et al. 1993).

Hydroxides

Several Fe and/or Mg hydroxides appear to be secondary meteorite minerals. A greater number of meteoritic hydroxides, not included in Table 1, are likely terrestrial weathering products: akagenéite [$\text{FeO}(\text{OH},\text{Cl})$; Buchwald 1977], böhmite [$\text{AlO}(\text{OH})$; Bevan et al. 2019], chlormagaluminite [$\text{Mg}_4\text{Al}_2(\text{OH})_{12}\text{Cl}_2 \cdot 3\text{H}_2\text{O}$; Ivanova et al. 2016], feroxhyte [$\text{Fe}^{3+}\text{O}(\text{OH})$; Buseck and Hua 1993]; goethite and lepidocrocite [both $\text{FeO}(\text{OH})$; Buchwald 1977; Noguchi 1994; Karwowski et al. 2015], hibbingite [$\text{Fe}_2(\text{OH})_3\text{Cl}$; Saini-Eidukat et al. 1994], hollandite [$\text{Ba}(\text{Mn}^{4+}\text{Mn}^{2+})\text{O}_{16}$; Ulyanov 1991], portlandite [$\text{Ca}(\text{OH})_2$; Okada et al. 1981], and pyrochlore [$(\text{Na},\text{Ca})_2\text{Nb}_2\text{O}_6(\text{OH},\text{F})$; Lovering et al. 1979].

Brucite [$\text{Mg}(\text{OH})_2$]. *SA brucite* of near end-member composition occurs as micrometer-scale grains in the matrices of the Orgueil CI carbonaceous chondrite (Boström and Fredriksson 1966) and in the Murchison and Mighei CM chondrites (Mackinnon 1980).

Amakinite [$\text{Fe}(\text{OH})_2$]. *SA amakinite*, the Fe^{2+} isomorph of brucite with <10 mol% $\text{Mg}(\text{OH})_2$, occurs in association with tochilinite and cronstedtite (replacing olivine and pyroxene) in CM chondrites (Pignatelli et al. 2016). Note that amakinite, which is likely a consequence of alteration by Fe-rich fluids, occurs in the alternating hydroxide-sulfide layers of Fe-rich ferrotuchilinite (Zolensky and McSween 1988).

Ferrihydrite [$\text{Fe}_2\text{O}_3 \cdot n\text{H}_2\text{O}$]. *SA ferrihydrite* has been reported from the fine-grained matrices of CV chondrites (Lee et al. 1996) as <8 nm diameter grains associated with interlayered serpentine and saponite in the Orgueil CI chondrite (Tomeoka and Buseck 1988). Keller and Buseck (1990b) found that ferrihydrite in the matrices of CO chondrites occurs both in granular masses that are probably pseudomorphs after framboidal magnetite, and in finely dispersed grains in a matrix, likely formed from the Fe in olivine and/or metal. Zolensky et al. (1993) suggest that meteoritic ferrihydrite in CI chondrites forms by oxidation of Fe^{2+} -bearing saponite, possibly as a terrestrial weathering phase.

Limonite [$\text{FeO}(\text{OH}) \cdot n\text{H}_2\text{O}$]. “Limonite” is a commonly used, though unapproved, name for fine-grained, mixed phase alteration products of Fe-bearing minerals—typically an intimate amorphous mixture of nanoscale iron oxides/hydroxides (e.g., goethite and hematite), often with intermixed clay minerals (<https://www.mindat.org/>, accessed 28 July 2020). Boström and Fredriksson (1966) reported *SA limonite* from the Orgueil CI chondrite matrix, which they ascribed to aqueous alteration on the meteorite’s parent body. Note, however, that others (Gounelle and Zolensky 2001; M. Zolensky, personal communication, 5 October 2020) suggest that all goethite-bearing assemblages formed through terrestrial weathering.

Silicates

Silicates, notably olivine, pyroxene, and feldspar group species, are major primary and secondary minerals in meteorites. As with other minerals, the distinction between primary and secondary silicates is not always obvious. Ideally, primary minerals formed by direct condensation from a vapor phase, by

crystallization from a cooling melt, or through solid-state reactions (e.g., reconstructive phase transitions, exsolution, element diffusion, or order/disorder transitions) during initial cooling. By contrast, secondary silicates formed by aqueous alteration, metasomatism, and/or thermal metamorphism of prior phases, therefore resulting in a new combination of chemical composition and atomic structure, in some instances accompanied by a change in oxidation state. Note, however, that many phases, including olivine, pyroxene, and feldspar group minerals, undergo gradual changes during aqueous alteration and/or thermal metamorphism. In such cases, distinctions between primary and secondary occurrences may become blurred.

Here we tabulate 52 silicates that are reasonably thought to represent secondary meteoritic minerals. Two additional hydrous silicates may form as secondary meteorite phases; however, these minerals are poorly described and as yet unconfirmed: Britholite-(Ce) [$(\text{Ce},\text{Y},\text{Ca})_5(\text{SiO}_4)_3(\text{PO}_4)_3(\text{OH},\text{F})$; MacPherson et al. 1988], if asteroidal in origin, would be the earliest known mineral with essential Ce. Pumpellyite [$\text{Ca}_2(\text{Mg},\text{Fe}^{2+})\text{Al}_2(\text{SiO}_4)_2(\text{Si}_2\text{O}_7)(\text{OH})_2 \cdot \text{H}_2\text{O}$] was provisionally identified by Gooding (1985) based on bulk composition, though Zolensky and McSween (1988) suggest that this phase is an Al-rich smectite. In addition, we do not include meteoritic hisingerite [$\text{Fe}_2\text{Si}_2\text{O}_5(\text{OH})_4 \cdot 2\text{H}_2\text{O}$], which is thought to be a terrestrial weathering product (Abreu 2016).

Quartz (SiO_2). Silica-rich phases may exsolve from highly reduced Fe-Ni-Si metal under thermal metamorphism and associated oxidation (Zanda et al. 1994). *SA quartz* occurs as a secondary phase in metamorphosed enstatite chondrites that were re-equilibrated at $T < 867^\circ\text{C}$ (Kimura et al. 2005).

Olivine group [$(\text{Mg},\text{Fe},\text{Ca})_2\text{SiO}_4$]. A significant effect of thermal metamorphism is the gradual equilibration of chondrules, manifest as disparate olivine compositions in adjacent chondrules in petrologic type 3.0 gradually become more uniform with an increase in metamorphic grade to type 3.9. At first, olivine becomes more strongly zoned; then uniform in composition (McCoy et al. 1991)—an effect shown dramatically by histograms of olivine compositions vs. petrologic type (Matsunami et al. 1990, their Figs. 7 and 8, see also their Tables 2 and 3). In spite of these significant compositional changes, equilibrated olivine in chondrules does not entirely conform to our definition of a secondary phase because the structure is unchanged. On the other hand, occurrences of olivine formed *de novo* from other phases (and thus clearly secondary) are not uncommon in meteorites, in which they span the range from nearly pure forsterite (Mg_2SiO_4) to nearly pure fayalite (Fe_2SiO_4). We lump all secondary members of the Mg-Fe solid solution into *SA olivine*.

Forsterite (Mg_2SiO_4) and fayalite (Fe_2SiO_4). The matrices of CO and CV chondrites contain sub-micrometer crystals of *SA olivine* that span the entire range from Mg- to Fe-dominant end-members (Brearley 1993a; Keller et al. 1994; Krot et al. 1995; Brearley and Jones 1998, their Figs. 144, 147, and 148). Olivine occurs as a secondary matrix mineral formed by the dehydration of phyllosilicates during thermal metamorphism in some chondrites (Tomeoka et al. 1989c; Akai 1990; Zolensky et al. 1991). This transition from clay to olivine often results in intermediate stages of partial transformation, with intimately mixed submicrometer-scale phases (Akai 1988, 1992).

Near end-member secondary forsterite, in association with

grossular, monticellite, and wollastonite, replaces åkermanite-rich melilite in altered CAIs of the Allende CV chondrite (Krot et al. 2007, 2020), whereas near end-member fayalitic olivine, Fe_2SiO_4 , occurs: (1) in rare silica-bearing chondrules with cristobalite and Ca-free pyroxene in ordinary chondrites (Brigham et al. 1986; Wasson and Krot 1994); (2) as rims around forsterite in type I chondrules of carbonaceous chondrites (Hua et al. 1988; Murakami and Ikeda 1994; Krot et al. 1995); and (3) as Fa_{88-99} grains to 100 μm diameter in association with magnetite, troilite, and pentlandite in CV chondrites (Hua and Buseck 1995). These occurrences have all been ascribed to secondary processes in chondrites (Krot et al. 1995, 1997a), though a few researchers suggest that fayalite rims could be primary as a consequence of condensation from an oxidized nebular gas (e.g., Hua et al. 1988; Weinbruch et al. 1990, 1994; Krot et al. 1997a). Fayalitic olivine is also one of several secondary phases found in opaque assemblages in the Allende CV carbonaceous chondrite (Kimura and Ikeda 1995) as well as in chondrules as replacement of low-Ca pyroxene (Kimura and Ikeda 1997). Varela et al. (2012) suggest that fayalite in the Allende CV chondrite may be the result of metasomatic exchange reactions between more forsteritic olivine and an Fe-rich fluid.

In the Belgica 7904 carbonaceous chondrite, matrix phyllosilicates were dehydrated and transformed to secondary olivine (in some cases Mn-bearing, up to 2.7 wt% MnO) that retained phyllosilicate textures, perhaps in a shock heating event (Kimura and Ikeda 1992, their Table 3a).

Doyle et al. (2015) employed ^{53}Mn - ^{53}Cr dating to demonstrate that fayalite formed as a secondary mineral by aqueous alteration in CM, CO, and CV chondrites within the first 2 to 5 Ma of nebular evolution.

Larnite (Ca_2SiO_4). *SA larnite*, the Ca end-member olivine, is a rare secondary phase found as inclusions with rankinite in andradite from the Bali CV chondrite (Ganino and Libourel 2017) and intergrown with calcite and wollastonite in the Allende CV chondrite (Krot et al. 2020).

Monticellite (CaMgSiO_4). Near end-member *SA monticellite* (with <6 mol% kirschsteinite) is a rare secondary mineral replacing melilite in CAIs from CV chondrites (Wark 1987; Krot et al. 1995; Brearley and Jones 1998, their Table A3.21).

Kirschsteinite (CaFeSiO_4). *SA kirschsteinite* is a rare secondary mineral in CAIs from CV chondrites (Krot et al. 1995). “Fe-rich monticellite” of approximate composition $[\text{Ca}(\text{Fe}_{0.7}\text{Mg}_{0.3}\text{SiO}_4)]$ is found in altered CAIs of CO chondrites (Greenwood et al. 1992; Kojima et al. 1995). Kirschsteinite is also observed in matrices of CV chondrites, where it precipitated from an aqueous fluid (MacPherson et al. 2017).

Garnet group [$\text{Ca}_3(\text{Al,Fe}^{3+},\text{V}^{3+},\text{Ti}^{4+})_2(\text{Si,Al})_3\text{O}_{12}$]. Calcic garnet group minerals are common alteration phases in both CAIs and the matrices of carbonaceous chondrites. Extensive solid solution among Al-, Fe^{3+} -, V^{3+} -, and Ti-rich end-members, at times with significant almandine ($\text{Fe}_3^+\text{Al}_2\text{Si}_3\text{O}_{12}$) and/or pyrope ($\text{Mg}_3\text{Al}_2\text{Si}_3\text{O}_{12}$) components, have been documented (Brearley and Jones 1998, their Tables A3.20 and A3.21).

Grossular ($\text{Ca}_3\text{Al}_2\text{Si}_3\text{O}_{12}$). *SA grossular*, often in association with anorthite and nepheline, occurs in altered CAIs in Allende and other CV chondrites (Fuchs 1974; Allen et al. 1978; MacPherson and Grossman 1984; Krot et al. 1995), and with

andradite, nepheline, sodalite, and hedenbergite in altered fine-grained inclusions in the Allende CV chondrite (Hashimoto and Grossman 1985). Grossular occurs in association with anorthite, spinel, and clinopyroxene as a secondary phase derived from thermal alteration of CAI melilite ($\sim 800^\circ\text{C}$) in CK carbonaceous chondrites (Chaumard et al. 2014).

Andradite ($\text{Ca}_3\text{Fe}_2^+\text{Si}_3\text{O}_{12}$). Essentially pure end-member *SA andradite* occurs in altered CAIs and their Wark-Lovering rims from the Allende and other CV carbonaceous chondrites (Fuchs 1971; Allen et al. 1978; Hashimoto and Grossman 1985; Krot et al. 1995). Andradite in association with diopside was recorded by Zolensky et al. (1996) from the matrix of the highly altered Kaidun polymict breccia.

Hutcheonite [$\text{Ca}_3\text{Ti}_2^+(\text{Si,Al})_2\text{O}_{12}$]. Ma and Krot (2014) reported *SA hutcheonite* with empirical formula $\text{Ca}_3(\text{Ti}_{1.5}^{4+}\text{Mg}_{0.25}\text{Al}_{0.17}\text{Fe}_{0.05}^{2+}\text{V}_{0.03}^{3+})(\text{Si}_{1.7}\text{Al}_{1.3})\text{O}_{12}$ (i.e., 75 mol% hutcheonite) as an alteration mineral that occurs in CAIs of the Allende CV chondrite. It occurs as crystals to 4 μm maximum dimension in association with grossular, monticellite, and wadalite.

Goldmanite [$\text{Ca}_3\text{V}_2^+(\text{Si,Al})_2\text{O}_{12}$]. The V-rich garnet, *SA goldmanite*, occurs as a secondary phase with taenite in an altered CAI from the Leoville CV3 chondrite. Simon and Grossman (1992, their Table 1) reported an average composition of $[\text{Ca}_3(\text{V}_{1.22}\text{Al}_{0.46}\text{Fe}_{0.18}\text{Ti}_{0.13})\text{Si}_3\text{O}_{12}]$, representing 61 mol% goldmanite in solid solution with 23 mol% pyrope.

Titanite (CaTiSiO_5). *SA titanite* (also commonly referred to as “sphene”), with ~ 15 mol% Al substituting for Ti, occurs as 5 μm diameter anhedral grains with nepheline in the Allende CV carbonaceous chondrite (McGuire and Hashimoto 1989). Titanite has also been reported from polymict ureilites (Delaney et al. 1984).

Adrianite [$\text{Ca}_{12}(\text{Al}_4\text{Mg}_3\text{Si})\text{O}_{32}\text{Cl}_6$]. Ma and Krot (2018) described *SA adrianite*, a secondary phase formed by alkali-halogen metasomatism of melilite, anorthite, perovskite, and/or fassaite from an altered CAI from the Allende CV chondrite. Adrianite, with an empirical formula $[(\text{Ca}_{11.7}\text{Na}_{0.2})(\text{Al}_{3.9}\text{Mg}_{2.9}\text{Si}_{7.2})\text{O}_{32}\text{Cl}_{5.8}]$, is isomorphous with the more Al-rich wadalite (see below), with which it coexists and may form a continuous solid solution. However, until more data are available, we list adrianite and wadalite as distinct Si- and Al-rich natural kinds, respectively.

Wadalite [$\text{Ca}_6\text{Al}_5\text{Si}_2\text{O}_{16}\text{Cl}_3$]. Ishii et al. (2010) reported *SA wadalite* from an altered CAI in the Allende CV chondrite. They suggest formation from melilite and anorthite precursors by metasomatism with a Cl-rich fluid. Wadalite, with an empirical formula $[(\text{Ca}_{11.6}\text{Na}_{0.1})(\text{Al}_{7.4}\text{Mg}_{1.3}\text{Si}_{5.4})\text{O}_{32}\text{Cl}_{5.7}]$ (Ma and Krot 2018), is isostructural with the more Si-rich adrianite.

Pyroxene group [$(\text{Ca,Mg,Fe})_2(\text{Si})_2\text{O}_6$]. Pyroxene group minerals are common as both primary and secondary meteorite minerals. In chondrules, diffusion rates in pyroxene are slower than in olivine, so compositional equilibration only occurs in higher petrologic grades compared to olivine. As with olivine, intermediate petrologic grades lead to zoning in clinopyroxene. As noted above, we list as secondary only those minerals with a distinctively new structure and composition that arise from aqueous and/or thermal alteration of precursor phases.

We distinguish pyroxene natural kinds based on three criteria: (1) the minerals are isostructural; (2) they are members of a continuous solid solution; and (3) they form by the same

paragenetic process. However, pyroxene compositions and structures are complex; therefore, it is not always obvious if two minerals can be lumped into one natural kind. Here we recognize three natural kinds of secondary Ca-Mg-Fe pyroxenes, including low-Ca, Mg-dominant *SA orthopyroxene* (space group *Pbca*; encompassing the orthoenstatite-ferrosilite solid solution), Ca-Mg-dominant clinopyroxene that we designate *SA diopside* (space group *C2/c*), and Ca-Fe²⁺-dominant clinopyroxene that we designate *SA hedenbergite* (also space group *C2/c*). Note that diopside and hedenbergite are known to form a continuous solid solution; however, near end-member diopside and hedenbergite coexist in some altered metamorphosed CV chondrites (e.g., Clayton et al. 1984). Therefore, *SA diopside* and *SA hedenbergite* are distinct natural kinds. In addition, Kimura and El Goresy (1989) mention a rare example of the Mn-rich orthopyroxene, *SA donpeacorite* (see below).

Orthoenstatite (MgSiO₃) and ferrosilite (FeSiO₃). *SA orthopyroxene* [usually with 12 to 28 mol% of the FeSiO₃ component; Brearley and Jones (1998), their Figures 185 and 186 and Table A3.35] commonly arises from thermal metamorphism and consequent inversion of clinoenstatite in ordinary and enstatite chondrites. Orthopyroxene subjected to intermediate stages of metamorphism (types 4 and 5) often displays partial inversion, whereas type 6 OC meteorites contain all orthopyroxene unless subsequently subjected to the shock inversion of ortho- to clinoenstatite (Brearley and Jones 1998, and references therein). Orthopyroxene is also minor phase in the matrices of CM chondrites (Müller et al. 1979). Note that while Mg-rich orthopyroxenes may be of primary or secondary origin, examples with a significant ferrosilite (FeSiO₃) content are invariably of secondary origin (Rubin and Ma 2021).

Diopside (CaMgSi₂O₆). *SA diopside*, at times with significant hedenbergite (CaFeSi₂O₆) and augite [i.e., Ca < (Mg+Fe)] components, occurs as a minor phase in equilibrated ordinary chondrites (Brearley and Jones 1998, their Table A3.36). Calcic clinopyroxene tends to display greater Mg/(Mg+Fe) contents with increasing metamorphic grade; however, even in type 6 and 7 examples that have been subjected to *T* > 1000 °C no new pyroxene phases occur. Note that some thermally metamorphosed clinopyroxenes are close to the diopside end-member. For example, Clayton et al. (1984) describe secondary diopside associated with hedenbergite, andradite, and wollastonite in altered CAIs from the Allende CV chondrite, while diopside associated with andradite was recorded by Zolensky et al. (1996) from the matrix of the highly altered Kaidun polymict breccia. Secondary diopside from altered CAIs may contain a significant aluminous kushiroite component (CaAl₂SiO₆), possibly approaching near end-member composition (A. Krot, personal communication, 11 October 2020). Sylvester et al. (1993) describe Al-diopside in association with anorthite, sodalite, calcite, and other phases from metasomatized refractory inclusions in CV chondrites, while Al-bearing diopside was produced by alteration of melilite in CAIs in the Allende CV chondrite (MacPherson et al. 1981; Hashimoto and Grossman 1985). Secondary diopside with a significant augitic component is found in association with anorthite in metamorphosed CK carbonaceous chondrites (Chaumard et al. 2014).

Hedenbergite (CaFeSi₂O₆). *SA hedenbergite*, in some instances close to end-member composition (Brearley and Jones

1998, their Table A3.22) occurs in association with wollastonite, diopside, and andradite as a product of metasomatism by Fe-rich fluids in the Allende CV chondrite and other carbonaceous chondrites (Clayton et al. 1984; MacPherson and Grossman 1984; Sheng et al. 1991; Krot et al. 1995).

Donpeacorite or kanoite [(Mn,Mg)MgSi₂O₆]. Kimura and El Goresy (1989) mention a rare example of an Mn-rich pyroxene, either orthorhombic *SA donpeacorite* or monoclinic *SA kanoite*, from the ALH 85085 CH chondrite. The empirical composition of this occurrence is [(Mn_{0.6}Mg_{0.4})MgSi₂O₆] (e.g., 60 mol% donpeacorite plus 40 mol% orthoenstatite).

Wollastonite (CaSiO₃). Essentially pure *SA wollastonite*, typically in acicular crystals 2 to 5 μm long and commonly associated with grossular, is likely a product of metasomatism of melilite in CAIs of the Allende CV carbonaceous chondrite (Fuchs 1971; Allen et al. 1978; Barber et al. 1984; MacPherson and Grossman 1984; Krot et al. 1995), though an origin as a primary condensate has also been invoked (Grossman 1975). Secondary wollastonite typically occurs in association with grossular, andradite, hedenbergite, and nepheline.

Amphibole group and related chain biopyriboles. Amphiboles and related mixed-chain biopyriboles display extremely complex solid solutions, as well as chain disorder, which complicates identifying valid mineral species as well as historical natural kinds—challenges amplified by their typical submicrometer-scale grain sizes and complex textures. Consequently, identities of most of these phases remain provisional until additional compositional and structural data become available. At least two anhydrous amphiboles, fluoro-richterite (Olsen et al. 1973; Rubin 1983) and kaersutite (Prinz et al. 1982) were listed as primary asteroidal minerals by Morrison and Hazen (2021). Here we list several hydrous amphiboles as secondary asteroidal phases.

Magnesio-arfvedsonite [NaNa₂(Mg₄Fe³⁺)Si₈O₂₂(OH)₂]. Ivanov et al. (2003) reported “arfvedsonite” in association with aenigmatite, fluorapatite, wilkinsonite, and clay minerals as inclusions in albite in unusual alkaline and subalkaline clasts from the Kaidun polymict breccia meteorite. The empirical formula of 20 μm diameter crystals is [(Na_{1.9}Ca_{0.7}K_{0.3})(Mg_{2.4}Fe²⁺Mn_{0.1})(Fe³⁺Al_{0.1})Si₈O₂₂(OH)₂], which corresponds to a complex solid solution with ~60 mol% *SA magnesio-arfvedsonite* (i.e., Na-Na-Mg-Fe³⁺) plus significant contents of arfvedsonite (Na-Na-Fe²⁺-Fe³⁺), a calcic amphibole [e.g., ferro-ferri-hornblende (□-Ca-Fe²⁺-Fe³⁺)], and a potassic amphibole [e.g., potassic-chloro-hastingsite (K-Ca-Fe²⁺-Fe³⁺)].

Anthophyllite [(Mg,Fe)₇Si₈O₂₂(OH)₂]. Brearley (1997a) reported a rare occurrence of *SA anthophyllite* in association with talc, magnesio-hornblende, and disordered biopyriboles (named jimthompsonite) from the Allende CV chondrite. The composition of anthophyllite was not recorded.

Magnesio-hornblende [□Ca₂(Mg,Fe²⁺)₄(Si₇Al)O₂₂(OH)₂]. *SA magnesio-hornblende* with Mg/(Mg+Fe) ~0.80 was reported by Brearley (1997a) in association with talc, rare anthophyllite, and disordered biopyriboles, which replace enstatite in chondrules from the Allende CV chondrite.

Winchite [□NaCa(Mg₄Al)Si₈O₂₂(OH)₂] and barroisite [□NaCa(Mg₃Al₂)Si₈O₂₂(OH)₂]. Dobricá and Brearley (2014) reported amphibole group minerals with variable composition corresponding to the winchite-barroisite solid solution from the

Tieschitz unequilibrated (3.6) ordinary chondrite. We lump these phases into *SA winchite*, the more magnesian of the two similar species. Amphiboles, rather than phyllosilicates, are the most abundant hydrous minerals in Tieschitz, suggesting the presence of aqueous fluids during thermal metamorphism.

Jimthompsonite [(Mg,Fe)₃Si₆O₁₆(OH)₂]. High-resolution transmission electron microscopy of complex, disordered biopyriboles from altered enstatite-rich chondrules in the Allende CV chondrite revealed a nanometer-scale region consisting of a triple-chain jimthompsonite layer three unit-cells thick, surrounded by double- and quadruple-chain regions (Brearley 1997a). Such occurrences have been related to incompletely reacted chain silicates converting to phyllosilicates (Veblen and Buseck 1979). We suggest that such disordered biopyriboles domains may represent a distinct and important natural kind in altered assemblages; however, we name this occurrence as *SA jimthompsonite*, with the proviso that this disordered meteorite occurrence is not equivalent to the crystalline species of that name.

Mica group. Isolated mica group minerals are scarce in meteorites, with five species each known from a single meteorite type—four of them from the Allende CV carbonaceous chondrite, the other from the Murray CM2 chondrite. However, mica may occur more commonly as intimate intergrowths with other phyllosilicates, for example, as mica-montmorillonite assemblages in altered CAIs of the Allende CV chondrite (Tomeoka and Buseck 1982b), and as a Na-rich 10 Å mica intergrown with 7 Å serpentine in the Mokoia CV chondrite (Tomeoka and Buseck 1990).

Aspidolite [NaMg₃AlSi₃O₁₀(OH)₂]. Krot et al. (1995, their Table 8) identified several meteorite occurrences of the trioctahedral mica, *SA aspidolite* (also known as “Na phlogopite”) with Na > K (though quantitative chemistry was not reported) from the altered chondrules or CAIs of several CV3 carbonaceous chondrites.

Phlogopite [KMg₃AlSi₃O₁₀(OH)₂]. Kimura and Ikeda (1996) described *SA phlogopite* in the groundmass of altered chondrules in the Allende CV chondrite, while Tomeoka and Buseck (1990) identified “Na-rich phlogopite” intergrown with serpentine in the Mokoia CV chondrite. Lacking more detailed compositional information, we cannot conclude with certainty if phlogopite and aspidolite in CV chondrites represent distinct natural kinds.

Paragonite [NaAl₂AlSi₃O₁₀(OH)₂]. Lee and Greenwood (1994, their Table 4) tentatively identified *SA paragonite* in an altered spinel-fassaite-perovskite CAI from the Murray CM2 carbonaceous chondrite in association with secondary calcite, sulfates, and clay minerals.

Clintonite [CaAlMg₂SiAl₃O₁₀(OH)₂]. *SA clintonite* with an empirical formula of [(Ca_{0.95}Na_{0.05})(Mg_{2.5}Al_{0.3}Fe_{0.05}Ti_{0.05}□_{0.1})Al_{2.4}Si_{1.6}O₁₀(OH)₂] occurs in the Allende CV chondrite as an alteration vein up to 40 nm wide and 500 nm long replacing grossular (Keller and Buseck 1991).

Margarite [CaAl₂Al₂Si₂O₁₀(OH)₂]. *SA margarite* was identified by Keller and Buseck (1991) as an alteration product of anorthite in the Allende CV chondrite. The Fe- and Mg-bearing margarite, with an empirical formula of [(Ca_{0.95}Na_{0.05})(Al_{1.65}Fe_{0.20}Mg_{0.15})Al_{2.2}Si_{1.8}O₁₀(OH)₂], occurs as lamellae 20 to 500 nm wide, with intergrown as 10 and 20 Å polytypes.

Clay minerals. Clay minerals are common in altered parts of chondrites, notably their fine-grained matrices, most often

as a consequence of parent-body processes, though at times possibly complemented by pre-accretionary hydration (e.g., Bischoff 1998; Zolensky et al. 1993; Krot et al. 1995, their Table 2; Ciesla et al. 2003). Clay minerals are an especially relevant mineral group when discussing the challenges of any classification system identifying species or kinds. In spite of the approval of more than 50 clay mineral species, as well as dozens of additional approved and unapproved varieties (Hazen et al. 2013, their Tables 2 and 3), clay minerals in nature display compositional variability, disordered mixed layering, and nanoscale structural heterogeneities that rarely conform to end-member idealized species.

Clay minerals are particularly common in the altered matrices of carbonaceous chondrites (especially CI and CM, but also CO, CV, and CR) as well as ordinary chondrites (Barber 1981, 1985; Zolensky and McSween 1988; Brearley 1993a, 1997b; Zolensky et al. 1993). Exact identification of these clay minerals is often problematic, especially lacking electron diffraction data and/or high-resolution imaging. Secondary phyllosilicates are often complexly interlayered and do not correspond to a single species. For example, saponite is often found interlayered with serpentine (Zolensky et al. 1993, 1996; Endress et al. 1994; Brearley 1995, 1997b). Similarly, Tomeoka and Buseck (1988) illustrated interlayering of serpentine and saponite, in some cases in close association with ferrihydrite, in the Orgueil CI chondrite. In CV carbonaceous chondrites, Tomeoka and Buseck (1982b) reported montmorillonite/mica and montmorillonite/K-feldspar intergrowths in altered CAIs of Allende; Keller et al. (1994) found intergrown saponite and dioctahedral mica in Bali, and Tomeoka and Buseck (1990) observed Na-rich 10 Å phlogopite intergrown with 7 Å serpentine in Mokoia. Lee and Greenwood (1994) describe 7 Å berthierine intergrown with chlorite or serpentine in altered CAIs of the Murray CM2 chondrite.

Of special interest are intergrowths of tochilinite (itself an ordered intergrowth of mackinawite- and brucite-type layers) with the Fe³⁺-dominant serpentine group mineral, cronstedtite (e.g., Tomeoka and Buseck 1985; Nakamura and Nakamura 1996; Marrocchi et al. 2014)—materials collectively referred to as tochilinite-cronstedtite intergrowths (TCIs).

Mesostasis in CO and CM chondrites is often altered to green phyllosilicates, sometimes termed “spinach” (Fuchs et al. 1973), composed of chlorite and/or berthierine (Richardson and McSween 1978; Ikeda 1983). However, in the Belgica 7904 carbonaceous chondrite, matrix phyllosilicates were dehydrated and transformed to secondary olivine that retains phyllosilicate textures, perhaps in a shock heating event (Kimura and Ikeda 1992, their Table 3b).

Yet another complication in dealing with “clay minerals” is the presence of “protophyllosilicates” that are amorphous at the scale of electron diffraction. For example, in the unequilibrated ALH A77307 CO3 carbonaceous chondrite a continuous range exists from amorphous regions to those with short-range order (on the scale of nanometers), to well-defined localized and/or intergrown 7, 10, and 14 Å domains of serpentine, smectite, and chlorite, respectively (Brearley 1993a, Table 2 therein).

In the following sections, we catalog confirmed phyllosilicates from the serpentine, talc, smectite-vermiculite, and chlorite groups, as well as related TCIs.

Serpentine group [(Mg,Fe²⁺,Fe³⁺,Al)₃(Al,Fe³⁺,Si)₂O₅(OH)₄]. Minerals of the serpentine group feature a trioctahedral (Mg,Fe²⁺,Fe³⁺,Al) sheet bonded to a tetrahedral (Al,Fe³⁺,Si) sheet in a 7 Å layer repeat with interlayer (OH⁻) groups. These layer silicates are found commonly in altered CM chondrites as the dominant matrix phase, with examples spanning the compositional range from Mg- to Fe-rich (Bunch and Chang 1980; Barber 1981; MacPherson et al. 1984; Greenwood et al. 1994; Lee and Greenwood 1994; Marrocchi et al. 2014; see Brearley and Jones 1998, their Fig. 117). According to Zolensky and McSween (1988), it is common to find three distinct coexisting serpentine phases in the matrices of individual CM chondrites, perhaps corresponding to cronstedtite, greenalite, and “ferroan antigorite.”

Magnesian serpentines are complicated by the occurrence of three distinct structural types—antigorite, chrysotile, and lizardite—related to long-range structural differences in the topology within and between layers. All three types have been invoked in the meteoritics literature, though antigorite and lizardite have not been convincingly demonstrated (Barber 1981; Akai and Kanno 1986; Zolensky and McSween 1988; Keller and Buseck 1990b). Because a continuous Mg-Fe²⁺ solid solution appears to exist in meteoritic serpentines and Fe-rich occurrences predominate, we lump all Fe²⁺-Mg-dominant, Al-poor serpentines into *SA greenalite*. Given the compositional complexity of serpentine group solid solutions, coupled with the common occurrence of nanometer-scale intergrowths of serpentine and other layer structure phases, it is difficult to define discrete mineral species or natural kinds. Here we list *SA greenalite*, *SA cronstedtite*, and *SA berthierine*, which appear to encompass most meteoritic occurrences.

Greenalite [(Fe²⁺,Mg)₃Si₂O₅(OH)₄] and **chrysotile** [(Mg,Fe²⁺)₃Si₂O₅(OH)₄]. Serpentine group minerals in the matrices of CM chondrites span a range of Fe²⁺/(Mg+Fe²⁺) from ~0.3 to >0.9 (e.g., Zolensky et al. 1993). We lump examples of Mg-Fe²⁺-rich serpentines with compositions near the chrysotile-greenalite solid solution into *SA greenalite*. Note, however, that some specimens contain significant Al, which likely represents an amesite [(Mg,Al)₃(Al,Si)₂O₅(OH)₄] and/or berthierine [(Fe²⁺,Mg,Fe³⁺)₃(Al,Si)₂O₅(OH)₄] component, as well as S and Ni, which may point to interlayering with tochilinite or another phase (Tomeoka et al. 1989a). Keller and Buseck (1990b) identified 7 Å Mg-rich, Fe-bearing phyllosilicate regions, likely corresponding to chrysotile, a few unit-cells thick by electron diffraction in matrices of the relatively unaltered Lancé CO3 chondrite associated with Fe-rich olivine and ferrihydrite. Examination of other more altered Kainsaz and Warrenton CO chondrites found ferrihydrite but no phyllosilicates, indicating that the phyllosilicates in Lancé may represent an earlier stage of aqueous alteration. In addition, Tomeoka and Buseck (1982a) described an unusual occurrence of a Fe-Ni-rich serpentine-group mineral, possibly owing to the incorporation of a significant brindleyite [(Ni,Al)₃(Si,Al)₂O₅(OH)₄] component in greenalite, from altered CAIs in the Allende CV chondrite.

Cronstedtite [(Fe²⁺,Fe³⁺)₃(Si,Fe³⁺)₂O₅(OH)₄]. *SA cronstedtite* is commonly found in the matrices of CM carbonaceous chondrites in nanoscale tochilinite-cronstedtite intergrowths (Marrocchi et al. 2014). Cronstedtite of close to end-member composition occurs in well-defined grains up to 10 μm maximum

dimension from the Cochabamba CM2 chondrite (Müller et al. 1979), though it always contains some Mg in solid solution, and commonly Al, as well (Bunch and Chang 1980). Palmer and Lauretta (2011) reported cronstedtite as a common alteration product of kamacite in CM chondrites in microenvironments with high Si and low S.

Berthierine [(Fe²⁺,Mg,Fe³⁺)₃(Al,Si)₂O₅(OH)₄] and **amesite** [(Mg,Al)₃(Al,Si)₂O₅(OH)₄]. Some serpentine group minerals in the matrices of CM and CO chondrites have been reported to be aluminous (Barber 1981; Ikeda 1983; Zolensky and McSween 1988; Lee and Greenwood 1994), perhaps at times with more than 50 mol% berthierine or amesite component. For example, *SA berthierine* has been invoked as a component of the distinctive green “spinach” of altered chondrules in CM chondrites (Richardson and McSween 1978; Ikeda 1983).

Talc group [(Mg,Fe²⁺)₃Si₄O₁₀(OH)₂]. Talc group minerals are characterized by a 10 Å octahedral layer sandwiched between two tetrahedral layers (i.e., T-O-T) with interlayer (OH) groups. *SA talc* is the only meteoritic example documented thus far.

Talc [(Mg₃Si₄O₁₀(OH)₂]. Brearley (1997a) documented *SA talc* in association with calcic amphibole and disordered biopyriboles replacing enstatite in the Allende CV chondrite. Ikeda (1992) described a “sodian talc” with significant Na-Al substitution for Mg in Yamato 82162, which is a highly altered carbonaceous chondrite.

Smectite and vermiculite group [(Na,Ca)_{0.3}(Mg,Fe²⁺,Al,Fe³⁺)₂₋₃(Si,Al)₄O₁₀(OH)₂·4H₂O]. Smectite and vermiculite group clay minerals feature the 10 Å T-O-T layer structure of talc and mica group minerals, but the interlayer regions incorporate significant H₂O molecules and alkali and alkaline earth cations that can cause the structure to expand reversibly perpendicular to the layers. These phases are among the most common phyllosilicates in meteorites that have experienced aqueous alteration. However, as with other clay mineral groups, extensive solid solution, sub-micrometer scale grains, and intergrowths of different layer types complicate efforts at definitive identifications. The occurrence of the trioctahedral smectites of the saponite-ferrosaponite solid solution are well established. However, isolated suggestions of dioctahedral Al and Fe³⁺ species, including montmorillonite, nontronite, and vermiculite are not accompanied by convincing structural and compositional data (see below) and their inclusion as meteorite minerals must for now remain tentative.

The use of unapproved names for some smectite-vermiculite group phyllosilicates also adds uncertainty. In particular, “smectite” is not an approved species name, but it is often used in the meteoritics literature to designate both dioctahedral and trioctahedral expandable 10 Å (Mg-Fe-Al) clay minerals. Given the extensive solid solution among these minerals, we were tempted to lump all of these phases into “SA smectite.” However, the trioctahedral species saponite and ferrosaponite appear to be distinct from dioctahedral montmorillonite and nontronite. Therefore, we tentatively identify four group members: *SA saponite*, *SA montmorillonite*, *SA nontronite*, and *SA vermiculite*.

Saponite and ferrosaponite [(Ca,Na)_{0.3}(Mg,Fe²⁺)₃(Si,Al)₄O₁₀(OH)₂·4H₂O]. We lump trioctahedral smectites that lie close to the Mg-Fe²⁺ solid solution (i.e., with minor Al and Fe³⁺) into *SA saponite*, because most examples are Mg-dominant. *SA*

saponite, in some cases Fe-rich (e.g., Alexander et al. 1989), occurs in the fine-grained matrices of CV chondrites as sub-micrometer layered regions in association with olivine, which it may replace (Keller and Buseck 1990a; Tomeoka and Buseck 1990; Keller et al. 1994; Lee et al. 1996). Keller and Buseck (1990a, their Table 1) report the average composition of Mg-rich, Al-poor saponite from the Kaba CV3 chondrite as $[(\text{Na}_{0.25}\text{K}_{0.03}\text{Ca}_{0.02})(\text{Mg}_{2.59}\text{Fe}_{0.25}\text{Al}_{0.02}\text{Ti}_{0.01}\square_{0.13})(\text{Al}_{0.44}\text{Si}_{3.56})\text{O}_{10}(\text{OH})_2 \cdot n\text{H}_2\text{O}]$. Tomeoka and Buseck (1988) illustrated interlayering of serpentine and saponite, in some cases in close association with ferrihydrite, in the Orgueil CI chondrite. Saponite is rarely found in CM chondrites, though Brearley (1995) reported saponite/serpentine intergrowths (as opposed to more typical cronstedtite/tochilinite intergrowths) in association with magnetite in the matrix of the unusual Bells CM2 chondrite. Of special note are remarkable 1 mm diameter saponite single crystals replacing olivine in a clast from the unusual Kaidun polymict breccia (Zolensky and Ivanov 2003, their Fig. 18).

A partial solid solution exists between saponite and Al- and/or Fe³⁺-bearing dioctahedral montmorillonite. For example, Tomeoka and Buseck (1990, their Table 3) report several analyses, including an approximate composition from the Kaba CV3 chondrite as $[(\text{Na}_{0.42}\text{K}_{0.05}\text{Ca}_{0.35})(\text{Mg}_{2.08}\text{Fe}_{0.44}\text{Al}_{0.23}\square_{0.25})(\text{Al}_{1.07}\text{Si}_{2.93})\text{O}_{10}(\text{OH})_2 \cdot n\text{H}_2\text{O}]$, i.e., with 66 mol% saponite plus significant nontronite and montmorillonite components. Some occurrences of so-called “Al-rich smectite,” “high-Al phyllosilicate” (HAP), or “sobokite” in altered CAIs of CV chondrites (e.g., Cohen et al. 1983, their Table 6) are likely similar examples that result from the alteration of feldspathic glass and/or Ca-rich clinopyroxene (Buseck and Hua 1993, their Tables 3, 4, and 5).

Montmorillonite $[(\text{Na},\text{Ca})_{0.3}(\text{Al},\text{Mg})_2\text{Si}_4\text{O}_{10}(\text{OH})_2 \cdot n\text{H}_2\text{O}]$. Montmorillonite is the proper name for dioctahedral smectites in which Al is the dominant cation in the octahedral layer. *SA montmorillonite* has been tentatively identified as a rare alteration phase of fassaite in CAIs of the Allende CV chondrite (Tomeoka and Buseck 1982b; Krot et al. 1995) and in the matrices of the Cold Bokkeveld and Nawapali CM chondrites (Zolensky and McSween 1988, and references therein). Note, however, that Tomeoka and Buseck (1982b) state that montmorillonite is “used as representing minerals having rather broad compositional ranges in these groups and not particular species.” We suspect that similar liberties in nomenclature apply to examples of nontronite and vermiculite (see below).

Nontronite $[\text{Na}_0\text{Fe}_2^+(\text{Si},\text{Al})_4\text{O}_{10}(\text{OH})_2 \cdot n\text{H}_2\text{O}]$. Nontronite is the general name for the Fe³⁺-dominant dioctahedral smectite. Hutchison et al. (1987) suggested that an Na-Ca-Fe-bearing smectite from the Semarkona ordinary chondrite is best fit by *SA nontronite*. However, detailed compositional information was not provided.

Vermiculite $[\text{Mg}_{0.35}(\text{Mg},\text{Fe}^{3+},\text{Al})_3(\text{Si},\text{Al})_4\text{O}_{10}(\text{OH})_2 \cdot 4\text{H}_2\text{O}]$. Zolensky et al. (1993) found a 100 nm thick grain of a phyllosilicate that they ascribed to *SA vermiculite* in the matrix of the Nagoya CM chondrite. However, this and other occurrences of meteoritic vermiculite are tentative because they may also represent saponite (e.g., Zolensky and McSween 1988).

Chlorite group $[(\text{Fe}^{2+},\text{Mg},\text{Al},\text{Fe}^{3+})_6(\text{Si},\text{Al})_4\text{O}_{10}(\text{OH},\text{O})_8]$. Chlorite group minerals incorporate alternating TOT and brucite-type layers in a 14 Å arrangement. A few instances of meteoritic

chlorite, all presumably trioctahedral Mg-Fe²⁺-dominant varieties, have been reported from carbonaceous chondrites. However, confusion arises for three reasons: (1) the name “chlorite” is sometimes used interchangeably with “septachlorite,” which is an unapproved name for 7 Å berthierine-type serpentines; (2) the name “chamosite” is now the official IMA name for the 14 Å, Fe²⁺-dominant chlorite, but it has also been used historically for 7 Å berthierine-type serpentines; and (3) 7 Å serpentine sometimes occurs in two-layer polytypes that yield a 14 Å X-ray or electron diffraction spacing that can be confused with true chlorite group minerals (Bunch and Chang 1980; Barber 1981). Consequently, several reported instances of meteoritic “chlorite” are undoubtedly not true chlorite group minerals. It is likely that a continuous Mg-Fe²⁺ solid solution occurs between clinochlore and chamosite; however, until more compositional data are available, we recognize *SA clinochlore* and *SA chamosite* as distinct natural kinds.

Clinochlore $[\text{Mg}_5\text{Al}(\text{AlSi}_3\text{O}_{10})(\text{OH})_8]$. Zolensky et al. (1993, their Table 4) characterized a 14 Å Mg-Al-rich phyllosilicate $[\text{Mg}/(\text{Mg}+\text{Fe}) \sim 0.83; 19.4 \text{ wt}\% \text{Al}_2\text{O}_3]$ from the Nagoya CM2 chondrite, which they identified as *SA clinochlore*. Nagoya clinochlore occurs in association with a more Fe-rich saponite.

Chamosite $[(\text{Fe}^{2+},\text{Mg},\text{Al},\text{Fe}^{3+})_6(\text{Si},\text{Al})_4\text{O}_{10}(\text{OH},\text{O})_8]$. Convincing evidence for a true Fe²⁺-dominant chlorite, probably *SA chamosite*, was presented by Brearley (1993a, Fig. 7 therein), who employed electron diffraction to illustrate 14 Å layered Fe-rich regions a few unit-cells thick in matrices of the ALH A77037 CO3 chondrite, in association with a 7 Å Mg-serpentine. In addition, Lee and Greenwood (1994, their Fig. 9b and Table 4) reported rare 14 Å chlorite-like regions associated with more abundant 7 Å aluminous serpentine in altered CAIs of the Murray CM2 chondrite. The average composition of these coexisting phyllosilicates is $[\text{Mg}/(\text{Mg}+\text{Fe}) \sim 0.47; \sim 25 \text{ wt}\% \text{Al}_2\text{O}_3]$. Gooding (1985) tentatively identified 14 Å ferroan chlorite, suggesting that the structure and chemistry of phyllosilicates in the matrices of the Nagoya CM2 chondrite and Semarkona LL3 ordinary chondrite match characteristics of chamosite, though he could not rule out a mica-like phase.

Tochilinite-cronstedtite intergrowths (TCIs). A variety of matrix phases, notably in CM chondrites, are poorly crystallized to amorphous and thus difficult to catalog with IMA protocols. Fuchs et al. (1973) termed a suite of these materials that incorporate Ni, S, Mg, Fe, and Si as “poorly characterized phases” or “PCPs,” though that designation has been replaced by TCIs (e.g., Vacher et al. 2019). Bunch and Chang (1980) characterized three compositional types (originally called PCP I, II, and III), which have subsequently been shown to represent a continuum of complexly interlayered mixtures of cronstedtite, an Fe-rich serpentine, and tochilinite, which is itself an ordered sequence of mackinawite- and brucite-type layers (Mackinnon and Zolensky 1984; Nakamura and Nakamura 1996). The lateral misfit among serpentine, brucite, and mackinawite layers, in some instances exacerbated by the compositional variability of these independent sheets, leads to various nanometer-scale topologies, including undulatory, kinked, and enrolled (and therefore fibrous) morphologies (e.g., Tomeoka and Buseck 1985). Given the common occurrence and distinctive characteristics of these complex mixed-layer phases, we recognize *SA TCI* as

a separate natural kind.

TCI (Fe-Ni-Si-S-O). *SA TCI* occurs as the most abundant phase in the altered matrices of many CM chondrites (Fuchs et al. 1973; Bunch and Chang 1980; Tomeoka and Buseck 1985).

Feldspar group [(Na,K,Ca)Al(Al,Si)₃O₈]. Feldspar group minerals of the albite-anorthite series commonly occur as primary meteorite minerals but also as alteration phases. Secondary feldspars arise through three processes: (1) thermal metamorphism and devitrification of glassy mesostasis; (2) metasomatism of calc-silicates by alkali-halogen-bearing fluids during metamorphism; and (3) thermal metamorphism of other Ca-bearing minerals, including augite and melilite. A wide range of secondary plagioclase compositions has been documented. Until more compositional data are available, we lump all examples into *SA plagioclase*. Note that we have not found reports of secondary potassic feldspars.

Anorthite (CaAl₂Si₂O₈) and albite (NaAlSi₃O₈). Anorthitic *SA plagioclase* (An₆₅₋₉₅) is a common secondary phase in metamorphosed ordinary chondrites, where it occurs as a consequence of devitrification of feldspathic glass (Brearley and Jones 1998, and references therein, Figs. 187 and 188, Table A3.37). Na-bearing albitic feldspar (An_{<20}) is found as crystallites in the devitrified mesostasis of some R group chondrites. These meteorites experienced both significant thermal and aqueous alteration, and the alkali feldspar is thus assumed to be of secondary origin (Rubin and Kallemeyn 1994; Schulze et al. 1994).

In thermally metamorphosed CK chondrites, secondary anorthite may form as a consequence of the breakdown of augite (Noguchi 1993). Anorthite (>An₆) forms as a secondary phase along fractures in melilite in some altered CAIs in CV chondrites (Allen et al. 1978; MacPherson and Grossman 1984) and in association with grossular, spinel, and clinopyroxene derived from melilite alteration in CAIs of CK carbonaceous chondrites (Chaumard et al. 2014). Noguchi (1994) described secondary plagioclase (An₄₅₋₉₅) in the Coolidge carbonaceous chondrite, which may have affinities with CV chondrites.

SA plagioclase also forms via hydrothermal activity in ordinary chondrites. Dyl et al. (2012) described an unusual secondary plagioclase-rich clast from the Villaberto de la Peña L6 ordinary chondrite, in which a short-lived hydrothermal event is estimated to have reached ~800 °C and 1 bar water pressure for a period of no more than 10 years. Coexisting feldspar grains range from albitic (~An₁₀) to intermediate (An₃₅), with Na-Ca and oxygen-isotope diffusion profiles that reveal the transient hydrothermal event. Lewis and Jones (2016) describe secondary plagioclase (An₂₋₈₈) in ordinary chondrites of types 4 to 6 and provide evidence that feldspar and coexisting phosphates experienced metasomatism by alkali-halogen-bearing fluids during metamorphism.

Celsian (BaAl₂Si₂O₈). *SA celsian* occurs as inclusions to 2 μm diameter in secondary Na-rich melilite in a metasomatically altered CAI from the Allende CV chondrite (A. Krot, personal communication, 11 October 2020).

Feldspathoid group. Three feldspathoid group minerals, *SA nepheline*, *SA sodalite*, and *SA marialite*, occur in carbonaceous chondrites that experienced thermal metamorphism.

Nepheline [Na₃(Al₄Si₄O₁₆)]. *SA nepheline* is commonly encountered in CV carbonaceous chondrites. It occurs as an

alteration product of melilite in CAIs from CV chondrites, often in association with anorthite, grossular, and sodalite (Marvin et al. 1970; Allen et al. 1978; MacPherson and Grossman 1984; Krot et al. 1995; Kimura and Ikeda 1997; Brearley and Jones 1998, their Table A3.21). In the Allende CV chondrite, fine-grained inclusions contain nepheline and sodalite associated with other secondary phases, including grossular, diopside, anorthite, and monticellite (Clayton et al. 1984; Hashimoto and Grossman 1985; Wark et al. 1987). In CO chondrites, nepheline may represent as much as 80 vol% of altered CAIs (Ikeda 1982; Tomeoka et al. 1992, their Table 3; Kojima et al. 1995). Plagioclase can also be altered to nepheline, for example, in chondrules of CO chondrites (Brearley and Jones 1998, their Fig. 35). A rare occurrence of a single grain of nepheline with significant substitution by K, Ca, Mg, and Fe in the thermally metamorphosed Rio Negro ordinary chondrite, was reported by Fodor et al. (1977).

Sodalite [Na₄(Si₃Al₃)O₁₂Cl]. *SA sodalite*, commonly in association with more abundant nepheline, is a secondary alteration phase in CAIs from the Allende and other CV chondrites (Blander and Fuchs 1975; MacPherson and Grossman 1984; Kornacki and Wood 1985; Kimura and Ikeda 1997). In addition, Krot et al. (1995) recorded sodalite as a product of metasomatism of both CAIs and mesostasis in CV carbonaceous chondrites. Sodalite occurs with nepheline in CO chondrites, as revealed by the presence of chlorine (Tomeoka et al. 1992).

Marialite (Na₄Al₃Si₉O₂₄Cl). A scapolite group mineral of empirical composition [(Na_{3.08}K_{0.15}Ca_{0.47}Fe_{0.39}Mg_{0.09})(Al_{3.46}Si_{8.54})O₂₄Cl], predominantly marialite, was reported by Alexander et al. (1987) from a thermally metamorphosed clast in the Bishunpur LL3 ordinary chondrite.

Zeolite group. Zeolite group hydrated framework silicates are rare secondary phases in meteorites. We record a tentative occurrence of *SA chabazite*. We do not include an unidentified Al-Si phase with minor K, Mg, and Ca, possibly a zeolite replacing melilite, which occurs in altered CAIs of the Leoville CV chondrite (Wlotzka and Wark 1982, their Table 1). Note, in addition, that a possible occurrence of meteoritic stilbite from the ALH A77296 weathered L6 ordinary chondrite (Gooding 1981) and an Na-rich zeolite from the Tieschitz ordinary chondrite (Alexander et al. 1986) are likely products of terrestrial weathering.

Chabazite-Na [(Na₃K)Al₄Si₆O₂₄·11H₂O]. (Zolensky and Ivanov 2003) report a single mm-long fragment of *SA chabazite-Na* from the unusual Kaidun polymict breccia. The chabazite crystal is zoned in Na-K, which is characteristic of heated material.

Other silicates

Roedderite [(Na,K)₂Mg₂(Mg₃Si₁₂)O₃₀] and merrihueite [(K,Na)₂(Fe,Mg)₅Si₁₂O₃₀]. Krot and Wasson (1994) described roedderite-merrihueite-bearing chondrules from ordinary chondrites, which they interpreted as arising from the reaction of silica with alkali-rich gas on a parent body. Minerals from the roedderite-merrihueite solid-solution series are also minor phases in the matrices of thermally metamorphosed enstatite chondrites (Fuchs et al. 1966; Rambaldi et al. 1986b; Ikeda 1989). We ascribe all such occurrences to *SA roedderite*, even though some examples may be slightly K-dominant. The formation mechanism of these occurrences is not certain, though Rambaldi et al. (1986b) describe a specimen from the Qingzhen EH3

chondrite with empirical composition $[(\text{Na}_{1.1}\text{K}_{0.9})_2(\text{Mg}_{4.7}\text{Fe}_{0.3})(\text{Si}_{11.8}\text{Al}_{0.1})\text{O}_{30}]$ that occurs in matrix-connected veins through oxide and silicate phases, thus pointing to fluid alteration. In previous contributions (Morrison and Hazen 2021; Hazen et al. 2021) we attributed at least some roedderite-merrihueite occurrences to primary chondrule mineralization. However, Rhian Jones (personal communication, 4 June 2020) suggests that all occurrences are secondary.

Wilkinsonite $[\text{Na}_4(\text{Fe}_3^+\text{Fe}_4^+)\text{O}_4(\text{Si}_{12}\text{O}_{36})]$. Ivanov et al. (2003) reported the closely related minerals aenigmatite and wilkinsonite in association with arfvedsonite and fluorapatite as inclusions in albite in unusual alkaline and subalkaline clasts from the Kaidun polymict breccia. Zolensky and Ivanov (2003) report the composition of wilkinsonite as $[\text{Na}_{4.0}(\text{Fe}_{7.2}^{2+}\text{Mg}_{0.4}\text{Ca}_{0.2}\text{Fe}_{3.8}^{3+}\text{Al}_{0.2})\text{O}_4(\text{Si}_{11.9}\text{O}_{36})]$.

Aenigmatite $[\text{Na}_4(\text{Fe}_{10}^{2+}\text{Ti}_2)\text{O}_4(\text{Si}_{12}\text{O}_{36})]$. Zolensky and Ivanov (2003) record the compositions of two *SA aenigmatite* grains in association with albite and fluorapatite, with average composition $[\text{Na}_{4.0}(\text{Fe}_{7.5}^{2+}\text{Mg}_{2.2}\text{Ca}_{0.2}\text{Mn}_{0.2}\text{Ti}_{2.0})\text{O}_4(\text{Al}_{0.2}\text{Si}_{11.8}\text{O}_{36})]$. Wark (1986, Table 2 therein) suggests that “a substituted aenigmatite” from an altered CAI in the Allende CV chondrite, with approximate composition $[\text{Ca}_4(\text{Mg},\text{Fe},\text{Ca})_{12}(\text{Al},\text{Si})_{12}\text{O}_{40}]$, is a fine-grained alteration product of melilite. If so, this Ca-rich mineral may represent a new compositional variant of the aenigmatite group.

Gehlenite $(\text{Ca}_2\text{Al}_2\text{SiO}_7)$ and **åkermanite** $(\text{Ca}_2\text{MgSi}_2\text{O}_7)$. A solid solution exists between gehlenite $(\text{Ca}_2\text{Al}_2\text{SiO}_7)$ and åkermanite $(\text{Ca}_2\text{MgSi}_2\text{O}_7)$. Melilite is most commonly observed as a primary mineral in Ca-Al-rich inclusions, but it also occurs as a secondary CAI metamorphic phase (*SA melilite*) in the form of zoned mantles on pyroxene formed by reaction with high-temperature Ca-rich fluids (Wark et al. 1987). In addition, secondary Na-bearing melilite (up to 7 wt% Na_2O) replaces primary igneous anorthite in Type B and C CAIs (M. Zolensky, personal communication, 5 October 2020).

Indialite $[\text{Mg}_2\text{Al}_3(\text{AlSi}_5)\text{O}_{18}]$. Fuchs (1969) reported “hexagonal cordierite” from the Allende carbonaceous chondrite. Mikouchi et al. (2016) subsequently described an occurrence of indialite, the high-temperature ($T > 1450$ °C) beryl-structured polymorph of cordierite, with composition $[\text{Na}_{0.19}\text{Mg}_{1.95}\text{Fe}_{0.02}\text{Al}_{3.66}\text{Si}_{5.19}\text{O}_{18}]$ from an Al-rich chondrule in Yamato 82094, which is an ungrouped carbonaceous chondrite. They suggested that prior reports of rare meteoritic cordierite were also indialite. We therefore lump all of these occurrences into *SA indialite*.

Dmisteinbergite $(\text{CaAl}_2\text{Si}_2\text{O}_8)$. Dmisteinbergite, a high-temperature polymorph of anorthite, has been reported as both a primary and secondary meteorite mineral, notably from CAIs in the Allende CV chondrite (Ma et al. 2013), which also contain zones of Ba-rich dmisteinbergite (up to 27 mol% $\text{BaAl}_2\text{Si}_2\text{O}_8$). Park et al. (2013) and Krot et al. (2020) characterized acicular Na-bearing *SA dmisteinbergite* in hydrothermally altered CAIs from the Allende CV chondrite. Fintor et al. (2014) presented evidence for the hydrothermal origin of dmisteinbergite associated with secondary nepheline, sodalite, and grossular, which are likely alteration products of melilite from a CAI in the NWA 2086 CV3 chondrite.

Rankinite $(\text{Ca}_3\text{Si}_2\text{O}_7)$. *SA rankinite* occurs with larnite as inclusions in andradite in the Bali CV chondrite—phases indicative of low silica activity (Ganino and Libourel 2017).

Tilleyite $[\text{Ca}_5\text{Si}_2\text{O}_7(\text{CO}_3)_2]$. The unusual silicate carbonate, *SA tilleyite*, was discovered in grains up to 15 μm maximum dimension as a minor phase in association with secondary wollastonite, grossular, and monticellite in a void space within a forsterite-bearing Type B CAI in the Allende CV chondrite (A. Krot, personal communication, 11 October 2020).

Organic minerals

Whewellite $[\text{Ca}(\text{C}_2\text{O}_4)\cdot\text{H}_2\text{O}]$. Fuchs et al. (1973) reported the calcium oxalate, *SA whewellite*, in proximity to olivine grains in a so-called “white inclusion” in the Murchison CM2 chondrite. Whewellite is found in association with olivine, calcite, and organic matter (kerogen). They suggest that this occurrence points to equilibration below 480 °C, at which temperature calcium oxalate decomposes to CaCO_3 and CO .

Amorphous phases

A wide range of amorphous phases occur in the fine-grained matrices and altered portions of chondrite meteorites. Silicate glass and “amorphous silicate matrices” (ASM) of varying compositions, including ferromagnesian, feldspathic, and silica-rich, are documented from numerous chondrite and achondrite meteorites (e.g., Brearley and Jones 1998, and references therein; Zolensky and Ivanov 2003; Hopp and Vollmer 2018). Origins have been ascribed to primary condensation in nebular environments, quenching of molten silicates, and impact melting/amorphization (Davoisne et al. 2006; Keller and Messenger 2011; Ruzicka 2014; Lunning et al. 2016; Rubin and Ma 2021). However, with the exception of the formation of quasi-amorphous layer silicates such as TCI (Tomeoka and Buseck 1985, 1990; see above) or the alteration of a preexisting amorphous phase by hydration and oxidation (Hopp and Vollmer 2018), these silicate phases have not been attributed to secondary processes and are not listed here.

Kerogen $(\text{C},\text{H},\text{O},\text{N},\text{S})$. Disordered carbonaceous material, here lumped under *SA kerogen*, is a significant amorphous component of many carbonaceous chondrites, with C comprising ~2.2 wt% of the Murchison CM2 chondrite (Fuchs et al. 1973) and more than 5 wt% of the Nagoya CM2 chondrite (Bunch and Chang 1980). The matrix of the Vigarano CV3 carbonaceous chondrite contains amorphous C-rich material that includes poorly graphitized carbon as well as kerogen-like material with O, N, and H. Kerogen may also contain significant fractions of polycyclic aromatic hydrocarbons (PAHs) and fullerenes (Buseck and Hua 1993). Organic matter in carbonaceous and enstatite chondrites holds evidence for significant thermal processing (Cody et al. 2008; Piani et al. 2012; Kebukawa et al. 2019). In some cases, as with regions of highly disordered X-ray- and electron-amorphous layered hydrous silicates (Brearley 1993a; Greshake 1997), a continuum may exist between crystalline, “poorly graphitized,” and non-crystalline volumes of carbonaceous material (Abreu and Brearley 2011).

IMPLICATIONS

Aqueous alteration and/or thermal metamorphism in planetesimals dramatically increased the chemical and structural diversity of the preterrestrial mineral kingdom. In Parts I through V, we have now tabulated 447 historical natural kinds of miner-

als representing 263 IMA-approved mineral species plus 18 as yet unapproved crystalline phases and 16 amorphous phases. Of this total of 297 diverse minerals, 119 phases (i.e., 40%) are new to Part V, including the earliest known examples of halides, arsenides, tellurides, sulfates, carbonates, hydroxides, and a wide range of phyllosilicates.

Secondary processes also dramatically increased the chemical diversity of minerals with 41 different essential (i.e., species-defining) elements, including the earliest known appearances of essential Co, Ge, As, Nb, Ag, Sn, Te, Au, Hg, Pb, and Bi. Nevertheless, as with earlier stages of mineral evolution, secondary meteorite minerals are dominated volumetrically by relatively few phases. We estimate that only 34 minerals of the 166 listed in Table 1 (see asterisked species in Table 1) occur widely or ever exceed 1 vol%. Those more common secondary minerals, furthermore, incorporate only 15 different essential elements, all of which are relatively abundant: H, C, O, S, P, Cl, Na, Mg, Ca, Fe, Ni, Al, Cr, Si, and Ti. By contrast, at least 94 of the 166 minerals in Table 1 are known as volumetrically trivial phases from only one or two meteorite groups. Thus, as in many other mineral-rich environments, relatively few mineral species are common, whereas most are rare (Hazen et al. 2015; Hystad et al. 2015a, 2015b; Hazen and Ausubel 2016).

Our studies of the evolving distribution and diversity of pre-terrestrial minerals, especially in the context of the numerous new secondary minerals formed by aqueous alteration and/or thermal metamorphism, raise several intriguing questions:

- What are the relative roles of temperature, pressure, bulk chemistry, and time in mineral diversification? For example, wet environments appear to display significant mineralogical diversity, with a greater number of essential chemical elements, compared to anhydrous environments. Is it possible to identify the relative influences on mineral diversity of water as a solvent that mobilizes and concentrates many elements, vs. the distinctive crystal chemical roles of OH⁻ and H₂O molecules?

- To what extent are high-temperature mineral assemblages, such as those of primary CAI condensates or primary igneous phases in chondrules, intrinsically less diverse than lower-temperature assemblages? And is there a similar effect related to pressure?

- Can we employ statistical methods of mineral ecology (e.g., Hystad et al. 2019) to predict the number, as well as the nature and contexts, of meteorite minerals that exist but have not yet been discovered and described?

- In spite of the increased number of essential elements represented in secondary meteorite minerals, several important mineral-forming elements, including Li, Be, B, Ga, Se, Sb, Rb, Sr, La, and rare earth elements, Th, and U are not represented as essential elements in any known pre-terrestrial mineral (though they are well documented as minor or trace elements in meteorites). Do these elements occur primarily in solid solution (i.e., Ga for Al in feldspar), or might they form nanoscale inclusions or concentrate in grain boundaries?

- A common characteristic of natural evolving systems, including the mineral kingdom, is congruent complexification—a logical sequence of processes that modify and expand diversity of historical natural kinds (Hazen et al. 2008; Hazen and Eldredge 2010). Can we document and visualize an increase in the average

chemical and structural complexity of minerals through these earliest stages of mineral evolution (e.g., Krivovichev et al. 2017)?

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