Carbon Mineralogy and Crystal Chemistry

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INTRODUCTION

Carbon, element 6, displays remarkable chemical flexibility and thus is unique in the diversity of its mineralogical roles. Carbon has the ability to bond to itself and to more than 80 other elements in a variety of bonding topologies, most commonly in 2-, 3-, and 4-coordination. With oxidation numbers ranging from −4 to +4, carbon is observed to behave as a cation, as an anion, and as a neutral species in phases with an astonishing range of crystal structures, chemical bonding, and physical and chemical properties. This versatile element concentrates in dozens of different Earth repositories, from the atmosphere and oceans to the crust, mantle, and core, including solids, liquids, and gases as both a major and trace element (Holland 1984; Berner 2004; Hazen et al. 2012). Therefore, any comprehensive survey of carbon in Earth must consider the broad range of carbon-bearing phases.

The objective of this chapter is to review the mineralogy and crystal chemistry of carbon, with a focus primarily on phases in which carbon is an essential element: most notably the polymorphs of carbon, the carbides, and the carbonates. The possible role of trace carbon in nominally acarbonaceous silicates and oxides, though potentially a large and undocumented reservoir of the mantle and core (Wood 1993; Jana and Walker 1997; Freund et al. 2001; McDonough 2003; Keppler et al. 2003; Shcheka et al. 2006; Dasgupta 2013; Ni and Keppler 2013; Wood et al. 2013), is not considered here. Non-mineralogical carbon-bearing phases

The International Mineralogical Association (IMA) recognizes more than 380 carbon-bearing minerals (http://rruff.info/ima/), including carbon polymorphs, carbides, carbonates, and a variety of minerals that incorporate organic carbon in the form of molecular crystals, organic anions, or clathrates. This chapter reviews systematically carbon mineralogy and crystal chemistry, with a focus on those phases most likely to play a role in the crust. Additional high-temperature and high-pressure carbon-bearing minerals that may play a role in the mantle and core are considered in the next chapter on deep carbon mineralogy (Oganov et al. 2013).

**SYSTEMATIC CARBON MINERALOGY**

Carbon, a non-metal that typically forms covalent bonds with a variety of other elements, is the most chemically adaptable element of the periodic table. In an ionic sense, element 6 can act as a cation with oxidation number +4, as in carbon dioxide (CO$_2$) or in the carbonate anion (CO$_3$$^{2-}$). Alternatively, carbon can act as an anion with oxidation number as low as −4, as in methane (CH$_4$) and in other alkanes. Carbon also frequently displays a range of intermediate oxidation number states from +2 in carbon monoxide (CO) or −2 in methanol (CH$_3$OH), as well as occurring in its neutral state (C) in a variety of carbon allotropes.

Carbon chemistry is also enriched by the ability of C to form single, double, or triple bonds, both with itself and with a wide range of other chemical elements. In many carbon compounds each C atom bonds to 4 other atoms by single bonds, as in methane and carbon tetrafluoride (CF$_4$). But carbon commonly forms double bonds with itself, for example in ethene (H$_2$C=CH$_2$), oxygen in carbon dioxide (O=C=O), or sulfur in carbon disulfide (S=C=S), and it can form triple bonds with itself, as in ethylene (commonly known as acetylene; HC≡CH), or with nitrogen, as in hydrogen cyanide (HC≡N). Carbon’s remarkably diverse mineralogy arises in part from this unmatched range of valence states and bond types.

In the following sections we review the systematic mineralogy of carbon, including carbon allotropes, carbides, carbonates, and minerals that incorporate organic carbon molecules.

**Carbon allotropes**

The element carbon occurs in several allotropes, including graphite, diamond, lonsdaleite, fullerenes (including buckyballs and carbon nanotubes), graphene, and several non-crystalline forms (Table 1). These varied carbon allotropes exhibit extremes in physical and chemical properties—variations that reflect differences in their atomic structures. Diamond and lonsdaleite are the hardest known substances, whereas graphite is among the softest. Transparent diamond is an exceptional electrical insulator while possessing the highest known thermal conductivity at room temperature, whereas opaque graphite is an electrical conductor and thermal insulator. Isotropic diamond is widely used as a tough abrasive, whereas anisotropic graphite is employed as a lubricant. Diamond possesses the highest nuclear density of any known condensed phase at ambient conditions (Zhu et al. 2011) and is predicted to have high-pressure phases of even greater density (Oganov et al. 2013), whereas some carbon nanogels feature among the lowest known nuclear densities.
<table>
<thead>
<tr>
<th>Name</th>
<th>Structure Description</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>Graphite</td>
<td>Hexagonal; stacked flat layers of 3-coordinated $sp^2$ C</td>
<td>Klein and Hurlbut (1993)</td>
</tr>
<tr>
<td>Diamond</td>
<td>Cubic; framework of 4-coordinated $sp^3$ C</td>
<td>Harlow (1998)</td>
</tr>
<tr>
<td>Lonsdaleite</td>
<td>Hexagonal; framework of 4-coordinated $sp^3$ C</td>
<td>Frondel and Marvin (1967a); Bundy (1967)</td>
</tr>
<tr>
<td>Chaoite (also “Ceraphite”)</td>
<td>A disputed form of shocked graphitic C from impact sites</td>
<td>El Goresy and Donnay (1968); McKee (1973)</td>
</tr>
<tr>
<td>Fullerenes</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Buckyballs</td>
<td>Closed cage molecules $sp^2$ C: C$<em>{60}$, C$</em>{70}$, C$_{76}$, etc.</td>
<td>Kroto et al. (1985)</td>
</tr>
<tr>
<td>Nano onions</td>
<td>Multiple nested buckyballs with closed cages</td>
<td>Sano et al. (2001)</td>
</tr>
<tr>
<td>Nanotubes</td>
<td>Cylindrical fibers of $sp^2$ C, single tubes or nested</td>
<td>Belluci (2005)</td>
</tr>
<tr>
<td>Nanobuds</td>
<td>Buckyballs covalently bonded to the exteriors of nanotubes</td>
<td>Nasibulin et al. (2007)</td>
</tr>
<tr>
<td>Buckyball and chain</td>
<td>Two buckyballs linked by a carbon chain</td>
<td>Shvartsburg et al. (1999)</td>
</tr>
<tr>
<td>Graphene</td>
<td>One-atom-thick graphitic layers with $sp^2$ bonding</td>
<td>Geim and Novoselov (2007)</td>
</tr>
<tr>
<td>Carbon Nanofibers (also known as “Graphite Whiskers”)</td>
<td>Graphene layers arranged as cones, cups, or plates</td>
<td>Hatano et al. (1985); Morgan (2005)</td>
</tr>
<tr>
<td>Amorphous Carbon</td>
<td>Non-crystalline carbon, with disordered $sp^2$ and $sp^3$ bonds. Sometimes inaccurately applied to soot and coal.</td>
<td>Rundel (2001)</td>
</tr>
<tr>
<td>Glassy Carbon (also “Vitreous Carbon”)</td>
<td>Dense, hard, non-crystalline 3-dimensional arrangement of $sp^3$ bonds, possibly related to a fullerene structure</td>
<td>Cowlard and Lewis (1967)</td>
</tr>
<tr>
<td>Carbon Nanofoam (also “Carbon Aerogel”)</td>
<td>Nano-clusters of carbon in 6- and 7-rings, forming a low-density three-dimensional web.</td>
<td>Rode et al. (2000)</td>
</tr>
<tr>
<td>Linear Acetylenic Carbon (also known as Carbyne)</td>
<td>Linear molecule of acetylenic C ($\cdots$C=C=C=\cdots)</td>
<td>Heimann et al. (1999)</td>
</tr>
</tbody>
</table>
Here we focus on the essential characteristics of the three most common minerals of native carbon—graphite, diamond, and lonsdaleite—all of which play roles in Earth’s subsurface carbon cycle. For more comprehensive reviews of the chemical and physical properties of these carbon polymorphs see Bragg et al. (1965), Deer et al. (1966), Field (1979), Davies (1984), Klein and Hurlbut (1993), Harlow (1998), and Zaitsev (2001).

**Graphite.** Carbon forms covalent bonds with itself in all of the carbon polymorphs, in which C adopts one of two coordination environments (Table 1). In graphite, graphene, buckyballs, nanotubes, and several types of amorphous and glassy carbon, the carbon atoms are in planar three-coordination. This trigonal coordination, with typical C-C distances of ~1.42 Å and C-C-C angles close to 120°, results from the hybridization of carbon’s electrons into three orbitals, known as the \( \text{sp}^2 \) bonding configuration because the 2s orbital mixes (or hybridizes) with two 2p orbitals to form three \( \text{sp}^2 \) orbitals.

The hexagonal layered structure of graphite (Fig. 1a; Animation 1) features electrically neutral, monoatomic, flat carbon layers, which bond to each other through van der Waals attractions and which are separated from each other by 3.41 Å. This layered structure leads to the distinctive, highly anisotropic properties of graphite (Fig. 1b). Individual carbon layers, which are known as graphene when meticulously separated from a graphite crystal or vapor deposited, are extremely strong and display unique electronic and mechanical properties (Geim and Novoselov 2007; Geim 2009). Weak van der Waals forces between these layers lead to graphite’s applications in lubricants and as pencil “lead.”

Natural graphite is a common crustal mineral that occurs most abundantly in metamorphic rocks in pods and veins as a consequence of the reduction and dehydration of sediments rich in carbon (e.g., Klein and Hurlbut 1993). Commercial metamorphic graphite deposits, notably from China, India, and Brazil, are exploited in the manufacture of pencils, lubricants, steel, and brake linings. Graphite is also found as an accessory mineral in some igneous rocks and as a micro- or nano-phase in a variety of meteorites. In these diverse lithologies graphite is commonly precipitated in veins from reduced C-O-H fluids (Rumble and Hoering 1986; Rumble et al. 1986). In addition, graphite can crystallize from the vapor state in the carbon-rich, expanding hot envelopes of energetic late-stage stars, including supernovas.

**Figure 1.** Graphite. (a) The graphite (native C) crystal structure (hexagonal, space group \( P6_3/mmc; a = 2.464 \text{ Å}; c = 6.736 \text{ Å}; Z = 4 \)) incorporates monoatomic layers of 3-coordinated carbon, linked by van der Waals interactions. (b) Natural graphite crystals reflect the hexagonal crystal structure. This 1 mm diameter crystal displays basal pinacoid and prism faces. The crystal, associated with calcite, is from the Crestmore quarries, Riverside Co., California. Photo courtesy of John A. Jaszczak. [Animation 1: For readers of the electronic version, click the image for an animation of the graphite crystal structure.]
**Diamond and lonsdaleite.** In contrast to the \(sp^2\) bonding environment of carbon in graphite, each carbon atom can bind to 4 adjacent C atoms in tetrahedral coordination, as exemplified by the diamond and lonsdaleite polymorphs (Figs. 2 and 3; Animations 2 and 3). The C-C distance in these minerals are \(\sim 1.54\) Å, while C-C-C angles are close to the ideal tetrahedral value of 109.5°. This tetrahedral bonding configuration reflects the hybridization of one 2s and three 2p orbitals from each carbon atom to form four \(sp^3\) orbitals.

The structures of cubic diamond and hexagonal lonsdaleite are similar: both forms of carbon feature tetrahedral coordination of C in a three-dimensional framework. However, given a flat layer of linked carbon tetrahedra with all vertices pointed in the same direction (layer A), there exist two ways to stack subsequent layers, in orientations described as B or C. The diamond structure represents a three-layer stacking sequence of \[\ldots ABCABC\ldots\] along the (111) cubic direction, whereas lonsdaleite stacking is two-layer \[\ldots ABAB\ldots\] along the (001) hexagonal direction. A similar stacking difference distinguishes the cubic and hexagonal forms of a number of topologically similar mineral pairs, such as the polymorphs of zinc sulfide (ZnS), sphalerite and wurtzite, respectively. Note also that the stacking sequence of these carbon polymorphs can incorporate errors, as is often the case with lonsdaleite that has formed by impact shock of graphite (Frondel and Marvin 1967).

**Figure 2.** (a) The diamond (native C) crystal structure (cubic, space group \(Fd\overline{3}m; a = 3.560\) Å; \(Z = 8\)) features a framework of tetrahedrally coordinated carbon atoms. (b) A natural diamond crystal reflects the cubic crystal structure. The semi-translucent diamond cube (ref. no. S014632) is \(24.3 \times 21.8 \times 21.7\) mm in size and weighs 156.381 carats (31.3 gm) and is shown with a 1-carat diamond for scale. It is represented to be from Ghana. Photo courtesy of Harold and Erica Van Pelt. [Animation 2: For readers of the electronic version, click the image for an animation of the diamond crystal structure.]

**Figure 3.** The lonsdaleite (native C) crystal structure (hexagonal, space group \(P6_3/mmc; a = 2.52\) Å; \(c = 4.12\) Å; \(Z = 4\)). [Animation 3: For readers of the electronic version, click the image for an animation of the lonsdaleite crystal structure.]
The framework structures of diamond and lonsdaleite lead to their superlative physical and chemical properties (Davies 1984). Notably, the exceptional hardness and strength of these phases arises from the strong three-dimensional network of C-C bonds. Recent theoretical studies on diamond and lonsdaleite suggest that diamond possesses the greater strength (Pan et al. 2009; Lyakhov and Oganov 2011).

The key to understanding the contrasting properties of the natural carbon allotropes is their different pressure-temperature stability fields. The deep origin of diamond was first recognized in the years following the widely publicized discovery of the 20-carat Eureka diamond by children playing in a dry central South African streambed in 1866, and the even more dramatic 83.5-carat Star of South Africa diamond two years later. A subsequent diamond rush brought more than 10,000 prospectors to the semi-desert region, and inevitably led to the recognition of diamond in their volcanic host rock, dubbed kimberlite (Davies 1984; Hazen 1999). The South African kimberlites’ cone-shaped deposits, which cut vertically through shattered country rock, spoke of violent explosive eruptions from great depth (Lewis 1887; Bergman 1987; Mitchell 1995).

The hypothesis that diamond comes from depth was reinforced by determination of the crystal structures of diamond versus graphite (Bragg and Bragg 1913; Hull 1917; Hassel and Mark 1924; Bernal 1924). The higher coordination number of carbon in diamond (4-fold) compared to graphite (3-fold), coupled with its much greater density (3.51 vs. 2.23 g/cm³), provided physical proof that diamond was the higher-pressure polymorph. Early determinations of the carbon phase diagram (Rossini and Jessup 1938), though refined in subsequent years (Bundy et al. 1961; Kennedy and Kennedy 1976; Day 2012; Fig. 4), revealed that diamond is the higher-pressure, lower-temperature form and that, under a normal continental geotherm, diamond likely forms at depths greater than 100 kilometers. By contrast, the steeper geothermal gradients of the oceanic crust and mantle preclude diamond formation, at least within the sub-oceanic upper mantle and transition zone.

Efforts to synthesize diamond under extreme laboratory conditions extend back to the early 1800s, long before the crucial role of high-pressure was recognized (Mellor 1924;
Among the most renowned 19th-century chemists to try his hand at diamond making was Frederick-Henri Moissan, who won the Nobel Prize for his risky isolation of the dangerous element fluorine. Moissan employed a novel electric arc furnace to generate record temperatures ~3000 °C. Moissan initially thought he was successful in synthesizing diamond, but is now known to have formed hard, transparent crystals of silicon carbide—a compound that he also discovered in nature, and what is now known as the mineral moissanite (Moissan 1904a; Hazen 1999; see below). Numerous other heroic efforts prior to 1950 also failed (Bridgman 1931, 1946; von Platen 1962; Coes 1962). In spite of the relatively accessible pressure-temperature regime of diamond stability > 4 GPa (approximately 40,000 atm or 40 kbar), the transition from graphite to diamond is kinetically inhibited. The keys to facile diamond synthesis—employing a liquid metal flux coupled with sustained pressures above 5 GPa and temperatures above 1200 °C—were not achieved until the post-World War II efforts by scientists at the General Electric Research Laboratory in Schenectady, New York (Bundy et al. 1955; Suits 1960, 1965; Wentorf 1962; Hall 1970; Strong 1989; Hazen 1999). The breakthrough of high-pressure diamond synthesis by Francis Bundy, Tracy Hall, Herbert Strong, and Robert Wentorf in December of 1954 led to what is today a multi-billion dollar industry that supplies annually hundreds of tons of industrial diamond abrasives.

Two other synthesis techniques have expanded the varieties of diamond available for commercial exploitation. Of special interest are efforts to generate diamond and lonsdaleite under shock conditions that mimic bolide impact events (DeCarli and Jamieson 1961; Beard 1988). In some meteorites lonsdaleite and/or diamond is found to replace graphite in crystallites that retain a hexagonal shape (Langenhorst et al. 1999; El Goresy et al. 2001; Langenhorst and Deutsch 2012). In this rapid, solid-state martensitic transition the flat, graphitic $sp^2$ planes of carbon atoms shift relative to each other and buckle to produce a $sp^3$ array of C layers that are rather disorderly in their stacking arrangement.

The subsequent discovery of techniques for diamond and lonsdaleite synthesis by vapor deposition at low-pressure conditions that mimic diamond formation in expanding stellar envelopes has greatly increased the potential for diamond use in science and industry (Angus et al. 1968; Derjaguin and Fedoseev 1968; Angus and Hayman 1988; Spear and Dismukes 1994; Irifune and Hemley 2012). These varied efforts in diamond synthesis have produced exceptional new materials, including isotopically pure diamonds with the highest recorded thermal conductivity, semiconducting diamonds, nano-crystalline polishing powders, lonsdaleite crystals that are harder than many natural diamond, and a range of deeply colored flawless synthetic gemstones up to 10 carats (Liang et al. 2009; Meng et al. 2012).

Intense research on natural diamond is also providing important insights regarding Earth’s geochemical and tectonic evolution. A number of recent studies focus on diamond’s rich and revealing suites of oxide, silicate, carbide, and sulfide inclusions from depths of up to perhaps 850 kilometers—mantle samples that provide evidence for aspects of geochemical and tectonic evolution over more than 3 billion years of Earth history (Shirey et al. 2002; McCammon et al. 2004; Sommer et al. 2007; Pearson et al. 2007; Gübelin and Koivula 2008; Shirey and Richardson 2011; Walter et al. 2011; Shirey et al. 2013).

**Carbides**

Carbides, which form when carbon bonds to a less electronegative element, are represented by dozens of synthetic compounds that have a range of industrial applications (Ettmayer and Lengauer 1994). The International Mineralogical Association has recognized 10 different naturally occurring carbide minerals (Tables 2 and 3; http://rruff.info/ima/). Although rare and volumetrically trivial as reservoirs of carbon in the crust, they may represent a significant volume of carbon in Earth’s deep interior, and thus may provide insight to the deep carbon cycle (Dasgupta 2013; Wood et al. 2013).
All metal carbides are refractory minerals; they have relatively high solidus and liquidus temperatures, with melting points typically above 2400 °C (Nadler and Kempter 1960; Lattimer and Grossman 1978). A number of these minerals are found in association with diamond and other high-pressure phases, as well as with assemblages of unusual reduced minerals, including native Al, Fe, Si, Sn, W, and more than a dozen other native metallic elements, as well as exotic sulfides, phosphides, and silicides (Table 4). Carbides, along with diamond, may thus represent Earth’s deepest surviving minerals, and may prove a relatively unexplored window on the nature of the deep mantle environment and the deep carbon cycle.

Moissanite. Moissanite (α-SiC), also known commercially as carborundum when sold as an abrasive (hardness 9.5), is the most common of the natural carbides. Produced synthetically for more than a century (Acheson 1893), moissanite is used in numerous applications, including automobile parts (e.g., brakes and clutches), bulletproof vests, light-emitting diodes, semiconductor components, anvils for high-pressure research and, since 1998, it has been marketed as inexpensive diamond-like artificial gemstones (Bhatnagar and Baliga 1993; Xu and Mao 2000; Madar 2004; Saddow and Agarwal 2004).

Since its discovery in 1893 by Henri Moissan in mineral residues from the Canyon Diablo meteor crater in Arizona (Moissan 1904b), natural moissanite has been found in dozens of localities, including meteorites, serpentinites, chromitites, ophiolite complexes, and in close association with diamond in kimberlites and eclogites (Lyakhovich 1980; Leung et al. 1990; Alexander 1990, 1993; Di Pierro et al. 2003; Lee et al. 2006; Qi et al. 2007; Xu et al. 2008; Trumbull et al. 2009; Shiryaev et al 2011; see also http://MinDat.org). While some occurrences of moissanite are apparently of near-surface origin, including sites of forest fires and contact metamorphism of silicate magmas with coal beds (Sameshima and Rodgers 1990), silicon carbide also represents one of the deepest mantle minerals known to reach the surface. The discovery of moissanite inclusions in diamond (Moore and Gurney 1989; Otter and Gurney 1989; Leung 1990; Gorshkov et al. 1997), combined with observations of native silicon and Fe-Si alloy inclusions in moissanite (Trumbull et al. 2009), may point to an origin in reduced mantle microenvironments. However, further research is required to determine the range of oxygen fugacities under which moissanite is stable at mantle pressures and temperatures.

Occurrences of silicon carbide in mantle-derived kimberlites and several ophiolite complexes reflect its stability at high pressure and very low oxygen fugacity (Mathez et al. 1995). Secondary ion mass spectrometric (SIMS) analysis shows that ophiolite-hosted

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**Table 2. Natural carbide minerals (data compiled from http://MinDat.Org locality and species databases).**

<table>
<thead>
<tr>
<th>Mineral (Key)</th>
<th>Formula</th>
<th>Paragenesis*</th>
<th>Locality register**</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cohenite (C)</td>
<td>(Fe,Ni,Co)₃C</td>
<td>1,2,4</td>
<td>G1, G2, M1, R1, R2</td>
</tr>
<tr>
<td>Haxonite (H)</td>
<td>(Fe,Ni)₂C₅</td>
<td>2</td>
<td>M1</td>
</tr>
<tr>
<td>Isovite (I)</td>
<td>(Cr,Fe)₃C₆</td>
<td>4</td>
<td>R3, R4</td>
</tr>
<tr>
<td>Khamrabaevite (K)</td>
<td>(Ti,V,Fe)C</td>
<td>2,4?</td>
<td>M1, T1, U1, U2</td>
</tr>
<tr>
<td>Moissanite (M)</td>
<td>SiC</td>
<td>2,3,4</td>
<td>C1-C6, M1, R5, R6, S1, T2, U2</td>
</tr>
<tr>
<td>Niobocarbide (N)</td>
<td>(Nb,Ta)C</td>
<td>4</td>
<td>R7</td>
</tr>
<tr>
<td>Qusongite (Q)</td>
<td>WC</td>
<td>1,3,4</td>
<td>C3, C4, C7</td>
</tr>
<tr>
<td>Tantalcarbide (Ta)</td>
<td>(Ta,Nb)C</td>
<td>4</td>
<td>R7</td>
</tr>
<tr>
<td>Tongbaite (To)</td>
<td>Cr₂C₂</td>
<td>4</td>
<td>C8, R3</td>
</tr>
<tr>
<td>Yarlongite (Y)</td>
<td>(Cr₂Fe₂Ni)C₄</td>
<td>4</td>
<td>C4</td>
</tr>
</tbody>
</table>

*1 = coal fire or intrusion in coal/graphite; 2 = meteorite; 3 = kimberlite; 4 = other ultramafic
**See Table 3 for locality key
Table 3. Natural carbide locality register, with associated reduced and/or high-P mineral species (data compiled from http://MinDat.Org locality and species databases). Several moissanite localities of poorly-defined paragenesis are not included.

<table>
<thead>
<tr>
<th>Code</th>
<th>Locality</th>
<th>Carbides</th>
<th>Type</th>
<th>Associated Phases</th>
</tr>
</thead>
<tbody>
<tr>
<td>C1</td>
<td>Jeffrey Mine, Quebec</td>
<td>M</td>
<td>Ser</td>
<td>Gra, Cu, Awa, IMA, Hea, Pyr, Sha</td>
</tr>
<tr>
<td>C2</td>
<td>Fuxian kimberlite field, Liaoning Prov.</td>
<td>M</td>
<td>Kim</td>
<td>Dia, Gra, Lon, Fe</td>
</tr>
<tr>
<td>C3</td>
<td>Mengyin kimberlite field, Linyi Prefecture</td>
<td>M, Q</td>
<td>Kim</td>
<td>Dia, Cr, Fe, Pb, W, Pen, Pol, Coe</td>
</tr>
<tr>
<td>C4</td>
<td>Luobusa ophiolite, Tibet</td>
<td>M, Q, Y</td>
<td>Oph</td>
<td>Dia, Gra, Al, Cr, Cu, In, Fe, Ni, Os, Pd, Rh, Ru, Si, Ag, Sn, Ti, W, Zn, Zan, Ala, Wus, Luo</td>
</tr>
<tr>
<td>C5</td>
<td>Picrite lava outcrop, Yunnan Prov.</td>
<td>M</td>
<td>Ser</td>
<td>Cu, Zn</td>
</tr>
<tr>
<td>C6</td>
<td>Dongjiashan Hill, Dabie Mountains</td>
<td>M</td>
<td>Ser</td>
<td>Dia, Si</td>
</tr>
<tr>
<td>C7</td>
<td>Yangliuping N-Cu-PGE Deposit, Sichuan Prov.</td>
<td>Q</td>
<td>UGr</td>
<td>Cr, Pt, Dan, Cub, Lau, Lin, Vio</td>
</tr>
<tr>
<td>C8</td>
<td>Liu Village, Henan Prov.</td>
<td>To</td>
<td>Ult</td>
<td>Cr, Cu, Fe, Pb, Alt, Awa, Bi2, Cub, Pen, Pyr, Vio,</td>
</tr>
<tr>
<td>G1</td>
<td>Bühl, Weimar, Kassel, Hesse</td>
<td>C</td>
<td>MCo</td>
<td>Fe</td>
</tr>
<tr>
<td>G2</td>
<td>Disko Island, Kitaa Prov.</td>
<td>C</td>
<td>Ult</td>
<td>Gra, Fe, Pb, Arm, Sha, Tro, Sch</td>
</tr>
<tr>
<td>M1</td>
<td>Various localities</td>
<td>C, H, K, M</td>
<td></td>
<td></td>
</tr>
<tr>
<td>R1</td>
<td>Khungtukun Massif, Khatanga, Siberia</td>
<td>C</td>
<td>MCo</td>
<td>Gra, Cu, Fe, Tro, Wus</td>
</tr>
<tr>
<td>R2</td>
<td>Coal Mine 45, Kopeisk, Urals</td>
<td>C</td>
<td>CoF</td>
<td>Gra, Tro</td>
</tr>
<tr>
<td>R3</td>
<td>Is River, Isovsky District, Urals</td>
<td>I, To</td>
<td>Pla</td>
<td>Os, Pt, Iso, Tet, Cup, Mal, Fer, Gup, Xif</td>
</tr>
<tr>
<td>R4</td>
<td>Verkhneivinsk, Neiva River, Urals</td>
<td>I</td>
<td>Pla</td>
<td>Os, Ru, Any</td>
</tr>
<tr>
<td>R5</td>
<td>Billeekh intrusion, Saha Rep., Siberia</td>
<td>M</td>
<td>Gab</td>
<td>Al, Sb, Cd, Cu, Fe, Pb, Sn, Zn</td>
</tr>
<tr>
<td>R6</td>
<td>Avacha volcano, Kamchatka</td>
<td>M</td>
<td>Pic</td>
<td>Dia</td>
</tr>
<tr>
<td>R7</td>
<td>Avorinskii Placer, Baranchinsky Massif, Urals</td>
<td>N, Ta</td>
<td>Pla</td>
<td>Jed, Ru</td>
</tr>
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<td>Monastery Mine, Free State Prov.</td>
<td>M</td>
<td>Kim</td>
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<td>T1</td>
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<td>K</td>
<td>Gra, Bi, Fe, Bis</td>
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<tr>
<td>U1</td>
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<td>K</td>
<td>??</td>
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<tr>
<td>U2</td>
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<td>Ala, Mav</td>
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</table>

*See Table 2 for key to carbides
**CoF = coal fire; GCo = gabbro intrusion in coal; Gra = granodiorite; Kim = kimberlite; Lam = lamproite; Oph = ophiolite; Pic = picrite; Pla = placer; Ser = serpentinite; UGr = ultramafic intrusion in graphite schist; Ult = unspecified ultramafic
***See Table 4 for key to associated phases
Table 4. Select reduced and/or high-\(P\) mineral species associated with natural carbides (data compiled from http://MinDat.Org locality and species databases). For locality key see Table 3.

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<th>Mineral (Key)</th>
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<td>Danbaite (Dan)</td>
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<td>Mavlyanovite (Mav)</td>
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<td><strong>Phosphides</strong></td>
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<td>R3</td>
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<td>Jedwabite (Jed)</td>
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<td>C1,G2,U1</td>
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<tr>
<td>Phosphorus (P)</td>
<td>(Fe,Ni)3P</td>
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moissanite has a distinctive $^{13}$C-depleted isotopic composition ($\delta^{13}$C from −18 to −35‰, $n = 36$), which is significantly lighter than the main carbon reservoir in the upper mantle ($\delta^{13}$C near −5‰). Alternatively, significant isotope fractionation between carbide and diamond has been observed in high-pressure experiments (−7 per mil at 5 GPa; Mikhail et al. 2010; Mikhail 2011), greatly complicating the potential for identification of carbon reservoirs through carbon isotope systematics of mantle-derived samples (Mikhail et al. 2011). It has been suggested that moissanite may also occur in the lower mantle, where the existence of $^{13}$C-depleted carbon is strongly supported by studies of extraterrestrial carbon (Trumbull et al. 2009).

Moissanite, like diamond, also forms by vapor deposition (Hough et al. 1997) and it is relatively common in space in the envelopes of carbon-rich AGB stars. It is subsequently carried as a pre-solar guest in carbonaceous chondrites. The origins of moissanite and other carbides have been inferred from unusual variations in both C-isotopes and N-isotopes (Daulton et al. 2003). Meteoritic carbides commonly contain dissolved nitrogen, while the comparable family of nitride minerals [e.g., osbornite (TiN)] contains some dissolved carbon. This mutual limited solubility of N-C in minerals persists through natural diamond (C), which also contains minor N, and in the future may be useful for understanding crystallization histories and source reservoirs.

Moissanite is also relatively widely distributed as micro-crystals in the ejecta from some meteorite impact craters formed in continental crust, and it may be associated with impact diamond and a variety of iron silicides like suessite (Ernston et al. 2010). Low-pressure SiC is also found in the KT impact layer (Hough et al. 1995; Langenhorst and Deutsch 2012), though unlike impact diamond, SiC is not ubiquitous in crustal impact deposits (Gilmour et al. 2003). Indeed, another possibility for deep SiC formation, given the antiquity of some kimberlite/diamond hosted SiC, might be residues from giant impact processes during formation of Earth’s Moon, since at that time materials from a cross-section through the upper mantle were violently exposed to the vacuum of space.

Ideal moissanite has a hexagonal structure closely related to that of lonsdaleite (and identical to wurtzite), in which every atom is tetrahedrally coordinated and corner-linked tetrahedral layers are stacked ideally in a two-layer [...]ABAB […] configuration (Fig. 5). The Si-C distance of 1.86 Å is appreciably longer than that of the carbon polymorphs because of the greater size of Si compared to C. Moissanite is known to recrystallize at temperatures between 1400 and 1600 °C to $\beta$-SiC, which is cubic and similar to the diamond structure.

More than 250 stacking polytypes of SiC have been documented, notably hexagonal forms with 4-layer (4H) and 6-layer (6H—the most common terrestrial polytype; Capitani et al. 2007) repeats {[…]ABAC[…] and […]ABCACB[…], respectively, and a rhombohedral form with a 15-layer (15R) sequence […]ABCBACABACBCACB…}. Hundreds of other polytypes with repeat sequences from dozens to hundreds of layers (i.e., 141R and 393R) have also

Figure 5. The moissanite (SiC) crystal structure (hexagonal, space group $P6_3mc$; $a = 3.081$ Å; $c = 5.031$ Å; $Z = 2$). Carbon and silicon atoms appear in blue and grey, respectively.
been characterized (Krishna and Verma 1965; Lee et al. 2006; Capitani et al. 2007; Shiryaev et al. 2011; see also http://img.chem.ucl.ac.uk/www/kelly/LITERATURESICWEB.HTM#4 and http://MinDat.org). Such complex polytypes have inspired a variety of models related to possible growth models, including screw dislocation or spiral growth mechanism (e.g., Frank 1949, 1951; Verma and Krishna 1966), the faulted matrix model (Pandey and Krishna 1975a, 1975b, 1978), one-dimensional disorder theory (Jagodzinski 1954a, 1954b), and the axial nearest neighbor Ising model (Price and Yeomans 1984).

A curious crystal chemical aspect related to silicon carbide is the apparent absence of Si-C bonding in other naturally occurring compounds (Nawrocki 1997; Franz 2007; Tran et al. 2011). Chemists have explored a rich landscape of synthetic organic silanes, silanols, and silicones, but natural examples of these potentially crystal-forming compounds have not yet been described.

Cohenite. The iron carbide cohenite \([(Fe,Ni,Co)\text{C}\)], also called cementite when it occurs as a bonding agent in steel, is second in abundance as a natural carbide only to moissanite. In nature cohenite is known primarily as an accessory mineral from more than a dozen iron meteorites (Brett 1967), but it also occurs occasionally with native iron in the crust, for example at Disko Island in central west Greenland, and the Urals in Russia. Though iron carbides are rare in nature, the low-pressure phase behavior of carbon in iron has been studied extensively by the steel industry (e.g., Brooks 1996).

Iron carbide occurs occasionally with native iron in the crust. For example, local occurrences of metallic iron with iron carbide ("cohenite") may result from thermal interaction and reduction of basalt with coal or other carbon-rich sediments (Melson and Switzer 1966; Pederson 1979, 1981; Cesnokov et al. 1998). Most famously, Fe-Ni carbide (Ni-poor cohenite) occurs in massive native iron, with schreibersite, sulfides and a variety of minerals in graphite-bearing glassy Tertiary basalts on and around Disko Island, Greenland (individual iron masses > 20 tons; Nordenskiöld 1872; Pauly 1969; Bird and Weathers 1977; Goodrich 1984; Goodrich and Bird 1985). A further 10 ton mass was discovered as recently as 1985, 70 km away from the original Disko iron location (Ulff-Moller 1986). Comparable massive and dispersed native iron containing not only iron carbide but also silicon carbide (H-6, moissanite), occurs together with a rich variety of more than 40 minerals including native metals (Al, Cu), in glass-bearing Permian doleritic sills on the Putorana Plateau, Siberia (Oleynikov et al. 1985). These occurrences are partly brecciated and, although superficially resembling meteoritic textures, they are considered to be terrestrial (Treiman et al. 2002). The origin of the Disko iron is still debated: detailed mapping of dispersed iron in regional basalts strongly favors large-scale interaction of carbon-rich sediments with volcanic lavas (Larsen and Pedersen 2009). However, their correlation with basal stratigraphic units on the Nussussuaq peninsula that preserve unambiguous Ir-bearing impact spherules has reintroduced the prospect for involvement of a meteorite impact in the origin of carbon-rich Disko iron (Jones et al. 2005) as originally invoked by its discoverer (Nordenskiöld 1872).

The structure of cohenite, (orthorhombic; space group \(Pbnm\), \(a = 4.518\, \text{Å}; \ b = 5.069\, \text{Å}; \ c = 6.736\, \text{Å};\) Hendricks 1930), has been extensively studied. Many samples display different cell parameters, potentially related to a variety of causes such as quenching rates. The structure is composed of regular trigonal prisms of iron atoms with carbon at the center (Fig. 6; Animation 4). (Note that in the first structure experiments it was reported that cohenite features a framework of near regular \(\text{CF}_{6}\) octahedra, each with 6 iron atoms surrounding a central carbon atom. However, the positions of the carbon atoms were not determined and the assumption of octahedral coordination was incorrect.)

New carbides from Chinese ultramafic rocks. Two natural occurrences of exotic carbides deserve special note, because they point to a possibly rich and as yet largely unexplored deep
Carbide mineralogy. The first localities are associated with ultramafic rocks in Central China: (1) in podiform chromitites of the Luobusha ophiolite complex, Autonomous Tibetan Region; (2) within the Mengyin kimberlite field, Linyi Prefecture; (3) within the Yangliuping N-Cu-PGE Deposit, Sichuan Province; and (4) at Liu Village, Henan Province. These deposits incorporate minor amounts of three carbides that are unknown from any other region: qusongite (WC; Fang et al. 2009; Shi et al. 2009), yarlongite [(Fe₄Cr₄Ni)C₄; Nicheng et al. 2005, 2008], and tongbaite (Cr₃C₂; Tian et al. 1983; Dai et al. 2004). The association of these carbides with moissanite, cohenite, and khamrabaevite [(Ti,V,Fe)C], as well as other dense, high-temperature phases such as diamond, coesite (a high-pressure form of SiO₂), and varied native metals, including Fe, Ni, W, Cr, Pb, and W, points to a high-temperature, high-pressure origin in a reduced mantle environment (Robinson et al. 2001; Shi et al. 2009).

New carbides from placer deposits of the Urals. A second enigmatic carbide region is found in the Ural Mountains of Russia, within both the Avorinskii Placer, Baranchinsky Massif, and in sediments of the Neiva River near Verkhneivinsk. Placer deposits have yielded <0.3 mm-diameter grains of isovite [(Cr,Fe)₂₃C₆; Generalov et al. 1998], as well as euhedral crystals from the complex nonstoichiometric solid solution between niobocarbide and tantalocarbide [(Nb,Ta)Cₓ₋ₓ; Gusev et al. 1996; Novgorodova et al. 1997]. All known specimens of the latter two minerals, however, were collected early in the 20th century and the exact location of the placer deposit is currently unknown. These two possibly related placer deposits produce an enigmatic suite of other unusual minerals, including native rhenium and osmium (both with melting temperatures > 3000 °C), anyuite [Au(Pb,Sb)₂], and jedwabite [Fe₅(Ta,Nb)₃]. The source lithologies and paragenesis of these minerals are not known, though the concentration of Nb and Ta suggests a possible association with carbonatitic magmas. In any case, they point to the potential diversity of rare carbides in unusual geochemical environments.

Rhombohedral carbonates

By far the most abundant carbon-bearing minerals, both in the number of different species and in their total crustal volume, are the carbonates, of which more than 300 have received IMA approval (http://rruff.info/ima/; Fig. 7). Several previous compilations have reviewed carbonate minerals in detail (Reeder 1983a; Klein and Hurlbut 1993; Chang et al. 1997). Here we summarize key aspects of the mineralogy and crystal chemistry of select carbonates (Tables 5 and 6). Almost all of these minerals incorporate near-planar (CO₃)²⁻ anions, with an equilateral triangle of oxygen atoms around the central carbon atom. Most C-O bond distances are between 1.25 and 1.31 Å and O-C-O angles are close to 120°. Rigid-body libration of these molecular anions contributes to distinctive characteristics of carbonate vibrational spectra, notably three prominent infrared absorption features at ~690-750, 840-900, and 1400-1490 cm⁻¹ (Adler and Kerr 1963; White 1974; Chang et al. 1997) and the strong symmetric stretching modes found near 1100 cm⁻¹ in Raman spectra (Rutt and Nicola 1974).
Figure 7. (caption on facing page)
Two types of rhombohedral carbonates—the calcite and dolomite groups—collectively represent by far the most abundant carbonate minerals in Earth’s crust, with calcite (CaCO\(_3\)) and dolomite [CaMg(CO\(_3\))\(_2\)] in massive sedimentary and metamorphic formations accounting for at least 90% of crustal carbon (Reeder 1983b). Orthorhombic carbonates in the aragonite group also play a significant role in Earth surface processes (Speer 1983), particularly through biomineralization (Stanley and Hardie 1998; Dove et al. 2003; Knoll 2003). These three groups are surveyed below.

**Calcite and the calcite group.** The most important carbonate minerals belonging to the calcite group (Table 5; Reeder 1983b) include calcite (CaCO\(_3\)), magnesite (MgCO\(_3\)), rhodocrosite (MnCO\(_3\)), siderite (FeCO\(_3\)), and smithsonite (ZnCO\(_3\)). It should be emphasized that these minerals seldom occur as pure end-members, but instead commonly form solid solutions with many divalent cations. The calcite structure (space group \(R\bar{3}c\); e.g., Bragg 1914; Effenberger et al. 1981; Chang et al. 1997) has a topology similar to that of NaCl, with each Ca\(^{2+}\) coordinated to 6 (CO\(_3\))\(^{2−}\) groups, and each (CO\(_3\))\(^{2−}\) group in turn coordinated to 6 Ca\(^{2+}\) cations. However, the orientations of the (CO\(_3\))\(^{2−}\) groups, while the same within each layer, are 180° out of phase in successive layers, thus doubling the repeat distance in the \(c\) axial direction relative to a sodium chloride analog. Note that this layer-by-layer alternation of (CO\(_3\))\(^{2−}\) group orientations results in an oxygen atom distribution that approximates hexagonal close packing, with C and Ca occupying 3- and 6-coordinated interstices, respectively (Megaw 1973). The flattened shape of this carbonate anion results in an obtuse rhombohedral angle of 101°55’ (Fig. 8; Animation 5).

Figure 7. The diversity of carbonate minerals. (a) Rhodochrosite (MnCO\(_3\)), rhombohedral crystals from the Home Sweet Home Mine, Mount Bross, Alma District, Park County, Colorado, USA. (b) Ankerite [CaFe\(^{2+}\)(CO\(_3\))\(_2\)] from Brownley Hill mine, Nenthead, Cumbria, England showing the typical curved saddle-shaped rhombohedral crystals. (c) Artinite [Mg\(_5\)CO\(_3\)(OH)\(_2\)·3H\(_2\)O], divergent sprays of clear colorless acicular crystals from San Benito County, California, USA. (d) Aurichalcite [Zn\(_5\)(CO\(_3\))\(_2\)(OH)\(_6\)·12H\(_2\)O], divergent sprays of light blue lathlike crystals from Bisbee, Cochise County, Arizona, USA. (e) Hydromagnesite [Mg\(_5\)(CO\(_3\))\(_2\)(OH)\(_2\)·4H\(_2\)O], colorless bladed crystals from Paradise Range, Nye County, Nevada, USA. (f) Jouravskite [Ca\(_3\)Mn\(^{4+}\)(SO\(_4\))(CO\(_3\))(OH)\(_6\)·12H\(_2\)O], yellow hexagonal prism associated with calcite from the Wessels mine, Kurumen, Kalahari Manganese fields, Cape Province, South Africa. (g) Stichtite [Mg\(_6\)Cr\(_2\)CO\(_3\)(OH)\(_{16}\)·4H\(_2\)O], an aggregate of contorted purple plates from Dundas, Tasmania, Australia. (h) Zaratite [Ni\(_2\)CO\(_3\)(OH)\(_4\)·4H\(_2\)O], green amorphous crust, intimately associated with népouite from Lord Brassy mine, Tasmania, Australia. All photos courtesy of the RRUFF project and irocks.com.

Figure 8. The structure of calcite (CaCO\(_3\)) (rhombohedral; space group \(R\bar{3}c\); hexagonal setting \(a = 4.989\) Å; \(c = 17.061\) Å; \(Z = 6\); rhombohedral setting \(a = 6.375\); \(\alpha = 46.1°\); \(Z = 2\)). Calcite crystal from Elmwood Mine, Carthage, Tennessee. Photo courtesy of Rob Lavinsky. [Animation 5: For readers of the electronic version, click the image for an animation of the calcite crystal structure.]
Table 5. Selected anhydrous carbonates, primarily species with more than 10 known localities (data compiled from http://MinDat.Org databases).

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<td>KNa₂₂(SO₄)₃(CO₃)Cl</td>
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<td>Bastnäsite Group</td>
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<td>Synchysite Group</td>
<td>Ca(REE)(CO₃)F</td>
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<td>Huangboite-(Cc)</td>
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<td>Schröckingerite</td>
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<td>Trona</td>
<td>Na₃(HCO₃)(CO₃)·2H₂O</td>
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<td>Lanthanite Group</td>
<td>(REE)₂(CO₃)₃·8H₂O</td>
<td>Tyrolite</td>
<td>Ca₆Cu₆(AsO₄)₃(CO₃)(OH)₁₁·11H₂O</td>
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Calcite is widely distributed in Earth’s crust; it appears most commonly within sedimentary rocks, where it occurs as the principal mineral of limestone, and as a natural cementing agent in many siliceous sandstone and shale units that were deposited under marine conditions. Calcite also dominates some metamorphic rocks such as marble and calcareous gneiss; occurs widely in hydrothermal systems, where it forms extensive vein networks; and is common in some unusual carbonate-rich igneous rocks such as carbonatites (Jones et al. 2013).

Although widely formed under Earth’s near-surface conditions, distributions of calcite and other rhombohedral carbonate minerals have varied significantly through Earth history, principally as a consequence of feedbacks between the geosphere and biosphere (Knoll 2003; Hazen et al. 2008; Hazen et al. 2013). At present, the majority of calcium carbonate deposition occurs as calcite precipitated within shallow marine settings. In these settings, magnesium—whose concentration is nearly 4× that of calcium in normal marine fluids—is readily incorporated into the calcite crystal lattice, with Mg concentrations of marine calcites equal to a few to nearly 20 mol% MgCO$_3$ (MacKenzie et al. 1983; Morse and MacKenzie 1990; Morse et al. 2006; Berner and Berner 2012). Such Mg-bearing calcites are commonly referred to as magnesian calcite or “high Mg calcite” (HMC) and are distinguished from calcites with low Mg concentrations (“low Mg calcite” or LMC). Magnesium is one of a suite of divalent ions that readily co-precipitate with calcium in the calcite lattice. Because ionic co-precipitation reflects a combination of the ionic availability, temperature, and lattice structure, and because it can substantially affect the solubility and rate of dissolution of the resultant calcite, differential co-precipitation of ions within calcite has been, and continues to be, a subject of intense investigation as a means of unraveling the geologic history of the oceanic system (Morse and Mackenzie 1990).

One of calcite’s remarkable and as yet largely unexplored features is its extraordinary range of crystal forms (e.g., Dana 1958). Habits range from the more common rhombohedral and scalenohedral crystal forms, to needle-like, platy, and equant shapes with expression of at least 300 different documented crystal forms (Fig. 9; see also specimen photographs on http://MinDat.org). Some of the most varied crystal habits are widely distributed in association with both biological skeletalization (Fallini et al. 1996; Dove et al. 2003) and speleogenesis (Frisia et al. 2000) and reflect a complex array of physical, chemical, and biological influences during crystallization. Because calcite crystal morphology is strongly affected by the kinetics of crystal growth, unraveling the differential effects of fluid saturation state, carbonate ion availability, ionic activity, the presence or absence of ionic inhibitors to nucleation and growth, and even the presence or absence of mineral catalyzing organic molecules (e.g., Cody and Cody 1991; Teng and Dove 1997; Teng et al. 1998; Orme et al. 2001), is critical to reveal as of yet untapped insights to Earth’s crustal evolution. This need to document connections between environment and crystal form may be true, in particular, for our understanding of distinct carbonate morphologies such as “herringbone” calcite (Sumner and Grotzinger 1996; Kah et al. 1999) and “molar-tooth” calcite (Pollock et al. 2006) that show distinct environmental distributions through Earth history (see Hazen et al. 2013).

**Other calcite group minerals.** The magnesium carbonate magnesite (MgCO$_3$) forms primarily through alteration of Mg-rich igneous and metamorphic rocks, commonly in association with serpentinite, as well as by direct precipitation from Mg-rich solutions and as a primary phase in mantle-derived carbonatites. Anhydrous magnesium carbonate commonly hydrates to form one of several secondary minerals (Table 6), including hydromagnesite [Mg$_4$(CO$_3$)$_3$(OH)$_2$·3H$_2$O], artinite [Mg$_2$CO$_3$(OH)$_2$·3H$_2$O], dyepingite [Mg$_5$(CO$_3$)$_4$(OH)$_2$·5H$_2$O], pokrovite [Mg$_2$CO$_3$(OH)$_2$], nesquehonite (MgCO$_3$·3H$_2$O), and landsfordite (MgCO$_3$·5H$_2$O).

Rhodochrosite, the manganese carbonate (MnCO$_3$), most commonly occurs as a vein-filling phase in hydrothermal ore districts. Limited solid solutions with Ca, Fe, and Mg end-members, as well as Zn, Ba, and Pb, are typical, as is partial alteration to manganese oxide.
Figure 9. Crystal forms of natural calcite. (a) Rhombohedral cleavage fragment of optical grade material from near Presidio, Texas, USA, University of Arizona Mineral Museum 16674 (b) Scalenohedrons collected in the early 1800’s from the Bigrigg mine, Cumbria, England. Bob Downs specimen; (c) Hexagonal prisms with rhombohedral terminations from Joplin, Missouri, USA UAMM 16545; (d) Herringbone growths of acicular crystals from the Southwest mine, Bisbee, Arizona, USA, UAMM 9499; (e) Hexagonal prisms with pyrite centers from Charcas, San Luis Potosi, Mexico, UAMM 1214; (f) Stalactite globules from Southwest mine, Bisbee, Arizona, USA, UAMM 9499; (g) Aggregate of bladed crystals from the Onyx cave, Santa Rita Mts, Arizona, USA, UAMM 5503; (h) Hexagonal prisms with pyramidal terminations from the Camp Bird mine, Imogene Basin, Ouray County, Colorado, USA, UAMM 6703. All photographs by Alesha Siegal, University of Arizona.
hydroxides. Rhodochrosite is typically pale pink in color, though relatively rare deep rose pink specimens occur occasionally and are highly prized as semi-precious gemstones.

The iron carbonate siderite (FeCO$_3$) occurs in massive beds as an important component of some Precambrian banded iron sedimentary formations (Klein 2005), as well as in hydrothermal veins associated with ferrous metal sulfides. Siderite commonly incorporates Ca, Mn, and Co, and it forms a complete solid solution with magnesite in a variety of lithological settings, as well as with smithsonite (ZnCO$_3$) in hydrothermal lead-zinc ore deposits. Siderite is only stable under conditions of relatively low $f_{O_2}$ (Hazen et al. 2013). It is metastable under ambient oxic conditions and typically decomposes to a suite of iron oxide-hydroxides such as goethite [FeO(OH)], and related hydrous phases [FeO(OH)-$n$H$_2$O]—reactions that are accelerated by chemolithoautotrophic microbial activity.

**Dolomite group.** The dolomite group (space group $R3$) is topologically identical to calcite, but in these double carbonate minerals two or more different cations occupy alternate layers perpendicular to the $c$ axis (Wasastjerna 1924; Wyckoff and Merwin 1924; Reeder 1983b; Chang et al. 1997; Fig. 10; Animation 6). Important end-member minerals in this group (Table 5) include dolomite [CaMg(CO$_3$)$_2$], ankerite [FeMg(CO$_3$)$_2$], kutnohorite [CaMn(CO$_3$)$_2$], and minnerecrdite [CaZn(CO$_3$)$_2$]. Note that in both the calcite and dolomite mineral groups the (CO$_3$)$_2^-$ anions lie perpendicular to the rhombohedral $c$ axis and they librate with a helical motion along this axis (Gunasekaran et al. 2006; Fig. 11; Animation 7).

The planar orientation of the CO$_3^{2-}$ anions in the calcite and dolomite group minerals results in many of their distinctive properties, for example in their extreme optical anisotropy (maximum and minimum refractive indices differ by ~0.2 in these minerals), which causes the familiar double refraction seen through calcite cleavage rhombohedra. The near-perfect [104] cleavage of calcite and dolomite group minerals also arises from anisotropies in bonding; this cleavage plane results in the minimum number of broken Ca-O bonds and no broken C-O bonds. Strong bonding in the plane parallel to the CO$_3^{2-}$ anions (i.e., the $a$ axis of the hexagonal setting) compared to weaker Ca-O bonds in the perpendicular direction (the $c$ axis of the hexagonal setting) leads to extreme anisotropy in calcite’s thermal expansion, as well. The $c$-axis thermal expansion is positive ~ $+3.2 \times 10^{-5} \degree C^{-1}$, whereas $a$-axis thermal expansion

**Figure 10.** Dolomite. (a) The structure of dolomite [MgCa(CO$_3$)$_2$] (rhombohedral; space group $R3$; hexagonal setting $a = 4.807 \text{ Å}; c = 16.00 \text{ Å}; Z = 3$; rhombohedral setting $a = 6.015, \alpha = 47.1^\circ; Z = 1$). Blue spheres are carbon atoms, with light green CaO$_6$ octahedra and dark green MgO$_6$ octahedra. (b) Dolomite crystals reflect their rhombohedral crystal structure. [Animation 6: For readers of the electronic version, click the image for an animation of the dolomite crystal structure.]
is negative $\sim -0.3 \times 10^{-5} \circ C^{-1}$ (Markgraf and Reeder 1985). Note, however, that in magnesite and dolomite the shorter, stronger Mg-O bonds result in thermal expansion that is positive in both $a$- and $c$-axial directions.

Dolomite is by far the most abundant species among the dolomite group minerals. It forms primarily within both sedimentary and metamorphic deposits through the diagenetic replacement of calcite during interaction with Mg-rich fluids. Phase relations in the $CaCO_3$-$MgCO_3$-$FeCO_3$ system, including the phases calcite, magnesite, siderite, and dolomite, as well as magnesian calcite, as summarized by Chang et al. (1997), reveal extensive regions in pressure-temperature-composition space of coexisting calcite group and dolomite group minerals—a topology that is borne out by the common association of calcite and dolomite in sedimentary rocks.

Ankerite and kutnahorite are $Fe^{2+}$- and $Mn^{2+}$-bearing dolomites, respectively, with near-continuous solid solutions observed among the Mg, Fe, and Mn end-members (Essene 1983). These phases occur most commonly as a result of hydrothermal alteration of calcite by reduced fluids rich in $Fe^{2+}$ and $Mn^{2+}$.

**The aragonite group**

A large number of $CaCO_3$ polymorphs enrich carbonate mineralogy (Carlson 1983; Chang et al. 1997). The aragonite group, which includes the aragonite form of $CaCO_3$ plus cerussite ($PbCO_3$), strontianite ($SrCO_3$), and wetherite ($BaCO_3$), prevail in carbonates that contain cations with ionic radii as large or larger than calcium. The crystal structure of aragonite (Speer 1983; Fig. 12; Animation 8), first determined by Bragg (1924), is orthorhombic with the standard space group $Pnam$ (with $c < a < b$). However, the structure is more conveniently described in a non-standard orientation with $a < c < b$, resulting in space group $Pmcn$. The structure in this orientation possesses alternating (001) layers of divalent metal cations and $(CO_3)^{2-}$ anions. Two types of $(CO_3)^{2-}$ layers ($C_1$ and $C_2$) alternate with two orientations of metal cations (A and B) in a stacking sequence $[\ldots AC_1 BC_2 \ldots]$ (Speer 1983). Each $(CO_3)^{2-}$ anion is coordinated to 6 divalent metal atoms, and each metal atom is coordinated to 9 oxygen atoms (Fig. 12a).

Aragonite is a denser polymorph of $CaCO_3$ than calcite (2.95 vs. 2.71 g/cm$^3$), and has been recognized as a mineral characteristic of relatively high-pressure, low-temperature metamorphic environments (McKee 1962; Coleman and Lee 1962; Ernst 1965). Aragonite, however, is also one of the predominant carbonate minerals—along with calcite and dolomite—that comprise the vast amount of carbon mineralization in Earth’s surface environments. Although aragonite is the least stable of these minerals, and undergoes rapid recrystallization to thermodynamically more stable forms, it has been recognized as a fundamental constituent of marine depositional environments since at least the Archean (Sumner and Grotzinger 2004), and represents a primary shallow-marine depositional facies throughout the Proterozoic (Grotzinger and Read 1983; Bartley and Kah 2004; Kah and Bartley 2011). Furthermore, since the onset of enzymatic
biomineralization, more than 500 million years ago, aragonite has been a primary constituent of the fossil record, forming metastable skeletons of some calcareous algae (Bathurst 1976) as well as the skeletal components of a variety of invertebrates, including a wide variety of molluscs, scleractinian corals, and some bryozoans (Cloud 1962; Rucker and Carver 1969; Knoll 2003; Dove 2010). Three other aragonite group minerals, the Sr, Ba, and Pb carbonates strontianite, witherite, and cerussite, respectively, are all found primarily in relatively low-temperature hydrothermal or supergene environments, commonly associated with sulfates and metal sulfide ores (Smith 1926; Mitchell and Pharr 1961; Mamedov 1963; Speer 1977; Dunham and Wilson 1985; Wang and Li 1991). Strontianite, in particular, has also been found in association with enzymatic biomineralization, with strontianite comprising nearly 40% of the skeletal carbonate in some scleractinian corals (McGreegor et al. 1997), although it is uncertain whether strontianite in this case is primary or an early diagenetic phase resulting from recrystallization along the metastable strontianite-aragonite solid solution (Plummer et al. 1992).

Phase diagrams for the aragonite polymorphs of CaCO$_3$ and other calcite group minerals underscore the effects of divalent cation coordination number on carbonate structure type. Several calcite group minerals transform to the aragonite structure at high pressure (Carlson 1983; Yoshioka and Kitano 2011); conversely, the aragonite group minerals strontianite and witherite transform to the calcite structure at high temperature (Chang 1965).

**Other anhydrous carbonates**

More than 90 anhydrous carbonates other than the above mentioned rhombohedral and orthorhombic species have been described, though only a handful of these diverse phases are
common (Table 5; http://rruff.info/ima/). Of these minerals, vaterite, huntite, and several minerals associated with carbonatites deserve special note.

**Vaterite.** Vaterite is a polymorph of CaCO$_3$ that is unstable under ambient conditions, yet plays an important role in biomineralization. Nanocrystalline vaterite rapidly converts to calcite, but can be stabilized indefinitely by a variety of hydrophilic organic molecules. Vaterite’s critical role in biomineralization arises from the low energy required for its conversion to other crystalline forms (Xu et al. 2006; Soldati et al. 2008; Wehrmeister et al. 2012). Therefore, it is the preferred mineral phase (along with amorphous calcium carbonate, or ACC) for storing material critical to skeletal growth. Typically, metazoans process nan-aggregates of vaterite (or ACC; Addadi et al. 2003) to form mesoscale syntaxial structures, which may provide an energetically favorable pathway to the construction of larger skeletal elements (e.g., single-crystal echinoderm plates).

Details of the vaterite crystal structure, which must be analyzed using nano-scale powders by X-ray or electron diffraction methods, remains in doubt. It was once thought to be hexagonal (space group $P6_3/mmc$; $a = 7.135$ Å; $c = 16.98$ Å; Kahmi 1963). However, Le Bail et al. (2011) found evidence for an orthorhombic structure with 3-fold cyclic twinning (space group $Ama2$; $a = 8.472$ Å; $b = 7.158$ Å; $c = 4.127$ Å; $Z = 4$). By contrast, Mugnaioli et al. (2012) propose a monoclinic unit cell ($a = 12.17$ Å, $b = 7.12$ Å, $c = 9.47$ Å, $\beta = 118.94^\circ$), which is a geometric transformation of the smallest hexagonal cell proposed by Kamhi (1963). All studies agree that vaterite features Ca in distorted 7 or 8 coordination, with an octahedron of 6 Ca-O bonds at ~2.4 Å, and two longer Ca-O bonds at >2.9 Å (Fig. 13; Animation 9). Thus, calcium coordination in vaterite is intermediate between that of calcite (6) and aragonite (9). Wang and Becker (2009) employed first-principles calculations and molecular dynamics simulations to elucidate details of the vaterite structure and CO$_3^{2-}$ group orientations. Previous studies had suggested rotational disorder among carbonate groups, but Wang and Becker (2009) demonstrate a more stable configuration with an ordered CO$_3^{2-}$ superstructure. Vaterite is commonly rotationally disordered when first crystallized, but it can achieve carbonate orientational order through annealing.

**Huntite.** Huntite [CaMg$_3$(CO$_3$)$_4$] occurs as a low-temperature mineral, both by direct precipitation from aqueous solutions enriched in Mg and as an alteration product of dolomite or magnesite (Kinsman 1967). Its hexagonal structure (space group $R32$; $a = 9.503$ Å; $c = 7.821$ Å) bears some similarities to rhombohedral carbonates. Magnesium is in octahedral

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Figure 13. The crystal structure of vaterite (CaCO$_3$). (hexagonal, space group $P6_3/mmc$; $a = 7.135$ Å; $c = 16.98$ Å; $Z = 16$). Blue spheres and green polyhedral represent carbon and calcium-oxygen polyhedral, respectively. [Animation 9: For readers of the electronic version, click the image for an animation of the vaterite crystal structure.]
coordination in all of these carbonates, but calcium in huntite is in trigonal prismatic coordination, as opposed to octahedral coordination in calcite, dolomite, and other species (Dollase and Reeder 1986; Fig. 14 [Animation 10]).

**Carbonatite carbonate mineralogy.** Finally, a number of exotic carbonate minerals are found associated with carbonatites, which are defined as igneous rocks (extrusive or intrusive) with greater than 50% carbonate minerals (Tuttle and Gittins 1966; Bell 1989; Jones et al. 2013). These rare mantle-derived magmas are stable over a wide temperature range and can erupt as remarkably cool surface lavas at ~500 to 600 °C (Dawson et al. 1990; Church and Jones 1995). Carbonatites encompass a range of compositions, including those dominated by calcite (Ca), dolomite (Ca-Mg), ankerite (Ca-Fe), and alkali (Na-K+Ca) carbonate minerals. Furthermore, these magmas are often enriched in an unusual suite of elements in addition to carbon: alkalis, alkaline earths, fluorine, phosphorus, rare earth elements, and niobium (Deans 1966). As a result of these chemical complexities, approximately 30 different carbonate minerals have been identified from these varied sources (Kapustin 1980; Chang et al. 1997).

Several of the rhombohedral carbonates are found commonly in carbonatites, including calcite, magnesite, siderite, rhodochrosite, and dolomite, as well as orthorhombic aragonite and strontianite (Garson and Morgan 1978; Kapustin 1980; Dziedzic and Ryka 1983). These familiar minerals are accompanied by a number of exotic double carbonates include alstonite and barytocalcite [both polymorphs of CaBa(CO$_3$)$_2$], norsethite [BaMg(CO$_3$)$_2$], and the rare earth carbonates burbankite [(Na,Ca)$_3$(Sr,Ba,Ce)$_3$(CO$_3$)$_5$], sahamalite-(Ce) [Ce$_2$Mg(CO$_3$)$_4$] and carbocernaite [(Ca,Na)(Sr,Ce,La)(CO$_3$)$_2$]. Note that reference is sometimes made to “breunnerite” [(Mg,Fe)CO$_3$], but this Fe-rich magnesite is not a valid mineral species.

The alkali carbonatites are exemplified by Oldoinyo Lengai in Tanzania (the only currently active carbonatite volcano), and feature several minerals with the general formula [(Na,K)$_2$Ca(CO$_3$)$_2$] (Dawson 1962, 1966; Chang et al. 1997). The two sodium-rich end-members are nyerereite (typically with Na/K ~4.5, though the official IMA chemical formula lacks K) and a possible higher-temperature polymorph zemkorite (with Na/K ~6.2; again, the official IMA-approved formula lacks K); potassium end-member species are fairchildite and a lower-temperature form, bütschliite (Dawson 1962; Mrose et al. 1966; McKie and Frankis 1977; Yergorov et al. 1988; Chang et al. 1997). These phases are also accompanied by the rare sodium carbonate gregoryite [Na$_2$(CO$_3$)], which can only form in alkali-rich, alkaline earth-poor systems.

![Figure 14](image.png)

**Figure 14.** The structure of huntite [CaMg$_3$(CO$_3$)$_4$]. (hexagonal; space group R32; $a = 9.503$ Å, $c = 7.821$ Å; $Z = 3$). The huntite structure is unusual in the trigonal prism coordination of calcium, coupled with octahedral coordination of magnesium. [Animation 10: For readers of the electronic version, click the image for an animation of the huntite crystal structure.]
Carbonates can also develop suites of rare earth element (REE) carbonate or carbonate-fluoride minerals, including the bastnäsite group [REE(CO$_3$)$_2$F, found with dominant REE = Ce, La, Nd, or Y], synchysite and parasite [both CaREE(CO$_3$)$_2$F, found with dominant REE = Ce, Nd, or Y], huangoite-(Ce) [Ba(Ce,REE)(CO$_3$)$_2$F], and the cordylite group [NaBaREE$_2$(CO$_3$)$_4$F, found with dominant REE = Ce or La]. REE-carbonates are often geologically associated with U and Th minerals (Ruberti et al. 2008; also see section on “Uranyl carbonates” below). Two of the world’s largest economic ore bodies for REE are carbonatites: Mountain Pass California (age 1.37 Ga; Olson et al. 1954; Jones and Willie 1986; Castor 2008) and Bayan Obo, China (age 1.35 Ga: Le Bas et al. 1992; Yang et al. 2011). Finally, several hydrous carbonates, including the uncommon REE phases of the ancyelite group [(Cu, Sr, Pb)(REE)(CO$_3$)$_2$(OH)-H$_2$O, found with dominant REE = Ce, La, or Nd], calcinite-(Ce) [(Ce,REE)$_2$(CO$_3$)$_2$-4H$_2$O], and the lanthanite group [REE$_2$(CO$_3$)$_2$-8H$_2$O, found with dominant REE = Ce, La, or Nd], along with two closely associated hydrated Mg-Al carbonates, manasseite [Mg$_6$Al$_2$(CO$_3$)(OH)$_{16}$-4H$_2$O] and hydrotalcite [Mg$_6$Al$_2$(CO$_3$)(OH)$_{4}$·4H$_2$O], have been reported from some carbonatites (Kapustin 1980).

**Hydrous carbonates**

Most carbon-bearing minerals—more than 210 of the approximately 320 IMA approved carbonate species—are hydrated or hydrous (Table 6; http://rruff.info/ima/). These diverse species, whose classification has been systematized by Mills et al. (2009), occupy numerous specialized near-surface niches but are for the most part volumetrically minor. Most of these diverse minerals have mixed anionic groups, including more than 30 carbonate-sulfates, more than 40 carbonate-silicates, and more than 20 carbonate-phosphates, plus carbonates with uranyl, arsenate, borate, and other ionic groups, which represent near-surface alteration products of other minerals (see Tables 5 and 6 for some of the more common representative examples).

**Malachite and azurite.** The common hydrous copper carbonates, malachite [Cu$_2$(OH)$_2$CO$_3$] and azurite [Cu$_3$(OH)$_2$(CO$_3$)$_2$], which form in the oxidized zone of copper deposits, are typical of a large number of near-surface carbonate phases, including more than 20 other hydrous copper carbonates. These colorful minerals are among the thousands of mineral species that Hazen et al. (2008) identified as potentially biologically mediated (Hazen et al. 2013).

**Uranyl carbonates.** At least 30 carbonates incorporate the (UO$_2$)$_{2+}$ uranyl cation group. These fascinating and colorful phases, which commonly form as alteration products of near-surface uranium ore bodies, include 4 anhydrous carbonates: rutherfordine [(UO$_2$)(CO$_3$)$_3$, 6H$_2$O, bayleyite [Mg$_2$(UO$_2$)(CO$_3$)$_2$(H$_2$O)$_{12}$-6H$_2$O], liebigite [Ca$_2$(UO$_2$)(CO$_3$)$_3$-11H$_2$O], and schrockingerite [NaCa$_3$(UO$_2$)(SO$_4$)(CO$_3$)$_2$F-10H$_2$O]. These minerals are all Ca or Mg carbonates that presumably formed by alteration of calcite or dolomite in a relatively oxidized subsurface environment—conditions that must significantly postdate the Great Oxidation Event (Hazen et al. 2008, 2009; Sverjensky and Lee 2010; Hazen et al. 2013).

**Hydrated calcium carbonates.** In addition to the mineralogically complex hydrated carbonates outlined above, hydrated phases of calcium carbonate—i.e., ikaite [CaCO$_3$·6H$_2$O] and its pseudomorphs [which have been given a number of unofficial varietal names such as thinolite, glendonite, jarrowite, and fundylite (Browell 1860; Dana 1884; Shearman and Smith 1985; Ito 1996; Swainson and Hammond 2001)], plus monohydrocalcite (CaCO$_3$·H$_2$O) and hydromagnesite [Mg$_2$(CO$_3$)$_4$(OH)$_2$-4H$_2$O]—play critical roles in the near-surface co-evolution of the geosphere and biosphere. Hydrated calcite is common in modern microbial systems...
and commonly represents an initial precipitate phase within microbially mediated carbonate deposits. Similarly, pseudomorphs after ikaite provide thermodynamic indicators of cold marine temperatures, and can potentially be used as a unique indicator of glaciated conditions in the geologic past, for example, as pseudomorphs after ikaite in the Mississippian of Alberta, Canada (Brandley and Krause 1997) and in glendonites in Neoproterozoic shallow shelf environments (James et al. 1998).

Minerals incorporating organic molecules

The IMA has recognized approximately 50 organic minerals, and it is likely that many more species remain to be identified and described (Table 7; http://rruff.info/ima). These diverse phases fall into three main categories—organic molecular crystals, minerals with organic molecular anions, and clathrates. Note that while many of these minerals are a direct or indirect consequence of biological activity, some minerals with organic molecules may have a non-biological origin. Consequently, Perry et al. (2007) have also introduced the term “organomineral” to designate “mineral products containing organic carbon,” but “not directly produced by living cells.” As examples they cite carbon-bearing siliceous hot-spring deposits, desert varnish,stromatolites, and a variety of trace fossils.

Organic molecular crystals. Organic molecular crystals encompass those carbon-bearing minerals in which electrically neutral organic molecules crystalize into a periodic arrangement, principally through van der Waals interactions (Table 7). This diverse group includes a number of species known only as accessory minerals associated with coal; for example, 9 different hydrocarbon minerals, such as kratochvilite (C_{12}H_{10}), fichtelite (C_{10}H_{34}), dinite (C_{20}H_{36}), and evenkite (C_{26}H_{48}); acetamide (CH_{3}CONH_{2}); and the ring-shaped nickel porphyrin mineral abelsonite, Ni(C_{5}H_{6}N_{3}O). Burning coal mines also produce molecular crystals through sublimation, including kladnoite [C_{6}H_{4}(COO)_{2}NH] and hoelite (C_{14}H_{16}O_{2}; Jehlička et al. 2007). Other molecular crystals are unique to fossilized wood [flagstaffite (C_{10}H_{22}O_{3})]; roots [refikite (C_{20}H_{12}O_{2})]; or bat guano deposits in caves, for example the purines uricite (C_{5}H_{12}N_{4}O_{3}) and guanine (C_{5}H_{5}N_{5}O), and urea [CO(NH)_{2}]. It is intriguing to note that average temperatures on Earth are too high for many crystals of small organic molecules to form, including crystalline forms of carbon dioxide (which has been observed near the poles of Mars; Byrne and Ingersoll 2003), methane, ethane (C_{2}H_{6}), ethylene (C_{2}H_{2}), and propane (C_{3}H_{8}). One can anticipate that these and many other such minerals await discovery in the cold, hydrocarbon-rich near-surface environment of Saturn’s moon Titan, as well as in the form of condensed phases in dense molecular clouds (Glein 2012).

Minerals with organic anions. Approximately 25 mineral species incorporate organic anions bonded to Ca, Mg, Cu, Na, and other metal cations (Table 7). Predominant among these organic salt minerals are more than a dozen oxalates with the (C_{2}O_{4})^{2−} anion. The most common oxalate is weddellite [Ca(C_{2}O_{4})·2H_{2}O], which is found in such varied environments as bat guano, sediments derived from lichens, human kidney stones, cactus (saguaro, Carnegiea gigantea), and the depths of the Weddell Sea. Other oxalates include caoxite [Ca(C_{2}O_{4})·3H_{2}O], glushinskite [Mg(C_{2}O_{4})·2H_{2}O], humboldtine [Fe(C_{2}O_{4})·2H_{2}O], lindbergite [Mn(C_{2}O_{4})·2H_{2}O], natroxalate [Na_{2}(C_{2}O_{4})], and oxammite [(NH_{4})_{2}(C_{2}O_{4})·2H_{2}O] (the latter a biomineral known exclusively from guano deposits).

Other organic anions in minerals include formate (HCOO)^−, for example in formicate [Ca(CHOO)_{2}] and dashkovite [Mg(HCOO)_{2}·2H_{2}O]; acetyl (CH_{3}COO)^− in hoganite [Cu(CH_{3}COO)_{2}·2H_{2}O] and calclacite [Ca(CH_{3}COO)Cl·5H_{2}O] (the latter mineral known only from specimens of limestone stored in wooden drawers); methyl sulfonate (CH_{3}SO_{3})^{−} in ensteinburkite [Mg(CH_{3}SO_{3})_{2}·12H_{2}O]; and thiocyanate (SCN)^− in julienite [Na_{2}Co(SCN)_{4}·8H_{2}O]. Finally, Rastsvetaeva et al. (1996) describe a Cu^{2+} succinate monohydrate phase that occurs as a consequence of washing copper mineral specimens with detergents.
**Clathrate silicates.** Clathrates comprise a third as yet poorly described group of minerals containing organic molecules. These minerals feature open three-dimensional framework structures that incorporate molecules in cage-like cavities. Chibaite and melanophlogite are silica clathrates with open zeolite-like SiO$_2$ frameworks that may contain hydrocarbons from methane to butane (C$_{n}$H$_{2n+2}$) as the guest molecule (Skinner and Appleman 1963; Momma et al. 2011). Melanophlogite with CO$_2$ and/or methane as the guest molecule has been found in natrocarbonatite lavas at Oldoinyo Lengai (Carmody 2012).

**Clathrate hydrates.** Water-based clathrates, also known as gas hydrates or clathrate hydrates, are remarkable crystalline compounds that form at low temperatures (typically < 0 °C) and elevated pressures (> 6 MPa). These materials have attracted considerable attention because of their potential for applications to energy storage and recovery applications (Max
Different gas clathrate hydrate structure types have a variety of cage sizes and shapes, which depend primarily on the size and character of the gas molecule.

The principal documented natural clathrate hydrate mineral is an as yet unnamed methane hydrate commonly known as “methane ice, which crystallizes in marine sediments of the continental shelves or in permafrost zones below a depth of ~130 (Hyndman and Davis 1992). Methane hydrate is structurally similar to the silica clathrates. Its cubic structure (space group \textit{Pm} \textit{3m}, \textit{a} = 12 \text{ Å}) features a three-dimensional H$_2$O framework with two types of cages partially filled with CH$_4$ molecules: a pentagonal dodecahedron (designated 5$^{12}$, or a cage formed by 12 interconnected 5-member rings of H$_2$O) and a tetrakaidecahedron (designated 5$^{12}$6$^2$), each with average radii ~4 Å, each holding one CH$_4$ molecule (Fig. 15; Animation 11). Although the methane content is variable, methane ice holds on average ~0.17 mole of methane per mole of water, corresponding to a density of ~0.9 g/cm$^3$ (Max 2003).

The geographic distribution of methane hydrate is extensive, with hundreds of confirmed deposits (Hyndman and Spence 1992; Kvenvolden 1995; Milkov 2004). The total methane storage in clathrates was estimated by Allison and Boswell (2007) as $2 \times 10^{16}$ m$^3$—a quantity orders of magnitude greater than that represented by all other natural gas reserves. Annual natural gas consumption in North America, by comparison, is $\sim 6 \times 10^{11}$ m$^3$ (Boswell 2009). In fact, the methane stored in clathrate hydrates may exceed the energy represented by known reserves of all other fossil fuels combined (Kvenvolden 1995; Grace et al. 2008).

In addition to this common phase, Guggenheim and Koster van Groos (2003) and Koster van Groos and Guggenheim (2009) have reported the synthesis of a possible new gas-hydrate phase that consists of a clay-methane hydrate intercalate. In addition, Chou et al. (2000) have reported other methane clathrate hydrate phases that occur exclusively at high pressures.

Mineral-molecule interactions

Finally, it should be noted that organic molecules often interact strongly with mineral surfaces, especially in aqueous environments (Hazen 2006; Jonsson et al. 2009; Bahri et al. 2011; Cleaves et al. 2011). Such interactions of organic species with mineral surfaces have received special attention for at least two reasons related to the biosphere. First, a number of

![Figure 15](image_url). Methane hydrate. (a) The structure of methane hydrate \([\text{H}_2\text{O}\cdot\text{CH}_4]\) (cubic; space group \textit{Pm} \textit{3m}; \textit{a} = 12 \text{ Å}). (b) Methane hydrate burns at room conditions, as the hydrate melts and methane is released. Photo courtesy of Jan Steffan, Expedition SO148, GEOMAR. [Animation 11: For readers of the electronic version, click the image for an animation of the methane hydrate crystal structure.]
minerals have been invoked as possibly playing key roles in the origins of life through the selection, concentration, protection, and templating of biomolecules, as well as the possible catalysis of biomolecules (Bernal 1951; Goldschmidt 1952; Orgel 1998; Lahav 1999; Schoonen et al. 2004; Hazen 2005, 2006). Specific hypotheses focus on the roles of hydroxides (Holm et al. 1993; Pitsch et al. 1995; Hill et al. 1998), quartz (Bonner et al. 1974, 1975; Evgenii and Wolfram 2000), feldspars and zeolites (Smith 1998; Parsons et al. 1998; Smith et al. 1999), carbonates (Hazen et al. 2001), phosphates (Weber 1982, 1995; Acevedo and Orgel 1986), borates (Ricardo et al. 2004; Grew et al. 2011), phosphides (Pasek et al. 2007), and sulfides (Wächtershäuser 1988, 1990, 1993; Russell et al. 1994; Russell and Hall 1997; Huber and Wächtershäuser 1998; Bebié and Schoonen 2000; Cody et al. 2000, 2001, 2004; Cody 2004). In this regard, clay minerals have received special attention for their potential ability to template and catalyze the polymerization of amino acids and nucleotides as steps in the origins of life (Cairns-Smith 1982, 2005; Cairns-Smith and Hartman 1986; Ferris et al. 1996; Ertem and Ferris 1996; Orgel 1998; Ferris 2005).

A second possible influence of mineral-molecule interactions on the biosphere also invokes clay minerals, which might have contributed significantly to the rise of atmospheric oxygen through the adsorption, concentration, and subsequent burial of significant amounts of organic matter from the terrestrial environment (Kennedy et al. 2006). This process of organic burial is one of the most efficient mechanisms for atmospheric oxidation (Berner et al 2000; Hayes and Waldbauer 2006; Hazen et al. 2013).

CONCLUSIONS: UNRESOLVED QUESTIONS IN CARBON MINERALOGY

More than two centuries of mineralogical research have revealed much regarding the varied C-bearing mineral phases in Earth’s near-surface environment. Nevertheless, much remains to be learned. Among the most fundamental mineralogical questions—one evident from any museum display of carbonate minerals—is what physical, chemical, and biological processes lead to the remarkable range of calcite crystal forms? No other crystalline phase exhibits such a wide range of morphologies. What environmental factors influence calcite crystal forms? And why don’t other rhombohedral carbonates display a similar variety?

Studies of minerals that incorporate organic molecules are in their infancy, and numerous other phases at the interface between the crystalline and biological worlds are likely awaiting discovery. Characterization of such phases may prove especially important in resolving debates regarding abiotic versus biotic origins of some deep organic molecules (Sephton and Hazen 2013; McCollom 2013). Furthermore, organic mineralogy may play a dominant role on Titan, as well as planets and moons in other C-rich star systems. Indeed, carbon mineralogy may provide one of the most sensitive geological indicators of the evolution and present state of other worlds.

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