Iron Formations: Their Origins and Implications for Ancient Seawater Chemistry

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References
9.18.1 Introduction

Giant hematite and martite–goethite iron ores (≥56% Fe) hosted in iron formations (IFs) are the principal source of iron for the global steel industry. Given their economic importance, IFs have been extensively studied, but many aspects of their origin remain enigmatic because modern analogues are unknown. IFs were deposited, albeit intermittently, for more than 3 billion years, but as the Earth system changed fundamentally, so did the style of IF deposition. Aspects of, and changes in, the Earth system that are most relevant to the deposition of IF include volcanism, evolution of the biosphere, and ocean composition (e.g., Bekker et al., 2010; Holland, 2005; Huston and Logan, 2004). In this chapter, interplays among these factors and their respective links to the deposition of IF are discussed. The late-stage alteration processes responsible for the transition of IFs to economic-grade iron ore are not discussed (for recent reviews of these processes see Beukes et al., 2008; Clout and Simonson, 2005; Evans et al., 2013; Morey, 1999; Rasmussen et al., 2007; Taylor et al., 2001).

The abundance of IFs in Precambrian successions was used in early studies to argue for a largely anoxic atmosphere and ocean system (e.g., Cloud, 1973; Holland, 1984). It is generally accepted that accumulation of such large masses of iron required the transport of Fe(II), because Fe(III) is essentially insoluble at circumneutral pH values in the presence of even traces (<1 μM) of dissolved oxygen. Although earlier studies invoked a continental source of iron for IFs (Borchert, 1965; James, 1954; Lepp and Goldich, 1964), the discovery of modern seafloor-hydrothermal systems shifted emphasis to the hydrothermal processes in the deep ocean as the most likely source (e.g., Isley, 1995). Although a biological role in iron precipitation was suggested over a century ago (e.g., Gruner, 1922; Harder, 1919; Leith, 1903), the importance of microorganisms began to receive greater acceptance only with the discovery of microfossils present in Paleoproterozoic IFs in the Animikie Basin of the Lake Superior region (e.g., Barghoom and Tyler, 1965; Cloud, 1965) and, more recently, as the understanding of their significance in the modern iron cycle increased dramatically. Cloud (1965, 1973) further suggested that the redox state of the atmosphere was buffered at low levels of free oxygen, primarily by the reducing potential of the oceans and continents, including continuous IF deposition. Subsequently, it was ascertained that the Animikie IFs were deposited at ~1.88 Ga after the Great Oxidation Event (GOE) at ~2.32 Ga (e.g., Bekker et al., 2004), and that many of the Gunflint-type microfossils, interpreted earlier as oxygenic photosynthesizers, were instead likely metabolic iron oxidizers (Golubic and Lee, 1999; Planavsky et al., 2009) on the basis of their morphology and geochemical data for host rocks.

Acquisition of precise geochronologic constraints for Precambrian sedimentary successions also helped challenge the earlier assumption that IFs were continuously deposited before the rise of atmospheric oxygen (e.g., James, 1983). It is now believed that deposition of large, economically important IFs was instead restricted in time and coincided with mantle plume breakout events, as recorded by the secular distribution of large igneous provinces (LIPs), dike swarms, and submarine-emplaced mafic volcanic rocks (e.g., Isley and Abbott, 1999). These events not only provided the dissolved ferrous iron for IF, but also tempered the oceanic redox state and its chemistry by increasing the seafloor-hydrothermal flux of reductants such as H₂ and H₂S. In addition, associated with the deposition of IF, higher oceanic spreading rates, increased submarine and subaerial volcanic activity, high sea level, greenhouse conditions, and an enhanced production of volcancogenic massive sulfide (VMS) deposits are predicted consequences of mantle plume breakout events (e.g., Barley et al., 2005; Condie et al., 2001; Isley and Abbott, 1999). Considering that the typical duration for emplacement of LIPs is on the order of 10 My (Ernst and Buchan, 2001), a similar duration for the deposition of individual IFs should be expected unless a number of unrelated LIPs were emplaced closely in time (superplume breakout event).

Emerging age constraints provide further insights into IF genesis. For example, if the oceanic and atmospheric redox states are a major control on iron transport and deposition, why then were a number of giant IFs deposited at ~2.45 Ga? Deposition of these IFs occurred shortly before the first significant rise in atmospheric oxygen – GOE, thus suggesting a genetic link. Tectonically, it also coincides with a time of supercontinent assembly (e.g., Barley et al., 2005). If atmospheric oxygen rose during the early Paleoproterozoic, then what factors gave rise to a second prominent peak in IF deposition at ~1.88 Ga after a significant gap in large IF deposition? This pulse of IF deposition seemingly occurred during a mantle plume breakout event and supercontinent assembly (Bekker et al., 2010; Ernst and Bell, 2010; Hamilton et al., 2009), again suggesting a link.

The disappearance of IFs at ~1.8 Ga has historically been explained by either complete ocean oxidation (Holland, 1984) or development of sulfidic conditions in the deep ocean (Canfield, 1998). Neither of these models fully addresses the mechanism(s) that caused the ocean redox state prior to 1.88 Ga to change back to anoxic ferruginous conditions, and, subsequently after ~1.88 Ga, to either oxygenated or sulfidic conditions. A suboxic redox state of the deep ocean after ~1.85 Ga was proposed by Slack et al. (2007, 2009) on the basis of Ce anomalies and abundant hematite and magnetite in VMS-related, deep-water, oxide-facies exhalites of late Paleoproterozoic and Mesoproterozoic age. A variable, both geographically and temporally, deep-ocean redox state, including the presence of anoxic but nonsulfidic waters, in the mid-Proterozoic has also been recently proposed (Planavsky et al., 2011; Poulton et al., 2010).

It has long been argued that Archean and Paleoproterozoic (e.g., 1.88 Ga) IFs were deposited in entirely different settings and have different mineral compositions and textures (e.g., Klein and Beukes, 1992). Almost all Archean IFs consist predominantly of interbanded iron- and silica-rich layers and were generally, but not universally, deposited in relatively deep-water settings, as they typically lack evidence for wave or storm action. A large portion of the ~1.88 Ga IFs was, in contrast, deposited close to, or above, storm- and fair-weather wave base, and commonly has granular textures. These differences reflect not only distinctions in depositional settings but also different mechanisms for Fe(II) oxidation and Fe(III) precipitation.

Following a gap between ~1.85 and 0.7 Ga, when only small IFs were deposited, large IFs reappeared at the end of the Neoproterozoic, apparently related to snowball Earth.
events (Kirschvink, 1992). These are mineralogically simple iron and silica oxide deposits that, in some places, are also stratigraphically associated with economic phosphorus and manganese deposits (Klein and Beukes, 1992). These Neoproterozoic IFs and Phanerozoic IFs, termed ironstones, are not typically voluminous, relative to the much larger Archean and Paleoproterozoic deposits. Ironstones appear to be temporarily linked to marine anoxic events in many cases and comprise iron silicates and oxides without the chert enrichment, which is characteristic of Precambrian deposits, but generally with appreciable phosphorus contents. Significantly, Archean, Paleo-
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proterozoic, and Phanerozoic IFs and ironstones are temporally associated with organic matter-rich black shales (e.g., Simonson, 2003; Van Houten and Arthur, 1989). Other Phanerozoic examples of Fe-rich rocks, commonly termed
umber and jasper, developed above volcanic rocks and likely formed through either diffuse seafloor-hydrothermal venting or direct precipitation from seawater as fallout from hydrothermal plumes (Alt, 2003; Elderfield et al., 1972; Grenne and Slack, 2005). A modern analogue of umber deposits has recently been described in relation to ultradiffuse hydrothermal venting at the base of Leïi Seamount, at 5000 m below sea level (Edwards et al., 2011a,b).

Recent research has also highlighted that tectonic processes, in addition to mantle plume events and changes in iron oxidation mechanisms through time, imposed a major control over the deposition and preservation of IFs. The growth of continents created crucial shallow-water depositional sites, and enhanced preservation of IFs in the geological record (Simonson, 2003). Before large landmasses developed considerable freeboard, IFs were likely deposited in close association with volcanic edifices and were often recycled into the mantle by subduction. Certain tectonic settings, such as isolated to semi-isolated back-arc, rift, and foreland basins, have been invoked to explain the iron source to, and basin-scale water column redox stratification in, basins, in which IFs were deposited (e.g., Beukes and Gutzmer, 2008; Ohmoto et al., 2006). Further, it is now possible with high-precision geochronology not only to correlate IFs of similar ages in different basins but also to establish that IF deposition in one basin coincided with the lack of iron enrichment in another. Nonetheless, it remains difficult in the case of Precambrian basins to separate basin and paleogeographic controls from those determined by ocean circulation and upwelling processes. Exhalites that formed distal to deep-water VMS deposits in open-marine settings help in evaluating ancient ocean redox states (Slack et al., 2007, 2009). This information can be directly compared with inferences from correlative IFs in order to constrain the redox conditions of the coeval global ocean.

Although tectonic processes exert a first-order control on Fe flux to the ocean, the marine redox state as established by oxygen content and oxidation state of sulfur and nitrogen determines whether iron and manganese can be transported in solution within the ocean. In addition, secular changes in seawater silica concentrations influenced the composition of IFs, specifically their silica and trace element contents. It is generally accepted that seawater silica contents were high during the Precambrian, but it is unclear whether silica in seawater declined dramatically at the beginning of the Phanerozoic (Siever, 1992), or if silica concentrations remained high until the Cretaceous when the emergence of diatoms removed most silica from seawater (Grenne and Slack, 2003).

In this chapter, we place the deposition of IFs in a framework of broader changes in tectonics, mantle plume activity, and oceanic and atmospheric redox states in order to assemble a new integrated model for their deposition. It is highlighted that no single parameter is solely responsible for IF deposition. Rather, complex give-and-take relationships among all of these parameters determined the time intervals and settings in which IFs were deposited throughout Earth history, as well as changes in their mineralogy and composition. Figure 1 shows the global distribution of large IFs (≥1000 Gt) and selected smaller deposits discussed below.

9.18.2 Definition of IF

The term ‘iron formation’ has often been restricted to stratigraphic units composed of layered, bedded, or laminated rocks (Figures 2–5) that contain 15 wt% or more iron, and where the iron minerals are commonly interlayered with quartz, chert, or carbonate (Gross, 1980; James, 1954). James (1954) defined four facies of IF: silicate, carbonate, oxide, and sulfide. Sulfide-facies IFs are pyritic carbonaceous shales or slates, and, as such, not typically considered IF in the strictest sense. Barren or mineralized, seafloor-hydrothermal, iron-rich exhalites and sulfidic cherts are in some cases also assigned to the IF category and host some important gold deposits in Archean terranes. The former, in many cases, represent facies of VMS deposits, whereas the latter could be either exhalites or hydrothermally replaced IF (Groves et al., 1987). As a result, many Archean (sulfide-facies) IFs described in the literature are not true IFs (Hofmann et al., 2003). All other facies are generally interbedded with variably recrystallized chert (Simonson, 2003). Oxide-facies IF consists predominantly of magnetite or hematite, whereas carbonate-facies varieties contain siderite or ankerite as major constituents. The mineralogy of silicate-facies IFs is more complex and depends to a large extent on the degree of metamorphism. Under relatively low-grade metamorphic conditions, at the biotite zone and below, greenalite, minnesotaite, stilpnomelane, chamosite, ripidolite (Fe-chlorite), riebeckite, and ferri-annite may be present. At higher grades, cummingtonite, grunerite, pyroxene, garnet, and fayalite can occur.

On the basis of interpreted depositional settings, IFs have classically been subdivided into Superior type and Algoma type (Gross, 1980; see also discussion in Bekker et al., 2012). Superior-type IFs were regarded as having been deposited in nearshore continental-shelf environments, because they typically are interbedded with carbonates, quartz arenite, and black shale, but only with minor amounts of volcanic rocks (Gross, 1980). Algoma-type IFs are generally hosted within volcanic rocks, and, in some cases, in graywacke, and apparently formed by exhalative-hydrothermal processes close to volcanic centers. Barrett et al. (1988a) argued, on the basis of geochemical signatures, that some Algoma-type deposits formed within restricted basins like the modern Red Sea. Algoma-type IFs are present in volcano-sedimentary sequences of greenstone belts ranging in age from Eoarchean to Phanerozoic (Goodwin, 1973; Huston and Logan, 2004; Isley and Abbott, 1999; James, 1983; Peter, 2003; Peter et al., 2003).
Although common in Superior-type IF, granular iron formations (GIFs) are generally absent within Algoma-type deposits, which are typically banded with chert and Fe oxide, silicate, or carbonate pairs on various scales (Figure 2(a)). GIFs older than 3.0 Ga appear to be predominantly of the Algoma type, which likely reflects the scarcity of preserved cratonic successions prior to this time. Mineralogically, Algoma- and Superior-type IFs are similar.

Algoma-type IFs are generally thinner and smaller in lateral extent relative to Superior-type IFs and rarely exceed 10^7 Mt ( Huston and Logan, 2004), although the former are more abundant in terms of numbers of deposits and geographic distribution ( Beukes and Gutzmer, 2008). Typically Algoma-type IFs are less than 50 m thick and rarely extend to more than 10 km along strike. However, these characteristics do not indicate that all Algoma-type IFs were originally smaller, as most had been affected by deformation and tectonic dismemberment, implying that their original size and extent are likely underestimated ( Gole and Klein, 1981). By contrast, Superior-type IFs are typically laterally more extensive and thicker than Algoma-type IFs. Some major, Superior-type sedimentary deposits (not orebodies) initially contained an estimated 10^8 Mt at 15 wt% Fe ( Isley, 1995; James, 1983).

Clear differentiation between Superior and Algoma types of IF is difficult in Archean successions affected by strong deformation and shearing that produced tectonic dismemberment or imbrication of genetically unrelated sequences. For example, IFs interbedded with quartz arenite and carbonate are locally interlayered with thick packages of mafic–ultramafic extrusive rocks. Although some of these sequences may represent primary stratigraphic units that were deposited in extensional continental or arc settings (e.g., Srinivasan and Ojakangas, 1986), others could have formed by the tectonic imbrication of cratonic cover sequences and overlying piles of mafic–ultramafic rocks (e.g., Dirks et al., 2002).

This distinction is further complicated by the full gradation between Superior- and Algoma-type IFs that emerged as studies advanced. For example, bimodal volcanic rocks, tuff beds, and stilpnomelane-rich shale are commonly associated with IFs in the Anikimik basin of North America and the Hamersley Group of Western Australia, both being typical examples of Superior-type IFs. On the other hand, some Paleoarchean and younger IFs in greenstone belts occur in sedimentary successions containing minimal recognizable volcanic material. However, the geochemistry and lithology of IFs and host rocks clearly indicate that both Algoma- and Superior-type IFs were deposited contemporaneously with submarine volcanism and intense hydrothermal activity. Therefore, the two types of IF can be considered as idealized end-members for precipitates ranging from proximal hydrothermal deposits to distal hydrothermal deposits that all precipitated from seawater coeval with seafloor-hydrothermal activity.
The gradient in hydrothermal influence on IF is also reflected in their geochemistry. Compositions of the Algoma-type end-member typically record local volcanic or hydrothermal conditions, rather than being representative of the large-scale chemistry of the oceans during their formation. In contrast, the deposition of Superior-type end-member reflects processes that probably acted on a global scale, and thus they are likely more representative of seawater composition (Huston and Logan, 2004). However, the potential influences by nearby cratonic areas also need to be considered (e.g., Alexander et al., 2009). In general, for discerning whether the geochemistry of IF reflects local or global seawater composition,
compositions, a simple distinction between Algoma and Superior types is insufficient. Instead, one has to constrain the tectonic setting and degree of isolation of the basin in which IFs were deposited, relative to the global ocean, before any inferences regarding composition and redox state of ancient seawater can be made.

**9.18.3 Mineralogy of IF**

IFs are defined by their unusual mineralogy, which includes mostly silica and a wide range of Fe-rich and Al-poor minerals. Most IFs comprise layers containing magnetite and/or hematite, which alternate on the scale of several millimeters with bands of microcrystalline silica, forming microbands (Figures 2 and 3). Well-banded IFs (Figures 2 and 3), known as banded iron formations (BIFs), are mostly restricted to Archean and early Paleoproterozoic sequences. Large portions of late Paleoproterozoic IFs from the Superior and Slave cratons in North America and the Capricorn Orogen in Western Australia comprise sand-sized grains that commonly are cross-bedded and lack the finely laminated textures of BIF; these are generally referred to as GIFs (Figure 5). GIFs are typically intercalated with well-laminated IFs, Fe-rich mudstone, mafic and felsic volcanic rocks, and carbonate and sandstone.

The mineralogy of BIF and GIF from the best preserved sequences is remarkably uniform, comprising mostly silica, magnetite, hematite, Fe-rich silicate minerals (stilpnomelane, minnesotaite, greenalite, and riebeckite), carbonate minerals (siderite,ankerite,calcite, and dolomite), and, less commonly, sulfides (pyrite and pyrrhotite).

Chert (and crystalline quartz in metamorphosed IFs) is ubiquitous in all types of IF. In BIF, chert layers are commonly banded, alternating with millimeter-thick laminae of Fe-rich silicate and carbonate minerals. Individual laminae are wavy to wrinkly and, locally, appear to truncate against overlying laminae. In places, chert forms precompaction nodules draped by compacted laminae, suggesting an early paragenesis for the nodular chert. In GIF, the chert ‘peloids’ show open packing, indicating precompaction lithification.

Magnetite is widespread in IFs, where it occurs as euhedral, fine- to coarse-grained crystals (Figures 2 and 3). It is particularly abundant in cherty Fe-rich layers as laminae comprising dense clusters of intergrown euhedra (Figure 3(a)–3(c)). Magnetite commonly is replaced by hematite (termed martite) and locally replaces carbonate minerals. Magnetite is clearly a secondary mineral that formed mostly during the late history of the

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**Figure 3** Photomicrographs from the Dales Gorge Member of the Brockman Iron Formation, Hamersley Province, Western Australia. (a) Banded iron formation comprising several larger clusters of magnetite euhedra in a band of laminated hematite. Field of view is 2.5 mm. (b) Two magnetite euhedra in a matrix of hematite, silica and carbonate. Field of view is 1.25 mm. (c) Alternating laminae of silica and iron oxides. Hematite cores are surrounded by magnetite. Field of view is 0.3125 mm. (d) Laminae of fine-grained hematite alternating with bands of chert. Field of view is 0.625 mm. (e) Band of laminated stilpnomelane and fine-grained siderite. Field of view is 0.625 mm. (f) Impact ejecta layer in the Dales Gorge S-band #4 containing melt spherules largely replaced by stilpnomelane. Field of view is 6 mm.

Hematite is the most common Fe-oxide mineral in IFs, where it typically occurs with magnetite in millimeter- to centimeter-thick layers (Figure 3). Together with magnetite, hematite defines the lamination in most chert layers. Hematite may also be present in some intercalated mudstones, but in this case, it is much less abundant than magnetite (or pyrite). The timing of hematite growth is texturally ambiguous, although rare hematite spheroids (Ayres, 1972) likely represent some of the earliest components of IFs (Figure 4(e) and 4(f)). If they did form very early, then the precursor phase probably was a form of ferric oxyhydroxide, such as ferrihydrite.

Stilpnomelane is an Fe- and K-rich, Al-poor, hydrous silicate mineral, with a composition that is similar to that of sedimentary nontronite, an iron-rich, alumina-poor smectite. Stilpnomelane is diagnostic of lower greenschist facies metamorphic conditions. It is the most common Fe-silicate in many IFs, where it occurs as highly pleochroic plates and fibers, forming solid bands interlaminated with secondary carbonate (Figure 3(e)), and as irregular mattes and sheaves. Stilpnomelane is the main constituent of most mudstones associated with IFs, and is also a common component of chert and tuff layers and impact ejecta layers (Figure 3(f)) intercalated with IFs. For example, it replaces volcanic glass in felsic tuff beds and, locally, fine-grained stilpnomelane-filled spheres occur in laminated chert beds (Ayres, 1972; Figure 4(a)). At higher temperatures (>400 °C), stilpnomelane is replaced by biotite.

Minnesotaite is an Fe- and Mg-rich hydrous silicate that typically is less abundant than stilpnomelane, although there are some exceptions in sections of the late Paleoproterozoic IFs in the Superior craton. Minnesotaite occurs as radiating plates

Figure 4  Samples from the Dales Gorge Member of the Brockman Iron Formation, Hamersley Province, Western Australia. (a) Numerous spherical structures composed of stilpnomelane in a chert matrix. Field of view is 0.625 mm. (b) Fine-grained sprays of minnesotaite needles arranged in typical 'bow-ties.' Field of view is 0.3125 mm. (c) Coarse euhedral ankerite rhomb enclosing crystals of hematite and riebeckite in a chert matrix. Field of view is 0.625 mm. (d) Back-scattered electron image of inclusion-rich apatite in a chert matrix. (e) Plane-polarized light image of numerous minute hematite crystals (opaque) preserved in a chert–hematite layer. Field of view is ~0.3 mm. (f) Reflected light image of (e) showing hematite crystals (white) containing numerous minute inclusions (speckled) in a matrix of chert (dark gray). Field of view is ~0.3 mm.
and needles, forming common ‘bow-tie’ texture (Figure 4(b)) and forms during metamorphism and late-stage hydrothermal alteration.

Riebeckite is a Na-rich amphibole that occurs in most BIFs from the Hamersley region in Western Australia and in the Kaapvaal craton of South Africa (Beukes, 1973; Klein and Gole, 1981). Riebeckite-rich layers are characteristically blue in hand specimen. Riebeckite forms clusters of randomly oriented fibers, which locally may be aligned to define a structural fabric. Paragenetic relations indicate that riebeckite formed late in the history of IFs. The fibrous form of riebeckite (crocidolite) forms veins that are a source of asbestos; these are overpressure veins that formed during regional folding (Krapež et al., 2003).

Greenalite is a pale green Fe-silicate that occurs in many IFs but typically is less abundant than stilpnomelane and minnesotaite. Greenalite is rare in the Hamersley and Transvaal BIFs, but where present occurs as a late-stage, secondary mineral. However, in GIF from the Superior craton, greenalite is more abundant and appears to be among the earliest minerals to have formed (Klein, 2005). Nevertheless, greenalite is not an original sedimentary mineral.

Siderite is the most common carbonate in BIFs, typically occurring as minute, single, globular crystals or massive layers of microcrystalline crystals. In some beds, siderite may comprise >50% of the rock. Small siderite nodules are common in some layers, where they display evidence of differential compaction, thus indicating that the nodules probably formed before deep burial. The texturally earliest siderite is present as minute (<20 μm) spheroids, which occur in some IFs and intercalated mudstone from the Hamersley Province (Ayres, 1972). This siderite appears to have replaced or overgrown an earlier phase, which occurs in the core of some crystals, indicating a diagenetic origin for the siderite. Sideritic BIF in many cases is not a primary sedimentary facies, because it can be closely linked to near-shore alteration.

Ankerite and ferroan dolomite occur in many chert layers typically as coarse, euhedral rhombs that overgrow and contain inclusions of chert, hematite, magnetite, and Fe-silicate minerals (Figure 4(c)), indicating that these carbonates were among the last minerals to form.

Trace minerals include pyrite, apatite, monazite, xenotime, zircon, ilmenite, and K-feldspar. Trace amounts of apatite have been reported from most IFs, forming euhedral crystals containing abundant inclusions of silica and other matrix minerals (Figure 4(d)). Apatite, even from very low metamorphic grade IFs, lacks significant carbonate and fluorine substitution into the crystal structure (e.g., Li et al., 2011), which would be expected if the mineral had formed during early marine diagenesis. Therefore, apatite is likely a burial diagenetic and metamorphic product. Apatite formation is probably associated with phosphorus exclusion during recrystallization of iron oxides or oxidation of buried organic matter (Li et al., 2011). Apatite is the most abundant phosphate mineral, but monazite and xenotime are also present in some IFs. Monazite forms minute (typically <100 μm), inclusion-rich aggregates in mudstone, whereas xenotime occurs mostly as overgrowths on zircon grains, which are likely diagenetic in origin. Zircon is rare in IFs, but can be locally abundant in millimeter- to centimeter-thick layers of felsic tuff (Pickard, 2002, 2003). Xenotime overgrowths can be dated in situ by the U-Pb SHRIMP method to obtain ages for IF upgrading (e.g., Rasmussen et al., 2007).

In conclusion, no unambiguously original sedimentary minerals occur even in the best-preserved IFs that only underwent very low-grade metamorphism. Microspherical and nodular textures appear to be among the earliest features of the IFs; however, it is highly unlikely that the minerals associated with those textures are original.

9.18.3.1 Precursor Sediments

Since all IFs have undergone significant modifications even during diagenesis and prehnite–pumpellyite facies metamorphism, their mineralogy reflects a combination of factors, including the original bulk composition of the precursor sediment, diagenetic and metamorphic conditions, and post-depositional fluid flow. Effects of increasing temperature and pressure have yielded a progressive change in mineralogy through replacement and recrystallization, increase in grain size, and obliteration of primary textures (Klein, 2005).

The alternating layers of magnetite and hematite are commonly interpreted to have formed from a ferric iron oxyhydroxide rain to the sediment pile. During early diagenesis, the ferric oxyhydroxide was apparently converted to hematite. It is possible that ferric iron and dissolved/absorbed ferrous iron combined to form a mixed-valence iron phase that later was converted to magnetite during late diagenesis and metamorphism. However, the magnetite could have also formed entirely during metamorphism, in cases with organic matter acting as the reductant (e.g., Perry et al., 1973; Tompkins and Cowan, 2001). Although the texture of most of the magnetite and hematite grains indicates a secondary, postdepositional origin, rare spheroids of hematite, about 5–20 μm in diameter, as mentioned above, have been considered to be among the earliest textures known in BIFs (Ayres, 1972). The spheroids are interpreted to represent original iron oxyhydroxides that were converted to hematite during diagenesis or regional metamorphism. Similar spheroidal textures comprising solid stilpnomelane and siderite are also present (Figure 4(a); Ayres, 1972).

Silica is widely considered to have been delivered to the sediments absorbed on iron oxyhydroxides, scavenged with organic matter, and precipitated from the water column at the sediment–water interface in colloidal form (e.g., Fischer and Knoll, 2009; Krapež et al., 2003; Grenne and Slack, 2005). An alternative interpretation is that the chert formed largely beneath the sediment–water interface as a replacement of a precursor sediment (Krapež et al., 2003).

Recently, Rosing et al. (2010) proposed that the ubiquitous coexistence of magnetite and siderite in Archean and Paleoproterozoic IFs could be used to constrain early Earth’s atmospheric composition. They suggested that atmospheric CO₂ and CH₄ concentrations have been consistently overestimated in modeling early Earth’s energy balance to explain the presence of liquid water on Earth at the time when, in the Sun’s youth, solar luminosity was greatly reduced. Rosing et al. (2010) suggested that the coexistence of siderite and magnetite in IFs represents an assemblage that is close to thermodynamic equilibrium with the atmosphere, thus constraining the partial pressures of CO₂ and H₂. However, both these minerals are of diagenetic and metamorphic origins. Further, even in the
unlike the case that IF mineral assemblages formed during early diagenetic conditions in an environment close to chemical equilibrium with seawater, they cannot be used to constrain atmospheric carbon dioxide levels. For instance, dissimilatory iron-reducing bacteria have been observed to generate magnetite even at very high aequous $\mathrm{CO}_2$ concentrations ($\sim 50$ mM; Behrends and Van Cappellen, 2007) and headspace $\mathrm{pCO}_2$ ($\sim 0.2$ atm; Roh et al., 2003). Therefore, $\mathrm{pCO}_2$ values greatly elevated above modern levels do not preclude the formation of magnetite at any stage in IF evolution. IF mineral assemblages thus cannot be used to place constraints on past atmospheric $\mathrm{CO}_2$ concentrations. More broadly, since almost all Fe-bearing minerals in IFs formed at different times during diagenesis or metamorphism, their chemical and isotopic compositions are unlikely to reflect chemical equilibrium with seawater. Accordingly, the use of mineral assemblages to infer composition of the environment in which they formed, or extrapolations regarding composition of the atmosphere–ocean system, is typically plagued with uncertainties.

### 9.18.3.1.1 Secular trend in Fe mineralogy of GIFs

The secular change in the mineralogy of the same sedimentary facies may, however, provide insight into environmental change. Specifically, the mineralogy of GIFs may record atmosphere–ocean redox state and evolution leading to deposition of IFs. GIFs first appear at $\sim 2.6–2.5$ Ga in South Africa, Western Australia, and Brazil (Beukes and Klein, 1990; Simonson and Goode, 1989; Spier et al., 2007). However, most granules and sand-sized grains in these older GIFs consist of minerals having reduced and mixed-valence iron (e.g., magnetite, Fe-silicates, and carbonates); only rarely have hematite granules been observed and never as coated grains or oolites (Beukes and Klein, 1990; Simonson and Goode, 1989; Spier et al., 2007). In contrast, the extensive GIFs of the Animikie basin that were deposited at $\sim 1.9$ Ga contain oolites, coated grains, and granules made of minerals having predominantly oxidized iron (e.g., hematite), although reduced or mixed-valence coated grains, and granules are also present. Similar to the Animikie basin, younger Proterozoic IFs and Phanerozoic ironstones having coated grains are predominantly composed of ferric iron.

The secular trend in the distribution of mixed-valence iron silicates is poorly established. Early work suggested that glauconite did not form in open-marine, shallow-water settings until the Phanerozoic (Cloud, 1955). However, this view was rebutted by Kimberley (1989), who described Proterozoic examples including the $\sim 2.3$ Ga Gordon Lake Formation (Chandler, 1986) and emphasized that Archean examples are missing from the rock record. Indeed, mixed-valence iron silicate minerals such as berthierite, chamosite, and glauconite are absent in shallow-water Archean successions, even though they became abundant during the GOE (e.g., ironstone of the Timeball Hill Formation; Dorland, 1999). We infer that the lack of these minerals in shallow-water, Archean settings reflects low oxygen concentration that inhibited formation of mixed-valence Fe-silicates. In contrast, in Archean deep-water settings, mixed-valence Fe-silicates such as the precursors to stipnomelane and greenalite are common, indicating an upside-down redox profile similar to that documented in the Transvaal basin by combined sedimentary facies and mineralogical study (Beukes and Klein, 1990). Walker (1984) linked this counterintuitive redox structure to decreasing organic carbon fluxes offshore. Thus, the study of iron oxidation in mixed-valence Fe-silicate minerals holds potential to reveal information about ocean oxidation in the past. Specifically, the marked absence of mixed-valence iron minerals prior to $\sim 2.3$ Ga is one of several lines of evidence pointing to the lack of a discrete redoxcline and predominantly anoxic marine conditions in the Archean.

### 9.18.4 Depositional Setting and Sequence-Stratigraphic Framework

Depositional environments of IFs range from deep water, beyond or above a continental slope for BIF to shallow water, above storm- and fair-weather wave base for GIF. Both BIF and GIF were deposited during periods of high or rising sea level (cf. Fralick and Pufahl, 2006; Krapež et al., 2003; Simonson and Hassler, 1996).

Detailed sedimentologic studies of IFs have been conducted in the Hamersley Province of Western Australia, and in the Transvaal and Griqualand West structural basins of South Africa. There is no evidence from these studies that depositional environments were restricted. For instance, in the Hamersley Province, basin architecture did not change conclusively during the deposition of BIF and associated non-BIF facies, but rather the change that did occur involved a higher hydrothermal flux of reduced iron ($\mathrm{Fe}^{2+}$) to the basin (Krapež et al., 2003). It is therefore likely that increased hydrothermal activity rather than changes in basin architecture was the first-order control on that flux.

Studies of BIF of the Hamersley Province have documented the presence of iron minerals and chert in layered pairs that vary from 0.2- to 2.0-mm-thick microbands to 10- to 50-mm-thick mesobands. Initially, Trendall and Blockley (1970) (see also Trendall, 1990) suggested that such microbands and mesobands could be correlated basin-wide, which led to the inference that BIFs are chemical varves (Morris and Horwitz, 1983; Trendall, 1973). More recent studies, however, have shown that only the chert mesobands can be correlated (Krapež et al., 2003).

Through facies and sequence-stratigraphic analyses, Krapež et al. (2003) and Pickard et al. (2004) concluded that all chert in BIF is diagenetic in origin. Moreover, they concluded that chert mesobands are siliceous equivalents of modern-day seafloor hardgrounds (Figure 2), in which silica replaced precursor sediment at or below the sediment–water interface. Three-dimensional and microscale lenticularity of chert and relics of precursor sediment within lamina sets and discontinuous bands (Figure 2(b) and 2(c)), as well as erosion surfaces on bedded cherts, show that chert has a replacement origin and formed during early diagenesis, prior to compaction.

Minute spheroids ($\sim 10$ m diameter) documented in the Brockman Iron Formation (Hamersley Group) were interpreted to record paragenetically early textures (Ayres, 1972). The spheroids are distributed along laminae in bedded chert, and along the chert laminae of BIF. Assuming that lamina sets in BIF originated from a process similar to that responsible for the lamina sets in lithofacies interbedded with BIF, they can be assumed to have a density current origin (Krapež et al., 2003). This density current interpretation is supported by the presence in BIF of two bedding styles of mesobands: microbanded and
tabular-bedded (Figure 2); the latter typically is massive or weakly laminated (Ewers and Morris, 1981; Morris, 1993). Massive mesobands grading upwards into microbanded mesobands resemble density-current intervals, and are preserved at various scales (Figure 2(e) and 2(f)); some of those massive intervals are internally graded and contain tabular, mm-scale detrital fragments of shale. Occurrences of erosional truncation of draped laminae (Krapež et al., 2003) also support the density-current interpretation (Figure 2(b) and 2(c)). Resedimentation occurred either by bottom currents or gravity-driven turbidity currents, and the resulting sediment bodies may have been contourite drifts (Krapež et al., 2003). The precursor sediments to BIFs were therefore microgranular in texture. Krapež et al. (2003) concluded that the precursor sediments could have been granular hydrothermal muds, composed of iron-rich smectite and particles of iron oxohydroxide and siderite that were deposited on the flanks of submarine volcanoes.

The sequence architecture of formations containing BIF is identical to that of clastic sedimentary rocks that accumulated beyond the continental slope, comprising lowstand fans of shelf-derived sediment overlain by condensed sections of pelagites, hemipelagites, or intrabasinal clastic sediment (e.g., Haq, 1991). Depositional sequences in the Dales Gorge Member of the Hamersley Province, documented by Krapež et al. (2003), comprise lowstand density-current deposits (dolostone, graded shale, and rare conglomerate) overlain by a condensed section of BIF. Bedded chert defines the top of each depositional sequence by showing: (1) sharp, erosional contacts with overlying dolostone or mudstone; and (2) gradational contacts into underlying BIF. In contrast, the contacts from lowstand mudstone to condensed-section BIF are transitional (Krapež et al., 2003).

Lowstand deposits change from dolostone-shale to shale-shale only toward the southwest in the Hamersley Province, indicating a paleoslope to the southwest (Simonson et al., 1993); an overall thinning of lowstand deposits accompanies this down-paleoslope change. Lowstand deposits in distal sections contain BIF units that are identical to BIF in the condensed sections. This pattern suggests that the precursor sediments to BIF characterized basin sedimentation not only during rising and high sea level, but also during some lowstands, beyond the limits of shelf-derived resedimented sediments. Isopachs of BIF presented by Trendall and Blockley (1970) show that thickness variations define mounds elongated across the paleoslope. These trends, combined with the sequence architecture, indicate that the source of precursor sediments for BIF was within the basin realm and that the depositional system was some form of contourite current. A contourite drift is a sedimentary deposit that accumulates along the continental slope, from density currents that follow the contours of the basin floor, possibly entering the basin from a distal submarine canyon. For the precursor sediments to BIF, the deep-sea currents may have traveled down canyons headed in volcanic complexes. The implication is, therefore, that some precursor sediments to BIF were hydrothermal clays or clays derived by submarine weathering of basalts that accumulated on the slopes of, and among, submarine volcanoes (Krapež et al., 2003).

This model for deposition of the Dales Gorge BIF is applicable to other deep-water BIF such as those in the correlative Transvaal (South Africa), Krivoy Rog (Ukraine), Kursk Magnetic Anomaly (KMA, Russia), and Quadrilátero Ferrífero (Brazil) successions (Appendix 1). It is, however, not applicable to shallow-water IFs, such as those in the Pontonga and Witwatersrand superfamilies (Beukes and Cairncross, 1991), where responses to rising and falling sea levels differed according to sequence stratigraphy. Detailed sedimentological models for the Algoma-type IFs or thin BIF interbedded with shallow-water deposits, which integrated a basin-scale and sequence-stratigraphic approach, are not available and therefore it is premature at this point to discuss their sedimentological setting relative to the predictive architecture of deep- or shallow-water depositional environments.

GIFs are clastic sedimentary rocks that are largely restricted, at least in terms of preservation, to continental basins of Paleoproterozoic age. Paleoproterozoic basins surrounding the Superior craton of North America constitute the type area, where two lithofacies were recognized for a long time: slaty and cherty GIF (Ojakangas, 1983). The so-called slaty lithofacies is iron-rich shale comprising alternating, millimeter-scale, parallel- and wavy-laminated layers of iron-oxides or -silicates and chert, interbedded with lenses of grainstone. Lamina sets are similar to those in BIF, being made up of a basal layer rich in iron oxides and an upper chert-rich layer; gently-dipping erosive truncations draped by mudstone are also present (Pufahl and Fralick, 2004).

The cherty lithofacies is a grainstone with a cherty cement; in situ and reworked stromatolites are common (Ojakangas, 1983; Planavsky et al., 2009; Pufahl and Fralick, 2004). This lithofacies comprises interconnecting lenses of trough cross-stratified grainstone. The largest lenses typically have a basal layer of intraformational breccia derived from reworking of the underlying iron-rich mudstone.

Millimeter- to centimeter-scale, grain-size grading in iron-rich shale indicates that the depositional process was gravity settling. Occurrence of a siliciclastic component to the mudstone/shale supports suspension deposition from density currents. Density-current settling is common along current-, wave-, and storm-dominated shores, such that micrograded sets of mudstone are essentially the ambient sedimentary style. Traction currents likely were key in the deposition of granular beds within the mudstone sequences. Sets are channel-shaped, showing internal trough cross-stratification. Evidence of wave- and current-formed sedimentary structures and hummocky cross-stratification is abundant (e.g., Ojakangas, 1983; Pufahl and Fralick, 2004), establishing that the depositional environment was a shallow-water shelf disturbed by storms and influenced by significant sea-level changes. The environment also was coeval with bimodal volcanism, indicating almost certainly a volcano-tectonic influence. In general, the stratigraphic pattern is one of alternating packages of storm-influenced event beds and background density-current deposits, passing upwards into shallow-water grainstones of the cherty lithofacies (Figures 5 and 6(a)). The sequence-stratigraphic architecture therefore is upwards thickening and coarsening, reflecting deposition during rising and high sea level on a shelf. In the shallowest parts of the Animikie basin, these shallow-water facies can be capped by exposure surfaces, indicating IF filled the accommodation space. Although contrasting with the sequence architecture of BIF of the Hamersley
basins (Krapež et al., 2003), the two styles are linked and are in agreement with a classic sequence-stratigraphic profile, defined by GIF on the shelf and BIFs on the deep-water basin floor. For GIF of the Superior craton, the transition between pairs of slaty and cherty lithofacies is a unit of convolute IF (Figure 6(a)).

Sequence-stratigraphic models for Phanerozoic (and some Mesoproterozoic) GIFs (ironstones) are slightly different because they comprise thin and locally discontinuous lithofacies within siliciclastic depositional sequences. Maynard and Van Houten (1992) suggested that oolitic ironstones were deposited after the peak of regression and prior to the peak of transgression (Figure 6(b)). The ironstones are capped by a hardground, which in the Mesoproterozoic oolitic IFs of northern Australia, is recognized by ooids, granules, and intraclasts floating in an early diagenetic chert matrix/cement (Harms, 1965). Cherty hardground records the peak of siliciclastic sediment starvation on the shelf, thereby representing a maximum flooding surface (Figure 6(b); cf. Fürsich et al., 1992; Pope and Read, 1997). Oolitic IF within siliciclastic depositional sequences therefore formed during transgression rather than still-stand. Not only would peak flooding be the time of high organic productivity and anoxia on the shelf, but it likely would also be the period of maximum ingress of basinal waters, again suggesting that a basinal supply of Fe(II) was a key aspect of IF.

9.18.4.1 Basin-Type Control on IF Deposition

After the rise of atmospheric oxygen during the GOE, isolated to semi-isolated basins favored the development of conditions necessary for Fe(II) transport. Restriction commonly results in less vigorous circulation and limits exchange between oxygenated surface waters and bottom waters. Additionally, a smaller flux of reductants was required to induce anoxia. Under these conditions during the Phanerozoic, small IFs and exhalites formed in arc-related basins in association with volcanic successions. Before the GOE, ocean redox state was not a limiting factor for the deposition of IFs. However, large hydrothermal fluxes were required to deliver iron from the site where it was released from volcanics to the

Figure 5  Middle Proterozoic GIFs. (a) Trough cross-bedded GIF of the c.1.88 Ga Temiscamie Iron Formation, Mistassini basin, Quebec. (b) Quartz-rich sandstone overlying ooidal ironstone and containing large rounded clasts of cemented ooidal ironstone and wavy heavy mineral bands composed of hematite; Train Range Member of the Mullera Formation, Constance Range, Northern Territory–Queensland border, Australia. (c) Graded bed comprising granules and ooides. Field of view is 2 cm. (d and e) Rounded granules comprising hematite, magnetite, chlorite, chert, and carbonate “floating” in a matrix of carbonate cement. Field of view is 6 mm in (d) and 2.5 mm in (e). (c–e) c.1.88 Ga Frere Formation, Earraheedy Group, Western Australia.
depositional site on continental margins and oceanic plateaus, where it could survive subduction.


Knowledge of sedimentation rates of IFs is important for understanding their iron sources and mechanisms of deposition, as well as depositional settings that are favorable for their formation. There have been many geochronologic studies (e.g., Altermann and Nelson, 1998; Arndt et al., 1991; Barley et al., 1997; Barton et al., 1994; Pickard, 2002, 2003) that discuss sedimentation rates of the precursor sediments to BIF. Inferred compacted sedimentation rates range from as low as 2–6 m My^{-1} to as high as 30–33 m My^{-1}. However, there has been limited attention paid to extremely pulsed nature of IF deposition. Krapež et al. (2003) showed that each sedimentation unit of Hamersley BIF is capped by a seafloor hardground of chert, and that many of those hardgrounds were eroded prior to deposition of subsequent sets. Therefore, it is difficult to determine the significance of average depositional rates for isolation, and submarine volcanic activity would have all favored IF development. This framework not only allows the screening of basins for their potential to have large IFs, but also justifies the use of known IFs as markers of condensed sections on a basin scale and for interbasinal correlation.

9.18.4.2 Sedimentation Rates
BIF when so much time within each sedimentation unit and depositional sequence had been taken up by non-depositional processes such as erosion and subseafloor silica replacement. Clearly, depositional rates so calculated significantly underestimate true depositional rates. Although all sedimentary rocks are characterized by pulsed deposition (Sadler, 1981), the sedimentary architecture of IFs suggests that they have experienced extreme (in frequency and duration) sedimentary hiatuses. Therefore, although previous estimates of depositional rates for IFs are similar to those for typical deep-sea sediments, there undoubtedly were pulses of rapid sedimentation similar to, or likely even more extreme than, those characteristic of near-axis, modern deep-sea sediments.

Based on the sequence analysis presented in Krapeč et al. (2003), and known geochronological constraints, each depositional sequence spans an average duration of about 1.5 My, roughly equal to the duration of third-order eustatic cycles possibly driven by pulses in spreading rates of mid-ocean ridges. The thickness of each depositional sequence is not constant, and neither is the proportion of diageneric lithofacies, further negating the meaning of IF sedimentation rates based on the ratio of averaged thickness to averaged time interval. It is therefore reasonable to infer that rates as high as 33 m My⁻¹ for the maximum compacted sedimentation rate are an underestimate, and that even higher sedimentation rates should be considered in modeling IF deposition.

9.18.5 IF: A Proxy for Ancient Seawater Composition

9.18.5.1 Trace Elements

IFs, as chemical deposits, are among the most obvious lithologies to use to investigate the composition of ancient oceans. Their precursor minerals precipitated directly from, or formed by, interaction with seawater. Moreover, many IFs contain low concentrations of crustally sourced elements such as Al, Ti, Zr, Th, Nb, and Sc, which supports an authigenic origin. Hence, secular changes in IF composition have long been used as proxies for the chemical evolution of seawater over time (e.g., Bau and Duslki, 1996; Bolhar et al., 2002; Jacobsen and Pimentel-Klose, 1988), and, most recently, to evaluate the types of nutrients that were available to ancient marine life (e.g., Bjerrum and Canfield, 2002; Konhauser et al., 2007a,b; Planavsky et al., 2010a,b). Four examples of how hydrogenous trace elements can track seawater composition are provided below.

9.18.5.1.1 Rare earth elements

Analysis of rare earth element (REE) patterns is a powerful tool to understand conditions under which IFs were deposited (e.g., Alexander et al., 2008; Bau and Duslki, 1996; Bau and Möller, 1993; Derry and Jacobsen, 1990; Frei et al., 2008; Fryer, 1977; Kato et al., 2006; Klein and Beukes, 1989; Planavsky et al., 2010a,b). Given a constant ionic charge, all of the REE should generally display similar behavior, with differences being linked to ionic radius. The most notable exceptions are the redox-related anomalies shown by Ce and Eu. Ce can be oxidized from the trivalent to tetravalent state under similar redox conditions to Mn(II) oxidation. In high-temperature (>250 °C) hydrothermal systems, Eu can undergo reduction from the trivalent to divalent state (Sverjensky, 1984), resulting in positive anomalies in hydrothermal fluids relative to neighboring lanthanide series elements (Sm and Gd). Fluid pH and ligand (sulfate, chloride, and fluoride) concentrations have an effect on REE patterns of hydrothermal fluids formed in back-arc basins such as the Manus basin (Cradock et al., 2010). Additionally, certain REE (foremost La and Gd) display anomalous bonding behavior linked to their f orbital configurations in low-temperature aqueous systems. This anomalous bonding behavior, referred to as the lanthanide tetrad effect, can be explained from a quantum mechanics perspective with refined, spin-pairing energy theory (Kawabe et al., 1999). Anomalous redox- and non-redox-controlled bonding behavior results in deviations from the pattern expected based exclusively on charge and radius-smoothed REE patterns when normalized to average shale (Byrne and Sholkovitz, 1996). Lastly, there are differences in burial fluxes of light versus heavy REE in aqueous systems. In marine systems, carbonate complexation of REE results in the light REE having a much higher sorption affinity. Therefore, deviations in REE patterns can be used to track high-temperature, hydrothermal, and low-temperature, aqueous geochemical processes.

REE studies of IFs build on the assumption that there is minimal fractionation of REE during adsorption onto ferric iron oxyhydroxide precipitates; IFs are inferred to trap an REE signature of seawater at the site of ferric iron precipitation. This assumption is based on both experimental studies and results from natural systems, and is likely valid for pH-buffered marine systems. For instance, Mn-poor hydrothermal plume particles essentially record a seawater REE pattern (e.g., Sherrell et al., 1999). REE behavior in modern seawater is relatively well understood and serves as a foundation for interpreting REE patterns of IF. There have been two main objectives in REE studies of IFs: (1) tracing Fe sources, and (2) using the redox-dependent properties of REE to decipher oxidation mechanisms responsible for iron deposition. Europium anomalies have been central in the use of REE to trace Fe sources. Europium enrichment in chemical sedimentary rocks that precipitated from seawater indicates a strong influence of hydrothermal fluids on the seawater-dissolved REE load (Derry and Jacobsen, 1988, 1990; Klinkhammer et al., 1983). The disparate behavior of Eu from neighboring REE in hydrothermal fluids is linked to Eu (III) reduction at high temperatures (>250 °C) and low Eh conditions (Klinkhammer et al., 1983; Sverjensky, 1984). It is generally assumed that Fe and REE will not fractionate during transport from spreading ridges or other exhalative centers, and therefore a large positive Eu anomaly indicates that Fe in the protolith of IF is hydrothermally derived (e.g., Slack et al., 2007). Nd isotope data further support a hydrothermal source of Fe to Precambrian IFs (e.g., Bau and Duslki, 1996; Derry and Jacobsen, 1990; Jacobsen and Pimentel-Klose, 1988).

Secular trends in the magnitude of Eu anomalies in large sediment-hosted IFs have historically been assumed to indicate variations in hydrothermal flux (e.g., Derry and Jacobsen, 1990), possibly linked to thermal history of the mantle. However, without independent constraints it is not possible to exclude a link between a long-term decrease in the magnitude of Eu anomalies to shifts in the continental delivery of REE, potentially related to either crustal growth or crustal emergence above seawater. Large positive Eu anomalies are a common
feature in Phanerozoic and Proterozoic distal hydrothermal sediments (e.g., Peter, 2003; Slack et al., 2007), but positive Eu anomalies in post-Paleoproterozoic sediments are generally assumed to indicate a local hydrothermal flux rather than hydrothermally dominated seawater composition of the global ocean. The lack of large Eu anomalies in IFs (e.g., ~2.22 Ga Hotazel Formation in South Africa; Tsikos and Moore, 1997) or any of the Neoproterozoic IFs associated with the ‘snowball Earth’ glaciations (e.g., Halverson et al., 2011; Klein and Ladeira, 2004) may indicate that during deposition of these IFs the oceans were not greatly influenced by a high-temperature (~250 °C) hydrothermal flux.

REE studies have also focused on redox-controlled, water-column REE behavior in modern anoxic basins. In general, oxygenated marine settings display a strong negative Ce anomaly when normalized to shale composites (Ce(SN)), whereas suboxic and anoxic waters lack large negative Ce anomalies (e.g., Byrne and Sholkovitz, 1996; German and Elderfield, 1990). Oxidation of Ce(III) to Ce(IV) greatly reduces Ce solubility, resulting in its preferential removal onto Mn–Fe oxyhydroxides, organic matter, and clay particles (Byrne and Sholkovitz, 1996). In contrast, suboxic and anoxic waters lack large negative Ce anomalies due to reductive dissolution of settling Mn–Fe-rich particles (Byrne and Sholkovitz, 1996; German et al., 1991). Similarly, light REE depletion and high Y/Ho ratios develop in oxygenated waters due to preferential removal of light versus heavy REE and of Ho relative to Y onto Mn–Fe oxyhydroxides and other particle-reactive surfaces. As a result, the ratio of light to heavy REE markedly increases across redox boundaries due to reductive dissolution of Mn–Fe oxyhydroxides (Byrne and Sholkovitz, 1996; German et al., 1991), whereas dissolved Y/Ho ratios decrease across redox boundaries. In many modern marine basins, the Ce anomaly and light to heavy REE ratio return to values near that of the shale composite across the Mn and Fe redox boundaries. In some basins, even positive Ce anomalies and light REE enrichment develop within anoxic and suboxic waters (e.g., Bau et al., 1997b; de Baar et al., 1988; De Carlo and Green, 2002; Schijf et al., 1995). Redox-induced shifts in REE patterns in some modern stratified basins have been directly linked to Mn-cycling in the suboxic zone (De Carlo and Green 2002; German et al., 1991).

In many Archean and early Paleoproterozoic IFs there is no deviation from trivalent Ce behavior (e.g., Alexander et al., 2008; Bau and Dulski, 1996; Bau and Möller, 1993; Bau et al., 1997a; Frei et al., 2008; Fryer, 1977; Prakash and Devapriyan, 1996), suggesting that the water column from which ferric oxyhydroxides precipitated was reducing with respect to Mn (cf. Bau and Dulski, 1996). In support of this model, in a recent survey, 18 different Paleoproterozoic and Archean IFs lack significant true Ce anomalies until after the GOE (Planavsky et al., 2010a,b; Figure 7(a) and 7(b)). There are several reported cases of Ce anomalies in Archean IFs (e.g., Kato et al., 2006). However, many – if not all – of these cases can be linked to analytical artifacts or analysis of samples that were affected by supergene alteration or weathering (Bekker et al., 2010; Braun et al., 1990; Planavsky et al., 2010a,b; Valeton et al., 1997).

There also appear to be differences in trivalent REE behavior in IFs before and after the rise of atmospheric oxygen. Archean and early Paleoproterozoic IFs are characterized by consistent light REE depletion and high Y/Ho ratios (Planavsky et al., 2010a,b). This pattern contrasts markedly with data for late Paleoproterozoic IFs that show significant ranges in light to heavy REE (Pr/Yb(SN)) and Y/Ho ratios both below and above the shale composite value (Planavsky et al., 2010a,b; Figure 7(c)). This range of light to heavy REE and Y/Ho ratios (Figure 7(c) and 7(d)) in late Paleoproterozoic IFs likely reflects variable fractionation of REE+/Y by Mn- and Fe-oxyhydroxide precipitation and dissolution. Such an interpretation implies deposition of late Paleoproterozoic IFs, at ~1.88 Ga, in basins having varying redox conditions and a strong redoxcline, which separated the upper oxic water column from the deeper-water, suboxic to anoxic waters (Planavsky et al., 2009). A similar Mn redoxcline was likely absent in the Archean oceans. Significant Ce anomalies are lacking in carbonates deposited in shallow-marine settings on Archean carbonate platforms, which is consistent with this model (Planavsky et al., 2010a,b).

Isotope ratios of some REE (e.g., Ce and Nd), in addition to REE concentrations, have been used to constrain REE and Fe sources to seawater and the time when REE systematics was established (Amakawa et al., 1996; Derry and Jacobsen, 1990; Hayashi et al., 2004; Shimizu et al., 1991; Tanaka and Masuda, 1982). Both Ce and Nd have short residence times in the modern ocean, 90–165 and 1000–1500 years, respectively, and heterogeneous isotope compositions (Amakawa et al., 1996). The Archean oceans were also likely strongly heterogeneous having $^{14}Nd$ values of +1 to +2 typical of deep waters dominated by hydrothermal Nd sources, and lower values down to −3 typical of shallow waters dominated by terrestrial Nd sources (Alexander et al., 2009). Similarly, $^{143}Ce$ values in IFs show a strong hydrothermal impact on seawater composition in the Archean (Shimizu et al., 1990, 1991). Further, the La–Ce geochronometry can be a valuable tool to constrain whether negative or positive Ce anomalies reflect seawater composition, diagenesis, or later metamorphic alteration, by dating when REE systematics was established and comparing this age with the independently known depositional age (Hayashi et al., 2004).

9.18.5.1.2 Phosphorus

IFs have the potential to track dissolved phosphate concentrations in ancient oceans (e.g., Bjerrum and Canfield, 2002). It is well established, based on work in modern hydrothermal systems, that phosphate sorption onto iron oxides follows a distribution coefficient relationship; the amount of solid-phase P in iron oxides scales with dissolved phosphate concentrations (e.g., Edmond and German, 2004; Feely et al., 1998). Importantly, however, this does not imply that P is simply adsorbed onto ferric oxides. In fact, there is evidence that P is co-precipitated with a Fe–Ca–P phase (Lilley et al., 1995). During early diagenesis, sediment phosphorus concentrations decrease slightly, but most phosphorus will be retained during Fe oxide recrystallization and secondary apatite precipitation (e.g., Poulton and Canfield, 2006). The process will not significantly vary in different depositional settings, because it is controlled largely by the surface chemistry of iron oxides. This simple framework opens a pathway for estimating the levels of this key nutrient in the ocean through Earth history (Bjerrum and Canfield, 2002).
Iron Formations: Their Origins and Implications for Ancient Seawater Chemistry

A complicating factor of this seemingly simple approach is that distribution coefficients can vary with different solution chemistry, potentially leading to dramatically different P/Fe ratios in IFs. Fortunately, phosphate outcompetes most anions for sorption sites on iron oxides. However, dissolved Si, at high concentrations, will outcompete phosphate for such sites (e.g., Konhauser et al., 2007a,b). Additionally, metal-silica coprecipitation should diminish the resulting particle’s point of zero net charge, rendering it less reactive to dissolved anions (e.g., Konhauser et al., 2007a,b). Therefore, higher dissolved Si levels should yield lower P/Fe ratios in iron oxides, at constant phosphate concentrations. This process is significant because the marine Si cycle and dissolved Si concentrations have changed dramatically through time with the proliferation of enzymatic Si precipitation, foremost with the radiation and evolution of diatoms in the Cretaceous (Maliva et al., 2008; Siever, 1992). More generally, dissolved Si concentrations in seawater have decreased through the Phanerozoic as the Si cycle became increasingly biologically controlled. A decrease in dissolved Si concentrations through time is likely to have affected the abundance of a wide range of elements in iron oxide-rich rocks, not just P concentrations. Interestingly, at low levels, variations in Si concentration appear to have little effect on anion sorption to iron oxides. In the North Atlantic, bottom-water Si is about 40 μM whereas in the Pacific the concentrations are much higher, ~170 μM (e.g., WOCE, 2002). One might expect the slope between dissolved P concentrations and P/Fe ratios to be different between the Atlantic and the Pacific oceans; in the Pacific, the particles should adsorb much less P. However, the trends in both oceans are similar (Edmonds and German, 2004; Feely et al., 1998). In the modern ocean, Si has no obvious influence on P/Fe ratios for the concentration range of 40–170 μM Si. This is not surprising, since at these relatively low Si concentrations, negligible Si becomes incorporated into ferric-oxhydroxide plume particles. Importantly, the range of variation in modern

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Figure 7  Secular trends in REE + Y characteristics in a set of 18 Archean and Paleoproterozoic iron formations. (a) Plot of CeSN and PrSN anomalies for a set of late Paleoproterozoic (●) and Archean and early Paleoproterozoic (□) iron formations. Positive Ce anomalies are only present in late Paleoproterozoic iron formations, whereas all iron formations shown lack true negative Ce anomalies. True negative Ce anomalies are defined by Ce/ Ce*SN (CeSN/0.5(PrSN + LaSN)) and Pr/Pr* SN (PrSN/(0.5CeSN + 0.5NdSN)) values above and below unity, respectively. This approach, first described by Bau and Dulski (1996), discriminates between positive La and true negative Ce anomalies. (b) Ce anomalies (CeSN/(0.5(PrSN + LaSN))). (c) Light to heavy REE ratios (Pr/YbSN). (d) Y/Ho ratios. Black bar indicates PAAS shale composite values. Archean iron formations are characterized by higher Y/Ho ratios and lower light to heavy REE ratios than those of the shale composite; late Paleoproterozoic iron formations have a large range of Y/Ho ratios and light to heavy REE ratios. Observed secular trend in REE + Y characteristics is best explained by the absence of an Fe-Mn chemocline in the Archean oceans prior to the rise of atmospheric oxygen (see text for details). Modified from Planavsky NJ, Bekker A, Rouxel OJ, Knudsen A, and Lyons TW (2010) Rare earth element and yttrium compositions of Archean and Paleoproterozoic iron formations revisited: New perspectives on the significance and mechanisms of deposition. Geochimica et Cosmochimica Acta 74: 6387–6405; Planavsky NJ, Rouxel OJ, Bekker A, et al. (2010). The evolution of the marine phosphate reservoir. Nature 467: 1088–1090.
dissolved Si concentrations is small compared to changes in seawater Si concentrations envisaged during biomineralization evolutionary events in the late Precambrian and Phanerozoic (100s–1000s of micromolar Si shifts; Maliva et al., 2005; Siever, 1992). However, this potential discrepancy warrants additional experimental work on phosphate sorption onto iron oxides at lower silica concentrations. It is possible that a threshold value exists at which the inhibitory effect becomes pronounced.

Additionally, iron-rich particles in modern, neutrally buoyant plumes and rising plumes contain P-rich organic matter and Fe as sulfides (Edmonds and German, 2004; Feely et al., 1998). The P/Fe ratios of hydrothermal plume particles are likely to be affected by these organic P and Fe(II) phases (even though the Fe(II) phases are oxidized to Fe(III) phases relatively rapidly). However, these effects are small enough that a single global phosphorus–Fe-oxide $K_d$ value can be delineated from modern hydrothermal particles (Feely et al., 1998). In summary, P/Fe ratios in modern hydrothermal plume particles are primarily controlled by the phosphate concentration in coeval seawater.

When viewed in light of the evolution of the Si cycle, P/Fe ratios in IFs offer a perspective on dissolved P concentrations in the oceans through time (e.g., Planavsky et al., 2010a,b; Figure 8). It appears that a relatively narrow range of variation existed in dissolved phosphate concentrations throughout the Phanerozoic, consistent with models of global biogeochemical cycles (e.g., Arvidson et al., 2006). In contrast, during the Precambrian, phosphate concentrations may have been significantly elevated relative to those of modern oceans. Foremost, Neoproterozoic IFs have very high P/Fe ratios, despite the high dissolved Si concentrations at that time, which would have inhibited phosphate sorption onto iron oxides. Similarly, in Archean and Paleoproterozoic IFs, P/Fe ratios are suggestive of elevated marine phosphate concentrations. This hypothesis is a deviation from the prevailing view of the early Precambrian phosphorus cycle (Bjerrum and Canfield, 2002). A large amount of phosphorus is currently removed from seawater during the oxic alteration of seafloor basalts by absorption of P onto basalt-derived iron oxyhydroxide particles (see Chapter 10.13). In largely anoxic early Precambrian oceans, however, this phosphorus flux would have been shut off, which likely partly

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**Figure 8** (a) P/Fe molar ratios through time in iron oxide-rich distal hydrothermal sediments and iron formations with low amounts of siliciclastic input. Open squares are individual samples. The P/Fe ratio reflects the size of the marine phosphate reservoir; phosphate sorption onto ferric oxyhydroxides follows a distribution coefficient relationship. The ratio is also influenced by the concentration of dissolved silica, because phosphate and silica hydroxides compete for sorption sites on ferric oxyhydroxides. (b) Phosphate concentrations were extrapolated from average P/Fe ratios for individual formations at dissolved seawater silica concentration of 0.67 mM (cristobalite saturation) and 2.2 mM (amorphous silica saturation). The compilation of P/Fe data suggests that there were elevated seawater phosphate concentrations in the Precambrian and a peak in phosphate level associated with the Neoproterozoic snowball Earth glaciations. This late Precambrian increase in dissolved phosphorus concentration may have stimulated high rates of organic carbon burial and a corresponding increase in atmospheric oxygen levels – paving the way for the metazoan diversification. Modified from Planavsky NJ, Rouxel OJ, Bekker A, et al. (2010). The evolution of the marine phosphate reservoir. *Nature* 467: 1088–1090.
explains elevated Precambrian phosphate concentrations in seawater. If this interpretation is quantitatively important, it provides an underappreciated feedback that may help constrain ocean productivity and, indirectly, surface oxidation from overly high levels. The Neoproterozoic IFs are closely tied temporally to snowball Earth glaciations, suggesting that glaciations or deglaciations could have played a complementary role to the high P levels inferred for Neoproterozoic seawater (cf. Hoffman and Schrag, 2002; Planavsky et al., 2010a,b).

9.18.5.1.3 Nickel
Nickel concentrations similar to those of phosphate follow a distribution coefficient relationship during precipitation of ferric oxyhydroxides. Therefore, the amount of Ni in oxide-facies IFs with limited contribution of detrital materials can be used to track first-order trends in dissolved Ni concentrations in seawater (Konhauser et al., 2009). Ni is also of special interest because it is a bioessential nutrient that in the modern ocean follows a nutrient-type profile. The nickel content (expressed as molar Ni/Fe) in IFs has changed dramatically over time, beginning with a drop in Ni availability in the oceans at \( /C242.7 \text{Ga} \) (Konhauser et al., 2009; Figure 9). Iron-normalized Ni concentrations in \( /C243.8–2.7 \text{Ga} \) IFs are greater than 0.0004 M, were about half that value between 2.7 and 2.5 Ga, and subsequently slowly approached modern values (<0.0001) by 0.55 Ga. This drop in seawater Ni availability would have had profound consequences for microorganisms that depended on it, particularly methane-producing bacteria – methanogens (Konhauser et al., 2009). These bacteria have a unique Ni requirement for their methane-producing enzymes, and a deficiency in this metal could have decreased their population. Crucially, these bacteria have been implicated in controlling oxygen levels on ancient Earth because the methane they produced was reactive with oxygen and kept atmospheric oxygen levels low (Zahnle et al., 2006). Methanogenic bacteria preferentially take up light Ni isotopes, driving residual Ni in seawater to positive Ni isotope values. This isotopic shift has been interpreted as a unique biological fractionation mechanism specific to methanotrophs (Cameron et al., 2009). However, recent studies by Gueguen et al. (in press) show that nonbiological and even high-temperature magmatic processes equally fractionate Ni isotopes, thus challenging the original interpretation of large biological fractionations in Ni isotopes.

9.18.5.1.4 Chromium
The enrichment of the redox-sensitive trace metal Cr can be used to track the terrestrial Cr flux to the oceans (Frei et al., 2009; Konhauser et al., 2011). Data for nearshore IFs cannot be used to infer bulk seawater composition, but they do offer insights into the supply and composition of continental drainage waters, and, by extension, the physical and chemical weathering processes on land at the time of their deposition. In a recent compilation of Cr concentrations in IFs (Figure 10(a) and 10(b)), Konhauser et al. (2011) argued that Cr was largely immobile on land until \( /C242.5 \text{Ga} \). After that time, Cr enrichments started to increase in shallow-water IFs and peaked essentially synchronous with the permanent loss at \( /C242.32 \text{Ga} \) of mass-independent fractionation of sulfur isotopes that defines the GOE (Bekker et al., 2004; Guo et al., 2009). This indicates that Cr was solubilized at a scale unrivaled in Earth history, yet muted Cr isotope fractionations at that time (Frei et al., 2009; see Figure 10(c)) argue against extensive oxidative Cr(VI) transport during the GOE. Instead, Cr must have been mobilized predominantly in reduced Cr(III) form and supplied to the oceans in solution, or adsorbed onto authigenic weathering products. Crucially, it is likely that

![Figure 9](https://example.com/figure9.png)

**Figure 9** Ni/Fe mole ratios for iron formations versus age. Ni/Fe ratios are proposed to track marine Ni reservoir, suggesting a significant decline in marine Ni concentrations after \( /C242.7 \text{Ga} \). The figure contains 1214 measurements, including literature data (circles), bulk (squares) and grain-by-grain laser ablation analyses (crosses). Modified from Konhauser KO, Pecoits E, Lalonde SV, et al. (2009) Oceanic nickel depletion and a methanogen famine before the Great Oxidation Event. *Nature* 458: 750–753.
only microbially catalyzed oxidation of crustal pyrite could have generated the degree of acidity required for appreciable Cr(III) solubilization (Rai et al., 1989). Today, aerobically respiring bacteria are essential to this process, catalyzing the continued oxidation of Fe(II) as pH values drop below the threshold for inorganic Fe(II) oxidation. Based on these constraints, it was suggested that the Cr(III) pulse beginning at ~2.48 Ga and peaking at ~2.32 Ga indicates that such bacteria began utilizing O₂ for the first time to oxidize a previously stable and abundant crustal pyrite reservoir (Konhauser et al., 2011). Sulfuric acid generated by this metabolism ultimately leached Cr from ultramafic source rocks and residual soils. This profound shift in weathering regimes constitutes Earth’s first acid continental drainage system (cf. Bekker and Holland,

Figure 10 Chromium in iron formations and ironstones through time. (a–c) Show secular trends in Cr concentrations (a), authigenic enrichment in Cr relative to Ti (b), and Cr isotope values replotted from Frei et al., 2009 (c). Increasing enrichments in Cr in iron formations near the Archean–Proterozoic boundary suggest a change in Cr cycling. However, muted Cr isotope fractionations during this time suggest transport of reduced rather than oxidized Cr to marine environments. In (a) and (b), squares denote bulk analyses, circles are laser-ablation analyses, black represents volcanic-hosted iron formations, red denotes sediment-hosted iron formations, blue indicates Proterozoic oolitic iron formations and Phanerozoic ironstones that formed in shallow marine waters, and green represents Phanerozoic hydrothermal and exhalative deposits. In (b), Cr/Ti ratios have been normalized to the evolving Cr/Ti ratio of upper continental crust (solid line) according to the restoration model of Condie (1993). In (c), solid and dashed lines represent the mean and 2σ values, respectively, for Archaean and Paleoproterozoic Cr isotope compositions from Frei et al. (2009). Modified from Konhauser KO, Lalonde S, Planavsky NJ, et al. (2011) Aerobic bacterial pyrite oxidation and acid rock drainage during the Great Oxidation Event. Nature 478: 369–373, with additional data from Rye and Holland (2000).
1992). Numerous siderite-rich IFs also have been analyzed in the United States and Canada (Perry et al., 1973; Winter and Knauth, 1992). Carbon isotopes have long been used as a tool to understand the genesis of IFs, fundamental constraints on Precambrian paleoenvironments, and the evolution of life (e.g., Baur et al., 1985; Beaumont and Roberts, 1999; Becker and Clayton, 1972; Goodwin et al., 1976; Hren et al., 2009; Shen et al., 2006; Thode and Goodwin, 1983; Walker, 1984). Following analytical advances in stable isotope geochemistry, new isotope tracers are now available, including for the two major elements in IFs, iron and silicon, as well as for trace elements such as chromium, nickel, molybdenum, germanium, and uranium. Although still in its infancy, the growing field of non-traditional stable isotope geochemistry will certainly open new avenues for investigations of IFs.

### 9.18.5.2 Stable Isotope Studies of IF

Light stable isotopes of oxygen and carbon and, to a lesser extent, hydrogen, sulfur, and nitrogen have been used widely to understand the genesis of IFs, fundamental constraints on Precambrian paleoenvironments, and the evolution of life (e.g., Baur et al., 1985; Beaumont and Roberts, 1999; Becker and Clayton, 1972; Goodwin et al., 1976; Hren et al., 2009; Shen et al., 2006; Thode and Goodwin, 1983; Walker, 1984). Following analytical advances in stable isotope geochemistry, new isotope tracers are now available, including for the two major elements in IFs, iron and silicon, as well as for trace elements such as chromium, nickel, molybdenum, germanium, and uranium. Although still in its infancy, the growing field of non-traditional stable isotope geochemistry will certainly open new avenues for investigations of IFs.

#### 9.18.5.2.1 Traditional light stable isotopes

Carbon isotopes have long been used as a tool to understand the genesis of IFs. Most carbon isotope studies have focused on the carbonate fraction, due in part to the generally low organic carbon content in IFs. The most extensive studies have been undertaken on the low metamorphic grade deposits of the late Neoarchean to early Paleoproterozoic Transvaal Supergroup in South Africa (e.g., Beukes and Klein, 1990; Fischer et al., 2009) and the ~2.5 Ga Brockman IF in Western Australia (e.g., Baur et al., 1985; Becker and Clayton, 1972; Kaufman et al., 1990), as well as the ~1.88 Ga Biwabik and Gunflint IFs in the United States and Canada (Perry et al., 1973; Winter and Knauth, 1992). Numerous siderite-rich IFs also have been analyzed for carbonate carbon isotopes (e.g., Ohmoto et al., 2004). Based on a recent compilation by Johnson et al. (2008a), carbonates associated with IFs tend to be isotopically light, with δ13C values ranging from −20.0 to +2.4‰ against Vienna Pee Dee Belemnite (V-PDB) and the majority of values clustered around −8 to −6‰ (Heimann et al., 2010). Organic carbon isotopes also are δ13C-depleted, with values as low as −41.4‰. Studies of carbonates from the Brockman IF (Baur et al., 1985) show that isotopically light carbon and oxygen isotope values correlate with concentrations of iron. The negative carbonate carbon isotope values have been interpreted as evidence for direct carbonate (siderite) precipitation from an iron-rich water column, stratified with respect to carbon isotope composition of total dissolved inorganic carbon and influenced by a deep-water, hydrothermal flux associated with submarine magmatic activity (e.g., Beukes and Klein, 1990; Winter and Knauth, 1992). Although a stratification of several per mil in the carbon isotope composition of dissolved inorganic carbon is present in the modern ocean (e.g., Kroopnick, 1985), a much smaller gradient is preserved in the early Precambrian rock record and is expected under the high pH12 conditions required to compensate for a lower solar luminosity during early Earth history (Hotinski et al., 2004). As another caveat to those interpretations, petrographic evidence (e.g., Ayres, 1972) indicates that almost all iron-rich carbonates in IFs formed during burial diagenesis and, therefore, are unlikely to have precipitated within the water column (see, e.g., how iron-rich carbonate in Figures 2(e), 2(g), and 3(e) occludes laminae). Consistent with these petrographic observations, some Fe-rich carbonates, in particular those in oxide and silicate facies IFs, tend to have more negative oxygen isotope values than coeval calcite and dolomite that precipitated from seawater (Heimann et al., 2010), suggesting that the Fe-rich carbonates formed during late diagenesis in sediments, possibly at several kilometers depth. Formation of δ13C-depleted Fe-rich carbonates also is commonly linked with a fermentative metabolism and anaerobic respiration in the anoxic water column and sediments. This interpretation was first suggested by Perry et al. (1973) and, later, by Walker (1984), who proposed that the markedly light carbon isotope values of siderite in IF reflect diagenetic precipitation of ferruginous carbonate linked to organic matter remineralization, with ferric oxides being the terminal electron acceptor. Heimann et al. (2010) placed isotopic mass balance constraints on this model. The case of dissipatory iron reduction (DIR) and complete C and Fe retention in sediments is described by the following equation, in which CH₂O represents the total organic carbon:

\[
4\text{Fe(OH)}_3 + \text{CH}_2\text{O} + 3\text{HCO}_3^- \rightarrow 4\text{FeCO}_3 + 3\text{OH}^- + 7\text{H}_2\text{O}
\]

This equation implies that three out of four carbon atoms in siderite were derived from seawater bicarbonate present in pore waters and the water column. At a lower contribution from seawater bicarbonate (e.g., at a greater depth in sediments and at a more pronounced isolation from the water column), some Fe would be lost from sediments to pore waters during DIR. A higher contribution of seawater bicarbonate would correspond to the case where the upward-diffusing iron in pore waters is bound at a higher level in sediments or at the water–sediment interface with seawater bicarbonate. Using average carbon isotope values for bicarbonate and organic carbon (0 and −30‰, respectively), the above equation requires the average carbon isotope value of siderite to be close to −7.5‰, if all iron reduced by DIR was retained in the sediments. Interestingly, as noted above, the carbon isotope values of siderite in IFs cluster close to this value, although some bias toward less-negative values is likely due to oversampling of massive beds formed at or just below the sediment–water interface. In this rather simplistic view of early diagenesis, IF carbon isotope data suggest that a significant amount, if not all, of the iron released by DIR was retained in the sediments and did not escape back into the water column.

Another plausible model links the light carbon isotope ratios in iron carbonates of IFs to iron-based methane oxidation. Recently, this process has been documented at modern marine methane seeps (Beal et al., 2009) and in an iron-rich lake (Crowe et al., 2011). These findings build from thermodynamic predictions, which suggest that bacteria are capable of linking methane oxidation to ferric iron reduction. Methano-genesis was likely a common metabolic pathway in organic matter-rich shales within iron formation-bearing sequences, and possibly in precursor sediments for Archean and Paleoproterozoic IFs (e.g., Konhauser et al., 2005). In this scenario, methane produced in sediments during early diagenesis diffused upwards and was biologically oxidized with ferric oxides,
producing both ferrous iron and bicarbonate that would then coprecipitate as iron-rich carbonate. The presence of markedly negative carbonate carbon isotope values as low as ~−20% relative to PDB is consistent with methane cycling having mediated carbonate precipitation, given that methane is much more isotopically depleted in 13C than is typical organic matter. This process could also create organic matter having extremely depleted δ13C values without aerobic involvement.

The oxygen isotope composition of chert in IFs, particularly coupled to other isotope proxies such as Si, H, and Fe, has been used to address issues such as Precambrian ocean temperatures (Hren et al., 2009; Knauth, 2005; Knauth and Lowe, 2003; Robert and Chaussidon, 2006) and seawater oxygen isotope compositions (Perry, 1967), the source of silica to IFs (Robert and Chaussidon, 2006; Steinhoefel et al., 2009; van den Boorn et al., 2007), and the degree of metamorphic overprint (Valaas Hyslop et al., 2008). Assuming that oxygen isotope values of Precambrian oceans were similar to those of ice-free recent oceans (Knauth and Lowe, 2003; Muehlenbachs, 1998) and that postdepositional isotope exchange was minimal, the bulk oxygen isotope composition of chert might be a useful paleothermometer because isotope fractionation between silica and seawater is dependent on temperature. Under this assumption, low oxygen isotope values of ~<+22‰ Standard Mean Ocean Water (SMOW) for 3.4–3.2 Ga cherts were interpreted as evidence for hot early oceans, implying seawater temperatures of 55–85 °C (Knauth and Lowe, 2003). This was later supported by the in situ ion microprobe study of cherts by Robert and Chaussidon (2006), revealing covariation between the silica and oxygen isotope records. Subsequent studies applying similar techniques and the modeling of diageneis and metamorphism established that some of the ~3.5–3.3 Ga Onverwacht Group cherts in South Africa were completely equilibrated with postdepositional gold-bearing fluids and, therefore, had lost their original seawater signal (Marin-Carbonne et al., 2011). Other studies have shown that chert in the 1.88 Ga Gunflint Formation of Canada precipitated at much lower (37–52 °C) temperatures (Marin-Carbonne et al., 2012). Similarly, a study of isotope compositions of the ~3.42 Ga Buck Reef Chert in South Africa (Hren et al., 2009) questioned previously inferred temperature constraints of Knauth and Lowe (2003). Hren et al. (2009) explored the temperature dependence of oxygen and hydrogen isotope fractionations in order to calculate δ18O values of ambient fluids during transformation of amorphous silica to microcrystalline quartz, finding that the Paleoarchean ocean was isotopically depleted in 18O relative to the modern ocean and thus was far cooler (<40 °C) than previously envisaged. In using these and similar arguments, the assumption has generally been made that chert represents a marine chemical precipitate, and that its isotopic composition directly reflects the composition and temperature of the Archean ocean at the time of deposition. This is, however, unlikely for Paleoarchean cherts within volcano-sedimentary sequences of greenstone belts, because their origin is closely linked to syndepositional, low-temperature hydrothermal processes on the seafloor (Hofmann and Bolhar, 2007; Hofmann and Harris, 2008). Temperature estimates obtained from stable isotope paleothermometry, therefore, reflect the temperature of chert precipitation as a result of mixing of hydrothermal fluids with cold seawater. The diagenetic origin of bedded cherts in BIF of the Hamersley Group (Krapež et al., 2003) also challenges the meaning of these isotope data.

Sulfur isotope compositions of organic matter-rich and sulfidic shales interbedded with Neoarchean and Paleoproterozoic IFs have been extensively studied to constrain biological sulfur cycling, sources of sulfur, and ocean redox structure (e.g., Cameron, 1983; Goodwin et al., 1976; Grassineau et al., 2001; Thode and Goodwin, 1983). On the other hand, super-heavy S isotope values in Neoproterozoic postglacial lithologies, including Fe and Mn formations, were linked to Rayleigh fractionation during pyrite formation under low seawater sulfate conditions (Liu et al., 2006). The results were used by Habicht et al. (2002) to infer a low (<200 µM) seawater sulfate content in the Archean ocean. An increase in the range of sulfur isotope values in ~2.7 Ga black shales interbedded with IFs has been interpreted as an expression of dissimilatory bacterial sulfate reduction or redox cycling in stratified Archean oceans (Goodwin et al., 1976; Grassineau et al., 2001). Studies of multiple sulfur isotopes have shown that the range of δ34S values alone in sediments deposited before the GOE cannot be used to support either of these two interpretations, because photochemical processes in an anoxic atmosphere significantly fractionate sulfur isotopes (Farquhar et al., 2000). Indeed, the same 2.7 Ga sedimentary units contain a large range of mass-independent fractionation in sulfur isotopes, thus arguing for the role of photochemical processes and against oxygenated surface environments. Although biological sulfur cycling was likely present in Archean oceans (e.g., Philippot et al., 2007; Shen et al., 2001, 2009), it is difficult to constrain its role using sulfur isotopes alone. Multiple sulfur isotope analyses of sulfur hosted in IFs also show a large range of mass-independent fractionations (Farquhar and Wing, 2005; Kaufman et al., 2007; Partridge et al., 2008), suggesting that small amounts of sulfur compounds derived from photochemically fractionated sulfur species co-precipitated with IFs.

Very little data are available for nitrogen isotope composition of IFs, mainly due to their low nitrogen contents. However, ammonium might substitute for potassium ion in the primary clay minerals (cf. Williams and Ferrell, 1991) within IFs, such as in the sedimentary precursor to stilpnomelane, making IFs at low metamorphic grade a prospective lithology for nitrogen isotope studies. From the N isotope data presently available for organic matter-rich shales, cherts, and IFs it appears as though a bimodal secular pattern exists with a change occurring across the Neoarchean–Paleoproterozoic boundary, interpreted as reflecting progressive oxidation of surface environments (Beaumont and Robert, 1999; Garvin et al., 2009; Godfrey and Falkowski, 2009; Shen et al., 2006; Thomazo et al., 2011). The older record is marked by a large range of negative and highly positive values, which are generally related to bacterial nitrogen fixation as a principal pathway for biogeochemical nitrogen cycling in anoxic oceans, where nitrification with oxygen as an electron acceptor was limited or absent (Farquhar et al., 2011). A shift to predominantly positive nitrogen isotope values in Neoarchean shales is related to an emergence of nitrification and denitrification (a process that requires dissolved nitrate), coeval with the appearance of dissolved oxygen in at least the upper part of the ocean water column (Garvin et al., 2009; Godfrey and Falkowski, 2009; Thomazo et al., 2011). Interestingly, 2.7 and 2.5 Ga
IFs, black shales, and cherts also contain nitrogen with highly positive isotope values (Beaumont and Robert, 1999; Jia and Kerrich, 2004; Shen et al., 2006). Considering that these lithologies were deposited in relatively deep-water (below storm wave-base) anoxic environments, it is highly unlikely that any nitrite or nitrate was exported into these environments from local oxygenated settings, where they could have gained positive nitrogen isotope values due to Rayleigh distillation associated with denitrification. On the other hand, it seems plausible that oxidized nitrogen species could have been produced during early diagenesis when organic matter and ammonium were biologically remineralized, with ferric iron acting as electron acceptors in the absence of oxygen. Such a process could lead to the formation of isotopically distinct nitrogen pools. There is tentative evidence for microbial ammonium oxidation to nitrite coupled to Fe(III) reduction in laboratory experiments (Sawayama, 2006) and under anaerobic conditions in wetland soils (Clement et al., 2005). Although further documentation of this process is needed, it is a thermodynamically feasible metabolic pathway, and it is likely common in modern Fe-rich systems.

9.18.5.2.2 Nontraditional stable isotopes

9.18.5.2.2.1 Fe isotopes

A number of recent iron isotope studies of IFs have been made with the aim of tracking the biogeochemical cycling of iron on early Earth (Beard et al., 1999; Craddock and Dauphas, 2011; Dauphas et al., 2004; Heimann et al., 2010; Johnson et al., 2003, 2008a,b; Planavsky et al., 2009; Rouxel et al., 2005; Steinhoefel et al., 2009). Importantly, this recent iron isotope work has bolstered evidence for a rain of ferric oxyhydroxides during IF deposition, for early diagenetic microbial iron cycling, and for a hydrothermal iron source for IFs (e.g., Anbar and Rouxel, 2007; Dauphas and Rouxel, 2006; Heimann et al., 2010; Planavsky et al., 2012). Iron isotopes are typically reported as $\delta^{56}$Fe values with a range of 5% in nature. Several recent reviews provide detailed information on iron isotope systematics and fractionations (Anbar and Rouxel, 2007; Beard et al., 2003; Dauphas and Rouxel, 2006; Johnson et al., 2008b). The application of iron isotopes to understanding the ancient iron cycle is based on a foundation of extensive experimental work that determined kinetic and equilibrium fractionation factors during redox reactions and mineral precipitation (Beard et al., 1999, 2010; Bullen et al., 2001; Butler et al., 2005; Croal et al., 2004; Crosby et al., 2005; Guibaud et al., 2011; Welch et al., 2003; Wiesli et al., 2004). In general, the largest iron isotope fractionations occur during redox reactions. For instance, during Fe(III) reduction and Fe(II) oxidation there is $\sim 1.5\%$ fractionation. The notable exception is during sulfide formation, which might be accompanied by a large ($>2\%$) fractionation (Guibaud et al., 2011; Rouxel et al., 2008a,b); since IFs typically lack sulfides, this pathway for isotopic fractionation is not particularly relevant. Additionally, a sound understanding exists of the isotopic composition of iron from different sources (Dauphas and Rouxel, 2006; Johnson et al., 2008a). Most importantly, iron derived from hydrothermal sources has slightly negative or near-zero iron isotope values (e.g., Rouxel et al., 2008a,b), whereas a benthic flux (dissolved iron derived from iron reduction during early sediment diagenesis on continental shelves and supplied to deep euxinic part of the basin) is likely to have a pronounced negative iron isotope value (e.g., Severmann et al., 2008). This isotopically light benthic Fe source has also been found

Figure 11  Secular variations in $\delta^{56}$Fe values for Precambrian iron formations. Data for magnetite and/or hematite (open diamonds) include bulk mineral and in situ (using laser-ablation MC-ICP-MS and ion-microprobe) analyses. Data for Fe-rich carbonates (gray squares) and hematite hosted in Mn-carbonates (Hotazel Formation) correspond to bulk mineral analysis. A total of 490 datapoints is shown including 83 datapoints for the $\sim 1.88$ Ga GIF from the Animikie basin, North America (Frost et al., 2007; Planavsky et al., 2009; Valaas Hyslop et al., 2008), 138 datapoints for the $\sim 2.5$ Ga Brockman Iron Formation, Western Australia (Johnson et al., 2008a,b), 58 datapoints for the $\sim 2.5$ Ga Transvaal Supergroup IF, South Africa (Johnson et al., 2003), 27 datapoints for the $\sim 2.7$ Ga Shurugwi and Belingwe greenstone belts IF, Zimbabwe (Rouxel et al., 2005; Steinhoefel et al., 2009), 137 datapoints for the Eoarchean Isua, Akilia, and Innersuartuut IF and metamorphic rocks (Dauphas et al., 2004, 2007; Whitehouse and Fedo, 2007), 12 datapoints for the Neoproterozoic Rapitan IF (Halverson et al., 2011), 52 datapoints for IF and Mn formation from the Late proterozoic Hotazel Formation, Transvaal Supergroup, South Africa (Tsikos et al., 2010), and 45 datapoints for other Late proterozoic and Archean IFs (Planavsky et al., 2012). Gray horizontal bar represents the average $\delta^{56}$Fe values for igneous rocks and hydrothermal sources (e.g., Beard et al., 2003; Rouxel et al., 2008a,b). For comparison, iron isotope values of jaspers and iron formations from Phanerozoic hydrothermal deposits are shown (from Planavsky et al., 2012); note that stromatolitic and oolitic-ooloidal ironstone of the Middle Eocene age from Egypt has $\delta^{56}$Fe values ranging from $+1.14$ to $+2.28\%$ (Salama et al., 2011).
to be transferred to the water column, notably in coastal settings (Rouzel and Auro, 2010).

Bulk samples of IFs commonly contain positive or near-crustal δ56Fe values (Cradock and Dauphas, 2011; Johnson et al., 2003, 2008a; Planavsky et al., 2012; Rouzel et al., 2005; Steinthoefel et al., 2009; Figure 11), which provide insight into iron-enrichment mechanisms. The two most commonly proposed iron sources for IFs, hydrothermal (Bau et al., 1997a,b; Bak et al., 2008), have negative (subcrustal) δ56Fe values. Hence, the presence of positive δ56Fe values in IF must reflect a fractionation during mineral precipitation. More specifically, positive δ56Fe values point toward a rain of ferric oxyhydroxides to the sediment–water interface (e.g., Dauphas et al., 2004). Ferric oxyhydroxides are fractionated by ~1.5% relative to dissolved ferrous iron (see Johnson et al., 2008b for review). This enrichment in heavy Fe isotopes contrasts with the isotope fractionations associated with siderite, ankerite, and green rust precipitation, which are heavy iron isotope-depleted relative to the ambient Fe(II)aq pool (Wiesli et al., 2004): the exact fractionation during precipitation of iron silicates is unknown. In general, positive iron isotope values in IFs indicate that Fe(III) delivery was the main process driving iron enrichments. Additionally, the expression of large Fe isotope fractionations implies partial Fe(II) oxidation, pointing toward oxidation at low Eh conditions (Planavsky et al., 2012). If Fe(II) oxidation took place during mixing of anoxic Fe(II)-rich and oxygenated marine waters, as was commonly envisaged in the past (Cloud, 1973; Holland, 1984), oxidation would have been essentially quantitative given the rapid oxidation kinetics of iron at neutral pH (e.g., Stumm and Morgan, 1995). This rapid and quantitative oxidation would have prevented any significant expression of iron isotope fractionations, which is the case with modern hydrothermal plume fallout (Severmann et al., 2004). Therefore, the presence of large iron isotope fractionations argues against oxidation having been quantitative.

The presence of positive δ56Fe values in IFs dominated by reduced or mixed-valence iron minerals may also provide strong evidence for microbial Fe(III) reduction in precursor sediments to the IFs (Cradock and Dauphas, 2011; Johnson et al., 2008a,b). For example, siderite with positive δ56Fe values and siderite with similar δ56Fe values as coexisting iron oxides (Cradock and Dauphas, 2011) indicate a lack of isotopic equilibrium between siderite and aqueous Fe(II) in ancient anoxic oceans (Johnson et al., 2008a). Siderite that precipitated from seawater should have a negative δ56Fe value, given the isotopic fractionation during siderite precipitation and expected iron isotope values for seawater. Therefore, siderite with positive δ56Fe values must have been derived by reductive dissolution of iron oxyhydroxides rather than having precipitated directly from seawater. In most cases, microbial Fe(III) reduction can be assumed to be driving the reductive oxide dissolution. This also implies quantitative reduction of iron oxide precursors, which raises an important question regarding the nature of the electron donors in organic matter-starved systems of IF. Although iron isotope data for IFs may point towards microbial Fe(III) reduction in the rock record, possibly dating back to the earliest known sedimentary rocks at ~3.8 Ga (Cradock and Dauphas, 2011), independent evidence is needed to exclude the possibility of siderite formation during metamorphic reaction of iron oxides and organic carbon, before positive δ56Fe values in siderite can be linked with microbial Fe(III) reduction. As stressed above, petrographic evidence for siderite precipitation from seawater is generally lacking.

Positive δ56Fe values in IFs are also consistent with a hydrothermal iron source. The common occurrence in bulk IF samples of δ56Fe values between 0.5 and 1.0‰ (e.g., Planavsky et al., 2012) is consistent with partial oxidation of hydrothermal Fe(II), because hydrothermally sourced iron should have an initial δ56Fe value between −0.5 and 0‰. These positive δ56Fe values are less likely to be linked to partial oxidation of benthic and diagenetically derived iron, since the latter fluxes are typically characterized by quite negative δ56Fe values (<−1.5‰). Iron isotopes do not provide definitive evidence for a hydrothermal iron source, but support this model, which is also strengthened by many other lines of geological and geochemical evidence (e.g., Bekker et al., 2010).

Iron having near-crustal or negative δ56Fe values in IFs could have been sourced from either a hydrothermal or a benthic source. The degree of water-column Fe(II) oxidation is expected to vary widely in Archean and Paleoproterozoic oceans, potentially accounting for a range of iron isotope values in the particulate iron oxyhydroxide flux for a given dissolved iron isotope value. Additionally, it is likely that when reducing marine conditions prevailed, as indicated for the Archean and Paleoproterozoic oceans by REE studies (e.g., Bau and Dulski, 1996; Planavsky et al., 2010a,b), partial Fe(II) oxidation could have resulted in a dissolved iron isotope gradient. Precipitation of isotopically heavy iron oxides would leave behind an isotopically light dissolved iron pool, which also could have been transferred to the IF rock record with additional oxidation (Rouzel et al., 2005). The co-occurrence of isotopically light carbonate carbon and iron isotope values has been proposed as evidence for the reductive origin of light iron isotopes in IFs, possibly with iron being sourced from within the sediment pile (Cradock and Dauphas, 2011; Heimann et al., 2010; Johnson et al., 2008a). This model builds from the earlier interpretations outlined above that light carbonate carbon isotope values may be derived by remineralization of organic matter (e.g., Baur et al., 1985; Becker and Clayton, 1972; Kaufman et al., 1990). The light iron isotope values similarly may record partial microbial reduction of iron oxyhydroxides during early diagenesis (Johnson et al., 2008a). However, negative carbonate iron isotope values can result from direct precipitation of seawater-derived Fe(II), whereas the light carbonate carbon isotope values are linked to organic matter remineralization. Stated more simply, iron isotopes can be controlled by the flux of iron to the sediment–water interface, whereas carbon isotopes provide a signal of early diagenetic microbial processes. IF carbonates having positive δ56Fe values also show markedly negative carbonate carbon isotope values (Cradock and Dauphas, 2011), consistent with the above-outlined model.

There has been extensive discussion on negative iron isotope values in IFs. Several conflicting interpretations exist for these values, such that independent geological and geochemical evidence should be used to discriminate these models (Johnson et al., 2008a; Tsikos et al., 2010). Further, light iron isotope values are not a silver bullet for DIR even if coupled with light carbonate carbon isotope values.
A compilation of bulk rock and mineral-specific $\delta^{56}$Fe values for Archean and Paleoproterozoic IFs (Figure 11) shows an overall range between $-2.5$ and $2.7\%_\text{o}$, which encomasses most of the natural variations in iron isotopes observed to date. Although these data reveal that many Archean and early Paleoproterozoic IFs were a sink for isotopically heavy iron, in contrast to later Proterozoic and Phanerozoic iron oxide-rich rocks, further work is needed to explore whether a secular change exists in the iron isotope composition of IF prior to the GOE. Notably, markedly low $\delta^{56}$Fe values are common in the $\sim 2.2$ Ga Hotazel Formation of South Africa and especially in manganese-rich samples. This distinctive feature may reflect the deposition of iron and manganese from hydrothermal fluid depleted in heavy iron isotopes, by progressive Fe(II) oxidation and precipitation in the deeper part of a redox-stratified basin that had a redox state intermediate between that required for iron and manganese oxidation (Tsikos et al., 2010).

The average iron isotope composition of different types of IF is a major unresolved question. In some cases, IFs have been estimated to have a near-crustal average $\delta^{56}$Fe value (Johnson et al., 2008a). However, as discussed above, many IFs display positive $\delta^{56}$Fe values and thus have been interpreted as a significant sink of isotopically heavy iron (Planavsky et al., 2012; Rouxel et al., 2005). Unfortunately, previous iron isotope studies were not ideally designed to estimate the average iron isotope composition of IFs. In order to attain the most accurate average estimate possible, thick sections of randomly selected drill core intervals should be homogenized prior to analysis. Although this method is tedious, if done for several major Archean IFs it could help constrain iron isotope mass balance during that era. More specifically, this work would serve as a test for the model in which deposition of isotopically heavy IFs in the deeper parts of basins created a pool of isotopically light dissolved iron that was then buried in shallow-water environments in the Archean (Rouxel et al., 2005).

### 9.18.5.2.2 Chromium isotopes

Frei et al. (2009) measured Cr isotope values for 23 IFs ranging in age from 3.8 to 0.635 Ga (see Figure 10(c)). Their data revealed a pronounced difference between Archean–Paleoproterozoic and Neoproterozoic IFs. The former have Cr isotope values within a small range of $-0.26$ to $0.28\%_\text{o}$, whereas the latter reach values up to $+4.9\%_\text{o}$. Based on the small positive Cr isotope excursions in IFs as old as 2.7 Ga, Frei et al. (2009) argued that Cr (VI) was mobilized from land to the oceans, where it was then quantitatively reduced to Cr(III) and incorporated into IF. The Cr (III) oxidative step that generates the Cr(VI) requires a MnO$_2$ catalyst, which importantly necessitates significant levels of atmospheric O$_2$ as much as 300 My before the GOE (e.g., Holland, 2005). However, consideration of the depositional setting questions this conclusion. The Archean IFs that display statistically significant positive $\delta^{53}$Cr values were proximal to submarine hydrothermal centers (Figure 10(c)) and analyses of some modern hydrothermal vent fluids suggest hydrothermal Cr flux, likely as a reduced, organically bound Cr(III) phase (Sander and Koschinsky, 2000). Some of the solubilized Cr(III) in hydrothermal fluids is expected to be removed in the subsurface or at the sediment–water interface together with hydrothermal sulfides. Although redox-independent processes typically result in extremely small isotope fractionations, it is possible that kinetic isotope effects in hydrothermal systems were amplified, similarly to what has been observed for other transition metals (e.g., Rouxel et al., 2008a,b). The end result would be a dissolved Cr pool with small positive $\delta^{53}$Cr values that could be recorded in some IFs proximal to volcanic centers. In this context, positive $\delta^{53}$Cr values from $\sim 2.7$ Ga IFs (Frei et al., 2009) may provide insights into hydrothermal processes in the Archean, but are unlikely to reflect oxidative continental weathering. The currently available IF Cr isotope results are intriguing, but given the uncertainties about Cr cycling in Archean submarine hydrothermal systems, these data are not a straightforward archive of atmospheric processes.

Slightly positive Cr isotope values as reported in the upper part of the $\sim 1.88$ Ga Gunflint IF (Frei et al., 2009), are also difficult to interpret in terms of terrestrial oxidative processes, because this section of the IF experienced meteoric alteration during the $\sim 30$ My hiatus that followed deposition of this unit (Burton and Fralick, 2007). Destruction of original sedimentary features, intense silicification and carbonatization and pronounced groundwater-derived vanadium enrichment characterize the upper part of this alteration zone (Burton and Fralick, 2007). Groundwater systems, similarly to hydrothermal systems, are likely to amplify small sorption isotope effects. In fact, modeling of reactive Cr transport in a groundwater plume suggests that there can be a tenfold amplification of a Cr sorption isotope effect (Johnson and Bullen, 2004). Chromium isotopes in sedimentary rocks altered by groundwater systems are therefore equally unlikely to provide insights into atmospheric processes. By contrast, highly positive values in the Neoproterozoic IFs (Frei et al., 2009) likely reflect redox cycling, since both the partial reduction and the oxidation of Cr should create a mobile Cr(VI) reservoir with positive Cr isotope values. It would appear that the onset of a rigorous, terrestrial Mn-oxide cycle occurred sometime between the Paleo- and Neoproterozoic, but the current data gap precludes precisely defining the timing of this event.

### 9.18.5.2.3 Silicon isotopes in chert and IF bands

In order to better constrain silica sources, a growing number of studies have investigated the Si isotope composition of Precambrian IFs and cherts (Abraham et al., 2011; Andre et al., 2006; Ding et al., 1996; Jiang et al., 1993; Robert and Chausidon, 2006; Steinhoefel et al., 2009, 2010; van den Boorn et al., 2007, 2010). Silicon isotope ratios are resistant to hydrothermal alteration and thermal effects, including high-grade metamorphism (Andre et al., 2006), due to the high abundance of Si in most rocks. Si isotope ratios are thus regarded as recording primary signatures. However, biogenic and abiotic, low-temperature amorphous silica precipitates are significantly enriched in light Si isotopes relative to dissolved Si source (e.g., Basile-Doelsch, 2006; Ding et al., 1996; Douthitt, 1982), resulting in residual dissolved Si depleted in light isotopes. Diatoms, which preferentially incorporate light Si isotopes, dominate the modern marine silica cycle. As a result, dissolved Si in modern seawater has positive $\delta^{30}$Si values (+0.6 to +2.2\%$, Figure 12; De La Rocha et al., 2000; Reynolds et al., 2006). The same fractionation pattern arises from chemical precipitation of silica near seafloor
modern ocean, although the values may have been higher (Fitoussi et al., 2009; Savage et al., 2010), hydrothermal sea-floor precipitates typically have more negative d values in the same range as mafic/ultramafic volcanic rocks (Si isotope composition in the same range as mafic/ultramafic hydrothermal vents. Whereas black smoker vent fluids have a Si isotope composition typical of a hydrothermal source of the silica and are similar to those of modern hydrothermal vents (Fitoussi et al., 2009; Savage et al., 2010), seawater and mid-ocean ridge hydrothermal fluids (De La Rocha et al., 2000), modern siliceous precipitates of seafloor hydrothermal systems (Ding et al., 1996; Wu, 1991), Paleoarchean cherts from the Pilbara craton (van den Boorn et al., 2007, 2010), the 3.8 Ga IS from the Isua greenstone belt (André et al., 2006), the 2.7 Ga IS from the Shurugwi greenstone belt (Steinhoefel et al., 2010) and North China craton (Ding et al., 1996; Jiang et al., 1993), and the 2.5 Ga IS from the Hamersley Group and Transvaal Supergroup (Steinhoefel et al., 2010).

Figure 12  δ30Si values of mafic to ultramafic volcanic rocks (Fitoussi et al., 2009; Savage et al., 2010), seawater and mid-ocean ridge hydrothermal fluids (De La Rocha et al., 2000), modern siliceous precipitates of seafloor hydrothermal systems (Ding et al., 1996; Wu, 1991), Paleoarchean cherts from the Pilbara craton (van den Boorn et al., 2007, 2010), the 3.8 Ga IS from the Isua greenstone belt (André et al., 2006), the 2.7 Ga IS from the Shurugwi greenstone belt (Steinhoefel et al., 2010) and North China craton (Ding et al., 1996; Jiang et al., 1993), and the 2.5 Ga IS from the Hamersley Group and Transvaal Supergroup (Steinhoefel et al., 2010).

The modern ocean has several iron sources: (1) river and groundwater discharge, (2) continentally derived atmospheric dust, (3) remobilization from coastal and shallow-water sediments, (4) sea ice, and (5) hydrothermal fluids (Boyd and Ellwood, 2010). Oxidizing conditions in modern seawater limit dissolved iron content in open oceans to ~0.02–2 nM (average 0.5 nM; Bruland and Lohan, 2006); the present residence time of iron in the oceans is 100–200 years (Johnson et al., 1997), which is much shorter than the global ocean circulation time of ~1000 years (Boyd and Ellwood, 2010).

During oxidative continental weathering, iron is retained in soils in the oxidized state. Iron-chelating and Fe(III)-reducing bacteria play an important role in iron release from soil horizons (Brantley et al., 2004). Rivers carry a significant load of iron in the form of organically bound complexes, colloids, and particulate matter. This continentally derived iron flux is largely deposited in estuaries due to flocculation of colloids induced by salinity (Boyle et al., 1977; Escober et al., 2009). Where groundwaters are delivered into subterranean estuaries, redox change as well as pH change during seawater mixing result in near-quantitative Fe(II) oxidative precipitation (Rouxel et al., 2008a,b). Outwash in glaciated environments typically has high loads of mixed-valence silicate species and colloidal iron, but low concentrations of dissolved Fe and dissolved organic carbon (Schroth et al., 2011). In contrast, the dissolved iron flux in rivers flowing over unglaciated settings consists of organically bound iron complexes and iron
oxyhydroxides. Therefore, rivers flowing through glaciated and unglaciated terrains deliver different Fe species to the oceans, which has important implications for Fe bioavailability in seawater.

Dissolved iron derived from aerosols amounts to 2–12 × 10^9 mol per year and corresponds to 30–70% of the iron flux associated with upwelling deep waters; resuspended sediments and the diagenetic flux from coastal and shallow-water sediments are other important components estimated to contribute 4.7–8.9 × 10^9 mol per year to the oceanic iron budget (Elrod et al., 2004). Anoxic to suboxic conditions in pore waters in shelf sediments can lead to an iron shuttle – the delivery of reactive iron from the shelf to the basin (Lyons and Severmann, 2006; Severmann et al., 2008). Nonsteady-state diagenesis in tropical muds deposited on the delta top and inner shelf along the Amazon–Guiana coast of South America provides another example of an environment where Fe(III) reduction is the predominant metabolic process responsible for organic carbon oxidation. As a consequence, significant amounts of Fe are cycled between the bottom waters and surface sediments (Aller et al., 1986, 2010).

Although most base metals and some iron released from hydrothermal vents at mid-ocean ridges and in back-arc systems are precipitated at or immediately above the seawater–oceanic crust interface, as hot reducing hydrothermal fluids mix with cold oxidized seawater, iron and manganese disperse via hydrothermal plumes as far as hundreds of kilometers before deposition as metal oxyhydroxide particles, organic complexes and metal sulfide clusters and nanoparticles (e.g., Boyle et al., 2005; Sander and Koschinsky, 2011; Toner et al., 2009; Yucel et al., 2011; also see Chapter 8.7). Wu et al. (2011) suggested a long-range (>500 km) transport of hydrothermal iron from the East Pacific Rise. The hydrothermal iron fluxes to the deep ocean are thought to account for 11–22% of the deep-ocean iron reservoir, with the total estimated hydrothermal iron flux being 9 × 10^9 mol per year (Bennett et al., 2009; Tagliabue et al., 2010). Yucel et al. (2011) estimated that nanoparticulate pyrite constitutes up to 10% of the dissolved Fe (<200 nm) flux from high-temperature (>300 °C) hydrothermal vents. In addition, Fe-rich smectite clays, such as nontronite and montmorillonite, preferentially form close to depressed oxygen concentrations, hydrothermally active areas at the seafloor, where metalliferous sediments are deposited (Cuadros et al., 2011; Taitel-Goldman and Singer, 2002).

Concentrations of metals in hydrothermal fluids, including Fe and Mn, primarily increase with elevated fluid temperatures and salinity. The composition of the volcanic and plutonic host rocks as well as magma degassing and subsurface sulfide dissolution/precipitation may also affect the concentration of metals in seawater hydrothermal fluids (Edmond et al., 1979; Mottl et al., 1979; also see Chapter 8.7). In contrast to mafic-hosted hydrothermal systems, those sited within ultramafic rocks developed at slow-spreading ridges tend to have high contents of dissolved reduced gases (e.g., H₂ and CH₄) and metals (Douville et al., 2002; Marbler et al., 2010). Furthermore, because ultramafic-hosted hydrothermal systems release less H₂S, a larger proportion of dissolved iron precipitates as iron oxyhydroxides (Marbler et al., 2010).

Serpenitization of ultramafic rocks under submarine conditions releases H₂ from the fluids and results in partial Fe(II) oxidation to yield magnetite precipitation (Holm and Charlou, 2001).

Several submarine deeps in the Red Sea contain hot, anoxic stratified brine pools derived from rift-hosted hydrothermal systems that at depth leach Cenozoic evaporites. Metalliferous sediments containing iron oxides, silicates, sulfides, and carbonates precipitate from these brines (Taitel-Goldman and Singer, 2002). Interestingly, whereas the deep-water brine deposits are characterized by hydrothermal signals in terms of Sr and Nd isotopes and REE patterns, shallower-water deposits of the Shaban Deep have Sr, Nd, and REE patterns dominated by a seawater component (Cocherie et al., 1994), which is likely a reflection of the efficiency of REE sorption onto iron oxides and the presence of a discrete, stable redoxcline.

Modern shallow-water, iron-rich sediments are restricted to areas strongly affected by hydrothermal circulation related to active volcanism, such as within the Santorini Caldera in the Aegean Sea, where biologically mediated, ferric hydroxide and opaline silica precipitate (Hanert, 2002; Puchelt, 1973), and in Lake Malawi, eastern Africa, where nontronite mud and peloids and limonite with opal ooids are forming (Müller and Förstner, 1973). In addition, iron ooids and pisoids composed of iron oxyhydroxides admixed with amorphous silica, with volcanic rock fragments in the center, have been described by Heikoop et al. (1996) from offshore Mabiangtang, Indonesia, in the photic zone; these could be modern analogues of GIF. Iron-silicate ooids and peloids were also found from Cape Malapaas cuca to El Fraile Point, Venezuela, in shallow (~35–40 m depth) waters in an exhalative system connected to ultramafic rocks via a transform fault zone (Kimberley, 1994). Iron- and silica-bearing ooids also occur in the bottom sediments of Lake Chad, West Africa, off the Chari Delta (Lemoalle and Dupont, 1973).

Modern analogues comparable in scale to BIFs appear to be unknown, although evidence presented above questions whether there is a depositional rhythmicity to silica and iron banding in ancient IFs. Accordingly, we are left to speculate on the primary compositional structure of BIF.

Low-temperature Si- and Fe-rich hydrothermal deposits generally form in areas of active venting along mid-ocean ridge axes (Corliss et al., 1978; Mills, 1995), at off-axis seamounts (Alt, 1988), hot spots, and arc/back-arc submarine volcanic complexes (e.g., De Carlo et al., 1983). These submarine vent deposits generally do not form laterally extensive and very thick Fe–Si-rich deposits similar to IFs. Nevertheless, local but relatively thick Fe–Si-rich hydrothermal deposits have been documented in the recent rock record and on the modern ocean floor (e.g., Edwards et al., 2011a,b; Hekinian et al., 1993; Juniper and Fouquet, 1988). For example, silica- and iron-rich deposits 20 m thick occur at Ocean Drilling Program (ODP) Site 801 within Jurassic oceanic crust in the western Pacific Ocean (Alt et al., 1992; Rouxel et al., 2003). Fossil hydrothermal Si–Fe deposits, comprisingumber, jasper, and IF, have been found in ophiolites as old as 490 Ma (Little et al., 2004; Robertson, 1975). Recently, a modern analogue of IF deposits has been described in relation to ultradiffuse hydrothermal venting at the base of Loihi Seamount, at
5000 m below sea level (Edwards et al., 2011a,b). This hydrothermal system is expressed as regional seafloor seepage with gelatinous iron- and silica-rich deposits, located between and over basalt pillows, in places over a meter thick. The genetic model for umbers has previously invoked deposition of iron and manganese oxides via water-column precipitation from hydrothermal fluids, followed by particle fallout and accumulation in local depressions in a ridge flank setting. However, the laminated Fe–Mn structures located between basalt pillows observed at the base of Loihi Seamount offer an alternative interpretation for umber genesis that is consistent with geological observations.

In summary, modern environments where iron deposition occurs are comparable neither in scale nor in extent to those of ancient IFs. These modern environments nevertheless do provide some insights into processes that could have operated in the Precambrian to form IFs.

9.18.6.1 Hydrothermal Pulses of Si Synchronous with Fe Addition to Seawater

There has been limited work done on the sources of silica in IFs relative to iron. As such, there is still only a nascent understanding of some basic aspects of silica deposition in IFs and of the Precambrian Si cycle. IFs typically contain 34–56 wt% SiO₂ (Klein, 2005) and, potentially, represent a major sink for dissolved Si in the geological past. Prior to detailed facies and sequence-stratigraphic analyses of BIF, these deposits were long considered ambient chemogenic pelagites of the world ocean. However, it is now known that BIFs are deposits associated with pulses of reduced iron into basins. Although it is generally accepted that the source of silica in BIF was ambient seawater during most of the Precambrian, when the biological sink for seawater silica was presumably absent (Siever, 1992), the source of silica to the oceans of that time period remains uncertain. Two potential sources, similar to that for iron, have been proposed: submarine hydrothermal fluids (Mortlock et al., 1993; Wang et al., 2009) and the continents (Frei and Polat, 2007; Hamade et al., 2003). Support for the latter proposition has come mainly from the study of Ge/Si ratios in IFs, as hydrothermal fluids and rivers have very different Ge/Si ratios (Froelich et al., 1985; Mortlock et al., 1993). On the basis of covariation of Ge/Si ratios with silica content in the ~2.5 Ga Dales Gorge Member of the Brockman Iron Formation (Hamersley Group, Western Australia), Hamade et al. (2003) proposed a decoupling of Fe and Si sources, with Si being predominantly derived from riverine waters having low Ge/Si ratios due to weathering of continental landmasses. Caution is required in this interpretation, however, due to the strong Ge fractionation relative to Si by sorption onto iron oxyhydroxides (Pokrovsky et al., 2006) or by quartz precipitation (Evans and Derry, 2002). In addition, release of Ge to pore waters may also affect Ge/Si ratios (Rouxel et al., 2006). In either case, Ge/Si ratios in cherts may not reflect seawater composition but instead record multiple, unrelated, and geologically protracted processes. On the other hand, as mentioned above, Si isotopes are consistent with a hydrothermal Si source, suggesting that there were hydrothermal pulses of Si synchronous with Fe addition to seawater that in part drove IF deposition.

Ferric oxyhydroxide particles are highly reactive toward dissolved silica (e.g., Konhauser et al., 2007b), implying that drawdown of Fe from seawater was accompanied by drawdown of silica (e.g., Slack et al., 2007). It is possible that the silica component was scavenged from seawater during particle sedimentation and later was mobilized during diagenesis (see Fischer and Knoll, 2009, for additional discussion). Consistent with this interpretation, in GIF there are Fe-bearing chert peloids, deposited from suspension, and Fe-free chert cement that was deposited shortly after deposition, because the peloids have floating contacts. On the other hand, the petrographic and field evidence indicates that chert in bands in BIF was not a direct chemical precipitate from seawater, but rather a pore-filling cement and replacement of sediment; that is, chert in IFs is not a diageneric replacement of silica that precipitated in the water column.

A relevant geological perspective to address this issue comes from occurrences of chert in sequences that record relative magmatic quiescence and that have a similar duration and sedimentation rate to those that were deposited during periods of enhanced hydrothermal activity leading to the deposition of IFs. That silica was present in seawater during deposition in all Hamersley basin sequences is shown by the universal presence within them of bedded cherts or seafloor hardgrounds. For instance, sedimentation units in black shale of the Bee Gorge Supersequence, which directly underlies the Brockman Supersequence (predominantly consisting of IF), are capped by bedded cherts in exactly the same manner as tops of sedimentation units in the overlying IFs. There are hundreds of Bee Gorge Member ‘cycles’ from shale to bedded chert at scales of 1 m and less. In contrast, only a few seafloor replacement chert units exist in the black shale condensed sections in the underlying Roy Hill Member of the ~2.6 Ga Jeerinah Formation, which is not associated with IFs or increased hydrothermal activity, but was deposited at a similar sedimentation rate. Monotonous shales containing rare interbedded chert only return at the base of the ~2.4 Ga Turee Creek Supersequence, just above the Boolgeeda Iron Formation.

A possible explanation for these observations is that Si concentration in the water column was much lower during deposition of the Roy Hill Member. If this interpretation is correct, it implies that sedimentary basins at the time when IFs were deposited also contained higher seawater silica concentrations. It seems likely, therefore, that the silica content of the Precambrian ocean was not constant and that secular variations existed in [Si], like in [Fe], linked with changes in hydrothermal activity.

9.18.6.2 Oxidation Mechanism: Biological versus Nonbiological

Basic conditions leading to the deposition of IFs in ancient oceans are generally agreed upon: the precursor sediments precipitated from seawater containing micromolar (<100 μmol) levels of ferrous iron (Holland, 1973, 1984). An amplified marine reservoir of dissolved iron was possible due to the presence of (1) a reducing atmosphere or one having a low oxidizing potential (Bekker et al., 2004; Holland, 1984), (2) low marine sulfate and sulfide concentrations (Habicht et al., 2002), and (3) a high hydrothermal flux of iron (Kump and Seyfried, 2005). It is generally agreed that IF deposition is linked with an oxidative
mechanism that converted dissolved Fe(II) into solid-phase iron oxyhydroxide particles that then settled and accumulated on the seafloor. The caveat is, however, that unequivocal original grains of ferric oxyhydroxide in IFs have not been conclusively documented.

Specific mechanisms involved in the deposition of IFs (Figure 13) remain poorly resolved despite more than a century of investigation (e.g., Beukes and Gutzmer, 2008; Harder, 1919; Klein, 2005; Ohmoto et al., 2006). Given that IF deposition spans major evolutionary changes in Earth’s
surface composition, from an early anoxic atmosphere dominated by CO₂ and CH₄ to one that became partially oxygenated and CO₂-rich (e.g., Bekker and Kaufman, 2007), it is likely that IFs formed via different mechanisms during Precambrian time. The three mechanisms most widely supported in the literature are briefly summarized below.

9.18.6.2.1 Oxidation of Fe(II) by cyanobacterial O₂

The classic model for IF deposition invokes ferric iron precipitation occurring at the interface between oxygenated shallow waters and reduced upwelling iron-rich waters (Cloud, 1965, 1973; Figure 13(a)):

\[ 2\text{Fe}^{2+} + 0.5\text{O}_2 + 3\text{H}_2\text{O} \rightarrow 2\text{Fe(OH)}_3 + 4\text{H}^+ \]

Oxygen is assumed to have been produced by planktonic oxygenic photosynthesizers. Historically, cyanobacteria were inferred to be the primary producers utilizing oxygenic photosynthesis in Archean oceans, because compelling evidence for an eukaryotic fossil record before ~1.9 Ga is absent (Han and Runnegar, 1992; Javaux, 2011; Knoll et al., 2006). Several lines of evidence tentatively suggest that oxygenic photosynthesis evolved during the Neoarchean (e.g., Buick, 1992; Buick et al., 2006; Scott et al., 2011); however, at present, direct ‘smoking gun’ evidence for the former existence of cyanobacteria during that time period remains lacking (e.g., Brooks, 2011).

9.18.6.2.2 Metabolic Fe(II) oxidation

Metabolic Fe(II) oxidation is the most commonly invoked mechanism of IF deposition (Figure 13(b)). The potential importance of this process has been recognized for almost a century (e.g., Harder, 1919). However, it was not until a surge in the amount of interest and research on microbial iron oxidation over the past 10–20 years that this model for IF deposition became prevalent.

Three main pathways of metabolic microbial iron oxidation exist: (1) microaerophilic, (2) anoxygenic photosynthetic, and (3) nitrate-dependent. Microaerophilic bacteria, such as Gallionella ferruginea, Leptothrix ochracea, and Martiropfundus ferroxydans, play an important role in Fe(II) oxidation in modern iron springs and seafloor hydrothermal vent systems. In addition to using ferrous iron as their electron donor, microaerophilic Fe(II) oxidizers use oxygen as their electron acceptor and take in carbon dioxide, which is then reduced to organic carbon through the process of chemooautotrophy:

\[ 6\text{Fe}^{2+} + 0.5\text{O}_2 + \text{CO}_2 + 16\text{H}_2\text{O} \rightarrow [\text{CH}_3\text{O}] + 6\text{Fe(OH)}_3 + 12\text{H}^+ \]

Presently known microaerophilic Fe(II) oxidizers belong to the phylum Proteobacteria, but the metabolic pathway appears to be nested within several different clades of this broad grouping of bacteria. Interestingly, microaerophilic Fe(II) oxidizers have been recently discovered to be widespread in marine systems. For example, a strain of bacteria that is morphologically indistinguishable but phylogenetically distinct from Gallionella was found to be abundant at the iron-rich hydrothermal vents on Loihi Seamount and other seafloor, iron-rich hydrothermal systems (Emerson and Moyer, 2002; McAllister et al., 2011). Under low oxygen conditions, microaerophilic microbial Fe(II) oxidizers can dominate the iron cycle because the rate of microbial Fe(II) oxidation can be more than 50 times faster than abiotic rates (e.g., Sogaard et al., 2000). Additionally, Fe(II)-oxidizing bacteria are present, and may be abundant, at the chemoflume in ferruginous lakes, where ferric iron-rich sediments are deposited (e.g., Pavin Lake in France; Lehours et al., 2007).

Anoxygenic photosynthetic oxidation – photoferrotrophy – is another metabolic Fe(II) oxidation pathway. This pathway was predicted to be common on early Earth and linked to IF deposition (e.g., Baur, 1979; Hartman, 1984) before organisms capable of this metabolism were first cultured in the early 1990s (Widdel et al., 1993). Since then, a variety of phylogenetically diverse strains of anoxygenic Fe(II)-oxidizing photosynthesizers have been recognized, including strains of purple sulfur, and purple nonsulfur and green sulfur bacteria. Anoxygenic photosynthesis uses Fe(II) instead of H₂O as an electron donor, producing Fe(III) rather than dioxygen (Ehrenreikh and Widdel, 1994), according to the following reaction:

\[ 4\text{Fe}^{2+} + 11\text{H}_2\text{O} + \text{CO}_2 \rightarrow [\text{CH}_3\text{O}] + 4\text{Fe(OH)}_3 + 8\text{H}^+ \]

In the past two decades, a number of experimental studies have confirmed that various purple and green bacteria can use Fe(II) as a reductant for carbon dioxide fixation (e.g., Heising et al., 1999; Straub et al., 1999; Widdel et al., 1993).

More recently, organisms capable of metabolically coupling iron oxidation to nitrate reduction have been discovered (e.g., Edwards et al., 2003; Straub et al., 1996), providing another possible microbially mediated mechanism for IF deposition:

\[ 10\text{Fe}^{2+} + 2\text{NO}_3^- + 24\text{H}_2\text{O} \rightarrow 10\text{Fe(OH)}_3 + \text{N}_2 + 18\text{H}^+ \]

Nitrate-dependent Fe(II) oxidation has been shown to be widespread in sediments (Straub and Buchholz-Cleven, 1998). Most of the described nitrate-dependent Fe(II)-oxidizing strains require an organic substrate (e.g., acetate), although lithoautotrophic, nitrate-reducing Fe-oxidizing strains (z- and γ-Proteobacteria) have been isolated in pure culture (Edwards et al., 2003). A chemolithoautotrophic Fe(II)-oxidizing, nitrate-reducing enrichment culture has been identified (Straub et al., 1996) that is capable of oxidizing Fe(II) with nitrate autotrophically. However, this culture consists of chemoheterotrophic nitrate-reducing bacteria and a novel chemolithoautotrophic Fe(II)-oxidizing bacterium (Blothe and Roden, 2009). The inability (thus far) in laboratory experiments to obtain a pure culture of nitrate-reducing iron-oxidizers suggests that a consortium of organisms is needed for nitrate-dependent Fe(II) oxidation.

There is a strong basis to believe that metabolic Fe(II) oxidation could have driven IF deposition. Simple modeling indicates that even modest populations of microaerophilic or photosynthetic iron oxidizers could account for deposition of IFs, even assuming rapid accumulation rates (Konhauser et al., 2002). Recent modeling indicates that photosynthetic Fe(II) oxidation can result in near-to-quantitative drawdown of upwelling ferrous iron under modest ocean mixing and circulation conditions (e.g., Kaplan et al., 2005). Furthermore, metabolic Fe(II) oxidation is currently driving the deposition of extremely iron-rich sediments in ferruginous Lake Matano.
(Crowe et al., 2008), and microaerophilic iron oxidizers are likely important in the production of IF-like sediments on the seafloor, such as the Loihi hydrothermal field (Emerson and Moyer, 2002). Metabolic iron oxidation thus appears to be common in modern iron-replete aquatic systems. It follows that these processes likely were common in iron-rich water columns in ancient oceans.

Numerous studies have documented a fossil record of microbially mediated iron oxidation in Proterozoic and younger, iron-oxide-rich sedimentary rocks (Grenne and Slack, 2003; Juniper and Fouquet, 1988; Lager, 2001; Little et al., 2004; Planavsky et al., 2009; Slack et al., 2007) but not in Archean IFs. Microaerophilic Fe(II) oxidizers produce distinctive oxide-coated structures that have been found in jaspers and IF dating back to the Paleoproterozoic (Lager, 2001; Planavsky et al., 2009; Slack et al., 2007). Gallionella-type or Mariprofundus-like oxidizers are bean-shaped cells that grow at the terminus of a helical structure called a stalk, which is composed largely of polysaccharides typically encrusted by ferricyanide (Comolli et al., 2011), whereas Leptothrix-type oxidizers produce an iron-encrusted sheath. The apparent absence of these fossils remains in the Archean, contrasted to their local abundance in younger iron-oxide-rich rocks, might be a signal of a temporal increase in the involvement of microaerophilic oxidizers. However, it has been demonstrated that G. ferruginea does not form a stalk under the lower range of oxygen conditions at which these organisms can thrive (Hallbeck and Pedersen, 1996). Therefore, generally more reducing conditions in the Archean may have precluded template-mediated oxide formation (i.e., formation of oxide-coated sheaths and helical structures), but might not necessarily have excluded microaerophilic oxidizers.

It may be possible to use geochemical tools rather than the fossil record to pinpoint oxidation mechanisms. For instance, the REE signature of IFs through time indicates the lack of a discrete redoxcline prior to the REE signature of IFs through time indicates the lack of a multibillion-year duration. This hypothesis, although not yet supported by either detailed modeling or empirical data, warrants serious consideration because it has the potential to explain the enigmatic association of Archean jasper and oxide-facies IF with some Cu-rich VMS deposits that formed in deep-water settings.

### 9.18.6.2.3 Ultraviolet photooxidation of Fe(II)

As an alternative to biological models for Fe(II) oxidation, Cairns-Smith (1978) proposed that ferrous iron was photooxidized by the high flux of ultraviolet (UV) photons that would have reached Earth’s surface prior to the rise of atmospheric oxygen and the subsequent development of a protective ozone layer (Figure 13(c)). This reaction proceeds readily in acidic waters exposed to wavelengths in the 200–300 nm range:

\[
2\text{Fe}^{2+} + 2\text{H}^{+} + h\nu \rightarrow 2\text{Fe}^{3+} + \text{H}_2
\]

Brateman et al. (1983) and Anbar and Holland (1992) further explored the viability of the photochemical oxidation model at circumneutral pH over a range of UV wavelengths (217–406 nm). Based on a quantum yield determined from proton flux, they suggested that at pH > 6.5, the presence of the dissolved ferrous iron species Fe(OH)\(^{2+}\) is important because it is oxidized by photons having \(\lambda \approx 300–450\) nm, a wavelength region where the solar flux is more intense and where seawater is more transparent as compared to \(\lambda < 300\) nm. The dissolved ferric iron formed is subsequently hydrolyzed and precipitated as ferric oxyhydroxide. Extrapolating from these experiments, a mean photochemical oxidation rate of 0.5 mol Fe(II) m\(^{-2}\) year\(^{-1}\) has been estimated at rapid upwelling rates (4000 m year\(^{-1}\)), indicating that this process alone could have accounted for deposition of as much as 1.8 \(\times 10^{14}\) mol Fe(III) annually (François, 1986). Other estimates place the total amount of Fe(II) photooxidized annually at 2.3 \(\times 10^{13}\) mol (Brateman and Cairns-Smith, 1986). These rates are much greater than the annual rates inferred during deposition of the largest Archean and Paleoproterozoic BIF (Pickard, 2002, 2003), although, as mentioned above, it is difficult to constrain maximum sedimentation rates for IFs.

Importantly, the earlier photochemical models focused on determining the rates of Fe(II) photooxidation at, or close to, thermodynamic equilibrium with 0.02 mM Fe and under rather simplistic geochemical conditions in which other ions were unavailable for reaction with dissolved Fe(II). In this regard, Konhauser et al. (2007a) performed a series of experiments designed to mimic conditions in a photic zone proximal to a seamount-type vent system effusing elevated concentrations of Fe(II) into seawater that was saturated with respect to amorphous silica and calcite. Under those conditions, the photochemical contribution to solid-phase precipitation was negligible compared to the formation of the ferrous silicate mineral, greenalite, or the ferrous carbonate, siderite. Many IFs are composed predominantly of iron carbonates or iron silicates, both of which have been widely suggested to be abiogenic marine precipitates (e.g., Ohmoto et al., 2004). However, as discussed above, based on petrographic and isotopic constraints, siderite in most IFs is an early diagenetic, or later, mineral phase. In experiments where Fe(II) was exposed to either photosynthetic Fe(II)-oxidizing bacteria or dioxygen, ferric oxyhydroxide formed considerably faster than by UV-photooxidation.

As an alternative to these classical models, Foustoukos and Bekker (2008) argued that some deep-water, volcanic-hosted IFs deposited in association with VMS deposits could have formed by oxidation during phase separation in the subsurface into vapor and brine, with hydrogen and HCl being removed into the vapor phase, generating oxidizing and alkaline conditions in the brine. During this process, transition metals would form chloro-complexes and would then be enriched in the brine, which would be expelled from magmatic chambers during large eruptions. This hypothesis, although not yet supported by either detailed modeling or empirical data, warrants serious consideration because it has the potential to explain the enigmatic association of Archean jasper and oxide-facies IF with some Cu-rich VMS deposits that formed in deep-water settings.

### 9.18.7 Secular Trends for Exhalites, IFs, and VMS Deposits

#### 9.18.7.1 Relationships among Mantle Plumes, IF Deposition, and VMS Mineralization

Secular trends in the distribution of Precambrian IFs and VMS deposits (Figure 14) have been discussed previously by many authors (e.g., Bekker et al., 2010; Groves et al., 2005; Huston and Logan, 2004; Huston et al., 2010; Isley and Abbott, 1999;
Although the deposition of IFs has been genetically linked to mantle plume breakout events and mafic volcanism (Isley and Abbott, 1999), the largest VMS deposits are instead generated in arc settings marked by bimodal volcanism (Franklin et al., 2005). Given that IFs were likely precipitated from hydrothermal plumes and that iron in hydrothermal plumes was released by submarine hydrothermal alteration of volcanic rocks in the deeper part of the ocean, large VMS deposits should have formed contemporaneously near the volcanic sources. Exhalites that formed in stratigraphic association with Cu-rich VMS deposits provide a unique record of deep-water ocean redox processes.

Figure 14 Secular distributions of mantle superplume breakout events and selected marine mineral deposits during the Precambrian. Distribution of mantle superplumes is from Abbott and Isley (2002) in which the y-axis (height) is the sum of Gaussian time series for high-Mg intrusive rocks and layered intrusions, flood basalts, and dikes. Distribution of marine Fe deposits is plotted as amount of iron formation in billion metric tons (Gt), integrated over time intervals of 50 Ma (data in Appendix 2); apparently small deposits that lack published tonnage data are assigned a size of 0.2 Gt; deposits having poor age control are included as pale orange bars. Distribution of marine Mn deposits is plotted in million metric tons (Mt) of Mn from Maynard (2010). Time periods dominated by banded iron formation (BIF), granular iron formation (GIF), and Rapitan-type iron formation are shown schematically (see text). Distribution of VMS deposits is similarly plotted in Gt for time intervals of 50 Ma (data in Franklin et al., 2005; Appendix A2). Secular facies variation of exhalites associated with deep-water, Cu-rich VMS deposits is from Slack and Cannon (2009). Modified from Bekker A, Slack J, Planavsky N, et al. (2010). Iron formation: The sedimentary product of a complex interplay among mantle, tectonic, oceanic, and biospheric processes. Economic Geology 105: 467–508, with data for Mn deposits added from Maynard, 2010. With permission from the Society of Economic Geologists.
state (Slack et al., 2007). The following sections explore secular trends in seafloor-hydrothermal exhalites and IFs to infer the ocean redox state during their deposition.

### 9.18.7.2 Secular Patterns in Precambrian VMS-Related Exhalites

There is consensus that the source of iron in Precambrian IFs was in hydrothermal fluids that vented into the deep ocean during submarine volcanism (Bekker et al., 2010, and references therein). Because the generation and transport of this iron are believed to be linked to hydrothermal processes and anoxic conditions, secular trends of IFs, VMS deposits, and the redox state of coeval deep seawater aid in understanding of episodic IF deposition during Precambrian time (Figure 14).

Insights into ancient deep-ocean redox states come from diverse proxies, one of which is the nature of exhalites that formed by precipitation from hydrothermal vents and plumes as parts of Cu-rich VMS systems. Cu-rich VMS deposit formation is restricted to settings with fluids having temperatures ≥300 °C and water depths of >850 m, which permit the precipitation of relatively abundant Cu (>1wt%) in sulfides on or near the seafloor. At shallower depths, hydrothermal fluids at this temperature, assuming present seawater salinity (3 M NaCl), boil in the subsurface and do not produce Cu-rich massive sulfide (Slack et al., 2007).

The exhalites occur as stratiform layers or lenses, up to several meters thick, typically in the hanging wall above VMS deposits or at the same stratigraphic level for hundreds of meters or more along strike (Spry et al., 2000). Given abundant evidence of exhalite mineralization on the modern seafloor (Hannington et al., 1995; also see Chapter 8.7), ancient exhalites are widely interpreted as precipitates from spatially associated VMS systems (e.g., Grenne and Slack, 2005; Isley, 1995; Peter, 2003). The mineralogy and geochemistry of exhalites in the geological record vary widely (Spry et al., 2000), but mainly consist of fine-grained quartz (chert) and one or more Fe-rich minerals including hematite, magnetite, greenalite, stilpnomelane, grunerite, siderite, ankerite, pyrite, or pyrrhotite; some exhalites have abundant Mn (in rhodochrosite, kutnahorite, and spessartine), Ba (in barite, hyalophane, and celsian), P (in apatite), F (in fluorite), or B (in tourmaline). In addition to the requirement of a Cu-rich mineralogy in related VMS deposits, several other limitations exist for using exhalites as a paleoredox proxy (see Bekker et al., 2010; Slack et al., 2007).

Mineralogical data for 45 Precambrian exhalites linked to Cu-rich VMS deposits (Figure 14; Appendix 2) show a seafloor pattern in which exhalites older than ~1.85 Ga mainly comprise Fe-silicates, sulfide, sulfidic shale, pyritic chert, or pyritic tuff. In contrast, exhalites younger than ~1.85 Ga typically have abundant iron oxides in jasper, hematitic IF, or magnetitic IF as well as Fe-silicates in some cases. Occurrence of the former group of reduced-facies exhalites, which predominates in the Archean and early Paleoproterozoic, suggests deposition on the seafloor under anoxic and possibly sulfidic bottom waters, based on geochemical and thermodynamic arguments (Huston and Logan, 2004; Ohimoto et al., 2006). Younger Precambrian exhalites that contain abundant hematite and magnetite have relatively high Fe(III)/Fe(II) ratios that rule out sulfidic bottom waters during mineralization, assuming no oxidation took place during postdepositional alteration, diagenesis, metamorphism, or weathering (Slack et al., 2009). The presence of hematite or magnetite in these exhalites requires at least suboxic conditions (<5 μM O₂) in coeval deep seawater, in order to permit the precipitation of ferric oxyhydroxide precursors as documented in modern seafloor-hydrothermal vents and plumes (Grenne and Slack, 2005; Slack et al., 2007). It is possible that magnetite in ancient exhalites had a mixed ferrous–ferric ‘green rust’ precursor such as Fe(OH)₂, which in theory could have formed in anoxic seawater (Murray, 1979). This precursor is unlikely, however, because Archean and Paleoproterozoic seawater, as well as diagenetic pore fluids, probably were saturated in FeCO₃ prior to Fe(OH)₂ saturation, owing to high atmospheric pCO₂ during this time period and resulting bicarbonate saturation in coeval seawater and shallow pore waters (Slack et al., 2009, and references therein). Fully oxidic exceptions for contemporaneous deep seawater are also discounted, based on REE data on late Paleoproterozoic hematite- and magnetite-rich exhalites that have small negative to small positive shale-normalized Ce anomalies, in contrast to the larger negative Ce anomalies that characterize modern iron oxyhydroxide deposits (Slack et al., 2007, 2009).

There are several exceptions to the patterns of Fe-silicate and sulfide exhalites that are genetically linked to Archean and early Paleoproterozoic Cu-rich VMS deposits (Figure 14). These exceptions, which include magnetite IF, hematite IF, or jasper, occur at the following deposits: 2960 Ma Scuddles, Western Australia; 2720 Ma Geco and Willroy, Ontario, Canada; ~2530 Ma Wutai, Shaxi Province, China; and ~1870 Ma Bend and Eisenbrey, Wisconsin, USA. The presence of abundant Fe(III) in hematite and/or magnetite within these >1850 Ma deposits (Slack and Cannon, 2009) should be further evaluated to constrain their origin.

The absence of Cu-rich VMS deposits during some periods of Earth history precludes the use of exhalites for evaluating the redox state of coeval deep oceans. For example, several VMS deposits have been documented in sequences >3000 Ma, including the oldest (3465 Ma) deposits at Big Stubby and Lennons Find in the Pilbara craton of Western Australia. These deposits contain abundant Zn and Pb with little or no elevated Cu (Franklin et al., 2005; Huston et al., 2002; Appendix 2), and hence could have formed at relatively shallow depths of a few hundred meters. The oldest Cu-rich VMS deposit with a genetically linked exhalite unit is the ~2.97 Ga Miranda deposit in South Africa (Slack and Cannon, 2009). A long gap of ~620 Ma in exhalite data exists from ~2530 to 1910 Ma, during which 16 VMS deposits are known (Franklin et al., 2005; Appendix 2), but none has a reported exhalite. Additional long gaps in the record of Precambrian deep-marine exhalite mineralization of this type are from ~1700–1400 Ma and ~1000–770 Ma. The emerging record of deep-ocean redox state from studies of exhalites associated with VMS deposits is entirely consistent with other geochemical and geological proxies for the rise of atmospheric oxygen in the early Paleoproterozoic and suboxic deep-ocean redox state in the late Paleoproterozoic and Mesoproterozoic.
9.18.7.3 Secular Patterns in Sedimentary Iron Deposits

In order to evaluate a secular pattern in sedimentary iron deposits, we divide Earth history into time intervals characterized by different redox states of the atmosphere–ocean system. The following discussion begins with the Eoarchean and progresses sequentially toward the modern iron cycle.

9.18.7.3.1 Eoarchean IFs

In the Isua greenstone belt of western Greenland, ~3.8 Ga chert and BIF at amphibolite facies metamorphic grade are tectonically interleaved with amphibolite that locally preserves pillow structures (Myers, 2001). The BIF consists predominantly of quartz and magnetite, with minor cummingtonite/graunerite, actinolite, hornblende, and calcite (Dymek and Klein, 1988). Polat and Frei (2005) observed significant enrichments of Fe in Isua pillow basalts, which they attributed to leaching from oceanic crust and precipitation on the seafloor from hydrothermal fluids, suggesting that high-temperature hydrothermal alteration of the immediate subsurface might have played an important role in deposition of the Isua BIF.

In the ~4.3–3.8 Ga Nuvvuagittuq greenstone belt of the northeastern Superior Province of Canada, quartz–magnetite–graunerite BIF, as much as 30 m thick, and other siliceous rocks are intercalated with metasomatized amphibolite (Dauphas et al., 2007; Młoszewska et al., 2012; O’Neil et al., 2007). The REE-Y profiles and Fe isotope compositions of this BIF are consistent with its origin as marine exhalate. Low Al₂O₃, TiO₂, and high field strength element (HFSE) concentrations show that it is relatively detritus-free. There are distinctly seawater-like REE-Y profiles and consistently positive Eu anomalies. These features suggest that the rocks preserved some of their primary composition despite metamorphic overprint (Młoszewska et al., 2012).

9.18.7.3.2 Paleoarchean IFs

Paleoarchean IFs are intercalated with bedded chert in ultramafic to felsic volcanic successions in greenstone belts of the Kaapvaal and Pilbara cratons. Bedded cherts consist, to a large extent, of silicified sedimentary and volcaniclastic rocks. Although silicification is common in modern and ancient low-temperature hydrothermal systems in both continental and oceanic settings, the abundance of chert in the Paleoarchean is striking. In contrast, IFs are relatively rare in the >3.0 Ga record, despite evidence for widespread leaching of iron from seafloor rocks during hydrothermal alteration (Hofmann and Harris, 2008). It is possible that the absence of widespread IF deposition during this time period reflects preservational bias. However, many cherts of this age range are uniformly underlain by zones of low-temperature seafloor alteration up to several tens of meters thick (Duchac and Hanor, 1987; Hofmann and Harris, 2008; Hofmann and Wilson, 2007), which suggests the predominance of low-temperature hydrothermal processes at this time. These zones are characterized by silicification and depletions of many elements, including Fe, Mg, and some transition and base metals, which were likely exhaled by hydrothermal systems to the Paleoarchean ocean, thus providing a source of dissolved iron (Hanor and Duchac, 1990; Hofmann and Harris, 2008).

The Orverwacht Group of the Barberton greenstone belt, South Africa, records a time interval of ~240 My, but contains only a single stratigraphic unit that includes IF. The ~3.4 Ga Buck Reef Chert is an unusually thick sequence of predominantly black and white banded chert that contains a 100-m-thick unit of banded ferruginous chert, in which siderite is regarded to be the main iron-bearing mineral at depth (Tice and Lowe 2004). Also in the Barberton greenstone belt, jaspilitic IF units, several tens of meters thick, are intercalated with ferruginous shale of the 3.26–3.23 Ga Fig Tree Group (Heinrichs, 1980; Hofmann, 2005). Felsic volcanic detritus and evidence for hydrothermal activity in rocks interlayered with the IF are widespread (Hofmann, 2011). Two thin jaspilitic units interbedded with ferruginous shale also occur at the top of fining-upward sequences within the ~3.23 Ga Moodies Group of the Barberton greenstone belt and in association with volcanic rocks (Eriksson, 1983).

The Iron Ore Group of the Singhbhum craton in India contains economically important BIF deposits that likely represent different stratigraphic units (e.g., Banerji, 1977). In the southern part of the craton, a low-metamorphic grade, 120-m-thick BIF overlies a sequence of basalts and felsic volcanic rocks dated at 3.51 Ga, possibly indicating a Paleoarchean age for this BIF (Mukhopadhyay et al., 2008), although contact relationships with the underlying dated succession are unclear. In the western part of the craton, BIFs are intercalated with shales that are likely Mesozo- to Neoarchean in age. Absolute age constraints are not available, highlighting the need for high-precision geochronological studies of the Iron Ore Group in this region.

Amphibolite- to granulite-facies grade BIFs associated with amphibolites, calc-silicate rocks (carbonate-altered mafic rocks), talc-carbonate schists, and metacherts are present in a number of Paleoarchean high-grade remnants of greenstone belts in South Africa (e.g., Assegaaï–De Kraalen granitoid-greenstone terrane; Saha et al., 2010), Zimbabwe (e.g., Sebakwean Group; Wilson, 1968), India (e.g., Sargur Group; Naqvi and Rogers, 1987), and elsewhere. Strong deformation and metamorphism make their paleoenvironmental significance unclear. The close spatial association with siliceous and carbonate-bearing metasomatized lithologies suggests a strong hydrothermal influence on the formation of these BIF.

9.18.7.3.3 Neoarchean to Mesozoic IFs

BIFs are a distinct component of Neoarchean greenstone belts. These BIFs are either intercalated with submarine volcanic rocks or associated with sedimentary strata of continental-shelf environments. Good examples of continental shelf BIF deposits are found on several cratons that experienced transient crustal stability prior to the global magmatic event at 2.7 Ga. The 2.83–2.70 Ga Manjeri Formation of the Zimbabwe craton is a fluvial to shallow-marine succession that was deposited unconformably on older greenstones and granitoids (Hofmann and Kusky, 2004; Hunter et al., 1998). The IF of this unit is intercalated with quartz arenite, shale, and carbonate strata, and lacks obvious links to coeval volcanic rocks. Lithologically similar sequences include the ~2.8 Ga Central Slave Cover Group (Bleeker et al., 1999) of the Slave craton, the ~2.7 Ga Steep Rock Group in the Wabigoon greenstone belt and correlative sequences in Canada (Stone, 2010; Tomlinson
et al., 2003; Wilks and Nisbet, 1988), and the ~2.7 Ga Bababudan Group of the Dharwar craton in India (Srinivasan and Ojakangas, 1986; Trendall et al., 1997).

IFs interlayered with volcanic rocks can be found in most Mesoarchean and Neoarchean greenstone belts. They are commonly thin, less than 20 m thick, and laterally discontinuous. The volcanic components typically consist of mafic–ultramafic or bimodal volcanic rocks that erupted in relatively deep-water, oceanic settings. In several cases, a direct genetic relationship has been documented between volcanic activity and deposition of IF, reflecting volcanic and hydrothermal activity both proximal and distal to volcanic centers (Chown et al., 2000; Fyon et al., 1992).

Mesoarchean IFs that developed in cratonic cover successions between 3.0 and 2.9 Ga are well documented in the Witwatersrand and Pongola basins in the southeastern part of the Kaapvaal craton in South Africa (Beukes and Cairncross, 1991; Smith, 2007). These IFs are laterally extensive, but thin (~10 m thick), and commonly are interbedded with ferruginous shales deposited during marine transgressions. There are no direct relationships with volcanic rocks or volcanism at that time, in accord with trace element and Nd isotope data on the oldest IF of the Mozaan Group in the Pongola basin (Alexander et al., 2008), which indicate a strong continental contribution to the trace element inventory of these IFs.

9.18.7.3.4 Neoarchean IFs

IFs of 2.7–2.45 Ga age are the most laterally extensive and economically valuable of any time in Earth’s history. The oldest extensive and thick sediment-hosted BIF is the ~2.60 Ga Marra Mamba Formation of the Hamersley Province in Western Australia. It was deposited in a deep-water basinal setting on a passive continental margin during sea-level highstand of the Marra Mamba Supersequence, and has an average thickness of about 210 m (Krapež et al., 2003; Trendall and Blockley, 1970). This BIF carries a pronounced positive Eu anomaly (Albert and McCulloch, 1993), suggesting a strong hydrothermal imprint on REE systematics of the global ocean during its deposition. Although the Hamersley and Transvaal successions are equivalent in age and are correlative based on sequence stratigraphy, only ankerite-banded chert of the Campbellrand succession in South Africa corresponds to the Marra Mamba BIF (Beukes and Gutzmer, 2008). A mantle plume breakout event comparable in age to that of the Marra Mamba Formation is unknown.

Only one IF (Bruno’s Band) occurs on the Pilbara and Kaapvaal cratons stratigraphically between the ~2.60 Ga Marra Mamba IF and the overlying major 2.50–2.45 Ga IFs (Beukes and Gutzmer, 2008; Krapež et al., 2003). This younger episode of IF deposition closely corresponds with volcanism during the 2.50–2.45 Ga series of mantle plume breakout events of global extent (e.g., Heaman, 1997) and immediately preceded a supercontinent assembly (Barley et al., 2005). Deposition of these IFs occurred on a reactivated continental margin (Krapež et al., 2003), during a rise in sea level. In addition to these well-dated, time-equivalent IFs, those in the Quadrilátero Ferrífero region in Brazil, Middleback Ridge (Gawler craton) in South Australia, Krivoy Rog area in Ukraine, and KMA region in Russia are broadly similar in age based on available geochronological and chemostratigraphic constraints, as well as on similar patterns of megabanding (Bekker et al., 2003; Kulik and Korzhnev, 1997; Priłutsky et al., 1992; Spier et al., 2007; Szpunar et al., 2011). These giant IFs were also deposited on reactivated continental margins and are separated from overlying Paleoproterozoic sequences by a prominent unconformity. The unconformities correspond to a long gap in sedimentation following supercontinent assembly at ~2.4 Ga. Assuming that these IFs are similar in age, more than 70 wt% of the total original iron resources in Precambrian IFs was deposited during the time interval 2.60–2.40 Ga. Surprisingly, a coeval peak in VMS deposition is absent from the geologic record (Figure 14); in fact, no VMS deposits are known to have formed within this age range (Franklin et al., 2005; Appendix 2). Texturally, these IF deposits differ from the older BIF because they contain intervals having granular textures as well as micro- and nanospheres of hematite (Ahn and Buseck, 1990; Ayres, 1972; Beukes and Klein, 1990; Simonson and Goode, 1989; Spier et al., 2007). However, most granules consist of iron silicates and carbonates with only rare hematite granules; no hematite-coated grains have been documented (Beukes and Klein, 1990; Simonson and Goode, 1989; Spier et al., 2007). Such features are important because deposition of these IFs directly preceded the GOE, and post-GOE GIFs predominantly contain hematite as the iron-bearing mineral within granules and on coated grains. This observation implies that processes responsible for IF deposition before and after the rise of atmospheric oxygen were different, requiring more than one model to explain their formation.

9.18.7.3.5 IFs deposited after the GOE and before ~1.93 Ga

Giant IFs were not deposited between ~2.4 and 1.88 Ga. Nonetheless, sediment-hosted and volcanic-hosted IFs are known from this time period. Shortly after the rise of atmospheric oxygen at ~2.4 Ga (e.g., Bekker et al., 2004), oolitic hematitic ironstone of the lower Timeball Hill Formation, South Africa, was deposited in shallow water above fairweather wave base (Dorland, 1999; Schweigart, 1965). Its deposition at ~2316 Ma (Hannah et al., 2004) may coincide with a magmatic event at ~2.32 Ga (e.g., Berman et al., 2005; Eriksson et al., 1994a,b; Fetter et al., 2000; Hartlaub et al., 2007). The significance of this event and whether it is related to mantle processes are unknown. Correlative IFs are not documented on other continents.

The slightly younger ~2.22 Ga Hotazel Formation, also in South Africa, contains IF interlayered with manganiferous sedimentary rocks; this is the largest manganese deposit in the world (Tsikos et al., 2003). The iron and manganese formation lies above, and is most likely genetically related to, the submarine-emplaced Ongeluk Lavas of mafic to intermediate composition that are coeval with a 2.22 Ga mantle plume breakout event (Ernst and Buchan, 2001). Although volcanic rocks and dikes related to this event are developed on nearly all continents, other iron and manganese formations of this age are unknown. Iron- and manganiferous-rich strata of the Hotazel Formation consist of three upward-shallowing sequences deposited in a slope environment (Schneiderhan et al., 2006). Significantly, the deposit lacks positive Eu anomalies, but has pronounced negative Ce anomalies, indicating an oxygenated state of at least the upper part of the ocean (Tsikos and Moore,
The absence of a positive Eu anomaly is also important because it indicates that the global ocean was not dominated by a high-temperature, hydrothermal flux at that time and that Fe and Mn were likely derived locally, within the basin, by low-temperature, shallow-water alteration of the underlying thick (≥1 km) sequence of volcanic rocks. Considering that the 2.22 Ga mantle plume breakout event coincides with a positive carbon isotope excursion in seawater composition, the Lomagundi Event, related to high relative burial rates of organic carbon (e.g., Bekker et al., 2001; Karhu and Holland, 1996; Schidlowski et al., 1976), this REE pattern may indicate that significant parts of the oceans were already oxygenated by that time and that Fe and Mn were soluble only in isolated to semi-isolated basins overwhelmed by a hydrothermal flux of reductants.

Hematitic oolites and hematite-rich sandstones continued to be deposited in shallow-marine environments during the Lomagundi carbon isotope excursion in South Africa (e.g., Silverton Formation; Schweigart, 1965) and on the Kola Peninsula, Russia (Kuetsjärvi Sedimentary Formation; Akhmedov, 1972a). IFs deposited during the Lomagundi excursion are developed within the Lomagundi Group of the ~2.2–2.1 Ga Maqongdi belt, Zimbabwe (Master, 1991). In West Africa, they are present in the Ijil Group, Mauritania (Bronner and Chauvel, 1979) and the Nigerian schist belts (Mücke, 2005), both in the 2.2–2.1 Ga Birimian basin. Iron- and manganese-rich strata are also developed in West Africa in the Francévillian basin, Gabon (Leclerc and Weber, 1980). Volcanic-hosted IFs deposited during the Lomagundi carbon isotope excursion are also known in Brazil (e.g., Aimbé Formation, Guarinos Group; Resende and Jost, 1995; Ipatucuru Complex of the Rio Ipatucuru greenstone belt; Dalton de Souza et al., 2003) and Norway (Iddjajav’ri Group, Karasjok greenstone belt; Often, 1985). Considering uncertainty in the ages of these IFs, it is difficult to relate them to a specific mantle plume breakout event. However, several such events occurred between 2.2 and 2.1 Ga (Ernst and Buchan, 2001) and thus it seems likely that the deep-ocean redox state was too high, rather than the strength of mantle plume events being too low, to form giant sedimentary iron deposits during this time interval.

Following the end of the Lomagundi carbon isotope excursion at 2.1–2.0 Ga, small, volcanic-hosted IFs were deposited in several basins in North America (e.g., Homestake Iron Formation, Black Hills, South Dakota; Frei et al., 2008) and Finland (Laajoki and Saikkonen, 1977; Paakola, 1971). Oolitic hematitic ironstone is also present in the Kolajoki Sedimentary Formation, Kola Peninsula, Russia (Akhmedov, 1972b), which was deposited during this time. Combined, these data suggest that dynamic ocean redox conditions were re-established in the aftermath of the GOE, characterized by periodic upwelling of iron into shallow-water settings above storm and wave base in association with mantle plume breakout events. A number of mantle plume events are recognized during this time interval, the most prominent being the ~2069–2053 Ma event that led to the emplacement of the Bushveld Complex in South Africa, and a number of mafic to ultramafic dikes and intrusions on the Superior and Sarmatia cratons (Chernyshev et al., 1998; Elming et al., 2010; Ernst and Bleeker, 2010). The absence of contemporaneous large accumulations of sedimentary iron is puzzling. However, considering that this event immediately followed the Lomagundi excursion, during which large amounts of oxygen were released to the atmosphere–ocean system, it is possible that the ocean redox state was buffered to change by a hydrothermal flux of reductants, preventing anoxia and significant Fe transport to sites favorable for IF accumulation.

9.18.7.3.6 C.1.93–1.85 Ga IFs coeval with large VMS deposits

Extensive and large IFs reappear after an approximately 500 My gap, at about 1.88 Ga. These are predominantly GIFs and are most common in North America at the margins of the Superior craton (Simonson, 2003), and in Western Australia (Goode et al., 1983; Rasmussen et al., 2012). They are coeval with emplacement of the ~1.88 Ga ultramafic LIP (Heaman et al., 1986, 2009; Hultert et al., 2005) that is potentially related to a mantle plume breakout event (Hamilton et al., 2009; see for a different view Heaman et al., 2009) during the early stage in the assembly of Laurentia. These IFs are now recognized to be correlative, based on high-precision geochronology (Findlay et al., 1995; Fralick et al., 2002; Machado et al., 1997; Schneider et al., 2002; Stott et al., 2010). IFs extend discontinuously for more than 3000 km along the southern and eastern margins of the Superior craton, from Minnesota, Wisconsin, and Michigan, to Quebec (Mistassini basin), and northward to the Labrador Trough. Correlative and texturally similar GIFs also occur in the center of the Superior craton in the Hudson Bay region (Richmond Bay and Belcher Islands) and in Sutton Inliers, NW part of Ontario (Chandler, 1984; Hamilton et al., 2009; Hawley, 1926; Stott et al., 2010). This group of IFs is considered to have been deposited in extensional settings with coeval submarine mafic volcanism (Fralick et al., 2002; Ricketts et al., 1982; Schulz and Cannon, 2007). A back-arc basinal setting for the IFs has been proposed by these workers, whereas a foreland basin setting was advocated by others (e.g., Hoffman, 1987; Ojakangas et al., 2001; Schneider et al., 2002).

An intriguing question is whether deposition of these IFs represents either local, basin-scale conditions or the composition and redox state of the global ocean. This is a critical issue because occurrences and ages of these IFs have been used by some workers to infer deep-water anoxic conditions in the coeval global ocean (e.g., Poulton et al., 2004; Slack and Cannon, 2009); if these IFs instead reflect more restricted, basin-scale conditions, our understanding of the global ocean redox state might be incorrect. Paleogeographic reconstructions are not adequately detailed to answer this question; however, tidal signatures have been observed in IFs and interbedded sedimentary rocks of this age in Minnesota and the Hudson Bay region (Ojakangas, 1983; Chandler, 1984), which are consistent with at least episodically open-marine conditions during IF deposition. Two independent questions can be asked as a basis to address this issue. First, do sedimentary successions of similar age on the margins of other cratons provide evidence for high concentrations of iron in seawater? Second, do iron-oxide exhalites occur in association with coeval, deep-water (Cu-rich) VMS deposits?

The ~1.88 Ga Frere Iron Formation along the northern margin of the Yilgarn craton of Western Australia (Goode et al., 1983; Rasmussen et al., 2012) and IF of the Gibraltar Formation in the Kahochella Group along the southeastern
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margin of the Slave craton in North America (Bowring et al., 1984; Hoffman et al., 2011; Roscoe et al., 1987) are granular and likely equivalent in age to the Animikie IFs. In addition, the presence of the ~1.88 Ga Rochford Formation along the eastern margin of the Wyoming craton, although poorly dated and not granular (Frei et al., 2008), supports synchronous deposition of IFs on several cratons. On the other hand, a ~20-m-thick magnetite–hematite oolitic iron formation in the middle member of the Waterton Formation (Hurwitz Group) on the Hearne craton of Canada (Miller and Reading, 1993) is most likely older than the 1.88 Ga IFs on the margins of the Superior craton. The age of this oolitic unit is not well constrained, but detrital zircon geochronology for several units within the Hurwitz Group, including the Waterton Formation, suggests that it is ~1.93 Ga (Davis et al., 2005). In contrast, the precisely dated (1882 ± 3.5 Ma) lower part of the Recluse Group in the Kilohigok basin on the Slave craton, northwestern Canada, lacks IF (Bowring and Grotzinger, 1992). Instead, organic matter-rich and sulfidic shales are present at this stratigraphic level (Bowring and Grotzinger, 1992). Iron speciation and sulfur isotope systematics of these shales might help constrain the local redox state. However, coeval euxinic and ferruginous conditions are expected when there is an anoxic deep ocean (e.g., Planavsky et al., 2011; Reinhard et al., 2009).

Deposition of IFs on the Superior craton was synchronous with a peak in tonnages of VMS deposits (Figure 14), some of which were generated in arcs adjacent to the craton. VMS deposits of this age are known in the hinterland to the south of the Animikie basin (Schulz and Cannon, 2007), in the Labrador Trough (Barrett et al., 1988), and in the Trans-Hudson orogen (Syme and Bailes, 1993). Recent geochronological studies of the host metavolcanic rocks to VMS deposits in the Pembine-Wausau terrane of northern Wisconsin indicate that these deposits formed at ~1875 Ma, contemporaneously with GIF of the Animikie basin (Schulz and Cannon, 2008). These data also suggest that the hydrothermal systems were the source of iron for the IFs, consistent with earlier models (e.g., Isley, 1995). However, laterally continuous iron-oxide exhalates are absent at or near these deep-water VMS deposits that formed presumably in open-marine conditions. This observation is unlikely to reflect preservational bias, because slightly younger 1.84, 1.79, and 1.78 Ga Cu-rich VMS deposits that similarly formed in arc settings contain abundant hematite and magnetite exhalates, jasper, and IF (Slack and Cannon, 2009; Slack et al., 2007). This analysis points toward an anoxic and ferruginous composition of deep waters in open-marine settings at ~1.93–1.88 Ga.

The Animikie basin of the Lake Superior region contains another stratigraphic level with regionally extensive IFs. This level is stratigraphically above the 1850 Ma Sudbury impact ejecta layer and is older than the ~1830 Ma regional metamorphic event related to the Penokean orogeny (Cannon et al., 2010). These IFs are mineralogically and texturally different from the ~1880 Ma IFs of the Animikie basin, and were likely deposited in deeper waters, below fair-weather and, probably, storm-wave base. They are present in the: (1) Marquette Iron Range, Michigan (~60-m-thick Bijiki Iron Formation Member of the Michigan Slate containing siderite, chert, iron oxides and silicates; Cannon et al., 2010; Ojakangas, 1994; Ojakangas et al., 2001), (2) Iron River-Crystal Falls Iron Ranges (~15-m-thick chert–siderite slate of the Stambaugh Formation; James et al., 1968), (3) Gogebic Iron Range, Wisconsin (~47-m-thick IF of the Tyler Formation consisting of chert and siderite; Cannon et al., 2008; Schmidt, 1980), and (4) Mesabi Iron Range, Minnesota (~27-m-thick chert–siderite IF in the lower part of the Virginia Formation; Lucente and Morey, 1983). Deposition of these IFs might be genetically linked with submarine mafic volcanism in the Animikie basin based on spatial association with, for example, the Badwater Greenstone, but this relationship has not been documented in detail. The IFs commonly are interbedded with, or overlain by, black sulfidic shales, which likely record the development of euxinic conditions in the basin (cf. Poulton et al., 2004). Despite poor exposure, these iron deposits are easily traceable by magnetic anomalies. Their deposition indicates that anoxic and ferruginous seawater redox conditions were re-established in the Animikie basin after 1850 Ma, in association with mafic volcanism, although evidence for a shallow redoxcline in this case is absent and duration of these conditions was likely short and the extent of iron deposition probably was limited.

Thinly banded, silicate-facies IF of similar age, or slightly older, is present in the Pipe Formation of the Thompson nickel belt, Manitoba (Bleeker and Macey, 1996). However, GIFs are absent along the western and northern margins of the Superior craton, whereas they are developed continuously along the eastern and southern margins. Considering the

Figure 15  Distribution of iron formations and magmatic occurrences of the ~1880 Ma Circum-Superior event on the Superior craton. CSb, Cape Smith belt with ~1870 Ma volcanics of the Chukotat Formation; SIs, Sleeper Island sills; Bl, Belcher Group; SI, Sutton Inlier with metasediments and sills; Tb, Thompson belt; GI, Gunflint Iron Formation with volcanics; MRs, Marquette Range Supergroup with iron formation, volcanics, and sills; MI, Upper Albanel Group of the Mistassini basin with iron formation. Red star and circle locate inferred center and 1000 km radius of underlying mantle plume ~1880 Ma. Paleoproterozoic basins that contain 1.88 Ga iron formation and volcanics are shown in green, those that contain only iron formation are in red, and those that contain only volcanic units of this age as well as dikes and sills of this age in the interior of the craton are shown in black. Modified from Ernst RE and Bell K (2010) Large igneous provinces (LIPs) and carbonatites. Mineralogy and Petrology 98: 55–76.
occurrence of similar facies IFs in the Hudson Bay area and Sutton Inliers, it is plausible that a significant part of the Superior craton was submerged and sloped to the south during deposition of the Animikie IFs, due to uplift caused by the mantle plume then centered on the northwestern corner of the Superior craton (cf. Ernst and Bell, 2010). In this interpretation, IFs in the Sutton Inliers and the Hudson Bay area are trending thin, shallow-water facies of the intracratonic basin extending to the southern and eastern margins of the Superior craton (Figure 15).

A key unresolved question is why the ~1.88 Ga magmatism led to extensive deposition of IFs on several continents given that the long time intervals preceding and following this event had minimal IF deposition despite being similarly affected by the plume activity. In terms of simple mass balance, it might be that the flux of hydrothermally derived reducants (e.g., Fe, H₂S, and H₂) at 1.88 Ga was capable of overwhelming the redox state of the ambient deep ocean. Therefore, the volume of magma and the rate at which it was added by mantle plumes to the oceanic realm at this time could be a critical factor. If so, the question can be rephrased: what processes were responsible for generating the unique strength of the ~1.88 Ga mantle plume event? We emphasize here that this event corresponds with the early stage in supercontinent assembly, a topic discussed in more detail below.

9.18.7.3.7 Proterozoic age gap in major IF deposition

It is generally assumed that after ~1.85 Ga, major sediment-hosted IFs were not deposited for approximately 1.1 Gy (Huston and Logan, 2004; Isley and Abbott, 1999; Klein, 2005; Slack and Cannon, 2009). This distinctive pattern in the secular trend for IFs has been explained by a change in the deep ocean redox state from anoxic to oxygenated (Holland, 1984) or sulfidic (Canfield, 1998) conditions. In contrast, it has been recently proposed that the deep ocean was anoxic but not sulfidic (Planavsky et al., 2011; Poulton et al., 2010), or suboxic (Slack et al., 2007, 2009). The earlier suggestion by Holland (1984) for oxygenated deep-ocean conditions after ~1.88 Ga has fallen out of favor, but (arguably) the model has not been ruled out. An alternative model, suggesting that deep-ocean conditions were predominantly sulfidic over ~1.1 Ga of the Middle Proterozoic history (e.g., Canfield, 1998; Poulton et al., 2004), also has lost favor recently because practically all evidence for this model can be satisfied with locally developed euxinic conditions on continental shelves and in intracratonic basins (e.g., Planavsky et al., 2011; Scott et al., 2008; Slack et al., 2007). The emerging consensus is that the redox state of the deep ocean was variable over the time interval of ~1.85–0.75 Ga, but distinctly more reducing than that characteristic of the Phanerozoic (Planavsky et al., 2011; Poulton et al., 2010; Slack et al., 2007). The terms ‘ferruginous’ and ‘suboxic’ are not ideal for defining the redox state of the deep ocean during this period since the first was widely used for the Archean and the second has been used in different ways for modern and ancient conditions (Canfield and Thamdrup, 2009). More importantly, higher hydrothermal fluxes of metals, including Fe and Mn, should be expected if the oceans were at a low oxidation state and had a higher content of dissolved organic carbon (cf., Rothman et al., 2003) to effectively ligate metals.

The record of this hydrothermal flux might be found in Fe and base metal contents of mid-Proterozoic shales and in their isotope signatures. Poulton et al. (2010) argued that euxinic shales deposited on continental margins (possibly in areas corresponding to modern oxygen-minimum zones) and in intracratonic basins could have been a major sink for the hydrothermal iron flux to the Middle Proterozoic oceans, and were thus responsible for the absence or scarcity of Middle Proterozoic IFs. At present, this suggestion has not yet been quantitatively evaluated, but it would require a higher total Fe content in average shale during this time interval. Data in Kump and Holland (1992) do show that the average Proterozoic shale has more Fe than the average Phanerozoic shale but less than the average Archean shale. Considering that giant IFs were also deposited during the Archean, this trend in average content of Fe seems to suggest that other factors besides the presence of marginal euxinia controlled the abundance of IFs in the Middle Proterozoic. It has also been proposed that a rise in seawater sulfate level after ~1.88 Ga led to a decrease in the flux of the hydrothermal iron – shutting off the deposition of major IF (Kump and Seyfried, 2005). However, seawater sulfate levels decreased rather than increased in the Middle Proterozoic after the ~2.22–2.1 Ga Lomagundi carbon isotope excursion (Bekker and Holland, 2012), thus challenging this model.

Although large sedimentary IFs during the Middle Proterozoic (1.85–0.75 Ga) are indeed absent, several small IFs and iron-rich lithologies in sedimentary rock-dominated successions are known outside of the Animikie basin (see Appendix 1). Magnetite and siderite IFs of the Aok Formation in the Neoproterozoic Shaler Supergroup in the Duke of York and Brock Inliers of Victoria Island, northern Canada, were deposited at ~840 Ma before the onset of the oldest Neoproterozoic glacial events (Rainbird et al., 1994; Young, 1981). Older examples within this age range include the ~1.70 Ga Freedom Formation of the Lower Baraboo Series, Wisconsin, which contains in the lower part banded ferruginous chert interlayered with sideritic and kaolinitic slates 60–160 m thick (Leith, 1935; Van Wyck and Norman, 2004; Weidman, 1904) and the likely correlative magnetite–chert IF of the Tomiko terrane in Ontario (Easton, 2005). Additionally, the Chuanlinggou IF on the North China craton, a classic GIF deposit, also appears to be latest Paleoproterozoic in age (~1.7 Ga; Dai et al., 2004; Wan et al., 2003).

Noteworthy examples of small iron deposits that formed during this time gap in shallow-water settings occur in northern Australia in the Sherwin Formation and the Munyi Member of the Corcoran Formation, both within the ~1.49 Ga Maiwok Subgroup of the Roper basin, Northern Territory (Abbott and Sweet, 2000; Canavan, 1965) and the correlative Constance Range ironstone of the Train Range Member of the Mullera Formation in the South Nicholson basin, Queensland (Harms, 1965; Jackson et al., 1999). These iron-rich units consist of oolitic, pisolithic, and peloidal, trough cross-stratified ironstone and ferruginous siliciclastic rocks as much as 23 m thick (Abbott and Sweet, 2000; Harms, 1965). Unweathered samples show grapestones and ooids as much as 8 mm in diameter (Figure 5(b)) composed of hematite and magnetite with nuclei of quartz or oolitic intraclasts; chamosite and greenalite are present locally in a secondary siderite matrix (Abbott and Sweet, 2000). Beds within the ironstone units are bounded by
erosional surfaces with sharp compositional contrast between ooidal ironstones and overlying quartz-rich sandstones. The presence of intraclasts and peloids of iron-rich sediment in quartz-rich sandstone and clasts of ironstone within intraformational conglomerate (Figure 5(b)) strongly suggests that the sediments were originally iron-rich. Many ironstone intraclasts have a pisolithic coating, testifying to the primary deposition of iron oxyhydroxides in the sedimentary environment. The ooids and peloids likely formed at the redox boundary between shallow, oxidized and deep, anoxic waters. Notably, the Sherwin Formation and Munyi Member of the Corcoran Formation enclose black shales of the Velkerri Formation that were used previously to infer a long-term sulfidic state of the coeval deep ocean, based on sulfur isotope composition, iron speciation, and molybdenum isotope composition (Arnold et al., 2004; Shen et al., 2003). Leaving aside arguments whether available geochemical and geological evidence for the ocean redox state during the Mesoproterozoic is consistent with a pervasively euxinic deep ocean (e.g., Meyer and Kump, 2008; Poulton et al., 2010; Slack et al., 2007, 2009), this intricate relationship between ironstones and euxinic black shales strongly suggests that ferruginous and euxinic conditions alternated in the Mesoproterozoic intracratonic Roper and South Nicholson basins on geologically short timescales.

Oolitic ironstones are also developed in the ~1.38 Ga Xiahuayuan Formation of the Xiahuayuan region in Hebei Province of North China (Su et al., 2010) and in the poorly dated, late Neoproterozoic (but preglacial) Nizhne–Angara Formation of the Angara-Pit area, Siberia, Russia. IFs deposited at ~1490, 1368, and 767 Ma (see Appendix 1) broadly – within analytical error – correspond with ages for emplacement of LIPs at ~1460, 1380, and either 780 or 755 Ma (Ernst et al., 2008). Poor age constraints presently available for other IFs deposited during this time interval make it difficult to test their possible relationship with mantle plume events.

In addition to shallow-water, sediment-hosted IFs, numerous shallow-water exhalites are associated with sedimentary exhalative (SEDEX) and VMS deposits that formed during the late Paleoproterozoic and Mesoproterozoic. These deposits contain jasper, hematite, and magnetite layers having positive Eu and negative Ce anomalies (e.g., Hatton and Davidson, 2004; Lottermoser, 1989; Lottermoser and Ashley, 1996; Strydom et al., 1987). Although these VMS and exhalite deposits commonly formed within isolated to semi-isolated basins and thus should not be used to infer the redox state of the coeval deep ocean, they do provide strong evidence that shallow-water settings were oxygenated at that time. In contrast, mineralogical and geochemical data for deep-water, open-marine exhalites provide strong evidence for a low-oxygen (suboxic) state of the deep ocean during their deposition (Slack et al., 2007, 2009). Interestingly and surprisingly, large Mn deposits are absent during this time period with the single exception of the ~1.44 Ga Wafangzı Mn deposit in NE China (Maynard, 2010; Su et al., 2010). This trend, if real, suggests that a strong oxic sink for Mn might not exist during this time period in the oceans.

Even though IFs of this age range are of minor economic importance, their existence suggests that from ~1.85 to 0.75 Ga hydrothermal iron from deep-water oceanic settings was episodically delivered, although in rare cases and in generally small amounts, to shallow-water environments for the deposition of IF, ironstone, and iron-rich shale. In addition, within isolated to semi-isolated basins having strong volcanic and hydrothermal activity, the flux of reductants was strong enough to locally overwhelm the water-column redox state, thus leading to iron transport on a local scale. Open-marine, deep-water environments were in general sufficiently oxidized for iron to precipitate from hydrothermal plumes as iron oxyhydroxide. Combined, this record of the iron cycle is very similar to that in the Phanerozoic. Although it does not provide rigorous quantitative constraints on the deep-ocean redox state, the record challenges models for persistently oxic, fully anoxic, or sulfidic conditions within open-marine, deep-water settings. In summary, a low and yet variable redox state of the deep ocean is most consistent with this record.

### 9.18.7.3.8 Neoproterozoic manganese deposits and IFs

Manganese deposits and IFs of Neoproterozoic age are extensively but discontinuously developed in association with glacial deposits. In addition, iron-rich shales are relatively common in contemporaneous, glacially influenced successions (e.g., Young, 2002). The latter relationship was reemphasized by Canfield et al. (2008) to infer a return during the late Neoproterozoic to Archean-style anoxic, ferruginous ocean conditions. Significant iron and manganese accumulations in association with glacial deposits are present in a number of Neoproterozoic successions worldwide, although for other Neoproterozoic IFs a clear stratigraphic or temporal relationship with ice ages has not been established (e.g., Cabral et al., 2011; see also Appendix 1).

Presently available geochronological constraints and stratigraphic correlations imply that IFs are at least temporally related to the Sturtian (~715 Ma) and Marinoan (~635 Ma) glacialities, although some uncertainty exists regarding a number of Neoproterozoic glacial events, their ages and duration, and chemostratigraphically based correlations (e.g., Kendall et al., 2009). The Rapitan IF and Franklin igneous event on Victoria Island in Canada are both ~715 Ma (Heaman et al., 1992; Macdonald et al., 2010), supporting a genetic relationship among volcanism, mantle processes, and deposition of IF at least on the basin scale. Neoproterozoic IFs are typically enriched in Mn and P and have high Co, Ni, and Cu contents (Halverson et al., 2011; Klein and Beukes, 1993; Planavsky et al., 2010a,b; see also Chapter 9.11). Highly positive sulfur isotope values in shales and manganese formations overlying Neoproterozoic glacial diamictites have been related to Rayleigh distillation in low-sulfate oceans, which were highly susceptible to the development of anoxic ferruginous conditions and hydrothermal delivery of iron and manganese into shallow basins during mantle plume breakout events (Liu et al., 2006). Importantly, IFs were deposited during deglaciation as indicated by the presence of dropstones in IFs and interlayering with tillites. Poor age constraints for other Neoproterozoic IFs do not warrant correlation with Neoproterozoic mantle plume events, although several such events occurred during the late Neoproterozoic (e.g., Ernst and Buchan, 2001). However, almost all Neoproterozoic IFs are spatially and temporally linked with submarine mafic and felsic volcanic units. Some basins hosting IFs of this age also contain coeval VMS deposits (e.g., Buh et al., 1992; Cabral et al., 2011), suggesting a proximal hydrothermal source of metals. Most of these IFs were...
Neoproterozoic IFs generally comprise laminated and nodular hematite, massive magnetite, hematitic mudstone, and jasper (Figure 16). Lenticular and nodular chert and jasper are less common, but are present within laminated jasper beds (Yeo, 1981). Banded cherts, similar to those in Archean and Paleoproterozoic IFs, are absent. Furthermore, the thickness of Neoproterozoic IFs is highly variable over relatively short distances. GIFs and intraformational conglomerates containing hematite pebbles developed locally at the top of the Rapitan Iron Formation and in the Jacadigo Group, Brazil, but coated grains are uncommon (Klein and Beukes, 1993a; Klein and Ladeira, 2004). REE patterns for Neoproterozoic IFs have either no or slightly positive Eu anomalies, and either no or slightly negative Ce anomalies (Derry and Jacobsen, 1990; Fryer, 1977; Halverson et al., 2011; Klein and Beukes, 1993a; Klein and Ladeira, 2004; Liu et al., 2006; Lottermoser and Ashley, 2000), likely indicating a high degree of dilution of locally derived, hydrothermal fluid by mildly oxidized seawater. Models that are generally accepted for Phanerozoic manganese deposits (see Chapter 9.11) are probably also applicable to the origin of Neoproterozoic IFs. These models infer anoxic conditions with enhanced submarine volcanism in the deeper parts of isolated to semi-isolated basins and manganese precipitation occurring at the redox boundary on the shallow margins of the basins. Another factor, in addition to mantle plume events and tectonics, which could have contributed to deposition of IF, is dramatic sea-level fall during ice ages. Sea-level fall would decrease the overlying hydrostatic pressure and shift equilibrium in the seafloor hydrothermal systems toward higher Fe/H₂S ratios, thus promoting larger Fe fluxes into the oceans (Kump and Seyfried, 2005). In sum, although the occurrence of Neoproterozoic IFs supports long durations of glaciations and the existence of redox-stratified basins, they do not require extreme snowball Earth conditions in order to form.

9.18.7.3.9 Phanerozoic ironstones, anoxic events, and VMS deposits

The most prominent peaks in deposition of Phanerozoic ironstones occur in the Ordovician–Devonian and Jurassic–Paleogene (Figure 17). The ironstones generally are only several meters thick, rarely reach 25 m in thickness, and extend for over 1000 km along ancient continental margins in Fennoscandia and the Himalayas where they demarcate areas of past upwelling (e.g., Garzanti, 1993; Sturesson, 2003). The ironstones are composed largely of iron oxide ooids (goethite and limonite), with smaller amounts of Fe-silicates (chamosite, berthierite, and glauconite) and Fe-carbonates ( siderite), typically as cements. In contrast to most Proterozoic IFs and early Paleozoic hydrothermal deposits, ironstones have very little chert, but commonly are

Figure 16  Rapitan Iron Formation at Cranswick River, Mackenzie Mountains, Northwestern Territories, Canada. (a) Nodular and banded jasper interlayered with hematite bands and overprinted by anastomosing hematite. (b) Jasper nodules and lenses in massive hematite. (c) Banded jasper–hematite iron formation with dropstone overprinted by anastomosing hematite. (d) Jasper nodules and lenses in massive hematite. Photographs are courtesy of E. Turner.
enriched in manganese and phosphorus. However, in some cases, ironstones are associated with chert deposits (Borchert, 1965; Franceschelli et al., 2000; Glenn and Arthur, 1990).

Many ironstone deposits contain stilpnomelane, volcanic glass shards, and other tuffaceous material implying contemporaneous volcanism. Typically, bulk analyses of oolitic ironstone, when normalized to shale composites or local shale, have minor negative to positive Ce anomalies, are enriched in light REE with respect to heavy REE, and lack significant Eu anomalies (e.g., Bhattacharyya and Cretar, 1993; Sturesson, 2003).

Ironstones are common to time intervals characterized by negative seawater Sr isotope excursions (Sturesson, 2003), peaks in the formation of VMS deposits (cf. Franklin et al., 2005; Peter, 2003), large volcanic- and sediment-hosted Mn deposits (Maynard, 2010), extensive ocean anoxia, greenhouse conditions, volcanic events, emplacement of LIPs, and quiescence in Earth’s magnetic field (e.g., Garzanti, 1993; Sturesson, 2003; Van Houten and Arthur, 1989). Ironstones are typically thought to have been deposited during periods with low sedimentation rates at the beginning of a sea-level rise, as suggested by their position in Transgressive System Tracts (Figure 6(b); Burkharter, 1995; Maynard and Van Houten, 1992; Taylor et al., 2002) and commonly are topped by hardgrounds, thought to represent the maximum

Figure 17  Secular distribution of Phanerozoic oolitic marine ironstones (from Petranek and Van Houten, 1997) compared to those of marine anoxic conditions and VMS deposits (from Bekker et al., 2010) and a number of sedimentary Mn deposits (from Maynard, 2010). Data for VMS deposits and for sedimentary Mn deposits are compiled for time intervals of 25 Ma. Marine anoxic conditions are shown for inferred global periods of anoxia by black bars and more limited periods of anoxia by gray bars; the latter reflecting less-widespread, regional black shale facies. Data for anoxic conditions during Cambrian through Jurassic periods are from Arthur and Sageman (1994), for Cretaceous from Leckie et al. (2002), and for Cenozoic from Jacobs and Lindberg (1998).
flooding surface. They are also common to upwelling areas where phosphates were deposited. Collectively, these diverse associations have been linked with mantle superplume events (e.g., Gazzanti, 1993). Furthermore, ironstones are related to time periods when seawater was saturated with calcite, rather than with aragonite or high-Mg calcite, and when the so-called ‘calcite seas’ developed, which correspond with times of rapid seafloor spreading and high rates of Mg removal from seawater (Maynard, 1986; Stanley and Hardie, 1998).

The formation of berthierite and siderite in shallow, well-agitated marine environments requires weakly reducing but non-sulfidic conditions in the water column and sediment pore waters. However, berthierite and hematite ooids co-occur in ironstones, even forming discrete laminae within individual ooids, thus indicating fluctuating redox conditions during either deposition or early diageneis, but prior to compaction. In addition, glauconite, an abundant sedimentary aluminosilicate mineral with Fe(III) substituting for Al(III) and lower Eh requirements for precipitation than berthierite, is virtually absent in oolitic ironstones (Maynard, 1986) and does not co-exist with chamosite but occurs in deeper-water facies equivalents (e.g., green shales; Hunter, 1970). Where sedimentary facies analysis has been performed, the following sequence of facies and mineralogical changes from shallow, oxidized waters to deep, reduced waters is evident: hematitic oolitic ironstone; chamositic oolitic ironstone; glauconitic, illitic, and chloritic green shales; and, finally, organic matter-rich, sulfidic shales.

The genesis of Phanerozoic ironstones and the iron source for these deposits remains controversial. Extreme degrees of weathering on peneplaned continents linked with high sea-level stands during dispersion of supercontinents are commonly inferred (e.g., Van Houten and Arthur, 1989). However, neither sea-level highstand nor supercontinent dispersal seems to be a prerequisite for the deposition of Phanerozoic ironstones (Van Houten, 1985). Furthermore, because iron is highly insoluble in the presence of even traces of dissolved oxygen, this model implies that iron was delivered from the continents in solution with slightly acidic river waters or groundwaters, in colloidal form, or complexed with organic ligands, and the iron was precipitated in deltas or along shorelines on mixing with seawater having higher pH and ionic strength (Castano and Garrels, 1950; Huber and Garrels, 1953). However, considering the fast rate of iron precipitation on mixing with seawater, a corollary of this model is that ironstone deposition should be centered on a point source as commonly found in subterranean estuaries showing a well-defined ‘iron curtain’ near the sediment–seawater interface (Charette and Sholkovitz, 2002; Rouxel et al., 2008a,b), rather than along the full extent of the shoreline with little variation in thickness. An alternative model relates ironstones to submarine weathering of volcanic ash (e.g., Sturesson, 2003), even though many ironstone-bearing units lack volcanic ash beds. Reworking of pedogenic ferruginous pisoids into the marine environment has also been suggested, but it does not explain their restricted stratigraphic position, global secular trend, low Al content in goethitic and hematitic oolites (e.g., Maynard, 1986), or εNd(t) values above those of proximal crustal sources (Sturesson et al., 1999), indicating some contribution to REE budget from juvenile sources.

Aller et al. (1986) developed a model of an early diagenetic iron source from anoxic sediments into the water column as originally advocated by Borchert (1965). Aller et al. (1986) proposed a model for ironstone deposition based on modern Amazon deltaic sediments in which high amounts of organic matter loading, anoxia at the sediment–water interface, and intense physical reworking promote the re-oxidation of pore-water iron. These processes in the Amazon River allow for significant burial rates of reactive iron phases, such as ferric and mixed-valence authigenic iron minerals, despite the presence of abundant dissolved sulfate (Aller et al., 1986). The model is consistent with the depositional setting of ironstones, although it does not explain their episodic secular distribution or much more pronounced iron enrichments relative to modern Amazonian deltaic sediments. Nonsteady diagenetic reworking of preexisting metal-enriched (anoxic) shales is a potential mechanism for ironstone deposition that has not been explored in any detail, but would provide an explanation for the close association of ironstones with periods of marine anoxia.

Cotter and Link (1993) were among the first workers to suggest that iron was supplied from deep-water, anoxic settings. Broad correspondence of the secular distribution of Phanerozoic ironstones with that of VMS and volcanic- and sediment-hosted Mn deposits (Figure 17) offers additional insight into ironstone deposition and suggests that the iron originated in coeval seafloor-hydrothermal systems. Along these lines, we suggest that deposition of major Phanerozoic ironstones is mechanistically linked to times of global ocean anoxia during mantle superplume events. Enhanced volcanic and hydrothermal activity led to extensive formation of VMS deposits, higher sea level, and contributed to short-term, ocean-wide anoxia. Under anoxic conditions, hydrothermal and diagenetic iron could have been transported by upwelling currents, together with phosphorus and manganese, onto the shelf where the iron was oxidized biologically or abiologically at a shallow-water dynamic redoxcline. Cretaceous anoxic events also have been genetically linked to extensive submarine magmatism based on geochemical evidence (Turgeon and Creaser, 2008) and geological arguments (Kerr, 1998; Sinton and Duncan, 1997). The model implies that even if sulfidic conditions persisted throughout the oceans, more hydrothermal iron than in the modern ocean was transported from the deep ocean, as either ligand bound or nanoparticulate phases. However, even under widespread anoxic conditions, sulfide formation in the deep oceans could have been limited by organic matter availability. Anoxic conditions with low sulfide levels could have allowed for transport of dissolved iron onto the continental shelves. This model is supported by iron isotope data that point toward hydrothermal iron delivery onto the portions of the upper slope in the Atlantic Ocean during Cretaceous OAE 2A (Owens et al., 2012).

9.18.8 Controls on IF Deposition

The above discussion allows us to discriminate the major controls on IF deposition. Although in some cases it is difficult to resolve whether an IF was deposited proximally or distally to submarine volcanism, IFs hosted in volcanic-dominated successions are in general more common during Earth’s early
history (>2.3 Ga) and probably reflect a higher mantle heat flux. As time progressed, relatively small sediment-hosted IFs appeared (or were preserved) in the geologic record starting around 3.2 Ga, but did not reach significant thickness and extent until 2.6 Ga. An intriguing cause-and-effect relationship underlies fundamental environmental changes between 2.5 and 2.3 Ga including emplacement of the first true LIP, a peak in the deposition of distal, sediment-hosted IFs, atmospheric oxidation, and Paleoproterozoic glaciations. The response of the biosphere to the superplume event may have led to changes in surface environments. Following atmospheric oxidation at ~2.4 Ga, iron and manganese deposition was localized in basins characterized by restricted circulation and intense submarine volcanism.

The later peak in IF deposition at ~1.88 Ga was also closely linked in time with a significant mantle plume breakout event that affected the Superior craton, among others. We infer that at least on the scale of basins that surrounded the Superior and Yilgarn cratons, the redox state and sulfate content of seawater were sufficiently low that the hydrothermal flux of iron and other reductants (e.g., Mn, H2S, and H2) overwhelmed the oxidizing potential of seawater. It is likely that very low marine sulfate concentrations at this time, similar to those during the Archean, would have increased the iron/sulfide ratio in emanating hydrothermal fluids, which also would have promoted iron-replete conditions in the ocean (Kump and Seyfried, 2005) and deposition of major IFs at the dynamic redoxcline in shallow waters (Planavsky et al., 2009). High atmospheric pCO2 and, consequently, surface temperatures during the mantle plume breakout event could have led to stagnant circulation and the decreased solubility of dissolved oxygen in surface waters, resulting in anoxic conditions within these basins, if not in the global ocean. In addition, mantle plume breakout events generate higher sea level (Condie et al., 2001), leading again to sluggish ocean circulation promoting anoxia on a basin to global scale.

Our analysis agrees with previous studies (e.g., Isley and Abbott, 1999) that the deposition of giant IFs corresponded closely in time with major mantle plume events. Therefore, rather than linking the termination of deposition of large, economic IFs at ~1.85 Ga and the expansion of sulfidic conditions to changes in surface conditions (e.g., Poulton et al., 2004), we propose that these secular changes, at least in the Animikie basin, were a consequence of several contemporaneous processes but foremost the waning of a mantle plume event and a decrease in the hydrothermal iron flux.

Recurrent associations in the Precambrian between superplume events and IF deposition have been known for some time (Barley et al., 2005; Condie et al., 2001; Isley, 1995). Recently, their secular relationship in the Precambrian and Phanerozoic with VMS deposits was also established (Bekker et al., 2010). The link is obvious: hydrothermal systems were more extensive, vigorous, reducing, and metal-rich during LIP emplacement. As a result, larger volumes of metals were leached from submarine volcanic rocks and vented to the ocean floor, where massive sulfide deposits formed; consequently, ocean redox state was lowered by an enhanced flux of hydrothermally derived reductants and Fe with Mn were delivered by plumes to shallow-water settings where they precipitated, forming iron- and manganese-rich sediments. Peaks in the tonnage of VMS deposits are typically associated with time intervals when supercontinents were assembled, because VMS mineralization hosted by bimodal volcanic rocks in back-arc basins has a high preservation potential in the rock record. Considering that mantle plumes are relatively common throughout Earth’s history, it remains uncertain why superplumes and associated peaks in diverse mineral deposits are genetically linked with the early stages in the supercontinent assembly. The association is, however, striking and repetitive at ~2.74–2.69, 2.5–2.45, 2.05–2.06, 1.88, 1.1, and 0.5 Ga when the Kenorland, Vaalbara, Nuna, Zimvaalbara–São Francisco, Rodinia, and Gondwana supercontinents were assembled, respectively (cf. Huston et al., 2010). It is proposed herein that external ocean closure during the early stage of supercontinent amalgamation dramatically changed the heat budget of the mantle, leading to mantle overturn, superplume events, and eventually the development of a new mantle convection pattern (Figure 18). As a result, superplume-initiated rifting at the time of external ocean contraction allows plate tectonics to persist without interruption in the aftermath of the supercontinent assembly.

Whereas Fe–Si compounds, the likely protolith for pre-GOE IFs, formed predominantly in the deeper parts of basins and were widely redistributed by density currents, post-GOE IFs record the upwelling and oxidation of ferrous iron, precipitation on the shelf and, later, transport by storm events and wave currents back into the basin. GIF first appeared at ~2.6 Ga or earlier, but their predominant mineralogy was not iron oxides until shortly after the GOE.

Whereas VMS deposits show a peak in global tonnage at ~1.88 Ga, coeval with the peak in deposition of GIF, VMS deposits are puzzlingly absent during the 2.50–2.45 Ga mantle plume event when large tonnages of IF accumulated (Figure 14). Potentially, long-term global tectonic processes played a role. The hydrothermal flux of iron likely was derived predominantly from mid-ocean ridges at that time and corresponding
VMS deposits that formed in this setting were likely recycled to the mantle. Differences between IFs deposited before and after the GOE, in terms of original textures and depositional settings, imply potentially different mechanisms for iron precipitation. Similarly, assembly of the supercontinent Rodinia at \( \sim 1.2–1.1 \) Ga, corresponding with a small peak in VMS tonnage and a massive mantle plume event that affected North America, southern Africa, and western and central Australia (Ernst et al., 2008), is not reflected in the record of IFs. Open-marine, sedimentary successions of this age are rare, which could at least partially explain the lack of IFs of this age in the rock record. As stressed above, the development of a more sulfate-rich ocean with respect to that during the Archean and early Paleoproterozoic would have lowered iron/sulfide ratios in hydrothermal fluids (Kump and Seyfried, 2005), decreasing the potential for deposition of massive iron deposits.

Reasons for the 1.1 Gy gap in the deposition of large IFs in the Proterozoic remain uncertain. Although we note several exceptions to this IF hiatus, these are small units that predominantly occur within restricted basins. If the temporary cessation of plate tectonics (cf., Silver and Behn, 2008) was responsible for the gap, why did major mantle plume events and, more importantly, VMS deposits formed during this time period (Figure 14)? Mantle plume events do not necessarily require plate tectonic processes, but VMS mineralization does need a submarine heat source in an oceanic environment, or at least marine basins that developed on rifted continental or oceanic crust (e.g., Franklin et al., 2005; Huston et al., 2010).

The secular distribution of IFs clearly deserves more attention. Although precise geochronology has been very successful in dating Precambrian successions, including host strata to IFs and VMS ores, some major economic iron deposits are still poorly dated; knowing their ages might critically influence our understanding of past oceanic and atmospheric redox states, as well as large-scale evolution of the Earth system. Among these poorly dated deposits (Appendix 1) critically in need of high-precision geochronology, are Paleoproterozoic examples including Krivoy Rog in Ukraine, the KMA in Russia, and the Caué iron Formation in Brazil; a number of Mesoproterozoic and Neoproterozoic IFs in Russia, China, and Australia; and Archean IFs in India.

Neoproterozoic IFs are interbedded with glacial deposits, but it is unlikely that snowball Earth conditions alone led to their deposition. Long-term ice cover likely played an important role in lowering the redox state of deep seawater and decreasing marine sulfate concentrations, by inhibiting or slowing down atmosphere–ocean exchange, continental pyrite oxidation, and riverine transport of sulfate to the oceans. However, the geological setting of the giant Rapitan iron Formation, as well as other Neoproterozoic IFs (Appendix 1), provides a strong case for temporal and genetic connection to submarine volcanism. Therefore, at least three factors were likely critically important in the genesis of Neoproterozoic IFs: (1) seawater redox state, (2) an enhanced hydrothermal flux of iron during mantle plume breakout events, and (3) basin configuration including at least partial physical separation from the global ocean.

Close relationships among IFs, VMS deposits, seawater anoxia, and mantle plume events during the Precambrian provide a key framework for interpreting the origin of Phanerozoic ironstones. In contrast to many of the previous studies of ironstones, we emphasize similarity in the secular trends of ironstones, VMS, and sediment-hosted Mn deposits, organic matter-rich shales, and intervals of enhanced submarine volcanism during mantle plume breakout events in the Phanerozoic (Figure 17). We propose that some Phanerozoic ironstones formed during anoxic events when hydrothermal iron was delivered to shallow shelf settings either in isolated basins or on open continental margins.

The biosphere also played an important role in the genesis of IFs. Not only was biotic activity involved in direct or indirect precipitation of iron, but it also influenced the seawater redox state and composition. For example, emergence and evolution of the biological silica sink during the Precambrian and Phanerozoic influenced scavenging efficiency of phosphorus and metals onto ferric oxyhydroxides. However, even precipitation and oxidation mechanisms for iron-rich compounds that were the precursor sediments of Archean IFs are not unequivocally established. The problem remains that it is neither obvious what the texture and mineralogical composition of those sediments were, nor where and how the original sediments were deposited prior to their resedimentation into the basin realm. Certainly, the widespread presence of graded beds within BIF supports a model in which some of the original sediments were granular, as it is seemingly impossible to produce graded beds from amorphous iron-rich clays, iron-rich gels, or chemically deposited varves – the traditionally inferred precursor sediments to BIF. Whereas recognition that classic BIF of the Dales Gorge Member, in the Hamersley Province of Western Australia, are density-current deposits is significant, that recognition has made interpretation of the precursor sediments even more difficult, because the original depositional site is unknown. The same problem exists for GIFs, because they are essentially clastic sedimentary rocks.

More broadly, our model for IF genesis provides a new perspective on the secular evolution of the ocean redox state. We suggest that submarine volcanism was episodically responsible for generating extensive ocean- and basin-scale anoxia by the hydrothermal venting of significant fluxes of reductants such as \( \text{H}_2, \text{H}_2\text{S}, \text{Fe}^{(II)}, \) and \( \text{Mn}^{(II)} \). This perspective applies to both Precambrian and Phanerozoic environments, and provides a framework for better understanding ocean evolution and the origin of sedimentary mineral deposits.

9.18.8.1 Influence of Hydrothermal Processes on Ocean Composition and Organic Productivity

Late Archean and Paleoproterozoic IFs are commonly interbedded with or adjacent to organic matter-rich, sulfidic shales. Similar shale facies appear to be underrepresented in the Archean record, beyond stratigraphic intervals that were deposited during mantle plume breakout events (Condie et al., 2001). This association to date has not been explained satisfactorily. During the Phanerozoic, the link between black shales and mantle plume
events commonly has been explained by sea-level rise and a higher flux of reductants into the ocean. The reductant flux is tied to anoxia, which favors organic matter preservation. This explanation does not apply, however, to the Archean oceans that were persistently anoxic (e.g., Planavsky et al., 2010a,b).

An important perspective is provided by considering the carbon isotope mass balance relevant to Archean surface anoxic conditions (e.g., Bekker and Holland, 2012). In the modern world, when the sedimentary cover is recycled to the oceans, organic carbon is oxidized and inorganic carbon is dissolved, making both available for biological processing. However, under anoxic surface conditions in the Archean, organic carbon was likely redeposited and hence was not biologically recycled. Including this aspect into the carbon isotope mass balance (cf. Bekker and Holland, 2012) requires that burial of ‘new’ organic carbon was less than 5% of the modern burial rate of organic carbon. As a result, the amount of oxygen released by the biological carbon cycle was also very low. We infer that restricted availability of critical nutrients, such as Mo, could have limited biological productivity and consequently organic carbon burial to these low levels.

In order to explain the origin of organic matter-rich shales associated with IFs, we infer high primary productivity that was driven by high nutrient fluxes, sourced from either the deep ocean or the continents. In nearly all modern aquatic systems, primary production of organic matter is limited by phosphorus or bioavailable nitrogen (e.g., Redfield, 1958), but trace metals can also colimit or even limiting. Phosphorus was asserted to be the ultimate limiting nutrient in the Archean oceans, given a high flux of iron oxides that removed phosphate (Bjerrum and Canfield, 2002). However, during the Archean and Paleoproterozoic, especially during peak times of IF deposition, high dissolved silica concentrations could have prevented iron oxyhydroxides from being a major dissolved phosphate sink (Konhauser et al., 2007b). Additionally, high levels of bicarbonate during the Precambrian (cf. Grotzinger, 1990) would have inhibited the precipitation of carbonate fluorapatite (CFA) by greatly increasing its solubility. The possibility of inhibited CFA formation is particularly important; in the modern oceans, this flux accounts for the vast majority (60–80%) of phosphorus burial (e.g., Ruttenberg and Berner, 1993). Furthermore, low burial rates of ‘new’ organic matter and high efficiency of its remineralization in an anoxic ocean lacking mechanisms for adequate export of organic carbon to sediments likely resulted in a low burial flux of phosphate with organic carbon. Combined, these factors point toward a dramatically different Archean than the modern phosphorus cycle and relatively higher, rather than lower, dissolved phosphorus concentrations in Archean seawater with respect to modern seawater (Planavsky et al., 2010a,b).

If phosphorus content was not the controlling factor for Archean primary productivity, could nitrogen be the culprit? In the Archean anoxic ocean, dinitrogen was seemingly fixed by diazotrophic cyanobacteria and ammonium assimilation was also likely common (Garvin et al., 2009; Glass et al., 2009; Godfrey and Falkowski, 2009). Both nitrogen fixation and ammonium assimilation are operated by enzymes, which are dependent on bioessential trace metals such as Fe, V, and Mo, as metal cofactors (Glass et al., 2009). Both the V and Mo dissolved loads were generally low or negligible in early oceans (Lalonde et al., 2011; Scott et al., 2008), because under an essentially anoxic atmosphere these metals are insoluble. Thus, Mo limitation was likely in the Archean oceans due to anoxic redox conditions on land, just as iron is locally limiting in the modern oceans because of a predominance of the oxic water column. Iron concentrations, in contrast, were high and iron was likely utilized by diazotrophs on early Earth for nitrogen fixation and ammonium assimilation (Glass et al., 2009). Iron availability enhances nitrogen fixation and photosynthesis in modern environments (e.g., Berman-Frank et al., 2001, 2007); however, Fe–Mo nitrogenase is 100 times as efficient as Fe–Fe nitrogenase in dinitrogen fixation (Zerkle et al., 2006). Therefore, nitrogen limitation (via trace metal colimitation) of primary productivity was much more important than phosphorus limitation in the early oceans. It is possible that nitrogen stress was partially alleviated during mantle plume breakout events when IFs were deposited, since bioessential trace metals (e.g., Mo and V) can be sourced from hydrothermal fluids (e.g., Kuhn et al., 2003; also see Chapter 8.7). It is difficult to quantitatively estimate an increased level of organic productivity due to enhanced hydrothermal metal delivery to the Archean oceans during mantle plume events, considering the absence of positive carbon isotope excursions in the Archean carbonate record.

Related changes in the atmosphere and terrestrial settings are also relevant. The volcanism that occurred during mantle plume events delivered large amounts of CO2 to the atmosphere, and enhanced chemical weathering under resulting greenhouse conditions would have generated a large flux of dissolved bicarbonate and nutrients, such as phosphorus and, to a lesser extent, nitrogen, to the oceans. This flux, by itself, might account for high organic productivity and high absolute burial rates of organic carbon, while the relative burial rate of organic carbon remained constant (cf. Aharon, 2005). Volcanic SO2 flux to the atmosphere was also scaled up during the mantle plume breakout events, and a larger sulfur flux was delivered to shallow-marine environments. As a result, iron oxyhydroxides precipitated in deeper-water environments and carbonate-facies IFs mark a transition zone to highly productive shelves on which organic matter-rich and sulfidic shales were deposited, utilizing reactive iron delivered by iron-depleted hydrothermal plumes (Figure 19; Bekker et al., 2009; Rouxel et al., 2005). These sulfidic shales could have scavenged seawater Mo that was delivered to the oceans hydrothermally or, possibly, through oxidative weathering, providing an explanation for Mo enrichments observed in the 2.5 Ga Mount McRae Shale (Anbar et al., 2007) and shales proximal to the ~2.7 Ga Kidd Creek VMS deposit (Watanabe, 2002). Some of the hydrothermally delivered Mo was likely coprecipitated with iron oxyhydroxides in IFs, contributing to Mo isotope fractionations observed in the Mount McRae Shale (Duan et al., 2010). The carbonate-facies IFs reflect diagenetic reactions in settings where microbial Fe(III) reduction was coupled with
remineralization of organic matter (Walker, 1984). Less productive shallow-water shelves in the Archean oceans were depositional sites for carbonates as well as silicate- and carbonate-facies GIFs during the mantle plume events (Beukes and Klein, 1990).

In contrast, post-GOE Paleoproterozoic IFs are predominantly granular, shallow-water, oxide-facies IFs that grade to finely laminated, sub-storm base (but still shelfal) Fe-silicate and carbonate-facies types. These extensive GIFs are related to rapid oxidation of iron delivered by upwelling iron-rich waters.
to nearshore, high-energy environments and to subsequent sedimentary reworking of these precipitates. Deeper, in areas of high organic productivity on shelf, organic matter-rich and sulfidic shales were deposited in association with GIF during mantle plume events after the GOE (Figure 19).

Combined, these data suggest that after the GOE, there was a dynamic redoxcline between continuously oxic shallow waters and ferruginous deeper waters on the continental shelves, which was the typical depositional site for IFs (Bekker et al., 2010). In contrast, pre-GOE IFs were predominantly deposited in deeper-water settings having either a dynamic chemocline separating anoxic ferruginous deep waters from anoxic sulfidic shallow waters where black shales were deposited on highly productive shelves or a gradual transition to shallow-water, reduced-facies GIF in low productivity areas around submerged cratons (Figure 19). This fundamental difference in the distribution of IF facies is linked to oxygenation of the shallow ocean and the formation of a discrete redoxcline. The difference between pre- and post-GOE IFs is recorded not only in their sedimentary facies, but also in a much larger range of iron isotope fractionations in pre-GOE IFs (Planavsky et al., 2012; Rouxel et al., 2005). Without a fully oxygenated surface layer, partial oxidation was common in upwelling water masses during the Archean, resulting in Rayleigh fractionation during Fe removal and consequently a much larger range of sedimentary iron isotope values.

### 9.18.8.2 Implications for Atmospheric Oxidation

The emerging view is one of coupled enhanced primary productivity and deposition of IFs initiated by mantle plume events. If this model is valid, then it has implications for the rise of atmospheric oxygen. Once extensive oxidative continental weathering began, it provided abundant Mo to the oceans to activate the much more efficient Mo-nitrogenase enzyme (with respect to Fe–Fe nitrogenase enzyme). In addition, an enhanced hydrothermal Mo flux was likely delivered to the oceans during deposition of IFs at ~2.48–2.45 Ga. With more Mo available, nitrogen fixation would have flourished, leading to increased primary productivity by photosynthetic cyanobacteria. In turn, higher net oxygen production not only would have provided the oxygen required to remove iron from seawater, but also would have increased the terrestrial flux of dissolved Mo to the oceans. Under these conditions, Fe–Mo-nitrogenase would have replaced the less-efficient Fe–Fe nitrogenase, removing nitrogen limitation and leading to irreversible atmospheric oxidation. Paradoxically, in this model the rise of atmospheric oxygen is brought on by an enhanced flux of reductants from the mantle. Although IFs were deposited and LIPs emplaced before then, both processes are not comparable in scale or extent with those at 2.5–2.45 Ga (Barley et al., 2005; Condie et al., 2001; Heaman, 1997; Isley and Abbott, 1999).

Consistent with the above model, the oldest positive carbon isotope excursion is found in carbonates of the ~2.48 Ga Tongwane Formation, South Africa, which sits above an IF (Bekker et al., 2001). Mass-independent sulfur isotope fractionations, a proxy for the atmospheric redox state, declined shortly after deposition of the ~2.5–2.45 Ga IFs. Coincident with this shift is the onset of glaciations related to the GOE and the oxidation of atmospheric methane, which was an important greenhouse gas during the Archean (Bekker and Kaufman, 2007).

### 9.18.9 Euxinic Conditions Induced by Shift in Dissolved Fe/S Ratio of Seawater due to Iron Oxidation

Although ferruginous conditions appear to have been dominant in Archean oceans, there are several documented cases of euxinia or free sulfide existing in the coeval water column (e.g., Anbar et al., 2007; Kendall et al., 2010; Reinhard et al., 2009; Scott et al., 2008, 2011). Evidence for euxinia indicates that total dissolved S/Fe ratios commonly were greater than the pyrite stoichiometry of two (FeS₂). These results seem surprising, given a general acceptance of low sulfate and high iron concentrations in Archean seawater. However, even at very low sulfate concentrations it might be possible to develop euxinia. Low-sulfate, iron-rich oceans having spatial heterogeneities in dissolved S/Fe ratios are prone to locally establish euxinic conditions. Euxinia can develop in such oceans where Fe(II) oxidation effectively strips dissolved Fe²⁺, creating a water mass with a relatively high S/Fe ratio. This water mass might retain a high dissolved S/Fe ratio when it later feeds a coastal setting, assuming the common case where rates of isopycnal mixing are much greater than those of diapycnal mixing. Because microbial Fe(III) reduction requires particulate contact, the reaction is expected to be very sluggish in the water column and therefore allows iron oxyhydroxides to sink. Ferric–ferrous iron interactions, Si coatings on Fe oxyhydroxides, and low Corg content in the water column also likely prevented Fe(III) reduction above the sediment–water interface. If a water mass in the photic zone has relatively high S/Fe ratios, it would become euxinic given a sufficient organic matter flux to fuel sulfate reduction. This scenario, in light of modern circulation patterns such as that feeding the Peru upwelling zone, could develop in an equivalent position to the modern OMZ euxinia in a low-sulfate, low-oxygen ocean regardless of the dissolved sulfate/Fe²⁺ ratios in deep waters. Although the Archean oceans are typically envisioned to be iron-replete and sulfide-poor, euxinia was likely relatively common in many settings having high rates of sulfate reduction. This is also consistent with the highly negative Fe isotope values in sulfidic black shales, which suggest extensive iron depletion through partial oxidation in the water column (Rouxel et al., 2005).

Changes in the atmosphere and in terrestrial settings linked with mantle plume events were also likely relevant to the local development of euxinic conditions in Archean oceans. Associated volcanism would have supplied large amounts of H₂S/SO₂ and CO₂ to the atmosphere and to marine environments. Enhanced chemical weathering under greenhouse conditions likely delivered nutrients to the oceans and, at the same time, sulfur compounds to shallow marine environments. As a result, iron oxyhydroxides would precipitate in deeper-water environments, whereas sulfidic conditions would develop in shallow-water environments where iron-depleted hydrothermal plumes delivered reactive iron (Figure 19a; Bekker et al., 2009; Rouxel et al., 2005). A dynamic chemocline would therefore exist between anoxic ferruginous deep waters and anoxic sulfidic shallow waters, persisting until hydrothermal iron and atmospheric/terrestrial sulfur fluxes subsided, which would have...
switched the ocean back to an unstratified, anoxic state. Expansion of the mixing zone between hydrothermally derived metal and atmospherically derived sulfur compounds during superplume events would have generated a high productivity zone ideally suited for increased burial of organic carbon.

**9.18.10 Research Perspectives and Future Directions**

Progress in research on IFs is closely linked to advancements in the understanding of surface and submarine-hydrothermal processes in the Precambrian and advances in geochronology, petrology, analytical geochemistry, and chemical oceanography. Early studies from the 1940s to the 1960s were directed mainly to map the distribution of this newly recognized iron resource in Precambrian terranes, and to provide age constraints and a stratigraphic framework. The discovery of Precambrian IFs immediately initiated speculations among geochemists about the mode of their origin based largely on application of thermodynamics to aqueous chemistry of the ancient oceans. Classical petrographic and metamorphic petrology studies focused on establishing paragenesis and mineral composition of the protolith sediments of IFs. Starting in the early 1970s, sedimentological facies analysis followed by basin analysis and sequence stratigraphy were incorporated into studies of IFs. The latter two aspects provide relatively recent and still underexplored directions for future research, with the goal of linking IF origin to plate tectonics and supercontinent cycles.

Progress since the late 1970s in the understanding of chemical oceanography and hydrothermal processes in the modern ocean has certainly influenced studies of IFs, although progress has been limited largely because direct comparison with modern hydrothermal analogues may not be directly applicable to the processes of IF deposition. Quantitative modeling of hydrothermal processes is needed in order to investigate the possible effects that these processes had on ancient oceans that lacked oxygen and sulfate.

Analyses of REE and Nd isotopes on bulk samples confirmed that hydrothermal processes dominated the chemistry of the early oceans. Once the role of hydrothermal processes in the origin of IFs and their links with submarine volcanism were appreciated, geochronology was employed to provide a test for synchronicity or diachronicity of IFs in unrelated basins on different cratons. This work helped establish a temporal and genetic link between IF deposition and mantle plume events. Although this work led to precise ages and revealed episodicity in deposition of major IFs, some giant IFs are still undated and their ages, once available, may dramatically impact our understanding of surface processes on early Earth.

Biological influences on the deposition of IFs were considered by Precambrian geologists and paleontologists for some time. However, only since the 1990s, with advances in the field of geomicrobiology, has the full potential of anaerobic photochemical iron oxidation as a driver for IF precipitation been realized. Models and experimental work on modern anaerobic iron-oxidizing communities have not yet been coupled fully with the rock record, in the sense that a direct proxy for this process in IFs is so far unknown. In fact, although models for iron oxidation have advanced significantly, it remains uncertain what signature in the rock record can discriminate among various mechanisms for iron oxidation.

Advances in analytical techniques have further broadened the field of geochemical investigation of IFs. New inductively coupled plasma mass-spectrometry (ICP-MS) instrumentation allows fast analysis of major and trace elements, including REE, as well as nontraditional stable isotopes (e.g., Fe, Cr, Si, and Ni). In situ laser-ablation techniques coupled with multicollector inductively coupled plasma mass-spectrometry (MC-ICP-MS) instrumentation and secondary ion mass spectroscopy (SIMS) provide an incredible level of spatial resolution (~10–20 μm) with a precision and accuracy approaching those of more time-consuming techniques using mineral separates and column chemistry, which provide limited spatial resolution. However, as discussed above, the absence of primary minerals in IF makes this direction of research, ultimately focused on the isotope and trace element composition of ancient seawater, likely to fail. Bulk sample analysis might provide a better way to characterize the composition of primary precipitates because the postdepositional redistribution of elements and intermineral isotopic fractionations will likely have a minimal effect on the scale of a hand specimen. However, in situ analysis still has a high potential when it is closely linked with detailed petrographic work. For example, the high preservation potential of primary textures in silicified layers and chert nodules is still largely unexplored, as are intra- and intergrain isotopic and trace element compositional heterogeneity in GIF. Data on these types of material might provide a better insight into seawater composition as well as mixing rates and element residence times in ancient oceans.

Field and detailed petrographic studies of Precambrian IFs rarely have been undertaken since the early 1990s and many IFs located in remote areas or areas having logistically difficult access were not investigated until the recent boom in iron ore prices. Industrial support during the last two decades was also mainly directed toward understanding how high-grade iron ores formed. As a result, recent state-of-the-art geochemical work has been done mainly on random or old grab samples without detailed geological control. Until now, no study has attempted to compare deep-water and shallow-water, or proximal and distal, sections of the same IF in the same depositional basin despite the obvious appeal of this approach.

Future progress in the study of IFs depends largely on careful integration of field-based and petrographic studies together with advanced geochemical analyses. Further, close links with geomicrobiology, chemical oceanography, and an understanding of submarine-hydrothermal processes are essential for deciphering the origin and evolution of IFs through time. The prospect is bright, but the burden of interaction and collaborative research is on participants, because each of these fields is in a rather advanced stage. However, as elsewhere in modern science, major breakthroughs are at the interface of fields and sciences rather than within established disciplines. Integration thus is needed among traditional, ‘old-fashioned’ disciplines and new advanced fields; the outcome of this research and training will likely lead to a well-rounded new generation of Precambrian geologists.
## Appendix 1  Precambrian Banded Iron Formations, Granular Iron Formations, and Rapitan-Type Iron Formations

<table>
<thead>
<tr>
<th>Formation/group/deposit</th>
<th>Location</th>
<th>Age (Ma)</th>
<th>Size (Gt)</th>
<th>Reference(s)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Maly Khinghan Formation</td>
<td>Far East, Russia</td>
<td>~560</td>
<td>10 000</td>
<td>4, 6</td>
</tr>
<tr>
<td>Yerbel Formation</td>
<td>Southeastern Uruguay</td>
<td>~560</td>
<td>Unknown</td>
<td>7, 8</td>
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<tr>
<td>Jacadigo Group (Urucum district)</td>
<td>Mato Grosso do Sul State, Brazil</td>
<td>~600</td>
<td>1000</td>
<td>4, 9–11</td>
</tr>
<tr>
<td>Bisokpabe Group</td>
<td>Togo, West Africa</td>
<td>~600</td>
<td>Unknown</td>
<td>12, 13</td>
</tr>
<tr>
<td>Chestnut Hill Formation</td>
<td>New Jersey Highlands, New Jersey, USA</td>
<td>~600</td>
<td>Unknown</td>
<td>14, 15</td>
</tr>
<tr>
<td>Holowilena Ironstone</td>
<td>South Australia, Australia</td>
<td>~650</td>
<td>Unknown</td>
<td>16–18</td>
</tr>
<tr>
<td>Braemar Iron Formation</td>
<td>South Australia, Australia</td>
<td>~650</td>
<td>Unknown</td>
<td>18–20</td>
</tr>
<tr>
<td>Vii′va and Koyva Formations</td>
<td>Western slope of Middle Ural Mountains, Russia</td>
<td>~650</td>
<td>Unknown</td>
<td>21–24</td>
</tr>
<tr>
<td>Bakeev (Tolparovo) Formation</td>
<td>Southern Ural Mountains, Russia</td>
<td>~650</td>
<td>Unknown</td>
<td>22</td>
</tr>
<tr>
<td>Dzhetymtau Suite</td>
<td>Middle Tian-Shan, Kyrgyzstz</td>
<td>~650</td>
<td>Unknown</td>
<td>25–28</td>
</tr>
<tr>
<td>UK Formation</td>
<td>Southern Ural Mountains, Russia</td>
<td>~700(?)</td>
<td>Unknown</td>
<td>22</td>
</tr>
<tr>
<td>Yamata Formation</td>
<td>East Siberia, Russia</td>
<td>~700(?)</td>
<td>Unknown</td>
<td>6</td>
</tr>
<tr>
<td>Lake Khanka Formation</td>
<td>Far East, Russia</td>
<td>~700(?)</td>
<td>Unknown</td>
<td>6</td>
</tr>
<tr>
<td>Rapitan Formation</td>
<td>Yukon Territory and Northwest Territories, Canada</td>
<td>715</td>
<td>1000</td>
<td>4, 29–32</td>
</tr>
<tr>
<td>Chuos Formation</td>
<td>Northern Namibia</td>
<td>~715</td>
<td>Unknown</td>
<td>4, 12, 33, 34</td>
</tr>
<tr>
<td>Tindir Group</td>
<td>Alaska, USA</td>
<td>715</td>
<td>Unknown</td>
<td>35, 36</td>
</tr>
<tr>
<td>Fulu Formation</td>
<td>Jiangxi Province, China</td>
<td>~741</td>
<td>1000</td>
<td>6, 37</td>
</tr>
<tr>
<td>Medvezhevo Formation</td>
<td>Patom uplift, Siberia, Russia</td>
<td>~700–750</td>
<td>Unknown</td>
<td>38, 39</td>
</tr>
<tr>
<td>Kingston Peak Formation</td>
<td>California, USA</td>
<td>~700–750</td>
<td>Unknown</td>
<td>40, 41</td>
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<tr>
<td>Numees Formation</td>
<td>Southern Namibia</td>
<td>~700–750</td>
<td>Unknown</td>
<td>12, 33, 34, 42</td>
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<tr>
<td>Mugur Formation</td>
<td>Tuva, Russia and Mongolia</td>
<td>767 ± 15</td>
<td>Unknown</td>
<td>6</td>
</tr>
<tr>
<td>Nizhne-Angara Formation</td>
<td>Angara-Pit area, Enisey Ridge, Siberia, Russia</td>
<td>~800</td>
<td>Unknown</td>
<td>43</td>
</tr>
<tr>
<td>Aok Formation (Shaler Supergroup)</td>
<td>Northwest Territories, Canada</td>
<td>~840</td>
<td>Unknown</td>
<td>44, 45</td>
</tr>
<tr>
<td>Xiamaling Formation</td>
<td>Xiahua yuan region, Hebei Province, North China</td>
<td>1368 ± 12 Ma</td>
<td>Unknown</td>
<td>46, 47</td>
</tr>
<tr>
<td>Roper Group (Corcoran and McMinn Formations)</td>
<td>Northern Territory, Australia</td>
<td>~1490</td>
<td>Unknown</td>
<td>48, 49</td>
</tr>
<tr>
<td>South Nicholson Group (Mullera Formation)</td>
<td>Queensland, Australia</td>
<td>~1500</td>
<td>Unknown</td>
<td>50, 51</td>
</tr>
<tr>
<td>Shoshong Formation</td>
<td>Eastern Botswana</td>
<td>~1600</td>
<td>1</td>
<td>4, 52, 53</td>
</tr>
<tr>
<td>Chuanlinggou Iron Formation</td>
<td>Hebei Province, China</td>
<td>~1650–1600</td>
<td>Unknown</td>
<td>54, 55</td>
</tr>
<tr>
<td>Pike’s Peak Iron Formation</td>
<td>Arizona, USA</td>
<td>~1728(57)</td>
<td>Unknown</td>
<td>57–59</td>
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<tr>
<td>Gibraltar Formation</td>
<td>Northwest Territories, Canada</td>
<td>~1880</td>
<td>Unknown</td>
<td>60–62</td>
</tr>
<tr>
<td>Frere Formation</td>
<td>Western Australia, Australia</td>
<td>~1890</td>
<td>10 000</td>
<td>4, 63–66</td>
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<tr>
<td>Alwar Group (North Delhi fold belt)</td>
<td>Rajasthan and Haryana Provinces, India</td>
<td>~1850–2000</td>
<td>Unknown</td>
<td>67–69</td>
</tr>
<tr>
<td>Lake Superior region</td>
<td>USA and Canada</td>
<td>10 000 (region total)</td>
<td>4</td>
<td></td>
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(Continued)
<table>
<thead>
<tr>
<th>Formation/group/deposit</th>
<th>Location</th>
<th>Age (Ma)</th>
<th>Size (Gt)</th>
<th>Reference(s)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Gunflint Iron Formation</td>
<td>Ontario, Canada</td>
<td>&lt;1878 ± 2 (71) to 1850 ± 1 (72)</td>
<td>(See above)</td>
<td>72–74</td>
</tr>
<tr>
<td>Negaunee Iron Formation</td>
<td>Michigan, USA</td>
<td>1874 to 1850/C6</td>
<td>1.3 100 000 (71) to &gt;1850 ± 1 (72)</td>
<td>4, 81, 82</td>
</tr>
<tr>
<td>Biwabik Iron Formation</td>
<td>Minnesota, USA</td>
<td>&lt;1880 to 1850/C6</td>
<td>(See above)</td>
<td>74, 75, 78</td>
</tr>
<tr>
<td>Ironwood Iron Formation</td>
<td>Michigan and Wisconsin, USA</td>
<td>&lt;1880 to 1850/C6</td>
<td>(See above)</td>
<td>75, 77, 79</td>
</tr>
<tr>
<td>Riverton Iron Formation</td>
<td>Michigan, USA</td>
<td>&lt;1880 to 1850/C6</td>
<td>(See above)</td>
<td>75, 77, 80</td>
</tr>
<tr>
<td>34. Sokoman Iron Formation</td>
<td>Labrador and Quebec, Canada</td>
<td>1877.8 ± 1.3</td>
<td>100 000</td>
<td>4, 81, 82</td>
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<td>35. Rochford Formation</td>
<td>South Dakota, USA</td>
<td>1884 ± 29</td>
<td>Unknown (small)</td>
<td>75, 83</td>
</tr>
<tr>
<td>36. Basile Formation</td>
<td>Northwest Territories, Canada</td>
<td>1930</td>
<td>100</td>
<td>84, 85</td>
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<tr>
<td>37. Liaohé Group</td>
<td>Liaoning Province, China</td>
<td>~1950–2050</td>
<td>(estimated)</td>
<td>86, 87</td>
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<td>38. Estes Formation</td>
<td>South Dakota, USA</td>
<td>2020–2010</td>
<td>Unknown (small)</td>
<td>75, 83</td>
</tr>
<tr>
<td>39. Pääkkö Iron Formation</td>
<td>Central Finland</td>
<td>2080 ± 45</td>
<td>100</td>
<td>4, 88, 89</td>
</tr>
<tr>
<td>40. Glen Township Formation</td>
<td>Minnesota, USA</td>
<td>~2100</td>
<td>Unknown (small)</td>
<td>75, 90</td>
</tr>
<tr>
<td>41. Lomagundi Group</td>
<td>Northern Zimbabwe</td>
<td>~2100–2200</td>
<td>Unknown (small)</td>
<td>91, 92</td>
</tr>
<tr>
<td>42. Caldeirão belt</td>
<td>Bahia State, Brazil</td>
<td>&gt;2078 to &lt;2687</td>
<td>Unknown (small)</td>
<td>93, 94</td>
</tr>
<tr>
<td>43. Malumfashi, Maru, Birnin Gwari, Kushaka, Muro, Egbe-Isanlu, and Obajana schist belts</td>
<td>Northern and southern parts of West-Nigeria</td>
<td>~2100</td>
<td>Unknown (small)</td>
<td>95</td>
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<td>44. Ijil Group</td>
<td>Western Mauritania</td>
<td>~2200</td>
<td>100</td>
<td>4, 96, 97</td>
</tr>
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<td>45. Nimba Itabirite</td>
<td>Liberia, West Africa</td>
<td>&gt;2100 to &lt;2615</td>
<td>10 000</td>
<td>4, 98–100</td>
</tr>
<tr>
<td>46. Hotazel Iron Formation</td>
<td>Northern Cape Province, South Africa</td>
<td>~2200</td>
<td>150 (calculated; 101)</td>
<td>102–105</td>
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<tr>
<td>47. Timeball Hill Formation</td>
<td>Gauteng and North West Provinces, South Africa</td>
<td>~2320</td>
<td>10</td>
<td>106–108</td>
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<td>48. Kursk Supergroup</td>
<td>Western Russia</td>
<td>~2450</td>
<td>300 000(109)</td>
<td>4, 110, 111</td>
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<td>49. Krivoy Rog Supergroup</td>
<td>Eastern Ukraine</td>
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<td>50 000</td>
<td>4, 110, 112, 113</td>
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<td>50. Transvaal Province</td>
<td>Northern Cape Province, South Africa</td>
<td>2431 ± 31</td>
<td>(See above)</td>
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<td>Griquatown Iron Formation</td>
<td>Northern Cape Province, South Africa</td>
<td>2465 ± 7</td>
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<td>Kuruman Iron Formation</td>
<td>Northern Cape Province, South Africa</td>
<td>2480 ± 6</td>
<td>(Included above; 118)</td>
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<td>Penge Iron Formation</td>
<td>North West Province, South Africa</td>
<td>2495 ± 6</td>
<td>(Included above; 121)</td>
<td>119, 120</td>
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<td>51. Hamersley Basin</td>
<td>Western Australia, Australia</td>
<td>2445 ± 5</td>
<td>19 000</td>
<td>122, 123</td>
</tr>
<tr>
<td>Boolgeeda Iron Formation</td>
<td>Western Australia, Australia</td>
<td>2449 ± 3</td>
<td>19 000</td>
<td>122–124</td>
</tr>
<tr>
<td>Weeli Wolli Formation</td>
<td>Western Australia, Australia</td>
<td>2461 ± 6</td>
<td>30 000</td>
<td>122–124</td>
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<tr>
<td>Brockman Iron Formation (Joffre Mbr)</td>
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<td>13 000</td>
<td>122, 124, 125</td>
</tr>
<tr>
<td>Brockman Iron Formation (Dales Gorge Mbr)</td>
<td>Western Australia, Australia</td>
<td>2495 ± 6</td>
<td>100 000(121)</td>
<td>4</td>
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<th>Formation/group/deposit</th>
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<th>Age (Ma)</th>
<th>Size (Gt)</th>
<th>Reference(s)</th>
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<td>Mt. Sylvia Formation</td>
<td>Western Australia, Australia</td>
<td>~2506</td>
<td>3000</td>
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<tr>
<td>Marra Mamba Formation</td>
<td>Western Australia, Australia</td>
<td>2597 ± 5</td>
<td>17 000</td>
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<tr>
<td>52. Caué Formation</td>
<td>Minas Gerais State, Brazil</td>
<td>~2450</td>
<td>100 000</td>
<td>4, 127, 128</td>
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<tr>
<td>53. Indian Creek Metamorphic Suite</td>
<td>Montana, USA</td>
<td>2470–2750</td>
<td>1</td>
<td>4, 129–131</td>
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<tr>
<td>54. Ruker Series</td>
<td>Prince Charles Mountains, East Antarctica</td>
<td>2450–2480</td>
<td>Unknown</td>
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<tr>
<td>55. Benchmark Iron Formation</td>
<td>South Dakota, USA</td>
<td>2480–2560</td>
<td>Unknown (small)</td>
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</tr>
<tr>
<td>56. Hutchison Group (Middleback Ranges)</td>
<td>Cleve domain, South Australia, Australia</td>
<td>~2500</td>
<td>1000</td>
<td>4, 135–137</td>
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<tr>
<td>57. Nemo Iron Formation</td>
<td>South Dakota, USA</td>
<td>2560–2890</td>
<td>Unknown (small)</td>
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<tr>
<td>58. Chitradurga Group</td>
<td>Karnataka Province, southern India</td>
<td>2614 ± 8</td>
<td>1000 (estimated)</td>
<td>138–140</td>
</tr>
<tr>
<td>59. Beardmore-Geraldton assemblage</td>
<td>Ontario, Canada</td>
<td>~2700</td>
<td>Unknown (small)</td>
<td>141, 142</td>
</tr>
<tr>
<td>60. Anshan Iron Formation</td>
<td>Liaoning Province, China</td>
<td>~2700</td>
<td>10 000 (estimated) (143)</td>
<td>144–146</td>
</tr>
<tr>
<td>61. Manjeri Iron Formation</td>
<td>Southern Zimbabwe</td>
<td>~2700–2830</td>
<td>Unknown (estimated)</td>
<td>147–149</td>
</tr>
<tr>
<td>62. Bababudan Group</td>
<td>Karnataka Province, southern India</td>
<td>2720 ± 7</td>
<td>1000 (estimated)</td>
<td>150–152</td>
</tr>
<tr>
<td>63. Central Slave Cover Group</td>
<td>Nunavut, Canada</td>
<td>2730–2920</td>
<td>Unknown (estimated)</td>
<td>153</td>
</tr>
<tr>
<td>64. Carajás Formation</td>
<td>Pará State, Brazil</td>
<td>~2740–2750</td>
<td>50 000 (estimated)</td>
<td>154, 155</td>
</tr>
<tr>
<td>65. West Rand Group</td>
<td>Gauteng Province, South Africa</td>
<td>~2960</td>
<td>10</td>
<td>4, 156–158</td>
</tr>
<tr>
<td>66. Pongola Supergroup</td>
<td>KwaZulu-Natal Province, South Africa</td>
<td>~2960</td>
<td>100</td>
<td>4, 159, 160</td>
</tr>
<tr>
<td>67. Jack Hills belt</td>
<td>Western Australia, Australia</td>
<td>&gt;3080 ± 20</td>
<td>Unknown (small)</td>
<td>161, 162</td>
</tr>
<tr>
<td>68. Moodies Group</td>
<td>Barberton, South Africa</td>
<td>~3230</td>
<td>Unknown (small)</td>
<td>163–165</td>
</tr>
</tbody>
</table>

*Most iron formations ~ 2.4 Ga and older consist largely of banded iron formation (BIF); those ~ 2.3–0.8 Ga are predominantly granular iron formation (GIF); Neoproterozoic Rapitan-type iron formation (0.80–0.56 Ga) associated with glacial deposits are in italics. Several ironstones of Mesoproterozoic age are included in this compilation (see text). Algoma-type iron formation, which occurs in or is closely associated stratigraphically with volumetrically abundant volcanic rocks (1–3), is excluded because this generally smaller type of iron deposit, relative to predominantly sediment-hosted iron formation, does not inform the redox state of deep seawater on a large scale (2). Sizes of BIFs are estimated amounts of originally deposited iron formation (4), and are much greater than those of reported ore reserves or resources, which are affected by extent of erosional preservation, mining practices, economics, and other factors. This compilation does not include numerous (mostly small) sediment-hosted iron formations in China and elsewhere in Far East Asia (5), for which geologic, geochronologic, and tonnage data are difficult to obtain.

56. This age is problematic because the contact between the dated felsic metavolcanic unit and the Pike's Peak Iron-Formation is a fault.
70. Radiometric age within iron formation.
71. Minimum age from occurrence of Sudbury impact layer at top of iron formation.
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101. Tonnage figure excludes Mn-rich units.


109. Size of original iron formation unit is estimated based on tonnage data for Fe (4).


114. Tonnage total for region includes Penge Iron Formation in North West Province, South Africa (4).


121. Regional total; tonnages for individual iron formations in the Hamersley Basin are estimated using relative stratigraphic thicknesses compared to the total tonnage for the basin.


### Appendix 2 Exhalites Associated with Precambrian Deep-Water (Cu-Rich) Volcanogenic Massive Sulfide Deposits

<table>
<thead>
<tr>
<th>VMS deposit (District)</th>
<th>Location</th>
<th>Age (Ma)</th>
<th>VMS (Mt)</th>
<th>Exhalite type(s)</th>
<th>Reference(s)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Keite Inlet</td>
<td>Alaska, USA</td>
<td>~554–600</td>
<td>&lt;1</td>
<td>Magnetite IF</td>
<td>9, 10</td>
</tr>
<tr>
<td>Jingtieshan</td>
<td>Gansu Province, China</td>
<td>586–670</td>
<td>&lt;1</td>
<td>Hematite–magnetite–siderite IF, jasper</td>
<td>11, 12</td>
</tr>
<tr>
<td>Um Samiuki</td>
<td>Eastern Desert, Egypt</td>
<td>~750</td>
<td>&lt;1</td>
<td>Magnetite IF</td>
<td>13–15</td>
</tr>
<tr>
<td>Matchless</td>
<td>Central Namibia</td>
<td>760–780</td>
<td>2.5</td>
<td>Magnetite IF</td>
<td>16–18</td>
</tr>
<tr>
<td>Otiwezwe</td>
<td>Central Namibia</td>
<td>760–780</td>
<td>15.4</td>
<td>Magnetite IF</td>
<td>16, 19, 20</td>
</tr>
<tr>
<td>Xiqu</td>
<td>Zhejiang Province, China</td>
<td>~900</td>
<td>31.0</td>
<td>Jasper</td>
<td>21–23</td>
</tr>
<tr>
<td>Altlén-Tape</td>
<td>Eastern Romania</td>
<td>~1000</td>
<td>10.9</td>
<td>Magnetite IF</td>
<td>24</td>
</tr>
<tr>
<td>Palmeiropolis</td>
<td>Tocontin State, Brazil</td>
<td>1170–1270</td>
<td>5.0</td>
<td>Magnetite IF</td>
<td>25, 26</td>
</tr>
<tr>
<td>Boksputs</td>
<td>Cape Province, South Africa</td>
<td>1275–17</td>
<td>1.7</td>
<td>Magnetite IF</td>
<td>27, 28</td>
</tr>
<tr>
<td>Itaberaba</td>
<td>São Paulo State, Brazil</td>
<td>1395 ± 10</td>
<td>None (29)</td>
<td>Magnetite and hematite IF</td>
<td>30, 31</td>
</tr>
<tr>
<td>Bon Ton</td>
<td>Colorado, USA</td>
<td>~1700</td>
<td>&lt;1</td>
<td>Magnetite IF</td>
<td>32, 33</td>
</tr>
<tr>
<td>Jones Hill</td>
<td>New Mexico, USA</td>
<td>1720 ± 15</td>
<td>11.0</td>
<td>Magnetite IF, silicate IF, ferruginous chert</td>
<td>34–36</td>
</tr>
</tbody>
</table>

- **Old Dick/Bruce (Bagdad)**: Arizona, USA; 1721 ± 6; ~2; Hematitic chert; 37–39
- **United Verde (Jerome)**: Arizona, USA; 1738.5 ± 0.5; >30; Jasper, hematite IF; 5, 40, 41
- **Copper Chief (Jerome)**: Arizona, USA; 1738.5 ± 0.5; <1; Jasper, magnetite, and hematite IF; 5, 40, 42
- **Miguela**; Eastern Bolivia; 1765–1690; 1.6; Magnetite and hematite IF; 43, 44
- **Gunnison (Gunnison)**: Colorado, USA; 1776 ± 7; <1; Magnetite IF, hematitic chert; 33, 45, 46
- **Encampment/Pearl**; Wyoming, USA; 1792 ± 15; <1; Magnetite IF, silicate IF, ferruginous chert; 47, 48

San Diego: Western Australia, Australia; 1843 ± 2; 2.3; Magnetite IF, sulfide–silicate IF; 49, 50

Sherritt Gordon: Manitoba, Canada; ~1860; 7.0; Sulfidic chert and schist; 51, 52

Crandon: Wisconsin, USA; 1869 ± 6; 63.5; Pyritic argillite/tuff, pyritic chert; 53–55

Eisenbrey/Thorntappe: Wisconsin, USA; ~1870; 3.0; Magnetite IF, silicate IF; 56, 57

Bend: Wisconsin; ~1870; 3.7; Jasper, magnetite, and hematite IF; 56, 58

Soucy: Québec, Canada; ~1880; 5.5; Sulfide–silicate IF; 59, 60

Rutan: Manitoba, Canada; 1883 ± 2; 82.8; Pyritic chert, sulfide IF; 61–63

Anderson Lake (Snow Lake): Manitoba, Canada; 1892 ± 3; 3.4; Sulfide IF; 64, 65

Spruce Point (Snow Lake): Manitoba, Canada; 1892 ± 3; 1.9; Sulfide IF; 65, 66

Bigstone (Snow Lake): Manitoba, Canada; 1892 ± 3; 1.5; Sulfidic tuff; 65, 66

Aijala: Southwestern Finland; ~1895; <1; Silicate IF, oxide IF; 67, 68

Flin Flon (Flin Flon): Manitoba, Canada; 1903 ± 7; 62.9; Sulfide IF, magnetite IF; 69, 70

Wutai: Shanxi Province, China; ~2530; Unknown; Sulfide IF, magnetite IF; 71, 72

Izok: Nunavut, Canada; 2614 ± 8; ~1; Pyritic IF; 73–75

Corbet (Noranda): Quebec, Canada; 2700 ± 2.6; 2.8; Pyritic IF; 77, 78

Milenbach (Noranda): Quebec, Canada; 2700 ± 2.6; 3.6; Sulfide IF; 78–80

High Lake: Nunavut, Canada; 2705 ± 1; 15.6; Pyritic IF, carbonate IF; 81

Gecco (Manitouwadge): Ontario, Canada; 2770 ± 2; 58.4; Magnetite IF, sulfide–silicate IF; 82–84

Willroy (Manitouwadge): Ontario, Canada; 2773 ± 2; 4.0; Magnetite IF, sulfide–silicate IF; 82–84

Bell Allard (Matagami): Québec, Canada; 2724.5 ± 1.8; 3.2; Sulfidic chert and tuff, silicate IF; 85–88

Orchon (Matagami): Québec, Canada; 2724.5 ± 1.8; 3.5; Sulfidic chert and tuff, silicate IF; 85–88

Selbaie: Québec, Canada; 2729 ± 3; 29.9; Pyritic IF; 89, 90

Sturgeon Lake (Sturgeon Lake): Ontario, Canada; 2735.5 ± 1.5; 4.0; Pyritic tuff; 91–94

Lyon Lake (Sturgeon Lake): Ontario, Canada; 2735.5 ± 1.5; 3.2; Pyritic tuff; 91–94

Scuddles (Golden Grove): Western Australia, Australia; 2960 ± 6; 10.5; Sulfidic chert, magnetite IF, jasper; 95, 96

Maranda (Murdochson): North West Province, South Africa; 2966.5 ± 1.6; <1; Pyritic chert; 97, 98

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*Exhalites are chemical sedimentary rocks that formed on the sea floor by precipitation from submarine-hydrothermal fluids, generally as fallout from nonbuoyant plumes either proximal or distal to volcanic massive sulfide (VMS) deposits (1–4). Occurrences compiled here are restricted to VMS deposits having >1% Cu, which implies formation from high-temperature (>300 °C) fluids in relatively deep seawater at depths of >850 m (5). Excluded from this compilation are exhalites that occur within the same volcanic or volcanosedimentary sequence as the VMS deposit(s), but at different (or unknown) stratigraphic levels (6). Exhalites associated with Zn–Pb deposits are also excluded, because they could have formed from low-temperature fluids in shallow restricted basins, within oxic seawater above the chemocline. Similarly excluded are magnetite-rich lenses that formed by subseaﬂoor replacement and not as true exhalites, such as in the Gossan Hill VMS deposit in Western Australia (7), and epigenetic magnetite-rich bodies belonging to the iron oxide–copper–gold (IOCG) class of deposits that formed in continental, not submarine, settings (8). With few exceptions, ages of the VMS deposits are based on high-precision U–Pb zircon geochronology of felsic metavolcanic host rocks. Abbreviations: Ma, million years; Mt, million metric tons; IF, iron formation.*

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**Note:** The table values are rounded to the nearest integer for simplicity.
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