Sediment-Hosted Stratabound Copper Deposit Model

Chapter M of
Mineral Deposit Models for Resource Assessment


U.S. Department of the Interior
U.S. Geological Survey
Annotated view of the outcrops and geology along the east side of Rock Lake, Sanders County, Montana, showing the setting of the discovery outcrops of the Montanore sandstone copper deposit. The view looks approximately due north. Stratigraphic unit abbreviations are: Ysr, Mesoproterozoic St. Regis Formation; Yru, Mesoproterozoic Revett Formation, upper member; Yrm, Revett Formation, middle member; A, Revett Formation, lower member, A quartzite beds; C, Revett Formation, lower member, C quartzite beds; and E, Revett Formation, lower member, E quartzite beds. The A and C beds of the lower member of the Revett are mineralized and dipping west toward the axis, beneath the lake, of a narrow syncline. The syncline formed as a drag fold between the Rock Lake Fault, approximately beneath the photographer, and the Snowshoe Fault, sketched in blue with the relative movement of the eastern (opposite) block shown to illustrate the cause of the fold. (Photograph by Timothy S. Hayes, U.S. Geological Survey).

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By Timothy S. Hayes, Dennis P. Cox, Nadine M. Piatak, and Robert R. Seal, II

Chapter M of
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Conversion Factors

SI to Inch/Pound

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<th>By</th>
<th>To obtain</th>
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</thead>
<tbody>
<tr>
<td>Length</td>
<td></td>
<td></td>
</tr>
<tr>
<td>centimeter (cm)</td>
<td>0.3937</td>
<td>inch (in.)</td>
</tr>
<tr>
<td>millimeter (mm)</td>
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<td>inch (in.)</td>
</tr>
<tr>
<td>meter (m)</td>
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<td>foot (ft)</td>
</tr>
<tr>
<td>kilometer (km)</td>
<td>0.6214</td>
<td>mile (mi)</td>
</tr>
<tr>
<td>Area</td>
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<td></td>
</tr>
<tr>
<td>square kilometer (km²)</td>
<td>0.3861</td>
<td>square mile (mi²)</td>
</tr>
<tr>
<td>square kilometer (km²)</td>
<td>247.1</td>
<td>acre</td>
</tr>
<tr>
<td>Mass</td>
<td></td>
<td></td>
</tr>
<tr>
<td>gram (g)</td>
<td>0.03527</td>
<td>ounce, avoirdupois (oz)</td>
</tr>
<tr>
<td>kilogram (kg)</td>
<td>2.205</td>
<td>pound avoirdupois (lb)</td>
</tr>
<tr>
<td>megagram, tonne (Mg,t)</td>
<td>1.102</td>
<td>ton, short (2,000 lb)</td>
</tr>
<tr>
<td>megagram, tonne (Mg,t)</td>
<td>0.9842</td>
<td>ton, long (2,240 lb)</td>
</tr>
</tbody>
</table>

Temperature in degrees Celsius (°C) may be converted to degrees Fahrenheit (°F) as follows:

°F=(1.8×°C)+32

Temperature in degrees Fahrenheit (°F) may be converted to degrees Celsius (°C) as follows:

°C=(°F–32)/1.8

Concentrations of chemical constituents in water are given either in milligrams per liter (mg/L) or micrograms per liter (µg/L).

Abbreviations

- ppm: parts per million
- kg/t: kilogram per tonne (1 kg/t = 1000 times 1 ppm)
- ppb: parts per billion
- µm: micron
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Abstract

This report contains a descriptive model of sediment-hosted stratabound copper (SSC) deposits that supersedes the model of Cox and others (2003). This model is for use in assessments of mineral resource potential. SSC deposits are the second most important sources of copper in the world behind porphyry copper deposits. Around 20 percent of the copper in the world is produced from this class of deposits. They are also the most important sources of cobalt in the world, and they are fourth among classes of ore deposits in production of silver. SSC deposits are the basis of the economies of three countries: Democratic Republic of Congo, Poland, and Zambia. This report provides a description of the key features of SSC deposits; it identifies their tectonic-sedimentary environments; it illustrates geochemical, geophysical, and geoenvironmental characteristics of SSC deposits; it reviews and evaluates hypotheses on how these deposits formed; it presents exploration and assessment guides; and it lists some gaps in our knowledge about the SSC deposits. A summary follows that provides overviews of many subjects concerning SSC deposits.

Summary of Sediment-Hosted Stratabound Copper Deposit Model

Description of Key Features

Sediment-hosted stratabound copper (SSC) deposits are bodies of disseminated, cementing, and lesser veinlet-hosted copper (Cu) minerals that are peneconformable with their sedimentary or metasedimentary host rocks. These SSC deposits are characterized by strong zoning of the ore minerals laterally along and across the bedding, from pyrite, to chalcopyrite, to bornite, to chalcocite, and to hematite. Chalcocite and bornite zones are typically the ore zones.

SSC deposits occur in three subtypes divided by host lithology and by the corresponding type of reductant that precipitated sulfur (S) and Cu from warm (75–220 °C), oxidized (hematite-stable), metals-transporting, sedimentary brines.

The three subtypes, their typical host lithologies, and their corresponding reductants are (1) reduced-facies type: black to gray to green shale, siltstone, mudstone, or carbonaceous dolosiltstone containing solid amorphous organic matter and pyrite; (2) sandstone-type: gray, well-sorted, fine- to coarse-grained sandstone containing petroleum, probably sour gas in most cases; and (3) red-bed type: gray, poorly to moderately sorted, and commonly conglomeratic fluvial sandstone containing carbonized plant fragments from silt-size up to whole logs. These black, gray, white, or green host rocks either lie atop or are enclosed within thick (greater than 300 m and commonly greater than 1 km) sequences of red beds. The association with red beds is an important distinction from other Cu deposits found in sedimentary rocks.

Reduced-facies and sandstone Cu deposits can be quite large (tens of millions to more than a billion tonnes of ore), whereas red-bed-type deposits rarely have as much as 10 million tonnes of ore. Outside of the Central African Copperbelt, Cu grades of reduced-facies-type deposits average 1.38 percent Cu; sandstone-type deposits average 1.08 percent Cu; and red-bed-type deposits average 2.09 percent Cu (Parks and others, U.S. Geological Survey, written commun., 2012). Average grades of deposits in the Central African Copperbelt are higher: 2.59 percent Cu for reduced-facies-type deposits and 2.20 percent for sandstone deposits. SSC deposits are an important source of other metals, ranking first among all deposit types in world cobalt (Co) production and fourth among all deposit types in silver (Ag) production. Twenty-five percent of SSC deposits contain Ag, grading between 2 and 200 g/t, and 14 percent of SSC deposits contain Co, grading between 0.05 and 0.5 percent in primary ores (Parks and others, U.S. Geological Survey, written commun., 2012). Those that contain byproduct Ag do not contain byproduct Co and vice versa; only the Kupferschiefer, which contains both byproduct metals, is an exception.

Deposits are confined to a limited package of sedimentary beds. Reduced-facies deposits have the geometry of sheets. Sandstone-type deposits are tabular to lens-like. Red-bed-type deposits are lenses of Cu-bearing rock. The lateral extent of entire mineralizing systems is enormous, and because the Cu source rocks, typically within the red-bed packages, have nowhere been fully distinguished and mapped, the total dimensions are not known.
Regional Environment

SSC deposits formed in sedimentary basins that now vary from virtually undeformed to intensely folded and thrust-faulted. These deposits are hosted by sedimentary-rock sequences that were deposited within 20 to 30 degrees of the paleo-equator, and thus associated with hot, dry climates that produced thick packages of red beds and commonly also produced evaporites (Kirkham, 1989). These thick packages of red beds were deposited in either proto-oceanic rifts or as post-collisional molasse. Ore-hosting lithologies for reduced-facies deposits are those that immediately mark the transgression in overall transgressive packages of rocks. Depositional environments of such beds include low sedimentation rate open marine shelves, coastal sabkhas, and subtidal carbonate shelves. Sandstone-type host rocks were deposited in subtidal nearshore, beach, eolian, and fluvial environments. The host rocks of red-bed-type deposits are fluvial or, in rare cases, deltaic sandstones. Faults at high angles to the host beds are found at many SSC deposits. Some of these faults may have been conduits along which warm metals-transporting brines moved upward across strata before spreading within permeable sandstone host beds or before spreading below impermeable reduced-facies host beds. Locations where reduced-facies host rocks onlap basement paleohighs are particularly favorable. Deposits at a few places appear to be within rocks that were deformed by strong halokinesis.

Hypogene Ore and Gangue Characteristics and Alteration

In unmetamorphosed SSC deposits, pyrite was replaced by chalcopyrite, which was replaced in turn by bornite, then by chalcocite. Further, pyrite was replaced by magnetite (rarely) or hematite (commonly). A fine dusting of authigenic hematite is found in parts of the chalcocite zone in all deposits. Cu sulfides also replaced carbonaceous materials in the sedimentary host rocks at essentially every scale. Where organic matter or organic carbon compounds were not directly replaced by the Cu sulfides, the organic materials were altered chemically and physically, changing from predominant alginate to predominant bitumen, or they were oxidized to dissolved carbonate species and precipitated as ore-stage carbonates. Cu minerals occur as disseminated 0.01-to-5 mm grains in clusters commonly following coarser grained laminae in the host sediment. Grains of Cu sulfide most commonly occur interstitially to detrital silicates or interstitially to carbonate allochems.

Some gangue minerals in SSC deposits are zonally arranged with their appearance boundaries parallel to the zonation of sulfides. As with the sulfides, most of these gangue minerals are cements of the sedimentary rocks and replacements of earlier detrital and cementing minerals. Pyrite is found in deposit-fringing zones. Iron (Fe)-rich calcite cements are found within pyritic zones of sandstone Cu deposits at

Supergene Ore and Gangue Characteristics

In the Central African Copperbelt, Cu probably undergoes locally significant supergene enrichment as chalcocite, covellite, and native copper (and probably djurleite, some anilite, and possible traces of low digenite) below a zone of leached capping typically 30 to 90 m thick extending from the surface and beneath a zone of oxide enrichment. The leached zone typically contains minor malachite and plancheite. The underlying oxide Cu zone has abundant malachite, azurite, other Cu oxides, and native copper, and extends to as much as a few hundred meters from the surface. Still deeper is a zone of secondarily enriched chalcocite-rich ore that may extend at least to 400 m depth and locally to as much as 800 m. This enrichment extends far below the modern water table. Although it is clear that supergene enrichment as chalcocite has occurred in central Africa, the appearance, unchanged from the time of Bateman (1930), is that “perhaps the major part” of the chalcocite is hypogene.

Co in Central Africa is enriched residually within the zone of weathering, a different behavior than Cu. “Cobalt caps” commonly extend from surface into the zone of Cu-oxide enrichment. Co minerals in this environment are heterogenite-3R, kolwezite, sphaerocobaltite, cobaltoan dolomite, and cobaltoan calcite.

Outside of Central Africa, in most climates Cu is fixed in the weathering environment as malachite, brochantite, beer-bottle limonite, and other Cu oxides, and supergene enrichment is insignificant.
Geochemical Characteristics

Major-element compositions of mineralized rocks and country rocks show major metasomatic effects from SSC mineralizing systems. The Konkola deposit, Zambia, and deposits of the Western Montana Copper Sulfide Belt, United States, record potassium (K) enrichment accompanying Cu deposition. Both of those deposits also record barium (Ba) enrichment. In the altered hematitic rocks where metals were mobilized, the Western Montana Copper Sulfide Belt system shows sodium (Na) enrichment and extreme calcium (Ca) leaching. The major element, magnesium (Mg), is anomalous in many SSC deposits. The minor element, boron (B), is anomalous in some of the SSC systems in Zambia. It appears that SSC deposit mineral zonations are comparable with chromatographic separations of elements so that small variations in solution concentrations, temperature, or host-rock conditions can shift the positions of a mineral or elemental front in one system relative to another or relative to a different place within the same system.

Many trace elements are found in anomalous concentrations in zonal arrangements near and in SSC deposits. Ag and Co are common byproduct metals from within zones of Cu ore. Ag is nearly totally confined to chalcocite- and bornite-bearing zones, but Co extends to within pyritic zones. Also certainly among the trace elements enriched in deposit-related zonations are antimony (Sb), arsenic (As), Ba, lead (Pb), Mn, mercury (Hg), rhenium (Re), and zinc (Zn). The trace elements, gold (Au), palladium (Pd), and platinum (Pt), are found in anomalous concentrations, and the rock constitutes ore many places in a zone that parallels the sulfide zonation in a gentle cross through bedding along the footwall of Kupferschiefer Cu deposits in Poland. U and V may reach ore grades, either separately or together, in some red bed subtype deposits, and U also reaches ore grades in certain deposits of the Central African Copperbelt. Ferroan and manganoan carbonates of sandstone Cu deposits may be an identifier of mineralizing systems that extend well outside of the ore zones themselves. Some of these carbonates formed at the downstream fringes of a mixing and reaction zone between introduced metalliferous brine and reservoirs of natural gas (Hayes and others, 2012), and it is likely that they mark the locations of coupled redox reactions between SO$_4^{2-}$ of the metals-transporting brine and carbon in the reservoired methane (CH$_4$). Such a reaction could be expressed as:

$$\text{Ca}^{2+} + 2\text{CuCl}_{0.04} + \text{SO}_4^{2-} + \text{CH}_4 = \text{Cu}_2\text{S}$$

The physical properties and compositions of hydrothermal fluids that formed SSC deposits were poorly known until recent years. From fluid-inclusion microthermometry studies, ore deposition in the Central African Copperbelt reduced-facies-type deposits took place in two stages. The first stage that deposited the main volume of disseminated and replacement sulfides produced only tiny liquid-vapor fluid inclusions with no solid daughter minerals; the second stage of veinlet-controlled sulfides produced large inclusions with multiple daughter minerals including halite and sylvite as well as several other daughters or trapped solids. At the early stage, saline fluids (typically in excess of 15 eq. wt. % NaCl) transported Cu at 130 to 220 °C. Deposition occurred partly by mixing with low-temperature less-saline groundwater, possibly just ordinary pre-ore groundwater, with salinity below 10 eq. wt. % NaCl. The result of the mixing was large ranges in homogenization temperatures and salinities. Later, probably during or after the Lufilian Orogeny, second-stage Cu mineralization took place from fluids with temperatures over 200 to as much as 415 °C and with salinities of 35 to 55 eq. wt. % NaCl. Sources of the fluids are not known for either stage, but with the extraordinary salinities of the second-stage (veinlet) fluids, dissolved evaporites are likely sources for that salinity. Second-stage (veinlet) mineralization probably does not contribute large portions of the total Cu endowment. From fluid-inclusion microthermometry studies of three sandstone Cu systems, mixing was more important than in reduced-facies deposits. Temperature and salinity ranges were similar to the earlier of the two stages in Central African reduced-facies deposits.

Compositions of main-stage fluid-inclusion volatiles are known for a single case. Quadrupole mass-spectrometric analysis of ore-stage inclusion fluids from the Spar Lake sandstone Cu deposit has demonstrated that mineralization took place where a metals-transporting brine at about 200 °C mixed and reacted with reservoired sour gas. The metals-transporting brine was an aequous fluid lacking important amounts of dissolved CO$_2$, CH$_4$, or H$_2$S. It entered the gas reservoir from below before spreading in the reservoir sandstones and reacting with cooler natural gas. Ore was deposited as the brine reacted with CH$_4$ and H$_2$S in the natural gas toward lower fO$_2$. Neither the metals-transporting brine nor the natural gas had any component of magmatic or subcrustal fluids.

Organic geochemical studies have been completed at two reduced-facies-type deposits. At White Pine, United States, Nonesuch Shale organic matter was thermally matured during mineralization. In the Kupferschiefer, Poland and Germany, various organic chemical measurements, organic petrologic studies, and the spectral fluorescence intensities of extracts indicate a high degree of oxidation as well as high thermal maturity in the basal section of mineralized shale, interpreted to be caused by maturation of organics in the shale by ascending, hot, oxidizing, metalliferous brine.

All studied reduced-facies-subtype deposits have large ranges of S-isotope values among their sulfides, and they have some sulfides that have light values of δ$^{34}$S reaching commonly below −10 per mil (%). This distribution is universally interpreted as evidence that S now found in Cu sulfides was bacteriogenically fractionated from seawater sulfate and deposited as Fe sulfides at a time shortly after host-rock sedimentation just below the sediment-seawater interface. This δ$^{34}$S distribution in Cu sulfides is probably because of replacement of early diagenetic pyrite without appreciable redistribution of S isotopes. Some studies have demonstrated addition of
heavy $\delta^{34}$S sulfides after precipitation of the early diagenetic pyrite, probably at ore-stage, and probably by thermochemical sulfate reduction (TSR).

Values of S isotopes from the Spar Lake sandstone Cu deposit, U.S.A., are remarkably similar to distributions through the full diagenetic history of petroleum producing basins (Hayes and others, 2012). The S-isotope distribution indicates an original derivation of all the S by bacterial sulfate reduction (BSR) of seawater sulfate, followed by sequestering of the sulfide in organic matter of petroleum source rocks. With burial, maturation of the organic matter produced petroleum with relatively homogeneous sulfide $\delta^{34}$S values $15 \pm 5\%_o$ less than the original seawater $\text{SO}_4^{2-}$. With further burial, thermal cracking produced H$_2$S with the same relatively homogeneous $\delta^{34}$S that migrated and was exchanged in pore-gas reservoirs. Much of the ore-stage Cu sulfides simply precipitated that H$_2$S, producing a favored small range of S-isotope reservoirs. Much of the ore-stage Cu sulfides simply precipitated that H$_2$S, producing a favored small range of S-isotope values $15 \pm 5\%_o$ lighter than the $\delta^{34}$S of original seawater sulfate. There was an additional contribution of sulfide from later Thermochemical Sulfate Reduction (TSR) operating on formation water and (or) on sulfate of the metals-transporting brine in redox reactions with reservoired hydrocarbons (eqn. 1).

The sulfate added by ore-stage TSR was typically isotopically heavier than the reservoired sour-gas H$_2$S.

Carbon-isotope studies of ore-associated carbonate minerals from reduced-facies and sandstone Cu deposits include some low $\delta^{13}$C values. The low-$\delta^{13}$C-carbonates reflect oxidation of solid organic matter or petroleum to provide C for deposition of the carbonates.

Integrated S-isotope, C-isotope, and organic geochemical studies along with elemental geochemical studies of the Kupferschiefer showed that total C$_{org}$, C/S ratios, and hydrogen indices of the organic materials are facies-related and do not correlate with Cu grade. Those parameters were established when pyrite was deposited, probably by BSR from seawater during earliest diagenesis. Fe-C$_{org}$-S relations and ore microscopy indicated that, in parts of the Kupferschiefer bed with grades of 0.1 to as much as 6 percent Cu, the pyrite was replaced by Cu-Fe-sulfides, and the isotopic composition of the replaced pyrite was inherited. With yet more intense mineralization in the lower part of the Kupferschiefer bed, there was (1) deposition of sparry calcite with $\delta^{13}$C values substantially lowered from the typical marine values found in the top of the shale, coupled with (2) reciprocal change of $\delta^{13}$C in kerogen and bitumen to heavier isotopic values, and (3) an addition of Cu sulfides with heavier $\delta^{34}$S deposited by TSR of ore solution $\text{SO}_4$ coupled with oxidation of the organic carbon to carbonate. Cu grades greater than 8 percent in the lowest layers of the shale are dominated by the thermochemically reduced S from late in the ore stage. At Spar Lake, the isotopically light distal-fringe carbonates are coupled with the sulfide that was thermochemically reduced at ore-stage from $\text{SO}_4$ in the metals-transporting brine, in redox reactions like equation 1 with the reservoired CH$_4$. That sulfide from ore-stage thermochemical reduction is identified where samples have the combination of heavy $\delta^{34}$S values in Cu (or Pb) sulfides together with very light $\delta^{13}$C values in ore-stage calcite.

The ore processes at Sangerhausen and Spar Lake are similar. They simply use different reductants: solid organic material for the Kupferschiefer, CH$_4$ for Spar Lake.

Isotopic age dating of SSC deposits has suffered from the lack of easily dateable minerals within the deposits themselves. A few direct dating attempts have been made with limited success. Re-Os ages of Cu sulfides from different Kupferschiefer deposits have varied from the same time as shale deposition (within isotopic age uncertainty), to 55 million years after shale deposition, to 19 million years after shale deposition at Lubin, Poland. The Re-Os age at Lubin is close to agreement with the paleomagnetic age of Rote Fäule alteration in that district. Re-Os dating of disseminated bornite and chalcopyrite from Konkola, Zambia, yielded an age of 816 ± 62 Ma. That age is probably correct, but the large uncertainty, which resulted from high concentrations of common osmium in the sulfides, limits the utility of the dating. At White Pine, several age-dating studies have closely constrained the sequence and timing of ore-related events. Rapid deposition of more than a kilometer of Copper Harbor aquifer red beds and the lower Nonesuch reduced-facies host rocks and calcite diagenesis (interpreted as accompanying main-stage Cu-sulfide mineralization) all took place over no more than 15 million years after deposition of an upper Portage Lake Lava andesite flow, probably within just 6 million years. Basin inversion, compressional tectonics, second-stage Cu mineralization at White Pine, and mineralization of the Keweenawan Native Copper district had all taken place within the next 30 to 70 million years after that. Aleinikoff and others (2012) obtained a $^{206}$Pb/$^{204}$Pb age of 1,409 ± 8 Ma on ore-stage xenotime overgrowths on detrital zircons from Spar Lake. In host rocks that were deposited after 1,469 ± 3 Ma and before 1,454 ± 9 Ma, the xenotime-indicated mineralization age, 40 to 60 million years after host-rock deposition, is reasonable for the time at which a subsurface mineralizing brine would have reacted with reservoired sour gas.

Petrology of Associated Sedimentary Rocks

Common minerals in the sequence of diagenetic changes to sedimentary rocks are, for the host rocks of SSC deposits, important ore-stage gangue minerals. Most are not unusual or anomalous minerals in sedimentary diagenesis. Anhydrite that was replaced by quartz, dolomite, and sulfides in Zambian Copperbelt host rocks is an example. Just as in the changes seen in the diagenesis of anhydrite, the diagenetic changes seen in sandstone Cu deposits are all familiar in ordinary diagenesis, except, mainly, for the addition of Cu minerals. Every authigenic gangue mineral that is zonally distributed in the Spar Lake sandstone Cu deposit, with the exceptions of barite and Ba mica, has been found elsewhere as the product of ordinary diagenesis of sandstones. Many of the authigenic gangue minerals arranged zonally at Spar Lake have been found in the same paragenetic sequence as they appear at the deposit but with greater and greater burial depth and temperature in the Tertiary sandstones of the Gulf Coast in Texas.
Petrology of Associated Metamorphic Rocks

Many SSC deposits were simply metamorphosed along with the rocks of their host sedimentary sequences. Metamorphic devolatilization reactions are of interest for SSC deposits and their host sections. One is simply, hematite = magnetite + O₂. This reaction has been demonstrated in the Belt Supergroup rocks, United States. The importance is that original red beds of SSC systems can be metamorphosed to magnetite-bearing quartzites. This reaction may have produced magnetite quartzites that are several kilometers thick in the Udokan Series in Russia. A second metamorphic devolatilization reaction is pyrite = pyrrhotite + S₂. Almost all SSC deposit-bearing sequences that are at amphibolite grade have some pyrrhotite, and a few have abundant pyrrhotite. This pyrrhotite probably is a metamorphic product, not an ore-stage precipitate nor a pre-ore phase.

Genetic Theory and Mineral Resource Potential Assessment

SSC deposits require six elements to form. Evaluating the potential for SSC deposits using these six elements is analogous with evaluating a petroleum exploration “play.” Petroleum plays are analyzed considering the elements: (1) source rocks, (2) maturation, (3) migration, (4) reservoir rocks, (5) trap, and (6) seal. In order to form a SSC deposit, there must be (1) Cu source rocks, (2) liberation (dissolution) of Cu from those source rocks by a hot (mature, typically greater than 100 °C) sedimentary brine, (3) migration of the Cu-bearing brine, (4) non-oxidized, typically pyrite-bearing host rocks, (5) seal rocks (confining beds for the upward seeking metals-transporting brine), and (6) traps. The traps are physical and redox chemical traps.

A basin that contains Cu source rocks and host rocks is the fundamental unit that defines a permissive tract for exploration or assessments for SSC deposits. (1) The basin must contain some minimum, perhaps at least 300 m, of oxidized, hematite-stable permeable source rocks, probably red beds, with a small Cu content. Subaerial basaltic or other volcanic rocks are a positive factor as alternative Cu source rocks. (2) The basin must have undergone flow of hot (greater than 100 °C, and some places greater than 220 °C), briny, oxidized groundwater from an extensive catchment basin, cross-stratally upward toward host rocks. Because the metals-transporting solutions were brines, it is a positive factor if the basin contains or contained evaporites, either continental or marine. Evaporite presence either upsection or downsection relative to Cu source rocks and host rocks is permissive. It would be favorable if the basin has a history of halokinesis. (3) There must have been a flow system for transport of Cu in the sedimentary brines from source rocks to host rocks. Various types of flow systems with various hydrologic drives may have served, but the aquifers, at the time of mineralization, were confined so that brines were upward-seeking relative to stratigraphy. The flows probably had directions commensurate with dewatering the basin. (4) The basin must contain either organic-rich poorly permeable sedimentary beds that could provide solid organic matter as a reductant, permeable beds with reservoired petroleum vapor or liquids to act as reductant, or permeable beds with carbonized plant fragments to act as a reductant. (5) The basin must have had sufficient structural deformation to create traps. (6) The basin must have confining beds that are the upper parts of the reduced-facies host units, that overlie petroleum reservoirs, or that overlie sandstones containing carbonized plant fragments.

A permissive tract for SSC deposits is outlined by delineating an area on a geologic map that includes chemically reduced host rocks of any of the three types and generally stratigraphically lower red beds or former red beds and (or) flood basalts greater than or equal to 300 m thick. The exact thickness of source rocks needed to generate a metals-transporting brine is not known. A thickness of 300 m is considered to be near the minimum needed, and greater thicknesses can be considered more favorable. The tract may include known SSC deposits and prospects of any of the three subtypes. The tract outline approximately parallels the outline of the sedimentary basin, as depicted on the scale of the geologic map available for tract delineation. The presence of evaporite beds in the sequence and faults at high angles to bedding are particularly favorable.

Environmental Geology and Effects of Mining

Chalcocite group minerals and bornite in SSC deposits are fairly stable under oxidizing conditions. Pyrite is found only in trace amounts or not at all in the high-grade zones, but its content increases outward as Cu grade decreases. Therefore, environmental issues associated with the oxidation of pyrite can be minimized if mining avoids pyritic zones. The calcite and feldspar (or clays) present in some host rocks will help mitigate against the development of acid mine drainage. However, minerals such as Pb-bearing galena, sphalerite, which commonly contains cadmium, and As minerals in such as tetrahedrite/tennantite and luzonite, found associated with these deposits, can be disturbed during mining or concentrated during ore processing and can potentially weather and release environmentally harmful trace elements.

Tailings from reduced-facies-subtype deposits consistently contain around 0.2 percent Cu, but these have not proven to cause any major environmental problems (Sracek and others, 2010; E&MI, 1986b; U.S. Environmental Protection Agency, 1994; Williams and others, 2002). Secondary minerals on waste dumps include various Cu-, Zn-, and Pb-sulfates, chlorides, and carbonates. Gypsum and poorly crystalline Fe-oxides are reported as secondary phases in tailings. For tailings samples from the White Pine deposit, no acid-generating capacity was detected (Williams and others, 2002). The tailings of sandstone-subtype deposits from Montana contain low concentrations of metals like Cd and Sb, contain some leachable Pb, and contain S as sulfide ranging from 0.01 to 0.08 wt. %, which results in acid production values of only 0.3 to 2.5 kg/t CaCO₃. The net neutralization potentials of ore, waste rock, and tailings from Montana sandstone Cu deposits are reported as variable but generally low, from 2.8 to 10 kg/t.
CaCO₃. The favorable acid base accounting results are because of the low amounts of sulfides and presence of calcite in the tailings (U.S. Forest Service and others, 2009).

Overall, the metal mobility in mine-drainage waters from these deposits is low to moderate. Waters draining mines and tailings generally has near-neutral to moderately alkaline pH and low dissolved-metal contents. However, mining of Kupferschiefer deposits in Germany has locally contaminated stream sediments, soils, and surface water and groundwater with metals such as Zn, Pb, Cu, As, and Cd. The metals and S came predominantly from flue dust and slag from smelting (Schreck and Glaesser, 1998; Strauch and others, 2001; Schreck and others, 2005). In addition to the dissolution of metals from the flue dust, metals are dispersed into the environment as fine particles blown by the wind from waste or ore piles (Matheis and others, 1999).

Mining in the Central African Copperbelt in Zambia has led to elevated metal and sulfate concentrations in water that drained the mines and processing plants (Pettersson and others, 2000; Norrgren and others, 2000; Pettersson and Ingri, 2001). Though elevated concentrations of Co, Ni, and Zn are present, the concentrations do not exceed drinking water guidelines. Ore from Montana sandstone Cu deposits leached low concentrations of Cu (0.02–0.04 mg/L) and Mn (0.03 mg/L) and resulted in neutral pH water. Groundwater from monitoring wells frequently contains less than detectable dissolved metals concentrations. The pH of the groundwater is usually near neutral or slightly acidic (U.S. Forest Service and others, 2009).

Smelters are clearly the greatest environmental risks associated with mining SSC deposits. Soils within 1 km of the smelters in Poland contained elevated concentrations of Cu, Pb, Cd, and Zn (Roszyka and Szerszen, 1988; Helios Rybkica and Jędrzejczyk, 1995). The concentrations of Pb and Cd in ground and surface water in the vicinity of the smelters exceed World Health Organization (WHO) guidelines for drinking water (World Health Organization, 2006).

Detailed ecological studies have been carried out along the Kafue River, which flows through the Central African Copperbelt in Zambia. Pb in a grass species, three species of fish, and the liver of the lechwe (antelope species) were all anomalously high. River sediment collected downstream from several mines and near a city with major ore-processing facilities was significantly more toxic to zebrafish and tilapia than sediment collected farther downstream (Mwase and others, 1998). Near the Glogow smelters of Kupferschiefer ores in Poland, the concentrations of metals in cultivated consumable plants correlated with nearness to the smelters and with the direction and strength of prevailing winds. Other studies identified high levels of metals in tissues, milk, and excrement of cattle in the region. Discharge from the White Pine mine area has not caused measurable effects on the basis of a lack of impact on the benthic community, minor increases in chloride from mine discharges, and a lack of detectable effects on local water intakes (U.S. Environmental Protection Agency, 1994).

Knowledge Gaps

The source(s) of the metals-transporting solutions for SSC deposits remain conjectural. Brine inorganic chemistry will help determine the source(s) of the metals-transporting solutions. LA-ICPMS (laser-ablation inductively coupled plasma mass-spectrometry) or PIXE (proton-induced X-ray emission) methods promise to be successful where fluid inclusions are large enough, and encapsulated crushing followed by leaching of ore-stage authigenic minerals can be used in other such studies. However, the most reliable way to define the source(s) of metals-transporting solutions for SSC deposits would be to sample the metals-transporting solutions from an ongoing (presently active) mineralizing system. An ongoing sandstone Cu mineralizing system can probably be found in a basin that has production of sour natural gas and that has petroleum source rocks overlain by more than 300 m (preferably over 1 km) of red beds, overlain by intervals that contain gray or bleached sandstones. An ongoing reduced-facies-type mineralizing system may occur in a young rifting environment at low latitudes where organic-rich sabkha dolostone beds onlap basement highs and overlie thick sections of red beds. The search might be most productive in a young aulacogen like the Afar Rift or a young trans-tensional basin like the Gulf of California.

Mapping of alteration pathways over broad areas would limit the possible groundwater flow systems that could have produced ore. Certainly, high-angle faults and zones where pillars (water-escape structures) are densely distributed, both exposed downsectionward from mineralized rocks, are targets for study. From current knowledge, sodic minerals (albite, analcime, glauberite, riebekite, Na-illite, possibly scapolite), hematite, and possibly barite will mark the paths that fluids took between Cu source rocks and deposits. Better than either alteration mapping or paleohydrology studies, if an active ore system is found, its hydrologic drive could be straightforwardly observed and measured.

Several studies of smaller scope would also be useful. In the Central African Copperbelt, it is still unclear whether most of the chalcocite dates from supergene enrichment or whether it is hypogene. Knowledge of supergene enrichment is far advanced since the problem was last approached there, so new methods can be brought to bear. Insufficient knowledge of the changes caused in amphibolite-grade metamorphism of SSC deposits can be addressed at places like Aynak, Afghanistan, and the Ablah Group, Saudi Arabia. Such studies are economically important because high-grade metamorphism apparently makes the rock packages more amenable to geophysical exploration by IP (induced polarization) or MT (magnetotelluric) techniques. The most immediate payback from new studies will likely come from studies of exploration methods. Two methods that need testing are natural source induced polarization and soil gas chemistry.
Introduction

This report is a review of the geological literature about SSC deposits written in order to serve the purposes of U.S. Geological Survey (USGS) assessments of mineral-resource potential. SSC deposits are concentrations of Cu minerals that are disseminated as cements, replacements of clasts and earlier cements, and lesser veinlets within black, gray, green, or white sedimentary rocks within or atop sequences of red beds, which are typically many hundreds to thousands of meters thick. Such deposits have been mined far back into the 4th millennium B.C. in the Middle East (Weisgerber, 2006) and since A.D. 1199 in the Kupferschiefer in central Europe (Beck, 1905, p. 492). These deposits continue to be mined at a large scale, today, in the Central African Copperbelt, Poland, Kazakhstan, China, and the United States, and a large deposit of this type in Russia will come into production soon.

The report has been written to be used for the Updated National Mineral Resource Assessment of 2012. It builds on earlier published models that were used in assessments of mineral-resource potential and mineral environmental assessments by the USGS. It is an update to the descriptive model(s) of the deposit type in Cox and others (2003). The report follows the organization and choice of subject matter determined by the leadership of the USGS National Mineral Assessment Program in order to be consistent with other ore-deposits models.

The SSC deposits rank second in Cu production among the geologic deposit types of the world, behind porphyry Cu deposits. These deposits also produce important byproduct metals. They are the world’s largest producing deposit type for Co. They are the fourth largest producing deposit type for Ag, behind epithermal precious metal, porphyry Cu, and sedex Zn-Pb-Ag-barite deposits. SSC deposits form the base of the national economy for the Democratic Republic of Congo, Poland, and Zambia.

SSC deposits have been found on every continent except Antarctica (fig. 1), and they probably also exist in Antarctica should that continent ever be explored. However, just two districts account for 82 percent of the currently known resources of Cu within SSC deposits (Parks and others, U.S. Geological Survey, written commun., 2012). These are the Central African Copperbelt in Zambia and the Democratic Republic of Congo (fig. 2) and the Kupferschiefer in Germany and Poland (fig. 3). Among the 47 SSC deposits in the world that contain more Cu than a median porphyry Cu deposit, 28 are in the Central African Copperbelt and 8 are in the Kupferschiefer in Central Europe. Nonetheless, the scattering of large SSC deposits worldwide can probably be taken as evidence that additional large deposits remain to be discovered outside the Kupferschiefer and the Central African Copperbelt.

Deposit Type and Associated Commodities

Synonyms

These deposits have been termed sediment-hosted stratiform Cu, sedimentary rock-hosted stratiform Cu, Cu shale, Cu sandstone, cupriferous sandstones, sandstone Cu, red-bed Cu, Kupferschiefer type Cu, marine paralic type Cu, reduced-facies Cu, and Revett Cu deposits.

Brief Description

The SSC deposits are sedimentary rock-hosted peneconformable deposits typically dominated by chalcocite but strongly zoned laterally and vertically to bornite, then to chalcopyrite, and then to pyrite (figs. 4 to 7) (Kirkham, 1989; Cox and others, 2003). Some deposits also have a zone of native copper adjacent to the chalcocite zone and away from the bornite (figs. 4 and 5). However, where these are present, it is not necessarily clear that the native copper is a part of the same mineralizing event as the sulfides (for example; Mauk and others, 1992a; Brown, 2006), and it is unclear that the native copper is hypogene rather than supergene. The SSC deposits are hosted by gray, black, green, or white sedimentary rocks in sections otherwise dominated by or underlain by red beds, and the sections commonly contain evaporites, pseudomorphs of evaporites, or other indicators of aridity. Unmetamorphosed deposits of the class have a characteristic replacement paragenesis of sulfides and oxides within which pyrite was replaced by chalcopyrite, which was in turn was replaced by bornite and magnetite in some deposits, then by chalcocite and hematite in all deposits. The Cu-bearing sulfides formed where an advancing metals-transporting solution in equilibrium with hematite reacted with pre-existing pyrite and other detrital or pre-ore authigenic components of the host rocks and mixed and reacted with indigenous, probably sulfidic, pore fluids (Rentzsch, 1974; Hayes and others, 1989; Sutton and Maynard, 2005; Hayes and others, 2012). Depositional temperatures of ore varied from less than 90 °C to as much as 220 °C (Touwaide, 1930; Brown, 1971; Lur’ye and Gablina, 1976; Nishioka, 1983; Morrison and Parry, 1986; Hayes and others, 1989; Annels, 1989; Hayes, 1990; Breit and Meunier, 1990; Arehart and others, 1990; Gray and others, 1996; Dewaele and others, 2006; El Desouky and others, 2009; Hayes and others, 2012). Many earlier papers describe deposits of this class or address aspects of deposit genesis. Hitzman and others (2005) provided a broad-ranging earlier review of SSC deposits.
Figure 1. Locations of sediment-hosted stratabound copper (SSC) prospects and deposits of the world presently known to the U.S. Geological Survey. Deposits that have greater than 1.056 million metric tons of contained copper metal, the amount of the metal in a world median tonnage and world median grade porphyry copper deposit (Singer and others, 2008), are plotted in red. Thus, the deposits in red represent the largest and richest of the SSC deposits. Note that these are widely scattered worldwide, though the Kupferschiefer in central Europe and the Central African Copperbelt in the Democratic Republic of Congo and Zambia contain 36 of the 47 largest known SSC deposits (figs. 2 and 3).
Figure 2. Locations of known sediment-hosted stratabound copper prospects and deposits of the Central African Copperbelt, Democratic Republic of Congo and Zambia. Those plotted in red have copper metal in excess of a world median grade and tonnage porphyry copper deposit.
Figure 3. Locations of known sediment-hosted stratabound copper prospects and deposits of the Kupferschiefer in central Europe. Those plotted in red have copper metal in excess of a world median grade and tonnage porphyry copper deposit.
Deposits of this class are typically recognizable even where they have been metamorphosed to amphibolite or higher metamorphic grades because they still have sheetlike to tabular geometry mainly conformable with layering and have a diagnostic zonation along and across the layering of bornite to chalcopyrite to Fe sulfides. At grades of post-ore metamorphism exceeding upper greenschist, pyrrhotite may exceed or entirely displace pyrite in the zonation, and bornite commonly exceeds chalcocite.

SSC deposits are associated with red beds or former red beds where metamorphism has changed those rocks. The deposits are stratabound. That is, they are restricted to a particular stratigraphic interval within a sedimentary-rock sequence. However, the deposits do not perfectly follow sedimentary bedding. Association with red beds and stratabound character distinguish deposits of this class from other classes of sediment-hosted Cu deposits. For example, sediment-hosted Cu deposits at Mt. Isa, Australia, although they are stratabound, are not associated with red beds, occurring instead within dolostones near a facies change to black shales (Hitzman, 1983). Although Cu skarns are sediment-hosted, they are spatially and genetically associated with intermediate to acidic porphyries, and they are not stratabound. The popular synonymous class name, sediment-hosted stratiform Cu deposits (Brown, 1989), is rejected, because in detail, SSC deposits are not perfectly stratiform. In detail, even the very sheetlike reduced-facies SSC deposits (see below) cut bedding, with the deposit boundaries defined by sulfide mineral-zone boundaries rather than by bedding planes or by the lateral limits of the sedimentary host rocks (Brown, 1971; Rentzch, 1974).
The universal association of the deposits with red beds also presents a constraint on the ages of deposits of this type. Red beds, and associated SSC deposits, occur in the geologic record only since the Great Oxidation Event at about 2,400 Ma (Chandler, 1988; Bekker and others, 2004). Correspondingly, there are no Archean SSC deposits.

Host rocks and their corresponding reductants were used by Cox and others (2003) to divide the SSC deposits into three subclasses. In the "reduced-facies type" subclass (Kupferschiefer-type of Kirkham and others, 1994), in situ solid organic matter is the principal reductant for pre-ore, early diagenetic pyrite and ore-stage Cu sulfides. Deposits of this subclass include those associated with the Kupferschiefer, the White Pine deposit in Michigan, and the ores of the Central African Copperbelt that are hosted by the Roan Group "ore shale" (Zambia) or by the D Strat, RSF, and SDB beds of the Roan (Democratic Republic of Congo). The host rocks of deposits are marine-shelf shales, siltstones, and (or) impure dolostones immediately overlying a transgressive flooding surface (Oszczepalski, 1989; Cailteux, 1994; Selley and others, 2005). Immediately adjacent rocks, up- and down-section, may also be mineralized to ore grade. In SSC deposits, the host rocks are commonly, but not everywhere, parts of thicker, transgressive, vertical sequences (Hayes, 1991; Hitzman and others, 2005), and they mark the change up-section from proto-oceanic, rift-fill, nonmarine deposits, which are typically red beds, to marine-shelf deposits of rift- and passive-margin sequences (Hoffman and others, 1974). In other places, the host rocks of reduced-facies SSC deposits are isolated gray or green beds within a thick stack of red beds (Smith, 1976). In "red-bed type" RBA-Cu deposits, the reductant is carbonized terrestrial plant matter, and the host rocks are fluvial sandstones or conglomerates (Cox and others, 2003). The reductant is more widely and diffusely distributed in "sandstone type" RBA-Cu deposits than in the red-bed type. Sandstone-type (often, hereafter, sandstone Cu) is synonymous with the Revett Cu subclass of Cox and others (2003). Much as the "Kupferschiefer-type" of Kirkham and others (1994) was replaced with the generic "reduced-facies-type," here, the Revett-type is replaced with the generic, "sandstone Cu-type," using a class name that first appeared in Russia in the late 1950s (Satpaeva, 1958; Domarev and Bogdanov, 1959). Nearshore marine and beach sandstones, as well as some sandstones deposited in fluvial or eolian environments, host sandstone Cu ores (Cox and others, 2003). Host rocks for sandstone Cu deposits are the topset beds of deltaic sedimentary packages, including nonmarine and nearshore marine sandstones. The reductant for the Spar Lake sandstone Cu deposit in Montana was a reservoir of sour gas (Hayes and others, 2012), and the reductant was probably natural gas or another form of petroleum for other deposits of the subclass (Lur’ye and Gablina, 1978; Gablina, 1981; Hayes and Einaudi, 1986; Gablina, 1997; McGowan and others, 2003; Thorson and MacIntyre, 2005; Syusyura and Tyugay, 2008).
Figure 6. Mineral zones of the Spar Lake sandstone copper deposit, Montana, United States, from Hayes and others (2012). Mineral-zone colors match generally with figures 4 and 5. The diagram illustrates the distribution of authigenic minerals only in space, not in time. Most changes in concentration of a mineral in space are gradual but are presented here as sharp to avoid confusion of this diagram with any paragenetic diagram.
Figure 7. Mineral zonation along the curving long axis of the Spar Lake sandstone copper deposit, Montana, United States, from Hayes and Einaudi (1986). Mineral-zone colors match with figure 6 and match generally with figures 4 and 5. A, The distribution of ore without vertical exaggeration.

B, The mineral zonation related to bedding with 10X vertical exaggeration from the same eight drill holes and two outcrop areas and with the assumption that all fault offset was post-ore. The assumption is true for the Cross Fault but not for the East Fault. However, only the southeastern end of this section would be affected, from drill hole 20 to the south outcrops. The change to correct the zonation section can be visualized. The bottoms of the high-grade ore intercepts, falling through section from southeast to north (black bars along the drill hole traces), were probably along a single gas/water interface of a gas reservoir at ore stage (Hayes and others, 2012).
In preparing this revision to Cox and others (2003), the three subtypes were statistically tested to determine if they were discrete from one another in grade and tonnage characteristics; results of the statistical tests determined that they are discrete. That work first required rigorous definitions of the three subtypes, and those definitions are provided in detail in a separate publication on grade and tonnage models. The larger broad type of SSC deposit is, however, clearly distinct from all other types of Cu-rich ore deposits everywhere in the world except, perhaps, in the Central African Copperbelt. In Central Africa, there appear to be multiple generations of economically important stratabound Cu deposits, the earliest of which have characteristics of SSC deposits, and the later ones with greater degrees of structural geologic control and, for some of them, distinct mineralogy (for example, Kansanshi and Lonshi; Broughton and others, 2002; Stephens, 2003; respectively). More is said on the distinction between early and late SSC deposits in a later section on Associated and Transitional Deposit Types.

The characteristic sulfide zonation of SSC deposits generally is accompanied by a parallel zonation of authigenic gangue minerals. Authigenic carbonates are generally peripheral to and extending into chalcocite-bearing rock from the pyritic zones, although the carbonates may be absent from highest grade ores where, instead, chlorite or illite are commonly present with authigenic hematite (Hamilton, 1967; Hayes, 1984; Michalik, 2001). Authigenic feldspars are present in many mineralizing systems, but their zonation is seldom documented. Authigenic K-feldspar is documented at deposits in Zambia (Annels, 1989; Sweeney and Binda, 1989; Selley and others, 2005) and at Spar Lake (Hayes, 1984; Hayes and Einaudi, 1986). Authigenic quartz also may accompany Cu-sulfide deposition in some deposits (Michalik, 2001, p. 50; Hayes, 1984). In some deposits, albite increases at the expense of K-feldspar in “upstream” directions for the metals-transporting solutions (for example, Oamites: Lee and Glenister, 1976; places in Zambia: Annels, 1989, p. 432 and p. 437–439; Sweeney and Binda, 1989; Selley and others, 2005; and Spar Lake: Hayes and Einaudi, 1986).

Similarly, the characteristic paragenesis of sulfides and oxides in SSC deposits is accompanied by parallel paragenesis of gangue minerals, many forming during the ore stage. Most of the ore and authigenic gangue minerals occur as cements (Hamilton, 1967; Anhaeusser and Button, 1973; Bartholomé, 1974; Hayes, 1984; Hayes and Einaudi, 1986; Hayes and others, 1989; Sawlowicz, 1990) and (or) they selectively replace certain clasts or earlier cements (Brown, 1971; Annels, 1974; Bartholomé, 1974; Banaś and others, 1982; Hayes and Einaudi, 1986). Replacement of early diagenetic pyrite occurred during deposition of ore-stage assemblages at many SSC deposits. Ore-stage cementation textures and preferential deposition of sulfides in coarser-grained beds or laminae in many deposits indicate that primary permeability was employed by ore solutions (fig. 8) (also, Hayes and Einaudi, 1986, fig. 5C; Borg and Maiden, 1989, p. 531–532 and 535; Richards and others, 1988a, p. 125).

Figure 8. Photomicrographs showing denser than background concentrations of copper sulfides in a silt-sized lamination and in a medium-crystalline dolomite nodule within dolomitic mudstone from the Kamoto Principal deposit, Democratic Republic of Congo. A, Transmitted light, plane polarized. B, Reflected light in air, plane-polarized. Bornite is lavender, and a chalcocite-group mineral is white.
**Associated and Transitional Deposit Model**

A wide variety of mineral deposits may be associated with SSC deposits, but only two of these (basalt-hosted native Cu deposits and volcanic red-bed Cu deposits) appear to have genetic association. Others appear to merely share a common sedimentary-tectonic setting with SSC deposits.

The most clearly recognizable associate is native copper deposits within flood basalts, with the best example being the Keweenaw Native Copper district of the Upper Peninsula, Mich. (White, 1968; Jolly, 1974; Livnat, 1983; Bornhorst and others, 1986; Brown, 2006; 2009). Such native copper deposits formed at temperatures from 150 °C to perhaps as much as 340 °C (Livnat, 1983, p. 75–96) from fluids that may be viewed as formation waters of flood basalts. The fluids redistributed Cu from higher temperature zones, where the basalts were altering or metamorphosing through dehydration reactions to epidote, into lower temperature zones where the native copper was deposited with chlorite, quartz, pumpellyite, laumontite, and analcime, and the wallrocks were altered to pumpellyite, laumontite, and analcime (Stoiber and Davidson, 1959; Jolly, 1974). Native copper precipitated in quartz veins, in amygdules and autobrecciated zones of basalt flow tops, and within interflow conglomerates (Stoiber and Davidson, 1959; White, 1968; Bornhorst and others, 1986). The White Pine SSC deposit is just 75 km from the nearest part of the main Keweenawan native copper district, and minor native copper deposits are found as close as 29 km from White Pine (White, 1968, p. 305). The Portage Lake lavas underlie the White Pine deposit, so, conceptually, it is possible that the same fluids that created the native copper ores also mineralized the lower part of the Nonesuch Shale at White Pine. However, geologic constraints seem to favor, instead, that White Pine second-stage Cu mineralization, which is more localized in veins and consists mainly of native copper, correlates in time with the mineralization of the Keweenawan Native Copper district (Mauk and others, 1992c).

Another SSC deposit where native copper in flood basalts has been found nearby is Witvlei, Namibia, where native copper is hosted in basalts of the Opdam or Doompoort Formation (Anhaeusser and Button, 1973; Borg and Maiden, 1989). A third deposit where native copper-bearing flood basalts occur nearby is Kurpandzhin, Russia (Ioganson, 1988).

Native copper deposits have been found in a number of additional basalts in the world, but without known association to SSC deposits. One such basalt is the ca. 1270 Ma Coppermine River continental flood basalt on the Arctic coast of Canada, which is associated with the Muskox layered mafic to ultramafic intrusion as part of the Mackenzie large igneous province (Gilbert, 1931; Lang and others, 1976; Day, 2008). Others include the Late Precambrian Volynian volcanic rocks of northwestern Ukraine (Kvansntsyva and others, 2001), the Tongchanghe deposit in the Late Permian Emeishan flood basalts of Chengdu, in south-central China (Zhang and others, 2006), and the middle Cretaceous Seachondong basalts in east-central South Korea (Lee and Kim, 1970).

Some of these same basalt sequences that host native copper deposits also host deposits of Cu sulfides having mineralogy and paragenesis that resemble those of SSC deposits. In Northwest Territories, Canada, with the Coates Lake (Redstone) SSC deposit less than 1,000 m upsection, the mafic lava unit of the Little Dal Formation commonly contains chalcopyrite disseminated in amygdules and locally has a decimeter-scale layer at the top consisting entirely of sand-sized chalcopyrite and detrital hematite grains together with lava pebbles and jasper and chaledony clasts (Ruelle, 1982, p. 710). The Korean deposits in the Seachondong basalts also have chalcocite (Lee and Kim, 1970). Cu-sulfide deposits have been found in the Keweenawan volcanic rocks (Bornhorst and others, 1986, p. 24). The Cu sulfides are fairly widespread, although typically paragenetically later than the native copper deposits. The largest accumulation of Cu sulfides in the Keweenaw is at Mount Bohemia and has zonation and paragenesis of chalcocite, bornite, chalcopyrite, and pyrite like that of the SSC deposits (Robertson, 1975). Recent exploration at the 47 Zone deposit has discovered potentially economic Cu-sulfide deposits in the Coppermine River basalts (Coronation Minerals, Inc., 2006) that are apparently quite similar to Mount Bohemia. The Sustut Cu deposit in central British Columbia, Canada, appears to be a related deposit type. It occurs in Triassic basaltic volcanioclastic rocks and consists of several tabular bodies each with a core of native copper zoned outward to chalcocite, then bornite, then chalcopyrite, and then pyrite (Wilson and Sinclair, 1988). The Kenneecott deposits in eastern Alaska are probably also related to the volcanic red bed type deposits like those above (MacKevett and others, 1997). The Kenneecott deposits, although having a relatively telescoped mineral zonation, have an identical paragenesis where pyrite was replaced by chalcopyrite, then by bornite, and then by chalcocite + djurleite. The Kenneecott deposits occur as veins and prism-shaped replacements within brecciated limestone immediately above a contact between limestone and the Nickolai Greenstone. The Nickolai is another Large Igneous Province, and in the Kenneecott vicinity, it is a sequence of subaerial basalt flows that is more than 1,800 m thick. Metalliferous fluids at Kenneecott are interpreted to have formed during prehnite-pumpellyite metamorphism of the greenstone, where native copper and Cu sulfides also precipitated in small concentrations (Price and others, 2004).

The above volcanic-hosted or volcanic-associated deposits all fall within the volcanic red bed Cu deposit class of Kirkham (1984b) and Lefebure and Church (1996). Those authors also place within that class some of the volcanic-hosted peneconformable “manto” Cu-Ag deposits of Chile, such as Mantos Blancos and El Soldado (Maksaev and Zentilli, 2002; Wilson and others, 2003). However, the Chilean manto deposits occur in the same region and generally within the same volcano-sedimentary sequence as iron oxide-copper-gold (IOCG) deposits such as Candelaria, Manto Verde, and Punta del Cobre, so it is possible, instead, that the association of the Chilean manto deposits is really with IOCG deposits.
In either case, there are many shared processes between the mantos and SSC deposits. The mantos have the same zonation from central chalcocite outward to bornite, chalcopyrite, and pyrite zones. They have the same paragenesis, where pyrite is replaced stepwise by chalcopyrite, bornite, and chalcocite. In addition, they have an abundance of pyrobitumen derived from degradation of oil where the petroleum was an ultimate reductant for S, and perhaps Cu, in forming the Cu sulfides (Wilson and Zentilli, 1999; Wilson and others, 2003). An apparent distinction is that the manto deposits were formed at higher temperatures of 200 to greater than 300 °C than are common for SSC deposits (Wilson and Zentilli, 1999).

Sandstone-hosted U deposits are a second deposit type associated with SSC deposits, particularly the tabular- or trend-type sandstone U deposits as are present in the Colorado Plateau. In fact, certain of the tabular U deposits in the Triassic Chinle Formation are hybrid types in that they are also red-bed-subtype stratabound Cu deposits (Hayes, 1982). Examples are in the Silver Reef district, southwestern Utah (Proctor, 1953), and include the Hide Out and Happy Jack deposits in the White Canyon district (Miller, 1955) and the Whirlwind deposit in the Monument Valley district (Isachsen and Evenson, 1955). Northrup and others (1990a, 1990b) demonstrated that the tabular V-U deposits of the Henry Basin, Utah, formed at an interface between fresh groundwater and a subsurface brine, where that interface crossed through fluvial sandstone host rocks with carbonized plant fossils. Hayes (1982) concluded that probably all other tabular- and trend-type sandstone-hosted V-U deposits were formed at subsurface groundwater interfaces where a freshwater/brine contact existed. Hayes further hypothesized that sandstone-hosted deposits that contain all three metals, Cu, U, and V, were formed where the brine was the metals-transporting fluid for SSC deposits.

U is present in considerable abundances in some of the deposits of the Central African Copperbelt. For example, the Malundwe and Chimwungo deposits in amphibolite-grade basement rocks underlying the Roan Group on the Mombezhi Dome have zones of recoverable U: Malundwe is estimated to have 9.9 Mt of ore containing 0.087 percent U₃O₈ among its 161.7 Mt of 0.89 percent Cu ore, and Chimwungo is estimated to have 2.2 Mt of ore containing 0.056 percent U₃O₈ among its 739.6 Mt of 0.66 percent Cu ore (Davis and others, 2008, p. 20 and 22). The U occurs as uraninite mainly at the hangingwall and footwall contacts of the Cu orebodies. The nearby Lubwe prospect also has substantial U grades. These, then, are a transitional type of deposit. Some of the reduced-facies-type SSC deposits of the Kolwezi klippe, Mutosho, for example, also had recoverable U (Francois, 1996).

Two deposits in the Central African Copperbelt are atypical in that their hypogene ores are dominated by structurally controlled veinlets and replacements of chalcopyrite. These are Kansanshi (Broughton and others, 2002) and Lonshi (Stephens, 2003). These can be classed as post-orogenic stockwork and replacement deposits in order to distinguish them from the main ores of the Copperbelt, which are pre-orogenic stratabound disseminations. Kansanshi consists of veinlets and large veins of quartz-carbonate-chalcopyrite with alteration envelopes of albite and ferroan dolomite, cutting phyllites, schists, and marbles. Deposits of the Central African Copperbelt all have at least minor amounts of Cu sulfides localized in veinlets (Dewaele and others, 2006; El Desouky and others, 2009), but, in most deposits, the veinlet-hosted mineralization provides far less of the total Cu than the earlier Cu sulfides that cement the sedimentary rock and selectively replace clasts and diagenetic minerals. Post-orogenic stockwork and replacement deposits are not known in other SSC districts, although most deposits have later stages of vein Cu mineralization (examples are: Hayes and Einaudi, 1986, figs. 14B and 16; Jowett, 1987; Mauk and others, 1992b, 1992c), and the Kupferschiefer deposits in Germany are cut by Co-Ni-arsenide veins, called rücken, that occupy fault zones (Beck, 1905). A third atypical Cu deposit in the Copperbelt is Frontier. Frontier has classic mineral zonation from pyrite, to chalcopyrite, to bornite, to chalcocite, and to hematite. However, the zonation is distributed surrounding a fault at a low angle to bedding and in atypical host rocks (D. Jack, First Quantum Corp., oral commun., 2010). The fault is a probable feeder of oxidizing Cu-transporting solutions to the deposit.

Kipushi-type carbonate-hosted Zn-Pb-Cu-Ge deposits were listed as SSC deposit associates by Cox and others (2003), although that applied almost solely to the Central African Copperbelt where the Kipushi, Kangere, and Lombe deposits are present. These deposits are sited along faults and in breccia of Nguba Group carbonate rocks that are upsection from the Roan Group that contains the bulk of the SSC ore. The observation by Cox and others (2003) has since been further strengthened because SSC mineralization has been identified at Sebembere and Mufukushi near the Kabwe (formerly Broken Hill) Kipushi-type deposit in central Zambia (Kampunzu and others, 2009). Also, the second-best example of a Kipushi-type deposit is the Tsumeb deposit in northern Namibia, and a sandstone Cu deposit has been identified at Tschudi in the rocks immediately overlying the host rocks of Tsumeb (Kamona and Günzel, 2007). Tschudi is just 26 km west of Tsumeb.

Hydrothermal U orebodies may also be added to the list of SSC associates because the Shinkolobwe U deposit occurred between the Mombezhi Dome and the SSC deposits of the Kolwezi Klippe. Shinkolobwe was a “hypothermal” breccia and stockwork of veinlets cutting the SSC Cu host beds and consisted of uraninite, nickel- and Co-bearing sulfides, mazonite, and molybdenite, with quartz and chlorite, within a magmatic alteration zone (Ridge, 1976). Other deposits along a west-trending belt from Shinkolobwe are U-rich post-orogenic stockworks (Francois, 1996).

Hovee and Quirt (1989) suggested that unconformity-type U deposits have a common diagenetic-hydrothermal origin with SSC deposits. However, to our knowledge, there are no examples of SSC deposits and spatially associated unconformity U deposits.
Sediment-Hosted Stratabound Copper Deposit Model

IOCG deposits have spatial and temporal association to SSC deposits in several regions including, as above, the Coastal Cordillera of Chile, where volcanic-hosted manto deposits are associated with SSC deposits. An alternate hypothesis for the occurrence of SSC deposits is that they are Ni-rich portions of ordinary SSC deposits, related to gabbroic intrusions. This is supported by the fact that the Chinei gabbroic complex lies between Udokan and Pravo Ingamakat, and the deposit is compared with “outer contact ore of the Chinei Massif” (Gongalsky and others, 2007); a suite of late Lufilian gabbros intrudes the metasedimentary rocks in the area of the Kalumbila deposit (Steven and Armstrong, 2003).

Last and possibly most important of all the mineral deposit types that are associated with SSC deposits are evaporite deposits including gypsum, halite, and potash. This association has long been recognized and has been discussed frequently, typically calling on dissolution of the evaporites to contribute to salinity and complexing capacity for a metals-transporting groundwater (Davidson, 1965; Rose, 1976; Thiede and Cameron, 1978; Hayes, 1983; Warren, 2000). Many different rock sequences that host SSC deposits also host evaporite deposits. The most familiar of these may be the Zechstein sequence in central Europe. The Kupferschiefer bed is the basal bed of the Zechstein sequence. Although millions of tons of salt (halite) have been produced from stratigraphically higher levels of the Zechstein, there is only a small area of the basin in central Germany where there was Kupferschiefer-hosted Cu production and production of salt from the overlying Zechstein evaporites. For the most part, the Cu-producing part of the basin is along its southern margin, and the salt-producing part is to the north nearer the basin center. Many of the salt beds of the central basin are lower in elevation than the Kupferschiefer bed along the southern basin margin, so there remains a hydrologic problem in getting saline Cu-transporting solutions from Zechstein evaporites for formation of the Kupferschiefer deposits. Brine is typically denser than other waters and would gravitate to the base of a pore water column, unless its density is lowered by anomalous heating. Some external hydrologic drive is required to move the brine to the higher basin edges (Hitzman and others, 2005).

The Central African Copperbelt deposits commonly contain anhydrite (Annels, 1974; Muchez and others, 2008), but far more evaporites may have existed in the Katangan Supergroup at some earlier time. Jackson and others (2003) have interpreted that the Roan Group of the Katangan Supergroup once had evaporites at four separate stratigraphic intervals. “Flakes, slices” (ecailles) of coherent, mineralized Mines Series rocks, 1 to 10 km scale, float within Roan 1 breccia, which may have once been predominantly halite and other salts that are now dissolved. Damesmacher and others (1963) and Jackson and others (2003) interpret that dêcollements formed along those four salt intervals in producing the thrust stack that now exists in the Copperbelt in Democratic Republic of Congo. Given that hypersaline fluids have been documented in the ores of the Katangan (Dewaele and others, 2006; Heijlen and others, 2008; El Desouky and others, 2009), there is strong evidence for the involvement of evaporites in ore genesis been directly tested, and the result indicates that
they were likely involved. At Lisbon Valley, Utah, Breit and Meunier (1990) measured strontium isotope ratios of SSC ore-related fluid inclusions to be very radiogenic. They interpreted that gypsum and anhydrite of the Paradox Member of the Hermosa Formation that underlies the Lisbon Valley deposits by about 1–2 km was dissolved, contributing to the strontium of the inclusion fluids. Those fluids presumably dissolved Cu from the Cutler Formation arkoses that lie between, enroute to the stratabound Cu deposits.

**Primary Commodities**

Cu is the primary commodity in all SSC deposits. Co and Ag reach co-product concentrations in some, and Au and platinoids are produced from deposits of the Kupferschiefer in Poland, along with several minor metals (see below).

**By-Product Commodities**

The SSC deposits are the world’s most important source of Co. Cobalt is produced from about 10 deposits in Zambia, from most, but not all of the deposits in the Democratic Republic of Congo, and as a byproduct from the Kupferschiefer deposits in Poland. Current plans for mining the Bofélo deposit in Baja California, Mexico, call for recovery of Co, which occurs there at an average grade of about 0.07 wt. %. In 1990, for example, the Democratic Republic of Congo and Zambia produced 26,086 t of Co from SSC deposits, whereas the rest of the world combined produced only 9,736 t, mostly from other deposit types (calculated from http://minerals.usgs.gov/minerals/pubs/commodity/Co/lem/stat/tb9.txt).

Ag is produced as a co-product metal from the Spar Lake deposit in Montana and as a byproduct metal from the Kupferschiefer deposits in Poland and the Dzhezkazgan deposits in Kazakhstan. As a Ag-producing deposit type, the SSC deposits rank fourth in the world, behind epithermal, porphyry Cu, and sedex Zn-Pb-Ag-barite deposits (as calculated from statistics presented in Brooks, 2010).

Au, platinum (Pt), and Pd are now recovered from the Kupferschiefer deposits in Poland. They occur disseminated in a layer along the footwall of the Cu ore that parallels the mineral zonation of the Cu deposit slightly discordant to the bedding (Piestrzynski and others, 2002). Piestrzynski and others (2009) reported that the Au and platinoids deposit subjacent to Kupferschiefer Cu ore in the western part of the Polkowice mine averages 22 cm in thickness at 2.25 g/t Au, 0.14 g/t Pt, 0.08 g/t Pd, and 0.025 percent Cu.

The Kupferschiefer deposits of Poland also yield Pb, Co, Mg, nickel, selenium, and S for sulfuric acid (Pieczonka and others, 2001).

U and V were typically more valuable than Cu or Ag in some of the Triassic-hosted red-bed-type SSC deposits of the Colorado Plateau, United States (Miesch, 1963; Hayes, 1982). U is likely to be produced from the Malundwe and Chimiwungo deposits in Zambia (Davis and others, 2008).

**Trace Constituents**

Mn, Zn, Pb, Ba, Re, Hg, and As are present at anomalously high concentrations in some SSC deposits (Bakun and others, 1966; Nimry, 1973; Rentzsch, 1974; Seyfullin and others, 1974; Hayes and Nuckolls, Asarco, Inc., written commun., 1983; Sutton and Maynard, 2005), and are elements that were probably enriched together with Cu at ore stage. Example concentrations of these elements are given in the later section on trace-element geochemistry. Titanium is anomalous in numerous SSC deposits, but it is likely enriched in a detrital component of the host sedimentary rocks (Dimanche and Bartholomé, 1976). B is anomalously high in some Zambian Copperbelt deposits (Darnley, 1960, p. 160 and 162), where it is concentrated in authigenic tourmaline (Selley and others, 2005). Darnley (1960) concluded that the enrichment factor was about six times the B background in ore in Zambia. Mg, found in chloride in many deposits, and in magnesite in deposits of the Democratic Republic of Congo, is anomalous in some SSC deposits, though not at Konkola, Zambia (Sutton and Maynard, 2005).

**Example Deposits**

Cox and others (2003) reported grade and tonnage data for 101 well-explored deposits worldwide and used these data in their sediment-hosted Cu deposit grade and tonnage models. In addition to Cu grades, their compilation included Co and Ag grades. The compilation by Cox and others included 682 additional deposits and SSC mineral occurrences that lack reliable grade and tonnage data. Parks and others (written commun., 2011) contains summaries of more than 1,000 deposits and occurrences that highlight the diverse characteristics and settings of SSC deposits. It includes deposits newly described since Cox and others (2003), but a number of deposits were removed from the Cox and others compilation on the basis that they did not meet stricter definitions of the deposit type. Summaries of representative sediment-hosted stratabound copper deposits are discussed in Appendix 1:

- **A.** Kupferschiefer, Lubin-Sieroszowice-Polkowice-Rudna-Glogow Giboki-Przemyslowy-Bytom Odrzanski Retkow-Gaworzysie-Radnawice Zachody, Poland;
- **B.** Kolwezi Klippe, Democratic Republic of Congo;
- **C.** Creta, Okla., United States;
- **D.** Udokan, Russia;
- **E.** Spar Lake, Mont., United States;
- **F.** Juramento, Argentina;
- **G.** Corocoro, Bolivia;
- **H.** Nacimiento, N. Mex., United States;
- **I.** Happy Jack, Utah, United States.
History of Exploitation of Sediment-Hosted Stratabound Copper Deposits and History of Genetic Concepts

Archeometallurgy of Copper and of Sediment-Hosted Stratabound Copper Deposits

SSC deposits were among the earliest types of metal deposits mined and processed at an industrial scale. Before mining occurred at an industrial scale, tools and jewelry items of smelted Cu were made at a household scale, and yet before that, native copper was used in similar ways. Archeometallurgy has competing claims for the oldest occurrences of each of these types of Cu usage, and Cu even has its own named archeological period, the Chalcolithic. This was the period when Cu was the most advanced material in use, between the Neolithic (new Stone Age) and the Early Bronze Age, when there first was a metals industry at a small industrial scale and when the first man-made alloys, tin-bronze and arsenic-bronze, appeared. Cu was the first metal used for casting tools, probably because of its low melting temperature of 1,083 °C.

Modern-day Turkey supplied the oldest known Cu artifacts, consisting of beads, pins, awls, and fishhooks dated to 8505 ± 175 B.C. from Çayönü Tepesi, Anatolia (Maddin and others, 1991; as recalculated in http://context-database.uni-koeln.de/). The source of the Cu for the Çayönü artifacts was almost certainly Ergani Maden, a Cyprus-type volcanogenic massive sulfide deposit (Yigit, 1999). A single 6900 B.C. Cu bead was found at Ali Kosh in southwestern Iran (Smith, 1969). At 6000–5900 B.C., native copper was used to make a heavy mace head at Can Hasan, again in modern-day Turkey (Yalçin, 2000), and, by that time, native copper was in use in at least five sites in Iran (Thornton, 2009).

The claim to the earliest true Cu industry comes from eastern Serbia at Prokuplje, where evidence for smelting of Cu oxide ores and casting of a chisel, a 2-headed hammer, and an axe date to 5500 B.C. (Baird, 2007). A bonafide Cu industry has been proven nearby at Belovode, Serbia, that dates confidently to 5000 B.C., and that includes irrefutable evidence of smelting of ores, not merely melting of native copper fragments, associated with production of a variety of tools and jewelry (Radiyojevic and others, 2010). The specific source of the manganiferous and Co-bearing ore for Belovode has not been identified, but Belovode and Prokuplje are within a belt of porphyry Cu and epithermal Cu-Au deposits (Berza and others, 1998). Perhaps spanning that same time is a small-scale Cu industry at Tal-i Iblis, Iran, dated from 5470 to 4250 B.C. (Evett, 1967; as explained in Thornton, 2009). Tal-i Iblis is about 125 km from the Sar Cheshmeh porphyry Cu deposit (Waterman and Hamilton, 1975), which could have supplied the Cu.

Also at some time from 5266 to 3994 B.C. (Joffe and Dessel, 1995), there was an artisanal use of Cu ore at Shiqmim, Israel, including crucible-scale smelting and casting. The ore for Shiqmim originated from SSC deposits near Feinan, Jordan, about 150 km to the south and east (Golden and others, 2001, p. 954). Ores used at Shiqmim are distinctive and identifiable as the “tile ores” of Feinan, from its Massive Brown Sandstone unit. Trace-element analyses also match the Shiqmim ore pieces with Feinan. Furthermore, a single 14C date for a “Pottery Neolithic-aged” site at Wadi Feinan, which, in contrast, doesn’t give evidence for a mining-centered lifestyle, is 5500 to 5270 B.C. (Hauptmann, 1989, p. 119), a near match to the earliest Shiqmim dates. Unless earlier sites are found, Feinan appears to be the first-used SSC deposit.

According to current archeological knowledge, Feinan is also the site of the earliest industrial-scale exploitation of a SSC deposit, with the actual demonstration being the smelting workshop village excavated at Wadi Fidan 4 with 14C dates of 3630 to 2920 B.C. inclusive of uncertainties (Hauptmann, 2000, 2007; Weisgerber, 2006). Note, though, that the apparent 1,600 year delay between artisanal and industrial-scale usage of the deposit, through a time interval during which industrial-scale Cu metal production and even early attempts at making tin-bronze and As-bronze alloys were proceeding in southeastern Europe, Anatolia, and Iran. The Feinan dates are, nonetheless, consistent with the appearance of objects made from Cu metal in Mesopotamia and Egypt. Written language appeared at nearly the same time. Cuneiform writing in Mesopotamia apparently arose around 3900 B.C. (Kramer, 1981), and hieroglyphic writing reached a useful development in Egypt at around 3300 B.C. (Mattenisch, 2002). Not long after Feinan, though, the SSC deposits in Permian sedimentary rocks along the West Ural Foreland (Lur’ye and Gablina, 1972) were mined and smelted at a large area of archeometallurgical sites collectively called Kargaly that carry 14C ages that cluster near 3000 B.C. (Chernykh, 2004; Chernykh and others, 2004). Ages like this (3600 to 3000 B.C.) in Iran (Thornton, 2009) and slightly younger ages marked the beginning of the Bronze Age across almost the entire Old World, including China (Linduff and Mei, 2009) and Pakistan/India (Hoffman and Miller, 2009). Feinan later became a large integrated mining and Cu/bronze metallurgical center that, along with Timna, Israel, which is probably a strike-slip offset segment of the same deposit on the Dead Sea Transform, and a little more than 100 km to the south (Freund, 1965, p. 193), provided Cu for the Early Bronze Age rise to prominence of Egypt’s Old Kingdom (Rothenberg and Merkel, 1995; Weisgerber, 2006), and probably also provided some Cu to the Sumerian civilization of present-day Iraq. Feinan continued producing into the days of Old Testament Biblical Judea, when it was known as Edom (Levy and others, 2004). It continued as a Cu mining and production center into the Roman era when it produced its largest tonnages (Weisgerber, 2006; Hauptmann, 2007). A considerable resource of ore remains present near Feinan in three separated prospect areas (Nimry, 1973; Rabb’a and Nawasreh, 2006), but the largest of those known resources is within Jordan’s Dana Nature Reserve.
The archeological dating shows that perhaps 3,500 years passed between man’s first uses of Cu tools made from native Cu and the advent of the first primitive metals industry. Another 1,500 years passed before Egypt and Mesopotamia entered the Bronze Age using Cu from Timna and Feinan.

Turning back to the northern area of Bronze Age exploitation of SSC deposits, Kargaly was mined increasingly from 3000 to 2000 B.C. and hit a zenith of production about 1500 B.C. apparently contributing Cu metal for an enormous area of the Middle Asian steppes. The mining and Cu smelting then mysteriously and apparently abruptly ended about 1400 B.C. and did not return until about A.D. 1745 (Chernykh, 2004). Attempting to identify the cause of collapse of Kargaly is the subject of ongoing archeological research (Chernykh, 2004; Chernykh and others, 2004).

**Evolution of Concepts of SSC Deposit Genesis**

Jowett (1991) wrote a thorough earlier review of the history of genetic concepts for SSC deposits focused on the deposits of the Kupferschiefer in Germany and the deposits of the Zambian Copperbelt. The following draws heavily on that paper.

Earliest consideration of the genesis of the Kupferschiefer Cu ores favored a syngenetic origin, synchronous with the deposition of the shale. Mining of the Kupferschiefer at Mansfeld began in 1199 (Beck, 1905, p. 492). The first paper to discuss genesis of the Kupferschiefer deposits suggested that precipitation of Cu took place on the sea floor, essentially the first expression of a syngenetic theory (Lehman, 1756). From then through most of the 19th century, most scientists accepted a syngenetic origin as further evidenced, they thought, by the curled-up forms of fish fossils in the Kupferschiefer that were said to be poisoned by the metals-bearing seawater (Freiesleben, 1807–1817).

Epigenetic theory began first with the observation that Cu content was not uniform, varying even to barren Kupferschiefer outside the Mansfeld area (Beck, 1905, p. 491). Cross-cutting faults containing Co and nickel sulfides (Rücken) were called upon as feeders to the shale-hosted ores, and it was shown that curled fish fossils were found in other shales that lacked metal enrichments and even in recent sediments. Beysschlag and others (1916) observed that ore grades decreased upward within the shale and that grades did not follow the Kupferschiefer bed in dip and strike, but rather crossed it at a small oblique angle. Thus, through the early decades of the 20th century, opinion shifted to favor an epigenetic origin for deposits of the Kupferschiefer.

Schneiderhöhn (1923) led the next wave of syngeneticians when he interpreted the chalcopyrite microspheres of less than 0.2 μm diameter in the Kupferschiefer as replaced bacteria. He then took his syngenetic theory to the Zambian Copperbelt, where comparisons of the sulfide paragenesis with Butte-style veins (Bateman, 1930) had earlier been used to support epigenetic theories. Schneiderhöhn (1932) observed that the Copperbelt granites were not intrusive into the Roan Formation host rocks, but rather they were in nonconformable contact, so the granites could not have provided epigenetic magmatic hydrothermal fluids to the Ore Shale. Garlick (1961b) developed his theory of syngenesis to produce the various sulfide zones with increasing distance from a paleoshoreline—chalocite nearest the shore, then bornite, chalcopyrite, and pyrite farthest into the basin, as would be predicted by the relative solubility of those sulfides if Cu entered from the continentward side dissolved in river water. Vertical zonation of pyrite over the Cu sulfides could then be explained by transgression of an offshore pyrite-forming environment above the earlier nearshore chalcocite-depositing one. The elegance of the concept gave it great popularity that has not totally disappeared to this day. However, increasingly detailed study of the rocks and the application of better analytical tools do not support syngenesis.

Careful petrologic studies, some that also used geochemistry, started the movement toward today’s preference for depositing SSC deposits during diagenesis. Darnley (1960) described metasomatic changes to the rocks accompanying ore in Zambia and their corresponding anomalous geochemical signatures. Bartholomé (1962) at Kamoto and Brown (1971) at White Pine recorded very detailed paragenetic sequences, and both concluded that mineralizing solutions had invaded the reduced-facies hosts from underlying red beds. Annels (1974) demonstrated that Cu-rich sulfides in shale ores of Zambia replaced anhydrite, also favoring a diagenetic timing. Rentzsch (1974) integrated a number of different methods and, most importantly, demonstrated that Rote Fäule alteration zones were correlated to ore deposits of the Kupferschiefer, as well as being related to basin margins for underlying Rotliegendes red beds rather than to early Zechstein shorelines. In support of basinal brines from red-bed packages as metals-transporting solutions in diagenetic or epigenetic ore genesis, Rose (1976) demonstrated that chloride brines in equilibrium with hematite at intermediate oxidation states were capable of dissolving sufficient concentrations of Cu to act as ore solutions even at low temperatures.

Studies published in the 1980s continued to move away from syngenesis and, by the end of the decade, few economic geologists still held to that theory. Jowett and others (1987) demonstrated with a paleomagnetic study of Rote Fäule rocks of the Kupferschiefer in Poland that the ore-related zone had formed at ca. 240 to 230 Ma, affecting host rocks that were deposited at about 256 Ma, a strong demonstration of a non-syngenetnic origin for those deposits. Polar wandering curves have since been revised, moving the age of the Rote Fäule to 255 to 245 Ma (Nawrocki, 2000), but dating of the deposition of the Kupferschiefer also improved, and it was deposited from 260.4 to 258 Ma (Menning and others, 2006; Słowakiewicz and others, 2009). At Spar Lake, Mont., Hayes and others (1989) established with S-isotope geothermometry that the mineral zones had formed across a temperature gradient of as much as 100 °C, with temperatures in chalcocite-depositing zones greater than 130 °C. If the sulfides formed during syngenesis,
then they at least did not form as Garlick (1961b) envisioned at ambient temperatures. Diagenetic or epigenetic addition of isotopically heavy sulfide was shown for White Pine and Kamoto (Hoy and Ohimoto, 1989). Also, Annels (1989) reported ore-stage fluid-inclusion homogenization temperatures that averaged 130 °C at Chambishi Southeast and 147 °C at Chambishi in Zambia. Ore-fluid salinities of 16 to 21 eq. wt. % NaCl were measured. Breit and Meunier (1990) demonstrated, using Sr isotopes of fluid-inclusion fluids, that the Paradox evaporites were the most likely source for ore-fluid salinity in the origin of the sandstone Cu deposits at Lisbon Valley, Utah. Hayes (1990) studied fluid inclusions in quartz overgrowths and manganiferous calcite cements at Spar Lake and noted homogenization temperatures that varied dramatically across the mineral zonation. Homogenizations varied from 74 to 130 °C in galena-bearing rock, and in chalcocite- and bornite-bearing ore zones, homogenizations occurred from 130 to 184 °C. The regularly varying homogenization temperatures and a large salinity range were interpreted to record mixing between hot, saline metals-transporting brines and cooler, dilute formation water. The salinities were from less than 5 eq. wt. % NaCl to greater than 20 eq. wt. % NaCl, increasing toward the chalcocite- and bornite-bearing ore zones.

Since Jowett’s (1991) summary of studies on genesis of SSC deposits, new studies have further helped to understand the origin of this deposit type. From the Central African Copperbelt, new fluid-inclusion microthermometric studies were reported by McGowan and others (2006) at Nchanga, Zambia; by Dewaele and others (2006) at Musonoi and Kamoto, Democratic Republic of Congo; by El Desouky and others (2008a) at Lufukwe prospect, Democratic Republic of Congo; by El Desouky and others (2008b) at Mwitapile prospect, Democratic Republic of Congo; and by El Desouky and others (2009) at Luwiswishi and Kamoto, Democratic Republic of Congo. Referring only to results that did not come from syn- or post-orogenic veinlets, all these studies found large ranges in homogenization temperatures and in salinities, but with considerable consistency of the ranges. Diagenetic quartz that probably replaced anhydrite and that is found with Cu- and (or) Co-bearing sulfides precipitated at temperatures from 100 °C to 220 °C. Large ranges in salinities should probably be interpreted as mixing of saline melliteferous brines with relatively dilute pre-ore formation waters. Organic geochemical studies addressed the Nonesuch Shale of the White Pine deposit and the Kupferschiefer (Hieshima and Mauk, 1990; Ho and Mauk, 1996; Pratt and others, 1991; Püttmann and others, 1987; Püttmann and others, 1989; Sun and Püttmann, 1997; Bechtel and others, 2002). Nonesuch Shale Pristane/n-C17 and Phytane/n-C18 ratios at White Pine are lower within ore than outside of it, suggesting thermal maturation of the lower Nonesuch during mineralization. In the Kupferschiefer, a number of measures were found to indicate a higher degree of oxidation in the basal section interpreted to be caused by maturation and oxidation of organics in the shale by an ascending, oxidizing, metalliferous brine.

The 1990s and early 2000s saw the first attempts to isotopically date SSC mineralizing systems (Bechtel and others, 1996, 1999; Selley and others, 2005). These met with limited success. Illite K-Ar dates on Kupferschiefer mineralization were interpreted to indicate that mineralization occurred between 216 and 190 Ma, a rather long interval much different than the ages (for example, 255–245 Ma) estimated from a paleomagnetic study of the Rote Fäule (Jowett and others, 1987). Other dates on illite (Michalík, 1997) yielded ages as young as 188 Ma and thus led to otherwise poorly supported hypotheses of multiple times of mineralization (Michalík and Sawłowicz, 2001). Isotopic dating in the Central African Copperbelt indicated veinlet-dominated mineralization formed after the Lufilian Orogeny at ca. 510 Ma (Richards and others, 1988b; Selley and others, 2005, p. 983), but the earlier main stage of disseminated mineralization yielded an imprecise Re-Os age of 816 ± 62 Ma (Selley and others, 2005, p. 983).

## Regional Environment

### Temporal and Secular Environment

Kirkham (1989) described the temporal and secular environment of SSC deposits. He showed that among 820 deposits and occurrences, rocks of the Middle and Late Proterozoic (110) and the Permo-Carboniferous (290) contain the greatest numbers of deposits and occurrences. The distribution of deposits is strongly controlled by paleolatitude, with most deposits and associated evaporites located within 20 to 30 degrees of the paleo-equator at the times of host-rock deposition.

Hitzman and others (2005) showed that, by tonnage of contained Cu, there are three maxima through geologic time in the distribution of ages of SSC deposit host rocks (fig. 9). Kirkham (1989) had earlier found three maxima in the number of deposits by age, duplicating the tonnage maximum from the Neoproterozoic but with maxima in the number of deposits in the Carboniferous and Cretaceous, rather than in the Permian and Tertiary as by contained metal. By comparison with continental drift maps (Smith and others, 1981; Scotese and McKerrow, 1990) and with knowledge of the sedimentary tectonic environments of SSC deposits (see below), the time distribution of host rocks is likely showing a correlation to the rifting apart of supercontinents, Pangea in the Permian and Rodinia in the Neoproterozoic. That correlation is further explained in the following section.

### Geotectonic Environment and Relations to Sedimentary Rocks

If thick sections of red beds are classified strictly into sedimentary-tectonic environments, they fall into only two environments of classical Wilson cycles (Wilson, 1966; Hoffman and others, 1974), which are sediments deposited in protoceanic rifts and in post-collisional molasse. The SSC
respective.

age of host rocks, from Hitzman and others (2005). Pt 1, 2, and 3, are Paleoproterozoic, Mesoproterozoic, and Neoproterozoic, respectively.

Figure 9. Relation between contained tonnes of copper metal in sediment-hosted stratabound copper deposits and the depositional age of host rocks, from Hitzman and others (2005). Pt 1, 2, and 3, are Paleoproterozoic, Mesoproterozoic, and Neoproterozoic, respectively.

The sandstone or carbonate grainstone host rocks of sandstone Cu deposits include coarse-grained rocks from a variety of depositional environments, although all can be generally grouped as being parts of deltaic topset beds. Hayes and Einaudi (1986) deduced that the lower quartzite beds of the upper member of the Revett Formation host rocks at Spar Lake were deposited in a prograding beach and bar nearshore environment and that the middle quartzite host rocks were deposited on subtidal bars and by laterally shifting subtidal channels. Dakota Formation host beds at the Lisbon Valley deposit were deposited in similar environments (Ridgley, 1990), but the underlying Burro Canyon Formation host rocks at Lisbon Valley were fluvial sandstones (Craig, 1982). Nearby, at the Cashin deposit, the Entrada Formation sandstone Cu deposits include coarse-grained rocks from the Rotliegendes red beds that are elsewhere below the Kupferschiefer evaporites upsection from the Kupferschiefer. The lacustrine evaporites and the salts among the shelf carbonates can locally reach great thicknesses, and halokinesis may have occurred. Jackson and others (2003) have interpreted pre-ore salt horizons at three major and two minor levels in the ore-hosting Roan Group of the Central African Copperbelt and indicated that the salt first underwent extensive diapirism, then subsequently the salt defined the thrust planes of décollements, and finally the salt dissolved entirely, leaving heterolithic breccia.

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Sediment-Hosted Stratabound Copper Deposit Model

Some carbonaceous wacke where ore grades are highest, are separated by thin beds of coastal sabkha deposits. The sabkha mudrocks grade laterally into stromatolitic reef carbonates where the sequences onlap paleohighs. The host sandstones themselves logically were coastal and nearshore sand deposits.

Red-bed-subtype deposits are hosted by fluvial sandstones (Malan, 1968; Woodward and others, 1974; LaPoint, 1976; Flint, 1989) and less commonly by sandstones from deltaic environments (Smith, 1976). For these red-bed-subtype deposits, the host beds are commonly the only non-red rocks within thick sequences of red beds.

The most important feature of SSC host rocks is their content of reductants. These include plant or animal (cyanobacterial) fragments for the red-bed-subtype and reduced-facies-subtype deposits. The reductants were liquid or gaseous petroleum mixtures for the sandstone-subtype deposits. These reductants react with cuprous chloride and cuprous sulfate ions in oxidizing metals-transporting brines (Rose, 1976), reducing the sulfur and causing deposition of Cu sulfides.

Structural Setting(s) and Controls

The most important geometric aspect of SSC deposits is that they are sediment-hosted and peneconformable. Thus, they seem to be controlled almost entirely by bedding and lamination of their sedimentary host rocks. For each district, only one limited sequence of beds is mineralized to ore grades; in other words, the deposits are stratabound. Increasingly, however, it is being recognized that there are also important structural controls on locations of SSC deposits.

Faults at high angles to the host beds are found at many deposits. Many of the faults, however, simply cut and offset mineralized rocks. The importance of certain faults in ore genesis has been debated for years (for example, the White Pine fault at White Pine; Ensign and others, 1968; White, 1968, particularly in sandstone Cu deposits (for example, the East fault at Spar Lake; Hayes and Einaudi, 1986, p. 1927; Hayes, 1990, p. 27; Hayes and others, 2012). It can commonly be proven that the host beds are offset by faults that are suspected...
to have been genetically involved, but the timing of the offset relative to ore genesis is unclear, and the possibility that the faults were feeders for metals-transporting solutions cannot be discounted.

In the Kupferschiefer and Zambian Copperbelt deposits, faults at high angles to the host beds are present. They cut and offset the basement rocks but do not cut the host packages. These faults were the margins of subjacent grabens or half-grabens (fig. 11).

**Relations to Igneous Rocks**

The SSC deposits are notable in their total lack of any clear genetic relation to igneous rocks. The sparse igneous rocks in the same sedimentary sequences with SSC deposits are typically mafic sills and rarely mafic lavas, and the economic Cu deposits have no relation to those igneous rocks. In the clearest example, from the Udokan deposit in Russia, the acidic igneous dikes or sills simply intrude the stratabound mineralization producing contact metamorphism of the ore, but only minor local depletion of the Cu grades in the sedimentary wallrocks (Bakun and others, 1966, p. 455; Bogdanov and others, 1966, p. 1312).

**Relations to Metamorphic Rocks**

A number of important SSC deposits are completely unmetamorphosed, but many SSC deposits have been metamorphosed along with their enclosing sedimentary-rock sequences. A simple conclusion drawn from that is that ore deposition tends to pre-date regional metamorphism. The deposits of the Central African Copperbelt are all at least lower greenschist facies and the metamorphic grade is upper greenschist for deposits from Nchanga southward in Zambia (Hitzman, 2000) reaching epidote-amphibolite facies in the Roan Basin (Mendelsohn, 1961b). Deposits of the Udokan Series are at lower greenschist grade, although the rocks have no schistocity, whereas the base of the series is upper greenschist grade and is schistose. Deposits in the Missoula and Ravalli Groups of Montana are at lower greenschist facies and at lower to upper greenschist facies, respectively. Greenschist facies metamorphism of SSC deposits has typically been noted to produce no important change to the deposits themselves.

A number of important deposits are at amphibolite or perhaps higher metamorphic grade. Aynak, Afghanistan, and other deposits along the enclosing belt of Neoproterozoic to Early Cambrian schist-, marble-, and quartzite-hosted deposits.
are at amphibolite grade (Peters and others, 2007, p. 276). The deposits of the domes region west of the Zambian part of the Central African Copperbelt are also at amphibolite grade (Benham and others, 1976). In the deposits of the Domes Region, the main host rock was earlier interpretively correlated with the Copperbelt Orebody Member. The mineralized rock is now graphitic muscovite-phlogopite-kyanite schist, with feldspar-biotite schists that are interleaved. Feldspar-biotite gneisses and schists downsection, toward the core of the dome, were speculatively correlated with the red beds of the Mindola Clastics Formation. Rocks correlated with Roan impure stromatolitic dolomites are now para-amphibolites, although rocks correlated with purer carbonates of the Mwashya Formation, upsection, are dolomite marbles. Primary chalcocite is not present, and the ore sulfides are bornite and chalcopyrite. Formerly pyritic peripheral zones now contain pyrite and pyrrhotite. More recently, Davis and others (2008) have revised correlations within the Domes Region, and they suggest that the host rock formed as sheared, thrusted, and possibly nappe-folded zones within the Middle Proterozoic basement rocks beneath the Roan Group. The Oamites SSC deposit in Namibia, probably once a sandstone-subtype deposit, is metamorphosed to upper greenschist to amphibolite grade. In it, sandstone ore host rocks were metamorphosed to feldspathic-biotite-quartzite, and impure carbonate beds were metamorphosed to actinolite-tremolite-biotite schists. Sulfide zonation appears to be almost completely preserved, and “sulfide facies boundaries are roughly parallel to the oxidized lithofacies/reduced lithofacies boundaries and cut across the lithozone contacts” (Lee and Glenister, 1976, p. 383).

### Physical Description of Deposits

#### Deposit Geometry

The SSC deposits are confined to a limited sequence of sedimentary beds of each basin where they are found. Lateral dimensions are large relative to deposit thickness in all three deposit subtypes, dramatically so for the reduced-facies deposits.

Reduced-facies deposits have the geometry of sheets. Sandstone Cu deposits are tabular to lens shaped. Red-bed Cu deposits are lenses of Cu-bearing rock.

Aerial extents and thicknesses of sediment-hosted Cu deposits have been described by Gustafson and Williams (1981). Reduced-facies and sandstone Cu deposits are the largest. Sizes of sediment-hosted Cu deposits are strongly affected by post-depositional faulting and erosion.

Reduced-facies-type deposits in Central Africa have strike lengths from 3,000 to 5,300 m, widths from 550 to 2,000 m, and ore thicknesses from 2 to 70 m. The vertical extent perpendicular to bedding of the economically mineralized rocks in reduced-facies deposits is more commonly between 1 and 15 m. The White Pine deposit occupies an area of 50 km² constituting a single continuous ore layer 1 to 7 m thick. Within its chalcocite-bearing zone, individual beds have grades that vary only slightly over a distance of 10 km laterally (fig. 12). In noneconomic occurrences, which are very numerous in mineralized parts of basins, the thickness is commonly just centimeters.

#### Dimensions and the Size of Hydrothermal Systems Relative to Extent of Economically Mineralized Rock

The lateral extent of entire mineralizing systems may be enormous. Measured from the oxidized zone on the side towards the source of the metals-transporting fluids, the Cu ore zone can be tens of kilometers wide (for example, 25 km at Lubin, Poland). In the adjacent basin is a pyrite-bearing zone several hundred kilometers wide. The lateral dimensions are 100 to as much as 10,000 times greater than the thickness. Ore-stage hematite alteration (Rote Fäule) also extends tens to hundreds of kilometers in central Europe. Certainly, the total extent of altered rock is much larger than the economic zones of deposits, which may themselves be huge.

The economic parts of sandstone Cu deposits are tabular to lens-shaped, with maximum thicknesses of about 100 m and lengths and widths of kilometers to tens of kilometers. Deposit lateral dimensions are from 20 to 100 times the thicknesses. The largest sandstone Cu deposit is Udokan, Russia, which occupies a zone 10 km by 2.5 km that contains multiple ore lenses as large as 3 km long, 700 m wide, and several tens of meters thick. The middle quartzite beds hosted the Troy orebody at Spar Lake, Mont., were 2,500 by 540 m in area and 15 to 30 m in thickness. However, over about 90 percent of the area of the orebody, the thickness was consistent at 21 to 23 m. For sandstone Cu deposits, the pyritic zones are as much as tens of kilometers parallel to bedding, but they probably do not reach 100 km in width. The extent of the hematite zones is uncertain but may also be 100 km in width. The ore-related hematite zone extends into red beds where it is difficult to distinguish and has nowhere been fully mapped.

Red-bed deposits are smaller, commonly with lengths of a few tens of meters and thicknesses of less than 10 m. For red-bed Cu deposits, the pyritic zones lacking Cu minerals are consistently less than 1 km parallel to bedding, and they are typically just tens of meters. Similar to the sandstone Cu deposits, the hematitic zones may reach 100 km and have nowhere been fully mapped.

#### Source Rocks

Cu source rocks are red-bed conglomerates, sandstones, siltstones, and mudstones of deltaic, fluviatile, eolian, lacustrine, or intertidal origin. The conglomerates and sandstones have cross bedding and cut and fill channels, and they had high original permeability. Detrital labile minerals, such as
Figure 12. Stratigraphic column and typical copper grades of each host unit, White Pine deposit, Michigan, United States, from Mauk and others (1992a). Within the chalcocite-bearing zone, copper grades are remarkably consistent laterally within each bed across more than 10 kilometers laterally, and they correlate inversely with grain size of the host beds and directly with the content of organic carbon. Both relations are typical of reduced-facies-type deposits.

<table>
<thead>
<tr>
<th>Member</th>
<th>Description</th>
<th>Lithology</th>
</tr>
</thead>
<tbody>
<tr>
<td>Harbor Fm</td>
<td><em>Upper</em> Shale</td>
<td><em>Upper transition</em>: Fine-grained sandstone with black shale partings</td>
</tr>
<tr>
<td>Lower Nonesuch Formation</td>
<td><em>Red and gray</em>: Widely laminated greenish and gray siltstone and red shale</td>
<td></td>
</tr>
<tr>
<td>19.5</td>
<td><em>Marker</em> bed: Black thinly laminated siltstone with abundant white calcareous blebs. Thickness about 3 meters</td>
<td></td>
</tr>
<tr>
<td>16.5</td>
<td>Massive black siltstone</td>
<td></td>
</tr>
<tr>
<td>15</td>
<td><em>Stripy</em>: Finely laminated black silty shale with abundant calcite cement</td>
<td></td>
</tr>
<tr>
<td>8.5</td>
<td>Massive gray siltstone in graded beds 10 to 50 centimeters thick. Contains coarse-grained sandstone beds toward the south</td>
<td></td>
</tr>
<tr>
<td>5</td>
<td><em>Red and gray</em>: Widely laminated greenish gray siltstone and red shale</td>
<td></td>
</tr>
<tr>
<td>4</td>
<td>&quot;Upper&quot; Nonesuch Formation</td>
<td></td>
</tr>
<tr>
<td>3</td>
<td><em>UZV</em> (Upper Zone of Values) Massive dark gray siltstone</td>
<td></td>
</tr>
<tr>
<td>2</td>
<td>&quot;Brown massive&quot;: Reddish brown to gray massive siltstone with calcite nodules near the base</td>
<td></td>
</tr>
<tr>
<td>1</td>
<td>&quot;Thinly&quot;: Thinly laminated dark gray siltstone and black shale</td>
<td></td>
</tr>
<tr>
<td>0</td>
<td>&quot;Upper&quot;: sandstone: Red to gray fine-to-coarse-grained sandstone. Locally contains conglomeratic lenses. Reddish shale and siltstone common in the middle part of the unit</td>
<td></td>
</tr>
<tr>
<td>0</td>
<td><em>Lower transition</em>: Sandstone with black shale partings</td>
<td></td>
</tr>
<tr>
<td><em>Lower</em> Nonesuch Formation</td>
<td><em>Lower</em>: sandstone: Red to gray fine-to-coarse-grained sandstone. Locally contains conglomeratic lenses. Reddish shale and siltstone common in the middle part of the unit</td>
<td></td>
</tr>
<tr>
<td>19.5</td>
<td><em>Lower sandstone</em>: Red to greenish gray, coarse-to fine-grained sandstone</td>
<td></td>
</tr>
</tbody>
</table>

Modified from J.L. Mauk and others, 1992.
amphibole, pyroxene, and plagioclase, probably acted as the ultimate sources of Cu (Walker and others, 1967; Walker, 1989; Lapoint, 1974). Zielinski and others (1983) demonstrated that in the process of early diagenetic reddening in red beds, the Cu, and probably other metals, are “stored” in the earliest diagenetic ferric oxides. In hotter, deeper, more advanced conditions of diagenesis, the metals may be released to mineralizing solutions as the hydroxylated ferric oxides dehydrate to hematite. Goethite dehydroxylates to hematite at slightly above 125 °C at atmospheric pressure (Posnjak and Merwin, 1919, p. 326–334). Observed mineralizing solutions are very saline (Annels, 1989; Hayes, 1990; Dewaele and others, 2006; El Desouky and others, 2009), so the process of releasing metals into a mineralizing solution is probably not simply because of aging and dehydration of the ferric-oxide grain coatings and is, instead, a chemical leaching by saline groundwater. The groundwater may have entered into the red-bed sequence from outside, or it may have been indigenous to the red beds as an originally connate water. The Cu source rocks have been firmly identified only from the Redstone copperbelt in Canada (Lustwerk and Rose, 1983) and from the Western Montana Copper Sulfide Belt (Hayes, 1990).

Flood basalt packages may be alternative or additional source rocks for Cu in some districts and in the volcanic red-bed type of related deposits. Alteration of amphiboles and pyroxenes to epidote in those rocks appears to accompany Cu mobilization (Livnat, 1983; Borg, 1991). For these, also, the source of the mobilizing groundwater is unclear.

Host Rocks

Reduced-facies host rocks are commonly thin-bedded to very thinly laminated carbonaceous shale, siltstone, or very fine-grained sandstone, or they are similarly thin-bedded to very thinly laminated impure carbonate rocks. The Kupferschiefer shale is the deepest water marine rock in its local section, being a discrete transgressive bed commonly just 30 to 70 cm thick, with a top that typically grades upward into the overlying Werra carbonate and a bottom that is sharply erosional and underlain by a few centimeters to very rarely as much as 2 m of Zechstein conglomerate. In medieval times, only the Kupferschiefer shale bed itself was mined, and it typically averaged greater than 5 percent Cu. Now, at places such as Lubin, Poland, and Sangerhausen, Germany, the Kupferschiefer bed is mined along with locally as much as 10 m of underlying mineralized Weissliegendes sandstones and with locally as much as 4 m of overlying mineralized Werra carbonates beds. The zone of ore-grade Cu mineralization cuts very gently across all three beds allowing various combinations to be economically mined but typically totaling 3 to 4 m in thickness at Lubin at an average grade of 2 percent Cu.

The hosts of sandstone Cu deposits contrast with the reduced-facies deposit host rocks. Hosts of sandstone Cu deposits are medium- to very thickly bedded sandstones (bed thickness terminology of Ingram, 1954). These are typically well-sorted feldspathic or arkosic sandstones. Where conglomeratic, they typically have only mud intraclasts and are not polymictic. The hosts for red-bed-type deposits are conglomeratic sandstone, typically feldspathic or arkosic. They are commonly polymictic.

The presence of plant or animal remains provides a carbonaceous host for Cu deposition for the reduced-facies and red-bed-type deposits. Where best studied in the Kupferschiefer, the organic remains were originally alginite, sporinite, liptode-trinite, and amorphous material, and they became bitumen and kerogen during mineralization (Nowak and others, 2001). In the Late Permian Creta Shale, the organic remains were Triletes spores, which were replaced pseudomorphically by chalcocite (Hagni and Gann, 1976). In red-bed Cu deposits, the organic remains were of higher vascular plants, and those were pseudomorphed by Cu sulfides, including chalcocite, bornite, and chalcopyrite (Tarr, 1910, p. 223; Richard, 1915, p. 640; Fath, 1915, p. 146; Miller, 1955, p. 161–162; Soulé, 1956, p. 16). Red-bed Cu deposits are well known for their Cu-sulfide-pseudomorphed logs (at Nacimiento, N. Mex., for example; Woodward and others, 1974).

Some associated rocks are evaporite or sabkha beds. These provide a source of metals-transporting brine.

Structural Setting(s) and Controls

The SSC deposits occur in many different present-day structural settings. Host basins vary from early transtensional basins (for example, Boleo, Baja California, Mexico), through aulacogens and other general rifted settings (for example, Spar Lake; Kupferschiefer, central Europe), to post-orogenic molasse basins (for example, Dzhezkazgan, Kazakhstan; Tschudi, Namibia), and to fold-and-thrust belts (for example, Democratic Republic of Congo part of the Central African Copperbelt; Aynak, Afghanistan). They share, however, only two original sedimentary tectonic settings from the time of host-rock deposition. These are early oceanic rifted margins (for example, Kupferschiefer; Central African Copperbelt) exclusive of post-orogenic veinlet-controlled deposits such as Kansanshi and Lonshi; Oklahoma and northern Texas post-aulacogen setting; Juramento, Argentina) and late- to post-orogenic molasse basins (for example, Dzhezkazgan; Tschudi, Lisbon Valley).

The mineralizing process, however, in SSC deposits is controlled mostly by sedimentary lithology and not by structure. However, faults may have been important in other ways. Areas where faulting or folding have caused favorable beds to be uplifted and exposed, but not destroyed by erosion, are highly prospective. For certain sandstone Cu deposits, faults or fault zones may have fed oxidizing metal-liferous basinal brines from red beds into favorable host rocks. This is suggested by the geometry of the Cashin deposit in the Paradox Basin, Colo. (fig. 13) (Thorson and MacIntyre, 2005; MacIntyre, 2006), and it may be the case for many other
sandstone Cu deposits. Those in the Revett Formation in the Belt Supergroup each have one or more possible feeder faults, such as the East Fault at Spar Lake. It is still uncertain whether the East Fault did or did not feed metalliferous basinal brines into the host beds, although it is certain that metalliferous fluids had to come upward across section into the ore zone before spreading within those permeable beds.

In Kazakhstan, geologists recognize two types of sandstone Cu deposits: Dzhezkazgan-type (for example, Dzhezkazgan, Zhaman-Aibat) and Zhilandy-type (for example, Itauz, Sary-Oba, Kipshakpai, Karashoshak). The Dzhezkazgan-type form concentric rings along the flanks of anticlinal structures, with pyritic zones interior to each ring and chalcocite being exterior. The Zhilandy-type instead has a concentric zonation centered on an elongate central chalcocite zone, which is generally a fault (B. Syusyura, oral commun., 2009). As defined above, Cashin, Lisbon Valley, Spar Lake, Rock Creek, and Montanore are all similar to the Zhilandy-type deposits.

Reduced-facies-subtype deposits may also be affected and localized by faults. At White Pine, Mauk and others (1992b) showed a number of relations between compressional fault features and second-stage Cu mineralization. First stage constituted most of the finely disseminated chalcocite-dominated mineralization that replaced pyrite in the Nonesuch Formation and that produced a chalcocite-bornite-chalcopyrite-pyrite fringe (Brown, 1971). Second stage included (1) bedding-plane seams of native copper, (2) bedding-plane seams of chalcocite, (3) stockwork veins

Figure 13. Cross section of the Cashin sandstone copper deposit, Colorado, United States, from Thorson and MacIntyre (2005). The section is perpendicular to the strike of the Cashin fault and to the long axis of the deposit. The zone of greater than 0.1 percent copper (within the heavy red dashed line) is an irregular zone distributed grossly symmetrically surrounding the fault, although with values controlled strongly along bedding within the Wingate Sandstone host rocks and decreasing away from the fault. Mineralizing fluids are inferred to have ascended along the fault before migrating outward from it along permeable beds.

<table>
<thead>
<tr>
<th>Contact</th>
<th>Copper mineralization &gt;0.1%</th>
</tr>
</thead>
<tbody>
<tr>
<td>Stratigraphic</td>
<td>Uncertain presence or uncertain continuation</td>
</tr>
<tr>
<td>?</td>
<td>Borehole (with borehole ID)</td>
</tr>
</tbody>
</table>

Copper assay values

0 10 20 30 METERS

0 25 50 75 FEET

Modified from J.P. Thorson and T.J. MacIntyre, 2005, by permission.
of chalcocite localized adjacent to thrust faults, (4) cementing native copper occurring together with pyrobitumen in the top of the Copper Harbor Conglomerate, (5) slickensided sheets of native copper along bedding, (6) bedding-plane seams of native copper that pass laterally into seams of chalcocite, (7) mineralogically zoned disseminations developed as halos adjacent to thrust faults, and (8) disseminations that are mineralogically zoned, chalcocite to bornite to chalcopyrite to pyrite over centimeters above and below bedding-plane seams of chalcocite. A feature that compellingly demonstrates Cu mineralization during compressional faulting is a thrust fault that offset bedding by 90 cm but that offset a bedding-plane vein of native Cu by only 60 cm (fig. 14). The bedding-plane vein contained native copper deposited after offset began on the thrust fault and was then, itself, offset by later movement on the thrust. Second-stage mineralization at White Pine is dominated by native copper, although it does include minor chalcocite, whereas first-stage mineralization is Cu-sulfide-dominated, although it does include minor native copper. With its relation to compressional faulting and native copper dominance, the second-stage mineralization seems likely to correlate with the time of native copper mineralization in the Portage Lake Lava Series to the northeast in the Keweenawan Native Copper district. Brown (2006, 2008) interpreted that the second-stage mineralization was from a brine similar to that from the first, disseminated stage, but the sulfide in the host rocks had been used by the first stage, such that the second was dominated, in the basal ore section, by native copper. Brown correlated each stage with a stage of highland recharge to the Cu source rocks, and he correlated the second stage, which included mineralized veins, amygdular and brecciated flow tops, and conglomerates in the Portage Lake lavas, to compressional features at White Pine caused by late Mesoproterozoic Grenvillian deformation.

Another frequent type of control of reduced-facies deposits by faults is where such deposits are localized above the flanks of basement horst blocks, a relation demonstrated by several different deposits in the Zambian Copperbelt (fig. 11). Zambia also has deposits where the mineralized arenite beds are truncated against faulted basement (fig. 15).

**Geophysical Characteristics**

**Regional-Scale Geophysics**

There has been an increasing recognition that SSC deposits occupy recognizable settings within basins. This lends itself to all geophysical methods that can help determine basin geometry.

Reduced-facies-subtype deposits tend to occur above the flanks of basement horsts or generally above the flanks of basins. Gravity and magnetic methods can map characteristics of basement rocks through sedimentary sections, which themselves have much lower specific gravity and much lower magnetic susceptibility than the crystalline basement. Airborne magnetic surveys are fast, inexpensive, and easily combined with additional electromagnetic methods that could be effective in limited circumstances at deposit-scale (see below, the discussion of induced polarization, IP, methods). Faults, especially those that affect basement but not the overlying sedimentary rocks, can be mapped on first- and second-derivative maps of the gravity and magnetic signatures.

Because halite and sylvite have low specific gravities (2.16 and 1.99 g/cm³, respectively), gravity maps and particularly microgravity maps can identify salt structures. These may be associated with SSC deposits (examples are Lisbon Valley, Utah, and Corocoro, Bolivia).

Obviously, either or both seismic-reflection and seismic-refraction surveys can help identify basement structure and overall basin geometry.

**Deposit-Scale Geophysics**

Earlier models for SSC deposits (Kirkham, 1984a; Cox, 1986; Cox and others, 2003) made no mention of any geophysical methods applicable to exploration for this type of deposit. Some methods are, however, effective, particularly where the sedimentary section generally lacks conductive rock types.
Vector induced polarization was first tested over the pre-mining Spar Lake sandstone Cu deposit (Van Voorhis, Asarco, Inc., written commun., 1970), and following successful demonstration of the utility of the method, it was widely applied in the area by Kennecott Exploration Services. Between 1970 and 1975, they conducted induced polarization (IP) surveys along 18 crisscrossing lines in the area of the deposit and to its north. In addition to the known deposit, three anomalous areas were found, although one of these was downsection from any Revett Formation potential ore host rocks. The better of the remaining two anomalies, the Fairway Creek prospect, was drilled and resulted in discovery of a zone of anomalous pyrite concentrations with minor galena disseminated in upper member quartzites of the Revett Formation. Lying along trend with the elongation of the Spar Lake deposit and also along the East Fault, the Fairway Creek prospect, which has no outcrop exposure, may be the northwest distal continuation of the Spar Lake mineral zonation (fig. 7).

Van Voorhis (Asarco, Inc., written commun., 1970) noted that the rocks of the Spar Lake area generally had extremely high resistivity so that even with less than 1 volume % total sulfides (and about 0.1 v % magnetite) in the orebody, there was a well-defined anomaly. For the dipole-dipole survey that passed over the orebody looking through about 300 m of cover, modeled resistivity of the ore was 2,600 to about 3,300 ohm-meters, whereas the surrounding country rocks generally exceeded 5,000 ohm-meters. Orebody calculated metal factors were 8 to 10 milliradians, whereas country rock metal factors were everywhere 6 or less milliradians and were 0 to 3 milliradians at all distances greater than 1 km from the deposit boundaries. Responses over the Fairway Creek prospect sulfide-anomalous body were nearly identical for a cover thickness that was also about 300 m.

IP has also proven very effective in recent prospecting near the Malundwe and Chimiwungo deposits (Lumwana mine) in Zambia. The setting there is grossly similar in having very resistive rocks everywhere surrounding the deposits. Those deposits occur in highly sheared “mylonitized, recrystallized muscovite-phlogopite-quartz-kyanite schists,” with quartz-feldspar-biotite gneisses above and below the sulfide-mineralized schists (Davis and others, 2008, p. 27–28). The schists are interpreted as zones of thrusting and possible nappe folding that were mineralized with bornite, chalcopyrite, and distal pyrite. The Chimiwungo North orebody was discovered in 2004 by drill testing an IP chargeability anomaly, which then led to completing greater than 150 km² of IP coverage nearby over the next few years. The work identified 27 IP chargeability anomalies. Two of those (Kanga and Kababisa prospects) had received initial drill testing by June 2008, and both drill programs had intersected significant Cu sulfides (Davis and others, 2008, p. 27 and 31–34).

Figure 15. Distribution of ore in the Nchanga deposit, Zambia, in a vertically exaggerated schematic section that looks approximately downdip along the bedding, from McKinnon and Smit (1961). Comparable with figures 10 and 11. The distribution of ore is controlled by the pinchouts of the Mindola Clastics Formation against the basement high, subjacent to the major ore host, the Copperbelt Orebody Member. Note that the sandstone pinchouts against basement would constitute an updip pinch-out stratigraphic trap in a petroleum province.
The success of IP for deposit-scale prospecting for SSC deposits will probably be limited to areas that lack other conductive rock types, such as pyritic black shales or graphitic schists. Nonetheless, the method could be effective in additional areas. The Udokan region of Transbaikalian Russia is similar to the Belt Supergroup, composed almost entirely of resistive rocks, so IP promises to be a useful deposit-scale prospecting technique in that area. In the Democratic Republic of Congo, most orebodies are found in the Roan-2 stratigraphic interval within stratigraphically coherent “ecailles” (French: flakes) floating within chaotic breccia. Additional ecailles orebodies might be effectively located with limited-area IP surveys. The IP can expect only limited success where the surveys go beyond the areas of outcropping Lower Roan Group and breccia, onto upsection areas underlain by rocks of the Dipeta and Mwashya Groups and the Grand Conglomerate. Those rocks upsection all have substantial intervals of black shale. But within the Lower Roan exposure, where ore-bearing “ecailles” may be covered beneath breccia only meters below the surface without any surface expression, IP promises to be effective.

Application of seismic methods was mentioned above with reference to helping determine basin geometry. Seismic studies have also been used in deposit-scale prospecting in the Chu-Sarysu Basin, Kazakhstan. The blind Zhaman-Aibat and Taskura deposits were discovered by drilling on the geologically mapped Zhaman-Aibat anticline beginning in the 1920s (Japan International Cooperation Agency and Metal Mining Agency of Japan, 1997), using the analogy of that structure to the Dzhezkazgan brachyanticline. Having concluded that the sandstone Cu deposits of the Chu-Sarysu Basin are associated with accumulations of hydrocarbons (Sysyura and Tyugay, 2008), explorationists have since focused on seismically indicated positive structural closures, with several additional prospects successfully identified in recent years (B. Sysyura, oral commun., 2009). Hayes and others (2012) have confirmed that sandstone Cu deposits form in association with hydrocarbon accumulations. They found that Spar Lake formed where oxidizing Cu-transporting brines mixed and reacted with a reservoir of sour gas. Obvious next steps are (1) to start prospecting for former stratigraphic and combination traps in productive basins such as the Chu-Sarysu; and (2) to apply the thinking and mineral-exploration methods to other basins that host known SSC-mineralizing systems and that have known sour gas accumulations and seismic data, such as central Kansas, western Oklahoma, and northern Texas, United States.

**Remote Sensing of Sediment-Hosted Stratobound Copper Deposits**

Two possible applications of remote sensing may be of use in exploration for SSC deposits. First, Cu clearings were well-known and widely searched for using air photos in the early days of exploration in Zambia and the Democratic Republic of Congo (Ellis, 1961, p. 175–177). Even 100 years after their first recognition, similar features may still await discovery. In other climates with other specific types of vegetation, different types of vegetation anomalies may accompany outcropping SSC deposits, and the vegetation anomalies may be visible using remotely sensed spectral data. Second, bleaching of red beds is easily mapped by remote sensing (Beitler and others, 2003) either using simple color mapping or mapping of Fe oxides using Landsat, AVIRIS (Airborne Visible/Infrared Imaging Spectrometer), or ASTER (Advanced Spaceborne Thermal Emission and Reflection Radiometer) imagery. Current thinking holds that sandstone Cu deposits can mineralize only the bleached sandstones of such systems (Thorson, 2005; B. Sysyura, oral commun., 2009). Assumedly, the red sandstones would not be mineralized because they were in equilibrium with the hematite-stable metals-transporting brines, whereas some of the bleached sandstones would contain petroliferous fluids that would be reactive with the metals-transporting brines. If true, then only bleached sandstones would be prospective in regions such as the Colorado Plateau, United States, the Chu Sarysu Basin, Kazakhstan, and the Western Montana Copper Sulfide Belt, United States.

**Hypogene Ore Characteristics**

**Grade**

Because of the high Cu content of chalcocite, digenite, bornite, and djurleite, which characterize SSC deposits, Cu grades of this class of deposits average 2.1 percent, compared to porphyry Cu deposits (0.44 percent) or Kuroko massive sulfide deposits (1.3 percent) where chalcopyrite is the main Cu mineral. Less than 20 percent of SSC deposits contain substantial Ag, grading between 2 and 200 g/t, or Co, grading between 0.15 and 0.5 percent in primary ores (Mosier and others, 1986, p. 207–208). Co, in particular, is present at much higher grades locally in supergene, residually enriched ores in the Central African Copperbelt.

A platinoid metal and Au orebody has been identified and described, paralleling the zonation of Cu-, Pb-, and Zn-bearing sulfides and lying from 0 to 0.5 m below the Cu ore of the Kupferschiefer mineralizing system on the Fore-Sudetic Monocline in Poland. The deposit of precious metals averages 0.22 m in thickness, with grades of 2.25 g/t Au, 0.138 g/t Pt, and 0.082 g/t Pd, although no tonnage has been announced (Piestrzynski and others, 2002, p. 517). However, economically recoverable ore does not form a continuous layer beneath the Cu ore. For example, about 30 percent of the area of the Polkowice deposit is underlain with a greater than 20 cm thickness of greater than 1 g/t Au. Given, however, that the deposits of the Fore-Sudetic monocline total an area of more than 410 km² (S. Osczepalski, oral commun., 2008), this may be the largest Au deposit in Europe. Unfortunately, the Au and platinoid concentrations were not identified until after large tonnages of Cu ore had already been mined, and many workings have collapsed, which may render the Au resource unprofitable to mine in those areas (Z. Sawlowicz, oral commun., 2001).
Ore Mineralogy and Assemblages

Sulfide mineralogy and its zonation are well-defined for SSC deposits. They were briefly described earlier in the report and illustrated in figures 4, 5, 6, and 7.

Most SSC deposits contain chalcocite, digenite, bornite, and chalcopyrite. Some, such as Boleo, contain monoclinic (pseudo-orthorhombic) chalcocite crystals (Touwaide, 1930, p. 134–135), demonstrating that the main ore zone of the deposit formed at less than 103.5 °C (Roseboom, 1966). Others, such as White Pine and Dzhezkazgan, contain hypogene djurleite (Brown, 1971, p. 556; Lur’ye and Gablina, 1976, p. 72; respectively). However, statements by Brown (p. 565) and Lur’ye and Gablina (p. 73), that the presence of djurleite limits the temperature of ore deposition to 93 °C or less are incorrect unless more conditions are met. The maximum thermal stability of djurleite is 93 °C (Roseboom, 1966). The djurleite composition can, however, be reached by exsolution on cooling from a higher temperature solid solution with composition within the high digenite solid solution (Roseboom, 1966, p. 648). Djurleite would be a demonstration of origin at less than 93 °C only if it occurred in its own euhehedral crystals (analogous with the monoclinic chalcocite of Boleo), or if it occurred in large, monomineralic masses in the ores (Roseboom, 1966). Also, the presence of X-bornite (Cu-deficient, S-rich bornite, “orange bornite”) does not limit the origin of Dzhezkazgan, Kazakhstan, ores to 75 °C or less as was claimed by Lur’ye and Gablina (1976, p. 76) and repeated several times since. X-bornites, when heated, exsolve to chalcopyrite and stoichiometric (“brown”) bornite. So presence of X-bornite at Dzhezkazgan in the local absence of both chalcopyrite and brown bornite was taken to indicate that Dzhezkazgan had formed below 75 °C, the temperature by which the exsolution took place in some X-bornites from red-bed-type deposits of the Colorado Plateau in heating experiments (Brett and Yund, 1964). However, X-bornites have varying thermal stabilities (Brett, 1962; Brett and Yund, 1964; Yund and Kullerud, 1966). Other X-bornites from Colorado Plateau deposits did not exsolve chalcopyrite and brown bornite until they were heated to 100 °C (Brett and Yund, 1964), and synthetic X-bornite did not exsolve until heated to greater than 140 °C (Yund and Kullerud, 1966). Lur’ye and Gablina would have had to do heating experiments with the Dzhezkazgan X-bornites, themselves, to establish a minimum temperature provided by the X-bornite there, or claim only the conservative temperature estimate, that the deposit formed at 140 °C or less. Native silver is present in a number of deposits within rocks that also contain chalcocite and (or) bornite (Woodward and others, 1974; Jung and Knitzschke, 1976; Hayes and Einaudi, 1986, p. 1910 and 1916–1917; Volodin and others, 1994, p. 18). Stromeyerite is present as well in the Kupferschiefer and at Spar Lake. In those SSC deposits where Ag is found in concentrations of tens of grams per tonne, that Ag is sited mostly in solid solution in chalcocite, digenite, and (or) bornite (for example, Witvlei: Anhaeusser and Button, 1973, p. 292; Spar Lake: Hayes, 1983, Appendix II; Kupferschiefer: Sawlowicz, 1990, p. 264; Udokanski: Volodin and others, 1994, p. 18). At Spar Lake, each of those three sulfides typically carries about 0.1 wt. % Ag.

Sulfide minerals in the fringing zones host recoverable metals in many SSC deposits. Some deposits in Zambia and the Democratic Republic of Congo contain carrollite (Darnley, 1960, p. 154; Oosterbosch, 1962, p. 118), and some in the Democratic Republic of Congo and Zambia also have Co-rich pyrite (Brown and Bartholomé, 1972; Bartholomé and others, 1976). Galena and sphalerite are common in the zones fringing the Cu orebodies, between ore and pyritic zones.

The dominant sulfides and associated oxides in the Kupferschiefer system vary from zone to zone and include (1) hematite with only scant traces of any sulfides (Rote Fäule); (2) covellite with idaite, chalcopyrite, and trace magnetite; (3) chalcocite with digenite, covellite, and pyrite; (4) chalcocite with bornite, digenite, and pyrite; (5) bornite with chalcocite, chalcopyrite, sphalerite, galena, and pyrite; (6) bornite with chalcopyrite, pyrite sphalerite, and galena; (7) chalcopyrite and pyrite with sphalerite, galena, and tennantite; (8) pyrite, sphalerite, galena, and chalcopyrite; (9) pyrite, sphalerite, and galena with chalcopyrite; and (10) pyrite with only scant traces of any base-metal sulfides (Jung and Knitzschke, 1976). There is a zonation of metals varying from a zone devoid of base metals other than Fe (#1 above), to Cu zones (#2–6), to a Pb zone (#7), to a Zn zone (#8), and to another zone nearly devoid of base metals other than Fe (#10) (Jung and Knitzschke, 1976; Oszczepalski and Rydzewski, 1997). Ore grades of Cu, typically greater than 1 percent, are developed only in zones with bornite, chalcocite, or covellite as a predominant sulfide. The single exception is in Central Africa, where the chalcopyrite zone can reach ore grades and is attractive, as well, because it may have high concentrations of Co. Examples are the Kambove West deposit, Democratic Republic of Congo (Cailteux, 1994) and the Nkana South orebody, Zambia (Brems and others, 2009).

In each one of the other deposits of the reduced-facies and sandstone subtypes, there is a similar sulfide zonation, which progresses from a chalcocite or a bornite side to a pyrite side, although with local variations. Spar Lake, for example, has a chalcopyrite-only zone that is 1–3 m wide between galena-bearing rock and bornite-bearing rock, and pyrite nowhere coexists with bornite or chalcoite. The galena-dominant zone at Spar Lake has, in places, a thickness of greater than 20 m and a width of several hundred meters. At White Pine, the transition from rock with chalcocite as the dominant sulfide to rock with pyrite as the dominant sulfide is just a few centimeters wide, where that “transition fringe” passes through fine-grained lithologies. The transition fringe at White Pine contains minor amounts of the cadmium-sulfide mineral greenockite (Brown, 1971). Greenockite is also present at Happy Jack and other red-bed-type Cu–U deposits, Utah (Weeks and others, 1959), and the Sustut, British Columbia, volcanic red-bed-type deposit (Wilton and Sinclair, 1988).
Many other sulfides have been identified, each with a limited extent. Tetrahedrite in small concentrations is found between rock with bornite and rock with galena at Spar Lake and at Juramento, Argentina (Durieux and Brown, 2007, p. 890), but is not typically present within Kupferschiefer system deposits. Tennantite is found, instead, in approximately the same zonal position. In Kupferschiefer deposits, linnaeite is present at places in rock with pyrite, galena, sphalerite, and chalcopyrite in about equal abundances. Linnaeite is found together with chalcopyrite and pyrite in deposits of the Central African Copperbelt, but galena and (or) sphalerite are extremely rare in Zambian deposits (Notebaart and Vink, 1972). Altogether, more than 115 different ore minerals have been identified in the Kupferschiefer deposits (Pieczonka and others, 2001), but most of those occur only as traces yet may still contribute to geochemical signatures (for example, see Bechtel and others, 2002, p. 17–24).

U- and V-oxide and silicate minerals may be present in some red-bed-subtype deposits, as in the Colorado Plateau region, United States, and in Argentina east of the Andes in Cretaceous host rocks (Brodtkorb and Brodtkorb, 1984; Ferreyra and Lardone, 1990). In red-bed-subtype deposits, Cu-sulfide zonation is typically telescoped relative to the other subtypes. There may be no rock with only chalocite, and there may be no rock with only pyrite. U in coffinite (tetragonal \( U[SiO_4,OH]_4 \)) is predominantly in pyritic assemblages that may entirely lack Cu minerals or contain chalcopyrite. V sited in either V illite or V chlorite is also present mostly in the pyritic assemblages. U as uraninite (isometric \( UO_2 \)) and V as montroseite (orthorhombic \([V,Fe]O[OH]\)) are more typically found in copper-bearing assemblages, particularly with chalcopyrite, and somewhat less commonly with bornite or chalcocite (Miller, 1955; Phillips, 1960; Tschanz and others, 1958). The result is that Cu-poor variants of the Cu-U-V sandstone-hosted deposits of the Colorado Plateau region are the ones that typically contain coffinite and V phyllosilicates whereas Cu-rich variants more typically have uraninite and montroseite (Hayes, 1982).

### Paragenesis and Zoning

Cu minerals in SSC deposits and their Cu to S molar ratio can be listed in their paragenetic order and zonation as follows:

- Chalcopyrite (\( CuFeS_2 \)) 0.5
- Bornite (\( Cu_9FeS_8 \)) 1.25
- Digenite (\( Cu_3S_2 \)) 1.80
- Djurleite (\( Cu_{3+}S_{10} \)) 1.94
- Chalcocite (\( CuS \)) 2.0

Minerals at the low-Cu end of the list form a zone next to the reduced, pyritic zone in the deposits. Minerals at the Cu-rich end are found in zones next to the hematitic rock on the side from where the high salinity, metals-transporting fluids entered the systems (for example, Bechtel and others, 2001).

In unmetamorphosed deposits, pyrite is replaced by chalcopyrite, which is replaced in turn by bornite and then by chalcocite (Brummer, 1955; Brown, 1971; Anhaeusser and Button, 1973; Bartholomé and others, 1976; Lur’ye and Gablina, 1976; Chartrand and others, 1989). Furthermore, pyrite is replaced rarely by magnetite or commonly by hematite within the zones of Rote Fäule associated with Kupferschiefer deposits (Jung and Knitschke, 1976; Schmidt, 1987), and hematite accompanies chalcocite in replacements of pyrite at White Pine (Brown, 1971, p. 556) and Witvlei (Anhaeusser and Button, 1973, p. 292).

Continuing centimeters to meters into the Rote Fäule of the Kupferschiefer mineralizing systems in the Fore-Sudetic monocline, Poland, some important additional minerals are deposited. Rote Fäule (literally translated: red rot) is rock that is spotted or streaked with hematite and that lacks any important concentrations of Cu sulfides. The important minerals in the near-ore edge of the Rote Fäule zone rocks are (1) native gold of high purity associated with hematite and covellite, and (2) mercurian electrum with an assemblage of chalcocite, digenite, bornite, and minor clausthalite (isometric PbSe), tetraaurcupride (tetragonal CuAu), spionkopite (rhombohedral \( Cu_9S_8 \)), yarrowite (rhombohedral \( Cu_9S_8 \)), and tieffmannite (isometric HgSe). Native platinum (Pt) and palladium (Pd) occur together with the above. The result is an Au- and platinoid-bearing orebody from 0 to 0.5 m below the Cu-Ag orebody (Pietrzynski and others, 2002).

Some deposits contain carrollite and Co-pyrite in the near-ore part of the pyrite zone. These may extend for hundreds of meters parallel to bedding or for meters perpendicular to bedding and, for carrollite, into the chalcocite-bearing zone in the Central African Copperbelt reduced-facies deposits.

Cu sulfides in the deposits have been observed to replace carbonaceous materials of every type in the sedimentary host rocks of SSC deposits at essentially every scale. At the Creata reduced-facies-subtype deposit, many chalcocite grains are replacements of 100 to 200 μm diameter ovoid spores of the genus, Triletes (Hagni and Gann, 1976). At the Nacimiento red-bed-subtype deposit, whole fossil logs were replaced by pyrite and later Cu sulfides, such that the deposit was essentially a replaced logjam within fluvial sandstone paleochannels (Woodward and others, 1974).

Where organic matter or organic carbon compounds were not directly replaced by the Cu sulfides in forming SSC deposits, the organics were altered chemically and physically, or they were oxidized to carbonate species and precipitated as ore-stage carbonates. In mineralized Kupferschiefer, ore-solution sulfate was thermochemically reduced to sulfide utilizing the carbon in organic matter as the reductant. This resulted in oxidative alteration of the solid organic matter,
whose carbon was then incorporated into epigenetic, ore-stage carbonate minerals with δ13C values that are shifted downward compared with the typical marine values of the carbonates just tens of centimeters upsection. Some of the relict organic matter was altered to solid bitumen in the process, and the bitumen, as well as the kerogen, in well-mineralized Kupferschiefer has δ13C values reciprocally heavier than the original organic matter (Bechtel and others, 2001). Toward and into the Rote Fäule, the rocks became increasingly calcified, with a higher Ca/Mg molar ratio. Dolostone of the Werra (Zechstein 1) carbonated was altered to limestone in the Rote Fäule zone (Rentzsch, 1974, p. 411), and the thin Boundary Dolomite at the base of the Kupferschiefer bed is also altered to limestone. Original liptinite group macerals of the Kupferschiefer bed, including alginite, sporinite, collinite, and amorphous organic macerals, were altered to increased concentrations of inertinite and solid bitumen (Nowak and others, 2001). The reflectance of Kupferschiefer vitrinite increased correspondingly, from R0 of 0.72–0.86 percent in others, 2001). The reflectance of Kupferschiefer vitrinite increased correspondingly, from R0 of 0.72–0.86 percent in Cu-poor reduced rocks, to 0.72–1.13 percent in Cu-rich rocks, and to 1.03–1.10 percent in Rote Fäule-altered Kupferschiefer (Oszczepalski and others, 2002). The various organic petrologic indicators show that Rote Fäule-altered Kupferschiefer has seen a maximum temperature of about 130 °C in the Fore-Sudetic monocline district in Poland, corresponding to source rock within the window of oil generation (Nowak and others, 2001). Accompanying the petrographic alterations to the Kupferschiefer organic matter, the chemistry of the organic matter was also substantially altered (Püttmann and others, 1987). The content of saturated hydrocarbons was diminished, and the content of heteroaromatic compounds containing oxygen and S was increased by reaction with the metals-transporting brine. Cu-mineralized and Rote Fäule-altered Kupferschiefer rocks are enriched in polycyclic aromatic hydrocarbons, such as dibenzofuran, dibenzothiophene, fluoranthene, pyrene, and chrysene. Non-methylated phenanthrene in the organic matter increased, an indication of a greater degree of oxidation, assumedly from reaction with metals-transporting brines and greater maturity. The organic matter became more carbonized by the process with a lowering of H/C ratios, albeit with an increase in O/C ratios, corresponding to an increase in rank from flamecoal to gascoal (Rentzsch, 1974, p. 409; Oszczepalski and others, 2002).

Forms and Textures

Cu minerals occur as disseminated 0.05 to 5 mm grains randomly scattered through the host sediment or commonly localized in the coarser part of individual sedimentary lamina-tions (fig. 8), or in any other coarser-grained sedimentary or pre-lithification feature. The typical Cu-sulfide grain occurs interstitially to detrital silicates or carbonates. The Cu sulfides are most consistently in grain contact with a syntaxial quartz overgrowth on detrital quartz or a syntaxial overgrowth of K-feldspar on detrital K-feldspar, with the overgrowths appearing to precede deposition of the sulfide, and with the sulfide appearing to form the ultimate filling of the former pore (fig. 16) (Hayes, 1984; Sweeney and Binda, 1989, fig. 6). However, given that K-feldspar and quartz overgrowths are, themselves, zonally more abundant in the zones richest in Cu sulfides (fig. 6), that apparent paragenesis possibly needs interpretation. If an epigenetic ore solution entered a clastic rock and was supersaturated with respect to chalcocite and K-feldspar and a substrate K-feldspar detrital grain is present, then the texture described above would form and could indicate simultaneous precipitation of chalcocite and the K-feldspar overgrowth.

Many unique textures occur among the disseminated sulfides, as well as among the zonally distributed authigenic gangue minerals of SSC deposits. A common texture in sandstone Cu deposits has been termed a bedform dissemination (Hayes and Einaudi, 1986), where the sulfide or authigenic gangue minerals are densely disseminated along a primary sedimentary structure, such as cross laminae (fig. 17). Sulfides along large-scale cross laminae at Mufulira were once (Garlick, 1972, p. 295) interpreted as a detrital distribution of sulfides, but at Spar Lake, careful grain-size analysis shows that the sulfide-rich bedform disseminations occupy coarser grained laminae than adjacent sulfide-poor laminae, simply repeating the generality (fig. 8) that every slightly coarser grained primary sedimentary feature becomes a locus of sulfide deposition, assumedly because metals-transporting fluid flux was proportional to permeability (T.S. Hayes, unpub. data, 1985).

A common special texture in reduced-facies subtype deposits is a coarser grained (than the background) sulfide nodule that is either spherical or slightly flattened within the bedding. This texture constitutes a substantial proportion of the disseminated Cu sulfides (perhaps 10–15 percent) in reduced-facies host rocks in the Kupferschiefer, in Zambia, in the Democratic Republic of Congo, and at White Pine. In the African examples, the nodule or lens is also the site of coarse-grained quartz or dolomite (fig. 8), which hosts the fluid inclusions interpreted as being from the initial ore stages (Annels, 1989; Dewaele and others, 2006; El Desouky and others, 2009).

Localization of ore sulfides in water-escape structures is also recognized in many SSC deposits. Lustwerk and Wasserman (1989) described the classification of water-escape structures and the localization of Cu sulfides within the structures at the Redstone, Canada, reduced-facies subtype deposit.

In red-bed-subtype deposits, the Cu sulfides may be concentrated surrounding plant remains, and they commonly replace the carbonized wood, preserving or pseudomorphing the wood cells in great detail. For example, at Nacimiento, N. Mex., the wood cells are filled with pyrite and later chalcocite, and the cell walls are replaced by bornite (Woodward and others, 1974, fig. 4). In these deposits, where the sulfide is pyrite, it is commonly the case that the cellular structure has not been collapsed by compaction. That may or may not be so
where a Cu sulfide fills the lumens. For uncollapsed lumens, some mineral, perhaps only pyrite, had mineralized the wood relatively soon after sedimentation. Where pyritized wood has uncollapsed lumens but Cu sulfide-bearing lumens are partially collapsed (Phillips, 1960, p. 47), the texture indicates that the Cu sulfides formed later than the pyrite, the usual paragenesis of SSC deposits.

**Hypogene Gangue Characteristics and Alteration**

Modern mines that exploit SSC deposits typically mine by bulk methods, generally open pit or underground room and pillar. With typical grades of 0.5 to 3 percent Cu, the bulk of the rock from these mines consists of waste minerals. The waste minerals include detrital components, such as quartz, feldspars, carbonates, and clays, and authigenic gangue minerals. Detrital components vary with the host depositional environments from clay-rich, to carbonate-rich, and to quartz-feldspar-rich. The authigenic gangue minerals are also an integral part of the mineralizing system, and unlike for vein deposits where vein-fill is easily separable from wallrock alteration, there are only mineral appearance/disappearance boundaries in SSC mineralized systems, and such boundaries are seldom macroscopically visible in outcrops or mine faces.

The following sections adopt a division in which “alteration” is restricted to those minerals distributed in space in zones up the hydrologic gradient from ore and extending into ore. For most SSC deposits, upgradient is in a downsectional direction, taken to indicate that the aquifers through which the Cu was transported were confined; that is, artesian pressure was driving the metals-transporting brines. By this division, gangue minerals are the other authigenic waste minerals found within orebodies. This convention views the “alteration” minerals as those that occur most abundantly in the rocks from which the Cu came, or in rocks through which Cu was transported en route to the deposits. The convention aids in distinguishing the conditions of Cu transport from the conditions of Cu deposition, and, as with other hydrothermal ores, the alteration minerals are useful in understanding metals-transporting and metals-depositing conditions (Meyer and Hemley, 1967; Einaudi, 1979).
Gangue Mineralogy, Paragenesis, and Zoning

Similar to the deposit-defining Cu-sulfide minerals, the gangue minerals of SSC deposits are zonally arranged; their appearance/disappearance boundaries parallel the zonation of the sulfides. The ore-stage and pre-ore gangue minerals are cements of the sedimentary rocks and replacements of earlier detrital and cementing minerals.

Pyrite is an important gangue mineral in downgradient deposit-fringing pyritic zones, although there are probably two generations of pyrite in many deposits. Pre-ore, first-stage pyrite is commonly framboidal in reduced-facies-type deposits (Brown, 1971, fig. 25; Oszczepalski, 1989, plate II, fig. 5). In coarser host rocks, the pre-ore pyrite is a cementing mineral. In red-bed-type deposits, pre-ore pyrite commonly replaces carbonized wood fragments (Woodward and others, 1974, fig. 4; Thaden and others, 1964, fig. 16). In the reduced-facies deposits in the Democratic Republic of Congo, a second-stage cobaltiferous pyrite is probably from the ore-stage (Bartholomé and others, 1971), and that pyrite typically contains 4 wt. % Co. In Zambian rocks, concentrations of as much as 42 wt. % Co in pyrite were measured (Riley, 1968; Brown and Bartholomé, 1972). Ore-stage, Co-anomalous pyrite, typically about 0.015 wt. % Co, is recognized, as well, in the Spar Lake sandstone Cu deposit (Hayes, 1984) where it coats earlier cementing pyrite, overgrows the cementing pyrite, and replaces detrital quartz and feldspar of the framework to form cubic crystals or clots of cubic crystals with mutually interfering growth (fig. 18).

Iron-rich calcite cements are found within pyritic zones of sandstone Cu deposits at Spar Lake and Dzhezkazgan (Gablina and Tsepin, 1975; Hayes, 1984, 1990). The Spar Lake ferroan calcite has as much as 3.5 mole % FeCO$_3$, and it typically has more than 2 mole %. Fe-rich carbonates are found in approximately the same zonal position in the Cashin and Lisbon Valley deposits in the Paradox Basin, United States.

All mineral zones of SSC deposits contain leucoxene that pseudomorphs former ilmenite in exsolution-created, trellis-textured ilmeno-magnetite detrital grains or the ilmenite plates of exsolution-created, stacked-plate-textured ilmeno-hematite detrital grains (fig. 19). The magnetite and hematite of the original detritus is destroyed, and sulfide minerals, which incorporate Fe that earlier may have been within the detrital magnetite, hematite, and ilmenite, are commonly in close association (Dimanche and Bartholomé, 1976; Hayes, 1983; Hayes, unpub. observations from Rock Creek, Mont.; Paoli, Okla.; Farwan no. 1 occurrence, Saudi Arabia; and Jabal Muryyii deposit, Saudi Arabia; 1980; 1988; 1999; and 2000, respectively).
Figure 18. A, Reflected-light photomicrograph in plane-polarized light of the margin of a compound pyrite clot, with a core of cementing pyrite surrounded by a continuous pyritic outer rim from the pyrite-calcite zone (fig. 6) at the Spar Lake sandstone copper deposit. The red-outlined box identifies the field of view of B and C. B, Part of the continuous outer rim of pyrite, also in reflected light. The section has been etched for 1 minute in concentrated HNO₃, followed by 10 seconds in concentrated HCl to enhance any internal textures within the pyrite. A cuspat internal shape is within one of the cubic pyrite grains forming the continuous outer rim of the clot. The cuspat shape is similar to pyrite grains found cementing quartz grains in the interior of the clot. C, “Map” of cobalt characteristic X-rays from the electron microprobe of the same cubic grain shown in B. The parts of the pyrite cube outside of the cuspat interior shape are anomalous in cobalt. The pyrite clot formed in two steps; first a cementation between detrital quartz grains by ordinary pyrite, and then replacement of detrital quartz in the outer rim by cobalt-rich pyrite, overgrowing the cementing pyrite to form a continuous outer pyrite rim. The cobalt-rich pyrite precipitated later than the ordinary pyrite.

(Sweeney and Binda, 1989, plates I, II, and III; Sutton and Maynard, 2005, figs. 4 and 7). Quartz overgrowth on detrital quartz grains is prominent at Spar Lake and has the same kinds of textural relations to sulfides and carbonates that the K-feldspar overgrowths do (see below discussion of barite).

In fine-grained dolomitic siltstones in the Democratic Republic of Congo and Zambia, authigenic quartz replaced crystals and nodules of anhydrite. Annels (1974) documented widespread replacement of anhydrite nodules by calcite and sulfides, by pyrite of peripheral non-ore zones and the Cu sulfides of the ores at Nkana North Limb. Annels showed direct proportionality between Cu grades and abundance of “carbonate lenticles,” that is, anhydrite nodules that had been replaced. In the Democratic Republic of Congo, quartz, sulfides, and dolomite have replaced anhydrite (Dewaele and others, 2006). Some of the quartz and dolomite is pseudomorphic after anhydrite nodules and anhydrite crystals, and anhydrite relicts are found at places within the authigenic minerals (Muchez and others, 2008). Additionally, given the fluid-inclusion microthermometric results from inclusions hosted in such quartz-after-nodular-anhydrite, with homogenization temperatures of 105 to 192 °C (Annels, 1989; Dewaele and others, 2006), some of such quartz is probably ore stage, not from early diagenesis.

Authigenic tourmaline locally makes up 4 percent of the rock in the Central African Copperbelt (Darnley, 1960, p. 147; Selley and others, 2005), but it is not known if it has a zonal distribution.

Hematite, Mn-bearing minerals, and barite accompany bornite in the sulfide assemblages at Spar Lake. A fine dusting of pore-lining and pore-filling hematite (fig. 19C) and fine pigmenting hematite have been described at White Pine (Brown, 1971, p. 555), Kamoto (Dimanche and Bartholomé, 1976, p. 195), Witvlei (Anhaeusser and Button, 1973, p. 292), and Seboruco, Venezuela (Dimanche and Bartholomé, 1976, p. 199).
This is ore-stage hematite and should not be confused with the early diagenetic hematite or other ferric oxyhydroxides formed in early diagenesis of red beds. This earliest variety of hematite common to all red beds is recognizable because it ubiquitously rims and outlines detrital grains of those red beds (Turner, 1980, p. 316–317). Multiple hematite dust rims are found in some bornite- and chalcocite-bearing ore zone rocks at Spar Lake, separated by overgrowth quartz or K-feldspar, and second-rim hematite is probably from the ore stage (fig. 21). Near the bornite appearance at Spar Lake, manganiferous calcite, with as much as 12 percent MnCO₃, is later than ferroan calcite (fig. 22) (Hayes, 1990). Spar Lake Cu sulfides typically appear to be ultimate pore fillings, deposited after quartz- and K-feldspar overgrowths and with paragenetic position similar to the calcite cements (fig. 16). Weathered rocks with greater than 20 percent Mn are present in SSC deposits at Feinan, Jordan. The Mn may be sited within carbonates in the unweathered equivalents. Hypogene Mn-bearing oxides are found in great abundance, together with chalcocite, in the mineralized layers of Boléo, Mexico (Conly and others, 2006). Barite generally appears near the same place that bornite does in Spar Lake ore. It occurs as a cement that fills pores apparently after quartz and K-feldspar have formed overgrowths on detritus (fig. 23). It shares former pores with chalcocite or bornite, with no clear paragenetic relation to the sulfides. It increases in abundance in the central chalcocite-chlorite zone. It is rarely found in rocks with galena or chalcopyrite but has not been found co-existing with pyrite. Barite has also been described...
at Konkola, Zambia (Sutton and Maynard, 2005), in several different red-bed-type deposits in New Mexico and Utah (Phillips, 1960), in Weissliegendes sandstones, some ore-bearing, in Poland (Michalik, 2001), and it probably occurs in many other deposits together with bornite and chalcocite. Barite and albite form the paragenetically latest pore-filling cements at Spar Lake, along with the Cu sulfides.

Chlorite occurs as a pore-lining cement towards the centers of former pores that have overgrowths of quartz or K-feldspar but with chalcocite, digenite, and bornite still more central to the former pores at Spar Lake. Abundant Fe-rich chlorite is probably ore stage at White Pine (Hamilton, 1967; A. Brown, École polytechnique de Montréal, written commun., 2011). Additionally, authigenic chlorite is found in ore at Witvlei (Anhaeusser and Button, 1973), at Kamoto (Bartholomé and others, 1976, p. 34), and from the deposits in Zambia (Darnley, 1960). Spar Lake chlorite is Fe-rich, as much as 39 wt. % FeO and typically greater than 30 wt. %. This requires siting Fe as both Fe\(^{2+}\) and Fe\(^{3+}\) (Foster, 1962; Brown and Bailey, 1962), and such Fe-rich chlorites are typically formed in advanced, deep burial diagenesis or by hydrothermal processes, not during metamorphism (Brown and Bailey, 1962; Curtis and others, 1985). At Spar Lake, the manganiferous calcite of the bornite-calcite zone disappears abruptly at an easily mappable mineralogical boundary (Hayes and Einaudi, 1986, fig. 17D, p. 1922) with the chalcocite-chlorite zone of that deposit. Amounts of carbonate minerals change from about 5 v% at the inner edge of the bornite-calcite zone to only 0.3 % disseminated ankerite inward of that boundary.

Distribution of chlorite and calcite is similar to that of Spar Lake at other deposits where both have been observed, although details of their distributions are not as well known. At White Pine, the rocks are calcareous through all the zones from pyritic into and including most of the chalcocite-bearing rocks, but the calcite is not present in parts of the Lower Sandstone (Copper Harbor Conglomerate) underlying the ore. Some of the footwall Copper Harbor rocks have calcite and chlorite as sandstone cements, and others have only chlorite (Hamilton, 1967).

**Figure 20.** Sequence of authigenic minerals in the galena-calcite zone (fig. 6) of the Spar Lake sandstone copper deposit. A, Photomicrograph in cross-polarized transmitted light. B, Cathodoluminescent photomicrograph of the same field of view. Potassium feldspar detrital grain cores are luminescing blue. C, Image made by drawing grain boundaries and identifying mineral grains on an overlay of the upper and middle images. Mineral abbreviations: Kd – K-feldspar detritus, Ko – K-feldspar overgrowth, gn – galena, Fe-cal – ferroan calcite, t – albite law twinning plane within K-feldspar. The apparent sequence is: (1) detrital quartz and K-feldspar, (2) syntaxial K-feldspar overgrowth on detrital cores, and (3) pore-filling cementation by galena and ferroan calcite. As discussed, a reasonable interpretation is that K-feldspar overgrowth, galena, and ferroan calcite are partly synchronous.

Distribution of chlorite and calcite is similar to that of Spar Lake at other deposits where both have been observed, although details of their distributions are not as well known. At White Pine, the rocks are calcareous through all the zones from pyritic into and including most of the chalcocite-bearing rocks, but the calcite is not present in parts of the Lower Sandstone (Copper Harbor Conglomerate) underlying the ore. Some of the footwall Copper Harbor rocks have calcite and chlorite as sandstone cements, and others have only chlorite (Hamilton, 1967).

**Alteration**

The increasingly albite-rich assemblages in the “upstream” direction for metals-transporting solutions at such places as Spar Lake and Oamites probably continue into rock where Cu is depleted. With no Cu sulfides and the presence of authigenic hematite, such rock is analogous with the altered tongue of sandstone-hosted roll-front U deposits. In the Revett Formation surrounding Spar Lake, such rock typically has less than 5 ppm Cu compared with 20 ppm within rock totally unaffected by any mineralizing system processes (Hayes, 1990). Such rock almost certainly represents the source for Cu in the stratabound deposits. It is actually, however, quite difficult to recognize as
altered rock, because it occurs distant from Cu-mineralized rock and because the albite is finely disseminated through the rock. The albite has the appearance of being detrital at hand specimen or hand lens scale, and it is only at the scale of the petrographic microscope that the albite is seen to replace all the detrital plagioclase and occur as a pore-filling cement (fig. 24).

Whole-rock composition, however, identifies this rock as dramatically unusual. In a (meta)clastic rock with 15 percent total feldspars, and 12 percent of that being plagioclase, these rocks typically have less than 300 ppm Ca after albitization. Albitic alteration in cross-stratal zones is widespread in the Central African Copperbelt (Selley and others, 2005) and in the Udokan sedimentary complex that hosts the giant Udokan deposit. However, the albitization in the Udokan complex has not been related directly to the deposits but has been shown to be zonally related to faults (Abramov and Chernyshova, 2001; Abramov, 2008).

Albite is the sodic mineral that can be directly correlated with Cu source zones near several SSC deposits, but other sodic minerals may signal metals mobilization and depletion. In the Redstone (Coates Lake) system in Canada, the footwall red beds contain abundant glauberite (monoclinic NaCa(SO₄)₂) (R. Lustwerk, oral commun., 1985). In rocks of the Salib Arkosic Sandstone Formation, which are the footwall to SSC mineralization in the Burj Formation at Feinan and Wadi Abu Kushaiba, Jordan, the sodic phase is probably paragonitic illite (T. S. Hayes, unpub. data, 2001). Near the Dzhezkazgan deposits in Kazakhstan, mudstones upsection from the mineralized Dzhezkazgan Formation have been massively altered to riebeckite (Na₃[Fe³⁺Mg]₀.₄Fe₃⁺₂Si₂O₇(OH)₂) (B. Syusyura, oral commun., 2009). Similar alteration is known from the area surrounding Tenke-Fungurume in the Democratic Republic of Congo (M. W. Hitzman, oral commun., 2010). In the Agadez area, Niger, where there are many red-bed-type copper occurrences, analcime (monoclinic Na[AlSi₃O₈]·H₂O), the sodium zeolite, is remarkably abundant in the coarse arkosic red beds underlying the three separated strata that host copper mineralization (Imreh and Nicolini, 1962). That rock is locally named “analcimolite.” Thus, sodic minerals characteristically mark Cu source rocks for SSC deposits.

A second mineral that characterizes metal source areas for SSC deposits is hydrothermal hematite, which is the direct analogy of the ferric oxides of the altered tongue of a sandstone-hosted roll-front U deposit. Hematite in such a zonal arrangement is best known from the mineralizing systems

\[ \text{Kd} – \text{detrital K-feldspar, Ko – K-feldspar overgrowth, qd – detrital quartz, qo – quartz overgrowth, mcal – manganiferous calcite, cc-dig – chalcocite-digenite. The apparent sequence is (1) detrital quartz and K-feldspar, (2) hematite dust rims on detrital cores, (3) quartz and K-feldspar overgrowths on detrital cores, (4) a second stage of hematite dust rims deposited on quartz overgrowths, (5) a second stage of quartz overgrowth, and (6) pore-filling copper sulfides and manganiferous calcite. The manganiferous calcite may have started growing any time after (1), as could have the chalcocite-digenite, but the final growth of both had to post-date the second generation of hematite and probably the final stage of quartz overgrowth.} \]
of the Kupferschiefer, where it is called Rote Fäule (figs. 4 and 5). Rote Fäule, where it is present in rocks that lacked hematite prior to the ore event, appears as red spots that are centimeters to tens of centimeters in scale (fig. 25). Because it occurs as discrete, large spots or streaks, it is easily identified within the Weissliegendes, Kupferschiefer, and Werra Carbonate stratigraphic units (fig. 25). There is no description of Rote Fäule alteration within the Rotliegendes Formation red beds that lie downsection from the mineralized stratigraphic interval, and the downsection Rotliegendes rocks have assumedly been red beds since earliest diagenesis, so a second, later stage of hematite from the ore-stage either does not exist or it may not be identifiable.

In the sandstone Cu deposit at Paoli, Okla., hematite is clearly identifiable in a zone through which metals have been transported to form an SSC deposit. The deposit, itself, has a roll front geometric form (Shockey and others, 1974). The altered tongue sandstone is mottled red and light gray, and the hematite lends a brick-red color to Garber Sandstone host rocks that are pastel green in color where they are Cu- and Ag-mineralized or non-oxidized containing trace pyrite. Mudstones that occur upsection and downsection from the mineralized sandstones are ordinary red beds, although they are pastel green for a few centimeters at their contacts against the sandstone beds. The pastel green host sandstones also have red-mud intraclasts within them that have green rims; the type B pillars (Lustwerk and Wasserman, 1989) consist of pale green sandstone cutting red mudstone. The sandstones lack any carbonaceous plant fossils. These observations suggest that the pastel green color is a secondary feature: the sandstones were reddened during early diagenesis, then bleached prior to the ore event, and, finally, oxidizing metals-transporting solutions infiltrated the sandstones and deposited the chalcocite and native silver, and produced the red hematitic mottling in the altered tongue.

It remains speculative as to whether barite also characterizes zones of metals mobilization (fig. 6). Calculations in Hayes (1983, p. 221–230) suggest that barite would be stable in such rocks, and there are occurrences of barite disseminated in red-bed sandstones in several places in Oklahoma, near SSC deposits such as Paoli and Creta, and even within the altered tongue sandstones of Paoli. Barite in the Garber Sandstone in this zonal position at Paoli forms barite “roses,” and barite roses are found over more than 100 km along the outcrop belt in the Garber in central Oklahoma (Ham and Merritt, 1944). However, it is unclear whether such rocks have lost Cu to the hydrothermal systems. Barite is also found in considerable abundances within the ore system at Konkola, Zambia (Sutton and Maynard, 2005), but the barite may not take a single regular place within the mineral zonation; it is found within ore, below ore, and (or) above ore, and it is found in footwall clastic rocks of the Copperbelt Orefield Member, or even in the shale with grit that overlies that member.
Textures and Grain Sizes of Hypogene Gangue

The reason that alteration is seldom discussed in the literature of SSC deposits is that the grain sizes and textures of the alteration and gangue minerals, such as albite and hematite, are not different than other minerals in these altered rocks. As such, this alteration is not distinguished in appearance except where it forms nearly total replacements of the other minerals of the rock or unless it occurs in disconformable bodies. In two mineralizing systems, however, there are specific rock textures that distinguish the hematization where Cu is depleted. Related to deposits of the Revett Formation, Montana, and to those of the Burj Formation at Wadi Dana and Wadi Abu Kusheiba, Jordan, there is an unusual, cross-laminar, swirled banding of hematite in sodically altered Cu source rocks (fig. 26). The banding has the appearance of Leisegang banding, but the bands are pervasive through the rock volume rather than confined to a joint or fracture plane. Microscopically, the hematite in this texture is not distinguishable from the hematite of other red beds; it forms rims outlining detrital grains and is enclosed in later cements, commonly occurring between quartz detritus and quartz overgrowths. The banding in the Salib Formation interpreted source beds in Jordan is well exposed in the archeological ruins at Petra (fig. 27).

Although swirled hematitic banding characterizes altered rock of these two systems, such banding occurs in many other sandstone units where it has no relation to SSC deposits. For example, many different sandstone units of the Colorado Plateau, United States, show such banding.

Lateral and Vertical Dimensions

The lateral and vertical dimensions of alteration and Cu source areas related to a SSC system is uncertain, in part because there are only a few systems in the world where the alteration is recognized. Nonetheless, it can be stated that the dimensions of whole systems are enormous, apparently nearly the scale of whole basins. In the Spar Lake and associated deposit area, only a zone about 100 km long and 50 km wide has any Revett Formation rocks that are suitable host rocks, those which are white, gray, or green. These rocks in the Western Montana Copper Sulfide Belt (Harrison, 1972) average 0.5 km in thickness. To the west and east of the Copper Sulfide Belt, rocks of the Revett Formation are lavender gray and are entirely albite-altered and depleted in Cu. Downsection rocks of the Burke Formation and upsection rocks of the St. Regis Formation, both within the Cu-sulfide belt and to its west and east, are almost entirely albite-altered and depleted in Cu. Albite-altered rocks of the Ravalli Group continue westward for about 70 km...
Figure 24. Sequence of authigenic minerals in the chalcocite-chlorite zone (fig. 6) of the Spar Lake sandstone copper deposit. A, Photomicrograph in transmitted, plane-polarized light. B, Photomicrograph in transmitted, cross-polarized light. C, Image drawn by tracing grain boundaries on the first two. The section has been etched for 90 seconds in the vapors above concentrated HF acid. It has then been stained sequentially by sodium cobaltinitrite and Amaranth solutions by the method of Gabriel and Cox (1929), as subsequently modified by Keith (1939), Chayes (1952), and Rosenblum (1956). Using this method, K-feldspar stains bright yellow, and calcium-bearing plagioclase stains magenta colors with intensity of the color increasing with calcium concentration in the plagioclase. Pure albite etches but takes no stain color. Mineral abbreviations are: Qd – detrital quartz, Qo – quartz overgrowth, bn – bornite, and cc- chalcocite. The apparent sequence is (1) detrital quartz, (2) quartz overgrowth, (3) chalcocite, bornite, illite, and albite, and (4) albite (the central pore-filling of albite probably formed after the bornite ceased growing).

It is unlikely that from the vast volumes of albitized Ravalli Group rocks to the west and east of the Western Montana Copper Sulfide Belt, the Cu moved hundreds of kilometers laterally to orebodies within the belt. Mineralizing solutions probably did not migrate laterally for more than 100 km and perhaps not even for tens of kilometers. It is likely, instead, that the Cu moved mainly upward across stratification. In the areas to the west and east of the Cu sulfide belt, the Cu probably was deposited in the laterally persistent, although thin “greenbed” occurrences described by Harrison (1974), such as the Blacktail Mountain occurrence at the contact of the Spokane and Empire Formations (Harrison and Reynolds, 1979).

Weathering and Supergene Ore and Gangue Characteristics

The weathering of SSC deposits has received little attention from geologists in recent years, although it was the subject of debate at earlier times. In part, the lack of recent attention is because changes to Cu grade because of supergene processes are not particularly great for most combinations of deposit subtype and weathering conditions. In part, the lack of attention is because the geometry of an SSC deposit is rarely conducive to much supergene enrichment. These deposits tend to be sheetlike to tabular, so they generally have limited areas exposed in outcrop, and only shallow bedding attitudes lend themselves to making an extensive enrichment “blanket.” Steep bedding attitudes can also lend themselves to enrichment, although not with a blanket geometry, so the enrichment is not easily detected. Supergene enrichment, however, may be important in some deposits, particularly in the Democratic Republic of Congo and Zambia. This section will first discuss the mainly earlier work on supergene enrichment for the deposits of the Central African Copperbelt. It will then turn to descriptions of mineralogy and associations in leached cappings, oxide Cu zones, and enriched Cu-sulfide zones of SSC deposits in central Africa and elsewhere.
Climate is always an important variable in the process of weathering and supergene enrichment. Supergene effects in the Central African Copperbelt may differ from the world’s other SSC districts because of its climate. The Copperbelt climate is warm and relatively humid, with 120 to 140 cm/yr of rain. Lateritic soils are developed locally along the Copperbelt, but the lateritization nowhere produces a soil profile with economic Fe or aluminum concentrations. Weathering is relatively deep over much of the area; average depths to bedrock are 4 m (Maree, 1961, p. 65). Among the world’s known SSC districts, only Seboruco, Venezuela, and Bahia, Brazil, share a similar latitudinal zone with the Central African Copperbelt, and even within a latitudinal zone, rainfall, a first-order control on supergene enrichment processes, can vary greatly.

Hitzman and others (2005, p. 627) stated that “supergene processes appear to be responsible for forming much of the chalcocite” of Zambian Copperbelt deposits and that “the high grade of many orebodies in Zambia and probably Congo, relative to other sediment-hosted stratiform Cu deposits may be due in large part to supergene alteration.” However, it is likely that where the host rock is finely laminated, carbonaceous dolosiltstone, as is the case for most of the deposits of the Democratic Republic of Congo, the chalcocite contained therein is mostly hypogene, because supergene chalcocite is deposited from acidic shallow groundwater solutions. At pHs of about 3.5, typical of those forming supergene chalcocite
Chalcocite group minerals, including chalcocite, djurleite, digenite, and anilite, though, are the only Cu-, Co-, or Ag-bearing minerals in these deposits whose hypogene versus supergene origin can be strongly questioned.

The origin of chalcocite in Zambian and Democratic Republic of Congo SSC deposits was the subject of lively debate in the 1920s through the 1950s (Bateman, 1930; Schneiderhöhn, 1932; Oosterbosch, 1951; Garlick, 1961a). The debate seemed to resolve itself with the widespread acceptance of Garlick’s (1961b) syngenetic hypothesis of deposit genesis, but with that hypothesis now mainly viewed as incorrect, the primary or supergene origin of most of the chalcocite needs to be better addressed. Certainly, some of the older observations still apply. The literature that addresses supergene or hypogene origin of chalcocite group minerals in Central Africa all dates from before 1965.

Bateman (1930) specifically addressed the problem and did so from extensive ore microscopy (“one hundred or so representative sulphide ore specimens,” p. 388). His most important conclusion was that there was both hypogene and supergene chalcocite, but that (p. 404) “perhaps the major part of the Rhodesian chalcocite is of hypogene origin.” Bateman’s work pre-dated much of the present-day knowledge of the phase relations among the chalcocite-group minerals. At the time, there was recognition from work on supergene blankets at porphyry Cu deposits in the American Southwest, of two phases, termed white chalcocite and blue chalcocite, on the basis of their apparent colors in reflected light where they occur together. The chalcocite with composition, Cu$_2$S, had a low-temperature polymorph that was thought to be probably orthorhombic, and the high temperature one was thought to be isometric.

Buerger (1942) found that the “blue chalcocite” was a separate mineral that he named digenite, that it was isometric, that it formed solid solutions at high temperatures over a range of compositions between Cu$_9$S$_5$ and Cu$_2$S, and that at low temperature it had a much more restricted composition range near Cu$_9$S$_5$ and had non-isometric crystallography. Djurle (1958) found another phase of composition Cu$_{1.96}$S that had optical properties near those of chalcocite, and djurleite was soon found in numerous natural assemblages, notably including supergene blankets and SSC deposits (Butte, Mont., and Tsumeb, Namibia: Roseboom, 1962; Kenneecott, Ala.: Roseboom, 1966; Copiapó district, Chile: Sillitoe and Clark, 1969; White Pine, Mich.: Brown, 1971). Roseboom (1966) defined most of the phase relations among these and found that chalcocite (pure Cu$_2$S) had high temperature hexagonal and high pressure tetragonal polymorphs, the latter being quite rare, as well as a low-temperature monoclinic, pseudo-orthorhombic polymorph. Morimoto and others (1969) identified anilite (orthorhombic Cu$_7$S$_4$) as yet one more phase intermediate in composition between chalcocite and covellite. Sillitoe and Clark (1969) almost immediately identified anilite as a key component in the “sooty” chalcocite, which is widespread in supergene enrichment blankets, together with covellite, djurleite, and monoclinic chalcocite.
Weathering and Supergene Ore and Gangue Characteristics

Morimoto and Kato (1970) then redefined the low-temperature phase relations among chalcocite, djurleite, digenite, anilite, and covellite, including finding a eutectic of 70 °C between anilite and djurleite below which temperature digenite would not form. This digression into sulfide phase relations is needed to evaluate which among Bateman’s (1930) findings about Zambian “chalcocite” remain valid and which do not.

Bateman (1930, p. 394–395) described “subgraphic” intergrowths of bornite and “chalcocite” (fig. 28) and noted that they are common in Zambian ores and that he had never seen them in chalcocite of supergene origin. With the understanding that the chalcocite of the intergrowths is probably digenite, the conclusion stands that these types of exsolution intergrowths are probably hypogene. In fact, where they have the phase proportions of 0.33 to 1.00 bornite component, these intergrowths mark the unmixing of a solid solution that was stable at temperatures certainly greater than 82 °C and, across most of that range of compositions, greater than 180°C (Grguric and others, 2000). Such a grain with molar proportions of 0.56 bornite and 0.44 digenite is prominent in figure 28. Bateman (1930, p. 396–401) also found (1) needle-thin octahedral intergrowths of bornite in “chalcocite,” (2) lamellar intergrowths of blue and white “chalcocite,” (3) triangular pseudo-eutectic textures of unmixed bornite and “chalcocite” within triangular arrays of “chalcocite” blades, and (4) triangular-patterned lattice intergrowths of bornite and “chalcocite” in Zambian ores. Bateman reasoned that all these marked a formerly isometric crystal system for the “chalcocite,” and he attributed the textures to hypogene origins at greater than 91 °C, which was thought to be the thermal maximum stability of low-temperature monoclinic, pseudo-orthorhombic chalcocite. In fact, each of the textures indicating a former isometric crystal system was an exsolution of digenite and either bornite or chalcocite starting from a higher temperature digenite solid solution (Roseboom, 1966), and all these did originate at higher temperatures, thus suggesting hypogene origin. They could have, however, been deposited as low-temperature composite sulfide grains indicating replacement, then have been ramped up to high temperature and homogenized in the digenite solid solution compositional field, and then exsolved as they returned to lower temperature. But if that was how they formed, then other textures (see below) would also have gone through that history. Finally, Bateman (1930) observed rim-pattern and mesh-pattern textures where chalcocite replaced bornite or chalcopyrite, commonly also containing traces of covellite, and these he attributed to supergene origin. In short, knowing now about digenite, djurleite, and anilite, Bateman’s (1930) conclusion should be accepted that Zambian chalcocite-group minerals are of both hypogene and supergene origin and, if his distribution of samples was representative, that “perhaps the major part” of the chalcocite is hypogene.

Oosterbosch (1951) described mineral distributions at Fungerume, Democratic Republic of Congo, well enough to start further conjecture about weathering and enrichment, yet came to the conclusion that all the “chalcocite” had the same supergene origin. He (p. 131) stated that there is a zone of malachite and Cu silicates from surface to 100 m depth, underlain by a zone of chalcocite and “green salts” to a depth of 250 m, underlain by sulfides only, although with only scant traces of chalcopyrite. He found limonite and local occurrences of native copper to the greatest depths they had then drilled, 990 m, and noted that the top of sulfides was only “slightly related” to the present water table. With digenite then known, he noted digenite only below 250 m, and he found evidence for pre-existing cubic structure in the chalcocite only at those levels. Oosterbosch wrote (p. 142):

“The upper sulfide zone [that lying stratigraphically above the RSC beds] in which the chalcocite dominates contains 6 to 8 percent copper; the surficial oxidation causes the grade to drop on the average to 5.5 to 6 percent. Marked impoverishment appears to start only above a depth of 100m [that is, impoverishment within the malachite and copper silicates zone as described above].”
Finally, Oosterbosch (1951) gave the distribution of grades of chalcocite orebodies stratigraphically below the RSC beds. Average grade at 100 m depth was 5.99 percent Cu, at the top of his non-cubic, low-temperature chalcocite lacking earlier cubic structure, with green salts zone, totally lacking digenite. At 400 m depth, it was 8.93 percent Cu, in a zone containing some digenite and formerly cubic chalcocite. At 680 m, it was 5.43 percent Cu, and at 715 m, it was 5.58 percent Cu, with both from zones having some digenite and formerly cubic chalcocite.

Together, these data probably indicate some supergene enrichment of copper as low temperature (monoclinic, pseudo-orthorhombic) chalcocite at depths extending from 100 m to somewhat below 400 m depth at Fungurume. Nearly all of the chalcocite between 100 and 250 m depth is supergene, because none in that interval has cubic (digenite solid solution field) parentage. Everywhere below 250 m, some hypogene chalcocite is certainly present and came from the cubic, high-temperature solid solution with digenite as further evidenced by the presence of digenite that is absent everywhere above 250 m.

For the case of Roan Antelope, Brummer (1955, p. 288–290) observed digenite, white chalcocite, and sooty chalcocite. He attributed the sooty chalcocite, which was more abundant in upper levels and was associated with cuprite and malachite, to supergene origin. He stated that sooty chalcocite does not furnish any appreciable tonnage of ore at that mine, which is now called the Luanshya deposit.

Weathering effects on the ores of the Zambian Copperbelt were discussed in Mendelsohn (1961a), and the various statements reflected the lack of consensus about the importance of supergene effects. First, in general, Mendelsohn wrote (p. 129):

“In the southern part of the Copperbelt west of the Kafue anticline, below a narrow layer of mixed sulphide and oxide with some enrichment there is clean sulphide ore with little or no enrichment. Farther north there has been more supergene activity. At Nchanga a considerable part of the mineralization is of supergene origin and there has probably been much enrichment; . . . At Bancroft [now called Konkola] there has been leaching and redeposition of the copper to a depth of many hundreds of feet. The areas in which the orebody was leached and those that were enriched are related to the fold structure. At Mufulira there was supergene enrichment in all three orebodies in the higher levels. . . . The main supergene minerals are malachite, chalcocite, cuprite, chrysocolla, and vermiculite, and lesser minerals are tenorite and native copper. The vermiculite is an alteration product of biotite, or possibly in places other micas, and contains copper both as inclusions between the plates of the mica and as part of the crystal lattice.”

Those who described individual deposits in 1961 found what they considered to be varying effects of weathering and supergene enrichment across the Copperbelt. Mendelsohn (1961a, p. 400) said that “most of the oxidation appears to consist in the fixing of the Cu in new minerals with little or no migration.”

Important but localized enrichment took place at Chibuluma. Winfield (1961, p. 339) wrote:

“A gossany sandstone forms the entire sub-outcrop of the orebody and extends to between 200 and 270 feet below surface. The sulphides in this zone have been leached out and replaced by limonite. Malachite and chrysocolla are absent except near the abrupt change to sulphide at the upper edge of the orebody. The ore below the gossany sandstone has been enriched down to the 350-foot level, chalcocite and covellite commonly replacing the primary sulfides.”

The ore continued to more than 2,500 ft below surface. Because most of the ore at Chibuluma was chalcopyrite, with bornite found only in western and eastern extremes, the supergene origin of the chalcocite and covellite was obvious below the leached capping and the thin malachite-chrysocolla (oxide Cu) zone.

At Nkana, the distribution of supergene versus hypogene chalcocite has never been clearly described. However, native copper is known to depths as great as 2,700 ft in the Porous Sandstone [that overlies the Copperbelt Orebody member], and sulphides extend to a depth of at least 6,000 ft (Jordaan, 1961, p. 323).

Farther north, Nchanga is one of the two places where supergene Cu-bearing vermiculites were first identified (Bassett, 1958). Cu-vermiculite mineralization extends down along the 20° dips in sandstones to depths of greater than 800 ft, but it gives out below that, apparently having formed in weathering from earlier phlogopitic biotite. Notably, the pre-mining water table was nowhere more than 40 ft below surface. Features like this and the ones at Fungurume have led workers to repeatedly suggest a Mesozoic to Tertiary period of deep supergene oxidation (Selley and others, 2005) assumedly dating from a time when the water table was much deeper than it presently is. McKinnon and Smit (1961) noted that masses of cuprite commonly form a border around chalcocite layers. They stated that sulfuric-acid solutions have reacted with the aluminous minerals leaving kaolinite as a residual product. The arkose in the footwall of the orebody is particularly kaolinitized.

For the Konkola area, although weathering effects were profound, Schwellnus (1961, p. 228) wrote that, “in any area, the total number of copper units will remain the same regardless of their distribution by supergene solutions.” This indicates that the migration of Cu for the most recent supergene minerals was over only small distances. On a map (p. 229), Schwellnus showed the zone of malachite with lesser chalcocite and chrysocolla extending irregularly to a
depth of nearly 1,900 ft, and with lesser malachite occurrence extending below probably 2,100 ft. He (p. 234) described heavily leached ore sections where carbonate minerals had been removed to produce “a porous mass of chalcocite, goethite, and Mn wad in cavities parallel to the bedding.” He also showed major-element data (p. 224) that documented prominent leaching of the carbonates from the ore horizon [Copperbelt Orebody Member] in upper levels of the mine, corroborating our earlier statement about the likelihood that dolomite would dissolve in and above zones where supergene chalcocite is deposited.

At Mufulira, the identification of a leached capping was entirely straightforward. Brandt and others (1961, p. 450) reported that leaching of sulfides of the three orebodies was general to a depth of 200 ft from surface and sporadic below that depth, extending within some Footwall quartzites and grits to 2,000 ft or more. Near the surface, the ore beds were leached and porous in contrast to the solid and fresh unweathered orebody quartzites, and they contained Cu as noneconomic malachite, chrysocolla, cuprite, and native copper. It was simply assumed that all the chalcocite was supergene in origin, probably a correct assumption for the small amounts of chalcocite with some cuprite and malachite in porous quartzites adjacent to the Cu orebody that elsewhere consists only of bornite and chalcopyrite. Where 200 ft of quartzite leached capping with subecononic Cu oxides is found, and the rock below contains ore grades as chalcocite, it follows that supergene enrichment as sulfides probably occurred. Notably, though, four different textures illustrated by Bateman (1930) indicating “chalcocite” that had formed earlier in isometric crystals came from Mufulira. As above, the “chalcocite” with isometric heritage was probably hypogene and originated as the high digenite solid solution.

Mineralogy and Mineral Associations in Weathered and Enriched Zones

From the above, in the Central African Copperbelt, Cu undergoes locally significant enrichment as chalcocite and covellite, and probably djurleite and anilite, below a zone of leached capping that is highly variable in thickness but appears to extend downward for a few to 90 m from surface. The capping is followed downward by a zone of oxide Cu minerals and native copper that appears to typically extend from the base of leached capping to as much as a few hundred meters from the surface. Further below is a zone of supergene enriched chalcocite ore that extends at least to 400 m depth (Fungurume) and probably more likely to at least 800 m depth (Nkana, Nchanga, and possibly Konkola). Yet, Bateman’s (1930) principal conclusions stand, and the statement of Hitzman and others (2005) does not appear well supported: there are “chalcocites” of both supergene and hypogene origin in the Central African Copperbelt, and the major part is probably hypogene.

Co in Central Africa has a different behavior than Cu in the zone of weathering. Co is residually enriched within the zone of weathering and oxidation. The behavior is known but has not been studied systematically. Gauthier and Deliens (1999) described a well-developed “cobalt cap” at the Musonoi deposit in the Kolwezi klippe and in the nearby K.O.V. deposit.

The following sections describe, as well as possible, the minerals in the leached capping, the oxide Cu zone, and the chalcocite enrichment zone in Central Africa and elsewhere. Obviously, there are still major gaps in our understanding of these.

Leached Capping

In the Central African Copperbelt, mineralized arenites, where they have been leached of their Cu content near the surface, are porous, friable, and “gossany,” and the only two minerals that are clearly formed in the zone of weathering are goethite and kaolinite, on the basis of the descriptions from Chibuluma (Winfield, 1961), Konkola (Schwellnus, 1961), and Mufulira (Brandt and others, 1961). The source of the Fe for the goethite could be earlier Cu- and Fe-rich sulfides, ferroan dolomite (Darnley, 1960, p. 148), or both. Original carbonate and anhydrite cements have been leached from the capping.

The Fe oxide described in Central African leached cappings is goethite and not hematite. The richest leached cappings overlving chalcocite enrichment blankets of southwestern United States porphyry Cu deposits contain hematite, specifically called red hematite because of its blood-red streak (Titley, 2007). Red hematite was first described as a distinct component of porphyry Cu deposit leached cappings by Anderson (1982), who noted that at temperatures of formation below the dehydration temperature of goethite, the stable oxide should be goethite, not hematite. He (p. 279) suggested that the specific oxidation mechanism of chalcocite somehow stabilizes the hematite structure at low temperatures. Hayes (unpub. data) has found red hematite in a second low-temperature environment, as the cement of upgradient ferricretes of exotic Cu deposits, wherein the hematite certainly did not form as an oxidation product of chalcocite, but did form from solutions that had a high concentration of dissolved Cu. So, it is notable if red hematite is not the Fe oxide found in leached cappings in central Africa.

Co caps, described by Gauthier and Deliens (1999) and Decrée and others (2010), where Cu has been preferentially removed leaving Co residually enriched in the product, are a special form of leached capping in central Africa, although typically the Co oxides are accompanied by minor malachite and other Cu oxides. The Co oxides and malachite are found to depths locally hundreds of meters below the surface and commonly tens of meters. Co minerals in this environment, in order of relative abundance,
are heterogenite-3R (rhombohedral CoO(OH)), kolwezite (triclinic [Cu,Co]CO_3(OH)_2), sphaerocobaltite (rhombohedral CoCO_3), cobaltoan dolomite (rhombohedral [Ca,Co]Mg[CO_3])_2, cobaltoan calcite (rhombohedral [Ca,Co]CO_3), combainite (rhombohedral Ni,Co,CO_3[OH])_2, 4H_2O, oursinite (orthorhombic [Co,Mg][UO_2]Si_2O_7·6H_2O), and juliënite (monoclinic Na_2Co(SCN)·8H_2O). Combainite and oursinite have been found only in the oxidized Cu- and Co-rich U ores of Shinkolobwe, and juliënite has been found only at the surface at Shamitumba, where its crystallization is known to be bacterially mediated and ongoing (Schoep and Billiet, 1935). Of these, heterogenite is by far the most important, occurring in black microgranular kidney-shaped nodules, in botryoidal masses with shiny black surfaces at places in concentric superposed layers in outcrop, as earthy encrustations, and as stalactites, draperies, and oolites. Implicit in those occurrences is open space in the vadose zone, which then demands that there has been major dissolution, assumedly of the host-rock dolomite, within the residually enriched Co-oxides. Kolwezite commonly occurs in some of the same rocks with heterogenite, and it forms pale tan, powdery to velvety reniform crusts and mammillarly surfaces. Sphaerocobaltite occurs mainly in the transition from carrollite-bearing or linnaeite-bearing primary sulfides to the zone of Co oxides, which, itself, is typically well down within the zone of Cu sulfides where the carrollite weathers much more readily than coexisting Cu sulfides. Sphaerocobaltite forms rose-red to purple layers directly replacing carrollite, it occurs as microscopic inclusions vividly coloring the dolomite, it occurs as drusy aggregates, and rarely it occurs as deep rose to deep-red crystals or rosettes associated with cobaltoan dolomite or calcite. Most of the cobaltoan dolomite occurs as vividly pink druses on joint faces, but some cobaltoan calcite occurs as large scalenohedrons or hexagonal prisms that are also vividly pink. Finally, some of the malachite of Democratic Republic of Congo oxide zones contains as much as 6 wt. % Co, thus forming a limited solid solution with kolwezite.

Except for central Africa, no known rock is associated with SSC deposits that would properly be termed leached capping. But, similar features probably occur in other places.

**Oxide Copper Zones**

The major minerals of the oxide Cu zone in the Central African Copperbelt are malachite, chrysocolla, cuprite, native copper, tenorite, Cu wad, and Cu vermiculite. Pseudomalachite [monoclinic Cu_3(PO_4)_2(OH)_2], libethenite, cornetite [orthorhombic Cu_3PO_4(OH)], diopside [rhombohedral CuSiO_3·H_2O], shattuckite [orthorhombic Cu_3SiO_3·OH], plancheite [orthorhombic Cu_3SiO_3·(OH)·H_2O], chalcanthite [triclinic CuSO_4·5H_2O], brochantite [monoclinic CuSO_4·5H_2O], and antlerite [orthorhombic Cu_3SO_4(OH)_2] are also important locally (Notebaart and Vink, 1972). Many additional secondary Cu minerals have been identified but only in trace quantities. Among the weathering-zone Cu minerals, there is a good deal of predictability in occurrence on the basis of the host-rock types and depth within the weathering and enrichment profiles. Some description is provided below.

Gangue minerals within the oxide Cu zones have nowhere been thoroughly described. They could provide some important information if they had been. Two minerals can be predicted in this zone. The obvious one is kaolinite, which is described in the overlying leached cappings and accompanying supergene chalcocite-enrichment profiles. Secondary Co- and Cu-bearing carbonates are present, so another mineral predicted in the oxide zone or possibly in the supergene sulfide zone of the profiles is calcite. It is fairly likely that “dedolomite” will occur. Dedolomite is known from other locations where surface recharge or other freshwaters alter dolostones.

Dedolomite is a replacement of the dolomite in dolostones occurring in a recognizable texture, namely that of medium-crystalline, mosaic-textured masses of pure calcite with intergranular residues of the trace elements that were found in the original dolomite (Braddock and Bowles, 1963, fig. 84.1–E, p. C97; Evamy, 1967; Lyle, 1977, fig. 9B, p. 431; Kenny, 1992, p. 140–141). Typical intergranular residues include Fe as goethite and Mn as pyrolusite, ramsdellite, birnessite, or cryptomelane. A study of the weathering and enrichment profiles in the Central African Copperbelt is likely to encounter dedolomite, which will then aid in identifying the base of the chalcocite enrichment.

Within the Cu-oxide zones, the available anions from the wallrocks provide an important control on the oxide minerals that form. In the Democratic Republic of Congo, where finely laminated carbonaceous dolostones are the host rocks, malachite is the predominant oxide Cu mineral. Although most of the malachite is finely disseminated, the massive botryoidal coatings of malachite that line open spaces are widely recognized among mineral collectors around the world. In more siliceous parts of the section, such as the RSC beds, chrysocolla is abundant, and plancheite, diopside, and shattuckite are also found. Chrysocolla is abundant at Nchanga, interlayered with chalcocite. These silicates are also more common in the oxide zones of arenite-hosted ores. Planchéite was identified at Nchanga and Mufulira (Notebaart and Vink, 1972). Contrasting with that, the secondary phosphates are more common in oxide zones of the Zambian shale-hosted ores. Pseudomalachite is common in the Chambishi open pit and at Mindola, although it is widespread in minor amounts in all Copperbelt oxide ores. Libethenite is widespread but not abundant and also best known from the shale-hosted ores such as Mindola (Notebaart and Vink, 1972). Cornetite is known from Bwana Mkubwa and Nchanga and was first described in specimens from L’Étoile, Democratic Republic of Congo (Cesáro, 1912).

Cuprite and native copper are widespread near the transition to sulfides, and, as above, these can be found to depths far below the present-day water table. Cuprite in Zambia and the Democratic Republic of Congo is far more abundant than cuprite in oxide zones of most SSC deposits. In mapping
of more than 1.8 km of underground workings of the Spar Lake deposit, through all of which the rocks were either fully weathered or weathered along several hundreds of fractures, cuprite was identified along only two fractures through sulfide-bearing wallrock (T. Hayes, unpub. data).

The Cu vermiculites of Central Africa’s oxide Cu zones are unique to that region. Bassett (1958) studied these and found hematite and goethite microscopically associated with them. He rejected the hypothesis that chlorite was actually interlayered in these on the basis of the differential thermal analyses and concluded that the Cu vermiculites formed from biotite in the unweathered rocks when Cu entered the exchangeable intersilicate layers. However, Jebson (Anglo-American Corp., Central Africa, Ltd., Lubumbashi, D.R.C., written commun., 1970) disagreed and concluded instead that the cupriferous micas were composed of regularly interstratified layers of biotite and cupriferous chlorite (Notebaart and Vink, 1972). Their supergene origin is fully agreed upon, though, and they have been found in abundance at Nchanga, Chingola, and Roan Antelope (now Luanshya) and in lesser quantities in the oxide zones of all the other Zambian deposits in the Copperbelt.

In other regions, oxide Cu zones constitute the outcrops of orebodies and extend, at most, 60 m into the subsurface. Malachite is typically the most common Cu mineral in the weathered rocks. In sandstone Cu deposits, the malachite commonly is disseminated throughout the host rock, but in red-bed-type deposits, it can be concentrated, with azurite, on the carbonized plant material. Azurite is subordinate but notable in oxide zones of sandstone Cu deposits. At Lisbon Valley, it forms spherical concretions up to 2 cm in diameter. In the weathered rocks in the sandstone Cu deposits of the Belt Supergroup, United States, and the Udokan, Russia, malachite is not the dominant Cu phase. The climate is temperate to cool and total precipitation is moderate in these two areas, and they have brochantite as the principal disseminated Cu-oxide mineral in outcropping ore (V. Chechetkin, oral commun., 2009; T. Hayes, unpub data). In Belt Supergroup deposits joint and fracture surfaces commonly contain malachite from hypogene carbonate-bearing zones, brochantite from hypogene carbonate-free zones, and black oxides that are probably Cu wad from either. The Cu wad forms prominent dendritic growths.

Deposits of the Belt Supergroup weather differently on the sunny southern exposures than they do on northern exposures. The Cu sulfides persist in outcrop on the northern exposures, but they rarely do on south-facing outcrops. Beer-bottle limonite, also called glassy limonite or Cu pitch, forms pseudomorphs of the finely disseminated Cu sulfides and is the most common Cu phase in south-facing outcrops. In these southern exposures, the Cu grade is typically decreased by about one-half from its unweathered values. The beer-bottle limonite forms after chalcocite, digenite, bornite, or chalcopyrite. Galena is generally leached from southern outcrops.

By comparison, malachite- or brochantite-bearing northern outcrops with some remnant sulfides commonly keep 75 to 90 percent of their unweathered Cu grade. Galena is perhaps the most resistant of all sulfides to weathering in north-facing outcrops.

In south-facing outcrops, pyrite is pseudomorphed by dense medium-to-dark brown goethite in cubes or in clots. Pyrite may persist or it may be similarly pseudomorphed in north-facing outcrops.

To a greater degree in south-facing than in north-facing outcrops, the Fe- and Mn-bearing carbonates of the primary ore in the Belt Supergroup are leached, although they leave distinctive pseudomorphs. The Fe-rich calcites may leave leached porous zones in the pattern of bedform disseminations, as well as a fairly dense intergranular network of dendritic-like chocolate-brown goethite (fig. 29). Even at depths of greater than 500 m, the chocolate-brown goethite-after-ferruginous carbonates can fill and envelope fractures. Ankerite of the central chalcocite-chlorite zone or the chalcopyrite-ankerite zone leaves discrete, randomly disseminated, tiny flecks of bright orange limonite (fig. 29). Partial weathering of the same ankerite sites may produce small amounts of calcite intergrown with the bright orange limonite. Ankerite sites from the unweathered rock do not effervesce in 1.5 percent HCl, but the same sites, where they are turned orange in the partly weathered rock, effervesce from the 1 mm limonitic fleck in the weak acid. Where fully weathered, carbonate is completely gone from the orange limonite. Similarly weathering carbonates occur in the Paradox Basin deposits.

An additional group of supergene products is present at the red-bed-type Cu deposits of Silver Reef, Utah. These are the silver halide minerals, such as chlorargyrite (cubic AgCl), bromargyrite (cubic AgBr), and embolite (AgCl,Br). These are generally spatially separated from but locally together with malachite and azurite. Chalcocite and pyrite were present below the water table, and an unidentified Ag-sulfide species was reported as well. As in other red-bed-type deposits, these ore minerals were found as replacements of carbonized plant fossils (Proctor, 1953). Although Ag grades were as high as those at Silver Reef in the sulfide-zone ores of Paoli, Okla. (Shockey and others, 1974), the Ag halides are not known in the weathered ore at Paoli, so a difference in climate is the likely explanation for the different weathering products.

Copper-Sulfide Enrichment Zone

In the Central African Copperbelt, distinguishing sulfide-enrichment zones from primary hypogene chalcocite zones is not possible without extensive ore microscopic study, except where there is accompanying native copper or covellite of supergene origin. Those two minerals almost certainly signal some degree of supergene origin for the chalcocite. Kaolinite probably also is an indicator of supergene processes. Macroscopic bleaching of the silt- and clay-size gangue assemblages indicates some uncertain mineralogical change, which could include oxidation and leaching of organic matter and changes to the phyllosilicate assemblage.
Montana and Idaho (Harrison and Grimes, 1970; Harrison and 1975 from the Burke and Revett Formations, Belt Supergroup, 110 samples collected over the period from 1965 through about is the best understood of the various data. The data are from 42a Copperbelt, and the Revett mineralizing system in the 42b country rocks have been determined for relatively few SSC 42c deposits. In fact, only the rocks of the Udokan mineralizing 42d system in Russia, the Kupferschiefer mineralizing system 42e in Germany and Poland, the Konkola deposit of the Central 42f and others (1981) identified four lognormal populations of Cu 42g concentrations (fig. 30). These were, in order of decreasing 42h Cu concentration, (1) a population of ore-mineralized rocks 42i with a geometric mean of 1,500 ppm Cu, (2) an uninterpreted 42j population of mildly enriched rocks with a geometric mean of 100 ppm Cu, (3) a background population of rocks with a 42k geometric mean of 22 ppm Cu that they interpreted had seen no effects at all from the mineralizing processes, and (4) a deple6ted population that had lost most of its Cu, with a geo- 42m metric mean of 2 ppm Cu. It was notable that over 70 per- 42n cent of the total number of samples analyzed fell within the “depleted” population. Connor and others (1981) interpreted that the relatively large volume of depleted rocks had sourced Cu now found in the small volumes of mineralized and enriched rocks. All four abundance populations were found in a 30-m section in an area of 2 km², so the entire ore process required to produce a greenbed deposit involved only a small total volume of rock. The thin, layer-form mineralized layers with interlayered depleted intervals several meters thick suggest the possibility of groundwater convection cells that are flat within the bedding with dominant lateral flow. A process where the composition of source rocks is changed by subtraction of one or more elements, and the composition of sink rocks is changed by the addition of those same elements, can be termed “bi-metasomatic.” Where there are bi-metasomatic distributions of Cu, it might be expected that there would also be bi-metasomatic distributions of major elements as well.

Geochemical Characteristics

Geochemical characteristics of SSC deposits are relatively straightforward. Description of geochemical characteristics can be categorized by the type of analyses performed: (1) major-element rock geochemistry, (2) minor and trace-element rock geochemistry, (3) mineral compositions, (4) fluid-inclusion geochemistry, (5) organic geochemistry, (6) stable-isotope geochemistry, and (7) radiogenic-isotope geochemistry.

Major-Element Rock Geochemistry

Major-element compositions of mineralized rocks and country rocks have been determined for relatively few SSC deposits. In fact, only the rocks of the Udokan mineralizing system in Russia, the Kupferschiefer mineralizing system in Germany and Poland, the Konkola deposit of the Central African Copperbelt, and the Revett mineralizing system in the United States have such data. Results from this limited amount of work show substantial major-element changes associated with ore-forming events.

The limited amount of major-element geochemical data from the Revett Formation sandstone Cu mineralizing system is the best understood of the various data. The data are from 110 samples collected over the period from 1965 through about 1975 from the Burke and Revett Formations, Belt Supergroup, Montana and Idaho (Harrison and Grimes, 1970; Harrison and others, 1986a, 1986b). The dataset included five (meta)sandstone samples that were visibly mineralized with Cu sulfides or mala- chite, including three samples from the Spar Lake sandstone Cu deposit and one from the Snowstorm deposit. The samples were otherwise regionally distributed and widely scattered.

Changes in major-element chemistry that occur related to the Revett sandstone Cu ore system are best recognized when related to the Cu geochemistry. For that reason, it is important to first understand some aspects of Cu geochemistry in Belt Supergroup SSC mineralizing systems. Some such understanding came from earlier work at Blacktail Mountain, one of the many “greenbed” Cu occurrences of Harrison (1974).

Blacktail Mountain Cu occurs at the contact of the Empire Formation with the underlying Spokane Formation. Over an area of several thousand square kilometers in the central Belt Basin, the transitional contact from the Spokane Formation to the overlying Empire Formation has at least a trace of stratabound Cu mineralization. The Empire Formation consists of about 200 m of lenticular bedded, white, calcitecemented silicate within green argillite. The Spokane is a sequence as much as 1 km thick of purple to red argillite and silicate with much lesser coarse-grained quartzite, all in medium to very thin beds. The Blacktail Mountain occurrence is within a transition zone between the Spokane and Empire Formations where green and purple beds alternate over a thickness of about 30 m, and each of the several green beds within the transition is Cu mineralized to varying degrees (thereby “green- bed” Cu occurrences). Green beds numbers 3 and 5 have grades at places over 1 percent Cu but only over thicknesses of 0.2 to 1.5 m. From more than 3,500 rock geochemical analyses for Cu at the Blacktail Mountain occurrence, Connor and others (1981) identified four lognormal populations of Cu concentrations (fig. 30). These were, in order of decreasing Cu concentration, (1) a population of ore-mineralized rocks with a geometric mean of 1,500 ppm Cu, (2) an uninterpreted population of mildly enriched rocks with a geometric mean of 100 ppm Cu, (3) a background population of rocks with a geometric mean of 22 ppm Cu that they interpreted had seen no effects at all from the mineralizing processes, and (4) a depleted population that had lost most of its Cu, with a geo- metric mean of 1.9 ppm Cu. It was notable that over 70 per- cent of the total number of samples analyzed fell within the “depleted” population. Connor and others (1981) interpreted that the relatively large volume of depleted rocks had sourced the Cu now found in the small volumes of mineralized and enriched rocks. All four abundance populations were found in a 30-m section in an area of 2 km², so the entire ore process required to produce a greenbed deposit involved only a small total volume of rock. The thin, layer-form mineralized layers with interlayered depleted intervals several meters thick suggest the possibility of groundwater convection cells that are flat within the bedding with dominant lateral flow. A process where the composition of source rocks is changed by subtraction of one or more elements, and the composition of sink rocks is changed by the addition of those same elements, can be termed “bi-metasomatic.” Where there are bi-metasomatic distributions of Cu, it might be expected that there would also be bi-metasomatic distributions of major elements as well.

Figure 29. Patterns of weathering of different carbonate minerals in the Revett Formation-hosted sandstone copper system (fig. 6), northwestern Montana and northern Idaho. On the left, ferroan and manganiferous calcite of the pyrite-calcite, galena-calcite (pictured), chalcopyrite-calcite, and bornite-calcite zones (fig. 6), where weathered completely, leave intergranular chocolate-brown goethite in a dendritic-looking pattern. Where most abundant, leached cavities, lined with the dark goethite, occur at the sites of the former calcite. On the right, ankerite of the chalcolite-chlorite and chalcopyrite-ankerite (pictured) zones leaves discrete, tiny pseudomorphs of bright orange limonite at each ankerite site.
Figure 30. Populations of copper abundance in more than 3,500 whole-rock analyses from the Blacktail Mountain greenbed copper occurrence from Connor and others (1981).

EXPLANATION

- Modeled distribution 1
  Geometric mean = 1.9 parts per million
  Geometric deviation = 1.81 parts per million

- Modeled distribution 2
  Geometric mean = 22 parts per million
  Geometric deviation = 2.0 parts per million

- Modeled distribution 3
  Geometric mean = 100 parts per million
  Geometric deviation = 2.11 parts per million

- Modeled distribution 4
  Geometric mean = 1,500 parts per million
  Geometric deviation = 2.42 parts per million

- Sum of modeled distributions 1 - 4
The dataset from Harrison samples (fig. 31) also appears to fit into the same four lognormal populations of Cu abundance that were expressed in the much larger dataset from Blacktail Mountain. From the additional descriptive information on the Harrison samples, ore mineralized and enriched rocks include no lavender (colored by pervasive hematite) samples at all, and background and depleted populations are dominated by lavender samples (former red beds) including quartzites (metasandstones), siltites, and argillites. The bulk of the hematite in lavender rocks occurs as dust rims on detrital grains. A bi-metasomatic process was suggested for the Revett Formation mineralizing system as well, and bi-metasomatic distributions of the major elements were suspected on the basis of the Cu distribution. However, drill-hole data demonstrated that flat groundwater convection cells were not likely in the Revett where sandbodies tens of meters thick are mineralized and where certain drill holes remained in enriched rock for thicknesses of more than 500 m (Hayes and Einaudi, 1986, fig. 4).

The relation between alkali metals and Cu concentrations confirmed that the metasomatic effects could be seen in Harrison’s major-element data (fig. 32). Detailed petrologic examination of more than 250 polished thin sections, mostly from the Spar Lake deposit in the Revett but including some from Harrison’s collection, lent explanation to the behavior of the alkali metals. The covariance of Cu and K is because of an anomalous abundance of K feldspar overgrowth on detrital K feldspar grains within the Cu-ore zones (figs. 6, 7, 20, 21, and 22). Albite, in contrast, is more abundant in certain carbonate-free lavender quartzites in areas far removed from the ore than within the ore zones (Hayes and Einaudi, 1986, fig. 15). In the ore itself, albite postdates the overgrowth K-feldspar and, in part, it postdates the Cu sulfides (fig. 24).

A variable was constructed from the Harrison data (Hayes, 1990) to be tested as a measure of the bi-metasomatism (fig. 33). The constructed metasomatic variable consists of the logarithm of K abundance plus the quantity, 1 + the log of Ba abundance, that sum divided by the logarithm of Na abundance. Ba, a minor element, was given sufficient weight in the variable by adding one order of magnitude to its raw value. That measures the increased barite in the ore zones (figs. 6, 7, and 23). Forming a ratio of the two elements metasomatically enriched in the rocks where Cu is enriched, K and Ba, divided by the element Na, that is enriched instead where Cu is depleted, has the effect of emphasizing contrasts in the major- and minor-element data. The graph of the results from the Ravalli Group rocks clearly shows the bi-metasomatism within the major- and minor-element geochemical data. This rock metasomatism is the inverse of changes that would have occurred in the metals-transporting solutions, and it likely is a reflection mainly of the temperature of those solutions. In ordinary deep burial diagenesis, as temperatures increase (with deeper and deeper burial), K feldspars dissolve and albite precipitates (Boles, 1978; Boles and Franks, 1979; Boles, 1982; see fig. 53). The K concentrations in the groundwater increase, the Na concentrations decrease, and the rocks are sodically altered. Ba (from the altering K-feldspar) is probably released to the groundwater as well. White (1965, p. 358–360) observed the variation in molar K/Na ratios in geothermal waters, and from this, he proposed a crude geothermometer where the measured ratio in produced geothermal water predicted the last temperature of equilibration of the water with feldspathic rock. That geothermometer was later improved upon by Fournier and Truesdell (1973) who added the effects known from Ca/Na exchange with changing temperature.

In the Revett bi-metasomatic system, Cu deposition is accompanied by local K- and Ba-metasomatism, now known to have occurred over a zone of falling temperature (shown below, in the section on Fluid-Inclusion Geochemistry; Hayes, 1990), whereas Cu depletion is accompanied by Na-metasomatism, occurring locally in the zone of greater temperature and interpreted regionally to indicate zones that reached higher temperature.

Sutton and Maynard (2005, figs. 4, 5, 6, and 9, tables 1 and 2) observed intense K-metasomatism accompanying mineralization at Konkola North. Their data have been used to construct figure 34, showing effects very similar to those observed in the Belt Supergroup. Of note, the sampled rocks from Konkola are notably different than those that provided Harrison’s data from the Belt Supergroup. The Konkola rocks were sampled from nine core holes along a fence of drill holes stretching just 8.2 km with the cores representing a stratigraphic interval of almost 4 km. The difference shows clearly in the relation between Na and Cu. The rocks sampled by Sutton and Maynard (2005) do not necessarily appear to include any from a zone of Cu mobilization (fig. 35) even though the analyses included many samples where Cu was below the detection limit of 2 ppm.

Datasets of major-element data from the Kupferschiefer system of mineralized rocks include Oszczepalski and others (2002) and Hammer and others (1990). However, the Hammer and others (1990) dataset reported only the ranges of values so it was not possible to crossplot elements sample by sample. The data from Oszczepalski and others (2002) do not seem to indicate K metasomatism in the Cu-enriched rocks like that seen in the Belt Supergroup and at Konkola. However, when the K/Na ratio is plotted in relation to Cu (fig. 36), the regression line is similar to the regression from the same plot from the Belt Supergroup (fig. 32), but the scatter among the data from the Kupferschiefer discourages much confidence in the result. Individual sample numbers from the Oszczepalski and others (2002) dataset are included on figure 36, which allows a little more understanding. Samples from the S2 sample subset (a Sierszowice area borehole fence) plot across the high-K part of the figure, and samples from the PZ19 sample subset (a Polkowice mine underground lateral section) plot across the low-K part of the figure. Trends of those subsets parallel the fitted linear regression of the total dataset with ore-grade Cu samples on the right (additionally labeled CuS), samples transitional from Cu-sulfide-rich to Rote Fäule in the middle.
Figure 31. Populations of copper abundance in 110 whole-rock analyses from scattered samples of the Revett and Burke Formations that host the sandstone copper system in the Revett Formation, northwestern Montana and northern Idaho. Analyses provided by J.E. Harrison, U.S. Geological Survey, written commun., 1985.
Figure 32. Relation between whole-rock copper and potassium/sodium in 45 samples of the Revett and Burke Formation quartzites, northwestern Montana and northern Idaho. Analyses provided by J.E. Harrison, U.S. Geological Survey, written commun., 1985. The plot is made from a subset of the 110 samples graphed in figure 31.

Figure 33. Relation between whole-rock copper and a derived “metasomatic” variable in 45 samples of the Revett and Burke Formation quartzites, northwestern Montana and northern Idaho. Analyses provided by J.E. Harrison, U.S. Geological Survey, written commun., 1985. A linear regression line has been fitted to the log-transformed elemental data. Three quartzite samples of identical grain size and primary sedimentary structure are identified with their mineral-zone names from figure 6. The divisions of “likely leached of copper” and “copper-mineralized” are from figure 31.
Geochemical Characteristics

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(Additionally labeled trans), and Rote Fäule samples on the left (RF), and each subset yields results quite comparable with the Belt Supergroup data. Viewed as individual areas, the two datasets are confidently comparable with the Belt Supergroup data. The total-alkali metals vary with the different areas of sampling. Each area is then affected by the metasomatism in like manner starting from the different total-alkali values.

The sample values on figure 36 provide another increment of understanding. On that plot, the highest K/Na values from the S2Pk57, 58, 59–61 data subset are from the transitional zone between ore-grade Cu samples and Rote Fäule samples, but the highest K/Na values from the Pz19 data subset are in the ore (sample Pz19Pk6 Cu,S). The Polowice dataset appears to be fully comparable with the Belt Supergroup data, but the Sieroszowice S2Pk 57–61 dataset is slightly different. The data of Hammer and others (1990) from near Sangerhausen, FDR, is slightly different from all three; it appears to indicate K-metasomatism of the Rote Fäule rocks, rather than K-metasomatism in the Cu ore or in the transitional rocks. It appears that SSC deposit mineral zonations are comparable with chromatographic separations of elements in which small variations in solution concentrations, temperature, or host-rock conditions can shift the relative positions of a mineral or elemental enrichment zone in one system relative to another system, or relative to a different place in the same system. Using the example of pyrite-chalcopyrite-bornite-chalcocite mineral zone boundaries, Ripley and others (1985) demonstrated that different mineral-zone arrangements could be the result of small differences in temperature or activity of any of several different components in solution including fugacity of oxygen and activity of Cu, Fe, and S. Furthermore, various mineral-zone arrangements could also be the result of varying flow rates of metals-transporting solutions, themselves controlled by permeability and head. Mineral dissolution can then produce yet more variability by causing variation in porosity, thus permeability.

Major-element chemical work on Udokan includes some old data generated using FSX methods (Bakun and others, 1966) and some more recent data for which the analytical methods were not specified (Volodin and others, 1994). The data from Volodin and others, though it is sparse, representing average analyses of quartzite groups with variable cementation, was treated the same as with the other major-element datasets (fig. 37). There is a relatively weak suggestion of potassium metasomatism accompanying ore deposition broadly similar to other systems.

Minor- and Trace-Element Geochemistry

Many minor and trace elements are found in anomalous concentrations in zonal arrangements near and in SSC deposits. The element signatures vary considerably from district to district, and they vary as well among the different deposit subtypes. The Central African Copperbelt stands apart in having high concentrations of Co in its ores, although some Co metasomatism occurred as well at Boleo, Mexico, and in the Kupferschiefer deposits. The Co enrichment is not uniform, though, even in central Africa. In Zambia, for the most part, only the ores of the Copperbelt Orebody Member

Figure 34. Relation between whole-rock copper and whole-rock potassium in 236 samples from in and near the Konkola, Zambia, reduced-facies sediment-hosted stratabound copper deposit. Data from Sutton and Maynard (2005). Compare with figure 32. A linear regression line has been fit to the log-transformed elemental data.
Figure 35. Relation between whole-rock copper and whole-rock sodium in 236 samples from in and near the Konkola, Zambia, reduced-facies sediment-hosted stratabound copper deposit. Data from Sutton and Maynard (2005). A linear regression line has been fit to the log-transformed elemental data. Compare with figures 32 and 33.

Figure 36. Relation between whole-rock copper and whole-rock potassium/sodium in 16 Kupferschiefer reduced-facies ore-system samples, Fore-Sudetic Monocline, Poland. Analyses from Oszczepalski and others (2002). A linear regression line has been fit to the log-transformed data (despite the great scatter among the points). Compare with figures 32 and 33 and see the text discussion.
(formerly the Ore Shale) produce byproduct Co (fig. 38), at average grades of about 2.74 percent Cu and 0.19 percent Co (weighted average calculated from the database of Parks and others, written commun., 2011). Attempts at explanation of the Co distribution in Zambian Copperbelt ores (Annels and Simmonds, 1984) first recognized several different possible positions in mineral zonation for Co-enriched rock, and recognized that Co was found at high grades locally even in sandstone deposits, albeit in a position outboard of bornite and chalcocite toward pyritic rock. In the sandstone deposits in Zambia, carrollite commonly co-exists only with chalcopyrite and pyrite, thus not in rock economic to mine for Cu. Annels and Simmonds (1984) eventually suggested that the source of Co most likely for Zambian ores was the mafic sills, now amphibolites, that lie upsection from the ores by at least 100 m. Such mafic sills are unavailable as source rocks at other deposits in the world like the Kupferschiefer and Spar Lake where cobalt is anomalous and in ore-stage mineral settings (fig. 18), so Annels and Simmonds conclusions are not sufficient explanation for the involvement of cobalt in SSC-mineralizing systems. In Democratic Republic of Congo as in Zambia, only the reduced-facies-type deposits produce Co, at average grades of about 3.39 percent Cu and 0.27 percent Co (weighted average calculated from the database of Parks and others, written commun., 2011). However, it is likely that the average grade of Cu and Co in the Democratic Republic of Congo reflects the effects of supergene enrichment, whereas the average grade for the Ore Shale belt deposits of Zambia does not. No other deposits known in the world possess such high Co grades. However, Co may be recoverable from the remaining ores at Boleo. The Co grade at Boleo is reported at 0.05 percent (Yeo and others, 2007). Co is recovered from the ores of Kupferschiefer system in Poland, but an average grade of Co has not been reported. The average Co grade in eight ore samples, averaging 5.67 percent Cu, reported by Oszczepalski and others (2002) was 0.005 percent (50 ppm). Seven of those samples were from the Kupferschiefer bed, typically considerably higher in Cu grade than overlying Werra carbonate ore or underlying Weissliegendes sandstone ore.

Co may be anomalous and even economic in reduced-facies-type SSC deposits but not generally in sandstone Cu deposits nor in red-bed-type Cu deposits. In a study of minor and trace elements of the Spar Lake deposit (Hayes and Nuckolls, Asarco, Inc., written commun., 1983), among 205 samples collected from all parts of the mineral zonation, only a single sample had greater than 10 ppm Co. The sample had 110 ppm Co, and it came from a rock with traces of chalcopyrite, galena, and pyrite in the galena-calcite zone (fig. 6). This relative lack of anomalous Co at Spar Lake came despite the fact that a generation of Co-anomalous pyrite is known from the galena-calcite and inboard part of pyrite-calcite zones of the deposit (fig. 18). Microprobe analyses indicate that 17 Spar Lake Co-anomalous pyrites have average concentrations of about 160 ppm Co. The Co-anomalous pyrite is not known in concentrations of as much as 0.1 v. % in any of the Spar Lake rocks. None of the rocks at Udokan were reported to have more than 100 ppm Co, and massive Udokan Cu ores were reported to have no more than 10 ppm Co (Bakun and others, 1966, p. 461).

Figure 37. Relation between whole-rock copper and whole-rock potassium in eight samples from the Udokan sandstone copper system. Analyses from Bakun and others (1966) and Volodin and others (1994). A linear regression line has been fit to the log-transformed elemental data. Compare with figure 34.
Figure 38. Stratigraphic distribution of ores in the Zambian Copperbelt from Annels and Simmonds (1984). Only the ores hosted by the Ore Shale (now called the Copperbelt Orebody Member) host economically recoverable concentrations of cobalt.
The other metal that is commonly economic in SSC deposits is Ag. The Ag occurs typically in native silver and in solid solution in chalcocite, digenite, and bornite (and perhaps others like djurleite and X-bornite). Scant traces of stromeyleirit occur in a few deposits. For 1769.7 Mt of the Kupferschiefer ores on the Fore-Sudetic Monocline in Poland, the average Ag content is 57.5 g/t (57.5 ppm) Ag at an average Cu grade of 2.02 percent (KGHM Polska Międz., S.A., 2009). Ores of the Spar Lake deposit average 46 ppm Ag at an average Cu grade of 0.63 percent. The most Ag-rich of all is Paoli, Okla., where the average of 46 unweathered trench samples was 156 ppm Ag at an average Cu grade of 0.68 percent (Thomas and others, 1991). These are among the most Ag-rich deposits, but many deposits have some recoverable Ag. White Pine has 40 ppm at an average Cu grade of 1.2 percent Cu (Kirkham and others, 1994). Udokan has 12.6 ppm Ag at an average Cu grade of 1.45 percent (Volodin and others, 1994). The belt of recently discovered deposits in Botswana typically have 15–20 ppm Ag at 1.2 to 1.5 percent Cu (Discovery Metals, Ltd., 2011), and Juramento, Argentina, has 21.8 ppm Ag at a Cu grade of 0.8 percent (Durieux and Brown, 2007). All three subtypes of SSC deposits may have anomalous Ag concentrations. The Nacimiento, N. Mex., red-bed-type deposit had 2.4 ppm Ag at 0.67 percent Cu, and Stauber, N. Mex., had 5.5 ppm Ag at 2.57% Cu. Red-bed-type deposits are not generally low in Ag–Corocoro, Bolivia, had 106 ppm Ag at 7.1% Cu from ore that was probably hand-sorted. However, Co-rich SSC deposits do not typically have economic concentrations of Ag, and Ag-rich SSC deposits do not typically have economic concentrations of Co. The exception to this is the Kupferschiefer, which has recoverable concentrations of both. Published data with analyses for Co and Ag are rare from Co-producing districts. Only the work of Darnley (1960) from the Zambian Copperbelt, Wilson (1955) from Boleo, Mexico, and Jung and Knitzschke (1976) for the Kupferschiefer in Germany are known. Darnley analyzed rocks from Mufulira, Mokombo, and Chibuluma mines in Zambia. All three of these are sandstone Cu orebodies in which most of the ore occurs in the “sericitic quartzite” class that Darnley established, and the majority of ore occurs in the enormous size of the Lubin-Sieroszowice-Polkowice-Rudna(-Glogow Giboki-Przemyslowy-Bytom Odrzanski-Retkow-Gaworzyn-Radnawice Zachody) deposit, there is certainly tens of millions of tonnes of this type of precious metals ore.

The Lubin-Sieroszowice-Polkowice-Rudna mines in Poland also produce Pb, Co, Mg, nickel (Ni), selenium (Se), and S (Pieczonka and others, 2001). Pb and Co grades were discussed above. Eight samples of Kupferschiefer ore that averaged 5.7 percent Cu also averaged 125 ppm Ni (Oszechalski and others, 2002). Jung and Knitzschke (1976) reported an average of 91 ppm Ni for Kupferschiefer bed Cu ore that averaged 1.15 percent Cu. They also reported an average of 23.7 ppm Se for the same Mansfeld and Sangerhausen deposits.

The other metal that is commonly economic in SSC deposits is Ag. The Ag occurs typically in native silver and in solid solution in chalcocite, digenite, and bornite (and perhaps others like djurleite and X-bornite). Scant traces of stromeyleirit occur in a few deposits. For 1769.7 Mt of the Kupferschiefer ores on the Fore-Sudetic Monocline in Poland, the average Ag content is 57.5 g/t (57.5 ppm) Ag at an average Cu grade of 2.02 percent (KGHM Polska Międz., S.A., 2009). Ores of the Spar Lake deposit average 46 ppm Ag at an average Cu grade of 0.63 percent. The most Ag-rich of all is Paoli, Okla., where the average of 46 unweathered trench samples was 156 ppm Ag at an average Cu grade of 0.68 percent (Thomas and others, 1991). These are among the most Ag-rich deposits, but many deposits have some recoverable Ag. White Pine has 40 ppm at an average Cu grade of 1.2 percent Cu (Kirkham and others, 1994). Udokan has 12.6 ppm Ag at an average Cu grade of 1.45 percent (Volodin and others, 1994). The belt of recently discovered deposits in Botswana typically have 15–20 ppm Ag at 1.2 to 1.5 percent Cu (Discovery Metals, Ltd., 2011), and Juramento, Argentina, has 21.8 ppm Ag at a Cu grade of 0.8 percent (Durieux and Brown, 2007). All three subtypes of SSC deposits may have anomalous Ag concentrations. The Nacimiento, N. Mex., red-bed-type deposit had 2.4 ppm Ag at 0.67 percent Cu, and Stauber, N. Mex., had 5.5 ppm Ag at 2.57% Cu. Red-bed-type deposits are not generally low in Ag–Corocoro, Bolivia, had 106 ppm Ag at 7.1% Cu from ore that was probably hand-sorted. However, Co-rich SSC deposits do not typically have economic concentrations of Ag, and Ag-rich SSC deposits do not typically have economic concentrations of Co. The exception to this is the Kupferschiefer, which has recoverable concentrations of both. Published data with analyses for Co and Ag are rare from Co-producing districts. Only the work of Darnley (1960) from the Zambian Copperbelt, Wilson (1955) from Boleo, Mexico, and Jung and Knitzschke (1976) for the Kupferschiefer in Germany are known. Darnley analyzed rocks from Mufulira, Mokombo, and Chibuluma mines in Zambia. All three of these are sandstone Cu orebodies in which most of the ore occurs in the “sericitic quartzite” class that Darnley established, and the results were only published graphically, not with individual analyses for single samples. Of these, only Chibuluma produced Co, at metal grades of 3.71 percent Cu and 0.21 percent Co (Selley and others, 2005). Among Darnley’s 30 analyzed sericitic quartzite samples from Zambia, 27 had 5,000 ppm Cu or greater. Among the same 30 samples, the Co concentrations averaged 331 ppm including 2 samples that had 5,000 ppm Cu or greater. Among the same 30 samples, Ag concentrations ranged from less than 1 to a high of 15 ppm (no mean Ag concentration was given). Wilson (1955) stated that the average Cu grade of all ore mined at Boleo from 1886 to 1947 was 4.81 percent. For 126 high-grade ore samples Wilson collected throughout the district, the average Co concentration was 1,200 ppm, considerably greater than the 500 ppm in the currently remaining ore, reported by Yeo and others (2007), above. For four high-grade ore samples, Wilson (1955) measured an average Ag concentration of just 9 ppm. Jung and Knitzschke (1976) give the grades of Kupferschiefer shale-hosted ores in the German Democratic Republic (formerly East Germany, dominated by the Mansfeld and Sangerhausen deposits) at 1.15 percent Cu, 68.5 ppm Co, and 72.7 ppm Ag (2.1 oz/t). Completing the comparison with some more Ag-rich ores, a set of 12 samples with greater than 0.4 wt. % Cu from the Revett Formation sandstone Cu system in Montana and Idaho had mean Cu of 1.13 percent, 4.5 ppm Co, and 70.3 ppm Ag (2.1 oz/t). Note the great similarity between Revett and Kupferschiefer Cu and Ag grades, whereas the Kupferschiefer shale Co grade is at least an order of magnitude greater than the Co in the Revett sandstone-type ores.

Recoverable Re occurs in the Dzhekzagan deposits, where it reaches concentrations in ore of about 100 ppm and occurs in the mineral dzhezkazganite (isometric ReMoCu2PbS6?) (Gustafson and Williams, 1981). Re is also economically recoverable in the Kupferschiefer, where it averages 21 ppm at Mansfeld and Sangerhausen (Jung and Knitzschke, 1976).

Many deposits have peripheral zones toward their pyritic sides with anomalous Pb as galena and Zn as sphalerite. Locally, these can reach ore grades, and at Boléo, Cu ore has Zn grades that are considered economic for recovery as a byproduct. The Zn grade at Boléo is reported at 0.62 percent in 277.1 million tonnes of remaining ore containing 0.70 percent Cu (Yeo and others, 2007, p. 3). Jung and Knitzschke (1976) reported an average of 0.57 percent Pb and 1.10 percent Zn in Kupferschiefer bed-hosted Cu ore from Mansfeld and Sangerhausen that had 1.15 percent Cu. Twenty-one samples from the peripheral Galena-Calcite zone quartzites of the Spar Lake sandstone Cu deposit averaged 2,176 ppm Pb. The same 21 samples averaged just 9.1 g/t Ag, whereas the Cu orebody at 6,300 ppm Cu averages 46 g/t Ag, showing the zonal segregation of Pb from Cu-Ag.

Au, Pt, and Pd are recovered from the Kupferschiefer SSC mines in Poland. These precious metals occur in the footwall of the Cu ore in a zone that parallels the mineral zonation of the deposit in a gentle cross through the bedding (Piestrznyski and others, 2002). Total tonnages and average grades of Kupferschiefer-associated Au-platinoid ores have not been released, but oxidized (Rote Fäule) and transitional near-Cu-ore samples reported by Oszechalski and others (2002) averaged 4.74 g/t (ppm) Au, 0.45 g/t Pt, and 0.29 g/t Pd. Given the enormous size of the Lubin-Sieroszowice-Polkowice-Rudna(-Glogow Giboki-Przemyslowy-Bytom Odrzanski-Retkow-Gaworzyn-Radnawice Zachody) deposit, there are certainly tens of millions of tonnes of this type of precious metals ore.

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Kupferschiefer ores. Clausthalite (isometric PbSe) and tie-mannite (isometric HgSe) were reported by Piestrzynski and others (2002) from Rote Fäule-altered rocks, but mineralogic siting of Se in non-hematitic rocks of the Kupferschiefer system has not been reported.

Among the trace elements associated with reduced-facies SSC deposits, the Kupferschiefer bed ores of Mansfeld-Sangerhausen also contain an average of 281 ppm V, 1,812 ppm As, 158 ppm molybdenum (Mo), 19 ppm cadmium (Cd), 9 ppm thallium (Tl), 8 ppm germanium (Ge), 3 ppm tellurium (Te), 15 ppm Sb, 3 ppm Hg, and 2 ppm bismuth (Bi) (Jung and Knitzschke, 1976). The V, Se, and Mo may not be enriched because of the ore system but could be, instead, a syngenetic lithogeochemical enrichment in the euxinic black shale. These elements are probably sited with the carbonaceous matter in the shale (Wedepohl, 1964). The As may or may not be ore-associated and could, instead, also be a product of the euxinic depositional environment. However, its concentration is strongly correlated with the concentration of Cu (Jung and Knitzschke, 1976), and As is anomalous in parts of other SSC systems (see below), so most likely the As is part of the ore system and inherently elevated in the black-shale environment. Each of the other elements may be ore-related. The Hg almost certainly is because Piestrzynski and others (2002) identified Hg-bearing electrum and tiemannite in the Au mineralized rock at Polkowice, and Piestrzynski and Pieczonka (1997) measured 0.20 to 0.69 wt. % Hg in Polkowice chalcocite. Finally, Hg is known to be anomalous in other SSC ore systems (see below).

In the Colorado Plateau region, United States, red-bed-subtype stratabound Cu deposits have locally economic concentrations of associated U and V. At other deposits with similar geometry and even some in the same stratigraphic host sandstones, there are U-V deposits lacking Cu or U deposits lacking both Cu and V. The Happy Jack mine in the Red Canyon district has produced about 600,000 short tons of 0.74 percent Cu, 0.25 percent U, 0.047 percent V, and 0.007 percent Cu (70 ppm), 0.38 percent U, 0.91 percent V, and 0.047 percent Cu (70 ppm), 0.38 percent U, and 0.91 percent V of a single mill shipment from Al Miesch (U.S. Geological Survey, retired, written commun., 1980). Hayes (1982) concluded that all these tabular sandstone-hosted deposits were formed by subsurface interaquifer mixing of two distinct types of groundwater, one saline and the other much less saline, and that the chemical composition of each of the two types of groundwater varied in geologic time depending on climate and rocks through which the groundwater traveled. The deposition from two groundwater types, one briny and the other less saline, was confirmed in the Henry Mountains U-V deposits by Northrop and others (1990a, 1990b) and Goldhaber and others (1990). Hot, arid climates of the Permian and pre-Permian yielded oxidizing briny groundwater that, later in the Mesozoic, at places, transported Cu up from below. This was the typical mineralizing groundwater for SSC deposits. However, in the subtropical but more humid Late Jurassic through Early Cretaceous, most of the groundwater moving upward was not Cu-bearing and yielded U-V deposits that almost completely lack Cu. In the time from Late Triassic through Early Cretaceous, relatively fresh shallow groundwater was not strongly oxidizing because of swampy organic-charged recharge areas and the shallow recharge transported the U and V to mix in the subsurface with Cu-bearing saline groundwater at some places but with Cu-poor saline groundwater at others. There appear to be red-bed-type and possibly sandstone-type Cu deposits in the Neuquén and Salta Basins, Argentina, that also have all three elements, Cu, U, and V, in deposits, all in Cretaceous sandstone host rocks, but with a variety of elemental ratios (Angelelli, 1956; Brodkorb and Brodkorb, 1984; Ferreyra and Lardone, 1990).

Trace and minor elements associated with sandstone Cu subtype deposits were defined at Spar Lake by Hayes and Nuckolls (1983). Using highly selective sampling of rock features in 205 samples from core holes from opposite sides of the mineral zonation, they defined a full suite of mobile elements concentrated across the zonation. The full suite included only Cu, Ag, Pb, Ba, Mn, and As. It probably also included Sb, but the XRF analyses had a detection limit of 20 ppm that produced no Sb detections even though the detected As anomalies were later traced to tetrahedrite (not tennantite nor enargite). The tetrahedrite occurs in a relatively narrow zone from the outboard bornite-calcite through the inboard galena-calcite zone (figs. 6 and 7). Cu and Ag are found in the ore-bearing chalcocite-chlorite and bornite-calcite zones predominantly in chalcocite, digenite, and bornite but also in traces of native silver and stromeyerite. Pb is sited in galena of the galena-calcite zone as well as within traces of tetrahedrite. Ba is in barite, Ba-rich K-feldspar overgrowths on detrital K-feldspar grains, and in trace cymrite (labeled Ba mica on fig. 6), found from the inboard galena-calcite zone through the chalcocite-chlorite zone. Mn is found in carbonates of the Spar Lake zonation. There are far more abundant carbonates within the pyrite-calcite, galena-calcite, chalcopyrite-calcite, and bornite-calcite zones of the deposit than within the chalcocite-chlorite, chalcopyrite-ankerite, and albite zones. Additionally, the calcite of the bornite-calcite is notably Mn-rich in the near vicinity of that mineral’s disappearance boundary against the chalcocite-chlorite zone (see below). Notably, though sphalerite was identified as part of the mineral zonation in the galena-calcite zone, Zn was not found in greater concentrations in that mineral zone, indicating probably that the element had taken a new mineralogical siting rather than being part of the transported elemental suite. Zn and several other elements varied only with lithology, being more abundant in argillaceous host rocks than in metasandstones. Exactly the same was true of cadmium, which was probably also sited in the sphalerite. The same was true of Fe despite the high abundance of ore-stage ferroan and manganiferous calcite found in the bornite-calcite, chalcopyrite-calcite, galena-calcite, and pyrite-calcite zones. Those elements were apparently sited in chlorite and ankerite in the chalcocite-chlorite and chalcopyrite-ankerite zones of the deposit. The study also identified a fully immobile suite of elements including zirconium, yttrium, and titanium, which all occur in heavy minerals within the (meta)sedimentary rocks.
Almost certainly also anomalous within Revett sandstone Cu deposits is Hg (fig. 39). Though the absolute concentrations are small (everywhere at 1 ppm or less), there is a strong positive correlation between Cu and Hg in the data collected from Harrison’s samples.

B metasomatism expressed in anomalous abundances of tourmaline was discussed earlier as demonstrated in the work of Darnley (1960) and Selley and others (2005). It is clear that B metasomatism accompanying ore processes in Zambia is intense. That contrasts strongly with results from Burnie and others (1972) at White Pine, the only other place where B contents were documented. They found that the Nonesuch Formation siltstones and shales at White Pine were generally low in B, averaging 47, 66, and 81 ppm in three different mineralized shale or silt beds, but they further analyzed the clay and the chlorite contents of their samples. From that they concluded that the clay component of the Nonesuch Formation is more B rich than typical marine sediments but that the overall low B values reflected a low clay content. They showed that the B and the chlorite content of the rocks increase upsectionward within the Nonesuch, which is generally the inverse of the variation in Cu content.

**Mineral Compositions**

In reduced-facies and sandstone Cu deposits, Ag is sited predominantly in chalcocite, digenite, and bornite. At the Spar Lake sandstone Cu deposits, each of those three sulfides averages greater than 0.1 wt. % Ag. Twenty chalcocites analyzed by electron microprobe had a range from 0.002 wt. % to 1.62 wt. % Ag, and a mean of 0.30 wt. % Ag. Twenty-five digenite grains contained from non-detectable concentrations to 0.625 wt. % Ag, with a mean of 0.14 wt. % Ag. Eight analyzed bornite grains contained from non-detectable concentrations to 0.61 wt. % Ag and averaged 0.22 wt. % Ag (Hayes, T.S., unpub. data, 1988). The Ag concentrations in the Cu sulfides of the Kupferschiefer are similar. Sawlowicz (1990) found a mean Ag concentration of 0.25 wt. % in 58 analyzed grains. From the corresponding Fe concentrations, these were all chalcocite-group minerals with Cu/S molar ratios from 1.78 (anilite) to 2.0 (chalocite). Piestrzynski and others (2009) reported a measured range of Ag concentrations in Cu sulfides from 0.15 to 7.11 wt. %, and they tabulated eight bornite analyses with mean Ag concentrations of 0.175 wt. %. Spar Lake and the deposits of the Kupferschiefer also contain lesser abundances of native silver and traces of stromeyerite (fig. 6).

The most notable other co-product metal of SSC deposits is Co. Most of the Co in deposits in Zambia and the Democratic Republic of Congo occurs in carrollite \([\text{Cu(Co,Ni)}_2\text{S}_4]\), which forms a solid solution with linnaeite \([\text{Co}_3\text{S}_4]\). Linnaeite has also been reported at a number of the deposits in Zambia (Notebaart and Vink, 1972). The carrollite in these deposits occurs in relatively coarse grains, in fact, typically in grains that are much larger than the disseminated pyrite or the disseminated Cu sulfides. Co also occurs in some pyrite in these deposits, and where unmetamorphosed, the Co-rich pyrite is a later stage than the earliest pyrite. At Kamoto, Democratic Republic of Congo, the Co-rich pyrite forms rims that complete idiomorphic crystals surrounding earlier generations of pyrite (Bartholomé, 1974).

**Figure 39.** Relation between copper and mercury in 61 samples from the Revett and Burke Formations in northwestern Montana and northern Idaho. Analyses provided by J.E. Harrison, U.S. Geological Survey, written commun., 1985. The plot is made from a subset of the 110 samples graphed in figure 31.
Ferroan and manganano carbonates surrounding sandstone Cu deposits may be found in 100s of meter- to kilometer-scale halos fringing the deposits, and these could be exploration indicators of the deposits. Ferroan calcites have been identified at Dzhezkazgan (Gablina and Tsepin, 1975) and Spar Lake (Hayes and Einaudi, 1986). Mn contents of the calcite and ankerite at Spar Lake also vary zonally across that deposit (fig. 22). The ferroan and particularly the manganano carbonates of the Spar Lake deposit have anomalously low δ13C attributable to organic carbon (see below). Because these carbonates were forming at the fringes of a mixing zone between a metalliferous brine and reservoired natural gas (Hayes and others, 2012), it is likely that they mark the locations of coupled redox reactions between SO4$^{2-}$ of the metal-transporting brine and carbon in the reservoired CH4. Such a reaction could be expressed as:

$$\text{Ca}^{2+} + 2\text{CuCl}_{\text{aq}} + \text{SO}_4^{2-} + \text{CH}_4 = \text{Cu}_2\text{S} + \text{CaCO}_3 + 2\text{Cl}^- + 2\text{H}^+ + \text{H}_2\text{O}$$  (1)

**Fluid-Inclusion Geochemistry**

The physical properties and compositions of hydrothermal fluids that formed SSC deposits were poorly known until recent years. Syngenetic theories of origin for this deposit type, which were popular until the 1970s or early 1980s, probably delayed the application of fluid-inclusion studies. There still remains an opportunity for major contributions from fluid-inclusion studies of SSC deposits, particularly for determining fluid-inclusion compositions in attempts to identify the source(s) of metals-transporting groundwater(s).

**Fluid-Inclusion Microthermometry**

On the basis of fluid-inclusion microthermometry studies, fluids that formed SSC deposits varied relatively narrowly in composition, from low-salinity vapor, to dilute liquid, and to high-salinity brine.

Fluid-inclusion studies in the Central African Copperbelt have shown the complexity of Cu-Co mineralization in the region. All reports noted the small size of the inclusions, generally less than 5 μm. The earliest study of fluid inclusions in the Democratic Republic of Congo was by Pirmolin (1970). He studied the fluid inclusions within dolomite “porphyroblasts” within the RSC beds, between the upper and lower orebodies, at Kamoto. The observed two-phase liquid-vapor inclusions were too small to work with by microthermometry at that time. Larger inclusions with diameters of 9 to 50 μm were observed to be of two types: randomly distributed (probably primary) and “parallel to cleavages” (probably secondary). The primary inclusions all had from one to four solid daughter minerals or trapped phases, none of which were opaque, as well as small gas bubbles commonly occupying 2 percent of the inclusion volume. More recent observations (Richards and others, 1988b; El Desouky and others, 2009) show that these large inclusions with multiple solid daughter minerals date from a second stage of Cu mineralization at Kamoto. In heating runs, five of the Pirmolin’s large, complex inclusions homogenized by disappearance of the gas phase at between 140 and 150 °C, with, as well, dissolution of some solid phases at varying temperatures. Some parallelepiped-shaped solids dissolved at temperatures as low as 55–60 °C, additional parallelepiped solids dissolved at temperatures between 120 and 145 °C, and spheroidal solids dissolved, finally, between 200 and 240 °C, by which temperature all five inclusions that had earlier lost their gas bubbles between 135 and 150 °C had decrepitated. All seven of the inclusions where the gas bubble did not disappear were noted to have undergone “slight opening of the cleavages around 130 °C.” That is, they leaked during the heating run.

Pirmolin (1970) also applied XRF studies that allowed x-ray beam penetrations of 25 to 30 μm into some inclusions. This remains, even to this day, the only direct measurement of inorganic compositions of the fluids involved in forming SSC deposits. Pirmolin identified Ca, K, Si, Cl, Cu, Fe, Pb, Mn, Sn, Co, Ti, and S in the inclusion fluids but could make no attempt at quantification. He attributed Cu, Co, and Pb to the mineralizing process and Ca, Fe, Mn, and Si were attributed to interaction with the siliceous dolomite host rocks. He concluded that the K, Cl, S, and some Na formed the solid daughter minerals, and from that he deduced that the solids included halite, sylvite, and anhydrite. With some further understanding of the petrology of the Democratic Republic of Congo host rocks, it is now known that the dolomite whose inclusions Pirmolin (1970) studied was an ore-stage or later pseudomorph of original anhydrite. It is that fact that validates the coarse crystalline quartz and dolomite in Zambian and Congolese ores as a host for ore-stage fluid inclusions and that validates fluid inclusion microthermometry results from Pirmolin (1970), Annels (1989), Dewaele and others (2006), and El Desouky and others (2009).

Returning to the study of Pirmolin (1970), and with the assumption that anhydrite was replaced in depositing the dolomite that hosted the inclusions, a highly saline system of NaCl, KCl, and water, with high concentrations of Cu and Co, is accepted as the mineralizing fluid. The three component NaCl-KCl-H2O system was illustrated by Roedder (1984, p. 245). With the temperatures and solid behaviors above, the dissolution of the first solid, definitely KCl, at less than 60 °C allows only fluid compositions with 15–20 mol. % KCl for the three inclusions that had two parallelepiped-shaped solids. With dissolution of the second solid at 120 to 130 °C, the NaCl molarity is also closely limited and must be between 28 and 30 mol. % NaCl. The fluid is, therefore, saline—15 to 20 mol. % KCl and 28 to 30 mol. % NaCl, and thus 43 to 50 eq. mol. % NaCl total. The behavior in heating runs indicates homogenization temperatures of 135 to 150 °C probably are minimum temperatures for the deposition of the dolomite at a time probably many millions of years later than main-stage Cu-Co-sulfide deposition.
Richards and others (1988b) studied veinlet-controlled mineralization accompanied by montmorillonitic vein envelopes and quartz-hematite-rutile-barite-calcite vein centers cutting footwall elastic rocks at Musoshi, Democratic Republic of Congo. Fluid inclusions in the vein quartz were studied microthermometrically. The inclusions had multiple daughters certainly including halite and sylvite and probably including anhydrite, hematite, and other solids in certain inclusions. These inclusions homogenized with a systematically different sequence than those observed by Pirmolin (1970), by dissolution of first, sylvite, then the gas bubbles, and finally, halite, at temperatures between 247 and 300 °C for distal-veinlet zones and between 391 and 413 °C for influx-zone veins. Minimum salinities were 39 wt. % NaCl plus 15 wt. % KCl, and thus 50.75 eq. wt. % NaCl total. Finally, Richards and others (1988b) Pb-Pb dated the rutile of the veinlets at 514 ± 2 Ma and thus interpreted the fluid as a late, post-ore overprint from the Lufilian Orogeny relative to the main disseminated ore-stage.

Annels (1989) studied fluid inclusions in coarse crystalline nodular quartz, probably pseudomorphic after anhydrite, and in vein quartz at Chambishi and Chambishi East. He measured 138 to 180 °C homogenization temperatures and determined salinities of 10.5 to 16 eq. wt. % NaCl for pyrite- and carrollite-associated inclusions at Chambishi, 137 to 170 °C homogenization temperatures and 16 to 21 eq. wt. % NaCl salinities in chalcopyrite-associated quartz, and 118 to 150 °C homogenization temperatures and 16 to 20 eq. wt. % NaCl salinities in bornite-associated quartz. These results were discussed by Annels (1989), who noted the shift in paragenetic time from high temperature and low salinity to lower temperature at greater salinity. In these fluids that all date from the main stage of disseminated sulfides, with salinities varying from 10.5 to 21 eq. wt. % NaCl, mixing of two solutions seems likely, although it was not discussed. Late-stage veins with homogenization temperatures from 297 °C to 425 °C with the same salinities were interpreted by Annels (1989) to be related to late intrusion of mafic sills. This work was the first to deal with inclusions from the early, main-stage disseminated mineralization for the Central African deposits. Its homogenization temperatures are generally lower than temperatures from the overprinted veinlet stage, as are its salinities.

McGowan and others (2006) at the Nchanga deposit in Zambia examined inclusions in quartz veins from the Mines Sequence but did not report on the smaller inclusions in the stratabound ores. They found homogenization temperatures ranging from 140 and 180 °C and salinities between 31 and 38 eq. wt. % NaCl. Compared with other studies, it is clear that McGowan and others (2006) recorded results only from second-stage mineralization probably related to the Lufilian Orogeny (by comparison with Richards and others, 1988b; El Desouky and others, 2009).

Dewaele and others (2006) studied small inclusions in coarse-crystalline quartz, which probably replaced anhydrite, from stratabound ores from the Musonoi and Kamoto deposits in the Copperbelt in the Democratic Republic of Congo. Homogenization temperatures were between 127 and 192 °C at Kamoto and 105 and 188 °C at Musonoi, with salinities between 9.4 and 15.3 equivalent (eq.) wt. % NaCl at Kamoto and 8.4 and 18.4 eq. wt. % NaCl at Musonoi. These, clearly, were all related to the pre-orogenic main stage of sulfide mineralization, and, as with Annels’ (1989) results, mixing of two solutions is suggested by the large ranges in temperatures and salinities.

El Desouky and others (2009) conducted petrographic and microthermometric studies of borehole samples from Kamoto and Luwiswishi deposits and found evidence for two distinct mineralizing phases. In the first, Cu and Co minerals are found in layers and nodules of dolomite accompanied by quartz (both minerals probably pseudomorphic after anhydrite). Inclusion homogenization temperatures ranged from 117 to 220 °C. Salinities were calculated from 11.3 to 20.9 eq. wt. % NaCl (fig. 40). In the second phase, ore minerals are found in stratabound veins and breccias, and fluid inclusions have homogenization temperatures between 270 and 385 °C and salinities from 35 to 45.5 eq. wt. % NaCl. El Desouky and others (2009) concluded that the early main phase of sediment-hosted ore was enriched by a synorogenic second phase at the later time of the Lufilian Orogeny.

El Desouky and others (2008b) studied fluid inclusions in the Mwitapile sandstone Cu deposit in the Lufilian Foreland in Democratic Republic of Congo and compared the results with earlier studies at Lufukwe sandstone Cu deposit of the same region that had been studied by El Desouky and others (2008a). At Mwitapile, arkosic, calcareous sandstone shows a first generation of quartz overgrowths. Dissolution of the calcite produced cavities occupied by pyrite, chalcopyrite, bornite, chalcocite, and a second generation of quartz.

Microthermometry on inclusions in the early overgrowth quartz yielded homogenization temperatures between 111 and 182 °C and salinities between 22 and 25.5 eq. wt. % CaCl2. These results were comparable with the results from Lufukwe (El Desouky and others, 2008a). Inclusions in the second-phase quartz associated with the main ore stage had temperatures between 120 and 280 °C and salinities between 2.4 and 19.8 eq. wt. % NaCl. The higher temperatures were associated with higher salinities. Mineralization was interpreted to be related to mixing of the high temperature, high salinity, metals-transporting fluid with the lower temperature, reduced, first-phase fluids. It is not clear whether or not the main stage at Mwitapile correlates with the main stage from deposits at Kamoto, Musonoi, and Luwiswishi (Dewaele and others, 2006; El Desouky and others, 2009).

In summary, ore deposition in the Central African Copperbelt SSC deposits took place in two separate stages. The first stage produced only small liquid-vapor fluid inclusions, with no solid daughter minerals; the second stage produced large inclusions, with multiple daughter minerals including halite, sylvite, and other possible solids. At the early stage, saline fluids, typically in excess of 15 eq. wt. % NaCl and greater than 20 eq. wt. % NaCl in a few locations,
Figure 40. Summary of fluid-inclusion microthermometry results from sandstone copper and reduced-facies sediment-hosted stratabound copper deposits of the Democratic Republic of Congo. Histograms compile the results from primary inclusions, only, from Dewaele and others (2006) from the Kamoto and Musonoi reduced-facies deposits, from El Desouky and others (2008a) from the Lufukwe sandstone copper prospect, from El Desouky and others (2008b) from the Mwitapile sandstone copper prospect, and from El Desouky and others (2009) from the Kamoto and Luiswishi reduced-facies sediment-hosted stratabound copper deposits. Graphs A and B are homogenization temperatures and salinities from the early disseminated stage, and graphs C and D are from the later veinlet stage. Assignment of certain inclusions from Mwitapile to the late stage by El Desouky and others (2008b) (their second authigenic quartz phase) may be erroneous. Results from those inclusions are very similar in homogenization temperatures and salinities to early-stage inclusions from the other four deposits. Rather than occurring in veinlets, this authigenic quartz occurs as overgrowths on detrital quartz grains that are encased within copper-sulfide clots. The results from quartz overgrowths at Mwitapile are demarcated by the arrowed brackets in C and D.
transported Cu at temperatures of about 130 to greater than 220 °C. Ore deposition occurred partly by mixing with lower temperature and less saline groundwater, probably just the pre-ore groundwater with a salinity less than 10 eq. wt. % NaCl. The result was large ranges in homogenization temperatures and salinity. Later, probably during or after the Luflilian Orogeny, a second stage of Cu mineralization took place from fluids with temperatures greater than 200 to 415 °C and with salinities of 35 to 55 eq. wt. % NaCl. The second-stage Cu mineralization was dominantly vein-controlled, whereas the early Cu mineralization was disseminated through host rocks. Sources of the fluids are not known for either stage, but particularly given the extraordinary salinities of the second-stage fluids, dissolved evaporites, likely even potash salts, are probably significant. Descriptions to date (Selley and others, 2005, p. 965–966; Dewaele and others, 2006, p. 455) suggest that the second-stage mineralization did not contribute large parts of the total Cu mineralization except at a few places, such as Kansanshi, Lubembe, and Frontier (Doug Jack, oral commun., 2010).

Fluid-inclusion studies were important in understanding the origin of the Spar Lake sandstone Cu deposit. Fluid-inclusion studies (Hayes, 1990) were aimed partly at understanding the zonal pattern of ore and gangue cements. Only primary inclusions from within quartz overgrowths on detrital quartz grains or from pore-filling calcite cements were studied by microthermometry, and these were all small, between 2 and 7 μm in greatest dimension, with most less than 5 μm. All were 2-phase liquid-vapor inclusions. Six inclusion homogenization temperatures from the galena-calcite zone were from 72 to 126 °C, four from the chalcopyrite-calcite zone were from 150 to 164 °C, seven from the bornite-calcite zone were from 130 to 172 °C, and eight from the chalcocite-chlorite zone were from 143 to 184 °C (fig. 41). Those results demonstrated increasing temperatures from pyrite-calcite toward chalcoite-chlorite, corroborating earlier S-isotope geothermometry (Hayes and others, 1989). Two homogenizations from chalcopyrite-ankerite zone rocks were 145 °C and 169.5 °C, and those, in being equal to or lower than the temperatures in the adjacent chalcocite-chlorite zone, demonstrated that the chalcopyrite-ankerite zone was not a feeder zone to the deposit. Salinity calculations ranged widely and correlated poorly with homogenization temperatures, although freezing runs generally indicated lower salinities in the peripheral chalcopyrite- and galena-bearing zones. The salinities varied from 5 to 22.5 eq. wt. % NaCl. Those results were interpreted to indicate mixing of a high temperature, high salinity, metals-transporting fluid with cooler, less saline pre-ore fluids.

**Chemical Compositions of Mineralizing Fluids**

Exclusive of Pirmolin (1970) described above, the only work that has determined compositions of inclusion fluids of any SSC deposit was a study of the Spar Lake deposit (Hayes and others, 2012). In that study, quadrupole mass spectrometry was used to determine fluid-inclusion volatile compositions. Inclusion fluids were released from samples of separated carbonates, separated Cu sulfides, or crushed whole rocks by thermal decrepitation at 180 °C for periods of as much as one hour in order to accumulate enough gas from the small and sparse inclusions in ore-stage authigenic cements. The low temperature was chosen to avoid decrepitating any higher-temperature inclusions trapped in detrital grains at times earlier in those grains geologic histories. The results demonstrated that mineralization at Spar Lake took place where a metals-transporting brine at greater than 200 °C mixed and reacted with sour gas (natural gas containing H2S) reservoired in sandstones. The metals-transporting brine was an aqueous fluid lacking any important amounts of dissolved CO2, CH4, or H2S (fig. 42). Systematic arrangement of fluid-inclusion volatiles across the ore deposit was reflected in H2S and other hydrocarbon gases, as well as in CH4/CO2 ratios. The metals-transporting brine entered the gas reservoir from below at a temperature of at least 200 °C and reacted with cooler natural gas. The brine was apparently oxidizing, and the ore was deposited as it reacted to lower fO2 conditions with the H2S and CH4 in the reservoired natural gas (fig. 43). Neither the metals-transporting brine nor the natural gas give any indication of involvement of magmatic or subcrustal fluids; Spar Lake fluid-inclusion volatiles also have low helium R/Rα values ranging from 0.236 to 0.995, also indicating no involvement of mantle or magmatic fluids.

**Organic Geochemistry**

The earliest studies of organic geochemistry at a SSC deposit were at White Pine (Eglinton and others, 1964; Meinschein and others, 1964; Barghoorn and others, 1965). Their purpose was limited, however, to demonstrating and analyzing the forms of early life in the Mesoproterozoic Nonesuch Shale host rocks. The studies, nonetheless, demonstrated the presence of in situ-generated oil; of macerals of plant, possibly algal, remains; of an odd-carbon-preference distribution of alkane compounds in organic matter extracts; of pristine and phytane typical of those produced from diagenesis of plant materials; and of vanadyl porphyrins in extracts from the rock organic matter probably derived from chlorophyll (Barghoorn and others, 1965).

More recent organic geochemical studies have addressed the Nonesuch Shale of the White Pine deposit and the Kupferschiefer, including a number of studies designed to glean understanding of the mineralizing process from the organics and to determine the effects of the mineralizing process on the organics (Hieshima and Mauk, 1990; Ho and Mauk, 1996; Pratt and others, 1991; Püttmann and others, 1987, 1989; Sun and Püttmann, 1997; Bechtel and others, 2002). Work at White Pine found that Nonesuch Shale Prisam/n-C17, and Phyr/n-C18 ratios are lower within ore than outside of it, suggesting thermal maturation of the lower Nonesuch during mineralization. Yet the temperatures experienced in the ore were insufficient to destroy the pristine and phytane. In the Kupferschiefer, in the bottom section of mineralized
Sediment-Hosted Stratabound Copper Deposit Model

Figure 41. Fluid-inclusion homogenization temperatures from the Spar Lake sandstone copper deposit from Hayes (1990). Gray filled square symbols are for inclusions in quartz overgrowths on detrital quartz substrates (figs. 20, 21, 22, and 23). Blue square symbols are for inclusions from ore-stage manganiferous or ferroan calcite cements (figs. 20, 21, and 22). There is a gradient in the homogenization temperatures of as much as 100 °C from the galena-bearing fringing zone to the core of the chalcocite-bearing ore zone. That gradient occurs over typically about 300 meters laterally and 50 meters vertically. Nonetheless, the ore-depositing system was clearly a geothermal anomaly. Using this and salinity data, Hayes (1990) interpreted that hot, saline metals-transporting groundwater mixed with cooler, dilute formation waters in depositing ore.

Figure 41. Fluid-inclusion homogenization temperatures from the Spar Lake sandstone copper deposit from Hayes (1990). Gray filled square symbols are for inclusions in quartz overgrowths on detrital quartz substrates (figs. 20, 21, 22, and 23). Blue square symbols are for inclusions from ore-stage manganiferous or ferroan calcite cements (figs. 20, 21, and 22). There is a gradient in the homogenization temperatures of as much as 100 °C from the galena-bearing fringing zone to the core of the chalcocite-bearing ore zone. That gradient occurs over typically about 300 meters laterally and 50 meters vertically. Nonetheless, the ore-depositing system was clearly a geothermal anomaly. Using this and salinity data, Hayes (1990) interpreted that hot, saline metals-transporting groundwater mixed with cooler, dilute formation waters in depositing ore.
shale, saturated hydrocarbons are diminished and heteroaromatic systems containing oxygen and S are substantially enriched, compared with the less-mineralized shale just tens of centimeters upsection. The ratio of phenanthrene/sum of methylphenanthrenes is greatest in the richly mineralized basal part of the shale, and there are increased spectral fluorescence intensities of extracts relative to samples higher in the shale. All these measurements indicate a higher oxidation state in the basal section, interpreted to be caused by maturation and oxidation of organics in the shale by an ascending, oxidizing, metalliferous brine. Probably most important of all were studies of the Kupferschiefer that combined organic geochemical studies with studies of stable isotope systematics (discussed below).

**Stable-Isotope Geochemistry**

There have been numerous studies of S isotopes in sulfides of SSC deposits, and there have been several studies of C isotopes in carbonates of SSC deposits. Measurements of C-isotopes in organic materials in the Kupferschiefer and at White Pine were also conducted in recent years. A few studies also document values of O-isotopes of the carbonate minerals. The first study to document S and O isotopes in ore-stage sulfates was Hayes and others (2012).

**Sulfur Isotopes**

All reduced-facies-subtype deposits that have been studied have large ranges of S-isotope values among their sulfides, and they have some sulfides that have $\delta^{34}S$ values much lower than $-10$ per mil (‰) Dechow and Jensen, 1965; Marowsky, 1969; Burnie and others, 1972; Sawlowicz, 1989; Branam and Ripley, 1990. The White Pine deposit is fairly typical of this (fig. 44), with a total range in $\delta^{34}S$ values of 70‰ and values as low as $-29‰$. This characteristic is evidence that sulfide now found in Cu sulfides was bacteriogenically fractionated from seawater sulfate. The Kupferschiefer has even lower values of $\delta^{34}S$ because the Kupferschiefer is in Permian host rocks, from a time when the $\delta^{34}S$ of seawater sulfate was at its lowest known value in geologic history (Claypool and others, 1980). Kupferschiefer $\delta^{34}S$ values are as low as $-44‰$.

Interpretations of the great spread in pyrite and Cu sulfide $\delta^{34}S$ values and some light values have varied. Marowsky (1969) interpreted that both types of values indicated “syngenetic” sulfides, although actually referring to early diagenetic sulfide formation just below the seawater-sediment interface. Burnie and others (1972) recognized that the distribution indicated either earliest diagenetic mineralization or an earliest diagenetic distribution of pyrite values followed by replacement by
Sediment-Hosted Stratabound Copper Deposit Model

Later workers have been far more definite in preferring replacement of pyrite without appreciable redistribution of S isotopes. Hoy and Ohmoto (1989) argued that Cu sulfides incorporating S with heavy δ\(^{34}\)S were added at ore-stage to White Pine and Kamoto (fig. 45). The δ\(^{34}\)S of the added sulfide at ore stage was about +1.5‰ at White Pine, but it was as heavy as +9‰ at Kamoto (Hoy and Ohmoto, 1989). The timing of that sulfide addition was constrained only as being sometime after the early diagenetic establishment of a wide distribution of values in pyrite.

The timing was better constrained in recent integrative studies of the Kupferschiefer at Sangerhausen of S isotopes of Cu sulfides, C isotopes of pre-ore and ore-stage carbonates, organic geochemistry, and C isotopes of organic matter (Bechtel and others, 2001). Those studies will be described below in the section on Integrated Isotopic Studies.

S-isotope values at sandstone Cu deposits differ dramatically from those at reduced-facies deposits in that they have no notably negative δ\(^{34}\)S values, although they share the characteristic of having broad distributions of values. Spar Lake sulfide values spread from +2.1 to +29.1‰ (fig. 46). From S-isotopic studies at Spar Lake, there was an approach to thermal S-isotope equilibrium between pyrite, galena, and chalcopyrite of the distal zones. Tie lines connecting those three minerals (fig. 46) generally parallel one another. However, S-isotope equilibrium between the base-metal sulfides and barite was not attained or approached. Tie lines connecting the base-metal sulfides with barite generally do not parallel one another (fig. 46) and many cross one another.

About one-half of the total sulfides at Spar Lake are in large seepage pillars, a type of water-escape structure, locally called ore rods, and all the ore-rods sulfides have δ\(^{34}\)S values from +3.9 to +10.7‰ (fig. 47). The δ\(^{34}\)S values of ore-stage barite vary from +18.8 to +28.9 (fig. 46), and together with the sulfide S-isotopes, the overall distribution is remarkably similar to the distribution of positive S-isotope values of sulfide minerals and petroleum in the Amu-Dar’ya petroleum-producing basin in Turkmenistan and Uzbekistan (Belenitskaya and others, 1981) (fig. 47). The Spar Lake deposit formed where Cu-transporting brines mixed and reacted with reservoired sour gas. The
The full history of the S found in Spar Lake sulfides is interpreted as follows and can be followed on the diagram of the similar history of the Amu-Dar’ya basin (fig. 47). The bulk of the Spar Lake S was derived by bacterial sulfate reduction of seawater sulfate having δ\(^{34}\)S slightly higher than +20‰, followed by sequestering of the sulfide in organic matter of down-section source rocks in the Prichard Formation. Maturation of the organic matter later produced petroleum with relatively homogeneous sulfide with δ\(^{34}\)S of about +6‰. Thermal cracking then produced H\(_2\)S with relatively homogeneous sulfide also having δ\(^{34}\)S closely distributed about +6‰ in sour gas. There was an additional contribution of sulfide from later TSR operating on formation water with δ\(^{34}\)S of about +20 to +29‰. That TSR probably occurred during the entire time interval from initial trapping of petroleum through ore deposition. Additional sulfide was thermochemically reduced at ore-stage from sulfate (SO\(_4\)) in the metals-transporting brine in redox reactions like equation 1 with the reservoired CH\(_4\). The SO\(_4\) in the metals-transporting brine was also deposited in ore-stage barite having that distribution of δ\(^{34}\)S values of about +19 to +29‰. Knowledge of the fate of S and its isotopes in petroleum was documented by Thode and Monster (1965, 1970) and Krouse (1977).

Thus, the overall difference between S isotopes found in reduced-facies-subtype deposits and those found in sandstone Cu deposits is explained by the S being resident in different reductants. The deposit subtypes share the addition of ore-stage, isotopically heavy sulfide from the thermochemical reduction of sulfate (SO\(_4\)) from the metals-transporting brines as will be shown below in the sections on Carbon Isotopes and Integrated Isotopic Studies.

### Carbon Isotopes

Markowsky (1969) analyzed 105 Kupfershiefer bed samples for δ\(^{13}\)C in dolomite or calcite and 103 samples for δ\(^{13}\)C in organic matter. The values in organic matter were typical of bituminous shale, with a unimodal distribution centered surrounding −27.5‰ relative to PDB and with a narrow range from only −25 to −30‰. The distributions of calcite and dolomite δ\(^{13}\)C values were also typical. The calcite had a mode at about +2.5‰, with a fairly large range from −5.5 to +4.5‰. The range of values in dolomite was also fairly large, from +6.5 to −2.5‰. Many of Marowsky’s (1969) samples came from areas of barren pyritic Kupferschiefer from northern Germany. He noted the occurrence of light carbonates of biogenic origin above zones of sulfide accumulation but interpreted that it indicated bacterial involvement in syngenetic precipitation of the sulfides.

Earliest reported C-isotope studies in Zambia were from the work of Annels (1989). Only nine samples were analyzed. Stromatolitic reef dolomites were relatively close to typical
Neoproterozoic marine values of $-1$ to $+10\%_{\text{oo}}$ $\delta^{13}\text{C}_{\text{org}}$ and $-4$ to $-10\%_{\text{oo}}$ $\delta^{18}\text{O}_{\text{org}}$ (Halverson and others, 2005; Santos and others, 2000; Chu and others, 2007; Corsetti and others, 2007; Giddings and Wallace, 2009). Two dolomite nodules associated with Cu- and Co-bearing sulfides had $\delta^{13}\text{C}$ of $-14.8$ and $-25.2\%_{\text{oo}}$. Two calcite nodules were $-14.8$ and $-19.4\%_{\text{oo}}$. These are extremely depleted values that indicate a large proportion of organic carbon in the minerals. Sweeney and Binda (1989) reported dolomite lenticles with $\delta^{13}\text{C}$ values of $-8.5$ to $-21\%_{\text{oo}}$ from the Ore Shale at Konkola, layered dolomite from unit C of the Ore Shale at $-12.5\%_{\text{oo}}$, and Ore Shale calcite at $-17\%_{\text{oo}}$. They interpreted these depleted values as related to fresh groundwater, and the heavier values in the footwall clastics were interpreted as related to marine transgression. Values from carbonates in the footwall clastic rocks varied from $-4.5$ to $-9.5\%_{\text{oo}}$. Sweeney and Binda (1989) stated these $\delta^{13}\text{C}$ values indicated that “organic matter was oxidized during carbonate formation.” The Ore Shale values are far too low to have been due merely to continental groundwaters. These low $\delta^{13}\text{C}$ values of ore-related carbonates are indicators of the oxidation of organic carbon accompanying sulfide mineralization. Hitzman and others (2005) suggested that low $\delta^{13}\text{C}$ in carbonates within specific rock volumes would serve as proxy for the presence of mobile reductants (that is, petroleum). The carbonates also had $\delta^{18}\text{O}$ values that were low compared with contemporary marine carbonate values and that could be interpreted to be because of either involvement of fresher, less saline water in their precipitation or to relatively high temperatures of precipitation. Obviously, given Annels’ (1989) fluid-inclusion homogenization temperatures from some of these same samples, the low values were because of elevated hydrothermal temperature.

Sandstone Cu deposits, such as Spar Lake, also reflect the oxidation of organic carbon-bearing reductants in ore-stage products. The isotopically lightest carbon of the entire Belt Supergroup measured to date is found in ore-stage calcite and ankerite cements from the distal fringes of the Spar Lake deposit (fig. 48).

### Integrated Isotopic Studies

The most valuable of the studies of stable isotopes are those that have integrated studies of both S and C isotopes or integrated isotopic studies with other geochemical methods.

Bechtel and others (2001), studying samples from Sangerhausen, Germany, integrated S-isotope, C-isotope, organic geochemical, and elemental geochemical data. They showed that total $C_{\text{org}}$, C/S ratios, and hydrogen indices of the organic materials were facies related and did not correlate with Cu-enriched beds. Values of these parameters indicate where pre-ore pyrite was deposited by bacterial sulfate reduction from seawater. Then, with Fe-$C_{\text{org}}$, sulfur isotope, and ore microscopy, they showed that in parts of the Kupferschiefer bed with grades of 0.1 to 6 percent Cu, the pyrite was replaced by Cu- and Fe-bearing sulfides, and the isotopic signature of the replaced pyrite was inherited. With higher grade (greater than 8 percent Cu) mineralization, in the lower part of the Kupferschiefer bed, there was (1) deposition of sparry calcite with $\delta^{13}\text{C}$ values substantially lowered relative to the typical marine values found in the top of the shale, (2) reciprocal changes of values of $\delta^{13}\text{C}$ in kerogen and bitumen to heavier isotopic values remaining in those organic residues, and (3) an addition of Cu sulfides with heavier $\delta^{34}\text{S}$ deposited by thermochemical reduction of ore solution $\text{SO}_4$ coupled with oxidation of the organic carbon.

In studying the Polish Kupferschiefer, Bechtel and others (2002) combined C-isotope, Rock-Eval pyrolysis, and elemental geochemistry data. Just as in Germany, as the $\delta^{13}\text{C}$ of epigenetic calcite shifts to lower values with higher Cu grades, the $\delta^{13}\text{C}$ of residual kerogen and bitumen shift to heavier isotopic values (fig. 49). The hydrogen indices of the same samples shift to lower, more matured values, and the pyrolysis $T_{\text{max}}$ values increase. Over the same trends, the oxygen indices increase, seemingly altering the organic matter to entirely different kerogen types and crossing sharply across the ordinary burial maturation pathways for organic matter (fig. 50).

At Spar Lake (Hayes and others, 2012), the isotopically light distal-fringe carbonates are coupled with the sulfide that was thermochemically reduced at ore-stage from $\text{SO}_4$ in the metals-transporting brine. The sulfide from ore-stage thermochemical reduction is identifiable where samples have heavy $\delta^{34}\text{S}$ values in Cu- or Pb-bearing sulfides and light $\delta^{13}\text{C}$ values in ore-stage calcite from the distal fringes of the deposit (fig. 51).
Figure 46. Histograms of $\delta^{34}S$ values of sulfides and sulfates across the mineral zonation at the Spar Lake sandstone copper deposit, from Hayes and others (2012).

**EXPLANATION**
- **One sulfide sulfur isotope value**
- **Ore rods sulfide sulfur isotope value**
- **Barite sulfate sulfur isotope value**

Modified from T.S. Hayes and others, 2012
Isotopic Dating and Radiogenic Isotope Geochemistry

Isotopic dating of SSC deposits has suffered from the lack of easily dateable minerals within the deposits. Thus, most age constraints have been established by dating interlayered volcanic rocks, tuffs, and the basement rocks beneath the sedimentary-host sections. A few direct dating attempts have been made with limited success.

Earliest dating attempts examined Pb-isotopes, particularly because galena was a part of the mineralization. Pb-isotope studies have been used in attempts to date SSC mineralization and to determine sources of the metals for these deposits. Wedepohl and others (1978) studied whole-rock (?) Pb-isotopes from Kupferschiefer samples scattered over a large area of central and northern Germany, as well as individual samples from Holland, England, and Poland. The accepted depositional age of the Kupferschiefer is from 260.4 to 258 Ma (Menning and others, 2006; Słowakiewicz and others, 2009), which sets a maximum age for mineralization. Eight of Wedepohl’s samples were from Germany; two were ores of Cu, five were from Pb-rich zones, and one sample was from the pyrite zone. The samples were fitted to a line on the cross plot of $^{206}\text{Pb}/^{204}\text{Pb}$ and $^{207}\text{Pb}/^{204}\text{Pb}$ that intersected the Stacey and Kramers 2-stage terrestrial Pb evolution curve at 250 Ma and 1750 Ma. The interpretation was that these defined a secondary isochron. The former age is obviously close to the depositional age of the Kupferschiefer, and Wedepohl...
Figure 48. Relation between $\delta^{13}C$ and $\delta^{18}O$ of carbonates related to the Spar Lake sandstone copper deposit, from Hayes and others (2012). Isotope values for carbonates about 50 kilometers northeast of Spar Lake are within the lowest part of the range of such values for the Helena Formation and for Mesoproterozoic carbonates elsewhere in the world. Values for the Helena Formation overlying the ore at Spar Lake are lower in $\delta^{13}C$ and $\delta^{18}O$, interpreted to indicate that those carbonates derive partly from the carbon in thermally cracked hydrocarbons from a Revett Formation-hosted natural-gas reservoir, below. Values of calcite and ankerite cements in the Revett ores and distal galena and pyrite zones are lighter in $\delta^{13}C$, particularly, including many values less than −12 per mil. These are interpreted to have employed carbon from ore-stage oxidation of methane of the pre-ore natural gas reservoir in redox reactions coupled with the reduction of ore-fluid sulfate to sulfide in precipitating the copper- and base-metal sulfides.
Concentrations of copper (Cu), zinc (Zn), lead (Pb), vanadium (V), molybdenum (Mo), nickel (Ni), cobalt (Co), uranium (U), organic carbon (C_{org}), and carbonate carbon (C_{carb}), Rock Eval pyrolysis temperature (T_{max}) and hydrogen index (HI), δ^{13}C of organic carbon, δ^{13}C of calcite, and δ^{18}O of calcite for a mineralized Kupferschiefer section from the Sieroszowice mine, Poland, from Bechtel and others (2002). Stratigraphic divisions (ascending order): S1 – Weisslilegendes sandstone, T1 – Kupferschiefer shale bed, Ca1 – Zechstein 1 (Werra) carbonate. Pb, U, V, and Zn are highest where C_{org} is highest within the Kupferschiefer black shale, but Co and Ni are highest where Cu is highest in this section. The distribution of U and V probably reflects only the black-shale depositional environment, but Co and Ni are clearly enriched in the sediment-hosted stratabound copper-mineralizing process. The Rock Eval T_{max} climbs steadily downsectionward, indicating more exposure to hot mineralizing brines in that direction. HI, however, correlates with Cu grade indicating hydrocarbon generation correlated with mineralization intensity. Solid-phase C_{org} increases steadily in δ^{13}C downward while calcite carbon does the reverse. With increasing exposure to hot metals-transporting brines downsectionward, C_{org} was oxidized to carbonate, which was precipitated as calcite with low δ^{13}C, leaving the residual C_{org} with higher δ^{13}C. Plotted points enclosed in parentheses contained less than 0.2 weight percent organic carbon at which concentration the analytical values are questionable.
stated that it indicated syngenetic origin of the mineralization. The latter age was interpreted to be the age of the source Pb derived either from Precambrian basement of northern Europe or from clasts derived from those basement rocks. Notably, the other three Kupferschiefer samples (Dutch, British, and Polish) plotted well off the secondary isochron. The 250 Ma age is difficult to interpret, including, as it does, the barren sample as well as any Pb within unreplaced early diagenetic pyrite, but the source age (1750 Ma) is reasonable and suggests origin of the Pb in the Precambrian continental basement rocks or from (Rotliegendes?) clasts derived from those basement rocks.

Bechtel and others (1996) attempted to indirectly date the Kupferschiefer mineralization using K-Ar on illite from Kupferschiefer shale samples collected from barren pyritic, Zn-Pb, Cu-Zn-Pb, Cu, and Rote Fäule zones. Initial results indicated a Middle Triassic age of mineralization in agreement with the paleomagnetic age from Jowett and others (1987). Bechtel and others (1999) attempted to further refine the illite K-Ar ages using only mineralized samples and extrapolating the arrays of K-Ar ages in illite to 0 percent of the 2M1 polytype of illite, of presumably detrital origin, trying to date only 1M and 1Md illite polytypes of probable diageneric, authigenic origin. Results varied from 216 to 190 Ma. The wide range in extrapolated ages renders the results uncertain and has led to hypotheses of multistage mineralization, where Weissliegendes sandstone-hosted ore formed at a time later than Kupferschiefer shale-hosted ore (Michalik and Sawłowicz, 2001). Multi-stage mineralization for the disseminated (non-veinlet) ore is not convincing because sulfide-zone boundaries cross gently and continuously with no steep perturbations through all the different rock types.

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**Figure 50.** Standard von Krevelin organics maturation diagram for the samples of a Kupferschiefer section from the Sieroszowice mine, Poland, from Bechtel and others (2002). With a position more and more proximal to the source of hot mineralizing solutions, lead/zinc zone to copper zone to Rote Fäule zone, the Rock Eval pyrolysis maximum temperature (Tmax) climbs steadily, but the hydrogen index (HI) and oxygen index (OI) cross sharply across ordinary organic maturation pathways. The mineralizing process differs from ordinary maturation most obviously in being a chemical oxidation process. Plotted points enclosed in parentheses contained less than 0.2 weight percent organic carbon at which concentration the analytical values are questionable.

**Figure 51.** Relation between δ34S sulphide and δ13C carbonate for 18 samples where both were analyzed from the Spar Lake sandstone copper deposit. Samples having the combination, high δ34S sulphide and low δ13C carbonate probably indicate ore-stage coupled redox processes where carbon from methane (natural gas) was oxidized, precipitating as calcite, coupled with reduction of ore solution sulfate to sulfide, precipitating as copper sulfides. Four such samples are labeled. Figure 48 showed that much of the δ34S sulphide in Spar Lake ore was probably from sour gas H2S, so δ34S values much higher than that (greater than 6 permil) reflect increasing contributions of heavy sulfur at ore stage by thermochemical sulfate reduction, made clearer yet where the high δ34S values are accompanied by carbonates with light (organic-sourced) carbon.
The Re-Os direct dating of Kupferschiefer mineralization has been tried at several places and the results have been inconsistent. Pätzold and others (2002) found an age of 204.3 ± 0.5 Ma for Cu-sulfides at Mansfeld and argued for an epigenetic origin of the mineralization. Brauns and others (2003) measured and reported an isochron from several samples from the Sangerhausen deposit with an age of 257 ± 1.6 Ma, leading them to suggest syngenic origin. Pašava and others (2007b) reported an age of 240 ± 3.8 Ma from whole-rock samples of Cu-rich mineralization from Lubin mine. Pašava and others (2007a) reported an “altered” age of 279 ± 14 Ma from the Au-Pt-Pd-rich zone within Rote Fäule rocks of the Polkowice mine. Details on samples and the materials separated and dated for these various studies are not available. Consistency of the methods can be questioned. All these results can be compared with the results from illite dating (above) and with the results from paleomagnetic dating of the Rote Fäule by Jowett and others (1987). The paleomagnetic dating of Rote Fäule hematite initially yielded dates of 250 to 220 Ma. Those dates were later refined to 255 to 245 Ma by Nawrocki (2000) on the basis of improved polar wander paths. Only the Re-Os age of 240 ± 3.8 Ma on Cu-rich rocks from Pašava and others (2007b) is close to agreement with the paleomagnetic dating.

It remains unclear what the age of the main stage(s) of Cu mineralization is (are) for Kupferschiefer deposits. Although it is possible that different ages of mineralization could apply to distantly separated Kupferschiefer districts, it seems unlikely that, across the basin, the principal mineralizing events could have been spread from the age of host rock deposition in the Late Permian all the way through the Early Jurassic in rocks that have never seen a post-ore orogeny.

The SSC deposit with the best-constrained mineralization age is White Pine. In the Midcontinent Rift-fill rocks, the Portage Lake lavas were extruded between 1,096 ± 1.8 and 1,094 ± 1.6 Ma (U-Pb on zircon; Davis and Paces, 1990). These were followed upward by the Copper Harbor Conglomerate, coarse clastic red beds, with an interleaved andesite flow that was dated at 1,087 ± 1.6 Ma (U-Pb on zircon; Davis and Paces, 1990). In the White Pine deposit, the Junior Limestone bed has been dated at 1,081 ± 9 Ma by a 207Pb/206Pb isochron (Ohr, 1993). The Junior Limestone of the deposit has greater concentrations of common Os in the sulfides. Richards and others (1988a) studied Pb isotopes of the Cu-sulfides from Musoshi in the Copperbelt in Democratic Republic of Congo. They found, from 15 analyses of 3 samples that were free of veinlets and least recrystallized, as well as free from coarse biotite, a secondary isochron on the 207Pb/206Pb versus 208Pb/206Pb plot defining an age of 645 ± 15 Ma. Other, more recrystallized samples and veinlets were scattered below the two-stage model age line. They recognized two possible interpretations: (1) the deposit formed at 645 ± 15 Ma, or (2) the Pb isotopes of the deposit homogenized at that time in a pre-existing orebody, by a process that also introduced U and thorium into the deposit. In either case, the age is a minimum age for the disseminated deposit. Because the result is 100 million years after the youngest age allowed within the uncertainties from Selley and others (2005) (above), who also used Konkola disseminated sulfides, the second appears to be the proper interpretation; that is, 645 Ma is the age of homogenization of the Pb isotopes and it post-dates the age of mineralization. The results entirely eliminated the possibility of a contribution of mantle- or mafic-rock Pb to the Cu sulfides. This negates the hypothesis of Annels and Simmonds (1984) that the Co of the Ore Shale belt of deposits in Zambia came from metagabbro intrusions along the belt.

Selley and others (2005) also reported a 7-point Re-Os isochron with a slope age of 576 ± 41 Ma for disseminated and veinlet chalcopyrite and bornite from the Nkana-Mindola and Chibuluma West deposits and from carrollite from the upper orebody at Nchanga. The age corresponds to the early stages of basin inversion and post-dates deposition of the Petit Conglomerate at about 635 Ma. The Nkana-Chibuluma-Nchanga isochron also falls within the range of ages for Roan Group metamorphic monazites from the Chambishi...
basin. Those metamorphic ages are 592 ± 22, 531 ± 12, and 512 ± 12 Ma (Rainaud and others, 2005), and the latter two of those overlap as well with ages from veinlet mineralization.

Dating the later vein/veinlet-controlled mineralizing events in the Central African Copperbelt has proven relatively simple. For example, Richards and others (1988b) obtained an age of 514 ± 2 Ma by U-Pb for rutile in quartz-hematite-(±microcline-biotite-rutile) veins, with late calcite, siderite, anhydrite, and barite that cut stratiform Cu ore at Musoshi. Barra and others (2004) measured an age of 525.7 ± 3.4 Ma by Re-Os on molybdenite from a chalcopryrite-bearing vein that cuts stratiform mineralization at Nkana-Mindola. Richards and others (1988a) confirmed the late overprinting vein event with a U-Pb date of 514 ± 3 Ma on uraninite from a veinlet that cut the Ore Shale at Musoshi. Postpeak-metamorphic veinlet Cu-U-Mo mineralization at Kansanshi is dated at 512–502 Ma (Torrealday and others, 2000).

Uplift and cooling in the Central African Copperbelt have also proven relatively simple to date (Rainaud and others, 2005). The 40Ar/39Ar ages from Luanshya biotite give an age of 586 ± 1.7 Ma, coinciding with the oldest metamorphic events in the Copperbelt. Those metamorphic ages are 592 ± 22, 531 ± 12, and 512 ± 12 Ma (Rainaud and others, 2005), and the latter two of those overlap as well with ages from veinlet mineralization. Barra and others (2004) measured an age of 525.7 ± 3.4 Ma by Re-Os on molybdenite from a chalcopryrite-bearing vein that cuts stratiform mineralization at Nkana-Mindola. Richards and others (1988a) confirmed the late overprinting vein event with a U-Pb date of 514 ± 3 Ma on uraninite from a veinlet that cut the Ore Shale at Musoshi. Postpeak-metamorphic veinlet Cu-U-Mo mineralization at Kansanshi is dated at 512–502 Ma (Torrealday and others, 2000).

The rocks of SSC deposits have sedimentologic and petrologic features common to the same kinds of sedimentary sequences and rock types that lack the ore minerals. That simple observation is what has made the application of ordinary methods of sedimentology and sedimentary petrology effective means of understanding the genesis of these deposits.

**Petrology of Associated Sedimentary Rocks**

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**Depositional Environments**

Various depositional environments are represented in the host rocks of SSC deposits. Reduced-facies types and some sandstone Cu types have host rocks that are parts of overall large-scale transgressive sequences that may total several hundreds of meters in thickness. They feature open-shelf marine carbonate rocks overlying coastal marine rocks that are overlying fluvial coarse clastic rocks.

The Copperbelt Orebody Member of Zambia, as well as D.Strat., RSF, and SDB host beds of Democratic Republic of Congo, are deposits of coastal sabkhas (Illing and others, 1965; Kinsman, 1969; Renfro, 1974; Shinn, 1983). These are coastal marine deposits of thinly laminated carbonaceous gypsiferous mudstone and siltstone or of thinly laminated carbonaceous gypsiferous dolostone. The sabkha beds overlie coastal eolian sandstones, fluvial sandstones, or intertidal mud flat siltstones and mudstones.

The most obvious of all the sabkha environment host rocks for SSC deposits are the Cu-bearing gray to green shale beds of the otherwise red Late Permain Flowerpot Shale in southwestern Oklahoma and the Texas panhandle. Each of the beds is overlain by rock gypsum beds in vertical sequences of lithologies and sedimentary structures of sabkha depositional environments (Smith, 1976). Gypsum textures in particular identify the coastal sabkhas (Shinn, 1983).

The Kupferschiefer bed is the deepest marine rock of its local section, deposited during a maximum transgression of Zechstein time and below storm-wave base on an open shelf. It is not a coastal sabkha deposit. It has massive, bioturbated (dysaerobic), and laminated (anaerobic) facies. At the southern margin of the Zechstein Basin, however, the Kupferschiefer bed changes facies to deposits 3 to 5 m thick of a shallower water carbonate bank, with medium and thin alternation of marl and fossiliferous limestone packstones. The overlying Zechstein limestone consists of three shoaling upward sequences, 0.2 to 20 m thick, with open marine, sub-wave base burrowed carbonate mudstone or wackestone at the base, changing upwards to ooidal packstones and grainstones, and locally upward again to small-scale hemispheroidal stromatolitic, but probably subtidal beds.

Underlying the Kupferschiefer are the Weissliegendes sandstone beds that are considered to be eolian at the base and shallow marine, wave-dominated at the top (Peryt, 1989). Laterally, the Weissliegendes sandstone beds change to different depositional lithofacies. Pryor (1971) rejected an eolian origin for the Weissliegenden in parts of Germany and concluded that they were deposited on a shallow, offshore marine sea floor beyond the surf zone. Those environments are similar to the environments of deposition of the favored host rocks for sandstone Cu deposits (see below).

Host rocks for sandstone Cu deposits were deposited in a variety of environments. Ores are hosted by subtidal, high-energy nearshore marine rocks (deposits of longshore bars or tidal channels) where the host-rock depositional environments are best known at Mufulira (van Eden, 1974) and Spar Lake (Hayes and Einaudi, 1986). These are fine- to medium-grained,
well-sorted sandstones. Beach and nearshore sandstones also host some ore at Spar Lake. Those rocks are very fine- to fine-grained, well-sorted sandstones with an abundance of heavy mineral laminae in their uppermost several meters.

However, the depositional environment of the lens-shaped bodies of the highest grade rock at Mufulira, termed carbonaceous wacke and mottled sandstone, remain enigmatic. These are poorly sorted, in sharp contrast with other Mufulira quartzites (above), and they have a wider range of grain sizes, with their modes finer than other footwall and orebody quartzites. The sand grains are also less rounded (van Eden, 1974). It is likely that many of the observed features of the carbonaceous wackes attributed by van Eden (1974) to a low-energy depositional environment are actually the product of diagenesis in a petroleum reservoir that was then overprinted by the Cu mineralization.

**Carbonate/Evaporite Diagenetic Petrology**

Sedimentary rocks record the diagenetic changes that occur through time in sequences of cements and replacements. Changes include those because of increased heat and pressure from burial and changes because of introduction of groundwater of different composition. The latter are far more important in genesis of an SSC deposit, although metals-transporting solutions were apparently also hot (see earlier sections on fluid inclusions). For SSC deposits, the diagenetic changes are paragenetic sequences of ore and gangue minerals that record changes in pore-fluid chemistry before mineralizing fluids arrive, when mineralizing fluids are present, and after mineralization.

Some minerals that are typically precipitated during diagenesis of sedimentary rocks are, for the host rocks of SSC deposits, one and the same as important ore-stage gangue minerals. For the most part, these are not unusual or anomalous minerals in sedimentary diagenesis. There is no better illustration of this than with the quartz and carbonate minerals that replace the crystallloblasts (Oosterbosch, 1951, fig. 16) and the nodules of gypsum and anhydrite in silty or muddy carbonate rocks, such as in the D.Strat, RSF, and SDB beds of the Copperbelt in the Democratic Republic of Congo (fig. 8) (Muchez and others, 2008).

There is extensive literature recording the diagenetic replacement of gypsum or anhydrite by silica minerals (Folk and Pittman, 1971; Levandowski and others, 1973; Milner, 1976; Siedlecka, 1976; Chowns and Elkins, 1974; Milliken, 1979; Young, 1979; Geeslin and Chafetz, 1982; Ulmer-Scholle and others, 1993; Luebking and others, 2001; Henchiri and Slim-S’himi, 2006). In most cases, morphology of former anhydrite nodules is recognized in nodules now consisting of quartz and (or) chaledony. In the midcontinent area of the United States, the knobby, irregularly shaped silica nodules had earlier been termed “cauliflower chert” (Chowns and Elkins, 1974). In some, coarse-crystalline calcite is also present in the pseudomorphed anhydrite. Relicts of anhydrite are locally present. In one case (Ulmer-Scholle and others, 1993), pyrite is encapsulated within the replacement quartz, and oil occurs as fluid inclusions. In one case (Milner, 1976), pyrite is found as a rim surrounding gypsum in unreplaced nodules in the subsurface, whereas in two cases, hematite interpreted as weathering-zone pseudomorphs of subsurface pyrite is found as rims surrounding silica-replaced anhydrite nodules (Milner, 1976; Young, 1979). In one place where silica minerals are not found as a replacement of anhydrite nodules, coarse calcite replaces the anhydrite (Shearman and Fuller, 1969).

Dewaele and others (2006) described the entire diagenetic sequence of minerals in the orebodies at Kamoto and Musonoi, making effective use of cathodoluminescent petrology, although their main purpose in doing so appears to have been to determine generations of fluid inclusions. Nonetheless, they identified that the earliest diagenetic minerals were pyrite, anhydrite, and a first generation of dolomite. These were followed by quartz, chalcopyrite, bornite, digenite, and chalcocite that replaced most of the anhydrite and replaced pyrite everywhere except at the tops of orebodies. The earliest generation of ore minerals was followed by luminescing and therefore probably Mn-rich, dolomite and additional chalcolite and digenite. Later veinlet mineralization included growth-zoned, strongly luminescing, coarse dolomite, with chalcocite, digenite, and hematite. All these were followed by supergene alteration with malachite, azurite, chrysocolla, cuprite, and heterogenite.

**Sandstone Diagenetic Petrology**

There is more detailed reference material on sandstone petrology because petroleum geologists have extensively studied the petrology of reservoir rocks. Just as in the changes seen in the diagenesis of sulfate-bearing evaporites, the diagenetic changes seen in sandstone Cu deposits are all typical of any diagenesis, except, mainly, for presence of the Cu minerals. Every authigenic gangue mineral that is zonally distributed in the Spar Lake sandstone Cu deposit, with the exceptions of barite and Ba mica, has been found elsewhere as the product of ordinary diagenesis of sandstones.

Much of the paragenesis of the authigenic gangue minerals at Spar Lake was illustrated earlier in the section on Hypogene Gangue Minerals. This section will be used to provide references about deposition of the same authigenic minerals in unmineralized sedimentary rocks remote from SSC deposits, followed by some further description of these minerals within SSC ores.

Many of the authigenic gangue minerals arranged zonally at Spar Lake (fig. 6) have been found in the same paragenetic sequence as at the deposit, but across a gradient in burial depth and temperature in the Tertiary sandstones of the Gulf Coast in Texas (fig. 52; Boles, 1978, 1982; Boles and Franks, 1979). It is not being suggested that the Spar Lake deposit formed at the same burial depths at which the various minerals appeared on the Gulf Coast. Instead, the suggestion is that fluids from such depths, in equilibrium with albite-ankerite-chlorite, migrated to lower-temperature environments to mix with cooler fluids to deposit ore.
The discussions of petrology in sandstone Cu deposits, below, follow the sequence of mineral groups in figure 6, top to bottom. The section makes liberal use of cathodoluminescent photomicrographs to show cementation sequences (ore-stage paragenesis), and that serve to demonstrate the utility of using the luminoscope in studies of sedimentary rocks.

Iron-Titanium Oxides

Iron-titanium (Fe-Ti) oxides are minor constituents of most siliciclastic sedimentary rocks, and their authigenesis in such rocks has been studied with reference to sandstone U deposits (Adams and others, 1974; Reynolds and Goldhaber, 1978), to heavy mineral sand deposits (Bailey and others, 1956; Lind, 1961; Temple, 1966), and to SSC deposits (Dimanche and Bartholomé, 1976). These papers that describe changes to Fe-Ti oxide minerals after sedimentation all draw fundamentally on knowledge of Fe-Ti oxides in the source areas for the sediments (Haggerty, 1976).

The Fe-Ti oxides are useful in studying SSC deposits because they are sensitive indicators of redox states and redox histories. Parts of the redox histories are common between all three of the deposit types mentioned above. Two types of Fe-Ti oxide grains are useful starting materials. The first are grains that were originally of the ulvöspinel-magnetite solid solution, which typically alter subsolidus, but still in mafic rocks, to grains of ilmeno-magnetite. The ulvöspinel-magnetite solid solutions exsolve on cooling, with ilmenite forming rod-shaped grains along the magnetite isometric crystal axes. Cuts through the grains have the appearance of trellises of ilmenite within magnetite (Haggerty, 1976) (fig. 53). The second useful starting materials are ilmeno-hematite grains typical, instead, within Fenner reaction series igneous rocks. They consist of platelets of ilmenite within hematite, the ilmenite occurring along {0001} rhombohedral planes in the hematite (fig. 54), and these ilmenite platelets are also typically just a few to 10 µm thick (Haggerty, 1976). Both types of common Fe-Ti exsolution grains typically survive weathering, transport, and deposition, and remain mostly stable through all red-bed diagenesis, although alteration to produce fine, red, hematite pigment is typical in red beds (fig. 53) (Turner, 1980). In red beds, these two types of Fe-Ti oxide clasts continue to be stable throughout burial diagenesis (Turner, 1980) (for example, fig. 54).

In non-red sediments, however, the diagenetic history of both types of Fe-Ti exsolution detrital grains can be complex, and, for the case of SSC-deposit host rocks, it is complex (Dimanche and Bartholomé, 1976; Hayes, 1983, 1984). In reducing early diagenesis, the Fe of the magnetite, hematite, and ilmenite dissolves. However, the Ti of the ilmenite does...
not. Instead the ilmenite needles or platelets become pseudo-
morphed by TiO₂ phases, typically leucoxene, but probably
in places, either rutile or anatase (figs. 19A and 19B). Trellis
or stacked-plate leucoxene aggregates are far too delicate to
have survived erosion and transport as they are found now.
Instead, they had to be eroded and transported as grains
that still consisted mainly of magnetite or hematite, and the
alteration of exsolved ilmenite to leucoxene happened during
diagenesis. There is typically also a hand-lens to microscopic
scale association of the leucoxene trellises or stacked-plate
aggregates with pyrite or another Fe-bearing sulfide. Dissolu-
tion and pseudomorphing can apparently take place early, the
typical case for reduced-facies host rocks, or may take place
somewhat later in sandstone Cu hosts, perhaps because of
reducing fluids, such as petroleum, during their migration into
reservoir rocks. In reduced-facies and sandstone Cu hosts, ore-
stage replacement of pyrite by Cu sulfides accompanied by an
ore-stage generation of hematite pigment (fig. 19B and 19C;
Dimanche and Bartholomé, 1976, fig. 11C) is a final step.

Carbonate Cements
Calcite is one of the commonest of all cementing minerals in all types of sandstones (Dapples, 1971; Folk, 1974;
Fuchtbauer, 1972; Jonas and McBride, 1977). It can occur in
many different places in a diagenetic sequence of cements and
temporally from early to late (fig. 55; Blanche and Whitaker,
1978). As a cement, it has micritic or sparry textures. Folk
(1974) emphasizes that the micritic cements typically precipi-
tated from saline pore waters, and the sparry cements typi-
cally precipitated from dilute groundwater. In marine settings,
calcite cements can precipitate early, such that grain-to-grain
contacts of detrital grains are rarely intersected by the plane of
the thin section.

The calcite of the Spar Lake sandstone Cu deposit and
probably that of Dzhezkazgan (Gabrina and Tsepin, 1975) is
not early. It likely replaces parts of adjacent detrital quartz
and feldspar grains. It is poikilotopic, enclosing a number of
individual detrital 0.1-mm silicate grains in optically continu-
ous crystals of several millimeters. Although it is sparry, it did
not precipitate from dilute groundwater, as demonstrated by
fluid inclusions with greater than 10 eq. wt. % NaCl. Pos-
sibly, the elevated temperatures and deep burial at the time
of its deposition has made it an exception to Folk’s (1974)
generalities about micritic versus sparry calcite cements. It has
unique composition with high Fe through the pyrite-calcite,
galena-calcite, and chalcopyrite-calcite zones and with high Mn near its disappearance at the boundary between bornite-calcite and chalcocite-chlorite zones (fig. 22). At many places at Spar Lake, Cu sulfides replace calcite. The interpretation is that a more distal, early ore-stage mineral was replaced by a more proximal, later ore-stage mineral. Ankerite, ferroan dolomite, or siderite is common in sandstones that have undergone relatively deep burial (Boles, 1978; Macaulay and others, 1993; Hendry and others, 2000) or that have undergone diagenesis under reducing conditions (Tye and Hickey, 2001).

**Clay Cements**

Authigenic clay cements are common in sandstones (Füchtbauer, 1974; Wilson and Pittman, 1977). The clay minerals that commonly form as cements in sandstone are illite, kaolinite, and chlorite, and smectite is common in graywackes and lithic sandstones. The first three minerals have diagnostic textures where they grow authigenically as sandstone cements (Wilson and Pittman, 1977). Kaolinite forms books or vermiciform books. Illite bridges pores in arching sheets. Chlorite coats detrital grains with crystals standing on edge, with the c-axis tangential to rounded grain boundaries and its cleavages “edge-on.” Only the smectites have poorly defined crystals.

The high birefringent phyllosilicates zonally distributed in quartzites at Spar Lake are found coating quartz or K-feldspar grains but do not coat every grain completely, allowing some to develop long contacts with adjacent quartz grains (fig. 22). Rarely, the mica occurs inside quartz or K-feldspar overgrowths, possibly originally as a clay coating on detritus that is now recrystallized. Hayes (1983, Appendix I) showed such grain-coating white micas are Barich within bornite-calcite and chalcocite-chlorite zones.

Chlorite is common as a sandstone cement in many settings (Wilson and Pittman, 1977; Curtis and others, 1985). Authigenic chlorite is present in a number of SSC deposits. The chlorite in Spar Lake’s chalcocite-chlorite zone has typical textures with the chlorite crystals edge-on forming grain coatings. However, everywhere quartz or K-feldspar overgrowths are present, the chlorite is not in direct contact with the detrital grains; rather, it is found farther toward the centers of former pores, coating overgrowths, not detritus. Chalcocite or bornite locally occurs along cleavages of the chlorite, either replacing chlorite or growing synchronously with it. The chlorite also partially replaces detrital muscovite in that zone of the deposit (fig. 57). As stated earlier, the Spar Lake sandstone-cementing chlorite of the chalcocite-chlorite zone is an extraordinarily Fe-rich chamosite, which is unlike the chlorite cement found in typical sandstone diagenesis and certainly not metamorphic.

**Figure 55.** Photomicrograph showing calcite cementing a feldspathic sandstone. Photomicrograph in transmitted, cross-polarized light. Quartz is various shades of gray in rounded to subrounded grains. A grain of twinned albite is at center. The first stage of cements is hematite (hem, reddish brown) rimming detritus and altering biotite (bio) along cleavages. The second is sparry calcite (cal) in a poikilotopic texture (large calcite crystal enclosing multiple detrital grains – calcite grain 1 encloses 13 detrital grains within the field of view). Sample from the Permian Cutler Formation, southeastern Utah.

**Figure 56.** Photomicrograph showing grain-rimming “illite” cement. Photomicrograph in transmitted, cross-polarized light. According to X-ray diffraction, the highly birefringent mineral that coats detrital grains is a 2M1 white mica, but it probably originated as smectite or illite clay coatings, prior to burial metamorphism. Note that where the detrital grains were coated, no quartz overgrowth occurred, but where they were not coated, syntaxial overgrowth on detrital quartz cores did occur. Sample from the Mesoproterozoic Revett Formation, northwestern Montana.
Authigenic Feldspars

Authigenic feldspars in sedimentary rocks are widespread and common in many different environments (Baskin, 1956; Kastner and Siever, 1979). Authigenic albite is more common in carbonates, and K-feldspar is more common in siliciclastics. The authigenic K-feldspar occurs most commonly as syntaxial overgrowth on detrital feldspar cores. Albite, instead, appears to commonly replace detrital Ca-bearing plagioclase or forms euhedral grains within carbonate rocks.

Albite increases at the expense of detrital K-feldspar and detrital Ca-plagioclase with depth in Gulf Coast sandstones in Texas (Land and Milliken, 1981; Boles, 1982). These changes parallel other changes at depth that include the increase in illite at the expense of smectite, increasing cementation of ankerite, increasing quartz overgrowths, and the appearance and breakdown of kaolinite to chlorite (fig. 52). For some of these mineral reactions, there is a local source of needed elements. For example, the smectite conversion to illite can provide Fe and Mg for replacing calcite with ankerite and for the authigenic chlorite, and it can provide silica for detrital quartz overgrowths. It can release aluminum for kaolinite and later chlorite formation. But other elements must be newly introduced by solution. For albite, introduced Na is needed, and possibly accompanying that, chloride is needed to convert the relatively fresh groundwater at shallow depth to briny groundwater at the depths of precipitation of authigenic albite, chlorite, and ankerite. Land and Milliken (1981) calculated that Na for the albitionization from seawater buried with the sediment was insufficient and suggest that dissolution of halite is needed.

Authigenic K-feldspar and authigenic quartz at Spar Lake occur as overgrowths on detrital K-feldspar and quartz grains, respectively, and their textures have been shown in figures 16, 20, 21, 22, and 23. Authigenic albite forms final pore-filling cements (fig. 24), and it replaces earlier detrital and overgrowth K-feldspar. Albite also accompanies other cements, including apatite and Cu sulfides, in replacing detrital Ca-bearing plagioclase (fig. 58). Albite or another authigenic mineral has replaced all the detrital Ca-plagioclase in Spar Lake rock (Hayes, 1983, Appendix I).

Barite Cement

Barite is common as a trace authigenic constituent in sedimentary rocks, most frequently observed in gray and black shales (Johnson and others, 2009). Barite as a sandstone cement is much less widespread but is known from Germany, Italy, England, northeastern Africa, Egypt, and Oklahoma, United States (Tarr, 1933). It has been best studied from Oklahoma, where it is abundant in the upper half of the Early Permian Garber Sandstone. The barite forms concretions in sandstone and most prominently forms “barite roses.” A barite rose is a sand-barite crystal, where sand grains are cemented by and enclosed in large (0.5 to 10 cm) cementing crystals with poikilotopic texture. The barite crystals are multiply twinned on the b axis, with as much as 20 individual crystals sharing the same b axis and with the crystal axes of individual crystals spraying in acute angles to one another. In the sand-barite crystals, barite has partially replaced quartz and other detrital minerals such that most detrital grains are now angular, whereas outside the sand-barite crystal, the detrital grains are moderately to well rounded, each with a hematite dust rim. Hematite dust rims within the barite rose, similar to the sand grains themselves, are partially replaced, such that the color of the inside of the barite rose is red-gray rather than red-orange (Tarr, 1933; Ham and Merritt, 1944).

Sandstones within the Garber are host to the Paoli and other nearby sandstone Cu occurrences (Shockey and others, 1974). Cu-Ag mineralization at Paoli occurs along mineral appearance fronts between mottled hematitic sandstone and pastel green sandstone, with ore grades as chalcocite and native silver developed in the pastel green sandstone adjacent to the

Figure 57. Photomicrographs of chlorite replacing muscovite along cleavages from the chalcocite-chlorite zone of the Spar Lake sediment-hosted stratabound copper sandstone deposit (fig. 6). A, Photomicrograph in transmitted, plane-polarized light. Chlorite is limy green. Muscovite appears yellowish. B, Photomicrograph in transmitted, cross-polarized light. Muscovite has high birefringence colors of yellow and red. Chlorite is very low birefringent with blue-gray appearance. A detrital muscovite grain was bent during loading and compaction of the sand. Chlorite partly replaced the muscovite along its cleavages.
Figure 58. A, Photomicrograph in plane-polarized transmitted light of a replaced detrital plagioclase grain from the bornite-calcite zone of the Spar Lake sediment-hosted stratabound copper sandstone deposit (fig. 6). B, Cathodoluminscent photomicrograph of the same field of view. Apatite has high relief in A and luminesces yellow in B. C, Interpretive overlay of the same field of view. A detrital plagioclase grain has been completely replaced by authigenic minerals including apatite, barite, barium-rich white mica, albite, bornite, and chalcocite.

lack carbonaceous plant material; those that host roll front U deposits in Wyoming are, instead, gray and contain conspicuous amounts of carbonized plant debris. The mottled gray and red sandstone interior to the paleochannels and between Cu-Ag mineralization fronts also contains barite roses.

The barite concretions and barite roses of Oklahoma relate spatially, at least, to RBA Cu occurrences and deposits in the Permian rocks of the region. They may be more specifically within the Cu source rocks, the “altered” rocks of the usage in this report.

Spar Lake barite occurs as pore-filling cement that has grown toward centers of former pores from quartz or K-feldspar overgrowths (fig. 23). Most sulfide grains show the same textural relation. However, if a solution was introduced into a sandstone and was simultaneously saturated with respect to barite and K-feldspar, and K-feldspar detrital grains are present, the K-feldspar would precipitate as a syntaxial overgrowth, probably at the same time that barite precipitated, yet resulting in the appearance that overgrowth preceded barite filling the pores. Thus, it is likely that the barite grew at the same time as the K-feldspar overgrowths. The same textural observation and the same logic hold for most bornite-digenite-chalcocite, chalcopyrite, galena, calcite, and chlorite. Cu sulfides and galena share former pores with barite, and there is no evidence that either replaced the other (fig. 23).

Barite cements have also been found (1) in the Mindola Clastics below ore and within the Copperbelt Orebody Member ore at Konkola, Zambia (Sutton and Maynard, 2005), (2) in a single sample of Weissliegendes Sandstone below mineralized Kupferschiefer on the Fore-Sudetic Monocline, Poland (Michalik, 2001), (3) within the Udokan deposits, Russia (Volodin and others, 1994), and (4) in red-bed-type deposits near Nacimiento and Tularosa, New Mexico (Lindgren and others, 1910). Barite is clearly an important gangue phase related to many SSC deposits.

Tourmaline

Among the most unexpected of the minerals found as an ore-stage product in SSC deposits is the metasomatic tourmaline described from Zambia (Darnley, 1960; Selley and others, 2005) and Democratic Republic of Congo (Oosterbosch, 1951). Tourmaline is not a mineral commonly formed during diagenesis. Yet, Gautier (1979) found authigenic, pore-occluding tourmaline crystals, as well as overgrowths on detrital grains, in the Late Cretaceous Eagle Sandstone reservoir rocks of the front. The mottled hematitic sandstone occupies the interiors of paleochannels of sandstone within the otherwise muddy Wellington Formation. Although comparable in zonal geometry, the mottled hematitic sandstone is not comparable in color with the altered tongues of roll front U deposits. The hematitic parts at Paoli are deep red, nearing brick-red in color, whereas roll front U deposit altered tongues are grayish yellow (Shirley Basin, Wyo.; Harshman, 1972) or pink to grayish pink (Powder River Basin, Wyo.; Davis, 1969; Dahl and Hagmaier, 1976). The host sandstones are also notably different. The sandstones that are host to Paoli Cu-Ag mineralization are pastel green, and they
Petrology of Associated Metamorphic Rocks

It is common that SSC deposits are metamorphosed along with the rocks of their host sedimentary packages. This allows the conclusion that, everywhere that that is observed, the SSC deposits are pre-metamorphic. Veins and stockworks that post-date metamorphism or accompanied metamorphism occur as well, particularly within the Central African Copperbelt, and those are described and modeled separately. They are described in the section on Associated and Transitory Deposit Types. All examples of that type of deposit, syn- or post-orogenic veinlet controlled, have been removed from the accompanying database and from the grade and tonnage models.

Features of metamorphism observed in silicate and carbonate assemblages are also found in metamorphosed sulfide-bearing rocks. Among these are metamorphic devolatilization reactions, crystalloblastic series, ‘textural series’ that include polygonization, annealing, and recrystallization, and retrograde assemblages.

Rock Names

Only a few terranes that contain SSC deposits cross sufficiently through metamorphic isograds to allow directly equating protolith sedimentary rock types and their metamorphosed equivalents. These include the Central African Copperbelt, the Kalahari Copperbelt of Botswana and Namibia, and the Ablah Group in southwestern Saudi Arabia. A number of other deposit-bearing belts cross between different metamorphic facies, but sufficient descriptions of those have not been found.

Lee and Glenister (1976) attempted correlation of the Damara Supergroup of the Oamites deposit area in Namibia with the Katangan Supergroup in Katanga (table 1). In these Precambrian rocks, all the correlations are lithostratigraphic, so the metamorphosed rock type and protolith equivalency that they propose has no fossil-age control and only broad absolute-age control. The correlation is on what they believe to be an equivalent of the Petite Conglomerate, the Mariano aged tillite, marking the separation, in Katanga, of the Nguba Group (formerly Lower Kundulungu) from the Upper Kundulungu Group, which is the uppermost correlated tillite of the table. Notable equivalents that they propose are (1) basal arenitic sections (correlated with Mindola Clastics Group) are metamorphosed to sericitic quartzite with lesser biotitic schist, phyllite, and metaconglomerate with schistose biotitic matrix; (2) carbonaceous argillitic dolomites correlate with biotitic schist and amphibolitic schist; (3) carbonaceous shale/argillite correlates with graphitic schist; and (4) dolomite correlates with dolomitic marble. Strong support for their sedimentary to metamorphic rock-type correlations comes from the Central African Copperbelt and the Ablah Group of Saudi Arabia.

The Katangan Supergroup of Democratic Republic of Congo is defined entirely by sedimentary rock names, although even the earliest workers recognized that all but the Upper Kundulungu rocks had been metamorphosed to lower greenschist facies or higher. Extrapolating from rocks with better described metamorphic minerals and fabric from the south would indicate that some chlorite and some white mica are of metamorphic origin in the rocks in Congolese deposits. Francois (1992) stated that, 90 km south of Kolwezi, metamorphism reaches “lower mesozoone” grade with the formation of garnet and green amphibole. The border between Zambia and Democratic Republic of Congo is just 10 km beyond that greenschist/amphibolite isograd. Farther eastward, in the Zambian part of the Copperbelt, Mendelsohn (1961b) described the metamorphism and stated that the metamorphic grade in Katangan rocks increased from northeast to southwest, from the Mokambo concession to the Roan-Muliashi Basin. Hitzman (2000) showed isograd boundaries extending from the northwest to southeast such that a lower to middle greenschist isograd passes between the Nchanga and Mimbula deposits (fig. 38 for locations), with all the Chambishi Basin at middle greenschist facies. He showed a middle greenschist to lower amphibolite facies isograd passing between Chibuluma and the Roan-Muliashi Basin. Correspondingly, the metamorphism at Konkola, farthest north of the belt of shale ores, is said to ‘rarely reach the chlorite facies’ (Schwellnus, 1961); the Ore Shale is described as a biotite-quartz-argillite at Chambishi (Garlick, 1961a); and the Ore Shale consists mainly of tremolite and biotite, with local scapolite and clinozoisite in the Roan-Muliashi Basin (Mendelsohn, 1961b). Mendelsohn labels the metamorphism as epidote-amphibolite facies and he notes the presence of epidote and actinolite in carbonate-rich quartzites. He states that tremolite is locally abundant near the base of the Ore Shale, a unit that is commonly also crenulation-folded with strong development of axial-plane cleavage. The scapolite is found only in the Ore shale and occurs as porphyroblasts with inter-inclusion textures of the silicates but, interestingly, not of the sulfides. Cu sulfides are expelled in scapelite porphyroblast growth and they rim each porphyroblast. Actinolite characterizes the impure dolomites, but tremolite occurs in the purer dolomites and in the basal section of the Ore Shale.

Rock-type equivalents from lowermost greenschist to upper amphibolite are well known in the Ablah Group of southwestern Saudi Arabia. The rocks can be walked in
Table 1. Stratigraphy and lithology of the Damara Supergroup (Oamites Area) Correlated with the Katangan Supergroup, Zambian Copperbelt.¹

<table>
<thead>
<tr>
<th>Damara Supergroup (Oamites Area)</th>
<th>Katangan Supergroup (Zambian Copperbelt)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Biotitic schist, quartzite, minor marble, amphibolitic schist, tillite</td>
<td>Shale, quartzite, tillite</td>
</tr>
<tr>
<td>Marble, quartzite, graphitic schist</td>
<td>Khomas Subgroup</td>
</tr>
<tr>
<td></td>
<td>Swakop Group</td>
</tr>
<tr>
<td>Biotitic schist, amphibolitic schist, gneiss, ore horizon quartzite and conglomerate, marble</td>
<td>Dolomite, shale, tillite</td>
</tr>
<tr>
<td>Red-brown quartzite, conglomerate</td>
<td>Hakos Subgroup</td>
</tr>
<tr>
<td>Sericitic quartzite, schist, phyllite, conglomerate, limestone</td>
<td>Carbonaceous shale, argillite</td>
</tr>
<tr>
<td></td>
<td>Upper Kundulungu Group</td>
</tr>
<tr>
<td></td>
<td>Lower Katungal Group</td>
</tr>
<tr>
<td></td>
<td>Mwashia Group</td>
</tr>
<tr>
<td></td>
<td>Upper Roan Group</td>
</tr>
<tr>
<td></td>
<td>Copperbelt Orebody Member</td>
</tr>
<tr>
<td></td>
<td>Kitwe Group</td>
</tr>
<tr>
<td></td>
<td>Mindola Clastics Formation</td>
</tr>
</tbody>
</table>

¹Katangan stratigraphic nomenclature updated with reference to Selley and others (2005).

nearly continuous outcrop exposure from lowest greenschist facies into upper amphibolite facies along several individual lithostratigraphic units along a distance of about 15 km. Equivalents are shown in table 2 from Sanders (1983). The (para-) amphibolite that Sanders lists is a conspicuous rock type. It has sprays of tremolite crystals commonly 2 to 3 cm long, with the crystal spray flattened within the schistocity, and the rock otherwise consisting of granoblastic mosaic of equant 0.5 mm grains of quartz, oligoclase, and biotite, with traces of relict dolomite (fig. 59).

Additional details about the Ablah Group rocks are worthwhile to the discussion of metamorphic rock types and their protoliths. The Al Mehdadah prospect was earlier hypothesized to be a volcanogenic massive sulfide occurrence in metavolcanic host rocks that were basement to the Ablah Group metasediments. Sanders (1983) indicated, instead, that Al Mehdadah was an amphibolite-grade metamorphosed SSC occurrence. Other lateral changes of note in rocks of the sequence are that red beds change laterally along the 15 km to magnetite-bearing, still hematite-bearing, flat-gray-colored phyllitic albitic quartzites, and eventually to quartz-oligoclase-orthoclase-muscovite-magnetite schist near Al Mehdadah. The para-amphibolite unit correlates with argillitic dolomitic phyllite in the lower greenschist zone and overlies the lowermost section of former red beds. The Al Mehdadah prospect, itself, has footwall rocks of intensely magnetite-rich quartzites that originated as red beds. Within these is a crosscutting zone of rocks composed of 75 v. % magnetite and 25 v. % quartz ending against the footwall of the Cu-sulfide-rich conformable layer, grading from 1 to 2 percent Cu (as chalcopyrite where cut in core holes). The magnetite-quartz rock is interpreted as a pre-metamorphic stockwork of hematite veinlets cutting footwall red beds, originally a hematitic feeder zone to the deposit.
Devolatilization Reactions

Prograde metamorphic reactions are commonly devolatilization reactions that liberate water (H₂O), carbon dioxide (CO₂), or other gases as temperatures and (or) pressures increase. Metamorphic reactions can illustrate, for example, the formation of tremolite in the dolomitic basal ore shale of Zambia or in the para-amphibolite of the Al Mehdadah vicinity, Saudi Arabia, as follows:

\[
dolomite + quartz + talc = tremolite + CO_2 \quad (Kerrick, 1974)
\]

\[
2CaMg(CO_3)_2 + 4SiO_2 + Mg_3(Si_4O_10)(OH)_2 = Ca_2Mg_5(Si_8O_22)(OH_2) + 4CO_2
\]

(2)

A couple of specific devolatilization reactions are of interest for SSC deposits and their host sections. These are a redox reaction and a desulfidation reaction.

In the Belt Supergroup, Jack Harrison (oral commun., 1979) observed that Radvalli Group rocks of the eastern Belt Basin were red and hematitic, but that in the western Belt Basin, many of the stratigraphically correlated Radvalli Group rocks were lavender gray and contained small (0.3–1.0 mm) octahedra of magnetite (Kleinkopf and others, 1972). Harrison (oral commun., 1979) proposed that the magnetite was a burial metamorphic product from consumption of hematite. Harrison observed that western Belt rocks reached biotite grade of regional burial metamorphism while the eastern Belt was at a maximum of chlorite grade (Harrison, 1972; Maxwell and Hower, 1967). The redox reaction would be:

\[
hematite = magnetite + O_2
\]

\[
6Fe_O_3 = 4Fe_2O_4 + O_2
\]

(3)

Nadoll and others (2012) have recently demonstrated that Harrison was correct, using oxygen-isotope geothermometry. Quartz-magnetite and calcite-magnetite mineral pairs from the Spar Lake deposit were shown to have formed at temperatures from 359 to 437 °C. Compared with temperatures derived from S-isotope studies (50 to 116 °C in galena-calcite zone; Hayes and others, 1989), fluid-inclusion microthermometry (74 °C in galena-calcite zone to 178 °C or more in chalcocite-chlorite zone; Hayes, 1990), and fluid-inclusion volatiles (165–200 °C in chalcocite-chlorite and bornite-calcite zones; Hayes and others, 2012), these oxygen-isotope temperatures cannot be temperatures of ore deposition. However, they correspond well with temperatures of burial metamorphism from studies of the biotite-forming reaction at Spar Lake (Hayes, unpub. data). The Spar Lake magnetite is also thus probably burial metamorphic.

Other metamorphosed SSC deposits and their host sections also show this metamorphic magnetite including the Ablah Group of Saudi Arabia and probably Udokan, Russia. In the Udokan series, the ore-bearing Upper Sakukan Formation is underlain by 1,000 to 1,700 m of “magnetite-carrier sandstones” of the Lower Sakukan Formations and by 1,100 to 1,400 m of magnetite-bearing sandstones and siltstones of the Talakan Formation (Bogdanov and others, 1966). Magnetite also occurs in the chalcocite- and bornite-bearing ore but not in the peripheral chalcopyrite and pyrite zones (Volodin and others, 1994). Little is made of it, but magnetite is common in the Zambian Copperbelt. It occurs predominantly in footwall sediments (the Mindola Clastics Formation), at Chibuluma (Winfield, 1961), in the Roan-Muliashi Basin (Lee-Potter, 1961), and at lower metamorphic grade at Luansobe (Groen, 1961, p. 406–407) and Mufulira (Brandt and others, 1961). In the lower greenschist facies rocks of Luansobe and Mufulira, there is also the possibility that the magnetite is detrital and typical of red beds (see fig. 53).

The other metamorphic devolatilization reaction type of interest is desulfidation. The one that is locally common is the desulfidation of pyrite to form pyrrhotite:

\[
pyrite = pyrrhotite + S_2
\]

\[
2FeS_2 = 2FeS + S_2
\]

(4)

With the exception of Baluba (Lee-Potter, 1961), every SSC deposit that is at amphibolite metamorphic grade and has published detailed information on sulfide assemblages contains some pyrrhotite. These include Oamites, Namibia (Lee and Glenister, 1976); Roan Antelope, Zambia (Mendelsohn, 1961b); Nkana North Limb, Zambia (Annels, 1974); Nkana South, Zambia (Notebaart and Vink, 1972; Brems and others, 2009); Al Mehdadah, Saudi Arabia (Sanders, 1983), and Malundwe and Chimiwungo, Zambia (Benham and others, 1976). Traces of pyrrhotite are reported at Spar Lake, at the biotite isograd of greenschist facies (Hayes, 1983), and pyrrhotite is present at Chambishi East, Chambishi West, and Pitanda, Zambia, at middle to upper greenschist facies (Annels, 1989). However, pyrrhotite is not reported from deposits in the Central African Copperbelt that are at lower greenschist facies, not even where the same (Copperbelt Orebody Member) host rock is mineralized (Richards and others, 1988a; Sutton and Maynard, 2005; McGowan and others, 2006). Pyrrhotite is, however, found in vein ores in the Democratic Republic of Congo and Zambia and is relatively abundant at Kansanshi (Broughton and others, 2002).

Because traces of pyrrhotite occur in unmetamorphosed Kupferschiefer ore and unmineralized host rocks in Germany (Jung and Knitzschke, 1976), it is not certain that all the pyrrhotite in metamorphosed SSC deposits is formed by devolatilization of pyrite.

Crystalloblastic Series

Sulfides of SSC deposits form parts of a crystalloblastic series, together with silicate and carbonate minerals that form a crystalloblastic series of their own. Only parts of the crystalloblastic series are known because of the probable disappearance of certain minerals from metamorphosed products. As an example, at White Pine, which is unmetamorphosed, pyrite relicts are found coated by chalcocite, digenite, or bornite (Brown, 1971, figs. 16, 22, 23). In contrast, at Spar Lake,
which has been post-ore metamorphosed to biotite grade of
greenschist facies, pyrite is not found in microscopic contact
with any Cu sulfide except chalcopyrite. Chalcopyrite would
be an intermediate composition between, for example, bornite
and pyrite. It is likely that, at ore-stage at Spar Lake, relics
of pyrite once occurred within bornite or chalcocite, but that
post-ore metamorphism homogenized the composition of the
composite sulfide grains to a chalcopyrite + pyrite grain. Simi-
larly, at Spar Lake, chalcopyrite has not been found in grain
contact with either primary digenite or primary chalcocite.

In the list below, the minerals (sulfides and others) are
listed top to bottom in sequence of their tendency to form
euhedral crystals with increasing metamorphic grade. Where the
exact positioning is uncertain, typically because the minerals do
not occur in grain contact, a line is placed between the two.

  Magnetite
  Pyrite, Garnet, Tourmaline
  Amphibole (Tremolite)
  Kyanite
  Ankerite, Ferroan Dolomite, Siderite
  Biotite, Muscovite, Chlorite
  Albite in Carbonate Rock
  Calcite
  Quartz, K-feldspar, Albite in siliciclastic rock
  Galena, Chalcopyrite, Bornite, Digenite, Chalcocite

The types of petrographic relations that were observed in
assembling the above list are shown in figure 60. The inter-
growth of digenite and bornite has neither mineral imparting
a euhedral boundary against the other. Ankerite clearly forms
a rhomb-shaped crystal with its euhedral boundaries imparted
against the two sulfides, but magnetite imparts euhedral
boundaries against the ankerite and the sulfides. Clots of
sulfides with recrystallized magnetite, carbonates, and silicates
are minor at Spar Lake. Most of the ore has not annealed
or recrystallized after diagenesis, and the majority of the
rock still has cementation textures like those in many earlier
photomicrographs.

Perhaps the most important observation of the above
crystalloblastic series, as it affects SSC deposits, is that all the
common Cu sulfides of these deposits have the lowest of all
crystalloblastic strengths. After metamorphism, the Cu sulfides
appear interstitial to all the minerals with higher crystalloblastic
strengths, which include all other mineral groups in the rocks.
Thus, a strongly metamorphosed SSC ore has all sulfides,
except pyrite, in sites interstitial between quartz, feldspar,
mica, and carbonate grains, and the Cu sulfides occur only in
anhedral grain shapes. They are anhedral particularly against
metamorphic minerals such as garnet, tremolite, and tourmaline.

Textural Series

Annealing and recrystallization textures are observed
in SSC deposits at upper greenschist and higher grades
facies, and they can be used to distinguish between miner-
als of metamorphic and detrital and diagenetic origin. Many
sedimentary and diagenetic mineral grains are distinguished by
subtle discontinuities. For example, detrital quartz and feldspar
grains are commonly crowded with fluid inclusions, whereas
grain-rimming syntaxial overgrowth cements are not. Among
carbonate minerals, diagenetic-aged cements commonly grow
with crystal terminations towards the centers of pores or former
pores. Where annealing and recrystallization have occurred,
such discontinuities are erased, and the product is a mosaic of
even-grained quartz-feldspar-mica-carbonate with or without
porphyroblasts. Triple-point textures are common. Where
sulfides other than pyrite were involved in the annealing and
recrystallization, and the sulfides are less than about 10 v. %, the
sulfides are interstitial to the other mineral groups and anhedral.
Where the Cu sulfides constitute more than about 10 percent of
the rock, they go through textural series with advancing defor-
mation and metamorphic grade. This includes grain growth,
dislocation, polygonization, and recrystallization (for example,
the Al Mehdadah prospect, Saudi Arabia).

Retrograde Assemblages

Retrograde metamorphic minerals are found in metamor-
phosed SSC deposits and do not differ from the retrograde
minerals of other metamorphic rocks of similar bulk composi-
tion. Chlorite that is retrograde after garnet or tremolite is a
common example.

Retrograde sulfide assemblages and textures, however,
are common in metamorphosed SSC deposits with some
interesting details. The incoherent exsolution texture of bornite
that unmixed from the high-temperature solid solution with digenite (fig. 28) is ubiquitous in metamorphosed deposits. The same incoherent exsolution of bornite and digenite is absent from unmetamorphosed deposits, such as the Polish Kupferschiefer, the Paoli deposit, Lisbon Valley, and White Pine. Additional details can be seen in samples from Spar Lake (fig. 28). First, the grain of esolved bornite and digenite grain has none of its own crystal boundaries. The grain boundaries are against detrital (rounded) and diagenetic quartz and calcite, rather than metamorphic phases. The calcite is porefilling relative to the detrital and overgrown quartz. Other Cu-sulfide grains have mostly cuspat grain boundaries, suggesting all originated as cements. The quartz-feldspar framework is detrital with cementation and remains unmetamorphosed. Individual sulfide grains or composites are cementing, yet one sulfide has a clearly retrograded texture where the higher temperature bornite-digenite solid solution has unmixed. A secondary electron image of the bornite-digenite exsolution grain (a gray-scale image) was determined to have mole proportions of 43.8 percent bornite and 56.2 digenite. Then, following the assumption that a grain of bornite-digenite solid solution of that composition exsolved as a closed system, and that the solid solution cooled to intersect the solvus of Grguric and others (2000), the solvus would have been intersected at about 177 °C. Other incoherent exsolution-textured bornite-digenite grains have mole proportions of as high as 91 percent bornite and 9 percent digenite. That composition intersects the solvus at 237 °C. This latter temperature is 37 °C higher than the highest temperature determined for the ore stage at Spar Lake by S-isotope geothermometry (Hayes and others, 1989), fluid-inclusion microthermometry (Hayes, 1990), or gas-phase equilibria using fluid-inclusion compositions (Hayes and others, 2012). It is 53 °C higher than the highest fluid-inclusion homogenization temperature determined from the bornite-calcite zone, the mineral zone that the sample comes from. The preferred explanation is that an earlier composite sulfide grain, perhaps chalocite + chalcopyrite, homogenized by melting to the $\text{bn}_{0.91}\text{dig}_{0.09}$ composition as a closed system, and then retrograded by unmixing as temperatures decreased from peak metamorphism at greater than 430 °C to less than 237 °C. The bornite-digenite textures of figure 60 are probably the same type of exsolution.

Bornite-digenite is one of at least three types of Cu-sulfide exsolution textures described from SSC ores. This incoherent exsolution type has been described from Spar Lake, Roan Antelope in Zambia (Bateman, 1930; Brummer, 1955, fig. 25), and Fungurume in Democratic Republic of Congo (Oosterbosch, 1951). Coherent exsolutions of digenite from chalcopyrite indicating isometric parentage have been illustrated from Mufulira, Zambia (Bateman, 1930), Roan Antelope (Brummer, 1955, fig. 30), and Witvlei, Namibia (Anhaeusser and Button, 1973, fig. 11). Coherent exsolutions of chalcopyrite and digenite along cubic crystallographic directions from bornite (fig. 61) are recognized at Spar Lake, the Snowstorm mine also in the Western Montana Copper Sulfide Belt, Fungurume (Oosterbosch, 1951, fig. 25), and White Pine (Brown, 1971, fig. 27).

### Theory of Ore Deposit Formation

There are six requirements in order for SSC deposits to form. Evaluating potential for SSC deposits using these six requirements is fully analogous with the thinking commonly applied to evaluate a petroleum exploration “play.” Petroleum plays are conceptually analyzed considering (1) source rocks, (2) maturation, (3) migration, (4) reservoir rocks, (5) trap, and (6) seal rocks (Ulmishek, 1986; Bird, 1986; Magoon, 1988). In order to form a SSC deposit, it is generally agreed that there must be (1) Cu source rocks, (2) liberation of Cu from those source rocks by a hot sedimentary brine, (3) migration of the Cu-bearing brine, (4) host rocks, (5) trap, and (6) seal rocks that are confining beds for the upward seeking metals-transporting brine. The trap must be a chemical redox trap and a physical trap.

Cu source rocks are generally red beds, and they should contain labile or chemically unstable minerals with a small amount of Cu. The labile minerals, such as hornblende or pyroxene, dissolve during early red-bed diagenesis. The Cu is relocated within ferric-oxide coatings, which hold the Cu until the coatings recrystallize to hematite or until brine moves into the source rocks and mobilizes the Cu. Alternative source rocks are flood basalts, such as with the Keweenawan section of northern Michigan and the Rotliegendes volcanics in the Kupferschiefer. Additional alternative source rocks can be the basement rocks of the basins containing the deposits.

![Figure 61. Retrograde (?) exsolution textures among bornite (pink), chalcopyrite (light yellow), and digenite (purplish gray). Chalcopyrite and digenite form exsolved laminae along bornite cubic crystal directions, and there is an outer continuous rim of digenite on the bornite grains. Royal blue color is later supergene covellite. Sample is from the Snowstorm sandstone Cu deposit, Idaho. Photomicrograph in reflected plane-polarized light in oil immersion. The identical texture has been observed at Fungurume, Democratic Republic of Congo, and White Pine. Grace and Putnis (1976) produced chalcopyrite and digenite from formerly homogeneous bornite by simple heating, and Kosyak (1969) produced similar chalcopyrite laminae in bornite on the electron microprobe adjacent to where he used high beam currents in analyzing formerly homogeneous bornite, so retrograde metamorphic origin is uncertain.](image-url)
Liberation by dissolution of Cu is by hot sedimentary brines. The Cu in the ferric-oxide coatings is mobilized as chloride complex ions (Rose, 1976). These brines have been observed in fluid inclusions in the Central African Copperbelt and at Spar Lake in ore-stage authigenic gangue minerals (Annels, 1989; Hayes, 1990; Dewaele and others, 2006; El Desouky and others, 2008a, 2008b, 2009). The brines had salinities of at least 18 to 25 eq. wt. % NaCl and transported Cu at temperatures greater than 220 °C to as low as perhaps 100 °C. Because the deposits are found associated only with thick packages of red beds, typically greater than 1 km, the heat is likely supplied by deep burial of the source rocks. There is no reason to think that magmatic heat is involved in any way; Hayes and others (2012) found that fluid inclusion volatiles from the ore stage at Spar Lake, particularly helium isotopes, HCl/HF ratios, and nitrogen/argon ratios, “give no evidence of magmatic or mantle involvement.” Rose (1976, 1989) suggested that the subsurface brines are hematite-stable albeit with intermediate oxidation states. The source(s) of salinity for the brine is known, with certainty, probably only for Lisbon Valley, where it was dissolved Paradox evaporites (Breit and Meunier, 1990). Elsewhere, the salinity sources are not known, although they too were probably dissolved evaporites. However, it is also possible that lacustrine connate brines, particularly the Great Salt Lake-type of Hardie and Eugster (1970), or Kupferschiefer-type of Eugster (1985, 1989), could act as sources of salinity for metals-transporting solutions. The Great Salt Lake-type brines are near saturation with respect to Na-sulfate salts such as glauberite or thenardite (Hardie, 1968). Brines with chemistry similar to these are the possible metals-transporting solutions for the Kupferschiefer and, particularly, for Redstone (Northwest Territories, Canada), where glauberite has been identified in footwall red beds (R. Lustwerk, oral commun., 1986). Lustwerk and Rose (1983) demonstrated the same footwall red beds, the Redstone River Formation, to be greatly depleted in Cu, with many samples having concentrations less than or equal to 2 ppm Cu.

Transport of the Cu occurs under artesian heads (that is, by confined aquifer flow), because zoning and paragenesis indicate that Cu-rich, hematite-stable brines have entered the host rocks from below. Almost every other aspect of fluid and metal migration remains conjectural. Hitzman and others (2005) listed possible causes for the migration as (1) purely compaction-driven groundwater flow, (2) topographically driven groundwater flow (Brown, 2005, 2009), (3) topographically driven flow with convection or forced convection, and (4) thermohaline (buoyancy)-driven free convection induced by salt dissolution (Hanor, 1987; Evans and Nunn, 1989; Gvirtzman and others, 1997). In addition, there is also the possibility of pure thermally driven free convection (Raffensperger and Garven, 1995a, 1995b). Thermally driven free convection induced by igneous intrusion (Norton and Knight, 1977) can probably be eliminated from consideration because there appears to be no association between SSC deposits and intrusive rocks. Also to be added to the above is the fluid flow system that has been proposed for the Kupferschiefer by Blundell and others (2003), which is termed “seismic pumping.”

Considerable disagreement exists on what type of groundwater flow system may have driven Cu-transporting groundwater to the deposits. An early paleohydrologic model for transporting Cu to the White Pine deposit was constructed by White (1971), and it was later supported by and expanded upon by Brown (2005, 2009). White (1971) favored topographically driven groundwater flow to the deposit site at the time of the inversion of the Midcontinent Rift basin about 30 to 50 million years after deposition of the Nonesuch Formation reduced-facies host rocks. Brown (2009) also called upon topographically driven groundwater flow, but he believes that it took place earlier in the geologic history, essentially at the time of diagenetic deposition of Fe oxides to produce the Copper Harbor red beds and within a few million years after deposition of the Nonesuch Formation host rocks. Jowett (1986a) suggested that thermally driven free convection operated over the Early to Middle Triassic to move Cu from the Rotliegendes source red beds to mineralize the Late Permian reduced-facies host rocks of the Polish Kupferschiefer. Alternatively, Blundell and others (2003) offered their seismic pumping model to drive Cu-transporting brines out of basement fractures toward the Polish Kupferschiefer deposits. Perhaps the best of the paleohydrologic models to date is that of Swenson and others (2004) for the White Pine deposit. They demonstrated that a compaction-driven groundwater flow system during sedimentational loading during the deposition of as much as 4 km of Freda Formation atop the Nonesuch Formation host rocks, focused by a zone of thinning of the Copper Harbor red-bed aquifer, could have produced the White Pine deposit. The likely timing would have been 10 to 15 million years after Nonesuch Formation reduced-facies host rock deposition essentially during the sag stage of the Keweenawan rift basin, and the Cu concentrations in the brines could have been less than or equal to 50 ppm Cu. Hitzman and others (2010) contended that the supergiant SSC districts, the Kupferschiefer, the Central African Copperbelt, and Udokan, were probably formed over long geologic periods with repeated groundwater flow and mineralization episodes. However, the model of Swenson and others (2004) for White Pine appears to succeed with only the compaction-driven single-pass groundwater flow event. Nonetheless, a compaction-driven system, such as that modeled at White Pine, would, in many other districts, be followed by changing tectonic setting with likelihood of additional groundwater flow systems that could transport and deposit additional Cu. In Central Africa, for example, an early compaction-driven system that mineralized the reduced-facies host rocks would logically be followed by deep burial and emplacement of sour gas in arenites, which would later be mineralized to form the sandstone Cu deposits of the belt. Subsequent compressional tectonics accompanying Lufilian Orogeny might have produced the veinlet-localized stratabound Cu deposits and increased the grade of the pre-existing deposits.
The conduits or pathways followed by hot, metals-transporting brines have rarely been recognized. At local scales, they certainly include various water-escape structures (pillars) (fig. 62). At deposit scale, they probably include faults, such as at Cashin (fig. 13) and possibly Spar Lake. For the largest reduced-facies-type deposits, conduits at the deposit scale have not been identified because they probably include large volumes of coarse, siliciclastic rocks lying along the entire footwall of the orebodies. The “conduit” rocks have not been differentiated from ordinary red beds. In order to be differentiated, the subtle alteration must be recognized and mapped.

Host rocks are relatively well known, and three major variants correspond with each of the reduced-facies, sandstone Cu, and red-bed subtypes. Reduced-facies-type hosts are black shale (Kupferschiefer), dark gray to black siltstone (White Pine), dark gray dolosiltstone (Roan 2-hosted ores of Democratic Republic of Congo and Ore Shale-hosted deposits of Zambia), gray shale (Creta, Okla.), or locally, green shale or siltstone (Belt Supergroup “green bed” occurrences). In reduced-facies hosts, the major reductant is probably solid organic material. A pre-ore step in most is to create early diagenetic pyrite, which becomes a second reductant and sulfide source at the expense of some organic material.

Sandstone Cu host rocks are typically well-sorted siliclastic sandstones from a variety of deltaic topset environments, from subtidal to coastal to eolian and to fluvial. Carbonate grainstones can act the role as well (for example, Juramento, Argentina; Durieux and Brown, 2007), and grain sizes vary from conglomerate to very fine sand. Critically, these rocks at the time of arrival of the metals-transporting brine were probably reservoir rocks for petroleum, and they were gray, green, or white in color but not, at that time, red. In many places, the petroleum was probably sour gas and, in general, the petroleum was the reductant during ore deposition (equation 1). Because petroleum in some form was present in the host rocks before arrival of metals-transporting brines, some pre-ore diagenetic pyrite had probably formed, and it acted as another reductant and second sulfide source for the ore.

Host rocks for red-bed-subtype deposits were fluvial sandstones, commonly conglomeratic and containing carbonized plant fragments sized from silty partings up to logs. The plant fragments were the reductant for this deposit subtype. Because the fluvial sandstones contained the plant fragments at deposition, their early diagenesis was probably reducing, and pyrite was probably formed in proximity to the plant fragments before arrival of the metals. The pyrite, once more, was an additional reductant and sulfide source during ore deposition.

SSC deposits are found at the sites of traps that are both chemical and physical. The host rocks for SSC deposits were the chemical redox traps. Reductants in the host rocks acted as a chemical barrier for Cu moving in hot, briny, oxidizing groundwater. Physical traps for some reduced-facies deposits were the pinchouts of the sub-ore red-bed aquifers (fig. 15), where the coarse clastics onlapped basement rocks and were confined beneath the argillaceous reduced-facies host rocks. For sandstone Cu deposits, Dzhezkazgan and other deposits of the Chu-Sarysu Basin demonstrate that anticlines are a favored type of trap. However, stratigraphic or combination traps could also localize sandstone Cu deposits. Maiden (1989) recognized additional types of traps for South Australian SSC deposits. For the red-bed Cu deposits, the host rocks are classically comparable with “shoestring” sands of popular oilfield terminology. In a “shoestring” sandstone reservoir, seal rocks and a structural closure or stratigraphic trap are required, and the same is probably true of the rocks that host red-bed Cu deposits.

Outflow zones include the galena, sphalerite, and, for some, pyrite zones of deposits, particularly where there is abundant pyrite or Co-bearing pyrite. For the sandstone Cu deposits and, perhaps, red-bed Cu deposits, there may also be former red beds that are now altered to gray, green, or white in the hanging walls of the deposits. Where such rocks contain pyrite, perhaps with sphalerite and galena, they are probably outflow zone rocks.

Seal rocks or confining beds are necessary for forming SSC deposits. The seal rocks for the reduced-facies deposits are commonly the ore host rocks themselves, or, in the Kupferschiefer, the overlying Zechstein I Anhydrite unit. Seal rocks for sandstone Cu deposits are typically shales or argillites. It is likely that these are now gray or green, but prior to petroleum entrapment, they may have been red. Seal rocks for red-bed Cu deposits are also typically shale, and the same color history may be probably true for them, but in several examples the confining shales are still red.
Flow of oxidized metals-transporting brine through any rocks that were not hematitic prior to ore stage causes oxidation and reddening by hematite deposition in the rock (for example, Rote Fäule). When the groundwater contacts the reductants in black, gray, green, or white beds, Cu is deposited as sulfides by replacing pre-ore pyrite and in coupled redox reactions of the transporting solution’s sulfate, and possibly dissolved oxygen, with the reductants of the host rock and pore fluid. With flow farther into the host rocks, accompanied by further reaction with the host rock reductants, the lateral- and vertical-zoned arrangement of chalcocite, then bornite, then chalcopyrite, and then pyrite is produced.

The preceding paragraphs in this section pertain only to the hypogene genetic features of red-bed SSC deposits. Supergene enrichment of SSC deposits as chalcocite group minerals certainly occurs, but with chalcocite occurring as a common hypogene sulfide in this class of deposits, it is at places difficult to separate supergene from hypogene features. It is less difficult to recognize supergene enrichment where it takes the form of residual enrichment in Co-rich oxide zones and probably Ag-chloride zones, for example, Silver Reef, Utah.

The SSC deposits have spatial affiliations with several other types of mineral deposits. The clearest affiliations are with native copper deposits in flood basalts. Other affiliations are with the volcanic red-bed-type Cu deposits of Kirkham (1984b), sandstone-hosted U deposits, evaporite deposits, Kipushi-type Zn-(Ge-Cu-Pb±Ag) deposits, certain types of hydrothermal U-Ni-Se deposits such as Shinkolobwe, and, in places with Fe oxide-Cu-Au deposits. Among the above, the associations with deposits of native copper in flood basalts, with volcanic red-bed-type Cu deposits, and with sandstone-hosted U deposits may be associations that share some aspects of deposit genesis. The association with evaporites is assumed to stem from dissolving the evaporites to produce the high salinities for metals-transporting brines. The associations with Fe oxide-Cu-Au, Kipushi-type, and U-Ni-Se deposits are, instead, probably just shared sedimentary tectonic settings.

Exploration and resource assessments have purposes that are partly in common, finding mineral resources or noting the probable existence of mineral resources. The objective of resource assessments by government is to identify prospective terrain (1) as an aid to exploration groups, (2) as a way to assess the potential mineral endowment of the country for strategic purposes, including whether the country will be dependent on external sources of critical and strategic minerals, and (3) for land-use planning that determines whether land should be held open for multiple uses including mineral exploration and development or used in ways not compatible with mineral development. The sole objective of mineral exploration is to make a discovery of a potentially economic deposit. Although these two objectives differ in detail, the techniques employed in the initial stages of mineral exploration are similar to the techniques of resource assessment.

Kirkham (2001) and Hitzman and others (2005, p. 635) have discussed the main factors in exploration and assessment. In our own analysis (above) a SSC system must have (1) source rocks, (2) mobilization by hot sedimentary brines, (3) transport of those brines, (4) host rocks, (5) traps, both chemical and physical, and (6) seal rocks to confine the upward migrating brine. All these factors relate to the stratigraphy and history of sedimentary basins. A basin that contains source rocks and host rocks is the fundamental unit that defines a permissive tract for exploration or assessment. (1) The basin must contain a minimum of perhaps 300 m of oxidized, hematite-stable permeable source rocks, probably red beds, with a small Cu content. Flood basalts or other volcanic rocks are a positive factor as alternative Cu source rocks. (2) The basin must have undergone flow of groundwater equilibrated with hematite from an extensive catchment basin, and the groundwater must have traveled upward across strata and toward host rocks. It is a positive factor if the basin contains or contained evaporites, either continental or marine. Having the evaporites downstream from the source rocks and the host rocks is clearly favorable (as at Lisbon Valley), but with the possibility of basin inversion, evaporite presence anywhere in the section relative to source rocks and host rocks is permissive. It would be favorable if an examination of the geologic history of a basin suggests a history of halokinesis. It would be more direct to be able to document phases that mark pathways of metalliferous brine migration by the presence of alteration phases, including hematite in rocks that were not earlier red, albite, analcime, Na-sulfate minerals, riebeckite, paragonite, epidote, perhaps barite, and perhaps chlorine-bearing illite (Michalik, 1997), and it would be particularly favorable if the rocks containing the alteration phases now had low Cu contents. (3) There must have been a groundwater flow system that drives the Cu-bearing sedimentary brines from source rocks to host rocks. Various types of flow systems with various hydrologic drives are possible, but the flow system apparently had to, at the time of mineralization, operate within confined aquifers so that brines were upward-moving relative to stratigraphy. Alternatively, the brines were sourced in rocks hot enough that the density of the brines was less than the density of the resident groundwater of the rocks immediately downstream from the host rocks. Flow probably had a directional sense of dewatering the basin. Basins where there were never flows that met those criteria are unfavorable for SSC deposits. Such infertile basins are probably common. (4) The basin must contain either organic-rich impermeable sedimentary beds that could provide solid organic matter as a reductant, permeable beds with reservoired petroleum vapor and (or) liquids to act as reductant, or permeable beds with carbonized plant fragments to act as a reductant. Obviously, a basin would be most favorable if some such rocks contained local Cu mineral occurrences. (5) The basin should have had sufficient structural deformation to create traps. Minimally, that might constitute sagging of basin or sub-basin centers such that an onlap of reduced-facies host beds onto a basement high would become a stratigraphic trap. The traps could vary from that up to complex combination traps produced by pre-ore folding and faulting. (6) The basin should have had confining beds that either constitute the reduced-facies host rocks or that stratigraphically, or perhaps structurally, overlie petroleum reservoirs. This last criterion is met by nearly every studied basin.
Sediment-Hosted Stratabound Copper Deposit Model

Geological Guides

Mapping of Rock Types, Structural Geology, Minerals, Alteration, and Zoning

Mapping is by far the most important procedure in assessing the location, volume, and attributes of altered and (or) mineralized rocks and in determining the relative position of exposures of those altered or mineralized rocks within a SSC system. In exploration and assessment, all available geologic mapping is used. Where prior mapping is not available, explorationists will commonly conduct the required geologic mapping at least at reconnaissance scale. Resource assessment may require new geologic mapping in order to produce any reasonably accurate assessment.

Reconnaissance-scale geologic mapping may suffice in initial stages of exploration or assessment, and the mapping may need only identify major lithostratigraphic packages. It can be guided by the purposes of identifying the necessary elements of a SSC mineralizing system: Cu source rocks; sources for hot metals-transporting brines; conduits for metals-transporting brine migration, such as basin margins, large-scale alteration pathways, and faults; Cu host rocks; traps; and seals.

Alteration mapping may become increasingly useful, although alteration may be extremely subtle. Where wholesale replacement by albite occurs, as it does at places in the Central African Copperbelt and the Udokan system in Russia, albite alteration is easily mapped. However, although the albite zone can now be mapped in the Western Montana Copper Sulfide Belt (fig. 6), this has not proved useful in exploration to date. In frontier exploration areas, with the concept of alteration marking the path of migrating metals-transporting brines, identification of the characteristics of alteration may be a useful early task, and alteration mapping could then be quite productive. Even in better-explored systems, identification of the characteristics of alteration along migration pathways may prove useful.

Mineral Properties

A few mineral properties have emerged that can be mapped at a reconnaissance exploration or assessment scale. A notable one for sandstone Cu deposits is carbonate compositions and abundances. There is an anomalous concentration of calcite in the distal-fringe zones of the Spar Lake and other Revett Formation-hosted stratabound Cu deposits. Those calcite cements are Fe-rich through the pyrite-calcite, galena-calcite, and chalcopyrite-calcite zones of deposits. An associated Cu deposit will already have been located by the time an explorationist narrows the search enough to observe the Mn-rich calcites of the bornite-calcite zone of a deposit. The bleached rocks of chalcopyrite-ankerite zone of mineralizing systems have very sparse disseminations of ankerite that weathers much differently than the Fe-rich calcites (fig. 29).

For deposits in sandstone or siltstone host rocks, carbonate abundance, not just composition, should also be a guide if the carbonates result from oxidation of reactant natural gas, as is suspected.

Weathering products from different hypogene sulfide zones of sandstone Cu deposits can also be used to narrow the search for locating deposits. Brochantite, as opposed to malachite, is a strong indicator of the central highest-grade parts of such systems. The occurrence of Cu in Cu wad in black oxides distributed on joints and fractures is an indicator of a mineralizing system from farther away, perhaps kilometers along bedding, from centers of mineralization.

Distribution of Sulfide Minerals

Sulfide minerals in fresh rocks or their pseudomorphs in weathered outcrops are the best mappable mineralogical indicators for exploration or assessment. The strong sulfide zonation of reduced-facies SSC deposits and sandstone Cu deposits makes the sulfides and their zoning useful. The sulfides of distal-fringe zones vary a little with the subtype of the deposits.

Sandstone Cu deposits have large volumes of bleached rock (Chalcopyrite-ankerite zone of fig. 6), which extend the greatest distances surrounding orebodies. The bleached rock haloes can extend several tens of kilometers from the ore. Because the bleaching may have been caused by unrelated events, such as petroleum migration and emplacement, bleaching alone is not a vector to ore. It may, though, be a necessary effect preparatory to later ore events. Sparsely disseminated pyrite in strata that are elsewhere red beds can yield a broad halo that may extend as much as 10 km laterally along bedding from ore. The sandstone Cu deposits also have large galena- and (or) sphalerite-bearing inner halos that extend for 1–2 km laterally from ore.

For reduced-facies SSC deposits, the lateral extent of the different sulfide zones is even greater. The Pb-Zn zones of the Kupferschiefer are halos several tens of kilometers wide adjacent to the target Cu zones and separating the Cu zones from barren, pyritic zones (fig. 4) (Oszczepalski and Rydzewski, 1997). If a favored reduced-facies host bed is identified, sulfide mineral zonation, or the trace-metal chemistry that results from the sulfide mineral zonation, can lead to the Cu deposits from very great distances. The Central African Copperbelt systems unfortunately do not appear to have Pb- and Zn-enriched zones, and Co is also fairly narrowly distributed. Co does, though, form an outermost distal-fringe halo that extends 1 km or more away from ore along bedding.

Geochemical Guides

Soil-Gas Chemistry

With knowledge that sandstone Cu deposits occur where oxidizing, hot, metals-transporting brines mix and react with H2S-bearing natural gas (Hayes and others, 2012), soil-gas chemistry might be a powerful exploration
method in the search for such deposits. Fluid inclusions in ore-stage authigenic minerals of the Spar Lake deposit contained high concentrations of CH₄ and hydrogen in the distal fringe, and they contained high concentrations of sulfur dioxide in rich ore. It is suspected that fluid inclusions in more distal minerals might contain even higher concentrations of CH₄, as well as anomalous concentrations of H₂S that constitute unreacted or only slightly reacted sour gas. These gases can be measured in soil gases (McCarthy and Reimer, 1986). Carbon disulfide (CS₂), which might occur at detectable concentrations within unreacted sour gas, and Hg vapor may also be substantial (fig. 39). Weathering of ores destroys the hypogene carbonate and sulfide minerals that host the gas-filled inclusions. Localization of ores along faults, with locally deep weathering to depths of greater than 600 m, provides conduits for escape of the gases liberated from the fluid inclusions.

Trace-Element Geochemistry

From geochemical surveys of soil and stream sediment, it is well known that Cu, itself, is an excellent pathfinder element for these deposits in many different weathering and climatic conditions (for example, Payne, 1971). Additional ore elements are Ag and Co. Ag is useful in the sandstone Cu mineralizing system of the Revett Formation, because anomalies with Cu and Ag are distinguishable as a SSC system as distinct from anomalies associated with other types of deposits (Harrison and others, 1986a). Other potential pathfinder elements include Pb, Zn, As, Hg, Mn, and Ba. Soil geochemical surveys have proven effective in Botswana even where the bedrock that hosts the deposits are covered by tens of meters of Kalahari sands (Van Der Heever and others, 2009).

Supergene Enrichment and Leached Capping Characteristics

In many climatic zones, supergene enrichment is negligible for SSC deposits, and either Cu-oxides or relict sulfides are found in weathering-zone outcrops of deposits. In central Africa, heterogenite with or without lesser malachite, forms “Co caps” (Gauthier and Deliens, 1999). In Co caps, the amounts of heterogenite make it unmistakable as an earthy to dense, botryoidal black oxide. Leached cappings in central Africa have been only poorly described, although it is clear, particularly where carbonates or anhydrite with Cu-sulfides formerly cemented sandstones, that highly porous, brown, goethitic (?) sandstone results from weathering. Red hematite and beer-bottle limonites are not described from this rock, but kaolinite is a major component.

Weathered deposits with rich concentrations of malachite are known from the Democratic Republic of Congo, and specimens of malachite from these occurrences are famous worldwide. Some such deposits do not overlie sulfide deposits. Some are known to have been below Roan 2-hosted stratiform Cu ore, but others occur below thrust faults with Roan 2-hosted ore within the allochthonous plate (for example, within Kundulungu beneath the Kolwezi klippe) or associated with extrusion faults (probably halokinetic). Some occur randomly within tectonic or halokinetic breccias stratabound (Francois, 1992). Supergene enrichment is obvious in these cases. Native copper and covellite are additional common supergene zone products in the Central African Copperbelt, but they probably do not constitute major enrichments to hypogene grades.

Cu-Oxide Minerals

Malachite is the most widespread Cu-oxide mineral from the weathering of SSC deposits. In middle-latitude temperate, relatively humid climates, brochantite predominates instead, as the weathering product of the richest ore zones. Chryscolla is generally present in low abundances with either of the above. Copper wad may be present in associated black oxides, discretely along joint faces, or as weathered veinlets. Beer-bottle limonite is the most common of Cu-oxide minerals in sunny exposures where it commonly directly pseudomorphs earlier grains of disseminated sulfide cements.

Geophysical Guides

Basin Structure

Basin margins, where reduced-facies host rocks onlap basement rocks (fig. 10), are favorable sites for ore deposition. These can be detected by magnetic and gravity surveys and, most reliably, by seismic surveys. Regional-scale magnetic and gravity surveys are first-order geophysical methods in exploration for reduced-facies-type deposits. They are also first-order data for assessments wherever target stratigraphic intervals are covered.

In basins that have not been dramatically deformed post-ore and where SSC systems are already recognized (for example, Chu-Sarysu Basin, Kazakhstan; Permain rocks of Kansas-Oklahoma-Texas, United States), seismic reflection or refraction surveys in search of anticlines, stratigraphic traps, and combination traps, combined with more commonly used mineral exploration methods, should be effective at finding additional sandstone Cu deposits.

Red Beds and Beaching

Bleaching of red beds detectable by remote sensing of colors using Landsat imagery or of ferric oxides using ASTER or AVIRIS imagery can narrow the search for sandstone Cu deposits or limit the areas for assessments. Only the bleached former red beds can be mineralized by later ore systems (MacIntyre, 2006).
Direct Geophysical Targeting

Induced Polarization (IP) may be effective in direct sensing of deposits. The conductive sulfide signature of sandstone Cu deposits can be found with IP for rocks that contain no carbonaceous or very pyritic shales. The key to success of the method is that the unmineralized, non-graphitic and non-sulfidic rocks will be extremely resistive, whereas areas of SSC, though they may not be truly conductive, will be relatively non-resistive. The IP can be effective for exploration in local situations, such as in the Democratic Republic of Congo part of the Central African Copperbelt, where the sulfide orebodies may be shallowly covered by heterogeneous breccias that crop out between exposures of Kundulungu Group. However, in areas with Kundulungu sediments in the subcrop, IP will likely be ineffective at seeing through the thick intervals of carbonaceous and pyritic shale cover.

Attributes Required for Inclusion in a Permissive Tract

At the broadest level, permissive tracts for SSC deposits can be defined to include the presence of carbonaceous and (or) pyritic beds, and (or) beds containing trapped petroleum, or beds that formerly contained trapped petroleum. These host rocks should be located upsection from red beds thicker than 300 m or a former red-bed unit and (or) flood basalts, or laterally within 10 km of that thickness of Cu source rocks. A permissive tract for SSC deposits is outlined by delineating a geographic area on a geologic map that includes chemically reduced host rocks and generally stratigraphically lower red beds or former red beds and (or) flood basalts that are a minimum of 300 m thick. A thickness of 300 m is considered the approximate minimum needed to generate the required metals-rich brine, and greater thicknesses can be considered more favorable. The tract may include known SSC deposits and prospects of any of the three subtypes. Current economics of metal production will define a maximum allowable thickness of cover rocks overtop the host rocks. The tract outline approximately parallels the outline of the sedimentary basin, as depicted on the scale of the geologic map available for tract delineation. The presence of evaporite beds in the sequence and faults at high angles to bedding are particularly favorable. Tracts should include only those areas underlain by chemically reduced host rocks, provided that there is good knowledge of any downsection source rocks that demonstrates that no bleached former red beds are present.

Factors That Enable Favorability Zonation of Permissive Tracts

Areas with known SSC deposits within a permissive tract define locations having distinctly better favorability than other parts of that tract. As above, faults at steep angles to sedimentary layering define more favorable parts of tracts. Any parts of a basin where reduced-facies host beds onlap basement rocks are favorable. In terrains that have known sandstone Cu deposits, anticlines, areas with known stratigraphic traps, and those with known combination traps are more favorable than other parts of the tract (for example, Hitzman and others, 2005, fig. 8; the Paradox Basin). Areas of natural-gas production are also more favorable within basins that have known sandstone Cu deposits.

Basin margins are more favorable than basin centers (see, for example, Hitzman and others, 2005, fig. 6B showing the Zechstein Basin). Structurally high areas within central basins are more favorable than their surroundings, such as the structural high with Rote Fäule alteration near the Pomeranian Coast of the Zechstein basin (Oszczepalski and Rydzewski, 1997, Plate XVI). This structurally high zone probably extends westward across southern Denmark and northernmost Germany as the Ringkobing-Fyn High, then into the North Sea to become the mid-North Sea High defined from petroleum exploration.

In basins with reduced-facies-type mineralization, areas where host beds contain only pyrite, or even contain Zn- or Pb-bearing sulfides, are unfavorable, although a transition from Pb-rich rock to Cu ore can occur over a short distance In basins with reduced-facies-type mineralization, areas where host beds are known to be secondarily hematized or otherwise altered in ways analogous with sandstone roll front U deposit alteration; that is, rocks where Cu is depleted relative to background are unfavorable. Again, however, a transition from altered rock to Cu ore can occur over a short distance.

Burial histories derived from petroleum exploration such as Lopatin modeling can help subdivide permissive tracts into more favorable and less favorable areas. Only those areas within and adjacent to parts of the tract that underwent rapid subsidence and burial are favorable. Parts of basins that subsided slowly and are distant from fast subsiding parts, the shallow platforms, are unfavorable.

Factors That Influence Undiscovered Deposits Estimates

Different basins typically have characteristic distributions of Cu grades in their deposits, especially for deposits of the same subtype. For example, Mesoproterozoic-hosted reduced-facies deposits in south-central China have grades that average 1.04 percent (six deposits). The ores hosted in Neoproterozoic shale of the Zambian Copperbelt average 2.11 percent Cu (13 deposits or prospects, all within the Copperbelt Orebody Member). It is unlikely a 2.1 percent shale-hosted orebody will be found in the south-central China province, except if considering only a small tonnage. Cu grades are consistent from deposit to deposit within a basin, although they are different from one basin to the next. Similarly, sandstone Cu deposits of the Udokan sequence are higher grade than sandstone Cu deposits of the Revett Formation in the Belt Supergroup. It cannot be expected that a new Revett-hosted orebody will have 1.3 percent Cu, which characterizes the Udokan. Together, Spar Lake, Rock Creek, and Montanore,
in the Revett, average 0.75 percent Cu (Boleneus and others, 2006). The same observations hold for byproduct metals, Co and Ag. It cannot be expected that the next-discovered Udokan Series-hosted orebody will have 40 g/t Ag. Udokan contains 12.6 g/t Ag (Vladimir Chechetkin, Committee on Geology and Utilization of Mineral Resources of the Chita Region, Russia, written commun., 2009). But, in the Revett, the Spar Lake, Rock Creek, and Montanore deposits have 46, 73, and 75 g/t, respectively, and together they average 68 g/t (Boleneus and others, 2006). It can be expected that the next-discovered economic Revett-hosted deposit will have an Ag grade of more than 40 g/t. Providing that existing grade data permits it, basin-unique grade distribution models should be used in assessments. The statements above hold only as deposit-scale observations. Within deposits, there is commonly a bed-by-bed grade variation. The average grade of the White Pine deposit is 1.2 percent Cu, but the average grade of the “Domino” and “Lower Transition” beds within the deposit is 2.5 percent (fig. 12). Deposit average grades, particularly for reduced-facies-type deposits, may reflect the need for dilution with waste rock in order to extract a minimum thickness that can be mined underground by a large-tonnage method. Actual mineralized intercepts may be considerably higher grade, but the average reflects blending that rock with nearly barren waste rock to attain a mineable thickness. Within reduced-facies-type deposits particularly, individual sample grades can be quite high, but the challenge may be to find a sufficient thickness to mine.

Tonnage distributions are not similarly basin-unique. These distributions are geologically controlled and not simply an artifact from reporting only open-pit resources. The tonnages probably have more geologic cutoffs, such as structural geologic boundaries and outcrop boundaries.

Deposit densities are nearly impossible to estimate for reduced-facies-type SSC deposits. Minor continuations of mineralization are so widespread that it becomes an almost totally arbitrary decision where one deposit ends and the next begins (see the maps of Oszczełapski and Rydzewski, 1997). Therefore, deposit-density modeling methods as a control on undiscovered deposit number estimates do not work with these layer-form ores.

Reported tonnages, particularly, are probably incompletely representative of entire deposits. Geological limits, such as structural geologic boundaries or grade cutoff boundaries, would result in larger tonnages than are reported for many active mines and for many prospects. Instead, many of the reported tonnages from Democratic Republic of Congo, in particular, are only the tonnages considered accessible in an open-pit mine. Geologically, the deposits continue to greater depth, which is beyond the economic limit of mining and where the continuity of the deposit is not known. Nonetheless, through time, reserve growth is likely. This holds as well for certain deposits of the Kupferschiefer, where the Lubin-Sierozowice mining complex has reported larger total resources through time by mining the same single deposit to greater and greater depth as new technology and Cu prices allowed. The Lubin-Sierozowice-Polkowice-Rudna mines have reached depths of 1,250 m without encountering any limitations on their room-and-pillar methods because of the depth, and there appear to be no important difficulties in continuing farther down-dip. Either wallrock temperature or the saline and corrosive groundwater may eventually limit the operation (A. Piętrzyński, oral commun., 2001).

The degree of structural disruption of the rocks is a critical geologic factor in exploring for SSC deposits. Even small-offset faults can substantially influence the economics, because, in these layer-form ores, the faults cause considerable tonnages of waste rocks to be mined, diluting the grades. Dips between about 15 and 40 degrees are problematic because rubber-tired underground equipment struggles or fails to climb the dips (Derek Feeback, oral commun., 2007), and the broken ore will not gravity-feed in a sublevel stoping or a block caving operation. Deposits that are nearly flat-lying and undisrupted are at a great economic advantage. The Creta reduced-facies-type deposit was an example of that. The ore was typically just 21 cm thick (Dingess, 1976) at an average grade of 1.9 percent Cu, but the beds dip less than 2° and the topographic relief is low. Also, blasting was seldom required in the generally soft sedimentary rocks. These factors resulted in an area of more than 8 km² being strip-mined before overburden thicknesses and Cu prices forced the closure of the mine.

**Geoenvironmental Features and Anthropogenic Mining Effects**

All SSC deposits contain zones of chalcocite and other Cu,S minerals, bornite, chalcopyrite, pyrite, and subordinate galena and sphalerite. The chalcocite and bornite in the high-grade zone are fairly stable under oxidizing conditions. Pyrite is typically found only in trace amounts in the high-grade zone, but its content increases outward as Cu grade decreases. Therefore, environmental issues associated with the oxidation of pyrite and the subsequent potential production of acid mine drainage and leaching of metals can be minimized if mining plans avoid the outboard pyrite-containing zones. Two types of host rocks are associated with these deposits: low-energy marine or lacustrine siltstones, shales, and carbonate rocks; and high-energy continental sandstones, arkoses, and conglomerates (Cox and others, 2003) corresponding to the reduced-facies and sandstone Cu subtypes. The calcite present in some host rocks will help mitigate against the development of acid mine drainage. In addition to pyrite as a potential environmental concern, minerals such as galena, sphalerite that commonly contains Cd substituting for Zn, and As-bearing minerals such as tennantite, enargite, luzonite, and arsenopyrite can be disturbed during mining or concentrated during ore processing and potentially weather and release harmful trace elements into the environment. The geoenvironmental characteristics of sediment-hosted Cu deposits have been previously addressed by Lindsey and others (1995).
Pre-Mining Baseline Signatures in Soil, Sediment, and Water

Mine permitting requires an estimate of the pre-mining natural background conditions to serve as a goal for post-mining reclamation. For all three subtypes of SSC deposits, soil and stream sediments may contain anomalous abundances of Cu, Pb, Ag, and possibly As, Hg, and Zn in the vicinity of the deposits (Lindsey and others, 1995). For example, in the vicinity of SSC deposits in the Democratic Republic of Congo and Zambia, soils are Cu-rich such that Cu-resistant and Cu-accumulating plants have replaced normal vegetation (Reilly, 1967; Reilly and Stone, 1971; Malaisse and others, 1978). Upper canopy trees cannot grow in the Cu-rich soils yielding “copper clearings.” Stream sediments collected in the Kafue River upstream from mining contain between 15 and 146 ppm Cu (specify the element), some of which are just below the probable effect concentration (PEC) of 149 ppm (table 3) (MacDonald and others, 2000; Pettersson and others, 2000; Pettersson and Ingri, 2001). Soil and sediment associated with some Revett deposits in Montana contain as much as 2,000 ppm Cu, greater than 150 ppm Pb, and greater than 500 ppb Ag (Cazes and others, 1981; Wells and others, 1981).

In contrast to some of the sediment and soils with anomalous metal concentrations, background surface waters of the Kafue River, upstream from the mining area in Zambia, contain low Cu (less than or equal to 3 µg/L) and other trace elements (table 4) (Norrgren and others, 2000; Pettersson and Ingri, 2001). Also, the draft environmental impact statement for the Montanore Project, a proposed mine in the Rock Creek-Montanore sandstone Cu-type deposit in Montana, states that surface-water quality in streams and lakes near the proposed mine is excellent. The concentrations of total suspended solids, total dissolved solids, major ions, nutrients, and metals are generally low and frequently at or below detection limits even downstream from the deposit. The surface waters generally contain Fe, Cu, and Ag in low concentrations, well below environmental guidelines. The surface waters tend to be slightly acidic likely from organic acids and dissolved carbon dioxide. The hardness values in streams are commonly less than 30 mg/L (U.S. Forest Service and others, 2009).

Past and Future Mining Methods and Ore Treatment

Most reduced-facies-subtype deposits are developed by underground mining methods, predominantly by room-and-pillar to depths of 1,000 m, although in a few old mines, longwall methods were used. Ore is processed by grinding and flotation and the concentrate is smelted (E&MJ, 1979a, 1986a, 1986b). In the Mansfeld district in Germany, the volatile compounds and fine particles produced by smelting ore were recovered by suspending the material in water, followed by filtering and dewatering (Matheis and others, 1999). The resulting flue dust material was called ‘Theisenschlamm,’ and at times, it was reprocessed to recover metals. In addition to underground mining, ore such as in Zambia has been mined by open-pit methods and processed by a combination of flotation-smelting and solvent-extraction electrowinning (SX-EX) methods (E&MJ, 1979a). As for underground mining, at least one mine in Zambia performs acid leaching of the ore underground (Sracek and others, 2010). Also, in situ bioleaching was proposed as an effective means of recovering Cu from chalcocite ores from White Pine and Kona, Mich. (Johnson and others, 1988). To date, the bioleaching of these ores has taken place only at lab-bench scale.

Large red-bed-subtype deposits are mined by open-pit methods, whereas smaller deposits are generally mined by underground methods via adits, inclines, and shafts. Ore is processed by grinding and flotation, and the concentrate is shipped to smelters (Soule, 1956; Talbott, 1974). Future processing of ore may occur by heap leach-SX-EW on site (Mining Record, 1995).

Sandstone Cu deposits are mined by underground room-and-pillar methods with ore processed by grinding and flotation near the mine site and concentrates shipped to smelters (E&MJ, 1979b, 1982; U.S. Forest Service and others, 1992). Deposits in the Lisbon Valley district in Utah have been mined by open-pit operations in addition to underground methods. Modern ore processing was done by SX-EW leaching (Krahulec, 2006).

Volume and Footprint of Mine Waste and Tailings

The volume of mine waste and tailings produced from a deposit depends on the deposit size, depth, and geometry; the ore grade; and mining method. Reduced-facies-type deposits are generally high tonnage (median, outside of the Central African Copperbelt, 27 million tonnes) with a high Cu grade (median 1.38 percent) compared to other subtypes and are mined mostly underground (Lindsey and others, 1995; Cox and others, 2003). The amount of waste produced by deposits is exemplified by the Mansfeld district in Germany. Each tonne of ore in the Mansfeld district yielded 35–38 kg of metal, 740 kg of silicate slag, 20 kg of flue dust, and approximately 1,500 m³ of furnace-top gas (Matheis and others, 1999). Waste rock and slag from the Mansfeld district were piled in large heaps up to 9,000,000 m³ (Schubert and others, 2003). Strauch and others (2001) estimated that approximately 2,000 heaps of mining and smelting waste of mainly low-grade ore, host rock, and slag occur in this former mining district. After metal recovery from the flue dust ceased in 1978, more than 300,000 t of the material were deposited on the heaps of slag and mining debris (Matheis and others, 1999). One disposal site for flue dust in the same district is approximately 225,000 m³ (Schubert and others, 2001, 2003). The flue dust is considered the most important source of pollution in the Mansfeld district because of its elevated concentrations of As, Cd, Cr, Cu, Pb, and Zn (table 5) (Strauch and others, 2001).
Table 3. Concentrations of metals in stream, wetland, and lake sediment downstream from sediment-hosted Cu deposits.

[RF, reduced-facies-type deposit; Al, aluminum; As, arsenic; Cd, cadmium; Co, cobalt; Cr, chromium; Cu, copper; Fe, iron; Ni, nickel; Pb, lead; S, sulfur; Zn, zinc; %, percent; mg/kg, milligrams per kilogram; -, no data; <, less than]

<table>
<thead>
<tr>
<th>Mining area (deposit model)</th>
<th>Source Description</th>
<th>Al (mg/kg)</th>
<th>As (mg/kg)</th>
<th>Cd (mg/kg)</th>
<th>Co (mg/kg)</th>
<th>Cr (mg/kg)</th>
<th>Cu (mg/kg)</th>
<th>Fe (%)</th>
<th>Ni (mg/kg)</th>
<th>Pb (mg/kg)</th>
<th>S (%)</th>
<th>Zn (mg/kg)</th>
<th>Reference</th>
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<tbody>
<tr>
<td>Chambishi mine, Zambia (RF)</td>
<td>stream upstream of mining area (Kafue River)</td>
<td>5.5–5.9</td>
<td>-</td>
<td>30–46</td>
<td>-</td>
<td>15–146</td>
<td>2.4–4.9</td>
<td>-</td>
<td>-</td>
<td>0.35–0.65</td>
<td>-</td>
<td>-</td>
<td>Pettersson and others (2000); Pettersson and Ingri (2001)</td>
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<td>9.0–9.7</td>
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<td>789–1,030</td>
<td>-</td>
<td>11,028–12,855</td>
<td>3.8</td>
<td>-</td>
<td>-</td>
<td>0.20–0.80</td>
<td>-</td>
<td>-</td>
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<td>374</td>
<td>-</td>
<td>1,250</td>
<td>1.9</td>
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<td>-</td>
<td>-</td>
<td>-</td>
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</tr>
<tr>
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<td>wetland within wetland (New Dam)</td>
<td>1.7–2.3</td>
<td>&lt;0.1–0.8</td>
<td>42–8,439</td>
<td>9–130</td>
<td>109–12,112</td>
<td>3–220</td>
<td>9–75</td>
<td>0.10–1.05</td>
<td>7–125</td>
<td>-</td>
<td>-</td>
<td>von der Heyden and New (2004b)</td>
</tr>
<tr>
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<td>stream downstream of Helbra smelter</td>
<td>-</td>
<td>487</td>
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<td>121</td>
<td>198</td>
<td>5.1</td>
<td>0</td>
<td>567</td>
<td>-</td>
<td>6,429</td>
<td>-</td>
<td>Schreck and others (2005)</td>
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<tr>
<td>Mansfield district, Germany (RF)</td>
<td>stream downstream of Eisleben smelter</td>
<td>-</td>
<td>56</td>
<td>-</td>
<td>92</td>
<td>1,054</td>
<td>2.9</td>
<td>56</td>
<td>1,034</td>
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<td>3,796</td>
<td>-</td>
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<td>lake downstream of mining and smelting district</td>
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<td>2–1,800</td>
<td>&lt;2–55</td>
<td>11–47</td>
<td>65–92</td>
<td>21–2704</td>
<td>2.2–3.6</td>
<td>36–71</td>
<td>27–4,017</td>
<td>0.4–2.3</td>
<td>88–35,500</td>
<td>Becker and others (2001)</td>
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<tr>
<td>Probable Effect Concentrations (PEC)</td>
<td>-</td>
<td>33</td>
<td>4.98</td>
<td>-</td>
<td>111</td>
<td>149</td>
<td>-</td>
<td>48.6</td>
<td>128</td>
<td>-</td>
<td>459</td>
<td>-</td>
<td>MacDonald and others (2000)</td>
</tr>
</tbody>
</table>

1Range in typical concentrations for samples between 0 and 510 centimeter depths.
Table 4. pH and dissolved metal content of waters from sediment-hosted copper deposits. Drinking water and aquatic ecosystem guidelines also listed.

[RF, reduced-facies-type deposit(s); SS, sandstone copper-type deposit(s); Al, aluminum; Cd, cadmium; Co, cobalt; Cr, chromium; Cu, copper; Fe, iron; Mn, manganese; Mo, molybdenum; Ni, nickel; Pb, lead; S, sulfur; Zn, zinc; μg/L, micrograms per liter; <, less than; -, not reported or not analyzed; m, meters; km, kilometers; mg/L, milligrams per liter; Al, Fe, and pH are non-priority pollutants]

<table>
<thead>
<tr>
<th>Mining area</th>
<th>Water type</th>
<th>Water source</th>
<th>pH</th>
<th>Al (μg/L)</th>
<th>Cd (μg/L)</th>
<th>Co (μg/L)</th>
<th>Cr (μg/L)</th>
<th>Cu (μg/L)</th>
<th>Fe (μg/L)</th>
<th>Mn (μg/L)</th>
<th>Mo (μg/L)</th>
<th>Ni (μg/L)</th>
<th>Pb (μg/L)</th>
<th>S (mg/L)</th>
<th>Zn (μg/L)</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>White Pine, Michigan, USA (RF)</td>
<td>surface</td>
<td>water, smelter slag, and mine mine</td>
<td>7.2</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>Lindsey and others (1995)</td>
</tr>
<tr>
<td>Spar Lake, Montana, USA (SS)</td>
<td>surface</td>
<td>water surface</td>
<td>7.2–7.6</td>
<td>&lt;1</td>
<td>10–280</td>
<td>&lt;10</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>U.S. Forest Service and others (1992)</td>
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<td>Spar Lake, Montana, USA (SS)</td>
<td>surface</td>
<td>water surface</td>
<td>7.5–8</td>
<td>&lt;1</td>
<td>&lt;10</td>
<td>&lt;10–20</td>
<td>-</td>
<td>-</td>
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<td>-</td>
<td>-</td>
<td>-</td>
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<td>U.S. Forest Service and others (1992)</td>
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<td>Montanore, Montana, USA (SS)</td>
<td>surface</td>
<td>settling pond</td>
<td>7.7–10.9</td>
<td>-</td>
<td>&lt;10–20</td>
<td>-</td>
<td>-</td>
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<td>-</td>
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<td>-</td>
<td>-</td>
<td>-</td>
<td>U.S. Forest Service and others (1992)</td>
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<tr>
<td>Spar Lake, Montana, USA (SS)</td>
<td>surface</td>
<td>tailings impoundment outflow</td>
<td>7.5</td>
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<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
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<td>U.S. Forest Service and others (2009)</td>
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<td>surface</td>
<td>tailings pile</td>
<td>7.5</td>
<td>-</td>
<td>12</td>
<td>6</td>
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<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>600</td>
<td>-</td>
<td>-</td>
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<td>surface</td>
<td>wetland inflow</td>
<td>7.3–8.5</td>
<td>-</td>
<td>&lt;10–470</td>
<td>&lt;10–380</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>&lt;30–510</td>
<td>-</td>
<td>-</td>
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<td>von der Heyden and New (2005)</td>
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</tr>
<tr>
<td>Chambishi mine, Zambia (RF)</td>
<td>surface</td>
<td>wetland outflow</td>
<td>7.2–7.8</td>
<td>-</td>
<td>&lt;10–940</td>
<td>&lt;10–50</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>100–290</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>von der Heyden and New (2005)</td>
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<tr>
<td>Chambishi mine, Zambia (RF)</td>
<td>ground</td>
<td>tailings basin</td>
<td>6.0–6.2</td>
<td>20–50</td>
<td>1–2</td>
<td>24–26</td>
<td>1–6</td>
<td>2–19</td>
<td>1,680–2,420</td>
<td>990–1,070</td>
<td>5–10</td>
<td>0.3–0.4</td>
<td>40–91</td>
<td>-</td>
<td>-</td>
<td>von der Heyden and New (2004a)</td>
</tr>
<tr>
<td>Copperbelt district, Zambia (RF &amp; SS)</td>
<td>surface</td>
<td>upstream of mining upstream of mining</td>
<td>7.2–8.1</td>
<td>3–8</td>
<td>0.2–2</td>
<td>1–3</td>
<td>78–254</td>
<td>6–17 ND</td>
<td>0.1–1</td>
<td>-</td>
<td>1–2</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>Pettersson and Ingri (2001), Pettersson and others (2000)</td>
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<tr>
<td>Copperbelt district, Zambia (RF &amp; SS)</td>
<td>surface</td>
<td>Kafue River downstream mining area</td>
<td>-</td>
<td>21</td>
<td>54</td>
<td>15</td>
<td>18</td>
<td>122</td>
<td>6.8</td>
<td>7.6</td>
<td>0.5</td>
<td>NR 53</td>
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<td>2.0 km from smelter</td>
<td>-</td>
<td>2–13</td>
<td>-</td>
<td>16–62</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>37–72</td>
<td>-</td>
<td>-</td>
<td>Pile and others (1999)</td>
</tr>
<tr>
<td>Glogow, Poland (RF)</td>
<td>ground</td>
<td>2.0 km from smelter</td>
<td>-</td>
<td>1–7</td>
<td>-</td>
<td>7–31</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>17–62</td>
<td>-</td>
<td>-</td>
<td>Pile and others (1999)</td>
</tr>
<tr>
<td>Glogow, Poland (RF)</td>
<td>ground</td>
<td>5.5 km from smelter</td>
<td>-</td>
<td>1–8</td>
<td>-</td>
<td>5–16</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>26–40</td>
<td>-</td>
<td>-</td>
<td>Pile and others (1999)</td>
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<tr>
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<td>mine</td>
<td>6.9–7.7</td>
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<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
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<td>-</td>
<td>18,000</td>
<td>3,300</td>
<td>3</td>
<td>19,000</td>
<td>&lt;50</td>
<td>16,000</td>
<td>-</td>
<td>2,400</td>
<td>2,111,000</td>
<td>-</td>
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<td>Schubert and others (2003)</td>
<td></td>
</tr>
</tbody>
</table>

Drinking water guideline
- - - 3 - 50 2,000 - 400 70 70 10 - - World Health Organization (2006)

Aquatic ecosystem
- - - 750 2 - 570 13 - - 470 65 - 120 U.S. Environmental Protection Agency (2006)

Aquatic ecosystem chronic toxicity
- - 6.5–9 87 0.25 - 74 9 1,000 - - 52 2.5 - 120 U.S. Environmental Protection Agency (2006)

1Dissolved concentrations were calculated from total concentration and percent suspended.

2Freshwater criterion for Cd, Cr (assume Cr (III)), Cu, Pb, Ni and Zn are hardness dependent and calculated based on a hardness of 100 mg/L CaCO₃.
Reduced-facies-type deposits. The mining of reduced-facies deposits has produced tailings, slag, and flue dust as waste products. Tailings from two deposits, Chambishi and Nkana, in Zambia in the Central African Copperbelt generally contain 1 to 4 wt. % Fe, 0.1 – 0.3 wt. % Cu, and 300 to 800 mg/kg Co. A hardpan layer in one tailings pile contains substantially higher concentrations of these three metals (Sracek and others, 2010). Tailings from the Rudna, Polkowice, and Lubin mines in Poland are all dumped in a common tailings pond. The tailings from the Rudna mine are reported to contain approximately 0.2 percent Cu (E&MJ, 1986b). Tailings from the White Pine mine in the United States contain comparable concentrations; Cu ranges from less than detection to 0.17 wt. % (U.S. Environmental Protection Agency, 1994; Williams and others, 2002). The concentrations of Cu for tailings from all three countries are consistent.

Slag and flue dust from smelting in the Mansfeld mining district contains significant concentrations of Cu, Pb, Zn, Cd, and As (table 5); the flue dust is highly enriched in As, Cd, Cu, Pb, and Zn compared to the slag (Jahn and others, 1997; Matheis and others, 1999; Wennrich and others, 2001). The flue dust is extremely fine-grained and contains major concentrations of metals with as much as 193,000 mg/kg Zn, 143,000 mg/kg Pb, 8,100 mg/kg Cd, and 6,674 mg/kg As (Wennrich and others, 2001; Schubert and others, 2003) (table 5). This material was reprocessed from 1904 to 1978 to extract Zn, Pb, Ag, Re, Se, Sn, Tl, and Ge (Matheis and others, 1999).

Waste slag from the Głogów smelters in Poland contains between 0.5 and 0.7 percent Cu (E&MJ, 1986a). These are higher concentrations than reported for the Mansfeld district slag (0.16 wt. 5, table 5). Kucha and Cichowska (2001) report 6 and 40 mg/kg Ag, and 0.008 and 0.009 mg/kg Au in shaft and converter slags from Legnica. Pt and Pd were not detected at detection limits of 0.002 mg/kg (Kucha and Cichowska, 2001).

**Sandstone Cu-type.**—Mine waste from deposits hosted by the Revett Formation in Montana has low concentrations of trace elements. The orebody contains very little pyrite, and levels of Cd, Sb, and other metals are low. Copper occurs predominantly as Cu and Cu-Fe sulfides and Ag occurs as native Ag and in solid solution in Cu minerals. There are broad halos of galena-rich rock, much greater than 100 m in width and several tens of meters thick, surrounding the ore and, farther outboard, pyrite. The Pb-rich rock has been shown to leach Pb in considerable concentrations but can be avoided when mining (E&MJ, 1989; U.S. Forest Service and others, 2009). Tailings for the Revett deposits contain low S concentrations, ranging from 0.01 to 0.08 wt. %, presumably mainly as Cu sulfides. However, if the S were in pyrite, it would equate to an acid-generating potential value of 0.3 to 2.5 kg CaCO₃/T (U.S. Forest Service and others, 2009).

**Table 5.** Concentrations of elements (in milligrams per kilogram) in slag and flue dust from smelters in the Mansfeld mining district (RF), Germany.

<table>
<thead>
<tr>
<th>Sample type</th>
<th>As</th>
<th>Cd</th>
<th>Cr</th>
<th>Cu</th>
<th>Fe</th>
<th>Pb</th>
<th>Zn</th>
<th>Source</th>
</tr>
</thead>
<tbody>
<tr>
<td>Slag</td>
<td>3</td>
<td>1</td>
<td>233</td>
<td>1,636</td>
<td>13,708</td>
<td>172</td>
<td>7,288</td>
<td>Jahn and others (1997)</td>
</tr>
<tr>
<td>Slag</td>
<td>5</td>
<td>&lt;3</td>
<td>230</td>
<td>1,600</td>
<td>NR</td>
<td>200</td>
<td>2,800</td>
<td>Mathies and others (1999)</td>
</tr>
<tr>
<td>Flue dust (fresh)</td>
<td>3,800</td>
<td>435</td>
<td>130</td>
<td>13,200</td>
<td>12,700</td>
<td>143,000</td>
<td>193,000</td>
<td>Weiss and others (1997)</td>
</tr>
<tr>
<td>Flue dust (weathered)</td>
<td>6,674</td>
<td>451</td>
<td>201</td>
<td>10,117</td>
<td>14,722</td>
<td>95,772</td>
<td>83,182</td>
<td>Jahn and others (1997)</td>
</tr>
<tr>
<td>Flue dust (mean concentration)</td>
<td>NR</td>
<td>8,100</td>
<td>NR</td>
<td>15,500</td>
<td>12,900</td>
<td>15,900</td>
<td>9,900</td>
<td>Schubert and others (2003)</td>
</tr>
<tr>
<td>Flue dust (ground)</td>
<td>4,080</td>
<td>360</td>
<td>890</td>
<td>14,100</td>
<td>30,850</td>
<td>163,000</td>
<td>163,000</td>
<td>Morency and others (1998)</td>
</tr>
<tr>
<td>Flue dust</td>
<td>6,300</td>
<td>980</td>
<td>&lt;10</td>
<td>9,200</td>
<td>NR</td>
<td>12,400</td>
<td>15,900</td>
<td>Matheis and others (1999)</td>
</tr>
<tr>
<td>Flue dust</td>
<td>215</td>
<td>450</td>
<td>18</td>
<td>10,100</td>
<td>NR</td>
<td>9,600</td>
<td>8,300</td>
<td>Matheis and others (1999)</td>
</tr>
</tbody>
</table>

In contrast to the Mansfeld district, the White Pine mine included 26.3 km² of tailings in three surface impoundments (Williams and others, 2002). In 1991, White Pine generated 5.0 million tonnes of tailings and 182,000 t of reverberatory furnace slag (U.S. Environmental Protection Agency, 1994). During the 1970s and during periods of low production rates in the 1990s, slag was crushed and underwent beneficiation to recover Cu (U.S. Environmental Protection Agency, 1994). Red-bed-subtype deposits are generally low tonnage (median 1.2 million tonnes) with intermediate Cu grade (median 1.2 percent) and are mined either on the surface or underground. The volume of mine waste and tailings is expected to be lower for the red-bed deposits compared to the other subtypes of deposits because of their overall smaller size.

The sandstone Cu deposits are intermediate tonnage (median, outside of the Central African Copperbelt, 7.2 million tonnes) but lower Cu grade (median 1.08 percent). The Troy mine in Montana produced 2.9 million tons of tailings each year (U.S. Environmental Protection Agency, 1994). The mine was in operation from 1981 to 1993, and it re-opened in 2004 and has operated continuously since then. Because of the intermediate tonnage of these deposits, the amount of waste produced is intermediate between the red-bed-subtype and the reduced-facies-subtype deposits.

**Mine-Waste Characteristics**

Common mine waste produced by the processing of sedimentary-hosted Cu deposits includes flotation mill tailings, flue dust concentrated by spraying water into the smelter process gases, and silicate slag. These types of waste can be characterized on the basis of their chemical composition, mineralogy, and acid-generating potentials.

**Chemistry**

*Reduced-facies-type.*—The mining of reduced-facies deposits has produced tailings, slag, and flue dust as waste products. Tailings from two deposits, Chambishi and Nkana, in Zambia in the Central African Copperbelt generally contain 1 to 4 wt. % Fe, 0.1 – 0.3 wt. % Cu, and 300 to 800 mg/kg Co. A hardpan layer in one tailings pile contains substantially higher concentrations of these three metals (Sracek and others, 2010). Tailings from the Rudna, Polkowice, and Lubin mines in Poland are all dumped in a common tailings pond. The tailings from the Rudna mine are reported to contain approximately 0.2 percent Cu (E&MJ, 1986b). Tailings from the White Pine mine in the United States contain comparable concentrations; Cu ranges from less than detection to 0.17 wt. % (U.S. Environmental Protection Agency, 1994; Williams and others, 2002). The concentrations of Cu for tailings from all three countries are consistent.

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Mineralogy

SSC deposits contain zones of chalcocite and other minerals near Cu,S in composition, as well as bornite, chalcopyrite, pyrite, and subordinate galena and sphalerite in order outward from the oxidized source of Cu. Native copper and native silver may also be present. The chalcocite and bornite in the high-grade zone are fairly stable under oxidizing conditions, but the oxidation of pyrite can generate acid and cause environmental issues depending on its abundance. The overall concentrations of sulfide minerals are typically low for these deposits and only make up less than or equal to 10 wt. % of the processed ore. Because of the overall low sulfide content, the mine-waste mineralogy is dominantly a reflection of the host-rock mineralogy, which does vary between the subtypes of SSC deposits. For reduced-facies-type deposits, host rocks are marine or lacustrine rocks and approximately 30 percent of deposits are hosted by limestone, dolomite, or marl according to Cox and others (2003). The tailings produced from processing these deposits are more likely to contain higher abundances of carbonates that can mitigate acid generation. In contrast, about 3 percent of red-bed-type deposits and 4 percent of sandstone-type deposits are hosted by carbonates, although nearly all deposits contain at least a minor amount of carbonates. The majority of the red-bed (85 percent) and sandstone Cu deposits (77 percent) are hosted by sandstone or conglomerate (Cox and others, 2003). Therefore, tailings generated by processing ores of the various subtypes will be composed of silicate, carbonate, and (or) oxide phases from the host rocks, with only minor to trace amounts of sulfides. This is consistent with the findings of Sracek and others (2010), who report the primary mineralogy of tailings from the Zambian Copperbelt as quartz, muscovite, amphibole tremolite, orthoclase, calcite, and, at some sites, dolomite. The sulfide S at various depths within the tailings piles in the Zambian Copperbelt is only between 0.03 and 0.5 wt. %; pyrite and chalcopyrite were identified by electron microscopy. Also, inorganic C, which likely equates to calcite or dolomite, ranges from 0.4 to 3.1 wt. % (Sracek and others, 2010).

Several studies have focused on characterizing the mineralogy of flue dust from Germany and slag from Poland, both from reduced-facies-type deposits. First, flue dust from the Mansfeld district consists mainly of amorphous sulfidic particles containing Pb, Zn, Cu, and Fe (Schubert and others, 2003). The predominant mineral phases are sphalerite/wurtzite and galena, with poorly developed lattice structures and extensive ionic substitutions and secondary anglesite (Weiss and others, 1997; Morency and others, 1998). Substantial amounts of amorphous material are also present (Weiss and others, 1997). The mineralogy of various types of slag (shaft, convector, and suspension) from the Legnica and Glogow smelters in Poland includes, as the main component, a glass phase that contains abundant small oval aggregates of metallic Cu, sulfides, Fe-Co alloys, Pb-Ag alloys, spinels, sphalerite, arsenate phases, and less commonly olivine, pyroxene, and amphibole crystals (Kucha and Cichowska, 2001). Silver generally occurs as small metallic inclusions in metallic Cu, metallic Pb, and sulfides. Au is generally found with metallic Cu, metallic Ag, metallic Pb, and Pb-Ag alloys (Kucha and Cichowska, 2001).

Secondary Minerals

Secondary minerals form on mine-waste dumps, slag heaps, and flue-dust piles. Data on the secondary mineralogy of mine waste from deposits are discussed below.

Gordaitse (hexagonal Na(Zn,Cu)(SO₄)(OH)₂Cl₆H₂O) has been identified on the surface of mine-waste dumps in the Mansfeld district occurring with connellite (hexagonal Cu₁₉(SO₄)₂(OH)₃Cl₃3H₂O), anglesite (orthorhombic PbSO₄), and chloride bromargyrite (cubic Ag(Br,Cl)) (Jahn and Witzke, 1999). Jahn and Witzke (1999) report secondary gypsum (CaSO₄.2H₂O) and malachite (monoclinic Cu₃(CO₃)₂(OH)₂), as well as an amorphous Zn Pb sulfate, on the surface of dumps. Matheis and others (1999) reported secondary minerals form from weathering of ores and flue dust as relatively soluble Zn-dominated amorphous phases and more stable Cu and Pb-bearing carbonates and sulfates. After mining stopped in the Mansfeld district in 1990, the water that covered the flue-dust piles dried up, allowing oxidation and formation of secondary minerals, particularly sulfates, that have various solubilities. When exposed to rainfall, soluble secondary minerals are washed into the surrounding environment (Schubert and others, 2001, 2003).

Gypsum, hematite, and poorly crystalline Fe oxides are reported as secondary phases in tailings from the Zambian part of the Central African Copperbelt, mostly forming hardpan layers within the piles. The sulfate S reaches nearly 5 wt. % at one location at depth because of the formation of a gypsum hardpan; the sulfate S contents of the remaining samples are generally less than 1 wt. % (Sracek and others, 2010).

The secondary mineral, gordaitse, was reported in small cavities together with sphalerite (ZnS), boléite (cubic KPB₉₂Ag₉Cu₂₄(OH)₄₈Cl₁₈), and gypsum within slags of the Mansfeld district (Witzke and Pöllmann, 1996). These secondary minerals found on waste dumps and slag heaps indicate the reactive nature of these materials.

Acid-Base Accounting

The abundance of carbonate minerals and minerals that weather at intermediate rates, principally chlorite- and epidote-group minerals, define the acid-neutralizing capacity of these deposits, whereas the acid-generating potential is defined by the abundance of sulfides such as pyrite and chalcopyrite. The net neutralization potential is the acid-neutralizing capacity minus the acid-generating capacity; the neutralization potential ratio is the acid-neutralizing capacity divided by the acid-generating capacity. Many deposits contain substantial carbonate minerals in the ore or host rock, which neutralize acidity generated by the sulfides. Nacimiento, N. Mex., is an exception among the red-bed deposits and contains as
much as 1 v. % carbonate minerals (LaPoint, 1974). As for acid-generating sulfides, the ore for all three subtypes contain low pyrite contents. Host-rock pyrite content varies with greater than or equal to 1 v. % pyrite in host-rock shale and siltstone of the reduced-facies subtype and approximately 0.1 to 0.2 v. % pyrite in the peripheral pyrite-bearing zones of sandstone Cu-type deposits in the Revett Formation (Hayes and Einaudi, 1986).

Sulfide minerals have zonal distributions in these ore deposits, which need to be considered when evaluating potential environmental effects. The high-grade ore zone contains chalcopyrite and bornite, which are fairly stable under oxidizing conditions. This zone contains pyrite in trace amounts or none at all. In contrast to the high-grade ore zone, the low-grade zone consists of pyrite and chalcopyrite; pyrite abundance increases outward as Cu grade decreases (Cox and others, 2003). Therefore, the low-grade zone has a higher potential to create acid mine drainage but is not the target of the mining. The acid-base accounting methods commonly used assume the source of S is acid-generating pyrite. This overstates the potential for acid generation by the Cu-sulfide minerals. Therefore, the acid-generating potential may be overestimated by this method.

Limited acid-base accounting data have been reported in the literature for SSC deposits. Acid-base accounting data for waste from three reduced-facies-type deposits are given below. For tailings, no acid-generating capacity is detected for samples from the White Pine mine (Williams and others, 2002) and high acid-neutralizing capacities and neutral to alkaline conditions are reported for samples from Zambia in the Central African Copperbelt. For the Zambian tailings, neutralization potential ratios are between 3.4 and 84 kg CaCO3/T; most samples at depth have values greater than 20. The highest values are for the leached surface materials (Stacek and others, 2010). The acid-neutralizing capacity of flue dust from the Mansfeld district was approximated at only 45 kg CaCO3/T (900 mmol H+/kg) (Paschke and others, 2001).

The net neutralization potentials of ore, waste rock, and tailings from the sandstone Cu deposits, including Montanore, Rock Creek, and Spar Lake (Troy mine), were summarized by the U.S. Forest Service and others (2009). Ore from the Montanore deposit has an average net neutralization potential of ~4 kg CaCO3/T, with values ranging from ~24 to 11 kg CaCO3/T. Values for Rock and Troy ore samples average 5.1 and 7.6 kg CaCO3/T, respectively. The mean net neutralization potentials for tailings from these sandstone Cu deposits range from 2.8 to 10 kg CaCO3/T. Humidity cell tests on tailings resulted in leachates with near-neutral pH and low levels of metals (U.S. Forest Service and others, 2009). The acid-base accounting results are because of low sulfide concentrations in the tailings. The mean net neutralization potentials for mine waste from these deposits range from 3.6 to 15 kg CaCO3/T, although the waste-rock net neutralization potential is highly variable. For example, net neutralization potentials range from ~20 to 54 kg CaCO3/T for the Burke and Prichard Formations, which would be penetrated by an adit in order to access the Montanore deposit, and from 3.2 to 6.0 kg CaCO3/T for the lower Revett Formation host rocks of the deposit. These ranges indicate highly variable potential to generate acid from various layers within the host stratigraphic section of a deposit.

### Element Mobility Related to Mining in Groundwater, Surface Water, and Stream Sediments

The potential for acid drainage and the release of metals associated with these deposits is minimized by the widespread presence of carbonate minerals in ore and waste rock and by low pyrite and chalcopyrite contents. Overall, the metal mobility from these deposits is low to moderate in water draining mines. Waters draining mines and tailings generally have near-neutral to moderately alkaline pH and low dissolved-metal contents (table 4). However, the concentrations of some metals, including As, Cd, Cr, Cu, Hg, and Pb, may be elevated downstream from mining and milling operations relative to upstream concentrations.

Metals from mining waste can be immobilized by natural processes such as precipitation, sedimentation, and sorption. The presence of carbonate rock can mitigate the acidic water that may drain from tailings and waste rock. Metals in surface water may be sorbed by Fe or Mn oxyhydroxides or may form less-soluble metal carbonates, hydroxides, and oxides. Lakes and wetlands can act as a trap for particulates. Metals may be removed by filtration through soil or by plant uptake.

### Reduced-Facies-Subtype Deposits

The mining of deposits in the Mansfeld district has contaminated stream sediments, soils, and surface water and groundwater with metals such as Zn, Pb, Cu, As, and Cd. The source of the metals and S has been shown to be predominantly low-grade ore from mining and flue dust and slag from smelting (Schreck and Glaesser, 1998; Strauch and others, 2001; Schreck and others, 2005). Strauch and others (2001) stated that the flue dust is the most important source of pollution in this mining district. Metals were recovered from the flue dust until 1978, after which the material was deposited in open basins in piles or on slag or mining debris (Matheis and others, 1999). The flue dust is extremely fine-grained being micrometer to submicrometer in size adding to the weathering potential of the material. Wennrich and others (2001) demonstrated on the basis of laboratory extraction procedures that most metals in the flue dust have a high mobilization potential under simulated natural conditions. Paschke and others (2001), using a standard leaching test with distilled water as well as tests that simulated drainage from sludge ponds and acid-rain deposition on flue dust, showed Zn, Pb, Ni, and Cd were leached in concentrations that exceeded environmental limits. Matheis and others (1999) performed outdoor
In addition to stream and lake sediment, wetland sediment can also become contaminated downstream from ore-processing facilities. At the Chambishi mine in the Zambia, effluent from ore processing eventually flows through a natural wetland. The ranges in concentrations of various elements in wetland inflow, which is downstream from a tailings impoundment and chemically comparable to tailings outflow, and wetland outflow in the Chambishi-mine catchment are reported in table 4 (von der Heyden and New, 2005). This natural wetland system substantially improves the quality of the water by retaining potentially toxic metals, such as Cu and Co, and diluting some conservative pollutants, such as sulfate (von der Heyden and New, 2005). Von der Heyden and New (2004b) estimated that the wetland has accumulated 150 tonnes of Cu and 100 tonnes of Co in the upper 1 m of wetland sediment. The wetland has been receiving metal-laden effluent for between 40 and 50 years. The range in concentrations of Cu and Co in wetland sediment is shown in table 3. Von der Heyden and New (2004b) suggest that the sediment is a potential source of pollutants to the downstream ecosystem. Their work demonstrates that under conditions of lowered surface-water metal concentrations, the metals in the sediment may be released to the water; this may be the condition once mining operations have ceased, and the wetland no longer receives effluent (von der Heyden and New, 2004b). Von de Heyden and New (2003) also stress that the benefit to the downstream ecosystem from the filtering effect of the wetland is offset by the severe pollution of the wetland itself.

Sandstone Copper Deposits

Several environmental impact statements for sandstone Cu deposits in Montana discuss the mobility of trace elements to surface water. Tailing effluent and mine-water chemistries for the Spar Lake (Troy) mine are given in table 4 (U.S. Forest Service and others, 1992, 2009). The concentrations of Cd, Cu, and Pb in the tailings outflow exceed the guidelines for aquatic ecosystems. The potential mobility of trace elements from tailings and waste rock generated from mining the Rock Creek and Montanore deposits was assessed on the basis of standard leaching tests. Ore from the Rock Creek and Montanore deposits leached low concentrations of Cu (0.02 – 0.04 mg/L) and Mn (0.03 mg/L) and resulted in neutral pH water on the basis of a humidity cell test. On the basis of the Toxicity Characteristic Leaching Procedure, tailings and waste rock leached low amounts of metals such as Ba, Cr, Cu, Fe, Pb, Mn, and Zn (U.S. Forest Service and others, 2009). Waste rock from some parts of the deposit has the potential to generate acid, whereas other parts do not. Groundwater from monitoring wells in tailings impoundment sites and disposal areas in Montana contain low total dissolved solids and frequently non-detectable dissolved metals. The pH of the water is typically near-neutral or slightly acidic (U.S. Forest Service and others, 2009).
Smelter Emissions

Ore from SSC deposits is either smelted on site or shipped off site. Smelters may emit particulates, metals, nitrogen oxide, sulfur dioxide, carbon monoxide, and (or) hydrocarbons. Atmospheric dusts collected in the vicinity of the Glogow smelters in Poland contain sulfides and arsenides of Cu, Pb, Zn, and Ni, and alloys of Pb, Pb-Cu, and precious metals (Muszer, 2004). Numerous studies on the soil surrounding the Glogow and Legnica smelters report high metal concentrations, particularly for Cu and Pb, because of dust emissions; most studies document that concentrations in soils depend on the dominant wind direction and distance from the smelter (table 6) (Roszyk and Szerszen, 1988; Helios Rybicka and Jędrzejczyk, 1995; Karczewska, 1996; Pilc and others, 1999; Grzebisz and others, 2001; Kabala and Singh, 2001). Soil samples 1 km from the smelters contained 250–10,000 ppm Cu, 90–18,000 ppm Pb, 0.3–10.9 ppm Cd, and 55–4,000 ppm Zn (Roszyk and Szerszen, 1988; Helios Rybicka and Jędrzejczyk, 1995). Kabala and Singh (2001) examined soils between 1 and 6 km from the smelters and reported distinctly lower concentrations of 97–426 ppm Cu, 65–130 ppm Pb, and 45–92 ppm Zn in soils. Grzebisz and others (2001) also found Cu concentrations (26–330 ppm) within 10 km of the smelter that were comparable to those reported by Kabala and Singh (2001) (table 6). Overall, the concentrations of Cu and Pb determined in these studies were substantially higher than typical values for agricultural soils (Kabata-Pendias and Pendias, 1992). According to Kabala and Singh (2001), the relative mobility of Cu, Pb, and Zn increased with an increase in their total-metal content and decreased with an increase in the clay content of the soil. Pilc and others (1999) examined soils 2.0 and 5.5 km from the smelters and found concentrations of Cu, Pb, and Zn that were comparable to those reported by Kabala and Singh (2001) (table 6). Pilc and others (1999) found the highest metal concentrations in soils that had been fertilized intensely with manure, suggesting natural fertilizers may also introduce substantial amounts of pollutants to the soils. The concentrations of metals in groundwater and surface water in the vicinity of the smelters are given in table 4 (Pilc and others, 1999). The concentrations of Pb and Cd in many of these waters exceed the World Health Organization guidelines for drinking water (World Health Organization, 2006). As with the soil samples, the substantially higher metal concentrations in the groundwater may result from fertilization with manure.

Flue dust from the Mansfeld district has been documented to release metals into the environment through deflation, erosion, and leaching (Schreck and Glaesser, 1998). The flue dust not only contains large quantities of metals such as Pb and Zn but contains substantial amounts of organic compounds (that is, polycyclic aromatic hydrocarbons, polychlorinated dibenzo-p-dioxin) and high concentrations of radionuclides. The extremely fine-grained nature of the material and high solubility for some of the substances promote the release of these materials into surface water, groundwater, and soils (Weiss and others, 1997). (See section on “Element Mobility Related to Mining”).

In 1990, the smelter at White Pine emitted approximately 900 short tons of particulates and 225 tons of metal. Metal outputs include 198 ton/yr Cu, 25 ton/yr Pb, 9 ton/yr As, 1.8 ton/yr Cd, and less than 1 ton/yr each of Cr, Hg, and Ni (Lindsey and others, 1995).

Ecosystem Issues

Potential impacts to aquatic ecosystems near a mine site may be grouped into the following categories: changes in sedimentation and water quantity, changes in water quality, changes in toxic metal concentrations in fish tissues, and effects on threatened, endangered, or sensitive species. First, sedimentation rates in streams may increase because of mine construction and mine activities. Aquatic habitats could be affected by changes in flow because of various mechanisms such as changes in stream base flow because of mine and adit inflows, infiltration from disposal sites into shallow groundwater systems and then downgradient streams, diversion of precipitation intercepted by tailings impoundments for mill use, increased storm-water runoff from mine facilities, and increased peak flows affected by vegetation clearing for roads and mining facilities (U.S. Forest Service and others, 2009). Aquatic life can also be affected by high concentrations of metals and nutrients in streams. The effects

### Table 6. Concentrations of metals (in milligrams per kilogram) in surface soils surrounding the Glogow and Legnica smelters, Poland.

<table>
<thead>
<tr>
<th>Smelters</th>
<th>Distance from smelters</th>
<th>Cd</th>
<th>Cu</th>
<th>Pb</th>
<th>Zn</th>
<th>Source</th>
</tr>
</thead>
<tbody>
<tr>
<td>Glogow</td>
<td>Within 1 km</td>
<td>0.3–10.9</td>
<td>250–10,000</td>
<td>90–18,000</td>
<td>55–4,000</td>
<td>Roszyk and Szerszen (1988)</td>
</tr>
<tr>
<td>Glogow</td>
<td>Within 6 km</td>
<td>ND</td>
<td>97–426</td>
<td>65–130</td>
<td>45–92</td>
<td>Kabala and Singh (2001)</td>
</tr>
<tr>
<td>Glogow</td>
<td>Within 5.5 km</td>
<td>0.75–1.58</td>
<td>88–510</td>
<td>55–255</td>
<td>68–207</td>
<td>Pilc and others (1999)</td>
</tr>
<tr>
<td>Glogow</td>
<td>Within 10 km</td>
<td>ND</td>
<td>26–330</td>
<td>ND</td>
<td>ND</td>
<td>Grzebisz and others (2001)</td>
</tr>
<tr>
<td>Glogow &amp; Legnica</td>
<td>&quot;near&quot; the smelter</td>
<td>ND</td>
<td>42–298</td>
<td>ND</td>
<td>ND</td>
<td>Grzebisz and others (1997)</td>
</tr>
<tr>
<td>Glogow &amp; Legnica</td>
<td>Within 2.5 km</td>
<td>ND</td>
<td>7.4–1,710</td>
<td>11–460</td>
<td>6.4–153</td>
<td>Karczewska (1996)</td>
</tr>
<tr>
<td>Legnica</td>
<td>0.2 km</td>
<td>4</td>
<td>7,400</td>
<td>1,960</td>
<td>675</td>
<td>Helios Rybicka and Jędrzejczyk (1995)</td>
</tr>
</tbody>
</table>

1Includes A and B soil horizons.
depend on the hardness of the water for metals such as Zn, Cu, Pb, and Cd. The hardness of waters in the vicinity of the Cu deposits can vary depending on the presence and abundance of certain host rocks such as limestone. The greater the hardness, the less toxic the metal is to the ecosystem. Also, air quality may be affected by emissions from smelting and other processing facilities, and surface disturbances may result from open pits and leaching facilities. Lake levels may be disturbed if underground workings intersect fractured rock beneath lakes.

Detailed ecological studies have been carried out along the Kafue River, which flows through the Zambian part of the Central African Copperbelt. Syakalima and others (2001) analyzed water, grass, fish, and lechwe (antelope) liver for Pb; these samples were collected downstream from the mining-impacted areas from two national parks. This study found concentrations of Pb in water between 0.29 and 0.36 mg/L compared to the World Health Organization drinking-water guideline of 0.01 mg/L. In a grass species commonly grazed by wildlife and livestock, Pb concentrations are between 26.0 and 48.0 mg/kg dry weight. Three species of fish contained between 23.0 and 33.0 mg/kg dry weight Pb, and the liver of the lechwe contained between 7.0 and 25.0 mg/kg dry weight Pb (Syakalima and others, 2001). Overall, this study reported that Pb has the potential to bioaccumulate in this ecosystem and may cause adverse effects (Syakalima and others, 2001).

Other studies focused on effects of metals in Kafue River on tropical fish toxicity. Stream sediment collected downstream from several mines and near a city that has a major ore-processing facility was substantially more toxic to zebrafish and tilapia than sediment collected further downstream (Mwase and others, 1998). The toxicity was directly related to the contamination of sediments by the mining activities (Mwase and others, 1998). The results of another study showed tilapia exposed in situ to Kafue River water for 2 weeks bioaccumulated trace elements (that is, Cd, Co, Cu, Cr, and Ni) in experiments downstream from mining activities and other industrial point sources (Norrgren and others, 2000).

Several studies have been carried out on plants growing in the vicinity of the Glogow smelters in Poland (Pilc and others, 1999; Grzebisz and others, 2001). Grzebisz and others (2001) studied nonconsumable crops and found higher amounts of Cu in flax plants than in oil-seed rape. The concentrations of metals in cultivated consumable plants, including corn, carrots, red beets, and beans, correlated with the distance from the source of smelter emission and with the direction and strength of winds blowing most frequently in the region, indicating the plant pollution is a result of the dust emissions and not the concentrations of metals in the soils (Pilc and others, 1999). Pilc and others (1999) referred to other studies that identified high levels of metals in blood, milk, liver, kidneys, bones, hair, and excrement of the cattle grazing the region. Other studies used sequential extraction schemes on soils to assess the distribution of metal species in operationally defined fractions to gain insight into potential mobility and bioavailability (Helios Rybicka and others, 1994; Grzebisz and others, 1997; Karczewska, 1996).

The growth of plants on waste material is an indication of ecosystem health. Vegetation establishment was attempted in the 1960s on tailings from the White Pine mine. In general, the vegetation died after 2 to 3 years because of factors including the sand-blasting effect of wind blowing the tailings, the physical nature of the material, and possibly the chemical and nutrient imbalances. Nearly all Cu in the tailings was loosely bound and is considered available for uptake by plants. In 1997, a reclamation goal was set to revegetate the tailings to help stabilize them. Greenhouse and field tests were conducted in 1999 and 2000; large test plots were treated. As reported in 2002, the treated areas were self-sustaining and green (William and others, 2002).

The results of a study to assess the local and long-term effects on Lake Superior from the discharge from the White Pine mine was conducted in 1991. The study showed that discharge from the mine area was not causing measurable effects on the basis of a lack of impact on the benthic community, insignificant increase in chloride from mine discharges, and a lack of detectable effects on local water intakes (U.S. Environmental Protection Agency, 1994). USEPA (1994) referred to the Surface Water Quality Division of the Michigan Department of Natural Resources saying that a diverse fish community exists in the nearby Mineral River that receives drainage from the mine. They reported a reduced macroinvertebrate community downstream from the mine drainage compared to upstream in the river (U.S. Environmental Protection Agency, 1994). This reduction may be because of major physical and natural stream-quality differences in the White Pine area (U.S. Environmental Protection Agency, 1994). Biomonitoring of mine drainage conducted in 1991 showed acute and chronic toxicity for discharge collected in August but not for discharge collected in April, June, or October (U.S. Environmental Protection Agency, 1994).

Climate Effects on Geoenvironmental Signatures

The effects of various climatic settings on the geoenvironmental signatures of these deposits are not well documented. In general, the environmental impact associated with sulfide-mineral-bearing deposits is greater in humid to temperate climates in comparison to arid climates. Acidity and total-metal concentrations in mine drainage are generally higher in arid environments because of evaporation, which results in the storage of metals and acidity in highly soluble metal-sulfate-salt minerals. However, the lack of surface-water drainage inhibits the mobility of the acidity and metals. In contrast, in more tropical environments, the leaching of metals is exacerbated (Lindsey and others, 1995).
Knowledge Gaps and Future Research Directions

There are many different knowledge gaps in studies of SSC deposits, offering outstanding opportunities for usefully directed research. Some of these are discussed below.

Metals-Transporting Brines

Inorganic Brine Chemistry

Although it is clear from places that have been studied that the ore-forming groundwater for SSC deposits was briny and it mixed with a second, more dilute solution, the source(s) of the metals-transporting solutions remain somewhat conjectural. In only two places, source rocks have been found that have been depleted of their original Cu (Coates Lake deposit, Northwest Territories, Canada: Lustwerk and Rose, 1983; Western Montana Copper Sulfide Belt: Hayes, 1990). As known to date, only Coates Lake has continuous exposure of rock between depleted rocks and the Cu deposit.

The research tools needed to answer questions about the brine source(s) are in place. Brine inorganic chemistry will go a long way toward determining the source(s) of the metals-transporting solutions. In general, two methods, either macro-chemical or micro-chemical, have been used on other ore-deposit types. Both might be employed in studies of the fluid inclusions of SSC deposits.

Viets and others (1996) conducted ion chromatography studies of fluid inclusions in sphalerite or massive ore from Mississippi Valley-type Pb-Zn deposits. With ionic compositions of the fluids, various ionic ratio cross plots were constructed and compared with seawater composition, seawater residual brine compositions from evaporite precipitation sequences, and re-dissolved evaporite compositions, allowing conclusions about the origin of the fluids and their pre-ore and ore-stage history. The ore-depositing system of MVTs is a broadly similar interaquifer groundwater mixing process to that of SSC deposits (Plumlee and others, 1994, 1995). In detail, it is doubtful that the two deposit types have metals-transporting brines that are of similar origin, but it does seem likely that the metals-transporting brines of both became strongly saline by dissolving evaporites. Ion chromatography (Viets and others, 1996) would also be the choice for analysis of the anions of SSC-deposit brines. This method would probably be best applied to pure sulfide mineral separates, although it might also provide interesting comparisons from co-existing carbonates and sulfates, for example, from Kupferschiefer ores. It would be best applied to an unmetamorphosed deposit. For fine-grained SSC ores, hand picking and crushing minerals would probably seriously contaminate the ore-stage signature with fluid-inclusion fluids from the detritus, but an approach that thermally decrepitates the inclusions could still succeed (Hayes and others, 2012).

Multiple new techniques for microanalyses of fluid inclusions include laser-ablation inductively coupled plasma mass-spectrometry (LA-ICPMS) (Heinrich and others, 2003), proton-induced X-ray emission (PIXE) (Kurosawa and others, 2003), and synchrotron x-ray fluorescence (SXRF) (Bassett and Brown, 1990). These methods have been successfully used to define composition of inclusion fluids in ore-deposit studies of porphyry Cu, molybdenum, and tungsten deposits (Heinrich and others, 1992; Vanko and others, 2001; Klemm and others, 2007, 2008), Fe oxide-Cu-Au deposits (Baker and others, 2008), epithermal precious-metals deposits (Wallier and others, 2006), volcanogenic massive sulfide deposits (Moura, 2008), intrusion-related Au deposits (Baker and others, 2006), banded Fe formations (Brown and Oliver, 2002), and Mississippi Valley-type Pb-Zn deposits (Stoffel and others, 2008). The LA-ICPMS or PIXE methods promise to be very successful in studies of SSC deposits but only if fluid inclusions are large enough. Most inclusions from SSC-deposit main stages are less than 10 µm in size, and such inclusions are too small for LA-ICPMS, PIXE, or SXRF study. Coarser grain rocks would generally yield coarser grain minerals from the main-stage ore (cements), and those ore-stage minerals might have larger fluid inclusions than finer grain ores, but that has not yet been demonstrated. The LA-ICPMS, PIXE, and SXRF methods are well-suited to studying second-stage veinlet-hosted stages of SSC deposits.

Sources of Brine Salinity

In only one place has the source of salinity in a metals-transporting brine been strongly constrained for an SSC deposit. Breit and Meunier (1990) determined the strontium-isotope ratios of barite cements, dolomite cements, and barite in feeder veins for the Lisbon Valley sandstone Cu deposit. Such isotope determinations can also constrain the sources of the brines and the sources of their salinity. The study used chemical leaching methods to free the strontium, and it used samples of fine crystalline sandstone cements from the sandstone Cu deposit and other sandstones nearby and coarse crystalline material from feeder veins. The chemical leaching method could be applied to ore-stage gangue minerals of any grain size, although petrographic and microthermometric examination of the fluid inclusions in the minerals should be performed to yield a clearer understanding of the inclusions that host the fluids with saline-sourced elements and isotopes. Because so many SSC deposits occur in stratigraphic sections that contain evaporites, there is little doubt that evaporites have contributed to the salinity of metals-transporting solutions in most districts (Kirkham, 1989). Details that could be valuable in understanding deposit genesis may, nonetheless, be available from additional studies focused on the sources.
of brine salinity. In a few districts (for example, Udokan, Belt Supergroup, Cambrian of Israel and Jordan) the connection to evaporites is not obvious.

**Find an Active Ore-Forming System**

The most reliable results in attempting to define the source(s) of metals-transporting solutions for an SSC deposit would be to sample the metals-transporting solutions from a present-day active mineralizing system. In years past, finding an active system would have been considered impossible, but this does not appear to be the case now.

**Sandstone Cu-type.**—An active sandstone Cu mineralizing system can probably be found in a basin that has petroleum source rocks overlain by greater than 300 m, and preferably greater than 1 km, of red beds, overlain by intervals that contain gray or bleached shales, and that has production of sour natural gas. Groundwaters with around 10 ppm Cu in that setting are the target brines. A sampling target may already be identifiable; Lebedev and Nikitina (1968, p. 181) sampled and analyzed a brine from “aquifer 11” of the Cheleken oil and gas field and iodine-bromine deposit that was a sodium-calcium-chloride brine with dissolved Fe$^{3+}$ exceeding Fe$^{2+}$ and with 8.34 ppm dissolved Cu. The geologic setting of the Cheleken field is remarkably similar to what the setting of the ore-stage Dzhezkazgan sandstone Cu district must have been. However, further details on Cheleken aquifer 11 are not available.

**Reduced-facies-type.**—An active reduced-facies-type stratabound Cu mineralizing system may be locatable but is not so straightforward to find as the sandstone Cu-type system. One should be looking in a young rifting environment at low geothermics so that the active system would have been considered impossible, but this does not appear to be the case now.

**Hydrologic Drive(s) of Metals-Transporting Groundwater Flow**

**Paleohydrologic Modeling of Metals-Transporting Groundwater Flow**

A number of attempts have been made to model the groundwater flow systems that produced SSC deposits. These were by White (1971), Jowett (1986a), Cathles and others (1993), Person and Garven (1994), Blundell and others (2003), Swenson and others (2004), and Brown (2005, 2009).

Person and Garven (1994) examined the general case of groundwater flow in the protoceanic rift and open-ocean sag stages of passive margins. At full development of the sag phase, rapid deposition near the basin axis would result in two groundwater flow systems separated by the fine-grained impermeable beds deposited in the transgression between the protoceanic rift- and the sag-basin stages. A gravity-driven flow system would set up within the coarse-grained beds deposited atop the transgressive beds, and a system driven toward basin margins would be established below those transgressive beds driven by compactive loading along the basin axis.

The Kupferschiefer in Poland has seen a succession of three modeling attempts. Cathles and others (1993) tested an earlier hypothesis offered by Jowett (1986a) that Polish Kupferschiefer ore formed when thermally driven free convection of groundwater in the sub-ore Rotliegendes red beds occurred, confined beneath the Kupferschiefer bed and Zechstein evaporites, and confined laterally by basement rocks of horst blocks. The hypothesized thermal drive was provided by a generally high geothermal gradient accompanying recurrent rift tectonics. Mass balances for such a flow system succeeded only if dissolved Cu concentrations were at least 127 ppm, a concentration higher than in any known sedimentary brine. Blundell and others (2003) started with the observation that normal faulting drives a pulse of groundwater outward away from extensional fractures. They considered deriving metals-transporting groundwater from basement fractures by that means during extensional tectonics through the Early and Middle Triassic. In their model, a localized heat-flow anomaly would provide buoyant fluid coming from the basement fractures, and permeable eolian Weissliegen-des sandstones would provide a connection and allow rapid groundwater flow upward and laterally along the footwall of the Kupferschiefer bed. With these conditions met, normal faulting with a 100- to 200-year recurrence interval could provide enough groundwater to create the Lubin-Sierszowice deposit with solution concentrations of 60 ppm Cu over a period of 12 million years. This model could be called seismic pumping. Its likelihood depends on localized thermal anomalies along the floors and flanks of the rift basin.

The White Pine reduced-facies deposit has also seen a succession of paleohydrologic modeling studies. White (1971) tested the feasibility of forming the deposit solely from
Brown suggested that only this early diagenetic timing allows the proper conditions of oxidation potential for large-scale Cu transport and for a large-scale flux of oxygenated groundwater.

The combination of paleohydrologic and solution chemical modeling by Swenson and others (2004) is considered to be the most realistic of such modeling to date. Importantly, that work demonstrates that a single pass of compactive groundwater flow is sufficient to create reduced-facies Cu deposits. It has long been recognized that reduced-facies deposits in Zambia are localized adjacent to basement highs (figs. 10, 11, and 15), and this modeling explains the hydrology of the localization. It also explains the expanses of trace-mineralized reduced-facies host rocks between the deposits.

The “dynamic footwall reddening” model (Brown, 2005, 2009) is a feasible alternative to compactive groundwater flow for the White Pine deposit. However, there are reasonable alternatives to dissolved oxygen as an oxidizing agent, so proximity to recharge from the surface really is not a serious limitation on Cu mobilization. S+6 in sulfate for example can yield conditions for cuprous Cu solubility. Evaporite brines are typically rich in chloride and sulfate. Maximum Cu transport temperatures that were generally greater than 180 °C for Kamoto and Musonoi, Democratic Republic of Congo (El Desouky and others, 2009), and greater than 200 °C for Spar Lake (Hayes and others, 2012) are notably higher than would be found in dynamic footwall reddening. Isotopic dating does not allow Cu mobilization by dynamic footwall reddening at Spar Lake, because the source red beds were already buried by more than 5,000 m of overburden when mineralization occurred (Aleinikoff and others, 2012).

Additional paleohydrologic modeling studies of SSC deposits would be valuable particularly if they are well constrained by isotopic and stratigraphic ages. Sandstone Cu deposits have not been modeled, and they may have formed generally later in basin histories than reduced-facies deposits. Later in the basin histories, gravity-driven groundwater flow seems more probable. The Paradox Basin has a wealth of oilfield data available and a generally well-known geologic history (Nuccio and Condon, 1996). Once ore deposition has been dated, it would be a good place to study sandstone Cu deposit paleohydrology.

Identifying Flow Paths for Metals-Transporting Brines

Although paleohydrologic studies have proven effective where they could be combined with isotopic dating of ore systems (for example in some Mississippi Valley-type deposits: Bethke, 1986; Brannon and others, 1992; and in the reduced-facies Cu deposit at White Pine: Swenson and others, 2004), the paleohydrology is only testable in theory. Although paleohydrologic modeling studies should be encouraged, studies that attempt to map rock features promise to be more conclusive to identifying the hydrologic drive(s) of mineralizing systems.
Metals transport can be mapped if alteration along the flow path can first be recognized and if exposure is adequate either in outcrops or drill holes. Mapping of alteration pathways could greatly limit the possible flow systems that have produced ore. Certainly, high-angle faults and zones where pillars are densely distributed, both that are exposed downsection from mineralized rocks, are specific targets.

**Alteration.**—From current knowledge, sodic minerals (albite, glauberite, analcime, riebeckite, paragonite, and possibly scapolite), hematite, and possibly barite will mark the paths taken between Cu source rocks and deposits. Epidote is a mineral associated with Cu mobilization from volcanic rocks (Livnat, 1983; Borg, 1991) and is to be expected as well in altered rocks with volcaniclastic components. Albite is possibly the most common alteration mineral, but glauberite is the one identified within Cu-depleted rocks of the Redstone River Formation below the Coates Lake deposit (R. Lustwerk, oral commun., 1985). Starting at a deposit in the Kupferschiefer, it should, with careful examination of rock sequences such as the Rotliegendes in central Europe, be possible to identify and then map alteration paths.

In young mineralizing systems, it would be expected that alteration might also be marked by an anomalously mature ferric-oxide assemblage in the sourcing red beds, such as well crystallized hematite coating the grains of Cu-depleted rocks versus amorphous ferric oxides or poorly crystallized hematite in rocks that have not lost their Cu. A study testing this is suggested near Boléo.

**Metals sources.**—The source rocks for Cu in SSC deposits are marked as rocks with Cu concentrations depleted relative to rocks that were never affected by the mineralizing systems. In both examples where depleted red beds were identified, the background Cu concentrations of the source rocks were about 20 ppm and depleted values were typically below 4 ppm (Lustwerk and Rose, 1983; R. Lustwerk, oral commun., 1985; Hayes, 1990). The source rocks for the Revett Formation-hosted deposits are also remarkably depleted of their Ca; quartzites with greater than 15 v. % total feldspars have less than 400 ppm Ca in whole rocks (T. Hayes, unpub. data, 1989). They are discernibly depleted in Pb, as well. Each element that is depleted in the altered rock is found elsewhere at anomalously high concentrations within the zoned mineralogic systems of the deposits. Cu-depleted rock can be expected to have authigenic minerals unique to a zone of mobilization and including some of the minerals listed above.

### Supergene Enrichment

In the Central African Copperbelt, it is still unclear whether most of the chalcocite is formed during supergene enrichment or whether it is hypogene. Since the times of Bateman (1930) and Oosterbosch (1951), much has been learned about the process of supergene enrichment of Cu deposits. Many new tools have been developed. Just one example is the isotopic age dating of supergene phases such as Mn oxides, which may be directly applicable to heterogenite in the Democratic Republic of Congo. But even if it is only possible to use detailed mineralogy and mapping, better understanding can be gained. The porous sandstones overlying Mufulira ore together with the chalcocite-rich parts of that deposit are a good target for study. Working downward from a Co cap into hypogene ore at a deposit in the Democratic Republic of Congo is a second good target. Understanding the process of enrichment in the climate of the Central African Copperbelt promises to quickly provide value in central Africa and the rest of the world.

### Changes to SSC Deposits with High-Grade Metamorphism

The knowledge of the mineralogic and textural changes in amphibolite-grade and granulite-grade metamorphism of SSC deposits is insufficient. There are also opportune places to address related questions, such as in Aynak, Afghanistan, and in the Ablah Group occurrences in Saudi Arabia. There is immediate value in increased understanding of the subject, because high-grade metamorphism may leave the rock sequences more amenable to geophysical exploration by induced polarization and magneto-telluric methods.

### Direct Sensing of Concealed Ore

The most immediate and highest payback on new studies will likely come from developing exploration methods. Methods that need testing are natural-source induced polarization and soil-gas chemistry. Both methods attempt to directly sense the presence of ore or ore-related features in the subsurface.

A test of the natural-source IP method was started with field data collection at the Rock Creek sandstone Cu deposit in the Western Montana Copper Sulfide Belt in 2009. Data analysis is ongoing.

Knowing that sandstone Cu deposits form where hot, metals-transporting brines mixed and reacted with H,S-bearing natural gas, a test using soil-gas chemistry as an exploration method for sandstone Cu deposits is needed. Obviously, a CH₄ anomaly can be hypothesized. But it is possible that an anomalous signature might be a combination of different pathfinder gases, including CH₄, carbon disulfide, and Hg (fig. 39). A test of soil-gas chemistry in exploration could take place in the Western Montana Copper Sulfide Belt over a known but unmined deposit or it could take place in the Paradox Basin where the Flying Diamond deposit, southeast of the Lisbon Valley deposit, is an outstanding candidate for further study. Flying Diamond was earlier studied by an integrated set of exploration techniques including CO₂-O₂ soil-gas geochemistry (Adkins and others, 2010).
Observable Vectors for Narrowing the Search for Ore

Further study is needed to find exploration guides that may narrow the search for new ore. One such guide may be composition of carbonate minerals of sandstone Cu systems. Work has started to see if there is a quantifiable relation between Fe and Mn contents in calcites and ankerites and their zonation surrounding sandstone Cu deposits of the Western Montana Copper Sulfide Belt (Hayes, unpub. data, 1986). Fe-rich calcites have also been described in association with the Dzhekazgan deposits (Gabrina and Tsepin, 1975). Together with these, there is a suggestion from the observation of clearly hydrothermal dolomite in association with the Paoli, Okla., sandstone Cu deposit (Thomas and others, 1991) that carbonate minerals of anomalous composition may occur in many systems.

Vectors that may narrow the search for new ore may exist at many scales. Because our current understanding suggests that only basins with thick packages of red beds that were deposited relatively quickly develop SSC systems, Lopatin time-temperature models (Lopatin, 1971; Waples, 1980) of basins could be a vectoring technique comparing one basin to another or comparing different parts of the same basin. Lopatin modeling is designed to address thermal maturity of petroleum source rocks (Davey, 1985). It calculates time-temperature indices (TTIs) on the basis of absolute age and burial histories of rock units. The minimum TTI necessary for the onset of oil generation, for peak oil generation, and for the end of oil generation are known from experience (Waples, 1980). The TTIs calculated for source rocks also constrain the time-temperature histories of red beds in the same basins that might source metals-transporting brines, and it is now known that such units probably reached temperatures of 120 °C or greater to source a SSC-mineralizing brine. Thus, Lopatin modeling may be able to distinguish prospective red-bed-bearing basins from non-prospective ones. It could identify large prospective areas within a red-bed-bearing basin. The data needed for Lopatin modeling are widely available from essentially all basins in the world that have undergone some level of exploration for petroleum. Stratigraphic analysis and Lopatin modeling can then guide the early steps of exploration, selecting basin-scale or sub-basin-scale targets. Alteration mapping and geochemical patterns within basins with identified systems might be a medium-scale vectoring technique.

Acknowledgments

An earlier draft of this report was reviewed by Murray W. Hitzman and A.C. Brown who both suggested many useful revisions. Professor Brown probably spent many days in this effort, and we thank him, especially. Both reviewers are experts on SSC deposits who are known worldwide, so we note, particularly, that any conclusions drawn herein are our own, and they do not necessarily reflect the views of Hitzman or Brown. The greatest help on this report came, however, from our colleague, Rich Goldfarb, who edited a second draft in great detail and otherwise worked long and hard in trying to make this model as useful as possible.

We also thank the following individuals for their discussions over many years: Rod Kirkham, Marco Einaudi, Jack Harrison, Dean Kleinkopf, Jeff Mauk, Brian White, Joe Whelan, Gary Landis, Bob Rye, Michael Zientek, Dave Leach, Murray Hitzman, Alan Wallace, Tom Chadwick, Russ Smith, Larry Appelgate, Nancy Joseph, Gregor Borg, Dave Broughton, and Jon Thorson.

A final acknowledgement is due John Wallis, our technical illustrator contracted to the Spokane office of USGS. He brought 59 of the 62 figures of this report admirably into compliance with the requirements for figures in a Scientific Investigations Report.

Subjects covered in this report were determined by the leadership of the Updated National Mineral Resource Assessment of 2012 Project in order to provide consistency with models of other ore-deposit types. For some subjects that were to be covered, for example, hypogene gangue characteristics and major-element rock geochemistry, the Spar Lake deposit was among only a few SSC deposits where the subject had been specifically addressed. An unanticipated effect of the standardized coverage of subjects then became that examples from Spar Lake were used in most sections. Spar Lake examples were at hand because that deposit has been the subject of studies by the first author and colleagues at several times over the past 30 years. We apologize for what may seem an over-emphasis on Spar Lake.

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Sediment-Hosted Stratabound Copper Deposit Model


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Appendix 1. Summary of Representative Sediment-Hosted Stratabound Copper Deposits

A. Name: Lubin-Sieroszowice-Polkowice-Rudna(-Glogow Giboki-Przemyslowy-Bytom Odrzanski-Retkow-Gaworzysze-Radjawice Zachody), Poland

Deposit subtype: Reduced-facies
Location: 51°24′N 16°08′E
Grade/tonnage: Oszczepalski and Speczik, 2011: 3612.499 Mt, 1.99 percent Cu, 57.5 g/t Ag
Associated deposits: Cut by sparse Rücken veins
Regional geologic setting: Southern flank of the Zechstein Basin

Regional tectonic setting: Extensional tectonic regime in homoclinal northeastward dipping sedimentary rock on the northeast flank of the Fore-Sudetic block, a northwest-trending horst with its highest parts composed of Variscan (Carboniferous and Early Permian) metamorphic rocks subcropping below Tertiary and Quaternary cover. Age of the Fore-Sudetic block is Tertiary, pre-Pliocene in age but may mark the position of a pre-Middle Permian basement high from pre-Kupferschiefer, Lower Rotliegendes time. However, the Fore-Sudetic block was not a topographic high in evidence at all at the time of Kupferschiefer deposition – Kupferschiefer lithofacies boundaries cross it trending east-west without perturbation.

Significant structural control (Y/N): N

Primary host rocks and their ages: In ascending order: Late Permian (Guadalupian) Weissliegendes light gray to white sandstones (well-sorted, well-rounded, fine to coarse), Kupferschiefer dark gray to black shale and mudstone, and Zechstein I (Werra) Carbonates (predominantly gray lime wackestones) beds.

Associated red beds: Subjacent Upper Rotliegendes Formation, 100–450 m thick beneath ore; as much as 650 m thick in Fore-Sudetic basin to the northeast though locally pinching out over basement highs: conglomerates less than sandstones greater than lacustrine mudstones; in areas tens of kilometers to the north and southwest, Lower Rotliegendes includes subaerial rhyolite greater than andesite greater than basalt.

Associated evaporites: In mining area, Zechstein I anhydrite 100–200 m thick overlies Werra carbonates; Zechstein II, III, and IV evaporites overlie that in cycles, totaling 50–100 m, including minor halite; in the Polish basin 200 km to the northeast, these evaporites total more than 5 km and are affected by halokinesis. Hundreds of kilometers to the northwest or north in northern Germany and Poland, Upper Rotliegendes beds downsectionward from barren Kupferschiefer are up to 1 km thick and also contain halite.

Age of mineralization: Uncertain. Depositional age of the Kupferschiefer bed is 258 to 260.4 Ma. Paleomagnetism of Rote Fäule bed alteration gives 255 to 245 Ma (latest Permian and earliest Triassic; late Zechstein to early Bunter time, a time of renewed continental rifting) (Jowett and others, 1987, as modified by Nawrocki, 2000). K-Ar on authigenic illite in Kf shale yields 216 to 190 Ma (Late Triassic to Early Jurassic; later Keuper to Poidonia Shale time—coincides with the Central Atlantic Magmatic Province flood basalts, Brazil, Morocco, Newark basin; mantle plume)(Bechtel and others, 1999). K-Ar on authigenic illite in mineralized Weissliegendes sandstone gives 188–159 Ma (Early Jurassic through Middle Jurassic) (Michalik, 1997). U-Pb on thulcitite (black, shiny, urnao-organic compound in Kf shale gives 175–180 Ma (early Middle Jurassic) (Kucha and Przybyłowicz, 1999). Re-Os gives 240 ± 3.8 Ma from whole-rock samples of Cu-rich mineralization from Lubin mine (Early Triassic, age of the Bunter Sandstone that was deposited in renewed continental rifting events) (Pašava and others, 2007b). Pašava and others (2007a) reported an “altered” age of 279 ± 14 Ma from the Au-Pt-Pd rich zone within Rote Fäule rocks of the Polkowice mine (probably comparable with the “J lead” of Mississippi Valley-type deposits. Probably indicates an epigenetic addition of platinoid metals with old radiogenic 187Re, coming from rocks older than the Kupferschiefer itself).

Hypogene ore characteristics and mineral/metal zoning: Mineral-zone boundaries cross gently downsectionward from southwest to northeast, crossing through the entire host sequence. From inboard (west-southwest) to outboard (east-northeast), the zones are (1) Rote Fäule (hematite); (2) Rote Fäule with native gold, native platinum, native palladium, and minor covellite; (3) chalcocite; (4) bornite; (5) chalcopyrite; (6) galena; (7) sphalerite; and (8) pyrite. Coincidence of chalcocite and bornite zones with the host sequence (above) defines an orebody that is 2–10 m thick more than 15 km northeast-southwest × 28 km northwest-southeast (and continuing northward beyond at depths greater than 1,300 m). Thus, overall, a thin, nearly continuous sheet geometry. Ag in solid solution in chalcocite, digenite, and bornite with minor native silver and stromeyerite. At the scale of the whole deposit, grades are highest in chalcocite zone (all lithologies) and in Kupferschiefer shale bed (mean value of approximately 8.5 percent Cu), but that bed is typically just 30–50 cm thick. About half of all sulfides are in Weissliegendes sandstones averaging about 1.5 percent Cu. Werra carbonate ore has mean values of about 1.3 percent Cu.
**Hypogene gangue characteristics and alteration:** Gangue zonation, other than hematite, is poorly documented. However, alteration to hematitic Rote Fäule is accompanied by dolomitization of the Kupferschiefer bed, which is silicic (as contrasted with argillitic) and calcareous in chalcocite zone, calcareous and marly in outboard zones in Poland. However, Rentzsch (1974, p. 411) shows, instead, dolomitic rocks in Pb, Zn, and pyrite zones and calcite alteration accompanying Rote Fäule alteration (opposite what the first author sampled in Poland), assumedly from Spremberg-Graustein deposit. Elemental analyses show K metasomatism in chalcocite zone, mineral residence unknown.

**Geochemical characteristics:** In Kupferschiefer shale, the following are anomalous and probably enriched in ore processes: As (182 ppm, in trace tennantite, enargite), Co (69 ppm, in linnaeite, pyrite), Hg (3–61 ppm, in native metal alloys and chalcocite), Ni (90–125 ppm, in bravoite, millerite), Sb (15 ppm, in tennantite), and U (39–62 ppm, in organic fraction). Anomalously in Kupferschiefer shale but probably attributable to its eucinic depositional environment: Mo (158 ppm, in castaingite and organic fraction), Se (24 ppm, in clausthalite and organic fraction), and V (182 ppm, in organic fraction). As above, chalcocite zone appears to be K-metasomatized.

**Petrology:** Michalik (2001) is the best. Many others are narrowly focused on sulfides. None have addressed Rotliegendes footwall red-bed petrology, particularly not to look for metals pathways signatures. Good organic petrology (Nowak and others, 2001) documents that liptinite group macerals (most typically, alginate, but also sporinite and lipotdetrinite) in the unmineralized Kupferschiefer are altered to more and more solid bitumen and amorphous unstructured organic matter across the mineralized zone and into the Rote Fäule. Vitinite reflectance typically increases from around 0.72 percent to around 1.13 percent in the same direction, suggesting 130 °C as the maximum temperature reached in the Rote Fäule, thus probably 150 °C and greater for most Cu transport.

**Style(s) of mineralization:** (1) Disseminated within Weissliegendes, Kupferschiefer, and Werra carbonate, (2) lenses flattened within bedding during compaction, within Kupferschiefer, (3) replacements of pyrite framboids and stepwise in the sequence, pyrite, chalcopyrite, bornite, chalcocite, hematite, (4) bedding plane and high-angle veinlets of Cu sulfides ± calcite, (5) special textures (for example, rhythmic layers of dense dissemination in Weissliegendes), mineralized pillars, (6) sparse Rücken veins of Co-Ni arsenides, crosscutting disseminated ore, in faults.

**Post-ore deformation:** Sparse extensional fracturing and veins.

**Other notable features:** Lubin-Sieroszowice is the largest single SSC deposit known in the world. It has been frequently studied. It is, nonetheless, an excellent choice among deposits for new work because it is totally unmetsomorphosed. Work has focused on the deposit itself. Still needed are (1) fluid-inclusion microthermometry work, best done in Weissliegendes sandstone ore authigenic phases (overgrowth quartz and coarse carbonate cements, or in sulfides using IR illumination) (this should allow recognition of appropriate inclusions to study in source and pathway rocks for metals-transporting solutions), (2) fluid-inclusion compositions via PIXE or LA-ICPMS if possible, by thermal decrepitation of an authigenic fraction followed by leach and ion chromatography if micro methods are not possible, to define the mixing that probably went on within Weissliegendes ore and to extrapolate compositions to a metals-transporting solution composition, (3) mineralogy, geochemistry, and mapping in the Rotliegendes footwall rocks and possibly basement rocks to the Rotliegendes to stratigraphic depths as great as possible with the purpose to determine alteration pathways to the deposit, then to determine transport conditions by fluid-inclusion microthermometry and fluid compositions via PIXE, LA-ICPMS, or decrepitate-and-leach within the pathway rocks to define origin of the mineralizing brines. (4) reliable isotope age dating of the mineralizing event(s). K-Ar or Ar-Ar on illite has provided results that lack certainty; it is unclear that the illites date the mineralizing event(s). They may be dating other events in the basin. Re-Os on consistently treated sulfides may succeed—avoid pyrite-containing rock. Try Rb-Sr on sphalerite. Look for xenotome overgrowths on detrital zircon or authigenic monazite in Weissliegendes ore, for U-Pb or Pb-Pb by SHRIMP. Upgrade and separate the heavy mineral content with Wilfley table.


**B. Name:** Kolwezi Klippe, Democratic Republic of Congo, including Dikuluwe, Mashamba West, Mashamba East, Kamoto Principal, Mupine, Musonoi, KOV, T17, Kananga, and Mutsho (Ruwe) mines.

**Deposit subtype:** Reduced-facies

**Location:** 10°43′S 25°24′E (@ Kamoto Principal shaft)

**Grade/tonnage:** 1200 Mt, 4.5 percent Cu, 0.40 percent Co

**Associated deposits:** Relatively nearby and in same host sequence, Shinkolobwe U deposit, Kipushi Zn-Cu-Pb-Ge carbonate-hosted vein-and-replacement deposit

**Regional geologic setting:** Within the Lufilian Arc in Neo-proterozoic Katangan (meta)sedimentary rocks.

**Regional tectonic setting:** Within fold-and-thrust belt; folded and thrust assumedly in the Lufilian orogeny (790–540 Ma with folding and thrusting principally...
parted from the adjacent ecaille, other places almost float within breccia, some places seeming to be merely though folded, upper RAT through Dipeta stratigraphy scales, flakes, slices) of many different sizes of coherent, dissolved evaporites. The effect is that ecaille (French: flake, scale) against polymictic breccias interpreted to be replaced evaporites.

**Primary host rocks and their ages:** Neoproterozoic Katangan, (post-877 ± 11 Ma) Supergroup, Lower Roan Group (Roan 2, R-2), Kamoto Dolomite (R-2.1): in stratigraphic order, the “lower orebody” 6–11 m thick stratigraphically; consisting of locally the topmost beds of Gray RAT (R-2.1.1)—gray chloritic siltstone—then the D.Strat. (stratified dolomite, R-2.1.2.1)—gray, very thinly laminated dolostone—then the RSF (R-2.1.2.2)—gray to black carbonaceous dolosiltite, locally cherty; overlain by 13–25 m of RSC beds (R-2.1.3)—partially silicified stromatolitic dolostone, typically mineralized but not ore grade; overlain by the “upper orebody” in the lower part of Dolomitic Shales Fm. (R-2.2), 8.5–15 m of the SDB (dolomitic shales, basal) beds—gray, silty dolostone—then BOMZ (black ore mineral zone)—thinly laminated black argillitic dolostone. Stratigraphically overlying beds are a continuation of dolomitic shales, carbonaceous dolomitic shales, stromatolitic dolostones, and thin units of massive dolostone, all generally sparsely pyritic.

**Associated red beds:** Subjacent Red RAT (R-1) of the lower Roan Supergroup, base of the sequence not locally known (cut out on tectonic contacts). Red RAT is dolomitic sandstone containing hematite pigment and (or) crystalline hematite overlain by chloritic dolomitic siltstone also with hematite pigment and (or) crystalline hematite. Typically only 5 to 15 m thicknesses exposed below the Kamoto Dolomite before reaching a tectonic contact.

**Associated evaporites:** Trace anhydrite and abundant pseudomorphs of anhydrite in Red RAT, Gray RAT, Kamoto Dolomite, and additional formations upsectionward. Three zones, stratigraphically, are everywhere unconformable and marked by breccias. These are below RAT (between RAT and basal conglomerates that are known only adjacent to basement exposures), between RAT (R-1) and Kamoto (R-2), and between Dipeta Group (R-3) and Mwashya (R-4). From Mwashya upward, stratigraphy continues conformable for several kilometers thickness. The breccias are inferred to mark zones of dissolved evaporites. The effect is that ecaille (French: scales, flakes, slices) of many different sizes of coherent, though folded, upper RAT through Dipeta stratigraphy float within breccia, some places seeming to be merely parted from the adjacent ecaille, other places almost randomly oriented. Ecailles with mineralized R-2 may be continuous for several kilometers but in large continuous ecaille, eventually grade laterally to merely pyritic R-2.

**Age of mineralization:** Two stages. First and predominant is disseminated and replaces anhydrite and is clearly pre-folding. Re-Os age of 816 ± 62 Ma from Konkola, Zambia, is reasonable but imprecise. Second stage is veinlet controlled and overprinted on first and likely postdates the peak of Lufilian orogeny. Similar veins have been dated to 514 ± 2 Ma at Musoshi and to 525.7 ± 3.4 Ma at Nkana-Mindola, Zambia.

**Hydrogeologic ore characteristics and mineral/metal zoning:** Upper orebody is consistently zoned chalcocite to bornite to chalcopyrite stratigraphically upwards at the top. In rare ecaille at Kambove West, the same lateral zonation was observed from orebody center to margin. Carrolite is the major primary Co mineral and is found from chalcocite-bearing rock into chalcopyrite zone. Copper pyrite present from there outward in zonation.

**Hydrogeologic gangue characteristics and alteration:** Gangue zonation, other than hematite and pyrite, is not documented. Quartz and dolomite, together with sulfides, replace early diagenetic anhydrite.

**Geochemical characteristics:** Elemental geochemistry not well documented but demonstrably different than Ore Shale ores in Zambia (Moine and others, 1986). RAT Group rocks in Democratic Republic of Congo are extremely magnesian and almost alkali-free, whereas Ore Shale ores in Zambia are intensely K-metasomatized with K-feldspar. Mg richness is probably not just because of evaporite depositional environment. An important metasomatic addition of Mg is probable.

$\delta^{34}$S isotope study shows an early, broad distribution of $\delta^{34}$Ss of Cu sulfides that replaced earliest diagenetic pyrite retaining the pyrite’s isotope signature, followed by a later, extremely strong addition of relatively heavy $\delta^{34}$S sulfide with values about +3 to +9‰ (Hoy and Ohmoto, 1989). By comparison with sulfur isotope values at the Spar Lake sandstone Cu deposit, this is probably thermochronically reduced S that is a direct reflection of S earlier stored in organic matter, then freed by cracking the organics (interpretation of Hayes).

Fluid-inclusion microthermometry shows two mineralization stages: (1) early cementation disseminated and anhydrite replacement with temperatures 105–220 °C and salinities of 8.4–20.9 eq. wt. % NaCl (Dewaele and others, 2006; El Desouky and others, 2009). Broad ranges of $T_n$ and salinity indicate mixing of extant formation waters of low T and salinity with hot metals-transporting brines, and (2) post-orogenic mineralization (veinlet-controlled) took place from hotter (200–415 °C) even more saline (35–55 eq. wt. % NaCl) brines that almost certainly signal dissolution of evaporites.
**Sediment-Hosted Stratabound Copper Deposit Model**

**C. NAME**: Creta, Okla., United States

**Deposit subtype**: Reduced-facies

**Location**: 34°39′N 99°21′W

**Grade/tonnage**: 5.4 Mt, 1.9 percent Cu, 5.5 g/t Ag

**Associated deposits**: Industrial gypsum

**Regional geologic setting**: Approximately central to the greater U.S. Permian Basin extending from near Big Bend National Park, Tex., north-northeastward to south-central Nebraska; the basin probably opened to the south-southwest.

**Regional tectonic setting**: On the south-southwest-dipping north flank of the Hollis Basin in southwestern Oklahoma. The basin sits on the south flank of the Oklahoma Aulacogen that had thick deposits in the early Paleozone Era and again in the Pennsylvanian. Permian thickness trends appear, instead, to cross the west-northeast trending aulacogen (Mudge, 1967). Creta is underlain by more than 1,000 m of Permian rock, mostly red beds. Relatively flat-lying Permian sediments at surface overlie Early Paleozoic rocks that are considerably folded and offset by major pre-Permian faults.

**Significant structural control (Y/N)**: N

Primary Host Rocks and their ages: uniquely, the Prewitt Copper Shale bed of the Flowerpot Shale, El Reno Group of Late Leonardian-Early Guadalupian Age, over lain by the Haystack Gypsum bed, underlain by Flowerpot red mudstone and siltstone. The Prewitt Copper Shale bed has a drill-demonstrated lateral extent between pinchouts in every direction, except northeastward toward its pre-mining outcrop belt, of about 75 km². Examining isopach trends, perhaps 25% of its original extent was eroded.

**Associated red beds**: Subjacent section more than 1,000 m thick of Permian dolostone, red siltstone and mudstone, and gypsum-anhydrite; red beds predominate (Dixon, 1967; MacLachlan, 1967). Next underlying Pennsylvanian rocks are mostly gray shales of a deeply subsiding basin, lesser coal measures to the east.

**Associated evaporites**: Gypsum and anhydrite are common within the local Permian section and halite appears tens of kilometers to the west particularly in the stratigraphically overlying and partly equivalent Blaine Fm. The greatest thicknesses of salt lie in slightly older and age-equivalent rocks on the opposite, northern side of the Wichita uplift, in the Anadarko Basin and northward into central Kansas (Wellington and Ninnescah Fm., respectively).

**Age of mineralization**: Not known

**Hypogene ore characteristics and mineral/metal zoning**: Zonation is broadly typical. Most abundant ore mineral is anilite (Hagni and Gann, 1985). Digenite, djurleite, pyrite also significant. Argentiferous chalcocite, geerite, yarrowite, chalcopyrite, bornite, covellite, stromeyerite, also present; carrollite, galena and native bismuth in traces (Hagni, 1988). Cu sulfides replace pyrite and replace Trilobites spores (Hagni and Gann, 1976). Minor bornite, chalcopyrite, and galena generally overlie chalcocite zones. In this and other Cu-shale beds in southwestern Oklahoma and the northeastern Texas Panhandle (includes Mangum, Okla., and Medicine Mounds, Tex.—Johnson, 1976b; Smith, 1976), the favorable gray, host shale has the geometry of a “puddle,” an ovoid perhaps 11 km × 7 km and less than 100 cm thick at its maximum. Only that rock probably had early diagenetic pyrite (or pyrrhotite; Hagni, 1988) within rocks that are otherwise red. Carbonized plant debris in sandstones in the same and underlying Permian section are also mineralized (red-bed-type deposits) but lack continuity needed for ore. Occurrence of monoclinic, pseudo-orthorombic chalcocite crystals in ore (Hagni and Gann, 1976) constrains temperature at ore stage to 103.5 °C or less.

**Hypogene gangue characteristics and alteration**: Trace barite with chalocite (Kidwell and Bower, 1976). Gangue minerals generally poorly described but include (semi-quantitatively by XRD) illite approximately 48 percent, quartz approximately 19 percent, chlorite approximately 16 percent, gypsum approximately 16 percent. Adjacent and nearby red shales have more quartz, less gypsum, and similar amounts of illite and chlorite (Lockwood, 1972).

**Geochemical characteristics**: CuO has the following correlation coefficients with other elements and parameters: With quartz XRD abundance −0.30, gypsum +0.15, illite −0.12, chlorite +0.11, Ag −0.49, Pb +0.36, V −0.20,
Co –0.07, Ni +0.64, Fe2O3 –0.31, K2O –0.13, CaO +0.47, Organic Matter –0.11, ZnO –0.13, U –0.65, and MoO3 –0.31 (Lockwood, 1972). The major elements above indicate that the mineralization occurs in gypsiferous gray shale, rather than in non-gypsiferous red shale. δ34S of sulfides range from –1 to –34; gypsum +4 to +6, indicating bacteriogenically fractionated seawater sulfur in early diagenetic pyrite was replaced—Cu sulfides replaced pyrite with little change to the S-isotope ratios. 

**Post-ore deformation:** Sparse extensional fracturing and veins

**Other notable features:** Unreplaced spores are flattened by compaction by at least 2.5×; anilite-replaced spores are not at all flattened by compaction. Replaced spores did not undergo pre-ore pyrite replacement. Thus, Cu-sulfide mineralization preceded compaction almost entirely (Huyck and Chorey, 1991).

**Important references:** Hagni and Gann (1976); Johnson (1976a, b); Kidwell and Bower (1976); Lockwood (1972); Hagni (1988); Huyck and Chorey (1991).

**D. Name:** Udokan, Transbaikalian Russia

**Deposit subtype:** Sandstone Cu

**Location:** 56°34′37″ N 118°30′47″ (@ the junction of Naminga R. and a major tributary within the vertical projection of the deposit)

**Grade/tonnage:** 1,528 Mt, 1.45 percent Cu, 12.6 g/t Ag (V. Chechetkin, oral commun., 2010)

**Associated deposits:** Sediment-hosted U (details unavailable). Deposits of Fe, Ti, V, Cu, and PGE associated with the Chenel layered mafic-ultramafic complex are nearby; appear to be younger than Udokan, include basal zones of pentlandite-chalcopyrite ores in Udokan Supergroup wallrocks.

**Regional geologic setting:** Miogeosynclinal sequence within synclinal “troughs” within Archean craton

**Regional tectonic setting:** Southern margin of the Archean Aldan Shield in “troughs” filled with metasediments of low metamorphic grade—middle greenschist

**Significant structural control (Y/N):** N

**Primary host rocks and their ages:** Upper Sakukan Formation of the Kemen Series, Paleoproterozoic Udokan Supergroup. Paleoproterozoic-age sedimentation based on a metatuff dated 2,180 ± 50 Ma (U-Pb, zircon) in the conformably underlying and similarly metamorphosed Chinei Group. The favored lithologic package (U. Sakukan) is of fine- to coarse-grained, well-sorted quartzite with calcareous cements and with mud intraclasts varying locally up to 30 v. % of the rock. Overlying (Naminga Formation) rocks are argillitic and considered transgressive over the deltaic environments of the Sakukan.

**Associated red beds:** Lower Sakukan Formation is 1,000–1,700 m of “magnetite carrier sandstone” and below that the Talakan Formation is 1,100–1,400 m also with much “magnetite carrier sandstone.” The origin of this magnetite, called detrital in the literature, is suspected to be burial metamorphic after hematite of former red beds similar to Belt Supergroup, United States (Nadoll and others, 2012).

**Associated evaporites:** None known locally. Carbonate-rich units downsection by more than 3 km have hopper casts. Overlying argillitic rocks have abundant desiccation cracks but no mention of evaporites or pseudomorphs.

**Age of mineralization:** Unknown

**Hypogene ore characteristics and mineral/metal zoning:** Strongly zoned pyrite to chalcopyrite to bornite to chalcocite (to hematite?)! The mineralogical zoning defines several anastomosing lens- to layer-form bodies of ore within the Upper Sakukan, with lengths up to 6.1 km, widths of typically 0.8 km but up to 3 km, and thickness 20 to 130 m (commonly 70 to 100 m). Complex inter-tonguing relations of sulfide mineral zones. Coarser lithology commonly bears chalcocite whereas finer lithology carries chalcopyrite and (or) pyrite. Ag is mostly in solid solution in chalcocite (10–3,000 ppm Ag) and bornite (10–300 ppm). Au should be recoverable at 0.5 g/t in concentrate.

**Hypogene gangue characteristics and alteration:** Includes (“idiomorphic, regenerated”) magnetite, probably calcite, trace barite, but not described to accompany specific sulfide zones. Albiteization in cross-stratal zones described throughout the Udokan Supergroup and interpreted to be localized along faults (Abramov, 2008). Albiteization described to accompany sediment-hosted U deposits nearby, also in the Udokan Supergroup (V. Chechetkin, oral commun., 2010), but no particular association of albite with mineralization described.

**Geochemical characteristics:** Anomalous in ore are Co (10–100 ppm), Ni (10–100 ppm), Ag (trace to 100 ppm), Pb (10–100 ppm), Mo (less than 10 to rarely 100 ppm), V (10–1,000 ppm), Ba (trace to 1 percent) (probably mostly in barite), and Mn (0.1 to 1 percent) (probably sited in calcite). A suite of elements probably sited in former ilmeno-magnetite, ilmeno-hematite, and heavy
minerals is also found including Ti (everywhere greater than 1,000 ppm), Zr, and Cr (Bakun and others, 1966, table 2). Organic carbon co-varies with Cu in mineralized rocks; Corg greater than 0.3 percent corresponds to Cu grades greater than 1 percent (Volodin and others, 1994, p. 10). No data on stable isotopes or fluid inclusions.

**Petrology:** Not described in literature available in English though hosts are feldspathic (10–15 percent total feldspars; plagioclase 10–30 percent, K-feldspar 5 percent) to sericitic quartzites (5–15 percent muscovite).

**Style(s) of mineralization:** (1) μ-scale sulfides enclosed within magnetite, calcite, and epidote, (2) sandstone cements, and (3) replacements of framework grains to form patchy disseminations, lenses, and laminae of sulfides varying up to massive replacements.

**Post-ore deformation:** Broadly folded into an open syncline with its south limb locally overturned to the north, cut and offset by high-angle faults, also cut by thick gabbro-diabase dikes, and by later, thin quartz porphyry and lamprophyre dikes.

**Other notable features:** Udokan is the largest known sandstone Cu deposit in the world.

**Important references:** Bakun and others (1966); Bogdanov and others (1966); Volodin and others (1994); Abramov (2008)

E. **Name:** Spar Lake, Mont., United States (also known as the Troy mine)

**Deposit subtype:** Sandstone Cu

**Location:** 48°13′49″ N, 115°54′19″ W

**Grade/tonnage:** 80.6 Mt, 0.63 percent Cu, 46 g/t Ag

**Associated deposits:** None, though others within the basin include sedex (for example, Sullivan, Sheep Creek) and mesothermal Ag veins (Coeur d’Alene). Laramide overprint of the Belt Basin includes porphyry Cu, porphyry Mo, and polymetallic veins.

**Regional geologic setting:** Host rocks are deltaic topset beds; within a belt where former red-bed host rocks are bleached or partly bleached in contrast to outside the belt where all are still red beds; the belt may have been a structural high as indicated by lesser deposition of Missoula Group sediments in the “post-Ravalli dome” (Harrison, 1972).

**Regional tectonic setting:** Within probable aulacogen; probable association with growth faults that may have been used as conduits of metals-transporting brines (fault offsets may have alternatively or also contributed to pre-ore natural gas trapping).

**Significant structural control (Y/N):** Y, the East Fault, a down-to-the-east normal fault. Deposit occurs as two connected lobes elongate parallel to and on opposite sides of the East Fault within Revett Formation, upper member, host sandstones. Zonal geometry suggests that metals-transporting groundwater came upward across stratigraphy, then spread laterally within sandstones that had already been offset, boundaries of lobes at places where confining beds were faulted into contact with sandstones.

**Primary host rocks and their ages:** Mesoproterozoic Belt Supergroup (greater than 1470—less than 1400 Ma deposition); Ravalli Group; Revett Formation (lithostratigraphic metasandstone-rich package); upper member; lower, middle, and upper quartzite beds; and lower member; C quartzite beds; depositional environments of host quartzites included beach-and-offshore bar and subtidal channels and bars.

**Associated red beds:** Burke Formation (1–2 km thick, lavender-gray, hematite- and magnetite-bearing siltite greater than argillite greater than quartzite) and laterally equivalent Revett, also lavender-gray. Burke is 500 m downsection from host beds, separated from it by beds that are trace-mineralized or ore-mineralized.

**Associated evaporites:** Nearest known are Missoula Group (greater than 4 km upsection); none known at or below the host level stratigraphically, though more examination is due certain breccias within Prichard Formation greater than 3 km downsection (Cressman, 1985, p. 44–51).

**Age of mineralization:** 1409 ± 8 Ma in rocks that were deposited between 1470 and 1454 Ma.

**Hypogene ore characteristics and mineral/metal zoning:** Nested tongues expand laterally within sandstone layers, contract within intervening silty argillitic layers. Pyrite to galena to chalcopyrite to bornite (+ magnetite) to digenite-chalcocite (+ hematite).

**Hypogene gangue characteristics and alteration:** Ferroan calcite with pyrite, galena, chalcopyrite. Manganiferous calcite with bornite-digenite. Ankerite, Fe-rich chlorite, hematite with digenite-chalcocite. Barite together with chalcocite, digenite, bornite, and rarely, chalcopyrite or galena. Intense potassium metasomatism as potassium feldspar overgrowth on detrital Kspar grains within galena-chalcopyrite zone, increasing into bornite-digenite-chalcocite zone. Silicification as quartz overgrowth on detrital quartz. Late, cementing and replacing albite increases in the direction of chalcocite-hematite.

**Geochemical characteristics:** Elementally, the outstanding characteristics are K metasomatism accompanying ore deposition and Ca metasomatism in the distal-fringe zones. Additional elements transported with Cu and deposited within the zoned array are Ag, Pb, Sb, As, Ba, Na, S.

Fluid inclusions T° and salinity indicate mixing of a greater than 175 °C metals-transporting brine with a cooler, less-saline formation water. Fluid-inclusion
volatile compositions indicate mixing a 200 °C metals-
transporting brine with reservoired sour gas in the host
sandstones. There are systematic arrangements of other
hydrocarbon volatiles, of H2S, probably of H2S. No indica-
tion of magmatic volatiles or of any mantle contributions.
C isotopes of carbonates are light in fringing zone of Mn-
and Fe-rich calcites (see below). O isotopes of carbon-
ates indicate either an elevated T brine or freshwater (the
latter unlikely).

S isotopes show a spread from +2 to +29 but no nega-
tive values. S-isotopic equilibrium was approached
between galena, chalcopyrite, and H2S at ore-stage and
with declining temperatures, 130 °C to less than 100 °C,
outward from ore across the galena fringe. History of S is
(1) bacterial sulfate reduction of seawater sulfate
that started with δ34S of about +20, (2) sequestering
and storage of the then-reduced S in organic matter of
downsection Prichard Formation, (3) maturation and
migration of hydrocarbons carrying the S, (4) thermo-
chemical sulfate reduction from formation waters and
addition to the migrating hydrocarbons, (5) entrapment
of sour gas with continued thermochemical sulfate reduction
from formation waters, (6) ore-stage reaction of metals
in hot brine with rapid deposition across a mixing zone
utilizing the H2S of the sour gas, and accompanied at the
distal fringes of the mixing zone by ore stage coupled
redox between SO4 of the metals-transporting solution
and the C in CH4 of the sour gas, depositing sulfides with
heavy δ34S and manganiferous calcite with light δ13C.
The sulfide thus formed was thermochemically reduced.
Direct deposition of barite from metals-transporting
solution SO4, not in thermal equilibrium with the metals-
transporting brine.

Petroleum: Ore minerals and ore-stage gangue are
dominantly cements and replacements. Earliest stage
diagenesis probably forms red beds (ferric oxide
grain coatings). At the stage of petroleum migration, red
beds were bleached and porosity increased from loss of
ferric-oxide grain coatings. Emplacement of sour gas in
reservoir rocks was accompanied/followed by deposition
of pyrite, possibly some Fe-rich calcite, possibly some
apatite, possibly some tourmaline overgrowth. Invasion
of the gas reservoir was accompanied by pyrite replaced
by galena-chalcopyrite, replaced by bornite, replaced
by digenite-chalcocite; all accompanied by K-feldspar
overgrowth, quartz overgrowth, manganiferous calcite
cementation, chlorite cementation, barite cementation, late
(in the ore-stage) hematite cementation and albite cen-
tement and replacement.

Style(s) of mineralization: Ore minerals and ore-stage
gangue are predominantly cements and replacements of
earlier cements and clasts. Many bedform dissemina-
tions and clots of the ore minerals and zonally distributed
gangue minerals. Notable mineralized large seepage pil-
lars with local-scale evidence of rising metals-transporting
solutions. Minor (less than 5 percent) ore sulfides in later,
post-metamorphic veinlets in brittle fractures with demon-
strable contribution of ore components to veinlets from
local wallrocks (nowhere the opposite – veinlets contrib-
uting ore components to the local wallrocks).

Post-ore deformation: Pure burial metamorphism to
the biotite isograd of greenschist facies with estimated
conditions of 425°C and perhaps 1.5 kilobars. Except
within local clots of anomalous composition, sulfides
melted into solid solutions within the confines of each
former pore, but enclosing carbonates and silicates did
not melt or react. Within former hematite-sulfide clots,
magnetite, ankerite formed by de-oxidation of hematite. In
retrograde metamorphism, sulfide solid solutions exsolved
to incoherent bornite-digenite, coherent bornite-digenite-
chalcopyrite, and others. Folding and thrusting in the
Mesozoic was not probably at metamorphic conditions
and was accompanied by reverse faulting. Relaxation of
Laramide compressional stresses accompanied by brittle
fracturing, veinlet formation, and (renewed, at places)
normal faulting.

Other notable features: Silver predominantly in solid
solution in chalcocite, digenite, and bornite with 0.1 wt. % Ag
typically found in each sulfide. Native silver, stromeyerite,
and argentite or acanthite rare to trace. Every aspect of
genesis of Spar Lake, except post-ore metamorphism and
tectonism, is matched by Zilandy-type deposits in the
Dzhezkazgan district, Kazakhstan (B. Syusyura, oral com-
umun., 2009).

Important references: Hayes and Einaudi (1986); Hayes
and others (1989); Hayes (1990); Boleneus and others
(2006); Hayes and others (2012)

F. Name: Juramento, Argentina

Deposit subtype: Sandstone Cu

Location: 25°13′S, 65°09′ W

Grade/tonnage: 44.7 Mt, 0.80 percent Cu, 21.8g/t Ag

Associated deposits: None known.

Regional geologic setting: Within belt of folded Mesozoic
sediments in the foreland between the Andean continental
arc and the Rio de la Plata craton

Regional tectonic setting: Within intracontinental rift of
Jurassic-Cretaceous age associated with the breakup of
South America from Pangean Africa

Significant structural control (Y/N): N

Primary host rocks and their ages: Late Cretaceous
(Maastrichtian) Lecho and Yacoraite Formations in
ascending order; white to cream calcareous sandstone
and oolitic, stromatolitic limestone with lesser calcareous
siltstone, respectively
**Associated red beds:** Late Cretaceous (Campanian) Pirgua Supergroup red coarse clastics with minor bimodal basalt-rhyolite; total thicknesses up to 3,000 m in the adjacent basin

**Associated evaporites:** Only evaporites nearby are those of the early Tertiary Tunal Olmedo Formation, hundreds of meters upsectionward from host rocks. Host limestones contained blades of early diagenetic gypsum in trace amounts, now replaced by Cu sulfides.

**Age of mineralization:** Unknown. Gravity-driven groundwater circulations were available in Early to Late Miocene and Plio-Pleistocene

**Hypogene ore characteristics and mineral/metal zoning:** Strongly zoned pyrite to sphalerite to galena to minor tetrahedrite to chalcopyrite to bornite to chalcocite (to hematite) both downsectionward across bedding and from north to south in steeply west-dipping sedimentary host rocks.

**Hypogene gangue characteristics and alteration:** No gangue minerals interpreted to date from ore-stage (illogical). Early meniscus and isopachous calcite cements and frambooidal and other pyrite followed by coarse dolomite, apparently followed by ore sulfides in the sequence galena, sphalerite, tetrahedrite, chalcopyrite, bornite, and chalcocite. Those apparently followed by quartz and K-feldspar. Followed by stylolytization. Followed by dead oil staining in relic pores and dead oil in veinlets.

**Geochemical characteristics:** Insufficient data but sufficient indication to state that As in tetrahedrite-tennantite is anomalous in a fringe zone. Zn and Pb anomalous in a more distant fringe.

S isotopes (n=12 samples) of sulfides range from δ34S of −42.9 to −8.0. Pyrite mean is −37.8. Base metal sulfide mean is −17.0. By comparison with Spar Lake sandstone −42.9 to −8.0. Pyrite mean is −37.8. Base metal sulfide mineralization with sequence sphalerite, galena, tetrahedrite, chalcopyrite, bornite, and finally chalcocite, (5) quartz and K-feldspar cements and replacements, (6) major compaction, carbonate neomorphism, and stylolytization, (7) hydrocarbons (now dead oil) coating relic pores followed by 3rd calcite cement (not further described), and “late” fracturing with filling by dead oil, and (8) uplift and supergene alterations. As described, this sequence is too rigorous, as above. Instead of rigorously “first this, then that,” numerous textures could alternatively have been interpreted “synchronously and later than.” (for example, Authigenic quartz and K-feldspar that cement ooid grainstones and partially replace ooids probably accompanied Cu-sulfide deposition, and it may have been happening while compaction with stylolytization was ongoing.)

**Style(s) of mineralization:** Cu sulfides are predominantly cements and replacements in carbonate grainstones in the Lecho sandstones and Yacoraite grainstones and siltstones. Replacement of gypsum by Cu sulfides.

**Post-ore deformation:** Mild. Open folding rarely with vertical or overturned dips. Relatively intense post-ore fracturing. Essentially no metamorphism.

**Other notable features:** SSC deposits occur, as well, within Middle- and Late-Cretaceous sandstones in the Neuquén Basin in west-central Argentina, several hundred kilometers south of Juramento. All occurrences there are classic sandstone Cu deposits mineralizing rock that was first red, then bleached and associated with petroleum, and finally, Cu mineralized (J. Thorson, private consultant, written commun., 2009; Giusiano and others, 2008).

**Important references:** Durieux and Brown (2007); Alexander Mining (2005).

**G. Name:** Corocoro, Bolivia

**Deposit subtype:** Red-bed Cu

**Location:** 17°10′09″ S, 68°27′04″ W

**Grade/tonnage:** 7.8 Mt, 7.1 percent Cu, 106 g/t Ag (These grades, reflect hand-sorting of ore. Average grades closer to those mentioned under Other Notable Features, below.)

**Associated deposits:** Salt diapirs

**Regional geologic setting:** Bolivian altiplano between eastern and western Andean Cordillera. Intermontane basin in hyperarid climate.

**Regional tectonic setting:** Strong halokinesis produced salt diapirs and probably the related anticlines and synclines, as well as high-angle reverse faults.

**Significant structural control (Y/N):** Y. Steep dipping dragfolded flanks of salt domes and halokineti...
Primary host rocks and their ages: In ascending order, Corocoro Group, Huayllamarca Formation (locally called Ramos Formation) red mudstone and lesser red sandstone and gypsum with local bleaching, and Totora Formation (locally called Vetas Formation) gray sublitharenites with lesser red mudstone.

Associated red beds: Within the locally 10 km-thick Corocoro Group. The group is dominated by red beds with no less than 2,700 m of red beds downsection.

Associated evaporites: Lower Tertiary or Upper Cretaceous Jalluma or Chuquichambi Formation salt; expressed as gypsiferous shale breccias at surface. Halite penetrated in drilling at 386 m.

Age of mineralization: Not known.

Hypogene ore characteristics and mineral/metal zoning: Native copper dominates in sandstone-hosted ores in bleached Ramos. Chalcocite replaces carbonized plant fragments and forms cements in Vetas gray sandstones. Multiple “manto” levels and also bedding plane and fault-filling veins of native copper. Minor bornite, covellite, chalcocytire, galena, sphalerite, stromeyerite, and tennantite are also locally present.

Hypogene gangue characteristics and alteration: No gangue minerals described as ore stage. Overgrowth quartz, authigenic K-feldspar, celestite, and barite likely date from ore stage. Those minerals cement sandstones and are replaced along with earlier cements (hematite dust rims, clays, gypsum, and albite) by chalcocite or native copper. Gangue also includes anhydrite, aragonite, and chalcedony. Pre-ore bleaching in Ramos speculatively caused by a “mobile reductant” (dissolved humic acids and their precipitates) (Flint, 1989, p. 394–395).

Geochemical characteristics: Little-described. Chalcocite or native copper is accompanied by trace domeykite (isometric Cu₃As). Ore is said to have 15 g/t Ag per 1 percent Cu.

No fluid-inclusion studies, but dissolved evaporites in the metals-transporting brines are likely with the occurrence of mantos in beds dipping steeply off flanks of salt domes and adjacent to related reverse faults.

Petrology: Earliest diagenesis of sandstone hosts (probably the Ramos) is hematite and clay dust rims and gypsum. Next is albite. These were followed apparently by quartz overgrowth, then K-feldspar, and celestite and barite. These were followed by generation of secondary porosity by destruction of hematite dust rims and dissolution of plagioclase in detritus. Followed by Cu sulfides and native Cu and slightly later native Ag. Followed by covellite, secondary Cu minerals, gypsum, and quartz. Minor supergene enrichment as native Cu, chalcocite, and covellite. (Like Juramento, this paragenetic sequence gives interpretations only as “first this, then that.” Some could have been interpreted “synchronous with or later than.” The likelihood is that quartz overgrowth, authigenic K-feldspar, and barite are all ore stage.)

Style(s) of mineralization: Sandstone cements predominate. Bedding plane veins important at places. In Vetas, replacement of carbonized plant debris important. Observation that basal zones of each sandstone are favored might be interpreted that these are the coarsest, most permeable parts of each fluvial fining-upward sequence.

Post-ore deformation: Little or none.

Other notable features: Grades reflect hand sorting of ores. Corocoro is the longest-mined, best known of more than 20 deposits and districts on a north-northwest trend of salt diapirs with related SSC deposits that extends 200 km. Others on the trend from north-northwest to south-southeast are Porvenir mine, Llallagua mine, Maria Elena-Victoria mine, Pisaqueri mine, American Unidas mine, Transvaal mine, (Corocoro district—8 mines), Veta Verde mine, El Hogar mine, Ne group, San Augustin mine, San Francisco prospect, San Miguel mine, Chacarilla district (Porfia, Congreso, Amigo, Borda, Esperanza Eurea, San Fermin, and Tiviña mines and prospects), and Chuquichambi district (Llanquera-San Miguel and Santa Maria prospects). Corocoro district production 1963–1976 was 1,457,000 t of 2.55 percent Cu. Reserves in 1992 were 1,186,000 t of 2.78 percent Cu.

Important references: Singewald and Berry (1922); Ljunggren and Meyer (1964); Entwistle and Gouin (1963–1976), followed by 1,186,000 t of 2.78 percent Cu. Reserves in 1992 were 1,186,000 t of 2.78 percent Cu.

H. Name: Nacimiento, N. Mex., United States

Deposit subtype: Red-bed Cu

Location: 35°59′41″ N, 106°53′54″ W

Grade/tonnage: 10.0 Mt, 0.67 percent Cu, 2.4 g/t Ag

Associated deposits: None

Regional tectonic and geologic setting: Flank between the Nacimiento Precambrian-cored uplift and the San Juan Basin.

Significant structural control (Y/N): Probably N

Primary host rocks and their ages: Middle to Late Triassic Chinle Formation, basal Agua Zarca Sandstone Member (correlative with Shinarump Conglomerate of the following example deposit).

Associated red beds: Underlying Early to Middle Permian Abo and Yeso Formations (red-brown mudstone and lenticular red-brown and greenish gray arkosic sandstone; and reddish-orange-buff very fine-grained quartzose sandstone, respectively), totaling 265 m thick locally (much more to the east, north, and northwest); overlying Salitral Shale and Upper Shale Members of the Chinle totaling 245 m thick (with intervening Poleo Sandstone Member of gray sandstone with carbonaceous plant debris).
**Associated evaporites:** Thick Permian sections to the east in the Permian Basin, particularly in southeastern New Mexico, and Pennsylvanian sections to the northwest in the Paradox Basin, southwestern Colorado and southeastern Utah. None locally.

**Age of mineralization:** Not known, but likely pre-Tertiary. Cellular structure of wood was not collapsed before the wood was mineralized (petrified), but the first mineralization of the wood was not necessarily the Cu sulfides found replacing it now.

**Hypogene ore characteristics and mineral/metal zoning:** Principally chalcocite replacing carbonized plant debris up to the size of logs. Individual logs have pyrite replaced by bornite replaced by chalcocite replaced by covellite. Native silver described from near the modern water table (probably supergene).

**Hypogene gangue characteristics and alteration:** None described. Host sandstones are 90–95 percent quartz, 2–5 percent feldspar, 1–5 percent chert, 1–2 percent magnetite, 1 percent muscovite, traces of zircon, tourmaline, chlorite, and biotite. Authigenic quartz overgrowth described but host rocks are friable and have clayey matrix (clay minerals not identified).

**Geochemical characteristics:** Not described.

**Petrology:** Carbonized logs have wood cell walls replaced by or edged by bornite and cells filled by pyrite that was later replaced by chalcocite. Relict wood found as films of carbonaceous materials in boundary lines between the former cells. Quartz overgrowth appears to precede chalcocite cements (but could be interpreted as synchronous, as above).

**Style(s) of mineralization:** Dominating replacements of carbonized wood (deposit could be described as a replaced log-jam in sandstone). Lesser sandstone cements. Supergene malachite and chrysocolla cements form halos around sulfidized logs and other sulfides in porous sandstones.

**Post-ore deformation:** Host rocks have been folded and post-ore offset by faults.

**Other notable features:**

**Important references:** Lindgren and others (1910); Soulé (1956); Elston (1967); Woodward and others (1974)

I. **Name:** Happy Jack, Utah, United States

**Deposit subtype:** Red-bed U-Cu

**Location:** 37°45′06″ N, 110°17′37″ W

**Grade/tonnage:** 0.573 Mt, ~0.74% Cu, 0.25% U, ~0.047% V₂O₅; tonnage and U grade from Chenoweth (1993, p. 21), Cu and V₂O₅ grades from splits of 5 uranium ore shipments to the Monticello, Utah, refining plant (Shoemaker and others, 1959, p. 27).

**Associated deposits:** 24 other small U-Cu deposits in Shinarump channels cut into the underlying Moenkopi Formation in the White Canyon district.

**Regional geologic setting:** Gently folded package of siliciclastic sedimentary rocks on the southwest flank of the Paradox Basin of Pennsylvanian-Permian age.

**Regional tectonic setting:** Colorado Plateau; a long-term stable region with only gentle folding, lying between the region of steeply folded, Precambrian-cored Rocky Mountain uplifts and the Sevier fold-and-thrust belt.

**Significant structural control (Y/N):** N

**Primary host rocks and their ages:** Upper Triassic Chinle Formation, basal Shinarump Conglomerate Member, very fine to very coarse grained and locally conglomeratic quartzose (approximately 5 percent K-feldspar) sandstone with lesser siltstone and minor mudstone and with conspicuous concentrations of carbonized plant debris. Occurs in channel complexes with bases eroded into the underlying dusky red Moenkopi siltstone/sandstone. The Happy Jack Channel is 230 m wide and cut 3 m down into the Moenkopi, trending westward into the subsurface (from a cliff). The channel is a limit on the tonnage of the deposit that sits entirely within it.

**Associated red beds:** Underlying, predominantly red-bed sandstone section including, in ascending order, Pennsylvanian-Permian Rico Formation, Permian Cutler Formation, and Lower Triassic Moenkopi Formation, totaling about 540 m thick; Chinle is predominantly red beds about 155 m thick interrupted by two gray sandstone units, at the base (Shinarump) and in the middle (Mossback Member).

**Associated evaporites:** Beneath the Rico: Paradox Formation evaporites of the Pennsylvanian Hermosa Group; consisting of 29 cycles of, in ascending order; black calcareous shale; bioturbated, silty, sucrosic dolomite; laminated anhydrite; halite with or without potash; nodular anhydrite; black calcareous shale. The Paradox evaporites, as much as about 3,000 m thick along the basin axis just southwest of the Uncompahgre uplift in southwestern Colorado, thin to the southwest, pinching out by facies change to shelf carbonate-clastic cycles about 40–50 km to the northeast of Happy Jack, so no known evaporites vertically downsection from ore. A second basin with Pennsylvanian evaporites lies more than 150 km to the south, the Holbrook Basin, Ariz.

**Age of mineralization:** Not known but definitely older than 23 ± 3 Ma (Miller and Kulp, 1963). Probably, it is of similar age to U deposits in the Shinarump in Lisbon Valley, Utah, which are 197 ± 15 Ma in host rocks that were deposited by about 210 Ma. Deposits are epigenic, though.

**Hypogene ore characteristics and mineral/metal zoning:** U-Cu ores include ore layers in siltstone or at contacts with sandstone, replacements of carbonaceous trash zones,
and replacements of larger carbonized wood fragments. Most of uraninite and Cu sulfides replace carbonized plant debris; less is cementing in the sandstone. Uraninite is commonly microscopically associated with chalcopyrite. Chalcocite is predominantly supergene, veinining and replacing bornite and chalcopyrite tens of meters laterally from outcrop with only minor grade enrichment. Large-scale zonation has not been recognized. Minor galena and sphalerite also replace carbonized wood. Paragenesis: pyrite is replaced by galena, then chalcopyrite, then bornite, then uraninite, then sphalerite (Chalcocite considered entirely supergene).

**Hypogene gangue characteristics and alteration:** Quartz overgrowth is abundant. Local poikilotopic calcite may or may not be ore-stage and hypogene. Minor barite cements and crystals growing into pores in carbonized wood.

**Geochemical characteristics:** The weighted average of four ore shipments totaling 210 short tons contained the following concentrations of elements by semiquantitative, 15-class, X-ray spectrography, in parts per million: Al—15,000; Fe—30,000; Mn—70; Ca—1,500; Mg—700; Ti—1,500; Na—300; Ba—150; Sr—70; Zr—150; Cr—30; Sc—15; B—15; Y—30; V=540 (by Quantitative Spectrography); Ni—30; Co—700; Cu—7,000; Pb—150; U—7,700 (by Fluorimetric); Mo—70; Ag—3; Be—1.5; Zn—1,500 (A. Miesch, U.S. Geological Survey, retired, written commun., 1981). Ilsemannite and erythrite are secondary minerals found locally, signaling anomalous Mo and Co, respectively. Black Mn oxides with greater than 10 wt. % Co are found locally (given the Mn and Co concentrations above, these are probably heterogenite).

**Petrology:** Uraninite preferentially replaces the carbonized wood cell walls whereas Cu sulfides, bornite, and especially chalcopyrite fill the cell lumens. (Note the contrast with the Cu-dominated system at Nacimiento where the cell walls were bornite-replaced, and the cell lumens were filled in the sequence pyrite, then chalcocite.) Pyrite and chalcopyrite form pore-lining and pore-filling sequences in sandstone pores at places. Sulfides replace detrital quartz and K-feldspar, and locally also, overgrowth quartz.

**Style(s) of mineralization:** Predominant replaced carbonized wood. Lesser cements in sandstones. Replacements of both detrital feldspar and quartz and overgrowth quartz.

**Post-ore deformation:** Rocks have been gently folded to dip about 2 to 3° to the southwest on the west flank of the Monument Valley Upwarp

**Other notable features:** Other sandstone-hosted U deposits in the Chinle on the Colorado Plateau have dramatically different concentrations of Cu and V (Hayes, 1982).

**Important references:** Trites and Chew (1955), Isachsen and Evensen (1955), Miller (1955), Thaden and others (1964), Chenoweth (1993).