Diagenesis #13. Origin of chert: Diagenesis of biogenic siliceous sediments

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Introduction
Highly silicified sedimentary rocks are generally called chert, but are also known under generic or local names such as radiolarite, porcellanite, flint, jasper, lydite, or novaculite. They may contain up to 95% or more silica. The origin of chert has been much debated in the past. Systematic studies in the last decade have advanced our knowledge of chert genesis significantly and have shed light on many facets of the so-called "chert-problem." The three basic aspects of the chert-problem, which are (1) the source of the silica, (2) the depositional environment of siliceous sediments and (3) the diagenesis of cherts and chert-bearing rocks are now understood in principle. Nevertheless, a historical perspective will be included in this review in order to give the reader some idea of the developments that led to our present understanding of silica diagenesis and to the problems on which current research is focussed. Although the emphasis of this review is on diagenesis, a brief outline of the first two aspects of the former chert-problem will be presented in the introductory sections because a discussion of silica diagenesis is not possible without frequent reference to these other subjects.

Terminology and concentration units for dissolved and solid SiO₂ phases and fine-grained siliceous sediments. To facilitate communication, a few of the terms and concentration units used for the various dissolved and solid species of SiO₂ as well as for fine-grained siliceous sediments will be defined at the outset:

\textit{Dissolved silicon} (or \textit{silicon}) is the term used for the undissociated monomeric silicic acid \(\text{Si(OH)}_n\) which is the dominant species of dissolved SiO₂ in most natural waters. Its anhydrous equivalent SiO₂ has a molecular weight of about 60. One ppm SiO₂ by weight (or 1 mg kg\(^{-1}\)) therefore corresponds to 16.6 microgram-atoms SiO₂ per litre (µg-at L\(^{-1}\); the concentration unit commonly used in the oceanographic literature) or 16.6 µM (where µM stands for micromole per litre). This in turn is equivalent to 0.467 mg L\(^{-1}\) Si. Conversely, 1 mM (= 1 millimole per litre) corresponds to 60 ppm SiO₂.

In its various solid phases, SiO₂ will be referred to as silica. These include amorphous silica or opal-A, cristobalite-tridymite or opal-CT, low-temperature cristobalite or opal-C, low-temperature tridymite, chalcedony, quartzine, lutecite and quartz. Physical and mineralogical characteristics of these species are discussed in appropriate sections of the paper.

Biogenic fine-grained siliceous sediments are called siliceous coazes. The term "siliceous mud" instead of ooze is used for those deposits which have a substantial detrital terrigenous component. Diatom ooze, radiolarian ooze and (sponge) spicule ooze are distinguished according to the main groups of organisms contributing siliceous shells to pelagic (or lacustrine) sediments. Their lithified equivalents are diatomite, radiolarite and spiculite.

The rock names porcellanite and chert are used to denote fine-grained siliceous sedimentary rocks according to their textual and physical properties. "Porcellanite" refers to a porous rock with dull or matte lustre similar to unglazed porcelain whereas "chert" denotes a dense vitreous, hard and brittle rock (Bramlette, 1946). The difference between the two is related to detrital content, especially clay content, which is higher (25-50%) in porcellanite than in chert (0-25%, Jones and Murchev, 1986).

Siliceous rocks with more than 50% clay are called siliceous claystone, mudstone, or argillite. Lydite is a dark-grey to black variety of argillaceous chert or siliceous argillite. Local names for various chert types include flint, jasper, and novaculite. Flint is a name widely applied to chert nodules in the Upper Cretaceous chalk of northwestern Europe (Schmid, 1986), which have been used in artifacts and tools made by early man (arrowheads, knives, drills, etc.). The term is synonymous with firestone or hornstone and has been applied in the common language in England for more than a thousand years to any very hard and dense rock with conchoidal fracture surfaces (Bates and Jackson, 1980). Generally, jasper is a red, brown or yellow variety of chert (also green or black) that derives its colour from iron-oxide impurities. It may be associated with iron-ores or soil-forming processes. Novaculite is very dense, light-coloured quartz-chert from the Ouachita Mountains in Arkansas and Oklahoma, which has undergone high-grade diagenesis and low- to intermediate-grade metamorphism. The word is derived from the Latin word for razor (novacula) and has been formerly used in England for rocks that served as whetstones or razor hoses (Goldstein, 1959).

Sources of Non-detrital Silica in Siliceous Sediments
There are three main sources of non-detrital silica in sediments: (i) siliceous tests and skeletal elements of organisms, (ii) weathering solutions in semi-arid climates, and (iii) silicon supplied in solution by hydrothermal-volcanogenic systems. The predominant source of Phanerozic cherts is biogenic, although there also exist hydrothermal-volcanogenic deposits or low-temperature chemogenic deposits (in alkaline lakes or arid to semi-arid soils). In the Precambrian, the cherty variety of banded iron formations represents a widespread type of chert almost unknown from younger geologic periods whose origin is still being debated.

Low
temperature chemogenic and/or hydrothermal-volcanogenic deposits are the dominant Precambrian types of chert. This review focuses on the diageneis of biogenic siliceous sediments, whereas another review (Hesse, in press) deals with other chert types.

The total present-day silica production by marine organisms such as diatoms, radiolarians, silicoflagellates or siliceous sponges is estimated at 2.5 x 10^14 g annually (g. a. -1) or about 25 times the input of silica to the oceans (10.8 x 10^14 g. a. -1), to which rivers contribute 4.3, pore-water reflux a potential 5.7, submarine weathering 0.8, and submarine volcanoism 0.05 x 10^14 g. a. -1, according to Heath (1974). More recent calculations by Edmond et al. (1979a) attribute an additional 1.9 x 10^14 g. a. -1 to hydrothermal influx, although this still would not balance input and output. The greater part of the output is primary production by diatoms which appear at the beginning of the marine food chain. The discrepancy between the figures for input and output, however, is not a real difference because of internal oceanic recycling which involves dissolution of siliceous tests and upwelling of the dissolved silica (Figure 1). According to estimates by different authors (e.g., Calvert, 1966; Berger, 1970; Hurd, 1973) between 90 and 95% of the silica extracted from surface seawater by shell-producing organisms (in the form of solid amorphous silica) redissolves before burial. Dissolution starts after death of the organisms when the tests settle through the water column, and continues while they rest on the sea floor, and particularly after burial in the sediment. A fraction of the silica dissolves during burial, however, returns to the ocean by means of diffusion. This is evident from the strong silica concentration gradient in the upper 20 cm of the sediment. DeMaster (1981) includes the figure for pore-water reflux quoted above (5.7 x 10^14 g. a. -1) with internal oceanic recycling. This is the diffusional flux across the sediment/water interface. Most of the dissolution during burial, however, provides the silicon required for diagenetic reprecipitation in the sediment, the mechanisms of which are the central topic of this review.

Depositional Environments of Siliceous Sediments

Biogenic siliceous oozes on the modern ocean floors. Biogenic siliceous sediments occur both in deep and shallow-water depositional environments. Siliceous ooze is a typical pelagic sediment which accumulates on the deep-sea floor in regions where silica production in surface waters is high. This is the case in zones of oceanic upwelling, where nutrients dissolved somewhere at depth in the oceans are returned to the surface. Upwelling is related to global or local oceanic circulation patterns and occurs (i) along equatorial divergences, (ii) in the vicinity of subpolar convergences, (iii) along

the west coasts of continents, and (iv) in certain marginal seas where surface currents diverge or are driven offshore. In the Pacific Ocean, regions of upwelling correspond to 5 major belts of silica accumulation on the sea floor (Figure 2): an equatorial belt (north of the equator), the sub-Antarctic belt (north of the Antarctic convergence), the sub-Arctic belt (south of the Arctic convergence), the Gulf of California (continental west coast/marginal sea upwelling) and the Okhotsk and Bering Seas (marginal sea setting).

In low-latitude areas of high primary surface production, pelagic carbonates are deposited together with siliceous tests leading to mixed calcareous-siliceous oozes. Pure siliceous ooze or high-silica ooze will only accumulate in areas where dilution by other components is inhibited or decreased. This is the case below the calcite compensation level (CCL), e.g., in the equatorial belt of siliceous ooze to the north of the mixed calcareous-siliceous belt. (An equivalent belt south of the equator does not exist, because the young, shallow oceanic crust has not yet subsided below the CCL). Temperature affects silica solubility more than pressure. Silica is therefore considerably more soluble in surface waters than at great depth (Figure 3). In the deep sea, below 1 or 2 km water depth, temperature decreases only a few degrees Celsius. The temperature effect thus becomes very small and can be offset by the effect of increasing pressure which raises silica solubility. Pressure-corrected curves for silica solubility (Figure 3) therefore have a minimum at about 1.5 km depth, below which solubility increases slightly. However, an equivalent of the CCL for silica has not been observed in the oceans. From the foregoing it follows that silica dissolution is predominant in the upper 1,000 m of the water column, in contrast with the dissolution of pelagic carbonates. In the Gulf of California, siliceous muds accumulate despite the supply of terrigenous detritus to the gulf, because the terrigenous input is seasonally and locally suppressed by the prevailing wind and current patterns (Calvert, 1966a, b).

Pelagic stratigraphy. The zonal distribution of pelagic sediments as described for the Pacific, when combined with plate motions and the thermal subsidence of ageing oceanic crust, gives rise to a distinct stratigraphic succession of pelagic sediments, the so-called "plate stratigraphy" (Berger and Winterer, 1974) or the "pelagic" stratigraphy described by Hesse et al. (1974). It is typically developed on Cretaceous-Cenozoic crust of the major oceans and consists (from bottom to top) of mid-ocean ridge basalt overlain by pelagic carbonates, then cherts and siliceous oozes and brown abyssal clay (Figure 4). Calcareous ooze is generally the first pelagic sediment deposited on newly created oceanic crust, at least since Late Jurassic time, because young oceanic crust at the crest of mid-ocean ridges is elevated and normally projects above the CCL. With age-related subsidence, the crust will eventually pass below the CCL, leading to the deposition of siliceous ooze and/or brown abyssal clay. In the Pacific, the sequence may be duplicated, if the northwestward motion of the Pacific plate carries it under the equator, where the CCL is depressed sufficiently to intercept the sea floor for a second time (Figure 4). Upon approaching the West and North Pacific island arc/deep-sea trench systems, the oceanic crustal section which carries this pelagic stratigraphy, collects an ultimate cap of hemipelagic and volcanogenic sediments before its history is terminated in a subduction zone. During subduction, some of the pelagic sediments may be scraped from the

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**Figure 1** Global dissolved silicon budget of the oceans. (From Riech and v. Rad, 1979, modified after Heath, 1974). All numbers are in 10^14 g SiO₂ per annum.
Ophiolites, the "Steinmann trinity", and chart in mountain belts. In the plate-stratigraphic succession, siliceous sediments typically occupy an intermediate position between pelagic carbonate and brown abyssal clay. In mountain belts, the association of radiolarian charts in an ordered sequence with other pelagic sediments and with mafic and ultramafic rocks of the ophiolite suite has been recognized for nearly a century. In the Alps, it was called the "Steinmann trinity"; after its discoverer Gustav Steinmann (1906, 1927). It was taken as a suite of rocks diagnostic for "eugeosynclines", whose equivalence with oceanic crust was only recognized with the advent of the plate tectonics concept (e.g., Chipping, 1971). Ophiolites and the Steinmann trinity

Figure 2 Distribution of pelagic sediment types in the bottom sediments of the Pacific Ocean (from Hesse et al., 1974). Note that siliceous sediments south of the Aleutian and Kurile Islands are mixed with volcanogenic material.
are thus today identified with the remnants of oceanic crust and mantle in mountain belts (Figure 5). The "petrogenetic problem", which Wenk (1949) saw in the association of ultrasiliceous sediments (radiolarians) with ultrabasic igneous rocks, is no longer a problem. The silica of the silicous pelagic sediments is not derived directly from silicic undersaturated basic and ultrabasic rocks by submarine exhalative volcanism, as previously thought (as recently as 1971; see Gibson and Towe, 1971), but through upwelling from the oceanic silica reservoir and biogenic precipitation in surface waters, as discussed above. The "geoelectonic significance" of chert, where it forms part of an ophiolite sequence, is not affected by this reinterpretation. However, since the distribution of silicous sediments and cherts is controlled by (paleo-)oceanographic conditions, they are therefore not restricted to oceanic crust. It is no surprise to find ancient cherts underlain by continental crust as, for example, in the Apennines (McBride and Folk, 1979), where they occur next to ophiolitic cherts (Folk and McBride, 1978). It should also be mentioned that in the Alpine - Mediterranean ophiolite terrains, the sequence "pelagic carbonate - chert - red abyssal shale" is generally reversed compared with the standard sequence found in the Pacific, with radiolarian cherts preceding the pelagic carbonates. This is probably due to a much shallower CCL in pre-Late Jurassic oceans, a consequence of the absence of widespread calcareous planktonic organisms in these times (Bosallini and Winterer, 1975). Although most ophiolite-associated cherts are not volcanogenic, but biogenic, in origin, as we know now, the geochemistry of some radiolarian cherts nevertheless reveals a metasomatic overprint which is due to igneous activity at a former mid-ocean ridge. Examples are the Jurassic Mediterranean bedded cherts just mentioned which directly overlie basalt of the oceanic crust. Their upward decreasing iron content and increasing manganese concentration with increasing stratigraphic distance from the basalt reflects the effects of hydrothermal fluid circulation in the sediments (Barrett, 1981).

**Formation of bedded chert.** Many ancient, bedded chert formations for which geological and sedimentological evidence suggests a deep-water origin, appear to owe their emplacement to redeposition of silicous pelagic sediment by turbidity currents (e.g., Imoto and Saito, 1973; Nisbet and Price, 1974; Robertson, 1977; McBride and Folk, 1979) or other ocean currents. Sedimentary structures and fabrics, such as flute casts and other sole marks (e.g., Imoto et al., 1974), common graded bedding, parallel lamination, faint cross lamination, starved ripples, grain fabrics such as parallel alignment of sponge spicules and radiolarians, shale clasts, and sharp contacts with interbedded shales, support a redeposition mechanism. Alternative interpretations include (i) diagenetic segregation of silica into chert-rich beds and silica-poor claystones from initially homogeneous siliceous muds, or (ii) variations in surface-water productivity, or (iii) variations in the input rate of the terrigenous component. These processes, however, do not adequately explain the sharp lithologic boundaries between the chert and interbedded shale layers which should have been obliterated by the activity of burrowing organisms, if these sediments were deposited through slow pelagic accumulation. Nearly all "bedded cherts" contain shale interbeds, often of red or green color, that comprise 5-40% of the sequence.

In some cases, the chert beds display none of the evidence for redeposition listed above, but still have sharp top and bottom contacts. This is the reason why some investigators prefer variations in biogenic productivity and/or diagenetic segregation as origin for these deposits (e.g., Dietsche 1980) for the well-bedded Upper Jurassic radiolarians of the Eastern Alps), although the lack of primary sedimentary structures may be due to other reasons (i.e., diagenetic obliteration due to recrystallization and/or uniform small grain size). There seems to be general agreement, however, that the bedded in radiolarian cherts cannot be ascribed to a single process (e.g., Baltuck 1983; Hein and Karl, 1983; Jenkyns and Winterer, 1982).

Recognition of symmetrical geochemical changes (from the centre of beds upward to the top and downward to the base) and symmetrical grading of radiolarian abundance (Mizutani and Shibata, 1983; Sano, 1983; Steinberg et al., 1983) lend new support to the diagenetic origin of some bedded cherts.

![Figure 3 Solubility of biogenic silica in the Central equatorial Pacific estimated as a function of temperature and pressure (dotted line, from Hurd, 1983, after Griffin, 1980)](image)

**Figure 4** Pelagic stratigraphy model for the Pacific Ocean showing duplication of the biogenic sediment sequence due to the westward passage of the Pacific plate under the equatorial high-productivity belt (from Hesse et al., 1974). Section oriented approximately NW-SE. The diagenetic compensation level (CCL) acts as a "snow line". The East Pacific Rise and other regions rising above it are covered with light-colored biogenic ooze. Areas below the CCL and outside the belts of siliceous ooze are characterized by red-brown abyssal clay (Figure 2).
sandstones and siltstones, on the other hand, are remarkably poor in concretionary silica (Dapples, 1967), in contrast with continental deposits. The Tertiary shallow marine examples of the Atlantic and Gulf coastal plains of the US (mentioned above) contain silica-cemented sandstones that appear to have received the silica from the associated opaline clays (Wermund and Moiola, 1966; Wise and Weaver, 1973; Weaver and Beck, 1977; Carver, 1980). In Paleozoic sequences, siliceous sponge spicules are often abundant, particularly in deep basinal deposits, but have also been considered as shoreline indicators (Cavaroc and Ferm, 1968; Lane, 1981). In a number of cases, spicules provide an adequate source of silica. Their diagenetic end-products are nodular cherts or chalk nodules which often occupy only a small fraction of the total volume of the carbonate or evaporite host rock. These nodular cherts are distinctly different from the ancient bedded cherts in the rock record and the biogenic siliceous oozes, muds and cherts on and under the modern ocean floors. Diagenesis of the later will be discussed in this paper. The diagenesis of the former may proceed under conditions considerably different from those of the deep-water cherts, and those of the siliceous deposits in alkaline lakes and in soils. The diagenesis of these and other types of chert including silicified wood will be discussed in another paper (Hesse, in press), ending with a note on the cherty varieties of Precambrian banded iron formations.

Stages of Burial Diagenesis in Biogenic Siliceous Sediments

During burial, biogenic siliceous sediments undergo a characteristic sequence of mineralo-logic transformations which may be subdivided into three stages, each defined by the predominance of one of the common low-temperature silica phases (Calvert, 1971): (1) the opal-A stage of siliceous oozes or muds; (2) the opal-CT stage of porcelanite, and (3) the microquartz or quartz stage of chert sensu stricto (quartz chert).

Silica recrystallization in petrified wood follows the same sequence, whereas some cherts in carbonates and some non-biogenic cherts may also follow different diagenetic pathways as discussed in Hesse (in press).

Porcelanite and chert do not simply reflect the mineralogical difference between opal-CT and quartz as a result of progressive diagenesis. As suggested by Isaacs et al. (1983), primary compositional differences, such as the ratio of clay minerals to silica minerals, are often more important in determining textural and physical properties on which the distinction of rock types is based. In the Monterey Formation, the most silicarich sediments (pure diatomites) form rocks that appear macroscopically as cherts although they may still consist entirely of opal-CT. Diatomaceous shale on the other hand, has the appearance of porcelanite, even after having reached the quartz stage. Because these rock names were originally designed for the field description of siliceous rocks (Bramlette, 1946), they will be used in this review in the sense of Isaacs et al. (1983). That is, the distinction between porcelanite and chert will be based primarily on detrital content. Often, however, porcelanite consists predominantly of opal-CT, and chert predominantly of (micro)quartz, this being the reason why Calvert (1971) first suggested a nomenclature of siliceous rocks strictly based on mineralogy, a usage largely followed by the Deep Sea Drilling Project (e.g., Riech and v. Rad, 1979). See however, Pisciotto (1981) for the alternative descriptive approach.

![Figure 5](image-url)  
**Figure 5** (a) Schematic representation of the "Steemann trinity" (= ophiolite stratigraphy) of ultrabasic rocks, altered basalts and pelagic sediments (radiolarian cherts and limestones); (b) origin of the ophiolite stratigraphy through ocean crust generation at mid-oceanic ridges; and (c) its destruction in subduction zones (from Chipping, 1971).
The characteristics of the different silica phases and the diagenetic transitions from one phase to the other will be described in the following paragraphs. Important questions discussed are: What are the physiochemical mechanisms involved in the transformations? Are the transformations of opal-A to opal-CT and of opal-CT to (micro-)quartz exclusively due to dissolution-reprecipitation reactions as favored by recent findings, or is there any evidence for the solid state reactions originally suggested by Ernst and Calvert (1969)? What are the rate-controlling factors? In particular, how does lithology influence transformation rates? Why is an intermediate state of metastable opal-CT required in the transition from opal-A to quartz?

Characteristics of the Silica Phases in Biogenic Siliceous Sediments

Opal-A: Physical characteristics, solubility, and polymerization. The tests of siliceous microfossil as well as silica precipitated inorganically under earth surface conditions, as in hot-spring areas or desert soils, consist of an X-ray amorphous substance named opal-A (Jones and Segnit, 1971), because it is mineralogically similar to precious opal. The X-ray pattern of this near-amorphous substance with a broad hump near 4Å (Figure 6a) resembles that of glass. This material is very porous and has a substantial water content (4-9% in precious opal). Opal-A dissolves easily in seawater, which is highly undersaturated with respect to opaline silica everywhere in the modern oceans. Seawater contains a fraction of a part per million to 10-15 ppm dissolved silicon (or up to 160-250 μM L⁻¹), i.e., considerably less than the equilibrium solubility of amorphous silica, which is between 60 and 130 ppm at 25°C and a pH < 9. Silica-secretting marine micro-organisms thus must extract dissolved silicon from seawater at concentrations well below the thermodynamic equilibrium solubility of opal-A, but will raise it to saturation under the catalyzing influence of enzymatic reactions. Concentrations of dissolved silicon in the ocean are lowest in surface seawater because of the extraction by organisms. They increase with water depth because of the dissolution of siliceous tests settling through the water column, leading to a mid-depth maximum. A subsequent decrease at greater depths is caused by the renewal of bottom waters from silica-poor surface-water sources in polar regions, especially in the North Atlantic. Bottom maxima in dissolved silicon, on the other hand, are associated with belts of siliceous sediments, particularly diatom ooze (Edmond et al., 1979b), with Antarctic Bottom Water, and areas of hydrothermal activity.

In inorganic systems, equilibrium solubility of silica depends on a number of factors besides temperature (Figure 7a), notably surface area and particle size of the solid silica phase (Figure 7b), to a lesser extent pressure as stated in a previous section (Figure 7c). It increases significantly for a pH greater than about 9 (Figure 7d, Williams et al., 1985; Williams and Cerara, 1985). Other ions in solution can have distinct effects on dissolution rates of solid silica. Dissolved Al and Fe which adsorb to the opal-A surface suppress opal-A solubility and dissolution rate (Iler, 1955, 1973; Lewin, 1961). Also, aging of the biogenic siliceous tests affects solubility through changes in surface area and crystal size in the shell wall (Hurd and Theyer, 1975). Equilibrium solubilities of the other common low-temperature silica polymorphs, opal-CT and quartz, are respectively one and two orders of magnitude lower than those of opal-A (Figure 7a), although for opal-CT no thermodynamic data exist.

Because of the significance of silica solubilities for the diagenetic transformations of the solid phases, a somewhat closer look at the chemistry and kinetics of the dissolution and precipitation processes, including polymerization, of silica is in order. In solutions of pH < 9, dissolved silicon is present predominantly in the form of undissociated silicic acid, H₄SiO₄⁻ (orthosilicic or monomeric silicic acid). The dissolution of solid silica can be written as

\[
\text{SiO}_2(s) + 2 \text{H}^+ = \text{H}_2\text{SiO}_4^{-}(aq)
\]

At equilibrium, the solubility product, K_{sp}, thus approximates the activity a(H₂SiO₄⁻). In solutions of pH > 9, orthosilicic acid dissociates (Figure 7d), in a first step into

\[
\text{H}_4\text{SiO}_4^{-}(aq) \rightarrow \text{H}_2\text{SiO}_4^{-} + \text{H}^+
\]

and for higher pH values (> 13) in a second dissociation step into

\[
\text{H}_2\text{SiO}_4^{-} \rightarrow \text{H}_2\text{SiO}_4^{2-} + \text{H}^+
\]

These dissociations raise the solubility significantly, because the total dissolved silicon concentration is the sum of the dissociated and undissociated species.

In supersaturated solutions, dissolved silicon polymerizes forming first oligomers (dimers, trimers and ring structures), then higher-molecular-weight polymers as siloxane bonds (Si-O-Si) develop through combination of silanol (Si-OH) groups:

\[
\begin{align*}
\text{OH} & \quad \text{OH} \\
\text{HO-Si-OH} & \quad \text{HO-Si-O-Si-OH} \quad \text{H}_2\text{O} \\
\text{OH} & \quad \text{OH}
\end{align*}
\]

Williams and Cerara (1985) emphasize the fact that low-molecular-weight oligomers also contribute to the total solubility in solutions at or near opal-A saturation, contrary to the assumption in most previous experimental work that all dissolved silicon is monomeric in such solutions. Oligomers may even persist in undersaturated solutions at temperatures below 130°C, although under such conditions they will gradually depolymerize (e.g., Krauskopf, 1956). At concentrations below opal-CT saturation, monomeric silicic acid is the only form present. On the other hand, if supersaturation persists, high-molecular-weight polymers (with molecular weights up to 10,000) form. Such polymers have colloidal dimensions (greater than 50Å) and may remain suspended as sols as long as pH remains relatively high and salinity low; otherwise they will form cross-links with neighbours and coagulate into gels (Figure 8a).

Silica polymers display a negative surface charge, which results in a point of zero charge (PZC, i.e., the pH at which the residual surface charge disappears) as low as 2 ± 0.5 (Parks, 1965). Silica colloids thus repel each other unless the surface charge is neutralized by other ions in solution such as metal hydroxides. Interestingly, the hydroxide most commonly used for silica precipitation in industrial applications, Mg(OH)₂ (Iler, 1979), is also thought to be instrumental in the nucleation of opal-CT, as discussed below.

The ultrastructure of precious opal shows a hexagonal or cubic closest packing of opal-A spheres that are remarkably uniform in size (Iler, 1979). Apparently, they are deposited from colloidal particle suspensions of uniform size that grow in supersaturated solutions in the absence of natural coagulants. The sphere diameter is in the appropriate size range (0.1-0.38 μm) to diffract visible light (in the same manner crystal lattices diffract X-rays) and to produce the indescent diffraction colours of gem opals. These spheres consist of several concentric shells of primary smaller spheres with diameters ranging from 100 to 500Å depending on the opal type (Figure 8b). Uniform size of the primary spheres apparently is achieved by the dissolution of smaller polymers and redeposition of the dissolved silica on the larger ones (Darragh et al., 1976).

Opal-CT: Ultrastructure. The "CT" in opal-CT stands for cristobalite/tridymite (Jones and Segnit, 1971) and denotes a modification of opal which has structural characteristics of both α-cristobalite and α-tridymite. Opal-CT thus is the low-temperature form (α-form) of cristobalite/tridymite formerly called flasslite (Mallard, 1890). In opal-CT, cristobalite alternates randomly with unidimensionally disordered tridymite layers. On X-ray powder diagrams, the main diffraction peak of opal-CT is a doublet at 4.1 and 4.3 Å (Figures 6b to 6i).

Wise et al. (1972) first observed the occurrence of opal-CT in the form of small spheres less than 5 μm in diameter in deep-sea drilling samples under the scanning electron microscope (SEM). These spheres, which consist of an interpenetrative growth of tiny
crystalblite/tridymite blades, were called opal-CT lepispheres (Figure 9a) by Weaver and Wise (1972). On high-resolution SEM photos, lepispheres in their incipient growth stage show sets of subparallel opal-CT blades which penetrate each other at angles between 70° and 71°. Individual blades are 2-5 μm long and 0.05-0.1 μm thick, and display ragged or rounded edges (Figure 9b). The penetration angle corresponds to the angle of 70°32' at which equivalent faces of a crystalblite octahedron intersect one another (Figures 9c,d). Flörke et al. (1976) give a plausible crystallographic interpretation for the peculiar twinning behavior of opal-CT in the lepispheres, which appears to be dictated by crystal growth kinetics. At more advanced growth stages the skeletons will coalesce to form lepispheres. The diameter of individual lepispheres usually does not exceed the length of the blades (about 5 μm) of which they are composed. Larger lepispheres are almost invariably aggregate forms (Figure 9e; see also Carver, 1980, figs. 1C, 1D, 3A, 3B). Relics of former opal-CT lepispheres in Paleozoic cherts which have been portrayed in the literature (e.g., Jones and Knauth, 1979; Meyers, 1977) are invariably too large for genuine lepispheres, unless they are aggregates.

Figure 7 Solubility diagrams for various silica phases as a function of (a) temperature (from Williams and Crerar, 1985; modified after Walther and Helgeson, 1977); (b) specific surface area (from Williams et al., 1985); (c) pressure for quartz at various temperatures between 25° and 600°C (from Williams and Crerar, 1985; modified after Walther and Helgeson, 1977); and (d) pH (from Williams and Crerar, 1985; after Volosov et al., 1972).
Low-temperature tridymite and cristobalite as separate, distinct mineralogic phases. The occurrence, under special geologic conditions, of low-temperature tridymite and cristobalite as separate mineralogic phases distinct from opal-CT was proposed by Tada and Iijima (1983). The disordered low-temperature tridymite (α-tridymite) is characterized by a relatively broad diffraction peak at 4.11 Å on X-ray powder diffractograms obtained from mixtures of α-tridymite, α-cristobalite and opal-CT. On the same diffractograms, the better ordered α-cristobalite (also called opal-C) produces a sharper diffraction peak at 4.04 Å. As will be shown later, both the 4.11 and 4.04 Å peaks have also been attributed to opal-CT at various stages of burial maturation (Murata and Larson, 1975). The justification given by Tada and Iijima (1983) for distinguishing a disordered tridymite from 4.11Å-opal-CT is that it did not recrystallize to a 4.04Å-phase upon heating (for 11 days to 1100°C) contrary to what was expected. In nature, this low-temperature tridymite is observed as a pore-filling cement precipitated from ground water or hydrothermal solutions in silicic volcanic rocks. The low-temperature α-cristobalite is also generally associated with volcanic and silicic volcaniclastic rocks. In deep-sea sediments it has been reported from volcanogenic sediments overlaying basalt in the West Pacific (Lau Basin, Griffin et al., 1972) and Central Pacific (Zemmel and Cook, 1973).

Nature of the Transformation Mechanisms

Opal-A to opal-CT transformation. Only a small percentage of the large amount of biogenic silica (opal-A) produced by planktonic organisms in the surface ocean reaches the sea floor, and only a fraction of this escapes dissolution during the first 50 meters of burial. Even this very small proportion of solution-resistant species will ultimately undergo dissolution at greater sub-bottom depths. The effects of progressive dissolution have been documented by systematic SEM studies of diatom oozes (e.g., Hein et al., 1978). Breakage of partially dissolved diatom valves accompanies and enhances dissolution, culminating in the complete destruction of the tests. Dissolved silicon profiles from DSDP holes in the pelagic realm reflect the dissolution of solid silica particles in the subsurface. These profiles show consistent increases in dissolved silicon with sub-bottom depth, not uncommonly exceeding 1 mM (or 60 ppm) at a few 100 m below the sea floor. Concentrations may fluctuate considerably with depth, but these fluctuations are generally superimposed on an overall trend of downward increasing silica concentrations (Figure 10). In many holes, however, the gradual downward increase is followed by an abrupt decrease below a certain depth (e.g., DSDP site 495, Harrison et al., 1982; site 570, Hesse et al., 1985) consistent with precipitation as opal-CT. In the examples mentioned, this is substantiated by the recovery of the first porcelain nodules from this depth. Below this depth, a second less pronounced maximum may be observed in holes with sufficient penetration, probably corresponding to the dissolution of opal-CT and the subsequent precipitation as quartz described in the next section.

Continued dissolution of opal-A in the sediment during burial is the result of slowly increasing temperature and pressure. Siliceous tests of diatoms and Radiolaria have large specific surface areas ranging from several tens to 450 m²·g⁻¹ (Kastner et al., 1977), compared with 0.1 m²·g⁻¹ of crushed quartz in the size range of 3-5µ (25-75µm) (Van Lier et al., 1960). The sieve structure of the porous test walls of these organisms (Figure 9f) is only partly responsible for these large specific surface areas. As Hurd et al. (1979) have shown, it is ultimately the size of the small opal-A domains (2.5-4.5Å diameter) in the tests which causes the high specific surface areas. Surface area can significantly affect solubility. When the surface area to volume ratio of a substance becomes large, small changes in pressure and temperature may then markedly influence its solubility. Suppression of solution inhibiting factors such as the removal of protective coatings of organic matter may further increase solubility. This explains the continuation of opal-A dissolution during burial, even though up to 99% of the opal originally produced may have already been dissolved during settling and initial burial.

Dissolution during sediment burial, in contrast with the earlier dissolution, occurs in a more or less closed system in which concentration levels may reach supersaturation before opal-A dissolution has gone to completion. In this case, dissolution will be interrupted by reincorporation of a less soluble non-biogenic opal-A which forms overgrowths on partially dissolved siliceous tests. These inorganically precipitated overgrowths are called opal-A'(Hein et al., 1978).

Figure 9 (opposite page) (a) Small opal-CT lepispheres (2-3 µm in diameter) growing on euhedral calcite crystals in a cavity in partially silicified Maestrichtian chalk, DSDP 14-144-3-2, 103-104 cm. (b) Opal-CT blades of a "juvenile" lepisphere displaying twinning angle of 70° corresponding to the intersection angle of the faces of a cristobalite octahedron as shown in (c) and (d) (from v. Rad et al., 1977). (e) Composite lepisphere 50 microns in diameter which resulted from coalescence of numerous individual lepispheres, DSDP 12-171-22C (from Flörke et al., 1976). (f) Sieve structure of diatom frustule. (SEM photos in Figures 9a, b, e, f courtesy U. v. Rad).

Figure 8 (a) Polymerization behaviour of dissolved silicon as a function of pH and ionic strength of the solution (from Williams and Crerar, 1985). (b) Formation of precious opal by deposition of large spheres of opal-A of uniform size in a hexagonally or cubically closest packed layer. Large spheres consist of concentric shells of smaller primary spheres (from Williams and Crerar, 1985; after Darragh et al., 1976).
Crystallite size calculated from X-ray diffraction data for opal-A is larger (20-27Å) than for biogenic opal-A (12-16Å). In individual DSDP holes, opal-A overgrowths have been found to occur over a narrow stratigraphic range of a few metres, indicating that they redissolve shortly after formation together with the remaining opal-A (Hein et al., 1978). Therefore they are rarely observed. Their presence, however, may explain some of the fluctuations seen in the concentration – depth profiles of dissolved silica (Figure 10).

Recognition of inorganically precipitated opal-A by Heath (1969) and Hein et al. (1978) demonstrates that the opal-A to opal-CT transformation is not a single-step process but involves a series of dissolution-reprecipitation reactions as postulated theoretically by the model of Williams et al. (1985). This model is based on surface-area effects (Figure 7b). The hypothetical SiO2–H2O system used is a closed system at constant temperature, pressure, pH and consists of the pure reactants. In this system, opal-A of a diatom species with a specific surface area, say of 250 m² g⁻¹ (i.e., *Thalassiosira decipiens*, Figure 7b), will have a solubility of about 100 ppm. If dissolution of the frustules of this species is fast relative to nucleation and growth of other new silica phases, then the solution will soon become supersaturated with respect to opal-A of a lower surface area (i.e., opal-A'). The solution is also supersaturated with respect to opal-CT of any given surface area to the left of point 3 in the diagram, as well as with respect to any quartz. The nucleation and growth rates of opal-CT will be slower than those of any opal-A, and those of quartz will be slower than those of opal-CT, because opal-CT and quartz are more highly ordered forms of solid silica compared to opal-A. Under these assumptions, an opal-A' with a surface area lower than that of opal-A will precipitate and the solution will evolve along the pathway from points 1 to 2 shown in Figure 7b – a process called "Ostwald ripening" (Iler, 1979; Williams et al., 1985). Near point 2, the effect of surface area on opal-A solubility becomes negligible. At this point, the solid silica phase with the next lower solubility, i.e., opal-CT may appear. Opal-CT is preferentially precipitated over quartz, because of its higher nucleation and growth rates. Depending on the dissolution rate of opal-A relative to the nucleation and growth rate of opal-CT, further evolution of the solution will follow pathways from points 2 to 4 along either curve "a" (fast nucleation rate) or curve "b" (slow nucleation rate). In the first case, rapid nucleation (relative to growth) leads to relatively large specific surface area, because the nuclei are small and numerous. Only when the silica removal rate exceeds the dissolution rate of the remaining opal-A, will silica concentration drop off and the surface area of the newly formed opal-CT decrease (curve 2-4a). If, on the other hand, the growth rate of opal-CT exceeds the nucleation rate early in the process, then the fluid should evolve along pathway 2-4b. Analogous considerations should apply to the opal-CT to quartz transformation discussed in a subsequent section, although the structural reorganization within the opal-CT, which precedes this transformation, is interpreted as a solid-state reaction by some authors.

The discovery of the opal-CT epiplaxes with the euhedral crystal shapes of the cristobalite/tridymite blades by Wise et al. (1972), Weaver and Wise (1972) and Berger and v. Rad (1972) provided the first proof that

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**Figure 10** Concentration of dissolved silicon in pore-water profiles of DSDP holes (L, DSDP leg; S, DSDP site).
the recrystallization of siliceous oozes to porcelainite occurs through a dissolution–reprecipitation mechanism. Lepidospheres develop only where crystallization takes place in open pore spaces such as the cavities of microfossils. More commonly, a densely felted mass of opal-CT forms which may impregnate the sediment and/or replace other mineral phases. The latter process may involve pseudomorphic replacement of opal-A by opal-CT in radiolarian tests which perfectly preserves the shape of the shell (Figure 11). The process is still a dissolution–reprecipitation mechanism, which probably proceeds on a grain-by-grain basis with local precipitation immediately following dissolution. Presence of a matrix of organic matter, which is not or only partially affected by dissolution, may serve as a template which helps preserve the original shape of the shell. The process may be similar to the silicification mechanism in ooids (Hesse, 1987) or to the petrification of wood. Where Radiolaria occur embedded in lutitic pelagic limestone, an opal-CT replaced test may simply represent a cast of the former opal-A shell. Conservation of the original shape during the replacement process cannot be used as evidence for a solid-state reaction in the opal-A to opal-CT transformation.

Opal-CT to quartz transformation. The opal-CT to quartz transformation was assumed to be a solid-state mechanism by Ernst and Calvert (1969) and Heath and Moberly (1971), but morphological evidence from scanning electron microscopy (Stein and Kirkpatrick, 1976) makes it more likely that this step is also a dissolution–reprecipitation reaction like the opal-A to opal-CT transformation. In their hydrothermal experiments, Ernst and Calvert (1969) heated opal-CT in distilled water to temperatures of 300°, 400°, and 500°C under 2 kbar pressure. Quartz formed in each experiment. At 500°C the conversion was complete after 30 hours, at 400°C after 500 hours (12.5 days), and at 300°C after 5,000 hours (7 months). For a given temperature the conversion rate seemed to be constant. The concentration of the reactant opal-CT had decreased during the experiment in a more or less linear fashion with time. On the basis of general reaction-rate laws, Ernst and Calvert concluded that the reaction was of zero order with respect to the number of participating reactants; it had to be a solid-state reaction.

Stein and Kirkpatrick (1976) re-examined the reaction products of these experiments under the SEM and found fibrous and euhedral prismatic quartz crystals among the reacted material clearly requiring a dissolved phase for crystallization. After re-evaluation of the experimental data (fraction of opal-CT transformed versus time) in the light of nucleation and growth theory (Christian, 1965; Nielsen, 1964), these authors concluded that a sigmoidal curve with slower reaction rates at the beginning and at the end of the conversion process would approximate the rate data better than the straight-line plots of Ernst and Calvert (1969).

Crystallographic structural changes of opal-CT and quartz in the porcelainite and quartz-chert stages. Sharpening and shift of the opal-CT (101) reflection. The important finding of a dissolution–precipitation step in the opal-CT to quartz conversion by Stein and Kirkpatrick in 1976 did not close the discussion about the nature of the transformation mechanism involved in this phase change. A year before, in 1975, Murata and Larson’s results of a high-precision X-ray diffraction study gave rise to new speculations about the possibility of low-temperature diagenetic solid-state reactions. This study revealed characteristic changes in the main diffraction peak of opal-CT (41A) with progressive burial diagenesis. In contrast to Tada and Iijima (1983), this peak is usually attributed to the (101) diffraction of α-cristobalite. In siliceous rocks of the Monterey Formation in the Temblor Range of California, a distinct shift of this peak from 4.11 to 4.04Å is observed with
increasing burial (Figure 6). (Slow scanning at a rate of 0.5°/2θ min⁻¹ permitted measurement of the peak position with a precision of ±0.001°). This decrease in the d-spacing is accompanied by a progressive sharpening of the same peak, a gradual disappearance of the α-tridymite (0001) peak (at 4.32-4.28 Å), which is replaced by the quartz (100) peak (at the opal-CT to quartz transition). Minor secondary cristobalite peaks at 31 and 2.8 Å also appear with increasing burial approximately in the middle of the opal-CT stage. While the shift in the d-spacing of the (101) peak is relatively rapid at the top and bottom of the opal-CT zone in the Temblor Range, it nevertheless proceeds continuously over the entire burial range of the opal-CT zone, albeit very slowly in the middle stages (Figure 12). In contrast, in the Santa Maria Valley, Pisciotta (1981) found the shift to become more rapid with increasing burial depth starting in the middle of the opal-CT zone. The bulk density of the sediments, on the other hand, does not show any systematic increase over the range of burial depths (730-2,030 m below reference level). It stays more or less constant at an average value of 1.66 g cm⁻³, but increases abruptly to 1.66 g cm⁻³ over an 80 m thick sediment interval in the opal-CT to quartz transition zone (Figure 12). Oxygen isotope data from the same burial depths display a similar kind of behavior: no systematic change within the opal-CT zone (average value of δ¹⁸O = +29.4‰ relative to SMOW), but a significant decrease by about 9‰ within a short distance below the porcelainite/chert boundary. Murata et al. (1977) concluded from these observations that the structural ordering of cristobalite, manifested in changes of the d(101) spacing, occurred through reactions in the solid state within the opal-CT stage, but involved complete dissolution – recrystallization at the stage boundary with the quartz zone.

As Isaacs et al. (1983) pointed out, however, "neither the structure of the mineral opal-CT nor the mineralogic significance of the d-spacing of opal-CT are sufficiently well understood to interpret the changes in the structure of opal-CT as an ordering process in the strict sense, although this is what most authors assume. For instance, the sharpening of the (101)-cristobalite peak with increasing burial may reflect growing crystallite size of the opal-CT which in turn may

Figure 13 (a) X-ray diffractionogram showing the quartz quintuplet at about 68°20', and the parameters a and b used to define the quartz crystallinity index CI of Murata and Norman (1976, fig. 2; for explanation see text).
(b) Tracings of diffractiongrams showing increasing quartz crystallinity from bottom to top. Asterisk marks (212) peak at 67.74° which is used for the determination of the index. CuKα radiation (from Murata and Norman, 1976, fig. 1).

Figure 14 (d) (101) spacing isolaplates for opal-CT porcelainite in the Temblor Range southwest of Taft (California). These provide an example of "isograding" mapping of low- to intermediate-grade diagenetic rocks. Note apparent congruence between structural features and diagenetic trends (Redrafted after Murata and Randall, 1975, fig. 4).
result from gradual dissolution of smaller crystals and redeposition of the dissolved silicon on larger ones (an Ostwald ripening process). This may be considered a reordering process in a general but not in a strict sense. High-precision X-ray diffractograms are sensitive enough to depict such minute structural changes, whatever their physical cause, and therefore record these modifications continuously with burial, whereas standard oxygen mass spectrometry or pycnometer methods may not pick up small but continuous changes with progressive burial. Williams and Crerar (1965) and Williams et al. (1965) suggest that some of the seemingly abrupt mineralogical or geochemical changes at the boundaries of the opal-CT stage may in fact be more gradual than they appear to be. Before reaching a certain threshold level characteristic of a diagenetic stage boundary, some of these changes occur so slowly that the less sensitive methods do not see them. Effects will only be seen after a significant portion of the rock has undergone transformation. Alternatively, the oxygen-isotope composition may in fact change very little within the opal-CT stage because the isotopic ratio of growing larger crystals may be inherited from dissolving smaller ones. Density may also change very little because the morphology of the opal-CT blades remains essentially unchanged. These considerations would make the problem amenable to interpretations entirely based on dissolution—precipitation reactions and would eliminate the need for solid-state reactions, which at the low temperatures of diagenericization are prohibitively slow.

The relationship between specific surface area and solubility schematically shown in Figure 7b also explains why opal-CT is required as an intermediate metastable phase in the sequence of diageneric silica transformations. Quartz cannot form directly from the dissolution of opal-CT, at least not from an equilibrium solution, because silicon solubility in equilibrium with opal-CT of any specific surface area is too high. At such high silicon concentrations the faces of any embryonic quartz crystal would be crowded by silanol groups which would not have time to be properly fitted into the crystal lattice. Quartz growth will be blocked and a less "well-ordered" phase, opal-CT, will form instead. Only when the "equilibrium solubility" of the phase (for which no precise thermodynamic data exist) has been lowered sufficiently through Ostwald ripening, will quartz crystallization become possible. Opal-CT is a classical example for Ostwald's step rule which states that the transformation of an unstable to a stable mineral phase (at earth surface conditions) often requires one or more intermediate metastable phases. This is an immediate consequence of the crystallization kinetics (Morse, 1988) and answers one of the questions raised at the beginning. However, under special circumstances (see section on reaction rates), it is possible that quartz may precipitate from solutions whose silicon was derived directly from opal-CT dissolution but had not reached opal-CT equilibrium concentration, or even (the lower) opal-CT concentration. 

Crystalinity index of quartz. Sharpening of the [101] crystalite peak with increasing maturation may be likened to the improvement in quartz crystalinity with progressive diageneric described by Murata and Norman (1976). The quartz crystalinity index of these authors is based on high-precision XRD measurements at slow scanning speeds (0.25°2θ·min⁻¹) of the high-angle region between 67° and 69°2θ (Cu K, radiation). In this 2° range, moderately well-crystallized quartz produces a quintuplet of XRD peaks of which the (212) peak at 67° measures the effects of the recrystallization of cryptocrystalline to microcrystalline authigenic quartz during diageneric and low-grade metamorphism. The quartz crystalinity index most probably is a measure of crystal size and is defined as

\[
\text{CI} = 10 \frac{\text{F}}{\text{B}}
\]

where A is peak height on the high-angle side, B is the total peak height (as shown in Figure 13a) and F is a scaling factor which varies with instrumental settings and adjusts the crystallinity index to a scale of 0 to 10. The crystallinity of authigenic quartz (e.g., in the Monterey Formation) is characterized by (2.0 to 3.2), even in quartz cherts sensu stricto. Cretaceous quartz cherts in the West Pacific at about 400 m sub-surface depth have crystallinities generally less than 1 (using a scaling factor of 1.36, Pisciotto, 1980). Franciscan cherts show considerably better crystallinities (Figure 13b), but reach high values (above 8.0) only in metamorphosed rocks (Murata and Norman, 1976). Opal-CT d(101) spacing and quartz crystalinity as mapping tools for diageneric grade. Variations in the d(101) parameter of opal-CT have been used to map regional maturation levels in relatively low-grade diageneric rocks (Murata and Randall, 1975).
comparable to isograde mapping in higher-grade metamorphic terrains. The contours of iso-d-spacings in the Monterey Formation in the Temblor Range of California generally conform to the geological structures (Figure 14). The lowest d-spacings (4.05 to 4.04 Å) straddle the axes of anticlines. However, as Mizutani (1977) noticed, the dip of the reconstructed iso-d-spacings in the cross-sections of Figure 15 is less than the structural dip. This probably indicates that structural deformation in the Temblor Range started when opal-CT maturation was still in progress (Figure 15c). Alternatively, as discussed below, discordant maturation levels with respect to tectonic structures could reflect lithological effects of the host rocks (Isaacs, 1982).

Keller et al. (1985) used quartz crystallinity based on measured apparent crystal size measured under the SEM to map diagenetic and metamorphic grade in the Ouachita Mountains of Arkansas and Oklahoma and in a contact metamorphic aureole on the Isle of Skye, Scotland. Crystal size ranges from less than 1 μm for cryptocrystalline, anhedral quartz in non-metamorphic terrains to more than 100 μm for polygonal, triple-point euhedral quartz in tremolite, diopside and forsterite-grade rocks. The contour maps outline the regional structural trends and depict areas of subsurface igneous activity.

Rate Controlling Factors of the Diagenetic Silica Transformations
Rates of the opal-A to opal-CT transformation in deep-sea environments: Temperature and time. Temperature is the dominant rate-determining factor for the diagenetic silica transformations as illustrated by the experiments of Ernst and Calvert (1969) for the opal-CT to quartz transformation (described above), or the experiments by Kastner et al. (1977) for the opal-A to opal-CT transformation (described below). The theoretical basis for this is the Arrhenius equation for the rate constant k of a chemical reaction

\[ k = A e^{(E_a/RT)} \]

where \( E_a \) is the activation energy, \( T \) is temperature, \( R \) is gas constant, and \( A \) is a frequency factor. During the diagenesis of deep-sea pelagic sediments or silica-bearing shallow-water sediments, pressure effects are generally negligible compared with temperature or other effects. Pressure is generally hydrostatic in these environments and varies only to a small extent over the burial intervals of a few hundred meters in which the first silica transformations take place.

What are the actual in-situ temperatures at which the opal-A conversion to opal-CT takes place? Downhole temperature measurements at DSDP sites 184 and 185 (leg 19) in the Bering Sea give temperatures of 35°-51°C at 500-600 m sub-bottom depths in the diatomaceous sediments where the bulk change occurs from opal-A to opal-CT (Hein et al., 1978). In the Temblor Range of California, Murata et al. (1977) obtained oxygen isotopic temperatures for opal-CT from porcelanites of the Monterey Formation in the range of 41° to 56°C (mean 48°C) using the fractionation factors of Clayton et al. (1972) and Labeyrie (1974) and assuming a δ18O of 0‰ (SMOW) for the pore water from which the opal-CT precipitated. At a nearby locality, an interstitial water δ18O of +3.7‰ had to be assumed to obtain the same range of temperatures. These estimates correspond well with a temperature of 50°C calculated on the basis of geothermal gradients by the same authors for an assumed subsurface depth of 700 m in the marine section of the Temblor Range where the transformation to opal-CT allegedly took place. A considerably lower range of temperatures from 9°-27°C was estimated by Pisciotto (1981). This range applies strictly to the top of the opal-CT zone (defined as the depth where opal-CT exceeds 5 weight percent (wt. %)) in the Monterey Formation in drill holes of the Santa Maria Valley in California and is an absolute minimum range. It is based on detailed considerations of the thermal and subsidence history of the region. Opal-CT formation started about 2 million years (m.y.) after deposition and reached a peak after 4 to 10 m.y. (the shorter time corresponding to higher heat flow). In the pelagic realm away from the mid-ocean ridges, the transformation occurs at shallow sub-surface depths of a few hundred metres where temperature rarely exceeds 20°-30°C. At these low temperatures, the transformations are very slow and the opal-A to opal-CT conversion takes some 10 m.y. to go to completion, and perhaps up to 30 or 50 m.y. for temperatures well below 30°C (Figure 16). The transformation can be traced regionally on seismic reflection profiles because it is associated with the transition from siliceous ooze to porcelanite, which is a major regional lithification event. In the Northwest Pacific, for example, this event is marked by the occurrence of an "opaque" seismic reflector in the subsurface, which first appears at the mid-crest of Hawaii (at approximately 155°W, Figure 17) in mid-Tertiary siliceous sediments (15-20 m.y. old) which rest on Upper Cretaceous oceanic crust (about 90 m.y. old). This reflector increases in thickness westward from less than 20 m to 200-300 m in the West Pacific where it represents the combined thickness of both siliceous rocks and pelagic limestones as numerous drill holes have shown (e.g., DSDP legs 7, 16, 17, 20, 32, 55, 61, 62, 89). The "opaque layer" can be deciphered on seismic profiles as a distinct interval because it is underlain by older, less brittle and relatively poorly consolidated clayey sediments which produce a "lower seis-
mically transparent layer” in the region. The latter sediments correspond to the lower horizon of brown abyssal claystone that formed when that part of the Pacific plate which is now north of the equator was still located south of the equator and had first subsided below the CCL (see section on pelagic stratigraphy model, p. 172). The “upper opaque layer” on North Pacific seismic profiles west of Hawaii thus records a diagenetic history that needed an activation time of at least 15 m.y. before it produced bulk physical changes. This illustrates the importance of the factor time which may play the major role in diagenesis, if temperatures are low.

Host-rock lithology as rate-controlling factor. In addition to temperature and time, there are other factors which may promote or retard the diagenesis of siliceous sediments. Since Bramlette’s classical work on the Monterey Formation, numerous exceptions from the general maturation sequence “siliceous ooze → porcelanite → quartz chert” have been recognized. For example, in the Monterey Formation, porcelanite and/or quartz chert beds occur within the zone of non-recrystallized diatomaceous mudstone and diatomite, and chert layers occur within the porcelanite zone. Apparently, the diageneric reactions leading to the silica transformations proceed faster in certain horizons than in adjacent ones. In other cases, the transition to quartz seemed to have occurred directly from opal-A without an intermediate opal-CT stage (e.g., Lancelot, 1973).

As observed during early legs of the DSDP (e.g., v. Rad and Rösch, 1972; Lancelot, 1973; Keene, 1975), lithology of the host rocks has a distinct influence on silica transformations. For example, in clayey sediment sections, silica maturation had reached the opal-CT stage of porcelanite, whereas in accompanying calcareous sediments of the same age and thermal maturation level, quartz chert had formed. This led to speculations that, if the host rock is calcareous, opal-A might convert directly to quartz (quartz-precipitation hypothesis of Lancelot, 1973) without the intermediate opal-CT stage.

Experiments by Kastner et al. (1977) provided insight into the role of host-rock lithology and related solution chemistry for the opal-A to opal-CT transformation. These authors performed hydrothermal experiments reacting diatom frustules and radiolarian tests at 150°C with seawater or distilled water in the presence of either calcareous ooze or montmorillonitic clay. The main results of these experiments can be summarized as follows:

1. In all experiments at 150°C, but not in all room-temperature experiments, corrosion of the siliceous tests took place after periods of one day to one month.
2. After 2 to 3 months, precipitation of opal-CT lepispheres was observed, but only in the experiments with seawater in the presence of calcite. In these experiments, embryonic opal-CT lepispheres could be detected after only one day. Precipitation of the lepispheres occurred preferentially on the surfaces of foraminifera.
3. In the presence of calcite, the concentration of dissolved silicon decreased compared with those without calcareous ooze, in which no opal-CT precipitation was observed.

Because of the known role of magnesium hydroxide as a coagulant in silica precipitation, particular attention was paid in these experiments on the effects of Mg(OH)₂ for the precipitation process. Apparently, the presence both of Mg-ions and hydroxy groups is essential for opal-CT crystallization, as shown by further results of Kastner et al. (1977).

4. In each of the experiments in which opal-CT crystallization occurred, the concentration of Mg²⁺ and (OH)⁻ was lowered in the ratio 1:2, parallel with the decrease in the concentration of dissolved silicon.
5. Newly formed embryonic opal-CT lepispheres always contained trace amounts of Mg.
6. In the absence of Mg (e.g., in experiments with artificial seawater lacking Mg), no opal-CT crystallization was observed during the experiment, even though calcite was present.
7. In experiments with initial borate alkalinity instead of bicarbonate alkalinity, opal-CT lepispheres formed in the same manner as in

![Figure 17 Thickness of seismic-stratigraphic units in the Pacific (from Wyllie, 1971; after Ewing et al., 1968). Isopach lines for upper transparent layer in tenths of seconds reflection time (two-way travel time, i.e., each contour represents approximately 100 m).](image-url)
the presence of bicarbonate indicating that it is the hydroxyl ion which is involved in the transformation and not the bicarbonate ion. 8. In experiments with montmorillonitic clays, dissolution of siliceous tests took place, but no reprecipitation of opal-CT occurred. However, small amounts of an Mg-rich clay formed.

The results highlighted the significance of a magnesium hydroxide compound (MHC) which attracts silanol groups with their high negative surface charge and causes them to precipitate and nucleate as opal-CT lepispheres. The MHC has been postulated to be the mineral sepiolite (Kent and Kastner, 1985). Opal-CT precipitation apparently is more rapid than sepiolite growth once the nuclei have formed, therefore opal-CT outdoes sepiolite. The role of calcium carbonate, which also dissolves in the process, is thought to resupply alkalinity consumed in the formation of the nuclei. Seawater supplies the necessary Mg. When the (OH)-groups have been exhausted, the rate of nucleus formation drops and the existing opal-CT embryos grow to well-developed lepispheres, (provided continued opal-A dissolution supplies the required dissolved silicon). In experiments with montmorillonitic clays, the clay minerals compete for the Mg and thus prevent the formation of the magnesium-hydroxide nuclei. Consequently, no opal-CT formed during the duration of the experiments (up to 6 months).

**Opal-CT precipitation as rate-limiting factor.** In a continuation of these hydrothermal experiments, Kastner and Gieskes (1983) determined the rate-limiting steps in the dissolution - reprecipitation process. The overall reaction rate is a function of the rates of dissolution, nucleation and growth of the silica phases involved. Which one of these mechanisms will be the rate-limiting factor depends on specific chemical and physical conditions. Amorphous silica of varying specific surface area was placed in aqueous solutions of MgCl₂ (0.03M) and NaHCO₃ (0.03M) and heated to temperatures of 50°C, 75°C, 100°C, 125°C, and 150°C for periods of 1 to 30 days (Figure 18). In all experiments, Mg and (OH) concentrations decreased in the 1:2 ratio previously established for the process, and opal-CT lepispheres or embryonic lepispheres formed. This confirmed that the MHC serves to nucleate the opal-CT. In experiments with low-surface-area silica (i.e., Eocene Radiolaria with a specific surface area of 5.3 m² g⁻¹), the opal-A dissolution rate apparently was the rate-controlling step at temperatures of 50°C, 75°C, and 100°C. At these temperatures, the concentration of dissolved silicon remained below the equilibrium solubility of β-cristobalite, the silica phase which most closely approximates a "poorly ordered opal-CT" and for which solubility data are available. Nuclei formation and growth of opal-CT apparently kept pace with opal-A dissolution at these temperatures. At 125°C and 150°C, however, opal-CT precipitation became the rate-limiting step after a few days, while the concentration of dissolved silicon rose above the equilibrium solubility of β-cristobalite (Figure 18). In experiments with higher-surface area silica (i.e., Ludox

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**Figure 18** Changes in dissolved silicon, magnesium and alkalinity as a function of time and temperature in hydrothermal experiments with Eocene Radiolaria (specific surface area of 5.3 m² g⁻¹). - , solubility of β-cristobalite; --A, solubility of opal-A (from Kastner and Gieskes, 1983, fig 1).

**Figure 19** Changes as in Figure 18, however for Ludox silica (specific surface area of 61.5 m² g⁻¹; from Kastner and Gieskes, 1983, fig. 2).
silica with a specific surface area of 61.5 m²·g⁻¹), the same behavior was found at all temperatures studied: β-cristobalite solubility was rapidly exceeded, but opal-A equilibrium solubility concentrations were not attained, again indicating that opal-CT precipitation was the rate-limiting process (Figure 19). These experiments also clearly illustrate the dominant role of temperature as a rate-controlling factor in the conversion of opal-A to opal-CT.

The opal-CT to quartz transformation: Temperature and rate. Temperature estimates for the opal-CT to quartz transformation based on oxygen isotope data and heat flow considerations for the Monterey Formation range from 55°C to 110°C (Murata et al., 1977; Murata and Larson, 1975; Pisciotto, 1981). The actual temperature at which the transformation occurs at a particular locality is a complex function of many factors, including the previous opal-CT maturation history. Pisciotto's (1981) temperature estimates for the base of the opal-CT zone in the Monterey Formation (35°-61°C), are considerably lower than those of other authors as they were for the top of this zone.

As discussed previously, opal-CT precipitation is the rate-limiting factor in the opal-A to opal-CT conversion, when dissolution of immature, high-surface-area opal-A is involved. Under such conditions, the concentration of silicon in solution will eventually approach the theoretical equilibrium solubility value for opal-CT, which is a function of the surface area of this phase for a given temperature. Lowering the equilibrium solubility for dissolved silicon through Ostwald ripening of the opal-CT phase counterbalances a solubility increase with rising temperature during burial. This is a prerequisite for quartz precipitation. A high-surface-area opal-CT at 50°C has a theoretical solubility of 130 ppm. The solubility of a low-surface-area opal-CT at 110°C is about 150 ppm (Fourrier, 1973), only slightly higher than that of the high-surface area opal-CT at 50°C. A solution “in equilibrium” with “ordered” opal-CT at 110°C, is therefore super-saturated only 1.5 times with respect to chalcedony or cryptocrystalline quartz. A solution “in equilibrium” with a high-surface-area (or “disordered”) opal-CT at 110°C on the other hand, would have a solubility of 290 ppm (Fourrier, 1973) or nearly 3 times the equilibrium solubility of chalcedony. Since the growth of a new (“more highly ordered”) silica phase such as chalcedony or quartz is favoured by low supersaturation, the “ordering” process during burial of opal-CT operates toward this end and ultimately facilitates quartz precipitation.

Lowered silicon concentrations are also required to understand the reduced d(101) spacing of the initial opal-CT in clayey host sediments. In the Monterey Formation, Isaacs (1982) observed that the first opal-CT to appear in a progressive burial sequence has a d(101) spacing which varies inversely with detrital mineral content of the host sediment (Figure 20). This observation confirms the results of Murata and Larson (1975) from the Chico Martinez Creek study that, at any given burial depth, the opal-CT d-spacing in (detritus-rich) porcellanite is 0.004-0.015 Å smaller than in associated (detritus-poor) quartz chert (Figure 12). This indicates retardation of opal-CT nucleation in clayey sediments because of the competition for MHC by clay minerals, consistent with the results of Kastner et al. (1977). Furthermore, it indicates delayed growth of opal-CT until silicon solubilities “in equilibrium” with a lower surface-area opal-CT (of a d-spacing of 10 or less) are encountered. Such reduced silicon concentrations in clay-rich host sediments can be accomplished most easily by adsorption of silica on detrital minerals (Siever and Woodford, 1973) and by the neof ormation of silica-rich clay minerals, provided their nucleation and growth rates are faster than those of the silica phases that could potentially form. These and other possible scenarios are discussed in more detail by Williams et al. (1985).

The findings of Isaacs (1982) summarized in Figure 20 also suggest that the detrital mineral content, while retarding the opal-A to opal-CT transformation, would actually enhance the opal-CT to quartz transformation, which is what one would expect from the discussion in the previous section. In sediments with more than 70% detrital minerals the initial opal-CT has a d(101) spacing of 4.08 Å or less. In these sediments, the transformation to quartz should occur earlier than in sediments with less than 30% detrital minerals with an initial d(101) spacing of 4.11 Å. This is because reduction of the opal-CT d-spacing to values below 4.07 Å (4.09 Å in certain Cretaceous cherts from the West Pacific (Pisciotto, 1980)) is a prerequisite for quartz precipitation. Such an inference, however, is at variance with the earlier observation (e.g., Lancelot, 1973) that the silica maturation sequence seems to be accelerated in calcareous host rocks, which leads to the relatively early formation of quartz chert.

This ambiguity might be resolved, if it is assumed that the initial retardation of the opal-A to opal-CT transformation in clayey sediments delays the entire silica maturation sequence to an extent which cannot be compensated for by later acceleration due to an abbreviated opal-CT stage. Although the opal-CT stage may in fact last longer in carbonates than in clay-rich siliceous sediments, contrary to Lancelot's (1973) assumption, this may be outweighed by a considerably earlier initiation of the transformation sequence in carbonates.

The field relationship between detrital content and opal-CT transformation rates

![Figure 20 Schematic summary of the role of host rock composition and temperature as controlling factors in the opal-CT maturation sequence. The initial d(101) spacing of opal-CT has been shown to vary inversely with detrital content. (From Isaacs et al., 1983; after Isaacs, 1982).](image-url)
reported by Isaacs (1982) from the Santa Barbara coast is not observed everywhere in the Monterey Formation. Pisciotto (1981), for example, found the appearance of opal-CT as well as quartz at lower temperatures in the Santa Barbara basin to be associated with lower detritus contents. The most recent study of the problem by Hinman (1987) investigated the role of organic matter for the transformation reactions. Her results suggest that in organic-matter rich sediments the rate of silica diagenesis should be reduced. Organic acids released by organic-matter maturation would dissociate and lower the pH thereby reducing carbonate alkalinity. This in turn would slow down the transformations by decreasing the silica polymerization rate. Although lowering of the pH will promote carbonate dissolution and thus buffer the pH, this effect may be nullified, if little carbonate is available or accessible for dissolution. In this case, no effect of carbonate content on the silica transformation rates may be seen, as in Isaacs’ (1982) study. The initial opal-CT formed in organic-matter rich sediments should have a lower d(101) spacing because of a slower nucleation and growth rate under decreased carbonate alkalinity. In so far as organic-matter content may be positively correlated with detrital mineral content, Hinman’s (1987) results are in accordance with Isaacs’ findings. However, Hinman (1987) did not find evidence that an initially lower d(101) spacing of opal-CT in organic-matter and detritus-rich sediments also accelerates the subsequent transition to quartz. In contrast, in the Santa Maria basin, quartz formation seemed to be slowed down because it occurred at higher temperatures in porcelanite than in chert.

It seems unlikely that the opal-CT stage can be bypassed entirely in the normal diagenetic silica maturation process, except under special conditions. The reasons why the diagenetic evolution path opal-A → opal-CT → quartz is observed in most situations have been outlined in the previous sections on reaction kinetics, and confirm the applicability of Ostwald’s step-rule for diagenetic silica transformations. Rare exceptions seem to be restricted to micro-environments such as the cavities of foraminifer (e.g., Keene, 1975) or Radiolaria in calcareous sediments. These are often filled with chalcedony, while the host rocks are calcareous opal-CT rocks or diatomaceous (opal-A) shales (Isaacs, 1982). This indicates anomalously early quartz (i.e., chalcedony) precipitation. Precipitation of the chalcedony occurred in spherical open cavities indicating limited overburden weight and very shallow burial. This chalcedony does not appear to have had an opal-CT precursor in the cavities, because elsewhere in the host rock opal-CT is still present which shows no sign that the transformation to quartz has started. Most likely, the chalcedony formed directly from opal-A, possibly because the concentration of dissolved silica in the microenvironment of the cavities, for some unknown reason, remained below α-cristobalite equilibrium solubility during the early stage of opal-A dissolution. Among the mechanisms discussed by Williams et al. (1985) to maintain low silica concentrations in pelagic carbonates, the early precipitation of calcium zeolites is a possibility, although it seems no specific studies have been directed at this problem.

Hinman (1987) suggested that direct quartz precipitation may be possible in the absence of Mg²⁺, if sufficient alkalinity is available. Mg²⁺ ions in solution appear to have an inhibiting effect on quartz precipitation. This behaviour is opposite to that of opal-CT crystallization, which is promoted in the presence of Mg²⁺ (and high alkalinity).

**Figure 21 (a, left)** Folded chert dyke in Lower Cretaceous radiolarian porcelanite, West Pacific. Porcelanite fragments in the dyke are cemented by microquartz and chalcedony which enclose considerable amounts of opal-CT sediment, except where the cement is very light-coloured (especially at the dyke margins). Folding of the microdyke was caused by differential compaction. The pure chalcedonic cement at the dyke margin was probably precipitated into pore space that was generated by shrinkage of the host sediment during ongoing compaction and was probably the latest cement phase. Height of thin-section approximately 2 cm. DSDP site 20-199A, sample 5cc, #1.

**Figure 21 (b, right)** Multiple fractures filled with injected opal-CT sediment and fragments which were cemented by microquartz and chalcedony. Margins and ends of the linear fractures are light-coloured and consist almost entirely of microquartz or chalcedony cement which probably was the last cement. Height of section approximately 2.5 cm. DSDP site 20-198A, sample 5cc, #4.
Summary: The "stability" fields of the various silica phases in deep-sea diagenetic environments. The distribution of the various silica phases in oceanic sediment sequences has been summarized by Riech and v. Rad (1979) on the basis of available DSDP data (Figure 16). There is considerable overlap on this depth/age graph between the fields for the three main silica phases. 

Opal-A has been found in sediments as old as 85 m.y. (Late Cretaceous), but only at shallow subsurface depths. With increasing depth (and temperature) its maximum survival time is shortened significantly. For example, at 1,000 m subsurface depth, opal-A is normally no longer present in sediments older than about 20 m.y.

Opal-CT takes a minimum of about 10 m.y. to first appear, and this requires elevated heat flow, otherwise its formation takes even longer. Exceptionally young opal-CT has been observed by Weaver and Wise (1973) in Pliocene sediments (less than 5 m.y.), where it occurred adjacent to igneous dykes and sills injected close to the sediment/water interface. On the other hand, opal-CT may still persist in sediments 100 to 120 m.y. old. Little new opal-CT forms, however; in pre-tertiary sediments (i.e., sediments older than 85 m.y.), because very little source material (opal-A) is left in those sediments. With age and increasing burial, opal-CT undergoes the progressive crystallographic structural changes in the d(101) plane described earlier. No opal-CT has been found in pre-Cretaceous sediments (older than 144 m.y.).

In Figure 16, the quartz field overlaps the opal-CT field, and to a slight degree the opal-A field, illustrating how local variations in heat flow, pore fluid chemistry, host-rock lithology and other factors can affect the rate of the diagenetic transformations. Quartz-chert cements need a minimum of 30 to 40 m.y. to form at burial depths of 500 m or more, and considerably longer at shallower depths. Genuine quartz cements, consequently, are rare in Conocoinclude pelagic sediments, but become predominant in Lower Cretaceous and older siliceous rocks. They are the exclusive lithology of silicious sediments in the Paleozoic and Precambrian.

Physical Diagenesis of Siliceous Sediments
The treatment of the diagenesis of siliceous sediments would be lacking an important aspect without a discussion of some phenomena of physical diagenesis in cherts. A step-wise compaction history is characteristic of siliceous sediments during burial, which contrasts with the more gradual compaction history of argillaceous sediments. This is implied in the constant bulk density of Figure 12 for the opal-CT stage and reflects the rapid crystallographic changes associated with the phase changes at the stage boundaries.

Other diagenetic features characteristic of chert formations are dykes, diapirs, and breccias which have been well described in the literature (e.g., Talafiero, 1934). They indicate differential compaction and variable rates of silica recrystallization and cementation. In a given sequence of siliceous sediments, some layers may already be lithified or semi-lithified while others are not. Shaking during an earthquake may rupture the lithified layers and cause injection of un lithified sediment into the fractures from adjacent un cemented layers (see numerous illustrations in Snyder et al., 1983), particularly if these had been overpressured. The opal-A to opal-CT and the opal-CT to quartz transformations are dehydration reactions. They may generate excess pore pressure and cause hydraulic fracturing. The examples shown in Figure 21 are Cretaceous radiolarian porcelanites from the West Pacific east of the Marianas deep-sea trench/island arc. The dykes probably reflect seismic activity associated with the Marianas subduction zone. The brittle host sediment is porcelaneous predominantly consisting of opal-CT with chalcedony-filled radiolarians. The fractures and dykes contain porcelaneous fragments and fine-grained opal-CT sediment cemented by microquartz and chalcedony. Some of the microquartz seems to have recrystallized in place from opal-CT. The folded dyke in Figure 21a behaved as a more competent layer, while the surrounding host sediment was still undergoing compaction. Similar mechanisms of dyke and vein formation are invoked by Stenitz (1970) and Snyder et al. (1983). Paris et al. (1985) ascribe chert breccias and veins in the Archean Barberton greenstone belt in South Africa to the injection of hot, high-pressure hydrothermal fluids related to volcanism. Some chert breccias, however, are of normal epigenetic, synsedimentary origin, such as those in the Burlington Limestone of Missouri which according to Carozzi and Gerber (1978) formed due to storm action on a Mississippian carbonate flat. They provide evidence for the early-diagenetic origin of some chert-nodules in limestones, which will be discussed in more detail in Hass (in press).

Conclusions
Our understanding of the origin of biogenic cherts, the main type of Phanerzoic cherts, has improved significantly during the last 15 years. Principal remaining open questions concern the mineralogical significance of the shift in the d(101)-spacing in opal-CT with maturation, which is closely related to the problem of solid-state reactions at low temperatures. To solve this problem, a better understanding of the crystallographic structure of the mineral opal-CT is required. The influence of host-rock lithology on the diagenetic silica transformations also requires further study. Results of Isaac's (1982) suggest that detrital content accelerates the opal-CT to quartz transformation while it has a retarding effect on the opal-A to opal-CT transformation (Kastner et al., 1977). However, the overall maturation seems to be faster in carbonates than in clayey sediments (Lancelot, 1973).

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References

General These cover the sections Introduction, Sources of non-detrital silica in siliceous sediments, Biogenic siliceous oozes, Pelagic stratigraphy, Formation of bedded chert, Shallow water chert, and Conclusions


Stages of Burial Diagenesis in Biogenic Siliceous Sediments - Characteristics of the Silica Phases in Siliceous Sediments


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