New Perspectives on Abiotic Organic Synthesis and Processing during Hydrothermal Alteration of the Oceanic Lithosphere

MURIEL ANDREANI AND BÉNÉDICTE MÉNEZ

15.1 Introduction

The main known organic compounds on Earth are biologically derived, whether they are direct products of biological activity or the result of thermal degradation of bio-derived material. While the synthesis of organic compounds from inorganic reactants is a common process in the chemical industry, it remains an unverified component of the deep carbon cycle on Earth and possibly on other planetary bodies. Abiotic organic synthesis is central to life emergence and sustainability, and possibly to “geo-inspired” resources. Intensive efforts are still needed to unravel the possible forms, sources, quantities, and formation mechanisms of abiotic carbon compounds under geologically relevant conditions. An improved knowledge of their processing within the lithosphere is also mandatory to better quantify their impact on biogeochemical cycles and their contribution to C fluxes between Earth’s external and internal envelops. Their presence in fluids and rocks may also affect the kinetics of fluid–silicate reactions and the fates of other elements, particularly the redox-sensitive ones (e.g. transition metals, S).

Abiotic organic compounds in the lithosphere can have two main origins: either rising from a deep volcanic source in the mantle or formed in situ in the upper lithosphere during hydrothermal processes from mantellic or seawater inorganic carbon compounds (see Refs. 1 and 2 for reviews). In the present chapter, we focus on lithospheric hydrothermal processes that include late magmatic stages and fluid–rock reactions.

In fluids, natural occurrences of recognized abiotic organic volatiles of hydrothermal origin include methane (CH₄), short-chain alkanes, and small organic acids. They have mainly been observed in geothermal systems or continental seepages within ophiolites and Precambrian shields³ and at hydrothermal vents near mid-ocean ridges⁴ and subduction forearcs.⁵ Hence, they are not necessarily associated with deep active volcanism, and they attest to the contribution of lithospheric hydrothermal processes to abiotic organic synthesis. In most cases, these occurrences of organic volatiles are associated with high concentrations of H₂ reached by reduction of water during aqueous alteration of ferrous iron-bearing minerals. The latter are particularly abundant in mantle and olivine-rich lithologies whose alteration process is known as serpentinization.
CH$_4$ and short-chain hydrocarbons have attracted scientific attention over recent decades, and different reactions have been proposed to explain their possible abiotic formation.$^{1,3}$ The ones occurring in the oceanic lithosphere include re-speciation and high-temperature (T) reactions (>400–500°C) of magmatic C–O–H fluids during cooling, carbonate decomposition to CH$_4$ (<800°C), and inorganic carbon reduction by H$_2$ at lower T (<400–500°C). The latter has been the most investigated pathway and may occur through Fischer–Tropsch-type (FTT) reactions under gaseous or aqueous conditions. However, the exact mechanism and the nature of potential metal catalysts are still debated (see reviews in Refs. 1 and 6). Up to now, experimental results have converged toward kinetic inhibition of CH$_4$ formation at low T (<300–400°C) according to the low methane yield achieved in most experiments (see Section 15.3.1). This is in agreement with recent work putting forward a dominant deep source for CH$_4$ venting at mid-ocean ridges.$^{7,8}$ This deep CH$_4$ may result from the entrapment and re-speciation at high T (>400°C) of mantle-derived CO$_2$ within fluid inclusions or vesicles in magmatic rocks,$^{9,10}$ the fluids being later released during hydrothermal alteration of oceanic rocks. Conversely, no consensus has yet emerged to explain the origin of CH$_4$ seepages observed in continental settings where fluid–rock reactions occur at even lower T (<100°C). Such settings may offer favorable environments for gas-phase reactions that are more efficiently catalyzed than aqueous ones.$^{11}$ However, most of these assumptions are based on CH$_4$ carbon isotopic signatures that can be highly similar to those produced by microbial methanogenesis under alkaline conditions at high levels of dissolved inorganic carbon.$^{12,13}$

Among organic acids, formate is the only one so far recognized as being of possible abiotic origin in nature, resulting from the equilibration in aqueous solution of CO$_2$ or carbonate ions and H$_2$ under neutral to alkaline conditions.$^{14-16}$ A variety of other (e.g. methanethiol, amino acids) and heavier (aliphatic and aromatic hydrocarbons, organic acids) organic compounds were also observed in hydrothermal vent fluids and chimneys, but they are often considered as biologically derived.$^{16-24}$ Should abiotic synthesis of such compounds occur, they are likely to be too diluted in hydrothermal fluids to be distinguished from background biological contributions.$^{24}$

Bulk analyses of hard rocks from the oceanic lithosphere describe up to 1500 ppm of total organic carbon (TOC) in serpentinized abyssal peridotites.$^{25-28}$ If some are present as volatile organic compounds adsorbed on minerals, a fraction is possibly present as solid carbonaceous matter (CM). The nature, diversity, and origin of such organic phases remain poorly constrained, and one cannot preclude a biological contribution to their formation$^{29-31}$ and a thermogenic origin as for kerogens.$^{30,31}$ Part of this controversy is due to the fact that occurrences were found in potentially colonized environments and that spectroscopic, chemical, and isotopic evidence lacks unequivocal criteria to discriminate between abiotic and biotic origins. In addition, most of the conditions and reactions leading abiotically to carbonaceous compounds are mostly unknown, including their relationship with methane. The experimental difficulty to produce sufficient CH$_4$ during hydrothermal synthesis compared to what is observed in nature suggests alternative organic products.
being preferentially formed and leaves the door open to the existence of metastable compounds in the shallow lithosphere.32–35

All of these data force one to consider new paradigms for abiotic organic synthesis in hydrothermal settings, not being CH₄ centered, but rather including the potential formation of organic carbon compounds with intermediate oxidation states, possibly mediated by rock-forming minerals. In this chapter, we discuss natural occurrences of carbonaceous compounds within the hard rocks of the oceanic lithosphere or analogs to complement the numerous studies and reviews focused on organic compounds in fluids discharged at oceanic hydrothermal vents or continental seeps (e.g. Refs. 3 and 20). Data on natural rocks affected by low-T (<400°C) and high-T (>400°C) hydrothermal reactions are compared to the most pertinent experimental and theoretical data currently available. Some resulting assertions remain hypothetical but provide new schemes for considering and investigating organic synthesis in lithospheric hydrothermal environments.

15.2 Carbonaceous Matter in Hydrothermally Altered, Mantle-Derived Rocks

Accumulations of CM can be found from the Proterozoic to the present time in volcanic and igneous rocks from a diversity of geodynamic contexts, including spreading zones, oceanic hot spots, island arcs, and continental rifts.36,37 We focus here on the organic carbon suspected to be nonbiological based on investigations of the most favorable rocky environments for abiotic organic synthesis (i.e. during hydrothermalism affecting mantle-derived rocks).

15.2.1 Bulk Rock Investigations

The first mention of the presence of CM in mafic and ultramafic rocks came from the vast Russian literature published since the 1950s. At that time, the so-called Russian–Ukrainian School had spent significant effort in exploring hydrocarbons in mafic and ultramafic rocks to support a deep mantle origin hypothesis for hydrocarbons deposits. We refer to Sephton and Hazen for a historical perspective and an inventory of achievements in this field. Although a large part of these studies remains untranslated, a review of the occurrences of putatively abiotic condensed naphtides showed a large diversity of compounds associated with volcanic and igneous rocks in a wide range of geodynamic contexts.37 These compounds include aromatic and aliphatic hydrocarbons and their N-, O-, and S-bearing derivatives. Notably, they mainly occur in mafic and ultramafic rocks. The first study targeting oceanic rocks has focused on the Rainbow (36°14’N) and Logatchev (14°45’N) hydrothermal fields (Mid-Atlantic Ridge (MAR)).38 In addition to the presence of low-molecular-weight alkanes and isoalkanes, this study reported the presence of CM in serpentinized rocks, in metalliferous sediments, and in secondary Fe–Ni sulfides. The Fe–Ni sulfides presented the highest concentrations and the broadest diversity of viscous and solid organic compounds depicted as resinous bitumen. Notably, a compositional
relationship was established between these resinous bitumen and hydrothermal crystalline hydrocarbons identified as karpatite (C\(_{24}H_{12}\)) and idrialite (C\(_{22}H_{14}\)) found in the same rock samples and thought to derive from serpentinization reactions.\(^{38}\)

Karpatite (Figure 15.1) and idrialite correspond to polycyclic aromatic hydrocarbon (PAH) minerals. PAH minerals are the most typical molecular organic minerals with well-defined chemical compositions and crystallographic properties.\(^{39}\) These molecular crystals can form thanks to the high stability of PAHs in hydrothermal fluids,\(^{40}\) allowing transport without thermal degradation and subsequent concentration and crystallization at the hydrothermal discharge zone.\(^{41,42}\) Coronene and phenanthrene, the precursors of karpatite and of a number of natural molecular organic minerals (Table 15.1), were shown to derive from the hydrothermal alteration of organic matter present in sediments,\(^{43}\) but can also be produced during basalt cooling (Section 15.3.4).\(^{44,45}\) Although the mechanisms of PAH concentration prior to crystallization is still unknown, occurrences of molecular organic

<table>
<thead>
<tr>
<th>Mineral</th>
<th>Chemical formula</th>
<th>Chemical nomenclature</th>
<th>Refs.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Karpatite</td>
<td>C(<em>{24}H</em>{12})</td>
<td>Coronene</td>
<td>48</td>
</tr>
<tr>
<td>Idrialite</td>
<td>C(<em>{22}H</em>{14})</td>
<td>Picene</td>
<td>49</td>
</tr>
<tr>
<td>Kratochvilite</td>
<td>(C(_4H_8))CH(_2(C_6H_4))</td>
<td>Fluorene</td>
<td>50</td>
</tr>
<tr>
<td>Simonellite</td>
<td>C(<em>{19}H</em>{24})</td>
<td>1,1-Dimethyl-7-isopropyl-1,2,3,4-tetrahydrophenanthrene</td>
<td>51</td>
</tr>
<tr>
<td>Evenkite</td>
<td>(CH(_3)) (CH(<em>2))(</em>{22}(CH_3))</td>
<td>n-Tetracosane</td>
<td>52</td>
</tr>
</tbody>
</table>

Figure 15.1 (a) A vein of karpatite (yellow crystals) surrounded by quartz (white crystals) and cinnabar (red spots). Scanning electron micrographs of (b) the broken surface of native karpatite and (c) its layering at the end of the layered structure. Reproduced with permission of Springer Nature, from Potticary et al. (2017), *Sci Rep*, 7, 9867, figures 1b, 2a,b.\(^{46}\)
minerals may attest to localized high concentrations in (poly)aromatic compounds in the oceanic lithosphere.

More recently, three studies characterized by bulk molecular approaches (1) serpentinized peridotites and gabbroic rocks recovered at the Atlantis Massif (30°N MAR) at both the Lost City hydrothermal field and Hole U1309D (Expeditions 304/305, Integrated Ocean Drilling Program (IODP)),26 (2) peridotites from the Ashadze (12°58'N) and Logatchev hydrothermal sites along the MAR,53 and (3) the fossil ocean–continent transition recorded in the Swiss Alps.54 All of these studies used gas chromatography and/or gas chromatography–mass spectrometry. Based on the presence of biomarkers (e.g. pristane, phytane, squalane, hopanes, and steranes) or higher relative abundances of \( n\)-C\(_16\) to \( n\)-C\(_{20}\) alkanes,26,54 amino acids, and long-chain \( n\)-alkanes53 identified in the solvent-extracted fraction, they all concluded that the organic carbon was of biological origin, as supported by TOC isotopic analysis of the oceanic rocks.25,28 However, biological contamination may have overprinted any possible abiotic geochemical signatures,26 since these oceanic basement rocks have been affected by long-lived hydrothermal alteration and present-day microbial ecosystems.29–31

15.2.2 In Situ Investigations at the Microscale

Development of in situ techniques currently allows imaging of rock-hosted CM along with their relationship with mineral parageneses. The scarce data for oceanic rocks report that the carbon phases exhibit a structural organization varying from amorphous to well-organized graphite-like material (Figure 15.2a and b). Organic compounds have also been detected as thin films or gel-like materials embedding minerals and filling microfractures (Figure 15.2c). As is detailed below, some have an origin that is still debated, depending on the geological setting and textural criteria.

The Hyblean basaltic diatreme (Sicily, southern Italy), recognized as a paleo-oceanic serpentinite-hosted hydrothermal system comparable to those found along slow-spreading ridge segments,57 is one of the most studied places where the presence of large amounts of putatively abiotic carbonaceous compounds has been related to hydrothermal activity. A series of papers reported the occurrence of heavy hydrocarbons in deep-seated xenoliths including metasomatic gabbroic xenoliths47 and highly serpentinized peridotite xenoliths.58 In the gabbroic xenoliths, the presence of saturated aliphatic and aliphatic–aromatic hydrocarbons was highlighted by bulk rock analysis using electron impact-direct pyrolysis mass spectra. These observations were supported by Fourier-transform infrared (FTIR) spectroscopy showing carbon-cored clayey vesicles and dull-black, tiny pellets associated with the clayey mat pervasively intruding into fractures of the host rocks.47 Organic crystalline phases were also detected using X-ray diffraction (Table 15.1), and thermal decrepitation allowed the analysis of hydrocarbons trapped in fluid inclusions. Similarly, in the extensively serpentinized and carbonated ultramafic xenoliths, microscopic accumulations of sulfur-bearing organic matter commonly occur between hydrothermal minerals
(i.e. secondary calcites and fibrous phyllosilicates), forming occasionally coarse bituminous patches. The organic matter was estimated to represent 3–6% of the whole rock. Micro-FTIR spectroscopy highlighted the presence of condensed aromatic rings with aliphatic tails consisting of a few C atoms, which is suggestive of asphaltene-like structures. Such structures were confirmed by solubility tests in toluene and n-hexane, thermogravimetric analyses and differential thermal analysis. X-ray photoelectron spectroscopy (XPS) also indicated minor S and O functional groups.

In these studies, it has been proposed that asphaltenes and high-molecular-weight hydrocarbons, respectively, derived from in situ aromatization or progressive polymerization and polycondensation reactions of light aliphatic hydrocarbons formed by FTT reactions. The hydrocarbon-bearing xenoliths possibly represent any level of the hydrothermal system, including its deepest and hottest parts at ~400°C. Nevertheless, both biotic and abiotic origins could have been possible. It is challenging to infer the possible processes and formation conditions giving rise to heavy hydrocarbon accumulations in such a complex geologic setting. Mantle xenoliths experienced transport and decompression during the formation of the diatreme, with possible consequences for CM that are unrelated to hydrothermal circulation.

Indeed, CM commonly occurs in basalts and mantle xenoliths, which were thoroughly investigated during the 1980s and 1990s. At that time, the aim was to assess the mantle carbon content by using a suite of techniques including electron microscopy, chemical imaging, XPS, and carbon isotopes, as well as thermal desorption surface analysis by laser ionization and low-energy electron diffraction. In these rocks, CM was
observed as discrete platy lumps of up to 20–200 µm in size or as thin, amorphous films of a few nanometers located on quench-produced crack surfaces, grain boundaries, and the walls of fluid inclusions. They consisted dominantly of graphite-intercalated compounds along with ill-defined complex mixtures of graphite-like compounds and organic material composed of C, H, and possibly N. All of these studies concluded that CM had an abiotic origin based on: (1) its preferential concentration on sulfide spherules attached to vesicle walls,\textsuperscript{60} where the sulfides may have played a catalytic role in organic compound formation and concentration as also observed for hydrothermal sulfides,\textsuperscript{38} or (2) the close association of carbon with Si, Al, alkalis, halogens, and/or transition metals, which are elements that were likely present in the volcanic gas at the origin of these carbon accumulations.\textsuperscript{62} Again, to account for the production of CM in basalts and mantle xenoliths, these studies invoked: (1) FTT reactions involving volcanic gases degassed of host lava and reacting with fresh and chemically active crack surfaces formed by thermal stresses during eruption, decompression, and cooling; and (2) subsequent evolution of the condensate during cooling. Although heterogeneous catalysis at the mineral surface was the favored hypothesis, organics may have been alternatively assimilated into the volcanic gases prior to eruption and deposited on cracks formed during eruption and cooling.\textsuperscript{62}

The possible role of mineral surfaces in the abiotic formation of CM in the oceanic lithosphere was pointed out in a series of recent studies. Various associations between minerals and condensed CM were documented using scanning electron microscopy (SEM) and Raman spectroscopy within magma-impregnated, mantle-derived serpentinites of the Ligurian Tethyan ophiolites, in a context of common occurrences in the lower oceanic crust (Figure 15.2a and b).\textsuperscript{55} Three distinct types of CM in paragenetic equilibrium with low-T mineralogical assemblages have been sequentially formed at decreasing T during the hydrothermal alteration of the rock assemblage. The first type corresponds to thin films of aliphatic chains, coating hydroandraditic garnets in bastitized pyroxenes. The second type forms micrometric aggregates associated with the alteration rims of spinel and plagioclase. The third and most massive type appears as large aggregates (up to 200 µm in size) bearing highly aromatic carbon and short aliphatic chains associated with hematite and Fe-saponite assemblages replacing the pseudomorphoses after plagioclase (Figure 15.2a and b). The systematic association of a given type of CM with a specific mineral paragenesis indicates that condensed CM precipitated simultaneously to the growth of the host mineralogical assemblage, overall in favor of an abiotic endogenesis. The mineral formation would have been accompanied by the production of H\textsubscript{2} able to reduce carbon species, as is the case when ferric hydrogarnets and ferric serpentines form\textsuperscript{64–66} or when spinel oxidizes to Cr-magnetite forming ferritchromite rims.\textsuperscript{67} This possible abiotic pathway was not considered in former studies performed on hydrogarnet-hosted carbonaceous compounds that suggested, based on their biological Raman and FTIR spectroscopy signatures, that this disordered CM may have resulted from the hydrothermal alteration of cryptoendolithic microbial ecosystems.\textsuperscript{30,31} Overall, while the involved mechanisms and reactants are still unknown, this recent study emphasizes a key role of local parageneses forming unique microenvironments prone to the synthesis of
The productivity could have also been controlled by the local presence of catalysts such as Cr$^{3+}$ in hydrogarnets or Fe-saponite in addition to specific redox conditions among the large redox potential gradients existing in serpentinizing systems down to the microscale (Figure 15.3). The catalytic role of phyllosilicates in abiotic organic synthesis has been suggested by the syngenetic link between phyllosilicates and organic compounds in several hydrothermal systems. The abundance of long-chain aliphatic and aromatic hydrocarbons (identified using FTIR spectroscopy) has been reported in a diapir of saponite-dominated clays intruded in a diatremic tuff-breccia deposit. The clays formed during the high-T (350–400°C) hydrothermal alteration of mafic and ultramafic lithologies in the Hyblean crustal basement. Based on the close association of hydrocarbons with hydrothermal clays, the lack of fossils, and the resemblance with the previously described xenolith-hosted organic compounds, a possible abiogenic origin was suggested for the hydrocarbons via FTT reactions mediated by the organoclay. A close association of organic compounds with clays minerals such as saponite has been previously described in interplanetary dust and in carbonaceous meteorites that have undergone significant aqueous alteration processes. A similar association was recently highlighted at a micrometric scale using FTIR spectroscopy in oceanic serpentinites collected at ~170 m depth below seafloor during the IODP Expeditions 304/305 targeting the Atlantic Massif.

Overall, all of these studies highlight that hydrothermally derived organic carbon trapped within the upper lithosphere as heavy and aromatic compounds can be chemically and structurally diverse, although formation mechanisms are not yet well-known. Whether aromatization occurs in situ from CO$_2$/CO and H$_2$ during serpentinization or derives from progressive polymerization and polycondensation reactions of light aliphatic ± aromatic hydrocarbons inherited from higher-T FTT reactions still needs to be addressed. This latter
scenario implies that reactants for the low-T (<400°C) hydrothermal synthesis of abiotic organic compounds that may occur during serpentinization might be more diverse than H₂ and CO₂/CO and account for aliphatic and aromatic compounds, as supported by the presence of molecular organic minerals in serpentinized rocks (Sections 15.2.1 and 15.2.2).³⁸,⁴⁷

15.2.3 Carbon in Fluid Inclusions Trapped in the Oceanic Lithosphere

Fluid inclusions are used to document the composition of high-T fluids (either of magmatic or seawater origin) circulating in the lower crustal component of hydrothermal systems. Hence, they can inform on the intermediate abiotic processes that may happen at higher T and then feed with diverse reactants the low-T hydrothermal reactions, including organic synthesis. Fluid inclusions are made of liquid and/or gas with or without tiny daughter minerals that are trapped within a crystal structure during either primary crystallization or secondary healing of fluid-filled cracks. They occur throughout the gabbroic and peridotitic plutonic crust⁷⁴ and have been vastly studied by Raman spectroscopy and microthermometry. In ophiolites, fluid inclusions formed in sub-seafloor rocks may be preserved during tectonic uplift and maintain their original chemical signatures. This has been shown for pure methane fluid inclusions occurring in olivine from partially serpentinized harzburgites and dunites from the Nidar ophiolite complex (eastern Ladakh, India).⁷⁵ In oceanic rocks, all studies have pointed to the presence of CH₄ ± H₂ ± CO₂ within fluid inclusions,⁹,¹⁰,⁷⁴,⁷⁶,⁷⁷ although propane and ethane were also sometimes reported in the gaseous phases.⁷⁸ Analyses of fluid inclusions in primitive olivine gabbros, oxide gabbros, and evolved granitic rocks recovered from the slow-spreading Southwest Indian Ridge at Ocean Drilling Program Hole 735B (Atlantis II fracture zone) recorded CH₄ concentrations 15–40 times those of hydrothermal vent fluids from sediment-poor environments and of basalt-derived volcanic gases.¹⁰,⁷⁴ These studies also frequently reported the presence of disordered graphite, carbonaceous compounds, and putative graphite coating inclusion walls.⁹,⁷⁴,⁷⁶

While several processes were proposed to explain the presence of organic compounds in high-T hydrothermal fluids,⁷⁹,⁸⁰ fluid inclusions are generally believed to represent evolved magmatic fluids dominated by CO₂ or CO.⁹,¹⁰,⁷⁴ Indeed, subsequent to devolatilization and re-equilibration, graphite precipitation is promoted by re-speciation of magmatic CO₂, and the formation of CH₄-enriched fluids is promoted by cooling from 800°C to 500°C of these H₂-rich fluids. Alternatively, re-speciation and reduction of entrapped CO₂-bearing magmatic fluids by diffusion of external H₂ into the inclusions during cooling was also considered.⁹,⁷⁴ Once trapped within crystals as fluid inclusions, the fluids can still evolve following reactions between the fluid and the host mineral. Direct evidence for in situ H₂ production and strongly reducing conditions within serpentinizing olivine fluid inclusions was provided for the Mineoka ophiolite complex considered as an analog to serpentinite-hosted hydrothermal vent systems.⁸¹ In the latter study, mineral-phase
equilibria indicated that CH₄–H₂-bearing fluids were trapped under equilibrium conditions at T below 300°C, and the absence of CO₂ was suggestive of extensive reduction of CO₂ to CH₄ within the inclusions. Whether CH₄ was also produced at higher T remains an open question. The presence of H₂ and CH₄ together with secondary mineral microinclusions was similarly reported in olivine and orthopyroxene crystals from a harzburgite from the northern Oman ophiolite considered to be similar to abyssal peridotite. In olivine, the mineral inclusions mainly consisted of lizardite and brucite with small amounts of magnetite, while in orthopyroxene they were made of talc and chromian spinel. The differential in situ production of reduced gas and secondary phases was related to the presence of either magnetite or a magnetite component in chromium spinels. Alternatively, both H₂ and CH₄ inherited from high-T magmatic processes could have hydrothermally evolved differentially depending on the reactivity of their mineral host (e.g. its capacity to produce in situ H₂ or to form mineral byproducts able to catalyze organic synthesis, as discussed in Section 15.2.2).

The presence of reduced carbon species in fluid inclusions within oceanic gabbros and mantle peridotites could represent a potentially important source of organic compounds in hydrothermal fluids. In particular, CH₄-rich aqueous fluids trapped in the oceanic lithosphere within fluid inclusions recently redrew attention as they could constitute the main source of methane venting at unsedimented mid-ocean ridge hydrothermal fields. Similar processes were invoked to explain elevated concentrations of both CH₄ and H₂ in fluids from the Menez Gwen (37°50’N, MAR), Lucky Strike (37°17’N, MAR), and Piccard (Mid-Cayman Ridge) vent fields usually referred to as basalt-hosted vents supposedly less rich in reduced gases compared to ultramafic rock-hosted vents, despite the possible production of non-negligible amounts of H₂ by diking-eruptive events.

Overall, in agreement with Section 15.2.2, studies on fluid inclusions emphasize the need to consider a larger range of reactants in addition to CO₂/CO and H₂ for low-T hydrothermal synthesis of abiotic organic compounds. This includes methane and light hydrocarbons, but also potentially reactive graphitic phases. Due to their tiny size and their entrapment in minerals, the latter are extremely challenging to analyze, and most of the previous studies seldom went further in characterizing the chemical diversity and degree of structural order of this CM. The precipitation of highly crystalline hydrothermal graphite from aqueous fluids containing CO₂ and CH₄ was reported at T as low as 500°C during the propylitic hydrothermal alteration of volcanic host rocks. Other studies, targeting different geodynamical contexts, have shown that precipitation of CM in fluid inclusions, notably on their walls, can lead to graphitic material with varying degrees of crystallinity and disorder. Disordering in graphite is principally caused by in-plane defects and/or heteroatoms (e.g. O, N, S). While crystalline graphite was shown to be highly refractory and chemically inert, the chemical reactivity of graphitic carbon increases with structural disorder and the abundance of heteroatoms and unsaturations. Hence, this possibly impacts the diversification of abiotic organic compounds observed in the altering oceanic lithosphere.
15.3 Comparison with Experiments and Thermodynamic Predictions

The heterogeneity of natural systems and the multistage character of hydrothermal alteration seem crucial for organic synthesis. This considerably complicates the identification of organic products, their origin, and the abiotic reaction sequences. A complementary experimental and theoretical approach is therefore mandatory to unraveling reaction paths and rates and determining the most favorable conditions that may lead to the abiotic formation of CM. The literature reviewed in the next sections highlights the gap between natural observations and experiments (Sections 15.3.1–15.3.3) and opens a wide field for future investigation, supported by thermodynamic predictions (Section 15.3.4).

15.3.1 Experimental Approach

Most of the experimental work dedicated to hydrothermal organic synthesis has focused on fluids, especially on the production of volatile organic compounds, using either monophasic (pure liquid) or biphasic (liquid + gas) systems. Solid mineral phases have not been systematically introduced in the experimental devices, and potential carbonaceous phases were rarely characterized after experiments. When present, minerals were either used as reactants, such as olivine to produce H₂, or as redox buffers such as the hematite–magnetite (HM), hematite–magnetite–pyrite (HMP), pyrite–pyrrhotite–magnetite (PPM), or quartz–fayalite–magnetite (QFM) assemblages. Minerals have also been introduced as potential catalysts of organic reactions. Unfortunately, they were never characterized after experiments, precluding the finding of CM occurrences.

Hence, the role of minerals on the formation of carbonaceous compounds under hydrothermal conditions, as emphasized from natural observations, has remained largely unexplored. This is partly explained by the analytical difficulty of detecting such a small organic fraction within the solid products (concentration close to or below detection limits) and of locating and characterizing it with high-resolution methods. In addition, contamination issues (from the experimental setup or deriving from organic compounds preexisting in the minerals used; e.g. trapped in fluid inclusions) remain central when looking for low levels of organic products whose nature is unknown and difficult to address.

The role of the experimental container (vessel or capsules) on CO₂ reduction or H₂ production reactions under hydrothermal conditions is potentially non-negligible while not clearly established. Stainless steel reactors have been shown to accelerate FTT reactions compared to quartz, glass, or Au and TiO₂ reactors. However, stainless steel reactor walls may be passivated after several hours of experimental runs. That is also the case for titanium, whose oxidation can be a source of H₂ if a preoxidation is not done prior to experiments. Gold, in the form of nanorods or particles, is well known to catalyze aqueous CO₂ reduction to CO during electrochemical experiments, but it is not usually considered for hydrothermal experiments, probably because CO was either not measured or not abundant in products, and also because the gold liner is seen as a smooth surface. Hence, vessels or capsules made of gold or oxidized titanium are usually thought to be the
most inert materials and are often preferred at high pressure (P) and T to stainless steel, platinum, alumina, or hastelloy (Ni–Fe-rich alloy). The effect of hastelloy may depend on the reaction of interest. It was infrequently used for FTT reactions, probably because of the potential catalytic properties of the Ni-rich metal alloy, but it was used for H₂ production. Its catalytic effect has not yet been demonstrated for such reactions. The use of Teflon™ especially may be a direct source of carbon contamination with increasing T. Concerning reactor permeability to gas, H₂ has a high diffusivity in most metals, but H₂ loss is expected to be relatively low at T < 400°C.

Hence, there is no homogeneity in the type of reactor used in the literature, which often precludes comparison between results, especially when the mineral effect has to be unraveled. In addition, the possible precipitation of a carbonaceous phase on the reactor wall has never been investigated.

### 15.3.2 Carbon-Bearing Reactants in Experiments

Oceanic hydrothermal systems developing near ridge axes have two main sources of inorganic carbon: (1) mantle-derived carbon delivered by magmatic activity in which CO₂ dominates in the C–O–H system, with very minor CO and CH₄; and (2) dissolved inorganic species that result from the equilibration of atmospheric CO₂ and seawater. Hence, aqueous CO₂ (CO₂(aq)) and bicarbonate and carbonate ions (∑CO₂⁻) are usually the most abundant species in solutions and are the preferential sources of carbon used in experiments and models designed to test abiotic organic synthesis under hydrothermal conditions mimicking natural systems.

Nevertheless, ∑CO₂ are not the only single-carbon compounds possibly available at equilibrium in hydrothermal fluids, especially if H₂ is available. Experiments have shown that the speciation of aqueous single-carbon compounds in the C–O–H system for T = 150–300°C and P = 35 MPa is controlled by reactions between ∑CO₂, CO, ∑HCOOH (HCOOH, formic acid + HCOO⁻, formate), CH₂O (formaldehyde), CH₃OH (methanol), and CH₄ (Figure 15.4). Indeed, the water–gas shift reaction (15.1) under aqueous hydrothermal conditions leads to the formation of ∑HCOOH as an intermediate product (Figure 15.4):

\[
CO + H₂O = CO₂ + H₂. \tag{15.1}
\]

∑HCOOH reaches a redox-dependent equilibrium with methanol, formaldehyde, and CH₄ within few days at T > 150°C, but may need years at T < 100°C. Formaldehyde and CH₄ were close to below detection limits in the experiments, suggesting kinetic inhibition at least for the P–T range tested. In the absence of CH₄, the relative concentrations of single-carbon compounds in fluids strongly depended on T, H₂ fugacity (fH₂), and pH. Under neutral and acidic conditions, CO₂ largely dominated, with minor amounts of CO and ∑HCOOH (several orders of magnitude < CO₂) at 350°C. At 150°C and similar
pH, methanol is predicted to be only one order of magnitude below CO$_2$ with minor CO and $\Sigma$HCOOH, but it could exceed CO$_2$ if the H$_2$ concentration increased to values similar to those measured at ultramafic rock-hosted hydrothermal vents (i.e. ~10 mM$^{102}$). Under alkaline conditions, the formate concentration can be equivalent to bicarbonate with decreasing T or increasing H$_2$ concentrations.

Direct formation of methanol (~1 mM max) from a H$_2$–CO$_2$-rich vapor phase can also occur, with a limited conversion rate (10$^{-4}$%) at 300–350°C (18 MPa), provided magnetite surfaces are available.$^{95}$ This could be realistic at mid-ocean ridges (e.g. following dike emplacement with volatile exsolution and migration in adjacent oxide gabbro or in serpentinized peridotites where magnetite is abundant). Magnetite shows decreasing surface reactivity with time and can be regenerated with increasing T, suggesting that it may serve as a catalyst during successive diking events, for instance. The authors do not preclude possible intermediates such as CO and $\Sigma$HCOOH in the redox reaction, but these compounds were not detected or measured, respectively, and no graphitic phase or alkanes were observed.$^{95}$

These works show that a large variety of single-carbon species can be available in fluids in addition to $\Sigma$CO$_2$ (mainly methanol and formate), especially in natural serpentinizing systems with highly variable pH and local H$_2$ levels.$^{64}$ This is particularly true in low-T serpentinizing environments that may be dominated by methanol or formate depending on pH, provided equilibrium is reached. This diversity reinforces the conclusions of Section 15.2 about the need to consider a wider range of carbon-bearing reactants for organic synthesis in natural systems and consequently in experiments. Natural systems also pointed to the possible availability of PAHs, CM, and deep magmatic CH$_4$ for low-T hydrothermal reactions. Except for formic acid, whose decomposition in aqueous fluid (Figure 15.4) is frequently exploited to experimentally produce H$_2$ and CO$_2$, these compounds have rarely (e.g. methanol$^{103}$) or never been used as reactants in geologically relevant experiments.

CH$_4$, which is abundant in hydrothermal fluids or cold seepages associated with the serpentinization of mantle rocks,$^{3,102}$ was usually considered as the reduction product of

Figure 15.4 Oxidation states of carbon in some single-carbon organic compounds. The water–gas shift reaction (15.1) is represented along with the successive reversible redox reactions that control the speciation of single-carbon compounds under hydrothermal conditions.$^{44,101}$
inorganic carbon sources rather than as a possible reactant. An important experimental effort was deployed to reproduce methane synthesis by FTT reactions under moderate to low-T conditions (\(\leq 500^\circ C\)), but systematically failed to produce abundant CH4. The natural H2/CH4 ratio measured in serpentinization-related oceanic hydrothermal environments (<30)\(^{102}\) remains much lower than experimental ones (>500; except values at 42\(^{104}\) and 17\(^{105}\) e.g. see McCollom\(^{106}\) for a review). This difference was proposed as an indication of the abiotic versus biotic origin of CH4 in natural systems, where intense H2 consumption and associated CH4 production can be attributed to biological activity.\(^{107}\)

This simplistic criterion was refuted because it cannot account for the complex processes occurring in the environment,\(^{108}\) and it also disregarded some other parameters. The H2/CH4 ratio may vary with T, decreasing as T increases from 200°C to 500°C at 300 MPa.\(^{109}\)

The effect of pressure has been seldom investigated, but a few studies have shown that increased pressure can account for a significant increase in the CH4 yield between 100 and 350 MPa.\(^{110}\) CH4 formation can also compete with CO2 carbonation depending on fH2, CO2 partial pressure, and T. At 200°C, CH4 formation by CO2 reduction is limited by H2 production during olivine alteration if the system is supersaturated with respect to carbonate because the kinetics of carbonate precipitation are faster. Finally, the micromolar levels of CH4 often found in experiments might not even be produced by in situ reactions, but instead may represent contamination. This is often difficult to assess since blank experiments are rarely provided and \(^{13}\)C-labeled experiments are scarce. Contamination issues are critical in very-low-T experiments (<100°C) where product levels are much lower and display contrasting values of H2 and CH4 despite similar protocols.\(^{13,67,111-115}\) The most complete investigation conducted so far at T \(\leq 100^\circ C\) led to the formation of low-molecular-weight organic acids (mainly formate and acetate); no CH4 was observed during H2 production by serpentinization.\(^{13}\) This suggests that at such low-T and moderately alkaline conditions (pH ~8), the inorganic carbon (CO2, CO, or bicarbonate) tends to equilibrate with organic acids instead of CH4 in a metastable assemblage as previously described for higher T (150–350°C).\(^{14,15,101}\)

Hence, the high H2/CH4 ratio observed in most experiments (e.g. run at ~300°C, 30–50 MPa) highlights the kinetic inhibition of CH4 formation from the reduction of dissolved CO2 species or formate under aqueous hydrothermal conditions.\(^{14,15}\) For this reason, the role of realistic catalysts (magnetite, hematite, chromite, Fe–Ni alloys, or sulfides) has been tested to circumvent this limitation. Most of them allowed very limited conversion of inorganic carbon to CH4 (<1% max) and even less for longer-chain alkanes (<0.1% max) under laboratory timescales (several months), with a final CO2/CH4 ratio ranging from 100\(^{116}\) to 1000.\(^{14,15,117-119}\) Lower values of CO2/CH4, as small as ~0.1–1.0 and 10, were obtained in studies using Fe–Ni alloys (porous awaruite)\(^{104}\) or Ni-sulfides\(^{119}\) and Co-bearing magnetite,\(^{105}\) respectively, pointing to some very specific catalysts possibly available in serpentinized peridotites. Conversely to CO2, hematite, magnetite, and Fe–Ni alloys did not affect the stability of formate and formic acid, at least for T of 170–260°C.\(^{15}\)

The presence of a gaseous phase in the system, which creates conditions closer to the industrial Fisher–Tropsch process, also seems to favor a higher yield of CH4 and the
formation of more complex hydrocarbons, even in the absence of a catalyst (if olivine and gold reactor walls are excluded). At very-low-T conditions, Ru-bearing chromitite exhibits efficient catalytic properties for CH4 production under gaseous conditions. The slow kinetics of the first step (CO2 to CO reduction) in a gas phase is already well known in the industry, and chemists have strived to improve catalyst efficiency, notably by increasing the local concentration of CO2 on the catalyst surface by altering the catalyst’s nature, shape, and size.

Overall, these experiments seldom report on the potential presence of heavier organic compounds or on the organic content of the liquid and solid phases, if any are present (see Section 15.3.1), leaving the possibility for other type of products and reaction mechanisms to account for low CH4 contents. Mineral reactants also need to be systematically introduced in future works for a more realistic approach that fits better with the conditions of natural systems (Section 15.2).

15.3.3 Experimental Occurrences of Carbonaceous Material

Very few experiments have investigated CM phases and their relationship with minerals using bulk or in situ methods such as SEM or transmission electron microscopy (TEM) with or without XPS, Raman, and FTIR spectroscopy. As for the characterization of natural occurrences, these methods are seldom used together, limiting comparisons, and the exact nature of CM is usually unidentified.

As described in Section 15.2.2, in the oceanic lithosphere, CM can be first deposited at high T (>400°C) on mineral surfaces and cracks during magmatic degassing at depth and subsequent migration and cooling of mantle-derived fluids. Experiments designed to reproduce this process simulated the sudden cooling of a C–O–H magmatic gas over freshly cracked olivines. Evidence for direct carbon precipitation on newly formed cracks has been reported despite thermodynamically unfavorable experimental conditions for graphite precipitation. In this study, it was proposed that freshly cracked surfaces of olivine, and possibly of other silicates, offer chemically active areas facilitating the heterogeneous nucleation of CM during interaction with C–O–H gases, at least for initial T between 400°C and 800°C. The deposited carbonaceous film displayed a more complex structure than graphite. Bonding was dominated by C–C and C–H species under the more oxidizing conditions tested, whereas the more reducing ones showed similar amounts of C–C, C–H, C–O, and metal–C species such as carbides (SiC or MgC). Carbides, mainly observed at 400°C, could have acted as reaction intermediates for FTT reactions, but this has not been proven since the analyses of companion gases could not be performed in this study.

The thermal decomposition of siderite under water vapor conditions (300°C) and saturated vapor pressure (Psal) also resulted in the formation of CM. Siderite alone provides both H2 from water reduction by Fe2+ oxidation and inorganic carbon transformed into a reduced carbon phase (15.2):

\[
6\text{FeCO}_3 \rightarrow 2\text{Fe}_3\text{O}_4 + 5\text{CO}_2 + \text{C}.
\]
Although siderite may not be abundant in the oceanic lithosphere, other carbonates are present and may locally decompose (e.g. during magmatic injections) and react if H₂ is available. After solvent extraction, the organic products were identified as dominantly alkylated and hydroxylated aromatic compounds, which considerably differs from the FTT products (aliphatic chains) that would have been catalyzed by the abundantly formed magnetite. In addition, discrepancies between the relative H/C ratios of reactants and products suggested the presence of an unidentified product with a low H/C ratio, which could be an insoluble C-rich phase remaining in the solid phase.

Under aqueous conditions more representative of low-T hydrothermal processes, two types of experiments report the formation of a poorly crystallized CM: (1) alteration of ferromagnesian silicates by a CO₂-enriched fluid; and (2) carbonate dissolution experiments.

Experiments involving the carbonation of olivine (400–500°C, 100 MPa, static capsules) and of a sandstone made of Fe²⁺-rich volcanic clasts (100°C, 10 MPa, flow-through reactor) reported the precipitation of a poorly crystallized graphitic phase (Figure 15.5a and b). In both experiments, the graphitic phase accounts for a non-negligible part of the reaction products in addition to phyllosilicates and carbonates. As in natural systems (Section 15.2.2), it is embedded in phyllosilicates, corresponding to serpentine (Figure 15.5a) or Fe-rich chlorite (chamosite; Figure 15.5b) here, closely associated with magnetite when present. CM formation was attributed to the reduction of CO₂ (or CO at high P–T) by H₂ initially present in solution (Eqs. (15.3) and (15.4)) or by Fe²⁺-bearing minerals.

\[
\text{CO}_2 + 2\text{H}_2 = \text{C} + 2\text{H}_2\text{O}. \tag{15.3}
\]

\[
\text{CO} + \text{H}_2 = \text{C} + \text{H}_2\text{O}. \tag{15.4}
\]

The two protocols described above resulted in different oxygen fugacities (fO₂). It was estimated to be close to the CCO buffer (C as graphite–CO) in the high-P–T runs, which prevented the formation of magnetite, in opposition to the low-P–T runs. This indicates that magnetite is not mandatory for the formation of graphitic material, whose nature remains poorly constrained, however. Organic volatiles have not been investigated in these studies.

Siderite dissolution experiments at 200°C and 300°C and 50 MPa described two types of carbonaceous products (Figure 15.5c and d). A poorly structured hydrated or hydrogenated CM occurred without spatial relationship with minerals in all runs (Figure 15.5c). At 200°C, a more ordered graphitic carbon formed on the surface of the neo-formed magnetite grains or near iron oxides at the siderite surface (Figure 15.5d). CO₂ was the only gas detected at 200°C, while additional small amounts of H₂ and CH₄ were detected at 300°C. A blank experiment showed that trace amounts (<mM) of dissolved organic compounds were present as contaminants. Although they were much less...
abundant than CO₂, there remain open questions regarding whether they contributed to the formation of low levels of CH₄ and the presence of carbonaceous phases. Whatever the case may be, the results of this work fit well with kinetic inhibition of CH₄ formation from CO₂ and H₂ at low T (<400°C; Section 15.3.2) and suggest a preferential precipitation of CM with decreasing T through reactions (15.3) and (15.5).

$$3\text{FeCO}_3 + \text{H}_2\text{O}_{(aq)} = \text{Fe}_3\text{O}_4 + 3\text{CO}_2_{(aq)} + \text{H}_2_{(aq)}.$$  \hspace{1cm} (15.5)
C + 2H₂ = CH₄.  \hspace{1cm} (15.6)

The inefficiency of magnetite as a FTT reaction catalyst is also pointed out here (see Section 15.3.2). Poisoning of the magnetite surface by carbon coatings (Figure 15.5d) at low T has been questioned; alternatively, the carbon coating may only form when CH₄ cannot (i.e. at low T). A syngenetic relationship between CM and CH₄ has also been proposed but not demonstrated. Equations (15.3) and (15.6) on the one hand and (15.7) and (15.8) on the other show two reversal genetic links. The second one (Eqs. (15.7) and (15.8)) appears less likely under the experimental conditions tested (≤300°C) according, again, to the kinetic inhibition hampering CH₄ formation (15.7).

CO₂ + 4H₂ = CH₄ + 2H₂O.  \hspace{1cm} (15.7)

CO₂ + CH₄ = 2C + 2H₂O.  \hspace{1cm} (15.8)

Equation (15.6), which suggests that carbon coating may serve as an intermediate step to CH₄ formation at least at 300°C where its absence would result from its full consumption, may be a more relevant hypothesis, as already mentioned in Section 15.2.3. Similarly, it was previously suggested that background carbon, possibly more reduced than graphite on the magnetite surface, facilitates CH₄ formation. \(^{118}\)

At lower T compatible with life, experimental data are lacking on the formation of CM, while this is the condition under which it is the most difficult to discriminate the origin of CM in natural systems (Section 15.2). In serpentinization experiments run at ≤100°C (see Section 15.3.2), no CH₄ was produced, and the strong decrease of CO₂ and H₂ after 3 months was not fully explained. \(^{13}\)

The limited data available to describe the association of or interplay between organic volatiles and CM limit the conclusions so far on their genetic relationship.

If all occurrences of CM in experimental solid products are due to in situ abiotic reactions from an inorganic source (not contamination), they suggest that at least small amounts of CM easily form under a wide range of hydrothermal conditions during short runs of a few days to a few months (100–500°C, 10–110 MPa, /O₂ between COO and QFM buffers). The rare available images report different aspects (spherules, film-like surficial layers, or porous accumulations) similar to some of those observed in natural systems (Figure 15.2). They seem to condense from gas or precipitate from fluid preferentially on the mineral surfaces where they possibly evolve. Minerals used in experiments (mainly magnetite) were not as diverse as in nature (Section 15.2), and other mineral surfaces such as hematite, sulfides, hydrogarnets, phyllosilicates, epidote, and quartz, not yet introduced in experiments, may allow the formation of carbonaceous compounds, possibly when the reducing conditions are not optimal. \(^{38,55,60,85,123}\)

In agreement with natural observations (Section 15.2.2), one experimental study has demonstrated that smectite clays, and particularly montmorillonites, can promote and preserve organic compounds formed under seafloor hydrothermal conditions (300°C, 100 MPa). \(^{103}\) While \(^{13}\)CH₄ was the dominant organic product in the gas-phase, aromatic
compounds, including PAHs (up to C_{20}), were extracted from the solid products that reacted for 6 weeks with labeled $^{13}$C-methanol (10.4 M). The absence of organic material within the illite sample compared to the smectite one showed that the clay interlayer properties exert a control on organic synthesis, notably the negatively charged layer of smectite. The exact reaction path was not constrained; the role of alkanes as intermediate reactants or products and the presence of methanol as a major reactant are worth noting. This study also highlighted the possible protection of organic compounds by clays from thermal alteration at 300°C, making them available for further reactions under lower-T conditions.

15.3.4 Thermodynamic Predictions

The nature of hydrothermal CM observed in natural and experimental works related to the alteration of mantle-derived rocks is not clarified yet. It ranges from graphite to very different forms in terms of structural order, nature and quantity of heteroatoms, unsaturations, and hence reactivity. There has been a considerable amount of work to quantify the thermodynamics of C–C bond formation in graphite and other phases.$^{32}$ We selected here the main results related to CM formation under hydrothermal conditions for comparison with previous sections.

The formation of graphite is predicted from C–O–H fluids under a wide range of hydrothermal conditions, provided there is no kinetic inhibition. Graphite can readily precipitate from the fluid; this is favored by a decrease in temperature or an increase in pressure along with a decrease in water content.$^{36,45,128,129}$ Both T decrease (deep fluid ascent) and water decrease (e.g. during hydration reactions) correspond well to natural oceanic scenarios in the deeper part of the lithosphere. Graphite formation results from redox reactions unless the reactant carbon is at the same oxidation state (i.e. zero). This does not necessarily require very strong reducing conditions, and graphite can be produced through either oxidation or reduction reactions (Figure 15.4).$^{44,130}$ In addition to graphite, CO$_2$ or CH$_4$ should be the dominant carbon species in hydrothermal fluids depending on temperature and oxygen fugacity. For a given oxygen fugacity, CO$_2$ should predominate at high-T conditions, while CH$_4$ is favored at the lower T$^{35,131}$ according to the equilibrium described in (15.7).

When methane and graphite formation is inhibited (i.e. when the kinetic barriers hamper access to a stable equilibrium,$^{35}$ as in most natural and experimental systems; Sections 15.2, 15.3.2, and 15.3.3), there is considerable potential for the hydrothermal synthesis of a wide range of metastable organic molecules provided that $f$H$_2$ is high enough.$^{33,44,45}$ The reversible character of several organic reactions in hydrothermal fluids contributes to this variety (e.g. hydration/dehydration or oxidation/reduction reactions).$^{132}$ Calculations have shown that organic synthesis of metastable phases is even possible in environments where the amount of H$_2$ is usually below those observed during the alteration of mantle-dominated environments;$^{44,45}$ as an example, the rapid cooling of volcanic gases
containing CO, CO₂, H₂O, and H₂ below ~250°C provides a thermodynamic drive for the abiotic synthesis of metastable hydrocarbons such as condensed n-alkane and PAHs, depending on the H/C ratio of the gas. Such products may relate to the complex CM and molecular organic minerals observed in natural samples and in CH₄-poor experiments. In more reducing environments, the energetic drive of metastable compound formation is increased and reaction temperatures can be shifted to higher values. Decreasing temperature allows for metastable mixtures of different condensed phases, in agreement with the variety of compounds found in natural rock textures (Section 15.2.2).

The potential abiotic formation of CM during serpentinization that is central to this chapter has been investigated by comparing experimental fluid compositions with natural hydrothermal compositions across a wide range of CO₂ concentrations. Predicted mineral assemblages consisted of serpentine and brucite with or without magnetite and carbonates depending on the relative activities of H₂ and CO₂. CO₂(aq) was considered at equilibrium with graphitic compounds that share the thermodynamic properties of graphite and a hydrogenated aromatic carbon compound (i.e. anthracene, a PAH made of three benzene rings) in order to mimic the experimental products (Figure 15.5c and d). Results show that serpentinization fluids can equilibrate with both the graphitic and the hydrogenated carbon when the formation of alkanes is prohibited (kinetic inhibition). As the precipitation of a hydrogenated carbon compound requires a higher hydrogen activity, its formation predicted after the serpentinization reaction significantly progresses (i.e. >70% complete), similarly to the classical serpentinization degree attained in oceanic peridotites (e.g. Andreani et al.).

15.4 Summary

According to natural, experimental, and theoretical results, the accumulation of CM within mantle-derived rocks of the oceanic lithosphere seems a very likely process, yet it is poorly considered during hydrothermalism. Even if CM formation under such conditions has been largely unexplored experimentally compared to volatile products, this outcome fits well with recent models that restrict CH₄ formation to high-T processes (>400°C) at depth, leaving open the possibility for various metastable carbon phases to form at shallower levels. Although the total amount of abiotic CM is not quantified yet, and might represent a small fraction of the TOC contained in the present-day oceanic lithosphere, a better understanding of its formation mechanisms, nature, and reactivity is of prime importance.

The large range of aspects, structural orders, and compositions of the CM trapped within rock textures, ranging from stable graphite usually at high T (>~500°C) to variably ordered, hydrogenated, O-, N- and S-bearing materials and PAHs, attest to the lithosphere’s organic wealth available for evolved chemical reactions and ecosystem development. The structure and functions of ecosystems inhabiting the shallow levels of the oceanic lithosphere indeed suggest that despite the general thought that CH₄ or inorganic carbon species represent the main sustainable feedstocks of carbon in those settings, they...
may not be the main sources of carbon for microorganisms that rather use formate or more diverse abiotic organic compounds, including PAHs. The variety of organic compounds reflects the characteristics of natural systems, which are dynamic and highly heterogeneous chemically, structurally, and hydrodynamically, therefore creating chemical and redox gradients that are able to drive metastable reactions over a wide range of physicochemical conditions. They also provide various mineral substrates that have been shown to facilitate CM accumulations and likely further transformation. Whether minerals can affect the nature and crystallinity of the CM remains to be addressed. According to the very different crystallo-chemistry and surface properties of mineral families, different types of organic compounds may even be expected within different mineral substrates, rendering them as micro-factories with high specificity as observed in natural serpentinites.

Observations force us to consider natural rocks as being built up of several far-from-equilibrium microenvironments that are expected to evolve with time and system fluctuations (H2, P, T, pH, and fluid transport and chemistry), each offering at a given stage a specific mineral assemblage and chemical conditions that are propitious to a given series of reactions. They considerably deviate from static, buffered, bulk mineral assemblages undergoing interactions with fluids near equilibrium, as is often considered. Such micro-environments may be the locations of organic reactions simply succeeding each other or competing together, but also inorganic reactions such as carbonate precipitation and biological activity, depending on the alteration history and fluid paths. Indeed, the oceanic crust is subjected to a progressive and multistage evolution of its organic pool, including both abiotic reactions and multiple recolonizations, all along the way from ridge to subduction, hence modifying the initial signature of a compound of interest.

We attempted to summarize the main stages of abiotic carbon processing in a heterogeneous oceanic lithosphere (Figure 15.6a) from deep to shallow levels in the sketch of Figure 15.6. We recommend going beyond the formation of CH4 and the FTT reaction by considering the heterogeneity and multistage character of the system at all scales. First, CM (or graphite) can directly precipitate on fresh mineral surfaces and in vesicles during cooling and re-speciation of CO2-rich magmatic fluids at depth (>400°C and up to 800°C) initially in the fluid or formed in situ and CM (or graphite) formed in situ (see Section 15.2.3), possibly helped by subsequent retrograde reactions (stage 1, Figure 15.6b). Then, provided relatively low fO2 is available, CM can continue to form at ≤400°C (stages 2 and 3, Figure 15.6b) among the new mineral assemblage from aqueous C-bearing hydrothermal fluids (magmatic or seawater derived) thought to carry a large variety of organic and inorganic carbon species formed at deeper lithospheric levels (see Sections 15.2.2 and 15.3.2). The strongest thermodynamic drive for the formation of most metastable organic molecules (see Section 15.3.4), including n-alkanes, organic acids, and PAHs, is below 200°C (stage 3, Figure 15.6b) for typical redox conditions occurring in the mafic component of the lithosphere (near the PPM buffer). In the dominant ultramafic component, conditions should be far more favorable to the abiotic formation of metastable products, and extending to higher temperatures (at least 300°C at the QFM buffer). Indeed,
Figure 15.6 Sketch depicting the main stages of carbon processing in the oceanic lithosphere during the multistage fluid–rock reactions\textsuperscript{133} recorded in such dynamic systems. (a) Geologic setting of a slow-spreading ridge where mantle is tectonically exhumed, simultaneously with localized and ephemeral magmatic injections. These environments appear to be the most favorable ones for abiotic organic synthesis according to the available natural observations, experiments, and thermodynamic calculations. (b) Summary of the three main stages of carbon processing in a column of the oceanic lithosphere shown in (a), illustrated through a magnification of the mantle rock textures along with the hypothetical nature of the percolating fluids. See the text for details. alk = alkanes.
serpentinization provides high levels of H₂ over a wide range of T and pH conditions (100°C, pH 9 to 350°C, pH 3)\(^{13,102}\) that favors reduction reactions while providing variable single-carbon reactants in the fluid, especially at low T < ~200°C (ΣCO₂, ΣHCOOH, methanol).\(^{101}\) A body of experimental work has shown that the most efficient production of H₂ from serpentinization occurs at ~300°C, notably with modified seawater after interaction with magmatic lithologies.\(^{138,139}\) Alteration of ultramafic rocks also releases specific transition metals such as Fe and Ni and is known to change sulfur speciation, leading to the formation of potential metallic–sulfide catalysts.\(^{140,141}\) The fH₂ of those systems can be highly variable down to the micrometer scale, ranging from QFM buffer conditions in magnetite-dominated domains to fH₂ values of several orders of magnitude higher to form Fe–Ni alloys such as awaruite near serpentinization fronts.\(^{141}\) Serpentinization is also characterized by the abundant formation of phyllosilicates, oxides, and hydroxides that are thought to facilitate CM precipitation, condensation, and transformation (stages 2 and 3, Figure 15.6b). The ephemeral magmatic activity associated with mantle-rock exhumation in a slow-spreading environment (Figure 15.6) may also drive fluctuating T or pH conditions and C–O–H fluid inputs. This can shift the local thermal regime from a cold-alkaline to a hot-acidic environment, and vice versa, as observed at the Rainbow massif.\(^{142}\) The tectono-magmatic activity near the ridge axis also affects transport processes that control the fluid residence time in the system, a determinant of C–O–H fluid speciation, especially at low T.\(^{101}\)

15.5 Limits to Knowledge and Unknowns

To achieve a better understanding of the mechanisms and processes leading to the abiotic formation of CM in the oceanic lithosphere and to carbon cycling under those environments in general, one of the major challenges future studies will have to face is to improve criteria for establishing the origin of organic carbon accumulations found within the habitable oceanic rocks. As highlighted in this chapter, graphitic compounds occur from deep levels where solely abiotic reactions can account for their generation, up to colonizable depths, where H₂ and abiotic organic compounds are valuable sources of energy and carbon for microbial communities\(^{27,143}\) below the T limit of life currently established at 122°C.\(^{144}\) This deep life indeed has the ability to capitalize on the steady stream of both inorganic and organic serpentinization by-products, converting C into biomass and competing against abiotic reactions of C reduction.\(^{23}\) These rock-hosted microbial ecosystems, along with the photosynthesis-derived organic compounds contained in seawater and injected into the hydrothermal conduits, experience hydrothermal degradation and can be transformed, similarly to sedimentary organic matter, into a kerogen-like or bitumen-like macromolecular material\(^{18,19,31}\) with structural and chemical similarities with the hydrothermally formed CM. Up to now, more attention has been paid to CM in meteorites\(^{145}\) than in hard rock samples from Earth’s lithosphere. Meteorites testify to the capacity of hydrothermal activity to drive complex abiotic organic synthesis.
Identifying CM origins will have to go through more systematic CM characterization in rocks from various geological settings. This may also contribute to the identification of new organic minerals (see mineralchallenge.net).

The role (in terms of kinetics and selectivity) of mineral phases in organic synthesis, particularly in CM formation, evolution, and preservation, should be a prime direction for future research, too. New experimental work is required in which minerals should be introduced and carefully characterized to possibly identify CM, which has been neglected so far, along with its reactivity and formation mechanisms. We encourage a more systematic investigation of the liquids, gases, and solids altogether in future hydrothermal experiments in order to unravel possible genetic relationships and to determine their respective roles in the organic synthesis factory.

Tracking reactions in situ in both natural and experimental solids requires maintaining the ongoing effort to develop microscale to nanoscale investigation methods and to identify biotic/abiotic criteria for use both in natural samples and in experiments that will have to face contamination issues. As detailed in Section 15.2, most of the evidence for CM abiogenicity was derived from the mineral assemblages and textural relationships between CM and mineral phases. Compared to bulk analysis, microscale characterization using high-resolution imaging techniques will allow a comprehensive description of the textural context in which organic carbon occurs and will provide constraints regarding whether or not the carbon compounds are indigenous. Microscale techniques can also capture the co-occurrence and relationships of CM with metals and minerals, and are thus particularly well suited to tracking organic matter production, evolution, and transport within the oceanic lithosphere.

Acknowledgments

The authors want to thank the Deep Carbon Observatory, the French CNRS (INSU, PIE, EPOV, and MITI Défì Origines programs), and the French National Research Agency (ANR-14-CE01-0008) for funding. This chapter is also the result of numerous fruitful discussions with several partners, notably the participants of the deepOASES project (PI B. Ménez).

Questions for the Classroom

1. Why is it important to better constrain the abiotic organic reactions on Earth?
2. Which impact (if any) could abiotic organic synthesis have on Earth today and have had in the past?
3. What is the main message of this chapter and what do you think of it?
4. Why is the oceanic lithosphere a propitious place to investigate abiotic organic synthesis?
5. Which other settings on Earth would be of interest, as well as elsewhere in the solar system? What would be the implications?
6 Which organic compounds are thermodynamically stable? How and why can metastable organic compounds be formed and subsist through time in rocks?

7 What are the main abiotic organic compounds formed through the hydrothermal alteration of the oceanic lithosphere? How do they vary as a function of temperature? What are the other important parameters?

8 What are the strengths of micro-imaging and *in situ* micro-spectroscopic approaches for unraveling the reaction paths leading to abiotic organic synthesis in rocks?

9 Can you think of potential markers for discriminating between the abiotic and biologic origins of carbonaceous compounds in rocks? Can they be addressed with available techniques of analyses or do they require technological development?

**References**


Abiotic Organic Synthesis and Processing


