Chapter 9: Stable Isotope Geochemistry

9.1 Introduction

Stable isotope geochemistry is concerned with variations of the isotopic compositions of elements arising from physicochemical processes rather than nuclear processes. Fractionation of the isotopes on an element might at first seem to be an oxymoron. After all, in the last chapter we saw that some of the value of radiogenic isotopes was that the various isotopes of an element had identical chemical properties and therefore that isotope ratios such as $^{87}\text{Sr}/^{86}\text{Sr}$ are not changed measurably by chemical processes. In this chapter we will find that this is not quite true, and that the very small differences in the chemical behavior of different isotopes of an element can provide a very large amount of useful information about chemical (both geochemical and biochemical) processes.

The origins of stable isotope geochemistry are closely tied to the development of modern physics in the first half of the 20th century. The discovery of the neutron in 1932 by H. Urey and the demonstration of variations in the isotopic composition of light elements by A. Nier in the 1930’s and 1940’s were the precursors to the development of stable isotope geochemistry. The real history of stable isotope geochemistry begins in 1947 with the Harold Urey’s publication of a paper entitled “The Thermodynamic Properties of Isotopic Substances”. Urey not only showed why, on theoretical grounds, isotope fractionations could be expected, but also suggested that these fractionations could provide useful geological information. Urey then set up a laboratory to determine the isotopic compositions of natural substances and experimentally determine the temperature dependence of these fractionations, in the process establishing the field of stable isotope geochemistry.

What has been learned in the fifty years since that paper was published would undoubtedly astonish even Urey. Stable isotope geochemistry, like radiogenic isotope geochemistry, has become an essential part of not only geochemistry, but the earth sciences as a whole. In this chapter, we will attempt to gain an understanding of the principles underlying stable isotope geochemistry and then briefly survey its many-fold applications in the earth sciences. In doing so, we add the final tool to our geochemical toolbox.

9.1.1 Scope of Stable Isotope Geochemistry

The elements interest in stable isotope geochemistry are H, Li, B, C, N, O, Si, S, and Cl. Of these, O, H, C and S are of the greatest interest. Most of these elements have several common characteristics:

1. They have low atomic mass.
2. The relative mass difference between their isotopes is large.
3. They form bonds with a high degree of covalent character.
4. The elements exist in more than one oxidation state (C, N, and S), form a wide variety of compounds (O), or are important constituents of naturally occurring solids and fluids.
5. The abundance of the rare isotope is sufficiently high (generally at least tenths of a percent) to facilitate analysis.

Geologically useful information has not been extracted (with some exceptions) from elements not meeting these criteria. For example, $^{48}\text{Ca}/^{40}\text{Ca}$ has a large relative mass difference, but calcium tends to form ionic bonds and occurs in a limited variety of lattice environments. Thus little fractionation has been observed. Mg is a light element, but in addition to its bonds being dominantly ionic, it is generally situated in same atomic environment (octahedrally coordinated by O). The elements of interest in radiogenic isotope geochemistry are heavy (Sr, Nd, Hf, Os, Pb), form dominantly ionic bonds, generally exist in only one oxidation state, and there is only small relative mass differences between the isotopes of interest. Thus fractionation of the isotopes of these elements are quite small and can generally be ignored. Furthermore, any natural fractionation is

*Fractionation refers to the change in an isotope ratio that arises as a result of some chemical or physical process.
corrected for in the process of correcting much larger fractionations that typically occur during analysis (with the exception of Pb). Thus is it that one group of isotope geochemists make their living by measuring isotope fractionations while the other group makes their living by ignoring them!

Stable isotope geochemistry has been applied to a great variety of problems, and we will see a number of examples in this chapter. One of the most common is geothermometry. Another is process identification. For instance, plants that produce 'C 4' hydrocarbon chains (that is, hydrocarbon chains 4 carbons long) as their primary photosynthetic product fractionate carbon differently than plants that produce 'C 3' chains. This fractionation is retained up the food chain. This allows us, for example, to draw some inferences about the diet of fossil mammals from the stable isotope ratios in their bones. Sometimes stable isotope ratios are used as 'tracers' much as radiogenic isotopes are. So, for example, we can use oxygen isotope ratios in igneous rocks to determine whether they have assimilated crustal material, as crust generally has different O isotope ratios than does the mantle.

9.1.2 Notation

9.1.2.1 The δ Notation

As we shall see, variations in stable isotope ratios are typically in the parts per thousand to parts per hundred range and are most conveniently and commonly reported as permil deviations, δ, from some standard.

Unfortunately, a dual standard has developed for reporting O isotopes. Geochemists working with substances other than carbonates reported O isotope ratios as permil deviations from SMOW (standard mean ocean water):

\[
\delta^{18}O = \left( \frac{(^{18}O/^{16}O)_{sample} - (^{18}O/^{16}O)_{SMOW}}{(^{18}O/^{16}O)_{SMOW}} \right) \times 10^3
\]

Those working with carbonates report \( \delta^{18}O \) relative to the Pee Dee Belemite (PDB) carbonate standard‡. This value is related to SMOW by:

\[
\delta^{18}O_{PDB} = 1.03086 \delta^{18}O_{SMOW} + 30.86
\]

A similar δ notation is used to report other stable isotope ratios. Hydrogen isotope ratios, δD, are reported relative to SMOW, carbon isotope ratios relative to PDB, nitrogen isotope ratios relative to atmospheric nitrogen (denoted ATM), and sulfur isotope ratios relative to troilite in the Canyon Diablo iron meteorite (denoted CDT). Table 9.1 lists the isotopic composition of these standards.

9.1.2.2 The Fractionation Factor

The fractionation factor, \( \alpha \), is the ratio of isotope ratios in two phases:

\[
\alpha_{A-B} \equiv \frac{R_A}{R_B}
\]

‡ There is, however, a good historical reason for this: analytical difficulties in the measurement of carbonate oxygen prior to 1965 required it be measured against a carbonate standard.
The fractionation of isotopes between two phases is also often reported as $\Delta_{A-B} = \delta_A - \delta_B$. The relationship between $\Delta$ and $\alpha$ is:

$$\Delta \approx (\alpha - 1)10^3$$

or

$$\Delta \approx 10^3 \ln \alpha$$

As we will see, at equilibrium, $\alpha$ may be related to the equilibrium constant of thermodynamics by

$$\alpha_{A-B} = (K/K_\infty)^{1/n}$$

where $n$ is the number of atoms exchanged, $K_\infty$ is the equilibrium constant at infinite temperature, and $K$ is the equilibrium constant written in the usual way (except that concentrations are used rather than activities because the ratios of the activity coefficients are equal to 1, i.e., there are no isotopic effects on the activity coefficient).

### 9.2 Theoretical Considerations

Isotope fractionation can originate from both kinetic effects and equilibrium effects. The former may be intuitively expected, but the latter may at first seem somewhat surprising. After all, we have been taught that the chemical properties of an element are dictated by its electronic structure, and that the nucleus plays no real role in chemical interactions. In the following sections, we will see that quantum mechanics predicts that the mass of an atom affects its vibrational motion therefore the strength of chemical bonds. It also affects rotational and translational motions. From an understanding of these effects of atomic mass, it is possible to predict the small differences in the chemical properties of isotopes quite accurately.

#### 9.2.1 Equilibrium Isotope Fractionations

Most isotope fractionations arise from equilibrium effects. Equilibrium fractionations arise from translational, rotational and vibrational motions of molecules in gases and liquids and atoms in crystals because the energies associated with these motions are mass dependent. Systems tend to adjust themselves so as to minimize energy. Thus isotopes will be distributed so as to minimize the vibrational, rotational, and translational energy of a system. Of the three types of energies, vibrational energy makes by far the most important contribution to isotopic fractionation. Vibrational motion is the only mode of motion available to atoms in a solid. These effects are, as one might expect, small. For example, the equilibrium constant for the reaction

$$\frac{1}{2} C^{16}O_2 + H_2^{18}O \rightleftharpoons \frac{1}{2} C^{18}O_2 + H_2^{16}O$$

is only about 1.04 at 25°C and the $\Delta G$ of the reaction, given by $-RT\ln K$, is only -100 J/mol (you’ll recall most $\Delta G$’s for reactions are in typically in the kJ/mol range).

#### 9.2.1.1 The Quantum Mechanical Origin of Isotopic Fractionations

It is fairly easy to understand, at a qualitative level at least, how some isotope fractionations can arise from vibrational motion. Consider the two hydrogen atoms of a hydrogen molecule: they do not remain at a fixed distance from one and other, rather they continually oscillate toward and away from each other, even at absolute zero. The frequency of this oscillation is quantized, that is, only discrete frequency values are possible. Figure 9.1 is a schematic diagram of energy as a...
function of interatomic distance in the hydrogen molecule. As the atoms vibrate back and forth, their potential energy varies as shown by the curved line. The zero point energy (ZPE) is the energy level at which the molecule will vibrate in its ground state, which is the state in which the molecule will be in at low temperature. The zero point energy is always some finite amount above the minimum potential energy of an analogous harmonic oscillator.

The potential energy curves for various isotopic combinations of an element are identical, but as the figure shows, the zero point vibrational energies differ, as do vibration energies at higher quantum levels, being lower for the heavier isotopes. Because of the difference in vibrational energies, the amount of energy that must be gained by the molecule to dissociate, i.e., the bond energy, will differ for different isotopic combinations. For example, 440.6 kJ/mole is necessary to dissociate a D₂ (²H₂) molecule, but only 431.8 kJ/mole is required to dissociate the ¹H₂ molecule. Thus the bond formed by the two deuterium atoms is 9.8 kJ/mole stronger than the H–H bond. The differences in bond strength can also lead to kinetic fractionations, since molecules that dissociate easier will react faster. We will discuss kinetic fractionations a bit later.

9.2.1.2 Predicting Isotopic Fractionations from Statistical Mechanics

Now let’s attempt to understand the origin of isotopic fractionations on a more quantitative level. We have already been introduced to several forms of the Boltzmann distribution law (e.g., equation 2.104), which describes the distribution of energy states. It states that the probability of a molecule having internal energy $E_i$ is:

$$P_i = \frac{g_i e^{-E_i/kT}}{\sum g_j e^{-E_j/kT}}$$

where $g$ is a statistical weight factor, $n_0$ is the number of molecules with ground-state or zero point energy, $n_i$ is the number of molecules with energy $E_i$, and $k$ is Boltzmann’s constant. The average energy (per molecule) is:

$$\bar{E} = \frac{\sum E_i g_i e^{-E_i/kT}}{\sum g_i e^{-E_i/kT}}$$

This factor comes into play where more than one state corresponds to an energy level $E_i$ (the states are said to be ‘degenerate’). In that case $g_i$ is equal to the number of states having energy level $E_i$. 
As we saw in Chapter 2, the partition function, $Q$, is the denominator of this equation:

$$Q = \sum g_i e^{-E_i/kT}$$  \hspace{1cm} (9.9)

The partition function is related to thermodynamic quantities $U$ and $S$ (equations 2.119 and 2.123). Since there is no volume change associated with isotope exchange reactions, we can use the relationship:

$$\Delta G = \Delta U - T\Delta S$$  \hspace{1cm} (9.10)

and equations 2.119 and 2.123 to derive:

$$\Delta G_i = -R \ln \left( \prod Q_i^v \right)$$  \hspace{1cm} (9.11)

and comparing to equation 3.98, that the equilibrium constant is related to the partition function as:

$$K = \prod Q_i^v$$  \hspace{1cm} (9.12)

for isotope exchange reactions. In the exchange reaction above (9.6) this is simply:

$$K = \frac{Q_{\text{CO}_2}^{18} Q_{\text{H}_2\text{O}}^{16}}{Q_{\text{CO}_2}^{16} Q_{\text{H}_2\text{O}}^{18}}$$  \hspace{1cm} (9.13)

The usefulness of the partition function is that it can be calculated from quantum mechanics, and from it we can calculate equilibrium fractionations of isotopes.

There are three modes of motion available to gaseous molecules: vibrational, rotational, and translational (Figure 9.2). The partition function can be written as the product of the translational, rotational, vibrational, and electronic partition functions:

$$Q_{\text{total}} = Q_{\text{vib}} Q_{\text{trans}} Q_{\text{rot}} Q_{\text{elec}}$$  \hspace{1cm} (9.14)

The electronic configurations and energies of atoms are unaffected by the isotopic differences, so the last term can be ignored in the present context. The vibrational motion is the most important contributor to isotopic fractionations, and it is the only mode of motion available to atoms in solids. So we begin by calculating the vibrational partition function. We can approximate the vibration of atoms in a simple diatomic molecule such as CO or O$_2$ by that of a harmonic oscillator. The energy of a ‘quantum’ oscillator is:

$$E_{\text{vib}} = (n + \frac{1}{2})h\nu$$  \hspace{1cm} (9.15)

where $\nu$ is the ground state vibrational frequency, $h$ is Plank’s constant and $n$ is the vibrational quantum number. The partition function for vibrational motion is given by:

$$Q_{\text{vib}} = \frac{e^{-h\nu/2kT}}{1 - e^{-h\nu/kT}}$$  \hspace{1cm} (9.16)$^\dagger$

For an ideal harmonic oscillator, the relation between frequency and reduced mass is:

\[ Q_{\text{vib}} = \prod_{i} \frac{e^{-h\nu_i/2kT}}{1 - e^{-h\nu_i/2kT}} \]  \hspace{1cm} (9.16a)

$^\dagger$ Polyatomic molecules have many modes of vibrational motion available to them. The partition function is calculated by summing the energy over all available modes of motion; since energy occurs in the exponent, the result is a product:

$$Q_{\text{vib}} = \prod_{i} \frac{e^{-h\nu_i/2kT}}{1 - e^{-h\nu_i/2kT}}$$  \hspace{1cm} (9.16a)
\[ v = \frac{1}{2\pi} \sqrt{\frac{k}{\mu}} \quad 9.17 \]

where \( k \) is the forcing constant, which depends on the electronic configuration of the molecule, but not the isotope involved, and \( \mu \) is reduced mass:

\[ \mu = \frac{1}{1/m_1 + 1/m_2} \quad \text{or} \quad \mu = \frac{m_1m_2}{m_1 + m_2} \quad 9.18 \]

Rotational motion of molecules is also quantized. We can approximate a diatomic molecule as a dumbbell rotating about the center of mass. The rotational energy for a quantum rigid rotator is:

\[ E_{\text{rot}} = \frac{j(j + 1)\hbar^2}{8 \pi^2 I} \quad 9.19 \]

where \( j \) is the rotational quantum number and \( I \) is the moment of inertia, \( I = \mu r^2 \), where \( r \) is the interatomic distance. The statistical weight factor, \( g \), in this case is equal to \((2j + 1)\) since there are 2 axes about which rotations are possible. For example, if \( j = 1 \), then there are \( j(j + 1) = 2 \) quanta available to the molecule, and \( (2j + 1) = 3 \) ways of distributing these two quanta: all to the \( x \)-axis, all to the \( y \)-axis or one to each. Thus:

\[ Q_{\text{rot}} = \sum (2j + 1)e^{-j(j+1)\hbar^2/8\pi^2\hbar^2 I T} \quad 9.20 \]

Because the rotational energy spacings are small, equation 9.20 can be integrated to yield:

\[ Q_{\text{rot}} = \frac{8\pi^2 I k T}{\sigma \hbar^2} \quad 9.21^* \]

where \( \sigma \) is a symmetry factor whose value is 1 for heteronuclear molecules such as \( ^{16}\text{O}^{18}\text{O} \) and 2 for homonuclear molecules such as \( ^{16}\text{O}^{16}\text{O} \). This arises because in homonuclear molecules, the quanta must be equally distributed between the rotational axes, i.e., the \( j \)'s must be all even or all odd. This restriction does not apply to heterogeneous molecules, hence the symmetry factor.

Finally, translational energy associated with each of the three possible translational motions (\( x \), \( y \), and \( z \)) is given by the solution to the Schrödinger equation for a particle in a box:

\[ E_{\text{trans}} = \frac{n^2\hbar^2}{8ma^2} \quad 9.22 \]

where \( n \) is the translational quantum number, \( m \) is mass, and \( a \) is the dimension of the box. This expression can be inserted into equation 9.8. Above about 2 K, spacings between translational energies levels are small, so equ. 9.8 can also be integrated. Recalling that there are 3 degrees of freedom, the result is:

\[ Q_{\text{trans}} = \frac{(2\pi mk T)^{3/2}}{\hbar^3} V \quad 9.23 \]

where \( V \) is the volume of the box \((a^3)\). Thus the full partition function is:

\[ Q_{\text{total}} = Q_{\text{vib}} Q_{\text{rot}} Q_{\text{trans}} = \frac{e^{-\hbar v/2kT}}{1 - e^{-\hbar v/kT}} \frac{8\pi^2 I k T}{\sigma \hbar^2} \frac{(2\pi mk T)^{3/2}}{\hbar^3} V \quad 9.24 \]

It is the ratio of partition functions that occurs in the equilibrium constant expression, so that many of the terms in 9.24 eventually cancel. Thus the partition function ratio for 2 different isotopic species of the same diatomic molecule, A and B, reduces to:

\[ Q_{\text{rot}} = \frac{8\pi^2(8\pi^3ABC)^{1/2}(kT)^{3/2}}{\sigma \hbar^3} \]

where A, B, and C are the three principle moments of inertia.

* Equation 9.21 also holds for linear polyatomic molecules such as CO \(_2\). The symmetry factor is 1 if it has plane symmetry, and 2 if it does not. For non-linear polyatomic molecules, 9.21 is replaced by:

\[ Q_{\text{rot}} = \frac{8\pi^2(8\pi^3ABC)^{1/2}(kT)^{3/2}}{\sigma \hbar^3} \]
Since bond lengths are essentially independent of the isotopic composition, this further reduces to:

\[
\frac{Q_A}{Q_B} = \frac{e^{-\hbar \nu_A / 2kT}}{1 - e^{-\hbar \nu_A / kT}} \frac{I_A \sigma_A m_A^{3/2}}{1 - e^{-\hbar \nu_B / kT}} \frac{e^{-\hbar \nu_B / 2kT}}{I_B \sigma_B m_B^{3/2}}
\]

Notice that all the temperature terms cancel except for those in the vibrational contribution. Thus vibrational motion alone is responsible for the temperature dependency of isotopic fractionations.

To calculate the fractionation factor \( \alpha \) from the equilibrium constant, we need to calculate \( K_\infty \).

For a reaction such as:

\[
aA_1 + bB_2 \rightleftharpoons aA_2 + bB_1
\]

where \( A_1 \) and \( A_2 \) are two molecules of the same substance differing only in their isotopic composition, and \( a \) and \( b \) are the stoichiometric coefficients, the equilibrium constant is:

\[
K_\infty = \frac{\sigma_{A_2} \sigma_{A_1}^a}{\sigma_{B_1} \sigma_{B_2}^b}
\]

Thus for a reaction where only a single isotope is exchanged, \( K_\infty \) is simply the ratio of the symmetry factors.

### 9.2.1.3 Temperature Dependence of the Fractionation Factor

As we noted above, the temperature dependence of the fractionation factor depends only on the vibrational contribution. At temperatures where \( T \ll \hbar / k \), the \( 1 - e^{-\hbar \nu / kT} \) terms in 9.16 and 9.26 tend to 1 and can therefore be neglected, so the vibrational partition function becomes:

\[
Q_{vib} = e^{-\hbar \nu / 2kT}
\]

In a further simplification, since \( \Delta \nu \) is small, we can use the approximation \( e^x = 1 + x \) (valid for \( x \ll 1 \)), so that the ratio of vibrational energy partition functions becomes

\[
\frac{Q_{vib}^A}{Q_{vib}^B} \equiv 1 - \hbar \Delta \nu / 2kT
\]

Since the translational and rotational contributions are temperature independent, we expect a relationship of the form:

\[
\alpha \equiv A + \frac{B}{T} \quad 9.29
\]
In other words, $\alpha$ should vary inversely with temperature at low temperature. At higher temperature, the $1 - \exp(-h\nu/kT)$ term differs significantly from 1. Furthermore, at higher vibrational frequencies, the harmonic oscillator approximation breaks down (as suggested in Figure 9.1), as do several of the other simplifying assumptions we have made, so that the relation

\[
\alpha \propto \frac{Q_{C^{18}O}Q_{O^{18}O}}{Q_{C^{16}O}Q_{O^{16}O}}
\]

Example 9.1. Predicting Isotopic Fractionations

Consider the exchange of $^{18}$O and $^{16}$O between carbon monoxide and oxygen:

\[
C^{16}O + ^{16}O^{18}O \leftrightarrow C^{18}O + ^{16}O_{2}
\]

The frequency for the C-$^{16}$O vibration is $6.505 \times 10^{13}$ sec$^{-1}$, and the frequency of the $^{16}$O$_2$ vibration is $4.738 \times 10^{13}$ sec$^{-1}$. How will the fractionation factor, $\alpha = (^{18}$O/$^{16}$O)$_{CO}/^{18}$O/$^{16}$O$_2$, vary as a function of temperature?

**Answer:** The equilibrium constant for this reaction is:

\[
K = \frac{Q_{C^{18}O}Q_{O^{18}O}}{Q_{C^{16}O}Q_{O^{16}O}} = 9.31
\]

The rotational and translational contributions are independent of temperature, so we calculate them first. The rotational contribution:

\[
K_{\text{rot}} = \left( \frac{Q_{C^{18}O}Q_{O^{18}O}}{Q_{C^{16}O}Q_{O^{16}O}} \right)_{\text{rot}} = \frac{\mu_{C^{18}O}\sigma_{C^{16}O}^{16} \sigma_{^{18}O}^{16}}{\mu_{C^{16}O}\sigma_{C^{16}O}^{16} \sigma_{^{18}O}^{16}} = \frac{1}{2} \frac{\mu_{^{18}O} \mu_{^{16}O}}{\mu_{^{16}O} \mu_{^{18}O}} = 0.9917
\]

Substituting $\mu_{^{16}O} = 6.857$, $\mu_{^{18}O} = 7.20$, $\mu_{^{18}O} = 8.471$, $\mu_{^{16}O} = 8$, we find: $K_{\text{rot}} = 0.9917/2 = 0.4959$

The translational contribution:

\[
K_{\text{trans}} = \frac{m_{C^{18}O}^{3/2}}{m_{C^{16}O}^{3/2}} \frac{m_{^{16}O}^{3/2}}{m_{^{18}O}^{3/2}} = 1.0126
\]

Substituting $m_{C^{16}O} = 28$, $m_{C^{18}O} = 30$, $m_{^{18}O} = 34$, $m_{^{16}O} = 32$ into 9.33, we find $K_{\text{trans}} = 1.0126$.

The vibrational contribution to the equilibrium constant is:

\[
K_{\text{vib}} = \frac{e^{-\frac{h(v_{c^{18}O} - v_{c^{16}O} + v_{^{18}O} - v_{^{16}O})}{kT}}}{(1 - e^{-\frac{h\nu_{c^{18}O}}{kT}})(1 - e^{-\frac{h\nu_{^{16}O}}{kT}})} = 9.34
\]

To obtain the vibrational contribution, we can assume the atoms vibrate as harmonic oscillators and solve equation 9.20 for the forcing constant, $k$ and calculate the vibrational frequencies for the $^{18}$O-bearing molecules. These turn out to be $6.348 \times 10^{13}$ sec$^{-1}$ for C-$^{18}$O and $4.605 \times 10^{13}$ sec$^{-1}$ for $^{16}$O$^{18}$O, so that 8.33 becomes:

\[
K_{\text{vib}} = \frac{e^{-5.580/T}(1 - e^{-3119/T})(1 - e^{-2208/T})}{(1 - e^{-3044/T})(1 - e^{-2272/T})}
\]

If we carry the calculation out at $T = 300$ K, we find:

\[
K = K_{\text{rot}}K_{\text{trans}}K_{\text{vib}} = 0.9917 \times 1.0126 \times 1.1087 = 1.0229
\]

To calculate the fractionation factor $\alpha$ from the equilibrium constant, we need to calculate $K_{\infty}$:

\[
K_{\infty} = \frac{(1/1)^{1}}{(2/1)^{1}} = \frac{1}{2}
\]

so that $\alpha = K/K_{\infty} = 2K$.

At 300 K, $\alpha = 1.0233$. The variation of $\alpha$ with temperature is shown in Figure 9.4.
between the fractionation factor and temperature approximates:

\[ \ln \alpha \propto \frac{1}{T^2} \]

Since \( \alpha \) is generally small, \( \ln \alpha = 1 + \alpha \), so that \( \alpha \propto 1 + 1/T^2 \). At infinite temperature, the fractionation is unity, since \( \ln \alpha = 0 \). This illustrated in Figure 9.3 for distribution of \( ^{18}O \) and \( ^{16}O \) between \( \text{CO}_2 \) and \( \text{H}_2\text{O} \). The \( \alpha \propto 1/T \) relationship breaks down around 200°C; above that temperature the relationship follows \( \alpha \propto 1/T^2 \).

It must be emphasized that the simple calculations performed in Example 9.1 are applicable only to a gas whose vibrations can be approximated by a simple harmonic oscillator. Real gases often show fractionations that are complex functions of temperature, with minima, maxima, inflections, and crossovers. Vibrational modes of silicates, on the other hand, are relatively well behaved.

9.2.1.4 Composition and Pressure Dependence

The nature of the chemical bonds in the phases involved is most important in determining isotopic fractionations. A general rule of thumb is that the heavy isotope goes into the phase in which it is most strongly bound. Bonds to ions with a high ionic potential and low atomic mass are associated with high vibrational frequencies and have a tendency to incorporate the heavy isotope preferentially. For example, quartz, \( \text{SiO}_2 \) is typically the most \( ^{18}O \) rich mineral and magnetite the least. Oxygen is dominantly covalently bonded in quartz, but dominantly ionically bonded in magnetite. The O is bound more strongly in quartz than in magnetite, so the former is typically enriched in \( ^{18}O \).

Substitution of cations in a dominantly ionic site (typically the octahedral sites) in silicates has only a secondary effect on the O bonding, so that isotopic fractionations of O isotopes between similar silicates are generally small. Substitutions of cations in sites that have a strong covalent character (generally tetrahedral sites) result in greater O isotope fractionations. Thus, for example, we would expect the fractionation between the end-members of the alkali feldspar series and water to be similar, since only the substitution of \( \text{K}^+ \) for \( \text{Na}^+ \) is involved. We would expect the fractionation factors between end-members of the plagioclase series and water to be greater, since this series involves the substitution of \( \text{Al} \) for \( \text{Si} \) as well as \( \text{Ca} \) for \( \text{Na} \), and the bonding of \( \text{O} \) to \( \text{Si} \) and \( \text{Al} \) in tetrahedral sites has a large covalent component.

Carbonates tend to be very \( ^{18}O \) rich because O is bonded to a small, highly charged atom, \( \text{C}^{4+} \). The fractionation, \( \Delta^{18}O_{\text{cal-water}} \), between calcite and water is about 30 per mil at 25°C. The cation in the carbonate has a secondary role (due to the effect of the mass of the cation on vibrational frequency). The \( \Delta^{18}O_{\text{carb-H}_2\text{O}} \) decreases to about 25 when \( \text{Ba} \) replaces \( \text{Ca} \) (\( \text{Ba} \) has about 3 times the mass of \( \text{Ca} \)).

Crystal structure plays a secondary role. The \( \Delta^{18}O \) between aragonite and calcite is of the order of 0.5 permil. However, is there apparently a large fractionation (10 permil) of C between graphite and diamond.

Pressure effects on fractionation factors turn out to be small, no more than 0.1 permil over 20 kbars. We can understand the reason for this by recalling that \( \partial \Delta G/\partial P = \Delta V \). The volume of an atom is entirely determined by its electronic structure, which does not depend on the mass of the nucleus. Thus the volume change of an isotope exchange reaction will be small, and hence there will be little pressure dependence. There will be a minor effect because vibrational frequency and bond length change as crystals are compressed. The compressibility of silicates is of the order of 1 part in \( 10^4 \), so we can expect effects on the order of \( 10^{-4} \) or less, which are generally insignificant.

9.2.2 Kinetic Isotope Fractionations

Kinetic isotope fractionations are normally associated with fast, incomplete, or unidirectional processes like evaporation, diffusion, dissociation reactions, and biologically mediated reactions. As an example, recall that temperature is related to the average kinetic energy. In an ideal gas, the average kinetic energy of all molecules is the same. The kinetic energy is given by:
Consider two molecules of carbon dioxide, $^{12}\text{C}^{16}\text{O}_2$ and $^{13}\text{C}^{16}\text{O}_2$, in such a gas. If their energies are equal, the ratio of their velocities is $(45/44)^{1/2}$, or 1.011. Thus $^{12}\text{C}^{16}\text{O}_2$ can diffuse 1.1% further in a given amount of time than $^{13}\text{C}^{16}\text{O}_2$. This result, however, is largely limited to ideal gases, i.e., low pressures where collisions between molecules are infrequent and intermolecular forces negligible. For the case where molecular collisions are important, the ratio of their diffusion coefficients is the ratio of the square roots of the reduced masses of $\text{CO}_2$ and air (mean molecular weight 28.8):

$$\frac{D_{^{12}\text{C}^{16}\text{O}_2}}{D_{^{13}\text{C}^{16}\text{O}_2}} = \frac{\sqrt{\mu_{^{12}\text{C}^{16}\text{O}_2}}} {\sqrt{\mu_{^{13}\text{C}^{16}\text{O}_2}}} = \frac{17.561}{17.406} = 1.0044$$

Hence we would predict that gaseous diffusion will lead to a 4.4‰ rather than 11‰ fractionation.

Molecules containing the heavy isotope are more stable and have higher dissociation energies than those containing the light isotope. This can be readily seen in Figure 9.1. The energy required to raise the $\text{D}_2$ molecule to the energy where the atoms dissociate is 441.6 kJ/mole, whereas the energy required to dissociate the $\text{H}_2$ molecule is 431.8 kJ/mole. Therefore it is easier to break bonds such as C-H than C-D. Where reactions attain equilibrium, isotopic fractionations will be governed by the considerations of equilibrium discussed above. Where reactions do not achieve equilibrium the lighter isotope will usually be preferentially concentrated in the reaction products, because of this effect of the bonds involving light isotopes in the reactants being more easily broken. Large kinetic effects are associated with biologically mediated reactions (e.g., photosynthesis, bacterial reduction), because such reactions generally do not achieve equilibrium and do not go to completion (e.g., plants don’t convert all CO$_2$ to organic carbon). Thus $^{12}\text{C}$ is enriched in the products of photosynthesis in plants (hydrocarbons) relative to atmospheric CO$_2$, and $^{32}\text{S}$ is enriched in H$_2$S produced by bacterial reduction of sulfate.

We can express this in a more quantitative sense. In Chapter 5, we found the reaction rate constant could be expressed as:

$$k = A e^{-E_b/kT} \quad (5.24)$$

where $k$ is the rate constant, $A$ the frequency factor, and $E_b$ is the barrier energy. For example, in a dissociation reaction, the barrier energy is the difference between the dissociation energy and the zero-point energy when the molecule is in the ground state, or some higher vibrational frequency when it is not (Fig. 9.1). The frequency factor is independent of isotopic composition, thus the ratio of reaction rates between the HD molecule and the H$_2$ molecule is:

$$\frac{k_{\text{D}}}{k_{\text{H}}} = e^{\left(\frac{E_b}{2kT}\right)}$$

or

$$\frac{k_{\text{D}}}{k_{\text{H}}} = e^{(\nu_{\text{H}} - \nu_{\text{D}})h/2kT} \quad 9.38$$

Substituting for the various constants, and using the wavenumbers given in the caption to Figure 9.1 (remembering that $\omega = c\nu$ where $c$ is the speed of light) the ratio is calculated as 0.24; in other words we expect the H$_2$ molecule to react four times faster than the HD molecule, a very large difference. For heavier elements, the rate differences are smaller. For example, the same ratio calculated for $^{16}\text{O}_2$ and $^{18}\text{O}^{16}\text{O}$ shows that the $^{16}\text{O}_2$ will react about 15% faster than the $^{18}\text{O}^{16}\text{O}$ molecule.

The greater translational velocities of lighter molecules also allow them to break through a liquid surface more readily and hence evaporate more quickly than a heavy molecule of the same composition. Thus water vapor above the ocean is typically around $\delta^{18}\text{O} = -13$ per mil, whereas at equilibrium the vapor should only be about 9 per mil lighter than the liquid.

Let’s explore this a bit further. An interesting example of a kinetic effect is the fractionation of O isotopes between water and water vapor. This is example of Rayleigh distillation (or condensation), and is similar to fractional crystallization. Let A be the amount of the species containing the major
isotope, \(H_2^{16}O\), and \(B\) be the amount of the species containing the minor isotope, \(H_2^{18}O\). The rate at which these species evaporate is proportional to the amount present:

\[
\frac{dA}{dt} = k_A A \tag{9.39a}
\]

and

\[
\frac{dB}{dt} = k_B B \tag{9.39b}
\]

Since the isotopic composition affects the reaction, or evaporation, rate, \(k_A \neq k_B\). Earlier we saw that for equilibrium fractionations, the fractionation factor is related to the equilibrium constant. For kinetic fractionations, the fractionation factor is simply the ratio of the rate constants, so that:

\[
\frac{k_B}{k_A} = \alpha \tag{9.40}
\]

and

\[
\frac{dB}{dA} = \alpha \frac{B}{A} \tag{9.41}
\]

Rearranging and integrating, we have:

\[
\ln \frac{B}{B^0} = \alpha \ln \frac{A}{A^0}
\]

or

\[
\frac{B}{B^0} = \left(\frac{A}{A^0}\right)\alpha
\]

where \(A^0\) and \(B^0\) are the amount of \(A\) and \(B\) originally present. Dividing both sides by \(A/A^0\)

\[
\frac{B/A}{B^0/A^0} = \left(\frac{A}{A^0}\right)\alpha^{-1}
\]

Since the amount of \(B\) makes up only a trace of the total amount of \(H_2O\) present, \(A\) is essentially equal to the total water present, and \(A/A^0\) is essentially identical to \(f\), the fraction of the original water remaining. Hence:

\[
\frac{B/A}{B^0/A^0} = f^{\alpha-1}
\]

Subtracting 1 from both sides, we have:

\[
\frac{B/A - B^0/A^0}{B^0/A^0} = f^{\alpha-1} - 1
\]

The left side of 9.45 is the relative deviation from the initial ratio. The permil relative deviation is simply:

\[
\Delta = 1000(f^{\alpha-1} - 1)
\]

Of course, the same principle applies when water condenses from vapor. Assuming a value of \(\alpha\) of 1.01, \(\delta\) will vary with \(f\), the fraction of vapor remaining, as shown in Figure 9.5.

Even if the vapor and liquid remain in equilibrium through the condensation process, the isotopic composition of the remaining vapor will change continuously. The relevant equation is:

\[
\Delta = \left(1 - \frac{1}{(1-f)/\alpha + f}\right)1000
\]

The effect of equilibrium condensation is also shown in Figure 9.5.
9.3 Isotope Geothermometry

One of the principal uses of stable isotopes is geothermometry. Like "conventional" chemical geothermometers, stable isotope geothermometers are based on the temperature dependence of the equilibrium constant. As we have seen, this dependence may be expressed as:

\[
\ln K = \ln \alpha = A + \frac{B}{T^2} \quad 9.48
\]

In actuality, the constants A and B are slowly varying functions of temperature, such that K tends to zero at absolute 0, corresponding to complete separation, and to 1 at infinite temperature, corresponding to no isotope separation. We can obtain a qualitative understanding of why this is so by recalling that the entropy of a system increases with temperature. At infinite temperature, there is complete disorder, hence isotopes would be mixed randomly between phases (ignoring for the moment the slight problem that at infinite temperature there would be neither phases nor isotopes). At absolute 0, there is perfect order, hence no mixing of isotopes between phases. A and B are, however, sufficiently invariant over a limited range temperatures that they can be viewed as constants. We have also noted that at low temperatures, the form of equation 9.48 changes to \( K \propto 1/T \).

In principal, a temperature may be calculated from the isotopic fractionation between any two phases provided the phases equilibrated and the temperature dependence of the fractionation factor is known. And indeed, there are too many isotope geothermometers for all of them to be mentioned here. Figure 9.6 shows some fractionation factors between quartz and other minerals as a function of temperature. Table 9.2 lists coefficients A and B for equation 9.48 for the oxygen isotope fractionation factor between quartz and other oxides and silicates.

Because of the dependence of the equilibrium constant on the inverse square of temperature, stable isotope geothermometry is employed primarily at low tempera-

Table 9.2. Coefficients for Oxygen Isotope Fractionation at Low Temperatures:

<table>
<thead>
<tr>
<th>( \Delta QZ - \phi )</th>
<th>A</th>
<th>B</th>
</tr>
</thead>
<tbody>
<tr>
<td>Feldspar</td>
<td>0</td>
<td>0.97 + 1.04b*</td>
</tr>
<tr>
<td>Pyroxene</td>
<td>0</td>
<td>2.75</td>
</tr>
<tr>
<td>Garnet</td>
<td>0</td>
<td>2.88</td>
</tr>
<tr>
<td>Olivine</td>
<td>0</td>
<td>3.91</td>
</tr>
<tr>
<td>Muscovite</td>
<td>-0.60</td>
<td>2.2</td>
</tr>
<tr>
<td>Amphibole</td>
<td>-0.30</td>
<td>3.15</td>
</tr>
<tr>
<td>Biotite</td>
<td>-0.60</td>
<td>3.69</td>
</tr>
<tr>
<td>Chlorite</td>
<td>-1.63</td>
<td>5.44</td>
</tr>
<tr>
<td>Ilmenite</td>
<td>0</td>
<td>5.29</td>
</tr>
<tr>
<td>Magnetite</td>
<td>0</td>
<td>5.27</td>
</tr>
</tbody>
</table>

* b is the mole fraction of anorthite in the feldspar. This term therefore accounts for the compositional dependence discussed above. From Javoy (1976).
tures, that is, non-magmatic temperatures. At temperatures in excess of 800°C or so, the fractionations are generally small, making accurate temperatures difficult to determine from them. However, even at temperatures of the upper mantle (1000°C or more), fractionations, although small, remain significant. Experimentally determined fractionation factors for minerals in the temperature range of 600° to 1300°C, which agree well with theoretical calculations, are given in Table 9.3.

Table 9.4 lists similar coefficients for the sulfur fractionation between H$_2$S and sulfur-bearing compounds. Recall that if phases $\alpha$ and $\gamma$ and $\alpha$ and $\beta$ are in equilibrium with each other, then $\gamma$ is also in equilibrium with $\beta$. Thus these tables may be used to obtain the fractionation between any two of the phases listed.

All geothermometers are based on the apparently contradictory assumptions that complete equilibrium was achieved between phases during, or perhaps after, formation of the phases, but that the phases did not re-equilibrate when they subsequently cooled. The reason these assumptions can be made and geothermometry works at all is the exponential dependence of reaction rates on temperature that we discussed in Chapter 5. Isotope geothermometers have the same implicit assumptions about the achievement of equilibrium as other geothermometers.

The importance of the equilibrium basis of geothermometry must be emphasized. Because most stable isotope geothermometers (though not all) are applied to relatively low temperature situations, violation of the assumption that complete equilibrium was achieved is not uncommon. We have seen that isotopic fractionations may arise from kinetic as well as equilibrium effects. If reactions do not run to completion, the isotopic differences may reflect kinetic effects as much as equilibrium effects. There are other problems that can result in incorrect temperature as well, for example, the system may partially re-equilibrate at some lower temperature during cooling. A further problem with isotope geothermometry is that free energies of the exchange reactions are relatively low; meaning there is little chemical energy available to drive the reaction. Indeed, isotopic equilibrium probably often depends on other reactions occurring that mobilize the element involved in the exchange. Solid-state exchange reactions will be particularly slow at temperatures well below the melting point. Equilibrium between solid phases will thus generally depend on reaction of these phases with a fluid. This latter point is true of ‘conventional’ geothermometers as well, and metamorphism, one of the important areas of application of isotope geothermometry, generally occurs in the presence of a fluid.

Isotope geothermometers do have several advantages over conventional chemical ones. First, as we have noted, there is no volume change associated with isotopic exchange reactions and hence no pressure dependence of the equilibrium constant. However, Rumble has suggested an indirect
pressure dependence, wherein the fractionation factor depends on fluid composition, which in turn depends on pressure. Second, whereas conventional chemical geothermometers are generally based on solid solution, isotope geothermometers can make use of pure phases such as SiO$_2$, etc. Generally, any dependence on the composition of phases involved is of relatively second order importance (there are, however, exceptions). For example, isotopic exchange between calcite and water is independent of the concentration of CO$_2$ in the water. Compositional effects can be expected only where it effects bonds formed by the element involved in the exchange. For example, we noted substitution of Al for Si in plagioclase affects O isotope fractionation factors because the nature of the bond with oxygen.

### 9.4 Isotope Fractionation in the Hydrologic System

As we noted above, isotopically light water has a higher vapor pressure, and hence lower boiling point, than isotopically heavy water. Let's consider this in a bit more detail. Raoult's Law (equ. 3.8) states that the partial pressure of a species

$$\ln \alpha_{Qz-H_2O} = -3.70 + 4.10 \times 10^6 \frac{T^2}{T^2}$$

and

$$\ln \alpha_{Mt-H_2O} = -3.10 + 1.9 \times 10^6 \frac{T^2}{T^2}$$

### Example 9.2: Oxygen Isotope Geothermometry

A granite-gneiss contains coexisting quartz, muscovite and magnetite of with the following $\delta^{18}O$: quartz: 11.2; magnetite: 1.9. Find the temperature of equilibration.

**Answer**: According to Bottinga and Javoy (1973), the fraction factors for these minerals can be expressed as:

$$1000 \ln \alpha_{Qz-H_2O} = -3.70 + 4.10 \times 10^6 \frac{T^2}{T^2}$$

and

$$1000 \ln \alpha_{Mt-H_2O} = -3.10 + 1.9 \times 10^6 \frac{T^2}{T^2}$$

At what temperature did these minerals equilibrate?

**Answer**: The fractionation factor $\alpha_{Qz-Mt}$ can be found as: $\alpha_{Qz-H_2O} / \alpha_{Mt-H_2O}$, and

$$1000 \ln \alpha_{Qz-Mag} = 1000 (\ln \alpha_{Qz-H_2O} - \ln \alpha_{Mag-H_2O}) = -3.70 + 4.10 \frac{T^2}{T^2} + 3.10 - 1.9 \frac{T^2}{T^2} = 0.6 + 2.20 \frac{T^2}{T^2}$$

Substituting $\Delta Qz-Mag$ for $1000 \ln \alpha_{Qz-Mag}$ and solving for $T$:

$$T = \sqrt{\frac{2.20 \times 10^6}{\Delta Qz-Mag - 0.6}}$$

We calculate $\Delta Qz-Mag$ as 9.2‰. Substituting this into 9.47, we find $T = 505$ K, = 232 °C.
above a solution is equal to its molar concentration in the solution times the partial pressure exerted by the pure solution. So for the two isotopic species of water (we will restrict ourselves to O isotopes for the moment), Raoult’s Law is:

\[ p_{H^2O^{16}} = p_{H^2O^{16}}^o \left[H_2O^{16}\right] \]

and

\[ p_{H^2O^{18}} = p_{H^2O^{18}}^o \left[H_2O^{18}\right] \]

Since the partial pressure of a species is proportional to the number of atoms of that species in a gas, we can define the fractionation factor, \( \alpha \), between liquid water and vapor as:

\[ \alpha_{lv} = \frac{p_{H^2O^{18}}}{p_{H^2O^{16}}} \]

Substituting 9.50a and 9.50b into 9.51, we arrive at the relationship:

\[ \alpha_{lv} = \frac{p_{H^2O^{18}}}{p_{H^2O^{16}}} \]

Thus interestingly enough, the fractionation factor for oxygen between water vapor and liquid turns out to be just the ratio of the standard state partial pressures. The next question is how the partial pressures vary with temperature. Thermodynamics provides the answer. The temperature dependence of the partial pressure of a species may be expressed as:

\[ \frac{d\ln p}{dT} = \frac{\Delta H}{RT^2} \]

where \( T \) is temperature, \( \Delta H \) is the enthalpy or latent heat of evaporation, and \( R \) is the gas constant. Over a sufficiently small range of temperature, we can assume that \( \Delta H \) is independent of temperature. Rearranging and integrating, we obtain:

\[ \ln p = \frac{\Delta H}{RT} + \text{const} \]

We can write two such equations, one for \([H_2O^{16}]\) and [one for \(H_2O^{18}\)]. Dividing one by the other we obtain:

\[ \ln \frac{p_{H^2O^{18}}}{p_{H^2O^{16}}} = A - \frac{B}{RT} \]

where A and B are constants. This can be rewritten as:

\[ \alpha = a e^{B/RT} \]

Over a larger range of temperature, \( \Delta H \) is not constant. The log of the fractionation factor in that case depends on the inverse square of temperature, so that the temperature dependence of the fractionation factor

\[ \delta^{18}O = 85^\circ S \]

Figure 9.8. Variation of \( \delta^{18}O \) in precipitation as a function of mean annual temperature.
can be represented as:

\[ \ln \alpha = A - \frac{B}{T^2} \]

Given the fractionation between water and vapor, we might predict that there will be considerable variation in the isotopic composition of water in the hydrologic cycle, and indeed there is. Figure 9.8 shows the global variation in \( \delta^{18}O \) in precipitation.

Precipitation of rain and snow from clouds is a good example of Rayleigh condensation. Isotopic fractionations will therefore follow equation 9.46. Thus in addition to the temperature dependence of \( \alpha \), the isotopic composition of precipitation will also depend on \( f \), the fraction of water vapor remaining in the air. The further air moves from the site of evaporation, the more water is likely to have condensed and fallen as rain, and therefore, the smaller the value of \( f \) in equation 9.46. Thus fractionation will increase with distance from the region of evaporation (principally tropical and temperature oceans). Topography also plays a role as mountains force air up, causing it to cool and water vapor to condense, hence decreasing \( f \). Thus precipitation from air that has passed over a mountain range will be isotopically lighter than precipitation on the ocean side of a mountain range. These factors are illustrated in the cartoon in Figure 9.9.

Hydrogen as well as oxygen isotopes will be fractionated in the hydrologic cycle. Indeed, \( \delta^{18}O \) and \( \delta^D \) are reasonably well correlated in precipitation, as is shown in Figure 9.10. The fractionation of hydrogen isotopes, however, is greater because the mass difference is greater.

### 9.5 Isotope Fractionation in Biological Systems

Biological processes often involve large isotopic fractionations. Indeed, for carbon, nitrogen, and sulfur, biological processes are the most important cause of isotope fractionations. For the most part, the largest fractionations occur during the initial production of organic matter by the so-called
primary producers, or autotrophs. These include all plants and many kinds of bacteria. The most important means of production of organic matter is photosynthesis, but organic matter may also be produced by chemosynthesis, for example at mid-ocean ridge hydrothermal vents. Large fractionations of both carbon and nitrogen isotopes occur during primary production. Additional fractionations also occur in subsequent reactions and up through the food chain as heterotrophs consume primary producers, but these are generally smaller.

9.5.1 Carbon Isotope Fractionation During Photosynthesis

Biological processes are the principal cause of variations in carbon isotope ratios. The most important of these processes is photosynthesis (a discussion of photosynthesis may be found in Chapter 14). As we earlier noted, photosynthetic fractionation of carbon isotopes is primarily kinetic. The early work of Park and Epstein (1960) suggested fractionation occurred in several steps. Subsequent work has elucidated the fractionations involved in these steps, which we will consider in slightly more detail.

For terrestrial plants (those utilizing atmospheric CO₂), the first step is diffusion of CO₂ into the boundary layer surrounding the leaf, through the stomata, and internally in the leaf. On theoretical grounds, a fractionation of $-4.4\%$ is expected, as we found in equation 9.36. Marine algae and aquatic plants can utilize either dissolved CO₂ or HCO₃⁻ for photosynthesis:

\[
\text{CO}_2(g) \rightarrow \text{CO}_2(aq) + \text{H}_2\text{O} \rightarrow \text{H}_2\text{CO}_3 \rightarrow \text{H}^+ + \text{HCO}_3^-
\]

An equilibrium fractionation of +0.9 per mil is associated with dissolution ($^{13}\text{CO}_2$ will dissolve more readily), and an equilibrium +7.0 to +8‰ fractionation occurs during hydration and dissociation of CO₂.

At this point, there is a divergence in the chemical pathways. Most plants use an enzyme called ribulose bisphosphate carboxylase oxygenase (RUBISCO) to catalyze a reaction in which ribulose bisphosphate carboxylase reacts with one molecule of CO₂ to produce 3 molecules of 3-phosphoglyceric acid, a compound containing 3 carbon atoms, in a process called carboxylation (Figure 9.11).

The carbon is subsequently reduced, carbohydrate formed, and the ribulose bisphosphate regenerated. Such plants are called C₃ plants, and this process is called the Benson-Calvin Cycle, or simply the Calvin Cycle. C₃ plants constitute about 90% of all plants today and include algae and autotrophic bacteria and comprise the majority of cultivated plants, including wheat, rice, and nuts. There is a kinetic fractionation associated with carboxylation of ribulose bisphosphate that has been determined by several methods to be $-29.4\%$ in higher terrestrial plants. Bacterial carboxylation has different reaction mechanisms and a smaller fractionation of about $-20\%$. Thus for terrestrial plants a fractionation of about $-34\%$ is expected from the sum of the individual fractionations. The actual observed total fractionation is in the range of $-20$ to $-30\%$. The disparity between the observed total fractionation and that expected from the sum of the steps presented something of a conundrum. This
solution appears to be a model that assumes the amount of carbon isotope fractionation expressed in the tissues of plants depends on the ratio of the concentration of CO$_2$ inside plants to that in the external environment.

The other photosynthetic pathway is the Hatch-Slack cycle, used by the C$_4$ plants that include hot-region grasses and related crops such as maize and sugarcane. These plants use *phosphoenol pyruvate carboxylase* (PEP) to fix the carbon initially and form oxaloacetate, a compound that contains 4 carbons (Fig. 9.12). A much smaller fractionation, about -2.0 to -2.5‰, occurs during this step. In phosphophoenol pyruvate carboxylation, the CO$_2$ is fixed in outer mesophyll cells as oxaloacetate and carried as part of a C$_4$ acid, either malate or asparatate, to inner bundle sheath cells where it is decarboxylated and refixed by RuBP (Fig. 9.13). The environment in the bundle sheath cells is almost a closed system, so that virtually all the carbon carried there is refixed by RuBP, so there is little fractionation during this step. C$_4$ plants have average $\delta^{13}C$ of -13‰. As in the case of RuBP photosynthesis, the fractionation appears to depend on the ambient concentration of CO$_2$.

A third group of plants, the CAM plants, has a unique metabolism called the *Crassulacean acid metabolism*. These plants generally use the C$_4$ pathway, but can use the C$_3$ pathway under certain conditions. These plants are generally adapted to arid environments and include pineapple and many cacti; they have $\delta^{13}C$ intermediate between C$_3$ and C$_4$ plants.

Terrestrial plants, which utilize CO$_2$ from the atmosphere, generally produce greater fractionations than marine and aquatic plants, which utilize dissolved CO$_2$ and HCO$_3^-$, together referred to as *dissolved inorganic carbon* or DIC. As we noted above, there is about an +8‰ equilibrium fractionation between dissolved CO$_2$ and HCO$_3^-$. Since HCO$_3^-$ is about 2 orders of magnitude more abundant in seawater than dissolved CO$_2$, marine algae utilize this species, and hence tend to show a lower net fractionation between dissolved carbonate and organic carbon during photosynthesis. Diffusion is slower in water than in air, so diffusion is often the rate limiting step. Most aquatic plants have some membrane-bound mechanism to pump DIC, which can be turned on when DIC is low. When DIC concentrations are high, fractionation in aquatic and marine plants is generally similar to that in terrestrial plants. When it is low and the plants are actively pumping DIC, the fractionation is less because most of the carbon pumped into cells is fixed. Thus carbon isotope fractionations between dissolved inorganic carbon and organic carbon can be as low as 5‰ in algae.

Not surprisingly, the carbon isotope fractionation in C fixation is also temperature dependent. Thus higher fractionations are observed in cold water phytoplankton than in warm water species. However, this observation also reflects a kinetic effect: there is generally less dissolved CO$_2$ available in warm waters because of the decreasing solubility with temperature. As a result, a larger fraction of the CO$_2$ is utilized and there is consequently less fractionation. Surface waters of the ocean are generally enriched in $^{13}C$ (relative to $^{12}C$) because of uptake of $^{13}C$ during photosynthesis (Figure 9.14). The degree of enrichment depends on the productivity: biologically productive areas show greater depletion. Deep water, on
the other hand, is enriched in $^{12}$C. Organic matter falls through the water column and is decomposed and "remineralized", i.e., converted to inorganic carbon, by the action of bacteria, enriching deep water in $^{12}$C and total DIC. Thus biological activity acts to "pump" carbon, and particularly $^{12}$C from surface to deep waters.

Essentially all organic matter originates through photosynthesis. Subsequent reactions convert the photosynthetically produced carbohydrates to the variety of other organic compounds utilized by organisms. Further fractionations occur in these reactions. These fractionations are thought to be kinetic in origin and may partly arise from organic C-H bonds being enriched in $^{12}$C and organic C-O bonds are enriched in $^{13}$C. $^{12}$C is preferentially consumed in respiration (again, because bonds are weaker and it reacts faster), which enriches residual organic matter in $^{12}$C. Thus the carbon isotopic composition of organisms becomes slightly more positive moving up the food chain.

9.5.2 Nitrogen Isotope Fractionation in Biological Processes

Nitrogen is another important element in biological processes, being an essential component of all amino acids, proteins, and other key compounds. The understanding of isotopic fractionations of nitrogen is much less advanced than that of carbon. There are five important forms of inorganic nitrogen ($N_2$, $NO_3^-$, $NO_2^-$, $NH_3$ and $NH_4^+$). Equilibrium isotope fractionations occur between these five forms, and kinetic fractionations occur during biological assimilation of nitrogen. Ammonia is the form of nitrogen that is ultimately incorporated into organic matter by growing plants. Most terrestrial plants depend on symbiotic bacteria for fixation (i.e., reduction) of $N_2$ and other forms of nitrogen to ammonia. Many plants, including many marine algae, can utilize oxidized nitrogen, $NO_3^-$ and $NO_2^-$, and a few (blue-green algae and legumes, for example) are able to utilize $N_2$ directly. In these cases, nitrogen must first be reduced by the action of reductase enzymes. As with carbon, fractionation may occur in each of several steps that occurs in the nitrogen assimilation process.

While isotope fractionations during assimilation of ammonium are still poorly understood, it appears there is a strong dependence on the concentration of the ammonium ion. Models of nitrogen isotope fractionation in plants imply a dependence of fractionation on the abundance of ammonium. Such dependence has been observed, as for example in Figure 9.15. The complex dependence in Figure 9.15 is interpreted as follows. The increase in fractionation from highest to moderate concentrations of ammonium reflects the switching on of active ammonium transport by cells. At the lowest concentrations, essentially all available nitrogen is transported into the cell and assimilated, so there is little fractionation observed.

The isotopic compositions of marine particulate nitrogen and non-nitrogen-fixing plankton are typically $-3\%$ to $+12\% \delta^{15}N$. Non-nitrogen fixing terrestrial plants unaffected by artificial fertilizers generally have a narrower range of $+6\%$ to $+13$ per mil, but are isotopically lighter on average. Marine blue-green algae range from $-2$ to $+4$, with most in the range of $-4$ to $-2\%$. Most nitrogen-fixing terrestrial plants fall in the range of $-2$ to $+4\%$, and hence are typically heavier than non-nitrogen fixing plants.

9.5.3 Oxygen and Hydrogen Isotope Fractionation by Plants

Oxygen is incorporated into biological material from $CO_2$, $H_2O$, and $O_2$. However, both $CO_2$ and $O_2$ are in oxygen isotopic equilibrium with water during photosynthesis, and water is...
the dominant source of O. Therefore, the isotopic composition of plant water determines the oxygen isotopic composition of plant material. The oxygen isotopic composition of plant material seems to be controlled by exchange reactions between water and carbonyl oxygens (oxygens doubly bound to carbon):

\[ \text{C}^16\text{O} + \text{H}_2^18\text{O} \rightleftharpoons \text{C}^{18}\text{O} + \text{H}_2^16\text{O} \]

Fractionations of +16 to +27‰ (i.e., the organically bound oxygen is heavier) have been measured for these reactions. Consistent with this, cellulose from most plants has $\delta^{18}O$ of +27±3. Other factors, however, play a role in the oxygen isotopic composition of plant material. First, the isotopic composition of water varies from $\delta^{18}O \approx -55$ in Arctic regions to $\delta^{18}O = 0$ in the oceans. Second, less than complete equilibrium may be achieved if photosynthesis is occurring at a rapid pace, resulting in less fractionation. Finally, some fractionation of water may occur during transpiration, with residual water in the plant becoming heavier.

Hydrogen isotope fractionation during photosynthesis occurs such that the light isotope is enriched in organic material. In marine algae, isotope fractionations of −100 to −150 ‰ have been observed, which is a little more than that observed in terrestrial plants (−86 to −120). Among terrestrial plants, there appears to be a difference between C₃ and C₄ plants. The former show fractionations of −117 to −121‰, while fractionations −86 to −109‰ have been observed in C₄ plants. However, little is known in detail about the exact mechanisms of fractionation.

As for oxygen, variations in the isotopic composition of available water and fractionation during transpiration are important in controlling the hydrogen isotopic composition of plants. This is illustrated in Figure 9.16.

### 9.5.4 Biological Fractionation of Sulfur Isotopes

Though essential to life, sulfur is a minor component in living tissue (C:S atomic ratio is about 200). Plants take up sulfur as sulfate and subsequently reduce it to sulfide and incorporate it into cysteine. There is apparently no fractionation of sulfur isotopes in transport across cell membranes and incorporation, but there is a fractionation of +0.5 to −4.5‰ in the reduction process, referred to as assimilatory sulfate reduction. This is substantially less than the expected fractionation of about −20‰, suggesting most sulfur taken up by primary producers is reduced and incorporated into tissue.

Sulfur, however, plays two other important roles in biological processes. First, sulfur, in the form of sulfate, can act as an electron acceptor or oxidant, and is utilized as such by sulfur reducing bacteria. This process, in which H₂S is liberated, is called dissimilatory sulfate reduction and plays an important role in biogeochemical cycles, both as a sink for sulfur and source for atmospheric oxygen. A large fractionation of +5 to −46‰ is associated with this process. This process produces by far the most significant fractionation of sulfur isotopes, and thus governs the isotopic composition of sulfur in the exogene. Sedimentary sulfate typically has $\delta^{34}S$ of about +17, which is
similar to the isotopic composition of sulfate in the oceans (+20), while sedimentary sulfide has a δ\(^{34}\)S of −18. The living biomass has a δ\(^{34}\)S of ≈ 0.

The final important role of sulfur is a reductant. Sulfide is an electron acceptor used by some types of photosynthetic bacteria as well as other bacteria in the reduction of CO\(_2\) to organic carbon. Unique among these perhaps are the chemosynthetic bacteria of submarine hydrothermal vents. They utilize H\(_2\)S emanating from the vents as an energy source and form the base of the food chain in these unique ecosystems. A fractionation of +2 to −18‰ is associated with this process.

### 9.5.5 Isotopes and Diet: You Are What You Eat

As we have seen, the two main photosynthetic pathways, C\(_3\) and C\(_4\), lead to δ\(^{13}\)C among the principal classes of autotrophs. Organic carbon with different carbon isotopic compositions. Terrestrial C\(_3\) plants have δ\(^{13}\)C values that average of about −27‰, C\(_4\) plants an average δ\(^{13}\)C of about −13‰. Marine plants (which are all C\(_3\)) utilized dissolved bicarbonate rather than atmospheric CO\(_2\). Seawater bicarbonate is about 8.5‰ heavier than atmospheric CO\(_2\), so marine plants average about 7.5‰ heavier than terrestrial C\(_3\) plants. In addition, because the source of the carbon they fix is isotopically more variable, the isotopic composition of marine plants is also more variable. Finally, marine cyanobacteria (blue-green algae) tend to fractionate carbon isotopes less during photosynthesis than do true marine plants, so they tend to average 2 to 3‰ higher in δ\(^{13}\)C.

Plants may also be divided into two types based on their source of nitrogen: those that can utilize N\(_2\) directly, and those that utilize only “fixed” nitrogen in ammonia and nitrate. The former include the legumes (e.g., beans, peas, etc.) and marine cyanobacteria. The legumes, which are exclusively C\(_3\) plants, utilize both N\(_2\) and fixed nitrogen, and have an average δ\(^{15}\)N of +1‰, whereas modern nonleguminous plants average about +3‰. Prehistoric nonleguminous plants were more positive, averaging perhaps +9‰, because the isotopic composition of present soil nitrogen has been affected by the use of chemical fertilizers. For both groups, there was probably a range in δ\(^{15}\)N of ±4 or 5‰, because the isotopic composition of soil nitrogen varies and there is some fractionation involved in uptake. Marine plants have δ\(^{15}\)N of +7±5‰, whereas marine cyanobacteria have δ\(^{15}\)N of −1±3‰. Thus based on their δ\(^{13}\)C and δ\(^{15}\)N values, autotrophs can be divided into several groups, which are summarized in Figure 9.17.

DeNiro and Epstein (1978) studied the relationship between the carbon isotopic composition of animals and their diet. They found that there is only slight further fractionation of carbon by animals and that the carbon isotopic composition of animal tissue closely reflects that of the animal’s diet. Typically, carbon in animal tissue is about 1‰ heavier than their diet. The small fractionation between animal tissue and diet is a result of the slightly weaker bond formed by 12C compared to 13C. The weaker bonds are more readily broken during respiration, and, not surprisingly, the CO\(_2\) respired by most animals investigated was slightly lighter than their diet. Thus only a small fractionation in carbon isotopes occurs as organic carbon passes up the food web. Terrestrial food chains usually do not have more than 3 trophic levels, implying a maximum further fractionation of +3‰; marine food chains can have up to 7 trophic levels, implying a maximum carbon isotope difference between primary producers and top predators of 7‰. These differences are smaller than the range observed in primary producers.
In another study, DeNiro and Epstein (1981) found that $\delta^{15}N$ of animal tissue reflects the $\delta^{15}N$ of the animal’s diet, but is typically $3$ to $4\%$ higher than that of the diet. Thus in contrast to carbon, significant fractionation of nitrogen isotopes will occur as nitrogen passes up the food chain. These relationships are summarized in Figure 9.18. The significance of these results is that it is possible to infer the diet of an animal from its carbon and nitrogen isotopic composition.

Schoeninger and DeNiro (1984) found that the carbon and nitrogen isotopic composition of bone collagen in animals was similar to that of body tissue as a whole. Apatite in bone appears to undergo isotopic exchange with meteoric water once it is buried, but bone collagen and tooth enamel appear to be robust and retain their original isotopic compositions. This means that the nitrogen and carbon isotopic composition of fossil bone collagen and teeth can be used to reconstruct the diet of fossil animals.

An interesting example of this is horses (Family Equidae), which have been around for 58 million years. Beginning in the early Miocene, a major radiation took place and the number of genera in North America increased from three at 25 Ma to twelve at 10 Ma. It subsequently fell at the end of the Miocene, and the last North American species became extinct in the Holocene. A major change in dental morphology, from low-crowned to high crowned, accompanied the Miocene radiation. For nearly 100 years, the standard textbook explanation of this dental change was that associated with a change in feeding from leaf browsing to grass grazing. Grasses contain enough silica to make them quite abrasive, thus a high crowned tooth would last longer in a grazing animal and would therefore be favored in horse’s evolution as it switched food sources. The change in horse diet was thought to reflect the evolution of grassland ecosystems (or biomes). This line of reasoning led to the conclusion that grasslands first became important biomes in the Miocene.

Carbon isotope ratios provide the first opportunity to test this hypothesis, since most grasses are C₄ plants whereas most trees and shrubs are C₃ plants. Wang et al. (1994) carried out such a test by analyzing the carbon isotopic composition of dental enamel from fossil horse teeth of Eocene through Pleistocene age. They found a sharp shift in the isotopic composition of the teeth consistent with a change in diet from C₃ to C₄ vegetation, but it occurred later than the change in dental morphology (Figure 9.19). The change in dental morphology begins in the mid-Miocene (about 18 Ma), while shift in $\delta^{13}C$ occurs at around 7 Ma. This leads to an interesting dilemma. Which change, that in morphology or that in carbon isotopic composition, actually reflects the appearance of the grassland biome?
Another recent study suggests a possible resolution: that grasslands appeared in mid-Miocene but only became dominated by C₄ grasses in the late Miocene. Quade et al. (1989) inferred that ecosystems in Pakistan shifted from C₃ dominant to C₄ dominant about 7 million years ago from δ¹³C of soil carbonate (Figure 9.20). They initially interpreted this as a response to the uplift of the Tibetan Plateau and the development of the Monsoon. However, other evidence, including oxygen isotope data from Pakistani soil carbonates, suggests the Monsoons developed about a million years earlier. Further, the synchronicity of the dominance of C₄ grasslands in Asia and North America suggests a global cause, while the Monsoons are a regional phenomenon. C₄ photosynthesis is more efficient at low concentrations of CO₂ that is the C₃ pathway. Thus the evolution of C₄ plants may have occurred in response to a decrease in atmospheric CO₂ suspected on other grounds (Cerling et al., 1993).

9.5.5.1 Isotopes in Archaeology

The differences in nitrogen and carbon isotopic composition of various foodstuffs and the preservation of these isotope ratios in bone collagen provides a means of determining what ancient peoples ate. In the first investigation of bone collagen from human remains, DeNiro and Epstein (1981) concluded that Indians of the Tehuacan Valley in Mexico probably depended heavily on maize (a C₄ plant) as early as 4000 BC, whereas archeological investigations had concluded maize did not become important in their diet until perhaps 1500 BC. In addition, there seemed to be steady increase in the dependence on legumes (probably beans) from 6000 BC to 1000 AD and a more marked increase in legumes in the diet after 1000 AD.

Mashed grain and vegetable charred onto pot sherds during cooking provides an additional record of the diets of ancient peoples. DeNiro and Hasdorf (1985) found that vegetable matter subjected to conditions similar to burial in soil underwent large shifts in δ¹⁵N and δ¹³C but that vegetable matter that was burned or charred did not. The carbonization (charring, burning) process itself produced only small (2 or 3‰) fractionations. Since these
Fractionations are smaller than the range of isotopic compositions in various plant groups, they are of little significance. Since pot sherds are among the most common artifacts recovered in archeological sites, this provides a second valuable means of reconstructing the diets of ancient peoples.

Figure 9.21 summarizes the results obtained in a number of studies of bone collagen and potsherds (DeNiro, 1987). Studies of several modern populations, including Eskimos and the Tlinglit Indians of the Northwest U.S. were made as a control. Judging from the isotope data, the diet of Neolithic Europeans consisted entirely of C₃ plants and herbivores feeding on C₃ plants, in contrast to the Tehuacan Indians, who depended mainly on C₄ plants. Prehistoric peoples of the Bahamas and Denmark depended both on fish and on agriculture. In the case of Mesolithic Denmark, other evidence indicates the crops were C₄, and the isotope data bear this out. Although there is no corroborating evidence, the isotope data suggest the Bahamas also depended on C₃ rather than C₄ plants. The Bahamians had lower δ¹⁵N because the marine component of their diet came mainly from coral reefs. Nitrogen fixation is particularly intense on coral reefs, which leads to ¹⁵N depletion of the water, and consequently, of reef organisms.

**9.6 Paleoclimatology**

Perhaps one of the most successful and significant application of stable isotope geothermometry has been paleoclimatology. At least since the work of Louis Agassiz in 1840, geologists have contemplated the question of how the Earth’s climate might have varied in the past. Until 1947, they had no means of quantifying paleotemperature changes. In that year, Harold Urey initiated the field of stable isotope geochemistry. In his classic 1947 paper (Urey, 1947), Urey calculated the temperature dependence of oxygen isotope fractionation between calcium carbonate and water and proposed that the isotopic composition of carbonates could be used as a paleothermometer. Urey’s students and post-doctoral fellows empirically determined temperature dependence of the fractionation between calcite and water as:

\[ T \, ^°C = 16.9 - 4.2 \Delta_{CaH_2O} + 0.13 \Delta_{CaH_2O}^2 \]  

For example, a change in \( \Delta_{C-W} \) from 30 to 31 permil implies a temperature change from 8° to 12°. Urey suggested that the Earth’s climate history could be recovered from oxygen isotope analyses of
ancient marine carbonates. Although the problem has turned out to be much more complex than Urey anticipated, this has proved to be an extremely fruitful area of research. Deep sea carbonate oozes contained an excellent climate record, and, as we shall see, several other paleoclimatic records are available as well.

9.6.1 The Marine Quaternary $\delta^{18}O$ Record and Milankovitch Cycles

The principles involved in paleoclimatology are fairly simple. As Urey formulated it, the isotopic composition of calcite secreted by organisms should provide a record of paleo-ocean temperatures because the fractionation of oxygen isotopes between carbonate and water is temperature dependent. In actual practice, the problem is somewhat more complex because the isotopic composition of the shell, or test, of an organism will depend not only on temperature, but also on the isotopic composition of water in which the organism grew, “vital effects” (i.e., different species may fractionate oxygen isotopes somewhat differently), and post-burial isotopic exchange with sediment pore water. As it turns out, the vital effects and post-burial exchange are usually not very important for late Tertiary/Quaternary carbonates, but the isotopic composition of water is.

The first isotopic work on deep sea sediment cores with the goal of reconstructing the temperature history of Pleistocene glaciations was by Cesare Emiliani (1955), who was then a student of Urey at the University of Chicago. Emiliani analyzed $\delta^{18}O$ in foraminifera from piston cores from the world’s oceans. Remarkably, many of Emiliani’s findings are still valid today, though they have been revised to various degrees. Rather than just the 5 glacial periods that geomorphologists had recognized, Emiliani found 15 glacial–interglacial cycles over the last 600,000 years. He found that these were global events, with notable cooling even in low latitudes and concluded that the fundamental driving force for Quaternary climate cycles was variations in the Earth’s orbital parameters.

Emiliani had realized that the isotopic composition of the ocean would vary between glacial and interglacial times as isotopically light water was stored in glaciers, thus enriching the oceans in $^{18}O$. He estimated that this factor accounted for about 20% of the observed variations. The remainder he attributed to the effect of temperature on isotope fractionation. Subsequently, Shackleton and Opdyke (1973) concluded that storage of isotopically light water in glacial ice was the main effect causing oxygen isotopic variations in biogenic carbonates, and that the temperature effect was only secondary.

The question of just how much of the variation is deep sea carbonate sediments in due to ice build-up and how much is due to the effect of temperature on fractionation is still debated. The resolution depends, in part, on the isotopic composition of glacial ice and how much this might vary between glacial and interglacial times. It is fairly clear that the average $\delta^{18}O$ of glacial ice is probably less than −15‰, as Emiliani had assumed. Typical values for Greenland

$$
\begin{align*}
\text{δ}^{18}O^\text{Glacial} & = -30 \\
\text{δ}^{18}O^\text{Interglacial} & = 0.0
\end{align*}
$$

Figure 9.22. Cartoon illustrating how $\delta^{18}O$ of the ocean changes between glacial and interglacial periods.
ice are -30 to -35‰ (relative to SMOW) and as low as -50‰ for Antarctic ice. If the exact isotopic composition of ice and the ice volume were known, it would be a straightforward exercise to calculate the effect of continental ice build-up on ocean isotopic composition. For example, the present volume of continental ice is $27.5 \times 10^6$ km$^3$, while the volume of the oceans is $1350 \times 10^6$ km$^3$. Assuming glacial ice has a mean $\delta^{18}O$ of -30‰ relative to SMOW, we can calculate the $\delta^{18}O$ of the total hydrosphere as -0.6‰ (neglecting freshwater reservoirs, which are small). At the height of the Wisconsin Glaciation (the most recent one), the volume of glacial ice is thought to have increased by $42 \times 10^6$ km$^3$, corresponding to a lowering of sea level by 125 m. If the $\delta^{18}O$ of ice was the same then as now (-30‰), we can readily calculate that the $\delta^{18}O$ of the ocean would have increased by 1.59‰. This is illustrated in Figure 9.22.

We can use equation 9.58 to see how much the effect of ice volume on seawater $\delta^{18}O$ affects estimated temperature changes. According to this equation, at 20° C, the fractionation between water and calcite should be 33‰. Assuming present water temperature of 20° C, Emiliani would have calculated a temperature change of 6° C for an observed increase in the $\delta^{18}O$ of carbonates between glacial times and present $\delta^{18}O$ of 2‰, after correction for 0.5‰ change in the isotopic composition of water. In other words, he would have concluded that surface ocean water in the same spot would have had a temperature of 14° C. If the change in the isotopic composition of water is actually 1.5‰, the calculated temperature difference is only about 2° C. Thus the question of the volume of glacial ice and its isotopic composition must be resolved before $\delta^{18}O$ in deep sea carbonates can be used to calculate palaeotemperatures.

Comparison of sealevel curves derived from dating of terraces and coral reefs indicate that each 0.011‰ variation in $\delta^{18}O$ represents a 1 m change in sealevel. Based on this and other observations, it is now generally assumed that the $\delta^{18}O$ of the ocean changed by about 1.5‰ between glacial and interglacial periods, but the exact value is still debated.

By now hundreds, if not thousands, of deep sea sediment cores have been analyzed for oxygen isotope ratios. Figure 9.23 shows the global $\delta^{18}O$ record constructed by averaging analyses from 5 key cores (Imbrie et al., 1984). A careful examination of the global curve shows a periodicity of approximately 100,000 years. The same periodicity was apparent in Emiliani’s initial work and led him to conclude that the glacial-interglacial cycles were due to variations in the Earth’s orbital parameters. These are often referred to as the Milankovitch cycles, after Milutin Milankovitch, a Serbian astronomer who argued in the 1920’s and 1930’s† that they caused the ice ages (Milankovitch, 1938).

Let’s explore Milankovitch’s idea in a bit more detail. The eccentricity (i.e., the degree to which the or-

†While Milankovitch was a strong and early proponent of the idea that variations in the Earth’s orbit caused ice ages, he was not the first to suggest it. J. Croll of Britian first suggested it in 1864, and published several subsequent papers on the subject.
bit differs from circular) of the Earth’s orbit about the Sun and the degree of tilt, or obliquity, of the Earth’s rotational axis vary slightly. In addition, the direction in which the Earth’s rotational axis tilts when it is closest to the Sun (perihelio) varies, a phenomenon called precession. These variations are illustrated in Figure 9.24. Though variation in these “Milankovitch parameters” has negligible effect on the total radiation the Earth receives, they do affect the pattern of incoming radiation (insolation). For example, tilt of the rotational axis determines seasonality and the latitudinal gradient of insolation. The gradient is extremely important because it drives atmospheric and oceanic circulation. If the tilt is small, seasonality will be reduced (cooler summers, warmer winters), and the average annual insolation gradient will be high. Precession relative to the eccentricity of the Earth’s orbit also affects seasonality. For example, presently the Earth is closest to the Sun in January. As a result, northern hemisphere winters (and southern hemisphere summers) are somewhat warmer than they would be otherwise. For a given latitude and season, precession will result in a ±5% difference in insolation. While the Earth’s orbit is only slightly elliptical, and variations in eccentricity are small, these variations are magnified because insolation varies with the inverse square of the Earth-Sun distance.

Variation in tilt approximates a simple sinusoidal function with a period of 41,000 yrs. Variations in eccentricity can be approximately described with period of 100,000 years. In actuality, variation in eccentricity is more complex, and is more accurately described with periods of 123,000 yrs, 85,000 yrs, and 58,000 yrs. Similarly, variation in precession has characteristic periods of 23,000 and 18,000 yrs.

Though Emiliani suggested that δ¹⁸O variations were related to variations in these orbital parameters, the first quantitative approach to the problem was that of Hayes et al. (1976). They applied Fourier analysis to the δ¹⁸O curve (Fourier analysis is a mathematical tool that transforms a complex...
variation such as that in Figure 9.23 to the sum of a series of simple sin functions). Hayes et al. then used spectral analysis to show that much of the spectral power of the $\delta^{18}O$ curve occurred at frequencies similar to those of the Milankovitch parameters. The most elegant and convincing treatment is that of Imbrie (1985). Imbrie’s treatment involved several refinements and extension of the earlier work of Hayes et al. (1976). First, he used improved values for Milankovitch frequencies. Second, he noted these Milankovitch parameters vary with time (the Earth’s orbit and tilt are affected by the gravitational field of the Moon and other planets, and other astronomical events, such as bolide impacts), and the climate system’s response to them might also vary over time. Thus Imbrie treated the first and second 400,000 years of Figure 9.22 separately.

Imbrie observed that climate does not respond instantaneously to forcing. For example, maximum temperatures are not reached in Ithaca, New York until late July, 4 weeks after the maximum insolation, which occurs on June 21. Thus there is a phase lag between the forcing function (insolation) and climatic response (temperature). Imbrie (1985) constructed a model for response of global climate (as measured by the $\delta^{18}O$ curve) in which each of the 6 Milankovitch forcing functions was associated with a different gain and phase. The values of gain and phase for each parameter were found statistically by minimizing the residuals of the power spectrum of the $\delta^{18}O$ curve in Figure 9.23. The resulting “gain and phase model” is shown in comparison with the data for the past 400,000 years and the next 25,000 years in Figure 9.25. The model has a correlation coefficient, $r$, of 0.88 with the data. Thus about $r^2$, or 77%, of the variation in $\delta^{18}O$, and therefore presumably in ice volume, can be explained by Imbrie’s Milankovitch model. The correlation for the period 400,000–782,000 yrs is somewhat poorer, around 0.80, but nevertheless impressive.

Since variations in the Earth’s orbital parameters do not affect the average total annual insolation the Earth receives, but only its pattern in space and time, one might ask how this could cause glaciation. The key factor seems to be the insolation received during summer at high northern latitudes. This is, of course, the area where large continental ice sheets develop. The southern hemisphere, except for Antarctica, is largely ocean, and therefore not subject to glaciation. Glaciers apparently develop when summers are not warm enough to melt the winter’s accumulation of snow in high northern latitudes.

Nevertheless, even at a given latitude the total variation in insolation is small, and not enough by itself to cause the climatic variations observed. Apparently, there are feedback mechanisms at work that serve to amplify the fundamental Milankovitch forcing function. One of these feedback mechanisms was identified by Agassiz, and that is ice albedo, or reflectance. Snow and ice reflect much of the incoming sunlight back into space. Thus as glaciers advance, they will cause further cooling. Any additional accumulation of ice in Antarctica, however, does not result in increased albedo, because the continent is fully ice covered even in interglacial times. Hence the dominant role of northern hemisphere insolation in driving climate cycles. Other possible feedback mechanisms include carbon dioxide and ocean circulation. The role of atmospheric CO$_2$ in controlling global climate is a particularly important issue because of burning of fossil fuels has resulted in a significant increase in atmospheric CO$_2$ concentration over the last 150 years. We will return to this issue in a subsequent chapter.

### 9.6.2 The Record in Glacial Ice

Climatologists recognized early on that continental ice preserves a stratigraphic record of climate change. Some of the first ice cores recovered for the purpose of examining the climatic record and analyzed for stable isotopes were taken from Greenland in the 1960’s (e.g., Camp Century Ice Core). Subsequent cores have been taken from Greenland, Antarctica, and various alpine glaciers. The most remarkable and useful of these cores has been the 2000 m core recovered by the Russians from the Vostok station in Antarctica (Jouzel, et al., 1987), and is compared with the marine $\delta^{18}O$ record in Figure 9.26. The marine record shown is the SPECMAP record, which is a composite record based on that of Imbrie et al. (1984), but with further modification of the chronology. The core provides a 160,000 year record of $\delta$D and $\delta^{18}O_{ice}$, as well as CO$_2$, and $\delta^{18}O_2$ in bubbles (the latter,
which was subsequently published and not shown in Figure 9.26, provides a measure of $\delta^{18}O$ in the atmosphere and, indirectly, the ocean, which corresponds to a full glacial cycle.

Jouzel et al. (1987) converted $\delta D$ to temperature variations after subtracting the effect of changing ice volume on $\delta D$ of the oceans. The conversion is based on a $6\%/{ }^\circ C$ relationship between $\delta D$ and temperature in modern Antarctic snow (they found a similar relationship using circulation models). The hydrogen isotopic fractionation of water is a more sensitive function of temperature than is oxygen fractionation. Their results, taken at face value, show dramatic $10^\circ C$ temperature variations between glacial and interglacial times.

Dating of the Vostok ice core was based only on an ice-flow model. Nevertheless, the overall pattern observed is in remarkable agreement the marine $\delta^{18}O$ record, particular from 110,000 years to the present. The record of the last deglaciation is particularly similar to that of the marine $\delta^{18}O$ record, and even shows evidence of a slight return trend toward glacial conditions from 12 kyr to 11 kyr BP, which corresponds to the well documented Younger Dryas event of the North Atlantic region. It is also very significant that spectral analysis of the Vostok isotope record shows strong peaks in variance at 41 kyr (the obliquity frequency), and at 25 kyr, which agree with the 23 kyr precessional frequency when the age errors are considered. Thus the Vostok ice core data appear to confirm the importance of Milankovitch climatic forcing. It is interesting and significant that even in this core, taken at $78^\circ S$, it is primarily insolation at $65^\circ N$ that is the controlling influence.

Figure 9.26. The upper curve shows $\delta D$ in ice from the Vostok ice core. The second curve shows the calculated temperature difference relative to the present mean annual temperature, and the lower curve shows $\delta^{18}O$ for seawater based on carbonate sediments. (From Jouzel, et al., 1987).
There are, however, significant differences between the Vostok record and the marine record. Some of these are probably due to inaccuracies in dating; others may reflect differences in northern and southern hemisphere response to orbital forcing.

9.6.3 Soils and Paleosols

As we found in Chapter 6, the concentration of CO$_2$ in soils is very much higher than in the atmosphere, reaching 1% by volume. As a result, soil water can become supersaturated with respect to carbonates. In soils where evaporation exceeds precipitation, soil carbonates form. The carbonates form in equilibrium with soil water, and there is a strong correlation between $\delta^{18}$O in soil carbonate and local meteoric water, though soil carbonates tend to be about 5‰ more enriched than expected from the calcite-water fractionation. There are 2 reasons why soil carbonates are heavier. First, soil water is enriched in $^{18}$O relative to meteoric water due to preferential evaporation of isotopically light water molecules. Second, rain (or snow) falling in wetter, cooler seasons in more likely to run off than that falling during warm seasons. Taking these factors into consideration, the isotopic composition of soil carbonates may be used as a paleoclimatic indicator.

Figure 9.27 shows one example of $\delta^{18}$O in paleosol carbonates used in this way. The same Pakistani paleosol samples analyzed by Quade et al. (1989) for $\delta^{13}$C (Figure 9.19) were also analyzed for $\delta^{18}$O. The $\delta^{13}$C values recorded a shift toward more positive values at 7 Ma that apparently reflect the appearance of C$_4$ grasslands. The $\delta^{18}$O shows a shift to more positive values at around 8 Ma, or a million years before the $\delta^{13}$C shift. Quade et al. interpreted this as due to an intensification of the Monsoon system at that time, an interpretation consistent with marine paleontological evidence.

Clays, such as kaolinites, are another important constituent of soil. Lawrence and Taylor (1972) showed that during soil formation, kaolinite and montmorillonite form in approximate equilibrium with meteoric water so that their $\delta^{18}$O values are systematically shifted by +27 ‰ relative to the local meteoric water, while $\delta$D are shifted by about 30‰. Thus kaolinites and montmorillonites define a line parallel to the meteoric water line (Figure 9.28), the so-called kaolinite line. From this observation, Sheppard et al. (1969) and Lawrence and Taylor (1972) reasoned that one should be able to deduce the isotopic composition of rain at the time ancient kaolinites formed from their $\delta$D values. Since the isotopic composition of precipitation is climate dependent, ancient kaolinites provide another continental paleoclimatic record.
There is some question as to the value of hydrogen isotopes in clays for paleoclimatic studies. Lawrence and Meaux (1993) concluded that most ancient kaolinites have exchanged hydrogen subsequent to their formation, and therefore are not a good paleoclimatic indicator. Others argue that clays do not generally exchange hydrogen unless recrystallization occurs or the clay experiences temperatures in excess of 60-80°C for extended periods (A. Gilg, pers. comm., 1995).

Lawrence and Meaux (1993) concluded, however, that oxygen in kaolinite does preserve the original δ18O, and that can, with some caution, be used as a paleoclimatic indicator. Figure 9.29 compares the δ18O of ancient Cretaceous North American kaolinites with the isotopic composition of modern precipitation. If the Cretaceous climate were the same as the present one, the kaolinites should be systematically 27‰ heavier than modern precipitation. For the southeastern US, this is approximately true, but the difference is generally less than 27‰ for other kaolinites, and the difference decreases northward. This indicates these kaolinites formed in a warmer environment than the present one. Overall, the picture provided by Cretaceous kaolinites confirm what has otherwise be deduced about Cretaceous climate: the Cretaceous climate was generally warmer, and the equator to pole temperature gradient was lower. A drawback with this approach, however, is the lack of good ages on paleosols, which have proven to be very difficult to date precisely. The ages of many paleosols used by Lawrence and Meaux (1993) were simply inferred from stratigraphy.

9.7 Hydrothermal Systems and Ore Deposits

When large igneous bodies are intruded into high levels of the crust, they heat the surrounding rock and the water in the cracks and pore in this rock, setting up convection systems. The water in these hydrothermal systems reacts with the hot rock and undergoes isotopic exchange; the net result is that both the water and the rock change their isotopic compositions.

One of the first of many important contributions of stable isotope geochemistry to understanding hydrothermal systems was the demonstration by Harmon Craig (another student of Harold Urey) that water in these systems was meteoric, not magmatic (Craig, 1963). The argument is based upon the data shown in Figure 9.30. For each geothermal system, the δD of the “chloride” type geothermal waters is the same as the local precipitation and groundwater, but the δ18O is shifted to higher values. The shift in δ18O results from high temperature (~< 300°C) reaction of the local meteoric water with hot rock. However, because the rocks contain virtually no hydrogen, there is little change in the hydrogen isotopic composition of the water. If the water involved in these systems were magmatic, it would not have the same isotopic composition as local meteoric water (it is possible that these systems contain a few percent magmatic water).

Acidic, sulfur-rich waters from hydrothermal systems can have δD that is different from local meteoric water. This shift occurs when hydrogen isotopes are fractionated during boiling of geothermal waters. The steam produced is enriched in sulfide. The steam mixes with cooler meteoric water, condenses, and the sulfide is oxidized to sulfate, resulting in their acidic character. The mixing lines observed reflect mixing of the steam with meteoric water as well as the fractionation during boiling.
Estimating temperatures at which ancient hydrothermal systems operated is a fairly straightforward application of isotope geothermometry, which we have already discussed. If we can measure the oxygen (or carbon or sulfur) isotopic composition of any two phases that were in equilibrium, and if we know the fractionation factor as a function of temperature for those phases, we can estimate the temperature of equilibration. We will focus now on water-rock ratios, which may also be estimated using stable isotope ratios.

### 9.7.1 Water-Rock Ratios

For a closed hydrothermal system, we can write two fundamental equations. The first simply describes isotopic equilibrium between water and rock:

\[
\Delta = \delta^{w}_f - \delta^{r}_f \tag{9.59}
\]

where we use the subscript \( w \) to indicate water, and \( r \) to indicate rock. The superscript \( f \) indicates the final value. So 9.59 just says that the difference between the final isotopic composition of water and rock is equal to the fractionation factor (we implicitly assume equilibrium). The second equation is the mass balance for a closed system:

\[
c^{w}_w W^{i} \delta^{i}_w + c^{r}_r R^{i} \delta^{i}_r = c^{w}_w W^{f} \delta^{f}_w + c^{r}_r R^{f} \delta^{f}_r
\]

where \( c \) indicates concentration (we assume concentrations do not change, which is valid for oxygen, but perhaps not valid for other elements), \( W \) indicates the mass of water involved, \( R \) the mass of rock involved, the superscript \( i \) indicates the initial value and \( f \) again denotes the final isotope ratio. This equation just says the amount of an isotope present before reaction must be the same as after reaction. We can combine these equations to produce the following expression for the water-rock ratio:

\[
\frac{W}{R} = \frac{\delta^{f}_w - \delta^{i}_r}{\delta^{i}_w - \delta^{i}_r} \frac{c^{r}_r}{c^{w}_w}
\]

The initial \( \delta^{18}O \) can generally be inferred from unaltered samples and from the \( \delta^D-\delta^{18}O \) meteoric water line and the final isotopic composition of the rock can be measured. The fractionation factor can be estimated if we know the temperature and the phases in the rock. For oxygen, the ratio of concentration in the rock to water will be close to 0.5 in all cases.

Equation 9.61 is not very geologically realistic because it applies to a closed system. A completely open system, where water makes one pass through hot rock, would be more realistic. In this case, we might suppose that a small parcel of water, \( dW \), passes through the system and induces an incremental change in the isotopic composition of the rock, \( d\delta \). In this case, we can write:
W. M. White

Chapter 9: Stable Isotopes

\[ Rc_\delta \delta = \delta_w - (\Delta + \delta_r) c_w dW \]  
9.62

This equation states that the mass of an isotope exchanged by the rock is equal to the mass of that isotope exchanged by the water (we have substituted \( \Delta + \delta_r \) for \( \delta_r \)). Rearranging and integrating, we have:

\[ \frac{W}{R} = \ln \left( \frac{\delta_i^f - \delta_i^w}{-\delta_i^r + \delta_i^w - \Delta} + 1 \right) \frac{c_w}{c_r} \]  
9.63

Thus it is possible to deduce the water rock ratio for an open system as well as a closed one.

Oxygen isotope studies can be a valuable tool in mineral exploration. Mineralization is often (though not exclusively) associated with the region of greatest water flux, such as areas of upward moving hot water above intrusions. Such areas are likely to have the lowest values of \( \delta^{18}O \). To understand this, let’s solve equ. 9.63 for \( \delta^f_r \), the final value of \( \delta^{18}O \) in the rock:

\[ \delta^f_r = (\delta^i_r - \delta^i_w + \Delta) e^{-\frac{Wc_w}{Rc_r}} + \delta^i_w + \Delta \]  
9.64

Assuming a uniform initial isotopic composition of the rocks and the water, all the terms on the r.h.s. are constants except \( W/R \) and \( \Delta \), which is a function of temperature. Thus the final values of \( \delta^{18}O \) are functions of the temperature of equilibration, and an exponential function of the \( W/R \) ratio. Figure 9.31 shows \( \delta^{18}O^f_r \) plotted as a function of \( W/R \) and \( \Delta \), where \( \delta^{18}O^i_r \) is assumed to be +6 and \( \delta^{18}O^i_w \) is assumed to be –13. In a few cases, ores apparently have precipitated by fluids derived from the magma itself. In those cases, this kind of approach does not apply.

Figure 9.32 shows an example of the \( \delta^{18}O \) imprint of an ancient hydrothermal system: the Bohemia Mining District in Lane Country, Oregon, where Tertiary volcanic rocks of the western Cascades have been intruded by a series of dioritic plutons. Approximately $1,000,000 worth of gold was removed from the region between 1870 and 1940. \( \delta^{18}O \) contours form a bull’s eye pattern, and the region of low \( \delta^{18}O \) corresponds roughly with the area of propylitic (i.e., greenstone) alteration. Notice that this region is broader than the contact metamorphic aureole. The primary area of mineralization occurs within the \( \delta^{18}O < 0 \) contour. In this area, relatively large volumes of gold-bearing hydrothermal solution cooled, perhaps due to mixing with groundwater, and precipitated gold. This is an excellent example of the value of oxygen isotope studies to mineral exploration. Similar bull’s eye patterns are found around many other hydrothermal ore deposits.

9.7.2 Sulfur Isotopes and Ore Deposits

A substantial fraction of base and noble metal ores are sulfides. These have formed in a great variety of environments and under a great variety of conditions. Sulfur isotope studies have been very valuable in sorting out the genesis of these deposits. Of the various stable isotope systems we will consider here, sulfur isotopes are undoubtedly the most complex. This complexity arises in part because there are four common valence states in which sulfur can occur in the Earth, +6 (e.g., \( \text{BaSO}_4 \)), +4 (e.g., \( \text{SO}_2 \)), 0 (e.g., \( \text{S} \)), –1 (e.g., \( \text{FeS}_2 \)) and –2 (\( \text{H}_2\text{S} \)). Significant equilibrium isotopic fractionations occur between these valence states. Each of these valence states forms a variety of compounds, and fractionations can occur between these as well (e.g., Figure 9.7). Finally, sulfur is im-
important in biological processes and fractionations in biologically mediated oxidations and reductions are often different from fractionations in the abiological equivalents.

There are two major reservoirs of sulfur on the Earth that have uniform sulfur isotopic compositions: the mantle, which has $\delta^{34}S$ of ~0 and in which sulfur is primarily present in reduced form, and seawater, which has $\delta^{34}S$ of +20* and in which sulfur is present as $SO_{4}^{2-}$. Sulfur in sedimentary, metamorphic, and igneous rocks of the continental crust may have $\delta^{34}S$ that is both greater and smaller than these values (Figure 9.33). All of these can be sources of sulfide in ores, and further fractionation may occur during transport and deposition of sulfides. Thus the sulfur isotope geochemistry of sulfide ores is remarkably complex.

At magmatic temperatures, reactions generally occur rapidly and most systems appear to be close to equilibrium. Below 200°C, however, sulfur isotopic equilibration is slow even on geologic time scales, hence equilibration is rare and kinetic effects often dominate. Isotopic equilibration between two sulfide species or between two sulfate species is achieved more readily than between sulfate and sulfide species. Sulfate-sulfide reaction rates have been shown to depend on pH (reaction is more rapid at low pH). In addition, equilibration is much more rapid when sulfur species of intermediate valence species are present. Presumably, this is because reaction rates between species of adjacent valence states (e.g., sulfate and sulfite) are rapid, but reaction rates between widely differing valence states (e.g., sulfate and sulfide) are much slower.

Mississippi Valley type deposits provide an interesting example of how sulfur isotopes can be used to understand ore genesis. These are carbonate-hosted lead and zinc sulfides formed under relatively low temperature conditions. Figure 9.34 shows the sulfur isotope ratios of some examples.

*This is the modern value for seawater. As we shall see in a subsequent chapter, though the sulfur isotopic composition of seawater is uniform at any time, this value has varied with time.
They can be subdivided into Zn-rich and Pb-rich classes. The Pb-rich and most of the Zn rich deposits were formed between 70 and 120° C, while some of the Zn-rich deposits, such as those of the Upper Mississippi Valley, were formed at temperatures up to 200° C, though most were formed in the range of 90° to 150° C.

Figure 9.35 illustrates a generalized model for the genesis of Mississippi Valley type deposits. In most instances, metals and sulfur appear to have been derived distant from sedimentary units, perhaps particularly from evaporites, by hot, deep-circulating formation water. In North America, most of these seem to have formed during or shortly after the late Paleozoic Appalachian-Ouchita-Marathon Orogeny. The direct cause of precipitation of the ore probably may have differed in the different localities. Oxygen and hydrogen isotope data suggest that mixing of the hot saline fluids with cooler low salinity ground water was probably the immediate cause of metal precipitation in some instances. In others, such as Pine-Point, local reduction of sulfate in the fluids to sulfide may have cause precipitation (Ohmoto, 1986). Local production of sulfide shortly before ore deposition may help to account for the lack of isotopic equilibrium in this deposit, since time is an element in the attainment of isotopic equilibrium. In other cases, mixing of separate S-bearing and metal-bearing fluids from different aquifers may have been the cause.

Figure 9.34. Sulfur isotope ratios in some Mississippi Valley-type Pb and Zn deposits. After Ohmoto and Rye (1979).

Figure 9.35. Cartoon illustrating the essential features of the genesis of Mississippi Valley sulfide ores. After Ohmoto (1986).
The sulfur isotope data suggest there are a number of possible sources of sulfur. In many cases, sulfur was apparently derived sulfate in evaporites, in others sulfur was derived from brines produced by evaporative concentration of seawater. In many cases, the sulfate was reduced by reaction with organic matter at elevated temperature (thermochemical reduction). In other cases, it may have been bacterially reduced. Other sulfur sources include sulfide from petroleum. In at least some deposits, sulfur appears to have been derived from more than one of these sources.

Sulfur isotope ratios in the S. E. Missouri district are quite variable and \( \delta^{34}S \) is correlated with Pb isotope ratios in galenas. This suggests there was more than one source of the sulfur. Based on a combined Pb and S isotopic study, Goldhaber et al. (1995) concluded that the main stage ores of the S. E. Missouri district were produced by mixing of fluids from several aquifers. They concluded that Pb was derived from the Lamont Sandstone (which directly overlies basement). Fluid in this aquifer was maintained at relatively high p\( \varepsilon \) by the presence of hematite, and hence had low sulfide concentrations. This allowed leaching and transport of the Pb. Isotopically heavy S was produced by dissolution of evaporite sulfate and thermochemical reduction and migrated through the Sillivan Siltstone member of the upper Bonneterre Formation, which overlies the Lamont Sandstone. Both Pb and isotopically light sulfur migrated through the lower part of the Bonneterre Formation. PbS precipitated when the fluids were forced to mix by pinchout of the Sillivan Siltstone.

9.8 Stable Isotopes in the Mantle and Magmatic Systems

9.8.1 Stable Isotopic Composition of the Mantle

The mantle is the largest reservoir for oxygen, and may be for C, H, N, and S as well. Thus we need to know the isotopic composition of the mantle to know the isotopic composition of the Earth. Variations in stable isotope ratios in mantle and mantle-derived materials also provide important insights on mantle and igneous processes.

9.8.1.1 Oxygen

Assessing the oxygen isotopic composition of the mantle, and particularly the degree to which its oxygen isotope composition might vary, has proved to be more difficult than expected. One approach has been to use basalts as samples of mantle, as is done for radiogenic isotopes. Isotope fractionation occurring during partial melting are small, so the oxygen isotopic composition of basalt should be similar to that of its mantle source. However, assimilation of crustal rocks by magmas and oxygen isotope exchange during weathering complicate the situation. An alternative is to use direct mantle samples such as xenoliths occasionally found in basalts. However, these are considerably rarer than are basalts.

Figure 9.36 shows the oxygen isotope composition of olivines and clinopyroxenes in 76 peridotite xenoliths analyzed by Mattey et al. (1994) using the laser fluorination technique. The total range of values observed is only about twice that expected from analytical error alone, suggesting the mantle is fairly homogeneous in its isotopic composition. The difference between co-existing olivines and clinopyroxenes averages about 0.5 per mil, which is consistent with the expected fractionation between these minerals at mantle temperatures. Mattey et al. (1994) estimated the bulk composition of these samples to be about +5.5 per mil.

Figure 9.37 shows the distribution of \( \delta^{18}O \) in basalts from 4 different groups. To avoid weathering problems, Harmon and Hoefs (1995) included only submarine basaltic glasses and basalts having less than 0.75% water or have erupted historically in their compilation. There are several points worth noting in these data.

MORB have a mean \( \delta^{18}O_{SMOW} \) of +5.7%, with a relatively smaller variation about this mean. Thus the depleted upper mantle appears to be a relatively homogeneous reservoir for oxygen. The homogeneity observed in MORB is consistent with that observed in mantle-derived xenoliths. The small difference between the means of the two groups (0.2%), may well reflect fractionation occurring during partial melting.
The other groups show significantly greater heterogeneity than either MORB or peridotite xenoliths. Oceanic island basalts, which presumably sample mantle plumes, have a mean identical to that of peridotite xenoliths, but are more variable. Subduction-related basalts (i.e., island arc basalts and their continental equivalents) are shifted to more positive $\delta^{18}$O values, as are continental intraplate volcanics.

Whether the variability in these groups reflects mantle heterogeneity, the melting process, assimilation, or merely weathering remains unclear. Hawaii is the one locality where high quality data exists for both basalts and xenoliths and comparisons can be made. Both Hawaiian basalts and olivines in Hawaiian xenoliths (Figure 5.36) have lower $\delta^{18}$O than their equivalents from elsewhere. This suggests the possibility, at least, that part of the variability in oceanic basalts reflects mantle heterogeneity. However, additional studies will be required to establish this with certainty.

9.8.1.2 Hydrogen

Estimating the isotopic composition of mantle hydrogen, carbon, nitrogen, and sulfur is even more problematic. These are all volatile elements and are present at low concentrations in mantle materials. They partitioning partially or entirely into the gas phase of magmas upon their eruption. This gas phase is lost, except for deep submarine eruption. Furthermore, $C$, $N$, and $S$ have several oxidation states, and isotopic fractionations occur between the various compounds these elements form (e.g., CO$_2$, CO, CH$_4$, H$_2$, H$_2$O). This presents two problems. First, significant fractionations can occur during degassing, even at magmatic temperatures. Second, because of loss of the gas phase, the concentrations of these elements are low. Among other problems, this means their isotope ratios are subject to disturbance by contamination. Thus the isotopic composition of these elements in igneous rocks do not necessarily reflect those of the magma or its mantle source.

Hydrogen, which is primarily present as water, but also as $H_2$ $H_2S$, and CH$_4$ can be lost from magmas during degassing. However, basalts erupted beneath a kilometer or more of ocean retain most of their dissolved water. Thus mid-ocean ridge basalts and basalts erupted on seamounts are
important sources of information of the abundance and isotopic composition of hydrogen in the mantle.

As Figure 9.38 indicates, MORB have a mean $\delta D_{SMOW}$ of about $-67.5\%$ and a standard deviation of $\pm 14\%$. How much of this variability reflects fractionation during degassing and contamination is unclear. Kyser (1986) has argued that mantle hydrogen is isotopically homogeneous with $\delta D_{SMOW}$ of $-80\%$. The generally heavier isotopic composition of MORB, he argues, reflects H$_2$O loss and other processes. Others, for example, Poreda et al. (1986), Chaussidon et al. (1991) have observed correlations between $\delta D$ and Sr and Nd isotope ratios and have argued that these provide clear evidence of that mantle hydrogen is isotopically heterogeneous. Chaussidon et al. (1991) have suggested a $\delta D$ value for the depleted upper mantle of about $-55\%$.

The first attempt to assess the hydrogen isotopic composition of the mantle materials was that of Sheppard and Epstein (1970), who analyzed hydrous minerals in xenoliths and concluded $\delta D$ was variable in the mantle. Since then, a number of additional studies of these materials have been carried out. As Figure 9.37 shows, phlogopites have $\delta D$ that is generally similar to that of MORB, though heavier values also occur. Amphiboles have much more variable $\delta D$ and have heavier hydrogen on average. Part of this difference probably reflects equilibrium fractionation. The fractionation between water and phlogopite is close to 0$\%$ in the temperature range 800°-1000°C, whereas the fractionation between water and amphibole is about $-15\%$. However, equilibrium fractionation alone cannot explain either the variability of amphiboles or the difference between the mean $\delta D$ of phlogopites and amphiboles. Complex processes that might include Rayleigh distillation may be involved in the formation of mantle amphiboles. This would be consistent with the more variable water content of amphiboles compared to

![Figure 9.37. $\delta ^{18}O$ in young, fresh basalts. Dashed line is at the mean of MORB (+5.7). After Harmon and Hoefs (1995).](image1)

![Figure 9.38. $\delta D$ in MORB and in mantle phlogopites and amphiboles. The MORB and phlogopite data suggest the mantle has $\delta D_{SMOW}$ of about $-60$ to $-90$.](image2)
phlogopites. There are also clear regional variations in \( \delta D \) in xenoliths that argue for large scale heterogeneity. For example, Deloule et al. (1991) found that \( \delta D \) in amphiboles from xenoliths in basalts in the Massif Central of France have systematically high \( \delta D \) (-59 to -28) while those from Hawaii can be very low (-125). At the opposite extreme, Deloule et al. (1991) also observed isotopic heterogeneity within single crystals, which they attribute to incomplete equilibration with magmas or fluids.

### 9.8.1.3 Carbon

Most carbon in basalts is in the form of \( \text{CO}_2 \), which has limited solubility in basaltic liquids. As a result, basalts begin to exsolve \( \text{CO}_2 \) before they erupt. Thus virtually every basalt, including those erupted at mid-ocean ridges, has lost some carbon, and subareal basalts have lost virtually all carbon (as well as most other volatiles). Therefore only basalts erupted beneath several km of water provide useful samples of mantle carbon, so the basaltic data set is essentially restricted to MORB and samples recovered from seamounts and the submarine part of Kilauea’s East Rift Zone. The question of the isotopic composition of mantle carbon is further complicated by fractionation and contamination. There is a roughly 4‰ fractionation between \( \text{CO}_2 \) dissolved in basaltic melts and the gas phase, with \( ^{13}\text{C} \) enriched in the gas phase. Carbonatites and diamonds provide alternative, and generally superior, samples of mantle carbon, but their occurrence is extremely restricted.

MORB have a mean \( \delta^{13}\text{C} \) of -6.5‰ (Figure 9.39), but the most \( \text{CO}_2 \)-rich MORB samples have \( \delta^{13}\text{C} \) of about -4‰. Since they are the least degassed, they presumably best represent the isotopic composition of the depleted mantle (Javoy and Pineau, 1991). Carbonatites also have a mean \( \delta^{13}\text{C} \) close to -4‰. Oceanic island basalts erupted under sufficient water depth to preserve some \( \text{CO}_2 \) in the vesicles appear to have similar isotopic compositions. Gases released in subduction zone volcanos and back-arc basin basalts (Figure 9.39), which erupt behind subduction zones and are often geochemically similar to island arc basalts, have carbon that can be distinctly lighter (lower \( \delta^{13}\text{C} \)), though most \( \delta^{13}\text{C} \) values are in the range of ~2 to ~4‰, comparable to the most gas-rich MORB. Carbon in mantle diopsides (clinopyroxenes) appears to be isotopically lighter on average, and the significance of this is unclear.

Diamonds show a large range of carbon isotopic compositions (Figure 9.39). Most diamonds have \( \delta^{13}\text{C} \) within the range of -2 to -8‰, hence similar to MORB. However, some diamonds have much lighter carbon. Based on the inclusions they contain, diamonds can be divided between peridotitic and eclogitic. Most peridotitic diamonds have \( \delta^{13}\text{C} \) close to -5‰, while eclogitic diamonds are much more isotopically variable. Most, though not all, of the diamonds with very negative \( \delta^{13}\text{C} \) are eclogitic. Many diamonds are isotopically zoned, indicating they grew in several stages.

Three hypotheses have been put forward to explain the isotopic heterogeneity in diamonds: primordial heterogeneity, fractionation effects, and recycling of organic carbon from the Earth’s surface into the mantle. Primordial heterogeneity seems unlikely for a number of reasons. Among these is the absence of very negative \( \delta^{13}\text{C} \) in other materials, such as MORB, and the absence of any

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**Figure 9.39.** Carbon isotope ratios in mantle (red) and mantle-derived materials (gray). After Mattey (1987).

<table>
<thead>
<tr>
<th>Material</th>
<th>Carbon isotopic compositions</th>
</tr>
</thead>
<tbody>
<tr>
<td>Diamonds</td>
<td>( n=1578 )</td>
</tr>
<tr>
<td>Carbonatites</td>
<td>( n=254 )</td>
</tr>
<tr>
<td>Diopsides</td>
<td>( n=32 )</td>
</tr>
<tr>
<td>MORB</td>
<td>( n=40 )</td>
</tr>
<tr>
<td>OIB</td>
<td>( n=16 )</td>
</tr>
<tr>
<td>BABB</td>
<td>( n=7 )</td>
</tr>
</tbody>
</table>

\( \delta^{13}\text{C}_{\text{PDB}} \)
evidence for primordial heterogeneity from the isotopic compositions of other elements. Boyd and Pillinger (1994) have argued that since diamonds are kinetically sluggish (witness their stability at the surface of the Earth, where they are thermodynamically out of equilibrium), isotopic equilibrium might not achieved during their growth. Large fractionations might therefore occur due to kinetic effects. However, these kinetic fractionations have not been demonstrated, and fractionations of this magnitude (20‰ or so) would be surprising at mantle temperatures.

On the other hand, several lines of evidence support the idea that isotopically light carbon in some diamonds had its origin as organic carbon at the Earth’s surface. First, such diamonds are primarily of eclogitic paragenesis, and eclogite is the high pressure equivalent of basalt. Subduction of oceanic crust continuously carries large amounts of basalt into the mantle. Oxygen isotope heterogeneity observed in some eclogite xenoliths suggests these eclogites do indeed represent subducted oceanic crust. Second, the nitrogen isotopic composition of isotopically light diamonds are anomalous relative to nitrogen in other mantle materials yet similar to nitrogen in sedimentary rocks.

9.8.1.4 Nitrogen

Figure 9.40 summarizes the existing data on the nitrogen ratios in the crust and mantle. There is far less data than there is for other stable isotope ratios because of the low concentrations and pervasive contamination problems. The solubility of N₂ in basalts is very limited, though much of the nitrogen may be present as NH₄⁺, which is somewhat more soluble. Hence of volcanic rocks, once again only submarine basalts provide useful samples of mantle N. There are both contamination and analytical problems with determining nitrogen isotope ratios in basalts, which, combined with its low abundance (generally less than a ppm), mean that accurate measurements are difficult to make. Measurements of δ¹⁵Nᵦ in MORB range from about −2 to +12‰. The few available analyses of Hawaiian basalts range up to +20. At present, it is very difficult to decide to what degree this variation reflects contamination (particularly by organic matter), fractionation during degassing, or real mantle heterogeneity. Perhaps all that can be said is that nitrogen in basalts appears to have positive δ¹⁵N on average.

Diamonds can contain up to 2000 ppm of N and hence provide an excellent sample of mantle N. As can be seen in Figure 9.40, high δ¹³C diamonds (most common, and usually of peridotitic paragenesis) have δ¹⁵N that range from -12 to +5 and average about -3‰, which contrasts with the generally positive values observed in basalts. Low δ¹³C diamonds have generally positive δ¹⁵N. Since organic matter and ammonia in crustal rocks also typically have positive δ¹⁵N, this characteristic is consistent with the hypothesis that this group of diamonds are derived from subducted crustal material. However, since basalts appear to have generally positive δ¹⁵N, other interpretations are also possible.
Fibrous diamonds, whose growth may be directly related to the kimberlite eruptions that carry them to the surface (Boyd et al., 1994), have more uniform $\delta^{15}N$, with a mean of about -5‰. Since there can be significant isotopic fractionations involved in the incorporation of nitrogen into diamond, the meaning of the diamond data is also uncertain, and the question of the nitrogen isotopic composition of the mantle remains an open one.

9.8.1.5 Sulfur

There are also relatively few sulfur isotope measurements on basalts, in part because sulfur is lost during degassing, except for those basalts erupted deeper than 1 km below sealevel. In the mantle, sulfur is probably predominantly or exclusively in the form of sulfide, but in basalts, which tend to be somewhat more oxidized, some of it may be present as SO$_2$ or sulfate. Equilibrium fractionation should lead to SO$_2$ being a few per mil lighter than sulfate. If H$_2$S is lost during degassing the remaining sulfur would become heavier; if SO$_2$ or SO$_4$ is lost, the remaining sulfur would become lighter. Total sulfur in MORB has $\delta^{34}$S$_{CDT}$ in the range of +1.3 to -1‰, with most values in the range 0 to +1‰. Sakai et al. (1984) found that sulfate in MORB, which constitutes 10-20% of total sulfur, was 3.5 to 9‰ heavier than sulfide. Basalts from Kilauea’s East Rift Zone have a very restricted range of $\delta^{34}$S of +0.5 to +0.8 (Sakai, et al., 1984) .

Chaussidon et al. (1989) analyzed sulfides present as inclusions in minerals, both in basalts and in xenoliths, and found a wide range of $\delta^{34}$S (−5 to +8‰). Low Ni sulfides in oceanic island basalts, kimberlites, and pyroxenites had more variable $\delta^{34}$S that sulfides in peridotites and peridotite minerals. They argued there is a fractionation of +3‰ between sulfide liquid and sulfide dissolved in silicate melt. Carbonatites have $\delta^{34}$S between +1 and −3‰ (Hoefs, 1986; Kyser, 1986). Overall, it appears the mantle has a mean $\delta^{34}$S in the range of 0 to +1‰, which is very similar to meteorites, which average about +0.1‰.

Chaussidon, et al. (1987) found that sulfide inclusions in diamonds of peridotitic paragenesis ($\delta^{13}$C ~ −4‰) had $\delta^{34}$S of about +1‰ while eclogitic diamonds had higher, and much more variable $\delta^{34}$S (+2 to +10‰). Eldridge et al. (1991) found that $\delta^{34}$S in diamond inclusions was related to the Ni content of the sulfide. High Ni sulfide inclusions, which they argued were of peridotitic paragenesis, had $\delta^{34}$S between +4‰ and −4‰. Low Ni sulfides, which are presumably of eclogitic paragenesis, had much more variable $\delta^{34}$S (+14‰ to −10). These results are consistent with the idea that eclogitic diamonds are derived from subducted crustal material.

9.8.2 Stable Isotopes in Crystallizing Magmas

The variation in stable isotope composition produced by crystallization of a magma depends on the manner in which crystallization proceeds. The simplest, and most unlikely, case is equilibrium crystallization. In this situation, the crystallizing minerals remain in isotopic equilibrium with the melt until crystallization is complete. At any stage during crystallization, the isotopic composition of a mineral and the melt will be related by the fractionation factor, $\alpha$. Upon complete crystallization, the rock will have precisely the same isotopic composition as the melt initially had. At any time during the crystallization, the isotope ratio in the remaining melt will be related to the original isotope ratio as:

$$\frac{R_i}{R_o} = \frac{1}{f + \alpha(1-f)}; \quad \alpha \equiv \frac{R_s}{R_l}$$

where $R_l$ is the ratio in the liquid, $R_s$ is the isotope ratio of the solid, $R_o$ is the isotope ratio of the original magma, $f$ is the fraction of melt remaining. This equation is readily derived from mass balance, the definition of $\alpha$, and the assumption that the O concentration in the magma is equal to that in the crystals; an assumption valid to within about 10%. It is more convenient to express 9.65, in terms of $\delta$:

$$\Delta = \delta_{\text{melt}} - \delta_0 \equiv \left[ \frac{1}{f + \alpha(1-f)} - 1 \right] \times 1000$$
where $\delta_{\text{melt}}$ is the value of the magma after a fraction $f\text{-}1$ has crystallized and $\delta_{\text{o}}$ is the value of the original magma. For silicates, $\alpha$ is not likely to much less than 0.998 (i.e., $\Delta = \delta^{18}\text{O}_{\text{tals}} - \delta^{18}\text{O}_{\text{melt}} >= -2$). For $\alpha = 0.999$, even after 99% crystallization, the isotope ratio in the remaining melt will change by only 1 per mil.

The treatment of fractional crystallization is analogous to Rayleigh distillation. Indeed, it is governed by the same equation:

$$\Delta = 1000(f^\alpha - 1)$$  \hspace{1cm} (9.46)

The key to the operation of either of these processes is that the product of the reaction (vapor in the case of distillation, crystals in the case of crystallization) is only instantaneously in equilibrium with the original phase. Once it is produced, it is removed from further opportunity to attain equilibrium with original phase. This process is more efficient at producing isotopic variations in igneous rocks, but its effect remains limited because $\alpha$ is generally not greatly different from 1. Figure 9.41 shows calculated change in the isotopic composition of melt undergoing fractional crystallization for various values of $\Delta$ ($= 1000(\alpha-1)$). In reality, $\Delta$ will change during crystallization because of (1) changes in temperature (2) changes in the minerals crystallizing, and (3) changes in the liquid composition. The changes will generally mean that the effective $\Delta$ will increase as crystallization proceeds. We would expect the greatest isotopic fractionation in melts crystallizing non-silicates such as magnetite and melts crystallizing at low temperature, such as rhyolites, and the least fractionation for melts crystallizing at highest temperature, such as basalts.

Figure 9.41. Plot of $\delta^{18}\text{O}$ versus fraction of magma solidified during Rayleigh fractionation, assuming the original $\delta^{18}\text{O}$ of the magma was +6. After Taylor and Sheppard (1986).

Figure 9.42. $\delta^{18}\text{O}$ as a function of $\text{SiO}_2$ in a tholeiitic suite from the Galapagos Spreading Center (GSC) (Muehlenbachs and Byerly, 1982) and an alkaline suite from Ascension Island (Sheppard and Harris, 1985). Dashed line shows model calculation for the Ascension suite.

Figure 9.42. $\delta^{18}\text{O}$ as a function of $\text{SiO}_2$ in a tholeiitic suite from the Galapagos Spreading Center (GSC) (Muehlenbachs and Byerly, 1982) and an alkaline suite from Ascension Island (Sheppard and Harris, 1985). Dashed line shows model calculation for the Ascension suite.
which suggests a typical fractionation during melting of 0.2‰ or less.

We can generalize the temperature dependence of oxygen isotope fractionations by saying that at low temperature (i.e., ambient temperatures at the surface of the Earth up to the temperature of hydrothermal systems, 300-400°C), oxygen isotope ratios are changed by chemical processes. The amount of change can be used as an indication of the nature of the process involved, and, under equilibrium conditions, of the temperature at which the process occurred. At high temperatures (temperatures of the interior of the Earth or magmatic temperatures), oxygen isotope ratios are minimally affected by chemical processes and can be used as tracers much as radiogenic isotope ratios are.

These generalizations lead to an axiom: igneous rocks whose oxygen isotopic compositions show significant variations from the primordial value (+6) must either have been affected by low temperature processes, or must contain a component that was at one time at the surface of the Earth (Taylor and Sheppard, 1986).

9.8.3 Combined Fractional Crystallization and Assimilation

Because oxygen isotope ratios of mantle-derived magmas are reasonably uniform (±1‰ of 5.6‰) and generally different from rocks that have equilibrated with water at the surface of the Earth, oxygen isotopes are a useful tool in identifying and studying the assimilation of country rock by intruding magma. We might think of this process as simple mixing between two components: magma and country rock. In reality, it is always at least a three component problem, involving country rock, magma, and minerals crystallizing from the magma. Magmas are essentially never superheated; hence the heat required to melt and assimilate surrounding rock can only come from the latent heat of crystallization of the magma. Approximately 1 kJ/g would be required to heat rock from 150°C to 1150°C and another 300 J/g would be required to melt it. If the latent heat of crystallization is 400 J/g, crystallization of 3.25 g of magma would be required to assimilate 1 g of country rock. Since some heat will be lost by simple conduction to the surface, we can conclude that the amount of crystallization will inevitably be greater than the amount of assimilation (the limiting case where mass crystallized equals mass assimilated could occur only at great depth in the crust where the rock is at its melting point to begin with). The change in isotopic composition of a melt undergoing the combined process of assimilation and fractional crystallization (AFC) is given by:

\[
\delta_m - \delta_0 = \left( \delta_a - \delta_i \right) + \Delta \times R \left( 1 - f \left( \frac{1}{R - 1} \right) \right)
\]

where R is the mass ratio of material crystallized to material assimilated, \( \Delta \) is the difference in isotope ratio between the magma and the crystals (\( \delta_{\text{magma}} - \delta_{\text{crystals}} \)), f is the fraction of liquid remaining, \( \delta_m \) is the \( \delta^{18}O \) of the magma, \( \delta_0 \) is the initial \( \delta^{18}O \) of the magma, and \( \delta_a \) is the \( \delta^{18}O \) of the material being assimilated. The assumption is made that the concentration of oxygen is the same in the crystals, magma and assimilant, which is a reasonable assumption. This equation breaks down at R = 1, but, as discussed above, R is generally significantly greater than 1. Figure 9.43 shows the variation of \( \delta^{18}O \) of a magma with an initial \( \delta^{18}O = 5.7 \) as crystallization and assimilation proceed.
Combining stable and radiogenic isotope ratios provides an even more powerful tool to examine assimilation. We shall consider this in Chapter 12.

9.9 Isotopes of Boron and Lithium

Although there are a few earlier works in the literature, there was little interest in the isotopic composition of boron and lithium until about 15 years ago. This is perhaps because both elements have low abundances in the Earth compared to the other elements we have discussed thus far. It is perhaps also because of the analytical problems: neither B nor Li form gaseous species that can be analyzed in the gas-source mass spectrometers used for analysis of other stable isotopes, and the fractionation produced in thermal ionization mass spectrometers, which are used for elements such as Sr, Nd, and Pb, would exceed the natural ones. Since the development of new analytical techniques that overcame this latter problem in the 1980’s, the field of boron isotope geochemistry has developed rapidly, and a range of about 9% in the $^{11}\text{B}/^{10}\text{B}$ ratio in terrestrial materials has been observed. Boron isotope geochemistry has been used to address a wide variety of geochemical problems. Most notable among these are: hydrothermal processes, the nature and origin of ore-forming fluids, the origin of evaporites and brines, pH of ancient oceans, the origin of subduction-related magmas, and geochemical evolution of the mantle. Lithium isotope geochemistry remains in its infancy.

Though both lithium and boron can occur as stoichiometric components of minerals, these minerals have limited occurrence, and these elements generally substitute for other elements in silicates. Boron is relatively abundant in seawater, with a concentration of 4.5 ppm. Lithium is somewhat less abundant, with a concentration of 0.17 ppm. In silicate rocks, the concentration of boron ranges from a few tenths of a ppm or less in fresh basalts and peridotites to several tens of ppm in clays. Lithium concentrations typically range from a few ppm to a few tens of ppm. As can be seen in Table 9.1, the $^{11}\text{B}/^{10}\text{B}$ is reported as per mil variations, $\delta^{11}\text{B}$, from the NBS 951 standard; the $^{6}\text{Li}/^{7}\text{Li}$ ratio is reported as per mil variation, $\delta^{6}\text{Li}$, from the NBS Li$_2$CO$_3$ (L-SVEC) standard.

In nature, boron is almost always bound to oxygen or hydroxyl groups in either triangular (e.g., BO$_3$) or tetrahedral (e.g., B(OH)$_4$) coordination (the only exception is boron bound to fluorine, e.g., BF$_3$). Since the bond strengths and vibrational frequencies of triangular and tetrahedral forms differ, we can expect that isotopic fractionation will occur between these two forms. This is confirmed by experiments which show a roughly 20‰ fractionation between B(OH)$_3$ and B(OH)$_4^-$, with $^{11}\text{B}$ preferentially found in the B(OH)$_3$ form.

In natural aqueous solutions boron occurs as both boric acid, B(OH)$_3$, and the borate ion, B(OH)$_4^-$. In seawater, about 80-90% of boron will be in the B(OH)$_3$ form. In fresh waters, B(OH)$_3$ will be more dominant; only in highly alkaline solutions, such as saline lakes, will B(OH)$_4^-$ be dominant. The most common boron mineral in the crust is tourmaline (Na(Mg,Fe,Li,Al)$_3$Si$_6$O$_{18}$(BO$_3$)$_3$(OH,F)$_4$), in which boron is present in BO$_3$ groups. In clays, boron appears to occur primarily as B(OH)$_4^-$, most likely substituting for silica in tetrahedral layers. The coordination of boron in common igneous minerals is uncertain, possibly substituting for Si in tetrahedral sites. As we noted in Chapter 7, boron is an incompatible element in igneous rocks and is very fluid mobile. It is also readily adsorbed onto the surfaces of clays. There is an isotopic fractionation between dissolved and adsorbed B of -20 to -30‰ (i.e., adsorbed B is $^{11}\text{B}$ poor), depending on pH and temperature (Palmer et al., 1987).

Figure 9.44 illustrates the variation in B isotopic composition in a variety of geologic materials. Spivack and Edmond (1987) found the $\delta^{11}\text{B}$ of seawater to be -39.5‰, and was uniform within analytical error (±0.25‰). Fresh mid-ocean ridge basalts typically have $\delta^{11}\text{B}$ of about -4‰. Oceanic island basalts (OIB) have slightly lighter $\delta^{11}\text{B}$ (e.g., Chaussidon and Jambon, 1994). The average B isotopic composition of the continental crust probably lies between -13‰ and -8‰ (Chaussidon and Abramède, 1992).

Perhaps the most remarkable aspect of B isotope geochemistry is the very large fractionation of B isotopes between the oceans and the silicate Earth. It was recognized very early that this difference
reflected the fractionation that occurred during adsorption of boron on clays (e.g., Schwarcz, et al., 1969). However, as we noted above, this fractionation is only about 30‰ or less, whereas the difference between the continental crust and seawater is close to 50‰. Furthermore, the net effect of hydrothermal exchange between the oceanic crust and seawater is to decrease the $\delta^{11}$B of seawater (Spivack and Edmond, 1977). The discrepancy reflects the fact that the ocean is not a simple equilibrium system, but rather a kinetically-controlled open one. Since all processes operating in the ocean appear to preferentially remove $^6$B from the ocean, seawater is driven to an extremely $^6$B-rich composition. In addition, it is possible that additional fractionations of boron isotopes may occur during diagenesis. Ishikawa and Nakamura (1993) noted that ancient limestones and cherts have more negative $\delta^{11}$B than their modern equivalents, calcareous and siliceous oozes, and suggested that further fractionation occurred during diagenesis.

Spivack and Edmond (1987) investigated the exchange of boron between seawater and oceanic crust. Boron is readily incorporated into the alteration products of basalt, so that even slightly altered basalts show a dramatic increase in B concentration and an increase in $\delta^{11}$B of 5 to 15‰ (Figure 9.44). During high temperature interaction between seawater and oceanic crust, Spivack and Edmond (1987) concluded that boron was quantitatively extracted from the oceanic crust by hydrothermal fluids. The isotopic composition of these fluids is slightly lower than that of seawater. They inferred that the B in these fluids is a simple mixture of seawater- and basalt-derived B and that little or no isotopic fractionation was involved. Analysis of hydrothermal altered basalts recovered by the Ocean Drilling Project generally confirm these inferences, as they are boron-poor and have $\delta^{11}$B close to 0 (Ishikawa and Nakamura, 1992).

The more positive $\delta^{11}$B values of island arc volcanics (IAV) relative to MORB must in part reflect the incorporation of subducted marine sediment and altered oceanic crust into the sources of island arc magmas (e.g., Palmer, 1991). This idea is consistent with a host of other data, which we will discuss in Chapter 12. There is, however, some debate as to the extent the more positive $\delta^{11}$B values in IAV might also reflect assimilation of sediment or altered oceanic crust during magma ascent (e.g., Chaussidon and Jambon, 1994).

The differences in $\delta^{11}$B between oceanic island basalts (OIB) and MORB is perhaps more problematic. Though no experimental or theoretical studies have been carried out, it seems unlikely that significant fractionation of boron isotopes will occur during melting of the mantle, both because the temperatures are high, and because the atomic environment of B in silicate melts is probably similar to that in silicate solids. Thus, as we found was the case for O isotopes, B isotope fractionation
probably occurs only at the surface of the Earth, and the difference between OIB and MORB must somehow reflect surface processes. Chaussidon and Marty (1995) argued that the boron isotopic composition of the mantle is that of OIB (-10‰) and that the higher $\delta^{11}B$ of MORB reflects contamination of MORB magmas by altered oceanic crust. This seems unlikely for several reasons. First, although there are still relatively few data available, MORB appear to be relatively homogeneous in their boron isotopic composition. This means assimilation would have to be systematic and pervasive and that all MORB magmas would have to assimilate roughly the same amount of material. Both of these seem highly improbable. Second, there is little or no other evidence for assimilation of oceanic crust by MORB magmas. Third, oceanic island basalts have an opportunity to assimilate not only altered oceanic crust, but also overlying sediment. Yet, according to the Chaussidon and Marty (1995) hypothesis, they are not systematically contaminated. Although they are not systematically contaminated, there is evidence of occasional assimilation of oceanic crust and/or sediment by oceanic island basalt magmas from both B and Os isotope geochemistry. This may explain some of the lower $\delta^{11}B$ values in OIB seen in Figure 9.43 (Chaussidon and Jambon, 1994).

The alternative explanation for the lower $\delta^{11}B$ in OIB is that they contain a component of material recycled into the mantle, through subduction, from the surface of the Earth. The idea that mantle plumes, and the OIB magmas they produce, contain material recycled from the surface of the Earth has been suggested on other grounds (Hofmann and White, 1982), and we shall discuss it further in Chapter 11. You (1994) points out that significant fractionation of B isotopes will occur during sediment dewatering at moderate temperatures during the initial stages of subduction. The fluid produced will be enriched in $^{11}B$, leaving the sediment depleted in $^{11}B$. Thus the effect of subduction zone processes will be to lower the $\delta^{11}B$ of oceanic crust and sediment carried into the deep mantle. Deciding between this view and that of Chaussidon and Marty (1995) will require further research.

One of the more interesting applications of boron isotopes has been determining the paleo-pH of the oceans. Carbonates biologically precipitated from seawater have $\delta^{11}B$ that is 10 to 25‰ lower than seawater (Figure 9.44). This fractionation results from the kinetics of B coprecipitation in CaCO$_3$, in which incorporation of B in carbonate is preceded by surface adsorption of B(OH)$_4^-$ (Vengosh et al., 1991; Heming and Hanson, 1992).

We noted above that boron is present in seawater both as B(OH)$_3$ and B(OH)$_4^-$. Since the reaction between the two may be written as:

$$ \text{B(OH)}_3 + \text{H}_2\text{O} \rightleftharpoons \text{B(OH)}_4^- + \text{H}^+ $$

The equilibrium constant for this reaction is:

$$ pK_{app} = \ln \left( \frac{[\text{B(OH)}_4^-]}{[\text{B(OH)}_3]} \right) - \text{pH} $$

The relative abundance of these two species is thus pH dependent. Furthermore, we can easily show that, the isotopic composition of these two species must vary if the isotopic composition of seawater is constant. From mass balance we have:

$$ \delta^{11}B_{sw} = \delta^{11}B_3F + \delta^{11}B_4(1 - F) $$

where F is the fraction of B(OH)$_3$ (and 1 - F is therefore the fraction of B(OH)$_4^-$), $\delta^{11}B_3$ is the isotopic composition of B(OH)$_3$, and $\delta^{11}B_4$ is the isotopic composition of B(OH)$_4^-$. If the isotopic composition of the two species are related by a constant fractionation factor, $\Delta_{3-4}$, we can write 9.70 as:

$$ \delta^{11}B_{sw} = \delta^{11}B_3F + \delta^{11}B_4 - \Delta_{3-4}F = \delta^{11}B_4 + \Delta_{3-4}F $$

Solving for $\delta^{11}B_4$, we have:

$$ \delta^{11}B_4 = \delta^{11}B_{sw} - \Delta_{3-4}F $$

Thus assuming a constant fractionation factor and isotopic composition of seawater, the $\delta^{11}B$ of the two B species will depend only on F, which, as we can see in equation 9.69, will depend on pH.
If the mechanism for incorporation of B in carbonate presented above is correct, the fractionation of $^{11}\text{B}/^{10}\text{B}$ between calcite and seawater will be pH dependent. The situation is slightly complicated because different carbonate-secreting species may fractionate B isotopes slightly differently, perhaps because they alter the pH of their micro-environment, or perhaps because $\text{B(OH)}_3$ is also utilized to varying degrees. Some care must therefore be exercised to exclude these “vital” effects. Beyond that, the isotopic composition of marine biogenic carbonate preserved in sediment cores should reflect the pH of the water from which they were precipitated.

Spivack et al. (1993) found that pore fluid in 620 m sediment core recovered at Ocean Drilling Program Hole 803D in the Indian Ocean had a constant B concentration and isotopic composition. From this they argued that the isotopic composition of the oceans has been constant of the past 21 Ma (the age of the lowest sediment recovered). The composition of foraminiferal calcite tests, however, varied significantly (Figure 9.45). The variation, they concluded, reflected an increase in the pH of the ocean between 15 and 7 million years ago.

In a similar study, Sanyal et al. (1995) compared the $\delta^{11}\text{B}$ of benthic and planktonic foraminifera deposited during the last glacial period to those in modern sediments. They found that $\delta^{11}\text{B}$ of both Atlantic and Pacific planktonic foraminifera were 1.8‰ higher during the last glacial period while $\delta^{11}\text{B}$ of Atlantic and Pacific benthic foraminifera were 2.5‰ higher. These differences correspond to increases in pH of 0.2 in the surface water and 0.3 in the deep water. Both Spivack et al. (1993) and Sanyal et al. (1995) pointed out that the pH of seawater depend on the concentration of atmospheric CO$_2$ (because CO$_2$ dissolves to form carbonic acid, H$_2$CO$_3$, which then dissociates to a H$^+$ ion and a bicarbonate ion). Thus the apparent changes in pH may reflect a long-term decrease in the concentration of atmospheric CO$_2$ during the Miocene period (which we mentioned earlier in this chapter), and short-term decreases in atmospheric CO$_2$ during glacial intervals.

Few studies have been made of Li isotopic variations in terrestrial materials. The available data are summarized in Figure 9.46. Seawater has a constant $\delta^6\text{Li}$ value of $-32.3\%_o$ (Chan and Edmond, 1988). As is the case for boron, it represents one extreme of the spectrum of isotopic compositions in the Earth. As for boron, this appears to reflect preferential removal of the light isotope by reaction with clays and other authigenic minerals. Chan et al. (1992) estimated the fractionation factor for clay-solution exchange at $-19\%_o$, which presumably reflects differences in the binding energies of $^6\text{Li}$ and $^7\text{Li}$ with water on the one hand and anions (O and OH) in the clay structure on the other.
References and Suggestions for Further Reading


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Chapter 9: Stable Isotopes


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Geochemistry

Chapter 9: Stable Isotopes


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CHAPTER 9: STABLE ISOTOPES


Problems

1. What would you predict would be the ratio of the diffusion coefficients of H₂O and D₂O in air?

2. A granite contains coexisting feldspar (3% An content) and biotite with δ¹⁸OSMOW of 9.2‰ and 6.5‰ respectively. Using the information in Table 9.2, find the temperature of equilibration.

3. Spalerite and galena from a certain Mississippi Valley deposit were found to have δ³⁴SCDT of +13.2‰ and +9.8‰ respectively.
   a. Using the information in Table 9.4, find the temperature at which these minerals equilibrated.
   b. Assuming they precipitated from an H₂S-bearing solution, what was the sulfur isotopic composition of the H₂S?

4. Glaciers presently constitute about 2.1% of the water at the surface of the Earth and have a δ¹⁸OSMOW of ≈-30. The oceans contain essentially all remaining water. If the mass of glaciers were to increase by 50%, how would the isotopic composition of the ocean change (assuming the isotopic composition of ice remains constant)?

5. Consider the condensation of water vapor in the atmosphere. Assume that the fraction of vapor remaining can be expressed as a function of temperature (in kelvins):
   \[ f = \frac{T - 223}{50} \]
   Also assume that the fractionation factor can be written as:
   \[ \ln \alpha = 0.0018 + \frac{12.8}{RT} \]
   Assume that the water vapor has an initial δ¹⁸OSMOW of -9‰. Make a plot showing how the δ¹⁸O of the remaining vapor would change as a function of f, fraction remaining.

6. Calculate the δ¹⁸O of raindrops forming in a cloud after 80% of the original vapor has already condensed assuming (1.) the water initial evaporated from the ocean with δ¹⁸O = 0, (2.) the liquid-vapor fractionation factor, α = 1.0092.
7. A saline lake that has no outlet receives 95% of its water from river inflow and the remaining 5% from rainfall. The river water has $\delta^{18}O$ of $-10\%$ and the rain has $\delta^{18}O$ of $-5\%$. The volume of the lake is steady-state (i.e., inputs equal outputs) and has a $\delta^{18}O$ of $-3$. What is the fractionation factor, $\alpha$, for evaporation?

8. Consider a sediment composed of 70 mole percent detrital quartz ($\delta^{18}O = +10\%$) and 30 mole percent calcite ($\delta^{18}O = +30\%$). If the quartz/calcite fractionation factor, $a$, is 1.002 at 500˚ C, determine the O isotopic composition of each mineral after metamorphic equilibrium at 500˚ C. Assume that the rock is a closed system during metamorphism, i.e., the $\delta^{18}O$ of the whole rock does not change during the process.

9. Assume that $\alpha$ is a function of the fraction of liquid crystallized such that $\alpha = 1 + 0.008(1-f)$. Make a plot showing how the $\delta^{18}O$ of the remaining liquid would change as a function of $1-f$, fraction crystallized, assuming a value for $\Delta$ of 0.5‰.

10. A basaltic magma with a $\delta^{18}O$ of +6.0 assimilates country rock that $\delta^{18}O$ of +20 as it undergoes fractional crystallization. Assuming a value of $\alpha$ of 1.002, make a plot of $\delta^{18}O$ vs. $f$ for $R = 5$ and $R = 10$ going from $f = 1$ to $f = 0.05$. 

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Chapter 9: Stable Isotopes

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