

# The carbon cycle and associated redox processes through time

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Earth's biogeochemical cycle of carbon delivers both limestones and organic materials to the crust. In numerous, biologically catalysed redox reactions, hydrogen, sulphur, iron, and oxygen serve prominently as electron donors and acceptors. The progress of these reactions can be reconstructed from records of variations in the abundance of <sup>13</sup>C in sedimentary carbonate minerals and organic materials. Because the crust is always receiving new CO<sub>2</sub> from the mantle and a portion of it is being reduced by photoautotrophs, the carbon cycle has continuously released oxidizing power. Most of it is represented by Fe<sup>3+</sup> that has accumulated in the crust or been returned to the mantle via subduction. Less than 3% of the estimated, integrated production of oxidizing power since 3.8 Gyr ago is represented by O<sub>2</sub> in the atmosphere and dissolved in seawater. The balance is represented by sulphate. The accumulation of oxidizing power can be estimated from budgets summarizing inputs of mantle carbon and rates of organic-carbon burial, but levels of O<sub>2</sub> are only weakly and indirectly coupled to those phenomena and thus to carbon-isotopic records. Elevated abundances of <sup>13</sup>C in carbonate minerals *ca* 2.3 Gyr old, in particular, are here interpreted as indicating the importance of methanogenic bacteria in sediments rather than increased burial of organic carbon.

**Keywords:** carbon cycle; carbon isotopes; atmospheric oxygen; methanogenesis; subduction; mantle

Together, biological and geological processes—oxygenic photosynthesis and the burial of organic carbon—get credit for producing and maintaining the O<sub>2</sub> in Earth's breathable atmosphere. This view of the carbon cycle as the engine of environmental evolution is based on sedimentary records. The disappearance of mass-independent fractionation of the isotopes of sulphur is the most reliable indicator for the accumulation and persistence of traces of O<sub>2</sub> in the atmosphere beginning at about 2.4 Gyr ago (Ga) ( $P_{O_2} \geq 10^{-5}$  atm; Pavlov & Kasting 2002; Farquhar & Wing 2003; Bekker *et al.* 2004). Biomarkers derived from lipids associated with cyanobacteria first appear in sedimentary rocks with an age of 2.7 Gyr (Summons *et al.* 1999; Brocks *et al.* 2003; Eigenbrode 2004). These provide strong evidence for oxygenic photoautotrophy at that time (i.e. for production of O<sub>2</sub> as opposed to its accumulation, persistence, and global distribution). Abundances of oxidized and reduced minerals in ancient soil profiles and sediments generally indicate a transition from weakly reducing to weakly oxidizing conditions at Earth's surface soon after 2.47 Ga (Bekker *et al.* 2004; Canfield 2005; Catling & Claire 2005).

But what about examining the engine itself? Records of the burial of organic carbon, which should be provided by abundances of <sup>13</sup>C in

sedimentary carbonates and organic material (Broecker 1970), could indicate the pace *and* the mechanism of oxidation. For events and processes during the past 500 Myr, carbon-isotopic records have been interpreted with considerable success (e.g. Garrels & Lerman 1981; Holland 1984; Berner 1991, 2004; Kump & Arthur 1999). The same approach has been extended to Precambrian records (e.g. Schidlowski *et al.* 1975; Hayes 1983, 1994; Knoll *et al.* 1986; Derry *et al.* 1992; Karhu & Holland 1996; Halverson *et al.* 2005), but conclusions have usually been qualitative rather than quantitative. The carbon-isotopic record can be described as 'consistent with' some postulated event or process, but understanding of the carbon cycle has not been complete enough to allow resolution of uncertainties or elaboration of details.

Two recent findings may change this. First, new evidence (Saal *et al.* 2002) has led to wide agreement (cf. Resing *et al.* 2004) on the rate at which C is delivered to the crust from the mantle at mid-ocean ridges. This significant reduction in uncertainties about the input allows a new approach to carbon budgets. Second, a previously overlooked output of carbon from the ocean has been recognized and quantified. Whereas earlier concepts limited the outputs to sedimentary carbonates and organic matter, the new view includes a very large flow of carbon that is taken up during the weathering of seafloor basalts. This changes the structure of the mass balances. The potential role of this production of 'carbonated basalts' in the carbon

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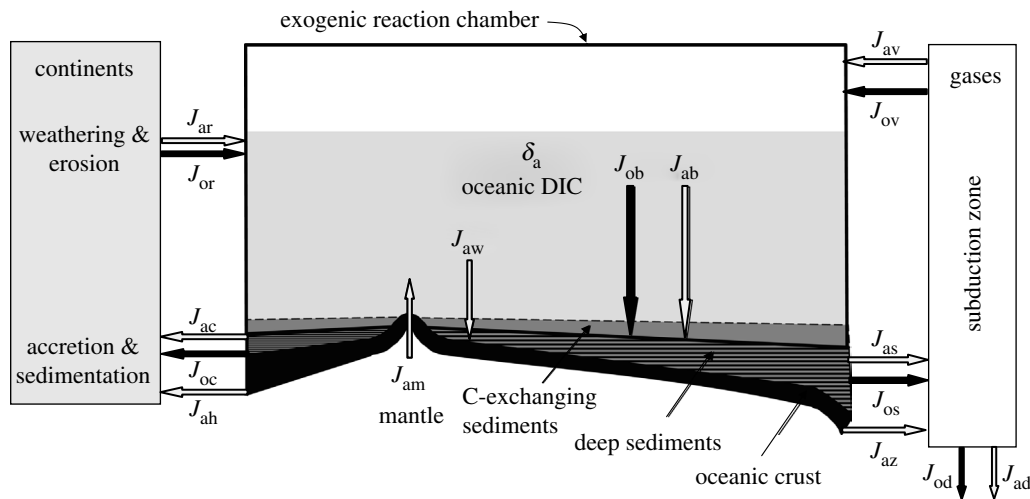


Figure 1. A schematic of the biogeochemical cycle of carbon. The arrows represent fluxes of carbon that are explained in the text.

cycle was first identified by Staudigel *et al.* (1989). Subsequently, Alt & Teagle (1999, 2003) have examined amounts and isotopic compositions of these carbonate minerals and discussed their importance in the modern carbon cycle. Walker (1990), Sleep & Zahnle (2001), and Nakamura & Kato (2004) have called attention to carbonation of submarine basalts as an important phenomenon during the Archaean. Bjerrum & Canfield (2004) have introduced a systematic treatment of this problem.

Our purpose here, based on those developments, is to demonstrate a new approach to studies of the carbon cycle and its effects on the global environment. Among those effects, we focus on oxidation of Earth's surface over the past 4 Gyr.

## 1. THE STRUCTURE OF THE CARBON CYCLE

A geochemical view of the carbon cycle is shown in figure 1. The scheme chosen highlights processes that control the isotopic composition of inorganic carbon dissolved in the ocean. That carbon pool is in rough equilibrium with the atmosphere and with carbonate minerals derived from seawater. It is the senior author of our best records of how the carbon cycle has operated over the course of Earth history. To interpret those records, we must consider the fluxes indicated in figure 1. They represent processes that are linked by balances of mass and electrons. The related equations are, at present, analytical tools rather than components of an elaborate model.

The reaction chamber in which isotopic variations are shaped is comprised of the atmosphere, hydrosphere, and C-exchanging sediments and soils. Carbon flows into that chamber from the mantle and by recycling of carbon within the crust. It leaves through burial in sediments and during weathering of seafloor basalts. If the amount of C in the reactor is constant, the inputs will be balanced by the outputs:

$$J_{am} + J_{ar} + J_{or} + J_{av} + J_{ov} = J_{ab} + J_{ob} + J_{aw}. \quad (1.1)$$

The terms in this equation represent fluxes of carbon in mol per time. The subscripts 'a' and 'o' refer to inorganic and organic carbon. In detail,  $J_{am}$  is the total

input of C from the mantle via outgassing of magmas at seafloor hydrothermal vents and at hot-spot and island-arc volcanoes;  $J_{ar}$  and  $J_{or}$  are, respectively, returns of carbonate and organic C by exposure and weathering of deposits on continents and shelves; and  $J_{av}$  and  $J_{ov}$  are returns, principally by arc volcanism, of carbonate and organic C remobilized during subduction. Among the outputs,  $J_{ab}$  and  $J_{ob}$  are the net burials of carbonate- and organic-C in marine sediments, deriving mainly from processes in surface waters but often bearing strong secondary imprints; and  $J_{aw}$  is carbonate being taken up at the seafloor during the weathering of basalts. These, and all additional definitions pertinent to this discussion, are summarized in table 1.

A second mass balance equates incoming and outgoing  $^{13}\text{C}$ . Its elaboration leads to a key indicator of variations in the operation of the carbon cycle. For simplicity and generality, it is often written in this form:

$$J_i \delta_i = J_{ab} \delta_{ab} + J_{ob} \delta_{ob} + J_{aw} \delta_{aw}, \quad (1.2)$$

where  $J_i$  represents the summed inputs to the exogenic reaction chamber,  $\delta_i$  represents their weighted-average isotopic composition and the remaining  $\delta$  terms represent the isotopic compositions of the indicated fluxes. In detail, a fully correct form would be

$$\begin{aligned} J_{am} \delta_{am} + J_{ar} \delta_{ar} + J_{or} \delta_{or} + J_{av} \delta_{av} + J_{ov} \delta_{ov} \\ = J_{ab} \delta_{ab} + J_{ob} \delta_{ob} + J_{aw} \delta_{aw}. \end{aligned} \quad (1.3)$$

Because values for many of the terms on the left side of this equation are inaccessible, analyses usually proceed from equation (1.2), incorporating the assumptions that  $J_i = J_{ab} + J_{ob} + J_{aw}$  and that  $\delta_i = \delta_{am}$ . When historical variations are considered, the latter has two components: (i) that mixing of recycling inputs eventually yields an unbiased sample and (ii) that  $\delta_{am}$  is constant. Over rock-cycle time-scales of 300 Myr or more, the first requirement is probably met. Second, the constancy and uniformity of  $\delta_{am}$  are well supported. Independent of age of emplacement or location, diamonds from peridotitic xenoliths have  $\delta = -5\text{‰}$  (Pearson *et al.* 2004). The same value is found in carbonatites and mantle-derived basalts (Kyser 1986; Mathey 1987).

Table 1. Definitions.

term	definition
main variables	
$A_{\text{Ox}}$	rate of accumulation of oxidants in crust (mol O <sub>2</sub> equivalent) per time
$J$	flux of carbon (mol per time)
$L$	flux of oxidant or reductant (mol O <sub>2</sub> equivalent) per time
$M$	molar quantity of C (no subscript: total crustal carbon, all forms)
$f$	fraction of C buried in organic form ( $=J_{\text{ob}}/(J_{\text{ab}}+J_{\text{aw}}+J_{\text{ob}})$ )
$\dot{j}_x$	flux of substance x (mol per time)
$\Delta_g$	isotopic difference between DIC in surface seawater and diagenetically stabilized carbonate minerals in sedimentary rocks (‰); see equation (1.4)
$\Delta_m$	isotopic difference between DIC in surface seawater and carbonate in weathered oceanic basalts (‰); see equation (1.6)
$\gamma$	fraction of crustal C recycling during $\tau$ . $\gamma = (1 - e^{-k\tau})$ , where $k = (\ln 2)/(\text{half-mass age})$
$\delta$	$\delta^{13}\text{C}$ relative to the Vienna PeeDee Belemnite standard (Zhang & Li 1990)
$\varepsilon$	isotopic fractionation between DIC in surface seawater and sedimentary organic carbon (‰); see equation (1.5)
$\varphi$	$\text{Fe}^{3+}/\Sigma\text{Fe}$
$\lambda$	fraction of buried carbonate accounted for by ocean-crustal carbonates ( $=J_{\text{aw}}/(J_{\text{aw}}+J_{\text{ab}})$ )
$\tau$	time-step in numerical integrations (10 <sup>7</sup> years); figures 5, 7 and 9
subscripts appended to $J$ , $L$ and $\delta$	
<i>first or only part</i>	
Ox	oxidant
Red	reductant
a	carbonate carbon
i	input to exogenic reaction chamber; see equation (1.3)
o	organic carbon
<i>second part, pertaining to a directional flux</i>	
b	burial, transfer from exogenic reaction chamber to deep sediments
c	transfer from deep sediments to continental crust
d	transfer from subduction zone to mantle
h	transfer from ocean crust to continental crust
m	transfer from mantle to exogenic reaction chamber
r	(returns) transfer from continent to exogenic reaction chamber
s	(subduction) transfer from deep sediments to subduction zone
v	(volcanism) transfer from subduction zone to exogenic reaction chamber
w	(weathering) transfer from exogenic reaction chamber to ocean crust
x	transfer from exogenic reaction chamber to space
z	transfer from ocean crust to subduction zone

The task now is to provide a useful approach to interpreting observed variations in  $\delta_{\text{ab}}$ , the carbon-isotopic composition of sedimentary carbonates. As has become conventional, we define the fraction of input C buried in organic form as  $f \equiv J_{\text{ob}}/J_i$ . Following Bjerrum & Canfield (2004), we define the fraction of total carbonate accounted for by ocean-crustal carbonates as  $\lambda \equiv J_{\text{aw}}/(J_{\text{aw}}+J_{\text{ab}})$ .

Our approach to the isotopic variables differs from previous expositions. As the reference point, we choose the isotopic composition of total dissolved inorganic carbon (DIC) in marine surface waters,  $\delta_a$ . The isotopic compositions of the outputs are related to  $\delta_a$  by the following expressions:

$$\delta_{\text{ab}} = \delta_a - \Delta_g, \quad (1.4)$$

$$\delta_{\text{ob}} = \delta_a - \varepsilon, \quad (1.5)$$

$$\delta_{\text{aw}} = \delta_a - \Delta_m, \quad (1.6)$$

where  $\Delta_g$  is the globally averaged isotopic difference between surface DIC and diagenetically stabilized sedimentary carbonates. At present, for example, comparison of pre-industrial  $\delta_a$  (Quay *et al.* 2003) and average sedimentary carbonate (Shackleton 1987)

indicates  $\Delta_g \approx 2\text{‰}$ . Local variations in  $\Delta_g$  can affect specific sedimentary records. The difference in isotopic composition between sedimentary organic carbon and DIC,  $\varepsilon$ , is principally (though not exclusively) due to isotopic discrimination during biotic carbon fixation. The sign chosen in equation (1.5), with  $\varepsilon > 0$  corresponding to depletion of  $^{13}\text{C}$  in biomass, is conventional in marine biogeochemistry. Values of  $\Delta_m$  will be positive when ocean-crustal carbonates are depleted in  $^{13}\text{C}$  relative to surface waters. Reports of  $\delta_{\text{aw}}$ , required to evaluate  $\Delta_m$ , are rare. Alt & Teagle (2003) find  $\delta_{\text{aw}} = 1.7 \pm 0.4\text{‰}$  for ocean-crustal carbonates that have formed during the past 160 Myr. The average value of  $\delta_{\text{ab}}$  during the same interval (Veizer *et al.* 1999) is 1.7‰. At present, therefore,  $\Delta_m \approx \Delta_g \approx 2\text{‰}$ . Other reports occasionally indicate lower values of  $\delta_{\text{aw}}$  and thus suggest larger values of  $\Delta_m$  but, based on associated sedimentary features, the authors uniformly attribute the depletion of  $^{13}\text{C}$  to infrequent additions of C derived from oxidation of organic material.

Substitution of equations (1.4)–(1.6) in equation (1.3) and simplification of the result yields

$$\delta_{\text{ab}} - \delta_i = f(\varepsilon - \Delta_g) + \lambda(1 - f)(\Delta_m - \Delta_g). \quad (1.7)$$

For  $\Delta_g=0$ , equation (1.7) is equivalent to equation (1.4) of Bjerrum & Canfield (2004). If, in addition, either or both  $\Delta_m$  and  $\lambda$  are zero, equation (1.7) becomes  $\delta_{ab}-\delta_i=f\varepsilon$ , the expression found in numerous prior discussions of isotopic fractionation in the carbon cycle. Does this expression indicate that  $\Delta_m$  and  $\Delta_g$  can be as effective as  $f$  and  $\varepsilon$  in controlling  $\delta_{ab}-\delta_i$ ? Probably not. Their leverage is relatively small. In most circumstances, the first term,  $f(\varepsilon-\Delta_g)$ , will be at least four times larger than the second,  $\lambda(1-f)(\Delta_m-\Delta_g)$ .

Rearrangement of equation (1.7) yields an expression for  $f$ , the fraction of carbon buried in organic form:

$$f = \frac{\delta_{ab} - \delta_i - \lambda(\Delta_m - \Delta_g)}{\varepsilon - \Delta_g - \lambda(\Delta_m - \Delta_g)}. \quad (1.8)$$

Commonly (e.g. Hayes *et al.* 1999),  $f$  is estimated from  $(\delta_{ab}-\delta_i)/(\delta_{ab}-\delta_{ob})$ . This is precisely equivalent to  $(\delta_{ab}-\delta_i)/(\varepsilon-\Delta_g)$ . The numerator and denominator in that fraction lack the correction terms,  $-\lambda(\Delta_m-\Delta_g)$ , which are prominent in equation (1.8). If  $\Delta_m \approx \Delta_g$ , as at present,  $\lambda(\Delta_m-\Delta_g)=0$  and the terms are inconsequential, independent of the importance of basalt carbonation. Such cases are summarized graphically in figure 2a. The vertical line at  $\delta_{ab}-\delta_i=6.5\text{‰}$  indicates  $\delta_{ab}=1.5\text{‰}$ , near Earth's observed, long-term average value. The lines correspond (left to right) to  $\varepsilon=22, 26, 30$  and  $34\text{‰}$ . The shaded area indicates that, depending on the value of  $\varepsilon$ ,  $\delta_{ab}=1.5\text{‰}$  corresponds to  $0.2 \leq f \leq 0.32$ . This range encompasses the values most frequently noted in previous discussions of the carbon cycle. In particular,  $\delta_{ab}-\delta_{ob}=28\text{‰}$ , corresponding to  $\varepsilon=30$  and  $\Delta_g=2\text{‰}$ , is representative of much of the Phanerozoic (Hayes *et al.* 1999).

The lower  $f$  values marked by shading in the other frames of figure 2 correspond to the same value of  $\delta_{ab}$ , but are based on different estimates of  $\Delta_g$ ,  $\Delta_m$  and  $\lambda$ . Negative values of  $\Delta_g$  are observed when sedimentary carbonates are strongly affected by methanogenic diagenesis (Irwin *et al.* 1977). In Phanerozoic strata, which have formed in the presence of relatively abundant  $O_2$  and  $SO_4^{2-}$ , this phenomenon is restricted to concretions or other zones, in which supplies of sedimentary organic matter have been large enough that methanogenesis has eventually become prominent. When concentrations of  $O_2$  and  $SO_4^{2-}$  in seawater were significantly lower, methanogenesis must have been more important. Accordingly, the effects of negative values of  $\Delta_g$ , corresponding to globally important levels of methanogenic diagenesis, are explored in figure 2b,c. Figure 2d shows that variations in  $\Delta_m$  are probably least important in affecting estimates of  $f$ . Under steady-state conditions, inversion of the oceanic  $^{13}C$  gradient—enrichment of  $^{13}C$  in bottom waters ( $\Delta_m=-5\text{‰}$ , figure 2d)—is practically required to produce  $\delta_{ab}-\delta_i < 0$ .

## 2. REDOX BALANCES IN THE CARBON CYCLE

Figure 3 duplicates the plan of figure 1, but depicts flows of oxidants and reductants generated by the carbon cycle. The biological cycle of production and respiration is at its centre. Redox partners for C are generalized as Red and Ox. The focus on *oxidants*

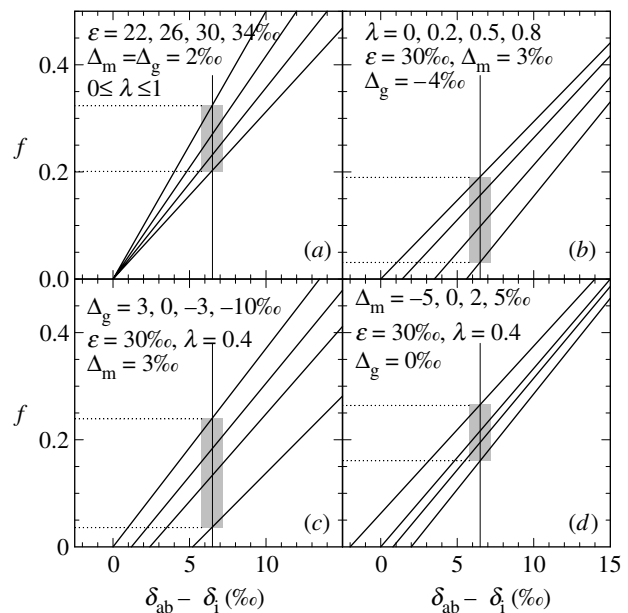


Figure 2. Graphs indicating relationships between  $f$  and  $\delta_{ab}-\delta_i$  (1.7). Slopes and intercepts vary in response to varying values of  $\varepsilon$ ,  $\lambda$ ,  $\Delta_m$ , and  $\Delta_g$ . In each frame, one of these has been assigned four different values and the others have been held constant. The values assigned are indicated in each frame. For the parameter that varies, the sequence of values corresponds to the lines as seen from left to right.

rather than *oxygen* is necessary. The carbon-burial flux provides information about the consumption of electron donors, but not about the identity of those donors. Over the course of Earth history, the electron-donating role of Red has been played by  $H_2$ ,  $Fe^{2+}$ ,  $S^{2-}$  and  $H_2O$  (at least). Correspondingly, the oxidized forms of these substances have accumulated in Earth's crust. The rate of accumulation will be set by the difference between the rate at which carbon-cycling produces oxidants and the rate at which those oxidants are consumed.

To quantify rates, we begin by viewing the  $J_o$  terms, introduced above as fluxes of carbon, as fluxes of reducing power. The  $L$  terms shown in figure 3 represent flows of oxidizing or reducing power carried by other elements or by mixtures of C and other elements, (mol  $O_2$  equivalent) per time. Because reduction of  $CO_2$  to organic carbon and oxidation of  $H_2O$  to  $O_2$  are both four-electron processes, values of the  $J_o$  and  $L$  terms are numerically equivalent. Production of 1 mol of organic carbon could be balanced by oxidation of 2 mol of  $H_2$ , 8 mol of  $Fe^{2+}$ , 0.5 mol of  $S^{2-}$  (to  $SO_4^{2-}$ ), or 2 mol of  $H_2O$ . All would be equivalent to 1 mol of  $O_2$ . To emphasize that it pertains to the net effect of multiple processes, rather than to a specific flux, we use  $A_{Ox}$  to designate the rate at which oxidants accumulate in the crust and exogenic reaction chamber.

To begin, the biological carbon cycle releases oxidants at the rate at which organic carbon is buried in deep sediments ( $J_{ob}$ ). The rate of accumulation is then moderated by effects of two types. Within the crust itself, a portion of the oxidants is consumed by the geological carbon cycle. Two pathways are shown in figure 3. The first is oxidative weathering of organic carbon exposed on and eroded from the continents

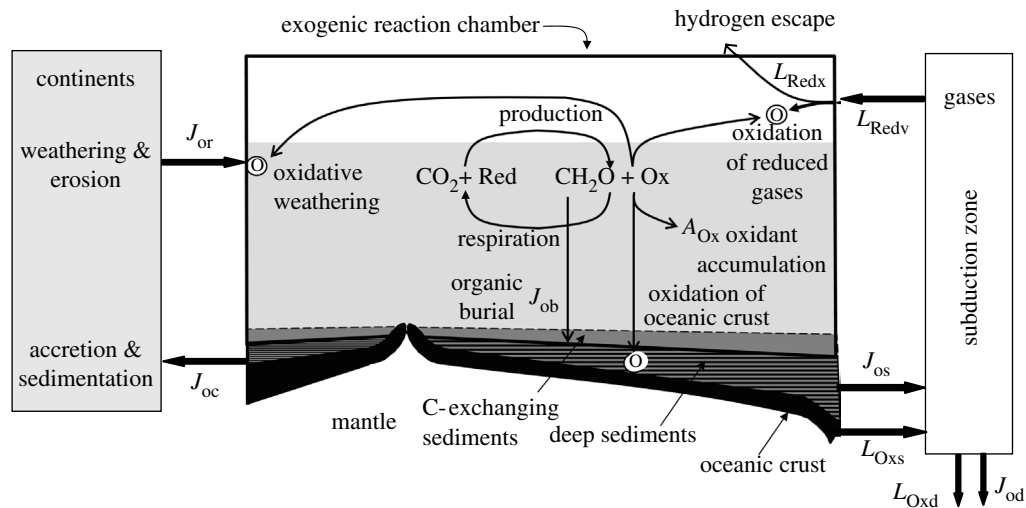


Figure 3. A schematic of flows of oxidants and reductants generated by the carbon cycle. Sites of oxidation are marked by the letter O.

( $J_{or}$ ). The second is oxidation of reduced gases produced at subduction zones. The corresponding flux is designated as  $L_{Redv}$ . It includes not only volcanic gases ( $H_2$ ,  $CO$ ,  $SO_2$ ), but also products ranging from methane to petroleum, which are delivered to the exogenic reaction chamber by thermal processes in subduction zones and deep basins.

The export of oxidants and reductants also moderates  $A_{Ox}$ . This occurs in subduction zones and at the top of the atmosphere. Within subduction zones, electrons are transferred between subducted organic materials ( $J_{os}$ ) and oxidants (metal oxides and  $SO_4^{2-}$ ,  $L_{Oxs}$ ) in the descending slab. Products will include not only those returned to the crust (noted above), but also unconsumed oxidants ( $L_{Oxd}$ ) and reduced carbon (diamond, graphite,  $J_{od}$ ) exported to the mantle. Subduction of carbonate decreases the mass of carbon in the crust, but, because carbon is supplied from the mantle as  $CO_2$ , does not export either oxidizing or reducing power from the crust. Subduction of sulphide is similarly inconsequential. Mantle S occurs as sulphide even under oxidizing conditions (e.g.  $\Delta FMQ \sim 2$ ; Luth 2004).

At the top of the atmosphere,  $H_2$  can be lost to space, thus exporting reducing power and effectively increasing  $A_{Ox}$ . This occurs when reduced gases of either volcanic or biological origin reach high altitudes. Depending on atmospheric conditions, a portion of the reducing power carried by these gases can be lost as hydrogen escapes to space (Catling *et al.* 2001).

Summation provides an estimate of the rate at which oxidants will accumulate:

$$A_{Ox} = (\text{net release of oxidants within crust by C cycle}) \\ + (\text{reductant export}) - (\text{net oxidant export}) \\ A_{Ox} \geq (J_{ob} - J_{or} - L_{Redv}) + L_{Redx} - (L_{Oxd} - J_{od}). \quad (2.1)$$

The second expression denotes  $A_{Ox}$  as a minimum value because the net release of oxidants by the C cycle may exceed  $J_{ob} - J_{or} - L_{Redv}$ . This would occur if a portion of the organic carbon returning from continents ( $J_{or}$ ) was not reoxidized but instead simply reburied. The efficiency of reoxidation will depend on

the nature of Ox (e.g.  $O_2$  versus  $SO_4^{2-}$ ) and the reactivity of  $J_{or}$  (e.g. graphitic kerogen versus hydrocarbons).

Equation (2.1) ignores a category of processes often considered in oxygen budgets. Sediments and oceanic crust assimilated by continents will incorporate portions of the Red and Ox from the exogenic reaction chamber. Oxidized and reduced forms of sulphur and iron are prime examples. Surging flows of these reactants to the sediments or from continents can, for example, significantly affect levels of  $O_2$  in the atmosphere and ocean. These and related phenomena, especially in the sulphur cycle, have already been elegantly treated by others (Bernier 2004; Canfield 2004).

Imbalances yielding a net release or consumption of oxidizing power by the carbon cycle are of two types. The more obvious are marked by isotopic signals indicating rapid variations of  $f$  and consequent departures from steady state. At such times, the system will evolve dynamically (Rothman *et al.* 2003). At other times, when  $f$  varies slowly, the system will evolve quasi-statically through a succession of steady states. The persistence of small imbalances between fluxes can lead to the accumulation of crustal inventories of carbon, chiefly on the continents. In this case, the fluxes designated in equation (2.1) should be represented as time-dependent variables. Then, at any time,  $t$ ,

$$\int_0^t A_{Ox}(t) dt = \int_0^t [J_{ob}(t) - J_{or}(t) - L_{Redv}(t)] dt \\ + \int_0^t L_{Redx}(t) dt - \int_0^t [L_{Oxd}(t) - J_{od}(t)] dt. \quad (2.2)$$

The first term on the right-hand side is the integrated difference between burial and reoxidation of organic carbon. It quantifies the accumulation of organic carbon in continents and marine sediments. The second and third terms quantify the effects of loss of  $H_2$  to space and of subduction of oxidized and reduced materials. Notably, this summation of oxidation has no isotopic dimensions.  $J_{ob}$  is related to  $f$  and thus to the isotopic record, but it is  $J_{ob} - J_{or} - L_{Redv}$  that matters,

and it is further altered by effects of subduction and escape of  $H_2$  to space.

The first term in equation (2.2) is closely related to a principle dating from the nineteenth century (J. J. Ebelmen's work from 1845 to 1855, reviewed by Berner & Maasch 1996) and elaborated in modern detail most influentially by Garrels (e.g. Garrels & Perry 1974) and Berner (2004 and earlier references cited therein). Specifically, the amount of organic carbon stored in the crust should balance the oxidizing power represented by the crustal inventories of  $Fe^{3+}$ ,  $SO_4^{2-}$  and  $O_2$ . To this, space science and plate tectonics have added the second and third terms.

It is difficult to reconstruct the histories of the variables in equation (2.2). A boundary value for the first integral, however, can be obtained by consideration of crust–mantle carbon budgets and variations in  $f$ , the organic-carbon burial fraction.

### 3. INPUTS OF MANTLE CARBON

Fluxes and inventories throughout the carbon cycle depend on inputs of carbon from the mantle. These occur at mid-ocean ridges, arc volcanoes, hotspots and in plume events. The present strengths of these sources will be considered sequentially. An estimate of variations over the course of Earth history will follow.

#### (a) *Mid-ocean ridges*

Each year,  $21 \text{ km}^3$  of basalt is added to the oceanic crust at spreading centres (Crisp 1984; also consistent with a plate-creation rate of  $3.4 \text{ km}^2 \text{ yr}^{-1}$  (Rowley 2002) and a plate thickness of 5–7 km (White *et al.* 1992; Kadko 1994)). Sampled after cooling at the seafloor, its  $CO_2$  content is dependent on the hydrostatic pressure. At the depths of mid-ocean ridges, the result is commonly about 200 p.p.m. (the routinely reported concentrations refer to weights of  $CO_2$ ). The question is how much  $CO_2$  was in the parent magma. The difference will have been transferred, by way of hydrothermal circulation, to the ocean.

Recently, Saal *et al.* (2002) have shown that, in undegassed MORB (mid-ocean ridge basalt), concentrations of  $CO_2$  vary with those of Nb. The weight ratio is  $CO_2 : Nb = 239 \pm 46$  ( $2\sigma$ ). Since Nb is not lost during degassing, the initial  $CO_2$  content of a sample of MORB can be estimated from its content of Nb. The average for normal MORBs (i.e. those in which trace-element abundances have not been affected by proximity to plumes or recently subducted continental materials) on the East Pacific Rise is 3.45 p.p.m. Nb (Su & Langmuir 2003). The estimated, average, initial content of  $CO_2$  is thus  $3.45 \times 239 = 825$  p.p.m. If 200 p.p.m. remain in the cooled basalt, the difference transmitted to the exogenic reservoir is 625 p.p.m. Given a rock density of  $2.8 \text{ g cm}^{-3}$ , this corresponds to  $0.8 \text{ Tmol C yr}^{-1}$  (Tmol = teramol =  $10^{12}$  mol). If the Nb average quoted for all MORBs (5.02 p.p.m.; Su & Langmuir 2003) is instead used as the basis for the calculation, the result is  $1.3 \text{ Tmol C yr}^{-1}$ . These values bracket a third estimate, namely  $0.9 \text{ Tmol C yr}^{-1}$ , reported in the original publication (Saal *et al.* 2002) and reached using a slightly different approach.

Numerous alternative estimates have been based on  $CO_2 : ^3He$  ratios in hydrothermal fluids and gases. These have been reviewed by Resing *et al.* (2004), who settle on a range of 0.5–2  $\text{Tmol C yr}^{-1}$ . Preferring the approach based on chemical analyses of the rocks, we adopt  $1 \text{ Tmol C yr}^{-1}$  as the present magnitude of the mid-ocean ridge component of  $J_{am}$ .

#### (b) *Arc volcanoes*

The annual magma volume is  $0.5 \text{ km}^3$  (Carmichael 2002). Carbon dioxide is abundant in the gases, but its isotopic composition often deviates from the mantle value and the  $CO_2$  is regarded as deriving from subducted sedimentary carbonates and organic carbon as well from the mantle (Sano & Marty 1995; Sano & Williams 1996; Shaw *et al.* 2003, 2004). The total flux of  $CO_2$  from arc volcanism is approximately  $1.6 \text{ Tmol yr}^{-1}$  (Hilton *et al.* 2002). Of this, approximately 13% is from the mantle (Shaw *et al.* 2003), yielding an arc-volcanic component of  $J_{am}$  of  $0.2 \text{ Tmol C yr}^{-1}$ . The remaining  $1.4 \text{ Tmol yr}^{-1}$  is recycling crustal C. Wallace (2005) obtains a similar result by a different method.

#### (c) *Oceanic islands and plumes*

In a collection of estimates, this is the most uncertain. The annual volume of magma is approximately  $3 \text{ km}^3$  (though possibly as small as  $1.9 \text{ km}^3$ ), combining igneous provinces and hotspot volcanoes on continents with those in the ocean (Crisp 1984). A more recent, separate tabulation of large igneous provinces by Marty & Tolstikhin (1998) finds a total of  $95.5 \times 10^6 \text{ km}^3$  in the past 250 Myr, for a rate of  $0.4 \text{ km}^3/\text{yr}$ . It is broadly agreed that these magmas are volatile-rich compared to MORB. Basing their estimate on  $^3He$  budgets, Marty & Tolstikhin (1998) suggest that the total output from oceanic islands and plumes 'is at best similar to that of spreading centres'. Given our estimate above, this suggests an input of somewhat less than  $1 \text{ Tmol C yr}^{-1}$ .

Since mid-ocean ridge magmas are roughly 10 times more voluminous, the estimated equal flux of mantle  $CO_2$  from oceanic islands and plumes calls for a 10-fold enrichment of  $CO_2$ , and thus perhaps of Nb, in the parent magmas. Observed enrichments of Nb (Hofmann 2004) range from at least 13-fold (Mangaia, Pitcairn, Tahaa) to 2-fold (Mauna Loa). Moreover, Pineau *et al.* (2004) have suggested that  $CO_2 : Nb$  ratios might range to values more than 3-fold higher than that found by Saal *et al.* (2002). In sum, the estimate of  $1 \text{ Tmol C yr}^{-1}$ , equal to that at the spreading centres, is plausible but highly uncertain.

Together, mid-ocean ridges, arc volcanoes and emissions at island volcanoes and during plume events provide an annual input from the mantle of approximately  $2.2 \text{ Tmol C}$ .

### 4. INVENTORIES AND ACCUMULATION OF CARBON

Total quantities of carbonate carbon in the crust, estimated from stratigraphic inventories (Holser *et al.* 1988; Wedepohl 1995; Hunt 1996; Des Marais 2001; Berner 2004; Arvidson *et al.* in press), commonly range

from 2800 to 6500 Emol (Emol = examol =  $10^{18}$  mol). The same reports provide estimates of the total quantity of organic carbon ranging from 675 to 1300 Emol. Only one of these (Arvidson *et al.* in press) includes carbonate associated with basalt. It also yields the lowest ratio of organic to total carbon, namely 0.13. The other reports yield organic fractions ranging from 0.15 to 0.20, and seem to be influenced by isotopic mass balances, which suggest higher relative quantities of organic carbon.

Wilkinson & Walker (1989) took an alternative approach and focused exclusively on sedimentary carbonates. Examining mass-age data, they found that the best fit could be provided by a constant-mass, constant-burial system with first-order recycling and including 9600 Emol carbonate carbon. An alternative fit emphasizing data from younger sequences yielded a result of 7900 Emol. If we arbitrarily adopt an organic-carbon fraction of 0.15, the corresponding inventories of organic carbon are 1700 and 1400 Emol, for total carbon inventories of 11 300 and 9300 Emol.

The most detailed inventory (Holser *et al.* 1988) provides total crustal C = 7640 Emol. Favouring the approach using mass-age data, particularly that based on younger sequences, we weight it equally with the stratigraphic compilations and estimate total crustal C = 8500 Emol.

The time required to accumulate this inventory depends on assumptions about the input flux. If the total flux of  $2.2 \text{ Tmol C yr}^{-1}$  estimated above were constant, the time required would be 3.86 Gyr. More commonly, it is believed that, earlier in Earth history, such fluxes were higher. Here, we follow Sleep & Zahnle (2001) and Lowell & Keller (2003). These authors scale input fluxes to estimates of high-temperature heat flow, chiefly at spreading centres. For the case in which continents grew episodically from 10 to 80% of current area between 3200 and 2500 Myr ago (Ma), high-temperature heat flow is calculated to decrease from  $9\times$  the current level at 3800 Ma, to  $4.9\times$  at 3200 Ma, to  $2.3\times$  at 2500 Ma, and then to decline exponentially. The resulting scaled carbon flux is shown in figure 4.

Integration of that flux beginning at 3800 Ma, the end of the late, heavy bombardment, yields the totals depicted graphically in figure 5. The sum exceeds 8500 Emol, the estimated present crustal inventory, after only 575 Myr, at 3225 Ma. In fact, with continents just beginning to form and crustal storage reservoirs thus sharply restricted, returns of carbon to the mantle would probably have become nearly equal to inputs from the mantle well before then.

Together, figure 5 and equations (1.1)–(2.2) provide a new context for considering the development of the carbon cycle. Before turning to the isotopic records, however, we must first review available information regarding the fates of carbon in subduction zones.

## 5. CARBON CYCLING AT SUBDUCTION ZONES

Processes in subduction zones are crucial to redox balances in the carbon cycle. Figure 6 shows flows of carbon (in oxidized and reduced form) and other

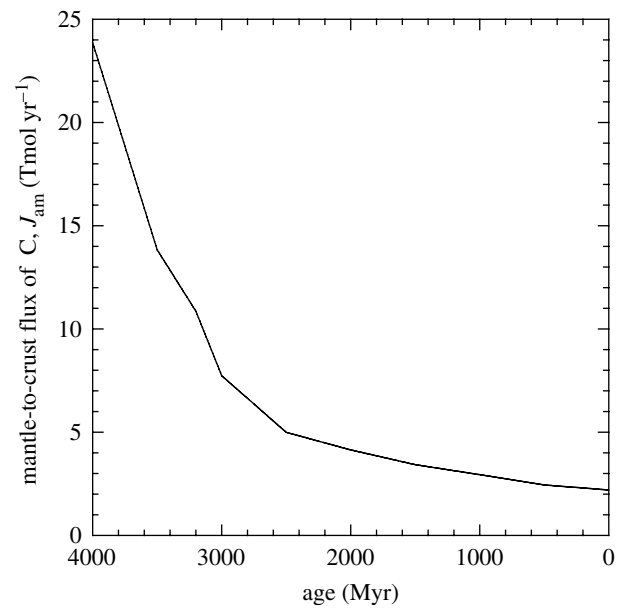


Figure 4. Estimated values of  $J_{am}$  as a function of time. The value at 0 Ma is documented in the text. The scaling relationship at earlier times derives from Sleep & Zahnle (2001) and Lowell & Keller (2003).

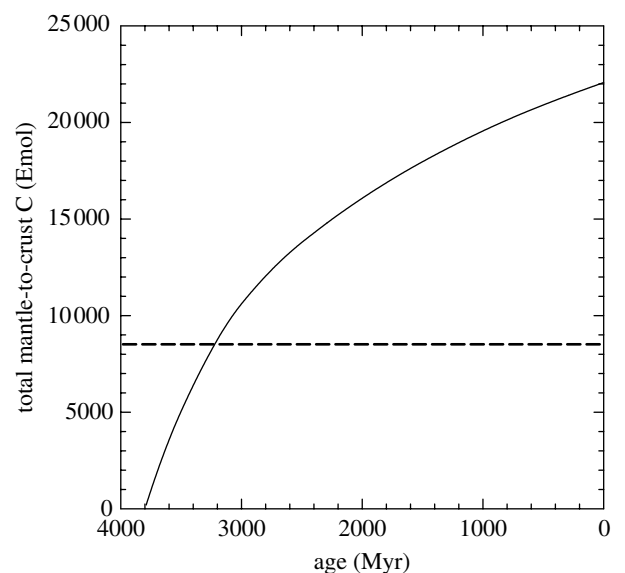


Figure 5. Total amount of mantle carbon delivered to the crust as a function of time. Specifically, total mantle-to-crust C =  $\sum_{3800}^t J_{am}(t)\tau$ , where  $t$  is the age, Myr, and  $\tau$ , the time-step, is  $10^7$  years. The broken line at 8500 Emol C represents the best estimate of the present crustal inventory of C (table 2).

products of carbon cycling into, through and out of subduction zones. Reducing power produced by the carbon cycle is carried into subduction zones by organic carbon ( $J_{os}$ ). Oxidizing power is carried by sulphate and by ferric iron. The crossing paths and redox reactions suggest the diverse transformations involved. While many different sequences of reactions are possible, the equations in figure 6 summarize the required balances of mass and electrons. All of the carbon that enters the subduction zone must be transferred in some form to either the crust or mantle. And the balance between inputs of reductants and oxidants must be reflected by materials leaving the subduction zone. Resolution and dissection of

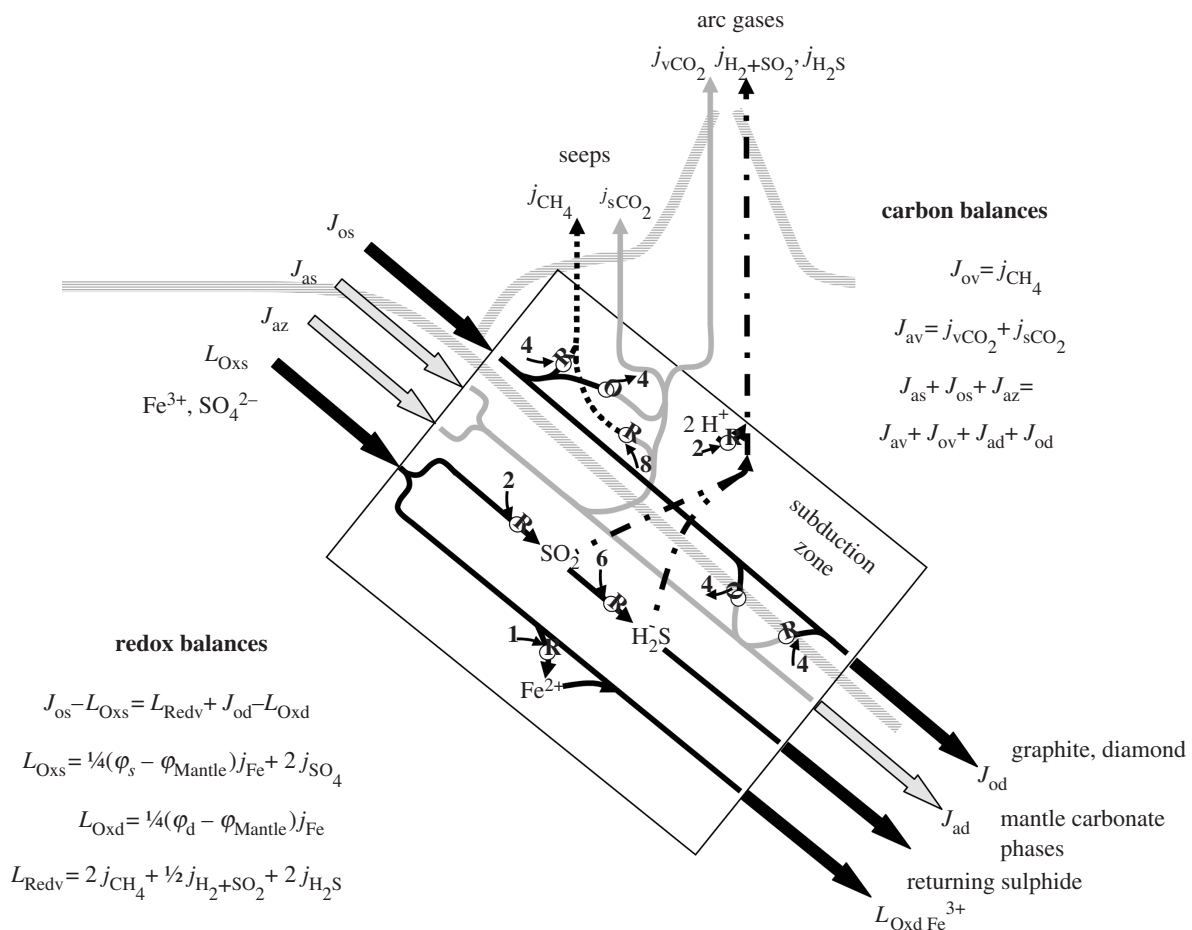


Figure 6. Schematic of fluxes of carbon and reducing power in a subduction zone. Terms denote  $Fe^{3+}/\Sigma Fe$  ( $\varphi$ ), and fluxes of carbon ( $J$ , mol per time), specific substances ( $j$ , mol per time) and oxidizing or reducing power ( $L$ , mol  $O_2$  equivalent per time). Subscripts are defined in table 1. Chemical reductions and oxidations are marked by circled letters (R or O). Adjacent numbers indicate number of electrons gained or lost.

processes within subduction zones are not currently possible, but available evidence bears on two key questions. When a slab is subducted, (i) what happens to the carbon and (ii) what happens to the reducing power carried by the organic matter?

#### (a) Carbon fluxes

The downgoing flux, ( $J_{ad} + J_{od}$ ), might be quantified directly if we knew how much material was being incorporated by the mantle and if samples of it were returned to the surface, so that their carbon contents could be determined. Although  $J_{od}$  and  $J_{ad}$  are sampled by diamond-bearing rocks and mantle xenoliths, it is not clear how representative these samples are, nor do they allow for quantification of the total downgoing C flux.

More headway can be made by considering the fluxes from arc volcanism and other volatile emissions from subduction zones and then estimating the downgoing flux from the difference between the trench input and surface output. The magnitudes of both are rather uncertain; here, we summarize the best estimates. The three inputs are the organic and carbonate components of subducted sediment ( $J_{os}$  and  $J_{as}$ ) and the carbonated basalt in the subducting crust ( $J_{az}$ ). The global subducting-sediment budget of Plank & Langmuir (1998) puts  $J_{as}$  at 0.9 Tmol C  $yr^{-1}$ . Estimates of  $J_{os}$  are

very poorly constrained, mostly because it is small in comparison to fluxes to and from the continents and shelves. For the modern C cycle, Holser *et al.* (1988) estimate  $J_{oc}/J_{os} \approx 40$ . We adopt their value,  $J_{os} = 0.2$  Tmol C  $yr^{-1}$ , suggesting a mean  $C_{org}$  content for subducting sediment of 0.13 wt%.

The third flux,  $J_{az}$ , likely delivers most of the carbon to subduction zones. It derives from  $J_{aw}$  for which estimates range over more than an order of magnitude from less than 1 to more than 3 Tmol C  $yr^{-1}$  (Bach *et al.* 2003). The alternate fate for carbon buried by basalt carbonation is represented by  $J_{ah}$ , the portion of carbonated basalt actually accreted onto continents. Since this is relatively small, it follows that carbonate can be stored in ocean crust along passive margins for hundreds of millions of years, as in much of the Atlantic basin today. The idea that  $J_{az}$  approaches  $J_{aw}$  incorporates an assumption that, following tectonic rearrangements, such material is eventually subducted. The resulting rough estimate of the total subduction input is 2–4 Tmol C  $yr^{-1}$ .

For the volatile outputs from arcs, the most complete budget has been compiled by Hilton *et al.* (2002), who considered subduction inputs to and volatile emissions from 26 arc systems worldwide. The authors calculated a global volcanic arc  $CO_2$  flux of 1.6 Tmol C  $yr^{-1}$  and emphasized the importance of



distinguishing between sources of volatiles. For  $\text{CO}_2$ , these are carbonate and organic C in the subducting slab and  $\text{CO}_2$  in the mantle wedge. Since the mantle component amounts to  $0.2 \text{ Tmol C yr}^{-1}$  (noted above),  $j_{\text{vCO}_2} = 1.4 \text{ Tmol C yr}^{-1}$ . The portion due to subducted carbonates can be estimated from the isotopic composition of the non-mantle component. For all of the arcs, it exceeded the input of sedimentary carbonate. In all cases but one, inclusion of carbonated basalt eliminated the shortfall. Notably, these results are at odds with calculated phase equilibria that indicate carbonated basalts should undergo little devolatilization along most subduction geotherms (Kerrick & Connolly 2001). The other component of  $J_{\text{av}}$  is  $j_{\text{sCO}_2}$ , emission of  $\text{CO}_2$  from seeps, especially in fore-arc and back-arc regions. This flux is unconstrained, and may be as large as  $j_{\text{vCO}_2}$  (Hilton *et al.* 2002). Ingebritsen & Manning (2002) have pointed to diffuse fluid flow through tectonically active crust as potentially a major flux of subduction-derived volatiles. This degassing pathway may be sufficient to reconcile the crust–mantle water balance, and could well constitute a significant return of slab-derived  $\text{CO}_2$  to the crust. Accordingly,  $J_{\text{av}}$  is between 1.4 and  $2.8 \text{ Tmol C yr}^{-1}$ .

The other return flux of carbon from subduction zones to the exogenic chamber is  $J_{\text{ov}}$ , the reduced carbon from high- and low-temperature seeps, primarily  $\text{CH}_4$ . The related geological forms are diverse, and include mud volcanoes and seeps associated with gas hydrates (Milkov & Etiope 2005; Milkov 2005). Their output of  $\text{CH}_4$ , estimated at  $2.1 \text{ Tmol yr}^{-1}$  (Milkov & Etiope 2005), derives from microbial methanogenesis and thermal processes in sediments and sedimentary rocks, as well as from subducted carbon. Since the carbon in  $J_{\text{ov}}$  ( $=\text{CH}_4$ ) has oxidation number equal to  $-4$  and the organic carbon in  $J_{\text{os}}$  has oxidation number equal to zero, redox balance provides the stronger constraint. If  $J_{\text{os}}$  is approximately  $0.2 \text{ Tmol C yr}^{-1}$ ,  $J_{\text{ov}}$  cannot be larger than  $0.1 \text{ Tmol C yr}^{-1}$ . Total volcanic and seep fluxes of carbon ( $J_{\text{av}} + J_{\text{ov}}$ ) are between 1.5 and  $2.9 \text{ Tmol C yr}^{-1}$ .

From inputs of  $2\text{--}4 \text{ Tmol C yr}^{-1}$  and recycling fluxes of  $1.5\text{--}2.9 \text{ Tmol C yr}^{-1}$ , we estimate that the fraction recycled is approximately 0.6. By difference, the flux returning to the mantle is  $0.8\text{--}1.6 \text{ Tmol C yr}^{-1}$ .

### (b) Reducing power

Any reduced C sent to the mantle—graphite, diamond, elemental C—leaves oxidant behind in the surface environment, thus contributing to  $A_{\text{Ox}}$ . It is also possible that organic C would be oxidized within the subduction zone by reaction with sulphate or an oxidized metal, such as  $\text{Fe}^{3+}$ . When this occurs, does the reduction product ( $\text{Fe}^{2+}$ , for example) become an organic-carbon proxy that also contributes to  $A_{\text{Ox}}$ ? No, the reaction instead amounts to a last-minute reversal of processes within the carbon cycle. Within the exogenic reaction chamber,  $\text{Fe}^{3+}$  will have been produced within the downgoing slab by hydrothermal alteration and by oxidation of  $\text{Fe}^{2+}$  at the expense of  $\text{O}_2$ . Or the sulphate will have been produced by processes within the carbon cycle. In either case, the oxidizing power will be represented by an equivalent

quantity of organic carbon. For the carbon cycle, therefore, the oxidation of organic carbon by oxidized metals or sulphur within subduction zones is a functional equivalent of biological respiration. Details follow.

If the  $\text{Fe}^{3+}$  or sulphate is a product of aerobic oxidation, it carries the oxidizing power of  $\text{O}_2$  produced during photosynthesis. The same organism that produced the  $\text{O}_2$  produced an equivalent amount of organic carbon. The oxidation of that organic material within the subduction zone amounts to a reversal of the overall process.

If the  $\text{Fe}^{3+}$  was produced at the expense of sulphate during hydrothermal alteration, that sulphate can similarly be traced to photosynthetic  $\text{O}_2$  and carbon. The chain of chemical events has an additional link, but the oxidation of organic carbon within the subduction zone is again simply a reversal of the process.

If the sulphate was produced by anaerobic, photosynthetic bacteria, those organisms will have produced an equivalent amount of organic carbon. The oxidation of that organic material within the subduction zone amounts to a reversal of the overall process.

Finally, if the  $\text{Fe}^{3+}$  was produced by serpentinization, an equivalent quantity of  $\text{H}_2$  will also have been produced and used by microbiota to produce an equivalent amount of organic carbon, either directly, through chemosynthesis, or indirectly, through consumption of  $\text{O}_2$ , photosynthetic production of organic matter, etc. Again, the oxidation of organic C within the subduction zone amounts simply to a reversal.

If the subducted organic carbon returned to the surface as  $\text{CO}_2$  after reducing an inorganic substance to some oxidation state lower than that of its input—if  $\text{Fe}^{3+}/\Sigma\text{Fe}$  in the downgoing slab were driven to values lower than  $\text{Fe}^{3+}/\Sigma\text{Fe}$  in unaltered MORB—that would contribute to  $A_{\text{Ox}}$ . Failing that,  $J_{\text{od}} > 0$  provides the only route by which processes in subduction zone can yield a net export of oxidizing power by the carbon cycle.

In magnitude,  $J_{\text{od}}$  is constrained to be less than  $J_{\text{os}}$ . Exchange of C between reduced and oxidized pools is not excluded. Some of the C in  $J_{\text{od}}$  might derive from  $J_{\text{as}}$  or  $J_{\text{az}}$ . Estimation of  $J_{\text{od}}$  requires knowledge of  $J_{\text{os}}$  and  $L_{\text{Redv}}$ . The rate of subduction of organic matter, as discussed above, is here taken as  $0.2 \text{ Tmol yr}^{-1}$ , though better estimates are clearly warranted.  $L_{\text{Redv}}$  has two components: the reduced gas flux from arc volcanoes, and the reduced gases from other seeps, represented by  $j_{\text{CH}_4}$ .

The reduced component of the volcanic gas flux is represented by  $j_{\text{H}_2+\text{SO}_2}$  and  $j_{\text{H}_2\text{S}}$ . In the present context, the question is how it relates to  $J_{\text{os}}$ , the reducing power delivered by subduction of organic carbon. The hydrogen abundance in volcanic gases is set by redox equilibrium with water, such that  $\text{H}_2:\text{H}_2\text{O}$  ratios are maintained near *ca* 0.01 at most eruptive temperatures (Giggenbach 1996). Taking the arc magmatic-water flux at  $17 \text{ Tmol yr}^{-1}$  (Wallace 2005) results in a hydrogen component of  $0.17 \text{ Tmol yr}^{-1}$ . The  $\text{SO}_2$  efflux from arcs, some of which may derive from sources other than reduction of subducted sulphate, is of similar magnitude,  $0.16\text{--}0.28 \text{ Tmol yr}^{-1}$  (Halmer *et al.* 2002; Wallace

2005). Other reduced species in arc gases are relatively minor. The ratio of  $\text{H}_2\text{S} : \text{SO}_2$  is generally between 1 and 0.05 (Halmer *et al.* 2002), and some of the hydrogen sulphide released at volcanic arcs likely derives from volatilization of sulphide in the downgoing slab rather than reduction of sulphate by organic carbon. The reducing power carried by other volatiles, such as  $\text{CO}$ ,  $\text{COS}$  and  $\text{CS}_2$ , is orders of lower magnitude. Since  $\text{H}_2$  and  $\text{SO}_2$  represent two-electron reductions, the total flux is halved to convert to moles  $\text{O}_2$  equivalent, whereas  $\text{H}_2\text{S}$  produced from sulphate represents eight electrons or 2 mol  $\text{O}_2$  equiv. The resulting estimate from gas chemistry is 0.2 Tmol  $\text{O}_2$  equiv.  $\text{yr}^{-1}$  from  $\text{H}_2$  and  $\text{SO}_2$  and 0.4–0.02 Tmol  $\text{O}_2$  equiv.  $\text{yr}^{-1}$  from  $\text{H}_2\text{S}$ . By itself, the flux of  $\text{H}_2$  and  $\text{SO}_2$  is already equivalent to the reducing power carried by subducted organic carbon. The obvious presence of additional reduced outputs, namely volcanic  $\text{H}_2\text{S}$  and hydrocarbons at seeps ( $j_{\text{CH}_4}$ ), shows that better knowledge of redox budgets for subduction zones is needed. With due regard for the uncertainties imposed by the present budgets, it also suggests that most or all reducing power carried by subducted organic carbon is returned to the exogenic reaction chamber and that  $J_{\text{od}}$  is small at present.

## 6. CYCLING OF CARBON AND ITS REDOX PARTNERS OVER TIME

Over time, the crust has accumulated carbon. Integrated inputs from the mantle have exceeded integrated returns to the mantle. As a means of exploring the balance, we can accept the fluxes and reservoirs proposed thus far as hypotheses and consider how 8500 Emol C might have accumulated and what electron donors were probably associated with the production of organic carbon.

### (a) *The crust–mantle carbon balance*

Values of  $(J_{\text{ad}} + J_{\text{od}})/J_{\text{am}}$  control the accumulation of crustal C. When they are less than 1, the crustal inventory will grow. Ideally, a geologic record would exist, but proxies for  $(J_{\text{ad}} + J_{\text{od}})$  are rare. The low  $\delta^{13}\text{C}$  values of some diamonds, particularly those of eclogite paragenesis, strongly suggest derivation from crustal organic carbon (Pearson *et al.* 2004), though this has been disputed (Deines *et al.* 2001). Recently, it has been suggested that organic carbon can be subducted beyond 250 km and contribute to sublithospheric diamonds (Tappert *et al.* 2005). The stability of carbonated eclogite at high pressures and temperatures (Dasgupta *et al.* 2004) indicates that eclogitization may be an important route for the subduction of carbon (both oxidized and reduced) into the mantle. This is particularly interesting in light of recent suggestions that eclogitization is a geologically recent phenomenon (Bjørnerud & Austrheim 2004). A hotter upper mantle earlier in Earth history would more efficiently devolatilize downgoing slabs at shallower depths, removing both carbon and water. Shallow decarbonation, combined with drying of the slab and consequent inhibition of the formation of eclogite, may have

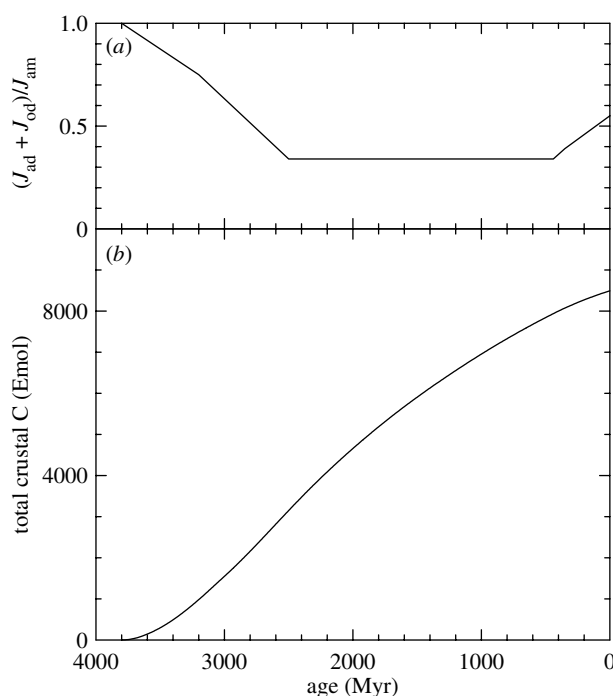


Figure 7. Growth of the crustal inventory of carbon over time. (a) Outputs to the mantle relative to inputs from the mantle as a function of time. The ratio is assumed to be near unity when continents were non-existent and to have declined as they increased in size. Values have been adjusted to provide the observed total of 8500 Emol C by 0 Ma. (b) The integral obtained by applying the return fraction specified in panel (a) to the fluxes shown in figure 4. In detail, total crustal C =  $\sum_{3800} J_{\text{am}}(t)[1 - ((J_{\text{ad}}(t) + J_{\text{od}}(t))/J_{\text{am}}(t))\tau]$ .

meant that  $J_{\text{od}}$  and  $J_{\text{ad}}$  were small early in Earth history (Des Marais 1985). Much depends on the tectonic regime, and when the present style of plate tectonics began, which has been the subject of much debate (Van Kranendonk 2004; Stern 2005).

One view takes the slow growth of continents together with the steep early declines in the rate at which carbon is delivered from the mantle (figure 4) as constraining variations in  $(J_{\text{ad}} + J_{\text{od}})/J_{\text{am}}$  quite strongly. The scenario associated with figure 4 provides 80% of the present continental area by 2.5 Ga. By comparison, Veizer & Mackenzie (2004) point to evidence suggesting that only 25% of continental crust accumulated between 4.0 and 2.6 Ga, another 35% in the interval to 1.7 Ga, and the final 40% thereafter. Because the continents house the major reservoirs of crustal carbon,  $(J_{\text{ad}} + J_{\text{od}})/J_{\text{am}}$  must approach 1 (no net crustal storage) in the early Archaean and decline to lower values only as continents grow. The time course of  $(J_{\text{ad}} + J_{\text{od}})/J_{\text{am}}$  shown in figure 7a fits these criteria while eventually yielding a crustal inventory of 8500 Emol C (figure 7b) and a modern  $(J_{\text{ad}} + J_{\text{od}})/J_{\text{am}} = 0.55$ . The latter value is in the middle of the range estimated above (0.36–0.73).

### (b) *The accumulation of organic carbon and oxidized electron donors*

Reconstruction of redox budgets requires estimates of  $f$ . These derive from interpretation of the carbon isotopic records, with due attention to the possible

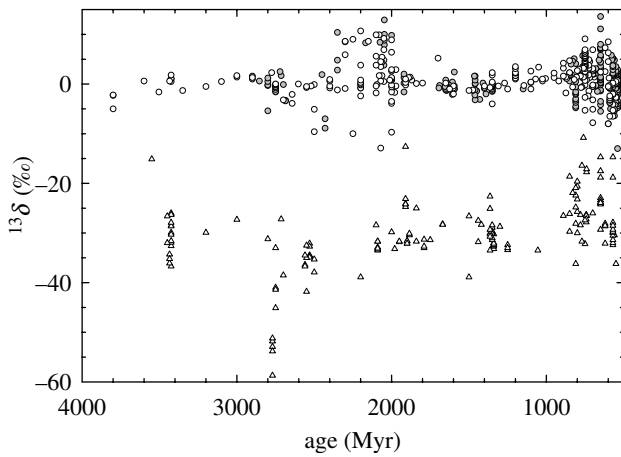


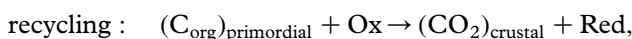
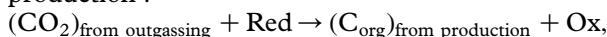
Figure 8. Isotopic compositions of sedimentary carbonates and of sedimentary organic carbon as a function of time. Each point represents the average isotopic composition of an entire stratigraphic unit. Carbonates (Shields & Veizer 2002) are represented by circles. Filled symbols represent geologic units which are well dated. Open symbols represent units for which the dates are approximate. Samples of organic material (Strauss & Moore 1992) are represented by triangles. The ages assigned in the original compilation have been revised to agree with those assigned by Shields & Veizer (2002).

importance of  $\Delta_m$ ,  $\Delta_g$  and  $\lambda$ . Figure 8 summarizes observations of  $\delta_{ab}$  and  $\delta_{ob}$ . The latter represent chemically isolated samples of kerogen (Strauss & Moore 1992). The ratios of H:C in Precambrian kerogens are frequently well below 0.5 (Hayes *et al.* 1983). Accordingly, the kerogens have been extensively dehydrogenated by processes that frequently involve the loss of  $^{13}\text{C}$ -depleted hydrocarbons and thus enrichment of  $^{13}\text{C}$  in the residual kerogen. Because our objective is to reconstruct probable isotopic compositions of organic carbon at the time it was removed from the exogenic reaction chamber (i.e. as it flowed through the arrow representing  $J_{ob}$  in figure 1), we will base our estimates of  $f$  mainly on the lower values of  $\delta_{ob}$  in each time interval.

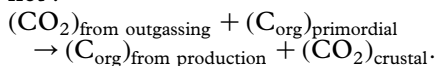
#### (c) 4400–3800 Ma

In parallel with the isotopic records summarized in figure 8, the integrations in figures 5 and 7 have begun at 3800 Ma. This point in time roughly marks the end of the late, heavy bombardment and the beginning of Earth's sedimentary record. Catling *et al.* (2001) have considered how earlier events might have set the stage for carbon cycling. In particular, they estimate that impacts of asteroids between 4400 and 3800 Ma probably delivered 1000 Emol of reduced carbon to Earth's surface. To whatever extent such material was not incorporated by the mantle, Earth's crust will have begun with an inventory of primordial organic carbon. When biological cycling of carbon began, the processes would have been

production :



net :



In these equations, Red and Ox are reduced and oxidized forms of a redox partner, such as Fe or S. Reactions between surviving, asteroidal organic material (termed 'primordial') and oxidants produced during carbon cycling might have been biologically catalysed and occurred within the exogenic reaction chamber or they might have been thermal and occurred only at depth within accumulating sediments. In either case, the resulting  $\text{CO}_2$  became part of the crustal inventory. Such reactions would have two effects: (i) an exchange of biotic organic carbon for primordial organic carbon and (ii) consumption of oxidants. Only when production acted to increase the inventory of  $\text{C}_{\text{org}}$  (here taken to begin at 3.8 Ga) would oxidants begin to accumulate.

#### (d) 3800–2800 Ma

For this interval, the isotopic record of sedimentary carbonates compiled by Shields & Veizer (2002) includes 24 stratigraphic units. With an average value of  $\delta_{ab} = -0.1\text{‰}$ , they are consistently enriched in  $^{13}\text{C}$  relative to mantle carbon. The standard deviation of the population is  $1.9\text{‰}$ . For the same interval, figure 8 yields an estimate of  $\delta_{ob} = -36\text{‰}$ . Nakamura & Kato (2004) measured values of  $\delta_{aw}$  in basalts from the Warrawoona Group (3425 Myr, Western Australia). Their average result,  $-0.3 \pm 1.2\text{‰}$ , does not differ significantly from  $\delta_{ab}$  in sedimentary carbonates in this time interval. Accordingly,  $\Delta_m \approx \Delta_g$ , the  $\lambda(\Delta_m - \Delta_g)$  terms in equation (1.8) are small, and  $f = (\delta_{ab} - \delta_i) / (\varepsilon - \Delta_g) = (\delta_{ab} - \delta_i) / (\delta_{ab} - \delta_{ob}) = 0.14$ . Observed values of  $\delta_{ab}$  and  $\delta_{ob}$  then indicate that 14% of the carbon being delivered to the exogenic reaction chamber by outgassing of  $\text{CO}_2$  from the mantle was being buried as reduced, organic carbon.

The net production of organic carbon during any increment of time can be estimated:

$$\begin{aligned} \text{net } \text{C}_{\text{org}} = & (\text{C}_{\text{org}} \text{ produced by reduction of } J_{\text{am}}) \\ & + (\text{C}_{\text{org}} \text{ produced by reduction of recycling C}) \\ & - (\text{C}_{\text{org}} \text{ oxidized during recycling}). \end{aligned} \quad (6.1)$$

The recycling carbon (i.e.  $J_{\text{or}} + J_{\text{ar}} + J_{\text{ov}} + J_{\text{av}}$ ) will be some portion of the total crustal carbon. If a representative half-mass age is taken as 300 Myr, the fraction recycling in a 10 Myr interval will be 2.3% (see table 1). In any 10 Myr interval, therefore, 2.3% of crustal carbon will be recycled, essentially mixing with the 10 Myr input of mantle carbon. Of that total, a portion controlled by  $(J_{\text{ad}} + J_{\text{od}}) / J_{\text{am}}$  will be returned to the mantle. Of the remainder, a fraction  $f$  will be buried in organic form. Assuming that all recycling organic carbon is oxidized, an integrable form of equation (6.1) is then

$$\text{net } \text{C}_{\text{org}} = f \left[ J_{\text{am}} \tau \left( 1 - \frac{J_{\text{ad}} + J_{\text{od}}}{J_{\text{am}}} \right) + \gamma M \right] - \gamma M_o, \quad (6.2)$$

where  $\gamma$  is the fraction of crustal carbon recycling in a time interval of length  $\tau$  (e.g. 0.023 for  $\tau = 10^7$  years and a half-mass age of 300 Myr), and, at the beginning of the time-step,  $M$  is the total crustal inventory of carbon in all forms (figure 7b) and  $M_o$  is the total inventory of organic carbon. Stepwise summation

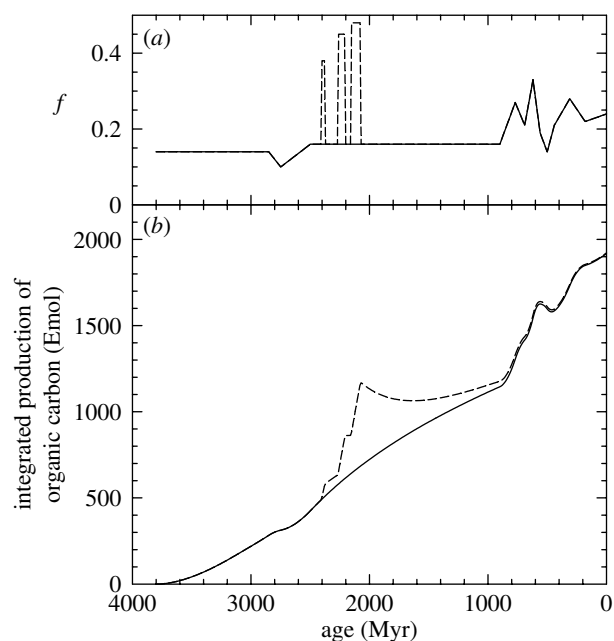


Figure 9. Integrated net production of organic carbon, equivalent to the net release of oxidizing power (Emol  $O_2$ ), over time. (a) Estimated values of  $f$ . For ages greater than 890 Myr, they are based on the isotopic compositions shown in figure 8. For younger ages, they are consistent with the isotopic compositions shown in figure 8 and are based in detail on records summarized by Hayes *et al.* (1999). The broken lines between 2080 and 2400 Myr represent isotopic excursions reviewed by Melezhik *et al.* (1999) and discussed in the text. (b) The integral obtained when the  $f$  values shown in figure 9a and the carbon-return ratios shown in figure 7a are substituted in equation (6.2), with  $\gamma=0.023$  and  $M=M_0=0$  at 3800 Ma.

provides an estimate of the integrated production of organic carbon, the first integral on the right-hand side of equation (2.2).

The calculation is based on (i) values of  $(J_{ad} + J_{od})/J_{am}$  chosen to provide the observed crustal inventory of C (figure 7a); (ii) values of  $J_{am}$  provided by figure 4; and (iii) values of  $f$  shown in figure 9a and estimated from the observations summarized in figure 8. Setting  $M=M_0=0$  at 3800 Ma and summing Net  $C_{org}$  yields the result shown in figure 9b. Because a sharp drop in  $\delta_{ob}$  suggests a major change in carbon cycling at approximately 2770 Ma (Hayes 1983, 1994), we will focus first on the interval 3800–2800 Ma. By 2800 Ma,  $M_0 = \text{oxidant release} = 300 \text{ Emol } O_2 \text{ equiv.}$

To place this result in context, consider the inventories summarized in table 2. For all of the materials considered, a range of estimates can be found in the literature. Those included here are representative high and low values. The value of 300 Emol  $O_2$  equiv. greatly exceeds the oxidizing power represented by  $O_2$  in the modern atmosphere and ocean and approaches the low estimate of the amount of oxidizing power represented by total crustal deposits of sulphate and sulphate dissolved in seawater. These comparisons decisively eliminate two otherwise-interesting possibilities, namely that O or S could have served as the dominant redox partner for C at this stage of Earth history. In the first case, levels of  $O_2$  would have risen high enough to inhibit the mass-independent fractionation of

Table 2. Crustal inventories of reduced and oxidized products of carbon cycling<sup>a</sup>.

product	estimated crustal total Emol $O_2$ equivalent		reference
	high	low	
$O_2$	37.2	37.2	Keeling <i>et al.</i> (1993)
$Fe^{3+b}$	1860	1020	Goldschmidt (1954) Ronov & Yaroshevsky (1976)
$SO_4^{2-}$	480	332	Garrels & Perry (1974) Holser <i>et al.</i> (1988)
total oxidants	2377	1389	
organic C	1280	675	Des Marais (2001) Arvidson <i>et al.</i> (in press)

<sup>a</sup> A similar compilation has been prepared by Catling *et al.* (2001). It includes an estimate of 1000 Emol organic C delivered to the crust by impacts of asteroids between 4400 and 3800 Ma. The entries above pertain to the present. To whatever extent primordial organic material has survived in the crustal inventory, it is included in the 'organic C' category.

<sup>b</sup> Calculated from data summarized by Lécuyer & Rickard (1999). Represents  $Fe^{3+}$  in excess over mantle  $Fe^{3+}/\Sigma Fe=0.12$  (Bezou & Humler 2005).

sulphur isotopes. In the second, concentrations of dissolved sulphate would have climbed to levels that lead to large differences between the  $^{34}S$  contents of sedimentary sulphides and sulphates, a signal that is not observed until later (Hayes *et al.* 1992).

The remaining oxidized product listed in table 2 is ferric iron. Its crustal inventory is large enough to offer plenty of headroom. Iron formations are, moreover, important components of the Archaean sedimentary record. The present context, however, requires that iron served as the electron donor in primary production while the environment remained strictly anaerobic. An organism capable of utilizing  $Fe^{2+}$  directly has recently been isolated (Jiao *et al.* 2005). In principle, we need look no further, but alternatives deserve consideration. These are: (i) the true value of  $f$  was really near zero; (ii)  $f$  was 0.14 but no oxidants accumulated; or (iii)  $f$  was 0.14 and the electron donor was  $H_2$ , for which the immediate oxidized product ( $H_2O$ ) was invisible, although iron was probably the ultimate source of the electrons.

The true value of  $f$  could be approximately zero if the basalt-carbonation correction had been wrongly excluded and the correct value of  $\lambda(\Delta_m - \Delta_g)$  was 4.9‰. If  $\lambda$  were 0.9, this would require that the initial average value of  $\delta_{aw}$  had been  $-5.5\text{‰}$  and that Nakamura & Kato (2004) found instead  $-0.3\text{‰}$ , because the isotopic compositions of all of their samples had been affected by post-depositional carbon-isotopic exchange. However, among 42 samples of four different types, the most negative value of  $\delta_{aw}$  is  $-3.4\text{‰}$  and there is no correlation between  $\delta$  and the abundance of carbonate. Evidence for the required alteration is therefore lacking, making it likely that  $f$  truly approached 0.14.

If  $f$  approached 0.14 but no oxidants accumulated ((ii), above),  $(J_{ad} + J_{od})/J_{am}$  must have been 1,

indicating rapid and quantitative return of all carbon to the mantle, even as 3.5 million km<sup>2</sup> of continental crust (Lowe 1992) formed prior to 3.2 Ga. Moreover, as substantial quantities of organic carbon were forming as required by  $f=0.14$  (and thus  $J_{\text{ob}}/J_{\text{am}} \geq 0.14$ ), the putatively non-accumulating oxidant, which could not be either O<sub>2</sub> or sulphate and which had to be present only at trace levels, was able to reoxidize all of the organic material completely even as the carbon itself returned quantitatively to the mantle. The required combination of circumstances is practically impossible.

The third alternative, in which  $f$  was approximately 0.14 and H<sub>2</sub> served as the electron donor, fits well with multiple lines of evidence. Anaerobic organisms capable of producing organic material from CO<sub>2</sub> and H<sub>2</sub> are abundant and include both chemoautotrophs (methanogens, acetogens) and photoautotrophs (photosynthetic bacteria; Tice & Lowe 2004 have already postulated that such organisms were important members of near-shore microbial communities 3.4 Ga). The potential versatility of microbial ecosystems based on such organisms is great enough to provide both producer and consumer assemblages and thus to stabilize carbon cycling and yield globally consistent isotopic fractionations. Fermentative consumption would remobilize organic C as a mixture of CO<sub>2</sub> and CH<sub>4</sub>, providing needed greenhouse warming (Kasting & Catling 2003) and setting the stage for the isotopic transient observed at 2.8–2.7 Ga (Hayes 1994).

The required net input of H<sub>2</sub> would be 600 Emol (= 300 Emol O<sub>2</sub> equiv.). Owing to the postulated declines in  $J_{\text{am}}$  (figure 4) and increases in  $[1 - (J_{\text{ad}} + J_{\text{od}})/J_{\text{am}}]$  (equation (6.2)), the required rate would rise only slowly from 0.5 Tmol yr<sup>-1</sup> at 3500 Ma to 0.9 Tmol yr<sup>-1</sup> at 2800 Ma. If, as is likely, conditions at the seafloor favoured serpentinization reactions (Sleep *et al.* 2004), a flux of 0.7 Tmol H<sub>2</sub> yr<sup>-1</sup> could be provided by alteration of less than 10<sup>15</sup> g Mg-rich (komatiitic) basalt per year. The present rate of generation of ocean crust is about 6 × 10<sup>16</sup> g yr<sup>-1</sup>, so the requirement poses no problem. In the overall sequence of electron transfers, it is only the immediate oxidized product of carbon fixation (H<sub>2</sub>O) which is 'invisible'. The reducing power represented by the organic matter is balanced by Fe<sup>3+</sup>, which is accumulating or being subducted.

To summarize, from both geological and biological points of view, there are highly plausible mechanisms by which substantial amounts of organic carbon could be produced and buried in accord with the isotopic records, with complementary releases of oxidizing power, without any requirement for generation of O<sub>2</sub> during the interval 3800–2800 Ga. The most likely redox partner for C during this interval is Fe, either directly via phototrophic oxidation of dissolved Fe<sup>2+</sup>, and/or indirectly, with H<sub>2</sub> shuttling electrons from Fe<sup>2+</sup> in seafloor basalts to phototrophic and chemoautotrophic producers.

#### (e) 2800–1800 Ma

The next billion years begin with the apparent onset of oxygenic photosynthesis (Hayes 1983, 1994; Summons *et al.* 1999), which includes a period during

which the abundance of <sup>13</sup>C in sedimentary carbonates was sometimes markedly enriched (Karhu & Holland 1996; Melezhik *et al.* 1999), and ends with the likely onset of sulphidic conditions in the deep sea (Canfield 1998).

Estimates of  $f$  derived from the isotopic records provide scant evidence that the development of oxygenic photosynthesis, for all its magnificence as a biochemical innovation, provided a significant increase in the net release of oxidizing power. If anything,  $J_{\text{ob}}$  initially declined, a feature marked by the notch visible at 2.8 Ga in figure 9a. By 2500 Ma,  $f$  appears to have risen to 0.16, a value only slightly higher than that estimated for the early Archaean. At 2450 Ma, when traces of O<sub>2</sub> quench the mass-independent fractionation of sulphur isotopes, the estimated minimal total release of oxidizing power by the C cycle is 460 Emol O<sub>2</sub> equiv. Although uncertainties in that total are very large, it is far short of the capacities of crustal Fe and S to supply electrons (table 2). Levels of O<sub>2</sub> in the environment would have been sharply limited by the strength of those sinks.

What, then, happened between 2400 and 2000 Ma? Contents of <sup>13</sup>C in carbonates are frequently elevated, occasionally exceeding +10‰ (figure 8). Accepting these compositions as representative of the global pool of oceanic DIC would imply a near-trebling of  $f$ . Writing three years after Karhu & Holland (1996) published 'Carbon isotopes and the rise of atmospheric oxygen', Melezhik *et al.* (1999) drew on additional data to define three apparently separate pulses of isotopic enrichment. That dissection of the record yields the three spikes in  $f$  that are represented by broken lines in figure 9a (alternative stratigraphic correlations yield only two spikes; Bekker *et al.* 2003). The isotopic signal has been associated with other lines of geochemical evidence indicating that the atmosphere became more oxidizing at about the same time. The ensemble has become known as 'The Great Oxidation Event' (Holland 2002).

One view of the scale of the event is provided by the broken line in figure 9b. If the isotopic signals represent pulses of carbon burial, the integrated production of organic carbon by 2000 Ma would be 600 Emol greater. As shown by table 2, that is more than enough to provide the present inventories of SO<sub>4</sub><sup>2-</sup> and O<sub>2</sub>. But there are problems with interpreting these isotopic variations in terms of carbon burial.

- (i) There is no parallel isotopic enrichment in the organic carbon, as would be expected if the carbonate represented the oceanic DIC that was the carbon source for primary producers. Karhu & Holland (1996) searched for a signal in the organic-carbon record and found its absence 'rather odd'. If the record were not so fragmentary and noisy (preservation is a much greater problem for organic carbon than for carbonate minerals) that the problem could be set aside (Melezhik *et al.* 1999), this alone would be fatal to the interpretation.
- (ii) Evidence is lacking for the large deposits of organic carbon that should have formed (Melezhik *et al.* 1999; Aharon 2005).

- (iii) The sequence of isotopic signals is reversed from that expected. If the oxygenation that cut off mass-independent fractionation of sulphur isotopes was due to an organic-carbon-burial event, the disappearance of the mass-independent fractionation of sulphur isotopes should not precede the first carbon-isotopic enrichments.
- (iv) It is difficult to envision supplies of nutrients great enough to sustain the levels of productivity required to supply the organic carbon. Aharon (2005) has discussed supplies of phosphate very insightfully, concluding that efficient stripping of P from organic matter prior to burial provides the only solution. To achieve the projected values of  $f$ , the C : P ratio in buried organic matter would have to be 4000 and the organic-preservation rate would have to be 6.6%, about fivefold higher than that observed in the Black Sea.
- (v) Stratigraphic correlations are not secure enough to demonstrate that the various isotopic excursions are coeval, and thus necessarily linked to variations in a global reservoir (Aharon 2005).
- (vi) Most of the units displaying the isotopic enrichments are dolostones with substantial isotopic variability. In their list of 12 'major localities,' Melezhik *et al.* (1999) report within-unit ranges of 3–13‰ and an average range of 7‰. Consistent with this, for the interval 2350–2000 Ma (inclusive), Shields & Veizer (2002) report 589  $\delta$ -values between –2.5 and +2.5‰ and 424 between +5 and +13‰. That is, 'normal' isotopic abundances are more common than elevated abundances.
- (vii) When complete chemical analyses of the carbonates are reported, they often include substantial concentrations of SiO<sub>2</sub> (to 40%; Melezhik *et al.* 1999). Stratigraphic columns indicate that some of the isotopically enriched carbonates are closely interbedded with shales (Buick *et al.* 1998), others are described as 'nodular' (Bekker *et al.* 2003).

The latter features (vi) and (vii) are more characteristic of diagenetic carbonates than of marine limestones faithfully carrying records of oceanic DIC. When it is recalled that values of  $^{13}\delta$  to +13‰ are common in much younger dolomites that have been affected by methanogenic diagenesis (Klein *et al.* 1999; Mazzullo 2000), that alternative interpretation demands attention. In fact, attribution of the isotopic signals to methanogenic diagenesis has already been favoured by several sets of authors (Yudovich *et al.* 1991; Dix *et al.* 1995).

In all likelihood, the diagenetic alternative has failed to win popularity because an alternative does not appear to be needed. Multiple, convergent lines of evidence—independent of carbon-isotopic signals—indicate that thresholds of environmental oxidation were crossed during this time interval (Bekker *et al.* 2004). When the carbon-isotopic record fits into this picture at least roughly, why not include it? Even more to the point, given the apparent reality of the oxidation,

how else should the carbon-isotopic enrichments be explained? They are temporally associated, dramatic and numerous (Bekker *et al.* 2003), globally distributed, and have the right polarity (enrichment rather than depletion).

#### (f) *Mechanisms of oxygenation*

The problem lies not in accepting the isotopic signals as markers of the oxidation but in assuming that they represent the *cause*. This is a key distinction. Either the steady functioning of the carbon cycle catalysed biological and ecological developments that shifted the relative strengths of oxygen sources and sinks (Holland 1978) or dramatic changes in carbon fluxes were required. If the former, the carbon-isotopic enrichments are environmental reporters comparable to the mass-independent fractionations of sulphur isotopes. If the latter, they point to increased burial of organic carbon but not necessarily to permanent changes in oxidation.

Flux-driven changes are subject to reversal. Buried organic material will be recycled by erosion and volcanism. As indicated by the downward trend of the broken line in figure 9b after 2000 Ma, oxidants will be consumed. Even if a rise in O<sub>2</sub> were driven by increases in  $J_{ob}$ , some additional change would be required to stabilize the transition and make it permanent. Geophysical phenomena such as rifting and increased sedimentation can increase  $J_{ob}$ , but are inherently cyclical. What they accomplish in one epoch will be undone in another. Increased burial of organic carbon is a half answer. It can push O<sub>2</sub> levels higher, but maintaining them will require some further change, a second half to the answer.

Unidirectional change, a permanent strengthening of the source of oxidants, is in the realm of evolutionary biology. Physiological and ecological changes could stabilize higher levels of O<sub>2</sub>. Do they serve as the second half of the answer? If so, they are the biological results of geophysical stimuli and we need an explanation for the linkage. Or are physiological and ecological changes answers in themselves? If so, advances in oxygenation have largely biological origins and combinations of disparate phenomena are not required.

Kinetic factors are also pertinent. In discussing the carbon cycle, it is common to distinguish between the *fast*, biological cycle of photosynthesis and respiration and the *slow*, geological cycle in which erosion, weathering and volcanism are balanced by carbon burial. In global chemical terms, O<sub>2</sub> is a highly reactive, transient intermediate that is produced and consumed within the exogenic reaction chamber. Its steady-state abundance must depend on the relative strengths of sources and sinks in that system. In contrast, sedimentary carbonates and organic materials are outputs from that system. Their relative abundances are controlled by geophysical factors that vary over relatively long time-scales.

#### (g) *Causes of the carbon-isotopic transient*

We can suggest a sequence of biologically driven environmental changes, associated with a rise in levels of O<sub>2</sub>, that would produce the observed isotopic signals without requiring enhanced burial of organic carbon.

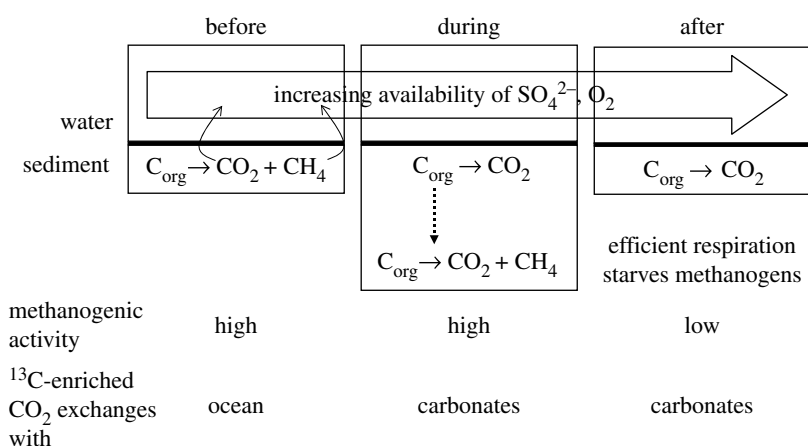


Figure 10. Summary of a sequence of conditions that could account for the abundance of  $^{13}\text{C}$ -enriched sedimentary carbonates 2400–2080 Myr ago.

They would also account for the absence of congruent variations in the organic-carbon isotopic record.

Especially prior to 2.3 Ga, methanogenic diagenesis must have been common. If electrons for biosynthesis were supplied mainly by  $\text{Fe}^{2+}$ , the insolubility of the oxidized product,  $\text{Fe}^{3+}$ , meant that electron acceptors were rare in most surface environments (Walker 1987). When respiring heterotrophs are excluded, the recycling of organic carbon in microbial communities will be managed by fermenters, with methanogens playing a vital role. This will have occurred in the same zones that are now occupied by aerobes. These methanogenic communities will have exchanged  $\text{CO}_2$  freely and directly with the oceanic water column, and, as a result, the isotopically enriched pools of  $\text{CO}_2$  that are characteristic of methanogenic diagenesis in modern environments will not have developed. This situation is summarized in the 'before' segment of figure 10.

The development of oxygenic photosynthesis by at least 2700 Ma began to change that situation, however slowly. Particularly near oxygen-producing communities (many of the isotopically enriched dolostones are associated with stromatolites; Melezhik *et al.* 1999), methanogens will eventually have been pushed to deeper levels in the sediment. When this occurred, apparently beginning at about 2400 Ma, isotopically enriched pools of porewater DIC will have developed. As carbon was exchanged with the carbonate minerals in the sediment (a process catalysed by the organic acids produced during fermentation), the isotopic signal characteristic of methanogenesis will have been recorded for the first time. Within about 400 Myr, a second event shut the signal off. Fundamentally, it must have been some weakening of the supply of organic material to methanogenic communities. As suggested in the 'after' segment of figure 10, it could have been caused by a further rise in the availability of electron acceptors. Alternatively, the tight association between stromatolitic producers and methanogenic consumers, indicated by many of the  $^{13}\text{C}$ -enriched sequences, might have been destabilized by environmental changes.

As in the Neoproterozoic, the isotopic excursions are associated with apparently extreme glaciations (Young *et al.* 2001; Tajika 2003). The effects of those

glaciations on global redox geochemistry, if any, are unclear. It has conversely been proposed that the glaciations were caused by global oxidation which destroyed a methane greenhouse *and* that this occurred promptly after the development of oxygenic photosynthesis (Kopp *et al.* 2005). The second part of this does not fit with biomarker records that place the advent of oxygenic photosynthesis at or before 2.7 Ga (Summons *et al.* 1999; Brocks *et al.* 2003; Eigenbrode 2004). It also does not fit with the inescapably slow effects of carbon cycling. Releases of oxidizing power have been continuous (figure 9b). But the oxidized products accumulated by 2.5 Ga cannot have accounted for more than a fraction of the inventory of ferric iron and almost none of the sulphate (cf. table 2). In those circumstances, by what mechanism did the development of oxygenic photosynthesis promptly sustain steady-state concentrations of  $\text{O}_2$  high enough to destroy the methane greenhouse? Until that question can be answered, the evidence for (i) a 400 Ma delay between the evolutionary event and its environmental impact and (ii) the continuing capacity of Fe and S as redox buffers should overrule speculation.

Oxidation of S by  $\text{O}_2$  would have produced sulphate in surface environments. However, by 1800 Ma, the estimated integrated release of oxidizing power (figure 9b) was only 820 Emol  $\text{O}_2$  equiv. This is still well below the low estimate of the amount required to build the crustal inventory of  $\text{Fe}^{3+}$  (table 2). As suggested by Canfield (1998), therefore, it is likely that production of sulphate mainly provided a route, via sulphate-reducing bacteria, to delivery of excess sulphide to deep ocean waters and the consequent cessation of deposition of banded iron formations.

#### (h) 1800 Ma–present

Carbon-isotopic records from the Neoproterozoic onward are good enough that values of  $f$  can be estimated with some confidence (Hayes *et al.* 1999). The results are crudely summarized in figure 9a. When these values of  $f$  are applied to the carbon fluxes estimated here, the projected final total release of oxidizing power is 1920 Emol  $\text{O}_2$  equiv. This is close to the average estimate of the total release represented by the sum of crustal  $\text{Fe}^{3+}$ ,  $\text{SO}_4^{2-}$  and  $\text{O}_2$ .

The agreement is notable, but discrepancies within the redox accounts are more impressive. Estimates of crustal inventories of oxidants vary by a factor of 1.7. Similarly, the estimated crustal inventories of organic carbon fall short (perhaps by two times) of the summed inventories of oxidized products and of the estimated total burial of organic carbon (figure 9b). At face value, this requires that reduced products have been exported from the crust more efficiently than oxidized products. Unless organic carbon could be exported to the mantle more efficiently than  $\text{Fe}^{3+}$ , we are practically required to attribute such an imbalance to loss of  $\text{H}_2$  from the top of the atmosphere. The redox imbalance and the likely importance of  $\text{H}_2$  loss have been incisively examined and elaborated by Catling *et al.* (2001) and by Catling & Claire (2005). They associate loss of  $\text{H}_2$  mainly with atmospheric  $\text{CH}_4$  and thus with the Archaean. By contrast, Tian *et al.* (2005) have noted the possible importance of the low exobase temperature of an anoxic atmosphere in limiting the rate of  $\text{H}_2$  escape from the early Earth. They contend that higher levels of  $\text{O}_2$  in the atmosphere after 2.4 Ga would have increased exobase temperatures and promoted escape of hydrogen. Resolution of the history of hydrogen escape will constrain the second integral on the right-hand side of equation (2.2), and be a key step in reconstructing the time course of  $A_{\text{Ox}}$ .

The dearth of reduced products becomes particularly notable when the likely integrated effects of subduction, represented by the third integral in equation (2.2), are considered. Lécuyer & Rickard (1999) contend that the *net* rate at which subduction is transferring  $\text{Fe}^{3+}$  from the crust to the mantle is presently  $7 \text{ Tmol yr}^{-1}$ , or  $1.8 \text{ Tmol O}_2 \text{ equiv. yr}^{-1}$ . If the continents are at steady state with respect to organic carbon, so that, in equation (6.2),  $f\gamma M + \gamma M_o = 0$ , the rate at which the carbon cycle is releasing oxidizing power is given by (cf. equation (6.2))

$$f[J_{\text{am}} - (J_{\text{ad}} + J_{\text{od}})] = 0.24(2.2 - 1.2) \\ = 0.2 \text{ Tmol O}_2 \text{ equiv. yr}^{-1}, \quad (6.3)$$

where the values substituted are centre points of the ranges estimated in earlier sections of this paper. Since the net rate of subduction of  $\text{Fe}^{3+}$  proposed by Lécuyer & Rickard (1999) greatly exceeds this value, it is either incorrect or represents a temporary situation. Attention to subduction of  $\text{Fe}^{3+}$  is, however, a very good idea. The continuing release of oxidizing power, coupled with the minimal variations in atmospheric levels of  $\text{O}_2$  during the Phanerozoic (Berner 2004) and balanced exchange of oxidizing power by the cycles of carbon and sulphur during the same interval (Canfield 2004), requires a continuing sink. It seems inescapable that this is supplied by oxidation of  $\text{Fe}^{2+}$  at the seafloor.

Bounds can be placed on the extent of this transfer over Earth history, since iron returning to the mantle with a higher  $\text{Fe}^{3+}/\Sigma\text{Fe}$  than it emerged with will result in oxidation of the upper mantle. There is a growing consensus that the redox state of the upper mantle, as measured by its oxygen fugacity, has not changed greatly over Earth history. The recent constraint of Li & Lee (2004), from the V/Sc systematics of MORBs,

indicates that the oxygen fugacity of the mantle source has increased by at most 0.3 log unit since 3.5 Ga. Using the relation of Kress & Carmichael (1991), this translates into a maximal 6% increase in the ferric iron content of the mantle, if all of that increase were attributable to addition of  $\text{Fe}^{3+}$ . Taking the mass of the upper mantle to be  $1.34 \times 10^{27} \text{ g}$  (Ballentine *et al.* 2002), the mantle Fe content to be 6.3 wt% (Palme & O'Neill 2003) and the mantle  $\text{Fe}^{3+}/\Sigma\text{Fe}$  to be 0.12 (Bezous & Humler 2005), this yields a maximal input of  $1.1 \times 10^{22} \text{ mol}$  of ferric iron since 3.5 Ga, or a maximum average input rate of  $3 \text{ Tmol Fe}^{3+} \text{ yr}^{-1}$  ( $0.75 \text{ Tmol O}_2 \text{ equiv.}$ ). This is comfortably within potential loads imposed by  $A_{\text{Ox}}$ .

The agreement between the projected release of oxidizing power and the average estimate of oxidized products is far from comforting. Because considerable amounts of  $\text{Fe}^{3+}$  must have been exported to the mantle, the total release of oxidizing power by the carbon cycle should exceed considerably the inventory of  $\text{Fe}^{3+}$  remaining in the crust. Part of any shortfall can be accommodated by recalling that the present estimates are minima. Finite values of  $J_{\text{od}}$  and a failure to reoxidize organic carbon in  $J_{\text{or}}$  will increase  $A_{\text{Ox}}$ .

The reburial of organic carbon can be treated quantitatively. If a portion,  $x$ , of  $J_{\text{or}}$  is simply reburied, two things will happen: (i) since it does not re-enter the exogenic system as  $\text{CO}_2$ , that C will be unavailable for reduction. The organic burial flux and thus the overall production of oxidizing power will be decreased by an amount  $f x J_{\text{or}}$ ; (ii) the oxidizing power consumed by oxidation of  $J_{\text{or}}$  will be reduced by an amount  $x J_{\text{or}}$ . The first term is a loss of oxidizing power, the second is a gain. The difference is  $(1-f)x J_{\text{or}}$ . Unfortunately, this term cannot simply be added to equation (6.2). Although it would account for the carbon and redox balances in a single increment of time, the proper treatment of subsequent increments would require that the reburied material be followed, with appropriate adjustments being made to sedimentary and continental inventories. Even during the Phanerozoic, the presence of detrital coal and kerogen in marine sediments (Sackett *et al.* 1974) shows that  $x$  is not zero. On the other hand, it is known that microorganisms in weathering profiles incorporate carbon even from refractory kerogens (Petsch *et al.* 2001) and that the great bulk of organic carbon in marine sediments is *not* recycled, detrital organic material. Under anaerobic conditions, however, reburial of organic carbon is likely to have been more important.

Can the carbon cycle operate at a point of redox neutrality, neither releasing nor consuming oxidants, even when isotopic compositions indicate significant rates of burial of organic carbon? The answer is in equation (6.3). Even if burial and recycling are balanced, redox neutrality still requires  $J_{\text{am}} = J_{\text{ad}} + J_{\text{od}}$ . In other words, inputs from the mantle are equal to outputs to the mantle. This is not a likely coincidence. Carbon flows into subduction zones not just from  $J_{\text{am}}$  but from multiple sources. If the portion of  $J_{\text{as}} + J_{\text{os}} + J_{\text{az}}$  equal to  $J_{\text{am}}$  is to be returned to the mantle while the balance flows to  $J_{\text{av}}$  and  $J_{\text{ov}}$ , a very clever demon will be required to operate the carbon valve in the subduction zone. Notably,  $f$  (and thus  $\delta_{\text{ab}} - \delta_i$ ) is not a measure of



$J_{\text{am}} - (J_{\text{ad}} + J_{\text{od}})$ . No particular value of  $\delta_{\text{ab}}$  can be recognized as indicating redox neutrality. This yields two fundamental insights. First, caution is required when interpreting the isotopic record in terms of redox variations. Second, the conditions required for redox neutrality are precise and improbable.

## 7. CONCLUDING REMARKS

We have taken a new approach to mass and redox balances involving the C cycle. Its principal strength is a realistic acknowledgement of the importance of inputs from and outputs to the mantle. This broader context has invited consideration of processes that may otherwise be overlooked or misconstrued. Its weakness is that our knowledge of some of the key processes is presently imprecise. We need to learn more about inputs of carbon, hydrogen escape and processes in subduction zones to better evaluate each component of  $A_{\text{Ox}}$ . Even now, however, the resulting uncertainties do not cripple the approach. The key finding is dramatic: the persistence of  $\delta_{\text{ab}}$  at values near zero rather than  $-5\%$  can be interpreted only in terms of continuous and substantial releases of oxidizing power by the carbon cycle. Two points follow.

The oxidation of the crust has been more continuous than episodic. Increases in steady-state levels of oxidants derive either from combinations of geological and biological changes or from biological changes alone. The scientific problem of reconstructing Earth's oxidation is more biological than geochemical.

Continuous sinks for oxidizing power are required, at least since atmospheric levels of  $\text{O}_2$  stabilized at current levels approximately 550 Ma. Since exchanges of oxidizing power between the carbon and sulphur cycles have been balanced during that same time interval, this can have been provided only by the oxidation of iron.

Together, these affect our view of the carbon cycle. It should not be seen as the competent and versatile manager of an electron-transfer market, effortlessly balancing offers and bids. Instead, it is a wrong-way passenger on a downgoing escalator. It continuously receives new  $\text{CO}_2$  from the mantle, relentless supplies of solar energy and nutrients leached from deposits that have retained their organic carbon. To hold its place, it must constantly produce new organic matter. While doing so, it releases oxidants that threaten to reverse its accomplishments. The stability of  $\delta_{\text{ab}}$  is an indicator of the success of its evolutionary strategies. The stability of  $P_{\text{O}_2}$  is evidence of an equally competent sink.

The seeds from which the ideas discussed have grown were planted when J.M.H. looked out of *Alvin's* viewports and observed the  $\text{H}_2$ -rich waters of the Lost City hydrothermal vent system. The generosity of Deborah Kelley and Jeffrey Karson, the co-chief scientists of the Lost City cruise, is greatly appreciated. Since then, we have benefited from helpful and stimulating exchanges with Stan Hart, Don Anderson, David Rowley and Alberto Saal. We are very grateful for critical reviews of this manuscript provided by David Catling and Jennifer Eigenbrode. J.M.H. receives support as a member of the Astrobiology team led by S. D'Hondt, University of Rhode Island. J.R.W. is supported by an NDSEG Graduate Fellowship from the Office of Naval Research.

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