The influence of air-sea exchange on the isotopic composition of oceanic carbon: Observations and modeling

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Abstract. Although the carbon isotopic composition of ocean waters after they leave the surface ocean is determined by biological cycling, air-sea exchange affects the carbon isotopic composition of surface waters in two ways. The equilibrium fractionation between oceanic and atmospheric carbon increases with decreasing temperature. In Southern Ocean Surface Waters this isotopic equilibration enriches δ¹³C relative to the δ¹³C expected from uptake and release of carbon by biological processes alone. Similarly, surface waters in the subtropical gyres are depleted in δ¹³C due to extensive air-sea exchange at warm temperatures. Countering the tendency toward isotopic equilibration with the atmosphere (a relatively slow process), are the effects of the equilibration of CO₂ itself (a much faster process). In regions where there is a net transfer of isotopically light CO₂ from the ocean to the atmosphere (e.g., the equator) surface waters become enriched in ¹³C, whereas in regions where isotopically light CO₂ is entering the ocean (e.g., the North Atlantic) surface waters become depleted in ¹³C. A compilation of high quality oceanic δ¹³C measurements along with experiments performed using a zonally averaged three-basin dynamic ocean model are used to explore these processes.

Introduction

Although it has been generally accepted that the distribution of δ¹³C of oceanic carbon is primarily controlled by biological cycling of ¹³C depleted organic matter (Kroopnick, 1985), only recently has attention been focused on the effects of air-sea exchange on oceanic δ¹³C (Oppo and Fairbanks, 1989; Charles and Fairbanks, 1990). Broecker and Maier Reimer (1992) and Charles et al. (1993) show how deep and intermediate water δ¹³C is influenced by air-sea exchange in the regions of water mass formation. The effect of air-sea exchange on surface ocean δ¹³C has been demonstrated using multibox models (Keir, 1991; 1993) as well as a general circulation model (Broecker and Maier-Reimer, 1992). It is important to fully understand the distribution of oceanic δ¹³C in the ocean because it is one of the few chemical signals from past oceans that is reliably recorded by foraminifera and preserved in the sediment record. As such, foraminiferal δ¹³C has been and will continue to be widely used to infer past states of the oceanic carbon cycle.

Here, we re-examine some previously published high-quality oceanic δ¹³C measurements in order to evaluate the relative strengths of air-sea exchange and biological cycling in various oceanic regimes. In addition, we use a two-dimensional ocean model which includes a carbon cycle (Wright and Stocker, 1992; Stocker et al., 1994) to examine the influence of air-sea exchange and to test our inferences about the effects of these processes in the real ocean. The two-dimensional model is ideal for such experiments in that it allows for the quick (relative to a full general circulation model) exploration of various scenarios which require up to 8000 years of integration for a steady state distribution of the carbon isotopic ratio, while retaining some essential aspects of the ocean circulation.

Controls on δ¹³C of Oceanic Carbon

Biological Control

During photosynthesis, organisms preferentially take up the lighter isotope of carbon (¹²C) increasing surface ocean δ¹³C. When the isotopically light organic material is remineralized, the δ¹³C of seawater dissolved inorganic carbon (DICO₂) decreases. This fractionation of about -19‰ for marine photosynthesis leaves the nutrient depleted surface ocean with a high δ¹³C, and the nutrient enriched deep waters with a low δ¹³C. If this process were the only one responsible
for the distribution of δ13C in the ocean, δ13C of oceanic CO2 would decrease by approximately 1.1 % for every 1 μmol kg⁻¹ increase in oceanic PO4 [Broecker and Maier-Reimer, 1992]. Since oceanic PO4 values have a range of about 3 μmol kg⁻¹, biological controls have the potential to produce a 3.3 % range in oceanic δ13C values. This linear relationship between PO4 and δ13C depends on a constant C/P ratio in organic material of 128, and a constant δ13C fractionation of -19 % relative to oceanic carbon. However, in reality, the discrimination against 13C during photosynthesis increases from about 19 % in the warm surface ocean to about 30 % in the Antarctic. The computed δ13C versus PO4 slope due to biologic processes alone in Antarctic Surface Waters would then be about -1.7 %/μmol kg⁻¹ instead of the average ocean slope of -1.1 %/μmol kg⁻¹. Rau et al. [1989] attribute the low δ13C of organic material in the Southern Ocean to the increased pool of aqueous CO2 in these cold waters. This means that at times in the past when atmospheric CO2 was different, the δ13C of organic material formed in the surface ocean could have changed [Rau et al., 1989; 1991], causing changes in the slope of δ13C versus PO4 relationship for uptake and decay of organic matter. The production of calcium carbonate shells by surface organisms and the subsequent dissolution of this shell material at depth will also affect the distribution of δ13C within the ocean. However, since this carbonate has an isotopic composition similar to the seawater in which it grew, the impact of this process on oceanic δ13C distributions is very small.

Equilibrium Fractionation During Air-Sea Exchange

Carbon isotope fractionation during air-sea exchange is also an important factor in determining the isotopic composition of carbon in surface seawater. If the CO2 in the atmosphere were in isotopic equilibrium with the dissolved inorganic carbon ([ΣCO2] = [HCO3⁻] + [CO3²⁻] + [CO2(aq)]) in the ocean, the dissolved inorganic carbon would be enriched in 13C relative to the atmospheric CO2 by about 8 % at 20°C [Inoue and Sagisura, 1985; Zhang et al., 1995]. This enrichment is the result of the dominance of HCO3⁻ (about 90 % of ΣCO2) with its enrichment relative to the atmosphere of 8.5 % [Mook et al., 1974] with smaller contributions from CO3²⁻ (about 10 % of ΣCO2) at 6 % enriched relative to the atmospheric value [Zhang et al., 1995] and still smaller contributions (about 1 % of ΣCO2) of CO2(aq) which is depleted by about 1 % relative to the atmosphere [Vogel et al., 1970]. All of these equilibrium fractionation factors depend on the temperature of equilibration, with the HCO3⁻ and, thus ΣCO2, becoming more enriched relative to the atmospheric value by about 0.1 % per degree of cooling [Mook et al., 1974]. If the surface ocean were in complete isotopic equilibrium with atmospheric CO2, the 30°C range in ocean temperatures would cause a 3 % range in oceanic δ13C, similar to the magnitude of δ13C change induced by biological processes. The relationships between the equilibrium isotopic fractionation factors for each dissolved species relative to the atmosphere is summarized in Figure 1.

For a 50-m deep mixed layer it would take about 10 years to equilibrate carbon isotopes between the atmosphere and the ocean (Figure 2), 10 times longer than for CO2 itself.

Figure 1. Equilibrium isotopic fractionation of seawater dissolved carbon species relative to atmospheric CO2. Fractionation factors are from Mook et al. [1974], Vogel et al. [1970] and Zhang et al. [1995].

[Broecker and Peng, 1974, Tans, 1980]. This is the result of the fact that ΣCO2 can equilibrate with the atmosphere through exchanges between the atmospheric CO2 and CO2 (aq) and CO3²⁻ pools (about 10 % of ΣCO2), but for carbon isotopic (δ13C of ΣCO2) equilibrium, all ΣCO2 must eventually exchange with the atmosphere. Because surface waters move about and are replaced on faster timescales than this, there is no region of the ocean where surface ocean carbon is in complete isotopic equilibrium with the atmosphere [Broecker and Maier-Reimer, 1992].

Figure 2. Results of a one-box mixed layer carbon cycle model which illustrates the effects of cooling the surface ocean on surface ocean δ13C and ΣCO2. The box is taken to be 50 m deep and exchanges only with the atmosphere. Model calculations for air-sea exchange are the same as those for the two-dimensional ocean circulation model. Alkalinity is held constant at 2300 µeq, atmospheric pCO2 is held at 280 µatm, and atmospheric δ13C is held constant at -6.5 %. Temperature is decreased from 20°C to 5°C at time = 0. Surface ocean δ13C takes about 10 times longer to reach the new equilibrium values than does surface ocean ΣCO2.
The influence of isotopic equilibration on the $\delta^{13}C$ of $\Sigma CO_2$ will depend not only on physical properties within the ocean but on biological processes as well. For example, if biological processes bring the $\delta^{13}C$ of $\Sigma CO_2$ closer to equilibrium with atmospheric carbon (e.g., biological drawdown in cold waters which will elevate $\delta^{13}C$), the subsequent impact of air-sea exchange on this water mass will be lessened. On the other hand, if biological processes tend to bring $\delta^{13}C$ farther from equilibrium with the atmosphere, the impact of air-sea exchange will be much greater. This process is illustrated by Zahn and Keir (1994) using a box model which includes biological cycling, but imposes isotopic fractionation for air-sea exchange that is independent of temperature. For this model, the $\delta^{13}C$ is lower than would be predicted from biological cycling alone in the low nutrient surface waters (low latitude and North Atlantic) where biological processes tend to raise $\delta^{13}C$, and higher than would be predicted for high nutrient surface waters (Southern Ocean) where biological cycling alone would cause low $\delta^{13}C$.

Effects of Invasion and Evasion of Atmospheric $CO_2$

The exchange of isotopically light (relative to oceanic $\Sigma CO_2$) atmospheric $CO_2$ and surface ocean $CO_2$ (aq) has the potential to leave oceanic carbon depleted in $^{13}C$ in areas of $CO_2$ invasion (e.g., high northern latitudes, low $pCO_2$ surface waters), and enriched in $^{12}C$ in areas of the ocean where $CO_2$ is outgassed (e.g., equatorial upwelling, high $pCO_2$ surface waters). As atmospheric $CO_2$ invades the ocean (Sigma $CO_2$ equilibration) and enters the $\Sigma CO_2$ (aq) pool, the isotopic composition of the $\Sigma CO_2$ becomes lower, because the increase in $\Sigma CO_2$ directly results from the increase in isotopically lighter $CO_2$ (aq). Similarly, it is the isotopically lighter $CO_2$ (aq) that leaves the oceanic carbon pool when $CO_2$ evades to the atmosphere, leaving the remaining $\Sigma CO_2$ enriched in $^{13}C$. This can be thought of as a Rayleigh-type distillation, which occurs when the phase leaving the bulk solution ($CO_2$ (aq)) has an equilibrium isotopic composition that is offset from the bulk solution ($\Sigma CO_2$). After $\Sigma CO_2$ has reached a new equilibrium concentration, the $\delta^{13}C$ of $\Sigma CO_2$ will eventually also re-equilibrate with the atmosphere, but as discussed above, this process takes about 10 times as long as $\Sigma CO_2$ equilibration. A simple one-box mixed layer model illustrates the effects of invasion/evasion of isotopically light $CO_2$ and the eventual return to isotopic equilibrium (Figure 3). As discussed above, the time surface waters spend at one temperature is short enough that carbon isotopic equilibration with the atmosphere is never achieved. So, in the areas where oceanic $pCO_2$ is greater than that of the atmosphere, that is, areas which lose $CO_2$ to the atmosphere (like the equator), $\delta^{13}C$ of $\Sigma CO_2$ will be enriched by this process, and in areas where oceanic $pCO_2$ is less than that of the atmosphere, that is, where the ocean gains $CO_2$ (like the North Atlantic), the $\delta^{13}C$ of $\Sigma CO_2$ should be correspondingly depleted.

The largest gain of atmospheric $\Sigma CO_2$ by surface waters due to the natural, unperturbed carbon cycle are found in the far North Atlantic (+100 $\mu\text{mol kg}^{-1}$), where nutrient (and biologically linked $\Sigma CO_2$) poor waters are cooled rapidly, increasing the solubility of $CO_2$ [Broecker and Peng, 1992]. Areas where carbon rich waters are upwelled at low latitudes, the equatorial Pacific and Arabian Sea, show losses of carbon ($\sim$25 $\mu\text{mol kg}^{-1}$) to the atmosphere. The gain of 100 $\mu\text{mol kg}^{-1}$ of carbon from the atmosphere in the North Atlantic would induce a change of -0.5% in surface water $\delta^{13}C$ ($\delta^{13}C$ of invading $CO_2$ is 10% lighter than surface ocean $\delta^{13}C$; invasion of $CO_2$ will cause an increase of 5% in size of the surface $\Sigma CO_2$ reservoir). The total range in the amount of carbon that has been gained or lost to the atmosphere throughout the ocean is about 125 $\mu\text{mol kg}^{-1}$ [Broecker and Peng, 1992], which would induce a range of about 0.6% in surface waters due to this effect. This is a small but significant effect when compared with the changes due to air-sea isotopic equilibration and biological cycling.

Both physical and biological processes within the ocean will influence the location and intensity of the invasion and evasion of carbon to/from the ocean and thus impact the changes in oceanic $\delta^{13}C$ due to this invasion/evasion. The
physics of ocean circulation will produce gradients in surface ocean pCO2 with higher pCO2 at the equator where waters are upwelled and warmed, and low pCO2 in polar regions where waters are cooled. These patterns are modified by biological processes which intensify the high pCO2 at the equator because the upwelled waters contain extra carbon from the organic material that has been re-mineralized into the water since it has left the surface. Similarly, the tendency for physical processes (temperature) to produce a negative pCO2 anomaly in the Southern Ocean is somewhat diminished due to the large amounts of carbon in this nutrient-rich water.

Kinetic Fractionation During CO2 Exchange

Because CO2 exchange between the ocean and atmosphere is not an equilibrium process, with some regions of the ocean degassing CO2 and others absorbing CO2, we must also consider the kinetic isotopic fractionation factors involved in these one-way transfers. The transfer of CO2 across the air-sea interface involves the processes of molecular diffusion. All else being equal, heavier CO2 molecules (those containing 13C) will diffuse more slowly than lighter molecules (those containing 12C). Theory [Siegenthaler and Münich, 1981] predicts that the CO2 leaving the ocean (for a sea-to-air transfer) will be depleted in the heavy isotope by about 0.5‰ in addition to the equilibrium fractionation, and similarly, the CO2 leaving the atmosphere (for an air-to-sea transfer) will be depleted in the heavy isotope by about 0.5‰ relative to the equilibrium fractionation. Experiments [Inoue and Sugimura, 1985; Wanninkhof, 1985; Jähne et al., 1987] suggest kinetic fractionation factors between -0.9 and -1.3 ‰. However, theory predicts kinetic fractionation for a diffusive transfer of gas between the liquid and gas phases, and the laboratory experiments are also performed under such diffusive conditions. In the ocean, where the gas transfer is more turbulent, the kinetic fractionation will be less, and in the extreme case of air-sea transfer via bubble injection, there will be no kinetic fractionation [Knox et al., 1992]. In either case, the effects of kinetic fractionation during invasion/evaporation are very small and will be masked by the effects of the temperature dependent equilibration and by the invasion/evaporation of CO2 which has a lower δ13C than oceanic carbon, 1ΣCO2.

Observed Oceanic δ13C Distributions

Surface Ocean Data

Although it has been shown that surface ocean δ13C lies somewhere in between the biologically predicted value and the value which would be at equilibrium with the atmosphere [Broecker and Peng, 1982; Broecker and Maier-Reimer, 1992], a more detailed examination of the distribution of surface ocean δ13C demands high-quality surface ocean δ13C and nutrient data in order to determine the relative influence of biological and air-sea exchange processes. Here we will use the δ13C data collected along 150°W and published by Kroopnick et al. [1977] along with surface PO4 and temperature data from this cruise. We also use the Southern Ocean Ettannin 1968 surface water δ13C transect along 135°E [Kroopnick, 1974], again along with surface PO4 and temperature data from this cruise. The Geochemical Ocean Sections Study (GEOSECS) Indian Ocean δ13C measurements with accompanying PO4 and temperature measurements were used, but the δ13C data sets from the Atlantic and Pacific GEOSECS expeditions were not included as they exhibited considerably more scatter than the Indian Ocean data set. We also include data from the Transient Tracers in the Ocean (TTO) program in the North Atlantic [Oistlund and Groll, 1987] supplemented by TTO δ13C data analyzed at Lamont-Doherty Earth Observatory (Table 1), as well as data from the Weddell Sea and South Atlantic Ocean reported by Mackensen et al. [1993]. We subtract 0.15 µmol kg⁻¹ from the Weddell Sea (PS16) PO4 measurements reported by Mackensen et al. [1993] to bring the PO4 values into agreement with the GEOSECS PO4 for these water masses. Location of transects are shown in Figure 4. The surface ocean δ13C, temperature, and PO4 data are plotted in Figure 5.

Air-sea exchange and surface water δ13C.

Broecker and Maier-Reimer [1992] show that if there were no air-sea exchange, the relationship between δ13C and PO4 in the ocean would be

\[ \delta^{13}C_{\text{eq}} = \frac{A_{\text{photo}}}{\Sigma CO_2} \delta^{13}C_{\text{P}} (\text{PO}_4 - \text{PO}_4^{\text{M,O}}) \]

When reasonable values are substituted (\( A_{\text{photo}} = 0.3 \), \( \Sigma CO_2 = 2200 \mu\text{mol kg}^{-1} \), C/P:Oeq = 128, PO4^{M,O} = 2.2 µmol kg⁻¹), the predicted relationship between δ13C and PO4 closely matches the relationship for waters in the deep Indian and Pacific Oceans (δ13C = 2.7 - 1.1 x PO4). This is to be expected, as the effect of air-sea exchange should be constant for these deep waters due to the homogeneity of source waters for the deep Indian and Pacific Oceans. We can determine the degree to which air-sea exchange processes (both the tendency toward isotopic equilibration and the effects of invasion/evaporation) have affected surface ocean δ13C by subtracting the δ13C value predicted from biological cycling alone from the actual δ13C:

\[ \delta^{13}C_{\text{obs}} = \delta^{13}C - (0.7 - 1.1 \times \text{PO}_4) \]

This is the same as \( \Delta \delta^{13}C \) defined by Broecker and Maier-Reimer [1992] but has been renamed \( \delta^{13}C_{\text{obs}} \) at the air-sea exchange signature, because \( \Delta \delta^{13}C \) has been frequently used in the paleoceanographic community to denote gradients of δ13C within the ocean. By definition, water with \( \delta^{13}C_{\text{obs}} \) of 0‰ has the same air-sea exchange signature as mean ocean deep water. A positive value of \( \delta^{13}C_{\text{obs}} \) simply means that the water has a higher δ13C than mean ocean deep water (more of an influence of air-sea exchange at cold temperatures/ climates at warm temperatures), whereas a negative δ13C value implies less δ13C enrichment due to air-sea exchange than for mean ocean deep water. The air-sea exchange signature, \( \delta^{13}C_{\text{obs}} \), should not be the same as predicted \( \delta^{13}C \), which will be determined both by the effects of air-sea exchange and biological cycling. This formulation for \( \delta^{13}C_{\text{obs}} \) will only represent the true effects of air-sea exchange if the assumptions about constant C/P ratios in organic material, constant δ13C of organic material, and reasonable biologically induced changes in oceanic carbon content are sufficiently accurate. In Antarctic Surface Waters where δ13C of organic material is substantially lower than for the rest of the ocean, \( \delta^{13}C_{\text{obs}} \) will overestimate the effects of air-sea exchange.
The data sets used in this study were collected over a span of 21 years (1968 to 1989) during which time the atmospheric \( \delta^{13}C \) has changed from -7.1 to -7.8 \% due to the addition of isotopically light fossil fuel \( \text{CO}_2 \) [Keeling et al., 1989]. Atmospheric \( \text{CO}_2 \) increased from 320 to 351 ppm during this same time period. So both via isotopic equilibration with an atmosphere which is constantly decreasing in \( \delta^{13}C \), and increased invasion of isotopically light atmospheric \( \text{CO}_2 \) into the surface ocean, we expect \( \delta^{13}C \) and also \( \delta^{13}C_{\text{as}} \) to decrease with time for the data sets we examine, and we expect that "natural" preanthropogenic \( \delta^{13}C \) would be higher than any of the data shown. However, the processes involved in determining the modern ocean \( \delta^{13}C \) should be the same as those which determined \( \delta^{13}C \) in the preindustrial ocean, and the study of these anthropogenically altered \( \delta^{13}C \) data sets should provide insight into these processes.

\( \delta^{13}C_{\text{as}} \) is highest in Antarctic Surface Water and in the North Pacific (Figure 5). This enrichment of \( \delta^{13}C \) is most likely due to exchange with the atmosphere at low temperatures. Although evasion of carbon from surface waters can also enrich the surface water \( \delta^{13}C \), in these areas more inorganic carbon has invaded from the atmosphere than is lost to it [Broecker and Peng, 1992], which would tend to decrease \( \delta^{13}C \). Although the residence time of Antarctic Surface Waters is not particularly long (about 2 years, [Gordon, 1988]), this is an area of high winds and thus high gas exchange rates. If we took into account the low \( \delta^{13}C \) of organic material formed in Antarctic Surface Waters, we would obtain a lower \( \delta^{13}C_{\text{as}} \) (by about 0.4 \% at the most) which would imply somewhat less intense air-sea exchange at low temperatures to account for the Antarctic Surface Water \( \delta^{13}C \). In all four Antarctic data sets we see high \( \delta^{13}C_{\text{as}} \) values in the vicinity of the Antarctic Polar Front, but in the Weddell Sea transect we also see lower \( \delta^{13}C_{\text{as}} \) values corresponding to the contribution of upwelled Circumpolar Deep Water (\( \delta^{13}C_{\text{as}} = 0 \% \)) in the Weddell Gyre. In the North Pacific, gas exchange at low temperatures results in the high \( \delta^{13}C_{\text{as}} \) in the northern end of the Hudson transect.

The lowest values of \( \delta^{13}C_{\text{as}} \) are found in the centers of the subtropical gyres. Surface waters circulate in these anticyclonic gyres for a relatively long period of time giving the oceanic carbon ample time to equilibrate isotopically with the atmosphere at high temperatures despite the relatively low rates of gas exchange. Intermediate \( \delta^{13}C_{\text{as}} \) values are found in the Subantarctic Surface Waters which result from a mixture of subtropical surface waters with low \( \delta^{13}C_{\text{as}} \) and Antarctic Surface Waters with high \( \delta^{13}C_{\text{as}} \) to the south.

The situation in the North Atlantic seems slightly more complex. The \( \delta^{13}C_{\text{as}} \) in the subpolar gyre, while higher than the subtropical gyres, is still not as high as cold waters in the North Pacific and Southern Ocean. Air-sea exchange at cold temperatures will tend to raise the \( \delta^{13}C_{\text{as}} \) of the northward moving upper waters but is unable to erase the signature of high temperature isotopic exchange from the subtropical gyres. However, also keeping the surface ocean \( \delta^{13}C_{\text{as}} \) low is the invasion of up to 160 \( \mu \text{mol} \ \text{kg}^{-1} \) of isotopically light \( \text{CO}_2 \) (both natural and fossil fuel) from the atmosphere (calculated from TTO \( \Sigma \text{CO}_2 \), S, \( \text{PO}_4 \), \( \text{NO}_3 \), and alkalinity data as by Broecker and Peng [1992]), which will tend to lower \( \delta^{13}C_{\text{as}} \) by 0.7 \%. In the far North Atlantic the \( \delta^{13}C_{\text{as}} \) remains low,

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Table 1. TTO \( \delta^{13}C \) Data Analyzed at Lamont-Doherty Earth Observatory

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\( \delta^{13}C \) is referenced to PDB
Biological processes and surface water $\delta^{13}C$.
The $\delta^{13}C$ of surface water is determined partly by the air-sea exchange processes discussed above, and partly by biological cycling of carbon isotopes. In the Southern Ocean, $\delta^{13}C$ increases from south to north all the way to the subtropical convergence as predicted by the decrease in $\text{PO}_4$ from south to north. However, the $\delta^{13}C$ rises by less than the full amount predicted by the $\text{PO}_4$ concentration, due to the influence of air-sea exchange at warmer temperatures to the north. The $\delta^{13}C$ trend also reflects the biologically predicted trend near the equator (between 10°S and 5°N on the Pacific transect), indicating that in this region, too, air-sea exchange cannot mask the effects of biological drawdown of isotopically light carbon. However, in the subtropical gyres the trend in $\delta^{13}C$ shows minima in the central $\text{PO}_4$ free portions of the gyre where biological drawdown alone would produce a $\delta^{13}C$ maximum. In these convergent gyres, air-sea exchange at high temperatures has sufficient time to mask the biological portion of the $\delta^{13}C$ signal.

Deep Ocean Data

The distribution of $\delta^{13}C$ of $\Sigma\text{CO}_2$ in the deep ocean primarily reflects the biological regeneration of isotopically light carbon, and, in general, is directly correlated to oxygen.
concentrations and inversely correlated to nutrient and $\Sigma$CO$_2$ concentrations [Deuser and Hunt, 1969; Craig, 1970; Kroopnick, 1985]. However, subsurface $\delta^{13}$C distributions are also controlled to a significant extent by the air-sea exchange signature imprinted at the surface. Broecker and Maier-Reimer [1992] suggest that North Atlantic Deep Waters (NADW) have lower $\delta^{13}$C by 0.4 % than would be predicted for an ocean with no air-sea exchange, and that Weddell Sea Bottom Water (WSBW) is enriched in $\delta^{13}$C by 0.2 %. Charles et al. [1993] use selected GESECS $\delta^{13}$C and PO$_4$ profiles to show that these air-sea exchange signatures are carried by deep water throughout the ocean. They also note that Antarctic Intermediate Water which is derived in large part from Antarctic Surface Waters is significantly enriched in $\delta^{13}$C due to air-sea exchange at low temperatures. Here we present a compilation of high quality deep water (below 2 km depth) $\delta^{13}$C and PO$_4$ data from all three oceans (Figure 6). This figure is almost identical to the relationship shown by Broecker and Maier-Reimer [1992], except that the WSBW is more enriched in $\delta^{13}$C by air-sea exchange ($\delta^{13}$C$_{se} = 0.4 \%$) and does not mix directly with NADW to form the North Atlantic mixing line. Instead, it is a shallower component of Circumpolar Deep Water, a mixture of WSBW, NADW, and returning deep waters from the Pacific, which mixes into the Atlantic Basin via the Vema Passage. As shown by Broecker and Maier Reimer [1992], the waters entering the deep Indian and Pacific Oceans derive from shallow (more NADW enriched) points along this North Atlantic mixing line via the Atlantis Passage and the Samoa Passage, respectively.

**Modeling Oceanic $\delta^{13}$C**

**Model Description**

In order to fully explore the effects of air-sea exchange and biological processes on oceanic $\delta^{13}$C, we use the two-dimensional dynamic ocean model of Wright and Stocker [1992], as modified and described by Stocker et al. [1994] (state A). The zonally averaged balances of momentum, mass, potential temperature, salt and tracers are solved in the three major basins of the world ocean with realistic surface area and volume (+/-5%). The geometry and the numerical grid agree with Stocker et al. [1994] except for an increased vertical resolution (21 levels).

The circulation is driven by east-west pressure differences that are parameterized in terms of the meridional density gradient. Fluxes of heat and freshwater at the surface are obtained by restoring sea surface temperature and salinity to the zonally averages of Levitus [1982]. To compensate for a shallower first layer (25 m instead of 50 m), the timescale for the relaxation of temperature and salinity to surface values was

![Figure 6. High quality deep water $\delta^{13}$C and PO$_4$ data from below 2000 m in the Atlantic (triangles), Pacific (squares), and Indian (crosses) Oceans. Data for the Atlantic Ocean includes the GESECS data analyzed at Scripps Institution of Oceanography [Kroopnick et al., 1972; Kroopnick, 1980], corrected by subtracting 0.1 % from $\delta^{13}$C values as suggested by Kroopnick [1980], data analyzed at Lamont Doherty Earth Observatory from the TTOI expedition (Table 1), and data from the Atlantic sector of the Southern Ocean [Mackensen et al., 1993]. Also plotted are GESECS Test Station I from the Pacific Ocean [Kroopnick et al., 1970] and all GESECS data from the Indian Ocean [Kroopnick, 1985]. Phosphate and $\delta^{13}$C data for the GESECS stations were obtained from the National Oceanic Data Center. Weddell Sea Bottom Water (WSBW) has high $\delta^{13}$C$_{se}$ reflecting the contribution of Antarctic Surface Waters. Circumpolar Deep Water (CDW) reflects the mixing of WSBW with NADW and returning flow from the Pacific and Indian Oceans. Atlantic data reflects mixing between North Atlantic Deep Water (NADW) and CDW which enters the Atlantic via the Vema Passage (VP). Deep waters in the deep Indian and Pacific follow the "Redfield Slope" (parallel lines) reflecting the regeneration of organic material into CDW which enters these oceans via the shallower Atlantis Passage (AP) and Samoa Passage (SP), respectively.](image-url)
lowered from 50 to 25 days. The effect of wind stress is included. Properties are exchanged between the ocean basins through the southern ocean where zonal mixing is instantaneous (timescales associated with the deep circulation are much longer than those of recirculation in the Antarctic Circumpolar Current (ACC)).

We further modify the model presented by Stocker et al. [1994], by shutting off the enhanced zonal mixing ($K_H = K_P$) not only south of 40°S but also north of 45°N as well. The enhanced zonal mixing is intended to simulate the zonal mixing by gyres, and should not be necessary across the subtropical/subpolar gyre boundary at 45°N. This modification influences the sea surface temperature field and yields better agreement between modelled and observed δ13C in the North Atlantic.

We use the inorganic carbon cycle described by Stocker et al. [1994]. Total carbon is carried within the ocean as a tracer. Oceanic pCO2 is determined in the surface layer using the chemical equilibria for the carbonate system. In this inorganic model, alkalinity and borate concentration are affected solely by evaporation and precipitation. They can therefore be obtained by scaling to the surface salinity value. The net flux of carbon ($F_\Sigma$) through air-sea interface is proportional to the difference between surface ocean and atmospheric pCO2:

$$F_\Sigma = F_{as} - F_{as} = k p\text{CO}_2\text{atm} - k p\text{CO}_2\text{ocean}$$

where $k$ is the gas exchange rate, $F_{as}$ is the flux of carbon from the atmosphere to the ocean, and $F_{as}$ is the flux of carbon from the ocean to the atmosphere. We modify the model described by Stocker et al. [1994] to include for a gas exchange rate which is a function of wind speed, $W$, using the relationship from Tans et al. [1994],

$$k = 0.016 \times (W/\text{m/s})^{-3}$$

We use zonally averaged wind speed from the climatology of Estensen and Kustner [1981], and the resulting exchange rates are shown in Figure 7. For the experiments which follow, atmospheric pCO2 is not interactive, but is imposed at a constant preanthropogenic value of 280 μatm.

In addition to total carbon concentration (which is taken here to be the same as 12C concentration due to the much greater abundance of this isotope), 13C is also shown as a tracer. The flux of 13C into the ocean ($F^{13}_\Sigma$) is determined as by Siegenthaler and Münchj [1981]:

$$F^{13}_\Sigma = \alpha_{as} F_{as} R_\Sigma - \alpha_{as} F_{as} R_S$$

where $R_S$ is the ratio of 13C/12C in the atmosphere and $R_\Sigma$ is the ratio of 13C/12C in oceanic carbon ($\Sigma$CO2). The fractionation factor for air-sea carbon transfer, $\alpha_{as} = \sigma_\gamma \sigma_\delta$, where $\sigma_\delta = 0.373 T(\text{°C}) + 1.00019$ [Vogel et al., 1970], is the temperature dependent equilibrium fractionation factor between atmospheric CO2 and surface ocean CO2 (aq), and $\sigma_\gamma = 0.9995$ [Siegenthaler and Münchj, 1981]. The kinetic fractionation factor for CO2 gas transfer across the air-sea interface. The fractionation factor for sea-air carbon transfer, $\alpha_{as} = \sigma_\gamma \sigma_\delta$. Here $\sigma_\gamma$, the temperature dependent equilibrium fractionation factor between surface ocean CO2 (aq) and total dissolved inorganic carbon ($\Sigma$CO2), is approximated by the equilibrium factor between oceanic CO2 (aq) and bicarbonate (HCO3-), $\sigma_{\text{HCO}_3^-} = 9.866/11(\text{K}) + 1.02412$ [Mook et al., 1974; Vogel et al., 1970]. For various model runs, as noted below, these fractionation factors were altered in order to explore the role of different air-sea exchange processes in determining surface ocean δ13C. Although the older fractionation factors referenced above are used in all model runs shown, substituting the fractionation factors presented by Zhang et al. [1995] changes model results only by a small amount (everywhere less than 0.3 %). Although only 12C and 13C are carried in the model, the carbon isotope ratio is expressed as δ13C (δ13C (‰) = (R/R_{PDB} - 1)×1000) for model output, where R is the ratio 13C/12C, and $R_{\text{PDB}}$ is the ratio for the reference value (PDB). The δ13C of atmospheric CO2 is specified at the preanthropogenic value of -6.48‰.

Model Results : Assessing the Role of Air-Sea Exchange

The inorganic carbon cycle model was run with no modifications (experiment 1, Table 2) in order to assess the effect of air-sea exchange on the distribution of δ13C.

![Figure 7. Gas exchange rates, $k$ (mol m^-2 y^-1 μatm^-1), used in experiments 1, 2, and 4. The constant gas exchange rate of 0.067 mol m^-2 y^-1 μatm^-1 used in experiment 4 is also shown for comparison.](image-url)

Table 2. Model Experiments

<table>
<thead>
<tr>
<th>Experiment Number</th>
<th>Modifications</th>
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<tbody>
<tr>
<td>1</td>
<td>No modifications (full temperature dependent and kinetic fractionation factors, wind speed dependent gas exchange), atmospheric pCO2 = 280, δ13C = -6.48</td>
</tr>
<tr>
<td>2</td>
<td>Anthropogenic increase in CO2 and δ13C prescribed for atmosphere</td>
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<tr>
<td>3</td>
<td>Constant gas exchange rate (no wind speed dependence), atmospheric pCO2 = 280, δ13C = -6.48</td>
</tr>
<tr>
<td>4</td>
<td>Fractionation between CO2 and δ13C constant (No temperature dependence), $(\alpha_{as} = 0.98840, \sigma_\gamma = 0.99840)$, atmospheric pCO2 = 280, δ13C = -6.48</td>
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Atmospheric CO$_2$ was fixed at the preindustrial value of 280 ppmv and the $\delta^{13}$C of atmospheric carbon at −6.48‰ [Friedli et al., 1986]. The model was integrated for 10,000 years until a steady state for $\delta^{13}$C was reached (Figure 8). The long time required to reach a steady state is due to the fact that carbon isotopes in the surface ocean are not completely equilibrated with the atmosphere. It is therefore necessary for the model to complete several ventilation cycles (at 1000 to 2000 years each) before the waters have been exposed at the surface for a sufficient amount of time for a steady state to be reached. For ease of comparison with modern oceanic $\delta^{13}$C$_{as}$ values, $\delta^{13}$C$_{as}$ was also calculated for the model by subtracting the model’s mean ocean value for $\delta^{13}$C from the model predicted $\delta^{13}$C. Because biological processes are not included in this model, the patterns of $\delta^{13}$C and $\delta^{13}$C$_{as}$ will be identical for each model run, but will be offset from one another by a constant amount.

As expected, cold surface waters have high $\delta^{13}$C, reflecting the equilibration at cold temperatures, and warm surface waters have low $\delta^{13}$C, with minima in the center of the subpolar gyres, reflecting the relatively long residence of these waters at the warm surface (Figure 9). However, in no area of the ocean is $\delta^{13}$C completely in equilibrium with the atmosphere (Figure 9, dashed line), as predicted based on the long time required for carbon isotopic equilibration at a given temperature (about 10 years). The $\delta^{13}$C$_{as}$ of subsurface waters reflects the $\delta^{13}$C$_{as}$ of the source region, with NADW having relatively low $\delta^{13}$C$_{as}$ and Antarctic Bottom Water (AABW) and Antarctic Intermediate Water with higher $\delta^{13}$C$_{as}$ (Figure 10). Before comparing model $\delta^{13}$C$_{as}$ with the air-sea exchange signature observed in the real ocean, we must quantitatively assess the impact of the rising atmospheric CO$_2$ and falling atmospheric $\delta^{13}$C due to land use changes and fossil fuel emissions over the last several hundred years.

Experiment 2 is started from the steady state preindustrial state (experiment 1) and atmospheric CO$_2$ and $\delta^{13}$C are specified to reflect the observed anthropogenic increase. Atmospheric CO$_2$ is prescribed as by Stocker et al. [1994], and $\delta^{13}$C is taken to be linearly related to the increase in CO$_2$ (Figure 11). The model is then integrated for 188 more years (1800 - 1988) to obtain the transient distributions of $\delta^{13}$C in the inorganic carbon cycle model. In order to calculate $\delta^{13}$C$_{as}$

![Figure 8](image.jpg)

**Figure 8.** Model values of $\delta^{13}$C of ΣCO$_2$ for the middepth North Pacific during model spin-up. A steady state is reached for $\delta^{13}$C of ΣCO$_2$ after the model is integrated for about 8000 years.

![Figure 9](image.jpg)

**Figure 9.** Steady state surface $\delta^{13}$C for experiment 1 (solid symbols) along with the $\delta^{13}$C predicted if the surface ocean were in complete isotopic equilibrium with the atmosphere (open symbols). This run includes full temperature dependent isotopic fractionation factors, and preindustrial atmospheric CO$_2$ and $\delta^{13}$C. The air-sea exchange signature (modelled $\delta^{13}$C minus modelled mean ocean $\delta^{13}$C for this inorganic model) for experiment 1 can be read from the right-hand axis.
Figure 10. Steady state $\delta^{13}$C$_{as}$ distributions for the Atlantic and Pacific Oceans for experiment 1 (temperature dependent fractionation factors and preindustrial atmosphere).

Figure 12. The $\delta^{13}$C$_{as}$ of surface water carbon for experiment 2, which includes the anthropogenic rise in atmospheric CO$_2$ and fall in $\delta^{13}$C of atmospheric CO$_2$ (solid lines and symbols). Also plotted is the $\delta^{13}$C$_{as}$ of surface water that results from the rise in atmospheric CO$_2$ alone (invasion effect), with atmospheric $\delta^{13}$C held constant (dashed lines, open symbols).

The patterns and ranges of $\delta^{13}$C$_{as}$ values for surface waters in the inorganic model with full isotopic fractionation factors (experiments 1 and 2) look very much like $\delta^{13}$C$_{as}$ calculated from observed $\delta^{13}$C and PO$_4$ (Figure 5d). Minimum values are in the subtropical gyres and maximum values are in the Antarctic and North Pacific. As with observed $\delta^{13}$C$_{as}$, $\delta^{13}$C in the inorganic model remains low in the North Atlantic despite the low surface temperatures. The relatively high $\delta^{13}$C$_{as}$ of Southern Ocean Surface Waters and low $\delta^{13}$C$_{as}$ of North Atlantic Surface Waters are reflected in the high $\delta^{13}$C$_{as}$ of AABW, and the low $\delta^{13}$C$_{as}$ of NADW (Figure 10), as in the real ocean (Figure 6).

In experiment 3, a constant gas exchange rate of .067 mol m$^{-2}$ yr$^{-1}$ atm$^{-1}$ is used instead of the zonally varying exchange rate of experiment 1. The pattern of surface ocean $\delta^{13}$C$_{as}$, predicted by this model run (Figure 13) is very similar to the model with variable gas exchange, and also replicates the air-sea exchange signature, $\delta^{13}$C$_{as}$ features observed in the real ocean (Figure 5d). The similarity between the distributions of surface ocean $\delta^{13}$C in the two model runs (experiments 1 and 4) implies that, to first order, the pattern of surface ocean $\delta^{13}$C$_{as}$ is determined primarily by the temperature and history.

Figure 11. The $\delta^{13}$C of atmospheric CO$_2$ for experiment 2 (1800-1988), is assumed to vary linearly with CO$_2$ from the preindustrial value of -6.48%o at 280 ppm [Friedli et al., 1986] to -7.74%o at 500 ppm in 1988 [Keeling et al., 1989]. This relationship is shown along with the rest of the pCO$_2$ and $\delta^{13}$C data from these publications.
of surface water masses, while geographic variations of gas exchange rates play a secondary role. The absence of enhanced gas exchange rates in the polar regions in this run does result in a slightly lower $\delta^{13}\text{C}_{\text{as}}$ gradient between low and high latitudes.

In experiment 4, the inorganic carbon cycle model was modified such that there is no temperature dependence to carbon isotope fractionation during air-sea exchange. This experiment demonstrates the effect of invasion and evasion of $\text{CO}_2$ on the distribution of carbon isotopes in the world ocean. These effects cause a spread of up to 0.6% in surface ocean $\delta^{13}\text{C}_{\text{as}}$ with high $\delta^{13}\text{C}_{\text{as}}$ in warm waters where the evasion of the isotopically lighter $\text{CO}_2$ (aq) enriches the surface water $\Sigma\text{CO}_2$ in $^{13}\text{C}$ (Figure 14). Polar surface waters in both hemispheres have similar low $\delta^{13}\text{C}$ values reflecting the invasion of $\text{CO}_2$ into these cold waters. Note that this differs from the case of the real ocean, where biological cycling will enhance the invasion of $\text{CO}_2$ into the North Atlantic where biological processes have depleted surface ocean carbon and mute the invasion of $\text{CO}_2$ into the Southern Ocean where already carbon-rich waters upwell. In general, the effects of invasion/evasion alone (high $\delta^{13}\text{C}$ in warm waters, low $\delta^{13}\text{C}$ in cold waters) are opposite in sign to the effects of temperature dependent equilibration with the atmosphere (high $\delta^{13}\text{C}$ in cold waters, low $\delta^{13}\text{C}$ in warm waters).

Although these model experiments give us a first order look at how air-sea exchange will affect the distribution of $\delta^{13}\text{C}$ in the ocean, in the real ocean biological processes will also influence air-sea exchange, and thus the air-sea exchange signature, $\delta^{13}\text{C}_{\text{as}}$, to some extent. When biological cycling is added to this model, we will be able to test the sensitivity of $\delta^{13}\text{C}_{\text{as}}$ to these processes. However, it is clear from the good comparisons between modelled and actual $\delta^{13}\text{C}_{\text{as}}$ that while biological processes are quite important in determining $\delta^{13}\text{C}$, they will have a secondary influence on the air-sea exchange signature, $\delta^{13}\text{C}_{\text{as}}$.

**Conclusions**

Although biological cycling seems to be the primary control on today's deep ocean $\delta^{13}\text{C}$ distribution, air-sea exchange plays an equally important role in determining the $\delta^{13}\text{C}$ of surface ocean carbon. Air-sea exchange controls $\delta^{13}\text{C}$
primarily due to the tendency toward isotopic equilibration between the ocean and atmospheric carbon reservoirs. The one-way transfer of isotopically light CO$_2$ to and from the oceanic $\Sigma$CO$_2$ reservoir can also significantly affect surface ocean $\delta^{13}$C. The importance of air-sea exchange on $\delta^{13}$C of surface waters can be seen in the departures from biologically predicted $\delta^{13}$C values in the surface ocean. Surface waters in the Southern Ocean have high $\delta^{13}$C anomalies due to isotopic exchange at low temperatures. Surface waters in the subtropical gyres have low $\delta^{13}$C anomalies due to isotopic exchange at high temperatures, while surface waters at the equator have high $\delta^{13}$C anomalies due to the evasion of CO$_2$. In the North Atlantic, surface waters have low $\delta^{13}$C anomalies reflecting the warm surface water history and invasion of isotopically light atmospheric CO$_2$. These anomalies are carried to deep sea from the region of water mass formation. The departure from biologically predicted $\delta^{13}$C can be quite large, and at times in the past the modern inverse relationship between deep water nutrient concentration $\delta^{13}$C could have changed if the oceans were filled with water masses originating in different surface regimes with different temperature and nutrient histories [Lynch-Stieglitz and Fairbanks, 1994]

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