Upper ocean carbon fluxes in the Atlantic Ocean: The importance of the POC:PIC ratio

Wolfgang Koeve
Marum, FB Geowissenschaften, Universität Bremen, Bremen, Germany

Received 20 November 2001; revised 13 May 2002; accepted 15 May 2002; published 15 October 2002.

The mean depth distribution of the POC:PIC ratio of sinking particles, measured with particle interceptor traps deployed in the Atlantic Ocean, is fitted by an exponential function (POC:PIC = 64.3Z^{-0.56}, r^2 = 0.69) The function is successfully evaluated by comparison with (a) estimates of the POC:PIC ratio of export production, computed from seasonal changes of nitrate and alkalinity and (b) estimates of the POC:PIC ratio of remineralization on shallow isopycnals. The basin mean POC:PIC ratio of export production is 4.2 to 4.37. The POC:PIC-depth function is combined with empirical relationships between the flux of particulate organic matter, primary production and depth, satellite derived primary production data sets, and the regional distribution of Ψ (the ratio of released CO2 precipitated carbonate during CaCO3 formation) in order to estimate the effective carbon flux (J_eff) in the Atlantic Ocean. Remineralization of organic carbon above the winter mixed layer (11–17%) and CaCO3 sequestration from the winter mixed layer (13–16%), which is the balance between CaCO3 production and shallow dissolution, are the two main processes which control the difference between export production (0.9 and 2.9 GT C yr^{-1}) and J_eff (0.64 and 2.2 GT C yr^{-1}) on the basin scale (65°N to 65°S). CaCO3 sequestration is the dominant process modulating effective carbon export in the tropics, while shallow POC remineralization dominates in temperate and polar waters. Observed regional patterns like polarward increases of the POC:PIC export ratio and of Ψ counteract each other largely when J_eff is computed.

INDEX TERMS: 4203 Oceanography: General: Analytical modeling; 4806 Oceanography: Biological and Chemical: Carbon cycling; 4863 Oceanography: Biological and Chemical: Sedimentation; KEYWORDS: carbon cycle, particle flux, Corg: Cinorg ratio, effective carbon export, biological pump, Atlantic Ocean


1. Introduction

The biological pump [Volk and Hoffert, 1985] drives a net flux of fixed organic carbon from the surface of the ocean into its interior. This flux, and the subsequent remineralization in the deep ocean produces a vertical gradient of total CO2, which controls the steady state CO2 level of the atmosphere [Shaffer, 1993]. To determine the effectiveness of this carbon flux is at the heart of carbon cycle studies in the ocean [SCOR, 1990; Fasham et al., 2001]. Export production [Eppley and Peterson, 1979], which is the sum of all organic carbon fluxes (particulate and dissolved organic carbon, POC and DOC) from the productive surface of the ocean, has often been referred to as a suitable measure of this effectiveness. However, a significant fraction of the carbon export is remineralized within the upper few hundred meters [Martin et al., 1987; Louanchi and Najjar, 2000] and nutrients and CO2 stemming from shallow remineralization can be recirculated to the surface ocean on short timescales of about a year and do not contribute to the vertical inner-ocean CO2 gradient. Additionally, a fraction of the organic carbon export is compensated by the export of particulate inorganic carbon (PIC, CaCO3). This latter compensation is due to the net effect of CaCO3 production and export on the surface ocean pCO2. During formation of one mole of CaCO3, alkalinity is reduced by two moles and total dissolved inorganic carbon decreases by one mole. In combination, these changes cause a shift in the carbonate system toward a higher CO2 concentration and hence a higher pCO2 [Broecker and Peng, 1982; Robertson et al., 1994; Frankignoulle et al., 1994; Buitenhuis et al., 2001]. Like for organic carbon, sequestration of particulate inorganic carbon to the deep ocean is the balance between CaCO3 production and shallow dissolution of CaCO3. Hence, new carbon production, shallow remineralization of PIC and the countereffective production, export and shallow dissolution of PIC must be considered to obtain a correct estimate of the carbon flux which is effective with respect to the steady state mean pCO2 of the surface ocean.
A useful measure of the effectiveness of the biological pump is the net export of carbon, which reaches into water masses which are isolated from exchange with the atmosphere on decadal to centennial timescales, here referred to as $J_{\text{eff}}$. To estimate $J_{\text{eff}}$ the export and the shallow remineralization and dissolution of POC and PIC need to be specified. However, we have a modest understanding of the large-scale patterns and integrals of these properties. Direct estimates of POC or PIC export from the euphotic zone, based on drifting particle interceptor traps [Zeitzschel et al., 1978; Asper, 1996], are sporadic, short term in nature (days to weeks; Martin et al. [1993]), and methodologically questionable [Gust et al., 1992; Kähler and Bauerfeind, 2001]. In particular, very few examples exist, which resolve the depth distribution of PIC fluxes in the upper ocean [Martin et al., 1993]. Model-based estimates can fill some of these gaps. Ocean carbon cycle models have provided estimates of new production [Najjar et al., 1992; Sarmento et al., 1993; Six and Maier-Reimer, 1996; Oeschlies et al., 2000], however, few such models [e.g., Bacastow and Maier-Reimer, 1990] included a formal description of the inorganic carbon cycle. Modeling of CaCO3-related processes has usually restricted itself to dissolution at and below the lysocline in the deep ocean and in the sediments [Archer, 1991, 1996; Sigman et al., 1998]. The production and fate of CaCO3 in the upper ocean have rarely been studied quantitatively using ocean carbon cycle models [Yamanaka and Tajika, 1996; Archer et al., 2000a]. Recently, one of these model studies [Yamanaka and Tajika, 1996] has challenged the paradigmatic idea that the POC:PIC ratio of the ocean is about 4–5 [Li et al., 1969; Broecker and Peng, 1982] and suggested values between 10 and 12.5. Solving this issue is of great importance for the understanding of suspected changes in the production of CaCO3 [Riebesell et al., 2000; Zondervan et al., 2001] and their feedback on the future net CO2-uptake of the ocean.

Linking maps of primary production [e.g., Berger, 1989; Longhurst et al., 1995; Antoine and Morel, 1996; Antoine et al., 1996; Behrenfeld and Falkowski, 1997] with empirical relationships between particle flux (J), primary production (PP), and trap deployment depth (Z) [Suess, 1980; Betzer et al., 1984; Hargrave, 1985; Pace et al., 1987; Berger et al., 1987 in the following referred to as J-PP-Z functions] provide an independent approach for basin scale estimates of POC fluxes in the ocean [Antia et al., 2001]. These basin-scale POC flux estimates can be combined with empirical relationship of the POC:PIC flux ratio and water depth for the estimation of $J_{\text{eff}}$ [Antia et al., 2001].

Recently Antia et al. [2001] and Koeve (W. Koeve, New production and shallow remineralization in the Atlantic Ocean—uncertainties and limitations of particle flux algorithms, submitted to Journal of Marine Research, 2001; in the following referred to as Koeve (submitted manuscript, 2001) estimated upper-ocean carbon fluxes for the Atlantic Ocean based on this empirical approach). A basin wide integrated carbon export production of 3.14 GT C yr$^{-1}$ (equivalent to a mean f-ratio of 0.33) was estimated from a newly derived J-PP-Z function [Antia et al., 2001]. POC fluxes were integrated over mixing cycles (i.e., from winter to winter) and corrected for trapping inefficiencies by applying Th-calibration [Scholten et al., 2001]. Remineralization above the winter mixed layer was about 5.7% of export production and compensation due to CaCO3 export from the winter mixed layer reduced the carbon flux by another 16.5%. Antia et al. [2001] estimate the effective carbon export of the Atlantic Ocean to be 2.47 GT C yr$^{-1}$. Koeve (submitted manuscript, 2001), using a slightly different approach, showed estimates of export production to be very sensitive to the choice of the primary production forcing data set. Basin-scale means of export production (and the respective export ratio, ER) ranged between 0.9 and 2.9 GT C yr$^{-1}$ (equivalent to an ER = 0.07–0.31). The fraction of shallow remineralization was better constrained and showed a relationship with the depth of winter-time mixing, which for extratropical regions explained 92% of the variation of this fraction. Remineralization above the winter mixed layer depth was up to 62% of export production in the North Atlantic subarctic province.

The overall aim of this paper is to describe the relative importance and the regional distribution of (a) shallow POC remineralization and (b) the CaCO3 balance (production and shallow dissolution) for the adjustment of the effective carbon flux (section 4). For this purpose, the empirical relationship between flux ratios of organic to inorganic carbon (POC:PIC flux ratio; “rain ratio”) from particle interceptor traps deployed in the Atlantic Ocean during the JGOFS decade [SCOR, 1990], and water depth is developed (section 3.1). This empirical relationship is evaluated by comparison with independent data. In particular, the POC:PIC export ratio (section 3.2), the vertical change of the POC:PIC ratio (section 3.4), and of the PIC flux (section 3.3) in the Atlantic are discussed during this comparison. In the following sections a short description of the data sets used and a definition of some fundamental terms are given.

2. Material and Methods

2.1. Data Sets, Depths Levels of Reference, Basin Scale Integration

Three-dimensional data sets of the organic carbon flux in the Atlantic Ocean are adopted from the study of Koeve (submitted manuscript, 2001). In that study a suit of eight comparative data sets of particulate organic carbon fluxes in the ocean was computed from maps of primary production [Antoine et al., 1996; Behrenfeld and Falkowski, 1997] and J-PP-Z flux functions. J-PP-Z fits were based on local estimates of primary production derived from the global primary production maps and Th$^{230}$-corrected [Scholten et al., 2001] POC flux observations made in the Atlantic Ocean in the 1990s as part of the JGOFS program [Antia et al., 2001]. A data set of PIC:POC flux ratios from particle flux studies in the Atlantic is also taken from the latter study. A detailed reference to the original particle flux data sets is given in the work by Antia et al. [2001] (their Table 1).

This study focuses on carbon fluxes at two depth boundaries. Export production of particulate organic carbon (JPOC$^{\text{exp}}$) and particulate inorganic carbon (JPIC$^{\text{exp}}$) is the flux of POC and PIC across $Z_{\text{exp}}$. Operationally, and for the sake of easy comparison with published model results, $Z_{\text{exp}}$ is defined as $Z = 125$ m in this study. The carbon flux into
mixed layer depth is defined as the change in density with respect to the surface by 0.125 kg dm$^{-3}$. JPOC$_{\text{exp}}$, JPIC$_{\text{exp}}$, JPOC$_{\text{sequ}}$, and JPIC$_{\text{sequ}}$ are two-dimensional data sets with a resolution of 1° latitude × 1° longitude.

### 2.2. Formal Description of Carbon Fluxes

Formally the depths dependence of the organic carbon flux $J_{\text{POC}}$ can be empirically described according to equation (1).

Sets of parameters $a$, $b$, and $c$ (Table 1), which were determined for eight combinations of different PP- and particle flux data sets and fitting techniques, are taken from the study of Koeve (submitted manuscript, 2001). Empirical descriptions of PIC fluxes were not directly estimated from trap observations but by combining equation (1) with empirical fits of the POC:PIC flux ratio (RR) according to equation (2)

$$J_{\text{POC}} = c PP^a Z^b \quad (1)$$

$$J_{\text{PIC}} = J_{\text{POC}} / RR \quad (2)$$

[12] As will be discussed in the next section, RR can be described in various ways as a function of the trap deployment depth. For these empirical fits the full non-polar data set (n = 76) of Antia et al. [2001] is used. This data set combines observations (n = 24) for which the trapping efficiency has been estimated ($\text{Th}^{230}$-approach; Scholten et al. [2001]) and data sets for which no $\text{Th}^{230}$ data were available. This procedure is justified by the observations that $\text{Th}^{230}$ is not selectively trapped by organic or inorganic particles [Scholten et al., 2001]. Estimating $J_{\text{PIC}}$ according to equation (2) has the advantage that a larger and more representative data set can be used, however, at the expense of combining empirical relationships which are based on different data sets.

[13] JPOC, JPIC, and RR are data sets on a three dimensional grid (latitude, longitude, depth). From JPOC, JPIC, RR, and $\Psi$, the ratio of released $\text{CO}_2$-precipitated carbonate during $\text{CaCO}_3$ precipitation [Frankignoulle et al., 1994; see section 4.3.], a property called the net carbon flux, $J_{\text{net}}$, can be defined as the carbon export that drives a potential draw-down of $\text{CO}_2$ from the atmosphere to the ocean (equation (3)). $J_{\text{net}}$ is defined on the same three dimensional grid as $J_{\text{POC}}$.

Finally, $J_{\text{eff}}$, the effective carbon flux is defined as the value of $J_{\text{net}}$ at $Z = Z_{\text{seq}}$. This indicates that only the net flux of carbon across the sequestration depths, which is estimated from the annual maximum mixing depth (see section 2.1), is regarded as being really effective in driving a draw-down of $\text{CO}_2$ from the atmosphere to the ocean.

Like JPOC$_{\text{sequ}}$ or JPIC$_{\text{sequ}}$, $J_{\text{eff}}$ is a two dimensional data set on a latitude/longitude grid. In this study $J_{\text{eff}}$ will be calculated according to equation (4), where $\Psi$ indicates an appropriately averaged value of $\Psi$ (see section 4.3)

$$J_{\text{net}} = J_{\text{POC}} - \Psi J_{\text{PIC}} \quad (3)$$

$$J_{\text{eff}} = J_{\text{POC}} - \Psi J_{\text{PIC}}$$

<table>
<thead>
<tr>
<th>Data set</th>
<th>Method</th>
<th>Parameter</th>
<th>Regression Analysis</th>
<th>n</th>
</tr>
</thead>
<tbody>
<tr>
<td>AM96</td>
<td>linear</td>
<td>1.69</td>
<td>1.008</td>
<td>24</td>
</tr>
<tr>
<td>BF97</td>
<td>linear</td>
<td>1.47</td>
<td>1.010</td>
<td>24</td>
</tr>
</tbody>
</table>

"These empirical relationships are established for primary production [Antoine et al., 1996; Behrenfeld and Falkowski, 1997] and particle flux [Antia et al., 2001] data sets from the Atlantic ocean (from Koeve, submitted manuscript, 2001; modified)."

"Linear: linear optimization; log: logarithmic optimization."

"The parameters of the exponential function $J_e = cPP^a Z^b$ (equation (1))."

<table>
<thead>
<tr>
<th>Data set</th>
<th>Method</th>
<th>Parameter</th>
<th>Regression Analysis</th>
<th>n</th>
</tr>
</thead>
<tbody>
<tr>
<td>AM96</td>
<td>log</td>
<td>1.29</td>
<td>0.993</td>
<td>24</td>
</tr>
<tr>
<td>BF97</td>
<td>log</td>
<td>0.77</td>
<td>0.998</td>
<td>24</td>
</tr>
</tbody>
</table>

"These empirical relationships are established for primary production [Antoine et al., 1996; Behrenfeld and Falkowski, 1997] and particle flux [Antia et al., 2001] data sets from the Atlantic ocean (from Koeve, submitted manuscript, 2001; modified)."

"Linear: linear optimization; log: logarithmic optimization."

"The parameters of the exponential function $J_e = cPP^a Z^b$ (equation (1))."
Table 2. Abbreviations Used in This Study

<table>
<thead>
<tr>
<th>Term</th>
<th>Units</th>
<th>Remarks</th>
</tr>
</thead>
<tbody>
<tr>
<td>$Z_{\text{exp}}$</td>
<td>m</td>
<td>reference depth of export production, in this study $Z_{\text{exp}} = 125$ m</td>
</tr>
<tr>
<td>$Z_{\text{sequ}}$</td>
<td>m</td>
<td>reference depth of carbon sequestration (i.e., the export of carbon into water layers which do not exchange with the atmosphere on timescales of $&gt;1$ yr)</td>
</tr>
<tr>
<td>$Z_{\text{wml}}$</td>
<td>m</td>
<td>climatological depth of the seasonal maximum of the mixed layer, winter mixed layer; in seasonal systems $Z_{\text{wml}}$ is a good approximation of $Z_{\text{exp}}$</td>
</tr>
<tr>
<td>JPOC&lt;sub&gt;exp&lt;/sub&gt;, JPOC&lt;sub&gt;125&lt;/sub&gt;</td>
<td>g C m&lt;sup&gt;-2&lt;/sup&gt; yr&lt;sup&gt;-1&lt;/sup&gt;</td>
<td>export production (also a measure of new carbon production); in this study JPOC&lt;sub&gt;125&lt;/sub&gt;, the flux of particulate organic carbon in 125 m depth is used as an approximation of JPOC&lt;sub&gt;exp&lt;/sub&gt;</td>
</tr>
<tr>
<td>JPOC&lt;sub&gt;sequ&lt;/sub&gt;</td>
<td>g C m&lt;sup&gt;-2&lt;/sup&gt; yr&lt;sup&gt;-1&lt;/sup&gt;</td>
<td>sequestration of organic carbon, i.e., the export of organic carbon across $Z_{\text{sequ}}$</td>
</tr>
<tr>
<td>JPIC&lt;sub&gt;125&lt;/sub&gt;</td>
<td>g C m&lt;sup&gt;-2&lt;/sup&gt; yr&lt;sup&gt;-1&lt;/sup&gt;</td>
<td>the export of particulate inorganic carbon across $Z_{\text{exp}}$</td>
</tr>
<tr>
<td>JPIC&lt;sub&gt;sequ&lt;/sub&gt;</td>
<td>g C m&lt;sup&gt;-2&lt;/sup&gt; yr&lt;sup&gt;-1&lt;/sup&gt;</td>
<td>the sequestration flux of particulate inorganic carbon across $Z_{\text{sequ}}$</td>
</tr>
<tr>
<td>$J_{\text{eff}}$</td>
<td>g C m&lt;sup&gt;-2&lt;/sup&gt; yr&lt;sup&gt;-1&lt;/sup&gt;</td>
<td>net carbon flux, see equation (3); $J_{\text{eff}}$ is defined at any depth</td>
</tr>
<tr>
<td>$r_{\text{R-POC}}$</td>
<td>mol:mol</td>
<td>the fraction of export production of organic carbon, JPOC&lt;sub&gt;125&lt;/sub&gt;, which is remineralized above $Z_{\text{sequ}}$; $r_{\text{R-POC}} = (\text{JPOC}<em>{125} / \text{JPOC}</em>{\text{sequ}}) / \text{JPOC}_{125}$</td>
</tr>
<tr>
<td>$r_{\text{R-PIC}}$</td>
<td>mol:mol</td>
<td>the fraction of export of particulate inorganic carbon, JPIC&lt;sub&gt;125&lt;/sub&gt;, which is remineralized above $Z_{\text{sequ}}$; $r_{\text{R-PIC}} = (\text{JPIC}<em>{125} / \text{JPIC}</em>{\text{sequ}}) / \text{JPIC}_{125}$</td>
</tr>
<tr>
<td>RR&lt;sub&gt;sequ&lt;/sub&gt;</td>
<td>mol:mol</td>
<td>the POC:PIC ratio (&quot;rain ratio&quot;) at $Z_{\text{sequ}}$</td>
</tr>
</tbody>
</table>

Abbreviations used in this study are summarized in Table 2.

3. POC:PIC Ratios

3.1. POC:PIC Flux Ratios From Particle Flux Observations

Two functionalities of the vertical profile of the POC:PIC flux ratio (RR) are analyzed, a linear regression (model I) of RR over ln (Z) [Antia et al., 2001] (equation (5)) and an exponential fit (equation (6))

$$RR = d \ln(Z) + e \quad (5)$$

$$RR = d Z^b \quad (6)$$

Figure 1. Vertical distribution of the POC:PIC flux ratio and PIC fluxes in the North Atlantic. (a) POC:PIC flux ratio (RR). +: observations, $n = 76$; Antia et al. [2001]. Solid line: exponential function, RR = 64.3 $Z^{-0.56}$; $r^2 = 0.693$ (this study). Dashed line: logarithmic fit, RR = 7.34 $- 0.827 \ln(Z)$; $r^2 = 0.664$ (this study). (b, c) Distribution of the normalized ((x_i-mean(x_i))/std(x_i)) and site independent relative vertical profile of JPIC<sub>*</sub>, named ($J_{\text{PIC}}/L$)*. Here L is a shortcut of the term cP<sub>*</sub> of equation (1) and b is the z-exponent of equation (1). (b) Estimates based on logarithmic RR-Z functions. Solid line: Antia et al. [2001]; $b = -0.68$, RR = 7.39 $- 0.83 \ln(Z)$; dashed line: this study; $b = -1$, RR = 7.34 $- 0.827 \ln(Z)$. (c) Estimates based on the exponential RR-Z function RR = 64.3$Z^{-0.56}$ from this study. Solid line: $b = -1.0$; dashed line: $b = -0.7$. Note the artificial increase in ($J_{\text{PIC}}/L$)* in the deep ocean in Figure 1b.
and RR = 64.3Z^{-0.56}[ln(RR) = -0.557 (±0.0516)ln(Z) + ln(4.16 ± 0.381)], respectively. The statistics for both functions are quite similar. Model-II regressions [Sokal and Rohlf, 1995] of the predicted versus the observed flux ratios yield $r^2$ values of 0.664 (RR-ln(Z)) and 0.693 (exponential fit). A major difference, however, shows up, when JPIC is estimated from JPOC and the POC:PIC ratio according to equation (2). In the case of the RR-ln(Z) function JPIC decreases only down to about 1500 m but increases again in the deep ocean (Figure 1b). This unprecedented increase of the PIC flux is not seen if the POC:PIC ratio is estimated from the exponential fit. Here JPIC continuously decreases with depth (Figure 1c).

### 3.2. The POC:PIC Ratio of Export Production

[19] In this section the POC:PIC ratio of export production (RR$_{exp}$) is estimated based on the particle flux algorithms discussed above (section 3.2.1), an analysis of data from process studies in the North Atlantic (section 3.2.2), and finally, based on climatological data (section 3.2.3).

#### 3.2.1. Estimates From Particle Flux Algorithms

[20] Using the exponential RR-Z function, a mean POC:PIC flux ratio in 125 m depth, RR$_{125}$, of 4.37 is estimated. Propagating the standard errors of the parameters d and e (see section 3.1) through the computation yields a range of 2.33–8.21 for RR$_{125}$. The logarithmic fit suggests an POC:PIC export ratio of 3.35 (±0.834).

#### 3.2.2. Estimates From Field Studies in the North Atlantic

[21] Few local estimates of the POC:PIC ratio of production have been published to compare the estimates from the particle flux algorithm with. In the following I will combine estimates of the POC:PIC ratio of net carbon production during a coccolithophorid bloom in the subpolar North Atlantic and from spring bloom studies in the temperate and subpolar North Atlantic [Robertson et al., 1994] with winter-time nutrient estimates [Koeve, 2001] to compute the range of the annually integrated POC:PIC ratio of new production in this region. Robertson et al. [1994] estimated the POC:PIC ratio from the relative change of potential alkalinity ($A_{pot} = A_t + NO_3;$ Fiadeiro [1980]) and total CO$_2$ over time. They found POC:PIC ratios of production of 1:1 for a coccolithophorid bloom and 6:1 during a diatom bloom. The annual mean POC:PIC ratio depends on the relative importance of diatoms, coccolithophorids, and non-calciifying flagellates (e.g., Phaeozystis) for annual new production. A first order estimate of the quantitative role of diatoms can be achieved from wintertime NO$_3$:Si(OH)$_4$ ratios and the mean nitrate:silicate uptake ratio of diatoms (about 1:1; Richards, 1958; Brzezinski [1985]). The decrease of wintertime NO$_3$:Si(OH)$_4$ ratios from 2.8 to 2 between 40°N and 60°N along 20°W [Koeve, 2001], converts to an increase of the relative contribution of diatoms to new carbon production from about 35–38% (at 40°N to 47°N) to 50–64% at 60°N ($f_{diamons}$ in Table 3). How this ratio

<table>
<thead>
<tr>
<th>Scenario</th>
<th>Latitude</th>
<th>[NO$_3$/Si(OH)$_4$] (Winter)$^c$</th>
<th>$f_{diamons}$</th>
<th>POC:PIC Ratio of Production</th>
<th>Remarks</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>47°N</td>
<td>2.6</td>
<td>0.38</td>
<td>2.9</td>
<td>assuming a POC:PIC ratio of 6:1 during the diatom bloom$^b$</td>
</tr>
<tr>
<td>2</td>
<td>47°N</td>
<td>2.6</td>
<td>0.38</td>
<td>4.0</td>
<td>like scenario 1, but assuming a POC:PIC ratio of 9:1 during the diatom bloom$^a$</td>
</tr>
<tr>
<td>3</td>
<td>60°–62°N</td>
<td>2.0</td>
<td>0.5</td>
<td>3.5</td>
<td>assuming summer nitrate depletion and the POC:PIC ratio during the diatom bloom of 6:1</td>
</tr>
<tr>
<td>4</td>
<td>60°–62°N</td>
<td>2.0</td>
<td>0.64</td>
<td>4.2</td>
<td>assuming summer silicate depletion,$^d$ summer nitrate concentrations$^e$ of about 3 μmol dm$^{-3}$, and a contribution of coccolithophorids to post spring bloom new production of 100%</td>
</tr>
<tr>
<td>5</td>
<td>60°–62°N</td>
<td>2.0</td>
<td>0.64</td>
<td>5.1</td>
<td>assuming summer silicate depletion,$^d$ summer nitrate concentrations$^e$ of about 3 μmol dm$^{-3}$, and a contribution of coccolithophorids to post spring bloom new production of 50%</td>
</tr>
</tbody>
</table>

**From Particle Flux Algorithms**

- 65°S–65°N: 3.4 RR$_{125}$, RR-ln(Z) fit; from$^f$
- 65°S–55°N: 4.4 RR$_{125}$, exponential fit of RR; from$^f$

**From Seasonal Changes of CO$_2$, Alkalinity, and Nitrate**

- NAST (27°–42°N): 3.9 province means of data presented in Figure 2; from$^f$
- NADR (42°–55°N): 4.3 $^{''-}$; from$^f$
- SARC/ARCT (>55°N): 6.1 $^{''-}$; from$^f$
- north of 40°N: 5.3–5.9 from$^f$
- 40°S–40°N: 4.9–5.1 from$^f$
- (4.2–4.4) ratio in brackets is calculated for data without the warming period: whole year scaling factor used by$^b$
- south of 40°S: 8.9–12.1 from$^f$

$^a$Koeve [2001].
$^b$Robertson et al. [1994].
$^c$Takahashi et al. [1990] as cited in the study by Robertson et al. [1994].
$^d$Veldhuis et al. [1993].
$^e$Sambrotto et al. [1993].
$^f$This study.
$^g$Antia et al. [2001].
$^h$Lee [2001].
translates to the annual POC:PIC ratio, depends on whether all post-diatom bloom nitrate is taken up over the growth season and which share coccolithophorids can take. If one assumes that all post-diatom bloom nitrate uptake is by coccolithophorids (with a POC:PIC ratio of 1:1, Robertson et al., 1994), an annual POC:PIC ratio of net production between 2.9 and 4.0 is calculated for 47°N, and slightly higher values are found for 60°N (3.5–4.2) (Table 3).

[22] These are minimum estimates for the POC:PIC ratio of production, since new production by non-calculating flagellates is ignored and since no correction for air sea exchange is taken into account. New production by non-calculating phytoplankton after the diatom spring bloom will reduce the pool of nitrate and phosphate available for coccolithophorids and thereby increase the annual POC:PIC production ratio. Air-sea exchange, which was prominent (up to 25% of the apparent total CO2 drawdown over 2–3 months) when the C:N ratios of new production was calculated from seasonal changes of total CO2 and nitrate [Körtzinger et al., 2001a], will act into the same direction.

In particular, it will increase the POC:PIC ratio during the diatom spring bloom, since air-sea pCO2 gradients and wind speed are higher during that season. Currently few process studies are available to describe the temporal and regional distribution of the POC:PIC export ratio in more detail. In the following section, I therefore present a first order estimate of the large scale distribution of the annual RRexp in the Atlantic from a climatological perspective.

3.2.3. Large Scale Distribution of RRexp

[23] In temperate and subarctic waters the surface mixed layer POC:PIC ratio of production (RRML) may be derived from the seasonal changes in salinity normalized total alkalinity and nitrate in the surface mixed layer and the C:N ratio of new production (equation (7)). Operationally it is assumed here that new carbon production (NP) which is exported via particles can be estimated from the seasonal drawdown of nitrate by multiplying with a constant C:N ratio of 6.625 [Redfield et al., 1963; Körtzinger et al., 2001a]. CaCO3 production (PCaCO3) is estimated from seasonal changes of alkalinity and nitrate according to the concept of potential alkalinity (see section 3.2.2). In this section, however, data from climatologies of nitrate, alkalinity, and salinity are used to compute RRML. Details of the computation are given in Appendix A. It is assumed that the mixed layer POC:PIC ratio is an appropriate proxy of the export POC:PIC ratio RRexp (i.e., RRexp ≈ RRML).

\[
RRML = \frac{NP}{PCaCO3} = \frac{\Delta NO3 \times C/N}{(\Delta A + \Delta NO3)/2} = \frac{\Delta NO3}{(\Delta A + \Delta NO3)} \times (C/N) \times 2
\]  

[24] In the subtropics and tropics the seasonal approach cannot be used since seasonal gradients of alkalinity and nitrate vanish. As a substitute I estimate RRexp from particle interceptor trap measurements [Lohrenz et al., 1992] at the Bermuda Time Series Station (BATS). RRexp at BATS is estimated to be 4.1–4.5 (see Appendix A for details).

[25] The major feature of the estimate of RRexp in temperate and subarctic North Atlantic is a polarward gradient, overlaid by a weaker east-west gradient with maximal values in the North East Atlantic (Figure 2). Minima of RRexp are seen at the boundary between the temperate and the subtropical domains (35°N and S) which is also the boundary between estimates from the seasonal change of potential alkalinity and estimates from shallow traps at BATS. The intensity of these minima is subject to the choice of the C:N ratio of sinking particles (see equation (7)) and the choice of the geographic position of the boundary itself. Province mean values of RRexp in the temperate and subarctic North Atlantic increase northward from 3.9 (NAST) to 6.1 (SARC/ARCT) (Table 3), the weighted mean for the temperate and subarctic North Atlantic (all three provinces) is 4.37, which compares well with RR125 from the exponential RR-Z fit.

[26] Recently, Lee [2001] presented an analysis of the global net community production and the global CaCO3 production based on an analysis of the annual cycle of surface water total CO2 and alkalinity. They estimated organic carbon:inorganic carbon uptake ratios of 5.3–5.9 for the temperate and subarctic Atlantic (40°N–70°N), slightly lower values for the subtropical and tropical band (40°S–40°N: RRML = 4.9–5.1) but significantly higher values for the southern Ocean (Atlantic south of 40°S: RRML = 8.9–12.1). Their Atlantic integral value was 6.1–6.9. The estimate of net carbon production from seasonal changes in total CO2 [Lee, 2001] does not differentiate whether carbon is stored and exported as particles or as DOC. One might use the difference between the RRML estimate from this study (4.2–4.37) and those from the study of Lee [2001] to approximate that on the mean 39–58% of net carbon production in the Atlantic is channeled through a seasonally accumulating pool of DOC [Kähler and Koeve, 2001] which eventually is exported from the mixed layer during winter time convection [Carlson et al., 1994] but is resired above the winter mixed layer depth or on shallow isopycnals [Doval and Hansell, 2000] and hence does contribute little to carbon sequestration. Regarding the involved uncertainties (about ±30% for the estimates based on data from the work of Lee, 2001), both studies support the concept of organic:inorganic carbon ratio of export production in the Atlantic is close to the canonical value of about 4–5 [Li et al., 1969; Broecker and Peng, 1982].

3.2.4. Polarward Increase of RRexp

[27] Estimates from this study (sections 3.2.2 and 3.2.3) suggest a polarward increase of RRexp in the Atlantic Ocean. This sounds counterintuitive since the subpolar North Atlantic, in particular waters south of Iceland, is well known for the frequent occurrence of coccolithophorid blooms [Holligan and Balch, 1991; Holligan et al., 1993; Fernandez et al., 1993; Balch et al., 1996] and one might therefore expect the POC:PIC ratio of export production to be low.

[28] In section 3.2.2 (and Table 3) it was shown that the northward increase of RRexp goes along with increases in the contribution of diatoms to annual new production, which were driven by a decrease of the winter NO3:Si(OH)4 ratio. Because of an overall deepening of the winter mixed layer from the subtropics to the subarctic North Atlantic winter time nutrient concentrations increase polarward [Glover and Brewer, 1988; Koeve, 2001]. Coccolithophorid blooms, i.e.,
high absolute rates of PIC production and the biomass of coccolithophorids, require high nitrate stocks after the diatom bloom has ended because of the depletion of silicate. Post-diatom bloom nitrate concentrations in the euphotic zone can be calculated to the first order from winter nitrate and silicate concentrations according to NO₃(post) = NO₃(winter) / C₀Si(OH)₄(winter) / C₂(N/Si)uptake ratio (like in section 3.2.2, a mean (N/Si)uptake ratio = 1 is assumed). Between 41°C and 59°C, the post-diatom bloom nitrate concentration increases from 2.9 to 6.5 mmol N m⁻³ at 59°C (calculated after data presented by Koeve [2001]). Simply by the availability of nitrate after the diatom bloom has ended there is an increasing potential for coccolithophorid blooms and CaCO₃ production toward the north. At the same time diatom (Table 3) and hence the POC:PIC production ratio increases northward. It is concluded, that the seemingly contradiction between increasing RRₑₓᵖₓ toward the northern North Atlantic (this study) and observations of huge coccolithophorid blooms south of Iceland can be explained by the north-south distribution of relative and absolute rates at which nitrate and silicate are supplied to the surface ocean during winter mixing.

3.3. Evaluation of the Change of the PIC Flux Over Depth

[29] One of the oldest paradigms in oceanography states that pelagic CaCO₃ dissolves exclusively below the lysocline and, due to high sinking speeds and slow dissolution, particularly within the sediments. Major evidence for this view is from the observation that vast areas and depth ranges of the ocean show a low corrosity for CaCO₃ (i.e., saturation state Ω > 0.8) and from experimental studies of CaCO₃ dissolution [Peterson, 1966; Keir, 1980]. Along the same line, sediment trap data suggested that the PIC flux does not decrease significantly below 1000 m depth [Tsunogai and Noriki, 1991; Milliman, 1993]. This is accentuated in the North Atlantic [Honjo, 1990; Honjo and Manganini, 1993], where the depth of the calcite-lysocline is found at about 4000–4500 m.

[30] Combining the J-PP-Z functions of Koeve (submitted manuscript, 2001) (Table 1) and the POC:PIC ratio fits from this study allows to estimate changes in the PIC fluxes with depth. All parameter sets predict a significant decrease of the PIC flux in the upper 1000 m. The ratio of PIC fluxes in 125 and 1000 m, JPIC₁₂₅/JPIC₁₀₀₀, varies between 1.4 and 2.5 (exponential RR fit; Figure 1c), indicating that 31–59% of the PIC export dissolves in the upper 1000 m. There are few data from (drifting) shallow sediment traps to compare this estimate with. The experiment of Martin et al. [1993] during the North Atlantic Bloom Experiment (NABE), in which floating VERTEX-type traps were deployed between 150 and 2000 m over the course of the spring bloom (6 weeks), supports a similar decrease in PIC fluxes of about

Figure 2. Regional distribution of RRₑₓᵖₓ in the Atlantic Ocean. North of 35°N and south of 35°S, RRₑₓᵖₓ is estimated from seasonal changes of potential alkalinity and nitrate. In the subtropical bands (10°–35°N, 10°–35°S) an estimate based on particle interceptor trap measurements at the Bermuda Time series Station (BATS) is assumed as a constant value. In the tropical band (10°S–10°N) an export flux ratio of 4.37 is used. For details see section 3.2.3 and the Appendix A.
50% in the upper 1000 m. Milliman et al. [1999] summarize the currently available evidence for shallow PIC dissolution and suggest that on average about 59% of the global carbonate production dissolves in the upper 1000 m. [31] It is in this depth horizon where organic carbon fluxes change most rapidly; the ratio of the organic carbon fluxes at 125 and 1000 m, JPIC125/JPIC1000, varies between 4.5 and 8.1, suggesting an organic carbon remineralization of 77–88% within the upper 1000 m. A possible mechanism driving CaCO3 dissolution above the lysocline is dissolution within “microenvironments” of high pHCO2, derived from intense remineralization of organic carbon. Here, the concentration of CO32− could decrease locally such that the concentration product of CO32− and Ca2+ becomes smaller than the dissolution product against the dissolution of CaCO3. Such microenvironments might be either fecal pellets or aggregates of particulate organic matter. Organic aggregates [Riley, 1963; “marine snow”], which are a very important vehicle for the transport of biogenic matter into the deep ocean [Fowler and Knauer, 1986], are known as sites of active metabolism [Pomeroy and Johannes, 1968; Turley, 1992] and may be sites of shallow and microscale CaCO3 dissolution. The build up of a pCO2 (and CO32−) gradient between the aggregate and the surrounding water would be a necessary prerequisite of a significant CaCO3 dissolution in such aggregates. There is, however, conflicting evidence whether the degradation products (nutrients, CO3, oxygen debt) accumulate in aggregates [Brzezinski et al., 1997] or not [Ploug and Joergensen, 1999]. Another possible process to promote shallow PIC remineralization is dissolution in the guts of zooplankton [Takahashi, 1975]. Critical to this issue is, whether the pH of grazer guts is high [Honjo and Roman, 1978] or may, like in the case of starved copepods [Pond et al., 1995], be low enough to support CaCO3 dissolution. Recently, laboratory feeding experiments comparing the amount of ingested and egested coccolithophors, indicated dissolution rates of 50% and more [Harris, 1994]. Milliman et al. [1999] provide a detailed and thorough overview of the current evidence for and the processes involved in shallow CaCO3 dissolution.

Below 1000 m, PIC dissolution decreases with increasing depth (7.5–10.5% of JPIC125 between 1000 and 2000 m, and 7–8% between 2000 and 4000 m). Overall, 46–77% of JPIC125 dissolves above 4000 m depth. This is in good agreement with the finding of Archer [1996] and Archer et al. [2000a] that the global production rate of CaCO3 required by the ocean surface/deep alkalinity contrast [Li et al., 1969] is higher by roughly a factor of three than the average CaCO3 rain rate to the seafloor required by sediment diagenesis models of Archer [1996], hence about 2/3 of calcium carbonate production dissolves in the water column.

### 3.4. POC:PIC Ratio of Remineralization

In this section the organic carbon:inorganic carbon remineralization ratio (RRrem) is estimated from the change of POC and PIC particle fluxes over depth, and from an analysis of the along-isopycnal change of the degradation products of POC remineralization and PIC dissolution in the interior of the ocean.

In the latter case, RRrem is calculated from $$RR_{rem} = \frac{\Delta \Sigma CO_2}{\Delta Ca} - 1$$ were Ca is computed from total alkalinity (TA) and nitrate, according to Ca = (TA + NO3)/2. ΔΣCO2 and ΔCa are estimated from regression slopes versus oxygen (ΔO2) on isopycnals or neutral surfaces. I use ΔO2/ΔCa ratios estimated by Takahashi et al. [1985] for shallow isopycnals just below the permanent thermocline (sigma-theta density levels σθ = 27.0 and σθ = 27.2; Table 4). Takahashi et al. [1985] report ΔO2/ΔΣCO2 ratios, which range between −1.7 ± 0.1 and −2 ± 0.1. Their analysis, however, did not correct for the time history of the invasion of anthropogenic CO2 into the ocean [Takahashi et al., 1985]. The temporal change in anthropogenic CO2 in the atmosphere causes a slight increase over time in the initial CO2 concentration when a water mass is formed. If this effect is not taken into account, the ΔΣCO2 due to remineralization along an isopycnal is underestimated and the ΔO2/ΔΣCO2
ratio of remineralization is overestimated. Correcting for the time history of anthropogenic CO₂ invasion, Körtzinger et al. [2001b] estimate a \( \Delta O_2/\Delta \Sigma CO_2 \) ratio of \(-1.3 \pm 0.1\) for isopycnals just beneath the permanent thermocline. \( R_{\text{remin}} \) on shallow isopycnals, thereafter, is estimated to range between 7.5–16.0 (North Atlantic) and 14.4–17.5 (South Atlantic) (see Table 4). These are mean estimates for the given ocean basins within which the specified isopycnals are encountered over a broad depth range of 250–700 m (\( \sigma_y = 27.0 \)) and 450–1000 m (\( \sigma_y = 27.2 \)).

Within this depth range mean \( R_{\text{remin}} \) estimated from particle flux data (exponential \( R_R-Z \) relationship) range from 6.8 to 14.7 and 3.1 to 6.7 (Figure 3, thick lines) for values of the \( Z \)-exponent of the \( J-PP-Z \) function of \( b = -0.7 \) and \( b = -1.0 \), respectively. This range of \( b \)-values was found to be representative for the particle flux data set used in this study (Table 1). If the \( R_{\text{remin}} \) estimate is based on the \( RR-\ln Z \) relationship (Figure 3, thin lines) much lower values are estimated (0.9–2.4). Clearly \( R_{\text{remin}} \) estimates from the exponential \( RR-Z \) relationship compare better with those from the logarithmic \( RR-Z \) relationship. The difference between \( R_{\text{remin}} \) estimates from the isopycnal analysis and from the exponential \( RR-Z \) particle flux relationship may be attributed to remineralization of DOC, which has been found to be significant on shallow isopycnals [Doval and Hansell, 2000].

3.5. POC:PIC Ratios: Some Conclusions

Two alternative mathematical functions with almost identical statistics can be used to represent the mean depth dependence of the POC:PIC ratio of particle flux in the Atlantic ocean (section 3.1). However, only one of the models, the exponential function \( RR = 64.3Z^{-0.56} \), yields (a) a consistent vertical distribution of the PIC flux (sections 3.1 and 3.3 and Figure 1c), (b) POC:PIC ratios of remineralization which match the distribution of breakdown products of POC remineralization and PIC dissolution on shallow isopycnals (section 3.4, Figure 3 and Table 4), and (c) POC:PIC export ratios \( (R_{\text{exp}}) \) which agree with basin scale means of \( R_{\text{ML}} \) estimates from seasonal changes of nutrients and alkalinity (section 3.2). On the contrary, the logarithmic \( RR-Z \) relationship which was adopted from the work of Antia et al. [2001] yields an unrealistic PIC increase in the deep ocean, too low POC:PIC ratios of remineralization, and also a too low mean POC:PIC export ratio.

4. Effective Carbon Flux

The calculation of the effective carbon flux is based on three quantities, the sequestration fluxes of organic and inorganic carbon \( (J_{\text{POC-seq}}, J_{\text{PIC-seq}}) \) and \( \Psi \), the integral ratio of released CO₂:precipitated carbonate (equation (4)). Two sets of \( J_{\text{eff}} \) calculations will be performed in section 4.4. In the first set one mean value for \( \Psi \) will be used together with estimates of \( J_{\text{POC-seq}} \) adopted from the study of Köve (submitted manuscript, 2001) and \( J_{\text{PIC-seq}} \) estimated after equation (2), applying the POC:PIC ratio estimates from the exponential \( RR-Z \) fit, i.e., with one, regionally invariable, vertical distribution of RR and a regionally constant value of \( R_{\text{exp}} \). In the second set a regionally variable data set of \( \Psi \) (section 4.3) will be combined with a data set of \( J_{\text{PIC-seq}} \) which is adjusted to the regional distribution of \( R_{\text{exp}} \) discussed in section 3.2.3 and shown in Figure 2 and with the estimates of \( J_{\text{POC-seq}} \) from the work of Köve (submitted manuscript, 2001). In the following three sections, I give a brief description of the distribution of \( J_{\text{POC-seq}} \) (section 4.1), \( J_{\text{PIC-seq}} \) (section 4.2), and \( \Psi \) (section 4.3). In section 4.4 the resulting data fields of \( J_{\text{eff}} \) are discussed.

4.1. Distribution of \( J_{\text{POC-seq}} \)

The distribution of organic carbon fluxes in the Atlantic is discussed in detail in the study of Köve (submitted manuscript, 2001) and is described here only in brief. Basin scale (65°N–65°S) integrals of export production computed from eight independent parameter fits (Table 1) of equation (1) varied between 0.9 and 2.9 GT C yr\(^{-1}\) (Table 5); the respective \( ER_{125} \) \( (J_{\text{POC125-PP}}) \), which is an estimate of the \( f \)-ratio, ranged from 0.07 to 0.31. Not only overall fluxes and ratios differ, but also their N-S distribution patterns and the relative importance of tropical versus “spring-bloom” biogeographical provinces (Figure 4a). The regional distribution of export production and \( ER_{125} \) was in particular sensitive to the choice of the primary production input data set (data presented by Antoine et al. [1996] versus data presented by Behrenfeld and Falkowski [1997]). The percent fraction of carbon remineralized within the winter mixed layer (Figure 4b) could be estimated more robustly. For the Atlantic Ocean as a whole remineralization within the winter mixed layer reduces the carbon export by 11–17% (Table 5). In the temperate and subpolar provinces of the North Atlantic, this
Table 5. Basin Wide Integrals for the North Atlantic

<table>
<thead>
<tr>
<th></th>
<th>AM96-Based,b</th>
<th>BF97-Based,c</th>
<th>Minimum Estimate</th>
<th>Maximum Estimate</th>
</tr>
</thead>
<tbody>
<tr>
<td>Primary production</td>
<td>9.64</td>
<td>12.88</td>
<td>0.92</td>
<td>2.95</td>
</tr>
<tr>
<td>Export production</td>
<td>2.17 ± 0.722</td>
<td>1.47 ± 0.416</td>
<td>0.25 ± 0.084</td>
<td>0.14 ± 0.35</td>
</tr>
<tr>
<td>Remineralization</td>
<td>0.26 ± 0.091</td>
<td>0.25 ± 0.084</td>
<td>0.77</td>
<td>2.60</td>
</tr>
<tr>
<td>$\psi$ JPIC&lt;sub&gt;sequ&lt;/sub&gt;</td>
<td>1.91 ± 0.631</td>
<td>1.21 ± 0.333</td>
<td>0.14</td>
<td>0.43</td>
</tr>
<tr>
<td>$\psi$ JPOC&lt;sub&gt;eff&lt;/sub&gt;</td>
<td>0.52 ± 0.102</td>
<td>0.21 ± 0.056</td>
<td>0.14</td>
<td>0.43</td>
</tr>
<tr>
<td>Respiration ratio</td>
<td>1.60 ± 0.529</td>
<td>1.00 ± 0.277</td>
<td>0.64</td>
<td>2.18</td>
</tr>
<tr>
<td>$r_{CaCO3}$&lt;sub&gt;b&lt;/sub&gt;</td>
<td>0.12 ± 0.006</td>
<td>0.16 ± 0.012</td>
<td>0.11</td>
<td>0.17</td>
</tr>
<tr>
<td>$r_{CaCO3}$&lt;sub&gt;f&lt;/sub&gt;</td>
<td>0.15 ± 0.002</td>
<td>0.14 ± 0.003</td>
<td>0.14</td>
<td>0.15</td>
</tr>
<tr>
<td>$r_{CaCO3}$&lt;sub&gt;g&lt;/sub&gt;</td>
<td>0.14 ± 0.004</td>
<td>0.14 ± 0.004</td>
<td>0.13</td>
<td>0.14</td>
</tr>
<tr>
<td>$r_{CaCO3}$&lt;sub&gt;h&lt;/sub&gt;</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

a Estimates are presented for the Atlantic Ocean between 65°N and 65°S; marginal seas like the Baltic, the Hudson Bay, and the Mediterranean are excluded. The integration is based on a 1° × 1° data set, grid points where the seasonal maximum of the mixed layer depth is equal or larger than the mean water depth at the grid point are ignored.
b AM96-based estimates use the primary production data set of Antoine et al. [1996].
c BF97-based estimates use the primary production data set of Behrenfeld and Falkowski [1997].
d All in GT POC-C-equiv. yr<sup>-1</sup>; ratios (PR – POC<sub>sequ</sub>, J/PIC<sub>125</sub>) are all in mol:mol.
e From Koeve (submitted manuscript, 2001).
f For the standard case: $\psi = 0.67$; RR<sub>exp</sub> = 4.37.
g With $\psi$ from Figure 6a, RR<sub>exp</sub> from Figure 2.
h With $\psi$ from Figure 6b, RR<sub>exp</sub> from Figure 2.
i With $\psi$ from Figure 6b, RR<sub>exp</sub> as in the standard case.

Reduction is equivalent to 37–61% of the export production. The sequestration of organic carbon, POC<sub>sequ</sub>, was estimated to range between 0.77 and 2.6 GT C yr<sup>-1</sup> (Table 5), the regional distribution of the province means of POC<sub>sequ</sub> is shown in Figure 4c.

4.2. Distribution of JPOC<sub>sequ</sub>

The distribution of the PIC export (JPIC<sub>125</sub>, Figure 5a) is basically a scaled plot of the export production, if the POC:PIC flux ratio (RR) is estimated as a function of depth only and RR<sub>exp</sub> = RR<sub>125</sub> = constant = 4.37. The estimate of the Atlantic wide integral of JPOC<sub>125</sub> ranges from 0.21 to 0.68 GT C yr<sup>-1</sup>, PIC dissolution within the winter mixed layer, as predicted from the change of RR over depth, accounts for 0.011–0.047 GT C yr<sup>-1</sup> or 4.6–9.4% of the export PIC flux. The north-south distribution of the ratio R<sub>R-PIC<sub>sequ</sub></sub> (=Remin<sub>PIC<sub>sequ</sub></sub>/JPOC<sub>125</sub>) (Figure 5b) is similar to the distribution of the fraction of POC remineralization to POC export production, R<sub>R-PIC<sub>sequ</sub></sub> (Figure 4b), however, the absolute values of R<sub>R-PIC<sub>sequ</sub></sub> are always smaller than R<sub>R-PIC<sub>sequ</sub></sub> JPOC<sub>sequ</sub> differs from JPOC<sub>125</sub> mainly in the North Atlantic (Figure 5a, c).

4.3. Released CO<sub>2</sub>: Precipitated Carbonate Ratio, $\psi$

4.3.1. Mean $\psi$

The ratio of released CO<sub>2</sub>:precipitated carbonate during CaCO<sub>3</sub> formation, $\psi$, is controlled by carbonate system chemistry and shows a weak relationship with salinity, a negative correlation with temperature, and a positive correlation with pCO<sub>2</sub> [Frankignoulle et al., 1994]. The value of $\psi$ in the surface ocean will mainly vary together with temperature and pCO<sub>2</sub>. For the early 1990s, the time period of sediment trap deployments used in this study, an integral value for $\Gamma = 0.67$ can be estimated. This value assumes a mean ocean equilibration temperature of about 15°C [Archer et al., 2000b], S = 35, and a mean surface ocean pCO<sub>2</sub> value of 350 µatm, and is estimated by interpolation of data presented in the study by Frankignoulle et al. [1994] (their Figure 2). This is a more realistic mean value than $\Gamma = 0.6$, which was applied by Antia et al. [2001], since that value requires the mean temperature of the ocean to be 25°C, for S = 35 and pCO<sub>2</sub> = 350 µatm [Ware et al., 1992].

4.3.2. Regional Distribution of $\psi$ in the Atlantic Ocean

In the following an estimate of the regional distribution of $\psi$ in the Atlantic is provided. Since these estimates of $\psi$ will be used to compute annual fluxes of J<sub>eff</sub> they need to integrate in a suitable way over the depth range between the surface and Z<sub>sequ</sub> and over the annual mixing cycle. Accordingly, I have chosen to estimate $\psi$ based on the local mean temperature of the mixed layer at the time of deepest mixing (i.e., during winter). CaCO<sub>3</sub> production might take place under conditions of significantly warmer temperatures during summer (i.e., with a smaller $\psi$). However, the net effect on the steady state surface ocean pCO<sub>2</sub> is fixed at the end of winter time mixing when surface waters, which were “impoverished” in CO<sub>2</sub> due to CO<sub>2</sub>-uptake during new production, have mixed with subsurface waters, which were enriched in CO<sub>2</sub> due to remineralization of organic carbon, and high wind speeds have brought the surface ocean, which during this season is characterized by low biological activity and low biogenic CO<sub>2</sub> fluxes, as close as possible to equilibrium with the atmosphere. A first estimate ($\psi_{350}$, Figure 6a) is based on the assumption that the surface pCO<sub>2</sub>
is in equilibrium with the atmosphere ($p_{\text{CO}_2}^{\text{atm}} = 350 \mu\text{atm}$). $\psi_{350}$ increases from low values (0.56–0.6) in the tropics to 0.7–0.82 in high latitudes. A second estimate of the distribution of $\psi$, in this case corrected for the deviation of the winter time surface $p_{\text{CO}_2}$ from equilibrium with the atmosphere $p_{\text{CO}_2}^{\text{insitu}}$ is given in Figure 6b. Both estimates agree within ±0.02 mol:mol north of about 40°S. It is concluded that the specific effect of CaCO$_3$ sequestration on $J_{\text{eff}}$ is sensitive to whether CaCO$_3$ is formed in cold or in warm (high- or low-latitude) waters. As I shall discuss in the next section in more detail, the polarward increase of $\psi$ tends to counteract the polarward increase of the POC:PIC export ratio discussed in sections 3.2.2 and 3.2.3.

Figure 4. North-South distribution of organic carbon export. Meridional integrals of characteristic open ocean biogeographical provinces, basically following the schema suggested by Longhurst [1998], are presented. Province definitions used in this study are as follows (numbers are as indicated on the lower x-axes): (2) Atlantic sectors of the Antarctic Province (ANTA; 65°S–55°S, 70°W–20°E), (3) Subantarctic Water Ring Province (SANT; 55°S–45°S, 70°W–20°E), (4) South Subtropical Convergence Province (SSTC; 45°S–35°S, 70°W–20°E), (5) South Atlantic Gyral Province (SATL; 35°S–10°S), (6) Western and Eastern Tropical Atlantic Province (WETRA; 10°S–10°N), (7) North Atlantic Tropical Gyral Province (NATR, 10°N–27°N, including the Caribbean Province), (8) North Atlantic Subtropical Gyral Province (NAST, 27°N–42°N, including the Gulf Stream Province), (9) North Atlantic Drift Region (NADR; 42°N–55°N), and (10) the combined Atlantic Subarctic and Arctic Provinces (SARC and ARCT; 55°N–65°N). The following properties are shown: (a) province means of the export production $J_{\text{POC}}^{125}$ (g C m$^{-2}$ yr$^{-1}$); (b) $r_{R-\text{POCsequ}}$, the fraction of POC-remineralization above the winter mixed layer ($R_{\text{POCsequ}}$): export production ($J_{\text{POC}}^{125}$); (c) province means of $J_{\text{POCsequ}}$, the sequestration flux of organic carbon, (g C m$^{-2}$ yr$^{-1}$); (d) the province mean of $Z_{\text{sequ}}$ (m). Solid lines/circles: means ± standard deviation of estimates based on primary production forcing data from the study of Antoine et al. [1996], dashed lines/squares: Behrenfeld and Falkowski [1997] based estimates. Modified after the work by Koeve (submitted manuscript, 2001).

Figure 5. North-south distribution of particulate inorganic carbon fluxes in the Atlantic Ocean. The following properties are shown: (a) province means of the PIC export ($J_{\text{PIC}}^{125}$, g C m$^{-2}$ yr$^{-1}$), (b) the fraction of PIC dissolution above the winter mixed layer depth, (c) province means of the PIC-sequestration flux ($J_{\text{PICsequ}}$, g C m$^{-2}$ yr$^{-1}$), and (d) $R_{\text{sequ}}$, the POC:PIC flux ratio at $Z_{\text{sequ}}$. For details see text. Solid lines/circles: estimates based on primary production forcing data from the study of Antoine et al. [1996], dashed lines/squares: estimates based on the work of Behrenfeld and Falkowski [1997]. For province captions see Figure 4.
Figure 6. Regional distribution of $\psi$, the integral ratio of released CO$_2$:precipitated carbonate during CaCO$_3$ sequestration in the Atlantic Ocean. (a) A $p$CO$_2$ of 350 $\mu$atm is assumed and $\psi$ is calculated from the relationship $\psi_{350} = 0.8 - 8.3 \times 10^{-3} \times T$ [Frankignoulle et al., 1994] where $T$ is the mixed layer temperature during the time of deepest mixing. Temperature data for this computation where taken from the monthly resolving analyzed temperature ($1^\circ \times 1^\circ$) data set of the WOA98 [Antonov et al., 1998]. The time of deepest mixing was estimated from the climatology of mixed layer depths ($\Delta \sigma = 0.125$ kg m$^{-3}$) taken from the World Ocean Atlas 1994 (Monterey and Levitus [1997]). (b) $\psi_{pCO_2_{\text{insitu}}}$ is $\psi$ at in situ $p$CO$_2$ at the time of deepest mixing, $p$CO$_2_{(w)}$. $p$CO$_2_{(w)}$ is taken from Koeve (unpublished data, 2001), basically it is calculated from a sea-air CO$_2$ difference climatology [Takahashi et al., 1995, 1999] and an estimate of the atmospheric $p$CO$_2$ at the time of mixing. The relationship between $\psi_{pCO_2_{\text{insitu}}}$ and $\psi_{350}$ for a given temperature is estimated from a logarithmic fit of data presented in the study by Frankignoulle et al. [1994].
4.4. Distribution of \( J_{\text{eff}} \)

[43] The effective carbon flux, \( J_{\text{eff}} \), is calculated according to equation (4) from estimates of \( \text{JPOC}_{\text{sequ}} \) (Figure 4c), \( \text{JPIC}_{\text{sequ}} \), and \( \Psi \). Several cases are considered. For the standard case (Figure 7a, b) a constant value for \( \Psi \) of 0.67 is combined with a \( \text{JPOC}_{\text{sequ}} \) data set (Figure 5c) which is computed by applying the POC:PIC ratio estimates from the exponential RR-Z fit, i.e., with one, regionally invariable, vertical distribution of RR. Sensitivity computations are carried out by applying estimates of \( \Psi \) shown in Figures 6a and b and a depth and regionally variable set of RR, which is computed from \( \text{RR}_{125} \) shown in Figure 2 and the \( z \)-exponent \( (e = -0.5) \) of the exponential RR-Z function. Throughout this section I will distinguish between estimates for which the calculation of the organic fraction (\( \text{JPOC}_{\text{sequ}} \)) is driven by primary production estimates derived by Antoine et al. [1996] (AM96 based estimates) and those by Behrenfeld and Falkowski [1997] (BF97 based estimates).

[44] Estimates of the basin scale integral of \( J_{\text{eff}} \) (Table 5) vary between 0.64 and 2.2 GT C m\(^{-2}\) yr\(^{-1}\), where the higher estimates are based on the AM96 primary production data set and the lower are based on BF97 primary production data. Most of this difference is due to high AM96-based fluxes in tropical and subtropical waters (Figure 7a, b). Elevated tropical and subtropical AM96 based fluxes are a consistent feature which \( J_{\text{eff}} \) shares with the distribution of primary production itself, \( \text{JPOC}_{125} \) (Figure 4a), \( \text{JPOC}_{\text{sequ}} \) (Figure 4c), and \( \text{JPIC}_{\text{sequ}} \) (Figure 5c). On the contrary BF97 based estimates of \( J_{\text{eff}} \), in particular the province mean values (Figure 7b), show little structure along the N-S axes. Maxima in temperate and subpolar waters in both hemispheres, which are evident for the export production (\( \text{JPOC}_{125} \), Figure 4a), have disappeared in the distribution of \( J_{\text{eff}} \). The major process that drives this change in the regional distribution is the shallow remineralization of organic carbon.

[45] The fraction of organic carbon export, which is compensated by the CaCO\(_3\) sequestration flux \( r_{\text{CaCO}_3} = \Psi \times \text{JPIC}_{\text{sequ}}/\text{JPOC}_{125} \) is almost constant everywhere in the Atlantic (Figure 7c). For the standard case a slight reduction can be seen in high latitudes of the North Atlantic due to the effect of shallow PIC dissolution (Figure 5b) in waters with deep winter mixing (Figure 4d). The fraction \( r_{\text{CaCO}_3} \) is more variable if \( \Psi \) and \( \text{RR}_{\text{exp}} \) are allowed to vary regionally (Figure 7d). The main feature here is a stronger gradient from tropical waters toward mid latitudes and from mid latitudes toward high latitudes in the North Atlantic. \( r_{\text{CaCO}_3} \) can be up to 50% larger, like in NAST, compared with the standard case. The basin scale effect of regionally variable values of \( \Psi \) and \( \text{RR}_{\text{exp}} \) however, is small (Table 5). This is mainly due to the fact that the basin scale averages of the data fields of \( \text{RR}_{\text{exp}} \) (4.15) and \( \Psi \) (\( \Psi_{350} = 0.66 \) and \( \Psi_{\text{pCO}_2,\text{in situ}} = 0.64 \)) are close to the a priori chosen (or from particle flux data estimated) mean values used in the standard case (\( \Psi = 0.67 \); \( \text{RR}_{\text{exp}} = 4.37 \)). In addition, poleward trends obvious for \( \Psi_{350} \) or \( \Psi_{\text{pCO}_2,\text{in situ}} \) and \( \text{RR}_{\text{exp}} \) or \( \text{RR}_{\text{sequ}} \) counteract each other and become insignificant on the basin scale. The use of constant values for \( \Psi \) and \( \text{RR}_{\text{exp}} \) for the basin-scale estimates of \( J_{\text{eff}} \) is justified by these findings.

[46] On the basin scale shallow remineralization accounts for 11–17% and CaCO\(_3\) compensation for 14–15% of export production. Overall \( \text{JPOC}_{\text{exp}} \) overestimates the carbon flux which is effective with respect to surface to deep CO\(_2\) gradients by about 30% for the Atlantic as a whole and for up to 80% in the subarctic North Atlantic.

5. Summary and Conclusions

[47] The POC:PIC ratio of particle flux in the Atlantic Ocean can be described by two empirical relationships with
almost identical statistics. It is found that a logarithmic fit of the POC:PIC flux ratio and \( Z \), which was adopted from the work of Antia et al. [2001], yields an unprecedented increase of PIC flux in the deep ocean and too low POC:PIC ratios of remineralization at intermediate water depths. An exponential fit of the POC:PIC flux ratio over \( Z \), on the other hand, yielded remineralization POC:PIC ratios and deep PIC fluxes which are consistent with independent estimates from the literature.

[48] The basin-scale mean POC:PIC ratio of export production (RR\text{exp}) in the Atlantic as estimated from the exponential fit is 4.37. Seasonal changes of nitrate concentrations and alkalinity from temperate and subarctic waters were used in combination with sediment-trap observations from the BATS to construct a map of RR\text{exp} which is independent of the deep particle-flux data used elsewhere in this study. Estimates of RR\text{exp} ranged between 2 and 9, but the basin-scale integrated value (4.2) agreed with the estimate inferred from deep traps (4.37).

[49] A ratio of organic carbon:inorganic carbon net production, which comprises POC and DOC production, of 6.1–6.9 was recently estimated for the Atlantic by Lee [2001] from an analysis of the seasonal changes of total CO\(_2\), alkalinity, and nitrate. The global estimate of the organic carbon:inorganic carbon production ratio of 8.3–9.8 found by Lee [2001], which is close to the range of 10–12.5 suggested from a modeling study by Yamanaka and Tajika [1996] is particularly due to high values in the subtropical Pacific (13.3–17.4) and the southern ocean (7.0–9.9). These ocean to ocean differences of primary organic: inorganic uptake ratios between the Atlantic and the Pacific resemble the distinction between a “carbonate ocean” (the Atlantic) and a “silicate ocean” (the Pacific) suggested earlier by Honjo [1996].

[50] A map which describes the regional distribution of the vertically and seasonally integrated released CO\(_2\):precipitated carbonate ratio (\( \psi \)) during CaCO\(_3\) formation is prepared for the early 1990s, the time when most of the particle flux studies used in this study where conducted. \( \psi \) varies between values around 0.56 in the tropics to values as high as 0.8 in high latitudes, which indicates that CaCO\(_3\) sequestration more effectively compensates POC sequestration in high latitudes than in low latitudes. However, the basin scale integral of \( \psi \) has about the same value (0.64–0.66) as the first guess choice for a regionally constant \( \psi \) (0.67), which was estimated to be consistent with the mean surface ocean \( pCO_2 \) equilibrium temperature of the contemporary ocean.

[51] Estimates of \( J_{\text{eff}} \) ranged from 0.66 to 2.18 GT C yr\(^{-1}\), this large range is in particular due to the propagation of regional differences of primary production maps [Antoine et al., 1996; Behrenfeld and Falkowski, 1997], which are used as forcing data to estimate the flux of organic carbon in this study. The integrated POC:PIC sequestration flux ratio for the Atlantic ocean is about 3.9, with much lower values in the subarctic North Atlantic. On the basin scale shallow remineralization of POC and CaCO\(_3\) compensation are about equally important in adjusting \( J_{\text{eff}} \). The effect of CaCO\(_3\) compensation on POC sequestration shows little regional variations and no correlation with the depth of winter-time mixing. The importance of shallow respiration for POC sequestration shows more distinct regional patterns. While in tropical and subtropical waters this process is negligible, up to about 61% of export production is remineralized in high latitude waters in the North Atlantic. The strong link between winter mixed layer depth and the relative effectiveness of the biological pump (\( \text{eff} = J_{\text{eff}}/J_{\text{POC}} \)) is summarized in Figure 8. About 92% of the variation of \( J_{\text{eff}}/J_{\text{POC}} \) can be explained by the depth of the winter mixed layer.

[52] With an anticipated change of ocean stratification due to global warming and the intensification of the water cycle [Manabe et al., 1991; Sarmiento et al., 1998], export production likely will change due to reduced nutrient fluxes to the surface ocean and a perhaps prolonged growth season [Bopp et al., 2001]. Much of this change is likely to become compensated by simultaneous changes of CO\(_2\) fluxes into the surface ocean [Maier-Reimer et al., 1996].

[53] The increase in the effectiveness of the biological pump (rising \( J_{\text{eff}}/J_{\text{POC}} \), Figure 8), however, constitutes a negative feedback process which can give rise to an imbalance in ocean carbon fluxes on timescales of the response of ocean circulation and thereby can increase the flux of anthropogenic CO\(_2\) flux into the ocean. An example of such a transient imbalance is described in a climate model study by Joos et al. [1999]. There the feedback of the biological pump around year 2100 increases the oceanic CO\(_2\)-uptake by about 10–15%, and thereby compensates for about 30–40%
of the reduction in CO₂ uptake from the combined effect of changes in the ocean circulation and the surface ocean temperature (i.e., higher ocean pCO₂ due to higher surface ocean temperature and hence a reduced CO₂ uptake). Model predictions of the regional distribution of changes in the mixed layer depth [Bopp et al., 2001] would indicate that temperate and subarctic waters, but not the tropics, will be the sites which show the most significant changes in terms of effective carbon fluxes into the deep ocean.

Appendix A: Estimating the Surface Mixed Layer POC:PIC Ratio of Production From Climatological Data

[54] Operationally it is assumed that the fraction of new carbon production (NP) which is exported via particles can be estimated from the seasonal drawdown of nitrate by multiplying with an appropriate C:N ratio of particles (equation (A1)). A C:N ratio of 6.625 [Redfield et al., 1963; Körtzinger et al., 2001] is used in this study. The seasonal nitrate drawdown (ΔNO₃) is estimated as the difference between an estimate of the winter nitrate concentration (NO₃wml) and the local minimal nitrate concentration (min(NO₃(seasonal)) observed in the seasonal nitrate climatology [NODC World Ocean Atlas, 1998 (WOA98); Conkright et al., 1998]. The winter nitrate concentration is calculated by applying the integration method of Glover and Brewer [1988] to the annual winter nitrate climatology from WOA98 (equation (A2))

\[ \text{NP} = \frac{C}{N} \times \Delta \text{NO}_3 = \frac{C}{N} \times [\text{NO}_3_{\text{wml}} - \text{min}(\text{NO}_3(\text{seasonal}))] \]  

(A1)

\[ \text{NO}_3(\text{W}) = \frac{1}{Z_{\text{wml}}} \times \int_{0}^{Z_{\text{wml}}} \text{NO}_3(\text{annual}) \, \text{d}z \]  

(A2)

[55] The production and export of CaCO₃ in the mixed layer, \( P_{\text{CaCO}_3} \), is estimated analogue to the approach of Robertson et al. [1994] from the seasonal change in potential alkalinity (equation (A3)). Here ΔNTA is the seasonal difference of the salinity normalized total alkalinity (NTA). NTA is estimated from regionally variable empirical relationships with temperature [Millero et al., 1998]. Monthly mean temperature values for the calculation of monthly NTA data fields are taken from WOA98 [Antonov et al., 1998]

\[ P_{\text{CaCO}_3} = \frac{[\Delta \text{NTA} + \Delta \text{NO}_3]}{2} \]  

(A3)

[56] Thereafter the mixed layer POC:PIC ratio of production, \( \text{RR}_{\text{mixed-layer}} \), is estimated as the ratio of NP/\( P_{\text{CaCO}_3} \).

[57] In the subtropics and tropics the seasonal approach cannot be used since seasonal gradients of alkalinity and nitrate vanish. As a substitute for the subtropical bands (10°S–35°N, 10°S–35°S) I estimate RRexp from mass flux and organic carbon flux data from drifting particle interceptor trap measurements (150 m depth) at the BATS [Lohrenz et al., 1992]. Since particle flux measurements at BATS do not include measurements of CaCO₃ fluxes, \( RR_{\text{exp}} \) is calculated assuming that the dry weight mass flux consists of organic matter (2.3POC), CaCO₃, opal, and a residual terrigenous component [Jickells et al., 1998], and that the opal to CaCO₃ ratio and the fraction of terrigenous matter can be extrapolated from deep trap observations from the close-by OFP-site. The long-term mean of the fraction of terrigenous matter in 3200 m at the OFP-site close to BATS is 18.7% [Conte et al., 2001] and the opal to CaCO₃ weight ratio during the early 1990s was 0.18 [Deuser et al., 1995]. Using the z-exponent of organic carbon flux decreases with depth from Koeve (submitted manuscript, 2001), the z-exponent of the CaCO₃ exponential flux function (this study) and a temperature dependent decay rate of opal [Gnanadesikan, 1999], a mean terrigenous fraction, and a mean [opal/CaCO₃]weight ratio in 150 m of 3% and 0.26, respectively, are estimated. Mass dry weight and POC data (available from the home page of the BATS Program at http://www.bbsr.edu) from traps deployed at 150 m between December 1988 and December 1997 (n = 106) are used. The median POC:PIC ratio in 150 m at BATS calculated from these data is 4.5, the time integrated annual mean POC:PIC ratio is 4.1. The latter value is used as an estimate of \( RR_{\text{exp}} \) in the subtropical band (10°S–35°S); in the tropics (10°S–10°N) an export flux ratio of 4.37 is used (Figure 2).

[58] Acknowledgments. Many people contributed to collecting the original data which form the basis of this study. Special thanks go to Klaus Kremling and Avan N. Antia, both at IM-Kiel, for their effort in guiding the German JGOFS particle flux synthesis group. The authors wish to acknowledge the use of the Ferret program for data analysis in this paper. Ferret (http://www.ferret.noaa.gov) is a product of NOAA’s Pacific Marine Environmental Laboratory. This work was supported by the German JGOFS synthesis program via a grant of the German BMBI-F to G. Wefer and W. Koeve. Fruitful discussion with A. Oschlies and P. Kähler, both at IM-Kiel, and helpful comments by two anonymous reviewers helped to clarify the ideas and the presentation.

References


Balch, W. M., K. A. Kilpatrick, P. Holligan, D. Harbour, and E. Fernandez, The 1991 coccolithophore bloom in the central North Atlantic, 2 Reftating...


---

W. Koeve, Zentrum für marine Umweltwissenschaften, (Marum), Fachbereich Geowissenschaften (FBS), Universität Bremen, Postfach 303440, D-28334 Bremen, Germany. (w.koeve@web.de; w.koeve@surfeu.de)