Magnitude of excess carbon sequestration into the deep ocean and the possible role of TEP

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ABSTRACT: It has been suggested that TEP (transparent exopolymer particles) may provide a vehicle for non-Redfield excess carbon export into the deep sea. Here, the hypothesis that organic carbon produced in excess of the Redfield C:N ratio is sequestered in the deep ocean is tested by a model-based mass balance approach. The model looks for a value of the C:N ratio of export production which is consistent with current knowledge of the rapid change of organic matter fluxes over depth, the remineralisation C:N ratio in the interior of the ocean, and the observed C:N ratios of sinking particles in the deep sea and at the sea floor. It is estimated that the contribution of excess carbon export into the deep ocean is equivalent to 3 to 5.6% (medians, depending on model assumptions; overall range: –16 to 21%) of the conventional Redfield biological pump (C:N = 6.6). Elevated C:N ratios of sinking particles in the deep ocean of 9 to 23, and their increase with depth, can be explained by C:N ratios of export production being only slightly larger than the vertically integrated C:N ratio of remineralisation in the interior of the ocean. The basin scale effect of this preferential nitrogen remineralisation, within the seasonal thermocline, on carbon sequestration is 1 order of magnitude lower compared with Redfield equivalent remineralisation or CaCO₃ sequestration. The often observed increase in the C:N ratio of sinking particles with depth does not require that the remineralisation C:N ratio increases with depth, but can also arise under conditions of constant C:N remineralisation ratios. It is concluded that only a small fraction of carbon overconsumption in the surface ocean is sequestered into the deep ocean. The majority appears to be remineralised in the upper twilight zone.

KEY WORDS: Carbon flux · Net community production · C:N ratio · Redfield · TEP

INTRODUCTION

The biological pump effects a net flux of organic carbon produced at the ocean's surface into its interior. This flux sustains a vertical CO₂ gradient, which contributes to the control of the mean state of the atmospheric pCO₂ in the unperturbed climate system. Conventional understanding has been that fluxes of carbon are largely related to fluxes of the major plant limiting nutrients (nitrate and phosphate) in a tight ratio. This view was based on the observation that the elemental ratios of remineralisation of biogenic matter in the deep ocean and those of plankton particles in the surface ocean are both similar to each other and relatively constant (Redfield et al. 1963).

This constant-stoichiometry approach has been frequently challenged. In particular, evidence has accumulated that the C:N ratio of net community production (NCP = gross primary production – community respiration) varies and is often higher than the Redfield C:N ratio of 6.6. NCP C:N ratios of up to 10 to 15 have been observed (Sambrotto et al. 1993, Körtzinger et al. 2001a). Biogenic net CO₂-uptake in excess of the Redfield equivalent of nitrate-based new production has been referred to as carbon overconsumption (Toggweiler 1993). Mechanistically, carbon overconsump-
tion can be related to the production and temporal accumulation of dissolved organic matter (DOM) low in nitrogen (Williams 1995, Kähler & Koeve 2001) or the preferential remineralisation and re-use of nitrogen in the euphotic zone (Thomas et al. 1999, Anderson & Pondaven 2001). Alternatively, unaccounted-for nitrogen sources (N$_2$-fixation, dissolved organic nitrogen) may falsely indicate excess uptake of carbon over Redfield equivalents of nitrogen uptake. Both N$_2$-fixation and the use of remotely produced (Williams & Follow 1998) dissolved organic nitrogen as nitrogen sources appear to be particularly important at low latitudes (Gruber & Sarmiento 1997, Hansell et al. 2004).

Most important for the understanding of the oceanic carbon cycle, however, is whether and how this excess carbon uptake is linked to carbon fluxes into the deep ocean. Only the fraction of carbon which escapes remineralisation in the seasonally mixed layer and is sequestered into waters which are cut off from contact with the atmosphere on time scales of more than 1 yr contributes to the surface to deep CO$_2$ gradient (Riebesell & Wolf-Gladrow 1992, Antia et al. 2001, Koeve 2002). At first glance, high C:N ratios of new and export production are difficult to reconcile with C:N ratios of remineralisation in the interior of the ocean, which largely conform with the Redfield ratio (Anderson & Sarmiento 1994). This apparent discrepancy may be resolved if carbon is preferentially, that is in excess of the already elevated C:N ratio of export production, exported to the deep ocean or to the sea floor. The increase of the C:N ratio in sinking particles over depth (e.g. Martin et al. 1987, Honjo & Manganini 1993, Conte et al. 2001) and elevated C:N ratios of freshly deposited detritus at the sea floor (Billett et al. 1983, Smith et al. 1998) may be quoted in support of this hypothesis.

Export production comprises the export of particulate and dissolved organic carbon, but only the seasonal DOM pool has been found to have an elevated C:N ratio (Williams 1995) and can, thereby, foster the export of excess carbon. However, the impact of DOM on long-term or deep carbon sequestration is restricted to regions of subduction of water masses, e.g. during thermocline ventilation (Doval & Hansell 2000). It is only particles which can effect diapycnal transport, thus effectively linking the surface ocean and the deep sea. Bulk particulate organic matter (POM) in the surface ocean, however, has C:N ratios close to the Redfield ratio (Copin-Montégut & Copin-Montégut 1983), as has marine snow (Allirdedge 1998, A. Allirdedge pers. comm.).

Recently, Engel & Passow (2001) and Engel (2002) have proposed that TEP (transparent exopolymer particles, Allirdedge et al. 1993) may provide a vehicle by which DOM from excess carbon uptake in the euphotic zone interacts with sinking particles and hence, contribute to particle export. TEP, which probably form from polysaccharides released by phytoplankton and bacteria, are very surface-active and easily coagulate among themselves and with other particles (Allirdedge et al. 1993). In addition to this mechanistic importance for particulate carbon fluxes in the ocean (Boyd & Stevens 2002, Passow 2002, 2004), it has been suggested that TEP, which have a high C:N ratio (Engel & Passow 2001, Mari et al. 2001), may lead to a preferred export of carbon from the surface ocean (Engel & Passow 2001). This process could provide a mechanism by which the total amount of carbon that is fixed in the ocean by phytoplankton and transported to the deep sea via the biological pump could well be more than calculated using the Redfield ratio (Engel 2002). Furthermore, a potential increase in TEP formation, resulting from rising pCO$_2$ (Engel 2002), may function as a feedback mechanism, counteracting rising atmospheric pCO$_2$ (Passow 2002), if it becomes sequestered into the deep ocean.

Testing this hypothesis is not straightforward, since our knowledge of upper ocean carbon and nitrogen export fluxes is very much limited by methodological problems of particle interceptor traps (Zeitzschel et al. 1978), which are most prominent in shallow to intermediate waters (Gardner 2000). Deep water particle flux data are believed to be more reliable since hydrodynamic as well as biological biases (trapping efficiency, swimmer artifacts) diminish. On average, deep ocean particle flux carbon to nitrogen ratios increase with depth by about 0.2 ± 0.1 units per 1000 m (Schneider et al. 2003), with average values of 7.9 to 8.2 (±0.5) at a depth of 4000 m. Near the sea bed, sediment trap data as well as observations of freshly deposited phytodetritus at the sea floor show C:N ratios ranging up to 10 or even 20 (Smith et al. 1998, Lampitt et al. 2001, Beaulieu 2002). However, deep ocean C:N ratios of sinking organic matter do not provide direct information on the export or sequestration C:N ratio. This is illustrated here with an example from laboratory-based batch (closed system) remineralisation experiments of non-axenic Chryptomonas sp. and Gymnodinium cultures (Verity et al. 2000). In both experiments, POM$_{C:N}$ ratios increased over time, with maximum values of 14 to 24 (Fig. 1), though the initial (‘export’) POM$_{C:N}$ was close to Redfield (5.7 and 6.3, respectively). The explanation of the increase is that the C:N remineralisation ratios (r$_{C:N}$, Fig. 1), which are computed here from the time course change of POC and PON, were just a little lower (initially 4.8 and 5.6, respectively) than the POM$_{C:N}$ at the beginning of the experiment. A little preferential N-remineralisation (Lee & Cronin 1984, Wakeham et al. 1984) during the period of rapid organic matter decay gives rise to a
strong relative carbon accumulation in the residual POM towards the end of the experiment.

The hypothesis of this paper is that the magnitude and distribution of deep ocean C:N flux ratios \(J_{\text{C:N}}\) can in a very similar way be explained by a small difference between initial C:N export ratios \(e_{\text{C:N}}\) and the \(r_{\text{C:N}}\) ratio in the water column, and hence are not an indication of significant excess carbon export into the deep ocean, for instance by means of TEP, or by other processes such as the stripping of ‘old’ (carbon-14 age) DOM from the water column by sinking particles (Druffel et al. 1996). In order to test this hypothesis, and considering the lack of reliable \(J_{\text{C}}\) and \(J_{\text{N}}\) data from the upper ocean, I chose to use a simple model-based mass balance approach. The model basically looks for a value of the C:N ratio of export production which is in agreement with our current knowledge of the rapid change of organic matter fluxes over depth, the remineralisation C:N ratio in the interior of the ocean, and the observed C:N ratios of sinking particles in the deep sea and at the sea floor. In the following section, I introduce the model and its basic assumptions. After presenting the results, I discuss the chosen parameters of the model in more detail against the background of our current understanding of flux and remineralisation of organic matter in the interior of the ocean. Finally, implications concerning the possible importance of TEP for excess carbon export and the coupling of surface new production and deep ocean fluxes are discussed.

METHODS

General outline. The 2 models used in this study apply the finding that the particle flux profile in the deep ocean can be described by a power-law function of export production due to particles \(E_P\) and depth \(Z\) following Eq. (1) (equations are given in Table 1; for acronyms, see Table 2) (Martin et al. 1987), where \(Z_{\text{EP}}\) is the reference depth of export production, formally the depth of the euphotic zone, and \(b\) is a vertical scaling parameter. This function, and similar power-law equa-

Table 1. Model equations. Analytical solutions (Eqs. 6, 7, 11 & 12) are valid if \(r_{\text{C:N}}\) is constant over depth. \(R\) is integrated between \(Z_{\text{EP}}\) and \(Z\). See Table 2 for definitions of parameters

\[
J_{\text{(Z)}} = E_P \times \left( \frac{Z}{Z_{\text{EP}}} \right)^b
\]

\[
E_P_{\text{C}} = E_P \times e_{\text{C:N}}
\]

Model 1

\[
J_{\text{(C,Z)}} = e_{\text{C:N}} \times E_P \times \left( \frac{Z}{Z_{\text{EP}}} \right)^b e\]

\[
R_{\text{(C,Z)}} = b \times e_{\text{C:N}} \times E_P \times Z^{b-1} / Z_{\text{EP}}^b
\]

\[
R_{\text{(N,Z)}} = \frac{R_{\text{(C,Z)}}}{r_{\text{C:N}}}
\]

\[
J_{\text{(N,Z)}} = E_P N - \int R_{\text{(N,Z)}} \times dZ
\]

\[
e_{\text{C:N}}^{\text{req}} = \left[ \left( \frac{Z}{Z_{\text{EP}}} \right)^b e \times \left( \frac{1}{J_{\text{C:N}}} - \frac{1}{r_{\text{C:N}}} \right) + \frac{1}{r_{\text{C:N}}} \right]^{-1}
\]

Model 2

\[
J_{\text{(N,Z)}} = E_P N \times \left( \frac{Z}{Z_{\text{EP}}} \right)^{b_n}
\]

\[
R_{\text{(N,Z)}} = b \times E_P N \times \frac{Z_{\text{EP}}^{b_n-1}}{Z_{\text{EP}}^{b_n}}
\]

\[
R_{\text{(C,Z)}} = R_{\text{(N,Z)}} \times r_{\text{C:N}}
\]

\[
J_{\text{(C,Z)}} = E_P C - \int R_{\text{(C,Z)}} \times dZ
\]

\[
e_{\text{C:N}}^{\text{req}} = J_{\text{C:N}} \times \left( \frac{Z}{Z_{\text{EP}}} \right)^{b_n} + r_{\text{C:N}} \times \left[ 1 - \left( \frac{Z}{Z_{\text{EP}}} \right)^{b_n} \right]
\]
tions relating deep ocean fluxes with surface ocean primary production (Suess 1980, Betzer et al. 1984, Antia et al. 2001, Koeve 2002), reflect the dramatic decrease of organic particle flux over depth. The Martin-power-law empirical function has been successfully used to describe the vertical distribution of particle flux at a set of 17 open ocean stations for which depth-resolving flux observations were available (Berelson 2001). The regional range of $b$-values from Berelson (2001) guided the choice of the $b$-value range used in this paper. The Martin-power-law function, again with regionally variable $b$-values, is also in excellent agreement with the global 3-dimensional distribution of nutrients, total CO$_2$ and oxygen (Schlitzer 2000, 2002).

Particle export into the deep ocean, as measured by particle interceptor traps, is mainly maintained by large fast sinking particles (Walsh & Gardner 1992), which descend with sinking speeds in the order of 10 to 100 m d$^{-1}$ towards the sea floor. Such marine snow particles are sites of intensive microbial activity (Smith et al. 1992, Ploug & Grossart 2000), with turnover times of organic carbon due to respiration of only 8 to 9 d (Ploug et al. 1999). The Martin-curve type of power-law functions used in this study are assumed to integrate over a spectrum of particles with different sinking speeds and decay rates (Boyd & Stevens 2002). Ocean flux data which are fitted into Martin-type curves are most credible in the deep ocean, where notorious problems of shallow traps, such as trapping efficiency (Buesseler 1991), contamination by swimming particles, is mainly maintained by large fast sinking particles (Walsh & Gardner 1992), which descend with sinking speeds in the order of 10 to 100 m d$^{-1}$ towards the sea floor. Such marine snow particles are sites of intensive microbial activity (Smith et al. 1992, Ploug & Grossart 2000), with turnover times of organic carbon due to respiration of only 8 to 9 d (Ploug et al. 1999). The Martin-curve type of power-law functions used in this study are assumed to integrate over a spectrum of particles with different sinking speeds and decay rates (Boyd & Stevens 2002). Ocean flux data which are fitted into Martin-type curves are most credible in the deep ocean, where notorious problems of shallow traps, such as trapping efficiency (Buesseler 1991), contamination by swimming particles, is mainly maintained by large fast sinking particles (Walsh & Gardner 1992), which descend with sinking speeds in the order of 10 to 100 m d$^{-1}$ towards the sea floor. Such marine snow particles are sites of intensive microbial activity (Smith et al. 1992, Ploug & Grossart 2000), with turnover times of organic carbon due to respiration of only 8 to 9 d (Ploug et al. 1999). The Martin-curve type of power-law functions used in this study are assumed to integrate over a spectrum of particles with different sinking speeds and decay rates (Boyd & Stevens 2002).

In the model, remineralisation of organic matter is computed from the change of organic matter flux over depth. Since the trapping efficiency of sediment traps varies with depth (Scholten et al. 2001, Yu et al. 2001), the ratio of carbon and nitrogen remineralisation and its possible change over depth cannot be assessed if only based on shallow trap data. This problem is most pronounced in the upper 500 to 1000 m of the ocean. As a substitute, the C:N ratio of remineralisation ($r_{\text{C:N}}$) can be estimated from the distribution of the remineralisation products, in this case nitrate, total CO$_2$ and oxygen, in the interior of the ocean. This approach looks at changes of these substances along well-chosen isopycnals or neutral surfaces (McDougall 1987) and includes corrections for isopycnal mixing of different end members (Takahashi et al. 1985), diapycnal mixing (Anderson & Sarmiento 1994) and the time varying invasion of anthropogenic CO$_2$ (Körtzinger et al. 2001b). By definition, remineralisation between the depth of the euphotic zone and the winter mixed layer cannot be estimated by this method, and it is assumed here that remineralisation ratios estimated from the permanent thermocline also apply to the seasonal thermocline. To reflect both regional and inter-study differences in $r_{\text{C:N}}$ and uncertainties of ratios within the seasonal thermocline, a range of $r_{\text{C:N}}$ values is applied (Table 2). This range is in good agreement with $r_{\text{C:N}}$ ratios estimated from decomposition experiments of phytoplankton-derived POM (Verity et al. 2000, Fig. 1).

The overall approach of this study was to estimate the value of the C:N ratio of export production ($r_{\text{C:N}}$) of particles which is required to sustain a given observed value of the C:N ratio of sinking particles ($J_{\text{C:N}}$) in the deep ocean under the constraint of specified values for $b$ and $r_{\text{C:N}}$. This particular value of $r_{\text{C:N}}$ is referred to as $r_{\text{C:N}}^\text{req}$ hereinafter. I tested how $r_{\text{C:N}}^\text{req}$ changes depending on the choice of deep ocean $J_{\text{C:N}}$, $b$ and $r_{\text{C:N}}$. The difference between $r_{\text{C:N}}^\text{req}$ and the classical Redfield C:N ratio (6.6) is used as a measure of excess car-

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Remark</th>
<th>Standard value (range)</th>
</tr>
</thead>
<tbody>
<tr>
<td>$Z_{\text{EP}}$</td>
<td>Depth of the productive surface layer (euphotic zone)</td>
<td>100 m</td>
</tr>
<tr>
<td>$EP_x$ ($EP_z$)</td>
<td>Export production of nitrogen (carbon) at depth $Z$</td>
<td>0.5 mol m$^{-2}$ yr$^{-1}$</td>
</tr>
<tr>
<td>$J_{\text{RZ}}$ ($J_{\text{CZ}}$)</td>
<td>Remineralisation of nitrogen (carbon) at depth $Z$</td>
<td></td>
</tr>
<tr>
<td>$b$</td>
<td>$Z$-exponent of the J-EP-Z curve (Eq. 1)</td>
<td>$-0.68$ ($-0.6$ to $-1.3$)</td>
</tr>
<tr>
<td>$r_{\text{C:N}}$</td>
<td>C:N ratio of remineralisation in the interior of the ocean</td>
<td>6.6 (5.6 to 7.6)*</td>
</tr>
<tr>
<td>$e_{\text{C:N}}$</td>
<td>C:N ratio of export production</td>
<td>5 to 15</td>
</tr>
<tr>
<td>$e_{\text{C:N}}^\text{req}$</td>
<td>Value of $e_{\text{C:N}}$ required to sustain observed values of $J_{\text{C:N}}$ for a given model and parameter set ($r_{\text{C:N}}$, $b$)</td>
<td></td>
</tr>
<tr>
<td>$J_{\text{C:N}}$</td>
<td>C:N ratio of sinking particles</td>
<td></td>
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*Also experiments with depth variable $r_{\text{C:N}}$ were performed

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bon export and is reported as a percentage of the Redfield equivalent carbon flux. It is computed as excess = \((e_{CN}^{\text{req}} - 6.6)/6.6 \times 100\).

There is no explicit model domain for this study. The 1-dimensional model may be applied everywhere in the open ocean. Parameters, initial conditions and constraints from observations, however, are biased towards the Atlantic Ocean. Remineralisation C:N ratios from isopycnal analysis average over large horizontal distances, time scales of years and, close to their outcrop and ventilation regions, also over depth. In a similar way, \(b\)-values are usually computed for annual flux data and, hence, reflect the annually averaged decay of organic matter over depth and the observed range of deep ocean \(J_{CN}\) (see ‘Results’) is used. Hence, this study intends to, and can only, look at the weighted average of excess carbon export into the deep ocean. There may be short periods, or specific regions, which are characterised by fluxes of C:N ratios in excess of what is calculated here. For excess carbon export to be significant and, via TEP, to provide a possible feedback process compensating for increasing atmospheric CO2 concentrations (Passow 2002), however, it must also be visible in average calculations of this type.

Model details. Standard runs of both models are used to explore the overall response of the model system to elevated C:N ratios of export production of particles. For the \(Z\)-exponent of organic matter decay, I use a value of \(b = -0.68\), which is the mean \(Z\)-exponent of particulate carbon flux estimated for the Atlantic Ocean by Antia et al. (2001), and \(Z_{DP} = 100\ m\). The model is initiated with a (particulate) nitrogen export production (\(EP_{N}\)) of 0.5 mol N m\(^{-2}\) yr\(^{-1}\), which is a typical value for the temperate and subarctic North Atlantic (Koeve 2001). Carbon export production of particles (\(EP_{C}\)) is computed as \(EP_{N} \times e_{CN}\), where \(e_{CN}\) is the C:N ratio of export production. Model runs are carried out with values of \(e_{CN}\) ranging from 5 to 15, reflecting the seasonal range of the C:N ratio of NCP, observed by Körtzinger et al. (2001a) in mesotrophic waters of the North Atlantic.

In Model 1, the carbon flux (\(J_{C}\)) profile is computed by applying the \(J\)-\(EP\)-\(Z\) function (Table 1, Eq. 1) to values of \(EP_{C}\) computed for any given value of \(e_{CN}\). Carbon remineralisation (\(R_{C}\)) is deduced from the first derivative of the \(J\)-\(EP\)-\(Z\) curve, and \(R_{C}\) nitrogen remineralisation, is computed as \(R_{N} = R_{C}/e_{CN}\). The standard value of \(R_{CN}\) is set to 6.6. Finally, the nitrogen flux (\(J_{N}\)) profile is estimated from the cumulative integral of \(R_{N}\) and \(EP_{N}\), and the resulting C:N ratio of particle flux is computed from \(J_{CN} = J_{C}/J_{N}\). Model 1 assumes that conventional methods for particulate organic carbon (POC) (Ehrhard & Koeve 1999) measure the total amount of particulate organic carbon sampled with sediment traps and, hence, that estimates of the \(Z\)-exponent \(b\) include, for example, the contribution of TEP-associated carbon.

Model 2 reflects the possibility that TEP-associated carbon is only partly included in standard POC analysis (Passow & Allerdge 1995, Engel & Passow 2001, see also ‘Discussion’ ‘Uncertainties of observed C:N flux ratios’) of sinking particles. It only applies the \(J\)-\(EP\)-\(Z\) function to the Redfield-associated carbon flux and treats any excess carbon flux independently of it. In fact, since the C:N remineralisation ratio is always close to the Redfield value, this method largely preserves excess carbon during its transit to depth. Technically, this model computes the nitrogen flux (\(J_{N}\)) profile from the \(J\)-\(EP\)-\(Z\) profile and estimates \(R_{N}\) from the first derivative of \(J_{CN}\). \(R_{C}\) is estimated from \(R_{C} = R_{N} \times e_{CN}\), and \(J_{C}\) is computed from integration of \(R_{C}\) over \(Z\) and from \(EP_{C}\), which is estimated as above. Model equations, including analytical solutions for \(i_{CN}\) = constant, are given in Table 1.

For both models, a series of sensitivity computations was performed using ranges of \(e_{CN}, i_{CN}\) and \(b\). For example, runs using constant \(i_{CN}\) values between 5.5 and 7.5 (for literature values see ‘Discussion: C:N ratios of remineralisation in the interior of the ocean’) are carried out, as well as experiments with depth variable \(i_{CN}\) values, computed from the function \(i_{CN} = 6.2 (Z/Z_{DP})^{0.13}\), adopted from Martin et al. (1987) and a function \(i_{CN} = 4.9786[\exp(0.000212Z)]\) fitted to data presented by Shaffer (1996). Values for \(b\) are probably not constant over the ocean. Analyzing data from 17 field experiments, Berelson (2001) found \(b\) to range between −0.6 and −1.28, and a positive correlation of \(−b\) with \(EP_{C}\). A range of −0.6 to −1.3 is adopted here for estimating the dependence of \(e_{CN}^{\text{req}}\) and \(b\).

A realistic description of POC fluxes including carbon associated with TEP will probably be somewhere between the 2 models used in this study, since some TEP will always be included in the standard analysis of POC.

RESULTS

The depth distribution of the C:N ratio of particle flux (\(J_{CN}\)) for standard experiments (\(i_{CN} = 6.6\)) with Models 1 and 2 is shown in Figs. 2 & 3, respectively. In both models, the modeled C:N flux ratios increase with depth if \(e_{CN}\) is larger than \(i_{CN}\). Even for marginally elevated \(e_{CN}\) ratios, between 6.6 and 7, Model 1 predicts deep water (4000 m) C:N ratios of sinking particles of 6.6 to 22. For higher \(e_{CN}\) ratios (8 to 15), the predicted C:N ratios of sinking particles increase dramatically already in shallow water, particularly due to the rapid nitrogen depletion in sinking particles computed by the model (Fig. 2 insert). Model 2 predicts a more moderate increase in C:N flux ratios since,
How does this compare with observed C:N flux ratios in the deep ocean? Observations with particle interceptor traps usually revealed higher than Redfield C:N ratios and also a slight trend (increase) with depth (Schneider et al. 2003). Mean annual flux ratios from a 9 yr time series observation at the OFP/BATS (Ocean Flux Program, Bermuda Atlantic Time Series, Conte et al. 2001) station increase from 7.8 ± 1.4 (500 m) to 9.2 ± 1.5 (1500 m) and 9.7 ± 1.4 (3200 m). Similar increases are seen at other sites in the North Atlantic. Honjo & Manganini (1993), for example, report annual C:N ratios of 7.8 to 8.3 between 2000 and 3800 m at 47°N, 20° W. Data compiled by Antia et al. (2001) from the Atlantic Ocean (n = 45) show a mean annual C:N flux ratio of 8.8 ± 1.7 and a small increase with depth (not shown). For the standard parameters \( b = -0.68, r_{C:N} = 6.6 \), the observed deep water C:N flux ratios of Honjo & Manganini (1993), Conte et al. (2001) and Martin et al. (1987) (11.5 at 4000 m) are reproduced by Model 1 for \( e_{C:N}^{req} \) values between 6.7 and 6.85 (Fig. 4).

For Model 2, the solutions for \( e_{C:N}^{req} \) range from 6.8 to 7.0. The respective excess carbon fraction to carbon export production ranges between 1.8 and 3.6% (Model 1), and 2.4 and 6.1% (Model 2). Using lower \( b \)-values in the models requires lower \( e_{C:N} \) input values to compute observed deep sea C:N flux ratios (Fig. 5). If a value of \( b = -1 \) (Koeve 2002) is assumed, the excess carbon export required to match the deep water C:N flux ratios is between 1 and 2% of the Redfield equivalent carbon export (open symbols in Fig. 4). Overall, the effect of \( b \) on \( e_{C:N}^{req} \) is moderate (Fig. 5).

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able Z-exponents are small compared to differences related to uncertainties of $r_{\text{C:N}}$. A depth-dependent distribution of $r_{\text{C:N}}$ has been suggested among others by Martin et al. (1987) and Shaffer (1996). Using such depth profiles of $r_{\text{C:N}}$ here yields $e_{\text{C:N req}}$ of 7.18 and of 5.8 to 6.2 for the Martin et al. (1987) and Shaffer (1996) equations, respectively, to match a deep water $J_{\text{C:N}}$ of 11.5 at 4000 m (Fig. 6).

Finally, an extreme assumption is made to reflect the possibility that TEP-associated carbon is severely underestimated in deep traps. Here, I assume that the C:N flux ratio at 4000 m is 23 (2 times the value measured by Martin et al. 1987 in their deep traps). Depending on the model assumptions, between 3 (Model 1, $b = -1$) and 21% (Model 2, $b = -0.68$) excess carbon export is required to yield such high deep water C:N ratios ($r_{\text{C:N}} = 6.6$, Fig. 4). Note the strong difference between the functional response of $e_{\text{C:N req}}$ over $J_{\text{C:N}}$ for Models 1 and 2.

### DISCUSSION

Results from this study revealed that an excess particulate carbon export from the euphotic zone, for example in the form of TEP, of only 1 to 6.1% of the Redfield equivalent carbon export is required to reproduce the observed $J_{\text{C:N}}$ of sinking particles in the deep ocean (8.3 to 11.5), when a remineralisation ratio of 6.6 (Redfield et al. 1963) is assumed (Fig. 4). A higher excess export demands that either (1) the deep ocean C:N flux ratio or (2) the Z-exponent of particle flux have been severely underestimated, or that (3) the C:N remineralisation ratio due to particles is much higher. In the following sections I discuss these 3 aspects separately.

### Uncertainties of observed C:N flux ratios

The estimation of the C:N flux ratio of particles sampled with sediment traps is prone to a number of uncertainties. For example, zooplankton, which is known to actively enter traps (‘swimmers’), may add organic matter with a low C:N ratio (5 to 6) and dilute the signal of sinking POM. Picking swimmers prior to POM measurements, a standard procedure of trap sample handling, would partly prevent this artifact; however, elevated concentrations of DOM have been found to accumulate in sample cups of sediment traps (Kähler & Bauerfeind 2001), originating from the dissolution of sinking particles, but likewise from swimmers. Kähler & Bauerfeind (2001) found an apparent C:N ratio of sinking particles, estimated from POC and PON analysis of trap samples, of 9.6 (annual integral) in a shallow (135 m) trap de-
ployed in the Greenland Sea. Significant concentrations of DOM with a low C:N ratio were measured in the supernatant of the trap cup. Including the DOC and DON fractions, which Kähler & Bauerfeind (2001) could assign to the passive particle flux, yielded a corrected C:N ratio of 3.8 (annual integral). Recognising that lipids may occasionally be completely lost from sediment traps (Kähler & Bauerfeind 2001) and assuming a lipid contribution to total plankton carbon of 22% (computed after Hedges et al. 2002), requires an upward correction and yields an annually integrated C:N ratio of (<)4.8. Obviously, the DOM corrected estimate of the C:N ratio of the passive flux of organic carbon entering the trap of 3.8 to 4.8 is very different from the original POC:PON based estimate of 9.6.

If the results of the study of Kähler & Bauerfeind (2001) apply in general to shallow traps, it causes doubt concerning elevated C:N ratios frequently reported from shallow sediment traps (e.g. Anderson & Pondaven 2003) and as summarised in Schneider (2003) and Schneider et al. (2003). Fortunately, both the swimmer and the DOM problem show a depth-dependence (e.g. Michaels et al. 1990, P. Kähler & IFM-Kiel pers. comm.), indicating that deep traps are much less affected (Honjo et al. 1995). Similarly, deep traps are also much less prone to hydrodynamic biases and trapping efficiency problems (Scholten et al. 2001, Yu et al. 2001) which add to the uncertainties of C:N flux ratios in shallow traps. Hence, I restricted myself to the use of data from deep traps as a constraint in the model experiments.

A problem, which may call the reliability of C:N flux ratios from deep traps into question, is whether TEP-associated carbon is fully included in the POC measurement of material from sediment traps. POM is conventionally filtered on glass fiber filters (Ehrhard & Koeve 1999), ‘depth filters’ consisting of a matrix of glass fiber structures, which have a nominal pore size of 0.7 μm (Whatman GF/F filters). Though the effective pore size may decrease over filtration time, it is possible that small particles (e.g. bacteria, viruses, TEP) are not retained quantitatively and hence, are not fully included in POC analysis. Using size-fractionated filtration, Passow & Alldredge (1995) observed that about 60 to 70% of TEP were lost when 0.6 μm membrane filters were used instead of 0.2 μm membrane filters. Pre-combusted glass fiber filters, however, may behave differently. Morán et al. (1999), for example, showed that glass fiber filters adsorb significant amounts of 14C-labeled freshly produced dissolved organic matter. I speculate that this may also apply to surface-reactive TEP and that TEP is more efficiently retained on GF/F filters compared to membrane filters of similar pore size.

As yet, little is known about the flux of TEP and TEP-associated carbon into the deep sea. A study from the Santa Barbara Channel off California suggests that the ratio of deep (500 m) flux to surface standing stock in the surface layer is similar for POC and TEP (about 1 to 2% d⁻¹ each) (Passow et al. 2001), indicating that Model 1 is more applicable. During that 2 yr study, the integrated overall TEP to POC ratio (roughly estimated from their Fig. 3) was about 0.5 mg Xanthan:mg POC. Combining these data with the carbon to TEP conversion relationship found for TEP produced from diatom cultures (Engel & Passow 2001), yields a ratio of about 0.4 (TEP-carbon:POC). Using this first order estimate, one may conclude that TEP-carbon either made up about 40% of the observed POC flux (compliant with the assumptions of Model 1) or that the true POC flux and the C:N flux ratio were underestimated by about 30% (compliant with the assumptions of Model 2) in the study of Passow et al. (2001). Model computation assuming a deep ocean J_{C:N} of 23 (2 times the observed 4000 m value of Martin et al. 1987) is equivalent to assuming true carbon fluxes to be underestimated by 100% due to unrecognised TEP, which is well beyond the upper bound estimate (30% underestimation) based on Passow et al. (2001). Hence, 2 × J_{C:N} experiments are not included in the envelop of the likely range of excess carbon export (shaded patch in Fig. 6).

Given the problems of trapping efficiency, POM dissolution in traps, swimmers and TEP-carbon analytics discussed above, is there independent evidence for the C:N ratio of sinking particles in the deep ocean? Samples of ‘fluff’, detritus on the sea floor, may provide such evidence. Such material has been observed to arrive within a few weeks after the surface ocean spring bloom in the deep sea (Billett et al. 1983) and has been sampled by multicorers or from submarines. Beaulieu (2002) reviews fluff observations and the chemical composition of the material. C:N ratios range from 6 to 24, with most of the observations having C:N ratios of lower than 12. The high end ones, partially originating from thin surface layers of the sediment (Smith et al. 1998, their Table 4), probably include older material, which has already been reworked in the sediments and, therefore, has a high C:N ratio. This is supported by the carbon-14 age of fluff (Druffel et al. 1998) being much more similar to sediment carbon-14 ages than to those from suspended or sinking POC, or from surface ocean DIC.

**Effect of b on e_{C:N}**

A range of values of the Z-exponents of the $J$-$EP-Z$ curve from −0.6 to −1.3 was tested (Fig. 5). Low end values ($b = −1.28$), estimated by Berelson (2001) for spring bloom conditions in the temperate North Atlantic, require carbon export to be about 5% (Model 1)
to 7% (Model 2) lower compared with high end \( b \)-values (\( b = -0.6 \)) to sustain observed deep ocean \( J_{\text{C:N}} \) values. Berelson (2001) suggests that low \( b \)-values correspond to regimes of higher export production (\( EP_{\text{b}} \)) and it may be that the regional distribution of deep water \( J_{\text{C:N}} \) reflects the variability in \( b \). This, however, is difficult to test since estimates of \( r_{\text{C:N}} \) (see next section) usually average over large horizontal distances. One may speculate that oligotrophic regions have higher \( e_{\text{C:N}}^{\text{req}} \) than eutrophic regimes. The overall effect of the \( Z \)-exponent of the \( J_{\text{EP}} \)-\( Z \) curve on \( e_{\text{C:N}}^{\text{req}} \), however, is small compared with the effect of \( r_{\text{C:N}} \) (Figs. 5 & 6). The remaining discussion, therefore, focuses on a brief review of \( r_{\text{C:N}} \), its vertical distribution and the possible impact of DOM remineralisation on \( r_{\text{C:N}} \).

**C:N ratios of remineralisation in the interior of the ocean**

The elemental ratios of remineralisation in the interior of the ocean have been estimated repeatedly from the change of total \( \text{CO}_2 \), \( \text{NO}_3 \), \( \text{PO}_4 \) and \( \text{O}_2 \) along isopycnals or on neutral surfaces. This method requires that the remineralisation signal is separated from changes in the tracer distribution due to hydrodynamic processes such as isopycnal and diapycnal mixing, and from effects due to denitrification and the invasion of anthropogenic \( \text{CO}_2 \). Most estimates of the mean ocean \( C \):\( N \) remineralisation ratio range between 5.6 ± 0.5 (Takahashi et al. 1985) and 7.3 ± 1.3 (Anderson & Sarmiento 1994). Evidence for vertical variations of these ratios is controversial. Several authors found no significant trend over depth (Broecker et al. 1985, Takahashi et al. 1985), in particular, when effects due to denitrification (Anderson & Sarmiento 1994) or from the time history of the invasion of anthropogenic \( \text{CO}_2 \) (Körtzinger et al. 2001b) have been corrected for. Other studies, on the contrary, found an increase over depth from shallow water \( C \):\( N \) remineralisation ratios of 5.1 (100 m) to 9.4 (3000 m) (Shaffer 1996) and from 4.5 ± 0.3 (\( \rho < 1027.7 \) \text{kg m}^{-3} \)) to 11.0 ± 0.5 (\( \rho > 1027.7 \) \text{kg m}^{-3} \)) in the subarctic North Atlantic (Thomas 2002). A similar increase of \( C \):\( N \) remineralisation ratios with depth has been inferred from the VERTEX particle flux data set (Martin et al. 1987).

The range of \( r_{\text{C:N}} \) used in this study is in good agreement with \( r_{\text{C:N}} \) estimates based on decomposition experiments of phytoplankton-derived POM (Verity et al. 2000; Fig. 1). The mean \( r_{\text{C:N}} \) (±SD) from both experiments is 5.4 ± 1.6 (\( n = 6 \)) and the standard deviation reduces to 0.5, if one only uses data from the time period of rapid POM change over time (\( n = 4 \)). If these biological estimates of \( r_{\text{C:N}} \) reflect POM degradation in the ocean more accurately than those from isopycnal analysis (see discussion on effects of DOM remineralisation on \( r_{\text{C:N}} \) below), they clearly point towards low excess carbon export into the deep sea (Fig. 6).

The spread of \( r_{\text{C:N}} \) values has been implemented in the sensitivity computations of this study. Constant \( r_{\text{C:N}} \) values between 5.5 and 7.5 require \( e_{\text{C:N}}^{\text{req}} \) ratios of 5.76 and 7.72 to sustain a deep ocean \( C \):\( N \) flux ratio of 11.5. Hence, \( r_{\text{C:N}} \) shows up to be the most significant determinant of the \( e_{\text{C:N}}^{\text{req}} \) to deep \( C \):\( N \) flux relationship.

Moreover, the shallow and intermediate water \( r_{\text{C:N}} \) largely determines the \( e_{\text{C:N}}^{\text{req}} \) due to the predominance of shallow waters for remineralisation. This is evident from experiments with depth variable \( r_{\text{C:N}} \) values. Adopting the \( r_{\text{C:N}} \) vertical distribution found by Shaffer (1996) yields \( e_{\text{C:N}}^{\text{req}} \) of 5.8 (Model 1) and 6.2 (Model 2) to sustain the deep water \( C \):\( N \) flux ratio of 11.5 (\( b = -0.68 \)). Depth integrated \( r_{\text{C:N}} \) from these experiments are 5.4 and 5.5, respectively, and are very close to the \( r_{\text{C:N}} \) ratio of 5.1 at a depth of 100 m (Shaffer 1996). The same is seen from the experiment with the \( r_{\text{C:N}} \) distribution adopted from Martin et al. (1987). Here, the 100 m value is 6.2, the depth integrated \( r_{\text{C:N}} \) is 7.0, and \( e_{\text{C:N}}^{\text{req}} \) is 7.18. Elevated deep water values of \( r_{\text{C:N}} \) had little impact on \( e_{\text{C:N}}^{\text{req}} \).

So far, the \( r_{\text{C:N}} \) ratio was used irrespective of whether it is due to the remineralisation of particulate or dissolved matter. In the deep ocean, almost 100% of remineralisation is due to particles, since the change of \( \text{DOC} \) (Hansell & Carlson 1998) along the deep water path of the conveyor belt is very small compared with the concomitant increase in total \( \text{CO}_2 \). On shallow isopycnals, however, DOM may contribute significantly to carbon remineralisation. Doval & Hansell (2000), for example report that remineralisation of \( \text{DOC} \) may account for up to 50% of apparent oxygen utilisation on shallow isopycnals of the subtropical thermocline in the South Pacific. No concurrent DON measurements are available from that study and hence, little is known about the \( C \):\( N \) remineralisation ratio of DOM on these shallow isopycnals. If the latter remineralisation ratio is elevated similarly to the \( C \):\( N \) ratio of seasonally accumulating DOM in the North Atlantic (Williams 1995, Kähler & Koeve 2001), the overall upper ocean \( r_{\text{C:N}} \) ratios may overestimate the remineralisation ratios related to the remineralisation of sinking particles.

**Implications**

Carbon export in excess of the Redfield (\( C \):\( N = 6.6 \)) equivalent of nitrogen export production ranging from about -16 to about 21% have been estimated in this study to be sufficient to sustain observed elevated deep water \( C \):\( N \) ratios of sinking particles. Assuming
that a considerable amount of carbon, for example in the form of TEP or carbon-rich DOM from particle dissolution in sample cups, has been overlooked analytically in deep sediment traps, yields an additional excess carbon export of between 3 (Model 1) and 15% (Model 2). The median excess carbon export amounts to 3 (n = 16, experiments with 2 \times J_{C:N} excluded) to 5.6% (n = 22, all experiments). It is concluded that, in agreement with C:N remineralisation ratios below the winter mixed layer, only a moderate fraction of carbon overconsumption is exported into the deep ocean.

This small amount of excess carbon export, or more specifically a C:N ratio of export production of particles which is a little larger than the depth integrated C:N ratio of remineralisation, is sufficient to explain elevated C:N ratios of sinking particles in the deep ocean of 9 to 11.5 (and up to 23 for 2 \times J_{C:N} experiments) and its increase over depth. This holds irrespective of whether the remineralisation C:N ratio is constant over depth or not. It is the preservation of a subtle amount of extra carbon in the export from the euphotic zone which becomes progressively significant during transit to depth when the flux of particulate organic matter rapidly decreases with depth, and particulate organic carbon and nitrogen are remineralised in a ratio close to that proposed by Redfield et al. (1963).

Evans & Fasham (1993) showed that preferential remineralisation of nitrogen from sinking POM will cause a decrease in CO₂ concentration in the surface ocean. Hence, what matters is the C:N ratio at the reference depth of carbon sequestration, that is of carbon export on longer than annual time scales (Riebesell & Wolf-Gladrow 1992). This reference depth is highly variable in the ocean. In temperate and higher latitude waters, it is the maximum depth of winter mixing; in upwelling regions, the lower boundary of the upwelling source water may be taken as such (Koeve 2002). For a brief estimate of the effect of preferential nitrogen remineralisation on the basin scale of the sequestration of organic carbon, I estimate the ratio of \( J_{C,N}\text{sequ}/e_{C,N} \) from formulas given in Table 1 using average values for \( Z\text{seq} \) for biogeographical provinces of the Atlantic ocean (Koeve 2002) and median values of \( e_{C,N} \) (6.79), \( r_{C,N} \) (6.6) and \( b \) (–1) from this study. Combining province means of \( J_{C,N}\text{sequ}/e_{C,N} \) with \( J\text{-POCsequ} \) (Koeve 2002) yields that preferential N-remineralisation is equivalent to 1 to 1.5% of the \( J\text{-POCsequ} \) without preferential N-remineralisation. Looking at the basin scale difference between carbon export production and effective carbon flux at the base of the winter mixed layer, the effects of Redfield equivalent remineralisation within the seasonal thermocline or the effect of CaCO₃ sequestration (Koeve 2002) are each about 10 times as large as the effect of preferential N-remineralisation.

Export of DOC from the euphotic zone during winter convection has been reported (Hansell & Carlson 2001) and can account for the export of a fraction of seasonal net carbon uptake. In addition, a rapid transfer of carbon from the surface mixed layer during summer has been estimated from a carbon and nitrogen budget approach (Körtzinger et al. 2001a). Dia- pycnal transport is required for this summer time carbon export and TEP has been invoked as a possible vehicle for it (Körtzinger et al. 2001a). In that study, carbon overconsumption was not associated with the diatom spring bloom (Koeve 2004) but occurred later into the summer system. Slow sinking speeds of TEP-rich aggregates (Engel & Schartau 1999) not ballasted with opal, calcium carbonate, or terrigenous matter and grazing (Dilling et al. 1998, Prieto et al. 2001) or disruption of aggregates by large zooplankton (Dilling & Allardyce 2000) may explain why such an export from the surface mixed layer during summer does not lead to an effective export of excess carbon into the deep sea.

The discrepancy between the range of \( e_{C,N}\text{req} \) from this study and observed ranges of the C:N ratio of net community production (NCP_{C:N}) may be interpreted as a measure of the coupling of total export production (POM + DOM route) and deep carbon fluxes. Since \( e_{C,N}\text{req} \) (Figs. 4 & 6) was found to be much lower than recently published NCP_{C:N} values (10 to 15, Sambrotto et al. 1993, Körtzinger et al. 2001a), I conclude that total export production and deep fluxes of carbon are only loosely coupled. A significant fraction of carbon overconsumption and its export from the euphotic zone may be related to a fraction of DOM poor in nitrogen (Williams 1995, Kähler & Koeve 2001) which, however, appears to be remineralised in the upper ocean (Ono et al. 2001). As far as the sequestration of carbon into the deep ocean, for example below the winter mixed layer, is concerned, there is no need to involve a significant extra flux of carbon due to transparent exopolymer particles, as recently suggested by Engel & Passow (2001) and Engel (2002).

Such a 'TEP-pump' appears, at least according to the results of this study, not to provide a significant mechanism to sequester carbon into the deep ocean in addition to the conventional Redfield biological pump, unless one can prove that either the remineralisation C:N ratio (\( r_{C,N} \)) or the C:N ratio of sinking particles (\( J_{C,N} \)) in the deep ocean are much larger than currently thought. For the computation of carbon fluxes into the deep ocean from J-EP-Z functions, \( E_{PC} \) may be reliably, that is within the error bounds of other uncertainties inherent to this approach, computed from nutrient-based estimates of export production and a constant, Redfield, C:N:P ratio.
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