Biogeochemical fluxes through mesozooplankton.

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Submitted to Global Biogeochemical Cycles 2005GB002511.

Abstract
Mesozooplankton are significant consumers of phytoplankton, and have a significant impact on the oceanic biogeochemical cycles of carbon and other elements. Their contribution to vertical particle flux is much larger than that of microzooplankton, yet most global biogeochemical models have lumped these two plankton functional types together. In this paper we bring together several newly available data syntheses on observed mesozooplankton concentration and the biogeochemical fluxes they mediate, and perform data synthesis on flux rates for which no synthesis was available. We update the equations of a global biogeochemical model with an explicit representation of mesozooplankton (PISCES). We use the rate measurements to constrain the parameters of mesozooplankton, and evaluate the model results with our independent synthesis of mesozooplankton concentration measurements. We also perform a sensitivity study to analyse the impact of uncertainty in the flux rates. The standard model run was parameterised based on the data synthesis of flux rates. The results of mesozooplankton concentration in the standard run are slightly lower than the independent databases of observed mesozooplankton concentrations, but not significantly. This shows that structuring and parameterising biogeochemical models based on observations without tuning is a strategy that works. The sensitivity study showed that by using a maximum grazing rate of mesozooplankton that is only 30% higher than the poorly constrained fit to the observations, the model mesozooplankton concentration gets closer to the observations, but mesozooplankton grazing becomes higher than what is currently accounted for. This is an indication that food selection by mesozooplankton is not sufficiently quantified at present. Despite the amount of effort that is represented by the data syntheses of all relevant processes, the good results that were obtained for mesozooplankton indicates that this effort needs to be applied to all components of marine biogeochemistry. The development of ecosystem models that better represent key plankton groups and are more closely based on observations should lead to better understanding and quantification of the feedbacks between marine ecosystems and climate.

Index terms: 1615, 1635, 4806, 4842, 4855
Key words: mesozooplankton, export production, synthesis of observations, global ocean
1. Introduction

Mesozooplankton are a group of multicellular organisms (size range: 200 µm to 2 mm) that feed on phytoplankton, microzooplankton, other mesozooplankton and detritus. Together with the specialised filter feeders (salps, pteropods), they are the largest organisms that still have a significant feedback interaction with primary production. Thus, they are the largest organisms that are of primary interest for global biogeochemical models of the marine carbon cycle.

One of the main impacts of ecosystem functioning on biogeochemistry is the distinction between microzooplankton grazing (associated with the microbial loop and regenerated production), and mesozooplankton grazing (associated with the classical food chain and export production). Global biogeochemical models tend to represent all zooplankton as one state variable, and until recently also represented all phytoplankton as one state variable. These NPZD models (Nutrient Phytoplankton Zooplankton Detritus), and the current Earth system models that incorporate these models, underestimate interannual variability of chlorophyll $a$ as observed by satellite (Le Quéré et al., 2005). This suggests that these biogeochemical models will also underestimate sensitivity to decadal to century scale climate variability. Our working hypothesis is that building models that more closely represent our current understanding of the marine ecosystem will help to solve this problem (Prentice et al., 2004).

The explicit representation of mesozooplankton in models should also improve the representation of algal blooms. The preferred prey size of copepods (the most abundant mesozooplankton) is larger than the preferred prey size of microzooplankton (Hansen et al., 1994). The combination of different maximum growth rates for the two zooplankton size classes and the different preferred prey sizes leads to an effective control of small phytoplankton by microzooplankton, so that phytoplankton blooms are predominantly made up of large cells. This positive correlation between cell size and primary production, and the higher sinking rate of mesozooplankton fecal pellets also lead to a positive correlation between primary production and export efficiency. From this, glacial/interglacial differences in the marine ecosystem have been shown to influence the atmospheric CO$_2$ concentration (Bopp et al., 2003).

Unlike microzooplankton (Sherr and Sherr, 2002), mesozooplankton have a maximum growth rate (Hirst and Bunker, 2003) that is considerably lower than that of phytoplankton (Eppley, 1972). Thus, they tend to dampen biomass increases of both phytoplankton and microzooplankton on a timescale of their doubling time (with a minimum of 2 days at 30 °C to 9 days at 0 °C, Hirst and Bunker, 2003). For a biogeochemical model this presents advantages. The closing term in biological models is the mortality term of the highest represented trophic level. Using mesozooplankton as the highest represented trophic level has the advantages of including their small, but significant direct grazing of primary production (about 10%, Calbet, 2001), and of representing the grazing mortality of microzooplankton in a dynamical way, rather than as a closure term, which would ignore significant trophic interactions. The lower growth rate of mesozooplankton suggests that representing mesozooplankton mortality as the model closure term, and thus ignoring the variability in mesozooplankton mortality caused by higher trophic levels, would have a smaller impact on the variability of phytoplankton and microzooplankton production.

Given the potential importance of mesozooplankton for the export efficiency (Honjo and Roman, 1978) in global biogeochemical cycles, we have reviewed the available data on mesozooplankton biogeochemistry. In recent years, several data syntheses of measurements have been generated for both mesozooplankton concentration (O’Brien et al., 2002; Finenko et al.,
2003; Piontkovski and Landry, 2003; Beaugrand et al., 2003) and flux rates (Straile, 1997, Ikeda et al., 2001; Hirst and Kiørboe, 2002; Hirst and Bunker, 2003). In addition, we present a new compilation of individual research papers for the other mesozooplankton flux rates (Kremer, 1977; Copping and Lorenzen, 1980; Torres et al., 1982; Small, 1983; Torres and Childress, 1983; Buskey, 1998; Steinberg et al., 2000; Besiktepe and Dam, 2002). With these observations we obtain all mesozooplankton equations and parameters except food preferences in the PISCES global biogeochemical model. This is part of an ongoing effort to improve the representation of ecosystem processes in global biogeochemical models that are more closely based on observations, both for model formulation and evaluation (Le Quéré et al., 2005).

2. Model

In this section we describe two versions of the biogeochemical model PISCES (Aumont et al., 2003). Both versions contain some changes relative to the last published version of Bopp et al. (2003). We will refer to the version with only those changes as PISCES, and the version that contains additional changes in the parameterisation and equations of the biogeochemical fluxes through mesozooplankton as PISCES-T.

2.1 PISCES

In the PISCES model, there are two phytoplankton size classes (diatoms and nanophytoplankton) and two zooplankton size classes (micro- and mesozooplankton). Phytoplankton nutrient limitation is calculated as the minimum of the potential growth rates on PO$_4$, Fe, and SiO$_3$ for diatoms. Nutrient limitation is multiplied by light limitation. Both zooplankton types eat both phytoplankton types, but microzooplankton prefer nanophytoplankton and mesozooplankton prefer diatoms. The model has 24 state variables. There are eight dissolved / nutrient components: dissolved inorganic carbon (DIC), Alkalinity, O$_2$, PO$_4$, SiO$_3$, Fe, DI$^{13}$C, and $^{18}$O$_2$. There are seven phytoplankton components: Nanophytoplankton (NAN), Diatoms (DIA), the iron content of the two phytoplankton, the chlorophyll content of the two phytoplankton, and the silica content of diatoms. There are two zooplankton components: Microzooplankton (MIC), and Mesozooplankton (MES). And there are seven dead particulate / detritus components: small particulate organic carbon (POC$_s$), large particulate organic carbon (POC$_l$), dissolved organic carbon (DOC), CaCO$_3$, particulate silica, and the iron content of the two size classes of POC. The model is forced by river input of PO$_4$ and SiO$_3$, and dust input of Fe and SiO$_3$.

Since the last published version, the particulate organic matter was split into two size classes, small POC (POC$_s$) and large POC (POC$_l$). The losses from nanophytoplankton and microzooplankton go to POC$_s$, while losses from diatoms and mesozooplankton go to POC$_l$. This helps the model to distinguish between a predominantly recycling microbial loop and a classical food chain with a higher ratio of export production to primary production. Both micro- and mesozooplankton feed on POC$_s$.

The model has variable Chl a /C ratios for the algae, variable Si/C ratios for diatoms and POC$_s$, variable Fe/C ratios for the algae and the two size classes of POC, and constant C/P and C/O ratios.

The model also has a full implementation of the penalty scheme of André (1990) when the mixed layer depth (MLD) is deeper than the euphotic depth (EuD). This is a parameterisation for the mortality that some algal groups suffer when they spend part of the time below the compensation light intensity. The penalty is 0 when MLD < EuD, maximum when the MLD > 2*EuD, and intermediate by linear interpolation in-between. The maximum penalty is set to 0.5
for nanophytoplankton and 0 for diatoms. This reflects their differing sensitivities to darkness, which possibly reflects a lower need for maintenance in diatoms because their silica frustule and lack of flagella lowers the energy needed to maintain the osmotic gradient across the plasma membrane (R. Geider, pers. comm.).

2.2 Mesozooplankton equations in PISCES-T

The change in the concentration of mesozooplankton (Figure 1) is calculated with the following equation:

\[
\frac{\delta \text{MES}}{\delta t} = \sum \text{grazing}_F \times \text{model growth efficiency} - \text{basal respiration} - \text{mortality}
\]

where MES is the mesozooplankton concentration. The food sources F for mesozooplankton are diatoms (DIA), microzooplankton (MIC), nanophytoplankton (NAN) and small particulate organic carbon (POC).

2.2.1 Grazing

The mesozooplankton grazing rate on any one food is described by the following equation:

\[
\text{grazing}_F = G_{0\,^\circ\text{C}}^\text{mes} \times b^T \times \frac{p_F^\text{mes} \times \text{MES}}{K_{1/2}^\text{mes} + \sum p_F^\text{mes} \times \text{MES}}
\]

Where \(G_{0\,^\circ\text{C}}^\text{mes}\) is the maximum grazing rate at 0 °C, \(b\) is the temperature dependence of growth rate \(b^{10} = Q_{10}\), \(T\) is the temperature, \(p_F^\text{mes}\) is the preference for food F, and \(K_{1/2}^\text{mes}\) is the half saturation constant for grazing.

2.2.2 Partitioning of grazing

In field and laboratory experiments, net mesozooplankton growth is given by:

\[
\text{net growth} = \text{grazing} - \text{egestion} - \text{respiration} = \text{grazing} \times \text{GGE}
\]

where GGE (gross growth efficiency) is the part of grazing that is incorporated into biomass, egestion is partitioned between particulate egestion to POC and dissolved egestion to DOC, and respiration produces dissolved nutrients and DIC. During experiments in which net growth occurs, egestion and respiration are typically proportional to grazing, which makes it possible to formulate a dimensionless partitioning of grazing:

\[
\text{GGE} + \text{unass} + \text{diss} + \text{resp} = 1
\]

where GGE, unass, diss and resp are the fractions of grazing that are partitioned to MES, POC, DOC and DIC. However, zooplankton also respire when they don’t feed, which makes it necessary to introduce a term that is independent of grazing: basal respiration. During net growth, basal respiration is part of respiration. It would be computationally inefficient to evaluate at every time and place if net growth is occurring. Therefore, we split respiration into feeding respiration and basal respiration, in which feeding respiration is proportional to grazing. The model always subtracts basal respiration from mesozooplankton biomass, and the fraction of grazing that is
partitioned to mesozooplankton is made proportionally larger and respiration proportionally smaller:

\[
\text{model growth efficiency + unass + diss + feeding resp} = 1 \tag{5}
\]

\[
\text{model growth efficiency} = \frac{GGE + \text{basal respiration}}{\Sigma \text{grazing}}_{\text{mes}} \tag{6}
\]

Further details of the partitioning were chosen based on the available experimental measurements and are given in Section 3.1.2 on the parameterisation.

2.2.3 Basal respiration

The basal respiration was calculated as:

\[
\text{basal respiration} = \text{resp}_{0 \degree C}^{\text{mes}} \times d^{T} \times \text{MES} \tag{7}
\]

where \(\text{resp}_{0 \degree C}^{\text{mes}}\) is the respiration rate at 0 °C, and \(d\) is the temperature dependence of respiration \((d^{10} \equiv Q_{10})\).

2.2.4 Mortality

The mortality rate was calculated as:

\[
\text{mortality} = \text{mort}_{0 \degree C}^{\text{mes}} \times c^{T} \times \text{MES} \tag{8}
\]

where \(\text{mort}_{0 \degree C}^{\text{mes}}\) is the mortality rate at 0 °C, and \(c\) is the temperature dependence of mortality \((c^{10} \equiv Q_{10})\). Most of the mortality of mesozooplankton is believed to be due to predation \((Hirst \text{ and Kiørboe, 2002})\), but for lack of higher trophic levels in the model, the mortality is added to POC\(_{l}\). We also tested a mesozooplankton concentration dependent mortality rate:

\[
\text{mortality} = \frac{\text{mort}_{0 \degree C}^{\text{mes}}}{\text{MES}_{\text{ave}}^{\text{power}}} \times c^{T} \times \text{MES}^{\text{1+power}} \tag{9}
\]

where \(\text{MES}_{\text{ave}}\) is the average mesozooplankton concentration. We will refer to Equation 8 as linear mortality and to Equation 9 with power = 1 as squared mortality.

2.2.5 Other fluxes

The other mesozooplankton mediated fluxes are:

\[
\delta \text{DOC/δt} = (1 - \text{inorg}) \times (1 - \text{unass - model growth efficiency}) \times \Sigma \text{grazing}_{\text{mes}} \times \text{MES} \tag{10}
\]

\[
\delta \text{POC/δt} = \text{unass} \times \Sigma \text{grazing}_{\text{mes}} \times \text{MES} + \text{mort}_{0 \degree C}^{\text{mes}} \times c^{T} \times \text{MES} \tag{11}
\]

\[
\delta \text{PO}_{4}^{3-}/\delta t = \text{inorg} \times (1 - \text{unass - model growth efficiency}) \times \Sigma \text{grazing}_{\text{mes}} \times \text{MES} + \text{resp}_{0 \degree C}^{\text{mes}} \times d^{T} \times \text{MES} \tag{12}
\]

where inorg controls the partitioning between respiration to DIC, PO\(_{4}\) & Fe, and dissolved egestion to DOC.
2.3 Differences between PISCES and PISCES-T

In PISCES-T the basal respiration rate is density dependent. In the observations, there is no indication that respiration rate is a function of mesozooplankton concentration (Ikeda, 1985; Ikeda et al., 2001). In PISCES-T the respiration rate is only temperature dependent, and respiration is the respiration rate times the mesozooplankton concentration.

In the standard simulation of PISCES-T, the mortality rate was changed to be independent of mesozooplankton concentration and an exponential function of temperature, so that the mortality flux becomes a linear function of mesozooplankton concentration. From the data it is obvious that copepod mortality is a function of temperature (Figure 2D). Since the dataset of Hirst and Köroboe (2002) does not include mesozooplankton biomass (it is a steady-state mortality estimate), there is no direct way to derive a squared mortality function as used in PISCES. We have therefore used a linear mortality function. In the sensitivity study we calculated mortality as a function of the square of the mesozooplankton concentration, in which the rate parameter was divided by the average observed mesozooplankton concentration (0.54 µM, Equation 9).

The POC degradation (both size classes) was changed to be an exponential function of temperature. We used the phytoplankton growth temperature dependence for this ($Q_{10} = 1.9$).

The sinking speed was changed to be independent of the nutrient concentration in the top 100 m.

These changes were made to make use of recently compiled databases of mesozooplankton flux rates. We did not add any significant computational complexity. Therefore, we will not address the issue of how much more complexity we may need to be able to match climate sensitivity of ocean biology in global biogeochemical models. This issue is explored in Le Quéré et al., 2005.

The parameters in PISCES-T were adjusted to give a reasonable ocean-atmosphere CO$_2$ flux (2.3 Pg·y$^{-1}$ in the 1980s and 2.4 Pg·y$^{-1}$ in the 1990s). This was done by increasing the POC degradation rate at 0°C to 0.18 d$^{-1}$ (PISCES 0.05 d$^{-1}$), increasing the sinking rates of POC$_s$ to 6 m·d$^{-1}$ (PISCES 3 m·d$^{-1}$) and of POC$_t$ to 100 m·d$^{-1}$ (PISCES 50 m·d$^{-1}$), and decreasing the half saturation constant for microzooplankton grazing to 6 µM (PISCES 18 µM). This PISCES-T simulation without nutrient restoring (see Section 2.5) will be presented in Carr (in press) and McKinley (in press).

2.4 Physical model

We use the OPA8.2 global general circulation model (Madec et al., 1999) with a horizontal resolution of 2° in longitude and on average 1.1° in latitude, and a vertical resolution of 10 m in the top 100 m, increasing to 500 m at 5 km depth. Since the model cannot calculate a meeting of the meridians on the north pole, the grid has been distorted in the northern hemisphere to have two “north poles” over northern Eurasia and North America (Timmermann et al., 2005). The model has a free surface height (Roullet and Madec, 2004), and is coupled to a thermodynamic sea ice model (Fichefet and Morales Maqueda, 1999). The vertical mixing is calculated at all depths using a turbulent kinetic energy model (Gaspar et al., 1990).

2.5 Model forcing

The model was forced by daily wind and precipitation from NCEP reanalysis (Kalnay et al., 1996) from 1993 to 2002. Sensible and latent heat fluxes are calculated with a bulk formula, using the temperature difference between the modeled sea surface temperature and the daily air
temperature from NCEP reanalysis. The latent heat flux also provides evaporation. At the end of each year the water balance is calculated. From this balance, a water flux correction is calculated that is applied over the course of the next year.

The model simulation was initialized with observations for DIC, Alkalinity, PO$_4^{3-}$, SiO$_3^{2-}$ and O$_2$ as in Le Quéré et al. (2003). Other tracers were initialised with the output of the previous model version. To keep the modeled nutrient distribution close to the observed distribution and facilitate the comparison of the results for mesozooplankton, chlorophyll $a$ and export at 100 m, to the evaluation data, we used nutrient restoring below the mixed layer depth. I.e., modeled PO$_4^{3-}$ and SiO$_3^{2-}$ were forced towards the seasonal World Ocean Atlas datasets (WOA01, Conkright et al. 2002) below the mixed layer depth, with a relaxation time of 30 days, both in the PISCES and the PISCES-T model. The WOA nutrient fields were linearly interpolated on a daily basis. Independent of the mixed layer depth, nutrients were never restored above 50 m depth and always restored below 100 m depth.

3. Data synthesis of mesozooplankton flux rates and concentrations
3.1 Parameterisation of the biogeochemical model

The parameters in the model were taken from several data syntheses or individual research papers (Table 1). If more than one paper was used, the model parameter was calculated as the weighted average, with the weight being the number of replicates that were measured in each paper.

3.1.1 Grazing

The grazing rate was calculated as the net growth rate divided by the gross growth efficiency. We calculated the net growth rate of the mesozooplankton using the extensive data compilation of Hirst and Bunker (2003). The data were refitted to the Michaelis and Menten (1913) kinetic equation with an exponential increase in growth rate with temperature (Figure 2A, B):

$$\mu_{\text{mes}} = \frac{\mu_{0^\circ \text{C}, \text{mes}} \times b^T \times \text{Chl} a}{K_{\frac{1}{2}, \text{chl}, \text{mes}} + \text{Chl} a}$$

This gave $\mu_{0^\circ \text{C}, \text{mes}} = 0.081 \text{ d}^{-1}$, $b = 1.059$ and $K_{\frac{1}{2}, \text{chl}, \text{mes}} = 0.057 \mu \text{g Chl} \cdot \text{L}^{-1}$. The $K_{\text{d}}$ for food was converted to carbon units by using the average C/Chl ratio of 54.8 [g/g] taken from the model.

We found no consistent observations of food preferences. From the data compilation of Straile (1997) on predator/prey size ratios we can conclude that mesozooplankton would prefer diatoms over nanophytoplankton. Likewise, Broglio et al. (2004) show that most copepods and half of the cladocerans prefer algae >5 $\mu$m over algae <5 $\mu$m. Neither database is structured in a way that allowed us to calculate preferences according to Equation 14. Preferences are typically chosen to add up to one. Doing this results in an effective $K_{\text{d}}$ that is much higher than the value of $K_{\text{d}}$ that is used in the grazing equation. This problem can be prevented (data not shown) by using preferences with a phytoplankton biomass weighted mean of 1:

$$\frac{(p_{\text{dia}, \text{mes}} \times \text{biomass}_{\text{dia}} + p_{\text{nan}, \text{mes}} \times \text{biomass}_{\text{nan}})}{\sum \text{biomass}_{\text{phytoplankton}} = 1}$$

$$p_{\text{dia}, \text{mes}} = p_{\text{nan}, \text{mes}} \times \text{relative preference}$$
The relative mesozooplankton preference for diatoms was 5 times that for nanophytoplankton. The phytoplankton biomasses were taken from a database based on accessory pigments over the world ocean (Uitz et al., pers. comm.). Since the $K_b$ was based on chlorophyll concentration, the preferences were calculated based on the phytoplankton biomass. The same preference was used for microzooplankton as for diatoms, and for small POC as for nanophytoplankton.

3.1.2 Partitioning of grazed food

The grazed material is partitioned between unassimilated fecal material, biomass increase, respiration that is associated with feeding (that is, which covers the cost of searching for and digesting food) to inorganic nutrients, and dissolved egestion to DOC. The unassimilated fraction was taken from Besiktepe and Dam (2002). As they note, their average is very close to the average of the extensive database compiled by Conover (1978). We did not use the database of Conover because it is based on a wide variety of older methods and various units that are often energy based rather than matter based, and it includes freshwater mesozooplankton. The GGE was taken from the metazoan GGE in Straile (1997, Figure 2C). This database includes freshwater mesozooplankton. The measurements we could find on dissolved egestion did not express this as a fraction of grazing, but as a fraction of respiration (Steinberg et al., 2000; Kremer, 1977; Copping and Lorenzen, 1980; Small et al., 1983). We did not convert this fraction to a fraction of grazing, but used it as measured. In the model, this results in a decrease of dissolved egestion when grazing decreases faster than basal respiration, due to an increase in the model growth efficiency (Equations 6, 10). There is some evidence to suggest that this is what happens (Ikeda and Dixon, 1982).

3.1.3 Basal respiration

We calculated basal respiration as routine respiration times the ratio between basal and routine respiration. We calculated routine respiration based on the database of Ikeda (Ikeda, 1985; Ikeda et al., 2001). The results were fit to Equation 7 (Figure 2E). In the respiration experiments, mesozooplankton were incubated without food for periods from 2.5 to 24 hours (references in Ikeda, 1985, and Ikeda et al., 2001). The respiration rate thus measured is expected to be close to the routine respiration rate that mesozooplankton show in their normal environment with intermittent swimming / feeding current activity (Ikeda et al., 2001). We consider swimming / generating a feeding current to be part of the feeding respiration, and have used the ratio between respiration at normal swimming speed and the respiration extrapolated to no swimming to represent the ratio between routine and basal respiration. From measurements of zooplankton respiration as a function of swimming speed the basal respiration rate was calculated by extrapolating to no swimming. The ratio between basal respiration and routine respiration of zooplankton with average swimming speeds was estimated to be $2/3$ (most measurements were done on euphausiids, Torres et al., 1982; Torres and Childress, 1983; Buskey, 1998).

3.1.4 Mortality

We calculated mortality based on the database of Hirst and Kiørboe (2002). The mortality rates of broadcast spawners were estimated using the same equation as for sac spawners (equation 1 in Hirst and Kiørboe, 2002) to derive a mortality rate for the total life history. The results for both spawning types together were fit to Equation 8 (Figure 2D).

3.2 Calculation of parameter uncertainties
The standard error that was calculated by the curve fitting program (Table 1) is not a realistic measure of the spread in the flux measurements (Figure 2A,B,D,E). Therefore, we calculated upper and lower bounds of the parameters by dividing each dataset in two parts. The high (low) part was taken as those flux measurements that were higher (lower) than the model equation fit to all the observations (the thick lines in Figure 2). We then fit the model equations to each low and high part of the datasets, which gives the parameters used in the sensitivity study in Table 2, and the thin lines in Figure 2. The $K_{s/2}$ calculated in the high part of the database for growth rate is lower than for the whole database, which is logical because it gives rise to higher growth rates at low food concentrations. Also note that the model equation for gross growth efficiency (GGE) does not include food concentration, which by definition gives horizontal lines in Figure 2C (when GGE is calculated as a linear function of food concentration, this explains only 1% of the variation in GGE). In a meta-analysis of the GGE of metazoans, GGE was found to be highly variable between < 10 and > 80 %, i.e. the measured range in GGE was as large as its theoretically possible range (Straile, 1997). GGE was neither strongly related to temperature, nor to predator-prey size ratio and food concentration. However, there is evidence that GGE decreases at high food concentrations due to a decrease in assimilation efficiency (Landry et al., 1984).

The uncertainty in the flux measurement data is highest for mesozooplankton growth rate (Figure 2A). The high estimate of net grazing at 0 °C is 8.8 times higher than the low estimate. This will have been influenced by using chlorophyll as a measure of the food concentration. Chlorophyll is the most widely available measure of food concentration (Hirst and Bunker, 2003), but a variable chlorophyll/C ratio in algae and a variable contribution of algae to total food concentration is probably the main contributor to the uncertainty. For further details on sources of variability see Hirst and Bunker, 2003. Respiration and GGE are better constrained than growth rate, but still show considerable uncertainty, with the high estimate of respiration at 0 °C being 3.3 times higher than the low estimate, and the high estimate of GGE being 3.2 times higher than the low estimate. The uncertainty in mortality is relatively low, with the high estimate being 1.6 times higher than the low estimate. An exponential increase with temperature fits the data better than a linear increase (Figure 2D).

3.3 Mesozooplankton concentration evaluation databases

To evaluate the model we compiled a dataset of mesozooplankton concentration by combining three datasets (Table 3). These three evaluation datasets are independent of the rate measurements that were used to calculate the model parameters. Both the measurements that were used to calculate the model parameters and the evaluation data were sampled within the top 200 meters of the ocean.

3.3.1 NMFS dataset

The first dataset is the Coastal and Oceanic Plankton Ecology, Production Observation Database (COPEPOD) from the National Marine Fisheries Service (NMFS). It has a global coverage and includes samples that were sampled over the top 200m. (O’Brien et al., 2002, http://www.st.nmfs.noaa.gov/plankton, with an update over the 2001 version, hereafter referred to as the NMFS dataset). Mesozooplankton were collected with nets of 200-333 µm mesh size (the majority were 300 µm nets, with 200 µm nets used primarily in the Antarctic region ). The NMFS dataset was composed primarily of total wet mass and total displacement volume measures of mesozooplankton concentration, with lesser amounts of total dry mass. While different methods of sample preservation were used, they were most often applied after the total
sample mass or volume was measured. The different concentration types were converted to µg C·L⁻¹ according to Cushing et al., (1958), and then divided by 12 g·mol⁻¹. This set of conversion factors was used in preference over newer ones because it was based on 330 µm mesh nets, and because it provides a set of conversion factors that is consistent between the different measures of mesozooplankton concentration present (see also Postel, 2000).

3.3.2 FSU dataset

The second dataset is from cruises by the Former Soviet Union in the tropical and subtropical Atlantic Ocean. Samples were quantified in two ways: total seston (Finenko et al., 2003) and species level measurements (Piontkovski and Landry, 2003). The combined data will hereafter be referred to as the FSU dataset. For the total seston data, mesozooplankton were collected over the top 100 m with vertical hauls, predominantly with 112- to 142-µm mesh size nets. Samples were stored in 4 % buffered formalin. The total seston data were based on night time (18:00 – 04:00) net hauls from 1968 to 1992. Most were subsequently analyzed for wet weight biomass or by displacement volume. Both measures were converted to µM C by multiplying 0.16 g dry weight/ g wet weight (Vinogradov and Shushkina, 1987), 0.31 g C/ g dry weight (Wiebe, 1988) and 1/12 mol/g C = 0.0042.

For the species level data, mesozooplankton were collected over various depth ranges (see below) with vertical or oblique hauls of 178 and 200 µm mesh size nets. Samples were stored in 4 % buffered formalin. For copepods, the biomass of individuals was based on the measured length of cephalotorax plus abdomen. This length was converted to weight using four conversion factors based on shape (between 0.017 mg·length [mm]^{3.056} and 0.088·length^{2.715}, Gruzov and Alekseyeva, 1970). Other mesozooplankton were converted based on species specific conversion factors.

Some of the hauls for the species level data were over a shallower depth range than 0-100 m (on average to 67 m). Since mesozooplankton concentration decreases with depth, it might be expected that these concentrations would be higher. However, for the times and places where both total seston and species level samples were taken, the latter were on average 65% of the former. This might be due to the inclusion of net-phytoplankton and detritus in the total seston samples and/or uncertainties in the abundance to biomass conversion in the species level samples. We therefore included the species level samples in the comparison to the model results from 0-100 m without an additional depth range correction, and used the average between seston and species level data where both were available. The average contribution of copepods to the total mesozooplankton was 52% in the Caribbean and 66% in the open ocean between 25°S and 25°N.

3.3.3 CPR dataset

The third dataset is from the Continuous Plankton Recorder survey in the North Atlantic Ocean (Beaugrand et al., 2003, hereafter referred to as the CPR dataset). Plankton was sampled on a continuously moving band of silk with an average mesh size of 270 µm, and preserved in 4% formalin. Data for the period 1958-1999 were used in this study. Total calanoid copepod biomass per CPR sample was calculated from the mean size of each calanoid copepod (minimum size of female, 108 species or taxonomic groups), their abundance (assuming a sampled volume of 3 m³ per sample, Warner and Hays, 1994), and an allometric relationship (Legendre and Michaud, 1998): MES [µM C] = 0.08 [kg·length[m]^{2.1} /12[g/mol]. The minimum size of females as adult females or copepodite stage V was chosen as they represent the majority of copepods caught in the samples. The contribution of copepods to mesozooplankton biomass decreases from about 70% in polar regions to about 50% in temperate regions (Longhurst, 1998).
No correction was made for this varying contribution, and the estimated copepod concentration was compared to modeled mesozooplankton concentration.

3.3.4 Combined evaluation data

Because of the different (ranges of) sampling depths for the evaluation data, we calculated the model/data ratio separately over the relevant depth ranges: for the NMFS dataset from 0-200 m, for the FSU dataset from 0-100 m, and for the CPR dataset at 0-10 m. Some of the species level FSU data were submitted to the NMFS dataset. These values were removed from the NMFS dataset we used. Where more than one evaluation concentration was available at the same location, each concentration was included and compared to the appropriate model depth range. The model protects mesozooplankton from extinction by setting a minimum concentration of 0.01 µM. For the data analysis, the evaluation data were also set to this minimum. The combined evaluation dataset contains 6260 mesozooplankton concentrations on the model grid.

3.3.5 Consistency among the mesozooplankton biomass evaluation databases

The average observed mesozooplankton concentration is 0.54 ± 0.75 µM C (Table 2). We calculated the relative consistency between the three datasets. To perform this comparison we need to correct for the different sampling ranges in the evaluation data. We calculate this correction factor from the model at the places where the three paired datasets overlap, using the appropriate depth ranges in the model (Table 3). To prevent the outliers from dominating the results, the data were log transformed, that is: $\text{ratio} = 10^{\text{average}}(\log(\text{shallow observations/deep observations} \times \text{deep model/shallow model}))$. The FSU and CPR datasets are considerably lower than the NMFS dataset (Table 3, comparison corrected). In the case of the FSU / NMFS comparison, this is apparently due to the lower conversion factors in the seston data, which forms the largest part of the FSU dataset. In the case of the CPR / NMFS comparison, this could be due to the fact that the CPR database only contains copepod biomass, while the FSU and NMFS datasets contain all mesozooplankton. In the case of the CPR / FSU comparison, these two issues seem to cancel out against each other. To calculate how well the three datasets agree, we calculated $10^{\text{average}}(\text{ABS}(\log(\text{shallow observations/deep observations} \times \text{deep model/shallow model})))$ for the three datasets together. On average, the three datasets differ 5.6 fold.

3.4 Data – model comparison

To compare the model results to the evaluation data for mesozooplankton concentration, we calculated both the average modeled mesozooplankton concentration (Table 2), and the average point by point match among the three evaluation datasets (Section 3.3.5) and between the evaluation data and the model (see Sections 4.2 and 5.2). Since the observations are spatially biased towards the coast we only used the model results at the locations and depth ranges for which evaluation data was available.

To calculate the variability of the evaluation data, the standard deviation was calculated, using all data irrespective of the depth range of sampling. Since most of the evaluation data is based on only a few measurements in time for each location, it would be misleading to compare the variability to the variability of the model after averaging at every timestep for 5 years. Therefore, we calculated variability after sampling each point for which evaluation data is available at a random time step over the same depth range as the evaluation data.

4. Model results
4.1 Standard simulation

By implementing all fluxes as derived from our compilation of *in situ* measurements into PISCES-T we get a standard simulation, the results of which match the evaluation data of mesozooplankton concentration fairly well. On average, the model underestimates mesozooplankton concentration (Table 2), particularly in the equatorial regions (Figure 3C,D, 4A,D).

The PISCES-T model can also reproduce the SeaWiFS measured chlorophyll *a* field fairly well (Figure 5A, D), except for the coastal zone. On the other hand, the SeaWiFS protocol has been derived for open ocean or case 1 waters, and overestimates chlorophyll *a* in coastal or case 2 waters due to the presence of Gelbstoff and suspended particulate matter (IOCCG, 2000). This is confirmed by lower concentrations in the coastal regions in the map of *in situ* chlorophyll *a* measurements (Figure 5B). Relative to PISCES, the modeled chlorophyll *a* in the standard simulation is lower and not as close to the observations (Figures 4A,B, 5C,D).

4.2 Sensitivity analysis

We performed a sensitivity analysis of the mesozooplankton rate parameters with the standard model. In addition, we have also calculated the impact of using squared mortality on mesozooplankton concentration, and the impact of POC degradation on sinking export at 100 m. depth (see below).

The sensitivity study showed that at a first approximation, the impact of changing the parameters that affect mesozooplankton concentration is consistent with how well they are constrained by the observations, that is, in the order of growth rate, GGE and mortality (Table 2). However, there are two exceptions to this general trend: the simulation with the high grazing rate and the simulations with changed respiration. The mesozooplankton concentration is not highest at the high grazing rate. At low grazing rate the mesozooplankton concentration increases with the grazing rate. As expected, this trend does not continue indefinitely, because eventually the mesozooplankton concentration decreases due to a decrease in phytoplankton (see model optimization below, Figure 6).

Respiration is less constrained than mortality, but the impact of changes in respiration rate on the yearly averages is very small. This is because the model growth efficiency is corrected for basal respiration, since the measured GGE includes basal respiration (Equation 6 in Section 2.2.2), so that basal respiration only has an impact when it is higher than grazing. On a seasonal scale, though, the simulations with no basal respiration (in which case model growth efficiency equals GGE) can be quite different from the standard simulation (both higher and lower). This is due to mismatches in seasonal variation of respiration with temperature and variation of grazing with food concentration (see the additional material, link given in Acknowledgements).

In most simulations of the sensitivity analysis (Table 2), the chlorophyll concentration is lower when grazing and mesozooplankton concentration are lower. This is probably due to a trophic cascade, in which lower mesozooplankton grazing on microzooplankton leads to higher microzooplankton grazing on phytoplankton. A trophic cascade becomes apparent when microzooplankton grazing on phytoplankton increases faster with a reduction in mesozooplankton grazing on microzooplankton than the concomitant reduced direct impact of mesozooplankton grazing on phytoplankton. On the other hand, the chlorophyll concentration is also lower when grazing is higher, which could be due to an increased direct impact of mesozooplankton grazing on phytoplankton, and/or due to increased export of nutrients from the surface ocean.
4.3 Model optimization

To bring the model closer to the evaluation data we optimised the least constrained parameter: the grazing rate. The grazing rate was changed between the low and high estimates that were used in the sensitivity analysis (Table 2), but without changing the $K_{1/2}$ or the $Q_{10}$ (Figure 6). Given the large range of grazing rates that is spanned by these low and high estimates, the optimum grazing rate lies quite close to the best fit to the observations (Figure 2A,B). At the optimum grazing rate, the model is closest to the observations of mesozooplankton concentration (Figure 6, Table 2). The agreement between the optimised model and the evaluation data of mesozooplankton concentration was calculated as $10^{\text{average}(|\log(\text{model/data})|)}$. On average, the optimized model differed 2.9 fold from the evaluation data. The exact grazing rate for the optimum depends on which rate is optimised: at a grazing rate of 0.37 d$^{-1}$ the mesozooplankton concentration is highest, at 0.4 d$^{-1}$ the grazing rate is highest and at 0.43 d$^{-1}$ the difference is smallest. The differences are really small, though. The results we show are for a grazing rate of 0.4 d$^{-1}$ (Table 2, Figures 3 & 4B).

4.4 Temperature dependence of POC degradation

Since bacterial activity is temperature dependent, we added a temperature dependence of POC degradation. To test the impact of using this temperature dependence, we compared it to a model run in which POC degradation was independent of temperature, and adjusted the degradation rate to give the same global export (Table 2). With the temperature dependence the model differs 2.3 fold from the evaluation data, while without the temperature dependence the difference was 2.4 fold. Although this is not a large difference, it is as good as the PISCES model (see also Figures 3E, F and 7B, C). PISCES used a sinking speed that depends on the nutrient concentration in the top 100 m. This reproduces the low export : primary production ratio in oligotrophic regions without representing the mechanism that connects the nutrient concentration to sinking speed, and therefore was a rather unrealistic parameterisation.

5. Discussion

5.1 Flux and biomass measurements

It has long been known that zooplankton fecal pellets can form a large part of the vertical material flux to the deep sea (e.g. Honjo and Roman, 1978), but the relative contribution to total export is still unresolved (Turner, 2002). The correlation between the mesozooplankton biomass databases and POC export at 100 m. (NMFS: $r=0.30$, n=5324; CPR: $r=0.35$, n=1568; FSU: $r=0.31$, n=763) is about the same as the correlation between SeaWiFS and export over the region where NMFS data is available ($r=0.29$, n=4885). Thus, we cannot make any a priori statement about the contribution of mesozooplankton to total export based on the mesozooplankton observations alone.

5.2 Data / model comparison

The area weighted average mesozooplankton concentration in the model (at the locations where evaluation data is available) could be brought into close agreement with the evaluation data (Table 2) by increasing the grazing rate by 30%, and otherwise using model parameters that were derived from independent flux rate measurements (Figure 2, Table 1). The point by point difference in average mesozooplankton concentration between the standard model and the evaluation data is 4.3 fold, while after optimization it decreases to 2.9 fold (Figure 6). This is smaller than the average point by point difference between the three evaluation datasets, which is 5.9 fold (Section 4.1). Thus, we cannot expect to have enough information to significantly
improve the model further by trying to minimize the data / model difference. This suggests that although at this point already several thousand measurements were used for both parameterisation and evaluation of the model, further improvement could be achieved by extending these databases. This is also suggested by the model / data ratio for the CPR data, which shows the highest differences at the edges of the sampled region (data not shown), where the number of observations is smallest.

5.3 The relative impact of simplifying mesozooplankton traits into model equations

As shown in the previous section, the difference between the standard model simulation and the evaluation data falls within the uncertainty range of the evaluation data, and can be further decreased by optimising the grazing rate within the uncertainty range of the observations. It is also possible that the mismatch in mesozooplankton concentration in the standard model is due to lacking processes and mesozooplankton behaviour, such as vertical migration, diapausing, different functional responses between the various taxonomic groups within the mesozooplankton, and reduced swimming activity during food shortage. For instance, the respiration rate of mesozooplankton (mostly *Calanus* and *Neocalanus* copepods) diapausing at depth (>500m) is 20-30% of that at the surface (*Ikeda et al.*, 2005).

In the model, the elemental ratios of mesozooplankton are fixed. For respiration, there is a substantial amount of data for C, N and P specific rates (*Ikeda*, 1985; *Ikeda et al.*, 2001). The Q_{10} values for the three elements are 2.5, 6 and 3.1, respectively. The rates for C and N are the same at about 30 °C, while for C and P they are the same at about 15 °C (data not shown). To calculate the model parameters, we used the specific rates of the three elements together (Figure 2E). At present, there is not enough data for all flux rates to simulate variable elemental stoichiometries of mesozooplankton.

The model also does not simulate the body mass of individuals, which can explain an important part of the variability in the flux rates for copepods (*Ikeda*, 1985; *Hirst and Bunker*, 2003). Copepods form a large fraction of the mesozooplankton biomass, and in some cases parameters are exclusively based on measurements of copepods (Table 1). Body mass of individual copepods shows an increasing trend towards the poles, so that some of this poleward trend will be attributed to temperature in the data fits we used. The spatial distribution of copepod species follows trends in SST (*Beaugrand and Reid*, 2003). Therefore, this (partial) attribution of body weight variability in the ocean to temperature variability in the model should be robust in the face of global change. If we were to assume that both the concentration observations and the flux rate observations are accurate, then the mismatch between the standard simulation and the evaluation data would be an indication of the relative importance of the missing processes in the model.

5.4 Sensitivity analysis

Grazing rate is the least constrained parameter of mesozooplankton. Possibly the main contributing factor to this is that there is no consistent data on the relationship between grazing and food quality. In the database we used, food concentration was expressed as chlorophyll *a* concentration. However, POC concentration would have been a better basis for defining material fluxes through mesozooplankton. Of the 2743 data points in the growth rate database, only 68 include a measurement of POC/N. In addition to variable chlorophyll/C ratios, the relative proportions of different species of live food and POC are important contributors to variability in grazing. Some studies have investigated the food selectivity of key dominant mesozooplankton and its influence on rates (e.g. *Broglio et al.*, 2004; *Koski et al.*, 1998), but no common
denominator is apparent from these studies, nor do they cover the full diversity of food items that is found in the ocean. This issue of food preference and grazing rate as a function of food quality is the single most important topic that needs to be quantified before we can make significant further progress in modeling mesozooplankton in global biogeochemical models.

The spatial distribution of export in the standard simulation is slightly better than without temperature dependence of POC degradation. It is as good as in the PISCES model, which has no temperature dependence, but a sinking rate of POC that depends on the nutrient concentration in the top 100m. This is not entirely surprising, as the most nutrient poor regions are found in the warm tropical gyres, so that in PISCES export is low because of the low sinking speed and in PISCES-T export is low because of the high degradation rate. Using a temperature dependence is a more reasonable mechanism though, that directly represents the increase of bacterial degradation with temperature.

5.5 Mortality as a square function of mesozooplankton concentration?

By using the average observed mesozooplankton concentration in Equation 9, the mortality in the simulation with squared mortality is decreased below the average observed mesozooplankton concentration and increased above it. Thus, modeled and observed mesozooplankton concentration are not independent in the simulation with squared mortality. Therefore, the fact that modeled mesozooplankton concentration in the simulation with squared mortality is closer to these same observations than the standard simulation (Table 2) can’t be used to decide whether Equation 9 is better than Equation 8.

In biogeochemical models it is common practice to use a mortality term for zooplankton that is a square function of their concentration. PISCES also used this approach. The numerical reasoning behind this is that the model becomes more stable by having a higher mortality rate at high zooplankton concentration and vice versa. In NPZD models it may serve to represent mesozooplankton grazing on microzooplankton. The ecological reasoning behind this is that carnivory becomes more important as the zooplankton concentration increases. Out of five tested sites, two showed evidence of carnivory of adults on eggs (Ohman et al., 2004). This suggests that the threshold for this to become significant is at or higher than typical densities found in the ocean. In addition to this uncertainty and the absence of life stages in our model (eggs, adults and all stages in between are treated as one biomass), there are a number of reasons why we chose to use a linear mortality function:

1) The variability of the model is closer to the observed variability when linear mortality is used: the standard deviation of the observations is 0.75 µM, whereas the standard deviation using the linear mortality term is 0.26 µM and using the squared mortality term it is 0.12 µM.
2) The mortality rate was calculated with a database that does not include mesozooplankton concentration (Hirst and Kiørboe, 2002). Therefore, we calculated the squared mortality by dividing by the average mesozooplankton concentration in the evaluation data (Equation 9), which is biased towards high productivity regions.
3) There is so little data even on egg and nauplii mortality from grazing by adults (e.g. Bonnet et al., 2004) that it is impossible to constrain the value of power in Equation 9.
4) There was no noticeable effect on model stability.

However, the mesozooplankton to chlorophyll a ratio decreases with chlorophyll on a log / log plot, both for the evaluation data (Figure 8A) and for lakes (Figure 2B in Leibold et al., 1997). This negative correlation is not evident in the standard simulation (Figure 8D), but is shown in the simulation with a squared mortality (Figure 8C). When using squared mortality, the mortality rate increases with the mesozooplankton concentration. Due to the correlation between
mesozooplankton concentration and chlorophyll concentration, this results in a negative correlation between the mesozooplankton to chlorophyll $a$ ratio and the chlorophyll $a$ concentration. This does not constitute proof that the negative correlation is caused by squared mortality in the evaluation data. Several other simulations in the sensitivity analysis with a linear mortality also show a negative correlation, most clearly so in the simulation at low mesozooplankton growth efficiency (Figure 8B). In the latter case, the total food to chlorophyll $a$ ratio decreases with chlorophyll $a$.

Another mechanism that could result in a negative slope is that average size of phytoplankton tends to increase with productivity, which probably leads to an increase in the body mass of mesozooplankton. Since mesozooplankton growth rate decreases with body mass (Hirst and Bunker, 2003) this could also lead to a negative correlation between chlorophyll $a$ and the mesozooplankton to chlorophyll $a$ ratio in the ocean, while this process is not included in the model.

5.6 Optimization towards evaluation data

From a statistical point of view there is little to choose between the standard simulation that is based on flux measurements and the optimised simulation that is based on concentration measurements. The simulated global mesozooplankton grazing rate in the optimised simulation is considerably higher than the standard simulation (Table 2). Hernández-León and Ikeda (2005) estimated a global grazing rate of 26 Pg C·yr$^{-1}$, which they argue is a conservative estimate. This estimate is based on the respiration data of Ikeda et al. (1985), which is the largest component of the respiration dataset we used in the model (Figure 2E), and thus is not a completely independent estimate.

The only other estimate of global mesozooplankton grazing that we are aware of reported only mesozooplankton grazing on phytoplankton (5.5 Pg C·yr$^{-1}$, Calbet, 2001). From literature data it is also possible to estimate microzooplankton production on phytoplankton. Microzooplankton production on phytoplankton was estimated to be 8.7 - 11.4 Pg C·yr$^{-1}$. This was calculated as the ratio of microzooplankton grazing to phytoplankton growth (57-75 %, Calbet and Landry, 2004) times the microzooplankton growth efficiency (30 %, Straile, 1997) times the primary production (50.7 Pg C·yr$^{-1}$, the average of the primary production estimates from Antoine et al., 1996; Behrenfeld and Falkowski, 1997; and Behrenfeld et al., 2005). There are two reasons why this estimate would be an optimistic estimate of the flux from phytoplankton through microzooplankton to mesozooplankton. First, Dolan and McKeon (2004) have reviewed the method that is used to determine the ratio of microzooplankton grazing to phytoplankton growth, and show evidence that at low chlorophyll concentrations phytoplankton loss rates are overestimated. And second, part of this microzooplankton production would be grazed by other microzooplankton. Leaving aside these uncertainties, mesozooplankton grazing on phytoplankton and on microzooplankton eating phytoplankton would be 14-17 Pg C·yr$^{-1}$. At this moment we know of no global estimates of mesozooplankton grazing on POC and mesozooplankton grazing on microzooplankton eating POC and bacteria.

The standard simulation gives a mesozooplankton concentration that is lower than the evaluation data (Table 2). In the optimised simulation it was possible to simulate an average mesozooplankton concentration that is much closer to the evaluation data for mesozooplankton concentration without an unreasonable change in the grazing rate. In this simulation mesozooplankton grazing constitutes 77 % of primary production, compared to the conservative estimate of 51 % of primary production that is calculated by Hernández-León and Ikeda, (2005). The sensitivity runs with a high GGE or a low mortality share this same feature of getting an
average mesozooplankton concentration that is close to the evaluation data but at relatively high grazing fluxes. To reconcile the grazing flux in our optimized model of 43 Pg C·y$^{-1}$ with the estimated grazing rates on phytoplankton plus microzooplankton production on phytoplankton of about 15 Pg C·y$^{-1}$, direct and indirect grazing on POC and bacteria would have to make up the largest part of mesozooplankton grazing. A more detailed global data synthesis of mesozooplankton grazing would be another significant step forward in constraining biogeochemical fluxes through mesozooplankton.

6. Conclusion

Analysis of decadal scale observations of mesozooplankton species distribution has shown a significant correlation with climate variability (Beaugrand and Reid, 2003). To represent such ecosystem responses and their feedbacks on greenhouse gases, NPZD models are not adequate. Thus, biogeochemical models will need to be developed to include more complete representations of the marine food web. This will increase the number of model parameters to the point where they can no longer be constrained by fitting models only to those observations that constrain the overall behaviour of the marine food web, such as air-sea gas exchange. Here, we show that it is possible to constrain models to give results that are consistent with observations of chlorophyll $a$, mesozooplankton biomass and export by using databases of independent measurements of flux rates to calculate model parameters. Since the responses and feedbacks between marine ecosystems and climate change are largely unknown, we feel it is imperative that models are built that rely on and are evaluated by as full a complement of observations as is available, and that show an adequate response to observable climate variability. Here, we have suggested that the representation of mesozooplankton would benefit most from addressing two additional issues: (1) food preference of mesozooplankton and grazing rate as a function of food quality, including the relative importance of phytoplankton and microzooplankton, and (2) a detailed, georeferenced, global data synthesis of mesozooplankton grazing rates.

Acknowledgements

We thank the staff at DKRZ for their excellent and friendly support for the use of the NEC-SX6 computer. We thank Keith Rodgers for providing the forcing files. We thank the Laboratoire d’Océanographie Dynamique et de Climatologie for making the code of the OPA model available. And we thank the members of the Dynamic Green Ocean Project, especially Colin Prentice and Sandy Harrison, for their input throughout this project. EB thanks the EU for financial support under contracts HPMD-CT-2000-00038 & EVK2-CT-2001-00134. The FSU database assembly was supported by the NSF grant # DEB-0203622 (to SP). We thank the reviewers for their comments. For details on the model equations, flux rate and mesozooplankton concentration observations, simulation results, and general model documentation see the additional information at: http://lgmacweb.env.uea.ac.uk/green_ocean/meso.html. Submissions of additional data to the databases are very welcome, also for microzooplankton and phytoplankton to EB: martinburo@email.com.

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# Tables

Table 1. Mesozooplankton and POC degradation rate parameters in the standard model

<table>
<thead>
<tr>
<th>Flux</th>
<th>Parameter (Figure 1)</th>
<th>Rate</th>
<th>St.err</th>
<th>Organisms</th>
<th>References</th>
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</thead>
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<tr>
<td>grazing</td>
<td>$G_0^{mes}$</td>
<td>0.31 d$^{-1}$</td>
<td>0.03</td>
<td>copepods</td>
<td>1,2</td>
</tr>
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<td>$Q_{10}$ grazing</td>
<td>$b^{*10}$</td>
<td>1.77</td>
<td>3%</td>
<td>copepods</td>
<td>1</td>
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<tr>
<td>half saturation grazing</td>
<td>$K_{0.5}^{mes}$</td>
<td>0.26 µM</td>
<td>0.02</td>
<td>copepods</td>
<td>1</td>
</tr>
<tr>
<td>gross growth efficiency</td>
<td>$GGE$</td>
<td>0.26</td>
<td>0.18</td>
<td>copepods, cladocerans, rotifers</td>
<td>2</td>
</tr>
<tr>
<td>particulate egestion</td>
<td>$unass$</td>
<td>0.31</td>
<td></td>
<td>copepod</td>
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<tr>
<td>inorganic fraction of excretion</td>
<td>inorg</td>
<td>0.68</td>
<td></td>
<td>meso- and macrozooplankton</td>
<td>4,5,6,7</td>
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<td>preference for nanphytoplankton</td>
<td>$p_{nan}^{mes}$</td>
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<td></td>
<td></td>
<td>13</td>
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<tr>
<td>preference for diatoms</td>
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<td></td>
<td></td>
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<td>mortality</td>
<td>$mor^{mes}$</td>
<td>0.053 d$^{-1}$</td>
<td>0.001</td>
<td>copepods</td>
<td>8</td>
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<tr>
<td>$Q_{10}$ mortality</td>
<td>$c^{*10}$</td>
<td>1.99</td>
<td>1%</td>
<td>copepods</td>
<td>8</td>
</tr>
<tr>
<td>routine respiration</td>
<td>$resp_{r}^{mes}$</td>
<td>0.012 d$^{-1}$</td>
<td>0.003</td>
<td>mesozooplankton</td>
<td>9</td>
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<tr>
<td>basal respiration / routine resp.</td>
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<td>0.67</td>
<td>0.13</td>
<td>euphausiids, copepods</td>
<td>10,11,12</td>
</tr>
<tr>
<td>$Q_{10}$ routine respiration</td>
<td>$d^{*10}$</td>
<td>3.16</td>
<td>10%</td>
<td>mesozooplankton</td>
<td>9</td>
</tr>
<tr>
<td>POC, &amp; POC$_s$ degradation</td>
<td>0.18 d$^{-1}$</td>
<td></td>
<td></td>
<td></td>
<td>13</td>
</tr>
<tr>
<td>$Q_{10}$ POC, &amp; POC$_s$ degradation</td>
<td>1.89</td>
<td></td>
<td></td>
<td></td>
<td>13</td>
</tr>
</tbody>
</table>

* estimated as 10 times the standard errors in the temperature dependencies b, c, and d.
* estimated from the within experiment errors in the O$_2$ measurements.

1) Hirst and Bunker 2003, n=2743
2) Straile 1997, n=382
3) Besiktepe and Dam 2002, n=272
4) Steinberg et al. 2000, n=46, fraction=0.76
5) Kremer 1977, n=37, fraction=0.62
6) Copping and Lorenzen 1980, n=7, fraction=0.55
7) Small et al. 1983, n=3, fraction=0.62
8) Hirst and Kiørboe 2002, n=1051
9) Ikeda 1985, Ikeda et al. 2001, n=2962
10) Buskey 1998, n=7, ratio = 0.54
11) Torres and Childress 1983, n=271, ratio = 0.77
12) Torres et al. 1982, n=28, ratio = 0.62
13) This paper
Table 2. Model evaluation and sensitivity analysis

<table>
<thead>
<tr>
<th>changed rate</th>
<th>parameter values</th>
<th>chlorophyll&lt;sup&gt;µg·L&lt;sup&gt;-1&lt;/sup&gt;&lt;/sup&gt;</th>
<th>mesozooplankton&lt;sup&gt;µM C&lt;/sup&gt;</th>
<th>PP&lt;sup&gt;Pg C·y&lt;sup&gt;-1&lt;/sup&gt;&lt;/sup&gt;</th>
<th>export&lt;sup&gt;Pg C·y&lt;sup&gt;-1&lt;/sup&gt;&lt;/sup&gt;</th>
<th>grazing&lt;sup&gt;Pg C·y&lt;sup&gt;-1&lt;/sup&gt;&lt;/sup&gt;</th>
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<tbody>
<tr>
<td>data</td>
<td></td>
<td>0.35</td>
<td>0.54</td>
<td>37-67*</td>
<td>9.6-11.1*</td>
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<tr>
<td>PISCES</td>
<td>Aumont et al. 2003</td>
<td>0.21</td>
<td>0.16</td>
<td>69.7</td>
<td>9.9</td>
<td>5.9</td>
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<tr>
<td>standard</td>
<td></td>
<td>0.16</td>
<td>0.26</td>
<td>67.4</td>
<td>8.2</td>
<td>19.0</td>
</tr>
<tr>
<td>sensitivity (thin lines in Fig. 2)</td>
<td></td>
<td></td>
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<td></td>
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<tr>
<td>grazing high</td>
<td></td>
<td>0.97 d&lt;sup&gt;-1&lt;/sup&gt;</td>
<td>0.08</td>
<td>36.8</td>
<td>11.3</td>
<td>34.3</td>
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<tr>
<td></td>
<td>K&lt;sub&gt;5&lt;/sub&gt;=0.18µM C</td>
<td></td>
<td>Q&lt;sub&gt;10&lt;/sub&gt;=1.61</td>
<td></td>
<td></td>
<td></td>
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<tr>
<td>grazing low</td>
<td></td>
<td>0.13 d&lt;sup&gt;-1&lt;/sup&gt;</td>
<td>0.12</td>
<td>55.2</td>
<td>2.2</td>
<td>0.2</td>
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<td></td>
<td>K&lt;sub&gt;5&lt;/sub&gt;=2.2µM C</td>
<td></td>
<td>Q&lt;sub&gt;10&lt;/sub&gt;=2.23</td>
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<td>GGE high</td>
<td></td>
<td>0.42</td>
<td>0.11</td>
<td>44.8</td>
<td>16.8</td>
<td>39.2</td>
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<tr>
<td>GGE low</td>
<td></td>
<td>0.13</td>
<td>0.12</td>
<td>58.4</td>
<td>2.7</td>
<td>2.8</td>
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<tr>
<td>mortality high</td>
<td>0.064 d&lt;sup&gt;-1&lt;/sup&gt;</td>
<td>0.14</td>
<td>0.09</td>
<td>63.0</td>
<td>4.5</td>
<td>7.8</td>
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<td>mortality low</td>
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<td>0.15</td>
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<td>56.5</td>
<td>15.2</td>
<td>42.2</td>
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<td>resp high</td>
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<td>0.26</td>
<td>67.8</td>
<td>8.2</td>
<td>19.0</td>
</tr>
<tr>
<td></td>
<td>Q&lt;sub&gt;10&lt;/sub&gt;=3.19</td>
<td></td>
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<tr>
<td>resp zero&lt;sup&gt;#&lt;/sup&gt;</td>
<td>0 d&lt;sup&gt;-1&lt;/sup&gt;</td>
<td>0.16</td>
<td>0.26</td>
<td>67.1</td>
<td>8.2</td>
<td>19.0</td>
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<tr>
<td>square mortality</td>
<td>mortality</td>
<td>0.098 (d·µM C)&lt;sup&gt;-1&lt;/sup&gt;</td>
<td>0.16</td>
<td>0.41</td>
<td>57.5</td>
<td>13.6</td>
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<tr>
<td>POC degr.</td>
<td>no T dep.</td>
<td>0.72 d&lt;sup&gt;-1&lt;/sup&gt;</td>
<td>0.18</td>
<td>69.0</td>
<td>8.2</td>
<td>20.3</td>
</tr>
<tr>
<td></td>
<td>Q&lt;sub&gt;10&lt;/sub&gt;=1</td>
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<td>0.30</td>
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<td></td>
<td></td>
</tr>
<tr>
<td>optimized</td>
<td>grazing rate</td>
<td>0.4 d&lt;sup&gt;-1&lt;/sup&gt;</td>
<td>0.15</td>
<td>55.7</td>
<td>15.4</td>
<td>42.8</td>
</tr>
</tbody>
</table>

<sup>0</sup> From SeaWiFS, area-weighted as Σ(concentration*area)/Σarea

<sup>*</sup> Model sampled where evaluation data is available, area-weighted like chlorophyll. See Section 3.3 for details.

<sup>§</sup> Total calculated as Σ(flux*area)

<sup>•</sup> Range of estimates of primary production from Antoine et al. (1996), Behrenfeld and Falkowski (1997), and Behrenfeld et al. (2005).

<sup>†</sup> Range of estimates of export at 100 m from Schlitzer (2004), and Laws et al. (2000).

<sup▼</sup> See section 5.5

<sup>#</sup> Low respiration: 0.0057 d<sup>-1</sup>, Q<sub>10</sub>=2.71
Table 3. Mesozooplankton concentration datasets

<table>
<thead>
<tr>
<th>Dataset</th>
<th>number of data</th>
<th>number of data regridded</th>
<th>region</th>
<th>depth [m]</th>
<th>intercomparison corrected see Section 4.1</th>
<th>intercomparison uncorrected see Section 4.1</th>
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<tbody>
<tr>
<td>NMFS</td>
<td>5834</td>
<td>4804</td>
<td>World</td>
<td>0-200</td>
<td>0.39 (n=258)</td>
<td>0.72</td>
</tr>
<tr>
<td>FSU</td>
<td>1666</td>
<td>795</td>
<td>(Sub)tropical Atlantic</td>
<td>0-100</td>
<td>0.91 (n=50)</td>
<td>1.04</td>
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<tr>
<td>CPR</td>
<td>176778</td>
<td>661</td>
<td>North Atlantic</td>
<td>7</td>
<td></td>
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</tbody>
</table>

* including the species level data in the FSU dataset
♥ excluding the species level data in the FSU dataset

<table>
<thead>
<tr>
<th>Dataset</th>
<th>filter size µm</th>
<th>maximum size [mm]</th>
<th>sample handling</th>
<th>measured quantity</th>
<th>Conversion to µM C</th>
<th>references</th>
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<tr>
<td>NMFS</td>
<td>200-333</td>
<td>5-10 mm</td>
<td>various</td>
<td>displacement volume [ml] wet weight [mg] dry weight [mg]</td>
<td>8</td>
<td>0.01 0.05</td>
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<tr>
<td>FSU seston</td>
<td>112-142</td>
<td>30 mm, no jellies</td>
<td>4% buffered formalin</td>
<td>displacement volume [ml] wet weight [mg]</td>
<td>4.2</td>
<td>0.0042</td>
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<tr>
<td>FSU species</td>
<td>178-200</td>
<td>10 mm</td>
<td>4% buffered formalin</td>
<td>countings, length [mm]</td>
<td>1.4·L^{3.056} - 7.3·L^{2.715}</td>
<td>Piontkovski and Landry 2003</td>
</tr>
<tr>
<td>CPR</td>
<td>~270</td>
<td>7 mm</td>
<td>4% formalin</td>
<td>countings, length [mm]</td>
<td>3.3·L^{4.4}</td>
<td>Beaugrand et al. 2003</td>
</tr>
</tbody>
</table>
Figure 1. Mesozooplankton mediated fluxes. Circles represent state variables, underlined texts represent fluxes, and italic texts represent parameters (see Table 1 for explanations and values).
Figure 2. Databases used for calculating parameter values. Bold lines are fit to all measurements. Thin lines are fit to the data that lie above and below the fit to all measurements, approximately corresponding to the high and low halves of the data. The dashed line in panels A & B is the net growth rate used in the optimised simulation. A) Copepod growth rate as a function of chlorophyll a concentration corrected to a constant temperature of 15 °C (Hirst and Bunker, 2003). Some high values (up to 321 µg·L⁻¹) are not shown but were included in the data fits. B) Copepod growth rate as a function of temperature corrected to a constant chlorophyll a concentration of 3.7 µg·L⁻¹ (Same data as A, the uncorrected data was fit to Equation 13). C) Gross growth efficiency. Horizontal lines are averages, not calculated as a function of food concentration (Straile, 1997). D) Copepod mortality as a function of temperature (Hirst and Kiørboe, 2002). E) Unfed routine mesozooplankton respiration rates as a function of temperature (Ikeda, 1985; Ikeda et al., 2001).
Figure 3. Zonal averages. Black lines) Observations, Green lines) PISCES, Red lines) PISCES-T standard simulation, Blue lines) PISCES-T optimised simulation, Shaded grey areas) range of simulations in sensitivity analysis (excluding PISCES). A, B) surface chlorophyll, C, D) surface mesozooplankton, E, F) export at 100 m. A, C, E) Atlantic Ocean B, D, F) Pacific and Indian Oceans.
Figure 4. Mesozooplankton concentration [µM C] A) observed (NMFS, FSU (roughly delimited by white line) & CPR (delimited by black line)), B) PISCES-T optimised simulation, C) PISCES, D) PISCES-T standard simulation, E) PISCES-T standard simulation (y-axis) vs. observed (x-axis) +: NMFS (18 values between 5 and 9 µM are not shown), □: FSU, ×: CPR. Model results are integrated over the same depth ranges as the observations (Table 3), and over the top 200m where no observations are available.
Figure 5. Surface chlorophyll $a$ concentration. The scale is irregular. A) satellite observed (SeaWiFS), B) measured in situ (WOA database averaged from 1955-1998), C) PISCES, D) PISCES-T standard simulation, (A, C & D averaged from 1998-2002).
Figure 6. Sensitivity analysis and optimization of the grazing rate. Arrow down: standard simulation, Arrow up: optimised simulation. A) left y-axis: Filled circles) mesozooplankton concentration [µM C], grey triangles) chlorophyll a concentration [µg·L⁻¹], right y-axis: open squares) point by point difference between model mesozooplankton concentration and evaluation data, (10^average(absolute(log(model/observation)))); B) Filled circles) mesozooplankton grazing, grey triangles) primary production, open circles) export at 100 m.
Figure 7. Export at 100 m. depth [mol·m$^{-2}$·year$^{-1}$]. A) Results from the inverse model of Schlitzer (2004), based on geochemical observations. B) no temperature dependence of POC degradation C) PISCES. D) PISCES-T standard simulation.
Figure 8. LOG(mesozooplankton/chlorophyll \(a\) [\(\mu M \cdot (\mu g \text{ Chl})^{-1}\)]) vs. LOG(chlorophyll \(a\) [\(\mu g \text{ Chl}\)])
A) observed (using World Ocean Atlas Chlorophyll \(a\), integrated over the appropriate depth ranges)
B) PISCES-T low GGE simulation, surface concentrations
C) PISCES-T squared mortality simulation, surface concentrations
D) PISCES-T standard simulation, surface concentrations.