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Diagnosis of Physical and Biological Controls on Phytoplankton Distribution in the Sargasso Sea

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Abstract The linkage between physical and biological processes is studied by applying a one-dimensional physical-biological coupled model to the Sargasso Sea. The physical model is the Princeton Ocean Model and the biological model is a five-component system including phytoplankton, zooplankton, nitrate, ammonium, and detritus. The coupling between the physical and biological model is accomplished through vertical mixing which is parameterized by the level 2.5 Mellor and Yamada turbulence closure scheme. The coupled model investigates the annual cycle of ecosystem production and the response to external forcing, such as heat flux, wind stress, and surface salinity, and the relative importance of physical processes in affecting the ecosystem. Sensitivity experiments are also carried out, which provide information on how the model bio-chemical parameters affect the biological system. The computed seasonal cycles compare reasonably well with the observations of the Bermuda Atlantic Time-series Study (BATS). The spring bloom of phytoplankton occurs in March and April, right after the weakening of the winter mixing and before the establishment of the summer stratification. The bloom of zooplankton occurs about two weeks after the bloom of phytoplankton. The sensitivity experiments show that zooplankton is more sensitive to the variations of biochemical parameters than phytoplankton.

Key words physical-biological coupled model; annual cycle; external forcing; bio-chemical parameter

1 Introduction

Photosynthesis, the conversion of solar energy to chemical energy, is a fundamental step in which inorganic carbon is fixed by algae and converted into primary production. Significant primary production can only occur in the well-lit euphotic zone. Hence, the animals which feed on the primary production can survive mostly within the mixed layer where a large quantity of food is available. Physical processes play an important role in marine ecosystem dynamics (Collins et al., 2009; Denman and Pena, 2002; Mann and Lazier, 1991) and can modify or limit biological production through the nutrient supply and mean irradiance field (e.g., McClain et al., 1990; Mitchell et al., 1991). This paper studies the linkage between physical and biological processes in the Sargasso Sea via the application of a physical-biological coupled model. The model is one-dimensional and designed to investigate the vertical structure of the upper ocean mixed layer. The depth of the mixed layer, the intensity of solar radiation penetrating into water column, and the vertical distribution of dissolved nutrients are some of the major factors regulating the biosystem of the sea. The seasonal variation in the atmosphere-ocean heat flux imparts a seasonal cycle to the depth of the mixed layer (Menzel and Ryther, 1960). The variation of wind stress also affects the depth of the mixed layer. According to Menzel and Ryther (1960), production off Bermuda in the Sargasso Sea is closely related to vertical mixing, high levels occurring when the water is well mixed to or near the permanent thermocline at 400 m depth and low levels when a seasonal thermocline is present in the upper 100 m.

Ecosystem models have been widely applied to different oceanic conditions (*e.g.*, Varela *et al.*, 1992; Radach and Moll, 1993; Sharples and Tett, 1994). A recent application of a similar coupled physical-biological model using the BATS data (Doney *et al.*, 1996) well reproduced the seasonal cycles of the upper water column temperature field, as well as of the chlorophyll and primary production.

The goals of this study are to investigate and understand the interplaying and relative importance of the physical processes and the vertical distributions of nutrients and biomass in the euphotic zone. The biochemical part comprises five components, *i.e.*, nitrate, ammonium, phytoplankton, zooplankton, and detritus (Oguz *et al.*, 1996). A case-study is carried out by applying the model to the Sargasso Sea oligotrophic region, using the site data from the U. S. Joint Global Ocean Flux Study (JGOFS) Ber-

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muda Atlantic Time-series Study (BATS). The coupling between the biological and physical model is accomplished via vertical mixing coefficients. The study begins with examining the seasonal response of the mixed layer physics and biology to external forcing, including windstress, heat flux, and surface salinity. Following that, sensitivity experiments of the model components to biochemical parameters are performed in order to understand how the internal biochemical parameters affect and what the impact of nutrients, light availability, and the interaction between the biochemicals and production is on the biological system.

2 The Method

The complete model includes the physical and biological submodels, which is one-dimensional and time-dependent. The vertical mixing process is parameterized by the level 2.5 Mellor and Yamada turbulence closure scheme (Mellor and Yamada, 1982). The biological submodel is intentionally kept simple to explore the basic biological interactions and mechanisms.

2.1 The Physical Model

The physical model is the one-dimensional version of the Princeton Ocean Model (Blumberg and Mellor, 1987). For a horizontally homogeneous and incompressible sea water body under Boussinesq and hydrostatic approximations, the horizontal momentum equation is expressed as

$$\frac{\partial \vec{V}}{\partial t} - f\hat{k} \times \vec{V} = \frac{\partial}{\partial z} \left[(K_m + v_m)(\frac{\partial \vec{V}}{\partial z}) \right], \tag{1}$$

where t is time, z is the vertical coordinate, \vec{V} is the horizontal velocity of the mean flow with the components (u, v), \hat{k} is the unit vector in the vertical direction, f is the Coriolis parameter, K_m denotes the coefficient for the vertical turbulent diffusion of momentum, and v_m represents its background value associated with internal wave mixing and other small-scale mixing processes.

The temperature T and salinity S can be determined from the transport equation,

$$\frac{\partial C}{\partial t} = \frac{\partial}{\partial z} \left[(K_h + \nu_h) \frac{\partial C}{\partial z} \right], \tag{2}$$

where *C* denotes either *T* or *S*, K_h is the coefficient for the vertical turbulent diffusion, and v_h is its background value. For simplicity, the solar irradiance which penetrates into the water column is not parameterized in the temperature equation. It is represented through the surface boundary condition together with other heat flux components. The density is a function of potential temperature, salinity, and pressure, written as a non-linear equation of state, $\rho = \rho(T, S, p)$ (Mellor, 1990).

The vertical mixing coefficients are determined from

$$(K_m, K_h) = lq (S_m, S_h),$$
 (3)

where l and q are the turbulent length scale and turbulent

velocity, respectively. S_m and S_h are the stability factors as in Mellor and Yamada (1982). In the level 2.5 turbulence closure, l and q are computed from the turbulent kinetic energy, $q^2/2$, and the turbulent macroscale equations. The turbulent buoyancy and shear productions are calculated by the vertical shear of horizontal velocity and vertical density gradient of the mean flow. K_h is the eddy coefficient for the vertical turbulent diffusion of biological variables.

The boundary conditions at the sea surface z=0 are

$$\rho_0 K_m \frac{\partial \vec{V}}{\partial z} = \vec{\tau}_s \,, \tag{4}$$

$$K_h \frac{\partial T}{\partial z} = \frac{Q_H}{\rho_0 c_p},\tag{5}$$

$$S = S_0 , (6)$$

where $\bar{\tau}_s$ is the wind stress vector at the sea surface, Q_H is the net sea surface heat flux, S_0 is the sea surface salinity, ρ_0 is the reference density, and c_p is the specific heat of water. The 400 m level is taken as the bottom boundary of the model. No stress, heat and salt flux conditions are specified at the bottom

$$\rho_0 K_m \frac{\partial \tilde{V}}{\partial z} = 0, \qquad (7)$$

$$K_h \frac{\partial C}{\partial z} = 0 , \qquad (8)$$

where C again denotes either T or S.

2.2 The Biological Model

Nitrogen plays a critical role in ocean biology as an important limiting nutrient, particularly in subtropical gyres, and is a natural currency for studying biological flows (Fasham *et al.*, 1990). Biological constituents in the coupled model are treated as equivalent concentrations of nitrogen (mmolN m^{-3}). Following the physical rules outlined above the advection and diffusion of biological scalars and the biological interactions are modeled as nitrogen flows between compartments. The investigation in ecosystem modeling focuses on identifying the appropriate types of compartments and the linkages among them.

Detailed models may produce better and more realistic simulation results, but at the expenses of added complexity, less interpretable solutions, and increasing number of free parameters. Therefore, an attempt has been made to keep the model as simple as possible without eliminating essential dynamics of the system. The simple, five-component system including phytoplankton P, zooplankton Z, nitrate N, ammonium A, and detritus D is outlined schematically in Fig.1.

The local changes of biochemical variables are described by

$$\frac{\partial B}{\partial t} = \frac{\partial}{\partial z} \left[(K_h + v_h) \frac{\partial B}{\partial z} \right] + F_B , \qquad (9)$$

where *B* represents one of the five biological variables, phytoplankton biomass *P*, herbivorous zooplankton biomass *Z*, pelagic detritus *D*, nitrate *N*, and ammonium concentration *A*. F_B is the biological interaction term, which, for the five biological variables, can be written as (*e.g.*, Wroblewski, 1977; Fasham *et al.*, 1990):

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$$F_P = \Phi(I, N, A)P - G(P)Z - m_p P, \qquad (10)$$

$$F_Z = \gamma G(P) Z - m_h Z - \mu_h Z , \qquad (11)$$

$$F_D = (1 - \gamma)G(P)Z + m_p P + m_h Z - \varepsilon D + w_s \frac{\partial D}{\partial z}, \quad (12)$$

$$F_A = -\Phi_a(I, A)P + \mu_h Z + \varepsilon D - \Omega A, \qquad (13)$$

$$F_N = -\Phi_n(I, N)P + \Omega A, \qquad (14)$$

where the parameters and their default values are given in Table 1.



Fig.1 Schematic of the five-compartment biological model for nitrogen.

Table 1 Parameters and model default values (Wroblewiski et al., 1988; Doney et al., 1996;Hurtt and Armstrong, 1996; Oguz et al., 1996)

Parameter	Definition	Value	Units
f	Coriolis parameter	10^{-4}	s^{-1}
g	Gravitational acceleration	9.81	ms^{-2}
$ ho_0$	Reference density	1000	$\mathrm{kg}\mathrm{m}^{-3}$
c_p	Specific heat of water	$4e^3$	$J(kg °C)^{-1}$
κ	Von Karman constant	0.4	-
σ_m	Maximum phytoplankton growth rate	0.75	d^{-1}
k_w	Light extinction coefficient for PAR	0.03	m^{-1}
k_c	Phytoplankton self-shading coefficient	0.03	$m^2 mmol N^{-1}$
R_n	Nitrate half-saturation constant	1	$mmolN m^{-3}$
R_a	Ammonium half-saturation constant	0.8	$mmolN m^{-3}$
R_g	Herbivore half-saturation constant	0.3	$mmolN m^{-3}$
a	Photosynthesis efficiency parameter	0.05	$(w/m)^{-1}$
m_p	phytoplankton death rate	0.05	d^{-1}
r_g	Herbivore maximum grazing rate	0.21	d^{-1}
m_h	Herbivore death rate	0.01	d^{-1}
μ_h	Herbivore excretion rate	0.05	d^{-1}
Е	Detrital remineralization rate	0.05	d^{-1}
Ω	Ammonium oxidation rate	0.03	d^{-1}
W_S	Detrital sinking rate	0.5	$m d^{-1}$
γ_h	Herbivore assimilation efficiency	0.8	-
P_{0}	Initial phytoplankton concentration	0.05	$mmolN m^{-3}$
H_0	Initial herbivore concentration	0.1	$mmolN m^{-3}$
D_{θ}	Initial detritus concentration	0.05	$mmolN m^{-3}$
A_0	Initial ammonium concentration	0.1	$mmolN m^{-3}$

The total production of phytoplankton, $\Phi(I, N, A)$, is defined by

$$\Phi(I, N, A) = \sigma_m \min[\alpha(I), \beta_t(N, A)], \quad (15)$$

where a(I) and $\beta_t(N, A)$ represent the light limitation and the total nitrogen limitation function of phytoplankton uptake, respectively. The empirical relationship between the maximum phytoplankton growth rate and temperature was used to assign the maximum phytoplankton growth rate σ_m (Eppley, 1972; Fasham *et al.*, 1990) and $\beta_t(N, A)$ is given in the form of

$$\beta_t(N,A) = \beta_n(N) + \beta_a(A), \qquad (16)$$

with $\beta_a(A)$ and $\beta_n(N)$ representing the contributions of

ammonium and nitrate limitations, respectively. These two terms are expressed by the Michaelis-Menten uptake formulation as

$$\beta_a(A) = \frac{A}{(R_a + A)},\tag{17}$$

$$\beta_n(N) = \frac{N}{(R_n + N)} \exp\{-\psi A\}, \qquad (18)$$

where R_n and R_a are the half-saturation constants for nitrate and ammonium, respectively. The exponential term of Eq. (18) represents the inhibiting effect of ammonium concentration on nitrate uptake, with ψ signifying the inhibition parameter (Wroblewski, 1977).

The individual contributions of nitrate and ammonium uptakes to the phytoplankton production are represented by (Varela *et al.*, 1992)

$$\Phi_n(I,N) = \sigma_m \min[\alpha(I), \beta_t(N,A)](\beta_n / \beta_t), \quad (19)$$

$$\Phi_a(I,A) = \sigma_m \min[\alpha(I), \beta_t(N,A)](\beta_a / \beta_t).$$
(20)

The light limitation is parameterized by Jassby and Platt (1976)

$$\alpha(I) = \tanh[aI(z,t)], \qquad (21)$$

$$I(z,t) = I_s \exp[-k_w + k_c P]z],$$
 (22)

where *a* is the parameter of photosynthesis efficiency controlling the slope of $\alpha(I)$ to the irradiance curve at low values of the photosynthetically active irradiance (PAR). I_s denotes the surface intensity of PAR, which is taken as 0.45 for the climatological incoming solar radiation according to the available data.

The zooplankton grazing ability is represented by the Michaelis-Menten formulation

$$G(P) = \sigma_g \frac{P}{(R_g + P)}.$$
 (23)

For phytoplankton, zooplankton, nitrate, and ammonium, the boundary conditions at the surface and bottom are given by

$$(K_h + \nu_h) \frac{\partial B}{\partial z} = 0$$
, at $z = 0$, $z = -D$. (24)

For the detritus equation the surface boundary condition is modified to include the downward sinking flux

$$(K_h + v_h)\frac{\partial D}{\partial z} + w_s D = 0, \text{ at } z = 0, z = -D.$$
 (25)

The same condition is also prescribed at the lower boundary of the model, which is well below the euphotic zone at 400 m depth. A relatively low sinking rate is specified ($w_s=0.5 \text{ m d}^{-1}$, Table 1). The advantage of selecting the bottom boundary at a considerable distance away from the euphotic layer is to allow the complete remineralization of detrital material before reaching the lower boundary of the model. The vertically integrated biological model is fully conservative.

2.3 Physical Forcing

The annual variations of wind stress and heat flux components are expressed by smoothed and climatological surface forcing functions (Doney *et al.*, 1996)

$$F = Mean + Amplitude.\cos(2\pi \frac{t}{365} - phase), \quad (26)$$

where the time, t, is given in days. The annual mean, seasonal amplitude, and phase, as shown in Table 2, are computed from climatological data sets (Esbensen and Kushnir, 1981; Isemer and Hasse, 1985) for the region of the BATS site (31°50′N and 64°10′W). According to Zhu

et al. (2002), the estimated non-solar heat fluxes at a high time resolution could have large errors even if observation errors are small, but with a lower time resolution the large errors could be avoided.

Table 2 Climatological physical forcing functions for the default case

Forcing factor	Units	Annual mean	Amplitude	Phase (°C)
Wind stress	$\mathrm{N}\mathrm{m}^{-2}$	0.081	0.040	60
Net longwave	$\mathrm{W}\mathrm{m}^{-2}$	-60.0	5.0	70
Sensible heat	$W m^{-2}$	-26.0	22.0	170
Latent heat	$W m^{-2}$	-162.5	90.0	170
Solar	$W m^{-2}$	198.7	_	_

The surface wind stress (Fig.2c) peaks at $1.2 \,\mathrm{dyn \, cm^{-2}}$ in March, and the annual mean heat loss from the nonsolar terms is $248.5 \,\mathrm{W \,m^{-2}}$ with a maximum of $365.5 \,\mathrm{W \,m^{-2}}$ in late December. Solar radiation is computed with a constant cloud fraction of 0.75, which leads to an annual mean solar heating rate of 198.7 W m⁻² that is within the reported climatological range of 180-200 W m⁻² (Esbensen and Knshnir, 1981). The required cloud fraction, however, is slightly higher than the climatological value of approximately 0.6 near Bermuda (Warren et al., 1988). The annual heat budget at Bermuda is not closed locally by air-sea exchange (the dashed line in Fig.2a), therefore, an excess heat flux at the surface is added in our model in order to run stable, multi-year integrations. The surface heat flux function we used to force the model is the solid line in Fig.2a.

Surface salinity was derived using linear interpolation of monthly mean CTD data at the top 8 meter of water column (World Ocean Atlas Levitus94). As shown in Fig.2b, salinity reaches the maximum value of 36.7 during winter and early spring, and the minimum value of 36.4 during summer. The variations in PAR (Fig.2d) were the climatological data from the World Ocean Atlas (Levitus94). The PAR is expressed as a harmonic function with an amplitude of 30 W m^{-2} and centered at 70 W m⁻² on February 28.

Temperature and salinity profiles are initialized with the Levitus94 September data as shown in Figs.3a and 3b, respectively. Biological simulations are initialized with a uniform nitrate concentration of 0.3 mmolN m⁻³ in the mixed layer (0–150 m), increasing linearly below that layer to 6.0 mmolN m⁻³ at 400 m (Fig.3c).

The model equations are solved using the finite difference procedure described by Mellor (1990). There are a total of 27 vertical layers for the 400 m of water column and the grid resolution increases toward the surface. An implicit scheme is used to avoid computational instabilities associated with small grid spacing and a time step of 10 min in numerical integration. The Aselin filter is applied at every time step to avoid time splitting due to the leapfrog scheme. The detailed solution steps are as follows: 1) The physical model is integrated for 5 years. An annual steady state cycle is reached after 3 years of integration. 2) The biological model is coupled with the fifth year solution of the physical model and integrated for 4 years, at the end of which the annual cycle of biological variables is obtained. The depth integrated total nitrogen content over the annual cycle, $N_t = N + A + P + Z + D$, should remain a constant if the equilibrium state is reached.



Fig.2 Annual variations in model surface boundary conditions. a) surface heat flux (solid line) and annual heat budget (dashed line) at Bermuda; b) salinity; c) wind stress; d) photosynthetic available irradiance.



Fig.3 Model initial conditions of a) temperature; b) salinity; c) nitrate.

3 Results

3.1 Response of Upper Layer Physical Structure to Physical Forcing

The annual response of the upper layer physical structure to physical forcing functions is shown in Fig.4. The winter structure is characterized by strong cooling and mixed layer deepening. In February and March the mixed layer depth can exceed 220m with a mixed layer temperature of about 19.5°C and a high eddy diffusivity (Fig.4c). After mid-April, the water column warms up gradually and the mixed layer depth decreases. Due to weak mixing, weak wind, and strong heating in summer, water surface temperature increases up to a maximum value of 27° C, the mixed layer shoals to a less than 10 m depth, and a sharp thermocline at the base of the mixed layer is developed. The characteristics of wind-related and shallow mixed layer are consistent with low values of eddy diffusivity shown in Fig.4c. The autumn period is characterized by a mixed layer depth of 50–75 m and temperature around 22°C, salinity around 36.575, which is followed by a deeper penetration of the mixed layer and cold water mass formation as a result of strong cooling in winter (January and February).



Fig.4 The time and depth variations in a) temperature, b) salinity, and c) eddy diffusion coefficient.

3.2 Response of the Upper Layer Biological Structure to Physical Forcing

The temporal and vertical distributions of five biochemical variables are shown in Fig.5. There are several phases of the biological structure within a year, agreeing with the physical structure of the upper ocean. Due to the deep convection in winter, the surface layer is rich with nutrients entrained from below. The mixed layer nitrogen concentration increases gradually to its maxima in April. As a result of nutrient enrichment and sufficient light availability, the phytoplankton bloom begins to develop in January and reaches the maximum level in March and April. During the same period, the water column is overturned completely and the deepest and coolest mixed layer is established due to strong vertical mixing. The spring phytoplankton grow during March and April until June. The summer and fall periods are characterized by nutrient depletion and low phytoplankton production in the mixed layer. The phytoplankton biomass is low because, with weak convection, the nutrient supply from the nutrient-rich water below the mixed layer is no longer available and the phytoplankton biomass is consumed by herbivore in surface waters. In summer, the stratification and the strong seasonal thermocline inhibit nutrient flux into the shallow mixed layer from below, and therefore prohibit the development of bloom during summer. Nitrate concentrations below the thermocline increase and, together with sufficient light availability, lead to the surface maximum of phytoplankton biomass in the layer between the seasonal thermocline and the base of the euphotic zone during July and August. Remineralization of particulate organic material following degradation of the spring bloom produces ammonium. Part of the ammonium is used in regenerated production and the rest is converted to nitrate through the nitrification process. The yearly distributions of zooplankton and detritus closely follow that of phytoplankton with a time lag of approximately two weeks. The maximum zooplankton concentrations correspond to phytoplankton blooms in spring as well as subsurface phytoplankton maximum in summer.



Fig.5 The time and depth variations in (a) nitrate, (b) ammonium, (c) phytoplankton, (d) zooplankton, and (e) detritus.

3.3 Dynamics of Phytoplankton Blooms

In this section, the main mechanisms controlling the initiation, development and degradation of blooms, as well as the subsurface maximum during summer, are briefly described. First, the relative roles of light and nutrient uptake are considered in the primary production process. The control of phytoplankton growth by either light or nutrient limitation during one year is shown in Figs.6a and 6b. The light limitation function has the opposite structure with values decreasing towards deeper levels (Fig.6a). The relatively high gradient region at about 50–100 m depths separates the low region near the

surface from the high nitrogen limitation region below during summer (Fig.6b). Therefore, the net growth function (Fig.6c), which has the minimum of light and nitrogen limitations, is generally governed by nitrogen limitation near the surface and by light limitation at deeper levels. A subsurface maximum is present at depths of about 50–100 m where both light and nitrogen limitations have moderate values. During summer, this is responsible for the subsurface phytoplankton production.

Fig.6c shows that the net growth function has the highest values within the upper 50 m layer during January and February. But the bloom develops around the end of March as shown in Fig.5c. There are two reasons for the



Fig.6 The time and depth variations in a) nondimensional light limitation function, b) nondimensional nutrient limitation function, and c) net limitation function in one year.



Fig.7 The time and depth variations in a) new production, b) regenerated production, c) zooplankton grazing, and d) time change of phytoplankton.

absence of the bloom generation in mid-winter. First, the amount of phytoplankton biomass at that time is not sufficient to initiate the bloom. Secondly, the relatively strong downward diffusion at surface layer (Fig.4c) counteracts against the primary production and therefore prevents the bloom development. However, as soon as the intensity of vertical mixing diminishes in April, a new balance is established. The time variation term of phytoplankton reaches the maximum at the surface at the beginning of April and the subsurface maximum is attained in the second half of April (Fig.7d). This new balance leads to an exponential growth of phytoplankton in the mixed layer. Soon after the initiation phase, zooplankton grazing (Fig.7c) begins to dominate the system and balance the primary production. This continues until nitrate stocks in the mixed layer are depleted and the nitratebased primary production (new production) (Fig.7a) weakens. At the same time, the rapid recycling of particulate material allows for the ammonium-based regenerated production (Fig.7b), and contributes to the bloom development. The bloom terminates abruptly towards the end of May when ammonium stocks are no longer sufficient for the regenerated production.

The downward diffusion process in the mixed layer is evident with K_h values of greater than $2 \text{ cm}^2 \text{ s}^{-1}$ from January to April (Fig.4c). The termination of convective mixing process in late April is indicated in Fig.4c by a sudden decrease of K_h . As shown further in Figs.4a and 5c, the period of high K_h values is identified with vertically uniform temperature structure of about 19.5°C and the phytoplankton structure of approximately 0.3 mmolN m⁻³. Following the termination of convective overturning, the subsurface stratification begins to establish. As the mixed layer temperature increases by about 0.5°C from 19.5°C to 20°C, the phytoplankton bloom reaches its peak amplitude of 3.5 mmolN m⁻³ within half month.

4 Comparison of Model Results with BATS Observations

The calculated temperature and salinity (Fig.4) compare well with the climatological data (1961–1970) (Fig.8) from Hydrostation S (WHOI and BBSR, 1988; Musgrave *et al.*, 1988) and with the model results of Doney *et al.* (1996). The results clearly reproduce the deep winter convective depth, shallow summer mixed layer, and sharp seasonal thermocline found in the data. The seasonal salinity cycle also generally agrees with climatology, showing the peak salinity during the winter convection period and the formation of a fresh surface layer during summer. A sub-surface salinity maximum S>36.6 appears in both the calculated results and the observations.

The biological model is driven with a uniform nitrate concentration of 0.3 mmolN m⁻³ in the mixed layer (0–150 m), increasing linearly below the mixed layer to 6.0 mmolN m^{-3} at the 400 m depth. A direct comparison between the coupled model and the data is difficult because the BATS dataset shows significant interannual variability and its available record is relatively short for

generating a true biological climatology. The climatological forcing has a likely effect on the model results of reduced variability in deep convection during winter, causing homogenization of properties through the winter mixed layer depth, and weakening individual bloom events driven by short-term variability. Monthly nitrate climatologies for the first four years of BATS (1988– 1992) are shown in Fig.9 (Knap *et al.*, 1991, 1992, 1993). The climatologies are useful for revealing the general characteristics of the model results, but quantitative comparisons should be limited to more robust features of biological seasonal cycle. The calculated nitrate field agrees reasonably well with the BATS data. Winter surface con-



Fig.8 Climatological (1961–1970) seasonal cycles of a) temperature and b) salinity at Hydrostation S (WHOI and BBSR, 1988; Musgrave *et al.*, 1988; Doney *et al.*, 1996).



Fig.9 Climatological seasonal cycle of nitrate for the first 4 years (1988–1992) of the BATS record (unit: mmol m⁻³) (Knap *et al.*, 1991, 1992, 1993).

centrations are about 0.2 mmolN m^{-3} and the summer nitracline depth is at about 100–125 m depths. The approximately uniform concentration in the deep winter mixed layer gradually increases through summer due to the remineralization of detritus. However, the calculated nitrate values are generally lower than the observed.

5 Sensitivity Studies

A series of experiments are carried out to analyze the model sensitivity to some adjustable parameters (Table 1). The experiments and the parameter values for each experiment are listed in Table 3. The experiments show that if one parameter affects phytoplankton distribution, it will have more influence on zooplankton. The important parameters that affect the structure of phytoplankton, and therefore that of zooplankton, are the maximum phytoplankton growth rate σ_m , the phytoplankton death rate m_p , the light extinction coefficient k_w , the nitrate half-saturation constant R_n , the maximum herbivore grazing rate r_g , the herbivore death rate m_h , the herbivore excretion rate μ_h , the herbivore assimilation efficiency γ_h , the herbivore half-saturation constant R_g , the detrital remineralization rate ε , and the detrital sinking rate w_s . The bloom structure is not sensitive to the phytoplankton self-shading coefficient k_c , the ammonium half-saturation constant R_a , the photosynthesis efficiency parameter a, and the ammonium oxidation rate. Three examples are presented below to examine how the biological parameters affect phytoplankton and zooplankton.

Table 3 Parameters for sensitivity experiments ('df' stands for the default value. The default is used if no value is shown in a box)

	σ_m	m_p	k_w	<i>k</i> _c	R_a	R_n	R_g	а	γ_g	m_h	μ_h	γ_h	Ε	Ω	Ws
df	.75	.05	.03	.03	.8	1	.3	.05	.21	.01	.05	.8	.05	.03	.5
A1	1.5														
A2	.375														
B1		.1													
B2		.025													
C1			.06												
C2			.015												
D1				.06											
D2				.015											
E1					1.6										
E2					.4										
F1						2									
F2						.5									
G1							.6								
G2							.15								
H1								.1							
H2								.25							
I1									.42						
12									.105						
J1										.02					
J2										.005					
K1											.1				
K2											.025				
LI												1.6			
L2												.4			
MI													.1		
MZ													0.25	06	
NI N2														.06	
N2														0.15	2
															5
02															0.25

5.1 Sensitivity to the Extinction Coefficient of PAR Value (Default $k_w=0.03 \text{ m}^{-1}$)

As shown in Table 3, two experiments were carried out with the PAR extinction coefficient of $k_w=0.06 \text{ m}^{-1}$ and $k_w=0.015 \text{ m}^{-1}$ in experiments C1 and C2, respectively. Increasing the default k_w value intensifies the distribution of phytoplankton and zooplankton towards the sea surface (Figs.10c and 10d). Decreasing its value, the distributions of phytoplankton and zooplankton are stretched into deeper water (Figs.11c and 11d). In the model, the phytoplankton growth rate depends on the minimum of nutrient limitation and light limitation. It is governed by the nitrogen limitation near the surface and by the light limitation at deeper levels. Comparing the light limitation in Fig.10a with Fig.6a, it can be seen that the light limitation in experiment Cl decreases in the entire water column except for the very near surface. The most striking difference is that in the default case, the 0.05 contour of light limitation is located between 100 and 150 m depths, while it is between 66 and 84 m in experiment Cl. The subsurface maxima of net limitation decreases and shifts towards the sea surface except in winter, which corresponds to the squeezing of the phytoplankton distribution towards the sea surface. The zooplankton, which feeds on phytoplankton, also moves about 50 m closer to the sea surface than in the default case. The dynamics in experiment C2 is opposite to that in experiment Cl.



Fig.10 The time and depth variations in Expriment Cl of a) the light limitation function, b) the net limitation function, c) phytoplankton, and d) zooplankton in one year.



Fig.11 The time and depth variations in Expriment C2 of a) the light limitation function, b) the net limitation function, c) phytoplankton, and d) zooplankton in one year.

5.2 Sensitivity to the Nitrate Half Saturation Coefficient (Default $R_n=1 \text{ mmolN m}^{-3}$)

If algae are placed in a nutrient medium, the concentration of nutrients decreases over time in the medium as they are incorporated into plant cells. The nutrient uptake rate of algae depends on nutrient concentration in the medium (Valiela, 1995). The nitrate or ammonium uptake rate of phytoplankton has a hyperbolic relationship with the nitrate or ammonium concentration in the environment (Eppley *et al.*, 1969). In the Michaelis-Menten equation, the half saturation constant reflects the relative ability of phytoplankton in using nutrients of low levels and thus may be of ecological significance. In the case of nitrate, nutrient uptake occurs in two steps. First, nutrients are taken into a phytoplankton cell at a rate determined by ambient nutrient concentration. Then, as the concentration inside the cell increases, nutrient is utilized in proportion to internal cellular concentration. If the nitrate uptake of nitrate maybe greatly underestimated because of the preference for ammonium by different algae. The half saturation constant is high in more euphotic and nutrient-rich waters, but is low in oligotrophic waters.

Two experiments were carried out with R_n =2mmolNm⁻³ and R_n =0.5 mmolN m⁻³ in experiments Fl and F2, respectively. Increasing R_n in expriment Fl increases the strength and duration of the phytoplankton spring bloom (Figs.12a and 12b). The subsurface maximum of phytoplankton now extends into July, while in the default case it extends into June. However, zooplankton has only weak distribution that spans the period from July to November in the upper 120 m. Opposite results were obtained when the value of R_n was decreased in case F2 (Figs.12c and 12d). The subsurface maximum of phytoplankton now only extends into May while the distribution of zooplankton is much stronger than in experiment F1, spanning the period form March to November.



Fig.12 The time and depth variations of a) phytoplankton and b) zooplankton in case Fl; c) phytoplankton and d) zooplankton in case F2.

5.3 Sensitivity to the Detrital Sinking Rate (Default $w_s=0.5 \text{ m d}^{-1}$)

The sinking rate of particulate organic matter, w_s , is one of the most critical parameters in the model. The appropriate value of w_s for the model is 0.5 m d^{-1} , which implies that the faster sinking and larger particles do not contribute to the processes taking place within the euphotic zone. Greater sinking values of detrital material decrease the detritus and subsequently nitrogen concentrations in the euphotic layer. Figs.13a and 13b show the results with the sinking rate of 3 m d^{-1} , and Figs.13c and 13d 0.025 m d^{-1} . The change in w_s alters the whole biological system drastically. In case O1, $w_s=3 \text{ m d}^{-1}$, and there exists only a weak bloom in April and May (Fig.13a), with almost no zooplankton biomass and detritus in the study area. The euphotic layer is depleted of both ammonium and nitrate accumulated at deeper levels. The case with $w_s = 0.025 \text{ m d}^{-1}$ allows a more than complete remineralization of detrital material before it reaches the lower boundary of the model. The concentrations of phytoplankton and zooplankton are higher with the decrease of w_s , than in the default case as shown in Figs.13c and 13d, especially during winter when the complete water column overturning provides rich supplies of nutrients in the euphotic zone.

6 Conclusions

In this paper, the interaction between physical and biological dynamics in the Sargasso Sea is studied. The response of a five-component ecosystem to the external forcing, including heat flux, wind, and salinity, and sensitivities of the ecosystem to biochemical parameters are investigated.

The model results compare quite successfully with the observations and the calculated results by Doney *et al.* (1996). The base simulation and the sensitivity experi-



Fig.13 The time and depth variations of a) phytoplankton and b) zooplankton in case O1; c) phytoplankton and d) zooplankton in case O2.

ments show a seasonal cycle of physics and biology. In summer, the shallow seasonal thermocline depth and the weak convection inhibit nutrient supplies from the mixed layer below, therefore the concentrations of all biochemical variables are low and limited to a thin surface layer. In winter, the strong vertical mixing and convection bring more nutrients to the euphotic zone. Hence, in March and April, right after winter, phytoplankton feeding upon nutrients reaches its spring bloom level. The bloom of zooplankton, which feeds on phytoplankton, follows the bloom of phytoplankton with a time lag of about two weeks. The results of the sensitivity experiments show that zooplankton is generally more sensitive to the variation of biochemical parameters than phytoplankton. The system is sensitive to all the parameters except the phytoplankton self-shading coefficient, the ammonium half-saturation constant, the photosynthesis efficiency parameter, and the ammonium oxidation rate. For example, a smaller detrital sinking rate and a higher detrital remineralization rate provide higher nutrient concentration in the euphotic zone, while a smaller light extinction coefficient corresponds to stronger and deeper penetrating solar radiation in water column. Both circumstances create a more intense bloom, deeper in space and lasting longer in time.

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