A biophysical model of population dynamics of the autotrophic dinoflagellate Gymnodinium breve

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ABSTRACT: A new model of dinoflagellate diel migratory behavior and population dynamics is presented using the Expanded Eulerian Method (Janowitz & Kamykowski 1999, Ecol Model 118:237–247) and adapting the concept of metabolism-influenced swimming orientation (Kamykowski & Yamazaki 1997, Limnol Oceanogr 42:1189–1202; Kamykowski et al. 1998a, in: Anderson et al. [eds] Physiological ecology of harmful algal blooms, Springer-Verlag, Berlin, p. 581–599; Yamazaki & Kamykowski 2000, Ecol Model 134:59–72). The model is constructed to simulate the observations in a 3 d laboratory mesocosm experiment (Kamykowski et al. 1998b, Mar Ecol Prog Ser 167:105–117; Kamykowski et al. 1998c, J Plankton Res 20:1781–1796) on autotrophic Gymnodinium breve (a red tide dinoflagellate species) under a nutrient-replete condition and in 12 h light:12 h dark cycle. A hypothesis of the acclimations of the Gymnodinium breve swimming orientation and speed to the internal biochemical and physiological state and external environmental conditions is proposed. A hypothesis proposed by Kamykowski et al. (1998b) on Gymnodinium breve reproduction strategy is tested in the model by considering the 2 daughter cells coming from a parent cell to differ in internal biochemical composition. The model simulations are in good agreement with the observations. Consistent with the observations, the model predicts the surface aggregation of a portion of the population during the light period with decreasing surface aggregation intensity over the 3 d period and approximately uniform vertical distribution of the population through the water column during the dark period as well as the diel convergence and divergence patterns of the mean internal cellular carbon and nitrogen between the surface cells and mid-column cells. As expected, Gymnodinium breve's internal biochemical and physiological states have a strong influence on its migratory behavior and consequently on its population dynamics. By comparison to a simulation with a reproduction strategy producing 2 identical daughter cells, it is shown that the reproduction strategy producing 2 daughters different in biochemical composition appears to be the one adopted by all or at least a large portion of the Gymnodinium breve population in the experiment.

KEY WORDS: Model · Population dynamics · Dinoflagellate · Behavior

INTRODUCTION

The dinoflagellate Gymnodinium breve has been the focus of research for decades, especially in recent years, because its intensive red tides in the Gulf of Mexico, particularly on the West Florida Shelf, have dramatic adverse impacts on the ecology, marine environment, and human activities. Available field and laboratory observations suggest that Gymnodinium breve (Odum et al. 1955, Steidinger & Ingle 1972, Steidinger & Joyce 1973, Heil 1986, Kamykowski et al. 1998b,c, Steidinger et al. 1998) like other dinoflagellates (Eppley et al. 1968, Harrison 1976, Eppley & Harrison 1979, Weiler & Karl 1979, Heaney & Furness 1980, Cullen & Horrigan 1981, Heaney & Eppley 1981, Levandowsky & Kaneta 1987, MacIntyre et al. 1997) exhibits a diel pattern of vertical migration. The vertical migration capability of Gymnodinium breve is speculated to contribute to its massive harmful algal blooms under certain biological and physical conditions (e.g. Heil 1986, Kamykowski et al. 1998b).
In the field, light intensities decrease with depth, and nutrient concentration usually increases with depth. Thus, Gymnodinium breve's observed basic swimming orientation pattern of ascending during the day light period and descending during the dark period indicates a clear survival strategy as related to natural light and nutrient gradients. However, the observations both in situ and in the laboratory have shown much more complex swimming patterns. Under nutrient-replete laboratory conditions (1) G. breve still continuously vertically migrates, aggregating at the surface during the day and dispersing through the water column at night (Heil 1986, Kamykowski et al. 1998b,c); (2) during the light period only a portion of a G. breve population aggregates at the surface, and surface aggregation varies with time and starts to decrease before the onset of the dark period; (3) during the dark period G. breve cells disperse through water column approximately uniformly, though not all the cells at the surface during the light period leave the surface; and (4) some cells deep in the water column start to ascend before the light period. Furthermore, a diel oscillation pattern in the internal cellular biochemical composition between the surface population and the population at depth has been displayed under nutrient replete laboratory conditions (Heil 1986, Kamykowski et al. 1998b).

In situ (1) maximum concentrations of G. breve have been observed at various depths in the water column under bloom conditions (Steidinger & Ingle 1972); (2) on many occasions, even throughout the light period, there are no high surface concentrations (Odum et al. 1955); and (3) persistent surface maxima have even been observed throughout the dark period during the bloom (Heil 1986).


The vertical migrating behavior of autotrophic dinoflagellates has also been studied by means of numerical modeling, but such an endeavor has not been conducted to model the migratory behavior of Gymnodinium breve. Most modeling approaches used in these studies are of the Lagrangian type (e.g. Woods & Onken 1982, Denman & Gargett 1995, Kamykowski & Yamazaki 1997), in which individual dinoflagellate cells are tracked to simulate their behaviors and responses to environmental stimuli. In theory, properly implemented Lagrangian and Eulerian models (see below) will yield the same results, though Lagrangian results would have to be converted to a Eulerian form for comparison with the mostly Eulerian data (taken at fixed positions and times). However, for large populations it becomes impracticable to track all cells present. As an alternative, subsets of the entire population as represented by functional groups may be tracked. The number of subsets can still be quite large. We will consider this point at the end of our model description. Eulerian approaches (e.g., Lande & Lewis 1989) thus far developed are only capable of using the mean cellular properties of cells at any particular spatial location to represent responses of all the cells present. Thus these cells can only act identically according to the mean characteristics, although they may in reality have totally different responses. A field population usually is composed of cells with different biochemical compositions and physiological states and these cells maybe exhibit totally different responses to the same environmental stimuli, depending on many non-linearly related internal cellular factors. Thus the traditional Eulerian approach is limited in application in modeling realistic dinoflagellate population migratory behavior and population dynamics. However, the Expanded Eulerian Method (EEM) developed by Janowitz & Kamykowski (1999) eliminates this limitation by increasing the number of independent variables beyond time and position and has the capability of simulating different behaviors of different cells at any particular spatial location. This approach is discussed in detail below.

Some generic Lagrangian metabolism-influenced orientation models have been developed to simulate migratory behavior controlled by internal cellular biochemical and physiological states (Kamykowski & Yamazaki 1997, Kamykowski et al. 1998a, Yamazaki & Kamykowski 2000). This concept of metabolism-influenced orientation can be incorporated to simulate the migratory behavior of Gymnodinium breve. The purpose of this article is to present a new numerical
model of population migratory behavior and population dynamics, using the EEM approach and adapting the concept of metabolism-influenced swimming orientation (Kamykowski & Yamazaki 1997, Kamykowski et al. 1998a, Yamazaki & Kamykowski 2000). The model is constructed to simulate _G. breve_’s migratory pattern, the dynamics of its internal cellular biochemical composition, and the spatial distribution of its populations under a nutrient replete condition, observed by Kamykowski et al. (1998b,c). The influence of internal cellular biochemical and physiological states on swimming behavior and consequently on the photosynthesis and cell reproduction is the focus of our present model. In this article, the characteristics of the temporal variations of a _G. breve_ population’s distribution and its average internal cellular biochemical distributions in a mesocosm observed in Kamykowski et al.’s (1998b,c) experiment is presented first. Then, the structure of the model itself is introduced. Finally, the simulation results will be presented and compared with the laboratory observations.

**THE LABORATORY EXPERIMENT**

Our model is run to simulate the laboratory experimental observations by Kamykowski et al. (1998b,c) on _Gymnodinium breve_ under a nutrient-replete condition. Because the biological submodels of our model and the associated parameters are constructed and determined particularly for this species according to the observations from this laboratory experiment, the laboratory experiment and its observational results are presented briefly in this section before the introduction of the model itself. This description is a brief version of that given by Kamykowski et al. (1998b,c).

The non-axenic, unialgal stock cultures of _Gymnodinium breve_ Davis (Wilson isolate) were used for the experiment. The experiment was conducted in a 225 l, nutrient-replete (150 µM NO₃-N) water column (155 cm deep) in a temperature-controlled room at 22°C. The mesocosm column was illuminated from above on a 12 h light:12 h dark cycle with ~350 µmol quanta m⁻² s⁻¹ PAR (photosynthetically active radiation) reaching the water surface.

The 3 d experiment started at 06:00 h (lights-on) on the first day and ended at 06:00 h on the fourth day. The experiment was conducted on the daughter cells at 2800 cells ml⁻¹ coming from the quantized parent cells, with the whole population dividing every third day and specifically during the night prior to the first experiment day. The cells were approximately uniformly distributed through the water column at the time of cell division. The quantized growth pattern, obtained by repeatedly isolating the subpopulation that aggregated at the surface in early afternoon, provides an opportunity to study cellular biochemistry using bulk chemical analyses (Kamykowski et al. 1998b) and also a good opportunity for numerical simulation.

During the 3 d experiment, a portion of the column population aggregated at the surface during the light period with the highest surface concentration occurring at noon each day. The strength of the surface aggregation decreased day by day. During the dark period, the column population dispersed approximately evenly through the water column.

The cellular biochemical composition of the cells collected from the surface and the mid-column (~0.75 m below the surface) of the 1.5 m mesocosm showed that the ‘quantized population apparently did not similarly synchronize biochemical composition over the light/dark cycle’ (Kamykowski et al. 1998b). Mid-column values of internal cellular carbon (carbohydrates and especially lipids) were consistently higher than surface values during the light period and nearly equal during the dark period. Mid-column values of internal cellular nitrogen (measured protein) were consistently higher than surface values during the light period, and there was less consistency in the surface to mid-column values during the 3 dark periods. Cellular concentrations of most constituents increased during the 3 d period (Kamykowski et al. 1998b).

Kamykowski et al. (1998b) speculated that the oscillatory pattern in biochemical composition was influenced by the diel biosynthetic patterns and the behavioral redistribution of cells in the water column. They also proposed a theory of a parent-daughter relationship to explain the observed behavioral and biochemical patterns (see Kamykowski et al. 1998b for details): the daughter and the parent may have the same or a different internal cellular biochemical composition, and the 2 daughters from a parent may also have the same or a different one. The different behavioral and biosynthetic patterns between different daughters, which initially may be the same or different, may be the reason for the biochemical divergence observed in the experiment.

**MODEL DESCRIPTION**

A spatially 1-dimensional (the vertical position), time-dependent model of _Gymnodinium breve_ population dynamics is developed here. In this model, the population dynamics are controlled by the migratory behavior and reproduction. Since most laboratory and field measurements are made at fixed points in space, an Eulerian modeling approach for population dynamics yields results which are directly comparable with measurements. The EEM developed by Janowitz & Kamy-
kowsi (1999) is adapted. In the traditional Eulerian approach, individual cells at a given time at any particular spatial location can be identified only by population mean for any cellular characteristic. In the EEM approach, in addition to the traditional time and space variables, other variables such as the biochemical internal states of a cell are added into the list of independent variables. Thus, at any time at any particular spatial location, the subpopulations of cells with different sets of internal states within the total population can be calculated. As a consequence, in the EEM approach, at any particular spatial location the responses to the same external cues of the cells with the different internal states can be different and the total response of the population is the collective response of the individual cells. Symbols and their definitions of variables used in the model are given in Table 1.

**General structure**

In the present model and the original EEM, the basic dependent variable to describe the population is the number of identical cells (i.e., cells having the same internal states) per unit volume, \( C \), at a spatial position \( (x,y,z) \) and at time \( t \). More precisely, we define

\[
C(t,x,y,z,X_{1},X_{2},...,X_{L})dX_{1}dX_{2}...dX_{L}
\]

(1)
as the number of cells per unit volume at \( (x,y,z) \) at time \( t \) with \( X_{l} - dX_{l}/2 < X_{l} < X_{l} + dX_{l}/2 \), where \( X_{l} \) is any internal cellular state to be considered in the model and \( L \) is the number of internal states. Thus, at time \( t \), the total cell concentration per unit volume of physical space, \( C_{\text{Tot}} \), at a particular position \( (x,y,z) \) is the sum of all cells of different internal states at that location,

\[
C_{\text{Tot}}(t,x,y,z) = \int C(t,x,y,z,X_{1},X_{2},...,X_{L})dX_{1}dX_{2}...dX_{L}
\]

(2)

An internal cellular state, \( X_{i} \), of any particular cell in a population may change with time according to its exposure to the environmental conditions and its internal biochemical and/or physiological processes.

The governing equation for the number of identical cells per unit volume in \( x, y, z, X_{1}, ..., X_{L} \) space, \( C \) is

\[
\frac{\partial C}{\partial t} + \frac{\partial (uC)}{\partial x} + \frac{\partial (vC)}{\partial y} + \frac{\partial (wC)}{\partial z} + \frac{\partial (V_{x}C)}{\partial X_{1}} + \frac{\partial (V_{y}C)}{\partial X_{2}} + \cdots + \frac{\partial (V_{L}C)}{\partial X_{L}} = + \text{Gains} - \text{Losses}
\]

(3)

In Eq. (3), \( u, v, \) and \( w \) are the fluid velocity components, \( V_{x} \) is the vertical swimming velocity of a dinoflagellate cell and \( V_{y} \) is the advection velocity (\( dX_{l}/dt \)) of the internal state of a cell, \( X_{l} \) i.e., the time rate of change of that state as experienced by a cell. For example, if the entire population had the value of \( X_{10} \) at some time and an uptake rate for \( X_{1} \) of \( dX_{1}/dt \) (=\( V_{1} \)), then a small time interval, \( dt \), later the value of this internal variable would be \( X_{10} + V_{1}dt \); the population would have moved, or been advected, along the \( X_{l} \) axis with speed \( V_{1} \). In the model, \( z \) is positive upwards with \( z = 0 \) at the surface. The gain term on the right hand side is the rate of increase or decrease in cell number due to reproduction. Mother cells are lost and daughter cells gained. The loss term is set to zero because no natural mortality or grazing pressure is considered in the model. Here, as we simulate a laboratory experiment, we shall neglect turbulence in the model and set \( u, v, w \) to zeros. The submodels used to predict \( dX_{l}/dt \) and their controlling factors will be described below.

In the model, cells having identical internal states are assumed to respond to the same external cues identically, while the cells having different internal state(s) may respond to the same external cues differently. A behavioral or physiological response of a cell, \( R \), can be a function of time \( (t) \), cell position \( (x,y,z) \), internal cellular states \( (X_{i}, i = 1,...,L) \), and external environmental fields \( (E_{j}, j = 1,...,M) \):

\[
R = f(t,x,y,z,X_{1},...,X_{L};E_{1},E_{2},...,E_{M}) = f(t,x,y,z,\vec{E})
\]

(4)

In our present model, the swimming behavior, internal carbon metabolism, nitrogen uptake, photoinhibition, sun-shade acclimation, and cell reproduction are the responses (see below for the details). The different responses of a cell may have their own subsets of \( (X_{i}, E_{j}) \). An external environmental condition, \( E_{p} \), experienced by a cell may vary with time due to both its own temporal change and the cell's motion (fluid motion and swimming) to different spatial locations. The total population response in a unit volume, \( R_{\text{Tot}} \), can be obtained from the individual responses by the integration

\[
R_{\text{Tot}}(t,x,y,z,E_{1},E_{2},...,E_{M}) = \int R(t,x,y,z,X_{1},...,X_{L};E_{1},E_{2},...,E_{M})C(t,x,y,z,X_{1},...,X_{L})dX_{1}dX_{2}...dX_{L}
\]

(5)

To simulate the laboratory mesocosm experiment, here we consider only a 1-dimensional (the vertical position) model and set the water velocity to zero. The governing Eq. (3) becomes

\[
\frac{\partial C}{\partial t} + \frac{\partial (V_{x}C)}{\partial X_{1}} + \frac{\partial (V_{y}C)}{\partial X_{2}} + \cdots + \frac{\partial (V_{L}C)}{\partial X_{L}} = + \text{Gains (reproduction)}
\]

(6)

A finite difference scheme is applied to construct our numerical model. Each independent field, \( X_{i} \), is discretized into a finite number of intervals \( (K_{i}) \) between its minimum and maximum value. Then, in the discrete approach, at time \( t \) the basic dependent variable, i.e.,
<table>
<thead>
<tr>
<th>Symbol</th>
<th>Definition</th>
<th>Value in the model</th>
</tr>
</thead>
<tbody>
<tr>
<td>a</td>
<td>Constant in the formula for $P_{in}$</td>
<td>46.0 µmol quanta m⁻² s⁻¹</td>
</tr>
<tr>
<td>b</td>
<td>Constant in the formula for $P_{in}$</td>
<td>17.0 µmol quanta m⁻² s⁻¹</td>
</tr>
<tr>
<td>C</td>
<td>Number of cells m⁻² in a discrete 'box' in x, y, z, X₁, ... Xₙ space at time t, i.e., C in discrete fashion</td>
<td>Variable, cells m⁻² box⁻¹</td>
</tr>
<tr>
<td>L</td>
<td>Number of cells having the same internal states per unit volume in x, y, z, X₁, ... Xₙ space at a spatial position (x,y,z) and at time t</td>
<td>Variable, cells per unit volume</td>
</tr>
<tr>
<td>CN</td>
<td>Internal cellular carbon (= X₁), not including $C_{protein}$</td>
<td>85.0 pmol C cell⁻¹ (90% of the range)</td>
</tr>
<tr>
<td>CN₂</td>
<td>Threshold of CN for swimming orientation control</td>
<td>87.3 pmol C cell⁻¹ (95% of the range)</td>
</tr>
<tr>
<td>CN₃</td>
<td>Threshold of CN for swimming orientation control</td>
<td>68.4 pmol C cell⁻¹ (60% of the range)</td>
</tr>
<tr>
<td>CN₄</td>
<td>Maximum internal cellular carbon</td>
<td>90.0 pmol C cell⁻¹</td>
</tr>
<tr>
<td>CN₅</td>
<td>Minimum internal cellular carbon</td>
<td>36.0 pmol C cell⁻¹</td>
</tr>
<tr>
<td>CN₆</td>
<td>Net photosynthetic carbon</td>
<td>Variable, pmol C cell⁻¹</td>
</tr>
<tr>
<td>CN₇</td>
<td>Carbon in cellular protein</td>
<td>Variable, pmol C cell⁻¹</td>
</tr>
<tr>
<td>CT</td>
<td>Cell concentration at a particular position (x,y,z) and at time t</td>
<td>Variable, pmol C cell⁻¹</td>
</tr>
<tr>
<td>CTₘ₇</td>
<td>Chlorophyll a concentration in the water</td>
<td>42.5 µg chl a cell⁻¹</td>
</tr>
<tr>
<td>CTₘ₈</td>
<td>Average cellular chlorophyll a concentration</td>
<td>1.60 m</td>
</tr>
<tr>
<td>D</td>
<td>Depth of the model domain</td>
<td>Variable, dimensionless</td>
</tr>
<tr>
<td>DC(j,i)</td>
<td>A cell’s normalized displacement along axis i at its interval j, relative to the interval length, L(j,i), in dt</td>
<td>Variable, the number of cell</td>
</tr>
<tr>
<td>d</td>
<td>A constant for determining swimming speed</td>
<td>0.28, dimensionless</td>
</tr>
<tr>
<td>dt</td>
<td>Time interval</td>
<td>300 and 600 s i.e., 5 and 10 min</td>
</tr>
<tr>
<td>Eₐ</td>
<td>Sun-shade photoacclimation variable (= Xₐ, 13 D PAR exposure)</td>
<td>Variable, [0,190] µmol quanta m⁻² s⁻¹</td>
</tr>
<tr>
<td>Eₐₑ</td>
<td>External environmental fields</td>
<td>Dependent on field</td>
</tr>
<tr>
<td>eₖ</td>
<td>Chlorophyll a-concentration-dependent total apparent light attenuation coefficient</td>
<td>Variable, [0,1], dimensionless</td>
</tr>
<tr>
<td>eₖₘ₇</td>
<td>PAR attenuation coefficient due to water alone</td>
<td>0.1 m⁻¹</td>
</tr>
<tr>
<td>Hₐ</td>
<td>Cumulative photoinhibition (= Xₐ)</td>
<td>Variable, [1, Kₐ]</td>
</tr>
<tr>
<td>Hₚₘ₇</td>
<td>Threshold of swimming orientation control</td>
<td>0.8, dimensionless</td>
</tr>
<tr>
<td>Hₚₘ₈</td>
<td>Instantaneous inhibition</td>
<td>Variable, [0,1], dimensionless</td>
</tr>
<tr>
<td>I</td>
<td>PAR (photosynthetically active radiation) intensity</td>
<td>Variable, 0 (night) or 350 (day) µmol quanta m⁻² s⁻¹</td>
</tr>
<tr>
<td>Iₚₘ₈</td>
<td>PAR-saturated acclimated threshold for inducing instantaneous photoinhibition</td>
<td>Variable, pmol quanta m⁻² s⁻¹</td>
</tr>
<tr>
<td>lₕ</td>
<td>Sun-shade acclimated saturation light intensity</td>
<td>350 µmol quanta m⁻² s⁻¹</td>
</tr>
<tr>
<td>lₚₘ₇</td>
<td>Maximum PAR intensity at the surface</td>
<td>17.5 µmol quanta m⁻² s⁻¹</td>
</tr>
<tr>
<td>lₚₘ₈</td>
<td>PAR threshold for swimming orientation control</td>
<td>20.0 µmol quanta m⁻² s⁻¹</td>
</tr>
<tr>
<td>j</td>
<td>Index of the intervals on the axis of depth z</td>
<td>0, 1, 2, ..., 80</td>
</tr>
<tr>
<td>j₀</td>
<td>Number of intervals on the axis of variable Xₐ, at time t</td>
<td>40, 40, 5, 5 for CN, N, Hₐ, Eₐ, respectively, dimensionless</td>
</tr>
<tr>
<td>j₁</td>
<td>Number of intervals on the depth z</td>
<td>8, dimensionless</td>
</tr>
<tr>
<td>j₂</td>
<td>Half-saturation constant</td>
<td>0.42 µM NO₃-N</td>
</tr>
<tr>
<td>jₙ</td>
<td>Minimum cellular nitrogen quota for protein synthesis to take place</td>
<td>0.75 pmol N cell⁻¹</td>
</tr>
<tr>
<td>L</td>
<td>Number of internal states</td>
<td>Dimensionless</td>
</tr>
<tr>
<td>L(j,i)</td>
<td>Length of the interval j on the axis of variable Xₐ or z</td>
<td>Variable, [6,32.32,3] pmol N cell⁻¹</td>
</tr>
<tr>
<td>N</td>
<td>Internal cellular nitrogen (= X₁)</td>
<td>8.069, dimensionless</td>
</tr>
<tr>
<td>Nₐ</td>
<td>Ratio of the N, and N in a cell</td>
<td>Variable, [5,20] pmol N cell⁻¹</td>
</tr>
<tr>
<td>Nₐmax</td>
<td>Maximum internal nitrogen</td>
<td>23.3 pmol N cell⁻¹</td>
</tr>
<tr>
<td>Nₐmin</td>
<td>Minimum internal nitrogen</td>
<td>6.32 pmol N cell⁻¹</td>
</tr>
<tr>
<td>[NO₃]</td>
<td>External nitrate concentration</td>
<td>150 µM NO₃-N</td>
</tr>
<tr>
<td>[NO₃]ₙ₈</td>
<td>Ambient nitrate concentration threshold for swimming orientation control</td>
<td>1.26 µM NO₃-N</td>
</tr>
<tr>
<td>P</td>
<td>Light-adapted production rate</td>
<td>Variable, pmol C cell⁻¹ h⁻¹</td>
</tr>
<tr>
<td>Pₚₘ₇</td>
<td>Sun-shade acclimated maximum photosynthetic rate</td>
<td>0.25 pmol C cell⁻¹ h⁻¹</td>
</tr>
<tr>
<td>Pₚₘ₈</td>
<td>A constant for determining $P_{in}$</td>
<td>0.67 pmol C cell⁻¹ h⁻¹</td>
</tr>
<tr>
<td>Pₚₙ</td>
<td>A constant for determining $P_{sh}$</td>
<td>0.25 pmol C cell⁻¹ h⁻¹</td>
</tr>
<tr>
<td>Pₚₖₐ</td>
<td>Maximum increment of diel photosynthesis variation</td>
<td>3.33 pmol C cell⁻¹ h⁻¹</td>
</tr>
<tr>
<td>Pₚₘ₉</td>
<td>Dark-adapted photosynthetic rate</td>
<td>Variable, pmol C cell⁻¹ h⁻¹</td>
</tr>
<tr>
<td>R</td>
<td>A behavioral or physiological response of a cell to external cues</td>
<td>Dependent on response or behavior</td>
</tr>
<tr>
<td>Rₚₘ₈</td>
<td>Dark carbon respiration rate</td>
<td>0.333 pmol C cell⁻¹ h⁻¹</td>
</tr>
<tr>
<td>S</td>
<td>Total population response in a unit volume</td>
<td>Variable, [1,10] m⁻³</td>
</tr>
<tr>
<td>S(j,i)</td>
<td>Sign of displacement D(j,i)</td>
<td>+1 or –1, dimensionless</td>
</tr>
<tr>
<td>S_C</td>
<td>Asymptotic swimming speed acclimated to a light intensity of 250 µmol quanta m⁻² s⁻¹</td>
<td>1.0 m h⁻¹ (≈ 278 µm s⁻¹)</td>
</tr>
<tr>
<td>Sₜₚₙ</td>
<td>Light-acclimated maximum swimming speed</td>
<td>Variable, m h⁻¹</td>
</tr>
<tr>
<td>S_Cₘₗ</td>
<td>Scale factor for $E_{in}$</td>
<td>1,85, dimensionless</td>
</tr>
<tr>
<td>Tₐₜ</td>
<td>Time scale of sun-shade acclimated parameters</td>
<td>3 d</td>
</tr>
<tr>
<td>Tₜₘ₈</td>
<td>Cell division time scale</td>
<td>3 d/dv</td>
</tr>
<tr>
<td>Tₜₘ₈</td>
<td>Time</td>
<td>3 d/dv</td>
</tr>
<tr>
<td>t₀</td>
<td>Length of the daylight period</td>
<td>12 h</td>
</tr>
<tr>
<td>u</td>
<td>Fluid velocity component in x coordinate</td>
<td>Variable, m h⁻¹</td>
</tr>
<tr>
<td>uₗₘ₇</td>
<td>Advection velocity of a cell along axis i at the interval j of the axis</td>
<td>Variable, dimensionless</td>
</tr>
<tr>
<td>v</td>
<td>Fluid velocity component in y coordinate</td>
<td>Variable, m h⁻¹</td>
</tr>
<tr>
<td>vₗₘ₇</td>
<td>Maximum nitrate uptake rate</td>
<td>Variable, m h⁻¹</td>
</tr>
<tr>
<td>w</td>
<td>Fluid velocity component in z coordinate</td>
<td>Variable, m h⁻¹</td>
</tr>
<tr>
<td>X</td>
<td>Any internal cellular state</td>
<td>Dependent on state</td>
</tr>
<tr>
<td>x</td>
<td>A horizontal coordinate</td>
<td>Variable, m</td>
</tr>
<tr>
<td>y</td>
<td>Horizontal coordinate orthogonal to x</td>
<td>Variable, m</td>
</tr>
<tr>
<td>z</td>
<td>Vertical coordinate, which is zero at the surface and positive upwards</td>
<td>Variable, m</td>
</tr>
<tr>
<td>Zₜₘ₈</td>
<td>Depth of the upper boundaries of the 3 D PAR exposure reference layers</td>
<td>0.0, 0.32, 0.64, 0.96, 1.28 m, and so</td>
</tr>
<tr>
<td>α</td>
<td>Initial slope of the curve of light acclimated swimming speed increment</td>
<td>0.55 µm² m⁻² pmol quanta⁻¹</td>
</tr>
<tr>
<td>Γ</td>
<td>Time scale of induction and recovery of the photoinhibition</td>
<td>1.0 h</td>
</tr>
<tr>
<td>ϕ</td>
<td>Phase making the maximum value occur at local noon</td>
<td>0 h</td>
</tr>
</tbody>
</table>
the number of the identical cells per unit volume at a spatial location \( \mathbf{C} \) in the governing Eq. (6) is replaced by the number of cells per unit horizontal area \( \mathbf{C} \) in a multi-dimensional ‘box’, a small region of \( z, X_i, \ldots, X_L \) space, constructed by the corresponding intervals of all the independent variables, including the depth,

\[
C(\mathbf{J}) = C(j_1, j_2, \ldots, j_L) = Cdzdx_Xdx_Y\ldots dx_L
\]

Here, \( i \) is the index of axes (i.e., the independent variables), \( j_i \) is the index of interval on axis \( i \), and \( j_z \) is the index of interval on the depth axis \( z \). The cells in a box are assumed to be distributed uniformly in each box and have the values of both the internal states and external conditions as the values at the center of the box. This means that the mid-point of each interval is the location of a grid point. Then, the advective velocity along axis \( i \) of a cell at the \( j \)th interval of the axis \( (j) \) is the function of its internal states and external conditions at the corresponding grid point and is denoted as

\[
V(\mathbf{J}, \mathbf{E}, j_i) = z_1, 2, \ldots, L
\]

and is abbreviated as \( V(j) \) below. The advective velocities must be specified and calculated before computing the cell redistribution along the corresponding axes at the next time step. The submodels for each internal state and its advective velocity are described below. Once the advective velocity of a cell at an interval of an axis, \( j_i \), is determined, the displacement, \( D_i(j_i) \), of the cell in the corresponding box relative to the corresponding width, \( L_i(j_i) \), of the box in a time interval \( dt \) is calculated by

\[
D_i(j_i) = V(j_i)dt/L_i(j_i) \quad j_i = 1, 2, \ldots, K_i \quad i = z, 1, 2, \ldots, L
\]

The total number of intervals, \( K_z \), and interval lengths can vary between axes, and the different intervals of an axis can be different. The change in cell number in the \( j \)th interval of the \( X_i \) axis due to advection, \( DC_i(j_i) \), in time interval \( dt \) is

\[
DC_i(j_i) = -S_i(j_i) \cdot D_i(j_i) \cdot C_i(j_i) + 0.5 \cdot \left[ S_i(j_i - 1) + 1 \right] \cdot C_i(j_i - 1) + 0.5 \cdot \left[ S_i(j_i + 1) - 1 \right] \cdot C_i(j_i + 1) \quad 1 < j_i < K_i
\]

\[
DC_i(1) = -0.5 \cdot \left[ S_i(1) + 1 \right] \cdot D_i(1) \cdot C_i(1) + 0.5 \cdot \left[ S_i(2) - 1 \right] \cdot D_i(2) \cdot C_i(2)
\]

\[
DC_i(K_i) = -0.5 \cdot \left[ S_i(K_i - 1) \right] \cdot D_i(K_i) \cdot C_i(K_i) + 0.5 \cdot \left[ S_i(K_i + 1) \right] \cdot \left[ \left. DS_i(K_i - 1) \cdot C_i(K_i - 1) \right] \right.
\]

\[
S_i(j_i) \text{ in the equations is the sign of the displacement } D_i(j_i) \text{, either +1 or -1. This numerical scheme is equivalent to the ‘upwind’ differencing, which is numerically stable. Eq. (10) is slightly more complex than the standard upwind differencing form as it allows for either positive or negative speeds. Any intention of the cells in the highest or the lowest interval of any axis to move across the upper or lower boundary of the model domain is forbidden, and these cells are retained in the corresponding interval until the advective direction changes. Thus, at time } t + dt \text{, the net change in cell number in a box is the summation of the advective changes related to that box along all the axes,}
\]

\[
C(t + dt,j_1j_2\ldots,j_L) = C(t,j_1j_2\ldots,j_L) + [DC_{1}(j_1) + DC_{2}(j_2) + \ldots + DC_{L}(j_L)]
\]

We note that to satisfy the Courant-Friedrichs-Levy (CFL) criterion for upwind differencing, \( dt \) must be small enough so that the sum of the magnitudes of the displacements is less than 1.

Finally, adding the increase in cell number generated by reproduction into the boxes receiving the fresh daughter cells and subtracting the number of the mother cells from the corresponding reproducing ‘boxes’, the final updated cell number in a box at time \( t + dt \) is obtained, and the cell concentration in the box is calculated by normalizing the box spatial volume to unit spatial volume. We note that in Janowitz & Kamykowski (1999) reproduction was not considered and the 2 internal variables used were exposures to the external nitrate and PAR fields over a 1 d time scale. These 2 variables were taken as proxies for the internal carbon and nitrogen pools. Here we shall eliminate the proxy status for carbon and nitrogen.

### Submodels for cellular biochemical and physiological states

In the Kamykowski et al. (1998b) experiment which we simulate here, the only gradient present in the external conditions is that due to the light intensity decreasing with depth, as the nutrient concentration will be taken as uniform. The observed complexity in the characteristics of the population distribution and especially of the biochemical divergence and convergence between the surface cells and the mid-column cells is impossible to explain by a diel migratory behavior controlled only by phototaxis and geotaxis. A dinoflagellate cell’s internal biochemical and physiological states are thus hypothesized to control its migratory behavior (Kamykowski & Yamazaki 1997, Kamykowski et al. 1998a), growth, and reproduction. Based on the laboratory observations on the relationship between *Gymnodinium breve*’s vertical distribution of internal cellular biochemical composition and population density (Kamykowski et al. 1998b), the model uses the metabolisms of the internal cellular carbon \( X_i \), not including the carbon in the cellular protein) and nitrogen \( X_3 \), the extent of cumulative photoinhibition \( X_3 \), and the cellular sun-shade acclimation state \( X_4 \) as the controlling factors and as the independent variables.
which determine a cell’s swimming behavior, growth, and reproduction. The internal cellular carbon and nitrogen contents are 2 primary and important cellular biochemical constituents. The extent of the cumulative photoinhibition and the cellular sun-shade acclimation state are 2 important physiological conditions determining a cell’s capability and efficiency in using solar energy in photosynthetic processes and thus growth.

We consider first the external fields. The value of PAR at the surface, \( I(0, t) \), is taken as specified. In our simulation we will take this surface value as a constant during the light period and zero during the dark period. The vertical attenuation of PAR is given by

\[
I(z, t) = I(0, t) \cdot \exp \left( - \int_0^z e_k(z', t) \, dz' \right)
\]  

(12)

Here \( I(z, t) \) is the PAR at depth \( z \) (positive upwards with \( z = 0 \) at the surface). To incorporate the self-shading effect in our model, the chlorophyll \( a \)-concentration-dependent total apparent light attenuation coefficient, \( e_k(z, t) \), given by Riley (1956) is used to calculate the light attenuation coefficient,

\[
e_k(z, t) = e_{k0} + 0.054 Chl(z, t)^{1/3} + 0.0088 Chl(z, t)
\]  

where \( e_{k0} \) is the PAR attenuation coefficient due to water alone, typically 0.04 to 0.40 \( \text{m}^{-1} \). A value of 0.1 \( \text{m}^{-1} \) is used in our model. \( Chl(z, t) \) is the chlorophyll \( a \) concentration in the water in units of mg chl a \( \text{m}^{-3} \).

From the observations by Kamykowski et al. (1998b), the range of cellular chlorophyll \( a \) in a cell is from 20.0 through 65.0 pg chl a cell\(^{-1}\). For the purpose of convenience and reducing the number of independent variables, in our model a constant average cellular chlorophyll \( a \) content, \( Ch_l \), of 42.5 pg chl a cell\(^{-1}\) is used for calculating \( Chl(z, t) \) from cell concentration in the water column. A *Gymnodinium breve* cell concentration between \( 10^4 \) and \( 10^6 \) cells \( \text{l}^{-1} \) can bring the second term in Eq. (13) to the order of 0.1 \( \text{m}^{-1} \), and a 1 order of magnitude higher *G. breve* concentration, i.e., between \( 10^5 \) and \( 10^6 \) cells \( \text{l}^{-1} \), can also bring the third term in the formula to the same order. These cell concentrations are within the range of the *G. breve* red tide cell concentration. The external nitrate field, \([\text{NO}_3]\), can be taken to be a specified function of depth and possibly time, though in our present simulation we will take it to be a constant, independent of both.

The first internal variable we shall discuss, though perhaps fourth in overall importance, is the 3 d PAR exposure of a cell, \( E_3 \), i.e., a cell’s sun-shade photoacclimation, which will be taken to determine the cell’s sun-shade acclimated parameters, including maximum photosynthetic rate, saturation light intensity, and PAR threshold for instantaneous photoinhibition induction. The exposure of a cell to the PAR on a decay time scale of 3 d (\( T_3 = 3 \text{ d} \)), \( E_3 (= X_3) \), is used as the cellular sun-shade acclimation state, based on Kamykowski & Yamazaki (1997). The deterministic equation of the 3 d PAR exposure for a cell is

\[
\frac{dE_3}{dt} = \frac{I - E_3}{T_3}
\]  

(14)

The range of the 3 d PAR exposure is between 0 and \( I_{\text{max}}/SC \). \( SC \) is the scale factor for \( E_3 \). For a periodic, sinusoidally time-dependent PAR,

\[
I(0, t) = I_{\text{max}} \sin[(t + \phi)\pi/td]
\]  

(15)

With all negative values of \( I(0, t) \) set to zero, the value of \( SC \) can be calculated by inserting Eq. (15) into Eq. (14), then dividing both sides of the resulting equation by the maximum PAR intensity, \( I_{\text{max}} \), requiring that the value at \( t = 1 \text{ day} \) equals the value at \( t = 0 \), and noting the maximum value of \( E_3 \) (Janowitz & Kamykowski 1999). The same method also applies to periodic PAR with constant intensity during the light period. In this latter case, \( SC \) will be 1.85, or the maximum unscaled value of \( E_3 \) is 0.54 \( I(0) \). A cell which remains at its initial depth and doesn’t swim will have at most an 8% change in \( E_3 \) in the course of a day, while a cell which swims from the bottom to the top of the simulated mesocosm and remains there for 3 d will have a change of 60% of the full range. Neither of these swimming behaviors will be realized in our simulation, and in a 3 d simulation the change in the value for \( E_3 \) for a cell will be a small fraction of the total range.

The scaled 3 d PAR exposure variable ranges from zero through 1, while PAR decreases (roughly) exponentially with depth. If \( N \) uniform intervals are chosen for the 3 d PAR exposure variable, all cells below a depth of \( \ln N/e_k \) would be taken to have the lowest value of this variable. To improve resolution of the domain we choose \( N \) non-uniform spaced contiguous intervals. The water column depth is taken to be split into \( N \) 3 d PAR exposure reference layers (note that the definition of these layers is different from the definition of the vertical position \( z \), which discretizes the water column), with the initial cell amount distributed uniformly with depth yielding a uniform value of \( e_k \), \( e_k \). The scaled 3 d PAR exposure variable is given the value of 1.0 at the top of the \( N \)th layer \( (z = 0) \) and a value of zero at the bottom of the lowest layer. The 3 d PAR exposure values at the top of layers 1 through \( N − 1 \) are then \( \exp(−ek \times ZT_i) \), where \( i = 1, ..., N−1 \), and \( ZT_i = [1 − (i/N)] \times \text{depth} \). These values along with zero and 1 define the boundaries of each interval, with the mid-point value taken as the average of the values at the top and bottom of each of the \( N \) layers. This approach gives equal weight to the variable through the vertical domain.

The second internal state we shall discuss, the cumulative photoinhibition, \( H_c (= X_4) \), is a measure of the
recent (previous few hours) PAR exposure for a cell, or more precisely how much this exposure has exceeded or lagged its $E_3$ value. The photoinhibition model is the same as the model used by Janowitz & Kamykowski (1991) except that the different time scales, $\Gamma$, of induction and recovery can be used. The governing equation for the cumulative photoinhibition, $H_c (= X_3)$, is

$$V_3 = \frac{dH_c}{dt} = \frac{H_t - H_c}{\Gamma}$$  \hspace{1cm} (16)

The time scales, $\Gamma$, for induction ($dH_c/dt > 0$) and recovery ($dH_c/dt < 0$) of photoinhibition may be different. The instantaneous inhibition, $H_c$, at time $t$, is given by

$$H_c = 0 \hspace{1cm} I < I_h$$  \hspace{1cm} (17)

$$H_c = 1.0 - \exp\{-(I - I_h)/I_h\}^3 \hspace{1cm} I \geq I_h$$  \hspace{1cm} (18)

Instantaneous photoinhibition is induced only when instantaneous PAR experienced by a cell exceeds its sun-shade acclimated PAR threshold for instantaneous photoinhibition induction, $I_h$. The ranges of $H_c$ and $H_t$ are both $[0,1]$. We shall take

$$I_h = 2E_3 + 5$$  \hspace{1cm} (19)

If a cell does not swim, it will be essentially uninhibited; the maximum value of photoinhibition for this cell is 0.003 out of 1.0. Cells which swim upwards can reach values of the photoinhibition of nearly 1.0 in a few hours.

The third internal state we will utilize is carbon (less that involved in protein synthesis). The time-dependent photosynthesis submodel is adopted from Kamykowski & Yamazaki (1997). The model incorporates 3 different time scales of photosynthetic response to the time-dependent irradiation, including the photoinhibition on the time scale of few hours, the diel endogenous circadian rhythm corresponding to the diel variation of the irradiation on the time scale of several hours, and the sun-shade photoacclimation on the time scale of few days. The equation for the net photosynthesis of a phytoplankton cell is taken from Denman & Marra (1986) for light-adapted photosynthetic rate,

$$\frac{dC_{\text{photo}}}{dt} = \left[P_{\text{sun}} + H_c\{P_t - P_{\text{sun}}\}\right]1.0 - \exp\left[-\frac{I}{I_h}\right]$$  \hspace{1cm} (20)

where $C_{\text{photo}}$ is the internal cellular carbon produced by the net photosynthesis, $P_{\text{sun}}$ and $P_t$ are the dark- and light-adapted production rate per cell, respectively; $I$ is the PAR, which is a function of time and depth; $I_h$ is the sun-shade acclimated saturation light intensity; $H_c$ is the cumulative photoinhibition, ranging from zero through unity.

The formula to determine $P_{\text{sun}}$ is adapted from Kamykowski & Yamazaki (1997) to include the diel endogenous circadian variation by making the instantaneous maximum photosynthetic rate ($P_{\text{sun}}$) a function of the sun-shade acclimated maximum photosynthetic rate ($P_{\text{ma}}$) increased by a sine-based, time-of-day variable that gives the highest value at local noon with the maximum increment $P_{\text{mc}}$,

$$P_{\text{sun}} = P_{\text{ma}} + P_{\text{mc}}\sin[(t + \phi)\pi/t_h]$$  \hspace{1cm} (21)

In the formula, $t_h$ is the length of the day and $\phi$ is the phase making the maximum value occur at local noon. The determination of $P_{\text{ma}}$, $H_c$, and $I_h$ and specification of the constants, $P_{\text{mc}}$ and $P_{\text{mc}}$, are given below.

Based on Kamykowski & Yamazaki’s (1997) determination method using the 3 d running average, a new formula is constructed here to determine $P_{\text{sun}}$,

$$P_{\text{sun}} = P_{\text{ma}} + P_{\text{mb}}\tanh[(E_3 - a)/b]$$  \hspace{1cm} (22)

where $P_{\text{ma}}$, $P_{\text{mb}}$, $a$ and $b$ are the constants to be determined.

A numerical experiment was conducted to determine the constants in Eqs. (16) & (20)–(22) for the Gymnodinium breve photosynthetic rate, using the data from the experiment by Kirkpatrick et al. (1997) and the experiment by Shanley & Vargo (1993). Although these experimental data were obtained for low-light adapted (60 µmol quanta m$^{-2}$ s$^{-1}$) $G$. breve cells under a low-light experimental condition (maximum PAR of about 270 µmol quanta m$^{-2}$ s$^{-1}$), these data are the only available time series of photosynthetic rate for $G$. breve. The constants determined from these data sets will be extrapolated for the high light regime in our model until further observations become available.

To best match the simulated photosynthetic rate with these observed rates, the constants were determined as follow: 0.25 pmol C cell$^{-1}$ h$^{-1}$ for $P_t$; 0.67 pmol C cell$^{-1}$ h$^{-1}$ for $P_{\text{ma}}$; 0.25 pmol C cell$^{-1}$ h$^{-1}$ for $P_{\text{mb}}$; 3.33 pmol C cell$^{-1}$ h$^{-1}$ for $P_{\text{mc}}$; 46.0 µmol quanta m$^{-2}$ s$^{-1}$ for $a$; 17.0 µmol quanta m$^{-2}$ s$^{-1}$ for $b$; $\Gamma$ of 1.0 h for both the induction and recovery; and

$$I_h = E_3$$  \hspace{1cm} (23)

Finally, the governing equation for the internal cellular carbon, $C_n (= X_3)$, not including the carbon in the cellular protein, is given during the light period by

$$V_l = \frac{dC_n}{dt} = \frac{dC_{\text{photo}}}{dt} - \frac{dC_{\text{protein}}}{dt}$$  \hspace{1cm} (24a)

and during the dark period by

$$V_l = \frac{dC_n}{dt} = -R_m - \frac{dC_{\text{protein}}}{dt}$$  \hspace{1cm} (24b)

where $R_m$ is the dark respiration rate of cellular carbon only in effect during the dark period and $C_{\text{protein}}$ is the internal cellular carbon built into the cellular protein.

The constant value of $R_m$ is chosen as 0.333 pmol C cell$^{-1}$ h$^{-1}$ (Shanley & Vargo 1993). Again, this value was obtained for low-light adapted Gymnodinium
breve cells under a low-light experimental condition with no data available for a high-light regime. In our model, once the protein is built, it, with the associated carbon, always remains as protein in a cell and is not lost out of the cell. Hence, the respiration does not apply to the carbon in the protein and it is excluded from $Cn$ (loss of protein and the carbon in the protein could be accommodated by a higher uptake rate, but was an unnecessary complication with no supporting data available). The range of the cellular carbon in our model is between the minimum 36.0 ($Cn_{\text{min}}$) and the maximum 90.0 ($Cn_{\text{max}}$) pmol C cell$^{-1}$ (Heil 1986). An estimate of the cost of motility to a dinoflagellate cell in terms of the total cell energy budget is less than 5% (Raven 1983), so it is neglected in our model. The expression for $dC_{\text{protein}}/dt$ is discussed below following the discussion on nitrogen uptake.

The final internal variable is the total nitrogen content of a cell. Assuming that the only external nitrogen resource is nitrate and that there is no nitrogen loss from the cell (again, loss of protein and the carbon in the protein could be accommodated by a higher uptake rate, but was an unnecessary complication with no supporting data available), the governing equation for the internal cellular nitrogen, $N (= X_2)$, is the Michaelis-Menten equation of nitrogen uptake,

$$V_2 = \frac{dN}{dt} = \frac{V_{\text{max}} \cdot [\text{NO}_3(z)]}{K_N + [\text{NO}_3(z)]} \quad (25)$$

Here, the half-saturation constant, $K_N$, of 0.42 µM N (Steidinger et al. 1998) is used; $[\text{NO}_3(z)]$ is the external nitrate concentration and can vary with depth, $z$. In this laboratory-based simulation, this depth variation of nitrate is not involved, but the capability is modeled for future application of the model to field situations (Liu 2000).

The range of the internal nitrogen pool of a cell not measured as protein, $N_i$, is approximately between 5 ($N_{i_{\text{min}}}$) and 20.0 pmol N cell$^{-1}$ ($N_{i_{\text{max}}}$) based on preliminary data collected by D.K. and the range of its nitrogen-in-measured-protein is between 1.32 and 3.30 pmol protein-N cell$^{-1}$ obtained by converting the range of the measured cellular protein (Kamykowski et al. 1998b) to the range of nitrogen-in-measured-protein using the fact that the weight of nitrogen in protein is about 20% of the weight of the protein. The estimated nitrogen-in-measured-protein is low compared to total cell nitrogen because of a conservative estimate of nitrogen:carbon by weight (DiTullio 1993) and because the measured protein under nutrient replete conditions probably represents only a portion of the total protein (Dortch et al. 1984, DiTullio 1993). Since this component does not critically affect the operation of the model, in the present model, modeled protein is scaled to the measured protein in Kamykowski et al. (1998b) although it is considered a low estimate compared to total protein. Hereinafter, the term protein is used to represent the scaled protein. As more information is gained on protein of Gymnodinium breve, the protein value can be easily adjusted. The sum of these 2 forms of nitrogen counts for all the internal nitrogen in the modeled cell with the range from 6.32 ($N_{\text{min}}$) to 23.30 pmol N cell$^{-1}$ ($N_{\text{max}}$). This generally agrees with the range of the internal cellular nitrogen measured by Heil (1986), which was approximately from 3.3 to 15.4 pmol N cell$^{-1}$. For convenience, the percentage of the nitrogen-in-protein in the total internal cellular nitrogen in our model is considered as a constant, and 13.1% is used to calculate nitrogen-in-protein from the total internal cellular nitrogen for the purpose of calculating the rate of internal carbon being built into cellular protein, which will be discussed below.

The value of $V_{\text{max}}$ is determined using the following method. Because the range of the internal cellular nitrogen pool is between 6.32 and 23.30 pmol N cell$^{-1}$, the poorest possible daughter cell has 6.32 pmol N cell$^{-1}$ nitrogen in the cell and the richest possible daughter cell has 16.98 pmol N cell$^{-1}$. Because the measured division interval ($T_{\text{div}}$) is 3 d for the quantized Gymnodinium breve cells in the experiment under nutrient-replete conditions and there is essentially no division of the Gymnodinium breve cells occurring during the 3 d experiment period, we assume that the poorest cells and the richest cells are all capable of filling up their internal nitrogen pool to be division-ready in 3 d. Dinoflagellates usually divide during a period of few hours prior to sunrise (D.K. pers. obs.); hence, in our model, a 3 h period prior to sunrise on each day is set as the reproduction period (see ‘Cell reproduction strategy’). The initial time of our model simulation is at 03:00 h, i.e., the starting time of the reproduction period, assuming that all the cells are fresh new daughters at that moment (see ‘Initial conditions and forcing of the simulation’). Thus, we assume that poorer cells have higher nitrogen uptake rate and richer cells have lower nitrogen uptake rate and that the poorest cells take 72 h to fill up their internal nitrogen pool to 23.30 pmol N cell$^{-1}$ (at the starting time of the reproduction period 3 d later) and the richest cells take 52 h to fill up the pool (1 h after the reproduction period 2 d later to prevent these cells dividing on the third day). Although light/dark modulation of nutrient uptake is likely (Cullen 1985) and nitrogen uptake rate may be higher during the light period and lower during the dark period (Flynn et al. 1996, Flynn & Flynn 1998), this feature is neglected here until more information on Gymnodinium breve is available. Under these assumptions, an exponential nitrogen uptake rate as a function of the internal cellular nitrogen is proposed here:

$$V_{\text{max}} = 5.46 e^{-0.186N} \quad (26)$$
Note that the units of $V_{\text{max}}$ given by this expression are pmol N cell$^{-1}$ h$^{-1}$. The average nitrogen uptake rate for the poorest daughter cells is about 0.236 pmol N cell$^{-1}$ h$^{-1}$ (i.e., the average rate for the cells to fill 16.98 pmol N cell$^{-1}$ deficit in 3 d) and the rate for the richest daughter cells is about 0.122 pmol N cell$^{-1}$ h$^{-1}$ (i.e., the average rate for the cells to fill 6.32 pmol N cell$^{-1}$ deficit in 3 d). The mean value of these 2 average nitrogen uptake rates is about 0.179 pmol N cell$^{-1}$ h$^{-1}$ and is close to the maximum nitrate uptake rate of 0.14 pmol N cell$^{-1}$ h$^{-1}$ reported by Steidinger et al. (1998) for G. breve under nutrient replete conditions.

Finally, the formula used for carbon lost to protein synthesis in Eq. (24) is

$$\frac{dC_{\text{protein}}}{dt} = 0.814 \cdot V_{\text{max}} \left( 1 - \frac{K_q}{0.869 \cdot N} \right)$$  \hspace{1cm} (27)

Since this protein is a constant percent of total nitrogen, $N$, the latter is used in model output comparisons to the laboratory data. See Liu (2000) for a detailed derivation of this equation.

In the model we use the mid-point value of any interval to represent the values of all the cells in that interval. Thus it is necessary to point out how we determine the minimum and the maximum value of each independent variable in a discrete fashion. For internal cellular carbon and nitrogen, we set the observed minimum and maximum values at the mid-points of the first and last intervals, respectively, on the corresponding variable axes. The minimum and maximum values of these variables in the model are set half a grid interval lower and higher than the observed minimum and maximum values, respectively. For photoinhibition, because the inhibition of the cells in the highest box and the lowest box need to be 1 and 0, respectively, we set the mid-point values of the first and the last intervals to zero and 1, respectively; thus the modeled minimum and the maximum values are smaller and higher than zero and 1, respectively. For 3 d PAR exposure, the lower boundary of the lowest interval is set to zero and the upper boundary of the highest interval is set to 1. This completes the specification of the internal advective velocities.

### Cell swimming behavior

The migratory behavior consists of 2 separate but interrelated components, swimming speed and swimming orientation. They are summarized in Table 2 with the descriptions and formulae given in Appendix 1. Unlike the advective velocities given previously, which were based on earlier independent experiments, our hypothesized swimming rules, while reasonable, are based to a great extent on the experiment we seek to simulate. Indeed, one objective of this work is to develop swimming rules which replicate the previously described experimental results. Thus we may view this numerical experiment as one in model calibration rather than model verification. The swimming

<table>
<thead>
<tr>
<th>Independent variable ranges for swimming orientation control</th>
<th>Swimming direction</th>
<th>Number of corresponding equation for swimming speed</th>
</tr>
</thead>
<tbody>
<tr>
<td>During the light period (i.e., between 06:00 and 18:00 h)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>If $H_c &lt; 0.8$ and</td>
<td></td>
<td></td>
</tr>
<tr>
<td>$C &lt; 0.6$</td>
<td>Up</td>
<td>(A2)</td>
</tr>
<tr>
<td>$0.6 \leq C &lt; 0.95$, and $I(z) &lt; 17.5$</td>
<td>Up</td>
<td>(A2)</td>
</tr>
<tr>
<td>$0.6 \leq C &lt; 0.8$, and $I(z) \geq 17.5$, $N \geq 0.9$</td>
<td>Rest</td>
<td>–</td>
</tr>
<tr>
<td>$0.8 \leq C &lt; 0.95$, and $I(z) \geq 17.5$, $N \leq 0.9$</td>
<td>Down</td>
<td>(A4)</td>
</tr>
<tr>
<td>$C \geq 0.95$</td>
<td>Down</td>
<td>(A4)</td>
</tr>
<tr>
<td>If $H_C \geq 0.8$</td>
<td>Down</td>
<td>(A3)</td>
</tr>
<tr>
<td>During the dark period between 18:00 and 00:00 h</td>
<td></td>
<td></td>
</tr>
<tr>
<td>All</td>
<td>Down</td>
<td>(A5)</td>
</tr>
<tr>
<td>During the dark period between 00:00 and 04:00 h</td>
<td></td>
<td></td>
</tr>
<tr>
<td>$[\text{NO}_3] &lt; 3.0K_N (= 1.26)$</td>
<td>Down</td>
<td>(A5)</td>
</tr>
<tr>
<td>$[\text{NO}_3] \geq 3.0K_N (= 1.26)$</td>
<td>Rest</td>
<td>–</td>
</tr>
<tr>
<td>During the dark period between 04:00 and 06:00 h</td>
<td></td>
<td></td>
</tr>
<tr>
<td>$C &lt; 0.1$ or $N &gt; 0.9$</td>
<td>Up</td>
<td>(A7)</td>
</tr>
<tr>
<td>All others</td>
<td>Rest</td>
<td>–</td>
</tr>
</tbody>
</table>

Table 2. Hypothesized swimming rules. The threshold values of $H_c$, $C$, and $N$ are fractions of the ranges between the minimum and maximum values of the respective variables. PAR intensity [i.e., $I(z)$] is expressed in µmol quanta m$^{-2}$ s$^{-1}$, and $[\text{NO}_3]$ and $K_N$ are expressed in µM NO$_3$-N.
rule parameters were tuned to give reasonable results, but we have not tried to finely tune these parameters. This was done as some of the observed variables, e.g., lipids and protein, are only measures of those predicted, e.g., internal carbon and nitrogen. Lipids are emphasized compared to carbohydrates (Cullen 1985) because they appear to be dominant storage component in Gymnodinium breve (Kamykowski et al. 1998b).

The formulas for determining the swimming speed in our model are developed based on the laboratory observations on Gymnodinium breve (Heil 1986, Kamykowski et al. 1998b,c). The formulations by Kamykowski & McCollum (1986) and Kamykowski et al. (1988) for several other dinoflagellate species, which are considered comparable to *G. breve*, are also utilized, as not all corresponding observations have been made on *G. breve* itself. The parameters in the formulas are expected to vary among species. The parameters obtained for Gyrodinium dorsum by Kamykowski & McCollum (1986) are used for *G. breve* in our model except that the maximum swimming speed of *G. breve* is taken as 1 m h\(^{-1}\) (Heil 1986). The swimming speed given by Kamykowski et al. (1988) model is temperature, light, and buoyancy acclimated and simplified to be incorporated into our model. Four different time regimes, each with different swimming rules are postulated. Here we give a qualitative description of the rules (more details and formulas are given in Appendix 1 and Table 2): (1) During the light period, cells which are very high in carbon or are highly photoinhibited descend. Cells which are not highly photoinhibited will ascend or descend, primarily based on carbon content. Cells low in carbon ascend, the lower the carbon content the higher the speed, as will cells at moderate content but at low light intensities. Cells at moderate carbon levels and high light intensities will rest if their nitrogen content is high but descend, seeking nutrients, if their nitrogen content is low. (2) During the second period, from ‘sunrise’ (lights-out) to midnight, all cells descend to find nutrients. However, the descent speed decreases for cells at higher internal nitrogen levels or higher external nitrate locations. The descent speed decreases most rapidly for small increases in nitrogen content when the nitrogen content is at 0.9 of the full range. (3) During the period from midnight to 04:00 h, cells at low external nitrate locations continue to descend while others rest. (4) During the final period from 04:00 h to ‘sunrise’ (lights-on), cells which are very low in carbon or very high in nitrogen ascend, while all others rest. The transition points, i.e., the values of the variables at which changes in behavior occur, were chosen to obtain reasonable results; this is consistent with the calibration role of this work.

**Cell reproduction strategy**

The final element of our model is reproduction. Kamykowski et al. (1998b) proposed 2 hypotheses (dynamic temporal mechanism and less dynamic multi-cell mechanism) to explain the possible reasons for the divergence in the internal biochemical composition between the cells at the surface and the cells at mid-column during the light period in their experiment. In the dynamic temporal mechanism, 2 daughter cells coming from a parent cell are the same biochemically, and all the parent cells are the same as each other. Time lags for different cells to reach the surface may cause the difference in cell behavior and consequently the divergence in the biochemical composition. On the other hand, in the less dynamic multi-cell mechanism, 2 daughter cells coming from a parent cell may be different and parent cells may be different from each other. The behavior difference between cells of different types may be the reason for the measured divergence in the biochemical composition. Kamykowski et al. (1998b) further stated, ‘in reality, a continuous spectrum of different types of cells may exist.’ Hence, 2 reproduction strategies are chosen for our model run: the least equal-daughter reproduction strategy and the equal-daughter reproduction strategy. By ‘the least equal daughter reproduction strategy,’ we mean one daughter (the poor daughter) gets the minimum amounts of internal carbon and nitrogen and the other (the rich daughter) gets the remainder of each constituent. By ‘the equal daughter reproduction strategy,’ we mean both daughters get the same amounts of these internal cellular biochemical constituents. The simulation results of the 2 different reproduction strategies will be compared with the observations.

Dinoflagellates usually divide during a period of few hours around sunrise. In our model, a 3 h period prior to sunrise is set as the reproduction period. Reproduction is taken to occur when a cell’s internal nitrogen is at its maximum, i.e., 23.30 pmol N cell\(^{-1}\) [\(N_{\text{max}}\)] and its internal cellular carbon pool exceed 90% of the range, i.e., 85 pmol C cell\(^{-1}\) [\(C_{\text{cell}}\)]. The division threshold of internal cellular carbon at 90% of the range is to allow the occurrence of the division of high carbon cells after 9 h overnight carbon loss due to dark respiration and protein synthesis. The daughter cells are assumed to keep the same values of the cumulative photoinhibition and the 3 d PAR exposure as their parents.

This completes the model description. Thus, given a concentration of cells at an initial time, Eq. (6) is solved by the algorithm given in Eqs. (9) & (10). The generalized velocities, which are functions of the external fields and internal independent variables are given in Eqs. (14), (16), (24) & (25) and in Appendix 1.
The above model can be used in a Lagrangian manner. Starting at its initial position and internal states, the changes in location and internal states of a single cell can be treated using the equations given above. However, as the experiment encompasses $4.5 \times 10^6$ cells, it is not practical to track each cell. A smaller number of cells could be tracked; each cell would represent a specified number of cells with nearly the same initial position and internal states of the tracked cell. Given the specified sensitivity of the swimming speed to internal states, the range of internal states that a single cell could represent would be quite small. Two cells starting at the same location and differing by only 1 pmol internal carbon might find themselves vertically 1 m apart at the end of 3 d. As this is on the order of the vertical extent of the mesocosm, the range in carbon represented by a tracked cell would have to be less than 1 pmol internal carbon. A series of numerical experiments would be necessary to find the appropriate number of cells to be tracked. The Eulerian approach utilized here is relatively easy to implement.

**INITIAL CONDITIONS AND FORCING OF THE SIMULATION**

The model configuration for simulating the 3 d experimental observations by Kamykowski et al. (1998b,c), which was briefly described earlier, is given in this section, and the model results and the comparison with the observations are presented in the next section. The model results presented here are for 3 different cases. In the first 2 cases, the model is run for 3 d using the 2 different reproduction strategies, first the least equal daughter reproduction strategy and second the equal daughter reproduction strategy, introduced in the last section. Because the laboratory observations by Kamykowski et al. (1998b,c) were over a 3 d period, our simulations also focus on a 3 d period. A comparison with the published data of Kamykowski et al. (1998b,c) will be undertaken. In the third case, a 10 d run using the least equal daughter reproduction strategy with the same initial conditions is conducted to investigate how the population and the average vertical distribution of its internal cellular biochemical composition may evolve over a longer period under the hypotheses developed in our model.

For the 3 d cases, the model run starts from 03:00 h on the first day and ends at 06:00 h on the fourth day. For the 10 d case, the model run starts from 03:00 h on the first day and ends at 06:00 h on the eleventh day. The initial time is set at 03:00 h instead of 06:00 h as in the experiment because our initial cells are assumed to all be fresh new daughter cells and the simulations (shown in the next section) show that the overwhelming majority of the reproducing cells divide at the beginning time of the 3 h reproduction period, i.e., at 03:00 h.

The external environmental conditions and initial conditions are set as closely as possible to the corresponding conditions in the experiment. The depth of the model domain is 1.60 m. The external nitrogen concentration is set to a constant 150 µM NO₃-N independent of depth and time. Under this external nitrate concentration, the calculated nutrient uptake rate is about 99.72% of the potential internal cellular nitrogen determined maximum uptake rate (from Eq. 25). A constant 350 µmol quanta m⁻² s⁻¹ PAR is on a 12 h light:12 h dark cycle, with light hours from 06:00 to 18:00 h on each day. Under 350 µmol quanta m⁻² s⁻¹ PAR, the light acclimated swimming speed, $S_{accli}$ is 1.009 m h⁻¹ and during the dark period it is 0.75 m h⁻¹. The initial cell concentration at 03:00 h on the first day is 2800 cells ml⁻¹ uniformly distributed through the water column. All these cells are the fresh daughter cells newly derived from the parent cells at a density of 1400 cells ml⁻¹. The parent cells’ internal nitrogen amounts are all at the maximum, and these parent cells are uniformly distributed over 90 through 100% of the internal carbon range. The initial biochemical compositions of these daughter cells are distributed from their parents according to the corresponding reproduction strategies considered in the simulations. The initial photoinhibition is set to zero for all the cells because the simulation starts at 03:00 h, when the PAR has been zero for 9 h.

In the model, equal grid spacing is used for all the axes except the 3 d PAR exposure. The numbers of grid points for the different axes are different: 8 grid points for depth, 40 for carbon, 40 for nitrogen, 5 for photoinhibition, and 5 for 3 d PAR exposure. All the grid points are set at the centers of the corresponding discrete intervals, and any cell in any interval is considered to have the state at its mid-point (the grid point). All the cells in any interval are uniformly distributed through the interval. The 3 d PAR exposure variable axis is discretized into 5 non-uniform intervals by setting the upper boundaries of the five 3 d PAR exposure reference layers at 5 evenly spaced depths, $Z \in [0, 0.32, 0.64, 0.96, 1.28]$ m, with an additional end point set at infinity below the surface. Then the mid-point values of these intervals are determined by the approach discussed in the section of model description earlier. The scale factor (SC) of the 3 d PAR exposure calculated by the method mentioned earlier is approximately 1.85 for the constant illumination of 350 µmol quanta m⁻² s⁻¹. The 3 d PAR exposure determined by this method is from zero though about 190 µmol quanta m⁻² s⁻¹ in our model configuration. The 3 d PAR exposure values at the 5 grid points and the related sun-
shade acclimated parameters are given in Table 3. The initial 3 d exposure values of all the cells in the population are set arbitrarily to the value of 28.70 µmol quanta m\(^{-2}\) s\(^{-1}\), which is the intermediate (i.e., the third interval) in the five 3 d PAR exposure intervals ranging from 4.10 µmol quanta m\(^{-2}\) s\(^{-1}\) to 138.0 µmol quanta m\(^{-2}\) s\(^{-1}\) (Table 3).

With these settings of the grid spacings and the corresponding advection speeds, to meet the CFL numerical stability criterion and to reduce the computing time, 2 time steps are used in our model. First a 5 min time step is used to update the population due to advective processes in nitrogen, photoinhibition, and swimming. Using this updated population, the advective change in population due to carbon and sun-shade acclimation is updated using a 10 min time step and the advective changes due to nitrogen, photoinhibition, and swimming are computed using another 5 min time step. The total advective changes due to all variables then define a new population at the end of the 10 min interval. Since there are 40,000 distinct internal states (or types of cells) in each of 8 layers, far too much data is accumulated to be retained. Therefore, each hour (every 6 full time steps) only the number of cells in each layer and for the cells in each layer, the population mean values and standard deviations of each internal state are retained in a computer file.

### SIMULATION RESULTS AND DISCUSSION

In this section we shall compare model predictions with the observations given in Kamykowski et al. (1998b,c). For a more complete discussion and more details see Liu (2000). We first briefly discuss the experimental data to which our simulated results will be compared. Fig. 1, redrawn from Kamykowski et al. (1998c), gives the relative fluorescence (proportional to cell density) as a function of depth and time. These data are based on fluorescence profiles performed every 2 h. Fig. 2, redrawn from Kamykowski et al. (1998b), gives cell density, lipid/cell, and protein/cell information based on 6-hourly samples at the surface and at 0.75 m depth. The lipid (measured protein) will be compared to simulated carbon (nitrogen) content. Observations will now be compared with the 3 d runs of Cases 1 and 2.

#### Case 1: A three-day simulation using the least equal daughter reproduction strategy

Initially, 5.6 \(\times\) 10\(^8\) cells are assigned to each of the 8 layers. The initial box numbers for the 3 d PAR exposure and cumulative photoinhibition, as discussed above, are \(j_{E_i} = 3\) and \(j_{H_{C_i}} = 1\) for all cells. One half the cells have \(j_{N} = 1\) and \(j_{C_{N}} = 1\), the poor daughters. One
eighth of the initial population in each layer is assigned to each of 4 boxes with $j_N = 26$ and $j_{Cn} = 15, 16, 17, 18$, the rich daughters. The range in carbon in the daughters corresponds to the range in carbon of the mothers, from 90 to 100% full. We now turn to the results.

Fig. 3 shows the simulated cell concentrations as a function of time and depth. The pattern in this figure may be compared with the fluorescence pattern in Fig. 1. Cell concentration values will be discussed below. Strong surface aggregations during the light period with the maximum surface cell concentration occurring around noon are present in the data and the model results, with diminishing maximum concentrations in both from the first to the second to the third day. This pattern is easy to understand from the perspective of our model. Ascent speeds are taken to decrease as the carbon content of the cell increases. Initially all cells are low in carbon, which produces relatively high ascent speeds in Day 1. Due to photosynthesis during Day 1, the carbon content of cells increases by the start of Day 2 and ascent speeds diminish; this process continues as we move to the light period of Day 3. Photoinhibition induced descent also plays a role in this day-to-day decrease of surface maximum concentrations. A model run (not shown) with photoinhibition induced descent turned off shows increased surface aggregation of 2, 18 and 48% over the Case 1 results at noon on Days 1, 2, and 3, respectively. Thus photoinhibition plays an increasingly important role as time goes by. This can be explained as follows: in Day 1, strong carbon deficit induced surface concentrations increase the near-surface values of the decay constant for PAR, $ek(z,t)$, and while ascending cells experience higher PAR values and get slightly photoinhibited. Their photoinhibition is less than it would be without self-shading. In subsequent light periods, the carbon-
induced surface aggregation diminishes as does the effect of surface aggregation on $e_k(x,t)$ and cells can become more photo-inhibited, leading to increased descent and reduced surface aggregation as time progresses. As in the observations, during the dark period the cell concentration (Figs. 3 & 4A) is fairly uniform through the water column from the surface through to the bottom; however, the number of cells remaining at the surface during the dark period increases over the 3 d. The simulated daily surface aggregation starts before the light period, as in the experiment, by ascent of very low carbon or very high nitrogen cells (see Appendix 1 for the thresholds). We turn next to a comparison of biochemical components at 2 depths.

Fig. 4 shows the simulated cell concentration, carbon content, and nitrogen content at 4 depths as a function of time. We shall focus on only 2 of these depths, the solid line (0.1 m), which we will compare to the open circles (surface) in Fig. 2, and the dashed line (0.7 m), which we shall compare to the filled circles (mid-column) in Fig. 2. First we consider cell concentration, the top panels in Figs. 2 & 4A. During the final 2.5 d both surface and depth data and calculations are in excellent agreement. During the first full day the predicted surface concentrations in the light period are too high (about 50%) and during the dark period are too low. The predicted concentrations at depth are too low compared with the data during the first light period but are in good agreement in the subsequent dark period and thereafter. These discrepancies may be associated with the uncertainties associated with prescribing the initial concentration and states of the cells. We have prescribed that there are no mother cells present initially and the 2 types of daughter cells are uniformly distributed with depth. Mother cells actually present and near reproduction would move more sluggishly and thus lower surface concentrations in Day 1 during the light period and raise concentrations at depth. Reproduction of these
cells during the first full dark period would bring their daughters' behavior in line with the remainder of the population. Further, the initial values of $E_3$ were taken to be uniform at the mid-range value, in the absence of any data; lower values would have increased photoinhibition and decreased surface concentrations. Aspects of the experimental procedures may also contribute to the discrepancies. In the experiment 2 l of fluid is withdrawn near the surface every 6 h and fluorescence profiles are obtained throughout the water column every 2 h. The sampling will differentially decrease the number of poor daughters in the column during the light period, and both processes may induce some mixing which might lower high concentrations, especially when concentration gradients are large, as in the first light period. The temporal decrease in the intensity of surface aggregation was also observed in laboratory monitoring of Gymnodinium breve in the water column, where only fluorescence profiling occurred. We next consider a comparison of the internal biochemical constituents.

We have no direct data on internal carbon and nitrogen from Kamykowski et al. (1998b) and use the measured cellular lipid and protein contents (the bottom 2 panels of Fig. 2) as proxies for these variables respectively. Note that quantitative comparison of the C and N budgets with the measured lipid and protein contents is not possible because the model quotas for C and N are based on Heil (1986) and the measured lipid and protein are based on Kamykowski et al. (1998b). The quantitative discrepancies may be due to different experimental conditions or to systematic losses in the protein determinations. Thus we can only compare patterns of fluctuation. Lipids at depth are consistently higher than surface values. The basic observed pattern is that divergence in properties between the surface cells and the mid-column cells occurs during the light period and that convergence occurs during the dark period, when the cells disperse approximately uniformly through the water column. Values of lipid at depth increase during the day then drop during the night. Surface lipids drop early in the light periods then rise through midnight and drop slightly until the onset of light. The same patterns of variation are seen in the simulated carbon content (Fig. 4B). Measured protein/cell is our only available proxy for total nitrogen/cell. As measured protein/cell is only one component for cellular nitrogen under nutrient replete conditions (Dortch et al. 1984) and may only account for 10 to 15% of the total nitrogen content of the cell as presently derived, the variations of measured protein may only partially represent the variations of total nitrogen content. Measured protein at depth is greater than measured protein at the surface in all but 3 of 13 observations (lower panel of Fig. 2). The predicted

nitrogen (Fig. 4C) at depth is greater than nitrogen at the surface about two-thirds of the run but primarily during the light periods. Unlike measured protein, nitrogen at the surface is greater than nitrogen at depth during the dark periods. This occurs in our model as cells high in nitrogen are prescribed to descend more slowly than nitrogen poor cells in the period between 18:00 h and midnight. This results in an increase of nitrogen/cell in this period as the low nitrogen cells leave the surface more rapidly. Thus the correspondence between nitrogen and measured protein is not as strong as we might like. The variation in the biochemical composition of Gymnodinium breve cells over depth during the light period were also observed in the laboratory experiment by Heil (1986): cells at depth have higher carbon and nitrogen than the surface cells, consistent with the present simulation. There was no observation available from Heil (1986) for the dark period.

While our model predicts no reproduction for most of the simulated period, some population increase is observed in the simulation at the end of the third full dark period, though less than might be expected. From a simple calculation, we can determine that the rich daughter cells, 50% of the initial population, should have been able to absorb sufficient nitrate and, in the presence of high PAR values and no photoinhibition, create enough carbon to reproduce at the end of the third dark period; however, our simulation indicates that only 12% do reproduce then. Even this diminished reproduction is sufficient to account for the sudden drop in properties/cell; the number of cells increases by 12% in a short time but the total amount of carbon and nitrogen does not. This small population increase results in a 11% decrease in properties/cell averaged over the entire mesocosm near the end of the simulation. The causes for a diminished reproductive population are 2-fold. First, surface aggregation, which results in large increases of ek near the surface, results in diminished PAR intensities from the surface to mid-depth compared with intensities which would prevail if the cells were dispersed. This directly diminishes carbon production as specified in Eq. (20). Further, cumulative photoinhibition values greater than 0.48 in Days 2 and 3 would reduce total carbon (relative to zero photoinhibition) to below the reproductive level. As discussed earlier, a significant fraction of the population reaches values of photoinhibition exceeding 0.8 on Days 2 and 3 as surface population are diminished by this effect. Thus a combination of photoinhibition and surface aggregation can diminish carbon production and hence the reproductive population at the end of the third full dark period. These model formulations require closer inspection in future experiments. For the comparison with the 10 d
simulation (shown below), we note that during the 3 d simulation the minimum PAR at the 1.5 m (the deepest grid point of the water column) is about 8.0 µmol quanta m\(^{-2}\) s\(^{-1}\) during the light period.

**Case 2: A three-day simulation using the equal daughter reproduction strategy**

This case differs from the preceding one because each of the 2 daughter cells is allocated half the carbon and nitrogen content of their mother cell. In this simulation, as before, \(5.6 \times 10^8\) cells are assigned to each layer in the vertical. One quarter of these cells were each assigned initially to boxes \(j_E = 3, j_H = 1, j_N = 28, j_C = 6, 7, 8, 9\). The results of this simulation are shown in Fig. 5, which shows cell concentration as a function of depth and time, and Fig. 6, which gives cell concentration, mean carbon content, and mean nitrogen content for 4 layers as a function of time.

The vertical population distribution pattern (Figs. 5 & 6A) is quite similar to the preceding case (Figs. 3 & 4A). However, the simulated divergence pattern in carbon (Fig. 6B) is much less significant for the 3 d period and even in the opposite divergent direction compared to the observations (Fig. 2) on the first day. The equal daughter reproduction strategy has lower initial maximum carbon values than the rich daughter cells of Case 1. The absence of the mid-column peak in carbon and the higher internal carbon at the surface than at mid-column on the first day (Fig. 6B) occur because no cells have enough carbon to descend during the first light period. In Case 1 the rich daughter cells do descend on Day 1, causing the carbon peak at mid-column. The smaller initial carbon difference between cells, which arises from the small difference between parent cells, may account for a pattern in carbon between the surface cells and the mid-column cells less divergent than in Case 1 over the whole simulation period.

The observed significant divergence pattern in nitrogen (Fig. 2) between the surface and mid-column cells during the light period is not reproduced for the first 2 d by the model with the equal daughter reproduction strategy (Fig. 6C). In the absence of numerical diffusion the cell population can be spread over 2 intervals along the nitrogen axis. For most of the 3 d simulation, all layers have nearly the same mean nitrogen/cell content. For the second full dark period, surface values of nitrogen/cell exceed these values at depth. At the start of this dark period, the nitrogen level is at 90% of the full range. At this value of internal nitrogen, the specified descent velocity is fortuitously most sensitive to small changes in nitrogen level. A 2.5% increase in nitrogen (1 interval) decreases the descent speed by 30%. This leads to cells low in nitrogen leaving the upper layer more rapidly than those higher in nitrogen, thus increasing the nitrogen/cell for those left in the surface layer. Numerical diffusion along the nitrogen axis also enhances this effect somewhat by extending the range of nitrogen by about 1 or 2 intervals in 40. In the equal daughter case, had the nitrogen level at the start of the dark period been 5% higher or lower than it actually was, differences in descent velocities would have been diminished, as would the differences in nitrogen/cell between layers. Thus, the predicted nitrogen/cell for this case is generally far more vertically uniform than for Case 1. As measured protein shows definite difference between the surface and mid-column, we might tentatively state the least equal daughter strategy is the one preferred by the cells; a mix between these 2 extreme reproduction strategies is also a possibility.

In this 3 d simulation, the effects of cell division on cell concentration and property/cell distribution have not been manifested, as only a small fraction of the cells divide at the end of the simulation. Further the effects of self-shading associated with surface aggregation, which diminishes the PAR values experienced by cells and hence carbon creation, and the impacts of reduced carbon production on reproduction cannot be ascertained in the 3 d simulation. We now turn to a 10 d simulation, which may give some insight into this process.
Case 3: A ten-day simulation using the least equal daughter reproduction strategy

To further investigate the effect of the self-shading, a 10 d run is conducted for the least equal daughter scheme; the simulations are shown in the Figs. 7, 8 & 9. The 10 d simulation shows that the population finally doubles in size on the tenth day (Fig. 8). The simulated final cell number reaches $1.01 \times 10^{10}$ cells in the water column. The self-shading effect becomes more important day by day as more and more cells aggregate at the surface during the 10 d simulation (Figs. 7A & 9). That more and more cells are retained in the upper reaches of the column where carbon/cell is lowest indicates that self-shading is having an important effect by decreasing carbon that is photosynthesized and hence slowing the reproductive rate. On the tenth day the highest surface cell concentration reaches $2.64 \times 10^7$ cells l$^{-1}$ at 11:00 h with a PAR attenuation coefficient of 15.81 m$^{-1}$ and a PAR intensity of only 72.00 µmol quanta m$^{-2}$ s$^{-1}$ at 0.1 m. The minimum bottom cell concentration drops to $1.92 \times 10^5$ cells l$^{-1}$ at 12:00 h. As a result of the increase in the PAR attenuation coefficient, the depth of the PAR threshold for swimming orientation control for ascent moves upwards in the water column to between 0.1 and 0.3 m on the tenth day. Thus more cells below this depth tend to move upwards from mid-depths. Consequently, even the lowest surface cell concentration is as high as $7.80 \times 10^6$ cells l$^{-1}$, about 29.60% of the previous day’s highest concentration, during the dark period. The highest bottom cell concentration only rises to $3.04 \times 10^6$ cells l$^{-1}$ at midnight between the tenth and eleventh days. The lowest PAR intensity at 1.5 m drops below 1.0 µmol quanta m$^{-2}$ s$^{-1}$ on the ninth day and further drops to 0.67 µmol quanta m$^{-2}$ s$^{-1}$ on the tenth day; this is well below the 3 d PAR exposure at the mid-point of the lowest interval of the five 3 d PAR exposure intervals, i.e., 4.10 µmol quanta m$^{-2}$ s$^{-1}$. Hence, a higher resolution for 3 d PAR exposure may be needed for the simulation though few cells are at these depths. The diel oscil-
Fig. 7. Least equal daughter reproduction strategy. Simulated (A) cell concentration, (B) average internal carbon, and (C) average internal nitrogen at 0.1, 0.7, 0.9, and 1.5m depth levels over a 10 d period.

CONCLUSION

As was our goal in this work, the model simulations reproduce the basic evolving patterns of population distribution and internal cellular carbon and (possibly) nitrogen distribution patterns observed in the Kamykowski et al. (1998b) experiment and also in the Heil (1986) experiment. Our model may capture the most important mechanisms which control Gymnodinium breve swimming behavior and population dynamics under nutrient replete conditions. Our simulation further demonstrates that internal biochemical and physiological states can play significant and determinative roles in G. breve swimming behavior and consequently in the population dynamics and that the least equal daughter reproduction strategy is likely a component of the reproduction strategy which G. breve adopts under nutrient replete conditions.

Ours is the first model to use the Expanded Eulerian Method (EEM) developed by Janowitz & Kamykowski (1999) to simulate observations; the simulation results demonstrate that the EEM approach is a very robust method for simulating phytoplankton population
At the current stage, our model only generically incorporates the metabolisms of the internal cellular carbon and nitrogen as the agents controlling the swimming behavior. More delicate controlling schemes, which look into the details of cellular biochemical composition, can be incorporated into the model in the future. For instance, the metabolisms of carbohydrate, lipid, NO\textsubscript{3}-N, NH\textsubscript{3}-N, free amino acid, DNA, RNA, protein, etc., can be considered to be included into the model individually, instead of just the 2 primary elements, cellular carbon and nitrogen. Lipid content is hypothesized to be highly correlated with cells’ geotaxis (Kamykowski et al. 1998b). Alternately, aspects of the coherent biochemical models like Geider et al. (1996, 1998) or Flynn et al. (1997) may be incorporated.

The hypotheses on the swimming orientation control and swimming speed acclimation proposed in our model can be applied to field populations and field environmental conditions, making the model easy to modify to simulate the field population dynamics under realistic environmental conditions, such as nutrient deplete conditions and the presence of the thermocline, pycnocline, and nutricline. Field PAR intensities at the surface are much higher than the PAR intensity in the Kamykowski et al. (1998b) and Heil (1986) laboratory experiments and vary with time during the day, but this can be readily incorporated into the model. However, data on photosynthesis at PAR levels higher than thus far utilized will be necessary to apply the model in higher light regimes.

Our model will be modified to investigate the red tide dynamics. Gymnodinium breve is common in the Gulf of Mexico at background cell concentrations of <1 \times 10^3 cells l\textsuperscript{-1} (Geesey & Tester 1993). In weeks, it can develop into a bloom of fish-killing concentration (1 to 2.5 \times 10^5 cells l\textsuperscript{-1}), depending on the environmental conditions (e.g. Steidinger 1973, 1975, Steidinger & Haddad 1981, Tester & Steidinger 1997). At 10^5 cells l\textsuperscript{-1} in surface waters, chlorophyll can be detected by satellite sensors, but it isn’t until 1 \times 10^6 cells l\textsuperscript{-1} that the human eye can detect discolored surface water. Cell concentrations have been recorded as high as 1 \times 10^7 cells l\textsuperscript{-1} in Florida water (Tester & Steidinger 1997). The highest surface cell concentration in our present model, reaching at least 9.48 \times 10^6 cells l\textsuperscript{-1} during the 3 d simulation, is a very significant bloom concentration.

Clearly, more observational data, both experimental and field, are required to verify the hypotheses proposed in our model, to further tune the model parameters, and to develop the model to simulate complex field population dynamics under in situ external environmental conditions.

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In the equation, \( S_{\text{accli}} \) (its internal carbon pool fills to certain high levels or until its content to ascend faster. A cell ascends or stays (for those used for cell reproduction strategy). These cells can continue to fill their carbon pool up to 100% full if, as they descend, they are exposed to sufficient PAR, which is the case in our simulations. These descending high-carbon cells may downgrade by losing their internal carbon to protein synthesis if they descend into depths with insufficient PAR intensity and change the orientation to upward direction again during the light period. This descending velocity is specified as

\[
V_z = -S_{\text{accli}} \left[ 0.2 + 0.8 \left( 1 - \frac{\left( \frac{C_n}{K_n} + \frac{[NO_3]}{[NO_3]_{\text{max}}} \right)}{\frac{C_n}{C_{\text{max}}'} - \frac{C_n}{C_{\text{min}}'}} \right) \right]^{1/2}
\]  

(A4)

Because the priority for these descending cells changes to nitrogen uptake, this descending speed is accelerated to the external nitrogen conditions. The higher the surrounding external nitrogen concentration, the slower the descending speed. The cells which have more internal cellular carbon descend faster because they have less demand for PAR for further photosynthesis.

According to the observation that the internal cellular carbon is higher at the mid-column than at the surface (Kamykowski et al. 1998b), a lower carbon threshold, \( C_{\text{thigh}} \), is set at 68.4 pmol C cell\(^{-1}\), which is at 60% of the range. The swimming orientation of the cells having internal carbon between \( C_{\text{thigh}} \) and \( C_{\text{bigh}} \) is controlled by a different set of rules. The internal carbon content of these cells is high but not high enough for cell division, so they still have the priority to build their internal carbon. Gymnodinium breve is a dinoflagellate species capable of having high photosynthetic capacity at low light and adapting to varying light environments (Shanley 1985, Garcia & Purdie 1992), so we assume there may be a common low PAR threshold which most G. breve cells may be comfortable with no matter if these G. breve cells are currently low-light adapted or high-light adapted. Hence, a PAR threshold, \( \theta_{\text{PAR}} \), is introduced for these cells. This PAR threshold in our model is set at 17.5 µmol quanta m\(^{-2}\) s\(^{-1}\) (equal to 5% of the

\[
V_z = -S_{\text{accli}} \left[ 0.2 + 0.8 \left( 1 - \frac{\left( \frac{C_n}{K_n} + \frac{[NO_3]}{[NO_3]_{\text{max}}} \right)}{\frac{C_n}{C_{\text{max}}'} - \frac{C_n}{C_{\text{min}}'}} \right) \right]^{1/2}
\]

(A4)

We note that in our model the index for \( z, \rho_z \) increases downwards so that the displacements along the increasing \( j_z \) axis in Eq. (10) have a sign opposite to the velocities given in Eqs. (A2)–(A7), e.g., a negative velocity corresponds to a positive displacement in the \( j_z \) direction. \( V_z \) in Eqs. (A2)–(A7) has the same meaning as \( V_z \) in Eqs. (3), (6), (8) & (9).

In the case where the cumulative photoinhibition of a cell exceeds the threshold, \( H_{\text{cum}} \), which is set at 0.8 out of 1.0 in our model, the cell descends to avoid a high intensity of PAR. The magnitude of this descending velocity is simply the light acclimated swimming speed given by Eq. (A1),

\[
V_z = -S_{\text{accli}} \left[ 0.2 + 0.8 \left( 1 - \frac{\left( \frac{C_n}{K_n} + \frac{[NO_3]}{[NO_3]_{\text{max}}} \right)}{\frac{C_n}{C_{\text{max}}'} - \frac{C_n}{C_{\text{min}}'}} \right) \right]^{1/2}
\]

(A4)

We note that in our model the index for \( z, \rho_z \) increases downwards so that the displacements along the increasing \( j_z \) axis in Eq. (10) have a sign opposite to the velocities given in Eqs. (A2)–(A7), e.g., a negative velocity corresponds to a positive displacement in the \( j_z \) direction. \( V_z \) in Eqs. (A2)–(A7) has the same meaning as \( V_z \) in Eqs. (3), (6), (8) & (9).
Appendix 1 (continued)

constant 350 µmol quanta m\(^{-2}\) s\(^{-1}\) PAR at the surface in the Kamykowski et al. [1998b] experiment) and is within the range of the light-adapted sun-shade acclimated saturation PAR intensity, \(I_0\), which was discussed earlier. Among these cells, those exposed to lower PAR intensities than the threshold continue to ascend until the PAR intensity exposure equals the PAR threshold, while those located at the depths exposed to higher PAR intensities descend until the depth with the threshold PAR intensity is reached, unless they have both their internal carbon pools below 80% of the range and their internal nitrogen pools above 90% of the range. These cells with both high internal nitrogen and relatively low internal carbon delay descent and stay at their current depths for higher PAR intensities than the threshold PAR intensity. The acclimated descending velocity for the cells in this category is also determined by Eq. (A4).

The ascending or descending cells are bounded by the surface and the bottom of the water column, respectively, and will be trapped at the boundaries until their swimming directions change.

During the dark period, nitrogen uptake holds the highest priority. Between sunset and midnight, cells descend with velocity

\[
V_s = -S_{\text{accl}} \left[ 0.2 + 0.8 \left( 1 - \frac{[\text{NO}_3(x)]}{K_N + [\text{NO}_3(x)]} \right) \right] \times \left( 0.515 \left[ 1 + \tanh \left[ 12 - 13.33 \left( \frac{N - N_{\min}}{N_{\max} - N_{\min}} \right) \right] \right] - 0.069 \right)
\]

(A5)

The external nitrogen concentration at a cell’s location again controls the cell’s descending speed. Cells having lower internal nitrogen content descend faster, and the cells having full internal nitrogen pool barely descend at all, because they have no demand for nitrogen uptake.

After midnight, all the cells stop descending and stay at the depths where they are located unless the external nitrogen concentrations at their depth is lower than a threshold which makes cells’ nitrogen uptake rates reach 75% of the potential maximum rate:

\[
\frac{[\text{NO}_3]_{\text{th}}}{K_N + [\text{NO}_3]_{\text{th}}} = 0.75 \quad \text{or} \quad \frac{[\text{NO}_3]_{\text{th}}}{K_N + [\text{NO}_3]_{\text{th}}} = 3.0 K_N
\]

(A6)

For the \(K_N\) of 0.42 µM NO\(_3\)-N in our simulation, \([\text{NO}_3]_{\text{th}}\) is 1.26 µM NO\(_3\)-N. Cells which are exposed to external nitrogen concentrations lower than the threshold continue to descend with the descending speed specified by Eq. (A5) until they find the depth with the appropriate external nitrogen concentration and stay there or until the time, which is set at 04:00 h (i.e., 2 h prior to lights-on), runs out. These controlling mechanisms may have important biological meaning for field populations because they prevent unnecessary descent during the night under adequate external nutrient conditions so that the cells can expeditiously ascend towards PAR intensities adequate for photosynthesis the next day. Hence the period between midnight and 04:00 h is named ‘the resting period’ in our model.

During the 2 h period prior to lights-on, any cells whose internal nitrogen contents are more than 90% of the range (nitrogen-rich) or internal carbon contents are less than 10% of the range (carbon-poor) ascend in anticipation of the next light period for the next cycle of the photosynthesis. Other cells stay where they are. The ascending velocity during this period is

\[
V_a = 0.25 S_{\text{accl}} \left( \frac{C_{\text{nmax}} - C_n}{C_{\text{nmax}} - C_{\text{pmin}}} \right)
\]

(A7)

After 06:00 h (lights-on), the next cycle begins.

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