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CHESAPEAKE BAY ENVIRONMENTAL MODELING PACKAGE

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Assessing a Ten-Fold Increase in the Chesapeake Bay Native Oyster Population

A Report to the EPA Chesapeake Bay Program

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Abstract

The Chesapeake Bay Environmental Model Package (CBEMP) was used to assess the environmental benefits of a ten-fold increase in native oysters in Chesapeake Bay. The CBEMP consists of a coupled system of models including a three-dimensional hydrodynamic model, a three-dimensional eutrophication model, and a sediment diagenesis model. The existing CBEMP benthos submodel was modified to specifically represent the Virginia oyster, *Crassostrea virginica*. The ten-fold oyster restoration is computed to increase summer-average, bottom, dissolved oxygen in the deep waters of the bay (depth > 12.9 m) by 0.25 g m\(^{-3}\). Summer-average system-wide surface chlorophyll declines by 1 mg m\(^{-3}\). Filtration of phytoplankton from the water column produces net removal of 30,000 kg d\(^{-1}\) nitrogen through sediment denitrification and sediment retention. A significant benefit of oyster restoration is enhancement of submerged aquatic vegetation. Calculated summer-average biomass improves by 25% for a ten-fold increase in oyster biomass. Oyster restoration is most beneficial in shallow regions with limited exchange rather than in regions of great depth, large volume and spatial extent.

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Chapter 1 Introduction

More than twenty years ago, grazing by benthos was implicated as a controlling process on phytoplankton concentration in tidal waters (Cloern 1982, Cohen et al. 1984). Officer et al. (1982) identified criteria for regimes in which benthic control is possible. They were:

1. Shallow water depths in the range of 2 to 10 m;
2. A large and widespread benthic filter feeding population;
3. Partially-enclosed regions of substantial size with poor hydrodynamic exchange;
4. Adequate nutrient supplies; and
5. Regions that show relatively constant and low phytoplankton levels.

A link between decimation of the oyster population and deteriorating water quality in Chesapeake Bay was proposed by Newell (1988). Newell calculated the 19th century oyster population could filter the entire volume of the bay in less than a week and suggested an increase in the oyster population could significantly improve water quality by removing large quantities of particulate carbon. Gerritsen et al. (1994) largely countered Newell’s suggestion. They noted that benthic filter feeders can be dominant consumers in shallow portions of the bay but are suppressed in deeper portions. Processes leading to suppression include hydrodynamic limits and hypoxia. Gerritsen et al. concluded that use of filter-feeding bivalves to improve water quality in large estuaries is limited by the depth and width of the estuary.

Recent research on the role of oysters in Chesapeake Bay has focused on processes by which oysters influence their immediate environment rather than on system-wide effects. Newell et al. (2002) provided experimental evidence that denitrification of nitrogen in oyster feces may enhance nitrogen removal in estuaries. They examined the effect of light on algal biomass and nutrient fluxes at the sediment-water interface and suggested that clarification of the water column by filter feeders may provoke a shift to an ecosystem dominated by benthic primary production. Porter et al. (2004) placed oysters in experimental mesocosms. Their work largely supported the suggestions by Newell et al. (2002). The found that oysters shifted processes to the sediment by decreasing phytoplankton biomass and increasing light penetration to the bottom. Increased
light penetration stimulated microphytobenthos, which diminished nutrient regeneration from the sediments. They found, however, that high bottom shear stress eroded the microphytobenthos and cautioned that, under high bottom shear conditions, nutrient regeneration from the sediments may increase. Most recently, Newell and Koch (2004) employed a model to examine the interactions between oysters, turbidity, and seagrass density. They predicted that restoration of oysters has the potential to reduce turbidity in shallow estuaries and facilitate efforts to restore seagrasses.

Our own interest in oysters stems from the “Chesapeake 2000” agreement. The agreement, signed by the executives of the Commonwealth of Pennsylvania, the State of Maryland, the Commonwealth of Virginia, the District of Columbia, the US Environmental Protection Agency, and the Chesapeake Bay Commission, rededicates the individuals and entities to the “restoration and protection of the ecological integrity, productivity, and beneficial uses of the Chesapeake Bay system.” The agreement sets specific goals including:

- Restore, enhance and protect the finfish, shellfish and other living resources, their habitats and ecological relationships to sustain all fisheries and provide for a balanced ecosystem.

The agreement lists methods to achieve this goal including:

- By 2010, achieve, at a minimum, a tenfold increase in native oysters in the Chesapeake Bay, based on a 1994 baseline.

and

- By 2004, assess the effects of different population levels of filter feeders such as menhaden, oysters and clams on Bay water quality and habitat.

The environmental effects of a ten-fold increase in population of native oysters were assessed by incorporating oysters into the Chesapeake Bay Environmental Model Package (CBEMP), a comprehensive mathematical model of physical and eutrophication processes in the bay and its tidal tributaries. This report is the primary documentation for the assessment.

**The Chesapeake Bay Environmental Model Package**

Three models are at the heart of the CBEMP. Distributed flows and loads from the watershed are computed with a highly-modified version of the HSPF model (Bicknell et al. 1996). These flows are input to the CH3D-WES hydrodynamic model (Johnson et al. 1993) that computes three-dimensional intra-tidal transport. Computed loads and transport are input to the CE-QUAL-ICM eutrophication model (Cerco and Cole 1993) which computes algal biomass, nutrient cycling, and dissolved oxygen, as well as numerous additional constituents and processes. The eutrophication model incorporates a predictive sediment diagenesis component (DiToro and Fitzpatrick 1993).
The first coupling of these models simulated the period 1984-1986. Emphasis in the model application was on examination of bottom-water anoxia. Circa 1992, management emphasis shifted from dissolved oxygen, a living-resource indicator, to living resources themselves. In response, the computational grid was refined to emphasize resource-rich areas (Wang and Johnson 2000) and living resources including benthos (Meyers et al. 2000), zooplankton (Cerco and Meyers 2000), and submerged aquatic vegetation (Cerco and Moore 2001) were added to the model. The simulation period was extended from 1985 to 1994.

Model improvements to address the issues raised by the Chesapeake 2000 Agreement started soon after the agreement was signed. The computational grid was further refined and plans were made to incorporate new living resources into the model. At the same time, regulatory forces were shaping the direction of management efforts. Regulatory agencies in Maryland listed the state’s portion of Chesapeake Bay as “impaired.” The US Environmental Protection Agency added bay waters within Virginia to the impaired list. Impairments in the bay were defined as low dissolved oxygen, excessive chlorophyll concentration, and diminished water clarity. Management emphasis shifted from living resources back to living-resource indicators: dissolved oxygen, chlorophyll, and clarity. A model recalibration was undertaken, with emphasis on improved accuracy in the computation of the three key indicators.

A revision of the CBEMP was delivered in 2002 (Cerco and Noel 2004) and used in development of the most recent nutrient and solids load allocations in the bay. This version of the model is used to examine the impact of the tenfold increase in native oysters. The 2002 CBEMP employs nutrient and solids loads from Phase 4.3 of the watershed model (Linker et al. 2000). (Documentation may be found on the Chesapeake Bay Program web site http://www.chesapeakebay.net/modsc.htm.) Nutrient and solids loads are computed on a daily basis for 94 sub-watersheds of the 166,000 km² Chesapeake Bay watershed and are routed to individual model cells based on local watershed characteristics and on drainage area contributing to the cell. The hydrodynamic and eutrophication models operate on a grid of 13,000 cells. The grid contains 2,900 surface cells ( .4 km²) and employs non-orthogonal curvilinear coordinates in the horizontal plane. Z coordinates are used in the vertical direction, which is up to 19 layers deep. Depth of the surface cells is 2.1 m at mean tide and varies as a function of tide, wind, and other forcing functions. Depth of sub-surface cells is fixed at 1.5 m. A band of littoral cells, 2.1 m deep at mean tide, adjoins the shoreline throughout most of the system. Ten years, 1985-1994, are simulated continuously using time steps of .5 minutes (hydrodynamic model) and .15 minutes (eutrophication model).

References


2 The Oyster Model

Introduction

The ultimate aim of eutrophication modeling is to preserve precious living resources. Usually, the modeling process involves the simulation of living-resource indicators such as dissolved oxygen. For the “Virginia Tributary Refinements” phase of the Chesapeake Bay modeling (Cerco et al. 2002), a decision was made to initiate direct interactive simulation of three living resource groups: zooplankton, benthos, and SAV.

Benthos were included in the model because they are an important food source for crabs, finfish, and other economically and ecologically significant biota. In addition, benthos can exert a substantial influence on water quality through their filtering of overlying water. Benthos within the model were divided into two groups: deposit feeders and filter feeders (Figure 1). The deposit-feeding group represents benthos that live within bottom sediments and feed on deposited material. The filter-feeding group represents benthos that live at the sediment surface and feed by filtering overlying water. The primary reference for the benthos model (HydroQual, 2000) is available on-line at http://www.chesapeakebay.net/modsc.htm. Less comprehensive descriptions may be found in Cerco and Meyers (2000) and in Meyers at al. (2000).

The benthos model incorporates three filter-feeding groups: 1) Rangea cuneata, which inhabit oligohaline and lower mesohaline portions of the system; 2) Macoma baltica, which inhabit mesohaline portions of the system; and 3) Corbicula fluminea, which are found in the tidal fresh portion of the Potomac. These organisms were selected based on their dominance of total filter-feeding biomass and on their widespread distribution. The distributions of the organisms within the model grid were assigned based on observations from the Chesapeake Bay benthic monitoring program (http://www.chesapeakebay.net/data/index.htm). Oysters were neglected in the initial application of the benthos model. The primary reasoning was that oyster biomass was considered negligible relative to the most abundant organisms.

Oysters

The oyster model builds on the concepts established in the benthos model. The existing benthos model was left untouched. The code was duplicated and one portion was modified for specific application to native
oysters, *Crassostrea virginica*. The original model assigned one of the three species exclusively to a model cell. In the revised model, oysters may coexist and compete with the other filter feeders. The fundamental state variable is oyster carbon, quantified as mass per unit area. The minimum area represented is the quadrilateral model cell, which is typically 1 to 2 km on a side. Oyster biomass and processes are averaged over the cell area. Oysters filter particulate matter, including carbon, nitrogen, phosphorus, silica, and inorganic solids from the water column. Particulate matter is deposited in the sediments as feces and pseudofeces. Respiration removes dissolved oxygen from the water column while excretion returns dissolved nitrogen and phosphorus.

Particulate carbon is removed from the water column by the filtration process. Filtration rate is affected by temperature, salinity, suspended solids concentration, and dissolved oxygen. The amount of carbon filtered may exceed the oyster’s ingestion capacity. In that case, the excess of filtration over ingestion is deposited in the sediments as pseudofeces (Figure 2). A portion of the carbon ingested is refractory or otherwise unavailable for nutrition. The unassimilated fraction is deposited in the sediments as feces. Biomass accumulation (or diminishment) is determined by the difference between carbon assimilated and lost through respiration and mortality. Respiration losses remove dissolved oxygen from the water column. Mortality losses are deposited to the sediments as particulate carbon.

The nutrients nitrogen and phosphorus constitute a constant fraction of oyster biomass. Particulate nitrogen and phosphorus, filtered from the water column, are subject to ingestion and assimilation. Assimilated nutrients that are not accumulated in biomass or lost to the sediments through mortality are excreted to the water column in dissolved inorganic form. All filtered particulate silica is deposited to the sediments or excreted to the water column. A fraction (~10%) of filtered inorganic solids is deposited to the sediments. The fraction is determined by the net settling velocity specified in the suspended solids algorithms. The remainder is considered to be resuspended.

The mass-balance equation for oyster biomass is:

$$\frac{dO}{dt} = \alpha \cdot Fr \cdot POC \cdot IF \cdot (1 - RF) \cdot O - BM \cdot O - \beta \cdot O \quad (1)$$

in which:

- $O =$ oyster biomass (g C m$^{-2}$)
- $a =$ assimilation efficiency ($0 < a < 1$)
- $Fr =$ filtration rate (m$^3$ g$^{-1}$ oyster carbon d$^{-1}$)
- $POC =$ particulate organic carbon in overlying water (g m$^{-3}$)
- $IF =$ fraction ingested ($0 < IF < 1$)
- $RF =$ respiratory fraction ($0 < RF < 1$)
- $BM =$ basal metabolic rate (d$^{-1}$)
- $\beta =$ specific mortality rate (d$^{-1}$)
- $t =$ time (d)
The assimilation efficiency is specified individually for each form of particulate organic matter in the water column. The respiratory fraction represents active respiratory losses associated with feeding activity. Basal metabolism represents passive respiratory losses.

**Filtration**

Filtration rate is represented in the model as a maximum or optimal rate that is modified by ambient temperature, suspended solids, salinity, and dissolved oxygen:

\[
Fr = f(T) \cdot f(TSS) \cdot f(S) \cdot f(DO) \cdot Fr_{max}
\]  

in which:

- \(f(T)\) = effect of temperature on filtration rate (0 \(\leq\) f(T) \(\leq\) 1)
- \(f(TSS)\) = effect of suspended solids on filtration rate (0 \(\leq\) f(TSS) \(\leq\) 1)
- \(f(S)\) = effect of salinity on filtration rate (0 \(\leq\) f(S) \(\leq\) 1)
- \(f(DO)\) = effect of dissolved oxygen on filtration rate (0 \(\leq\) f(DO) \(\leq\) 1)
- \(Fr_{max}\) = maximum filtration rate (m\(^3\) g\(^-1\) oyster carbon d\(^-1\))

Bivalve filtration rate, quantified as water volume cleared of particles per unit biomass per unit time (Winter 1978), is typically derived from observed rates of particle removal from water overlying a known bivalve biomass (Doering et al. 1986, Doering and Oviatt 1986, Riisgard 1988, Newell and Koch 2004). Since particle retention depends on particle size and composition (Riisgard 1988, Langdon and Newell 1990), correct quantification of filtration requires a particle distribution that represents the natural distribution in the study system (Doering and Oviatt 1986). Filtration rate for our model was based primarily on measures (Jordan 1987) conducted in a laboratory flume maintained at ambient conditions in the adjacent Choptank River, a mesohaline Chesapeake Bay tributary that supports a population of native oysters. These were supplemented with laboratory measures conducted on oysters removed from the same system (Newell and Koch 2004). Jordan reported weight-specific biodeposition rate as a function of temperature, suspended solids concentration and salinity. The biodeposition rate represents a minimum value for filtration since all deposited material is first filtered. Filtration rate was derived:

\[
Fr = \frac{WBR}{TSS}
\]  

in which:

- \(WBR\) = weight-specific biodeposition rate (mg g\(^-1\) dry oyster weight hr\(^-1\))
- \(TSS\) = total suspended solids concentration (mg L\(^-1\))

Filtration rate was converted from L g\(^-1\) DW h\(^-1\) to model units based on a carbon-to-dry-weight ratio of 0.5.

The observed rates indicate a strong dependence of filtration on temperature (Figure 3) although the range of filtration rates observed at any
temperature indicate the influence of other factors as well. The maximum filtration rate and the temperature dependence for use in the model are indicated by a curve drawn across the highest filtration rates at any temperature:

\[ Fr = F_{r\text{max}} \cdot e^{-KTg \cdot (T-T_{opt})^2} \]  

(4)

in which:

\( F_{r\text{max}} \) = maximum filtration rate (0.55 m\(^3\) g\(^-1\) oyster carbon d\(^-1\))

\( K_{Tg} \) = effect of temperature on filtration (0.015 o\(^C\)-\(^2\))

\( T \) = temperature for optimal filtration (27 o\(^C\))

**Suspended Solids Effects.** The deleterious effect of high suspended solids concentrations on oyster filtration rate has been long recognized although the solids concentrations induced in classic experiments, 10\(^2\) to 10\(^3\) g m\(^-3\) (Loosanoff and Tommers 1948), are extreme relative to concentrations commonly observed in Chesapeake Bay. We formed our solids function by recasting Jordan’s data to show filtration rate as a function of suspended solids concentration (Figure 4). The experiments indicate three regions. Filtration rate was depressed when solids were below \( \sim 5 \) gm m\(^-3\) and above \( \sim 25 \) gm m\(^-3\), relative to filtration rate when solids were between these two levels. The observations suggest oysters reduce their filtration rate when food is unavailable or when filtration at the maximum rate removes vastly more particles than the oysters can ingest. We visually fit a piecewise function to Jordan’s data (Figure 4) supplemented with an approximation of Loosanoff and Tommers’ results:

\[ f(TSS) = 0.1 \text{ when } TSS < 5 \text{ g m}^3\]

\[ f(TSS) = 1.0 \text{ when } 5 \text{ g m}^3 < TSS < 25 \text{ g m}^3\]

\[ f(TSS) = 0.2 \text{ when } 25 \text{ g m}^3 < TSS < 100 \text{ g m}^3\]

\[ f(TSS) = 0.0 \text{ when } TSS > 100 \text{ g m}^3\]

**Salinity Effects.** Oysters reduce their filtration rate when ambient salinity falls below \( \sim 20\% \) of the oceanic value (Loosanoff 1953) and cease filtering when salinity falls below \( \sim 10\% \) of the oceanic value. The form and parameterization of a relationship to describe these experiments is arbitrary. We selected a functional form (Figure 5) used extensively elsewhere in the CBEMP:

\[ f(S) = 0.5 \cdot (1 + \tanh (S - K_{Hsoy})) \]  

(5)

in which:

\( S \) = salinity (ppt)

\( K_{Hsoy} \) = salinity at which filtration rate is halved (7.5 ppt)

**Dissolved Oxygen.** Hypoxic conditions (dissolved oxygen < 2 g m\(^3\)) have a profound effect on the macrobenthic community of Chesapeake Bay. Effects range from alteration in predation pressure (Nestlerode and Diaz 1998) to species shifts (Dauer et al. 1992) to near total faunal depletion (Holland et al. 1977). In the context of the benthos model, effects of hypoxia are expressed through a
reduction in filtration rate and increased mortality. The general function from the
benthos model (Figure 6), based on effects from marine species, was adapted
unchanged for the oyster model:

\[
f(DO) = \frac{1}{1 + \exp\left(1.1 \cdot \frac{DO_{hx} - DO}{DO_{hx} - DO_{qx}}\right)}
\]

in which:

\[
DO = \text{dissolved oxygen in overlying water (g m}^{-3}\text{)}
\]

\[
DO_{hx} = \text{dissolved oxygen concentration at which value of function is one-half (1.0 g m}^{-3}\text{)}
\]

\[
DO_{qx} = \text{dissolved oxygen concentration at which value of function is one-fourth (0.7 g m}^{-3}\text{)}
\]

This logistic function has the same shape as the tanh function used to quantify
salinity effects (Figure 5). The use of two parameters, \(DO_{hx}\) and \(DO_{qx}\), allows
more freedom in specifying the shape of the function than the tanh function,
based on the single parameter \(KH_{soy}\), allows.

**Ingestion**

Oyster ingestion capacity must be derived indirectly from sparse
observations and reports. In the report on his experiments, Jordan (1987) states
“at moderate and high temperatures and low seston concentration (< 4 mg/L)
nearly all biodeposits were feces” (page 54). This statement indicates no
pseudofeces was produced; all organic matter filtered was ingested. Elsewhere in
Jordan (1987) we find that \(\sim 75\%\) of seston is organic matter and the filtration
rate at 4 g seston m\(^{-3}\) is \(\sim 0.1\) m\(^{-3}\) g\(^{-1}\) oyster C d\(^{-1}\) (Figure 4). The ingestion rate
must be at least the amount of organic matter filtered. Conversion to model units
indicates an ingestion rate of:

\[
\frac{4 \text{ g seston}}{m^{-3}} \cdot \frac{0.75 \text{ organic}}{\text{total}} \cdot \frac{g \text{ C}}{2.5 \text{ g seston}} \cdot \frac{0.1 \text{ m}^3}{g \text{ C d}} = \frac{0.12 \text{ g C ingested}}{\text{g oyster C d}}
\]

Tenore and Dunstan (1973) present a figure showing feeding rate and
biodeposition. The difference between feeding and deposition must be ingestion.
The largest observed difference is 19 mg C g\(^{-1}\) DW d\(^{-1}\) or 0.038 g C ingested g\(^{-1}\) oyster C d\(^{-1}\) (utilizing a carbon-to-dry-weight ratio of 0.5). No pseudofeces was
produced during their experiments so the derived ingestion rate is not necessarily
a maximum value.

In reporting on the removal of algae from suspension, Epifanio and
Ewart (1977) noted that large amounts of pseudofeces were produced when algal
suspensions exceeded 12 µg mL\(^{-1}\). These results indicate the amount removed
from the water column when algal suspensions were less than 12 µg mL\(^{-1}\), \(\sim 4\) to
17 mg algal DW g\(^{-1}\) oyster total weight d\(^{-1}\), was ingested. The 15 g total weight
oysters in Epifanio and Ewart’s experiments has a dry weight of 0.27 g (Dame 1972). The minimum ingestion rate is then:

\[
\frac{4 \text{ mg algal DW}}{g \text{ oyster TW}} \cdot \frac{15 \text{ g TW}}{0.27 \text{ g DW}} \cdot \frac{g \text{ oyster DW}}{0.5 \text{ g oyster C}} \cdot \frac{2500 \text{ mg DW}}{2500 \text{ g oyster C ingested}} \cdot \frac{g \text{ algal C}}{TWg} = 0.18 \text{ g C ingested g oyster C d}^{-1}
\]

Analogous unit conversions yield 0.76 g C ingested g⁻¹ oyster C d⁻¹ for a removal rate of 17 mg algal DW g⁻¹ oyster total weight d⁻¹.

Summary of these analyses indicates the order of magnitude for ingestion rate is 0.1 g C ingested g⁻¹ oyster C d⁻¹. The value 0.12 g C ingested g⁻¹ oyster C d⁻¹ was employed in the model based on our evaluation of Jordan’s experiments.

Assimilation

The fraction of ingested carbon assimilated by oysters depends on the carbon source. The assimilation of macrophyte detritus can be as low as 3% (Langdon and Newell 1990) while the assimilation of viable microphytobenthos is 70% to 90% (Cognie et al.). Tenore and Dunstan (1973) observed that oysters assimilated 77% to 88% of a mixed algal culture. Specification of assimilation for the oyster model is shaped by the nature of the eutrophication model. The eutrophication model considers three forms of particulate organic carbon: phytoplankton, labile particulate organic carbon, and refractory particulate organic carbon. Assimilation of phytoplankton is specified as 75%, based on citations above. The labile and refractory particulate organic carbon are detrital components. These are mapped to three G classes of organic matter (Westrich and Berner 1984) employed in the sediment diagenesis model (DiToro 2001). The G1, labile, class has half-life of 20 days. The G2, refractory, class has a half-life of one year. The G3 class is inert within time scales considered by the model. Model labile particulate organic carbon maps to the G1 class and is assigned an assimilation efficiency of 75%, corresponding to phytoplankton. Model refractory particulate organic carbon combines the G2 and G3 classes and is assigned an assimilation efficiency of zero.

Respiration

Two forms of respiration are considered: active respiration, associated with acquiring and assimilating food, and passive respiration (or basal metabolism). This division of respiration is consistent with models of predators ranging from zooplankton (Steele and Mullin 1977) to fish (Hewett and Johnson 1987). Active respiration is considered to be a constant fraction of assimilated food. Basal metabolism is represented as a constant fraction of biomass, modified by ambient temperature:

\[
BM = BMr \cdot e^{KT_{BMR} (T - T_r)}
\]

in which:

BM = basal metabolism (d⁻¹)
BMr = basal metabolism at reference temperature (d⁻¹)
T = temperature (°C)
Tr = reference temperature (°C)
KTbmr = constant that relates metabolism to temperature (°C⁻¹)

The rate of basal metabolism depends on organism biomass (Winter 1978, Shumway and Koehn 1982). The average oyster in Jordan’s (1987) experiments, upon which our filtration rates are based, is 2.1 g DW. Allometric relationships (Shumway and Koehn 1982) indicate basal metabolism for a 2.1 g DW oyster at 20 °C is 0.002 to 0.005 d⁻¹, depending on salinity. A graphical summary presented by Winter (1978) indicates metabolic rate for a 2 g DW oyster at 20 °C is 0.009 d⁻¹. Winter noted a 1 g DW mussel requires 1.5% of its dry tissue weight daily as a maintenance ration. Based on these reports, the value 0.008 d⁻¹ was employed for basal metabolism at a reference temperature of 20 °C. Parameter KTbmr was assigned the value 0.069 °C⁻¹, equivalent to a Q10 of 2, typical of measured rates in oysters (Shumway and Koehn 1982).

The respiratory fraction was assigned through comparison of computed oxygen consumption with metabolism in active oyster reefs (Boucher and Boucher-Rodoni 1988, Dame et al. 1992). The value RF = 0.1 was determined. A comparable value of 0.172 (specific dynamic activity coefficient) was assigned to herbivorous fish in Chesapeake Bay (Luo et al. 2001).

**Mortality**

The model considers two forms of mortality. These are mortality due to hypoxia and a term that considers all other sources of mortality including disease and harvest. Although bivalves incorporate physiological responses that render them tolerant to hypoxia, extended periods of anoxia result in near-extinction (Holland et al. 1977, Josefson and Widbom 1988). Casting the results of experiments and observations into a relationship that quantitatively relates mortality to dissolved oxygen concentration incorporates a good deal of uncertainty in functional form and parameterization. The effect of hypoxia on oyster mortality, adopted from the benthos model, employs two concepts. The first is the time to death under complete anoxia. This time to death is converted to a first-order mortality rate via the relationship:

\[
hmr = \frac{\ln(1/100)}{ttd}
\]

in which:

- hmr = mortality due to hypoxia (d⁻¹)
- ttd = time to death for 99% of the population (14 d)

The mitigating effect on mortality of dissolved oxygen concentration greater than zero is quantified through multiplication by \(1 - f(DO)\) in which \(f(DO)\) is the logistic function that expresses the effects of hypoxia on filtration rate (Equation 6). This functionality increases mortality as dissolved oxygen concentrations become low enough to affect filtration rate (Figure 6). When dissolved oxygen is depleted, filtration rate approaches zero and mortality is at its
maximum. As parameterized in the model, effects on filtration and mortality are negligible until dissolved oxygen falls below \( \sim 2 \) g m\(^{-3}\) (Figure 6). The time to death for 99% of the population exceeds 90 days when dissolved oxygen exceeds 1.4 g m\(^{-3}\) (Figure 7). Under this scheme, some fraction of the oyster population can survive an entire summer of hypoxia provided dissolved oxygen exceeds 1.4 g m\(^{-3}\). No significant portion of the oyster population will survive summer hypoxia for dissolved oxygen concentrations below 1.4 g m\(^{-3}\).

Mortality from all other sources, primarily disease and harvest, is represented by a spatially uniform and temporally constant first-order term. Magnitude of the term is specified to produce various system-wide population levels with the model. The order of magnitude can be derived from Jordan et al. (2002) who reported the 1990 total mortality of “market stock” oysters in northern Chesapeake Bay was 0.94 yr\(^{-1}\) (or 0.0026 d\(^{-1}\)). Of this total, 0.22 yr\(^{-1}\) (or 0.0006 d\(^{-1}\)) was natural mortality. The balance was fishing mortality.

**Nutrients**

Model oysters are composed of carbon, nitrogen, and phosphorus in constant ratios. In the original benthos model (HydroQual 2000), the carbon-to-nitrogen mass ratio of bivalves was set at 5.67:1; the phosphorus-to-carbon mass ratio was 45:1. Composition data for bivalves is not abundant. Calculations by Jordan (1987), based on earlier work by Kuenzler (1961) and Newell (1982), yield a carbon-to-nitrogen mass ratio between 4.8:1 and 6.9:1 and a phosphorus-to-carbon mass ratio of 66:1. The nitrogen composition values encompass the value used in the model. The phosphorus composition value differs from the model but no context exists to judge if the difference is significant.

The oyster model differs substantially from the original benthos model in the way nutrients are assimilated and processed. In the original model, nutrients are assimilated and excreted in constant ratios equivalent to the oyster composition. If assimilated carbon is in excess relative to assimilated nitrogen or phosphorus, the excess carbon is converted to feces and the bivalves are effectively nutrient limited. Computed bivalve growth is:

\[
G = \min \left[ \text{Cassim}, \text{Nassim} \cdot \text{SFCN}, \text{Passim} \cdot \text{SFCP} \right] \tag{9}
\]

in which:

- \( G = \text{bivalve biomass accumulation (g C m}^{-2} \text{ d}^{-1}\) \\
- \( \text{Cassim = carbon assimilation rate (g C m}^{-2} \text{ d}^{-1}\) \\
- \( \text{Nassim = nitrogen assimilation rate (g N m}^{-2} \text{ d}^{-1}\) \\
- \( \text{SFCN = bivalve carbon-to-nitrogen ratio (g C g}^{-1} \text{ N}) \) \\
- \( \text{Passim = phosphorus assimilation rate (g P m}^{-2} \text{ d}^{-1}\) \\
- \( \text{SFCP = bivalve carbon-to-nitrogen ratio (g P g}^{-1} \text{ N}) \) \\

If the carbon-to-nitrogen ratio in assimilated food, Cassim/Nassim, exceeds the ratio in bivalve composition, SFCN, then biomass accumulation is proportional to the rate of nitrogen assimilation. Similarly, when the ratio Cassim/Passim > SFCP, biomass accumulation is proportional to phosphorus assimilation. The
algal phosphorus-to-carbon ratio in the eutrophication model (Cerco and Noel 2004) is 57:1 for spring diatoms and 80:1 for other algae. Since these ratios exceed SFCP, growth of bivalves feeding on algae will be limited by the phosphorus content of the algae rather than the amount of carbon assimilated.

Algal composition does not provide a complete picture of the tendency for nutrient limitation of bivalve growth since modeled bivalves utilize detritus as well as algae. Initial applications of the oyster model indicated, however, that phosphorus limitation of oyster growth did occur. Nutrient limitation was eliminated through two methods. First, oyster phosphorus composition was thinned out; carbon-to-phosphorus ratio was increased to 90:1. More significantly, a mass balance approach to nutrient utilization and excretion was adopted. Biomass accumulation was modeled as carbon assimilation less respiration loss while nutrient excretion was calculated as the amount of assimilated nutrients not required for biomass accumulation.

**Model Parameters**

Parameter values for the oyster model are summarized in Table 1.
<table>
<thead>
<tr>
<th>Parameter</th>
<th>Definition</th>
<th>Value</th>
<th>Units</th>
</tr>
</thead>
<tbody>
<tr>
<td>Frmax</td>
<td>maximum filtration rate</td>
<td>0.55</td>
<td>m³ g⁻¹ oyster carbon d⁻¹</td>
</tr>
<tr>
<td>Topt</td>
<td>optimum temperature for filtration</td>
<td>27</td>
<td>°C</td>
</tr>
<tr>
<td>Ktg</td>
<td>constant that controls temperature dependence of filtration</td>
<td>0.015</td>
<td>°C⁻²</td>
</tr>
<tr>
<td>KHsoy</td>
<td>salinity at which filtration rate is halved</td>
<td>7.5</td>
<td>ppt</td>
</tr>
<tr>
<td>BMR</td>
<td>base metabolism rate at 20 °C</td>
<td>0.008</td>
<td>d⁻¹</td>
</tr>
<tr>
<td>KTbmr</td>
<td>constant that controls temperature dependence of metabolism</td>
<td>0.069</td>
<td>°C⁻¹</td>
</tr>
<tr>
<td>Tr</td>
<td>reference temperature for specification of metabolism</td>
<td>20</td>
<td>°C</td>
</tr>
<tr>
<td>RF</td>
<td>respiratory fraction</td>
<td>0.1</td>
<td>0 &lt; RF &lt; 1</td>
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<tr>
<td>DOₘ₅₀ₓ</td>
<td>dissolved oxygen concentration at which value of logistic function is one-half</td>
<td>1.0</td>
<td>g m⁻³</td>
</tr>
<tr>
<td>DO₃₅₀ₓ</td>
<td>dissolved oxygen concentration at which value of logistic function is one-quarter</td>
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<td>g m⁻³</td>
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<tr>
<td>ttd</td>
<td>time to death for 99% of the population</td>
<td>14</td>
<td>d</td>
</tr>
<tr>
<td>aₐₚₖ</td>
<td>assimilation efficiency for phytoplankton</td>
<td>0.75</td>
<td>0 &lt; a &lt; 1</td>
</tr>
<tr>
<td>aₐₖₒ</td>
<td>assimilation efficiency for labile organic matter</td>
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<td>0 &lt; a &lt; 1</td>
</tr>
<tr>
<td>aₐₖᵣ</td>
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</tr>
<tr>
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<td>maximum ingestion rate</td>
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</tr>
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<td>SFCN</td>
<td>carbon-to-nitrogen ratio</td>
<td>6</td>
<td>g C g⁻¹ N</td>
</tr>
<tr>
<td>SFCP</td>
<td>carbon-to-phosphorus ratio</td>
<td>90</td>
<td>g C g⁻¹ P</td>
</tr>
</tbody>
</table>
References


HydroQual. (2000). “Development of a suspension feeding and deposit feeding benthos model for Chesapeake Bay,” produced by HydroQual Inc. under contract to the U.S. Army Engineer Research and Development Center, Vicksburg MS.


Figure 1. Benthos model schematic.

Figure 2. Processes affecting filtered material.
Figure 3. Effect of Temperature on filtration rate.

Figure 4. Effect of suspended solids on filtration rate.
Figure 5. Effect of salinity on filtration rate.

Figure 6. Effect of dissolved oxygen on filtration and mortality rates.
Figure 7. Effect of dissolved oxygen on time to death for 99% of population.
3 Biomass Estimates

Introduction

The Chesapeake 2000 Agreement calls for a tenfold increase in native oysters in the Chesapeake Bay, based on a 1994 baseline. At the commencement of this study, no estimate of the baseline oyster population existed. Evaluation of the existing population and its distribution was required before the effects of proposed increases could be examined. Since our model is based on mass balance, population estimates took the form of mass rather than number of individuals. We use the terms “biomass” to indicate total weight of oysters e.g. kg C and “density” to indicate weight per unit area e.g. g C m⁻².

Existing Biomass

Virginia

Density estimates for Virginia were provided by Dr. Roger Mann, of Virginia Institute of Marine Science (VIMS), in October 2003. Estimates were based on patent tong surveys. The EPA Chesapeake Bay Program Office (CBPO) provided VIMS with model grid coordinates. Patent tong samples were averaged for each model cell and results were provided as g DW/m². Number of samples per cell varied from 4 to more than 50. Estimates were provided for one to five individual years in the interval 1998-2002. The coefficient of variation (CV, defined as standard deviation/mean) for inter-annual density estimates in individual cells (one or two km on a side) ranged from 0.11 to 1.67 with a median value of 0.69. The CV of the inter-annual total biomass was 0.088. The area of cells containing oysters was 377 km².

Maryland

Biomass and spatial distribution for Maryland were based on the recommendation of Dr. Roger Newell of the University of Maryland Center for Environmental Science. Dr. Newell recommended recent biomass estimates (Jordan et al. 2002) should be uniformly distributed across the historical oyster habitat denoted in the “Yates’ surveys (Yates 1911). The areas and locations of named oyster bars were obtained by the CBPO and bar areas were assigned to model cells. Total area of named oyster bars was 1330 km². Mean biomass for the period 1991-2000, 5.7 x 10⁸ g DW, was obtained from Jordan et al. (2002). A mean density of 0.43 g DW m⁻² (total biomass / total area) was assigned to the
bar area in each model cell. Since the bar area was usually less than the cell area, cell density was adjusted so that biomass per cell matched biomass of bars within the cell. The area of cells containing oysters was 3696 km$^2$.

**Other Filter Feeders**

Examination of the effects of oyster restoration requires consideration of existing filter feeders. Observations from the bay-wide benthic database (http://www.chesapeakebay.net/data/index.htm) were analyzed by HydroQual (2000) as part of the initial benthos modeling effort. The analysis indicates suspension feeding bivalves are distributed primarily in the upper bay and tributaries (Figure 1). Average bivalve densities in the upper bay are commonly an order of magnitude or more greater than the present density of oysters. The arithmetic densities computed by HydroQual are perhaps influenced by a few large density values; median densities might present a more realistic picture. Still, the data support the decision to neglect Maryland oysters in the original benthos model. In the lower bay, the existing oyster density is substantial relative to other bivalves in the lower Rappahannock River and in a limited portion of the James River. The decision to neglect existing oysters in these rivers should be revisited. Recent research (Thompson and Schaffner 2001) indicates polychaete filter feeders, with reported densities $\sim 6$ g C m$^{-2}$, may also play a substantial role in the lower bay.

**Summary**

The oyster density and distribution are distinctly different in the Maryland and Virginia portions of the bay (Figure 2). In the northern, Maryland, portion, lower densities are distributed over a wide area. In the southern, Virginia, portion, high densities are concentrated in limited areas, primarily in the lower James and Rappahannock Rivers. Oyster biomass in Virginia is five times the biomass in Maryland (Table 1) but distributed across an order of magnitude less area. We were puzzled by the limited distribution in Virginia, especially since maps and other information we obtained indicated a wider distribution of lease holdings and restoration areas. We were assured by Dr. Roger Mann that much of the leased area is unproductive and that biomass outside the areas reported to us is negligible. Our estimate of Maryland biomass is roughly half the biomass from two other independent estimates (Table 1). Our estimate of Virginia biomass is three times the biomass from an alternate independent estimate (Table 1).

**Modeled Biomass**

Model oyster density is dynamically computed based on environmental conditions including temperature, dissolved oxygen, salinity, and food supply. The densities are not specified as model inputs. Rather, they must be calculated as a function of model parameters and computed conditions. The calculation, rather than specification, of density ensures that oysters are not placed where conditions do not support their specified density. We initially attempted to calculate target oyster densities through dynamic variation of the mortality function. Mortality in each model cell was adjusted upwards or downwards as
Table 1

<table>
<thead>
<tr>
<th>Source</th>
<th>Maryland, kg C</th>
<th>Virginia, kg C</th>
<th>Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td>This study</td>
<td>287,000</td>
<td>1,170,000</td>
<td>Maryland from Jordan et al (2002). Virginia from Roger Mann (personal communication).</td>
</tr>
<tr>
<td>Newell (1988)</td>
<td>550,000</td>
<td>400,000</td>
<td>Year 2000 exploitable biomass based on skipjack catch per effort</td>
</tr>
<tr>
<td>Uphoff (2002)</td>
<td>570,000</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

calculated density exceeded or fell below specified levels. This process successfully capped density at target levels but many cells would not support existing density or a tenfold increase. The problem originated with the attempt to calculate target densities within individual cells. The calculated conditions in many cells would not support the target densities. Consequently, we switched to a strategy in which a bay-wide target biomass was specified. A uniform bay-wide mortality rate was prescribed that produced the target biomass. The mortality rate was obtained through a trial-and-error process in which various rates were prescribed and the calculated biomass was examined.

The spatial distributions of biomass and density are conveniently examined through aggregation of individual model cells into Chesapeake Bay Program Segments (CBPS). Program segments are subdivisions of the bay determined by mean salinity, natural boundaries, and other features. Our analysis is based on the original (circa 1993) segmentation (Table 2, Figure 3) in which the bay is divided in 35 segments with a median area of 150 km².

Computed density and biomass vary on intra-annual and inter-annual bases (Figure 4). Variations within the annual cycle are largely driven by temperature. Highest densities are computed in late summer and in fall, after a season of filtering at peak rates (Figure 5). Variations from year to year (Figure 6) are largely driven by runoff. Variations in runoff may enhance or diminish computed biomass, depending on local factors. Years with high runoff coincide with large nutrient loads that result in high phytoplankton abundance. The advantages produced by abundant food may be offset, however, by increased anoxia and by sub-optimal salinity.

Baseline Estimates

First-order estimates of the density and biomass of existing bivalve filter feeders can be obtained from the latest application of the CBEMP (Cerco and Noel 2004). This benthos component of this model was originally calibrated to match the observed density in the bay-wide benthic database (HydroQual 2000). Subsequent review (Schaffner et al. 2002) indicated the model tends to over-predict suspension-feeding density in the lower to mid-bay (where density is low) and under-predicts or approximates suspension-feeding density in the upper bay.
and tributaries (where density is high). Still, the model biomass is a useful baseline, especially in the absence of alternate bay-wide abundance estimates.

### Table 2

<table>
<thead>
<tr>
<th>CBPS</th>
<th>Designation</th>
<th>State</th>
</tr>
</thead>
<tbody>
<tr>
<td>CB2</td>
<td>Upper Chesapeake Bay</td>
<td>Maryland</td>
</tr>
<tr>
<td>CB3</td>
<td>Upper Central Chesapeake Bay</td>
<td>Maryland</td>
</tr>
<tr>
<td>CB4</td>
<td>Upper Middle Chesapeake Bay</td>
<td>Maryland</td>
</tr>
<tr>
<td>CB5</td>
<td>Lower Chesapeake Bay</td>
<td>Maryland - Virginia</td>
</tr>
<tr>
<td>CB6</td>
<td>Western Lower Chesapeake Bay</td>
<td>Virginia</td>
</tr>
<tr>
<td>CB7</td>
<td>Eastern Lower Chesapeake Bay</td>
<td>Virginia</td>
</tr>
<tr>
<td>EE1</td>
<td>Eastern Bay</td>
<td>Maryland</td>
</tr>
<tr>
<td>EE2</td>
<td>Lower Choptank River</td>
<td>Maryland</td>
</tr>
<tr>
<td>EE3</td>
<td>Tangier Sound</td>
<td>Maryland - Virginia</td>
</tr>
<tr>
<td>ET4</td>
<td>Chester River</td>
<td>Maryland</td>
</tr>
<tr>
<td>ET5</td>
<td>Choptank River</td>
<td>Maryland</td>
</tr>
<tr>
<td>ET6</td>
<td>Nanticoke River</td>
<td>Maryland</td>
</tr>
<tr>
<td>ET7</td>
<td>Wicomico River</td>
<td>Maryland</td>
</tr>
<tr>
<td>ET8</td>
<td>Manokin River</td>
<td>Maryland</td>
</tr>
<tr>
<td>ET9</td>
<td>Big Annemessex River</td>
<td>Maryland</td>
</tr>
<tr>
<td>LE1</td>
<td>Lower Patuxent River</td>
<td>Maryland</td>
</tr>
<tr>
<td>LE2</td>
<td>Lower Potomac River</td>
<td>Maryland</td>
</tr>
<tr>
<td>LE3</td>
<td>Lower Rappahannock River</td>
<td>Virginia</td>
</tr>
<tr>
<td>LE5</td>
<td>Lower James River</td>
<td>Virginia</td>
</tr>
<tr>
<td>RET1</td>
<td>Middle Patuxent River</td>
<td>Maryland</td>
</tr>
<tr>
<td>RET2</td>
<td>Middle Potomac River</td>
<td>Maryland</td>
</tr>
<tr>
<td>WE4</td>
<td>Mobjack Bay</td>
<td>Virginia</td>
</tr>
<tr>
<td>WT6</td>
<td>Magothy River</td>
<td>Maryland</td>
</tr>
<tr>
<td>WT7</td>
<td>Severn River</td>
<td>Maryland</td>
</tr>
<tr>
<td>WT8</td>
<td>South River</td>
<td>Maryland</td>
</tr>
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</table>

Autumn is the season when individual oysters attain maximum biomass and when most population surveys, on which our estimates are based, are conducted. For comparison with estimates of existing oysters, we averaged the calculated autumn (September – November) bivalve density and biomass from ten years (1985 – 1994). The density comparisons are averaged across total bottom area in each CBPS. The resulting densities are less than individual observations or averages across oyster bars since area not suited for bivalves is included in the average. In most portions of the bay, the calculated density of existing bivalve filter feeders vastly exceeds the estimated density of oysters (Figure 7). Notable exceptions are in the Rappahannock (LE3) and James (LE5) where existing oysters exceed other bivalve filter feeders. Oysters also predominate in two Eastern Shore tributaries (ET8, ET9) and in the lower
western shore of the mainstem (CB6). These segments are characterized by the virtual absence of other bivalves rather than by abundant oysters, however. Biomass comparisons (Figure 8) reflect the density comparisons. Oyster biomass exceeds other bivalve biomass in the lower Rappahannock and James Rivers. Oysters are virtually the only bivalves in the two noted Eastern shore tributaries (ET8, ET9) and in the lower western shore of the bay (CB6).

These comparisons have implications for the overall modeling effort and for the present work. As noted previously, the decision to ignore oysters in the model, until now, was a valid one, with the exception of the lower Rappahannock and James Rivers. For the present study, the model runs with no oysters provide an acceptable baseline for comparison with tenfold population increase since the oysters comprise only a small fraction of filter-feeding biomass throughout most of the bay.

**Tenfold Increase**

The model run for examination of the tenfold population increase, called for in the Chesapeake 2000 Agreement, was determined through a recursive process in which mortality rate was varied until the desired biomass was obtained. Intra- and inter-annual variations in computed biomass made an exact multiplier of existing oyster biomass impossible to obtain. We settled on comparison of computed autumn (September – November) biomass with population estimates since most surveys are conducted in the fall. We compared the mean of ten computed years, 1985-1994, with the estimates of existing population. We settled on a first-order mortality rate of 0.015 d\(^{-1}\), which produced a mean biomass 13-times the estimated existing biomass (Table 3). Biomass in individual years varied by roughly 50% above and below the mean. We refer to this run as the “tenfold increase” although the magnitude and spatial distribution of the increase varies. The southern, Virginia, portion of the bay receives only a fourfold biomass increase while the northern, Maryland, portion increases nearly 50-times. The disparity in multipliers reflects the disparity in initial biomass distribution. An implication of this model run is that, under existing conditions, the northern portion of the bay suffers higher mortality from harvest and disease than the southern portion since imposition of a uniform mortality rate results in greater biomass in the north than in the south. Estimates of the present population indicate the opposite trend. With the tenfold increase, oysters become the dominant filter feeders in the system (Figures 9, 10) although other bivalves predominate in a few segments that provide marginal oyster habitat. Also worth noting is a decline in bivalve biomass, as much as 50%, throughout much of the bay (Figure 11).

**Historical Biomass**

As one part of sensitivity analyses, we computed the biomass of oysters with no mortality from harvest or predation. Limitations to biomass in this run were food availability, respiration, and mortality from hypoxia. The computed biomass (Table 3) that resulted approached the pre-1870 biomass estimated by Newell (1988). This run is documented as an example of improvements that could result from full restoration of historic oyster biomass.
Table 3  
Estimated and Modeled Oyster Biomass, kg C

<table>
<thead>
<tr>
<th></th>
<th>Maryland</th>
<th>Virginia</th>
<th>Total</th>
</tr>
</thead>
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<tr>
<td>Existing, estimated</td>
<td>287,005</td>
<td>1,099,339</td>
<td>1,386,344</td>
</tr>
<tr>
<td>Historic (Newell 1988)</td>
<td></td>
<td></td>
<td>94,000,000</td>
</tr>
<tr>
<td>Tenfold, model</td>
<td>14,107,500</td>
<td>4,374,953</td>
<td>18,482,453</td>
</tr>
<tr>
<td>Historic, model</td>
<td>69,749,506</td>
<td>17,165,230</td>
<td>86,914,736</td>
</tr>
</tbody>
</table>

Equivalent Settling and Removal Rates

The influence of oysters on the environment is a function of their density, filtration rate, and local geometry. The product of density and filtration rate has units of length/time (velocity) and is denoted here as “Equivalent Settling Rate”:

\[
Woys = \frac{1}{A} \cdot \int O \cdot Fr \, dA \quad (1)
\]

in which:

- \(Woys\) = equivalent settling rate (m d\(^{-1}\))
- \(A\) = area over which rate is computed (m\(^2\))
- \(O\) = oyster density (g C m\(^{-2}\))
- \(Fr\) = filtration rate (m\(^3\) g\(^{-1}\) oyster carbon d\(^{-1}\))

The equivalent settling rate can be viewed as the velocity at which particles are transferred from the water column into the oyster bed. Higher velocities indicate more rapid removal. However, the distance to be covered (depth) affects removal as well as velocity. Geometry is brought into the characterization through calculation of “Equivalent Removal Rate”:

\[
Roys = \frac{1}{A} \cdot \int \frac{1}{D} \cdot O \cdot Fr \, dA \quad (2)
\]

in which:

- \(Roys\) = equivalent removal rate (d\(^{-1}\))
- \(D\) = local depth (m)

The equivalent removal rate can be viewed as a decay rate of material in the water column. High removal rates indicate the bivalves clear the water column rapidly. The inverse of the equivalent removal rate is an “Equivalent Residence Time”: the time required for the bivalves to filter the water column once.

Under existing conditions, highest settling rates are in smaller tributaries; lower settling rates prevail in the mainstem bay and in the portions of major western tributaries that adjoin the bay (Figure 12). The tenfold biomass increase
(Figure 13) and the historic biomass (Figure 14) shift the highest settling rates to the lower portions of the western tributaries and to the upper mainstem of the bay. Median settling velocity increases by an order of magnitude from present modeled conditions to historical conditions (Table 4).

Under existing conditions, the ranking of residence times corresponds to the ranking of settling rates (Figure 15). Shortest residence times (highest turnover rates) are in tributaries. More lengthy residence times prevail in the lower portions of western tributaries and in the mainstem bay. The effects of geometry influence the rankings under conditions of oyster restoration (Figures 16, 17). Several of the large-volume segments which rank high in terms of settling rate rank lower when their depth is incorporated into the index of potential bivalve influence. Overall, the median residence time of individual CBPS’s diminishes from 18 days under computed existing conditions to less than three days under historic oyster densities (Table 4).

<table>
<thead>
<tr>
<th>Table 4</th>
<th>Median Settling Rates, Removal Rates, and Residence Times</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Settling, m d⁻¹</td>
</tr>
<tr>
<td>Existing Conditions</td>
<td>0.15</td>
</tr>
<tr>
<td>Tenfold Oyster Increase</td>
<td>0.62</td>
</tr>
<tr>
<td>Historic Conditions</td>
<td>1.44</td>
</tr>
</tbody>
</table>

References


HydroQual. (2000). “Development of a suspension feeding and deposit feeding benthos model for Chesapeake Bay,” produced by HydroQual Inc. under contract to the U.S. Army Engineer Research and Development Center, Vicksburg MS.


Figure 1. Density of existing bivalve filter feeders (from HydroQual 2000)
Figure 2. Present oyster density in Chesapeake Bay
Figure 3. Chesapeake Bay Program Segments
Figure 4. Calculated oyster density in the lower Choptank River, 1985-1994

Figure 5. Seasonal-average calculated oyster density in the lower Choptank River
Figure 6. Calculated autumn oyster density in lower Choptank River

Figure 7. Estimated density of existing oysters and bivalve filter feeders
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Figure 8. Estimated biomass of existing oysters and bivalve filter feeders

Figure 9. Calculated density of oysters and bivalve filter feeders under the nominal tenfold increase in oyster biomass
Figure 10. Calculated biomass of oysters and bivalve filter feeders under the nominal tenfold increase in oyster biomass

Figure 11. Effect of tenfold increase in oyster biomass on biomass of other bivalve filter feeders
Figure 12. Equivalent settling rate from bivalve filter feeders under existing conditions

Figure 13. Equivalent settling rate from oysters and bivalve filter feeders under the tenfold increase in oyster biomass
Figure 14. Equivalent settling rate from oysters and bivalve filter feeders under historic conditions

Figure 15. Time for bivalves to filter the water column under existing conditions
Figure 16. Time for oysters and bivalves to filter the water column under the tenfold increase in oyster biomass

Figure 17. Time for oysters and bivalves to filter the water column under historic conditions
4 Oyster Effects on Water Quality

Introduction

Oysters affect the environment on a variety of spatial scales ranging from their immediate surroundings outwards to the entire water body. The effects are considered here on three scales. The first is the smallest that can be resolved in the model, the model cell. Cell areas are \( \approx 10^6 \text{ m}^2 \), an order of magnitude larger than typical Maryland oyster bars. Since modeled oysters are uniformly distributed within cells, however, the processes in cells occupied by oysters are comparable to processes in bars containing similar densities of oysters. The second spatial scale is the regional scale represented by Chesapeake Bay Program Segments (CBPS). Program segments (Figure 1) are subdivisions of the bay determined by mean salinity, natural boundaries, and other features. Median area is \( \approx 1.5 \times 10^8 \text{ m}^2 \), of which only a fraction is occupied by oyster bottom. The third scale is system-wide, an area of \( 1 \times 10^{10} \text{ m}^2 \), as represented by the model grid.

We selected three of the 35 CBPS for detailed examination of oyster effects on the regional scale. The selected segments (Figure 1) provide a range of geometry (Table 1) and environmental conditions. CB4 is a mainstem bay segment with the greatest volume, surface area, and depth of the selected segments. Due to the depth, only 70% of the area is suitable for oyster habitat, as determined by the historic Yates surveys. Perhaps the most significant characteristic of the segment is the regular occurrence of summer bottom-water anoxia. EE2 is an eastern embayment that encompasses the mouth of the Choptank River. Volume is an order of magnitude less and depth is half of the selected mainstem segment. Virtually all of EE2 is suitable oyster habitat. Minimum dissolved oxygen concentration in bottom water occasionally falls below 3 g m\(^{-3}\) but persistent anoxia does not occur. Segment ET9 is the Big Annemessex River, located on the Maryland eastern shore. Despite the name, the Big Annemessex is the smallest of the three selected segments, separated by an order of magnitude in volume and area from EE2. Average depth is roughly half the depth in the lower Choptank River. Virtually all the segment provides suitable oyster habitat and minimum dissolved oxygen concentration exceeds 6 g m\(^{-3}\).
### Table 1
Regional Characteristics

<table>
<thead>
<tr>
<th>Region</th>
<th>Volume, $10^9$ m³</th>
<th>Area, km²</th>
<th>Mean Depth, m</th>
<th>Fraction oyster bottom</th>
</tr>
</thead>
<tbody>
<tr>
<td>CB4</td>
<td>10.8</td>
<td>966</td>
<td>11.2</td>
<td>0.71</td>
</tr>
<tr>
<td>EE2</td>
<td>1.8</td>
<td>334</td>
<td>5.3</td>
<td>1.00</td>
</tr>
<tr>
<td>ET9</td>
<td>0.1</td>
<td>33</td>
<td>2.8</td>
<td>0.8</td>
</tr>
</tbody>
</table>

### Local Effects

#### Biomass-Specific Effects

Effects on the local scale can be normalized by oyster biomass or by surface area. Biomass-specific results allow comparisons to published rates in Chesapeake Bay and elsewhere. For examination of biomass-specific effects, we selected a cell at a depth of 6.7 m within the lower Choptank River, CBPS EE2 (Figure 1). This region supports a viable oyster population and represents the environment from which oysters were drawn for the experiments of Jordan (1987) and Newell and Koch (2004).

Biomass-specific filtration rates, computed within the model based on the simulated environment, agree closely with the experiments on which the rates were based as well as with other independent measures and calculations (Table 1). Order-of-magnitude similarity prevails between modeled and measured respiration and ammonium excretion (Table 1). An interesting contrast occurs with carbon deposition (Table 1). The model agrees well with Jordan’s measures but departs from other reports. The modeled and measured filtration and respiration measures are comparable across systems because these are primarily functions of oyster physiology. Carbon deposition is influenced by local organic carbon concentration as well as by physiological processes and, consequently, can only be compared when local organic carbon concentrations are similar.

#### Areal-Based Effects

The regional and system-wide effects of oyster restoration are best understood by first isolating the local impacts of oysters. This is accomplished by examining sediment diagenetic processes and fluxes between the bottom sediments, oysters, and water column for a range of oyster densities. The basis for comparison is the 2002 version of the model (Cerco and Noel 2004), which included no oysters. This is compared to multiple model runs with oysters, conducted at various mortality rates, that produced a range of oyster densities. Three cells are considered, one each from CB4, EE2, and ET9. All values are annual averages across the ten simulated years.

**Benthic Algae.** Benthic algae (Figure 2) are non-existent in the CB4 (3.7 m depth) and EE2 (6.7 m depth) cells in the absence of oysters. The shallow ET9 cell (2.1 m depth) supports viable benthic algae at zero oyster density. Density of benthic algae increases in all cells concurrent with oyster density as oysters clear the water column of suspended solids. The enhancement of benthic algae is consistent with experimental results (Newell et al. 2002, Porter et al. 2004)
although only the ET9 cell sustains algal density we calculate is sufficient to influence nutrient exchange at the sediment-water interface (Cerco and Noel 2004). The model state variable is algal carbon. Most observations are of

### Table 1
**Modeled and Observed Biomass-Specific Oyster Effects**

<table>
<thead>
<tr>
<th>Property</th>
<th>Rate</th>
<th>Source</th>
<th>Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td>Filtration rate, m² g⁻¹ oyster C d⁻¹</td>
<td>0.24</td>
<td>Model</td>
<td>Summer average</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Jordan (1987)</td>
<td>Mean value, T ≥ 20 °C</td>
</tr>
<tr>
<td></td>
<td>0.26</td>
<td>Newell and Koch (2004)</td>
<td>Average of measures at 20 and 25 °C</td>
</tr>
<tr>
<td></td>
<td>0.027 to 0.33</td>
<td>Epifanio and Ewart (1977)</td>
<td>For algal suspensions &gt; 1 g C m⁻³</td>
</tr>
<tr>
<td></td>
<td>0.27</td>
<td>Risgård (1988)</td>
<td>Calculated for a 2.1 g DW oyster at 27 to 29 °C</td>
</tr>
<tr>
<td>Respiration rate, g DO g⁻¹ oyster C d⁻¹</td>
<td>0.04</td>
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<tr>
<td></td>
<td></td>
<td>Boucher and Boucher-Rodini (1988)</td>
<td>Spring and summer rates</td>
</tr>
<tr>
<td></td>
<td>0.017</td>
<td>Dame et al. (1992)</td>
<td>Annual average</td>
</tr>
<tr>
<td></td>
<td>0.02</td>
<td>Dame (1972)</td>
<td>1 g DW oyster at 20 to 30 °C</td>
</tr>
<tr>
<td>Ammonium excretion, mg N g⁻¹ oyster C d⁻¹</td>
<td>1.43</td>
<td>Model</td>
<td>Summer average</td>
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<tr>
<td></td>
<td>&lt; 0.1</td>
<td>Hammen et al. (1966)</td>
<td>Ammonium plus urea</td>
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<tr>
<td></td>
<td>2.8 to 3.88</td>
<td>Boucher and Boucher-Rodini (1988)</td>
<td>Spring and summer rates, includes urea</td>
</tr>
<tr>
<td></td>
<td>0.8</td>
<td>Srna and Baggaley (1976)</td>
<td>1 g DW oyster at 20 °C</td>
</tr>
<tr>
<td></td>
<td>4.8 to 7.9</td>
<td>Magni et al (2000)</td>
<td><em>Ruditapes and musculista</em></td>
</tr>
<tr>
<td>Carbon deposition, g C g⁻¹ oyster C d⁻¹</td>
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<td>Model</td>
<td>Summer average</td>
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<tr>
<td></td>
<td>0.099</td>
<td>Jordan (1987)</td>
<td>Mean value, T ≥ 20 °C</td>
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<td>0.03</td>
<td>Haven and Morales-Alamo (1966)</td>
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<tr>
<td></td>
<td>0.002 to 0.012</td>
<td>Tenore and Dunstan (1973)</td>
<td>Depends on C concentration, range is 0.1 to 0.7 g C m⁻³</td>
</tr>
</tbody>
</table>
chlorophyll. Using a carbon-to-chlorophyll ratio of 50 (Gould and Gallagher 1990) indicates annual-average computed benthic algal chlorophyll is 30 to 40 mg m$^{-2}$ in the ET9 cell.

**Carbon and Oxygen Fluxes.** The introduction of oysters results in biodeposition of carbon to the sediments (Figure 3). Carbon deposition due to gravitational settling (Figure 4) is simultaneously diminished as particulate carbon that previously settled is instead filtered. Total carbon deposition (Figure 5) is diminished by the introduction of oysters indicating that the minimum computed density is sufficient to reduce net production of particulate carbon in the water column. The amount of carbon removed by filtering (Figure 6) levels off as oyster densities increase beyond the initial value. Several cells indicate diminished filtration at the highest oyster densities. We attribute the level filtration to an equilibrium between carbon supplied, through transport and production, and carbon removed. As oyster density increases, biodeposition decreases. At higher densities, larger fractions of the carbon filtered are lost through respiration or retained as biomass. Total carbon deposition, through settling and biodeposition, decreases continually in response to increased oyster density.

Increasing oyster densities are accompanied by continual increases in respiration (Figure 7) and decreases in diagenetic sediment oxygen consumption (Figure 8). As noted in the biomass-specific results, respiration is largely a function of oyster density, independent of location. The increased respiration is more than offset by decreased sediment oxygen consumption so that total oxygen consumption decreases as oyster density increases (Figure 9).

**Nitrogen.** Fluxes of particulate nitrogen reproduce the pattern established for carbon. The introduction of oysters produces biodeposits to the sediments. As oyster density increases, both biodeposition and settling decrease. Biodeposition decreases because a greater fraction of nitrogen filtered is lost through respiration or retained as biomass. Settling decreases because formation of particulate nitrogen in the water column, through algal activity, is diminished by oyster predation.

The introduction of oysters diminishes the release of diagenetically-produced sediment ammonium (Figure 10). Diminished ammonium release is partially offset by excretion from oysters but the net impact of oysters is reduced net release to the water column, especially at highest densities (Figure 11). Two processes contribute to the reduction in diagenetic ammonium release. The role of reduced nitrogen deposition is obvious. Enhanced sediment nitrification to nitrate is also apparent, as evidenced by enhanced sediment denitrification of nitrate to nitrogen gas (Figure 12). Denitrification is also enhanced by the flux of nitrate from the water column into the sediments; nitrate no longer used in algal production diffuses into the sediments instead. The net effect of oysters on total nitrogen is removal from the water column via enhanced denitrification and retention in the sediments (Figure 13).

**Phosphorus.** Oyster effects on particulate phosphorus follow the pattern established for carbon and nitrogen. Introduction of oysters results in biodeposition, which is partially offset by diminished gravitational settling. As
oyster density increases, both biodeposition and settling decrease. Biodeposition decreases because a greater fraction of phosphorus filtered is lost through respiration or retained as biomass. Settling decreases because formation of particulate phosphorus in the water column, through algal activity, is diminished by oyster predation.

The net effect of oysters on dissolved phosphorus contrasts with nitrogen and is site-specific. At two sites, release of diagenetically-produced phosphorus diminishes as oyster density increases while at the third site release of diagenetic phosphorus is largely independent of oyster density (Figure 14). The two sites at which release diminishes support the largest densities of benthic algae so interception of diagenetic phosphorus release is suggested. At the site with least benthic algae, EE2, oyster phosphorus excretion adds to the constant diagenetic flux so that net release of dissolved phosphorus to the water column increases (Figure 15) and net retention in the sediments decreases (Figure 16). At the other two sites, excretion offsets algal uptake so the net flux is nearly constant and retention in the sediments increases as a non-linear function of oyster density.

Regional Effects

Three model runs are considered: 1) no oyster restoration, derived from the 2002 version of the model; 2) a tenfold increase in oyster biomass; and 3) historic oyster density. Quantities selected for analysis include:

- Summer-average bottom dissolved oxygen,
- Summer-average surface chlorophyll,
- Summer-average light attenuation,
- Summer-average SAV biomass,
- Annual-average surface algal carbon,
- Annual-average net primary production,
- Annual-average particulate carbon deposition,
- Annual-average sediment oxygen demand,
- Annual-average surface total nitrogen,
- Annual-average particulate nitrogen deposition,
- Annual-average sediment diagenetic ammonium flux,
- Annual-average net nitrogen removal (denitrification plus burial),
- Annual-average surface total phosphorus,
- Annual-average particulate phosphorus deposition,
- Annual-average sediment diagenetic phosphorus release, and
- Annual-average net phosphorus removal

Our convention for surface concentration is the average over the upper 6.7 m of the water column, roughly the depth of the surface mixed layer in the mid-bay. Bottom dissolved oxygen is represented by all waters below 12.9 m in CB4 and below 6.7 m in EE2. Due to shallow depth, the surface mixed layer coincides with the bottom in ET9. Results are averaged across the entire regional area and across all model years.
Water quality standards in Chesapeake Bay are based on dissolved oxygen, chlorophyll, and water clarity. The ten-fold oyster increase improves summer-average, bottom, dissolved oxygen in this mainstem segment by less than 0.5 g m\(^{-3}\) (Figure 17). Simulation of historic oyster densities improves dissolved oxygen by roughly 1 g m\(^{-3}\). Computed surface chlorophyll is reduced by 30% for a ten-fold increase in oyster density and is halved when oysters are restored to historic densities (Figure 18). Light attenuation is reduced by roughly 20% for a ten-fold increase in oyster densities and by roughly 40% when oysters are restored to historic densities (Figure 19).

The improvements in dissolved oxygen and chlorophyll are effected by reductions in net primary production (Figure 20). A 20% reduction in production accompanies the ten-fold increase in oyster density. A reduction of nearly 40% results from restoration of historic densities. The water clarity improvements, effected by removal of phytoplankton and other solids from the water column, produce increases in computed SAV biomass of 33% to more than 100% (Figure 21).

Restoration of oysters increases net nitrogen removal (Figure 22), through denitrification and sediment retention, by 20% to 50% although the reduction in surface total nitrogen concentration is only 10% to 15% (Figure 23). When averaged over the region, the effect of oyster restoration is increased phosphorus retention in the sediments (Figure 24). Net removal increases by a third for a ten-fold increase in oyster density and doubles when oysters are restored to historic densities. Phosphorus concentration in the water column corresponds with net removal rates more closely than nitrogen (Figure 25). Surface total phosphorus concentration is reduced by 20% to 40%.

Improvements in summer-average, bottom, dissolved oxygen at the mouth of the Choptank are consistent with the mainstem segment: less than 0.5 g m\(^{-3}\) for a ten-fold increase in oyster density and roughly 1 g m\(^{-3}\) for restoration to historic densities (Figure 26). Percentage reductions in surface chlorophyll (Figure 27) and light attenuation (Figure 28) also correspond closely with the adjacent mainstem segment as do the reductions in net primary production (Figure 29) and improvements in SAV (Figure 30).

Computed dissolved oxygen concentration in the eastern shore embayment declines by 0.5 g m\(^{-3}\) as a consequence of oyster restoration (Figure 31). The decline in dissolved oxygen reflects diminished dissolved oxygen production associated with the 40% to 60% reduction in net primary production (Figure 32). Reductions in summer surface chlorophyll exceed the reductions in annual net production (Figure 33). The ten-fold increase in oyster density induces a 60% decrease in summer surface chlorophyll while restoration to historic densities induces a greater than 70% decrease. Light attenuation in this region decreases by a third to nearly a half (Figure 34). Corresponding increases...
in SAV greatly exceed the responses in other segments (Figure 35). SAV biomass nearly triples for a ten-fold increase in oyster density and increases by greater than a factor of four for restoration to historic oyster densities.

**Regional Budgets**

Nutrient budgets were constructed for each of the regions for the three subject model runs. Results are annual averages across all model years. Terms in the budgets are:

- **Point Source** – Direct inputs from municipal and industrial facilities
- **Distributed** – Loads to the region from the adjacent watershed
- **Atmospheric** – Loads to the water surface
- **Transport** – Net loads from the upstream region. For CB4, this is adjacent mainstem region CB2. For EE2, this is the Choptank River segment ET5. No upstream segment exists for ET9.
- **Net Removal** – Accumulation in the bottom sediments plus denitrification
- **Incremental** – Increase in net removal due to oysters

Nitrogen transport down the mainstem of the bay dwarfs all other sources and sinks in CB4 (Figure 36). In view of the enormity of nitrogen transported in relative to the amount removed by oysters, the ability of oyster restoration to impact this segment at all is remarkable. This budget suggests the impact of oysters on phytoplankton is through direct grazing rather than through nutrient removal that results in limits to phytoplankton growth. Although nutrient removal can be viewed as an ecosystem service, direct grazing should be regarded as the primary service. More phosphorus is removed in CB4 than flows in from upstream and local sources (Figure 37). The deficit is made up by net phosphorus transport from downstream, as indicated by our earliest model (Cerco and Cole 1994) and by bay nutrient budgets (Boynton et al. 1995). As with nitrogen, the incremental nutrient removal by oysters is small relative to the net transport along the bay axis.

Incremental nutrient removal by oysters in EE2 is significant relative to other regional sources and sinks. Under the restoration scenarios, net nitrogen (Figure 38) and phosphorus (Figure 39) removal exceed the local sources indicating nutrient import from the adjacent mainstem segment.

Nitrogen loading and net removal in segment ET9 are closely balanced under existing conditions (Figure 40). As with EE2, enhanced removal via oyster restoration results in nitrogen import from the adjacent Tangier Sound. This segment imports phosphorus under existing conditions (Figure 41). Net import is enhanced under conditions of oyster restoration.

**System-Wide Effects**

The methods, properties examined, and budgeting from the regional analyses are extensible to the entire system. We consider the system to extend from the fall lines of major tributaries to the mouth of the bay. We were
requested to make two supplementary model runs for the sponsor. These combined the 2002 model (Cerco and Noel 2004) with the nutrient and solids loads from the recent allocation. One run was completed without oysters. The second run incorporated the ten-fold oyster restoration. Since the results of those runs have not been documented, we summarize them here.

Summer-average dissolved oxygen concentration is considered for all portions of the bay greater than 12.9 m depth. Dissolved oxygen increases by 0.25 g m\(^{-3}\) for the ten-fold oyster restoration and by 0.8 g m\(^{-3}\) for restoration to historic levels (Figure 42). By way of comparison, the dissolved oxygen improvement attained by the allocation loads exceeds the improvement attained by oyster restoration to historic levels. Allocation loads combined with ten-fold oyster restoration provide the greatest level of improvement, more than 1 g m\(^{-3}\) over current levels. System-wide, summer, surface chlorophyll concentration declines by more than 1 mg m\(^{-3}\) for a ten-fold increase in oyster biomass and by 2.5 mg m\(^{-3}\) for restoration to historic levels (Figure 43). As with dissolved oxygen, the allocation loads provide greater benefit than oyster restoration with improved benefits from both load reductions and oyster restoration.

The water clarity improvements that accompany oyster restoration (Figure 45) produce increases in computed system-wide SAV biomass of 25% to more than 60% (Figure 46). The historic levels of oysters result in the greatest improvements in SAV, suggesting local solids removal can be more effective than indirect controls on organic solids effected through nutrient controls. Still, the allocation loads produce larger improvements than the proposed ten-fold increase in oyster biomass.

Load reductions produce greater reductions in total nutrients than oyster restoration. The allocation loads diminish system-wide surface total nitrogen by 0.27 g m\(^{-3}\) (Figure 47) and total phosphorus by 0.011 g m\(^{-3}\) (Figure 48) with marginal additional reductions accomplished by load reductions combined with oyster restoration. The maximum nutrient reductions accomplished by oyster restoration are 0.11 g m\(^{-3}\) total nitrogen and 0.009 g m\(^{-3}\) total phosphorus. These results contrast the different strategies for phytoplankton control. The allocation loads reduce phytoplankton through nutrient reductions. Oyster restoration controls phytoplankton by direct grazing; nutrient reductions are a by-product of algal removal.

System-wide nutrient budgets can be constructed that parallel the regional budgets. In this case, transport is the net flux at the mouth of the bay. Negative transport indicates nutrient loss to the ocean; positive transport indicates nutrient import from the ocean. Ten-fold oyster restoration removes 30,000 kg d\(^{-1}\) total nitrogen from the system (Figure 49). Oysters at historic levels remove 54,000 kg d\(^{-1}\). Ten-fold oyster restoration removes 4,000 kg d\(^{-1}\)
total phosphorus from the system (Figure 50). Oysters at historic levels remove 5,000 kg d\(^{-1}\). By way of comparison, the ten-fold restoration removes loading roughly equivalent to direct atmospheric deposition. These are 25,000 kg d\(^{-1}\) total nitrogen and 1,900 kg d\(^{-1}\) total phosphorus.

**References**


Figure 1. Chesapeake Bay program segments.
Figure 2. Effect of oysters on benthic algae.

Figure 3. Effect of oysters on particulate carbon biodeposition.
Figure 4. Effect of oysters on gravitational settling of particulate carbon.

Figure 5. Effect of oysters on total carbon deposition.
Figure 6. Effect of oysters on particulate carbon filtration.

Figure 7. Effect of oysters on areal respiration.
Figure 8. Effect of oysters on sediment oxygen demand.

Figure 9. Effect of oysters on total benthic oxygen consumption.
Figure 10. Effect of oysters on sediment-water ammonium flux.

Figure 11. Effect of oysters on net benthic dissolved nitrogen flux.
Figure 12. Effect of oysters on sediment denitrification.

Figure 13. Effect of oysters on net sediment nitrogen removal.
Figure 14. Effect of oysters on sediment-water dissolved phosphorus flux. Positive flux is release to the water column.

Figure 15. Effect of oysters on net benthic dissolved phosphorus flux.
Figure 16. Effect of oysters on net sediment phosphorus removal.

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Figure 23. Effect of oysters on annual-average, surface, total nitrogen in CB4.
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Figure 25. Effect of oysters on annual-average, surface, total phosphorus in CB4.
Figure 26. Effect of oysters on summer-average, bottom, dissolved oxygen in EE2.

Figure 27. Effect of oysters on summer-average, surface, chlorophyll in EE2.
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Figure 39. Effect of oysters on phosphorus budget in EE2.
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Figure 43. Effect of oysters on system-wide, summer-average, surface chlorophyll.
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Figure 45. Effect of oysters on system-wide, summer-average, light attenuation.
Figure 46. Effect of oysters on system-wide, summer-average, SAV biomass.

Figure 47. Effect of oysters on system-wide, annual-average, surface, total nitrogen.
Figure 48. Effect of oysters on system-wide, annual-average, surface, total phosphorus.

Figure 49. Effect of oysters on system-wide nitrogen budget.
Figure 50. Effect of oysters on system-wide phosphorus budget.
5 Discussion and Conclusions

Analysis of the oyster modeling is like peeling the proverbial onion. There’s always another layer to be examined. Every insight produces two more questions. Sufficient model runs have been conducted to resolve the oyster issue raised by the Chesapeake Bay 2000 Agreement:

By 2004, assess the effects of different population levels of filter feeders such as menhaden, oysters and clams on Bay water quality and habitat.

Additional examination of the runs can be conducted and fruitful insights remain to be obtained. The production of this report is motivated by the need to produce tangible, citable, documentation of the work completed to date.

Oyster restoration will, no doubt, benefit the bay environment. Our analyses indicate the chief benefit will be restoration of SAV, brought about by filtration of solids from the water column. The most significant conclusion from our work, however, is that oyster restoration is no panacea for the host of environmental problems that plague the bay. Oyster restoration should be viewed as one of many contributions to remediation of the bay’s problems.

Our work did not target specific regions of the bay with specific levels of restoration. Rather, target levels for system-wide biomass were attained and the spatial distribution of oysters was calculated dynamically based on computed environmental factors including salinity, suspended solids, and available food. Potential spatial distribution was limited to historic oyster beds. As a result of our approach, the modeled ten-fold increase in oyster biomass multiplied oysters in the Maryland portion of the bay by 50 times while the Virginia portion of the bay received only a four-fold increase, primarily in the lower James and Rappahannock Rivers. Consequently, our ten-fold increase probably exaggerates the benefits to be obtained by ten-fold increases in local oyster densities in the northern bay.

Our work indicates a ten-fold oyster increase will improve summer-average, bottom, dissolved oxygen by $\sim 0.3 \, \text{g m}^{-3}$ in the portion of the mainstem plagued by the worst anoxia. Oyster restoration alone is not likely to bring the deep channel of the mainstem into compliance with dissolved oxygen standards. A dissolved oxygen increase of $0.3 \, \text{g m}^{-3}$ has economic value when traded off against the costs of nutrient controls. Some portions of the bay that marginally violate dissolved oxygen standards will marginally meet the standards when improved by $0.3 \, \text{g m}^{-3}$. System-wide, the combination of oyster restoration
and the recent nutrient allocations are calculated to increase summer-average, bottom, dissolved oxygen by \( \sim 1.1 \text{ g m}^{-3} \).

Multiple reasons can be offered for the absence of more significant dissolved oxygen response to oyster restoration. The obvious explanation is that oysters are found in the shoals rather than over the deep trench. Phytoplankton production over the trench remains free to settle to bottom waters and contribute to anoxia. A more subtle explanation lies in the origins of mainstem anoxia. Oxygen depletion in the upper bay does not originate solely with excess production in the overlying waters. Rather, oxygen depletion is accumulated as net circulation moves bottom water up the channel from the mouth of the bay. This mechanism was originally proposed by Kuo et al. (1991) for the Rappahannock River and has been shown to apply to the mainstem bay as well (Cerco 1995). Improvement in upper bay dissolved oxygen requires reduction in lower bay oxygen demand. The oyster restoration strategy does nothing to diminish oxygen demand in the lower bay and, consequently, has limited impact on the upper bay.

Our work indicates oyster restoration removes both nitrogen and phosphorus from the bay water column. Nitrogen removal is more significant than phosphorus removal since nitrogen is the nutrient that contributes to excess algal production in the portions of the bay occupied by oysters (Fisher et al. 1992, Malone et al. 1996). We calculate the ten-fold increase in oyster biomass removes 30,000 kg d\(^{-1}\) total nitrogen from the system via enhanced denitrification and retention in the sediments. This removal can be put into perspective by noting the Susquehanna River provides \( \sim 150,000 \text{ kg d}^{-1}\) total nitrogen to the mainstem while point sources in the Baltimore vicinity provide \( \sim 15,000 \text{ kg d}^{-1}\) (Cerco and Noel 2004). Oyster restoration may substitute for a major upgrade in point-source controls but does not offset the larger distributed loading from the watershed.

The comparison above does not address timing. Loads from the watershed arrive largely during spring runoff and occasionally as autumn tropical storms. Removal via oysters occurs during the warm months concurrent with peak algal production. This issue introduces the question of primary “services” provided by oysters. We suggest the primary service is direct grazing on algae. Rather than quantifying the amount of nitrogen removed by oysters, we should ask what load reductions produce reductions in algal biomass equivalent to the reductions from grazing. Nutrient removal is a byproduct of grazing. In order for nutrient removal to have value, it must be shown that the removal enhances limits to algal production. The model can provide insights in this regard and additional examination is warranted.

Our model provides unique capability to address oyster restoration in the bay. We believe ours is the first approach to combine detailed representation of the bay geometry with mechanistic representations of three-dimensional transport, water-column eutrophication processes, sediment diagenetic processes, and dynamic computation of oyster biomass. Due to the large number of computed interactions, exact quantification of benefits such as SAV biomass improvement involves uncertainty. We believe, however, our basic findings regarding the nature and magnitude of restoration benefits are valid. Our results
are consistent with the earlier findings of Officer et al (1992) and Gerritson et al. (1994) and with the recent findings of Newell and Koch (2004). Benthic controls of algal production are most effective in shallow, spatially-limited regions. In these shallow regions, oyster removal of solids from the water column enhances adjacent SAV beds. The ability to influence deep regions of large spatial extent is limited by the location of oysters in the shoals and by exchange processes between the shoals and deeper regions.

The potential improvements obtained by oyster restoration are also limited by factors not considered in the model. Disease is an obvious limitation. Habitat destruction has also been suggested as an impediment (Rothschild et al. 1994). We recommend that oyster restoration be targeted to specific areas with suitable environments and that resulting environmental improvements be viewed on similar, local scales.

References


Evaluating Ecosystem Effects of Oyster Restoration in Chesapeake Bay

A Report to the Maryland Department of Natural Resources

September 2005

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Abstract

The Chesapeake Bay Environmental Model Package (CBEMP) was used to assess the environmental benefits of oyster restoration in Chesapeake Bay. The CBEMP consists of a coupled system of models including a three-dimensional hydrodynamic model, a three-dimensional eutrophication model, and a sediment diagenesis model. Restoration levels up to fifty times the 1994 base biomass were examined. Examination of results emphasized dissolved oxygen, chlorophyll concentration, and water clarity. Within Virginia, the improvement in summer, bottom-water dissolved oxygen at the maximum biomass investigated was 0.2 mg/L. Within Maryland, the improvement was doubled, more than 0.4 mg/L. Within Virginia, the range of oyster densities investigated reduced summer-average surface chlorophyll by up to ~ 0.7 µg/L, roughly 10% of the 1994 base concentration. Corresponding reductions in Maryland were up to ~ 2.3 µg/L, more than 25% of the 1994 base. Within Virginia, the range of oyster densities investigated reduced summer-average light attenuation by up to 8%, from 1.05 m⁻¹ at base levels to 0.97 m⁻¹ for a fifty-fold increase in oyster biomass. Following a pattern established for other benefits, improvements in Maryland exceeded Virginia. Summer-average light attenuation diminished by up to 13%, from 1.39 m⁻¹ under base conditions to 1.21 m⁻¹ for a fifty-fold increase in oyster biomass.

Ecosystem services performed by oysters include nitrogen removal and SAV restoration. The range of oyster densities investigated removed up to 24,600 kg d⁻¹ nitrogen in Maryland and up to 5,100 kg d⁻¹ in Virginia. Relative improvements in SAV biomass were greater than corresponding reductions in light attenuation. Percentage increases in summer SAV biomass in Virginia were up to 21%. Computed SAV biomass increased from 5,627 tonnes C under base conditions to 6,830 tonnes for a fifty-fold oyster restoration. In Maryland, improvements in SAV biomass were up to 43%. Computed summer SAV biomass increased from 5,227 tonnes C under base conditions to 7,486 tonnes C under maximum restoration.

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1 Introduction

Oyster biomass and harvest in the Chesapeake Bay system have been declining exponentially since the nineteenth century (Rothschild et al. 1994, Kirby and Miller 2005). A link between decimation of the oyster population and deteriorating water quality in Chesapeake Bay was proposed by Newell (1988). Newell calculated the nineteenth-century oyster population could filter the entire volume of the bay in less than a week and suggested an increase in the oyster population could significantly improve water quality by removing large quantities of particulate carbon. While Newell’s proposition was not universally accepted (e.g. Gerritsen et al. 1994), the idea that managing the natural resource can improve water quality has fascinated scientists and managers since the proposition was advanced.

The potential links between living resources and water quality are central to the “Chesapeake 2000” agreement, signed by the executives of the Commonwealth of Pennsylvania, the State of Maryland, the Commonwealth of Virginia, the District of Columbia, the US Environmental Protection Agency, and the Chesapeake Bay Commission. The agreement sets specific goals including:

- Restore, enhance and protect the finfish, shellfish and other living resources, their habitats and ecological relationships to sustain all fisheries and provide for a balanced ecosystem.

The agreement lists methods to achieve this goal including:

- By 2010, achieve, at a minimum, a tenfold increase in native oysters in the Chesapeake Bay, based on a 1994 baseline.

- By 2004, assess the effects of different population levels of filter feeders such as menhaden, oysters and clams on Bay water quality and habitat.

The environmental effects of a ten-fold increase in population of native oysters were assessed by incorporating oysters into the Chesapeake Bay Environmental Model Package (CBEMP), a comprehensive mathematical model of physical and eutrophication processes in the bay and its tidal tributaries (Cerco and Noel 2005).
The decline of the native oyster, *Crassostrea virginica*, has been attributed to overfishing (Jordan and Coakley 2004), disease (Andrews 1965, Andrews 1988), and habitat destruction (Rothschild et al. 1994, Kirby and Miller 2005). The intractable problem of disease has led to the proposal to introduce a disease-resistant exotic oyster, *Crassostrea ariakensis*, to the Chesapeake Bay system. The Maryland Department of Natural Resources (DNR) and other organizations have initiated a wide range of studies to evaluate the environmental impact of oyster restoration and of *C. ariakensis* introduction. These studies fall under the heading of Ecological Risk Assessment (ERA) and will be summarized in a Programmatic Environmental Impact Statement (EIS). This report provides information for the ERA and EIS by evaluating several endpoints related to ecosystem impacts of the oyster restoration effort. This work addresses, among other factors, oyster impact on dissolved oxygen, algal biomass, light penetration, and submerged aquatic vegetation (SAV) abundance.

**The Chesapeake Bay Environmental Model Package**

Three models are at the heart of the CBEMP. Distributed flows and loads from the watershed are computed with a highly-modified version of the HSPF model (Bicknell et al. 1996). These flows are input to the CH3D-WES hydrodynamic model (Johnson et al. 1993) that computes three-dimensional intra-tidal transport. Computed loads and transport are input to the CE-QUAL-ICM eutrophication model (Cerco and Cole 1993) which computes algal biomass, nutrient cycling, and dissolved oxygen, as well as numerous additional constituents and processes. The eutrophication model incorporates a predictive sediment diagenesis component (DiToro and Fitzpatrick 1993) as well as living resources including benthos (Meyers et al. 2000), zooplankton (Cerco and Meyers 2000), and submerged aquatic vegetation (Cerco and Moore 2001).

A revision of the CBEMP was delivered in 2002 (Cerco and Noel 2004) and used in development of the most recent nutrient and solids load allocations in the bay. This version of the model was used to examine the impact of the tenfold increase in native oysters (Cerco and Noel 2005). The same version is used here to examine ecological effects of a wider range of restored oyster biomass. The 2002 CBEMP employs nutrient and solids loads from Phase 4.3 of the watershed model (Linker et al. 2000). (Documentation may be found on the Chesapeake Bay Program web site http://www.chesapeakebay.net/modsc.htm.) Nutrient and solids loads are computed on a daily basis for 94 sub-watersheds of the 166,000 km$^2$ Chesapeake Bay watershed and are routed to individual model cells based on local watershed characteristics and on drainage area contributing to the cell. The hydrodynamic and eutrophication models operate on a grid of 13,000 cells. The grid contains 2,900 surface cells (.4 km$^2$) and employs non-orthogonal curvilinear coordinates in the horizontal plane. Z coordinates are used in the vertical direction, which is up to 19 layers deep. Depth of the surface cells is 2.1 m at mean tide and varies as a function of tide, wind, and other forcing functions. Depth of sub-surface cells is fixed at 1.5 m. A band of littoral cells, 2.1 m deep at mean tide, adjoins the shoreline throughout most of the system. Ten years, 1985-1994, are simulated continuously using time steps of .5 minutes (hydrodynamic model) and .15 minutes (eutrophication model).
Critical Assumptions

Baseline rules and critical assumptions were made at the commencement of the study. These were forced by available knowledge (or lack thereof) and by the requirement to produce a valid product within a reasonable time frame.

Mass-Balance Based Model

Our approach models oysters from a mass-balance perspective. Oyster biomass is computed as a function of food availability, respiration, and mortality. Environmental effects on life processes are explicitly considered so that filtering capacity is consistent with environmental conditions. Our approach emphasizes the spatial and temporal distributions of filtering capacity and the environmental effects of filtering and deposition. Population processes including recruitment and larval setting are not considered. The demographic modeling conducted as a part of the larger EIS effort includes population effects not considered here.

Equivalence of *C. virginica* and *C. ariakensis*

The oyster model incorporated into the CBEMP considers market-sized native oysters and is parameterized to the greatest extent possible with local observations. Insufficient information exists to distinguish *C. ariakensis* from *C. virginica* within the model. Available information suggests model parameters adopted for native oysters apply to the exotic oysters as well. Preliminary laboratory experiments (National Research Council 2004) indicate size-specific filtration rates for *C. ariakensis* are similar to those of *C. virginica*. Mann (2005) concluded there is no reason filtration rates should differ significantly between the two species. These findings are consistent with conclusions of Powell et al. (1992) that size-specific filtration rates are similar for most marine bivalve species.

Historical Spatial Distribution

Our approach restricts oysters to their historical locations. This approach is reasonable in view of oysters’ affinity for specific bottom types. More elaborate restoration schemes including the creation of new habitat or the construction of rafts can be readily modeled but are left for future investigations.

Spatially-Uniform Mortality Rates

The model combines mortality from harvest, predation, and disease into a single first-order mortality term. In the absence of any information, this term is considered to be spatially uniform throughout the system. Non-uniform mortality rates can be added to the model as a future effort.

References


2 The Oyster Model

Introduction

The ultimate aim of eutrophication modeling is to preserve precious living resources. Usually, the modeling process involves the simulation of living-resource indicators such as dissolved oxygen. For the “Virginia Tributary Refinements” phase of the Chesapeake Bay modeling (Cerco et al. 2002), a decision was made to initiate direct interactive simulation of three living resource groups: zooplankton, benthos, and SAV.

Benthos were included in the model because they are an important food source for crabs, finfish, and other economically and ecologically significant biota. In addition, benthos can exert a substantial influence on water quality through their filtering of overlying water. Benthos within the model were divided into two groups: deposit feeders and filter feeders (Figure 1). The deposit-feeding group represents benthos that live within bottom sediments and feed on deposited material. The filter-feeding group represents benthos that live at the sediment surface and feed by filtering overlying water. The primary reference for the benthos model (HydroQual, 2000) is available on-line at http://www.chesapeakebay.net/modsc.htm. Less comprehensive descriptions may be found in Cerco and Meyers (2000) and in Meyers et al. (2000).

The benthos model incorporates three filter-feeding groups: 1) Rangea cuneata, which inhabit oligohaline and lower mesohaline portions of the system; 2) Macoma baltica, which inhabit mesohaline portions of the system; and 3) Corbicula fluminea, which are found in the tidal fresh portion of the Potomac. These organisms were selected based on their dominance of total filter-feeding biomass and on their widespread distribution. The distributions of the organisms within the model grid were assigned based on observations from the Chesapeake Bay benthic monitoring program (http://www.chesapeakebay.net/data/index.htm). Oysters were neglected in the initial application of the benthos model. The primary reasoning was that oyster biomass was considered negligible relative to the most abundant organisms.

Oysters

The oyster model builds on the concepts established in the benthos model. The existing benthos model was left untouched. The code was duplicated and one portion was modified for specific application to native
oysters, *Crassostrea virginica*. The original model assigned one of the three species exclusively to a model cell. In the revised model, oysters may coexist and compete with the other filter feeders. The fundamental state variable is oyster carbon, quantified as mass per unit area. The minimum area represented is the quadrilateral model cell, which is typically 1 to 2 km on a side. Oyster biomass and processes are averaged over the cell area. Oysters filter particulate matter, including carbon, nitrogen, phosphorus, silica, and inorganic solids from the water column. Particulate matter is deposited in the sediments as feces and pseudofeces. Respiration removes dissolved oxygen from the water column while excretion returns dissolved nitrogen and phosphorus.

Particulate carbon is removed from the water column by the filtration process. Filtration rate is affected by temperature, salinity, suspended solids concentration, and dissolved oxygen. The amount of carbon filtered may exceed the oyster’s ingestion capacity. In that case, the excess of filtration over ingestion is deposited in the sediments as pseudofeces (Figure 2). A portion of the carbon ingested is refractory or otherwise unavailable for nutrition. The unassimilated fraction is deposited in the sediments as feces. Biomass accumulation (or diminishment) is determined by the difference between carbon assimilated and lost through respiration and mortality. Respiration losses remove dissolved oxygen from the water column. Mortality losses are deposited to the sediments as particulate carbon.

The nutrients nitrogen and phosphorus constitute a constant fraction of oyster biomass. Particulate nitrogen and phosphorus, filtered from the water column, are subject to ingestion and assimilation. Assimilated nutrients that are not accumulated in biomass or lost to the sediments through mortality are excreted to the water column in dissolved inorganic form. All filtered particulate silica is deposited to the sediments or excreted to the water column. A fraction (~ 10%) of filtered inorganic solids is deposited to the sediments. The fraction is determined by the net settling velocity specified in the suspended solids algorithms. The remainder is considered to be resuspended.

The mass-balance equation for oyster biomass is:

\[
\frac{dO}{dt} = \alpha \cdot Fr \cdot POC \cdot IF \cdot (1 - RF) \cdot O - BM \cdot O - \beta \cdot O \quad (1)
\]

in which:

- \(O\) = oyster biomass (g C m\(^{-2}\))
- \(\alpha\) = assimilation efficiency (0 < \(\alpha\) < 1)
- \(Fr\) = filtration rate (m\(^3\) g\(^{-1}\) oyster carbon d\(^{-1}\))
- \(POC\) = particulate organic carbon in overlying water (g m\(^{-3}\))
- \(IF\) = fraction ingested (0 < \(IF\) < 1)
- \(RF\) = respiratory fraction (0 < \(RF\) < 1)
- \(BM\) = basal metabolic rate (d\(^{-1}\))
- \(\beta\) = specific mortality rate (d\(^{-1}\))
- \(t\) = time (d)
The assimilation efficiency is specified individually for each form of particulate organic matter in the water column. The respiratory fraction represents active respiratory losses associated with feeding activity. Basal metabolism represents passive respiratory losses.

**Filtration**

Filtration rate is represented in the model as a maximum or optimal rate that is modified by ambient temperature, suspended solids, salinity, and dissolved oxygen:

\[ Fr = f(T) \cdot f(TSS) \cdot f(S) \cdot f(DO) \cdot Fr_{\text{max}} \]  

(2)

in which:

- \( f(T) \) = effect of temperature on filtration rate (0 ≤ \( f(T) \) ≤ 1)
- \( f(TSS) \) = effect of suspended solids on filtration rate (0 ≤ \( f(TSS) \) ≤ 1)
- \( f(S) \) = effect of salinity on filtration rate (0 ≤ \( f(S) \) ≤ 1)
- \( f(DO) \) = effect of dissolved oxygen on filtration rate (0 ≤ \( f(DO) \) ≤ 1)
- \( Fr_{\text{max}} \) = maximum filtration rate (m\(^3\) g\(^{-1}\) oyster carbon d\(^{-1}\))

Bivalve filtration rate, quantified as water volume cleared of particles per unit biomass per unit time (Winter 1978), is typically derived from observed rates of particle removal from water overlying a known bivalve biomass (Doering et al. 1986, Doering and Oviatt 1986, Riisgard 1988, Newell and Koch 2004). Since particle retention depends on particle size and composition (Riisgard 1988, Langdon and Newell 1990), correct quantification of filtration requires a particle distribution that represents the natural distribution in the study system (Doering and Oviatt 1986). Filtration rate for our model was based primarily on measures (Jordan 1987) conducted in a laboratory flume maintained at ambient conditions in the adjacent Choptank River, a mesohaline Chesapeake Bay tributary that supports a population of native oysters. These were supplemented with laboratory measures conducted on oysters removed from the same system (Newell and Koch 2004). Jordan reported weight-specific biodeposition rate as a function of temperature, suspended solids concentration and salinity. The biodeposition rate represents a minimum value for filtration since all deposited material is first filtered. Filtration rate was derived:

\[ Fr = \frac{WBR}{TSS} \]  

(3)

in which:

- \( WBR \) = weight-specific biodeposition rate (mg g\(^{-1}\) dry oyster weight hr\(^{-1}\))
- \( TSS \) = total suspended solids concentration (mg L\(^{-1}\))

Filtration rate was converted from L g\(^{-1}\) DW h\(^{-1}\) to model units based on a carbon-to-dry-weight ratio of 0.5.

The observed rates indicate a strong dependence of filtration on temperature (Figure 3) although the range of filtration rates observed at any
temperature indicate the influence of other factors as well. The maximum filtration rate and the temperature dependence for use in the model are indicated by a curve drawn across the highest filtration rates at any temperature:

\[ Fr = F_{r \text{max}} \cdot e^{-K_{tg} \cdot (T - T_{opt})^2} \]  

in which:

\( F_{r \text{max}} \) = maximum filtration rate (0.55 m\(^3\) g\(^-1\) oyster carbon d\(^-1\))

\( K_{tg} \) = effect of temperature on filtration (0.015 \( ^\circ \)C\(^-2\))

\( T \) = temperature for optimal filtration (27 \(^\circ\)C)

**Suspended Solids Effects.** The deleterious effect of high suspended solids concentrations on oyster filtration rate has been long recognized although the solids concentrations induced in classic experiments, 10\(^2\) to 10\(^3\) g m\(^-3\) (Loosanoff and Tommers 1948), are extreme relative to concentrations commonly observed in Chesapeake Bay. We formed our solids function by recasting Jordan’s data to show filtration rate as a function of suspended solids concentration (Figure 4). The experiments indicate three regions. Filtration rate was depressed when solids were below ~5 gm m\(^-3\) and above ~25 gm m\(^-3\), relative to filtration rate when solids were between these two levels. The observations suggest oysters reduce their filtration rate when food is unavailable or when filtration at the maximum rate removes vastly more particles than the oysters can ingest. We visually fit a piecewise function to Jordan’s data (Figure 4) supplemented with an approximation of Loosanoff and Tommers’ results:

\[ f(TSS) = \begin{cases} 
0.1 & \text{when } TSS < 5 \text{ g m}^{-3} \\
1.0 & \text{when } 5 \text{ g m}^{-3} < TSS < 25 \text{ g m}^{-3} \\
0.2 & \text{when } 25 \text{ g m}^{-3} < TSS < 100 \text{ g m}^{-3} \\
0.0 & \text{when } TSS > 100 \text{ g m}^{-3} 
\end{cases} \]

**Salinity Effects.** Oysters reduce their filtration rate when ambient salinity falls below ~20% of the oceanic value (Loosanoff 1953) and cease filtering when salinity falls below ~10% of the oceanic value. The form and parameterization of a relationship to describe these experiments is arbitrary. We selected a functional form (Figure 5) used extensively elsewhere in the CBEMP:

\[ f(S) = 0.5 \cdot (1 + \tanh (S - K_{Hsoy})) \]  

in which:

\( S \) = salinity (ppt)

\( K_{Hsoy} \) = salinity at which filtration rate is halved (7.5 ppt)

**Dissolved Oxygen.** Hypoxic conditions (dissolved oxygen < 2 g m\(^3\)) have a profound effect on the macrobenthic community of Chesapeake Bay. Effects range from alteration in predation pressure (Nestlerode and Diaz 1998) to species shifts (Dauer et al. 1992) to near total faunal depletion (Holland et al. 1977). In the context of the benthos model, effects of hypoxia are expressed through a
reduction in filtration rate and increased mortality. The general function from the benthos model (Figure 6), based on effects from marine species, was adapted unchanged for the oyster model:

\[
f(\text{DO}) = \frac{1}{1 + \exp\left(1.1 \cdot \frac{\text{DO}_{hx} - \text{DO}}{\text{DO}_{hx} - \text{DO}_{qs}}\right)}
\]

(6)

in which:

\(\text{DO} = \) dissolved oxygen in overlying water \((\text{g m}^{-3})\)

\(\text{DO}_{hx} = \) dissolved oxygen concentration at which value of function is one-half \((1.0 \text{ g m}^{-3})\)

\(\text{DO}_{qs} = \) dissolved oxygen concentration at which value of function is one-fourth \((0.7 \text{ g m}^{-3})\)

This logistic function has the same shape as the tanh function used to quantify salinity effects (Figure 5). The use of two parameters, \(\text{DO}_{hx}\) and \(\text{DO}_{qs}\), allows more freedom in specifying the shape of the function than the tanh function, based on the single parameter \(\text{KH}_{soy}\), allows.

**Ingestion**

Oyster ingestion capacity must be derived indirectly from sparse observations and reports. In the report on his experiments, Jordan (1987) states “at moderate and high temperatures and low seston concentration (< 4 mg/L) nearly all biodeposits were feces” (page 54). This statement indicates no pseudofeces was produced; all organic matter filtered was ingested. Elsewhere in Jordan (1987) we find that ~75% of seston is organic matter and the filtration rate at 4 g seston m\(^{-3}\) is ~0.1 m\(^{-3}\) g\(^{-1}\) oyster C d\(^{-1}\) (Figure 4). The ingestion rate must be at least the amount of organic matter filtered. Conversion to model units indicates an ingestion rate of:

\[
\frac{4 \text{ g seston}}{\text{m}^{-3}} \cdot \frac{0.75 \text{ organic}}{\text{total}} \cdot \frac{\text{g C}}{2.5 \text{ g seston}} \cdot \frac{0.1 \text{ m}^3}{\text{g C d}} = \frac{0.12 \text{ g C ingested}}{\text{g oyster C d}}
\]

Tenore and Dunstan (1973) present a figure showing feeding rate and biodeposition. The difference between feeding and deposition must be ingestion. The largest observed difference is 19 mg C g\(^{-1}\) DW d\(^{-1}\) or 0.038 g C ingested g\(^{-1}\) oyster C d\(^{-1}\) (utilizing a carbon-to-dry-weight ratio of 0.5). No pseudofeces was produced during their experiments so the derived ingestion rate is not necessarily a maximum value.

In reporting on the removal of algae from suspension, Epifanio and Ewart (1977) noted that large amounts of pseudofeces were produced when algal suspensions exceeded 12 μg mL\(^{-1}\). These results indicate the amount removed from the water column when algal suspensions were less than 12 μg mL\(^{-1}\), ~4 to 17 mg algal DW g\(^{-1}\) oyster total weight d\(^{-1}\), was ingested. The 15 g total weight
oysters in Epifanio and Ewart’s experiments has a dry weight of 0.27 g (Dame 1972). The minimum ingestion rate is then:

$$\frac{4 \text{ mg algal DW}}{g \text{ oyster TW}} \times \frac{15 \text{ g TW}}{0.27 \text{ g DW}} \times \frac{g \text{ oyster DW}}{0.5 \text{ g oyster C}} \times \frac{2500 \text{ mg DW}}{18.0 \text{ lg}} = \frac{0.18 \text{ g C ingested}}{g \text{ oyster C d}}$$

Analogous unit conversions yield 0.76 g C ingested g$^{-1}$ oyster C d$^{-1}$ for a removal rate of 17 mg algal DW g$^{-1}$ oyster total weight d$^{-1}$.

Summary of these analyses indicates the order of magnitude for ingestion rate is 0.1 g C ingested g$^{-1}$ oyster C d$^{-1}$. The value 0.12 g C ingested g$^{-1}$ oyster C d$^{-1}$ was employed in the model based on our evaluation of Jordan’s experiments.

**Assimilation**

The fraction of ingested carbon assimilated by oysters depends on the carbon source. The assimilation of macrophyte detritus can be as low as 3% (Langdon and Newell 1990) while the assimilation of viable microphytobenthos is 70% to 90% (Cognie et al.). Tenore and Dunstan (1973) observed that oysters assimilated 77% to 88% of a mixed algal culture. Specification of assimilation for the oyster model is shaped by the nature of the eutrophication model. The eutrophication model considers three forms of particulate organic carbon: phytoplankton, labile particulate organic carbon, and refractory particulate organic carbon. Assimilation of phytoplankton is specified as 75%, based on citations above. The labile and refractory particulate organic carbon are detrital components. These are mapped to three G classes of organic matter (Westrich and Berner 1984) employed in the sediment diagenesis model (DiToro 2001). The G1, labile, class has half-life of 20 days. The G2, refractory, class has a half-life of one year. The G3 class is inert within time scales considered by the model. Model labile particulate organic carbon maps to the G1 class and is assigned an assimilation efficiency of 75%, corresponding to phytoplankton. Model refractory particulate organic carbon combines the G2 and G3 classes and is assigned an assimilation efficiency of zero.

**Respiration**

Two forms of respiration are considered: active respiration, associated with acquiring and assimilating food, and passive respiration (or basal metabolism). This division of respiration is consistent with models of predators ranging from zooplankton (Steele and Mullin 1977) to fish (Hewett and Johnson 1987). Active respiration is considered to be a constant fraction of assimilated food. Basal metabolism is represented as a constant fraction of biomass, modified by ambient temperature:

$$BM = BMr \cdot e^{KT_{bmr} (T - T_r)}$$  \hspace{1cm} (7)

in which:

- $BM = \text{basal metabolism (d}^{-1})$
- $BMr = \text{basal metabolism at reference temperature (d}^{-1})$
The rate of basal metabolism depends on organism biomass (Winter 1978, Shumway and Koehn 1982). The average oyster in Jordan’s (1987) experiments, upon which our filtration rates are based, is 2.1 g DW. Allometric relationships (Shumway and Koehn 1982) indicate basal metabolism for a 2.1 g DW oyster at 20 °C is 0.002 to 0.005 d⁻¹, depending on salinity. A graphical summary presented by Winter (1978) indicates metabolic rate for a 2 g DW oyster at 20 °C is 0.009 d⁻¹. Winter noted a 1 g DW mussel requires 1.5% of its dry tissue weight daily as a maintenance ration. Based on these reports, the value 0.008 d⁻¹ was employed for basal metabolism at a reference temperature of 20 °C. Parameter KTbmr was assigned the value 0.069 °C⁻¹, equivalent to a Q10 of 2, typical of measured rates in oysters (Shumway and Koehn 1982).

The respiratory fraction was assigned through comparison of computed oxygen consumption with metabolism in active oyster reefs (Boucher and Boucher-Rodoni 1988, Dame et al. 1992). The value RF = 0.1 was determined. A comparable value of 0.172 (specific dynamic activity coefficient) was assigned to herbivorous fish in Chesapeake Bay (Luo et al. 2001).

Mortality

The model considers two forms of mortality. These are mortality due to hypoxia and a term that considers all other sources of mortality including disease and harvest. Although bivalves incorporate physiological responses that render them tolerant to hypoxia, extended periods of anoxia result in near-extinction (Holland et al. 1977, Josefson and Widbom 1988). Casting the results of experiments and observations into a relationship that quantitatively relates mortality to dissolved oxygen concentration incorporates a good deal of uncertainty in functional form and parameterization. The effect of hypoxia on oyster mortality, adopted from the benthos model, employs two concepts. The first is the time to death under complete anoxia. This time to death is converted to a first-order mortality rate via the relationship:

\[ hmr = \frac{\ln(1/100)}{ttd} \]  

in which:

hmr = mortality due to hypoxia (d⁻¹)

\( ttd \) = time to death for 99% of the population (14 d)

The mitigating effect on mortality of dissolved oxygen concentration greater than zero is quantified through multiplication by \((1 - f(\text{DO}))\) in which \( f(\text{DO}) \) is the logistic function that expresses the effects of hypoxia on filtration rate (Equation 6). This functionality increases mortality as dissolved oxygen concentrations become low enough to affect filtration rate (Figure 6). When dissolved oxygen is depleted, filtration rate approaches zero and mortality is at its...
maximum. As parameterized in the model, effects on filtration and mortality are negligible until dissolved oxygen falls below ~ 2 g m\(^{-3}\) (Figure 6). The time to death for 99% of the population exceeds 90 days when dissolved oxygen exceeds 1.4 g m\(^{-3}\) (Figure 7). Under this scheme, some fraction of the oyster population can survive an entire summer of hypoxia provided dissolved oxygen exceeds 1.4 g m\(^{-3}\). No significant portion of the oyster population will survive summer hypoxia for dissolved oxygen concentrations below 1.4 g m\(^{-3}\).

Mortality from all other sources, primarily disease and harvest, is represented by a spatially uniform and temporally constant first-order term. Magnitude of the term is specified to produce various system-wide population levels with the model. The order of magnitude can be derived from Jordan et al. (2002) who reported the 1990 total mortality of “market stock” oysters in northern Chesapeake Bay was 0.94 yr\(^{-1}\) (or 0.0026 d\(^{-1}\)). Of this total, 0.22 yr\(^{-1}\) (or 0.0006 d\(^{-1}\)) was natural mortality. The balance was fishing mortality.

Nutrients

Model oysters are composed of carbon, nitrogen, and phosphorus in constant ratios. In the original benthos model (HydroQual 2000), the carbon-to-nitrogen mass ratio of bivalves was set at 5.67:1; the phosphorus-to-carbon mass ratio was 45:1. Composition data for bivalves is not abundant. Calculations by Jordan (1987), based on earlier work by Kuenzler (1961) and Newell (1982), yield a carbon-to-nitrogen mass ratio between 4.8:1 and 6.9:1 and a phosphorus-to-carbon mass ratio of 66:1. The nitrogen composition values encompass the value used in the model. The phosphorus composition value differs from the model but no context exists to judge if the difference is significant.

The oyster model differs substantially from the original benthos model in the way nutrients are assimilated and processed. In the original model, nutrients are assimilated and excreted in constant ratios equivalent to the oyster composition. If assimilated carbon is in excess relative to assimilated nitrogen or phosphorus, the excess carbon is converted to feces and the bivalves are effectively nutrient limited. Computed bivalve growth is:

\[
G = \min\left[\frac{\text{Cassim} \cdot \text{Nassim} \cdot \text{SFCN} \cdot \text{Passim}}{\text{SFCP}}\right] \quad (9)
\]

in which:

- \(G\) = bivalve biomass accumulation (g C m\(^{-2}\) d\(^{-1}\))
- \(\text{Cassim}\) = carbon assimilation rate (g C m\(^{-2}\) d\(^{-1}\))
- \(\text{Nassim}\) = nitrogen assimilation rate (g N m\(^{-2}\) d\(^{-1}\))
- \(\text{SFCN}\) = bivalve carbon-to-nitrogen ratio (g C g\(^{-1}\) N)
- \(\text{Passim}\) = phosphorus assimilation rate (g P m\(^{-2}\) d\(^{-1}\))
- \(\text{SFCP}\) = bivalve carbon-to-nitrogen ratio (g P g\(^{-1}\) N)

If the carbon-to-nitrogen ratio in assimilated food, \(\text{Cassim}/\text{Nassim}\), exceeds the ratio in bivalve composition, \(\text{SFCN}\), then biomass accumulation is proportional to the rate of nitrogen assimilation. Similarly, when the ratio \(\text{Cassim}/\text{Passim} > \text{SFCP}\), biomass accumulation is proportional to phosphorus assimilation. The
algal phosphorus-to-carbon ratio in the eutrophication model (Cerco and Noel 2004) is 57:1 for spring diatoms and 80:1 for other algae. Since these ratios exceed SFCP, growth of bivalves feeding on algae will be limited by the phosphorus content of the algae rather than the amount of carbon assimilated.

Algal composition does not provide a complete picture of the tendency for nutrient limitation of bivalve growth since modeled bivalves utilize detritus as well as algae. Initial applications of the oyster model indicated, however, that phosphorus limitation of oyster growth did occur. Nutrient limitation was eliminated through two methods. First, oyster phosphorus composition was thinned out; carbon-to-phosphorus ratio was increased to 90:1. More significantly, a mass balance approach to nutrient utilization and excretion was adopted. Biomass accumulation was modeled as carbon assimilation less respiration loss while nutrient excretion was calculated as the amount of assimilated nutrients not required for biomass accumulation.

**Model Parameters**

Parameter values for the oyster model are summarized in Table 1.
### Table 1
Parameters for Oyster Model

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Definition</th>
<th>Value</th>
<th>Units</th>
</tr>
</thead>
<tbody>
<tr>
<td>( F_{\text{max}} )</td>
<td>maximum filtration rate</td>
<td>0.55</td>
<td>( \text{m}^3 \text{ g}^{-1} \text{ oyster carbon d}^{-1} )</td>
</tr>
<tr>
<td>( T_{\text{opt}} )</td>
<td>optimum temperature for filtration</td>
<td>27</td>
<td>( \circ C )</td>
</tr>
<tr>
<td>( K_{\text{tg}} )</td>
<td>constant that controls temperature dependence of filtration</td>
<td>0.015</td>
<td>( \circ C^{-2} )</td>
</tr>
<tr>
<td>( K_{\text{hsoy}} )</td>
<td>salinity at which filtration rate is halved</td>
<td>7.5</td>
<td>ppt</td>
</tr>
<tr>
<td>( \text{BMR} )</td>
<td>base metabolism rate at 20 ( \circ C )</td>
<td>0.008</td>
<td>( \text{d}^{-1} )</td>
</tr>
<tr>
<td>( K_{\text{Tbmr}} )</td>
<td>constant that controls temperature dependence of metabolism</td>
<td>0.069</td>
<td>( \circ C^{-1} )</td>
</tr>
<tr>
<td>( T_r )</td>
<td>reference temperature for specification of metabolism</td>
<td>20</td>
<td>( \circ C )</td>
</tr>
<tr>
<td>( RF )</td>
<td>respiratory fraction</td>
<td>0.1</td>
<td>( 0 &lt; RF &lt; 1 )</td>
</tr>
<tr>
<td>( DO_{hx} )</td>
<td>dissolved oxygen concentration at which value of logistic function is one-half</td>
<td>1.0</td>
<td>( \text{g m}^{-3} )</td>
</tr>
<tr>
<td>( DO_{qx} )</td>
<td>dissolved oxygen concentration at which value of logistic function is one-quarter</td>
<td>0.7</td>
<td>( \text{g m}^{-3} )</td>
</tr>
<tr>
<td>( t_{\text{td}} )</td>
<td>time to death for 99% of the population</td>
<td>14</td>
<td>( \text{d} )</td>
</tr>
<tr>
<td>( a_{\text{phy}} )</td>
<td>assimilation efficiency for phytoplankton</td>
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<td>( I_{\text{max}} )</td>
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<td>0.12</td>
<td>( \text{g prey C} \text{ g}^{-1} \text{ C d}^{-1} )</td>
</tr>
<tr>
<td>( S_{\text{FCN}} )</td>
<td>carbon-to-nitrogen ratio</td>
<td>6</td>
<td>( \text{g C} \text{ g}^{-1} \text{ N} )</td>
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<tr>
<td>( S_{\text{FCP}} )</td>
<td>carbon-to-phosphorus ratio</td>
<td>90</td>
<td>( \text{g C} \text{ g}^{-1} \text{ P} )</td>
</tr>
</tbody>
</table>
References


HydroQual. (2000). “Development of a suspension feeding and deposit feeding benthos model for Chesapeake Bay,” produced by HydroQual Inc. under contract to the U.S. Army Engineer Research and Development Center, Vicksburg MS.


Figure 1. Benthos model schematic.

Figure 2. Processes affecting filtered material.
Figure 3. Effect of Temperature on filtration rate.

Figure 4. Effect of suspended solids on filtration rate.
Figure 5. Effect of salinity on filtration rate.

Figure 6. Effect of dissolved oxygen on filtration and mortality rates.
Figure 7. Effect of dissolved oxygen on time to death for 99% of population.
3 Biomass Estimates

Introduction

Estimates of the current oyster biomass and distribution were prepared for the native oyster study (Cerco and Noel 2005). Since our model is based on mass balance, population estimates took the form of total mass rather than number of individuals. The present study employs alternate estimates of current biomass, provided by the sponsor, but retains the spatial distribution determined for the preceding study. This chapter reviews the initial estimates and presents the biomasses employed in the present study. We use the terms “biomass” to indicate total weight of oysters e.g. kg C and “density” to indicate weight per unit area e.g. g C m$^{-2}$.

Distribution of Native Oysters

Density estimates for Virginia were provided by Dr. Roger Mann, of Virginia Institute of Marine Science. Estimates were based on patent tong surveys. Patent tong samples were averaged for each model cell and results were provided as g DW m$^{-2}$. Number of samples per cell varied from 4 to more than 50. Estimates were provided for one to five individual years in the interval 1998-2002. The area of cells containing oysters was 377 km$^2$.

Mean Maryland biomass, for the period 1991-2000, was obtained from Jordan et al. (2002). This biomass, $5.7 \times 10^8$ g DW, was uniformly distributed across the historical oyster habitat denoted in the “Yates” surveys (Yates 1911). The areas of named oyster bars were assigned to model cells. Total area of named oyster bars was 1330 km$^2$. A mean density of 0.43 g DW m$^{-2}$ (total biomass / total area) was assigned to the bar area in each model cell. Since the bar area was usually less than the cell area, cell density was adjusted so that biomass per cell matched biomass of bars within the cell. The area of cells containing oysters was 3696 km$^2$.

The oyster density and distribution are distinctly different in the Maryland and Virginia portions of the bay (Figure 1). In the northern, Maryland, portion, lower densities are distributed over a wide area. In the southern, Virginia, portion, high densities are concentrated in limited areas, primarily in the lower James and Rappahannock Rivers. Our estimated oyster biomass in Virginia is five times the biomass in Maryland (Table 1) but distributed across an order of magnitude less area. We were puzzled by the limited distribution in Virginia, especially since maps and other information we obtained indicated a
wider distribution of lease holdings and restoration areas. We were assured by Dr. Roger Mann that much of the leased area is unproductive and that biomass outside the areas reported to us is negligible.

Although the leased area is unproductive, information provided by the sponsor, attributed to the Virginia Marine Resources Commission, indicates 30% of the leased bottom is suitable for larval settling. This suitable area, 84 km² (20,866 acres), is significant relative to the area of public oyster bars that presently support oysters, 46 km² (11,366 acres). Model scenarios that consider oyster restoration in portions of Virginia lease holdings and restoration areas would be a worthwhile addition to the scenarios considered thus far.

**Modeled Biomass**

Computed density and biomass vary on intra-annual and inter-annual bases (Figure 2). Variations within the annual cycle are largely driven by temperature. Highest densities are computed in late summer and in fall, after a season of filtering at peak rates (Figure 3). Variations from year to year (Figure 4) are largely driven by runoff. Variations in runoff may enhance or diminish computed biomass, depending on local factors. Years with high runoff coincide with large nutrient loads that result in high phytoplankton abundance. The advantages produced by abundant food may be offset, however, by increased anoxia and by sub-optimal salinity.

Target values for baywide total biomass were provided by the sponsor. These were approximately matched during the model simulations (Table 1). Exact matching is not possible due to the intra-annual and inter-annual variability. We initially attempted to calculate target oyster densities through dynamic variation of the mortality function. Mortality in each model cell was adjusted upwards or downwards as calculated density exceeded or fell below specified levels. This process ensured that target density was not exceeded but in many cells target density could not be achieved. The problem originated with the attempt to calculate target densities within individual cells. The calculated conditions in many cells would not support the target densities. Consequently, we switched to a strategy in which a bay-wide target biomass was specified. A uniform bay-wide mortality rate was prescribed that produced the target biomass. The mortality rate was obtained through a trial-and-error process in which various rates were prescribed and the calculated biomass was examined.

Autumn is the season when individual oysters attain maximum biomass and when most population surveys are conducted. Modeled biomasses reported here (Table 1) are the average calculated autumn (September – November) biomass from ten years (1985 – 1994). The modeled biomasses are interspersed with estimates from various sources. Model run OYS30 is in close agreement with the sponsor’s 1994 baseline estimate. Run OYS31 corresponds to a ten-fold increase over the sponsor’s 1994 baseline. Runs OYS28 and OYS34 bracket the sponsor’s estimate for the 1920-1970 period. The run with the highest calculated biomass, OYS33, represents only 25% of the pre-1870 biomass, however.
## Table 1
### Oyster Biomass Estimates

<table>
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<tr>
<th>Run Code</th>
<th>Mortality Rate, 1/d</th>
<th>Maryland, kg DW</th>
<th>Virginia, kg DW</th>
<th>Total, kg DW</th>
<th>Notes</th>
</tr>
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<tr>
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<td>0.0280</td>
<td>981,434</td>
<td>239,680</td>
<td>1,221,114</td>
<td>1,000,000 baseline provided by sponsor</td>
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<tr>
<td></td>
<td></td>
<td>1,100,000</td>
<td>800,000</td>
<td>1,900,000</td>
<td>1988 biomass from Newell (1988)</td>
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<tr>
<td></td>
<td></td>
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<td>2,772,688</td>
<td>Year 2000 exploitable biomass from Uphoff (2002)</td>
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<tr>
<td>OYS26</td>
<td>0.0255</td>
<td>3,867,648</td>
<td>699,594</td>
<td>4,567,242</td>
<td>Existing Biomass from Cerco and Noel (2005)</td>
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<td>OYS31</td>
<td>0.0236</td>
<td>8,509,914</td>
<td>1,785,170</td>
<td>10,295,084</td>
<td>Ten-fold increase estimated by sponsor</td>
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<tr>
<td>OYS32</td>
<td>0.0216</td>
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<td>2,808,368</td>
<td>15,290,664</td>
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</tr>
<tr>
<td>OYS28</td>
<td>0.0190</td>
<td>18,477,200</td>
<td>4,218,842</td>
<td>22,696,042</td>
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</tr>
<tr>
<td>OYS34</td>
<td>0.0175</td>
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<td>5,054,288</td>
<td>27,892,878</td>
<td>1920-1970 period provided by sponsor</td>
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<td>OYS33</td>
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<td>40,593,208</td>
<td>8,583,890</td>
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</tr>
<tr>
<td></td>
<td></td>
<td>120,000,000</td>
<td>68,000,000</td>
<td>188,000,000</td>
<td>Pre 1870 biomass from Newell (1988)</td>
</tr>
</tbody>
</table>
References


Figure 1. Present oyster density in Chesapeake Bay (Cerco and Noel 2005)
Figure 2. Time series of calculated daily oyster density in the lower Choptank River, 1985-1994

Figure 3. Seasonal-average calculated oyster density in the lower Choptank River
Figure 4. Calculated autumn oyster density in lower Choptank River
4 Ecosystem Services Provided by Oyster Restoration

Introduction

Oyster restoration can provide a variety of benefits classified under the heading “ecosystem services.” Water quality standards for Chesapeake Bay are based on dissolved oxygen, chlorophyll, and water clarity (U.S. Environmental Protection Agency 2003). Ecosystem services described here are focused on improvements in the water quality standards. Oysters affect their environment on multiple spatial scales ranging from the oyster reef outwards to the entire system. Examinations of oyster impacts on local, regional, and system-wide scales were conducted as part of the study of native oyster restoration (Cerco and Noel 2005). Analyses here divide the bay into two states, Maryland and Virginia. This division was prompted by the sponsor’s request for an estimation of nitrogen removal by state.

Chlorophyll

Oysters effect improvements in the environment by filtering phytoplankton and other suspended solids from the water column. Aside from direct removal, reductions in phytoplankton, quantified as chlorophyll concentration, may also occur via an indirect process: nutrient limitation induced through removal of nutrients, primarily nitrogen. Although phytoplankton require phosphorus and silica (for diatoms) as well, nitrogen limitation is the most significant influence on algal production in the interval when temperature-dependent oyster filtration is greatest (Fisher et al. 1992, Malone et al. 1996).

Within Virginia, the range of densities investigated reduce summer-average surface chlorophyll by up to ~ 0.7 µg/L, roughly 10% of the 1994 base concentration (Table 1). Corresponding reductions in Maryland are up to ~ 2.3 µg/L, more than 25% of the 1994 base. The disparity between the two states reflects the widespread distribution of oysters in Maryland relative to Virginia. Averaged over the area contained within each state, oyster densities are three to four times greater in Maryland than Virginia for any level or restoration (Table 1).

The range of densities investigated reduced surface total nitrogen concentration by up to 0.05 mg/L in Virginia (Table 1). The maximum reduction
was nearly identical in Maryland, 0.06 mg/L. Under base conditions, net nitrogen removal in Maryland, on an areal basis, is greater than in Virginia, 27 mg N m\(^{-2}\) d\(^{-1}\) versus 16 mg N m\(^{-2}\) d\(^{-1}\). The higher base rate in Maryland reflects deposition of particulate nitrogen below the major fall lines and diffusion of nitrate into bed sediments where it is subsequently denitrified. The difference between the two regions increases with the level of oyster restoration, attributable to the higher densities in Maryland. At the greatest densities examined, oyster restoration removes 4 mg N m\(^{-2}\) d\(^{-1}\) in Maryland versus 1 mg N m\(^{-2}\) d\(^{-1}\) in Virginia. Multiplication by bottom area in each state yields removal rate in mass terms: up to 24,600 kg d\(^{-1}\) additional nitrogen removal in Maryland versus up to 5,100 kg d\(^{-1}\) additional removal in Virginia (Table 1).

These removal rates can be put in perspective by examining some of the other loads to the system, derived from the 2002 model used for the recent load allocations (Cerco and Noel 2004). The Maryland removal rate corresponding to a fifty-fold increase in oyster biomass is roughly equivalent to the point-source nitrogen load to the Potomac basin (Table 2). The equivalence in loading should not be extended to equivalence in effects, however since the majority of the Potomac load enters in the tidal freshwater reach far removed from oyster habitat. The amount of nitrogen removed by Maryland oyster restoration to 1920 – 1970 levels is equivalent to direct atmospheric loading to the water surface; nitrogen removal from a ten-fold oyster restoration is half this amount.

The Virginia removal rate corresponding to a fifty-fold increase in oyster biomass is only half of direct atmospheric loading to the water surface (Table 2). Removal rates associated with restoration of oysters to 1920 – 1970 levels and with ten-fold oyster restoration are only small fractions of identifiable loads to the Virginia portion of the bay.

Additional perspective is gained by comparing the nitrogen removal via oyster restoration to nutrient reduction targets (Linker 2005). Recent allocations call for a 24,900 kg d\(^{-1}\) reduction in Maryland nitrogen loading. The allocation corresponds to nitrogen removal from a fifty-fold increase in oyster biomass (Table 2). The Virginia allocation calls for a 34,800 kg d\(^{-1}\) reduction in nitrogen loading. This allocation exceeds any feasible reduction from oyster restoration. The system-wide allocation calls for a 124,500 kg d\(^{-1}\) reduction in nitrogen loading. This allocation also exceeds any feasible reduction from oyster restoration. Nitrogen removal via oyster restoration can be a valuable supplement to alternate methods of nutrient control but is no substitute for conventional nutrient controls.

**Dissolved Oxygen**

Bottom-water hypoxia originates with excess algal production in the surface waters of the bay. Algae and detritus settle to the bottom where they undergo decay that generates oxygen demand and consumption. Density stratification prevents replenishment of oxygen-depleted waters with atmospheric oxygen from the surface.
Within Virginia, the range of oyster densities investigated reduced annual-average net algal production by up to 10%, from 0.68 g C m$^{-2}$ d$^{-1}$ at base levels to 0.62 g C m$^{-2}$ d$^{-1}$, for a fifty-fold increase in oyster biomass (Table 3). Corresponding reductions were greater in Maryland. Annual average net algal production was reduced up to 20%, from 0.74 g C m$^{-2}$ d$^{-1}$ at base levels to 0.59 g C m$^{-2}$ d$^{-1}$, for a fifty-fold increase in oyster biomass. Under base conditions, annual-average surface algal carbon concentration was equivalent in Maryland and Virginia, 0.5 g C m$^{-3}$ (Table 3). The maximum potential reduction attainable in Maryland, 0.07 g C m$^{-3}$, was double the potential gain in Virginia, however.

Oxygen improvements are considered for summer-average at depths greater than 12.9 m. This period and depth isolates the time and location of bottom-water hypoxia. Within Virginia, the improvement in bottom-water dissolved oxygen at the maximum biomass investigated was 0.2 mg/L (Table 3). Within Maryland, the improvement was doubled, more than 0.4 mg/L.

**Water Clarity**

Improvements in water clarity are effected by removal of both organic and inorganic solids from the water column. Water clarity is quantified in the model as the coefficient of diffuse light attenuation. The light attenuation coefficient is inversely proportional to water clarity. Lower light attenuation implies higher water clarity. We examined summer-average light attenuation since summer is the critical period for growth of submerged aquatic vegetation (SAV).

Within Virginia, the range of oyster densities investigated reduced summer-average light attenuation by up to 8%, from 1.05 m$^{-1}$ at base levels to 0.97 m$^{-1}$ for a fifty-fold increase in oyster biomass (Table 4). Percentage increases in summer SAV biomass were greater, up to 21%. Computed SAV biomass increased from 5,627 tonnes C under base conditions to 6,830 tonnes for a fifty-fold oyster restoration. Following a pattern established for other benefits, improvements in Maryland exceeded Virginia. Summer-average light attenuation diminished by up to 13%, from 1.39 m$^{-1}$ under base conditions to 1.21 m$^{-1}$ for a fifty-fold increase in oyster biomass. Corresponding percentage improvements in SAV, up to 43%, again exceeded improvements in attenuation. Computed summer SAV biomass increased from 5,227 tonnes C under base conditions to 7,486 tonnes C under maximum restoration.

**Discussion**

Results from these model runs were compared to runs conducted for the native oyster study (Cerco and Noel 2005). Results from all runs form a consistent body when compared on identical spatial scales e.g. model cell or system-wide. The reader is cautioned regarding nomenclature, however. Results from both studies were reported based on various levels of restoration including existing, ten-fold restoration, and historic levels. The existing oyster biomass provided by the sponsor of this study is less than the existing biomass derived by us for the native oyster study (Chapter 3, Table 1). Consequently, the ten-fold increase computed in this study represents lower biomass than the ten-fold
increase computed for the native oyster study. In the native oyster study, historical biomass refers to pre-1870 levels while the sponsor of this study uses “historical” to represent the 1920 – 1970 period. We recommend that results from both studies be summarized on identical spatial scales and presented as a function of target biomass rather than restoration levels.

Our work indicates the maximum improvement expected in deep-water summer dissolved oxygen is 0.2 (Virginia) to 0.4 (Maryland) mg/L. These effects are averaged over large expanses of the bay. Greater and lesser improvements will be found in specific locations. Still, oyster restoration alone is not likely to bring the deep channel of the mainstem, where complete anoxia may occur, into compliance with dissolved oxygen standards. Multiple reasons can be offered for the absence of more significant dissolved oxygen response to oyster restoration. The obvious explanation is that oysters are found in the shoals rather than over the deep trench. Phytoplankton production over the trench remains free to settle to bottom waters and contribute to anoxia. A more subtle explanation lies in the origins of mainstem anoxia. Oxygen depletion in the upper bay does not originate solely with excess production in the overlying waters. Rather, oxygen depletion is accumulated as net circulation moves bottom water up the channel from the mouth of the bay. This mechanism was originally proposed by Kuo et al. (1991) for the Rappahannock River and has been shown to apply to the mainstem bay as well (Cerco 1995). Improvement in upper bay dissolved oxygen requires reduction in lower bay oxygen demand. Oysters in the lower bay are concentrated in the western-shore tributaries, however. The oyster restoration strategy does little to diminish oxygen demand in the lower bay and, consequently, has limited impact on the upper bay.

Our model provides unique capability to address oyster restoration in the bay. We believe ours is the first approach to combine detailed representation of the bay geometry with mechanistic representations of three-dimensional transport, water-column eutrophication processes, sediment diagenetic processes, and dynamic computation of oyster biomass. Due to the large number of computed interactions, exact quantification of benefits such as SAV biomass improvement involves uncertainty. We believe, however, our basic findings regarding the nature and magnitude of restoration benefits are valid. Our results are consistent with the earlier findings of Officer et al. (1992) and Gerritson et al. (1994) and with the recent findings of Newell and Koch (2004). Benthic controls of algal production are most effective in shallow, spatially-limited regions. In these shallow regions, oyster removal of solids from the water column enhances adjacent SAV beds. The ability to influence deep regions of large spatial extent is limited by the location of oysters in the shoals and by exchange processes between the shoals and deeper regions. We recommend that oyster restoration be targeted to specific areas with suitable environments and that resulting environmental improvements be viewed on similar, local scales.
<table>
<thead>
<tr>
<th>Designation</th>
<th>VA oyster density g C m⁻²</th>
<th>MD oyster density g C m⁻²</th>
<th>VA Chl² ug/L</th>
<th>MD Chl² ug/L</th>
<th>VA total N³ mg/L</th>
<th>MD total N³ mg/L</th>
<th>VA N removal kg/d</th>
<th>MD N removal kg/d</th>
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<td>1994 base</td>
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<td>0</td>
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<td>0.87</td>
<td>473</td>
<td>2,812</td>
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<td>0.86</td>
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<td>15-fold increase</td>
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<td>0.52</td>
<td>0.85</td>
<td>2,344</td>
<td>6,918</td>
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<td>0.81</td>
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1 Annual average across state portion of the system

2 Summer (June – Aug.) average within surface mixed layer

3 Annual average within surface mixed layer

4 Incremental annual average removal compared to 1994 base
## Table 2: Nitrogen Loads and Incremental Removal Rates

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<thead>
<tr>
<th>Location</th>
<th>Virginia kg/d</th>
<th>Maryland kg/d</th>
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<td>James River Point Source</td>
<td>27,101</td>
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<td>James River Fall Line</td>
<td>20,455</td>
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<td>Distributed Loads</td>
<td>18,580</td>
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<td>Other Fall Line</td>
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<td>Atmospheric</td>
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<td>Other Point Source</td>
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<td>1920-1970</td>
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<td>2,344</td>
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<td>ten-fold</td>
<td>1,575</td>
<td>6,434</td>
</tr>
<tr>
<td>five-fold</td>
<td>473</td>
<td>4,754</td>
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<td></td>
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</table>
### Table 3  Ecosystem Benefits Associated with Dissolved Oxygen

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<tr>
<th>Designation</th>
<th>VA oyster biomass kg DW</th>
<th>MD oyster biomass kg DW</th>
<th>VA net production g C m⁻² d⁻¹</th>
<th>MD net production g C m⁻² d⁻¹</th>
<th>VA algal C g m⁻³</th>
<th>MD algal C g m⁻³</th>
<th>VA bottom DO mg/L</th>
<th>MD bottom DO mg/L</th>
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</thead>
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<tr>
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<td>981,434</td>
<td>0.68</td>
<td>0.74</td>
<td>0.50</td>
<td>0.50</td>
<td>4.68</td>
<td>2.14</td>
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<td>3,867,648</td>
<td>0.67</td>
<td>0.72</td>
<td>0.49</td>
<td>0.49</td>
<td>4.70</td>
<td>2.18</td>
</tr>
<tr>
<td>ten-fold increase</td>
<td>1,785,170</td>
<td>8,509,914</td>
<td>0.66</td>
<td>0.70</td>
<td>0.49</td>
<td>0.48</td>
<td>4.72</td>
<td>2.22</td>
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<td>12,482,296</td>
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<td>0.68</td>
<td>0.48</td>
<td>0.47</td>
<td>4.75</td>
<td>2.27</td>
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<td>1920 - 1970 level</td>
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<td>18,477,200</td>
<td>0.64</td>
<td>0.66</td>
<td>0.48</td>
<td>0.46</td>
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<td>25-fold increase</td>
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<td>22,838,590</td>
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<td>0.64</td>
<td>0.48</td>
<td>0.45</td>
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<td>0.47</td>
<td>0.43</td>
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<td>2.57</td>
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1 Autumn (Sept. – Nov.) average
2 Annual average net phytoplankton primary production
3 Annual average in surface mixed layer
4 Summer (June – Aug.) average in depth > 12.9 m
### Table 4  Ecosystem Benefits Associated with SAV

<table>
<thead>
<tr>
<th>Designation</th>
<th>VA oyster biomass 1 kg DW</th>
<th>MD oyster biomass 1 kg DW</th>
<th>VA light attenuation 2 1/m</th>
<th>MD light attenuation 2 1/m</th>
<th>VA SAV biomass 3 tonnes C</th>
<th>MD SAV biomass 3 tonnes C</th>
</tr>
</thead>
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<td>239,680</td>
<td>981,434</td>
<td>1.05</td>
<td>1.39</td>
<td>5,627</td>
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<td>five-fold increase</td>
<td>699,594</td>
<td>3,867,648</td>
<td>1.05</td>
<td>1.38</td>
<td>5,637</td>
<td>5,368</td>
</tr>
<tr>
<td>ten-fold increase</td>
<td>1,785,170</td>
<td>8,509,914</td>
<td>1.03</td>
<td>1.36</td>
<td>5,985</td>
<td>5,691</td>
</tr>
<tr>
<td>15-fold increase</td>
<td>2,808,368</td>
<td>12,482,296</td>
<td>1.02</td>
<td>1.33</td>
<td>6,169</td>
<td>5,973</td>
</tr>
<tr>
<td>1920 - 1970 level</td>
<td>4,218,842</td>
<td>18,477,200</td>
<td>1.02</td>
<td>1.30</td>
<td>6,113</td>
<td>6,332</td>
</tr>
<tr>
<td>25-fold increase</td>
<td>5,054,288</td>
<td>22,838,590</td>
<td>1.00</td>
<td>1.28</td>
<td>6,480</td>
<td>6,562</td>
</tr>
<tr>
<td>50-fold increase</td>
<td>8,583,890</td>
<td>40,593,208</td>
<td>0.97</td>
<td>1.21</td>
<td>6,830</td>
<td>7,486</td>
</tr>
</tbody>
</table>

1 Autumn (Sept. – Nov.) average

2 Summer (June – Aug.) average

3 Summer (June – Aug.) average
References


5 Information for Risk Assessment

Introduction

The project work plan calls for “…a discussion of uncertainty associated with model results and the best available quantitative estimation of uncertainty in results.” The ability to distinguish and quantify uncertainty varies with the nature of the model outputs. The uncertainty in quantities that are directly calculated by the model and regularly observed can be readily quantified. Uncertainty in derived model outputs or in quantities for which insufficient observations are available can be difficult or impossible to quantify although some qualitative description of uncertainty may still be possible. The quantities reported to the sponsor are presented in Table 1. Uncertainty in quantities that are part of the Bay Program monitoring program is quantified using the statistics described below. Uncertainty in the remaining quantities is described based on available information and the modelers’ experience.

Statistical Summaries

Statistics can be a valuable aid in assessing model performance. A wide variety of statistics is available and no standard suite exists. Neither are there definite criteria available for judging the success of model computations. We use a suite that has been applied to the succession of Chesapeake Bay applications and to other CE-QUAL-ICM applications. Use of these statistics allows for consistent interpretation of model performance and provides a database of comparable statistics from alternate model applications. Our standard statistics are:

Mean Difference

\[ MD = \frac{1}{N} \sum_{n=1}^{N} (P_n - O_n) \] (1)

in which:

N = number of observations
O_n = n^{th} observation
P_n = computation corresponding to n^{th} observation
**Absolute Mean Difference**

\[
AMD = \frac{1}{N} \cdot \sum_{n=1}^{N} |P_n - O_n| \quad (2)
\]

**Relative Difference**

\[
RD = 100 \cdot \frac{|MD|}{\left(\sum O_n/N\right)} \quad (3)
\]

The mean difference describes whether the model over-estimates (MD > 0) or under-estimates (MD < 0) the observations, on average. The mean difference can achieve its ideal value, zero, while large discrepancies exist between individual observations and computations. The absolute mean difference is a measure of the characteristic difference between individual observations and computations. An absolute mean difference of zero indicates the model perfectly reproduces each observation. The relative difference (%) is the absolute mean difference normalized by the mean concentration. Relative difference provides a statistic suitable for comparison between different variables or systems.

Performance statistics were computed based on the calibrated model used in the recent Chesapeake Bay nutrient allocations (Cerco and Noel 2004a). This is the same model to which oysters were added for the present study. Observations were selected to conform to the reported quantities e.g. surface observations when surface quantities are reported. Observations are from 42 stations examined in the model calibration (usually one station from each Chesapeake Bay Program Segment). These stations were sampled once or twice per month during the calibration period, 1985 – 1994. Algal carbon was not distinguished in the sampling so particulate organic carbon is substituted. Results are presented in Table 2.

**Zooplankton**

Mesozooplankton and microzooplankton are monitored but not at the same frequency and spatial density as water quality analyses. The mesozooplankton database consists of oblique vertical tows are from roughly 25 stations concentrated in the mainstem and larger tributaries. Vertical average values are derived from the model for comparison with the observations (Table 2). Microzooplankton observations are from 13 stations in the Maryland portion of the bay only. Microzooplankton samples are composites from “above pycnocline” or “below pycnocline.” Comparable quantities are derived from the model for comparison with the observations (Table 2).

**Submerged Aquatic Vegetation (SAV)**

The primary data base for calibration of the SAV model (Cerco and Moore 2001) was a time series of annual maximum abundance (tonnes C) by community type. The use of abundance observations is the reason SAV...
abundance is the primary quantity reported as model output. The model was also compared to living-resource criteria, primarily light attenuation. The following verbiage was used to describe model performance: “Comparison of model results with time series of observed community abundance indicates the model represents correctly the relative abundance in each community. Inter-annual variability and trends are not well represented, however. The median absolute difference between computed and observed bay-wide annual abundance, by community type, is 30% of observed values, with a range from zero to 240%.”

**Net Primary Production**

Development of primary production algorithms was the subject of special emphasis in the present Chesapeake Bay Environmental Model Package (Cerco and Noel 2004b). The model was calibrated against a data base of more than 160 observations collected throughout the bay from 1987 to 1994. Use of a paired t-test to compare individual observations with model calculations indicated the mean difference between computed and observed net production could not be distinguished from zero (p < 0.01). Regression was used to compare individual computations with observations. Results for the regression of computed versus observed net primary production were:

- Slope = 0.57 (95% CI = 0.08)
- Intercept = 0.32 (95% CI = 0.10)
- $R^2 = 0.26$
- $p < 0.0001$

**Benthic Algae**

The benthic algae model was developed for the Delaware Inland Bays (Cerco and Seitzinger 1997) and adapted to the Chesapeake Bay (Cerco and Noel 2004a). No local observations of benthic algal biomass exist for comparison with the model. Computed benthic algal biomass, up to 3 g C m$^{-2}$, was found to be consistent with biomass observed in a variety of systems. The model was checked for consistency with observed properties of benthic algae and their effects. The primary determinant of benthic algae is light at the sediment water interface. Algal density increases or decreases as illumination increases or decreases. We can conclude that algal biomass computed by the model is order-of-magnitude correct and responds correctly to environmental influences.

**Deposit and Filter Feeding Benthos**

Benthic deposit feeders and bivalve filter feeders (other than oysters) were added to the model as part of the Virginia Tributaries Refinements phase (HydroQual 2000). Computed benthos were compared to observations collected at various locations throughout the system. Observations showed a large degree of heterogeneity. Variations of two to four orders of magnitude in benthic biomass were commonly observed over the multi-year course of the sampling program. The variability made conventional comparisons of computations and observations (e.g. time series) difficult to evaluate. Probability plots were constructed that compared the distributions of observations and corresponding
computations at various sampling stations. Emphasis in evaluation was placed on the median observed and computed values. At some locations, the medians were within a few percent of each other. At other locations median observations and computations were separated by one to two orders of magnitude.

<table>
<thead>
<tr>
<th>Quantity</th>
<th>Observed in Monitoring Program?</th>
<th>Units</th>
</tr>
</thead>
<tbody>
<tr>
<td>Bottom Dissolved Oxygen</td>
<td>yes</td>
<td>g DO m⁻³</td>
</tr>
<tr>
<td>Surface Total Nitrogen</td>
<td>yes</td>
<td>g m⁻³</td>
</tr>
<tr>
<td>Surface Total Phosphorus</td>
<td>yes</td>
<td>g m⁻³</td>
</tr>
<tr>
<td>Surface Dissolved Inorganic Nitrogen</td>
<td>yes</td>
<td>g m⁻³</td>
</tr>
<tr>
<td>Surface Dissolved Inorganic Phosphorus</td>
<td>yes</td>
<td>g m⁻³</td>
</tr>
<tr>
<td>Submerged Aquatic Vegetation Biomass</td>
<td></td>
<td>tonnes carbon</td>
</tr>
<tr>
<td>Surface Total Suspended Solids</td>
<td>yes</td>
<td>g m⁻³</td>
</tr>
<tr>
<td>Surface Chlorophyll</td>
<td>yes</td>
<td>mg m⁻³</td>
</tr>
<tr>
<td>Surface Algal Biomass</td>
<td>yes, as POC</td>
<td>g C m⁻³</td>
</tr>
<tr>
<td>Net Primary Production</td>
<td></td>
<td>g C m⁻³ d⁻¹</td>
</tr>
<tr>
<td>Light Attenuation</td>
<td>yes</td>
<td>m⁻¹</td>
</tr>
<tr>
<td>Oyster Biomass</td>
<td></td>
<td>g C m⁻²</td>
</tr>
<tr>
<td>Benthic Algae</td>
<td></td>
<td>g C m⁻²</td>
</tr>
<tr>
<td>Mesozooplankton</td>
<td>yes</td>
<td>g C m⁻³</td>
</tr>
<tr>
<td>Microzooplankton</td>
<td>yes</td>
<td>g C m⁻³</td>
</tr>
<tr>
<td>Benthic Deposit Feeders</td>
<td></td>
<td>g C m⁻²</td>
</tr>
<tr>
<td>Other Benthic Filter Feeders</td>
<td></td>
<td>g C m⁻²</td>
</tr>
<tr>
<td>Constituent</td>
<td>MD</td>
<td>AMD</td>
</tr>
<tr>
<td>----------------------------------------</td>
<td>--------</td>
<td>--------</td>
</tr>
<tr>
<td>Bottom Dissolved Oxygen</td>
<td>-0.296</td>
<td>1.651</td>
</tr>
<tr>
<td>Surface Total Nitrogen</td>
<td>-0.004</td>
<td>0.333</td>
</tr>
<tr>
<td>Surface Total Phosphorus</td>
<td>-0.013</td>
<td>0.033</td>
</tr>
<tr>
<td>Surface Dissolved Inorganic Nitrogen</td>
<td>0.063</td>
<td>0.266</td>
</tr>
<tr>
<td>Surface Dissolved Inorganic Phosphorus</td>
<td>0.007</td>
<td>0.018</td>
</tr>
<tr>
<td>Surface Total Solids</td>
<td>-1.282</td>
<td>9.293</td>
</tr>
<tr>
<td>Surface Chlorophyll</td>
<td>0.453</td>
<td>7.355</td>
</tr>
<tr>
<td>Surface Particulate Organic Carbon</td>
<td>0.317</td>
<td>0.683</td>
</tr>
<tr>
<td>Light Extinction</td>
<td>0.040</td>
<td>0.677</td>
</tr>
<tr>
<td>Mesozooplankton</td>
<td>-0.002</td>
<td>0.018</td>
</tr>
<tr>
<td>Microzooplankon</td>
<td>-0.009</td>
<td>0.017</td>
</tr>
</tbody>
</table>

References


Guide to Model Outputs

Introduction

Model runs were conducted according to the workplan dated September 1, 2004. The workplan specified seven runs with biomass targets. One run was to be with the demographic model. Two runs were to be named at a later date. Results from the seven runs with biomass targets are included on the attached CD. For several reasons, the biomass targets can be satisfied only approximately. One factor is the model formulation. In the model, a mortality rate (representing primarily disease, harvest, and predation) is specified and the biomass is calculated. A mortality rate can be specified that will result in a biomass close to the target but the target value cannot be input exactly. Additional considerations that confound exact achievement of a target include the intra- and inter-annual variation in computed oyster biomass. Biomass varies on an annual cycle due to effects of temperature, dissolved oxygen, and salinity on filtration and respiration. Biomass varies from year to year due to variations in runoff that affect salinity, suspended solids, food availability, and other factors. Table 1 presents the biomass target and biomass achieved for each model run. The achieved biomass is taken as the ten-year average fall (September – November) computed biomass.

<table>
<thead>
<tr>
<th>Serial Number</th>
<th>Mortality, 1/d</th>
<th>Target Biomass, 10⁶ kg DW</th>
<th>Modeled Biomass, 10⁶ kg DW</th>
</tr>
</thead>
<tbody>
<tr>
<td>OYS30</td>
<td>0.028</td>
<td>1.0</td>
<td>1.22</td>
</tr>
<tr>
<td>OYS26</td>
<td>0.026</td>
<td>5.0</td>
<td>4.56</td>
</tr>
<tr>
<td>OYS31</td>
<td>0.0236</td>
<td>10.0</td>
<td>10.3</td>
</tr>
<tr>
<td>OYS32</td>
<td>0.0216</td>
<td>15.0</td>
<td>15.3</td>
</tr>
<tr>
<td>OYS28</td>
<td>0.019</td>
<td>20.0</td>
<td>22.7</td>
</tr>
<tr>
<td>OYS34</td>
<td>0.0175</td>
<td>25.0</td>
<td>27.9</td>
</tr>
<tr>
<td>OYS33</td>
<td>0.012</td>
<td>50.0</td>
<td>49.2</td>
</tr>
</tbody>
</table>

Model run OYS30 is an approximation of the 1994 baseline biomass. OYS28 is an approximation of the 1920 – 1970 biomass level.

Output Format

Results are presented for each of 71 segments (Figure 1) presently delineated by the EPA Chesapeake Bay Program. A pdf of the segmentation is included on the CD. Half these segments (Table 2) presently or potentially support oysters. Computations from each model cell are aggregated spatially to represent the Chesapeake Bay Program Segment (CBPS). Results are further aggregated on three time bases:

- Seasonal Results – Computations for each model time step (fifteen minutes) are aggregated into each of the forty seasons (four seasons x ten years) represented in the model run.
Seasonal Averages – Seasonal results are aggregated into four seasons each representing the average of ten computed seasons.

Annual Averages - Computations for each model time step are aggregated into each of the ten years represented in the model run.

<table>
<thead>
<tr>
<th>Table 2</th>
</tr>
</thead>
<tbody>
<tr>
<td>CBPS that Support Oysters</td>
</tr>
<tr>
<td>BIGMH  │ CHOHH  │ MANMH  │ POTMH</td>
</tr>
<tr>
<td>CB2OH  │ CHSMH  │ MOBPH  │ POTOH</td>
</tr>
<tr>
<td>CB3MH  │ CRRMH  │ NANMH  │ RHDHM</td>
</tr>
<tr>
<td>CB4MH  │ EASMH  │ PATMH  │ RPPMH</td>
</tr>
<tr>
<td>CB5MH  │ FSBMH  │ PAXMH  │ SEVMH</td>
</tr>
<tr>
<td>CB6PH  │ HNGMH  │ PAXOH  │ SOUMH</td>
</tr>
<tr>
<td>CB7PH  │ JMSMH  │ PIAMH  │ TANMH</td>
</tr>
<tr>
<td>CHOMH1 │ LCHMH  │ POCMH  │ WICMH</td>
</tr>
<tr>
<td>CHOMH2 │ MAGMH  │ POCOH  │ WSTMH</td>
</tr>
</tbody>
</table>

Seasons are defined as follows:

- Winter – December through February
- Spring – March through May
- Summer – June through August
- Fall – September through November

Several reported quantities are most relevant when reported for a specific location in the water column e.g. surface chlorophyll or bottom dissolved oxygen. In keeping with long-established model convention, “surface” is defined as all model cells within a layer that extends 6.7 m down from the air-water interface. This length scale approximates the surface mixed layer. The definition of “bottom” depends on local depth. For the deepest segments, the bottom includes all model cells greater than 12.8 m down from the air-water interface. Water at this depth is “below pycnocline.” For segments with no cells below 12.8 m, bottom includes all cells greater than 6.7 m down from the air-water interface. For segments with no cells below 6.7 m, bottom includes all cells less than 6.7 m down from the air-water interface and coincides with surface.

Results are provided in graphical and numerical formats on the CD. The CD is divided into folders that correspond with run serial numbers (Table 1). Within the folders are pdf’s of graphical results and multiple subfolders. The subfolders contain text files of the material presented in graphical format. Each CBPS is represented by two text files. (The use of two files makes these “printer friendly.”) Each column in the file contains a header consisting of a parameter code and the CBPS designation. For the “Seasonal” text files, the column headed “year” represents the decimal years into the model run at which the designated season ends. Reported quantities, parameter codes, and units are presented in Table 3.
Table 3
Reported Quantities

<table>
<thead>
<tr>
<th>Quantity</th>
<th>Parameter Code</th>
<th>Units</th>
</tr>
</thead>
<tbody>
<tr>
<td>Bottom Dissolved Oxygen</td>
<td>botdo</td>
<td>g DO m$^{-3}$</td>
</tr>
<tr>
<td>Surface Total Nitrogen</td>
<td>tn</td>
<td>g m$^{-3}$</td>
</tr>
<tr>
<td>Surface Total Phosphorus</td>
<td>tp</td>
<td>g m$^{-3}$</td>
</tr>
<tr>
<td>Surface Dissolved Inorganic Nitrogen</td>
<td>din</td>
<td>g m$^{-3}$</td>
</tr>
<tr>
<td>Surface Dissolved Inorganic Phosphorus</td>
<td>dip</td>
<td>g m$^{-3}$</td>
</tr>
<tr>
<td>Submerged Aquatic Vegetation Biomass</td>
<td>savbi</td>
<td>tonnes carbon</td>
</tr>
<tr>
<td>Surface Total Suspended Solids</td>
<td>tss</td>
<td>g m$^{-3}$</td>
</tr>
<tr>
<td>Surface Chlorophyll</td>
<td>chl</td>
<td>mg m$^{-3}$</td>
</tr>
<tr>
<td>Surface Algal Biomass</td>
<td>alcar</td>
<td>g C m$^{-3}$</td>
</tr>
<tr>
<td>Net Primary Production</td>
<td>npp</td>
<td>g C m$^{-2}$ d$^{-1}$</td>
</tr>
<tr>
<td>Light Attenuation</td>
<td>ke</td>
<td>m$^{-1}$</td>
</tr>
<tr>
<td>Oyster Biomass</td>
<td>oys</td>
<td>g C m$^{-2}$</td>
</tr>
<tr>
<td>Benthic Algae</td>
<td>benal</td>
<td>g C m$^{-2}$</td>
</tr>
<tr>
<td>Mesozooplankton</td>
<td>meso</td>
<td>g C m$^{-3}$</td>
</tr>
<tr>
<td>Microzooplankton</td>
<td>micro</td>
<td>g C m$^{-3}$</td>
</tr>
<tr>
<td>Benthic Deposit Feeders</td>
<td>dfeed</td>
<td>g C m$^{-2}$</td>
</tr>
<tr>
<td>Other Benthic Filter Feeders</td>
<td>ofeed</td>
<td>g C m$^{-2}$</td>
</tr>
</tbody>
</table>
Figure 1. Chesapeake Bay Program Segments