1-1-1997

Prediction of Benthic Impact for Salmon Net-Pens Based on the Balance of Benthic Oxygen Supply and Demand

R. H. Findlay

Les Watling
University of Maine - Main, watling@maine.edu

Follow this and additional works at: https://digitalcommons.library.umaine.edu/sms_facpub

Repository Citation
https://digitalcommons.library.umaine.edu/sms_facpub/98

This Article is brought to you for free and open access by DigitalCommons@UMaine. It has been accepted for inclusion in Marine Sciences Faculty Scholarship by an authorized administrator of DigitalCommons@UMaine. For more information, please contact um.library.technical.services@maine.edu.
Prediction of benthic impact for salmon net-pens based on the balance of benthic oxygen supply and demand

Robert H. Findlay1,*, Les Watling2

1Department of Biochemistry, Microbiology and Molecular Biology and Center for Marine Studies, Darling Marine Center, University of Maine, Walpole, Maine 04573, USA
2Darling Marine Center, University of Maine, Walpole, Maine 04573, USA

ABSTRACT: The ratio between oxygen supply and oxygen demand was examined as a predictor of benthic response to organic enrichment caused by salmon net-pen aquaculture. Oxygen supply to the benthos was calculated based on Fickian diffusion and near-bottom flow velocities. A strong linear correlation was found between measured carbon sedimentation rates and rates of benthic metabolism. This relationship allowed an estimation of oxygen demand based on sedimentation rates. Comparison of several production sites in Maine (USA) coastal waters showed that for sites where oxygen demand exceeded supply benthic impacts were high and for sites where oxygen supply exceeded demand benthic impacts were low. These findings were summarized in the form of a predictive model that should be useful in siting salmon production facilities.

KEY WORDS: Aquaculture · Benthic metabolism · Marine pollution

INTRODUCTION

As salmonid net-pen culture has become increasingly prevalent in marine near-shore environments so has concern for potential environmental damage. Numerous studies have demonstrated alterations of the benthos beneath and proximal to net-pens (Table 1). Effects are generally restricted to within 30 m of the pens and the magnitude of the effects has been categorized from 'no gross impact' (e.g. Frid & Mercer 1989) to 'highly eutrophicated' (e.g. Hall et al. 1990). Although effects have been documented, interpretation has been on a site by site basis due, in part, to the failure of linear models to relate benthic loading to benthic impacts, as well as the assumption that a single benthic assimilative capacity is applicable to all sea bottoms (e.g. Silvert 1992). It is clear that the first 7 effects documented in Table 1 are either direct or indirect consequences of increased benthic metabolism. Recent developments in our understanding of benthic-pelagic coupling suggest these responses can be directly linked to increases in carbon flux to the benthos (Nixon 1981, Hargrave 1984, Sampou & Oviatt 1991, Kemp & Boynton 1992, Hatcher et al. 1994).

The 2 main processes associated with organic matter degradation in shallow-water marine systems are the delivery of particulate organic matter from the water column to the benthos, and benthic decomposition and resulting solute fluxes. Recent developments in sediment trap technologies have greatly improved our ability to measure benthic carbon flux. Cylindrical traps should have the appropriate diameter to match the flow characteristics of the environment and aspect ratios >3.5 to minimize trap sampling bias (Butman 1986, Butman et al. 1986, Baker et al. 1988). In shallow-water systems resuspension followed by settling can confound measurements of deposition, but a comparison of the organic and inorganic content of sediments can be used to correct for the effect of resuspension on deposition rates (Kemp & Boynton 1992).
Table 1 Effects of fish net-pen culture on benthos beneath and proximal to net-pens

<table>
<thead>
<tr>
<th>Effect</th>
<th>Citation</th>
</tr>
</thead>
<tbody>
<tr>
<td>Increased sediment carbon inventories</td>
<td>Brown et al. (1987), Rosenthal et al. (1988) and citations within, Hall et al. (1990),</td>
</tr>
<tr>
<td></td>
<td>Weston (1990), Hansen et al. (1991), Ye et al. (1961), Hargrave et al. (1993)</td>
</tr>
<tr>
<td>Decreased sediment redox potential</td>
<td>Brown et al. (1987), Gowen &amp; Bradbury (1987), Rosenthal et al. (1988) and citations within,</td>
</tr>
<tr>
<td></td>
<td>Wildish et al. (1988), Lumb &amp; Fowler (1989), O'Connor et al. (1989), Wildish et al. (1990),</td>
</tr>
<tr>
<td></td>
<td>Hargrave et al. (1993)</td>
</tr>
<tr>
<td>Increased benthic metabolism measured as O₂</td>
<td>Gowen &amp; Bradbury (1987), Rosenthal et al. (1988) and citations within,</td>
</tr>
<tr>
<td>or CO₂ flux rates</td>
<td>Hall et al. (1990), Hansen et al. (1991), Holmer &amp; Kristensen (1992),</td>
</tr>
<tr>
<td></td>
<td>Hargrave et al. (1993), Findlay et al. (1995)</td>
</tr>
<tr>
<td>Increased H₂S and methane flux from the</td>
<td>Brown et al. (1987), Gowen &amp; Bradbury (1987), Rosenthal et al. (1988) and citations within,</td>
</tr>
<tr>
<td>sediments</td>
<td>Lumb &amp; Fowler (1989), Hall et al. (1990), Weston (1990), Hansen et al. (1991),</td>
</tr>
<tr>
<td></td>
<td>Holmer &amp; Kristensen (1992)</td>
</tr>
<tr>
<td>Increased nutrient flux from sediment</td>
<td>Gowen &amp; Bradbury (1987), Rosenthal et al. (1988) and citations within, Hansen et al.</td>
</tr>
<tr>
<td>Altered microbial community structure</td>
<td>Brown et al. (1987), Gowen &amp; Bradbury (1987), Rosenthal et al. (1988) and citations within,</td>
</tr>
<tr>
<td>Altered macrofauna abundance and</td>
<td>Brown et al. (1987), Gowen &amp; Bradbury (1987), Rosenthal et al. (1988) and citations within,</td>
</tr>
<tr>
<td>community structure</td>
<td>O'Connor et al. (1989), Weston (1990), Lim (1991), Ye et al. (1991),</td>
</tr>
<tr>
<td></td>
<td>Findlay et al. (1995)</td>
</tr>
<tr>
<td>Increased carbon flux to the benthos</td>
<td>Hall et al. (1990), Ye et al. (1991), Findlay et al. (1995), Tsutsumi (1995)</td>
</tr>
</tbody>
</table>

In several manipulative experiments, marine sediments enriched with organic matter showed increases in total benthic metabolism measured as CO₂ flux from and O₂ flux to the benthos (Anderson & Hargrave 1984, Keplay & Andersen 1985, Schwinghamer & Keplay 1987, Sampou & Oviatt 1991, Schwinghamer et al. 1991). In most of these experiments water movement was restricted and enrichment was sufficient to generate an oxygen demand greater than the supply deliverable by diffusion. Decomposition was mediated by anaerobic processes and virtually all oxygen consumption could be related to the reoxidation of reduced alternative electron acceptors. Under these conditions natural sediments will be devoid of macrofauna and sulfide-oxidizing bacteria will form mats at the water-sediment interface.

We reasoned that if the ability of the benthos to oxidize organic matter aerobically could be determined and compared to the organic flux originating from the net-pens, a prediction of benthic impact would result. As the State of Maine, USA (site of our study) has chosen the occurrence of aoxic conditions or the development of Beggiatoa sp. mats as defining an unacceptable benthic impact (Sowles et al. 1994), we chose these criteria as the endpoints of our model and focused on the factors that control the balance between aerobic and anaerobic metabolism in marine sediments. We broke the problem of siting net-pens into 2 halves: (1) predicting the impact of a known benthic carbon flux and (2) predicting a priori the flux from a net-pen system. Both are very significant problems and we have chosen to deal with the first in this manuscript. The purpose of this paper is to describe the theoretical basis of our model and the relationship between benthic organic matter loading and benthic metabolism as it relates to sediments proximal to several salmon net-pens in Maine coastal waters. We have developed our model of benthic impact associated with net-pen aquaculture using 2 parameters—oxygen supply to the sediments and sediment oxygen demand. We assumed that benthic metabolism, and therefore maximum sediment oxygen demand, would be limited by the maximum rate of oxygen delivery to the sediments. We calculated oxygen supply by assuming delivery by Fickian diffusion, using 3 key environmental variables: (1) flow velocity, (2) temperature, and (3) oxygen concentration of the bottom water. Sediment oxygen demand was predicted from measured carbon sedimentation rates as they related to benthic respiration. Our results, if coupled with a predictive model of carbon flux from a net-pen systems (for examples see Gowen & Bradbury 1987, Silver 1992), will yield a tool that can potentially minimize the environmental impacts of salmonid net-pen culture through regulation during the siting process.

MATERIALS AND METHODS

Study sites. We studied 3 salmon net-pen facilities within Maine coastal waters (Fig. 1). The sites were: (1) 2 individual net-pens located on a single lease site in Toothatcher Cove, Swans Island, (2) an erosional site in Eastport and (3) a depositional site in Eastport.
highway' constructed on the sea floor using braided nylon line that led to distal mooring blocks placed such that the ropes were parallel to the main current flows (currents at all sites were predominately bi-directional). Sampling stations were established at regular intervals along this transect. Sediment traps (2 each at the pen edges and at the ambient stations) were deployed at each sampling station and at the time of their recovery sediment cores for benthic metabolism (3 each) and microbial biomass (3 each) were taken. At selected intervals, cores for macrofaunal analysis were also taken. Samples reported in this study were taken directly beneath the down-current edge of a pen or pen complex (proximal sediments) and 100 m from the net-pen or pen complex (ambient or distal sediments). When possible, ambient stations were located perpendicular to the main current flow with respect to the net-pen or pen complex.

**Sampling.** We attempted to sample monthly throughout the year, but rarely were able to complete sampling during the winter months. All sediment traps and sediment cores were taken by SCUBA assisted divers. Cores used for benthic metabolism and microbial biomass were made from clear 6.5 cm ID polycarbonate tubing. Cores were carefully inserted into the sediments to a depth of approximately 10 cm and the top capped with a rubber stopper. The diver then dug a small hole along the outside of the core and slipped a similar stopper under the core. The top stopper was loosened slightly (to allow pressure relief) and the bottom stopper firmly inserted into the core. The top stopper was then also firmly inserted. Cores were

<table>
<thead>
<tr>
<th>Measure</th>
<th>Eastport depositional</th>
<th>Eastport erosional</th>
<th>Swans Island*</th>
</tr>
</thead>
<tbody>
<tr>
<td>Depth (MLW, m)</td>
<td>11.9</td>
<td>11.4</td>
<td>14.3</td>
</tr>
<tr>
<td>Sediment type</td>
<td>Muddy sand</td>
<td>Poorly sorted gravel</td>
<td>Muddy sand</td>
</tr>
<tr>
<td>Current velocities(^b)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Maximum (cm s(^{-1}))</td>
<td>35.8</td>
<td>52.3</td>
<td>17.6</td>
</tr>
<tr>
<td>Average (cm s(^{-1}))</td>
<td>12.6</td>
<td>21.5</td>
<td>2.1</td>
</tr>
<tr>
<td>Minimum 2 h velocity (cm s(^{-1}))</td>
<td>3.1</td>
<td>9.0</td>
<td>0.0</td>
</tr>
<tr>
<td>&gt;25 cm s(^{-1}) (% time)</td>
<td>8.7</td>
<td>61.7</td>
<td>0.0</td>
</tr>
<tr>
<td>Mass accumulation rate (g m(^{-2}) d(^{-1}))</td>
<td>7.29 ± 1.29</td>
<td>14.45 ± 1.70</td>
<td>3.87 ± 0.99</td>
</tr>
<tr>
<td>Maximum carbon flux to the benthos (mmol m(^{-2}) d(^{-1}))</td>
<td>1075.0</td>
<td>253.0</td>
<td>555.0/671.0</td>
</tr>
<tr>
<td>Benthic Index score(^c)</td>
<td>3</td>
<td>1</td>
<td>4/4</td>
</tr>
</tbody>
</table>

\(^a\)Two individual circular pens were studied at this site
\(^b\)Measured 1 m above the sea floor
\(^c\)As determined by the presence of *Begaigaia*-type mats, azonic conditions or other alterations of the benthic environment. '4' represents *Begaigaia*-type mats and/or azonic conditions, '3' modification of benthos and '1' little measurable impact (after Sowles et al. 1994)
tion rates were corrected for sediment resuspension by physical production data. Recording current meters (Endeco Model 105) were moored 1 m from the bottom at all sites and refreshed monthly. These meters provided an average current velocity and direction every 30 min. The minimum current velocity detected by these meters was typically 0.15 to 0.30 cm s\(^{-1}\). Meters were calibrated before each use. Water depths, sediment granulometry, temperature data, the number and weight of fish per pen, the number, size, configuration and orientation of the pens in use, and the size and type of food being fed were (when possible) obtained from the firms involved and corroborated, when possible, by diver observations.

Organic matter deposition. Carbon deposition was measured using sediment traps. Trap construction incorporated the latest technological suggestions (Butman 1986, Butman et al. 1986, Baker et al. 1988) and were designed to minimize sampling bias. Butman has clearly shown that the critical parameter is the relationship of trap diameter to current velocity. As such, we constructed traps from clear polycarbonate tubing, and matched trap diameter to the current velocity at each study site. Traps were either 37.5 cm in height and 10 cm in diameter (aspect ratio 3.75) and, at low-flow study sites, produced an appropriate Reynolds' number \( (R_t = 1 \times 10^4) \) or were 37.5 cm in height and 6.5 cm in diameter (aspect ratio 5.77) and, at high-flow study sites, produced a similar Reynolds' number. Traps were placed in swivel-mounted holders with the trap mouth 1 m above the sediment-water interface. Due to the long trap deployment times (~30 d), material collected in the traps was preserved with salt (TX-10, Akzo Salt, Inc.). The trapped material was sieved (500 μm) to remove animals (e.g. copepods) that swam into the traps and were killed. Sieving also removed detritus >500 μm. The removal of the >500 μm fraction decreased measured ambient and pen carbon flux rates by 10 to 100% (depending on time of year), but the vast majority of this fraction was dead zooplankton. No significant differences in the carbon and nitrogen content of the >500 μm fraction from pen and ambient traps were detected. The amount of material collected in the traps was determined gravimetrically, corrected for any remaining salts, and total organic carbon and nitrogen content determined using a Carlo Erba elemental analyzer. Sediment samples were first treated with fumes of hydrochloric acid to remove any inorganic carbon (usually present as calcium carbonate). Carbon deposition rates were corrected for sediment resuspension by assuming that all ash present within the trap originated from seabed sediment. The amount of carbon entering the trap due to resuspension of sediment was then estimated as:

\[
\text{Carbon}_{\text{resuspended}} = (\text{Trap ash content} \times \text{Seabed sediment carbon content}) \times (\text{Seabed sediment ash content})
\]

The component of variation for duplicate traps was typically less than 10%.

Benthic metabolism. Benthic metabolism was measured as the rate of oxygen consumption and benthic \( \Sigma \text{CO}_2 \) flux. The rate of oxygen consumption was estimated by measuring the loss of oxygen from the overlying water and benthic \( \Sigma \text{CO}_2 \) flux was estimated by measuring accumulation of \( \Sigma \text{CO}_2 \) in the overlying water over time. Undisturbed sediment cores were collected, immediately transported to shore-side facilities, the overlying water removed and replaced with ambient sea water and the cores sealed with a air-tight cap such that approximately 400 ml of ambient sea water overlay the sediment. The underside of the sealing cap was fitted with a small, caged magnetic stir bar and the overlying water was stirred at approximately 5.0 cm s\(^{-1}\). Cores were incubated at ambient temperature for 1 to 4 h (depending on temperature) in the dark. Oxygen concentrations declined 20 to 50% during the incubation period. At the end of the incubation period the cores were unsealed and samples removed for oxygen and \( \Sigma \text{CO}_2 \) determinations and the volume of the remaining water measured. Oxygen concentration was determined by microWinkler titration performed on site and \( \Sigma \text{CO}_2 \) concentration was determined using the potentiometric Gran titration and the method of Edmond (1970) as modified (where appropriate) by Bradshaw et al. (1981). Water samples used for \( \Sigma \text{CO}_2 \) determinations were fixed with HgCl\(_2\) and titrations were performed in the laboratory within 2 wk of collection. The component of variation for triplicate determinations of oxygen uptake rates was typically less than 20% and for triplicate \( \Sigma \text{CO}_2 \) determinations was typically less than 25%.

Data presentation and analysis. Oxygen uptake, \( \Sigma \text{CO}_2 \) production and carbon deposition rates are reported in the literature using a variety of units. To facilitate comparison between the various rates we have reported all rate measurements in units of mmol m\(^{-2}\) d\(^{-1}\). The relationships between \( \text{O}_2 \) uptake, \( \Sigma \text{CO}_2 \) production and carbon deposition were determined by linear regression analysis. As \( \text{O}_2 \) consumption and \( \Sigma \text{CO}_2 \) production were both measured during the determination of benthic metabolism, data from individual cores were compared. The relationship between carbon flux to the sediments and benthic metabolism was also determined by linear regression, but using mean values. The carbon flux to the sediment was determined using sediment traps (mean of 2 traps) deployed for 10 to 30 d prior to the determination.
of benthic metabolism. This value was compared to the mean (3 replicate analyses) O_2 consumption or ΣCO_2 production rates for that station at the date of the recovery of the sediment traps. Only 6 data points out of a possible 11 (4 were lost and 1 excluded) were used for the comparison of carbon sedimentation rate to ΣCO_2 production due to losses of ΣCO_2 production samples and exclusion of stations when Beggiatoa-type mats were present (chemosynthetic activity strongly skewed ΣCO_2 fluxes).

**Oxygen supply to the benthos.** We predicted maximum rates of benthic oxygen supply using Fick's first law:

\[ J = D \frac{(C_w - C_0)}{Z_d} \]  

where \( D \) is the molecular diffusion coefficient of the solute at the ambient temperature (in m^2 h^{-1}); \( C_w \) and \( C_0 \) are the solute concentrations in the bulk water and at the interface, respectively (in mmol m^{-3}); and \( Z_d \) is the thickness of the diffusion boundary layer (in m). The average thickness of the diffusive boundary layer \( (Z_d) \) for current velocities between 0.3 and 7 cm s^{-1} and the O_2 molecular diffusion constant at 15°C \( (D) \) were obtained from the literature (Jorgensen 1989, Broecker & Peng 1974, respectively). Oxygen concentrations at 15°C at saturation and zero were used for \( C_w \) and \( C_0 \), respectively.

**RESULTS**

**Oxygen supply to the benthos**

Field measurements supported the assumption of O_2 saturation for the bulk water. The relationship between flow velocity and the predicted maximum rates of benthic oxygen supply by molecular diffusion was non-linear with greatest increases in flux occurring at the lowest current velocities (Fig. 2). We consider this prediction accurate within a narrow range of key environmental parameters but an underestimation for most sediments. We consider it valuable because many sediments proximal to net-pens are constrained such that the prediction is accurate, at least during key periods of the tidal cycle.

**Benthic carbon deposition rates and benthic metabolism**

Carbon sedimentation rates at ambient sites ranged from 73.4 to 195.8 mmol m^{-2} d^{-1}. Carbon sedimentation rates at sites proximal to net-pens ranged from 108.0 to 1075.0 mmol m^{-2} d^{-1}. Losses during sampling and processing decreased the range of carbon sedimentation rates for which measurements of benthic metabolism were made to between 73.4 and 554.9 mmol m^{-2} d^{-1}. Benthic metabolism was measured in 2 ways in this study: O_2 flux rates and ΣCO_2 production rates. Measured oxygen uptake rates ranged from 29.0 to 100.0 mmol m^{-2} d^{-1} in ambient sediments and 107.9 to 581.8 mmol m^{-2} d^{-1} in sediments proximal to net-pens. Carbon flux rates from sediments distal to the pens ranged from 42 to 167 mmol C m^{-2} d^{-1} while flux rates from sediments proximal to salmon net-pens ranged from 152 to 384 mmol C m^{-2} d^{-1}. There was a strong linear relationship \( (r^2 = 0.965) \) between organic carbon flux to sediment and benthic oxygen consumption (Fig. 3). The molar ratio of carbon flux to the benthos to O_2 flux was near unity and this relationship was dominated by net-pen sediments. There was a strong 1:1 molar ratio \( (slope = 1.084, r^2 = 0.936) \) between the average daily carbon flux to the sediments and the benthic metabolism as estimated by ΣCO_2 production if sediments covered with Beggiatoa-type mats were not included in the analysis (Fig. 4).

Measurements of oxygen uptake and ΣCO_2 production showed the most complex relationship. If sediments both proximal and distal to net-pens with O_2 fluxes of ≤150 mmol O_2 m^{-2} d^{-1} were considered, then a strong linear relationship \( (r^2 = 0.697) \) between these 2 parameters was observed (Fig. 5a). The average respiratory quotient for these stations was 1.41. If all sediments proximal to the net-pens except those covered with Beggiatoa-type mats were included in
the analysis, a linear relationship was again observed \((r^2 = 0.883)\) but the respiratory quotient dropped to 0.98 (Fig 5). Sediments covered with Beggiatoa-type mats showed respiratory quotients ranging from 0.12 to 0.38 and \(\Sigma CO_2\) flux from the sediments decreased with increasing \(O_2\) uptake rates.

**Benthic impact as a function of flow velocity and carbon flux to the benthos**

The 4 net-pen locations were compared on the basis of their maximum carbon sedimentation rates and minimum 2 h flow velocities (Fig 6). The 1:1 molar ratios between carbon deposition and \(O_2\) and \(\Sigma CO_2\) flux (Figs. 3 & 4) observed suggest that waste salmon food and feces are readily degradable and that sedimentation rates are a good predictor of the ultimate oxygen demand generated by the sediments. Minimum 2 h flow velocities were chosen as the key parameter to describe oxygen supply to the benthos because current velocities varied with tidal stage, and 2 h exposures to reduced oxygen and elevated hydrogen sulfide concentrations causes permanent damage to the gill tissues of sensitive infauna (Theede et al. 1969). The loss of macrofauna will shift apparent sediment oxygen diffusion coefficients toward the molecular coefficient — under these conditions Fickian diffusion constrains the maximal rate of oxygen delivery.

The circular net-pens located within Toothacher Cove had minimum 2 h flow velocities of 0.0 cm s\(^{-1}\) and moderate maximum carbon sedimentation rates. Their level of impact, however, was quite high with Beggiatoa-type mats forming beneath both pens (Findlay et al. 1995, authors' unpubl. obs.). In contrast, the Eastport depositional site had a much higher maximum carbon sedimentation rate; yet the benthos maintained
Table 3. Comparison of carbon deposition and sediment metabolic rates measured in this study to those reported for other salmonid production sites. All values reported as mmol m\(^{-2}\) d\(^{-1}\)

<table>
<thead>
<tr>
<th>Parameter</th>
<th>This study</th>
<th>Literature range</th>
<th>Citations</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sediment O(_2) uptake</td>
<td>108–582</td>
<td>20–443</td>
<td>Hall et al. (1990), Hansen et al. (1991), Hargrave et al. (1993), Wu et al. (1994)</td>
</tr>
<tr>
<td>(\Sigma)CO(_2) production</td>
<td>152–384</td>
<td>32–619</td>
<td>Hall et al. (1990), Holmer &amp; Kristensen (1992)</td>
</tr>
</tbody>
</table>

*Calculated assuming a 50% carbon content for particulate organic matter caught in sediment traps.

a modified but aerobic community. Minimum 2 h flow velocities were 3.1 cm s\(^{-1}\). The benthic macrofaunal community beneath this pen system was abundant and diverse (Watling unpubl. data). The Eastport erosional site showed the highest flows (including minimum 2 h flow velocities) and the lowest carbon sedimentation rate. Using a suite of geochemical and biological indices, no benthic impact was detected at this pen system (authors' unpubl. data).

Two sites, Eastport depositional and Eastport erosional, showed carbon deposition rates (in mmol m\(^{-2}\) d\(^{-1}\)) approximately equal to or less than the predicted oxygen supply rate (Fig. 6). These sites showed minimal impacts compared to other Maine sites (Table 2) and most sites from around the world (e.g. Hall et al. 1990, Holmer & Kristensen 1992, Hargrave et al. 1993). The 2 net-pens located in Toothacher Cove showed higher levels of impact than the Eastport sites even though their carbon deposition rates were approximately one-half to two-thirds the rate measured at the Eastport depositional site. We believe that the higher level of impact is due to the fact that oxygen demand exceeded supply. Oxygen supply at the Toothacher Cove locations was constrained by no-flow conditions not present at the Eastport sites.

**DISCUSSION**

Oxygen is delivered to sediments in shallow-water marine environments by a variety of mechanisms—molecular diffusion, bioturbation, and flow and wave advection. We have based our model on the delivery of oxygen by molecular, or Fickian, diffusion alone. We do so based upon the conditions encountered at many salmonid production sites that experience the azoic sediments/Beggiatoa-mat endpoint. Most, if not all, of the sites have minimum flow velocities of <3.0 cm s\(^{-1}\) (for many minimum flow is 0.0 cm s\(^{-1}\)). At these velocities pressure gradients are insufficient to generate advection (Huttel & Gust 1992, Huttel et al. 1996). Wave advection, while possible in 10 to 15 m water depth, is a highly variable phenomenon in the sheltered coves and bays typical of net-pen aquaculture. Bioturbation will decrease in importance as the macrofaunal community experiences organic enrichment (moving from 'reference' through 'peak of opportunists' to 'azoic' sensu Pearson & Rosenberg 1978), and, by definition, will not be a significant factor in azoic sediments. Bioroughness, a factor in both molecular diffusion and advection, will also be quite low at the azoic sediments/Beggiatoa-mat endpoint. Detailed drawings of Beggiatoa mats (Møller et al. 1985) indicate bioroughness on the order of 0.1 mm. This is significantly less than the observed roughness of ~3.0 mm for a photosynthetic microbial mat studied by Jørgensen & Des Marais (1990) in which surface relief was found to

![Fig. 6. Comparison of oxygen supply, as predicted from minimum flow velocity at each site and Fickian diffusion, to oxygen demand predicted from maximum carbon sedimentation rates.](image)
increase diffusive flux by 49%. Studies in shallow or intertidal sediments where apparent sediment diffusion coefficients for oxygen are compared to the molecular coefficients typically note apparent diffusion coefficients 2 to 3 times higher than the molecular coefficient (e.g. Andersen & Helder 1987, Hofman et al. 1991). But these studies were also typically conducted using sediments with healthy macrofauna populations. Deep sea sediments are constrained in a fashion similar to salmonid production sites that experience the azoic sediment/Beggiatoa-mat endpoint (low to no flow, little bioturbation, low roughness); in these cases comparisons of apparent and molecular diffusion coefficients were in good agreement (Reimers et al. 1984, Reimers & Smith 1986).

For our model to be applicable to salmonid production sites around the world, the rates measured in this study should be comparable to those at other sites or, if they are not, then critical biogeochemical processes should operate similarly over the range of rates reported for all sites. Carbon deposition rates measured in this study were within the range reported in the literature but maximum values were 6-fold lower than the maximum value reported by Hall et al. (1990). In spite of this difference in benthic carbon flux, both sediment O₂ uptake and ΔCO₂ production rates measured in this study were comparable to those reported for other salmonid production sites (Table 2). As previously noted, none of the sites used in this study showed an accumulation of wastes beneath the net-pens, while this is commonly observed at sites with higher carbon flux rates (e.g. Hall et al. 1990, Hansen et al. 1991, Hargrave et al. 1993).

The observed correlation in the present study between O₂ flux and sedimentation does not imply that aerobic decomposition is directly responsible for degradation of organic matter at these sites. Aerobic oxidation of organic carbon with a similar redox state as carbohydrate (e.g. CH₂O) will produce 1 mole of CO₂ for every mole of O₂ consumed and a respiratory quotient (mole CO₂ produced/O₂ consumed) of 1.0. Respiratory quotients varied between sites and within sites depending on the rate of carbon flux to the benthos. For sediments, both proximal and distal, receiving organic carbon (fluxes of ≤150 mmol m⁻² d⁻¹), the respiratory quotient averaged 1.4 (Fig. 5a). These values are within the range of respiratory quotients reported for sediments with active benthic fauna from around the world (Hargrave & Phillips 1981, Mackin & Swider 1989, Kristensen et al. 1991, Boucher et al. 1994). In marine sediments, respiratory quotients >1 have been interpreted to indicate active sulfate reduction with reduced sulfur retained at depth within the sediments (as S⁰ or FeS₂, often temporarily, if considered on an annual basis). Under this interpretation aerobic respiration plays a direct role in decomposition of organic matter (Jørgensen 1977, Sørensen et al. 1979, Jørgensen & Sørensen 1985). Recent work suggests that this may not be the case (e.g. Mackin & Swider 1989, Ditoro et al. 1990) and that chemical oxidation of reduced metabolic endproducts produced by anaerobic decomposition of organic matter (frequently referred to as chemical oxygen demand) may account for the majority (>90%) of benthic O₂ fluxes. Holmer & Kristensen (1992) and Hargrave et al. (1993) reported conditions beneath net-pen facilities exhibiting the azoic sediments/Beggiatoa-mat endpoint where chemical oxygen demand was sufficient to account for benthic O₂ flux rates. Oxidation of H₂S, produced during the anaerobic decomposition of organic matter by sulfate-reducing bacteria, is thought to be the major source of the chemical oxygen demand in marine sediments.

Regardless of the pathway of organic matter decomposition, if the supply of oxygen is sufficient then hydrogen sulfide will not accumulate within surface sediments. This lack of accumulation may be the key to maintaining an active macrofaunal community and ultimately maintaining the assimilative capacity of the benthos. Conversely, the azoic sediments/Beggiatoa-mat endpoint is an indication that H₂S is present at the sediment surface (Jørgensen & Revsbech 1983). As sulfide accumulates, a variety of metabolic processes may become inhibited and carbon preservation within the sediments enhanced (Hargrave et al. 1993). Measurements of CO₂ production suggest that sediment metabolism is constrained in sediment impacted by wastes from salmonid production facilities. Maximum rates reported were 387 mmol m⁻² d⁻¹ (this study), 619 mmol m⁻² d⁻¹ (Holmer & Kristensen 1992) and 553 mmol m⁻² d⁻¹ (Hall et al. 1990). These rates were all measured in sediments from environments with periods of zero flow but which differed by an order of magnitude in their rates of carbon sedimentation. If sulfate reduction rates are limited by the toxicity of H₂S to the anaerobic, fermentative bacteria that produce the substrates utilized by sulfate-reducing bacteria, then accumulation of organic matter under net-pen facilities would also be governed by oxygen supply and demand. Measurements of maximum CO₂ production rates suggest that benthic accumulation of carbon would occur at sites with periods of no flow at benthic carbon flux rates of approximately 600 mmol m⁻² d⁻¹.

Most salmonid aquaculture sites around the world report current ranges of 0 to 10 cm s⁻¹. This suggests that these sites would experience significant periods of time with little or no flow and that they would be constrained by oxygen supply as were the Tofthager Cove sites. As such, a sedimentation rate in excess of 200 to 400 mmol C m⁻² d⁻¹ (extrapolated rate at 0.0 cm
and the rate at 0.3 cm s\(^{-1}\) is predicted as the threshold value at which the azoic sediments/Beggiatoa-mat endpoint would be observed. In 2 cases where sedimentation rates exceeded 400 mmol C m\(^{-2}\) d\(^{-1}\) and no-flow conditions were reported (Hall et al. 1990, Hansen et al. 1991), this endpoint was observed. Hargrave (1995) reported sedimentation rates of 100 to 117 mmol m\(^{-2}\) d\(^{-1}\) at the pen edge and 650 mmol m\(^{-2}\) d\(^{-1}\) beneath the pens at the L'Etang Inlet site (New Brunswick, Canada). Impacts beneath the pens were consistent with the azoic sediments/Beggiatoa-mat endpoint but at the pen edge they were less severe (Hargrave et al. 1993). Ye et al. (1991) reported carbon sedimentation rates in excess of 400 mmol m\(^{-2}\) d\(^{-1}\) but only documented low to moderate benthic impacts at their site. No-flow conditions were not reported and average current velocities would have maintained oxygen supply in excess of oxygen demand as predicted from sedimentation rates.

**Model summary**

We have developed our model of benthic impact associated with net-pen aquaculture using 2 parameters: oxygen supply to the sediments and sediment oxygen demand (both expressed as mmol m\(^{-2}\) d\(^{-1}\)). The relationship between these 2 parameters can be examined in 2 ways, either graphically as in Fig 6 or by the calculation of an index of impact (I) given as:

\[
I = \frac{O_2\text{ supply}}{O_2\text{ demand}}
\]  

(2)

We calculate oxygen supply by assuming delivery by Fickian diffusion, using Eq. (1) and 3 key environmental variables: (1) flow velocity as the determinate of diffusive boundary layer thickness, (2) temperature, and (3) oxygen concentration of the bottom water. For 15°C and saturated bottom water we estimate:

\[
O_2\text{ supply (mmol m}^2\text{ d}^{-1} = 736.3 + 672.5 \log(v) \]

(3)

where \(v\) is flow velocity in cm s\(^{-1}\) (Fig. 2).

Sediment oxygen demand is predicted from measured carbon sedimentation rates. Linear regression analysis of data from Maine production facilities yielded the equation used to convert carbon sedimentation rates to sediment oxygen demand, that is:

\[
O_2\text{ demand (mmol m}^2\text{ d}^{-1} = 1.07x - 32.6
\]

(4)

where \(x\) is carbon flux to the benthos in mmol m\(^{-2}\) d\(^{-1}\) (Fig. 3).

It is this balance between supply and demand that produces the prediction of impact. If the supply of oxygen is in excess of demand, then \(I > 1.0\), and impacts are minimal. If supply and demand are near unity, then \(I = 1.0\), and impacts are moderate. If demand is in excess of supply, then \(I < 1.0\), and the sediments exhibit the azoic sediment/Beggiatoa-mat endpoint. We suspect, but were unable to investigate since Maine does not have sites with accumulated organic matter, that \(I\) will be much less than 1.0 at sites where wastes form distinct deposits beneath the pens.

**Conclusions**

Flow velocities have long been recognized as important in predicting benthic impact associated with net-pen aquaculture, but only as a mechanism by which waste feed and feces are distributed within the environment. Our study clearly demonstrates the pivotal role of flow in reducing the benthic impact of net-pens. Increased flow velocity not only reduces carbon sedimentation rates, but also results in increased sediment oxygen delivery, in turn allowing greater benthic degradation of the waste stream. We hypothesize that the non-linear increase in oxygen delivery with increasing flow was the major cause of the inability to predict impacts across a variety of sites.

As \(O_2\) supply is derived from an assumption that impacted sediments are constrained to oxygen delivery by Fickian diffusion and that this rate is under the control of a few key environmental variables, it is likely that the calculation of \(O_2\) supply will be robust across a variety of geographical or hydrological regions. This may or may not be the case for \(O_2\) demand. The calculation of \(O_2\) demand is based on an empirical relationship developed for a series of sites from coastal Maine and none of the sites used showed significant accumulation of organic wastes beneath the pens. Data from Hansen et al. (1991) suggests that a different (\(-0.5:1\)) molar oxygen demand may be needed for long-established facilities with visible waste accumulation. Regardless, if an accurate a priori prediction of benthic carbon flux can be made for proposed net-pen sites, then this model will be a useful tool for predicting levels of benthic impact. Studies of \(\Sigma CO_2\) flux rates suggest that oxygen supply to the sediments and its role in regulating sediment sulfide concentrations might be useful in predicting carbon sedimentation rates at which accumulation of waste deposits would occur. This model may have general applicability to other types of organic enrichment if correlative studies linking carbon sedimentation rates to benthic \(O_2\) flux rates can be performed (as was done in this study) or if organic matter decay rate constants are known or can be determined for the introduced organic material.

This model synthesizes the understanding that sediment oxygen delivery and sediment oxygen consumption both play a role in determining benthic commu-
nity structure and, if these 2 parameters are measured and compared, then the level of impact can be predicted for net-pen aquaculture. Key parameters become benthic carbon flux and oxygen delivery rates; comparison of these rates reveal whether the available carbon will be consumed aerobically with minimal environmental impact or anaerobically inducing environmental impact.

Acknowledgements. This research was supported in part by grants from EPA Office of Exploratory Research (Assistance ID number R-817196-01-0), EPA District 1 (Assistance ID number X002387-01-0), National Marine Fisheries Service (U.S. Department of Commerce, National Oceanic and Atmospheric Administration, award number NA16FL0067-01), the UMI/UNH Sea Grant College Program (U.S. Department of Commerce, National Oceanic and Atmospheric Administration, project numbers R/FMD-215 and R/FMD-225), the Marine Aquaculture Innovation Center (project numbers 99-6, 91-13 and 93-08) and the PADI Foundation. We gratefully acknowledge the assistance of a great number of people whose efforts helped bring this project to fruition. These include E. Harrison, R. Doering, T. Sawyer, J. B. Pelletier, L. Klippel, L. McCann, M. Tarrentino, W. Tripp, J. Anderson, J. Fay, D. Meisenheimer, and C. Heimng and B. Tarbox of Inter-tide Corp. for assistance with diving and field operations. M. Tarrentino and J. B. Pelletier ably served as dive safety officers and T. Sawyer directed field operations in the absence of R.H.R. Doering, K. Hardy and L. Schick performed the CHN analyses and K. Hardy, L. Klippel and J. B. Pelletier processed the sediment trap samples.

LITERATURE CITED


This article was submitted to the editor


Manuscript received: September 10, 1996
Revised version accepted: June 5, 1997