Review

Benthic microalgal production in the Arctic: applied methods and status of the current database

Ronnie N. Glud¹,²,*, Jana Woelfel³, Ulf Karsten³, Michael Kühl² and Søren Rysgaard⁴

¹ Dustaffnage Marine Laboratory, Scottish Association of Marine Sciences, Oban, Scotland PA37 1QA, UK, e-mail: Ronnie.Glud@sams.ac.uk
² Marine Biological Laboratory, Department of Biology, University of Copenhagen, Strandpromadenen 5, 3000 Helsingør, Denmark
³ Institute of Biological Sciences, Applied Ecology, University of Rostock, Albert-Einstein-Str 3, 18051 Rostock, Germany
⁴ Greenland Institute of Natural Resources, Box 570, 3900 Nuuk, Greenland

* Corresponding author

Abstract

The current database on benthic microalgal production in Arctic waters comprises 10 peer-reviewed and three unpublished studies. Here, we compile and discuss these datasets, along with the applied measurement approaches used. The latter is essential for robust comparative analysis and to clarify the often very confusing terminology in the existing literature. Our compilation demonstrates that i) benthic microalgae contribute significantly to coastal ecosystem production in the Arctic, and ii) benthic microalgal production on average exceeds pelagic productivity by a factor of 1.5 for water depths down to 30 m. We have established relationships between irradiance, water depth and benthic microalgal productivity that can be used to extrapolate results from quantitative experimental studies to the entire Arctic region. Two different approaches estimated that current benthic microalgal production in the Arctic is between 1.1 and 1.6×10⁶ tons C year⁻¹. Climate change is expected to increase the overall primary production and affect the balance between pelagic and benthic productivity in the Arctic. It is therefore imperative to get better quantitative understanding of the relationship between increased freshwater run-off, shrinking sea-ice cover, light availability and benthic primary production to assess future impact on the Arctic food web and trophic coupling.

Keywords: Arctic; benthic microalgae; benthic primary production; photosynthesis.

Introduction

The Arctic oceans, including all marine water bodies at latitudes above the Arctic Circle, cover >20×10⁶ km² and encompass ~25% of the global coastal region (areas with water depths <200 m) (Menard and Smith 1966, Jakobsson et al. 2008). The Arctic coastal regions cover ~5.8–6.1×10⁶ km² with an average water depth of 80 m (Gattuso et al. 2006, Jakobsson et al. 2008) and a significant fraction of this area can be expected to accommodate benthic primary production. Gattuso et al. (2006) estimated that on average, 25% of the Arctic coastal seabed receives >1% of the surface down welling irradiance during the five summer months, with a much larger fraction expected to occasionally experience irradiances of this magnitude.

An increasing number of studies suggest that benthic microalgae contribute significantly to subtidal coastal ecosystem production (Charpy-Roubaud and Sournia 1990, Meyercordt et al. 1999, Nelson et al. 1999, Jahnke et al. 2000, Jahnke 2005, Glud et al. 2008). Furthermore, studies from lower latitudes have documented the importance of benthic microphytes for supporting shallow water food-webs (e.g., Middelburg et al. 2000). In an extensive review of 85 studies, Cahoon (1999) concluded that previous estimates on benthic microalgal productivity had markedly underestimated their importance for marine ecosystem production – especially in oligotrophic systems. Overall, this author provided an annual global estimate of 5×10⁸ tons C for the neritic microalgal primary production. Cahoon (op. cit.) also showed that, whereas some areas were relatively well studied (i.e., temperate intertidal and eutrophicated systems), others were grossly under-sampled, especially the Arctic region for which at that time there were only two studies! The polar coastal areas differ from the better studied temperate regions in being exposed to ice-cover and darkness during extensive periods of the year. Furthermore, they are pristine and experience relatively low temperatures. The seabed is often exposed to ice-mediated erosion and erratic, massive inputs of erosion material during spring (Rachold et al. 2004, Zacher et al. 2009). Therefore, findings from lower latitude systems cannot be extrapolated a priori to polar ecosystems.

Camera-based mapping of the sea-bed over larger areas in the Arctic often shows an extensive and dense cover of microalgae that is intensively grazed by macrofauna (Figure 1). One striking observation provided by such photo documentation is the pronounced small-scale patchiness, which must be accounted for when evaluating data on benthic productivity. Closer inspection of sea-bed samples shows that the microalgal cover is often dominated by pennate diatoms and in some cases dinoflagellates or Cyanobacteria; the first group appears to dominate Arctic microphytobenthos (Vetrov and Romankevich 2004 and references therein). In the follow-
Figure 1  *In situ* photographs of benthic microalgal coverage. (A) in NW Greenland (72°22.53′ N, 55°33.37′ W) at 18 m of water depth; (B) in NE Greenland (74°18.59′ N, 20°14.48′ W) in 20 m water depth and (C) in NE Greenland (74°18.59′ N, 20°14.74′ W) in 30 m water depth. In all cases the brown patches indicate the presence of pennate diatoms (the lower panel is dominated by the occurrence of stones covered with coralline red algae). Each image covers an area of 40×30 cm.

Photograph made available by P.B. Christensen, M. Sejr, and P. Batty and reproduced by kind permission of the owners.
labeling in the microenvironment of the active microalgae. Intact microphytobenthic communities are characterized by extensive temporal and spatial variability and by steep concentration gradients of solutes and light. This makes it very difficult to determine the specific DIC labeling experienced by the active microalgae, which is most certainly not constant during a given incubation (Revsbech et al. 1981, de Beer et al. 1997). There is currently no direct way to determine the specific labeling of inorganic carbon pool at the relevant scale, and estimating an average value of this crucial parameter essentially relies on qualified guessing. Due to this problem, the $^{14}$C approach is nowadays rarely used to assess benthic productivity.

In principle, the $^{14}$C approach quantifies the gross primary production, but depending on the incubation time, some of the fixed $^{14}$C can be respired by the microalgae themselves or by heterotrophic bacteria taking advantage of leaking photosynthetic products and exudates. It is very difficult to assess the extent to which this takes place during a given incubation, but it is well-established that heterotrophic activity in natural benthic communities is markedly stimulated by light due to a rapid turnover of freshly produced organic material (Epping and Jørgensen 1996, Kühl et al. 1996, Fenchel and Glud 2000). Therefore, although dependent on the incubation time, the $^{14}$C approach generally underestimates true gross photosynthetic activity. By extending the incubation to 24 h during a natural light/dark cycle it could be argued that the approach expresses the daily net primary production. However, most (and all Arctic) incubations apply an incubation of ~3–5 h and the gross primary production is calculated after subtracting any $^{14}$C fixation rate of parallel dark incubations (Matheke and Horner 1974). In the following, we abbreviate gross primary production as determined by this approach as GPPt ("t" for tracer).

While the benthic $^{14}$C incubation technique is rarely used anymore, many studies in Arctic waters are based on this approach and represent a significant contribution to the available database. We have identified seven Arctic studies that have applied the $^{14}$C approach to in situ benthic chambers deployed over a depth range of 3–25 m (Table 1). Despite the very limited database and the inherent scatter, maximal rates occurred at very shallow water depths (<6 m), with little activity at the deepest measuring sites (20–25 m), presumably as light availability approached the limit of microalgal photosynthetic activity.
Table 1 Benthic microalgal gross primary production in Arctic ecosystems estimated from in situ chamber incubations enriched by 14C-labeled DIC (GPPt).

<table>
<thead>
<tr>
<th>Activity range (mmol C m⁻² day⁻¹)</th>
<th>Mean (mmol C m⁻³ day⁻¹)</th>
<th>Chl a (mg m⁻²)</th>
<th>Depth (m)</th>
<th>References</th>
</tr>
</thead>
<tbody>
<tr>
<td>37.6–114.0</td>
<td>75.6</td>
<td>~130–380</td>
<td>~5</td>
<td>Matheke and Horner (1974)</td>
</tr>
<tr>
<td>0.0–0.2</td>
<td>0.1</td>
<td>1–28</td>
<td>7</td>
<td>Horner and Schrader (1982)</td>
</tr>
<tr>
<td>2.0–101.2</td>
<td>32.8</td>
<td>Nil</td>
<td>3</td>
<td>Kuznetsov (1991)</td>
</tr>
<tr>
<td>0.2–100</td>
<td>30.0</td>
<td>Nil</td>
<td>10</td>
<td>Kuznetsov (1991)</td>
</tr>
<tr>
<td>0.0–95.2</td>
<td>19.8</td>
<td>Nil</td>
<td>15</td>
<td>Kuznetsov (1991)</td>
</tr>
<tr>
<td>3.6–19.6</td>
<td>12.0</td>
<td>18–72</td>
<td>20</td>
<td>Kuznetsov (2002)</td>
</tr>
<tr>
<td>0.6–4.1</td>
<td>NI</td>
<td>40–940</td>
<td>1–25</td>
<td>Bondarchuk (1974, 1980)</td>
</tr>
<tr>
<td>Ni</td>
<td>4.6</td>
<td>2–640</td>
<td>5–25</td>
<td>Kuznetsov et al. 1998</td>
</tr>
</tbody>
</table>

The studies are all seasonal, but the table includes data only from the sea ice-free period. The Chl a data represent measurements for the uppermost centimeter, but in some instances, only mean values were assessable. Nil, no information is available.

Total exchange rates of O₂ and DIC

Generally speaking, the most widely applied procedure to assess the benthic production (or degradation) of organic material is the “whole core incubation” approach, wherein chambers or core liners holding an intact sediment core are either incubated in the laboratory under controlled light conditions or in situ taking advantage of natural irradiance (Cahoon and Cooke 1992, Jahneke et al. 2000, Glud et al. 2008). The net primary production during illumination can then be inferred from the gradual O₂ accumulation in the overlying water and/or the concurrent decline in the DIC concentration. The approach is relatively simple, and when larger sediment areas are enclosed, the inherent small-scale patchiness in microalgal biomass can, at least to some extent, be accounted for. One concern is, however, that chambers or cores change the local hydrodynamics. Efficient stirring or mixing of the enclosed water volume during incubations is thus essential and has, in various studies, been accommodated by pumps, impellers, discs or stirrers (Pamatmat and Fenton 1968, Cahoon 1988, Huettel and Gust 1992). It is beyond the scope of this review to discuss pros and cons of various chamber and stirring designs or in situ vs. laboratory-based measurements (for recent reviews, see Tengberg et al. 2005, Glud 2008), but the effect of any imposed stirring on the measured solute exchange rate must be acknowledged.

In benthic communities with a high O₂ production (or consumption) rate, the thickness of the diffuse boundary layer (DBL) can significantly affect the net exchange of O₂ (Jørgensen and des Marais 1990, Larkum et al. 2003, Glud et al. 2007) and, therefore, affect the derived net photosynthesis. For example, in a benthic microalgal mat, the O₂ exchange increased 5–6-fold as the measured DBL thickness was decreased by a factor of ~4 (Figure 3).

Total O₂ or DIC exchange rates measured during illumination quantify the net-photosynthesis (NPPc, c for chambers) of the benthic community, including oxygen consumption related to fauna and microbial respiration. The concurrent gross photosynthetic activity can thus be substantially higher. A common procedure for estimating the gross photosynthesis is to add the exchange rates measured in darkness (or in parallel incubations with opaque chambers) to that of the light incubations (e.g., Eyre and Ferguson 2002). This is a pragmatic approach, but the values should be interpreted with caution for at least two reasons: 1) the infauna of shallow water sediments frequently exhibit a daily variation in their activity level and feeding mode that affects the benthic O₂ consumption rate (Rosenberg and Lundberg 2004, Wenzhöfer and Glud 2004), and 2) microbial respiration is typically enhanced in light (Epping and Jørgensen 1996; see also below). Therefore, this approach generally underestimates the true gross photosynthetic activity.

Most studies based on chamber or core incubations infer benthic primary production from change in the O₂ concentration. However, electrons released from the light-induced splitting of water (i.e., the O₂ formation) can be channeled to a range of other processes in cell-housekeeping, rather than CO₂ fixation (Raven and Bear-dall 1981, Badger et al. 2000, Wagner et al. 2006). Provided any precipitation or dissolution of carbonate can be accounted for, a more direct procedure to quantify primary production could be to measure the daily (24 h) DIC consumption rate. But as O₂ is far easier to measure than DIC, the O₂ exchange measurements...
remain the preferred procedure and O₂ exchange is subsequently transformed into a CO₂ fixation rate assuming a photosynthetic quotient (PQ). Most measurements of PQ for integrated benthic communities, quantified as the ratio between the exchange of O₂ vs. DIC, range between 0.9 and 1.3, largely depending on the light and nutrient availability (Cammen 1991, Cahoon and Cooke 1992, Longphuirt et al. 2007, Taddei et al. 2008). Generally a value of 1.2 is applied to benthic communities, i.e., 1.0 carbon atom is fixed per 1.2 O₂ molecule produced. Based on 50 incubations of sediment cores collected over a water depth range of 5–30 m and exposed to light levels between 6 and 100 µmol photons m⁻² s⁻¹, the community PQ of a high Arctic sediment averaged 1.19±0.48 (Glud et al. 2002). Based on these measurements, and in order to be consistent with most studies conducted at lower latitudes, we have chosen to apply a conversion factor of 1.2 to derive the benthic gross primary production (GPPc) from the sum of O₂ exchange measured in darkness and light with benthic incubation chambers [the respiratory quotient (RQ) is thereby assumed to be the reciprocal of PQ].

We have identified two in situ studies (Kuznetsov and Strogaya 1989, Kuznetsov 2005) from the Arctic that used chambers for quantifying benthic microphytic production (Table 2). The values align with the GPPt data of Table 1, but rather than a straightforward depth relationship, maximal phototrophic activity occurred at intermediary water depths.

One published High Arctic study used laboratory core incubations under well-defined irradiance to quantify the productivity of benthic microalgae (Glud et al. 2002). These data are plotted against irradiance along with some instances only mean values were available for assessment. NI: no information is available.

The Chl a data represent measurements for the uppermost centimeter: in some instances only mean values were available. A photosynthetic quotient (PQ) is thereby assumed to be the reciprocal of PQ.

We have identified two in situ studies (Kuznetsov and Strogaya 1989, Kuznetsov 2005) from the Arctic that used chambers for quantifying benthic microphytic production (Table 2). The values align with the GPPt data of Table 1, but rather than a straightforward depth relationship, maximal phototrophic activity occurred at intermediary water depths.

One published High Arctic study used laboratory core incubations under well-defined irradiance to quantify the productivity of benthic microalgae (Glud et al. 2002). These data are plotted against irradiance along with some instances only mean values were available. A photosynthetic quotient (PQ) is thereby assumed to be the reciprocal of PQ.

We have identified two in situ studies (Kuznetsov and Strogaya 1989, Kuznetsov 2005) from the Arctic that used chambers for quantifying benthic microphytic production (Table 2). The values align with the GPPt data of Table 1, but rather than a straightforward depth relationship, maximal phototrophic activity occurred at intermediary water depths.

One published High Arctic study used laboratory core incubations under well-defined irradiance to quantify the productivity of benthic microalgae (Glud et al. 2002). These data are plotted against irradiance along with some instances only mean values were available. A photosynthetic quotient (PQ) is thereby assumed to be the reciprocal of PQ.

We have identified two in situ studies (Kuznetsov and Strogaya 1989, Kuznetsov 2005) from the Arctic that used chambers for quantifying benthic microphytic production (Table 2). The values align with the GPPt data of Table 1, but rather than a straightforward depth relationship, maximal phototrophic activity occurred at intermediary water depths.

One published High Arctic study used laboratory core incubations under well-defined irradiance to quantify the productivity of benthic microalgae (Glud et al. 2002). These data are plotted against irradiance along with some instances only mean values were available. A photosynthetic quotient (PQ) is thereby assumed to be the reciprocal of PQ.

We have identified two in situ studies (Kuznetsov and Strogaya 1989, Kuznetsov 2005) from the Arctic that used chambers for quantifying benthic microphytic production (Table 2). The values align with the GPPt data of Table 1, but rather than a straightforward depth relationship, maximal phototrophic activity occurred at intermediary water depths.

One published High Arctic study used laboratory core incubations under well-defined irradiance to quantify the productivity of benthic microalgae (Glud et al. 2002). These data are plotted against irradiance along with some instances only mean values were available. A photosynthetic quotient (PQ) is thereby assumed to be the reciprocal of PQ.
Table 3  Benthic microalgal gross primary production in open-water Arctic ecosystems as estimated from total O₂ exchange measurements (GPPc) of laboratory-based core incubations.

<table>
<thead>
<tr>
<th>Activity range (mmol C m⁻² day⁻¹)</th>
<th>Irradiance (µmol photons m⁻² s⁻¹)</th>
<th>“In situ” mean* Chl a (mg m⁻²)</th>
<th>Depth (m)</th>
<th>References</th>
</tr>
</thead>
<tbody>
<tr>
<td>36–90</td>
<td>95</td>
<td>112</td>
<td>136</td>
<td>5</td>
</tr>
<tr>
<td>16–46</td>
<td>95</td>
<td>25</td>
<td>30</td>
<td>10</td>
</tr>
<tr>
<td>10–36</td>
<td>95</td>
<td>10</td>
<td>11</td>
<td>15</td>
</tr>
<tr>
<td>10–46</td>
<td>95</td>
<td>6</td>
<td>8</td>
<td>20</td>
</tr>
<tr>
<td>19–26</td>
<td>95</td>
<td>2</td>
<td>2</td>
<td>30</td>
</tr>
<tr>
<td>15–21</td>
<td>95</td>
<td>29</td>
<td>34</td>
<td>5</td>
</tr>
<tr>
<td>19–25</td>
<td>95</td>
<td>16</td>
<td>19</td>
<td>10</td>
</tr>
<tr>
<td>1–3</td>
<td>95</td>
<td>1</td>
<td>1</td>
<td>15</td>
</tr>
<tr>
<td>1–3</td>
<td>95</td>
<td>1</td>
<td>1</td>
<td>20</td>
</tr>
<tr>
<td>0–5</td>
<td>95</td>
<td>&lt;1</td>
<td>&lt;1</td>
<td>30</td>
</tr>
<tr>
<td>18–34</td>
<td>140</td>
<td>36</td>
<td>3</td>
<td>1</td>
</tr>
<tr>
<td>40–54</td>
<td>200</td>
<td>42</td>
<td>NI</td>
<td>2</td>
</tr>
<tr>
<td>1–21</td>
<td>21–91</td>
<td>34</td>
<td>75</td>
<td>5</td>
</tr>
<tr>
<td>1–24</td>
<td>6–69</td>
<td>24</td>
<td>66</td>
<td>10</td>
</tr>
<tr>
<td>1–43</td>
<td>12–87</td>
<td>12</td>
<td>186</td>
<td>20</td>
</tr>
<tr>
<td>1–27</td>
<td>11–53</td>
<td>5</td>
<td>92</td>
<td>30</td>
</tr>
</tbody>
</table>

The Chl a data represent measurements for the uppermost centimeter; in some instances only mean values were available for assessment. NI: no information is available.

* The actual laboratory-based measurements were extrapolated to estimate average in situ irradiance at the sampling sites using recorded light levels and light extinction coefficients of the water column during the open-water period.

Oxygen microsensor measurements

The prime advantage of using chambers or cores (and eddy correlation) for quantifying benthic exchange rates is that their application is relatively simple and averages out small scale variations. However, such methods represent a “black-box” approach and only provide limited insight into the vertical and horizontal activity distribution or into the microenvironment in which the microbenthic phototrophic activity takes place. Oxygen microelectrode measurements allow a very detailed characterization of the distribution, production and consumption of O₂ at a given point in time (Revsbech et al. 1981, Epping et al. 1999) and multiple measurements thereby complement chamber incubations well. From measured microprofiles, the diffusive export of O₂ from the photic zone can be quantified as the sum of the upward and downward flux, and when converted into carbon equivalents, this represents the net primary production of the community (NPPm) (Figure 5). The upward flux is essentially what is quantified in the NPPc measured by chamber or core incubations and generally accounts for 70–90% of the NPPm (Epping and Jørgensen 1996, Kühl et al. 1996, Wenzhöfer et al. 2000, Christensen et al. 2003).

Oxygen microsensors also allow quantification of gross primary production (GPPm) by the light-dark-shift technique (Revsbech and Jørgensen 1983). Basically, this method calculates gross photosynthesis from the rate at which the O₂ concentration at a given point declines immediately after onset of darkness. The approach assumes that the O₂ gradients and respiration remain unaffected during the light/dark shift, and detailed investigations have proven that this is essentially correct, as long as the O₂ decline is determined within the first second of darkness (Glud et al. 1992, Lassen et al. 1998). This approach actually resolves the true gross primary production of the community, provided the applied PQ reflects the relationship between O₂ production and DIC fixation under the given conditions.

Figure 5 shows an example of the level of detail that can be obtained by microsensor measurements and degree of complexity in deriving an estimate of benthic productivity from O₂ exchange measurements with ben-

Figure 5  Oxygen microprofiles measured in a diatom-domi-

nated community of benthic microalgae. Black symbols represent measurements in darkness, while open symbols are measurements at a down-welling irradiance of 450 µmol photons m⁻² s⁻¹. Gray bars represent the gross photosynthesis as measured by the “light-dark-shift” technique. Jₚ represents the upward-directed O₂ flux as calculated from (Jₚ=Do dC/dz), where Do is the molecular diffusion coefficient for O₂ under the experimental conditions, and C is the O₂ concentration at depth Z within the diffuse boundary layer (DBL); the same approach is applied to estimate O₂ consumption from the microprofile measured in darkness). The downward O₂ flux is quantified by the same relation, but this time applying the tortuosity-corrected diffusion coefficient of the sediment matrix. The sum of Jₚ and Jₙₚ provides the net photosynthetic activity of the photic zone.
thic chambers or whole cores. The benthic O₂ consumption in darkness amounted to 13.0 mmol m⁻² day⁻¹, while the O₂ efflux from the photic zone (the upper 1.6 mm) during light equaled 81.7 mmol m⁻² day⁻¹, of which 74% (60.1 mmol m⁻² day⁻¹) diffused upwards into the overlying water and the residual 21.6 mmol m⁻² day⁻¹ sustained O₂ consumption in the deeper sediment layers (Figure 5). The concurrent depth-integrated gross production of O₂, measured by the light-dark shift technique, was 117.7 mmol m⁻² day⁻¹, meaning that the respiration within the illuminated photic zone (the upper 1.6 mm) was 36.0 mmol m⁻² day⁻¹ (117.7–81.7). The benthic O₂ consumption during light thus amounted to 57.6 mmol m⁻² day⁻¹ (36.0+21.6), which is >4 times higher than O₂ consumption in darkness. The stimulated activity during light was caused by two factors i) an increased O₂ consumption within the photic zone and ii) a deeper oxic penetration zone within which O₂ was consumed. If we apply the calculation procedure that is generally applied to chamber incubation, the “GPPc” is the sum of the O₂ consumption in darkness and the upward O₂ release in light (converted into C equivalents), which sums to 60.9 mmol C m⁻² day⁻¹ [(13+60.1)/1.2]. In reality, the NPPm is 68.0 mmol C m⁻² day⁻¹ (81.7/1.2) and the GPPm is 98.0 mmol C m⁻² day⁻¹ (117.7/1.2). This example illustrates how the widely-applied chamber approach underestimates actual benthic productivity and how confusing use of the terms net and gross primary production is in the existing literature. The example also illustrates the detailed insights that can be provided into O₂ turnover at a given spot using microsensor measurement. The approach is, however, very time consuming – a set of microprofiles resolving the O₂ distribution and the GPPm at a given spot for a single level of irradiance typically takes about 40–50 min, and given the natural variability of the sea-bed it is a non-trivial task to extrapolate the findings from one or a few measurements to a larger area. This remains the prime limitation of the microsensor measuring approach.

To the best of our knowledge, there are only two microsensor studies on benthic microalgal activity in the Arctic (Glud et al. 2002, Hancke and Glud 2004). In the first, sediment cores were recovered from 10 to 30 m water depth and were then incubated under similar light, temperature and salinity conditions in the laboratory for two days. Subsequently, O₂ microprofiles were measured in the diatom cover, and the compiled data exhibited a good PE-relationship, without any light-inhibition at the irradiances applied (Figure 6). Benthic microalgal communities rarely exhibit light inhibition because i) a downward migration of motile cells counteracts inhibiting light levels at the surface due to the extremely steep light gradients in such communities (Kühl et al. 1997), and ii) a gradual expansion of the photic zone with increasing irradiance compensates for any potential activity decline at the surface (Epping and Jørgensen 1996, Kühl et al. 1996, Christensen et al. 2003).

The fitted Platt equation of Figure 6 shows a photosynthetic capacity (Pₘ) of 67 mmol m⁻² day⁻¹ and an “index of light adaptation” (Eₜ) of 27 μmol photons m⁻² s⁻¹ for net photosynthesis (NPPc). When comparing with the GPPc-Platt relation (Figure 4), it is important to realize that the two relationships are based on different techniques and, as explained above, this causes the otherwise inconsistent observation that Pₘ for gross primary production (GPPc as derived from core incubations) is lower than the Pₘ for the net primary production (NPPm as derived from microprofile measurements). The light compensation point, where the benthic O₂ consumption balances the O₂ production (i.e., the light level at which the net O₂ exchange is zero mmol m⁻² day⁻¹) amounted to 4.7 μmol photons m⁻² s⁻¹. This value essentially defines the threshold irradiance required to sustain net primary production of the communities investigated at the given experimental conditions. The data in Figure 6 were measured in patches of benthic microphytes and ignored patches of bare sediment. Accounting for in situ coverage of benthic microalgae and mean daily light availability at the respective water depths, NPPm for the study was calculated and is presented in Table 4.

The only other Arctic microsensor study on benthic microalgae was performed on sediment cores recovered below snow covered sea-ice with very low phototrophic biomass (the sediment surface Chl a level was 2.7 mg m⁻²; Hancke and Glud 2004). This study focused on short-term temperature effects on respiration and photosynthesis of benthic communities dominated by diatoms rather than on quantifying benthic productivity. It was shown that both respiration and photosynthesis

---

Table 4 Benthic microalgal net primary production in an open-water Arctic ecosystem as estimated from microsensor measurements (NPPm) in the laboratory and extrapolated to in situ activity using information on light availability and benthic microalgal cover (BMC).

<table>
<thead>
<tr>
<th>Mean (mmol C m⁻² day⁻¹)</th>
<th>Depth (m)</th>
<th>In situ BMC (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>32.0</td>
<td>5</td>
<td>23</td>
</tr>
<tr>
<td>31.3</td>
<td>10</td>
<td>26</td>
</tr>
<tr>
<td>14.8</td>
<td>20</td>
<td>73</td>
</tr>
<tr>
<td>9.3</td>
<td>30</td>
<td>36</td>
</tr>
</tbody>
</table>

Glud et al. (2002).
increased with temperature, but that a stronger $Q_{10}$ response of heterotrophic activity gradually lead to reduced net benthic productivity as the temperature increased (Hancock and Glud 2004). The extent to which this observation is the result of the short-term nature of the experiment (days to weeks), and whether seasonal increases in temperature can shift shallow-water Arctic sediments into a more heterotroph-dominated status in late summer remain to be investigated.

Discussion

Relative importance of benthic vs. pelagic microalgal productivity

Growing directly at the sediment surface, benthic microalgae can exploit nutrients released by the underlying biogeochemical mineralization processes and can thus deprive the pelagic community of nutrients. In contrast, pelagic photrophs can better exploit the down-welling irradiance as compared to communities constrained to a narrow zone on the sediment surface. Thus, nutrient availability often regulates the relative importance of pelagic vs. benthic microalgal productivity. Eutrophic settings favor pelagic productivity, while oligotrophic settings favor benthic productivity (Charpy-Roubaud and Sournia 1990, MacIntyre et al. 1996). Even though rivers can carry nutrient-enriched water and induce plumes of stimulated pelagic production that reduce the benthic light availability locally (Parsons et al. 1988, Springer and McRoy 1993), Arctic coastal waters are pristine with low nutrient levels. Consequently, they can be expected to host a relatively large benthic productivity.

Several of the original benthic studies in Tables 1–4 performed parallel measurements of the pelagic productivity and have provided estimates on the relative importance of pelagic vs. benthic microalgal productivity. Eutrophic settings favor pelagic productivity, while oligotrophic settings favor benthic productivity (Charpy-Roubaud and Sournia 1990, MacIntyre et al. 1996). Even though rivers can carry nutrient-enriched water and induce plumes of stimulated pelagic production that reduce the benthic light availability locally (Parsons et al. 1988, Springer and McRoy 1993), Arctic coastal waters are pristine with low nutrient levels. Consequently, they can be expected to host a relatively large benthic productivity.

Table 5 Relative proportion of benthic microalgal primary production vs. pelagic primary production in Arctic coastal environments.

<table>
<thead>
<tr>
<th>Benthic contribution (%)</th>
<th>Water depth (m)</th>
<th>Study</th>
</tr>
</thead>
<tbody>
<tr>
<td>76</td>
<td>5</td>
<td>Matheke and Horner (1974)</td>
</tr>
<tr>
<td>3</td>
<td>3</td>
<td>Horner and Schrader (1982)</td>
</tr>
<tr>
<td>75</td>
<td>3</td>
<td>Kuznetsova (1991)</td>
</tr>
<tr>
<td>45</td>
<td>10</td>
<td>Kuznetsova (1991)</td>
</tr>
<tr>
<td>40</td>
<td>15</td>
<td>Kuznetsova (1991)</td>
</tr>
<tr>
<td>8</td>
<td>0–20</td>
<td>Kuznetsova (2002)</td>
</tr>
<tr>
<td>66</td>
<td>0–20</td>
<td>Kuznetsova et al. (1998)</td>
</tr>
<tr>
<td>96</td>
<td>5</td>
<td>Glud et al. (2002)</td>
</tr>
<tr>
<td>92</td>
<td>10</td>
<td>Glud et al. (2002)</td>
</tr>
<tr>
<td>72</td>
<td>20</td>
<td>Glud et al. (2002)</td>
</tr>
<tr>
<td>51</td>
<td>30</td>
<td>Glud et al. (2002)</td>
</tr>
</tbody>
</table>

Extrapolating benthic microalgal productivity in time and space

Extrapolations of the limited database on productivity of Arctic benthic microalgae to regional scales requires establishment of a relation to simple parameters that are measured more frequently than benthic primary production. The number of studies quantifying benthic Chl a concentration is much larger than the number of studies quantifying microalgal activity, and Chl a has often been used as an indication of the activity of benthic microalgae (Cahoon 1999, Vetrov and Romankevich 2004). All the original studies of Arctic benthic microalgae (Tables 1–4) provided estimates on biomass expressed as Chl a concentration in the sediment surface layer (mostly 0–1 cm). Minimum or maximum values of Chl a and benthic productivity do correlate (Tables 1–4), but overall there is no clear relationship between these two parameters. A simple linear relationship of the compiled data set expresses an $R^2$ value of only 0.23, and elimination of some obvious outliers does not markedly improve the relationship (data not shown). However, this may not be too surprising as the photic zone typically extends only a few mm into the sediment, and Chl a concentration averaged over the uppermost cm therefore only poorly represents the active phototrophic biomass (Kühl et al. 1997, Kühl 2005). Benthic Chl a concentrations may also be confounded by inactive degradation products originating from pelagic microphytes, senescent or saprothetic living microphytes or spores (Sun et al. 1994). Furthermore, it is well-established that one light-adaptive strategy of benthic microalgae is to regulate their cell-specific Chl a concentration (Blanchard and Montagna 1992). Clearly extract-

The relative importance of various photosynthetic communities has also been assessed using diver-operated pulse amplitude modulation (PAM) fluorometry (Schreiber 2005, Diving-PAM, www.walz.com). Measurements in an ice-covered coastal ecosystem off Hokkaido, Japan concluded that benthic microalgae were responsible for 13–66% of the ecosystem production at water depths ranging from 3–9 m (McMinn et al. 2005). However, such indirect approaches involving interpretation of variable chlorophyll fluorescence measurements are based on many assumptions about the photosynthetic apparatus and the coupling between electron transport and carbon fixation of the different organisms – assumptions that are difficult to test or justify with natural samples (Kühl et al. 2001), and, therefore, such studies can often at best be regarded as indicative.

The observations in Table 5 are in contrast to findings of Kuznetsov (2005), who estimated that the annual pelagic productivity exceeded the benthic microalgal productivity by factors of 1.3–2.8 (average 1.7) when integrated for sediments residing at 0–20 m depth in areas off the Kola Peninsula, Franz Josef Land, eastern Svalbard, West Novaya Zemlya and in the Pechora Sea. These values were, however, presented in abstract format without any explanation on how measurements and extrapolations were performed.

Extrapolations of the limited database on productivity of Arctic benthic microalgae to regional scales requires establishment of a relation to simple parameters that are measured more frequently than benthic primary production. The number of studies quantifying benthic Chl a concentration is much larger than the number of studies quantifying microalgal activity, and Chl a has often been used as an indication of the activity of benthic microalgae (Cahoon 1999, Vetrov and Romankevich 2004). All the original studies of Arctic benthic microalgae (Tables 1–4) provided estimates on biomass expressed as Chl a concentration in the sediment surface layer (mostly 0–1 cm). Minimum or maximum values of Chl a and benthic productivity do correlate (Tables 1–4), but overall there is no clear relationship between these two parameters. A simple linear relationship of the compiled data set expresses an $R^2$ value of only 0.23, and elimination of some obvious outliers does not markedly improve the relationship (data not shown). However, this may not be too surprising as the photic zone typically extends only a few mm into the sediment, and Chl a concentration averaged over the uppermost cm therefore only poorly represents the active phototrophic biomass (Kühl et al. 1997, Kühl 2005). Benthic Chl a concentrations may also be confounded by inactive degradation products originating from pelagic microphytes, senescent or saprothetic living microphytes or spores (Sun et al. 1994). Furthermore, it is well-established that one light-adaptive strategy of benthic microalgae is to regulate their cell-specific Chl a concentration (Blanchard and Montagna 1992). Clearly extract-
able Chl a concentration is often a poor proxy for extrapolating benthic productivity to wider areas.

Given the fact that benthic microalgae are generally well supplied with nutrients, it seems reasonable to assume that community productivity is light-limited and that light availability could serve as a good proxy for benthic primary production.

The minimum light requirement for benthic microalgae is not well defined, but communities of obligate benthic diatoms have been encountered down to almost 200 m water depth where the maximum light availability was <0.2 μmol photons m⁻² s⁻¹ (McGee et al. 2008). However, a visual coverage of benthic microalgae is rarely observed in water depths >40 m (Cahoon 1999), and the minimal light intensities at which polar and subpolar microphytobenthic activity have been recorded range between 0.5 and 2.5 μmol photons m⁻² s⁻¹ (Palmisano et al. 1985, Grant 1986, Karsten et al. 2006).

Assuming PAR-extinction coefficients of coastal waters ranging between 0.12 and 0.16 m⁻¹ for clear and turbid water, respectively (Jerlov 1970), and assuming an average down-welling irradiance during midsummer in the high Arctic of 414 μmol photons m⁻² s⁻¹ (Glud et al. 2002), a diel average of 0.5 μmol photons m⁻² s⁻¹ would reach down to 42 and 52 m in “turbid” and “clear” waters, respectively. The corresponding values for 2.5 μmol photons m⁻² s⁻¹ would be 30 and 46 m. This simple calculation does not account for any changes in spectral composition, but demonstrates that benthic microalgae can be photosynthetically active down to significant water depth in the Arctic region – especially in offshore, clear waters.

Based on remote sensing-derived estimates of light extinction coefficients and PAR distribution in surface waters in the Arctic, Gattuso et al. (2006) proposed that the relative proportion (S) of the coastal seabed receiving light above a given threshold (E, mol photons m⁻² day⁻¹) during the open-water period in summer on average could be estimated as:

\[ S = 16.0 - 13.6 \log (E) + 1.5 \log (E)^{q} + 0.7 \log (E)^{r} \]

The relationship accounts for the estimated relative distribution of case 1 water (light attenuation due to presence of phytoplankton), which cover ~66% of the investigated area, and case 2 water (light attenuation due to phytoplankton, suspended particles and matter; see Morel and Prieur [1977]). The relationship predicts that, on average, 35% (~2.1×10⁶ km²) of the coastal Arctic seabed receives a daily average irradiance >0.5 μmol photons m⁻² s⁻¹ during the open-water period. Correspondingly, 25%, 17%, and 5% of the coastal Arctic seabed would receive average irradiances above 2.5, 10 and 100 μmol photons m⁻² s⁻¹, respectively, during the open-water period. Deriving benthic light availability on larger scales from remote sensing is an innovative approach, but it is obviously associated with a number of shortcomings. Measurements can only be performed when cloud and ice cover allow, the spatial and temporal resolution is limited and the distribution of case 1 and case 2 waters can be only crudely assessed at present. Furthermore, the importance of nepheloid layers and high turbidity water, which will both increase light extinction, is poorly defined (Gattuso et al. 2006). Nevertheless such approaches are thus far the best tools available for a first estimation of benthic microalgal productivity over larger scales in the Arctic.

Using the PE relation in Figure 4 and the equation for light availability derived by Gattuso et al. (2006), we can extrapolate the estimated gross primary production (GPPc) derived by chamber/core incubations to the entire Arctic region. In essence, the same calculation could be made for the PE relation of NPPc, derived from microsensor measurements. However, as these data only represent conditions within well developed microalgal patches, the small scale variations in biomass have to be accounted for during extrapolating – and, as seen in Figure 1, this is a non-trivial task. Such an exercise has, however, been carried out and discussed in a confined area of Young Sound, NE Greenland (Glud et al. 2002).

On average, the daily GPPc extrapolated to the Arctic coastal region amounts to an average of 1.8×10⁷ t C day⁻¹ during the open-water period. The average open-water period for the coastal Arctic (water depth 0–50 m) can be estimated from remote sensed sea-ice concentrations available at the NSIDC (National Snow and Ice Data Center, USA). The average value amounts to 120 days using a grid size of 12.5 km and a 25% threshold of sea-ice concentration (Phil Hwang, unpublished data). However, as grids holding any coastline are masked out by such procedures, areas with land-fast ice are under-represented, and 120 days of open-water period must represent a maximum value. On the basis of selected publications, Cahoon (1999) estimated the average open-water period for the Arctic to be 90 days. Using this value, the annual benthic microalgal primary production amounted to 1.6×10⁹ t C year⁻¹ in the Arctic. A longer open-water period would increase the estimated primary production proportionally. This estimate does not include potential contributions during sea-ice cover, but even though some reports have documented “shade-adapted” benthic photosynthesis below sea-ice, reflectance and absorbance in snow-covered sea-ice is so high that such contributions must be marginal (Palmisano et al. 1985, Glud et al. 2007b). The integrated benthic primary production may seem marginal as compared to the existing estimates on the pelagic productivity of the Arctic oceans ranging from 21 to 42×10⁹ t C year⁻¹ (Subba-Rao and Platt 1984, Pabi et al. 2008). However, most of the benthic primary production is confined to regions with water depths shallower than 30–40 m, regions that account for only ~10–14% of the Arctic oceans, and in those areas, the relative benthic contribution to the total community primary production is correspondingly higher as also reflected in Table 5.

Light availability generally declines exponentially with water depth. Although we used different procedures for quantifying benthic productivity and for extrapolating the data to seasonal time scale, our compiled data (Tables 1–4) show a quasi exponential decline with increasing water depth (Figure 7). The inherent scatter in the data set is partly a result of compiling data obtained with different measuring procedures, but it can also be related to variations in the light extinction coefficients between
the different study areas. The natural variability between study areas apparently overrides any systematic bias related to the fact that different measuring procedures have been applied and that the dataset consist of a mixture of NPP and GPP estimates. However, our simple relationship offers another way to extrapolate the limited database to a regional scale by simply multiplying the productivity-depth relation \((42e^{-0.31t})\) to the bathymetry of the coastal Arctic oceans. Such calculation gives an average benthic microalgal primary production of \(1.2 \times 10^5\) t C day\(^{-1}\) for open-water periods, which under the assumption of an average open-water period of 90 days (Cahoon 1999) translates into an annual benthic microalgal primary production in the Arctic coastal ocean of \(1.1 \times 10^7\) t C year\(^{-1}\). This figure is ~30% lower than our estimate of \(1.6 \times 10^7\) t C year\(^{-1}\) using the light relationship above (Figure 4), but acknowledging the different procedures and the limited database, the estimates are surprisingly consistent.

Concluding remarks

The limited database (Tables 1–4) clearly suggests that benthic microalgae contribute significantly to the coastal ecosystem production in Arctic waters. In fact, our compilation indicates that benthic microalgal productivity is of similar magnitude or even exceeds the pelagic productivity in coastal areas with water depths <30 m. Extrapolation from the current data base using empirical relations between light availability, water depth and benthic microalgal activity estimates a contribution of \(1.1\)–\(1.6 \times 10^7\) t C year\(^{-1}\) to the coastal Arctic ecosystem.

The entire Arctic region is, however, grossly undersampled and further studies on benthic primary production should be encouraged. As in temperate regions, certain coastal areas are especially under-explored – these include rocky and sandy sediments, and no measurements have been performed in shallow off-shore areas. The introduction of the eddy correlation technique to aquatic biology (Berg et al. 2003) provides an opportunity to improve this situation – especially if linked to benthic observatories – by facilitating in situ, non-invasive measurements of large-scale net photosynthesis of benthic algae. Combined with more traditional measuring approaches this could provide an improved insight into benthic microalgal activity and the environmental controls that regulate it. The rapid developments in remote sensing of aquatic light distribution and bathymetric mapping will also facilitate more precise regional extrapolation of specific case studies in the coming years.

The sea-ice cover in the Arctic is rapidly declining (Serreze et al. 2007) and given the tight coupling between sea-ice cover and marine primary production (Rysgaard et al. 1999), this is expected to increase Arctic productivity. Arrigo et al. (2008) estimated that the pelagic productivity of the Arctic oceans has increased by 5–6% annually in recent years as a consequence of the increased light availability. Increased light availability is expected to further increase the competition for nutrients and we speculate that benthic primary production may consequently be stimulated significantly more than the pelagic production in the low-nutrient Arctic coastal region. On the other hand, a predicted increase in precipitation and permafrost thawing will increase the nutrient-enriched, turbid freshwater run-off and may locally counteract the expected increase in coastal light availability. The net outcome is hard to predict and can only be elucidated by giving priority to long-term observatory-based measurements of coastal primary production, which forms the foundation for the present and the future Arctic coastal food web.

Acknowledgements

We sincerely acknowledge Olga Kimmins for help translating the Russian literature that was made available to us. The work was supported by the Danish Natural Science Research Council, National Environmental Research Council (UK – NE/F012691/1 & NE/F020406/1) and the Deutsche Forschungsgemeinschaft for financial support (UK899/12-1/2/3). JW and UK also thank the crew at the AWIPEV-base at Ny Ålesund (Svalbard), and the German dive team (under supervision of Max Schwanzit) for assistance in the field. We thank Dr Phil Hwang for the calculation on periods of open-water in the coastal Arctic and two anonymous reviewers for constructive criticism improving the manuscript.

References


Received 23 February, 2009; accepted 8 June, 2009; online first 30 October, 2009