**Microbial communities of subtidal shallow sandy sediments change with depth and wave disturbance, but nutrient exchanges remain similar**

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**ABSTRACT:** Along 3 replicate transects, sediments were sampled from a subtidal sandbank in Cockburn Sound, Western Australia, at 4 depths: 1.5, 4 and 8 m and at 14 m on the flat at the base of the bank. Pulse amplitude modulated (PAM) fluorescence, fluxes of oxygen and inorganic nutrients, N2 fixation and denitrification were measured and sediments analysed for granulometry, pigments, fatty acids, neutral lipids, organic C and total N. There were 2 functional depth zones: 1.5 ~ <4, and ≥4 m. At 1.5 m, chl a concentration was 42.3 mg m–2 (1.83 SE, n = 12), sediments were net heterotrophic, and there were effluxes of inorganic nutrients in the light and uptake in the dark. The 2 intermediate depths had benthic microalgae (BMA) biomass around 88 mg m–2 chl a, and mean gross primary productivity of 2.23 mmol O2 m–2 h–1. At 14 m, chl a concentration was 75 mg m–2, and sediments were net autotrophic. Sediment–water exchanges of inorganic nutrients were dominated by NH4, with maximum efflux from the sediment (1044 μmol m–2 d–1) at 8 m and maximum uptake (539 μmol m–2 d–1) at 4 m. At 1.5 m depth, there was a marked discontinuity in most parameters as the microbial community metabolism and cycling of nutrients between the sediment and water column were altered in conditions of more frequent wave disturbance. At depths ≥4 m, we observed greater amounts of biomass and more primary productivity, but net exchanges of inorganic nutrients were remarkably consistent at all depths from 1.5 to 14 m.

**KEY WORDS:** Microphytobenthos · Benthic microalgae · Biomarkers · Lipids · Pigments · Biogeochemistry · Resuspension

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**INTRODUCTION**

Sandy sediments occupy large areas of coastal waters (Emery 1968), and can be productive over large areas of continental shelves (Cahoon et al. 1994). The benthic microalgae (BMA) in these sediments contribute to food webs and have important functions in nutrient cycling in coastal ecosystems (MacIntyre et al. 1996). The waters of the west coast of Australia are subject to the warm, oligotrophic Leeuwin Current (Pearce 1991). There is little or no upwelling along this coast and negligible freshwater input in summer, resulting in conditions of low productivity (Pearce et al. 2006). During summer, the water column dissolved inorganic nitrogen (DIN) concentrations are extremely low, <0.2 μ mol l–1 (Thompson & Waite 2003), and primary productivity on the continental shelf is hypothesised to be limited by the resupply of DIN from sediments (K. Wild-Allen & M. Feng unpubl.).

Hydrodynamic energy of the water column can have important effects on the benthic microbial community. For example, resuspension of sediments frequently causes loss of microalgal biomass by removal of cells (Baillie & Welsh 1980, Blanchard et al. 2001). Conversely, shelter from physical disturbance results in increased BMA biomass (Plante et al. 1986). The sediments...
ment grain size is generally proportional to the degree of physical disturbance (de Jong & de Jonge 1995), and an inverse relationship between grain size and biomass of diatoms has been shown (Cahoon et al. 1999, Watermann et al. 1999). Coarse sediments with strong advective porewater flows can trap organic matter (OM) and rapidly degrade it, leading to high rates of turnover (Rasheed et al. 2003). Bacteria are a vital part of the sediment microbial community and are influenced by water movement. In flume experiments, the heterotrophic fraction of the sediment community was found to be more easily removed by water flows than BMA (Shimeta et al. 2002). Similarly, the numbers of bacteria were found to be inversely related to the grain size of the sediments on a tidal flat in Nova Scotia, Canada (Dale 1974).

The uptake of nutrients from the water column is affected by the ratio of the autotrophic to the heterotrophic fraction of the sediment microbial community (Nilsson et al. 1991, Engelsen et al. 2008). There is a tendency to greater uptake with autotrophic dominance, and release with greater heterotrophy. The ratio is a function of irradiance, autotrophic biomass and OM concentration (Rizzo et al. 1996), and shifts in the ratio may occur at different depths in different environments. In the productive waters of a Swedish fjord, a light gradient resulted in a transition from autotrophic to heterotrophic dominance at a depth of only 5 m (Sundbäck et al. 2004). In the clearer waters off North Carolina, USA, sediments became net heterotrophic at 15 m, although BMA biomass extended to a depth of 41 m (Cahoon & Cooke 1992).

Much of the work on the ecology and biogeochemistry of subtidal sediments has been done in cool (Cahoon et al. 1993, Böer et al. 2008), more eutrophic (Lukatelich & McComb 1986) and more sheltered waters (e.g. Blanchard 1990), and the coast of Western Australia provides a useful contrast. Here we examined the effect of a depth gradient on subtidal sandy sediments from near the surface to 14 m depth in a relatively exposed, warm and oligotrophic environment. We hypothesised that variation in light and wave energy would affect BMA biomass in these shallow sandy sediments. In addition, we hypothesised that the ratio of microbial autotrophs to heterotrophs would decrease with depth, with the decreasing autotrophic dominance resulting in a decline in net primary productivity (NPP) and a reduction in the uptake of inorganic nutrients.

**MATERIALS AND METHODS**

**Location.** Cockburn Sound is a sandy basin on the west coast of Western Australia (Fig. 1). The sound is 16 km long by 9 km wide, and the depth of the central flat ranges from 16 to 22.1 m (Australian Hydrographic Service 2001). It is enclosed to the east and south by a coastline of carbonate sand beaches and limestone headlands. The western border is formed by subtidal sandbanks, rocky reefs, Garden Island and the causeway that joins the island to Cape Peron. Transects for this study were on the southern side of the subtidal Parmelia Bank (32° 8′ S, 115° 43′ E), which joins Woodman Point to make the northern border of the sound. Parmelia Bank sediments are comprised of at least 50% calcium carbonate (shell fragments, mainly foraminifera and molluscs), quartz and minor amounts of feldspar (Brachmanis 2002). The area is microtidal (range 0.1 to 0.9 m), and water temperatures range from ~15° to 24°C over an annual cycle (Western Australia Department of Environmental Protection 1996).

**Collection of samples.** Three transects, 50 m apart running north–south down Parmelia Bank (Fig. 1), were sampled at depths of 1.5 (top of bank), 4, 8 and 14 m (sandflat at the base of the bank) by SCUBA divers. Intact sediment cores (50 mm diameter, 3 depth–1 transect–1) were collected for ex situ incubations to measure fluxes of oxygen and inorganic nutrients, fluorescence, N₂ fixation and denitrification. Sample collection was carried out in summer, on 2 and 9 February 2004. Equipment and manpower constraints required 2 visits. All samples and cores were collected on the first occasion except for the cores used to measure denitrification. Weather and sea conditions were similar between the 2 trips.

For flux measurements, sediment cores were carefully collected without visibly disturbing the sediment surface and transported to the laboratory in cooled insulated containers. In the laboratory, cores were kept submerged in recirculated, filtered (2 μm) site water, at an in situ temperature of 24°C and photosynthetically active radiation (PAR) of 500 μmol of photons m⁻² s⁻¹ during the simulated in situ photoperiod of 13.5/10.5 h light/dark. We chose 500 μmol of photons m⁻² s⁻¹, as it was measured at intermediate depths in this region, and it was considered sufficient to saturate photosynthesis based on pulse amplitude modulated (PAM) flu-
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orescence measurements (Forehead 2006). Cores were very gently stirred with individual Teflon-coated magnetic stirrer bars (25 mm long), suspended ~50 mm above the sediment and driven by an external magnet rotating at 60 rpm, a rate used in studies of these processes elsewhere (Dalsgaard et al. 2000, Cook et al. 2004b). This speed was sufficient to circulate the water without causing any visible disturbance of the sediment surface. Cores were allowed to equilibrate in the laboratory for at least 6 h before flux measurements commenced. During the flux measurements, dissolved oxygen (DO) concentrations in capped cores were allowed to deviate by no more than 20% from initial values. Measurements of fluxes across the sediment–water interface in illuminated cores were carried out during in situ daytime and in darkened cores at night, to capture any existing endogenous diurnal patterns. Water samples were immediately filtered through pre-combusted Whatman GF/F filters into 10 ml polypropylene tubes, and frozen at –20°C. Nutrient analysis was carried out within 8 mo of sample collection.

**Site measurements.** PAR was measured with a Li-Cor LI-192SA π (180°) sensor attached to a LI-250A PAR meter (Li-Cor Biosciences). Salinity, DO and temperature were measured with a Yeo-Cal model 611 Intelligent Water Quality Analyser (Yeo-Kal Electronics).

**Grain sizing.** Sediments from each site were dried at 50°C and weighed before wet sieving through an 11-sieve stack (2360 to 45 μm). The sieves containing the sediments were then dried at room temperature, and the sediment fractions were weighed. Median grain size was calculated from the cumulative weight plot using the 11 size classes on the x-axis and % of total on the y-axis. The degree of sorting was calculated using the inclusive graphic quartile deviation index (QDI; Giere et al. 1988):

\[
QDI = \frac{\phi_{84} - \phi_{16}}{4} - \frac{\phi_{95} - \phi_{5}}{6.6}
\]

where \(\phi = -\log_2 \text{ (grain size in μm)}\), \(\phi_x\) is the \(\phi\) value of the \(x\)th percentile of the cumulative weight plot. The results for sediment grain size at different depths were classified into 1 of 7 categories based on their degree of sorting.

**Organic carbon (OC), nitrogen.** Sediment samples were freeze-dried and ground to a powder with a mortar and pestle. For OC analysis, 4 mg samples were weighed into aluminium cups (Elemental Microanalysis); for N analysis, sample weight was 40 mg and tin cups (Elemental Microanalysis) were used. Before analysis for OC, samples were sequentially acidified with sulphurous acid to remove carbonate (Nieuwenhuize et al. 1994) and dried in an oven at 50°C. Sediment samples for nitrogen analysis were weighed into tin cups. Samples were analysed for nitrogen and carbon contents using a Carlo Erba NA1500 CNS analyser interfaced via a Conflo II to a Finnigan Mat Delta S isotope-ratio mass spectrometer operating in the continuous-flow mode. Combustion and oxidation were achieved at 1090°C and reduction at 650°C. Where necessary (because of high carbon contents) the carbon signal was quantitatively diluted with helium.

**Lipids.** Sediment samples were extracted 3 times by a 1-phase dichloromethane–methanol–water mixture (3:6:1 v:v:v) according to a modified version of the method of Bligh & Dyer (1959). An aliquot of the total extract was saponified with 3 ml of 5% KOH in methanol:water (80:20) and heated at 80°C for 2 h. The neutral lipid fraction was extracted into hexane-chloroform (4:1) (this mixture of solvents was used for the remainder of the extractions), then acidified and the fatty acid (FA) fraction extracted. The neutral (minus FAs) fraction was treated with bis(trimethylsilyl) trifluoroacetamide (BSTFA), 100 μl at 60°C for 2 h to con-
vert hydroxylated compounds such as sterols and alcohols to their trimethylsilyl (TMSi) ethers. The FA fraction was treated with MeOH:HCl:CHCl₃ (10:1:1) at 80°C for 2 h, and the resulting FA methyl esters were extracted. Initial gas chromatography (GC) was performed using a Varian CP 3800. Analysing the residual neutral fraction, the GC was equipped with a 50 m x 0.32 mm i.d. cross-linked 5% phenyl-methyl silicone (HP5, Hewlett Packard), fused-silica capillary column. The FA fraction (as methyl esters) was analysed on the same instrument, except that a septum programmable injector (SPI) was used with the capillary column (Hewlett Packard HP1: fused-silica, 50 m x 0.32 mm i.d. cross-linked 1% phenyl-methyl silicone). Sterol and FA fractions were analysed using a flame ionisation detector (FID), with 5β(H)-cholestan-24-ol as the internal standard for sterols and the methyl ester of tricosanoic acid as the internal standard for FA. Peak identifications were based on retention times relative to authentic and laboratory standards and subsequent combined GC and mass spectrometer (GC-MS) analysis. The detection limit for individual sterols and FAs was ~0.2 mg m⁻² of 5 mm deep sediment. Individual sterols and FAs were identified by GC-MS identities performed on a Thermoquest/Finnigan GCQ-Plus bench-top mass spectrometer fitted with a direct capillary inlet and an automated on-column injector. Data were acquired in scan acquisition or selective ion monitoring. The nonpolar column (HP5) and operating conditions were the same as that described above for the GC-FID analyses, except that helium was used as the carrier gas.

Samples of Halophila ovalis were collected from near Cockburn Sound and frozen in liquid N₂ until analysis. Leaves were cut into pieces and macerated with a mortar and pestle in the same dichloromethane–methanol–water mixture that was used for the sediments. We considered 15:0 FA separately from the other bacterial FAs because of its large concentrations and potentially ambiguous origins (mostly bacterial, Sass et al. 2002, but also in diatoms, Dunstan et al. 1993), so it was not included in estimates of bacterial biomass. FAs used to calculate total BMA FAs were: 14:0, 16:4, 16:1ω9, 16:1ω7, 16:0, 18:2ω6, 18:4ω3, 18:1ω9, 20:5ω3, 22:6ω3; diatoms: 14:0, 16:0, 16:1ω7, 20:5ω3. These latter assignations were made based on the ratios in which they presented (particularly the ratio of 16:0 to 16:1ω7) and in conjunction with other biomarkers and microscopy. Quantities of some other BMA FAs were probably also from diatoms, but their proportions were not known. The chlorophyte (green algae) FAs could also be found in some other classes, so assignations to that class were made in conjunction with pigments (chl b, lutein; Jeffery et al. 1997).

Sterols are referred to by their trivial names for ease of reading, i.e. cholesta-5,22E-dien-3β-ol as 22-dehydrocholesterol; cholest-5-en-3β-ol as cholesterol; 24-methylcholesta-5,22E-dien-3β-ol as brassicasterol; 24-methylcholesta-5,24(28)-dien-3β-ol as 24-methylenecholesterol; 24-methylcholesta-5-en-3β-ol as campesterol; 24-ethylcholesta-5,22E-dien-3β-ol as stigmasterol; 24-ethylcholesta-5-en-3β-ol as sitosterol; and 4α,23,24-trimethyl-5α-cholesta-22E-en-3β-ol as dinosterol.

The conversion of stenols to stanols (hydrogenation of the delta 5 bond) is an early step in the process of diagenesis of stenols. Thus we used the stanol:steno ratio as an index of bacterial reprocessing and grazing by metazoans (Gagosian et al. 1980, Jaffe et al. 2001).

**Pigments.** Sediments were extracted twice in 100% acetone at 4°C (18 and 4 h). Following extraction, water was added to give a ratio of 9:1 acetone:water by volume, and filtered (0.2 μm membrane filter; Whatman Anatop) before analysis by a Waters high performance liquid chromatograph (HPLC), comprising a model 600 controller, 717 plus refrigerated auto-sampler, and a 996 photodiode array detector. Pigments were separated as described by Wright et al. (1991), detected at 436 nm, and identified against standard spectra. Concentrations of chl a, chl b, β,β-carotene, and β,ε-carotene in sample chromatograms were determined from standards, and all other pigment concentrations were determined from standards of purified pigments isolated from algal cultures. Information for the assignation of microalgal marker pigments was drawn from Jeffery et al. (1997).

The phaeopigments: chlorophyll ratio has been used to indicate the physiological or grazing state of a BMA community (Bidigare et al. 1986, Mundree et al. 2003). In this study, phaeophytin a and pyropheophytin a were the only degraded pigments that could be distinctly and reliably separated; therefore, the ratio of (phaeophytin a + pyropheophytin a) to chl a was used.

**Nutrient fluxes.** Incubations ranged from 2 to 3 h, depending on the rates of change of DO concentrations (to keep within the limit of a 20% increase or decrease). Inorganic nutrients were analysed on a Lachat
8000QC flow injection system. \(\text{NO}_2^-, \text{NO}_3^-\) and \(\text{PO}_4^{3-}\) were analysed by Lachat standard methods. \(\text{NH}_4^+\) was analysed using o-phthalaldialdehyde (OPA) derivatisation and fluorescence detection as described by Watson et al. (2005), except that the oven temperature was set at 50°C. Reference standards were from Ocean Scientific International (Petersfield, UK).

Fluxes were calculated as:

\[
\text{rate} = \frac{\Delta[n] \times v}{A}
\]

where \(\Delta[n]\) is the change in concentration of inorganic nutrient(s) over the incubation period, \(v\) is the volume of the water column in the core, and \(A\) is the cross-sectional area of the sediment surface. Hourly flux rates were extrapolated to 24 h rates as follows: fluxes in the light were multiplied by the interval over which PAR was at or above optimal irradiance for diatoms (using \(I_{\text{opt}}\) values measured with PAM fluorometry; see below). These intervals were adjusted for depth by multiplying PAR by the depth in m and a locally determined attenuation factor \((K_d)\) of 0.18 m\(^{-1}\) (Thompson & Waite 2003); the resulting periods were 13.4 h at 1.5 m, 13.3 h at 4 m, 12.4 h at 8 m, and 9.8 h at 14 m. PAR through the day was modelled by a sinusoidal function, based on irradiance data from nearby Hope Valley (Mountford & Greico 2002). Flux rates from darkened cores were multiplied by the remainder of the 24 h. A value of \(K_d = 0.108\) (P. Thompson unpublished data) was used to estimate the light climate in deeper waters.

**Oxygen fluxes.** Cores for oxygen flux analyses were measured for dissolved oxygen (DO) with a calibrated WTW Oxi 440 probe (precision ± 5%). The tops of the cores were raised above the surrounding water level and capped with watch glasses, and any bubbles were carefully removed. Measurements of [DO] were made at the beginning and end of incubations, and the rate of oxygen increase or decrease was calculated as:

\[
\text{rate} = \frac{\Delta[O_2] \times v}{A}
\]

Symbols are the same as for nutrient flux calculation. The rate of oxygen evolution during the day was used as a measure of NPP; oxygen uptake in the dark was used for community respiration (CR). Gross primary productivity (GPP) was calculated as the sum of NPP and CR.

To test for linearity, 4 measurements were made through the course of incubations for different sediments, and the fit to a straight line was consistently good \((R^2 \geq 0.98)\). Rates of NPP over 24 h were calculated by the same method used for nutrient fluxes (see above).

**PAM fluorometry.** Fluorescence parameters of BMA communities were measured from intact cores with a Phyto PAM 4 channel fluorometer (Heinz Walz GmbH) fitted with a Phyto-EDF fibre-optic attachment, designed for use on biofilms. The 4-channel fluorometer allows simultaneous measurements from 3 taxa with substantially different pigments. Microalgal cultures from the CSIRO Collection of Living Microalgae were used to calibrate the taxonomic discrimination of the measuring software (Phyto-Win v1.46, Heinz Walz). The cultures used were: brown, *Navicula jeffreyi* (CS-46); green, *Dunaliiella* sp. (CS-353); and blue-green, *Oscillatoria* sp. (CS-52). Given the reference spectra used in the Phyto PAM, ‘browns’ are approximately equivalent to any species with chl c (referred to hereafter as browns), greens would contain chl b (‘greens’) and cyanophytes would contain zeaxanthin (‘blue-greens’). All fluorescence terminology is consistent with Büchel & Wilhelm (1993). Measurement light was supplied at 25 Hz or 12 μs pulses at 4 wavelengths (470, 520, 645, 665 nm) to determine \(F_o\) (the initial minimal fluorescence yield). Saturating irradiance pulses of 0.2 s duration at 2600 μmol m\(^{-2}\) s\(^{-1}\) were applied in conjunction with increased actinic light at 1, 32, 64, 90, 150, 240, 405, 480 and 610 μmol m\(^{-2}\) s\(^{-1}\) to provide estimates of relative electron transport rates (ETR) which were used to construct rapid light curves and fit to the model of Eliers & Peeters (1988) to estimate photosynthetic parameters \(\text{ETR}_{\text{max}}\) (maximum electron transport rate), \(I_{\text{opt}}\) (optimal irradiance = that which saturated ETR), and alpha (\(\alpha = \text{initial slope of irradiance versus ETR curve}\)). The average instrument gain was 20.3 and ranged from 17 to 24. The active fluorescence \((\Delta F)\) is the difference between the fluorescence yield from a repeated saturating pulse (maximum yield, \(F_m\)) and the yield in the near dark \((F_d)\) after Brack & Frank (1998),

\[
\Delta F = (F_m - F_d)
\]

and provides a precise measure of relative biomass in the 3 different taxa.

The effective quantum yield of photochemical energy conversion in photosystem II (PSII) was defined after Genty et al. (1989) as:

\[
\theta_{\text{PSII}} = (F_m - F_i)/F_m
\]

**Denitrification.** Denitrification was measured by the isotope pairing method (Nielsen 1992). A time series of 4 (independent) cores was used for illuminated (as for oxygen fluxes) and dark (wrapped in aluminium foil) incubations for each depth. In the same manner as the nutrient fluxes, cores were stirred by magnets suspended ~50 mm above the sediment rotating at ~60 rpm. Experiments commenced with the addition of stock \(^{15}\text{NO}_3^-\) to a final concentration of ~60 μM in the water column over the sediment. This concentration was chosen after a series of experiments (as described by Rysgaard et al. 1995) had shown that
concentrations of $^{15}$NO$_3^-$ above 40 μM gave constant values of denitrification. Samples were taken for analysis of total NO$_3^-$ before and after the addition of $^{15}$NO$_3^-$ in order to calculate final $^{15}$N enrichment. Cores were then capped and left for 2 h to allow the added $^{15}$NO$_3^-$ to diffuse into the denitrification zone and attain equilibrium. Cores were sacrificed over a time span that allowed the DO to decrease by no more than 20% below saturation. They were sacrificed as follows: 1 ml of 50% ZnCl$_2$ was added to the water overlying the sediment, and the sediment was gently homogenised with the water column using a metal rod; coarser particles were allowed to settle for about 1 min before a sample of about 40 ml was taken using a gas-tight syringe. The sample was then placed in a 12 ml Exetainer (Labco) to which 250 μl of 50% w/v ZnCl$_2$ had been added. A headspace of He was introduced into the Exetainers within 2 wk, and the gas samples were subsequently analysed for the isotopic composition of the N$_2$ gas within several months. The gas was analysed using a Carlo Erba NA1500 CNS analyser interfaced via a Conflo II to a Finnigan Mat Delta S isotope-ratio mass spectrometer, in turn interfaced with a Hewlett Packard 5890 GC. Denitrification rates were calculated according to the isotope pairing equations of Dalsgaard et al. (2000).

**N$_2$ fixation.** N$_2$ fixation was measured by the acetylene reduction assay (Capone 1993). Cores were raised until the tops were clear of the water surface and water was gently removed until a depth of 50 mm remained over the sediment–water interface and sufficient to cover the stirrer bars. The cores were capped, using airtight clear lids, and acetylene was injected via a small septum to give a concentration of >20%. For each incubation, a time series of 3 gas samples was collected via the septa into 5 ml evacuated glass tubes (Vacutainer). Four hour incubations were performed, illuminated (500 μmoles of photons m$^{-2}$ s$^{-1}$) during the day and in the dark overnight, to capture any diurnal patterns of N$_2$ fixation. Two cores were incubated for each depth and transect: 1 in the light, 1 in the dark. The concentration of ethylene in the gas samples was measured within 1 wk using a Hewlett Packard 5890 GC equipped with an Alltech AT Alumina column (30 m, 0.53 mm i.d.) and a FID. A conversion factor of 4:1 was used to convert ethylene reduction to potential N$_2$ fixation (Capone et al. 1992).

**Statistics.** We used analysis of variance (ANOVA) or the general linear model routine followed by Tukey’s Honestly Significant Difference test (post hoc) in SPSS or Dunnett’s test in Sigma Stat (Systat Software). Other statistical procedures included paired t-tests and Pearson’s correlations. Where necessary, data were natural log transformed to stabilise variance.

**RESULTS**

**Site description**

The sand at 1.5 m had ripples ~8 cm high with a spacing of ~15 cm, and was even-coloured pale grey. At the 4 m site, there were ripples ~5 cm high, but none were seen at 8 m. Patches of seagrass, mainly *Halophila ovalis*, were observed at the 4 and 8 m sites, but not at 1.5 or 14 m. Sediments at both of the intermediate depths were unevenly coloured golden-brown; this colouring was commonly marked by broad (about 12–15 cm) trails of pale-grey coloured sand which were made by numerous large sand dollars *Peronella lesueuri*, present at densities of up to 6 ind. m$^{-2}$. Sand dollars burrowed down to 20 mm deep in sediments, and at 4 and 8 m deep sites on all 3 transects but not at any 1.5 or 14 m sites. At 14 m depth, the bottom appeared dark brown, and there was a layer of material on the surface that divers could easily resuspend by gentle water movement, without visibly disturbing the grey sediment beneath. There were a few scattered, partially decomposed blades of seagrass.

Irradiance (PAR) at the sea surface was 2366.5 μmoles of photons m$^{-2}$ s$^{-1}$. Using the January average daytime PAR from nearby Hope Valley, of 1756 μmoles of photons m$^{-2}$ s$^{-1}$ (Mountford & Greico 2002), resulted in a conservative estimate of 165 μmoles of photons m$^{-2}$ s$^{-1}$ PAR in the deepest (22.1 m, Australian Hydrographic Service 2001) part of the sound at midday. Using the value of PAR measured at the surface, 2300 μmoles of photons m$^{-2}$ s$^{-1}$, the 0.1% isobath is at 63 m, with PAR of 2.5 μmoles of photons m$^{-2}$ s$^{-1}$.

Sediment grain size (median ± SE) at 1.5 m (309 ± 12 μm, medium sand) was around double that at 14 m (155 ± 1.44 μm, fine sand), the 8 and 4 m sites were of intermediate grain size and not significantly different from each other (Fig. 2A). Sediments at 14 m were significantly (p < 0.05) less well sorted than those at 4 and 8 m. The greatest degree of sediment sorting was at the intermediate depths (Fig. 2B), though not significantly more than at 1.5 m. Observations and granulometry indicated that the sediments of Parmelia Bank can be divided into 3 zones: a zone of highest wave disturbance at around 1.5 m; the 4 and 8 m sites which were virtually indistinguishable from each other in terms of particle size and sorting; and apparently more stable depositional flat at 14 m.

**Organic carbon and nitrogen**

Concentrations of OC were greatest at 14 m (Fig. 3A), significantly greater than the other depths. Total nitrogen concentrations were also greatest at
14 m (Fig. 3B), and least at 1.5 m; the 2 intermediate depths were intermediate and similar to each other. The C:N ratio was significantly greater at the 1.5 m sites than at the other depths (Fig. 3C). The ratio of OC to chl \(a\) was greater at the 14 and 1.5 m sites than at the intermediate depths (Fig. 3D).

**Pigments**

In this study, fucoxanthin was used as a diatom marker. Fucoxanthin is widely regarded as a marker pigment for diatoms, but it also occurs in prymnesiophytes, dinoflagellates and raphidophytes (Jeffrey et al. 1997). In our results, the absence of chl \(c_3\) and peridinin make the first 2 classes unlikely sources. Raphidophytes could not be ruled out by pigment or lipid data, but the only reference that could be found to their occurrence in sediments was as resting stages (Stahl-Delbanco & Hansson 2002). In the marine environment, chl \(b\) is found in green algae, including prasinophytes, euglenophytes, chlorarachniophytes and chlorophytes. The sediments also contained the chlorophyte pigment lutein, and the correlation between lutein and chl \(b\) was strong and consistent \((p < 0.001)\). The mean ratio of lutein:chl \(b\) was 0.5, similar to the ratio calculated for chlorophytes in water 0 to 25 m deep (Mackey et al. 1998). Hence in this study, chl \(b\) was attributed to chlorophytes.

The 1.5 m sediments had significantly \((p < 0.001)\) lower concentrations of chl \(a\) than the other depths: 42.3 ± 1.83 mg m\(^{-2}\) \((n = 12;\) Fig. 4A). The greatest mean concentration was found at 4 m (89.13 ± 6.05 mg m\(^{-2}\)). The ratio of phaeopigments (phaeophytin \(a\) plus pyropheophytin \(a\)) to chl \(a\) at 1.5 m (0.008) was around an order of magnitude less than at the other depths (Fig. 4E).

The accessory pigments of the sediments were dominated \((95 ± 1\%)\) by chl \(c_1\) & \(c_2\), fucoxanthin, diadinoxanthin, diatoxanthin. The ratio of fucoxanthin:chl \(a\) was only significantly different to other depths at 1.5 m \((p = 0.005;\) Fig. 4B). Cyanobacteria (zeaxanthin) were a minor component of the community at all depths (Fig. 4C). Using a zeaxanthin:chl \(a\) ratio of 0.59 (Gibb et al. 2001), cyanobacteria made contributions to biomass ranging from 3.6 ± 0.1% of the chl \(a\) at 1.5 m to 2.3 ± 0.5% at 14 m. The chlorophyte fraction of the BMA community increased with depth (Fig. 4D). Assuming a chl \(b\):chl \(a\) ratio of 0.57 (Gibb et al. 2001), chlorophytes comprised around 1% of chl \(a\) at 1.5 m, but increased to 9.0 ± 3% at 14 m. The ratio of zeaxanthin:fucoxanthin was 12.9 ± 0.6% at 1.5 m, but only 3.1 ± 0.9% at 4 m, 3.6 ± 1.2% at 8 m and 4.6 ± 0.4% at 14 m.

**Lipids**

Phytol concentrations (Fig. 5A) at 4, 8 and 14 m were similar to each other, but concentrations were reduced by about 50% at 1.5 m. The amount of total neutral lipids (Fig. 5B) was significantly reduced at 1.5 m \((F_{3,8} = 41.5, p < 0.001)\) and was greatest at 14 m \((134 ± 5 \text{ mg m}^{-2})\). The stanol:stenol ratio (Fig. 5C) was also least at 1.5 m \((F_{3,8} = 33.7, p < 0.001)\) and greatest at 14 m \((0.19 ± 0.01)\). The values of the ratio changed with depth in a similar way to the ratio of phaeopigments:chl \(a\)
The ratio of sitosterol to brassicasterol was less at 1.5 m than at 14 m ($F_{3,8} = 32.6, p < 0.001$); the ratio was not significantly different between the other depths (Fig. 5D). At 1.5 m, concentration of 24-methylene cholesterol, a sterol found particularly in centric diatoms, was ~1/10 of that found at the other depths (Fig. 5E), and increased at a reduced rate between 4 and 14 m. The ratio of cholesterol (a sterol often used as a marker of grazers) to total sterols was not significantly different between depths ($F_{3,8} = 2.48, p = 0.135$).

BMA and bacterial FAs (for methods of allocation see Table S1) dominated the fatty acids in the sediments (Fig. 6A,B). Relationships between bacterial and algal markers were strong and significant, a correlation of the total bacterial FAs against algal FAs gave a Pearson’s $R$ of 0.917 ($p < 0.001$).
had an R value of 0.899 (p < 0.001). The ratio of algal:total FAs (linear regression: R = 0.740, p = 0.006) decreased with depth, while the ratio of bacterial:total FAs increased (linear regression: R = 0.893, p < 0.001; Fig. 6B). Though not significantly different from 8 or 14 m, the greatest mean concentrations of algal (185 ± 21 mg m–2) and bacterial (56 ± 3 mg m –2) FAs were found at 4 m. The BMA fatty acids averaged 68 ± 2% of total FAs across all depths.

The \textit{Halophila ovalis} leaves contained the following concentrations of sterols (μg g –1 dry weight of leaf): cholesterol 1.17, campesterol 1.99, stigmasterol 1.89, sitosterol 12.11. The ratio of sitosterol to phytol in the leaves was 0.17, campesterol:phytol 0.03, and stigmasterol:phytol 0.03. In Parmelia Bank sediments, the nearest value of the sitosterol:phytol ratio was 0.16 ± 0.02 (n = 3) at 4 m depth, possibly due to the abundance of \textit{H. ovalis}; the campesterol:phytol ratio was 0.09, and stigmasterol:phytol was 0.10.

**Oxygen fluxes**

Both NPP and CR were least at 1.5 m (Fig. 7A). Oxygen fluxes at the other depths were not significantly different from each other. The maximum value of NPP was 2.4 ± 1.0 mmol O₂ m⁻² h⁻¹ (n = 6) measured at 4 m; the greatest CR was 1.05 ± 0.06 mmol O₂ m⁻² h⁻¹ (n = 6), measured at 14 m. The productivity/respiration (P:R) ratio, calculated as GPP:CR, was 0.19 ± 0.09 at 1.5 m, an order of magnitude less than at the other depths (Fig. 7B). Using an average of the ratio of NPP:ETR from the other depths (117.4 ± 14.0), based upon measured rates of ETR at 1.5 m (see below), NPP at 1.5 m was 0.02 mmol m⁻² h⁻¹, the oxygen uptake in the light was around 2 mmol O₂ m⁻² h⁻¹, or more than double that in the dark (0.716 ± 0.099 mmol O₂ m⁻² h⁻¹). Estimated in situ NPP over a full day was not significantly different from 0 at 1.5 m, rose sharply to 32 ± 13 mmol O₂ m⁻² d⁻¹ (n = 6) or 0.34 ± 0.11 mmol O₂ mg chl a⁻¹ (n = 6) at 4 m and then decreased steadily (though not significantly) to 22 ± 9 mmol O₂ m⁻² d⁻¹ (n = 6) or 0.30 ± 0.02 mmol O₂ mg chl a⁻¹ (n = 4), or ~22 mmol C m⁻² d⁻¹.

**Fluorometry**

As determined by fluorescence, the biomass of all BMA taxa was lowest at 1.5 m (Fig. 8A) and greatest at the intermediate depths. ΔF, a high precision proxy for biomass, was up to an order of magnitude greater for diatoms than for cyanophytes and chlorophytes (Fig. 8A). The relative proportions of ΔF for diatoms and cyanophytes varied with depth, such that the ratio ΔF\textsubscript{D}:ΔF\textsubscript{C} was lower at 1.5 m than at the other depths, indicating the greatest dominance of diatom activity over cyanophytes at the shallowest sites. ΔF for chlorophytes was similar to that of cyanophytes, around 2 orders of magnitude less than for diatoms. Thus the...
community composition determined by fluorescence was primarily browns, ranging from 90% at 1.5 m to 83% at 14 m; contributions by cyanophytes and chlorophytes were around an order of magnitude less. Assuming the browns were largely diatoms, these results are consistent with those determined from other biomarkers.

Photosynthetic performance measured as the maximum electron transport rates (ETR$_{\text{max}}$), the irradiance required to saturate ETR (I$_{\text{opt}}$) and the initial slope of the irradiance versus ETR curve ($\alpha$ = alpha) varied significantly with depth (p = 0.007, <0.001, <0.001, respectively). The irradiance required to saturate electron transport (I$_{\text{opt}}$) was greatest at 1.5 m (Fig. 8B), and least at 14 m. For example, I$_{\text{opt}}$ for browns was 93 μmoles of photons m$^{-2}$ s$^{-1}$ at 1.5 m and 63 μmoles of photons m$^{-2}$ s$^{-1}$ at 14 m. I$_{\text{opt}}$ was measurable for cyanophytes and browns at all depths and showed consistent and significant decreases with depth (Fig. 8B). The I$_{\text{opt}}$ irradiance measured for cyanophytes at 14 m depth was 86.2 μmol of photons m$^{-2}$ s$^{-1}$, which was greater than the optimal intensity for diatoms and chlorophytes at that depth (Fig. 8B). ETR$_{\text{max}}$ and $\alpha$ were least at 14 m and increased at 4 and 8 m before declining again at 1.5 m (Fig. 8C). The effective quantum yield ($\theta_{\text{PSII}}$) also varied significantly with depth (p < 0.001)

nutrient fluxes

Fluxes of inorganic nutrients were, generally, highly variable between replicates at each depth (Fig. 9A–D). As a general result, the flux rates of inorganic nutrients were not significantly influenced by depth or irradiance. When net (24 h) fluxes were averaged over the 4 depths, NH$_4$ was the dominant inorganic nutrient (efflux of 170 ± 326 μmol m$^{-2}$ d$^{-1}$). There was uptake of Si (74 ± 79 μmol m$^{-2}$ d$^{-1}$) and efflux of PO$_4$ (14 ± 27 μmol m$^{-2}$ d$^{-1}$); NO$_x$ efflux was just 2 ± 4 μmol m$^{-2}$ d$^{-1}$. The N:P ratio of the sediment to water column exchange of dissolved inorganic nutrients averaged over 1.5 to 14 m was 13.

The 1.5 m sediments were the exception, as there was efflux of Si, PO$_4$ and NH$_4$ in the light and uptake in the dark, with the differences being significant (p < 0.01). NO$_x$ fluxes at 1.5 m were out of the sediment both in the light and in the dark, but were 2 orders of magnitude smaller than NH$_4$ fluxes; net fluxes over

Fig. 8. Fluorescence parameters of benthic microalgal communities measured in intact cores from 4 depths (n = 18). (A) Active fluorescence ($\Delta F$), the difference between the fluorescence yield from a repeated saturating pulse (maximum yield, $F_m$) and the yield in the near dark ($F_o$) after Brack & Frank (1998) and provides a precise measure of relative biomass in the 3 different taxa. (B) I$_{\text{opt}}$, the irradiance that saturates electron transport rates (ETR). (C) The photosynthetic parameter ETR$_{\text{max}}$ (maximum ETR). Error bars are SE; different letters above bars denote significant difference by Tukey’s HSD, α < 0.05. See Materials and Methods for details of the calculations
24 h were near 0 for Si, PO₄ and NH₄. At 4 m there was uptake of Si (11.3 ± 3.5 μmol m⁻² h⁻¹) and NH₄ (19 ± 15 μmol m⁻² h⁻¹) in the light, but effluxes of PO₄ and NOₓ; there were no significant differences between fluxes in illuminated and dark incubations. At 8 m, there was large variability in fluxes; Si fluxes were near 0, PO₄ and NOₓ were taken up in the dark and released in the light, and NH₄ was released in both the light and dark. At 14 m, fluxes of DIN were near 0, Si was taken up in the light and released in the dark and PO₄ was released in the light and taken up in the dark. Over 24 h, fluxes of inorganic nutrients (Fig. 9E–H) were dominated by NH₄; the greatest fluxes occurred at 4 m (uptake of 539 ± 207 μmol m⁻² d⁻¹) and 8 m (efflux of 1040 ± 750 μmol m⁻² d⁻¹) depths. Atomic ratios of the inorganic nutrient fluxes rarely neared Redfield values (Redfield 1958). The closest approximations were an N:Si ratio (Redfield = 1:1) of 1.6:1 in illuminated sediments from 4 m, N:P (Redfield = 16:1) of 24.3 in illuminated sediments from 8 m and N:Si of 1.9:1 in darkened sediments from 1.5 m.

**Denitrification**

Measurable denitrification only occurred in darkened cores at depths of 8 and 14 m. Rates were around 2 orders of magnitude smaller than those of dissolved inorganic nitrogen (DIN). N₂ effluxes were 0.08 ± 0.04 μmol m⁻² h⁻¹ at 8 m and 0.09 ± 0.03 μmol m⁻² h⁻¹ at 14 m depth. No denitrification was detected in any of the illuminated cores, or in samples from 1.5 or 4 m depths.

**N₂ fixation**

N₂ fixation only occurred at 4 of the 12 sites, rates were less than the other DIN fluxes by around 2 orders of magnitude, but of a similar size to denitrification. In darkened cores, fixation was measured at 1 of the three 14 m sites (transect c: 0.13 ± 0.01 μmol ethylene m⁻² h⁻¹) and at 1 of the 4 m sites (transect a: 0.29 ± 0.01 μmol ethylene m⁻² h⁻¹). In illuminated cores, fixation was measured at two 14 m sites (transect a: 0.11 ± 0.01 μmol ethylene m⁻² h⁻¹ and transect c: 0.29 ± 0.01 μmol ethylene m⁻² h⁻¹). The greatest of these rates (0.29) only equates to 0.07 μmol N₂ m⁻² h⁻¹ (using a ratio of 4 moles of acetylene reduced per mole of N₂ fixed; see Materials and Methods).

**Summary of results**

The sampled depths could be divided into 3 zones by physical parameters and 2 by biological parameters. The 1.5 m sites had wave-washed, coarse sediments with lowest BMA biomass, high OC content and a dominantly heterotrophic metabolism. The middle zone, comprising the 4 and 8 m sites, was heavily grazed and disturbed by sand dollars. The granulometry of the 2 intermediate sites was nearly indistinguishable. The 14 m sites were in a flat, depositional zone, with the smallest grain size and least-sorted sediments. The deepest sites had the highest biomass by concentrations of OC, total N and sterols. The 14 m sediments also had the highest ratio of bacterial to BMA FAs. Fluxes of inorganic nutrients were dominated by NH₄; net 24 h rates were near 0 at 1.5 and 14 m depths, but there was uptake at 4 m and efflux at 8 m.
DISCUSSION

The dominance of the BMA community by diatoms was apparent from both pigment and lipid markers. Diatoms are the dominant taxon of BMA in many parts of the world (Cahoon 1999). Nutrient competition experiments carried out on mixed benthic communities (Sommer 1996) showed that diatoms out-competed cyanophytes and chlorophytes over a range of N:P ratios as long as Si was not limiting. The consistent dominance by diatoms of the BMA community on Parmelia Bank suggests that Si was not limiting at any depth, and the greater uptake of Si at 4 m suggests strong diatom growth at that depth. The correlation between lutein and chl b suggests the presence of chlorophytes (Brots & Plante-Cuny 2003) ranging from 1% of BMA at 1.5 m to 9% at 14 m. Concentrations of the cyanophyte accessory pigment zeaxanthin were low, suggesting the BMA were 3.6% cyanophytes. The fluorescence-based estimate of cyanophyte biomass was of up to 17% of BMA, possibly greater because of differences in the pigment ratios between the species used to calibrate the fluorometer and those in growing in the sediments, or measurement of cyclic electron flow around Photosystem I (Fallkowski & Raven 2007). The low biomass of cyanophytes and absence of N2 fixation implies that concentrations of DIN remained adequate for diatom requirements even throughout summer.

The BMA chl a biomass data from 17 temperate sites over the depth range of 5 to 20 m averaged 84 ± 204 mg m–2 (Cahoon 1999), which is very similar to the 75 to 89 mg m–2 found in these moderately exposed sediments from the southwest coast of Australia over the depth range of 4 to 14 m. The results from 1.5 m depth contrast sharply with this general accord with international observations; chl a was 42 mg m–2, or only 30% of the average 128 mg m–2 reported by Cahoon (1999) at similar depths in 16 temperate zone locations. On the local scale, the BMA biomass in terms of lipids, chl a (measured by fluorescence and HPLC), and total N was 50% less at 1.5 m than at other depths.

A range of mechanisms might explain the pronounced decline in BMA in shallow waters. Of these, we suggest that grazing played a relatively minor role, since the phaeopigment:chl a ratio was lowest at 1.5 m. In addition, the concentration of cholesterol, a sterol marker of eukaryotic heterotrophs and grazing, was not significantly different between 1.5 m and the other depths. It is well known that there is an inverse relationship between grain size and biomass of benthic diatoms (Cahoon et al. 1999, Watermann et al. 1999). The primary factor determining sediment grain size is the degree of physical disturbance (de Jong & de Jonge 1995). The grain size of the sediments at 1.5 m on Parmelia Bank was significantly greater than at the other depths. Therefore, the abrupt reduction in biomass represents a decrease in BMA retention because of the more frequent, wave-driven resuspension of the shallow sediments (Barranguet et al. 1998). The contrast between the disturbed 1.5 m sites and more protected 4 m sites was similar to the negative relationship between wave disturbance and BMA biomass in southwestern Sweden (Sundbäck 1984), the Westerschelde Estuary in The Netherlands (Barranguet et al. 1998) and the Huon Estuary, Australia (Cook et al. 2004a). We suggest that much of the 250% site variation reported by Cahoon (1999) is due to variation in physical dispersion.

Despite the 1.5 m sediments having the smallest biomass, as measured by chl a, phytol and total neutral lipids, the OC concentration of the sediment was similar to that at the 4 and 8 m sites. The biomarker data did not suggest that there had been an accumulation of OM with an elevated C:N ratio (such as seagrass debris). The concentration of the major sterol of Halophila ovalis, sitosterol, and its ratio to the diatom sterol brassicasterol, was the lowest at 1.5 m. The C:N ratio at 1.5 m (16.1 ± 2.6, n = 6) was nearly double that of the other depths, and well above the Redfield value of 6.6 (Redfield 1958). The ratio of BMA to total FA was greatest in 1.5 m sediments. A potential cause of elevated C:N ratios, particularly in shallow sediments, is the presence of carbohydrates secreted by BMA (Decho 1990). The small effluxes of inorganic nutrients from the 1.5 m sediments suggests that porewater concentrations were relatively low, consistent with more wave flushing of these coarser sediments (Janssen et al. 2005). Environmental conditions of high irradiance and low nutrient availability promote the production of carbohydrates by BMA (Staats et al. 2000). Combined with a larger proportion of attached (epipsammic) diatoms that cannot migrate away from excess light (Admiraal 1984, Barranguet et al. 1997), the rates of photosynthesis at 1.5 m were likely to periodically exceed those required for balanced growth. Hence in the shallow sediments at 1.5 m, washed by oligotrophic waters and subject to high levels of PAR, secretions of carbohydrates by BMA are a likely cause of the elevated C:N ratio.

Bacteria comprised the smallest fraction of the sediment community at 1.5 m, presumably in response to the more energetic hydrodynamic conditions. In a flume experiment that gradually increased current speed over sediments, heterotrophic bacteria were resuspended before BMA (Shimeta et al. 2002). In spite of the low bacterial biomass at 1.5 m, the BMA biomass was 42 mg m–2 and ETRMAX was similar to other depths, yet the rate of GPP was negligible, suggesting a very high rate of oxygen consumption in illuminated
sediments. Generally, ETR is proportional to the photosynthetic rate, although it can underestimate C fixation in strong PAR conditions (Carr & Bjork 2003). Only at 1.5 m were the sediments net heterotrophic and the P:R ratio was an order of magnitude less than the other depths. The efflux of inorganic nutrients in the light was consistently and significantly greater than in the dark, suggesting that bacterial rates of OM breakdown exceeded the capacity of BMA for assimilation. The dark uptake of Si by diatoms at 1.5 m (12.7 ± 0.8 μmol m⁻² h⁻¹) suggests that diatoms were responsible for >10% of total benthic respiration, or 84 μmol O₂ m⁻² h⁻¹ (using the Redfield ratio of Si:C of 1:6.6, respiratory quotient of 1). Respiration dominated oxygen exchanges at 1.5 m and signs of bacterial activity were greatest there, yet the ratio used to infer rates of degradation of BMA (phaeopigments:chl a) was lowest by nearly an order of magnitude, and the more general index of diagenesis (stanol:sterol ratio) was reduced by half. The lipid ratio also implied that the degradation of BMA biomass by bacteria was significantly less important at 1.5 m than in the deeper, more sheltered sediments. We therefore suggest that much of the respiration in the 1.5 m sediments was due to consumption of the carbohydrates secreted by BMA. The carbohydrates secreted by BMA are known to stimulate bacterial growth (Murray et al. 1986) and are a substrate for microheterotrophs (Goto et al. 2001). These results suggest that the BMA drove the metabolism of these shallow, wave-disturbed sediments by secreting large amounts of carbohydrates, possibly 2 mmol C m⁻² h⁻¹. This rate was similar to those found in a study of shallow (0.5–2.0 m) subtidal sediments off the coast of Germany (Boer et al. 2008).

The sediments were strongly autotrophic at 14 m, and GPP was not significantly correlated with depth, making it difficult to estimate at which depth the sediments became net heterotrophic (GPP = 0). At this latitude, on the west coast of Australia, light attenuation (k) varies with depth from 0.14 m⁻¹ at 25 m to 0.07 m⁻¹ at 100 m (P. Thompson unpublished) falling to 0.04 m⁻¹ farther offshore (Thompson et al. 2007). Using the value of PAR measured at the surface, 2300 μmoles of photons m⁻² s⁻¹, and a k value of 0.1 m⁻¹, the 0.1% isobath (PAR of 2.3 μmoles of photons m⁻² s⁻¹) is at 63 m, similar to the 60 m estimated by Cahoon (1999). At 63 m, the photic zone extends to more than 50 km off shore along much of the west coast of Australia.

Grain size, sorting, BMA biomass, the exchanges of gases and inorganic nutrients at 14 m were similar to those at 4 and 8 m. The relatively flat area at 14 m was an area of deposition and accumulation, presumably due to the reduced hydrodynamic energy. It had the greatest concentrations of total neutral lipids and concentrations of OC and total N around double that at the other depths. The C:chl a ratio was greater than at any other depth, more than double that at 4 or 8 m. There were elevated concentrations of sitosterol, the major sterol (70% of total sterols) of the seagrass Halophila ovalis, suggesting that tissue from higher up on Parmelia Bank had contributed to the sediment OM. OM from seagrasses is refractory to degradation compared to microalgae, and can form a fraction of sediment OC that turns over slowly, or is buried (Jaffé et al. 2001), while chl a persists for less than 1 d in non-living matter (Sun et al. 1993). Concentrations of settled phytoplankton (24-methylene cholesterol) also increased with depth; a similar result was reported in a Swedish study (Wulff et al. 2005).

The macrofaunal grazer Peronella lesueuri (sand dollar) was only observed at intermediate depths, a habitat preference that may relate to the grain size (Pomory et al. 1995). Based on the concentrations of bacterial FA, cholesterol and ratios of degradation products, however, grazing was similar at 4, 8 and 14 m sites. The apparent absence of sand dollars from these deeper sediments suggests that microheterotrophs or small invertebrates were responsible for the relatively high levels of degradation products and cholesterol (associated with grazing and heterotrophs). This is similar to the Westerschelde Estuary (Barranguet et al. 1998), where relatively sheltered sediments had greater ratios of degradation products than more disturbed sandy sediments.

There were a number of changes in the sediment community with depth, but fewer changes in the resultant metabolism and biogeochemistry. The major shift occurred with a decrease in wave disturbance between 1.5 and 4 m, where high levels of bacterial activity gave way to an autotrophic dominance. During this study in January (mid-summer) 2004, the major inorganic nutrient flux was NH₄. Half of the net (24 h) inorganic nutrient fluxes measured (4 nutrients × 4 depths) were not significantly different from 0, suggesting an efficient recycling of OM between heterotrophs and autotrophs. When averaged over depth and 24 h, the sediments were a net source of DIN and DIP, and a sink for Si. The averaged release of DIN into the water column was 170 μmol m⁻² d⁻¹, a contribution of 0.02 μmol l⁻¹ in a 7 m deep water column. The absence of denitrification in the illuminated cores and the generally small rates were consistent with other studies reporting that photosynthesis inhibits denitrification in autotrophic sediments in oligotrophic waters (Rysgaard et al. 1995, Sundbäck et al. 2004). The small role played by N₂ fixation further implies that N was not a limiting nutrient in these sediments. The magnitudes of the fluxes of denitrification and of N₂ fixation suggest that these 2 fluxes may approximately balance in this environment.
SUMMARY

BMA biomass was dominated by diatoms. In contrast to other studies, BMA biomass did not correlate with irradiance or depth, probably because of the relatively narrow range of depths (1.5 to 14 m) investigated. Physical disturbance by waves in this exposed environment may also contribute to the relatively homogeneous distribution of BMA biomass. The shallow site (1.5 m) had the lowest BMA biomass, presumably due to the highest rate of wave-induced physical disturbance. Sediments from 4 to 14 m were net autotrophic. With the exception of 1.5 m, the flux rates of inorganic nutrients were not markedly influenced by depth or irradiance. Net NH₄ and PO₄ fluxes, albeit relatively small, were from the sediments into the water column at a molar N:P ratio of 13, suggesting efficient recycling of OM across a range of conditions.

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