Effects of reduced UV radiation on a microbenthic community during a microcosm experiment

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ABSTRACT: A microcosm experiment was conducted over a 77 d period (from 6 July to 21 September 1999) to investigate the response of a microbenthic community to the long-term reduction of ambient ultraviolet-B (UVBR, 280 to 320 nm) and ultraviolet-A (UVAR, 320 to 400 nm) radiation. Intact sediment cores were harvested from a muddy intertidal flat in the lower St. Lawrence Estuary (Quebec, Canada) and placed in an outdoor flow-through system under 3 treatments: ambient solar radiation, reduced UVBR and reduced UVR (280 to 400 nm). Both autotrophic (primary production, microalgal biomass, accessory pigments, diatom composition and abundance and UVR absorbing compounds) and heterotrophic (meiofaunal abundance) components were investigated. It was found that neither nematode nor ontogenic stages of harpacticoid copepod abundances were affected by the treatments. The temporal variations of the nauplius, copepodite and copepod populations correspond to the normal ontogenic evolution of harpacticoids. The microphytobenthic community, dominated by small (≤15 μm) pennate diatoms, presented significantly higher carbon fixation under UVR protected treatments on Days 14 and 21, suggesting a simultaneously detrimental but weak effect of UVBR and UVR on photosynthesis. Occurring in very low concentrations in the sediment, UVR absorbing compounds did not seem to be a major UVR-attenuation mechanism within the microbenthic community. The screening of UVBR and UVR also resulted in a stepwise decrease of microalgal biomass on Days 7, 21, 28, 35 and 42, and of diatom abundance on Day 35. The simultaneous accumulation of pheopigments likely reflected an enhanced grazing pressure under both UV-shielded treatments, presumably by microfaunal grazers. The fact that these noticeable changes were observed between the 3 treatments implies that both UVBR and UVR influence the microphytobenthic community. There was a lack of treatment effect during the second half of the experiment when UVR fluxes showed their normal seasonal decrease. Thus, our results indicate that effects of ambient UVR on microbenthos occur only under high solar irradiance. During this period, both ambient UVBR and UVAR exerted a controlling effect on the microalgal standing stock in muddy intertidal flats in the lower St. Lawrence Estuary.

KEY WORDS: Intertidal mudflat · Microphytobenthos · Meiofauna · St. Lawrence Estuary · UV radiation A and B

INTRODUCTION

Solar radiation is a fundamental ecological factor in aquatic and terrestrial ecosystems. The ultraviolet-B component (UVBR, 280 to 320 nm) of the solar spectrum is mainly absorbed by stratospheric ozone. In the past years, a decrease of the ozone layer has been discovered over Antarctica (Farman et al. 1985, Hofmann 1996, Randel & Wu 1999), the Arctic (Müller et al. 1997, Rex et al. 1997, Randel & Wu 1999) and temperate latitudes (Blumthaler & Ambach 1990, Kerr & McElroy...
The resulting enhanced UVBR has long been recognized as a significant and pervasive selective force on aquatic organisms (Vincent & Roy 1993) and therefore on ecosystems (e.g. Mostajir et al. 2000).

The last decade has stressed a great deal of progress in the understanding of the biological effects of UVBR. Less research has been directed toward the effects of ultraviolet-A radiation (UVAR, 320 to 400 nm). However, since this component of the solar spectrum comprises a substantial proportion of the total ultraviolet radiation (UVR, 280 to 400 nm) energy, UVAR could be found to be equally damaging as UVBR. Solar UVAR has been shown to have a wide range of harmful effects on phytoplankton photosynthesis (Hiebling et al. 1992, Arrigo 1994, Moeller 1994) and bacterioplankton activity (Sommaruga et al. 1997).

Biologically effective levels of solar UVR penetrate the water column to significant depths (Kirk 1994). Although shallow-water benthic communities can be particularly sensitive to UVR, their response is not uniform and varies as a result of temporal and spatial disparity of both UVR exposure and organism sensitivity (Hill et al. 1997). UVBR exposure can be responsible for a decrease in periphytic biomass (Vinebrooke & Leavitt 1996) and benthic primary production (Sundbäck et al. 1997, Odmark et al. 1998), and a shift in the species composition of algal communities (Santas et al. 1997, Vinebrooke & Leavitt 1999). UVAR, in addition to UVBR, is also known to disrupt the balance between primary producers and consumers in streams (Bothwell et al. 1994).

Investigations dealing with the UV-response of natural, intact microphytobenthic communities inhabiting intertidal flats are scarce. Yet, benthic microalgae colonizing these ecosystems serve crucial ecological function (MacIntyre et al. 1996, Underwood & Kromkamp 1999). They play a key role in sediment stability and geochemistry and provide a major food source for invertebrate grazers, particularly meiofauna (Miller et al. 1996). The aerial exposure of sediment at low tide permits UVR to penetrate at significant depths in sediments. For example, Wulff et al. (1999) reported a potential light penetration depth (1%) in dry sediment of 1100 to 1200 µm for photosynthetically active radiation, PAR (at 650 nm), 500 to 800 µm for UVA (365 nm) and 400 to 600 µm for UVB (310 nm). In addition, high photosynthetic rates under solar exposure can lead to the production of reactive oxygen species, provoking additional indirect damage (Garcia-Pichel & Bebout 1996). During a 6 wk long microcosm experiment, Nozais et al. (1999) showed that the overall intertidal meiobenthic community of the St. Lawrence Estuary was sensitive to ambient levels of UVBR. However, conclusions about the long-term ecological impacts of UVR cannot be forecast from short-term results (Bothwell et al. 1993). The present study aimed to determine the long-term (77 d) effects of reduced UVBR and reduced UVR (reduced UVBR and UVAR) on the microphyto- and meiozoobenthos inhabiting a muddy intertidal flat. The specific objectives were to: (1) assess the sensitivity of microphytobenthos (primary production and biomass, accessory pigments and diatom composition and abundance) to UVR; (2) examine the photoprotection level (UVR-absorbing compounds) of the community; and (3) investigate the direct effects of UVR on meiofauna abundance. We focussed on nematodes and ontogenic stages of harpacticoid copepods as these 2 taxonomic groups account for more than 95% of the total abundance of meiofauna at the sampling site (Tita 1999). Conducted over nearly 3 mo, the experiment was expected to be long enough to detect effects of UVR reduction on the dynamics of the microphytobenthic community and associated meiofauna.

**MATERIALS AND METHODS**

**Experimental set-up.** The experiment was carried out at the Station aquicole de Pointe-au-Père of the University of Quebec at Rimouski, on the south shore of the lower St. Lawrence Estuary, Canada (48° 31' N, 68° 28' W) from 6 July to 21 September 1999. A microcosm approach was adopted using an outdoor flow-through system consisting of 9 plastic microcosms (44 l) and a 800 l plastic tank containing filtered seawater (using a serial filtering of cotton cartridge filters of 50, 5 and 1 µm). A pump inside the tank (Little Giant Pump Co., Oklahoma City, OK; flow rate of 74 l h⁻¹) distributed the filtered seawater within every microcosm. A semi-diurnal tide traced from the natural tidal cycle allowed the seawater to increase from 1 cm (low tide) to 10 cm (high tide) above the sediment surface. The filtered seawater in the tank was changed every day, just before the slack of the natural low tide. Consequently, the water entering into the microcosms during the flood tide was completely renewed. Water temperature and salinity in the microcosms were recorded during the water renewal. Salinity was determined with a hand held salinity refractometer (Vista Series Instruments A366ATC).

Three light treatments (control, reduced UVBR and reduced UVR) were achieved by covering microcosms with UVR-transparent acrylic sheets (OP-4, Plastic Alto Inc., Quebec City, Canada; 4.76 mm thick), Mylar-D films (Cadillac Plastic Inc., Montreal, Canada; 0.13 mm thick) and UF-4 filters (Plastique Alto Inc., Quebec City, Canada; 4.76 mm thick) respectively. Spectral transmission properties of all these filters are presented in Fig. 1. Mylar-D sheets were changed every...
week in order to avoid modification in transmission properties due to photodegradation. Treatments were carried out in triplicate.

**Light measurements.** Throughout the experiment, an IL 1700 radiometer (International Light, Newburyport, MA) recorded incident downwelling solar irradiance next to the experimental device. UVBR, UVAR and PAR (400 to 700 nm) were measured using SUD240/SPS300/W, SUL033/UV/A/W and SUL033/PAR/W/QNDS1 flat photodetectors, respectively. The unweighted spectral irradiance at the water surface was calculated under the 3 treatments. For this calculation, we used incident irradiance data measured every nanometer from 280 to 400 nm with an OL 754 spectroradiometer (Optronic Laboratories, Orlando, FL; see Whitehead et al. 2000 for more details). To estimate the biological importance of UVR reduction on photosynthesis, irradiance was also weighted by the biological weighting function (BWF) for inhibition of photosynthesis in *Phaeodactylum* of Cullen et al. (1992).

**Study area and sampling methods.** The sampling site was located in the intertidal zone of Anse à l’Orignal in the Parc national du Bic (48° 20’ N, 68° 48’ W), 30 km west of the Station aquicole de Pointe-au-Père. The intertidal flat extends over 2 km² and faces north-east. Hydrodynamic processes depend on the semi-diurnal tide (M₂; Godin 1979) that provides a mesotidal regime. In summer, the maximum water height and the mean water clarity (as determined by transmissivity) in this cove varied from 71 to 311 cm and from 61 to 80%, respectively (Thorin et al. 1998). The colonized layer of the sampling area (i.e. the layer overlying a Godthwait clay deposit, Dionne 1977) consists of a sandy mud with 2% of organic matter (Tita 1999), covered with a 1 to 2 mm thick cohesive microbial mat. On 5 July, 108 intact sediment cores were collected using transparent plexiglas corers (inner diameter 68 mm, height 80 mm) and then randomly placed inside the microcosms (12 cores per microcosm). Using intact sediment cores permits the maintenance of the complex trophic network of the microbial benthic community. A careful daily inspection of sediment cores allowed us to remove macrofauna (e.g. the gastropods *Hydrobia minuta* and polychaetes *Nereis virens*) occurring at the sediment-water interface. On each sampling date, that is every week from 6 July, a core was randomly removed from each microcosm and sediment samples were taken with circular syringes of 9 mm inner diameter. Samples for primary productivity, microalgal biomass and meiofauna were collected every week whereas those for particulate organic matter, diatom composition and abundance, accessory pigments and UVR-absorbing compounds were taken 3 times (Days 0, 35 and 77). The top 2 to 3 mm of the syringes were retained for laboratory analysis, except that meiofauna abundance was quantified from the upper 2 cm since 90% of meiofauna are concentrated within this depth (Tita 1999).

**Inorganic nutrients and total organic carbon.** Concentration of ammonium (NH₄⁺), nitrate and nitrite (NO₃⁻ + NO₂⁻), silicic acid (Si(OH)₄) and phosphate (PO₄³⁻) were measured every week from samples collected in the overlying water of each microcosm. Samples were immediately filtered on precombusted Whatman GF/F glass-fiber filters before being deep-frozen (~80°C). NH₄⁺ concentration was determined according to the method of Solórzano (1969) as described by Parsons et al. (1984). The other inorganic nutrients were measured using a Technicon Autoanalyzer II (Strickland & Parsons 1972). For total organic carbon (TOC) measurements, duplicate samples were collected from the common water exit of the microcosms in 4 ml glass storage vials with Teflon-lined caps previously cleaned following the protocol of Burdige & Homstead (1994). The samples were acidified to ~pH 2 with 50% v/v H₃PO₄ (5 µl ml⁻¹). TOC was determined on a TOC-5000 analyzer (Shimadzu, Kyoto, Japan) using the analysis procedure given in Whitehead et al. (2000).

**Particulate organic matter.** One sample was taken from each core and immediately stored at −20°C for later determination of particulate organic carbon (POC) and nitrogen (PON). Samples were dried at 60°C for 24 h and pelletized before quantification with a Model 2400 CHN analyzer (Perkin-Elmer, Wellesley, MA).

**Microphytobenthos.** For the determination of accessory pigments and UVR-absorbing compounds, 2 separate samples were immediately stored at −20°C after sampling. All subsequent steps were done at dim light and room temperature. Extraction of pigments and UVR-absorbing compounds was performed in 98% methanol/2% ammonium acetate. The extract was sonicated (40 kHz, 5 min) and centrifuged (3200 rpm, 20 min).

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**Fig. 1.** Transmission spectra of OP-4, Mylar-D and UF-4 filters fixed over microcosms.
The extraction procedure was repeated once more. Both supernatants were pooled before being filtered on 0.2 µm Acrodisc® filters. A test showed that this technique extracts about 99% of the sediment pigment and UVR-absorbing compounds. Pigments were analyzed by reversed-phase high-performance liquid chromatography (HPLC) according to Wright et al. (1991) as modified by Roy et al. (1996), while the presence of UVR-absorbing compounds was determined using a Lambda 12 spectrophotometer (Perkin Elmer).

Immediately after sampling, 2 samples were pooled in 2% formaldehyde and stored at 4°C for the later enumeration and identification of the microalgae. The samples were diluted, and diatoms (empty and chloroplast-containing cells) were enumerated using the inverted microscope method (Lund et al. 1958). Diatoms were grouped into size and shape classes, and then identified when possible to genus level.

Microalgal biomass was based on chlorophyll a (chl a) concentration. After sampling, 2 syringes from each core were immediately frozen at −20°C. All subsequent manipulations were done at room temperature and dim light. The day after sampling, the frozen sediment was dried at 60°C for 24 h. Chl a extraction was performed in 10 ml 90% acetone and kept at 4°C in the dark for 18 h (Riaux-Gobin & Klein 1993). Fluorescence was measured before and after acidification with 5% HCl on a R010 Turner Designs Fluorometer, for the determination of chl a and pheopigment concentrations.

Benthic microalgal productivity was measured by the ¹⁴C-assimilation method. Two samples from each core were pooled in 50 ml of filtered seawater (Whatman GF/F). The sample was mixed and 0.5 ml of NaH¹⁴CO₃ (10 µCi ml⁻¹) was added. Three 15 ml subsamples were distributed in 3 borosilicate scintillation vials (1 dark and 2 light bottles) and then incubated in a downward position for 2 h (12:00 to 14:00 h) in a microcosm covered with a UVR-transparent acrylic sheet (OP-4). The dark bottle contained 50 µl 10⁻⁵ mol l⁻¹ of EcoLume (ICN Biochemicals®) scintillation cocktail (OP-4). The daily primary production rate was obtained by dividing the total particulate carbon uptake by the fraction of daily PAR received during the incubation.

**Meiozoobenthos.** Three samples from each core were immediately preserved in 4% formaldehyde, with subsequent addition of Rose Bengal. After a few months, each sample was passed firstly through a 500 µm sieve, to exclude macrofauna, and then through a 63 µm sieve to isolate the meiofauna. Meiofaunal organisms were sorted from sediment by centrifugation using Ludox-TM (Heip et al. 1985) and counted under a dissecting microscope.

**Statistical analyses.** Before undertaking the parametric statistical tests, the different variables were tested for normality of distribution (Kolmogorov-Smirnov test) and homogeneity of variance (Hartley test). When needed, data were In-transformed (Underwood 1981). To reveal the effects of treatment, a nested ANOVA was run separately for each sampling week, using 2 or 3 replicate cores nested in the factor treatment (Underwood 1981). For primary productivity and diatom abundance, treatment effects were tested by a one-way ANOVA. A multiple comparison test of means (Tukey test) was used when significant differences (p ≤ 0.05) were found between treatments. To infer relationships between variables, Spearman’s rank correlations (rₛ) were also calculated.

**RESULTS**

**Light, temperature, salinity and nutrients**

UVR intensities under the various filters were measured during a sunny day around noon (Fig. 2A). Mylar-D and UF-4 filters reduced UVBR irradiance by 80.7 and 99.9%, respectively, when compared to control (OP-4) whereas the relative reduction of UVAR under these 2 filters reached 18.5 and 86.6%, respectively. The biological weighting function of Cullen et al. (1992) shows the biological significance of the wavelengths between 310 and 340 nm (Fig. 2B). Compared to the control, Mylar-D and UF-4 filters removed the biologically weighted UVBR by 86.7 and 99.9% and that of UVAR by 24.1 and 99.3%, respectively.

The UVBR flat collector stopped transferring the incident UVBR data to the computer at the beginning of the experiment. To estimate the daily incident UVBR irradiance, daily incident UVAR was multiplied by the daily incident UVBR:UVAR ratio of 0.036 measured by Nozais et al. (1999) for the same site between 8 July and 19 August 1997. Since the Environment Canada monitoring station located in Montreal, Quebec (World Ozone and UV Radiation Data Centre, Environment Canada, Downsview, Ontario, Canada) provided evi-
dence of a similar state of ozone layer over Montreal in the summers of 1997 and 1999, one can assume that the UVBR fluxes reaching the experimental enclosures were similar for these 2 summers. The weekly UVBR irradiance given here represents only an estimation of the weekly UVBR flux during the experiment.

During this 11 wk long experiment, high variations in downwelling irradiance occurred. Maximal incident irradiance over the experimental enclosures at noon on sunny days ranged between 1700 and 1900 µE m⁻² s⁻¹ for PAR and 32 to 36 W m⁻² for UVAR. During the sunny days, daily incident PAR reached 55 E m⁻², while incident UVAR and estimated UVBR attained 1121 and 44 kJ m⁻² d⁻¹, respectively. During overcast days, incident irradiances drastically declined to 9 E m⁻² d⁻¹ for PAR, 146 kJ m⁻² d⁻¹ for UVAR and 6 kJ m⁻² d⁻¹ for UVBR. Weekly irradiances of PAR, UVAR and UVBR under the various filters decreased gradually during the experiment as a result of increasing cloud cover (especially during the last 3 wk) and decreasing day length as per seasonal trends (Fig. 3). The PAR:UVAR ratio remained relatively constant throughout the experiment.

During the whole experiment, the average water temperature, and therefore that of the sediment, was around 20°C, varying from 32°C at noon in July to 9°C in early morning in September. The average water temperature during the period of incubation for primary production measurements ranged between 13 and 20°C. Salinity varied from 18 to 30, with a mean value of 25.

The concentration of inorganic nutrients in the overlying water of the microcosms varied greatly throughout the experiment (Table 1). Starting with high initial values (NO₂⁻ + NO₃⁻ = 10.9 µM, NH₄⁺ = 2.1 µM, Si[OH]₄ = 14.5 µM and PO₄³⁻ = 0.90 µM), nutrient concentration decreased drastically to reach low values that correspond to a generally nutrient-depleted sea surface in summer. Molar ratios of dissolved inorganic nitrogen (DIN) to PO₄³⁻ and DIN to Si[OH]₄ also ex-

<table>
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<th>Day</th>
<th>NO₃⁻ + NO₂⁻ (µM)</th>
<th>NH₄⁺ (µM)</th>
<th>Si(OH)₄ (µM)</th>
<th>PO₄³⁻ (µM)</th>
<th>DIN:PO₄³⁻ (mol:mol)</th>
<th>DIN:Si(OH)₄ (mol:mol)</th>
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hibited great temporal variability, with values ranging from 0.9 to 14.6 and from 0.02 to 0.95, respectively (Table 1). There was no significant relationship between inorganic nutrients and the other measured biological variables.

TOC and particulate organic matter

The concentration of TOC in the common water of all the microcosms ranged between 103 and 187 µM, with an average value of 161 µM. POC concentrations in the top 2 to 3 mm of the sediment remained rather constant throughout the experiment, ranging from 2682 to 2726 mgC m⁻². The POC:PON molar ratio stayed around 11.5. No significant treatment effect on the concentrations of POC and nitrogen was found.

Microphytobenthos

HPLC pigment analysis revealed, in addition to chl a and degradation pigments, chlorophyll c₁+₂ (chl c₁+₂) and the carotenoids fucoxanthin, diadinoxanthin, diatoxanthin, β,β-carotene and 19'-hexanoyloxyfucoxanthin. In each treatment, ratios of fucoxanthin to chl a and that of chl c₁+₂ to chl a increased significantly over time, particularly during the second part of the experiment (Table 2). The different light treatments did not affect any accessory pigment to chl a ratios during the whole experiment, except that on Day 77 when reduced UVR resulted in an enhanced fucoxanthin:chl a ratio compared to the UVR shielded treatment.

Microscopical examination revealed that diatoms were the main algal group in the sediment. Flagellates and cysts of Chrysophyceae were also observed. The diatom community was dominated by small (≤15 µm) unidentified pennate diatoms (Fig. 4). The main identified diatoms belonged to the genera Cocconeis (≤25 µm), Navicula (20 to 55 µm) and Amphora (12 to 34 µm).
From approximately $21 \times 10^9$ cells m$^{-2}$, the total diatom abundance exhibited a rapid increase in all treatments to reach values ranging between 41 and $59 \times 10^9$ cells m$^{-2}$ on Day 77. During the experiment, there was a significant ($p \leq 0.05$) treatment effect on diatom abundance. Both shielded treatments resulted in a significant decline (by 42\%) in the total abundance of diatoms on Day 35 due to the lower number of small unidentified pennates. On Day 77, the total diatom abundance was 43\% higher with UVR screened off than under control.

Over the whole experiment, microalgal biomass (Fig. 5A) and the percentage of pheopigments (ratio of pheopigments to total pigments, i.e. chl $a$ + pheopigments; Fig. 5B) in the control averaged 33 mg chl $a$ m$^{-2}$ and 45\%, respectively. The microalgal biomass was inversely correlated with the percentage of pheopigments ($r_s = -0.43$, $p \leq 0.001$). No other correlation was found with the other biological or physical variables. Result of nested ANOVA revealed significant changes in both the microalgal biomass and the percentage of pheopigments, mainly during the first part of the experiment (Table 3). UVBR screening significantly altered microalgal biomass on Days 21, 35 and 42 (Fig. 5A), but to a lesser extent than did UVR reduction (Table 3). Conversely, the percentage of pheopigments in UVBR-shielded cores increased on Days 14 and 21 as much as under reduced UVR. Microalgal biomass was significantly less (22 to 35\%) under UVR reduction than under control on Days 7, 21, 28, 35 and 42 (Fig. 5A, Table 3). In contrast, screening of UVR caused a 19 to 28\% enhancement in the percentage of pheopigments on Days 14, 21 and 70. Microalgal biomass was significantly lower under reduced UVR treatment compared to the reduced UVBR treatment on Day 28 (Fig. 5A, Table 3). In contrast, the percentage of pheopigments was significantly higher (16 to 33\%) in the reduced UVBR treatment on Days 28, 35 and 42.

During the experiment, the benthic particulate primary production under control varied from 16 to 77 mg C m$^{-2}$ h$^{-1}$ (Fig. 6A) corresponding to 80 to 243 mg C m$^{-2}$ d$^{-1}$. Primary production was directly correlated to both the water temperature ($r_s = 0.88$, $p \leq 0.001$) and incident PAR prevailing during the incubation period ($r_s = 0.77$, $p \leq 0.001$). A multiple regression test showed that these 2 environmental factors accounted for 79\% of the variance in primary production. Throughout the experiment, less than 3\% of the total (particulate + dissolved) primary production was released to seawater as DOC. The particulate produc-

Table 3. Percent variation of microalgal biomass (chl $a$) and percentage of pheopigments (ratio of pheopigments to total pigments, i.e. chl $a$ + pheopigments) between the 3 treatments. Only significant values at $p \leq 0.05$ are presented.

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<td>Pheopigments (%)</td>
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Fig. 4. Composition and abundance of the diatom community of the microbial mat (A) exposed to ambient solar radiation, (B) shielded from UVBR, and (C) shielded from UVR (reduced UVR). Mean (+SE) of n replicate cores (n = 9 for Day 0, n = 3 for the other days) are shown. *Significant difference in cell abundance from the control (ambient solar radiation) ($p \leq 0.05$)
The abundance of the other taxa (turbellarians and foraminiferans) was negligible. Temporal variations in the abundance of nematodes and ontogenetic stages of harpacticoid copepods in the different light treatments are presented in Figs 7 & 8. Nematode abundance increased gradually from $840 \times 10^3 \text{ m}^{-2}$ on Day 0, to around $1600 \times 10^3 \text{ m}^{-2}$ under control on Day 77 (Fig. 7). After a gradual increase up to Day 28, reaching $265 \times 10^3 \text{ m}^{-2}$ under control, the harpacticoid nauplius abundance declined on Day 35, then peaked on Day 49 before finally decreasing for the remainder of the experiment (Fig. 8A). Copepodite abundance increased from $33 \times 10^3 \text{ m}^{-2}$ on Day 0 to $167 \times 10^3 \text{ m}^{-2}$ under control on Day 49 before gradually decreasing for the remainder of the experiment (Fig. 8B). Adult harpacticoid copepod abundance showed a pattern similar to that of copepodites, however the maximum abundance under control occurred on Day 63 (Fig. 8C).

There was some occasional effect of the light regime on meiofauna abundance (Figs 7 & 8); however, no consistent pattern was observed. No correlation was found between meiofauna abundance and the other determined variables.

**Meiozoobenthos**

At the beginning of the experiment, nematodes were the most numerous group of meiofauna (74%) followed by nauplii (12%), copepods (6%), copepodites (4%), oligochaetes (3%) and polychaete larvae (1%).

The abundance of the other taxa (turbellarians and foraminiferans) was negligible. Temporal variations in the abundance of nematodes and ontogenetic stages of harpacticoid copepods in the different light treatments are presented in Figs 7 & 8. Nematode abundance increased gradually from $840 \times 10^3 \text{ m}^{-2}$ on Day 0, to around $1600 \times 10^3 \text{ m}^{-2}$ under control on Day 77 (Fig. 7). After a gradual increase up to Day 28, reaching $265 \times 10^3 \text{ m}^{-2}$ under control, the harpacticoid nauplius abundance declined on Day 35, then peaked on Day 49 before finally decreasing for the remainder of the experiment (Fig. 8A). Copepodite abundance increased from $33 \times 10^3 \text{ m}^{-2}$ on Day 0 to $167 \times 10^3 \text{ m}^{-2}$ under control on Day 49 before gradually decreasing for the remainder of the experiment (Fig. 8B). Adult harpacticoid copepod abundance showed a pattern similar to that of copepodites, however the maximum abundance under control occurred on Day 63 (Fig. 8C).

There was some occasional effect of the light regime on meiofauna abundance (Figs 7 & 8); however, no consistent pattern was observed. No correlation was found between meiofauna abundance and the other determined variables.
Throughout the experiment, the absorbance of methanol extracts presented similar patterns, with 2 absorbance peaks around 270 and 340 nm (Fig. 9). DNA likely caused the first peak since DNA absorbs wavelengths between 230 and 310 nm (Buma et al. 1996). The second one is attributed to mycosporine-like amino acids (Dunlap & Shick 1998). Although no statistical analysis was attempted, the absorbance peaks showed no observable difference between treatments and over time.

**DISCUSSION**

**The microbenthic community**

There is a long history of investigations into sediment-associated microalgae colonizing intertidal flats in North America and in Europe (MacIntyre et al. 1996, Cahoon 1999 and references therein). In the St. Lawrence Estuary and Gulf (Canada), only the epilithic component of the microphytobenthos has been studied through an exhaustive taxonomic list (e.g. Cardinal et al. 1984, 1986, Poulin et al. 1984, 1990, Bérard-Therriault et al. 1986, 1987) and through the influence of environmental factors on the community structure (Fludon & Bourget 1983) and on the photosynthetic properties (Lamontagne et al. 1986, 1989) of epilithic diatoms. In contrast, sediment-associated microalgae colonizing soft sediment in this region have never been characterized. This study assesses for the first time the summer dynamics of the microphytobenthic community of a muddy intertidal flat in the St. Lawrence Estuary.
Although microphytobenthos in intertidal flats present successional patterns throughout the year (Pinckney et al. 1995, Barranquet et al. 1997), these areas are usually dominated by diatoms (Leach 1970, Cammen & Walker 1986, Gould & Gallagher 1990, Sundbäck et al. 1996c, Barranquet et al. 1997). In the present work, HPLC-based accessory pigment signature clearly established the dominance of diatoms within the algal community (Table 2). The carotenoid fucoxanthin was the predominant accessory pigment found along with chl c1, c2, diadinoxanthin, diatoxanthin and β,β-carotene. These pigments are all found in Bacillariophyceae (Stauber & Jeffrey 1988). Microscopical examination confirmed the occurrence of benthic diatoms in significant numbers, particularly small (≤15 µm) pennates (Fig. 4). The main genera identified were Amphora, Navicula and Cocconeis. Flagellates found in small number were likely Prymnesiophyceae since 19’-hexanoyloxyfucoxanthin and the above pigments commonly occur in these taxa (Gieskes & Kraay 1986, Wright & Jeffrey 1987).

Benthic microalgal primary production and biomass values can be compared with those of other intertidal flats on the East coast of North America and in Europe (MacIntyre et al. 1996, Cahoon 1999 and references therein). Benthic primary production exhibited large temporal variations (from 16 to 77 mg C m⁻² h⁻¹, corresponding to 80 to 243 mg C m⁻² d⁻¹) as a result of changes in cloud cover and water temperature during the incubation stage (Fig. 6A). These rates are consistent with those published (from 8 to 175 mg C m⁻² h⁻¹) for the East coast of the United States and in Northern Europe (Cahoon 1999). A the end of the experiment, the microalgal production and biomass at the sampling site were 44 mg C m⁻² h⁻¹ and 17 mg chl a m⁻² respectively, similar to those measured in the microcosms. Under natural light conditions, microalgal biomass in the top 2 to 3 mm of the sediment varied from 24 to 45 mg chl a m⁻² (Fig. 5A). These values are within the concentration range (15 to 220 mg chl a m⁻²) previously reported in North America (Cahoon 1999).

In coastal environments, the growth of shallow-water microphytobenthos is generally not limited by inorganic nutrients because of high rates of remineralization within the sediment (Admiraal et al. 1982, Sundbäck et al. 1996a, but see also Nilsson et al. 1991). In the present study, the nutrient concentration of the overlying water of the microcosms probably did not limit microphytobenthic growth. The high initial values were probably due to the release of inorganic nutrients by the intact sediment cores when placed into the microcosms. Nutrient concentrations rapidly reached values corresponding to a nutrient-depleted sea surface in the lower St. Lawrence Estuary during summer (Levasseur & Therriault 1987). Ratios of DIN to Si(OH)₄ and DIN to PO₄³⁻ were always lower than the critical values of Redfield et al. (1963) of 1 and 16 respectively, suggesting that DIN was the element in shortest supply relative to nutrient requirements of the microalgae (Table 1). However, since the P:B ratio was not correlated with DIN concentration, nutrient limitation probably did not occur in the microcosms. This idea is also supported by the fact that the POC:PON molar ratio in the top 2 to 3 mm of sediment remained constant (around 11.5) throughout the experiment.

Significant changes over time, irrespective of treatment, were found in the structure of the copepod community. The similar and delayed temporal variations of nauplii, copepodite and copepod abundance corresponded to the normal ontogenic evolution of harpacticoids (Fig. 8). Nauplius abundance increased from Day 0 to 28 as a result of the production of eggs by mature adult copepods and the lack of predators (i.e. polychaetes) which were manually removed. In the Parc du Bic, predation on harpacticoids by Nereis virens is substantial (Tita et al. 2000). Shrimps (Gregg & Fleeger 1998) and juvenile fishes (Morris & Coull 1992) can also be voracious predators of nauplii. The sudden decrease in nauplius abundance on Day 35 was probably related to recruitment into the copepodite stage, leading to enhanced copepodite and copepod abundance in the middle and end of the experiment, respectively. The nauplius abundance peak on Day 49, 21 d after the first peak on Day 28, probably reflected the occurrence of a second nauplius cohort (Fig. 8A). Palmer & Coull (1980) found, from a laboratory study, that the development time for the nauplius stage of Microarthridion littorale lasts from 30 d at 15°C to 20 d at 25°C. Yet, a second peak in copepodite abundance was not observed, which suggests that an increase in the nauplius mortality, as a result of food limitation in the microcosms, prevented the appearance of another abundance peak in copepodite population. As consumers, harpacticoid copepods feed on both planktonic and benthic diatoms, autotrophic flagellates, bacteria and detritus (Hicks & Coull 1983). It is known that they can regulate their behavior to maximize the intake of food (Montagna et al. 1995). Evidence of feeding differences among species (Pace & Carman 1996) and ontogenic stages (Decho & Fleeger 1988) suggests that meiobenthic copepods require a rather specialized food source. Therefore, a qualitative impoverishment of food for harpacticoid notably nauplii, was certainly possible since there was no external food contribution in the microcosms.

**Treatment effects**

Results of investigations into the effects of UVR on benthic fauna inhabiting shallow waters vary consider-
ably because of temporal and spatial disparity in both UVR exposure and organism sensitivity (Hill et al. 1997). Many studies have demonstrated that grazers are unaffected by UVR exposure (DeNicola & Hoagland 1996, Sundbäck et al. 1996b, 1997, Hill et al. 1997, Vinebrooke & Leavitt 1999). Our study agrees with these earlier experiments since we did not find any significant pattern of treatment effect on meiofauna, neither on nematode nor on ontogenic stages of harpacticoid copepod abundances. The burrowing activity of both nematodes (Heip et al. 1985) and harpacticoids (Hicks & Coull 1983) within the sediment likely contributed to reducing the UVR exposure. Conversely, Nozais et al. (1999) reported that the larval stages (nauplii and copepodites) of harpacticoid copepods could be susceptible to ambient levels of UVBR in the St. Lawrence Estuary. The same pattern did not occur in the present study. This could be attributed to the fact that sampling the top 2 cm of the sediment might have masked the effects of UVR on nauplii and copepodites living at the water-sediment interface. Another possible explanation is related to the UVBR levels. Due to technical problems, we did not record UVBR values and were forced to use the UVBR to UVAR ratio calculated by Nozais et al. (1999) to estimate UVBR. These levels may have been lower during the present experiment, perhaps resulting in the absence of UVR effects on larval stages of harpacticoid copepods.

A number of studies have demonstrated that ambient ultraviolet radiation inhibits photosynthesis of both planktonic (Smith et al. 1992, Mostajir et al. 1999b) and benthic (Sundbäck et al. 1997, Odmark et al. 1998) microalgae. The present experiment corroborates these earlier reports since UVR reduction resulted in an enhanced P:B ratio of microphytobenthos on 2 consecutive sampling days in July (Fig. 6B). This suggests a detrimental but weak effect of ambient UVR on photosynthesis. The fact that both UVBR and UVAR were simultaneously responsible for the declining chl a concentration under ambient solar radiation compared to UVBR reduction. Bothwell et al. (1994) also found that algal biomass accumulated in streams exposed to ambient solar radiation compared to reduced UVR. They attributed this periphyton accrual to the elimination of macrograzers. The fact that the 2 shielded treatments provoked a differential treatment response in chl a accrual and in the percentage of pheopigments suggests an enhanced grazing pressure in shielded sediment cores. The abundance of meiofauna was not disrupted by treatments. As discussed above, the sampling depth for studying meiofauna may have masked UVR effects on nauplius and copepodite abundance and therefore grazing rates. Besides, it is known that benthic microfauna (e.g. heterotrophic flagellates and ciliates) feed on microphytobenthos. For example, up to 10% of the total daily benthic primary production can be grazed by ciliates (Epstein et al. 1992). In the lower St. Lawrence Estuary, abundance and grazing rates of planktonic ciliates have been shown to be enhanced under reduced UVR conditions (Mostajir et al. 1999a). Along the same line, we hypothesize that sediment-associated microfauna were also affected by UVR during our study. Future research on the effect of UVR on benthic systems should include measurements of grazing rates of both microfauna and meiofauna.

Significant changes in the abundance of diatoms were found throughout the experiment. The decrease in the total diatom abundance in shielded cores on Day 35 supports the idea of an enhanced grazing pressure under these treatments (Fig. 4). However, these results contrast with those observed on Day 77. The enhanced diatom abundance on Day 77 in protected cores may reflect the enhanced P:B ratio at the end of the experiment. Although we did not find such effects statistically, the last 4 wk of the study were characterized by tendencies of increased microalgal P:B ratio under reduced UVR treatment (Fig. 6B). The fact that only the small unidentified pennate diatoms were affected by UVR reduction on Day 35 suggests, as formerly demonstrated by Santas et al. (1997), different UVR susceptibility within the diatom community.

**Attenuation of UVR damage**

The different strategies used by organisms for the minimization of UVR-induced damage are described in Roy (2000). Benthic microalgae can avoid UVR fluxes by migrating downward in the sediment since vertical migration offers substantial protection for cyanobacteria against UVR exposure (Nadeau et al. 1999). Although this behavioral process probably applies for diatoms (Sundbäck et al. 1996b), its long-term efficiency is doubtful (Underwood et al. 1999). Attenuation of UVR-induced impairments can also be achieved through the synthesis of various cellular com-
ponents absorbing in the UVR band. Our methanol extracts revealed UVR-absorbing compounds probably containing MAAs since they absorb wavelengths between 310 and 360 nm (Karentz et al. 1991) (Fig. 9). The occurrence of very low concentrations of UVR-absorbing compounds in our samples is consistent with the fact that diatom-dominated microbial mats usually accumulate very small amounts of UVR-absorbing compounds (Sundbäck et al. 1996b, 1997). Instead, microalgae may rely on other UVR-protective mechanisms such as an efficient UVR-induced damage repair system, as suggested by Peletier et al. (1996). Carotenoid pigments are also involved in photoprotection. For example, β,β-carotene acts as a quenching system against free radicals produced by UVR exposure (Burton & Ingold 1984). Underwood et al. (1999) noted that benthic diatoms increased their β,β-carotene concentration when faced with enhanced UVBR exposure. We found no significant difference between treatments for the β,β-carotene to chl a ratio (Table 2). Nor did we find a significant difference in the ratio of photoprotective pigments (diadinoxanthin + diatoxanthin) to chl a between treatments. Although the de-epoxidation of diadinoxanthin to diatoxanthin can play a photoprotective role when algal cells are exposed to excessive PAR light (Demers et al. 1991), its protection against UVBR and UVR has not been demonstrated in diatom-dominated microphytobenthic communities (Sundbäck et al. 1996b, Wulff et al. 1999, 2000). Finally, the total organic material in the seawater overlying the sediment might have provided some protection for the epibenthic organisms, especially at high tide. During our experiment, the concentration of total organic carbon (TOC) in the water column averaged 161 µM. UVR absorption by the chromophoric dissolved organic matter (CDOM) component of TOC can be high in the lower St. Lawrence Estuary (Whitehead et al. 2000).

Treatment effect and nutritional status

In contrast to planktonic communities, effects of nutrient supply on the response of benthic communities to changes in UVR have been poorly investigated. In a marine microtidal area of Sweden, Wulff et al. (2000) observed that nutrient limitation increased the sensitivity of benthic microalgae to ambient UVBR. The fact that we observed UVR effects on the microalgal P:B ratio despite the probable availability of inorganic nutrients implies that nutrient limitation is not a necessary condition to witness UVR effects in the St. Lawrence Estuary. Likewise, the UVR-susceptibility of some periphytonic communities inhabiting Canada’s streams was not related to their nutritional status (Bothwell et al. 1993).

UVBR versus UVAR effects

Both UVBR and UVAR were simultaneously studied in this experiment, thereby allowing for distinction between the effects of these 2 solar UV components. The enhanced microalgal P:B ratio on Days 14 and 21 in UVAR-protected cores (Fig. 6B) indicates that both solar ultraviolet components are simultaneously responsible for the treatment effect. In addition, very noticeable changes in chl a accrual and in the percentage of pheopigments were observed between reduced UVBR treatment and control, and between both protected treatments. This implies that UVBR and UVAR are separately responsible for these changes in the microphytobenthic community. UVBR is well recognized as an important stress factor for many benthic communities (Nozais et al. 1999, Wulff et al. 2000). Similarly, UVR can provoke direct inhibition of algal growth and photosynthesis in Canada’s streams (Bothwell et al. 1994). UVAR can also affect microalgae through the consumer level (‘the solar cascade’, Williamson 1995). Indeed, Bothwell et al. (1994) argued that UVR can enhance periphyton accrual through elimination of macrograzer populations (e.g. larval chironomids). In contrast, Wulff et al. (1999) reported from a 4 mo field experiment that ambient UVR did not exert any selective pressure on the overall microbenthic community in a shallow water of a microtidal sandy bay on the west coast of Sweden.

Short-term versus long-term effects

Compared to the first part of the experiment (early summer), a noticeable lack of UVR effects were noted during the second half of the study (late summer). We hypothesize that the high UVR fluxes at the beginning of the experiment enhanced the rate of photodamage relative to that of photorepair, resulting in the decreasing microalgal P:B ratio on Days 14 and 21. Then, as the summer progressed, ambient UVR fluxes decreased, resulting in a greater proportion in the rate of photorepair which eliminated the detrimental effect of UVR on the photosynthetic apparatus. Studying a planktonic community, Furgal & Smith (1997) concluded that the inhibition of photosynthesis by ambient UVBR was proportional to the seasonal variation of UVBR fluxes. Similarly, the decreasing seasonal UVR fluxes may also have softened the UVR-susceptibility of herbivorous grazers.

CONCLUSIONS

In summary, the reduction of UVR resulted in an enhancement of primary production normalized to
chi a on 2 consecutive sampling days in July, when daily incident irradiances were at their seasonal maximum. This suggests a detrimental but weak direct effect of UVR on photosynthesis. In addition, the decreased chi a concentration and diatom abundance in UVR-shielded treatments and the simultaneous increase in the percentage of pheopigments suggest an enhanced grazing pressure under these treatments. The lack of treatment effects during the second half of the experiment may be due to the decreasing UVR daily fluxes as per seasonal trends. Our results finally indicate that UVAR plays a role as important as UVBR on microphytobenthic photosynthesis and biomass, and presumably on microfauna grazing activity. Future experiments should be designed to test the influence of UV radiation on the grazing activity of microzoobenthos. Natural and seminatural experiments considering the macrozoobenthic compartment are also needed in order to improve our understanding of how microphytobenthos will respond to future changes in ambient UVBR fluxes and in the spectral balance (ratio of UVBR over PAR + UVAR + UVBR).

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