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Efficiency of photoprotection in microphytobenthos: the role of vertical migration and the xanthophyll cycle against photoinhibition

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Abstract

The capacity of estuarine microphytobenthos to withstand the variable and extreme conditions of the intertidal environment, prone to cause photoinhibition of the photosynthetic apparatus, has been attributed to particularly efficient photoprotection mechanisms. However, little is known regarding its actual photoprotection capacity or the mechanisms responsible for the protecting against photoinhibition. This study addressed these questions by (i) quantifying the photoprotection capacity and the extent of photoinhibition under high light exposure, (ii) estimating the contribution of vertical migration and the xanthophyll cycle to overall photoprotection and (iii) evaluating the effects of photoacclimation state. A new experimental protocol was developed, combining (i) chlorophyll fluorescence imaging, for the simultaneous measurement of replicates and experimental treatments, (ii) specific inhibitors for vertical migration and for the xanthophyll cycle, to quantify the relative contribution of each process, and (iii) recovery kinetics analysis of photosynthetic activity during light stress-recovery experiments, to distinguish reversible downregulation from photoinhibition. The results showed a high photoprotective capacity in both studied periods, May and October, with photoinhibition rates remaining below 20%. A clear change in photoacclimation state was observed, following the seasonal change in solar radiation, with acclimation to lower irradiances in autumn being associated with higher susceptibility to photoinhibition. Also the relative importance of vertical migration and the xanthophyll cycle varied between the sampling periods. While the two processes displayed a similar role in spring/summer, vertical migration became the dominant photoprotective process in autumn. However, the contribution of the two processes to overall photoprotection
reached only ca. 20%, suggesting the participation of other photoprotective mechanisms.

Running head: Photoprotection and photoinhibition in microphytobenthos

Key index words: microphytobenthos; photoinhibition; photoprotection; xanthophyll cycle; vertical migration; non-photochemical quenching; chlorophyll fluorescence; diatoms
INTRODUCTION

Benthic microalgae inhibiting estuarine intertidal flats are exposed to extreme and highly variable environmental conditions. Particularly during low tide, the sedimentary environment is characterized by the exposure to high levels of solar irradiance (Serôdio & Catarino 1999), including UV radiation (Waring et al. 2007, Mouget et al. 2008), extreme temperatures and salinities (Brotas et al. 2003, Rijstenbil 2005), intense rates of desiccation (Coelho et al. 2009), supersaturated oxygen concentrations (Chevalier et al. 2010), and nutrient and carbon depletion (Miles & Sundbäck 2000, Cook & Røy 2006). Being potentially damaging to the photosynthetic apparatus when acting individually, the combined effects of all these factors likely concur to the photoinhibition of photosynthesis of microphytobenthos microalgae. Of particular importance is the exposure to direct sunlight, which can result in excessive reductant pressure and in the formation of intracellular reactive oxygen species (ROS; Roncarati et al. 2008, Waring et al. 2010). High levels of ROS cause the permanent inactivation of photosystem II (PSII) protein D1, negatively impacting on photosynthetic yield and on primary productivity (Nishiyama et al. 2006).

Despite these harsh conditions, microphytobenthos of intertidal flats typically exhibit high growth rates, forming dense and diverse sedimentary biofilms, and are recognized as a major contributor to ecosystem-level carbon fixation and primary productivity (Underwood & Kromkamp 1999). Furthermore, the apparent lack of photoinhibition in microphytobenthic biofilms has been repeatedly reported (Kromkamp et al. 1998, Underwood 2002, Blanchard & Cariou-LeGall 1994, Blanchard et al. 2004, Underwood et al. 2005, Van Leeuwe et al. 2008). This success in coping with high light stress may be explained by the combined operation of two processes, the xanthophyll cycle and vertical migration, which could result in an overall particularly
efficient photoprotection (Serôdio et al. 2008, Perkins et al. 2010). In diatoms, the group
of microalgae that typically dominate in microphytobenthos assemblages, the
xanthophyll cycle has been reported to provide an exceptionally high photoprotective
capacity (Lavaud 2007, Brunet & Lavaud 2010, Goss & Jakob 2010). This is
particularly true for microphytobenthos in situ (Serôdio et al. 2005, Van Leuwee et al.
2008, Jordan et al. 2010, Chevalier et al. 2010). To this also seems to contribute the
activation of the xanthophyll cycle in the dark, attributed to chlorespiratory activity,
which has been considered as potentially advantageous during prolonged periods of
darkness (Jakob et al. 2001, Cruz et al. 2011), a situation common in the sedimentary
environment.

On the other hand, the negative phototactic behavior of benthic diatoms, mostly
raphid pennates, under high light has long been interpreted as a form of avoidance of
excessive light levels that would otherwise cause photoinhibition (Admiraal 1984,

This subject has attracted substantial attention in recent years, particularly
centered on the effects of vertical migration on biofilm photophysiology (Consalvey et
Cartaxana et al. 2011), and became facilitated by the introduction of a diatom motility
inhibitor (Cartaxana et al. 2008). However, these studies have been focused on the
response of photosynthetic activity during (Waring et al. 2007, Perkins et al. 2010) or
shortly after light stress (Mouget et al. 2008), mostly through in vivo measurements of
electron transport rate of PSII (ETR) or non-photochemical quenching (NPQ) of
chlorophyll fluorescence (PAM fluorometry, see below; Table 1) (Perkins et al. 2011).
Perhaps surprisingly, none of these studies has actually evaluated the efficiency of the
photoprotection provided by these two processes or compared their role against
photoinhibition in microphytobenthos biofilms. The distinction between photoprotection and photoinhibition processes from chlorophyll fluorescence cannot be inferred from the decrease in ETR or formation of NPQ under high light, but requires the analysis of the recovery kinetics of photosynthetic activity following exposure to high light stress (Horton & Hague 1988, Walters & Horton 1991, Müller et al. 2001). In diatoms, a rapid (within minutes) component of this recovery can be attributed to the reversal of the xanthophyll cycle ($q_E$, or ‘energy-dependent quenching’) while photoinhibitory effects ($q_I$, or ‘photoinhibitory quenching’) can be quantified from a second, much slower (within hours) component (Müller et al. 2001, Lavaud 2007). The $q_T$ (state-transition related quenching) component of NPQ recovery, which shows intermediate relaxation kinetics, does not exist in diatoms (Owens, 1986). As such, questions like ‘How efficient are photoprotective processes in preventing photoinhibition in microphytobenthos biofilms?’ ‘What is the relative contribution of migration and the xanthophyll cycle for overall photoprotection?’ or ‘To what extent does photoinhibition occur in microphytobenthos?’ are mostly unanswered.

This study was set out to address these questions, for which a new experimental protocol was designed, based on the combination of (i) chlorophyll fluorescence imaging, to allow the simultaneous measurement of a large number of samples and experimental treatments, (ii) the use of specific inhibitors for vertical migration and for the xanthophyll cycle, to quantify the relative contribution of each process to overall photoprotection, and (iii) the analysis of the recovery kinetics of photosynthetic activity following light stress, to distinguish downregulation due to the xanthophyll cycle and photoinhibition. This approach was further used to test the influence of photoacclimation state on photoprotection capacity and susceptibility to photoinhibition in microphytobenthic biofilms inhabiting a temperate intertidal mudflat.
**MATERIALS AND METHODS**

**Sampling and sample preparation.** Sediment samples were collected in the upper zone of an intertidal mudflat in the Baie de l’Aiguillon (46°15’18” N, 01°08’33” W), France, in late spring (May) and autumn (October) 2010, expected to show contrasting photoacclimation states following the seasonal variation in solar radiation (see below). The sampling site is composed of fine muddy sediments (< 63 µm) where microphytobenthic biofilms are largely dominated by diatoms (Herlory et al. 2004). During low tide, samples of the surface layers of sediment (approximately the top 1 cm) were collected using a spatula. In the laboratory, the sediment was sieved through a 500-µm mesh, to remove the mud snails *Hydrobia* sp. and other meio- and macrofauna, and was thoroughly mixed and spread in 4 cm deep plastic trays. The sediment was covered by water collected in the sampling site and left undisturbed overnight. In the next morning, at the start of the photoperiod, the slurries were again homogenized and identical portions of the resulting slurry were transferred to 24-well plates using a small spatula, filling the wells completely (ca. 3 ml). The well plates were exposed to homogeneous light field provided by two LED panels (equal contribution of red, far-red, blue and white LEDs; FloraLEDs panels, Plant Climatics, Germany) delivering a constant irradiance of 70 µmol quanta m$^{-2}$ s$^{-1}$ at the sample surface, in order to induce the upward migration of microalgae and the formation of the biofilm. Daily global solar radiation were obtained from a Meteo-France weather station located approximately 9 kms southwest from the sampling site, for two-week periods preceding the sampling dates, 15-30 May and 5-20 October 2010.

**Fluorescence measurements.** Chlorophyll fluorescence was measured using an imaging-PAM fluorometer (Maxi-PAM M-series, Walz GmbH, Effeltrich, Germany).
The measuring area of the fluorometer covered each entire well plate, so that up to a total of 24 sediment samples could be monitored simultaneously. All experiments were carried out after biofilm formation. This was determined by measuring the fluorescence level $F_s$, taken as a proxy for surface microalgal biomass, in a replicated set of samples exposed to constant low light of 55 µmol quanta m$^{-2}$ s$^{-1}$. Experiments were started after $F_s$ reached a plateau following the initial rise after the onset of the light period which typically took 2-3 hours of low light exposure. For each sample, the fluorescence signal was calculated by averaging the values of all pixels included in an area of ca. 63.6 mm$^2$ (area of interest), which corresponded to ca. 1500 pixels, centered inside each well. This area is smaller than the total area of each well (95.0 mm$^2$), the difference being due to the exclusion of the edge of each sample, often not representative of the rest of the biofilm. To minimize sample heating during prolonged exposure to high light, the experiments were carried out in a temperature-controlled room, at 20 ºC, and the fluorometer Perspex hood was maintained open at all times.

**Photoacclimation: light-response curves.** The photoacclimation state of the samples was characterized by measuring light-response curves of ETR and of NPQ in the two sampling periods. Light-response curves were generated by sequentially exposing the samples to 7 levels of actinic light, up to 700 µmol quanta m$^{-2}$ s$^{-1}$. Samples were exposed to each light level for 3 min (a period previously confirmed allowing for reaching a steady-state), after which a saturation pulse was applied and fluorescence levels $F_s$ and $F_m'$ were recorded. Six replicated measurements (on six different wells) were made for each light level. For each irradiance level, $E$, the relative ETR was calculated from the product of $E$ and the PSII effective quantum yield, $\Delta F/F_m'$ (Genty et al. 1989):
ETR vs $E$ curves were quantitatively described by fitting the model of Eilers & Peeters (1988), and by estimating the parameters $\alpha$ (the initial slope of the curve), $ETR_m$ (maximum ETR) and $E_k$ (the light-saturation, or photoacclimation, parameter):

$$ETR(E) = \frac{E}{a E^2 + b E + c}$$

where

$$\alpha = \frac{1}{c}, \quad ETR_m = \frac{1}{b + \sqrt{ac}} \quad \text{and} \quad E_k = \frac{c}{b + \sqrt{ac}}$$

Due to the unavoidable confounding effects of vertical migration on the measurement of $F_m$, NPQ was calculated using the adapted index, based on the relative difference between the maximum fluorescence measured during the construction of the light curve, $F_{m,m}$, and upon exposure to light, $F_{m}$ (Serôdio et al. 2005):

$$NPQ = \frac{F_{m,m} - F_{m}}{F_{m}}$$

NPQ vs $E$ curves were described by fitting the model of Serôdio & Lavaud (2011), and by estimating the parameters $NPQ_m$ (maximum NPQ), $E_{50}$ (irradiance corresponding to half of $NPQ_m$) and $n$ (sigmoidicity parameter):
\[
\text{NPQ} (E) = \text{NPQ}_0 \frac{E^n}{E_{50} + E^n}
\]  
(5)

These models were fitted using a procedure written in MS Visual Basic and based on MS Excel Solver. Model parameters were estimated iteratively by minimizing a least-squares function, forward differencing, and the default quasi-Newton search method. The model was fitted to individual light-response curves. Estimates of model parameters were compared using the Student’s \( t \)-test. The standard errors of the parameter estimates were calculated following Ritchie (2008).

**Photoprotection vs photoinhibition: light stress-recovery experiments.** The photoprotection capacity of microphytobenthos biofilms was estimated by quantifying the recovery of \( \Delta F/F_m' \) following a prolonged exposure to supersaturating irradiance. Three replicates were sequentially exposed to: (i) low light level of 55 \( \mu \text{mol quanta m}^{-2} \text{s}^{-1} \), for a minimum of 15 min, to ensure full light-activation of the photosynthetic apparatus and to determine pre-stress reference levels of \( \Delta F/F_m' \); (ii) supersaturating light level of 1200 \( \mu \text{mol quanta m}^{-2} \text{s}^{-1} \) for 3 hrs, to potentially induce photoinhibitory effects; (iii) low light (55 \( \mu \text{mol quanta m}^{-2} \text{s}^{-1} \)) for a minimum of 15 min to record the recovery kinetics. During the whole experiment, \( \Delta F/F_m' \) was measured by applying saturating pulses every 90 s. The recovery of \( \Delta F/F_m' \) upon the return to low light conditions was described by fitting an exponential function, adapted from a first-order kinetics model derived for describing the kinetics of NPQ (Olaizola & Yamamoto 1994, Serôdio et al. 2005):

\[
\frac{\Delta F / F_m'}{\Delta F / F_{m,rec}}(t) = \frac{\Delta F / F_{m,rec}}{\Delta F / F_{m,rec} (0) - \Delta F / F_{m,rec}} e^{-kt}
\]  
(6)
where $t$ is the time during recovery, $\Delta F/F_m'(0)$ and $\Delta F/F_m',rec$ represent the PSII quantum yield levels at the start of the recovery period and after full recovery (associated to $q_E$), and $k$ is the rate constant of $\Delta F/F_m'$ recovery. The values of $\Delta F/F_m'$ estimated by the model for $t = 10.5$ min, expressed as a percentage of the pre-stress levels, were used for estimating the effective photoprotective capacity of the biofilm. The remaining relative difference between pre- and post-stress levels of $\Delta F/F_m'$ was used as an estimate of the photoinhibitory effects imposed by high light.

The photoprotective roles of vertical migration and of the xanthophyll cycle were studied by applying specific inhibitors of the two processes. Vertical migration was inhibited by the diatom motility inhibitor Latrunculin (Lat) A, shown to effectively inhibit cell motility without causing appreciable effects on the photosynthetic activity (Cartaxana et al. 2008). To inhibit the activity of the xanthophyll cycle, the inhibitor of the diadinoxanthin de-epoxidase (DDE) dithiothreitol (DTT) was used. DTT is commonly used to inhibit the conversion of the pigment diadinoxanthin (DD) into the photoprotective form diatoxanthin (DT) (Lavaud et al. 2002a). DTT was applied in combination with Lat A, in order to ensure that the cells having the xanthophyll cycle inhibited remained exposed to high light.

The contribution of vertical migration to overall photoprotection capacity of the biofilm was estimated by the difference between the levels of $\Delta F/F_m'$ recovery in control (free moving cells) and Lat A-treated (vertical migration inhibited) samples. The contribution of the xanthophyll cycle was estimated by comparing the levels of $\Delta F/F_m'$ recovery in the samples treated with Lat A (only vertical migration inhibited) and in those treated with both Lat A and DTT (both vertical migration and the xanthophyll cycle inhibited). The inhibitor solutions were added after biofilm was fully formed, in a total of 200 μL for both the Lat A and the Lat A + DTT solutions. The same volume of
filtered seawater was added to the control samples. The solutions were added carefully
to minimize biofilm disturbance, by pipetting small volumes onto the sediment surface.
A minimum of 30 min was given for the inhibitors to diffuse and for the biofilms to
stabilize before measurements were started.

**Inhibitor preparation and effective dosage.** Solutions of Lat A of different
concentrations, ranging from 5 to 15 μM, were prepared from a concentrated solution (1
mM) prepared from dissolving purified Lat A (Sigma-Aldrich) in dimethylsulfoxide.
The minimum effective dosage of Lat A to induce inhibition of vertical migration was
determined following Cartaxana & Serôdio (2008). Samples treated with different
concentrations of Lat A (final volume, 200 μL) were darkened close to the time
expected for tidal flood, known to induce a rapid downward migration. The degree of
migration inhibition was estimated from the decrease in surface biomass following
darkening, as estimated from dark-adapted fluorescence level, $F_o$. Three replicated
samples were tested for each Lat A concentration.

DTT (BDH-Prolabo) was prepared fresh as in Lavaud et al. (2002a). A stock
solution of 300 mM (in ethanol) was diluted in filtered seawater to prepare working
solutions of concentrations ranging from 3.3 to 15 mM. The minimum effective dosage
of DTT was determined by measuring NPQ development upon exposure to 400 μmol
quanta m$^{-2}$ s$^{-1}$ for 30 min in samples treated with increasing concentration of DTT (final
volume, 200 μL). Three replicated samples were tested for each DTT concentration. For
the light stress experiments, samples were treated with 200 μL of a combined solution
of Lat A and DTT, prepared using the concentration of each inhibitor determined from
the effective dosage tests (see Results).

**Taxonomic composition.** In one of the trays, microalgae were collected by
covering the sediment with two layers of a 100 μm-mesh. The trays were exposed to
low indirect natural light from a north facing window (< 200 µmol quanta m$^{-2}$ s$^{-1}$) during the day following the sampling. The upper mesh was removed at the time of middle emersion period and it was washed with filtered (0.2 µm) natural sea water. The samples were fixed in Lugol and preserved at 4°C until their analysis. Diatom species were identified and counted using definitive mounts in Naphrax after cleaning the cells by cremation (2 h, 450°C) (Méléder et al. 2007). Taxonomic determination was performed by microscope on the basis of morphological criteria. A total of ca. 300 diatom frustules were counted to determine specific abundances.

RESULTS

Taxonomic composition

In both sampling periods, the microphytobenthic assemblages were dominated by long biraphid diatoms (length > 30 µm). In May, the assemblages were mainly dominated by Navicula cf. spartinentensis (61%, n = 350). Staurophora salina represented less than 20% of the assemblages but this species was two times longer than N. cf. spartinentensis (22 µm and 44 µm long, respectively). In October, the assemblages were co-dominated by Plagiotropis seriata (22%, n = 335) and Staurophora salina (19%); the size of P. seriata (190 µm long) was four times the one of S. salina one (44 µm long) strengthening its dominance in terms of biovolume. A third species, Pleurosigma strigosum (300 µm length) represented more than 10% of the assemblage abundance.

Photoacclimation
Significant differences were found between the light-response of ETR measured in May and October. In comparison with the ETR vs $E$ curves measured in May, the ones measured in October presented significantly higher values of $\alpha$ ($+26.7\%$, $t$-test, $p < 0.001$) and lower values of $ETR_m$ ($-41.5\%$, $t$-test, $p < 0.001$) (Fig. 1A). As a consequence, the photoacclimation parameter $E_k$ was significantly lower in October than in May ($-53.5\%$; $t$-test, $p < 0.001$). Regarding NPQ, significant differences were found between the light-response curves measured in the two periods (Fig. 1B). NPQ vs $E$ curves measured in May reached lower levels within the range of applied irradiances (on average, 2.19 and 3.25 at 700 $\mu$mol quanta $m^{-2}$ $s^{-1}$, in May and October, respectively), although the values of NPQ$_m$ were not significantly different ($t$-test, $p = 0.425$). The light-response curves were more sigmoid in May than in October ($t$-test, $p = 0.001$), the largest differences being found regarding the light level required for induction of NPQ, indicated by the parameter $E_{50}$, which was significantly lower in October than in May ($-38.5\%$; $t$-test, $p = 0.003$).

The light conditions in the region of the sampling area varied greatly between the two-week periods preceding the sampling periods, with global solar radiation reaching a daily average of 2369 J cm$^{-2}$ in May, more than double the value observed in October, 1008 J cm$^{-2}$.

**Inhibitor dosage**

Vertical migration was strongly inhibited for most of the Lat A concentrations tested, with an inhibition level above 75% being obtained with only 5 $\mu$M (Fig. 2). The inhibitory response to the increase in Lat A concentration presented a clear saturation-like pattern, with the increase from 10 to 15 $\mu$M resulting in an increase in inhibition of
only 8.5%. Considering that 10 μM was enough to inhibit vertical migration by more than 90%, and the small increase obtained by applying the higher concentrations, solutions of 10 μM Lat A were used in all experiments.

The response of NPQ to the increase in DTT also showed a saturation-like pattern, characterized by a strong decrease for concentrations up to 5 mM, and a virtually constancy for concentrations above this value (NPQ decreased by 19% between 5 and 15 mM; Fig. 3). However, even when the highest DTT concentration was applied, NPQ was never completely eliminated, remaining above 1.0. In all further experiments, a concentration of DTT of 10 mM was used.

**Light stress exposure and recovery**

Figure 4 exemplifies the variation of ΔF/F_m' during a light stress-recovery experiment. On control samples, exposure to high light induced an immediate and marked decrease in ΔF/F_m' from ca. 0.63 to values slightly below 0.1 (Fig. 4). ΔF/F_m' further decreased to values close to zero during the first 15 min of exposure, after which it gradually recovered, stabilizing at values around 0.1 after 90 min and until the end of the high light period. On inhibitor-treated samples, ΔF/F_m' also decreased to values close to zero upon the start of high light exposure, but, as opposed to control samples, never showed any appreciable recovery, remaining below 0.05 (Fig. 4). However, ΔF/F_m' levels were usually higher in Lat A-treated samples than in those treated with both inhibitors (Figs. 4, 5). Following the transition to low light, a clear recovery response was observed for all samples, with ΔF/F_m' reaching in all cases over 60% of initial values after 15 min. Treatment with Lat A effectively inhibited vertical migration during the whole experiment, as indicated by the small variation in F_s in Lat A-treated
samples (on average, -12.1\% for samples treated with Lat A and Lat A+DTT) as compared with the controls (-43.5\%; Fig. 6). The effects of inhibitors were particularly evident during recovery under low light, during which $\Delta F/F_{m}'$ followed the negative exponential pattern described by Eq. (6), the fit of which was very good in all cases ($r^2 > 0.91$; Fig. 7). Control samples recovered more rapidly than those treated with inhibitors, so that after 3 min after return to low light, $\Delta F/F_{m}'$ of non-inhibited samples was over 70\% and 60\% higher than on samples treated with Lat A in May and October, respectively. In both periods these differences were gradually reduced during exposure to low light, but after 10.5 min the percentage of recovery was significantly different among treatments and sampling periods (two-way ANOVA, $p < 0.001$ for both factors). In both May and October, the recovery of $\Delta F/F_{m}'$ was higher in the controls than in the Lat A-treated samples (Control vs Lat A; Tukey’s post-hoc test, $p = 0.043$ and $p = 0.010$, respectively), which was in turn higher than in samples treated with Lat A and DTT (Lat A vs Lat A+DTT; Tukey’s post-hoc test, $p = 0.042$ and $p = 0.030$, respectively). The percentage of recovery was in all cases significantly higher in May than in October (Tukey’s post-hoc test, $p < 0.05$), with the exception of samples treated with both inhibitors (Tukey’s post-hoc test, $p = 0.107$).

**Photoprotection efficiency and extent of photoinhibition**

Depending on the species, the full recovery of the xanthophyll cycle after a transition from high to low light mainly occurs after 6 min to 15 min (Goss et al. 2006, Lepetit & Lavaud, pers. obs.). Considering the intermediate period of 10.5 min, the recovery of $\Delta F/F_{m}'$ at this time was used as an estimate of the photoprotection capacity and to calculate the extent of photoinhibition occurred. The results indicate that the
microphytobenthos biofilms had a large photoprotective capacity in both periods, with a correspondingly low percentage of photoinhibition below 25%, although higher in May than in October (87.7 and 78.0%; Fig. 8, Table 2). From the reduction in the photoprotection capacity measured in samples treated with inhibitors, the contribution of vertical migration and of the xanthophyll cycle to overall photoprotection were estimated to reach a combined value only slightly above 20% (Table 2). While in May the two processes had a comparable contribution to photoprotection, the relative importance of the xanthophyll cycle was reduced to 7.2% in October.

**DISCUSSION**

**Photoacclimation and susceptibility to photoinhibition**

Comparatively to May, samples collected in October appeared acclimated to lower light levels, showing the pattern typically associated to ‘shade-acclimation’: a combination of higher values of $\alpha$ and of lower values of $\text{ETR}_m$, resulting in lower values of $E_k$, usually taken as an indication of photosynthesis saturating at lower irradiances. This change in photoacclimation state between May and October was consistent with the observed seasonal change in solar light conditions preceding the two sampling periods (i.e. global solar radiation more than two times higher in May than in October). These results also generally confirmed previous observations on the seasonal variability of microphytobenthos photosynthetic performance, showing patterns of acclimation to higher light levels during spring/summer and to lower levels in autumn/winter (Blanchard et al. 1997, Migné et al. 2004, Serôdio et al. 2006). They were also
consistent with the photoacclimation response of benthic diatoms grown in culture exposed to low and high-light regimes (Perkins et al. 2006, Schumann et al. 2007, Cruz & Serôdio 2008). Increases of α, as the observed from May to October, are commonly attributed to an increase in the cellular content of light-harvesting pigments, increasing the fraction of incident light that is intercepted and absorbed for photosynthesis; decreases in ETR$_m$ are typically associated with the decrease of the activity of the electron transport chain or the Calvin cycle, limiting factors of light-saturated photosynthesis (Henley 1993, MacIntyre et al. 2002, Behrenfeld et al. 2004).

A change in light response was also noticeable regarding NPQ, with the samples collected in October showing NPQ activation starting at lower light levels (lower $E_{50}$) and higher values of NPQ for most irradiances (higher NPQ$_m$). As with ETR, the observed variation in the NPQ vs $E$ curves was consistent with the previously reported for microphytobenthos (Serôdio et al. 2006) or for benthic diatoms acclimated to different light regimes (Cruz & Serôdio 2008).

However, while changes in the light-response of ETR may be interpreted and related to underlying physiological processes in a relatively straightforward manner, the physiological meaning of changes in NPQ levels is more difficult to ascertain. This is because the two components of NPQ, $q_E$ (photoprotection) and $q_I$ (photoinhibition) can only be distinguished through the analysis of the recovery kinetics after exposure to high light, but not from NPQ light curves. In this study, the light stress-recovery experiments allowed to conclude that the observed change in the NPQ light-response curves was due to a decrease in the $q_E$ component and a concomitant increase in the $q_I$ component. In the absence of information from NPQ recovery kinetics, similar increases in NPQ vs $E$ curves in autumn/winter periods have been, perhaps wrongly,
interpreted as being due to an increase in photoprotective capacity (Serôdio et al. 2005, 2006).

Furthermore, the results from the light stress-recovery experiments revealed an association between photoacclimation status and photoprotection efficiency, not shown before for these communities. Whatever the cause (see below), the acclimation to high light levels in summer was associated to a high photoprotection capacity and the low light-acclimation in autumn to a general loss in photoprotection and a higher susceptibility to photoinhibition.

**Photoprotection vs photoinhibition**

A central finding of this study is that photoinhibition was in all cases considerably low (ca. 20%), indicating photoprotection to be particularly efficient in the studied microphytobenthos biofilms. Despite the general view that these assemblages are largely immune to photoinhibition (Blanchard et al. 2004, Waring et al. 2007, Mouget et al. 2008), this process has been shown to occur under *in situ* conditions (Serôdio et al. 2008). Curiously, the rates of photoinhibition estimated in the cited study, reaching up to ca. 18%, are similar to the values here reported, despite the fact that they were estimated from hysteresis patterns observed during entire low tide exposure periods. The results of the present study therefore confirm that the photoprotective mechanisms available to benthic diatoms are not completely efficient in preventing some degree of photoinhibitory damage. However, it should be stressed the difficulty in comparing the measured rates of photoinhibition with results published for other habitats, or for other estuarine primary producers such as phytoplankton, seagrasses or macroalgae. Apart from the light history and the species-specific
differences, the extent of photoinhibition is directly related to light dosage, determined
by light intensity and duration of exposure, both largely variable amongst the different
experimental protocols used in different laboratory and field studies.

A number of unaccounted factors may have contributed for the measured low
values of $q_I$. Firstly, the well-known effect of depth-integration of subsurface
fluorescence (Forster & Kromkamp 2004, Serôdio 2004). This effect is caused by the
fact that only the cells at or near the surface are actually exposed to measured levels of
high light, and that the fluorescence signal measured at the surface also accounts for
cells positioned deeper in the photic zone and exposed to lower light levels. The
expected effect is a light-dependent overestimation of biofilm-level $\Delta F/F^\prime_m$ relatively to
the inherent, physiological values of the cells at the surface, which is then expected to
cause a systematic overestimation of $q_E$ and the underestimated of $q_I$ (Serôdio 2004).

However, besides this static effect, also dynamic effects can be expected. During
prolonged exposure to high light, the downward migration of microalgae to less
illuminated layers is likely to induce a gradual increase of $\Delta F/F^\prime_m$ (as measured at the
surface), independently of any photophysiological changes, thus causing the
overestimation of $q_E$. It is also conceivable that these types of effects may affect the
measurement of $\Delta F/F^\prime_m$ during the recovery under low light, due to upward migration as
a response to the decrease in incident irradiance. This, however, seems less likely due to
the relatively short time of this period and to the fact that a transition from high to low
light is a weaker stimulus for vertical migration, especially if coinciding with the end of
the low tide period (Coelho et al. 2011).

A second factor that might explain the low values of $q_I$ is the light doses applied
during the light stress-recovery experiments in the laboratory. Because these (3 hrs,
1200 µmol quanta m$^{-2}$ s$^{-1}$) were likely lower than the ones received during a typical
period of exposure at low tide (up to 8-10 hrs, 1500-2000 µmol quanta m\(^{-2}\) s\(^{-1}\)), larger, but still ecologically relevant, light doses could have been applied which would likely induce larger cumulative photoinhibitory effects. The light exposure conditions applied in this study, both regarding light intensity and duration, resulted from a compromise between inducing measurable effects, instrument limitations (maximum PAR irradiance provided by the imaging fluorometer) and minimizing uncontrollable experimental conditions (excessive sample heating and desiccation caused by the fluorometer LED panel). Despite these limitations, mostly instrument-related, the laboratory experimental approach used in this study has the advantage over studies carried out under in situ conditions (e.g. Serôdio et al. 2008; Perkins et al. 2010) of allowing applying controlled and reproducible conditions, making it possible to directly compare the migratory and physiological responses of samples collected in different places and occasions.

The estimation of \(q_E\) and \(q_I\) is also directly affected by the type of analysis made on the recovery kinetics in order to distinguish the two components of NPQ. For higher plants, \(q_E\) and \(q_I\) are distinguished on the basis of the recovery rate of \(F_v/F_m\), typically 10-15 min, assumed to correspond to the full reversal of the xanthophyll cycle (Horton & Hague 1988, Ruban & Horton 1995). Following the common practice for the distinction of \(q_E\) and \(q_I\), in this study these two components of NPQ were estimated based on a relaxation time of the xanthophyll cycle of 10.5 min. However, to evaluate the possible effects of considering different times for the reversal of the xanthophyll cycle on the relative magnitude of \(q_E\) and \(q_I\), a sensitivity analysis was performed, consisting on the re-calculation of these estimates when considering 6 and 15 min, values matching the range of relaxation times of the xanthophyll cycle expectable for diatoms (Gross et al. 2006, Lepetit & Lavaud, pers. obs.). The use of these different recovery periods did not alter significantly the general findings of the study, including
high levels of recovery and low photoinhibition rates, the increase in photoinhibition
levels from May to October, and a relatively low (< 30%) combined contribution of
vertical migration and xanthophyll cycle to overall photoprotection (Table 2).
Nonetheless, this analysis shows some effects, although largely expected from the
asymptotic pattern of $\Delta F/F_m'$ recovery during the considered period: the use of a shorter
period resulted in the estimation of lower rates of recovery, leading to a likely
overestimation of photoinhibition rates; conversely, longer periods resulted in larger
rates of recovery and probably overestimated levels of photoprotection (Table 2).
Moreover, due to the different relaxation patterns of samples exposed to different
treatments, the evaluation of the relative importance of vertical migration and the
xanthophyll cycle was also affected by the time period considered, with shorter and
longer recovery periods resulting in a higher apparent contribution of vertical migration
and of the xanthophyll cycle, respectively. These effects, however, did not affect
substantially the overall pattern of variation of the role of the two photoprotective
processes between the two sampling periods.

Recently, more sophisticated methods, based on the mathematical modeling and
deconvolution of the recovery curve, were proposed to trace the recovery of each
individual component of NPQ (Roháček 2010). This method could not be applied in this
study because of the particularities of the xanthophyll cycle in diatoms, which may not
verify the assumptions of the method. Firstly, the lack of $q_T$ (the state-transition
quenching) in diatoms (Owens 1986, Lavaud 2007, Goss & Jakob 2010), which called
for the modification of this model to a two-component NPQ. Secondly, the
impossibility of using changes in $F_s/F_m$ in biofilms as an indication of photoinhibition,
as this requires the darkening of the samples, known to induce changes in $F_m$ levels due
to vertical migration. Furthermore, in benthic diatoms, dark adaptation often causes the
$F_m$ level to decrease to values below $F_m'$ levels measured under low light (Serôdio et al. 2006). These reasons also prevented the use of other recently proposed methods to quantify the components of NPQ (Ahn et al. 2009, Guadagno et al. 2010).

The formation of DT in the dark and thus anoxic subsurface layers of the sediment, known to occur in diatoms (Jakob et al. 2001), and especially in benthic assemblages (Serôdio et al. 2006), is a likely explanation for the apparent impossibility to completely eliminate NPQ by applying the xanthophyll cycle inhibitor DTT (Fig. 3). The DT thus formed would remain present despite the treatment with DTT, which prevents new conversion of DD to DT, but does not induce the reversed reaction. Upon exposure to high light, the oxygenation of DT-rich subsurface layers would allow for the observed rise in NPQ, as the formation of NPQ from DT is known to be inhibited by anoxia (Cruz et al. 2011).

**Photoprotection: vertical migration vs xanthophyll cycle**

The use of specific inhibitors for vertical migration and for the operation of the xanthophyll cycle allowed estimating the relative contribution of each of these processes to overall photoprotection of the biofilm. The results showed a change with season and photoacclimation state of their relative importance. While in May the two processes seemed to contribute similarly to biofilm photoprotection, the loss of photoprotection capacity from May to October was associated to a decrease in the contribution of the xanthophyll cycle, so that vertical migration became the dominant photoprotective process. The observed change in the species composition of the microphytobenthic assemblage may explain this difference as the activity of the xanthophyll cycle can differ from a species to another (Lavaud et al. 2004, Goss et al.)
2006). It may be also hypothesized that this difference is related to the decrease in rates of enzymatic conversion between DD and DT associated to photoacclimation or due to acclimation to lower temperatures (Van Leuwee et al. 2008), an effect that is also species-related (Salleh & McMinn, 2011). Nevertheless, these results indicate that behavioral photoprotection seems able to maintain the overall photoprotection capacity, compensating for the decrease in the contribution of the xanthophyll cycle during the winter season.

The change in species composition, involving a dominance of larger cells in October, could also have affected the migratory response of the assemblages to high light. However, although some studies have shown a relation between migratory cell size and migratory behaviour in sediments (Hay et al. 1993, Underwood et al. 2005), there is no evidence that cell size is an important factor regarding the migratory response to light stress.

Vertical migration and the xanthophyll cycle have been considered as the main photoprotective mechanisms in microphytobenthos biofilms (Serôdio et al. 2005, Jesus et al. 2006, Mouget et al. 2008, Serôdio et al. 2008; Perkins et al. 2010). A perhaps surprising result of this study is the relative low contribution of these two processes to global photoprotection. This calls for the potential role of other processes responsible for the observed low rates of photoinhibition. Likely candidates include the cyclic electron flow around PSII (Lavaud et al. 2002b, 2007), the efficient scavenging of reactive oxygen species (Roncarati et al. 2008, Waring et al. 2010) or high turnover rates of the PSII protein D1 (Wu et al. 2011).

Use of inhibitors on microphytobenthic biofilms
An aim of this study was the introduction of a new experimental protocol to estimate photoprotection efficiency and the extent of photoinhibition in microphytobenthos biofilms. This involved the combination of: (i) the use of specific inhibitors for different photoprotective processes, applied alone and in combination with each other, allowing the estimation of the relative contribution of each process to overall photoprotection, and (ii) the use of imaging fluorometry on replicated samples in well plates, taking advantage of the self-forming nature of microphytobenthos biofilms from homogenized sediments, which allowed for adequate replication and low variability between replicates, and for the simultaneous testing of different treatments.

Some potential pitfalls exist regarding the use of inhibitors on biofilms and the interpretation of results. Firstly, it must be noted that when comparing controls (no inhibitor added) with Lat A-treated samples, it is likely that the differences in fluorescence parameters observed over time may not be attributed only to changes in cell physiological conditions but also to changes in cell composition in the upper layers of the sediment. This is because in the controls, as opposed to Lat A-treated samples, cells initially at the surface likely migrated down into layers below the photic zone, therefore changing the contribution to the fluorescence signal measured at the surface. As a consequence, any observed differences are expected to represent mainly changes at the biofilm (i.e., community)-level, and not only changes in the physiology of individual cells. This also explains the need to combine Lat A and DTT if the effect of inhibiting the xanthophyll cycle is to be evaluated in the same microalgal assemblage. By adding DTT to samples treated with Lat A, it is ensured that the same cells remain in the photic zone of the sediment and that measured changes in fluorescence are due to changes in their physiological status and not to changes in community composition. If only DTT is applied (Perkins et al. 2010), only biofilm-level effects can be evaluated, as
many cells will likely respond to high light by migrating downward and become
unobservable (Oxborough et al. 2000).
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Lavaud L, Kroth PG (2006) In diatoms, the transthylakoidal proton gradient regulates the photoprotective non-photochemical fluorescence quenching beyond its control on the xanthophyll cycle. Plant Cell Physiol 47:1010-1016


Table 1. Notation

\( \alpha \), initial slope of the ETR vs E curve

\( \alpha, \beta, \gamma \), parameters of the Eilers and Peeters (1988) model

DTT, dithiothreitol

\( \Delta F/F_m' \), effective quantum yield of PSII

DD, diadinoxanthin

DT, diatoxanthin

DDE, diadinoxanthin de-epoxidase

E, PAR irradiance

\( E_{50} \), Irradiance level corresponding to 50% of NPQ_m in a NPQ vs E curve

\( E_k \), photoacclimation parameter of an ETR vs E curve

ETR, PSII electron transport rate

ETR_m, maximum ETR level in an ETR vs E curve

\( F_o, F_m \), Minimum and maximum fluorescence of a dark-adapted sample

\( F_s, F_m' \), Steady state and maximum fluorescence of a light-adapted sample

Lat A, Latrunculin A

n, sigmoidicity coefficient of the NPQ vs E curve

NPQ, non-photochemical quenching
NPQ<sub>m</sub>, maximum NPQ level in a NPQ vs E curve

PAR, photosynthetically active radiation

PSII, photosystem II

q<sub>E</sub>, energy-dependent quenching

q<sub>I</sub>, photoinhibitory quenching

t, time during recovery following light stress

XC, xanthophyll cycle
Table 2. Extent of photoinhibition and efficiency of photoprotection (%), calculated as percentage of $\Delta F/F_{m'}$ recovery after 10.5 min. Relative contributions of vertical migration and of the xanthophyll cycle to overall photoprotection (%), as calculated from the reduction of the $\Delta F/F_{m'}$ recovery in samples treated with Lat A and with Lat A and DTT, respectively, relatively to control samples. Mean and standard error of three replicates. Numbers within parenthesis indicate results obtained when considering 6 and 15 min of recovery, respectively.

<table>
<thead>
<tr>
<th></th>
<th>May</th>
<th>October</th>
</tr>
</thead>
<tbody>
<tr>
<td>Photoinhibition</td>
<td>12.3 ± 0.55</td>
<td>22.0 ± 2.79</td>
</tr>
<tr>
<td></td>
<td>(16.6, 11.7)</td>
<td>(33.7, 17.0)</td>
</tr>
<tr>
<td>Recovery</td>
<td>87.7 ± 0.55</td>
<td>78.0 ± 2.79</td>
</tr>
<tr>
<td></td>
<td>(83.4, 88.3)</td>
<td>(66.3, 83.0)</td>
</tr>
<tr>
<td>Vertical migration</td>
<td>10.6</td>
<td>14.3</td>
</tr>
<tr>
<td></td>
<td>(24.0, 3.6)</td>
<td>(17.0, 11.7)</td>
</tr>
<tr>
<td>Xanthophyll cycle</td>
<td>10.1</td>
<td>7.2</td>
</tr>
<tr>
<td></td>
<td>(6.4, 13.2)</td>
<td>(7.3, 5.8)</td>
</tr>
<tr>
<td>Others</td>
<td>79.3</td>
<td>78.5</td>
</tr>
<tr>
<td></td>
<td>(69.5, 83.2)</td>
<td>(75.7, 82.5)</td>
</tr>
</tbody>
</table>
Fig. 1. Light-response curves of ETR (A) and NPQ (B) measured in May and October 2010. Mean of six replicates. Vertical bars: one standard error. Numbers represent the mean values of model parameters estimated for light-response curves measured for each individual sample.

Fig. 2. Variation of migration inhibition with the concentration of added Lat A solution. Mean of three replicates. Vertical bars: one standard error. Line represents the fit of exponential decay model.

Fig. 3. Inhibition of NPQ as a function of concentration of added DTT solution. NPQ induced upon exposure to 400 µmol quanta m⁻² s⁻¹. Mean of three replicates. Vertical bars: one standard error. Line represents the fit of exponential decay model.

Fig. 4. Light stress-recovery experiment. Variation of PSII quantum yield, ΔF/ΔF_m', during sequential exposure to low light (pre-stress, 55 µmol quanta m⁻² s⁻¹), light stress under high light (high light, 1200 µmol quanta m⁻² s⁻¹, 180 min) and recovery under low light (recovery, 55 µmol quanta m⁻² s⁻¹, 10.5 min) for controls and for samples treated with migration inhibitor Lat A and with migration and xanthophyll cycle inhibitors (Lat A + DTT), collected in May 2010. Mean of three replicates. Vertical bars: one standard error.

Fig. 5. Images of PSII quantum yield, ΔF/ΔF_m' (false color scale), as measured in the sediment samples used in the light stress experiment described in Fig. 4 at the end of
first low light exposure (pre-stress, 55 µmol quanta m$^{-2}$ s$^{-1}$), at the end of high light exposure (high light, 1200 µmol quanta m$^{-2}$ s$^{-1}$, 180 min) and following recovery under low light (recovery, 55 µmol quanta m$^{-2}$ s$^{-1}$, 10.5 min). Three replicated areas of interest for each treatment.

Fig. 6. Images of fluorescence level $F_s$ (false color scale) as measured in the sediment samples used in the light stress experiment described in Fig. 4. at the end of first low light exposure (pre-stress, 55 µmol quanta m$^{-2}$ s$^{-1}$), and after recovery following end of high light exposure (high light, 1200 µmol quanta m$^{-2}$ s$^{-1}$, 180 min; recovery, 55 µmol quanta m$^{-2}$ s$^{-1}$, 10.5 min). Three replicated areas of interest for each treatment.

Fig. 7. Recovery of PSII quantum yield, $\Delta F/F_{m'}$, during relaxation following light stress for control samples and for samples treated with migration inhibitor Lat A and with migration and xanthophyll cycle inhibitors (Lat A + DTT). Lines represent the exponential model described by Eq. (6) fitted to average $\Delta F/F_{m'}$ values. Detail of Fig. 4.

Fig. 8. Efficiency of photoprotection, as percentage of recovery after 10.5 min low light following high light exposure, in May and October, for controls and inhibitor-treated samples. Mean of three replicates. Vertical bars: one standard error.
Figure 2

Migration inhibition (%) vs. Lat A concentration (μM)

- Data points for different Lat A concentrations are plotted.
- The curve shows a trend of increasing migration inhibition with increasing Lat A concentration.

43
Figure 3
Figure 4
Figure 5
Figure 6
Figure 7

![Graph showing the change in \( \Delta F/F_m \) (% initial) over time (min) for different conditions: Lat A + DTT, Lat A, and Control.](image-url)
Figure 8

![Graph showing recovery (%) for different conditions: Control, Lat A, and Lat A + DTT. The graph compares May and Oct samples.](image)