Different carbon isotope fractionation patterns during the development of phototrophic freshwater and marine biofilms

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Abstract. Natural phototrophic biofilms are influenced by a broad array of abiotic and biotic factors and vary over temporal and spatial scales. Different developmental stages can be distinguished and growth rates will vary due to the thickening of the biofilm, which is expected to lead to a limitation of light or mass transport. This study shows that variation in CO2(aq) availability leads to a fractionation shift and thereby affects δ13C signatures during biofilm development. For phototrophic freshwater biofilms it was found that the δ13C value became less negative with the thickening of the biofilm, while the opposite trend was found in marine biofilms. Modeling and pH profiling indicated that the trend in the freshwater system was caused by an increase in CO2(aq), limitation resulting in an increase of HCO3− as C-source. The opposite trend in the marine system could be explained by a higher heterotrophic biomass and activity causing a higher carbon recycling and thereby lower δ13C values. We conclude that δ13C was more related to the net areal photosynthesis rate and carbon recycling, rather than to the growth rate of the biofilms.

1 Introduction

Phototrophic biofilms are surface-associated microbial communities, in which light is the ultimate source of energy and biomass originates mainly from microalgae and bacteria. They thrive on submerged biotic or abiotic substrata in light-exposed aquatic environments and exo-biopolymers (mostly polysaccharides) provide adhesion and cohesion to the microbial consortia in these biofilms (Decho, 2000). During the initial phase of biofilm development the maximum growth rate is determined by the incident irradiance, since at that stage the biofilm thickness does not limit diffusive transport of any of the substrates. Eventually, with the thickening of a biofilm both light attenuation and diffusive transport will progressively become limiting factors for growth. Diffusion limitation will increase during the development of a biofilm due to an increased biovolume-to-surface ratio, while the potential volumetric enzymatic rates remain equal. At high irradiiances mass transfer eventually determines the maximum thickness of a biofilm.

Phototrophic biofilms grow in all aquatic systems ranging from freshwater to hyper saline and all oxygenic phototrophs use RUBISCO in their photosynthetic apparatus for carbon fixation, and the inorganic carbon source used by RUBISCO is CO2(aq). One difference between freshwater and seawater chemistry is that seawater has higher inorganic carbon contents and stronger carbonate buffering (Stumm and Morgan, 1995). This results in a higher pH and a dominance in bicarbonate concentration relative to CO2(aq). In marine systems the dissolved inorganic pool consists of dissolved CO2 (<1%), HCO3− (~95%), and CO32− (~5%). Due to the low CO2(aq) concentration and a slow chemical conversion rate of HCO3− to CO2, the supply of CO2(aq) may be considered as a potentially limiting factor. On the other hand, high respiration rates of heterotrophic and phototrophic organisms present in biofilms may elevate the CO2(aq) levels and as a result lower the pH.
The inorganic carbon pool available for phototrophs consists of carbon atoms with different atomic weights, mainly 12C (~99%) and 13C (~1%) (Raven, 1998). Organic carbon in phototrophic organisms is generally depleted in 13C relative to the carbon source. This depletion is caused by a biological fractionation due to an enzymatic discrimination against 13C in the photosynthetic process (Hayes, 1993).

Under non-limiting conditions, fractionation by RUBISCO results in a δ13C ranging from −29‰ in plants to −21‰ in cyanobacteria (Roeseke and O’Leary, 1985); eukaryotic algae have δ13C values intermediate of these (Lewis et al., 2000). However, the overall fractionation of δ13C in phototrophic biofilms is not only the result of the biochemical properties of RUBISCO, but is also affected by CO2(aq) limitation within the biofilm, which will lower the effective enzymatic fractionation of RUBISCO for two reasons: First, the competition between both substrates 13CO2(aq) and 12CO2(aq) for the binding sites in RUBISCO becomes less as the diffusive transport becomes more important for the CO2 binding rate to the enzyme; it can be assumed that the diffusive transport rate is equal for both substrates. Second, discrimination against 13C will lead to a 13C enrichment of the remaining CO2(aq) pool. A shift in isotopic composition of the dissolved inorganic carbon (DIC) pool will result in an additional change in δ13C of the organic matter since fractionation (ε) is independent of the δ13C from the source (Lewis et al., 2000; Werne and Hollander, 2004; Hayes, 1993).

Many algae can increase the conversion rate of HCO3− to CO2(aq) by the enzyme carbonic anhydrase (CA) (Tortell and Morel, 2002; Cassar et al., 2004) and thereby increase the availability of CO2 for RUBISCO. This enzyme catalyzes the hydration and dehydration of CO2. The presence of CA activity in phototrophic biofilms will influence the δ13CO2(aq) and can increase the biomass δ13C up to 10‰ (Goericke et al., 1994).

Most studies on δ13C in photosynthetic microorganisms focus on phytoplankton species and it is found that δ13C values of cultured marine phytoplankton species vary from −30‰ to −18‰, while freshwater phytoplankton δ13C ranges between −40‰ to −25‰. The δ13C values do not only vary amongst species (Lewis et al., 2000), they also vary within species due to environmental factors such as growth rate, pH of the medium (i.e. CO2 availability) and irradiance (Johnston et al., 2004; Swansburg et al., 2002). δ13C values are frequently used as proxies for food sources in food web studies (delGiorgio and France, 1996; March and Pringle, 2003). Since phototrophic biofilms can be important food sources in aquatic ecosystems (Bott, 1996; Charlebois and Lamberti, 1996; France, 1996; Boschker et al., 2005) it is important to understand the processes that cause variations in δ13C within such surface-associated communities. Little is known about the factors that influence the δ13C of phototrophic biofilms but the values are reported to vary with growth rate, irradiance, turbulence, and flow velocity (Trudeau and Rasmussen, 2003; France, 1995a).

In this study we investigated the development of marine and freshwater phototrophic biofilms grown under defined environmental conditions from natural inoculates. We measured the δ13C values at the different stages in the biofilm development along with a range of water chemistry and biofilm parameters. The aim of the study was to analyze the variation of the δ13C during biofilm development and identify key biotic and abiotic processes for carbon isotope fractionation in phototrophic biofilms.

2 Methods

Freshwater and marine phototrophic biofilms were grown on removable transparent polycarbonate slides in an incubator (Zippel and Neu, 2005), allowing the development of algal-dominated biofilms at defined irradiance and flow regimes. The incubator contains 4 physically separated incubation lanes, each with a specific irradiance. Every lane had its own growth medium reservoir. Incubators were inoculated with homogenized phototrophic biofilm material. The inoculum for the freshwater biofilms grew on surfaces in the sedimentation tank of the waste water treatment of Fumicino airport, Rome (Congestri et al., 2005). The marine inoculum originated from biofilms growing on continuously submerged surfaces in flowing Oosterschelde water, the Netherlands. Incubation material was mechanically homogenized and then frozen in order to kill fauna and prevent top-down control of the biofilm as much as possible. The freshwater medium was a modified BG 11-medium (http://www.pasteur.fr/recherche/banques/PCC/Media.htm), with 20 µM silicate to allow diatom development. The marine biofilms were grown in medium prepared with commercially available aquarium sea salt (HW Meeressalz Professional, Wiegand, Germany) with additional silicate (20 µM), nitrate (0.35 mM), and phosphate (24 µM).

The medium (4 L per lane) was continuously circulated at flow rates of 25 or 100 l/h and was replaced twice a week. The water level was ~3 mm above the biofilm, resulting in a water velocity of ~0.5 or 2 m/s. The pH of the medium was measured with a pH electrode after its preparation. The pH of renewed freshwater medium was 7.7 while it was 8.1 for the marine medium. It was found that the pH changed during the residence period of 3 or 4 days of the medium in the incubator. Biofilms were grown at a range of different growth conditions (Table 1) and the development of the biofilm was followed by continuous recording of the light absorbance using 9 light sensors glued on the bottom of several slides along each flow lane. Absorbed light absorbance was used as a proxy for the phototrophic biomass at the given growth conditions (Zippel et al., 2007). Growth rates were estimated by fitting a Richards logistic growth equation (Sidorkewicz et al., 1999) through the light absorption values. The fit quality of the logistic growth model through the measuring points was high (average r²=0.975±0.038, lowest r² value was 0.820).
Table 1. Growth conditions and relative growth rates at the different incubations. Relative growth rates were calculated at a biomass 50% light absorption using the fitted parameters from the logistic growth equation. The fit quality of the growth model trough the measuring points was high (average $r^2=0.975+0.038$).

| medium   | temperature | Flow m/s | Relative Growth rate at 50% absorbance |
|----------|-------------|----------|----------------------------------------|
|          |             |          | Irradiance 15 | 30 | 60 | 120 |
| freshwater | 20          | 0.5      | 0.54           | 1.55 | 1.56 | 1.78 |
|           |             |          | 2             | 0.65 | 0.86 | 1.14 | 2.11 |
| salt water | 25          | 0.5      | 0.34           | 0.52 | 1.08 | 1.44 |
|           |             |          | 2             | 0.66 | 1.18 | 2.40 | 3.31 |

For sampling the biofilm samples were scraped off the slide and the wet weight was determined for each sample. The samples dry weight was measured and samples were stored at $-80^\circ$C until $\delta^{13}$C analysis. The wet weight of the biofilms was found to increase linearly with light absorption values of up to 85–90% (data not shown). The $r^2$ was 0.38 ($n=34$) and 0.55 ($n=44$) for the pooled freshwater and marine incubations respectively (both regression lines were significant; $P<0.01$).

The $\delta^{13}$C values were determined at 3 different growth stages during biofilm development. A first sample was always taken 10 days after inoculation (initial phase), the second sample was taken at the first working day after $\sim$50% light absorption was reached (exponential growth phase) and the third sample was taken at the first working day after $\sim$90% light absorption was reached (mature or stationary phase). Sampling started close to the outlet of the incubator to prevent disturbance of biofilms growing on the other slides. Sample slides were replaced by slides with no biofilm to prevent additional turbulence caused by height differences. There were not enough slides in the incubator for (pseudo) replicate sampling per irradiance per growth stage. However, a test run on freshwater medium revealed $\delta^{13}$C values of $-23.33\pm0.48$, $-24.55\pm0.33$, $-27.66\pm1.07$ and $35.17\pm0.02$ for the lanes with irradiances of 120, 60, 30 and 15 $\mu$mol photons m$^{-2}$ s$^{-1}$ respectively. Standard deviations of the average values ($n=3$) were low (SD<4% of average value) and therefore a randomly sampled slide may be considered as representative for the entire lane.

2.1 Stable isotope analysis

Biofilm samples were analyzed for natural $\delta^{13}$C abundance by a total combustion elemental analyzer coupled to an isotope ratio mass spectrometer (IRMS) (Finnigan, Germany). Samples were combusted at 1010°C and transported with a helium (5.0 purity) carrier gas flow to the IRMS for determining the isotopic $^{13}$C/$^{12}$C ratio of carbon. Stable isotope ratios were calculated as:

$$\delta^{13}C = \left( \frac{\delta^{13}C_{\text{sample}}}{\delta^{13}C_{\text{standard}}} - 1 \right) \times 1000$$ (1)

where the standard is the C-isotope ratio of Vienna PeeDee Belemnite (0.0112372).

Samples for total dissolved inorganic carbon (DIC) were taken weekly per lane as well as of new made medium. The concentration of DIC in the medium was determined by acidifying 50ml medium with 500 $\mu$l pure phosphoric acid in a closed container (Crimp Seal, Chrompack, the Netherlands) to convert all HCO$_3^-$ and CO$_3^{2-}$ into CO$_2$. The samples were stored at room temperature to reach equilibrium with the gas phase (5 ml). 500 $\mu$l of the gas phase was injected into a GC with a Poraplot Q column which was linked to the IRMS (Finnigan, Germany) for $\delta^{13}$C analysis. The CO$_2$ (aq) concentration was calculated using a calibration curve.

Fractionation ($\varepsilon$) was calculated for each growth phase as the difference between the average $\delta^{13}$DIC and the $\delta^{13}$C of the biofilm sampled at that phase. Fractionation was calculated according to the equation:

$$\varepsilon = (\delta^{13}\text{DIC}_{\text{biofilm}} - \delta^{13}\text{C}_{\text{biofilm}}) / (1 + \delta^{13}\text{C}_{\text{biofilm}} / 1000)$$ (2)

(Freeman and Hayes, 1992). Fractionation was calculated directly for the initial phase biofilms. However, during the exponential and mature phase, fractionation was calculated from the average $\delta^{13}$DIC during each specific development stage and the $\delta^{13}$C of the newly formed biomass. The $\delta^{13}$C of the newly formed biomass was calculated according to

$$\delta^{13}C_{t+1} = \frac{b_{t+1} \delta^{13}C_{t+1} - b_t \delta^{13}C_t}{(b_{t+1} - b_t)}$$ (3)
where \( b_{t+1} \) denotes the C-biomass at the phase of sampling and \( b_t \) the C-Biomass in the previous phase.

2.2 PLFA-determination

Phospholipid-derived fatty acids (PLFA) originating from cell membranes were used as group specific biomarkers to determine the relative abundance of heterotrophic and phototrophic biomass. PLFA extraction and derivatization was done by an adapted Blyer and Digh protocol (Boschker, 2004). Analysis of the methylated forms of PLFA was done by gas chromatography-flame ionization detection (GC-FID, Interscience, Belgium) using a polar analytical column (Scientific Glass Engineering BPX-70). Biomass contribution of heterotrophic components was estimated by Chemtax as described in Dijkman et al. (2006)

2.3 pH profiling

Three adjacent slides covered with biofilm were removed from the incubator and positioned into an external flow chamber with fresh medium. The environmental conditions in the external flow chamber (flow, temperature, irradiance) were identical to the respective conditions in the biofilm incubator. Glass pH microelectrodes (Glud et al., 1992) were used to measure depth-profiles of pH in the phototrophic biofilms. The pH microelectrode and a standard calomel reference electrode (Radiometer, Denmark) were connected to a high-impedance millivoltmeter (Keithley, USA). The microsensor was mounted on a motorized micromanipulator (Unisense A/S, Denmark) and depth-profiles were automatically recorded on a PC with a data acquisition system (Profix, Unisense A/S, Denmark). The pH microsensors were calibrated in standard pH buffer solutions of pH 7 and pH 10 (Radiometer, Denmark) and exhibited almost ideal Nernstian response characteristics and a response time of <60 s. The surface position where the pH microsensor touched the biofilm surface was estimated by visual inspection with a dissection microscope while approaching the sensor tip to the biofilm surface. In order to ensure steady state conditions, the biofilm samples were left in the external flow chamber for 30 min in the light before pH measurements were initiated. Similarly, light was switched off for 30 min before the pH profiles in darkness were acquired.

2.4 Biofilm modeling

In order to explain the variation in \( \delta^{13}C \) values found in the freshwater biofilms, we constructed a model describing the chemical and biological processes in a developing freshwater biofilm. Growth and relevant metabolic and geochemical processes in the biofilm were modeled with the PHO-BIA biofilm kinetic model programmed in Aquasim 2.1 (details in Wolf et al., 2007). The model is a multi-species and multi-substrate mechanistic biofilm model, which has been developed based on the general one-dimensional mathematical biofilm model (Reichert, 1998). It contains kinetics that describes the interactions between photoautotrophic, heterotrophic and chemoheterotrophic (nitrifying) functional microbial groups. The biological processes in the model include biomass growth, biomass inactivation and lysis, substrate and nutrient conversion. Growth is estimated as maximum growth rate multiplied by a limitation term, based on the most limiting substrate at the given time points. Light is considered as an energy source and light dependent carbon fixation by phototrophs is modeled via the Eilers and Peeters relationship (Eilers and Peters, 1988), which accounts for light saturation and photoinhibition. Biofilm-specific phenomena are taken into account, such as extracellular polymeric substances (EPS) production by phototrophs as well as gradients of substrates and light in the biofilm. Acid-base equilibria, in particular carbon speciation, are explicitly accounted for, allowing for the calculation of pH profiles and profiles of the different abiotic carbon species across the biofilm based on chemical acid-base-equilibriums as well as consumption and production terms. The model distinguishes between the usage of different inorganic carbon sources by phototrophs, i.e. \( CO_2^{aq} \) and bicarbonate and combines a number of kinetic mechanisms specific to phototrophic microbial communities, such as internal polyglucose storage under dynamic light conditions, phototrophic growth in the darkness using internally stored reserves, photoadaptation and photoinhibition. We used the same model parameters as Wolf et al. (2007). The settings of the model were based on the measured DIC concentrations and biofilm photosynthesis/respiration rates within the incubator (data not shown).

2.5 Statistical considerations

Due to some lacking data points and the impossibility of replication it was impossible to use multiple component statistical analysis tests. Therefore, the results have been tested per parameter using the single factor ANOVA test. If variation of an incubation parameter (i.e. flow rate, temperature, irradiance, medium type) did not yield significant differences in the measured growth parameters, we considered it acceptable to pool data of that specific parameter for further analysis.

3 Results

Phototrophic biofilm development depended mainly on irradiance (Fig. 1). High incident irradiance resulted in a shorter lag time as well as in a faster biofilm development due to a higher growth rate. Relative growth rates at 50% light absorption, as calculated from the fitted logistic growth curve (Table 1), varied significantly with irradiance (\( P < 0.05 \)) within the growth conditions applied. We found no differences in the relative growth rates (at 50%
light absorption) between marine or freshwater biofilms, nor did we find an effect of the flow rate on the growth rate \((P > 0.05)\). The temperature seemed to have some effect on the growth rate, but within the experimental design this was not significant \((P > 0.05)\).

The \(\delta^{13}C\) values of the different biofilms were measured at three developmental stages. For the freshwater biofilms it was found that \(\delta^{13}C\) values became less negative with an increase in biomass (Figs. 2a–d), similar trends in \(\delta^{13}C\) values were also found in biofilms grown outdoors in a helophyte filter, see Fig. 1 in supplementary information – http://www.biogeosciences.net/4/613/2007/bg-4-613-2007-supplement.pdf). Changes in \(\delta^{13}C\) were strongest for biofilms grown at the highest irradiances. However, the increase was not related to the actual growth rate of the biofilm. The biofilm development was described well by a logistic growth model. A logistic growth model per definition depicts a decrease in net growth rate as soon as biomass values reach a value above 50% of the maximum biomass in the system. Considering the good fit quality of the model through the measured growth curves (average \(r^2=0.975\pm0.038 n=24\) curves, lowest \(r^2\) value was 0.820), it implies that the actual growth rate was highest during the initial and exponential phase, whereas the net growth rate decreased thereafter with biofilm thickening (see also Fig. 2 in supplementary information – http://www.biogeosciences.net/4/613/2007/bg-4-613-2007-supplement.pdf). At the point where the \(\delta^{13}C\) value increased, the average growth rate was below the maximum growth rate.

The \(\delta^{13}C\) values showed a contrasting trend during the development of the marine biofilm (Figs. 3a–d). The \(\delta^{13}C\) values during the initial phase were higher (\(\sim 9\%_c\)) than in the freshwater biofilms, but instead of increasing with the thickening of the biofilm, the values became more negative over time.

The DIC concentrations were 0.34±0.05 mM and 1.5±0.4 mM in the freshwater and marine medium, respectively. The \(\delta^{13}C\) in the media were measured for the two 30°C freshwater runs and for all marine runs. The average initial \(\delta^{13}C\) value in freshwater was \(-10.9\pm0.8\%_e\) while it was \(-6.6\pm1.2\%_e\) in salt water medium. For the incubations of which \(\delta^{13}C\) values were measured it was possible to calculate the fractionation \((\varepsilon)\) for each growth phase (Fig. 4). The fractionation was significantly lower in the marine \((\varepsilon=13.1\pm1.5\%_e, n=16)\) compared to the freshwater incubations \((\varepsilon=17.5\pm1.5\%_e, n=8)\) \((P < 0.05)\) during growth in the initial phase. No effect of irradiance on fractionation was found during this phase for both marine and freshwater incubations \((P_{\text{freshwater}}=0.267 \text{ and } P_{\text{marine}}=0.42)\). For the freshwater runs a trend of decreasing fractionation with increasing irradiances was found during the exponential growth phase. This trend was stronger during mature growth. Fractionation values did not differ significantly between the irradiances regime during growth in these phases \((P > 0.05)\). The marine incubations showed an increase in fractionation at all irradiance regimes during the exponential phase. In the mature phase this increase continued at 60 \(\mu\)mol photons m\(^{-2}\)s\(^{-1}\). An opposite trend was found for the incubations at 120 \(\mu\)mol photons m\(^{-2}\)s\(^{-1}\). At this irradiance fractionation decreased during the mature phase. Fractionation values differed significantly \((P < 0.05)\) between both irradiances in mature marine biofilms.

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**Fig. 1.** Biomass development of phototrophic biofilms grown in freshwater and marine growth mediums. (a) Typical example of biomass development at 4 different irradiances in a freshwater growth medium, incubated at 30°C and 0.5 m/s flow. (b) Typical example of biomass development at 4 different irradiances in a marine growth medium, incubated at 15°C and 0.5 m/s. (c) Average relative growth rates at 50% absorbance for the four different irradiances. One overall average growth rate per irradiance was calculated from the growth rates of all incubations. The error bars represent the standard deviation on the average value. The relative growth rates were estimated using a logistic growth model.
Fig. 2. Development of $\delta^{13}C$ values in the bulk biomass in developing freshwater phototrophic biofilms. Figures (a–d) show the relationship of $\delta^{13}C$ values with biomass for each treatment (temperature and flow rate are indicated in the graphs). Biomass is expressed as light absorption. (e) $\delta^{13}C$ value samples taken at the end of the run to overcome the effects in changes in pH in the medium. Symbols indicate different irradiances $120 \mu$mol photons $m^{-2} s^{-1}$ (closed squares), $60 \mu$mol photons $m^{-2} s^{-1}$ (open circles), $30 \mu$mol photons $m^{-2} s^{-1}$ (closed triangles) and $15 \mu$mol photons $m^{-2} s^{-1}$ (open triangles).

For one freshwater run and one marine run, additional $\delta^{13}C$ samples were taken at the end of the run. The reason for the additional sampling was that the pH in the medium changed during the incubation (Fig. 5) affecting the $CO_2(aq)/HCO_3^-$ ratio in the overlying water, which may explain the observed variations in $\delta^{13}C$. The change in pH was more pronounced at high biofilm biomass and high irradiance.

For this additional sampling, we used slides that were put in the incubator as a replacement of the slides sampled during the course of the run. Some of these slides (replacements for the mature samples) have only been for 5–8 days in the incubator and only contained little biomass (equivalent to the initial phase). The difference with the normal incubations is that these low biomass samples have been incubated, in medium at the moment that the pH change were most pronounced. Despite this difference in the pH history,
the δ¹³C values in these additional experiments showed the same trend with biomass development (Figs. 2e and 3e) as was found in the normal freshwater and marine experiments. However, the effect of light was more pronounced than observed under the normal sampling procedure. Unfortunately we were not able to collect samples from the marine 120 µmol photons m⁻² s⁻¹ for this experiment, since growth on these replacement slides was so fast that they were all above the 85–90% absorption value for longer than one week.

Besides the pH of the medium, pH depth profiles were measured in biofilms, both in the exponential and the mature growth phase (Fig. 6). Biofilm thickness varied considerably at each growth stage. This is indicative for a large spatial heterogeneity of the biofilm structure but the profiles show a consistent and representative trend of typical in situ pH characteristics for each treatment. In darkness, the pH in the freshwater biofilm matrix did not vary significantly from the pH in the overlaying water, whereas in light the pH clearly increased with depth. This can be taken as an
Table 2. Percentage of heterotrophic biomass relative to the total biomass present in the different treatments based on chemtax analysis of the PLFA determination (NA = not analyzed due to a low amounts of biomass).

| Medium     | Temperature | Flow (m/s) | Irradiance $\mu$mol photons m$^{-2}$ s$^{-1}$ |
|------------|-------------|------------|---------------------------------------------|
|            |             |            | 120  60  30  15                               |
| Freshwater | 20          | 0.5        | 7  5  15  8                                  |
|            |             | 2          | 2  8  15  NA                                  |
|            | 30          | 0.5        | 2  6  6  NA                                  |
|            |             | 2          | 5  2  2  NA                                  |
| Average Freshwater |           |            | 4  5  10  8                                |
| Saltwater  | 15          | 0.5        | 6  11  11  NA                                |
|            |             | 2          | 18  15  17  19                              |
|            | 25          | 0.5        | 15  23  19  58                              |
|            |             | 2          | 22  36  44  NA                                |
| Average marine |           |            | 15  21  23  39                              |

indication of a high CO$_2$ consumption due to photosynthesis. The increase in pH with depth was most pronounced in the mature biofilms. In addition, a higher irradiance resulted in a more pronounced increase in pH. In marine biofilms it was found that in the dark the pH in the biofilm decreased with depth. In the light an increase was found in the upper part of the biofilm, while at deeper parts the pH decreased again and/or remained stable. A decrease in pH can be caused by a net production of CO$_2$ resulting from high respiratory activity. The increase in pH in the upper part of the biofilm was linked to photosynthetic activity and was strongest at high irradiance.

PLFA analyses showed that the heterotrophic bacterial biomass was always highest in the marine runs. The average contribution of bacteria in the marine biofilms was almost twice the value of the freshwater biofilms (Table 2).

3.1 Modeled differences in CO$_2$ consumption vs. HCO$_3^-$ consumption

The model describes the pH in the biofilm based on charge balance, including the speciation of the different forms of inorganic carbon due to transport, chemical and biological conversion processes. The speciation of the different inorganic C pools was calculated with the model for a freshwater biofilm of 500 $\mu$m thickness (Fig. 7a). It was found that at an irradiance of 120 $\mu$mol photons m$^{-2}$ s$^{-1}$, most of the CO$_2$ was consumed in the upper 150 $\mu$m of the biofilm. The HCO$_3^-$ pool was much larger than the CO$_2_{(aq)}$ pool and its concentration decreased relatively less with depth. Within the model, CO$_2$ is preferred above HCO$_3^-$ as carbon source and therefore the CO$_2$ consumption rate was highest in the top layer, while in that region almost no HCO$_3^-$ was consumed (Fig. 7b). The significance of HCO$_3^-$ as C-source increased with depth and maximal HCO$_3^-$ consumption was found at 350 $\mu$m depth. Below this depth, irradiance became limiting and determined the photosynthesis rate rather than the availability of the different inorganic C-pools. With the same model, an estimate of the depth integrated consumption rate of the different inorganic carbon pools during the biomass development was calculated (Figs. 8a and b). It was found that initially CO$_2$ was the most important C-source for carbon fixation. With increasing biofilm thickness of the the relevance of CO$_2$ for photosynthesis decreased. For thick biofilms (>400 $\mu$m), HCO$_3^-$ eventually became the most important C-source for photosynthesis (Fig. 8b).

4 Discussion

We will start the discussion with the freshwater system, where we observed that $\delta^{13}$C values increased with the development of phototrophic biofilms. In the initial growth phase no clear correlation was found between $\delta^{13}$C values and the growth rate or irradiance, indicating that the biofilm growth rate did not affect isotopic fractionation rates at this stage. This is in contrast with relationships found for isotope fractionation in phytoplankton, where several studies showed that the growth rate (Fry and Wainright, 1991; Laws et al., 1995; Rau et al., 1996) and irradiance (Rost et al., 2002) had an inverse relationship with isotopic fractionation in different phytoplankton species. The rationale behind this expected inverse correlation is that at high growth rates RUBISCO becomes transport limited for CO$_2$. This would result in under-saturation of RUBISCO and therefore a lower fractionation, as well as a shift from CO$_2$ to HCO$_3^-$ as the most important C-source. Both will lead to increasing $\delta^{13}$C values.
Fig. 4. Average fractionation ($\varepsilon$) per irradiance during the three growth phases throughout the development of phototrophic biofilms. Biofilms were grown at incident photon fluxes of 120 (closed squares), 60 (open circles), 30 (closed triangles) and 15 (open triangles). The average values and error bars were calculated over successively 2 (freshwater, (a)) and 4 (marine, (b)) separate runs.

It can be assumed that irradiance determines growth during the initial stage, and that no other substrate than light is limiting at this stage. A non-linear relationship between the relative growth rate at 50% absorbance and irradiance was found (Fig. 1 and Table 1); logistic growth models describe that growth rates are highest during the initial phase. During this phase, the $\delta^{13}$C values were lowest. As soon as the availability of any substrate becomes limiting for growth, the net biofilm growth rate will decrease during further development. After this onset of limitation, we found increasing $\delta^{13}$C values and a decrease in fractionation in the freshwater biofilms indicating that the diffusive transport of CO$_2$(aq) became limiting in the biofilm during the later developmental stages.

In free-living phytoplankton, population-based results may closely reflect the average of individual cells, but in our biofilm communities we could only measure the depth integrated value. Our model showed that the different C-sources are not homogenously distributed with depth and that the depth integrated net-C fixation becomes increasingly CO$_2$(aq) limited with biofilm thickness, especially since heterotrophic biomass was lower in the freshwater biofilms. Low heterotrophic biomass cannot provide intensive carbon recycling of photosynthetic products during the light period.

Variations of $\delta^{13}$C values in biofilms may be due to changes in the C-source rather than being solely the result of limited CO$_2$(aq) availability. We found a fractionation of $\sim$13–17‰ for initial biofilm growth, and the modeled C-concentrations and volumetric C-fixation rates were relatively high, both indicating that CO$_2$(aq) was not limiting...
at that stage of biofilm formation (Fig. 7). During further thickening of the biofilm, model calculations showed that the depth integrated areal C-fixation rate increased, despite decreasing biofilm growth rates and a decrease in the average volumetric C-consumption rate. This resulted in a shift from CO$_2$ towards HCO$_3^-$ as the most important C-source for photosynthesis. This shift explains the progression towards less negative $\delta^{13}$C values observed in thick freshwater biofilms.
biofilms. In addition, the model showed that the top layer (<70 µm) was not limited by CO$_2$(aq) at an irradiance of 120 µmol photons m$^{-2}$s$^{-1}$ in a 500 µm thick biofilm.

Fig. 7. (a) Modeled distribution of the C-species: CO$_2$ (dashed line), HCO$_3^-$ (dotted line), CO$_3^-$ (dash dotted line) and total inorganic C (solid line) over depth at 120 µmol photons m$^{-2}$s$^{-1}$ in a biofilm of ∼470 µm thick. The pH is also expressed (closed circles). (b) Modeled carbon consumption rate with depth in a biofilm of ∼470 µm thickness, at an irradiance of 120 µmol photons m$^{-2}$s$^{-1}$. Total inorganic carbon consumption is separated in a consumption rate of CO$_2$(aq) (dashed line) and HCO$_3^-$ (dotted line). The water-biofilm surface at a depth of 0 µm is indicated with a line.

Fig. 8. Relationship between depth integrated C-consumption with the biomass development (biofilm thickness) based on a mechanistic model describing phototrophic biofilm growth (a) net-total inorganic C consumption (squares), CO$_2$ consumption (circles) and HCO$_3^-$ consumption (triangles). The lower graph (b) indicates the shift in contribution from CO$_2$ (closed circles) to HCO$_3^-$ (open triangles) to the total C-consumption.

We also found that the difference in $\delta^{13}$C between the initial and stationary biofilm growth phase became larger at higher irradiances (Fig. 2). The opposite trend was found for fractionation at the different irradiances (less fractionation at higher irradiances, Fig. 4). Biofilm thickness was approximately similar (light absorption determined at the moment of sampling) for all sampled irradiances at each given sampling event. Therefore we conclude that for stationary-phase biofilms (i.e. no net growth) a relationship can be expected between depth integrated photosynthesis rate and $\delta^{13}$C fractionation.

Our data showed no correlation between the $\delta^{13}$C value and flow rate, although relationships between fractionation and mass transport or flow velocity are well described for phototrophic biofilms (France, 1995a; Larned et al., 2004; France, 1995b). However, these studies may have different flow regimes. Our system was developed to have as little turbulence as possible (Zippel et al., 2007), which was different from the system described by France (1995a) and Larned.
et al. (2004). Turbulence might affect biofilm development differently. Furthermore, the measured pH profiles indicated that the diffusive boundary layers were very thin, 50–150 μm and 50–100 μm for 0.5 and 2 m/s flow respectively. This difference may not be enough to result in clear differences in δ13C values for both flow rates.

Another reason for the absence of the relationship with flow velocity may be that in the initial, or exponential growth phase of biofilms, mass transfer limitation is not an issue. Differences in diffusive boundary layer thickness, as a result of higher flow rates or turbulence levels during these early phases will not affect fractionation efficiencies for 13CO2 by RUBISCO since CO2(aq) availability is not a limiting factor for fractionation. From the same line of reasoning as was used for the growth rate (see above), it can be argued that a correlation between flow rate or turbulence and fractionation will only be found under diffusion limited conditions, i.e. late exponential and mature biofilms. So it may be that the correlation found in literature is caused by sampling preferences for mature biofilms. Developmental stages have not been taken into account in these studies. In natural systems it will be difficult to distinguish and sample from the different stages of the biofilm development, since heterogeneity in developmental stages are present on a small spatial scale, due to sloughing, grazing, etc (Biggs, 1996; Havens et al., 1996).

Initial δ13C values were less negative in the marine biofilms, while their relative growth rates did not differ from the freshwater biofilms. The difference in δ13C values can be partially explained by the δ13DIC, which was 4–5‰ less negative than in the freshwater medium, but part of the difference must be due to differences in fractionation during the initial phase. Lower fractionation can result from the lower CO2(aq) concentration in the marine medium. However, neither the initial δ13DIC, nor the lower CO2(aq) explain the decrease in δ13C values with the thickening of the biofilms. It was calculated that the initial CO2(aq) concentration in the marine medium was approximately 3–4 times lower than the freshwater medium despite the higher DIC concentrations. Therefore, it seems likely that the marine biofilms were more limited in CO2(aq) than the freshwater biofilms. However, the high DIC may select for species with a high affinity for bicarbonate uptake from the start, and thus led to a relative high uptake of bicarbonate in the initial phase. Therefore there may not such a big shift from CO2 to bicarbonate as the primary C-source. A decrease in δ13C values may also be explained by the expected decrease in growth rate with the thickening of the biofilm, as has been shown for phytoplankton species (Fry and Wainright, 1991; Laws et al., 1995; Rau et al., 1996). A prerequisite for this mechanism would be that the decrease in growth rate is caused by another factor than carbon limitation. However, the same amounts of micro-nutrients have been added to both types of media.

In our experiments, one clear difference between the freshwater and the marine biofilms was that heterotrophic biomass was twice as high in the marine biofilms. Higher heterotrophic biomass is assumed to result in a higher recycling of carbon within the biofilm and as a result increase the availability of CO2(aq). A lowering of the pH with depth in the dark, indicative for a high respiratory activity, was indeed found in the marine biofilms. A decrease in pH with depth in the dark was almost absent in the freshwater biofilms and as a result we conclude that a much higher heterotrophic activity was present within the marine biofilms. This was also confirmed from oxygen micro profile measurements (data not shown). Carbon cycling within a biofilm may reduce the δ13C value. It has been shown that as soon as ecosystems shift from phototrophic to heterotrophic systems this will lower the δ13C values (Schindler et al., 1997; Bade et al., 2004). High respiration rates increase the level of heterotrophy of the biofilm and elevate the CO2 cycling and its availability for the phototrophic organisms. Moreover, respiration enriches mostly the 12CO2 pool since its source (biomass) is lighter, relative to the DIC pool.

In conclusion, we found that the δ13C value depends on the development phase of the phototrophic biofilm, and fractionation seems to be controlled by the balance between C-demand (net depth-integrated photosynthesis rates) and mass transfer limitation controlled by the biofilm thickness in combination with the depth integrated photosynthesis rate rather than by the actual biofilm growth rate. The freshwater biofilm model shows that initially our freshwater biofilms were not limited by CO2(aq) in the light. CO2(aq) only becomes limiting with the thickening of the biofilm, which results in a increase in importance of HCO3 as inorganic carbon source for photosynthesis.

However, the direction in which the δ13C value develops during biomass accretion seems to depend on: the net depth-integrated C-fixation rate, changes in the utilization of the different inorganic C-sources and the recycling rate of C driven by heterotrophic activity. In the marine incubation, most likely a higher heterotrophic activity in the biofilm combined with a low CO2(aq) in the medium resulted in the completely opposite trend in δ13C values, when compared to the freshwater incubations. We cannot conclude whether heterotrophic recycling is always more important in marine systems than in freshwater systems, but in our system it was. This difference was found despite different, independently grown inoculums have been used to seed biomass in the different runs. Since surface-associated microalgae and cyanobacteria are amongst the most successful and efficient primary producers in benthic aquatic environments, and are considered a main source of energy for higher trophic levels in natural systems, such trends in δ13C value with biofilm developmental stages need to be taken into account if δ13C values of biofilm are used for food web studies.

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