Carbon biogeochemistry of the Betsiboka estuary (north-western Madagascar)

Olivier Harifidy Ralison\textsuperscript{a,1}, Alberto Vieira Borges\textsuperscript{b}, Frank Dehairs\textsuperscript{a}, J.J. Middelburg\textsuperscript{c}, Steven Bouillon\textsuperscript{a,c,d,*}

\textsuperscript{a} Vrije Universiteit Brussel, Department of Analytical and Environmental Chemistry, Pleinlaan 2, B-1050 Brussels, Belgium
\textsuperscript{b} Université de Liège, Chemical Oceanography Unit, Institut de Physique (BS), B-4000 Liège, Belgium
\textsuperscript{c} Netherlands Institute of Ecology, Centre for Estuarine and Marine Ecology (NIOO-KNAW), Yerseke, The Netherlands
\textsuperscript{d} Katholieke Universiteit Leuven, Department of Earth and Environmental Sciences, Kasteelpark Arenberg 20, B-3001 Leuven, Belgium

\begin{abstract}
Madagascar’s largest estuary (Betsiboka) was sampled along the salinity gradient during the dry season to document the distribution and sources of particulate and dissolved organic carbon (POC, DOC) as well as dissolved inorganic carbon (DIC). The Betsiboka was characterized by a relatively high suspended matter load, and in line with this, low DOC/POC ratios (0.4–2.5). The partial pressure of CO\textsubscript{2} (pCO\textsubscript{2}) was generally above atmospheric equilibrium (270–1530 ppm), but relatively low in comparison to other tropical and subtropical estuaries, resulting in low average CO\textsubscript{2} emission to the atmosphere (9.1 ± 14.2 mmol m\textsuperscript{2} d\textsuperscript{−1}). Despite the fact that C\textsubscript{4} vegetation is reported to cover >80% of the catchment area, stable isotope data on DOC and POC suggest that C\textsubscript{4} derived material comprises only 30% of both pools in the freshwater zone, increasing to 60–70% and 50–60%, respectively, in the oligohaline zone due to additional lateral inputs. Sediments from intertidal mangroves in the estuary showed low organic carbon concentrations (<1%) and \delta\textsuperscript{13}C values (average −19.8‰) consistent with important inputs of riverine imported C\textsubscript{4} material. This contribution was reflected in \delta\textsuperscript{13}C signatures of bacterial phospholipid derived fatty acids (i + a\textsubscript{15:0}), suggesting the potential importance of terrestrial organic matter sources for mineralization and secondary production in coastal ecosystems.
\end{abstract}

\section{1. Introduction}
Rivers and estuaries are highly active areas in terms of organic matter and nutrient processing, where riverine transported organic matter can undergo significant biogeochemical modifications \citep[e.g.,][]{Aufdenkampe2007, Cole2007}. Despite the high riverine organic matter flux in the tropics \citep{Ludwig1996a, Ludwig1996b, Schlunz2000}, data on the biogeochemistry and nutrient cycling in tropical estuaries are still scarce in comparison to the temperate zone, and to the best of our knowledge, no biogeochemical studies on Madagascan estuaries exist. Recent data from east African estuaries \citep{Bouillon2007a, Bouillon2007b} suggest that fluxes of particulate organic carbon (POC) could be much higher than predicted by empirical models of riverine carbon transport for this zone \citep{Ludwig1996b}, and that inorganic carbon export might be higher than POC export, again in contrast to model predictions. While a number of datasets are available on the origin of POC in tropical rivers and estuaries, much less is known on the composition and origin of dissolved organic carbon (DOC) and its relation to POC. Particulate and dissolved pools of organic matter are often...
compositionaly distinct, and experimental studies indicate that preferential sorption of nitrogen rich compounds are at least partially responsible for such differences (Auffdenkampe et al., 2001 and references therein). Moreover, during riverine transport and retention in estuaries, partitioning of organic matter between dissolved and particulate organic matter pools may take place, with total suspended matter loads playing a critical factor in the overall partitioning (Middelburg and Herman, 2007).

Many of the larger east African river systems drain catchment areas where the vegetation consists of plants using C3 and C4 photosynthetic pathways, with a variable relative contribution in the overall land cover. In the more arid regions, C3 vegetation is often concentrated along rivers as floodplain or fringing forests. These systems therefore offer an ideal opportunity to study the effects of landscape scale vegetation patterns on riverine carbon pools and possible differential biogeochemical processing of C3 and C4 derived organic matter. Earlier studies suggest that the composition of the dominant vegetation is indeed a major determinant of riverine organic matter composition (Bianchi et al., 1994; Martinelli et al., 1999; Coynel et al., 2005), and that recent land use changes (i.e., over the last few decades) can be rapidly reflected in riverine organic matter composition (Bernardes et al., 2004). Recent data from tropical estuaries with significant C4 vegetation in the catchment areas further suggest that dissolved and particulate organic carbon pools can have variable relative contributions of different organic matter sources, and that C3 and C4 derived organic matter undergo different degrees of processing in the estuarine zone (e.g. Bianchi et al., 2004; Bouillon et al., 2007a). However, data from a range of estuaries are needed in order to generalize whether the reactivity of C3 and C4 derived organic matter is consistently different across different estuaries, and which implications this might have for the characteristics of the organic matter transported to the coastal zone in such systems. Mineralization of organic matter and its subsequent efflux as CO2 to the atmosphere has recently been shown to be an important link in coastal carbon budgets, and represents a significant but previously underestimated direct bypass of CO2 to the atmosphere in the global carbon budget (Borges, 2005). Given the lack of such data from tropical estuaries, field data from a variety of tropical systems are needed to better constrain fluxes and cycling of carbon in these systems.

In this study, we report data on the basic physico-chemical properties, organic and inorganic carbon pools and their stable isotope composition along the salinity gradient of the Betsiboka estuary (Madagascar), in order to present a first description of the biogeochemical characteristics of Malagasy estuaries. The Betsiboka is the main river of Madagascar in terms of length, basin area, as well as discharge (Berthois and Crosnier, 1965). Although there is ongoing discussion on the extent of original forest cover in the highlands of Madagascar (see e.g. Klein, 2002; Vagen et al., 2007 and references therein), it is clear that intensive deforestation has taken place in the catchment area, and that this has resulted in a very high erosion rate. During the rainy season, red lateritic soil is eroded and transported from the highlands (“Hauts plateaux”) by tributaries towards the estuary, and finally ends up in the Mozambique Channel. The Betsiboka estuary has therefore evolved to a high sedimentation zone and is characterised by one of the world’s fastest coastal dynamics (Lebigre, 1990). Drainage basins with steep elevation gradients and high erosion rates can be disproportionately important in sediment and organic carbon delivery to the ocean, i.e., provide higher inputs of sediment and organic carbon per unit of surface area than other types of drainage basins, as has been shown for islands in the southwestern Pacific Ocean (Milliman, 1995; Lyons et al., 2002).

2. Materials and methods

2.1. Study area

The Betsiboka (Fig. 1) is the main river system in Madagascar, draining an area of ~49,000 km² in the north-western part of the island (Randrianarjaona, 1983; Paulian, 1984). The Betsiboka River has its source in the Angavo cliffs, north of Antananarivo at an altitude of 1755 m. Major tributaries are Mahajamba, Isandrano and Ikopa rivers. Vanden Bossche and Bernacsek (1990) estimated its total length at about 525 km, and the discharge rate is reported to be in the order of 271 m³ s⁻¹ (i.e., 35 km³ y⁻¹, Vorosmarty et al., 1998). The Betsiboka discharges in the Mozambican channel near the town of Mahajanga, in the Boina region. The external part of its mouth called Bombetoka Bay is situated at 15°5′ S and 46°18′ E. The large size of the drainage basin associated with huge discharge (Berthois and Crosnier, 1965) causes important settling of sediment within the estuary, ranging from 15 × 10⁵ to 50 × 10⁴ t y⁻¹ (Randrianarjaona, 1983; Paulian, 1984; Ritter, 1986). The destruction of the original forests during the last 50 years intensified erosion, with erosion rates in the highlands in the order of 25–250 t ha⁻¹ y⁻¹ (Randrianarjaona, 1983). Furthermore, various anthropogenic factors are likely to influence the ecology and biogeochemistry of the estuary: new developments have been constructed behind the mangrove forest which faces the Bombetoka Bay. Shrimp aquaculture ponds have been implanted in the area since 1999 and at the time of sampling covered an area of ~200 ha. Mangroves are an important component of the estuary and cover approximately 46,000 ha. The climate is characterised by the alternation of a warm and wet season from November to April, and a well-marked dry and cool season from May to October. The sampling was conducted at the end of the dry season, when the Betsiboka was at its lowest level of discharge. The tidal regime is semi-diurnal, with an average tidal amplitude of 3.5 m.

2.2. Sampling and analytical techniques

Water samples were collected along the salinity gradient in the main estuary, and in a few bordering tidal mangrove creeks (Fig. 1) in October 2005 (dry season). Surface water for field measurements of dissolved O₂, pH, temperature and salinity were taken with a Niskin bottle ~0.5 m below the surface. Oxygen saturation level (Ο₂%) was measured immediately after collection with a polarographic electrode (WTW Oxi-340) calibrated on saturated air, with an accu-
Accuracy of ±1%. pH was measured using a Ross type combination electrode (ORION) calibrated on the NBS (US National Bureau of Standards) scale, as described by Frankignoulle and Borges (2001), with a reproducibility of ±0.01 pH units. Wind speed was measured in the field with a hand-held anemometer. Samples for total suspended matter (TSM) were taken by collecting a known volume of surface water (300–1000 ml), which was passed through pre-combusted (12 h at 450 °C) and pre-weighed 47 mm diameter glass fibre filters (Whatman GF/F). Subsequently, the filters were dried at 60 °C for 12 h prior to packing in Petri dishes.

Samples for the analysis of carbon stable isotope ratios of dissolved inorganic carbon (δ13CDIC) were taken from the same Niskin bottle by gently overfilling 12 ml glass headspace vials, poisoned by addition of 20 μl of a saturated HgCl2 solution, and closing tightly with a septum lined screw cap. δ13CDIC was measured as described in Gillikin and Bouillon (2007). Samples for the determination of total alkalinity (TA) were obtained by pre-filtering 100 ml of water through pre-combusted Whatman GF/F filters, subsequent filtration through 0.2 μm cellulose acetate syringe filters, and stored in HDPE bottles until analysis. TA was determined by automated titration with 0.01 M HCl. The pCO2 and DIC concentrations were computed from pH and TA measurements using the thermodynamic constants described in Frankignoulle and Borges (2001), with an accuracy of computed DIC and pCO2 values estimated at ±5 μmol kg⁻¹ and better than ±2%, respectively. The exchange flux of CO2 (F, mmol m⁻² s⁻¹ and converted to units of mmol m⁻² d⁻¹) across the water–air interface was calculated according to $F = k_a \Delta pCO_2$ where $k$ is the gas transfer velocity (m s⁻¹), $a$ the solubility coefficient for CO2 (mol m⁻³ atm⁻¹), and $\Delta pCO_2$ (atm) represents the difference in pCO2 between water and air (see Liss and Slater, 1974). For consistency with previous studies in tropical estuaries and mangrove creeks, we used the $k$-wind parameterization proposed by Raymond and Cole (2001) for estuarine environments.

Samples for DOC and δ13CDOC were prepared similar to those for TA, but poisoned with 50 μl of H3PO4, and stored in glass vials (40 ml) with a Teflon lined septum and screw cap. DOC concentrations and δ13C signatures were determined using a modified Thermo HiperTOC interfaced to a Thermo Delta + XL IRMS (Bouillon et al., 2006). Although (pre)filtration on glass fibre filters may result in a certain degree of sorption of DOC to the filter, this effect is likely to be minimal (Kaplan, 1994) and glass fibre filters are therefore commonly used for sample filtration or pre-filtration for DOC analysis (e.g., Abril et al., 2002; Sharp et al., 2002). Samples for determination of Chlorophyll a (hereafter Chl a) were obtained by filtering a known volume of surface water on pre-combusted 47 mm GF/F filters. These were stored in a
screw cap vial, placed in a cool box with ice, and stored frozen until later analysis. Pigments were extracted for approximately 12 h in 15 ml of 90% acetone at 4 °C and analyzed with a Turner TD-700 Fluorimeter. The accuracy of Chl a analysis was estimated to be ± 4%. Samples for determination of POC and δ13CPOC were filtered on 25 mm GF/F filters (pre-combusted at 450 °C for 4 h), dried at 60 °C overnight, and stored in Petri slides. These were later decarbonated under HCl fumes, re-dried and packed in Ag cups. δ15N values of particulate nitrogen (PN) were determined on subsamples of the 47 mm GF/F filters used for TSM determination, wrapped in Ag cups, and analyzed on the abovementioned EA-IRMS setup. It should be noted that due to either sampling errors or analytical problems, some stations have missing data for some parameters, so that the number of data reported in the figures does not always correspond to the number of sampling stations indicated on Fig. 1.

Intertidal mangrove sediments were sampled in different stations in the estuarine zone, for determination of %OC, and δ13C signatures of bulk sediment organic matter and bacteria specific PLFA (phospholipid fatty acids). Sediment cores were collected with a 3.8 cm diameter Plexiglas corer, and sliced into 0–1, 1–2, 2–4 and 4–10 cm sections. These were stored frozen, transported back to the home lab on dry ice, and freeze dried. Samples for %C and δ13C of bulk sediment organic carbon were acidified with 1 M HCl in silver cups to remove carbonates, and analyzed on a Thermo Flash EA1112 – Delta + XL setup. Extraction, derivatization, and δ13C analysis of bacterial PLFA was performed as described in Bouillon and Boschker (2006).

3. Results

The salinity along the Betsiboka main estuary ranged between 0 and 32.4, while the tidal mangrove creeks had brackish salinities between 19 and 26. TSM concentrations generally ranged between 30 and 100 mg l⁻¹ in the estuary, with two higher values recorded at salinities 14.5 and 33.1 (148 and 286 mg l⁻¹, respectively). TSM concentrations in the tidal creeks were highly erratic, ranging between 60 and 849 mg l⁻¹ (data not shown).

POC concentrations generally ranged between 0.5 and 2.0 mg C l⁻¹ in the main estuary, with some values ~3 mg C l⁻¹ in the meso- and polyhaline zone (Fig. 2A). In the tidal creeks, POC values were distinctly higher and ranged between 2.3 and 6.1 mg C l⁻¹ (Fig. 2A). Concentrations of DOC were in the order of 0.5 mg l⁻¹ in the freshwater and marine end members, but showed a clear non-conservative pattern with highest values in the mesohaline part of the estuary (up to ~1.5 mg C l⁻¹), and also relatively high DOC levels in the tidal creeks (0.9–1.4 mg C l⁻¹, Fig. 2B). Stable isotope signatures of DOC and POC ranged between −23.7‰ and −17.7‰, and between −23.6‰ and −16.9‰, respectively (Fig. 2C and D). In the POC pool, the estuarine gradient in δ13C was somewhat erratic, with no clear pattern observed, and similar values in the tidal mangrove creeks (Fig. 2C). For DOC, however, there is a clear pattern of more 13C enriched values in the mesohaline part of the estuary (Fig. 2D) coinciding with the net inputs of DOC in this zone (Fig. 2B), and relatively 13C depleted DOC in the tidal mangrove creeks (Fig. 2D). DOC/POC ratios ranged between 0.26 and 2.45 in the estuary, and were −0.4 in the tidal creeks (only 2 data points for the latter). Lowest values were associated

Fig. 2. Distribution and carbon isotope composition of organic carbon pools along the Betsiboka estuary (black diamonds) and tidal mangrove creeks (open diamonds): (A) POC, (B) DOC, (C) δ13CPOC, and (D) δ13CDOC.
with higher levels of TSM, and the DOC/POC ratios observed in the Betsiboka are in the lower range of values reported for estuarine and coastal systems in the Western Indian Ocean region (Fig. 3).

Chl a concentrations were generally low in the estuary proper (0.9–7.2 µg l⁻¹), but distinctly higher in some of the tidal creeks where values between 15 and 25 µg l⁻¹ were recorded (Fig. 4). POC/Chl a ratios ranged overall between 172 and 836, and were generally high in both the estuary proper (388 ± 194, n = 17) and the tidal creeks (587 ± 276, n = 5), indicating that the POC pool is dominated by detrital organic matter.

In the freshwater part of the estuary, δ¹⁵N of PN was relatively high (9.1‰), and increased further in the mesohaline part of the estuary, reaching values as high as 19.8‰ (Fig. 5). At salinity >12, however, δ¹⁵N values dropped markedly and ranged between –3.6‰ and 5.8‰, with data from the tidal creeks falling within this range (Fig. 5).

Along the salinity gradient, little variation in %O₂ levels was observed, with a slight oversaturation on average (107 ± 8%, Fig. 6A). The tidal creeks, in contrast, were generally undersaturated (74–94%), with one unusually high %O₂ level of 238‰, see above). pCO₂ values were generally between 270 and 725 ppm, with the exception of one exceptionally low value (117 ppm) in one of the creeks (note that this data point corresponds to the unusually high %O₂ level of 238‰, see above).

The distribution of DIC (Fig. 6C) shows a nearly conservative pattern along the estuarine gradient (with one high value at salinity 14.5), with low DIC concentrations in the freshwater end (5.37 mmol kg⁻¹). For δ¹³C DIC, the conservative mixing scenario was calculated according to (Mook and Tan, 1991):

\[ \delta^{13}C = \frac{Sal(DIC_{Sal}^{13}C_0 - DIC_{Sal}^{13}C_M) + SalDICf^{13}CM - SalDICf^{13}C_F}{Sal(DICf - DIC_M) + SalDICf_M - SalDICf_F} \]  

where Sal is the sample salinity; DICf, δ¹³Cf are the DIC concentration and stable isotope composition at the freshwater or least saline end-member; DICM, δ¹³CM are the DIC concentration and stable isotope composition at the marine end-member.

The corresponding δ¹³C DIC gradient deviates only slightly from that expected for conservative mixing (Fig. 6D), ranging between ~5.4‰ in the freshwater part to ~0.4‰ at salinities >30. Similar to what was observed in the pCO₂ and DIC data, the samples collected at salinity 14.5 deviated from this pattern, with a much more negative δ¹³C DIC signature of ~6.9‰ (Fig. 6D). In the tidal mangrove creeks, DIC is generally similar to or higher than that observed in the estuary for similar salinities (Fig. 6C), with δ¹³C DIC values consistently lower than at similar salinities in the estuary (~12.8‰ to ~1.8‰, Fig. 6D).

Sediments in the intertidal mangrove forests along the estuary and creeks showed relatively low organic carbon contents (0.5–1.1%) and TOC/TN ratios (8.4–11.7) (Table 1). δ¹³C signatures of bulk organic carbon (~19.8 ± 1.6‰, Table 1) were much enriched relative to the dominant mangrove vegetation (with typical values of ~30‰ to ~26‰), see Fig. 7. All sites showed highest %OC in the top layers, most pronounced in site 3 where %OC decreased from 1.10 in the 0–1 cm section to 0.58 in the 4–10 cm section, accompanied by an increase in δ¹³C from ~23.9‰ to ~19.7‰. δ¹³C signatures of the bacterial PLFA i + a15:0 (~24.0 ± 1.6‰, see Table 1 for data per site) were distinctly enriched relative to what would be expected for a dominant reliance on mangrove carbon, assuming a fractionation of ~3.7 ± 2.1‰ between the substrate and i + a15:0, and a global average δ¹³C signature for mangrove litter at ~28.2 ± 2.1‰ (see Bouillon and Boschker, 2006).

δ¹³C signatures of the expected C substrates correlated well with the bulk sediment organic carbon δ¹³C values (Fig. 8).
4. Discussion

The Betsiboka catchment experiences some of the highest reported erosion rates (Randrianarijaona, 1983). In line with this, TSM concentrations were generally high when compared to data from other estuaries and tidal creeks draining into the western Indian Ocean (Fig. 3), although it must be stressed that our sampling campaign was held during the dry season, and that it can be expected that TSM levels could even be much higher during periods of high water flow, when tropical rivers generally exhibit the highest particulate matter load due to runoff and high erosion rates (e.g., Coynel et al., 2005; Hung and Huang, 2005). As has been observed on a wider scale, high TSM levels are generally associated with a relatively low contribution of DOC to the total water column OC pool, i.e., with relatively low DOC/POC ratios (e.g., Ittekkot and Laane, 1991; Abril et al., 2002). This trend is confirmed when comparing the data gathered here with data from estuaries and tidal creeks from the western Indian Ocean region (Fig. 3), and is thought to result from dynamic exchange and repartitioning reactions (see Middelburg and Herman, 2007). DOC/POC ratios for the freshwater end are in line with those predicted by Ludwig et al. (1996b) for rivers opening in the Indian Ocean between 12° and 16°S, based on an empirical model. However, while the latter model predicts a DIC:TOC ratio (where TOC is the sum of POC and DOC) of ~0.34 for this region, our data for the freshwater part indicate a DIC:TOC ratio of 5.0, i.e., almost 15 times higher. Although such a discrepancy may be due to local factors, it should be noted that other field studies in tropical east African estuaries found a similar difference between observed and predicted DIC:TOC ratios (Bouillon et al., 2007a,b).

Table 1
Organic carbon (OC) and total nitrogen (TN) concentrations, elemental ratios (TOC/TN, atom), and δ13C signatures of organic carbon and bacteria specific PLFA i + a15:0 in intertidal mangrove sediments in the Betsiboka estuary

<table>
<thead>
<tr>
<th>Site</th>
<th>%OC</th>
<th>%N</th>
<th>TOC/TN</th>
<th>δ13C-TOC</th>
<th>δ13C i + a15:0</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>0.86 ± 0.03</td>
<td>0.09 ± 0.00</td>
<td>9.3 ± 0.5</td>
<td>-19.5 ± 0.7</td>
<td>-24.4 ± 1.3</td>
</tr>
<tr>
<td>2</td>
<td>0.90 ± 0.08</td>
<td>0.10 ± 0.01</td>
<td>9.1 ± 0.5</td>
<td>-19.1 ± 0.7</td>
<td>-23.5 ± 0.6</td>
</tr>
<tr>
<td>3</td>
<td>0.83 ± 0.34</td>
<td>0.07 ± 0.03</td>
<td>11.0 ± 0.7</td>
<td>-21.4 ± 2.4</td>
<td>-26.1 ± 1.8</td>
</tr>
<tr>
<td>4</td>
<td>0.84 ± 0.06</td>
<td>0.08 ± 0.01</td>
<td>10.1 ± 0.8</td>
<td>-18.8 ± 0.2</td>
<td>-23.4 ± 0.3</td>
</tr>
</tbody>
</table>

Values are averages ±1 s.d. of 4 different depth sections (0–1, 1–2, 2–4, and 4–10 cm), except for site 4 where data for the 0–1 cm section are missing.

Fig. 5. Distribution of nitrogen stable isotope signatures of particulate N along the Betsiboka estuary (black diamonds) and tidal mangrove creeks (open diamonds).

Fig. 6. Profiles of (A) oxygen saturation levels, (B) pCO2, (C) DIC concentrations, and (D) δ13C-DIC along the Betsiboka estuary (black diamonds) and tidal mangrove creeks (open diamonds). Line on panel (D) represents the conservative mixing scenario (see text for details).
The majority of rivers and estuaries act as net sources of CO2 to the atmosphere (Cole and Caraco, 2001; Borges, 2005; Duarte and Prairie, 2005; Cole et al., 2007). Although the number of data on tropical estuaries is still extremely limited (Borges et al., 2005), some estuaries along the western Indian Ocean were found to have high pCO2, consistent undersaturation in O2, and high resulting CO2 fluxes to the atmosphere (e.g., Bouillon et al., 2007ab). Similarly, in the freshwater part of tropical rivers consistent CO2 oversaturation and CO2 effluxes have been reported (e.g., see data in Cole and Caraco, 2001; Paquay et al., 2007), and water-atmosphere CO2 fluxes in the Amazon basin have been suggested to represent a quantitatively important component in the overall carbon budget of the basin (Richey et al., 2002).

Our pCO2 data on the Betsiboka, however, do not show very high levels of oversaturation, and are even undersaturated in a few instances (Fig. 6B). This results in an estimated average CO2 efflux (9.1 ± 14.2 mmol m^-2 d^-1, Fig. 9) which is markedly lower than that observed in many other tropical estuaries so far (Sarma et al., 2001; Bouillon et al., 2007a; Zhai et al., 2007). However, a similar range of pCO2 and low overall CO2 exchange was observed earlier in the subtropical Godavari estuary during a dry season sampling campaign (Bouillon et al., 2003), and it is not unlikely that in both cases, CO2 dynamics might be markedly different during high water periods.

The DIC profile shows near conservative behavior along the estuarine gradient, with the exception of a single station at salinity 14.5 (Fig. 6C). The latter also coincided with a distinctly higher pCO2 (Fig. 6B) and lower δ13C_DIC (Fig. 6D) indicating that this was not due to a measurement artifact. We speculate that the particularly low water column depth at this station (<0.5 m) could have resulted in disturbance of the sediment–water interface during sampling, and thus may have been influenced by sediment porewater or groundwater inputs enriched in DIC and pCO2 and depleted in δ13C_DIC due to respiration. The generally conservative DIC profile is consistent with the overall low pCO2, since mineralization and/or lateral inputs of high DIC water would be expected to lead to a substantial increase in DIC levels along the estuarine gradient (e.g., Bouillon et al., 2007b). The pattern in δ13C_DIC, however, does indicate that the apparently conservative behavior of DIC is not absolute, since δ13C_DIC is slightly lower (by ~1% at most) than expected for conservative mixing of seawater and freshwater end members, in particular in the mesohaline zone (Fig. 6D). The fact that this excursion is towards lower δ13C signatures indicates that this is due to inputs from mineralization, since other potentially influencing factors (outgassing, primary production, and carbonate dissolution) would all be expected to result in more positive δ13C_DIC signatures.

4.1. Sources of organic matter in the Betsiboka estuary

Based on the data on the global distribution of C3 and C4 vegetation presented in Still et al. (2003), the relative cover of C4 vegetation in the Betsiboka catchment area was estimated to be ~80–90%. This is consistent with the more detailed vegetation maps presented in Rasambainarivo and Ranaivoarivelo (2003) which show a predominance of non-wooded savannah, and with the dominant grass species mentioned all following the C4 photosynthetic
pathway. We had therefore expected that C4-derived organic carbon would dominate the riverine DOC and POC pools, but our $\delta^{13}C$ data do not confirm this hypothesis. Using end-member values of $-12.8\%$ and $-28.2\%$ for C4 and C3 derived carbon, respectively (see Bouillon et al., 2007a), the contribution of C4 vegetation to the POC and DOC pool in the freshwater end of the estuary can be estimated at $\sim35\%$ and $45\%$, respectively. Note that these estimates assume that algal inputs (e.g., from within-stream phytoplankton production) can be neglected, which would appear valid considering the relatively low Chl $a$ levels and high POC/Chl $a$ ratios observed (Fig. 4A and B). This much lower input of C4 derived carbon than expected based on vegetation cover suggests that there is differential mobilization of carbon from C4 vegetated areas and forested areas. Although it has been established that vegetation cover has a major influence on the composition of riverine organic carbon pools, Martinelli et al. (1999) similarly found that the $\delta^{13}C$ signatures of various size classes of POC in the Piracicaba Basin (Brazil) was more negative than would be expected based on the relative C4 cover in the basin. The latter authors found the highest C4 contribution in the fine POC fraction during periods of high water, and suggested that this was caused by high surface runoff from sugarcane and pasture fields during this period. Additionally, Coynel et al. (2005) found that forested sub-basins of the Congo River showed approximately threefold higher area specific fluxes of organic carbon than in savannah dominated basins. Such a mechanism would be consistent with the data of Martinelli et al. (1999) and those presented here, and imply that the relative contribution of C4 inputs to riverine organic carbon pools is consistently less than its relative vegetation cover.

An alternative mechanism to explain the low contribution of C4 material to the aquatic organic carbon pool is differential lability of C3 and C4 derived organic matter, i.e., C4 derived organic matter could be preferentially mineralized in the aquatic environment. This was suggested by Mayorga et al. (2005) who found that the $\delta^{13}C$ signature of respiratory CO$_2$ in rivers in the Amazon basin was considerably enriched in $^{13}C$ relative to dissolved and particulate organic carbon fractions. For the Mississippi River, in contrast, Bianchi et al. (2004) suggested a stronger association of C4 derived carbon with fine mineral fractions, which would lead to selective preservation of C4 derived material. Our data, however, show that at all salinities $<18$, the POC pool was $^{13}C$ depleted relative to DOC (by on average $\sim1.7\%$, see Fig. 2C and D), indicating a more important association of C3 derived OC to the particulate pool in the river and upper estuary. Thus, both differences in landscape-level mobilization of organic carbon and differences in lability could contribute to explain the observed contribution of C3 and C4 derived organic matter to the riverine organic carbon load.

Differences in $\delta^{13}C$ signatures of POC and DOC were minor on average, but showed a large variability (Fig. 2C, D, and Fig. 10). At salinity levels $<18$, $\delta^{13}C_{DOC}$ values were consistently higher than $\delta^{13}C_{POC}$, by up to 4.0$\%$; whereas at higher salinity and in the mangrove creeks, DOC showed $\delta^{13}C$ signatures consistently $^{13}C$ depleted relative to POC (by up to 4.4$\%$). This confirms recent observations from other (sub)tropical estuarine systems (e.g., see Fig. 10) that dissolved and particulate organic carbon pools may differ significantly in the relative contribution of different organic matter inputs. Very few studies have addressed the mechanisms causing such a different composition of DOC and POC and the changes that these undergo during their transit in the aquatic system, but the advent of more accessible techniques for stable isotope analyses on DOC will no doubt be an important step in generating the data needed to develop further understanding.

In the mesohaline part of the estuary, there were clear net inputs of POC and DOC (Fig. 2A and B), and this coincided with a marked increase in $\delta^{13}C$ values of both pools (Fig. 2C and D), indicating inputs of $^{13}C$ enriched organic matter. In view of the presence of a number of small tributaries draining predominantly grasslands, this pattern suggests important inputs of C4 derived organic matter in this part of the estuary. This salinity range also shows higher DOC/POC ratios (1.2–2.5) than in the freshwater part (0.5–0.9), indicating that these lateral inputs provide relatively more DOC than POC. Hence, due to these lateral inputs, the contribution of C4 material to POC and DOC pools was found to be highest in the salinity range 5–12, where it reaches local maxima of $\sim55\%$ and $75\%$, respectively (Fig. 2C and D).

The observed $\delta^{15}N_{PON}$ profile shows a surprisingly strong gradient in comparison to other tropical estuaries (e.g., Cifuentes et al., 1996; Jenererjahn et al., 2004), with high values of up to $+19.8\%$ in the mesohaline part of the estuary (Fig. 5). Such mid-estuarine maxima have been reported earlier for a number of temperate estuaries with relatively long residence times and typically with high NH$_4$ levels in their freshwater inflow (see Middelburg and Herman, 2007 and references therein). Such a pattern is thought to be the result of intense N processing (in particular, nitrification), which leads to a $^{15}N$ enriched DIN pool due to preferential removal of ‘light’ DIN species and hence, higher $\delta^{15}N$ values after incorporation of this enriched DIN by microbial communities (e.g., Caraco et al., 1998; de Brabandere et al., 2002). Whether this
mechanism is also the cause of the observed $\delta^{15}$N$_{PAP}$ profile in the Betsiboka, however, cannot be resolved without either detailed DIN profiles and/or specific N process rate measurements.

Although our sampling scheme focussed on the main estuary, the limited number of data collected in tidal mangrove creeks point out that they can differ markedly in their biogeochemical signatures when compared to the adjacent estuarine waters. First, although for some of the parameters only a limited amount of data are available, the tidal creeks showed higher POC and DOC concentrations (Fig. 2A and B), with slightly lower $\delta^{13}$C signatures (Fig. 2C and D), and most of the creeks also had significantly higher Chl a concentrations (Fig. 4A). Secondly, most of the mangrove creeks showed distinctly lower $%\text{O}_2$ levels, higher pCO$_2$ and DIC and lower $\delta^{13}$C$_{DIC}$ signatures. Such differences have been observed previously in other mangrove systems (e.g. Bouillon et al., 2003, 2007a) and are a result of more intense mineralization, either within the aquatic system or by the interaction with the intertidal area, where benthic mineralization occurring over large areas can result in a typical signature of low O$_2$, high pCO$_2$, and DIC, and lower $\delta^{13}$C$_{DIC}$ signatures. Such $\delta^{13}$C signatures in the water column (Fig. 2), the exclusion of algal sources is unlikely to significantly bias these estimates. The $\delta^{13}$C data on bacterial PLFA clearly follow the variations in bulk sediment $\delta^{13}$C (Fig. 8), and when correcting for fractionation effects using an empirically determined average offset of $-3.7 \pm 2.1\%e$ (Bouillon and Boschker, 2006), suggest that bacterial communities show little or no selectivity: $\delta^{13}$C signatures estimated for bacteria are only slightly more negative than those found in bulk TOC (Figs. 7 and 8), but given the uncertainty on the fractionation factor used, this falls within the range that can be expected when no selectivity would occur. This implies that riverine suspended matter, with a significant pool of C4 derived organic carbon, is an important source of organic carbon to these intertidal forests. The contribution of C4 derived carbon to benthic mineralization and bacterial secondary production in these systems is consistent with our earlier observations in a northern Kenyan delta with C4 inputs (Bouillon et al., 2007a). In the latter study, porewater DOC was found to be predominantly of mangrove (i.e., C3) origin, but unfortunately, comparative data from our Betsiboka estuary sediments are not available.

In summary, this first report on the biogeochemistry of a Malagasy estuary demonstrated that the organic carbon load consists of a mixture of C3 and C4 derived organic matter, with a markedly smaller contribution of C4 derived carbon than would be expected based on the relative vegetation cover in the catchment area. Nevertheless, sedimentation of this C4 derived organic matter in the intertidal mangrove areas in the estuary resulted in a significant contribution of terrestrial C4 carbon to bacterial mineralization in these sediments. At least during the dry season, the Betsiboka estuary was a relatively minor source of CO$_2$ to the atmosphere.

**Acknowledgements**

This work was supported by the Research Foundation – Flanders (FWO-Vlaanderen, contracts G.0632.06, G.0395.07, and 1.5.070.05). S.B. was supported by a post-doctoral mandate by the FWO-Vlaanderen, and A.V.B. is a research associate at the FNRS (Fonds de la Recherche Scientifique). We thank D.P. Gillikin for useful suggestions on a draft of this manuscript, and E. Krull and an anonymous referee who provided constructive comments to an earlier version of this paper. This is publication 4246 of the Netherlands Institute of Ecology (NIOO-KNAW).

**References**


Borges, A.V., 2005. Do we have enough pieces of the jigsaw to integrate CO$_2$ fluxes in the Coastal Ocean? Estuaries 28, 3–27.

