Dynamics of dissolved inorganic carbon and aquatic metabolism in the Tana River basin, Kenya

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Abstract. A basin-wide study was conducted in the Tana River basin (Kenya) in February 2008 (dry season), September–November 2009 (wet season) and June–July 2010 (end of the wet season) to assess the dynamics and sources of dissolved inorganic carbon (DIC) as well as to quantify CO₂ fluxes, community respiration (R), and primary production (P). Samples were collected along the altitudinal gradient (from 3600 to 8 m) in several headwater streams, reservoirs (Kamburu and Masinga), and the Tana River mainstream. DIC concentrations ranged from 0.2 to 4.8 mmol L⁻¹, with exceptionally high values (3.5 ± 1.6 mmol L⁻¹) in Nyambene Hills tributaries. The wide range of δ¹³CₐDIC values (−15.0 to −2.4 ‰) indicate variable sources of DIC, with headwater streams recording more positive signatures compared to the Tana River mainstream. With only a few exceptions, the entire riverine network was supersaturated in CO₂, implying the system is a net source of CO₂ to the atmosphere. pCO₂ values were generally higher in the lower Tana River mainstream compared to headwater tributaries, opposite to the pattern typically observed in other river networks. This was attributed to high suspended sediment in the Tana River mainstream fuelling in-stream community respiration and net heterotrophy. Diurnal fluctuations of biogeochemical processes were examined at three different sites along the river continuum (headwater, reservoir and mainstream), and were found to be substantial only in the headwater stream, moderate in the reservoir and not detectable in the Tana River mainstream. The pronounced diurnal fluctuations observed in the headwater stream were largely regulated by periphyton as deduced from the low chlorophyll a in the water column.

1 Introduction

Rivers are becoming increasingly more recognised as important not only in the transport of carbon (C) between the terrestrial and marine environments but also in terms of C
storage and processing. River systems are typically a source of CO$_2$ to the atmosphere, partly due to receiving CO$_2$-rich groundwater and partly due to heterotrophic processes within the system. Thus, dissolved inorganic carbon (DIC, comprising HCO$_3^-$, CO$_3^{2-}$ and CO$_2$) is an important component in river biogeochemistry. The dynamics of riverine DIC are primarily controlled by watershed inputs and in-stream processes, which lead to either addition or removal of carbon from the DIC pool and which also change the $\delta^{13}$C$_{DIC}$ values of riverine water (Fig. 1). The input of terrestrial organic matter (and the relative contribution of C3 over C4 plants to this), the weathering of carbonate/silicate minerals within the bedrock and the ratio of groundwater discharge to surface runoff are the main watershed processes controlling the dynamics of riverine DIC and its $\delta^{13}$C signature. Additional in-stream processes impacting DIC dynamics include CO$_2$ exchange with the atmosphere, in-stream dissolution or precipitation of carbonate/silicate minerals, primary production ($P$) and community respiration ($R$) (McConnaughey et al., 1994; Amiotte-Suchet et al., 1999; Abril et al., 2003; Finlay and Kendall, 2007). The strength of each of these processes depends on the size of the river and its productivity (Finlay, 2003) as well as the geology and hydrology of the basin (Bullen and Kendall, 1998), and varies seasonally as a function of temperature and discharge. The complexity of all these processes and their interplay makes it difficult to constrain the sources of riverine DIC.

Riverine carbon undergoes a high degree of transformation and some degree of retention as it cycles from the terrestrial biosphere to the ocean (Cole et al., 2007; Aufdenkampe et al., 2011). In most cases, this results in supersaturation of CO$_2$ in fluvial systems and significant CO$_2$ evasion to the atmosphere (Kempe 1984; Telmer and Veizer, 1999; Cole and Caraco, 2001; Richey et al. 2002; Cole et al., 2007; Aufdenkampe et al., 2011). Quantitative assessment of riverine CO$_2$ degassing in tropical rivers is critical in the context of carbon cycling, particularly considering that tropical systems account for $\sim 70\%$ of global riverine carbon fluxes, partly owing to their large areal extent, varying climatic conditions and diverse land-use covers (Milliman and Farnsworth, 2011).

In recent papers (Aufdenkampe et al., 2011; Butman and Raymond, 2011), the global freshwater systems CO$_2$ evasion to the atmosphere has been estimated to be of the order of $\sim 3.3$ Pg C yr$^{-1}$, three-fold greater than total carbon transport ($\sim 0.9$ Pg C yr$^{-1}$) to the global ocean. Such high CO$_2$ evasion necessitates improved understanding of DIC dynamics in both local and regional systems as well as underlying drivers of carbon cycling. The above CO$_2$ evasion estimate is prone to large uncertainty, and a range of estimates has been proposed, with Cole et al. (2007) giving a much more conservative number of 0.75 Pg C yr$^{-1}$. This is partly due to the lack of sufficient data to provide a representative and correct scaled flux, particularly in tropical systems (Regnier et al., 2013).

In recent years, the application of isotopic proxies has become invaluable in understanding riverine carbon cycling. Based on the premise that various carbon sources have distinct $\delta^{13}$C values, $\delta^{13}$C$_{DIC}$ signatures have been used in a wide range of fluvial systems to constrain the biological and geological sources of DIC (Amiotte-Suchet et al., 1999; Finlay, 2003; Doctor et al., 2008; Brunet et al., 2005, 2009). For instance, riverine DIC predominantly derived from oxidation of organic matter will have $\delta^{13}$C values of about $-28$ and $-13\%_o$ for by C3 and C4 plants respectively, or $-8\%_o$ if derived from atmospheric air exchange. On the other hand, CaCO$_3$ dissolution will shift $\delta^{13}$C signatures to heavier values ($\delta^{13}$C value of carbonate is close to $0\%_o$ if from marine origin) (Finlay, 2003; Brunet et al., 2005, 2009).

In order to quantify the predominant riverine metabolic activity, application of both the $\delta^{18}$O$_{DO}$ technique (DO=dissolved oxygen) and light and dark bottle incubation experiments have effectively been employed in past studies (Sinada and Karim, 1984; Cole and Caraco, 2006; Gazeau et al., 2007). The global average $\delta^{18}$O of atmospheric oxygen is $+23.5\%_o$, but fractionation during gas exchange across the air–water interface results in DO that is $\sim 0.7\%_o$ heavier than atmospheric oxygen, and thus DO in isotopic equilibrium with air will have $\delta^{18}$O of $+24.2\%_o$ (Nimick et al., 2011). Aquatic photosynthesis lowers the isotopic composition of source water since it produces DO that is similar in isotopic composition to source water, while aquatic respiration causes a kinetic isotopic fractionation of molecular oxygen, and hence enriching water with heavier

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![Fig. 1](https://www.biogeosciences.net/10/6911/2013/) Conceptual model showing the main watershed and in-stream biogeochemical processes controlling riverine DIC concentration and $\delta^{13}$C$_{DIC}$ signatures.
isotope (Parker et al., 2005). $\delta^{18}$O$_{DO}$ can be used to quantify the rates of photosynthesis and respiration in aquatic systems. In addition, monitoring of diurnal fluctuations in riverine biogeochemical proxies can equally provide quantitative information on river metabolism (Odum, 1956; Staehr et al., 2010).

According to the river continuum concept (RCC; Vannote et al., 1980), rivers show a predictable downstream variation in the linkage with the terrestrial environment. Thus, forested headwater streams (1st–3rd order) would be predominantly heterotrophic since metabolism is fuelled by allochthonous subsidies, whereas mid-order (4th–6th) streams would be predominantly autotrophic without significant riparian canopy cover. Large rivers (> 7th order) revert to heterotrophy since $R$ would be limited by increased turbidity and depth; thus $R$ exceeds $P$. Riverine $P$ and $R$ rates are rarely quantitatively assessed at basin-wide scale despite their significance in understanding the functioning of riverine systems, yet it can be safely stated that most large rivers do not conform with the original tenets of the RCC due to altered lateral exchange with floodplains and flow discontinuities (Maiolini and Bruno, 2007). Since the RCC, a variety of alternative concepts to describe matter and energy flow have been developed, as reviewed by Bouwman et al. (2013). The metabolic balance between riverine $P$ and $R$ defines whether an ecosystem is net heterotrophic ($P: R < 1$) or net autotrophic ($P: R > 1$) (Odum 1956; Venkiteswaran et al., 2007). Most heterotrophic freshwater aquatic systems are sustained by subsidies of organic materials input from terrestrial biome (Cole and Caraco, 2001), which decompose within the riverine system, and as such influence carbon concentrations and their isotopic composition. The predominant metabolic activity thus depends on the magnitude of allochthonous inputs and the subsequent fraction respired versus autochthonous production (Cole and Caraco, 2001).

With this background, the present study examines the riverine DIC dynamics and $\delta^{13}$C$_{DIC}$ values in the Tana River basin (Kenya), based on three basin-wide sampling campaigns covering the dry season, wet season and end of the wet season in order to constrain main sources of DIC. To quantify the strength of the riverine CO$_2$ sources, we estimated both $p$CO$_2$ and CO$_2$ fluxes in an attempt to identify the primary CO$_2$ evasion hotspots within the Tana River basin. In addition, the study also examines the longitudinal changes in the magnitude of riverine $P$ and $R$ rates. We further examined to what extent pelagic respiration sustained the CO$_2$ evasion. Finally, we monitored biogeochemical parameters during diurnal cycles at three different sites along the river continuum representing headwater streams, reservoirs and the Tana River mainstream.

2 Materials and methods

2.1 Study area

The Tana River is the longest river in Kenya (~1100 km), with a total catchment area of ~96 000 km$^2$ (Fig. 2a). The river system can be separated into two main parts, here referred to as “Tana headwaters” and the “lower Tana mainstream”. The Tana headwaters consists of numerous tributaries, which originate from the Aberdare Range in the central highlands of Kenya, the highlands around Mt Kenya, and the Nyambene Hills in eastern Kenya (Fig. 2a). The lower Tana mainstream encompasses the section downstream of the Nyambene Hills, where the river flows southeast for about 700 km through semiarid plains. Along this stretch, tributaries only discharge in short pulses during the wet season. As a result, the lower Tana mainstream forms a single transport channel during the dry season, delivering material to the Indian Ocean (Maingi and Marsh, 2002). The lower Tana River has extensive floodplains between the towns of Garissa and Garsen, but floodplain inundation has been irregular in recent decades due to the river flow regulation by five hydroelectric dams upstream (Maingi and Marsh, 2002). The associated reservoirs have a combined surface area of ~150 km$^2$, and a significant amount of sediment is trapped by these dams (Dunne and Ongweny, 1976; Brown and Schneider, 1998). Masinga is the largest reservoir (commissioned in 1981), and has a surface area of ~120 km$^2$ and a storage capacity of ~1560 x 10$^6$ m$^3$, while the Kamburu Reservoir (the third largest, commissioned in 1974) has a surface area of ~15 km$^2$ and a storage capacity of ~123 x 10$^6$ m$^3$ (GOK, TARDA, 1982). The Tana River basin includes different ecological zones experiencing different rainfall patterns, decreasing from the headwaters (altitude > 3050 m, annual precipitation ~1800 mm yr$^{-1}$), upper highlands (altitude 2450–3050 m, annual precipitation ~2200 mm yr$^{-1}$) and mid-altitude catchments (altitude 1850–900 m, annual precipitation between 900 and 1800 mm yr$^{-1}$) to the lower semiarid Tana catchment (altitude 900–10 m, annual precipitation between 450 and 900 mm yr$^{-1}$) (Brown and Schneider, 1998, Fig. 2b). The basin experiences a bimodal hydrological cycle, with long rains between March and May, and short rains between October and December, which also leads to a clear bimodal pattern in the river discharge (Fig. 3). The mean annual river discharge at the Garissa gauging station is 156 m$^3$ s$^{-1}$ over the period 1934 to 1975 (daily data from the Global River Discharge Database, available at http://daac.ornl.gov/RivDIS/rivdis.shtml). The high-altitude headwaters (Aberdare Range, Mt Kenya) are characterised by mountainous forest vegetation and moorlands at the highest elevations, giving way to more intense agricultural activities in mid-altitude regions. The semiarid lower Tana is dominated by open to wooded savannah grassland, with some riverine gallery forests along the Tana River.
Fig. 2. (a) Digital elevation model (DEM) of the Tana River basin, which consists of two main geographical units – the Tana headwaters and the lower Tana mainstream. The 57 sampling sites are indicated by black dots. (b) Profile of the Tana River from headwaters to Tana mouth. Sampling stations for the lower Tana River are indicated, as well as a selected number of headwater sampling sites to show their overall position. Note: Chania Stream, Masinga Dam (close to intake) and Tana-Bura are 24 h cycle sampling sites.

2.2 Sampling and analytical techniques

Sampling was carried out during the three campaigns of February 2008 (dry season), September–November 2009 (wet season) and June–July 2010 (end of the wet season) (Fig. 3). The average river discharge measured at Garissa station during the wet season (209 m$^3$s$^{-1}$) was 1.7 times higher than during the dry season (123 m$^3$s$^{-1}$) and 1.4 times higher than the end of the wet season (145 m$^3$s$^{-1}$) (Fig. 3). Samples were taken throughout the river basin (Fig. 2a, Supplement Table 1); sampling sites included a subset of small streams in the headwater regions, an approximately equidistant set of locations along the lower Tana River mainstream, and depth profiles at two of the five hydroelectric reservoirs (Masinga and Kamburu). The dry season campaign in February 2008 only covered a subset of these field sites; these data have already been presented in Bouillon et al. (2009), and are used here for comparison purposes. In both the 2009 and 2010 campaigns, an extensive basin-wide survey was carried out. In addition, three sampling sites (Chania Stream in the Aberdare Range, Masinga Reservoir and the Tana River at Bura) were selected for a 24 h cycle sampling (Fig. 2a).

During the dry season and the wet season campaigns, water temperature and DO were measured in situ with a polarographic electrode (WTW Oxi-340) calibrated on saturated air, with an accuracy of 0.1 %, while pH was measured using a combined potentiometric electrode (Metrohm Thermo Orion model 230). A YSI ProPlus multimeter was used during the end of the wet season campaign. The pH probe was calibrated using United States National Bureau of Standards buffer solutions (4 and 7). Water samples for the analysis of $\delta^{13}$C$_{\text{DIC}}$ and $\delta^{18}$O$_{\text{DO}}$ were taken with a Niskin bottle at $\sim0.5$ m below the water surface by gently overfilling 12 mL glass headspace vials, and poisoned with 20 µL of a saturated HgCl$_2$ solution. For the analysis of $\delta^{13}$C$_{\text{DIC}}$, a 2 mL helium (He) headspace was created, and H$_3$PO$_4$ was added to convert all DIC species to CO$_2$. After overnight equilibration, part of the headspace was injected in the stream of an elemental analyser – isotope ratio mass spectrometer (EA-IRMS: Thermo Finnigan Flash HT and Thermo Finnigan Delta V Advantage) for $\delta^{13}$C measurements. The obtained $\delta^{13}$C data were corrected for the isotopic equilibration between gaseous and dissolved CO$_2$ as described in Gillikin and Bouillon (2007). Measurements were calibrated with
certified reference materials LSVEC and either NBS-19 or IAEA-CO-1. For $\delta^{18}$O, a similar headspace was created, after which samples were left to equilibrate for 2 h. $\delta^{18}$O was then measured using the same EA-IRMS setup by monitoring $m/z$ 32, 33 and 34 and using a molecular sieve (5 Å) column to separate N$_2$ from O$_2$. Outside air was used as the internal standard to correct all $\delta^{18}$O data.

Water samples for total alkalinity (TA) were obtained by prefiltering surface water through pre-combusted GF/F filters (0.7 µm), with further filtration through 0.2 µm syringe filters in high-density polyethylene bottles. TA was analysed by automated electro-titration on 50 mL samples with 0.1 mol L$^{-1}$ HCl as the titrant (reproducibility estimated as better than ±3 µmol kg$^{-1}$ based on replicate analyses). The partial pressure of CO$_2$ ($p$CO$_2$) and DIC concentrations were computed from pH and TA measurements using the carbonic acid dissociation constants for freshwater from Millero et al. (1979). The accuracy of computed DIC and $p$CO$_2$ values was estimated at ±5 µmol L$^{-1}$ and ±5 ppm, respectively. Water samples for major elements and nutrient (NO$_3^-$) were obtained in 20 mL scintillation vials in the same way as TA samples, but were preserved with 20 µL of ultra-pure HNO$_3$ and 20 µL of a saturated HgCl$_2$ solution for major elements and nutrients, respectively. The concentrations of major elements (Ca, Mg, Na, Si) were measured using inductively coupled plasma-emission spectroscopy (Iris Advantage, Thermo Jarrel-Ash), while nutrients were measured using standard chromatographic techniques (ICS-900, Dionex).

Water–air CO$_2$ fluxes were measured using a custom-made floating chamber with a surface area and volume of approximately 0.045 m$^2$ and 5.2 L, respectively, coupled with an infrared CO$_2$ analyser (Licor-820) and data logger. The floating chamber was placed on the surface of the water and kept stationary for ~10 min. The slope of the linear regression between $p$CO$_2$ and time was obtained and used to calculate the CO$_2$ flux per unit area and per unit time based on the equation

$$F_{CO2} = (dpCO2/dt)(V/RTS),$$

where $dp$CO$_2$/dt is the slope of the CO$_2$ accumulation in the chamber (atm s$^{-1}$), $V$ is the chamber volume (L), $T$ is air temperature (K), $S$ is the surface area of the chamber at the water surface (m$^2$) and $R$ is the gas constant (L atm K$^{-1}$ mol$^{-1}$).

Samples for phytoplankton pigment analysis were obtained by filtering a known volume of surface water on pre-combusted 47 mm GF/F filters, which were immediately packed into cryotubes and stored in liquid N$_2$. Upon return to the laboratory, the samples were stored at −20°C until further analysis. Pigments were extracted in 10 mL of an acetone: water mixture (90:10), and a subsample was separated by high-performance liquid chromatography on a C18 reverse-phase column. Calibration was performed with working standards prepared from commercially available pure compounds.

The $R$ rates were determined by quantifying the decrease in DO (using the polarographic electrode WTW Oxi-340) of river water incubated in a dark, cool box filled with water to retain ambient temperature for approximately 24 h using 250 mL bottles (2–3 replicates) with airtight polyconic caps. The $P$ rates were quantified by determining the uptake of DIC after short-term (1–3 h) in situ incubations of river water during the day using 1 L polycarbonate bottle spiked with $^{13}$C-labelled sodium bicarbonate (NaH$^{13}$CO$_3$). A subsample for $\delta^{13}$CDIC of the spiked water was sampled for the purpose of computations. Samples for analysis of particulate organic carbon (POC) and $\delta^{13}$CPOC were obtained at the start (natural abundance values) and at the end of the incubation by filtering a known volume of surface water on pre-combusted (overnight at 450°C) 25 mm GF/F filters (0.7 µm). Filters were decarbonated with HCl fumes for 4 h, re-dried and then packed into Ag cups. POC and $\delta^{13}$CPOC were determined on a Thermo elemental analyser – isotope ratio mass spectrometer (EA-IRMS) system (Flash HT with Delta V Advantage), using the TCD signal of the EA to quantify POC and by monitoring $m/z$ 44, 45 and 46 on the IRMS. Quantification and calibration of $\delta^{13}$C data were performed with IAEA-C6 and acetanilide that was calibrated against international standards. Reproducibility of $\delta^{13}$CPOC measurements was typically better than 0.2%. The P data here refer only to surface water (~0.5 m deep) measurements (not depth integrated). Calculations to quantify the rates were made as described in Dauchez et al. (1995).

3 Results

3.1 In situ measurements, dissolved inorganic carbon and $\delta^{13}$CDIC

Generally, temperature, pH and specific conductivity increased consistently downstream (Pearson correlation: $r^2 = 0.90$, 0.41 and 0.47 for temperature, pH and specific conductivity, respectively; $p < 0.05$; $n = 132$) with water temperature ranging from 10.6 to 32.1°C, pH from 6.61 to 8.71, specific conductivity from 11 to 582 µS cm$^{-1}$ and DO from 41 to 134 % during the 2008 dry season, 2009 wet season and 2010 end of the wet season campaigns (Supplement Table 1, Supplement Figs. 1 and 2). DIC concentrations ranged from 0.2 to 4.8 mmol L$^{-1}$ with an overall mean of 1.1 ± 1.0 mmol L$^{-1}$. Exceptionally high values (3.5 ± 1.6 mmol L$^{-1}$) were systematically recorded in Nyambene Hills tributaries during all three campaigns (Fig. 4). DIC concentrations increased consistently downstream (Pearson correlation: $p < 0.01$, $r^2 = 0.67$; $n = 108$) when excluding values from Nyambene Hills tributaries. The DIC concentrations within different tributaries were similar (paired t test, $p > 0.05$) during the three campaigns. In contrast, DIC concentrations in the Tana River mainstream were significantly higher (ANOVA, $p < 0.05$) during the 2009 wet
season campaign (1.3 ± 0.2 mmol L⁻¹) and lowest during the 2010 end of the wet season campaign (1.1 ± 0.2 mmol L⁻¹).

δ¹³CDIC signatures ranged from −20.8 to −2.4 % during the three campaigns. δ¹³C DIC values within different tributaries were similar (paired t test, p > 0.05) during all three campaigns, but the Tana River mainstream recorded relatively depleted values (−11.2 ± 1.8 %) during the 2009 wet season campaign compared to the 2008 dry season and 2010 end of wet season campaigns combined (−8.7 ± 0.7 %). Overall, δ¹³C DIC showed a negative correlation with DIC concentrations (Pearson correlation: p < 0.01, r² = 0.49; n = 122) during all three campaigns, with tributaries from the Aberdare and Mt Kenya regions recording the lowest DIC concentrations (0.6 ± 0.3 mmol L⁻¹) and highest δ¹³C DIC signatures (−6.9 ± 2.7 %) and the Tana River mainstream intermediate DIC concentrations (1.2 ± 0.2 mmol L⁻¹) and δ¹³C DIC signatures (−9.7 ± 1.7 %), whereas Nyambene Hills tributaries recorded the highest DIC concentrations (3.5 ± 1.6 mmol L⁻¹) and the lowest δ¹³C DIC signatures (−11.3 ± 2.4 %). Substantially higher δ¹³C DIC values were observed at Masinga Reservoir (−4.5 ± 0.1 %) during the dry season, but this decreased rapidly to −8.9 % less than 5 km downstream.

3.2 Dynamics of riverine pCO₂ and CO₂ fluxes

The whole river network, with a few exceptions, was supersaturated with CO₂ (pCO₂ values between 110 and 5204 ppm) with respect to atmospheric equilibrium (385 ppm), with an overall median of 1002 ppm. Higher values were recorded during the 2009 wet season campaign (median 1432 ppm) compared to the 2010 end of the wet season (median 1002 ppm), whereas the 2008 dry season was two-fold lower (median 579 ppm, Fig. 5). Tributaries from the Aberdare and Mt Kenya regions recorded lower pCO₂ concentrations (median 867 ppm) compared to those from Nyambene Hills tributaries and the Tana River mainstream (median 1220 ppm, ANOVA p < 0.05, Fig. 6b). During the 2009 wet season campaign, tributaries from Mt Kenya region recorded higher pCO₂ concentrations (t test, p < 0.05), particularly those in mid-altitude catchments (1850–900 m), characterised by intensive agricultural practices. During the 2009 wet season campaign, sampling at Kamburu Dam was done shortly before the actual onset of the wet season. Surface waters were highly undersaturated in CO₂ during both the wet (pCO₂ = 118 ppm) and end of the wet season campaigns (pCO₂ = 305 ppm). Surface waters of Masinga Reservoir were similarly undersaturated in CO₂ during the 2008 dry season campaign (pCO₂ from 313 to 443 ppm, Bouillon et al., 2009) but relatively oversaturated in CO₂ during the end of the wet season campaign (pCO₂ from 2629 to 3389 ppm). However, pCO₂ was elevated downstream of the Masinga Reservoir exit (2303 to 5204 ppm at Tana River–Masinga Bridge) during all three campaigns, but the concentrations decreased rapidly downstream (Supplement Table 1). pCO₂ showed a negative correlation with δ¹³C DIC values (Pearson correlation: p < 0.01, r² = 0.52; n = 125; Fig. 6a) during all three campaigns.

CO₂ fluxes between the river and the atmosphere based on floating chamber measurements ranged from −12.4 to 639.0 mmol m⁻² d⁻¹ during the wet season campaign and from −57.1 to 813.7 mmol m⁻² d⁻¹ during the end of the wet season campaign. The CO₂ fluxes were similar during the two seasons (Paired t test, p > 0.05). Similar to pCO₂, disproportionately high water–air evasion was recorded during the end of the wet season campaign downstream of Masinga Reservoir exit (Masinga Bridge, 600 mmol m⁻² d⁻¹). On average, Nyambene Hills tributaries recorded the highest CO₂ evasion (−4.2 to 617 mmol m⁻² d⁻¹) during both seasons (Fig. 7) corresponding with higher pCO₂ values (806 to 4915 ppm). Generally, CO₂ fluxes showed a positive correlation with pCO₂ (Pearson correlation: p < 0.01, r² = 0.61; n = 76).
Nitrates concentrations ranged from 0.4 to 146 µmol L\(^{-1}\) during all three seasons (Supplement Table 2). The concentrations increased consistently downstream during both the wet season (Pearson correlation: \(p < 0.01, r^2 = 0.75; n = 50\)) and end of the wet season (Pearson correlation: \(p < 0.01, r^2 = 0.50; n = 50\); Fig. 8b), but during the dry season, nitrate did not show a systematic pattern with altitude. The highest mean concentrations (78.6 to 96.3 µmol L\(^{-1}\)) were recorded in Nyambene Hills tributaries during all three seasons.

Chlorophyll \(a\) concentration ranged from 0.18 to 9.60 µg L\(^{-1}\) with means of 2.74 ± 2.52, 1.94 ± 1.94 and 2.20 ± 2.75 µg L\(^{-1}\) during the dry season, the wet season and end of the wet season respectively (Supplement Table 2). The chlorophyll \(a\) concentration increased consistently downstream (Pearson correlation: \(p < 0.01, r^2 = 0.55; n = 117\); Fig. 9b), and values during different seasons were not significantly different (paired \(t\) test, \(p > 0.05\)). Elevated chlorophyll \(a\) concentrations (3.2 to 20.6 µmol L\(^{-1}\)) were recorded in the Kamburu and Masinga reservoirs during all three sampling campaigns.

The primary production rates ranged from 0.004 to 7.38 µmol C L\(^{-1}\) h\(^{-1}\) with means of 0.42 ± 0.46 and 0.83 ± 1.58 µmol C L\(^{-1}\) h\(^{-1}\) for the wet and the end of the wet season campaign, respectively. The primary production rates increased consistently downstream (Pearson correlation: \(r^2 = 0.55, p < 0.01; n = 84\); Fig. 9a) during both campaigns. Disproportionately high values ranging between 10.28 and 18.28 µmol C L\(^{-1}\) h\(^{-1}\) were recorded in the Kamburu and Masinga reservoirs (Supplement Table 2). The means for different seasons were similar (paired \(t\) test, \(p > 0.05\)), but Mt Kenya and Nyambene Hills tributaries recorded significantly higher values (0.19 ± 0.22 and 0.79 ± 0.45 µmol C L\(^{-1}\) h\(^{-1}\), respectively) during the 2009 wet season campaign (paired \(t\) test, \(p < 0.05\)) compared to the 2010 end of the wet season campaign (0.07 ± 0.07 and 0.26 ± 0.28 µmol C L\(^{-1}\) h\(^{-1}\), respectively).

Riverine \(R\) rates were measured only during the 2009 wet season campaign and ranged from 0.20 ± 0.07 to 3.64 ± 0.03 µmol O\(_2\) L\(^{-1}\) h\(^{-1}\) with a basin-wide mean of 1.43 ± 0.91 µmol O\(_2\) L\(^{-1}\) h\(^{-1}\) (Supplement Table 2). The \(R\) rates increased downstream (Pearson correlation: \(p < 0.01, r^2 = 0.38; n = 33\); Fig. 8a), with Aberdare and Nyambene Hills tributaries recording the lowest mean (0.84 ± 0.84 µmol O\(_2\) L\(^{-1}\) h\(^{-1}\)) and Mt Kenya tributaries an intermediate one (1.37 ± 0.90 µmol O\(_2\) L\(^{-1}\) h\(^{-1}\)), while the lower Tana River mainstream stretch had the highest (1.92 ± 0.72 µmol O\(_2\) L\(^{-1}\) h\(^{-1}\)). The \(R\) rates also showed a positive correlation with temperature (Pearson correlation: \(r^2 = 0.46, p < 0.01; n = 33\)).
3.4 Diurnal biogeochemical processes

During the diurnal cycles, water temperature fluctuated by 9.7 (6.9–16.6 °C), 2.1 (21.8–23.9 °C) and 1.5 °C (24.6–26.1 °C), while pH fluctuated by ∼0.58 (7.67–8.25), 0.14 (6.98–7.12) and 0.05 units (7.83–7.88) at Chania Stream, Masinga Dam and Tana River mainstream, respectively (Table 1). The DO saturation fluctuated by 34.6 (89.4–124.0 %), 10.6 (63.0–73.6 %) and 3.2 % (96.7–99.9 %) at Chania Stream, Masinga Dam and Tana River mainstream, respectively (Table 1). The in situ field measurements of pH, temperature and DO increased steadily during daytime and decreased at night at Chania Stream (Fig. 10), while no discernible trends were observed at Masinga Dam and the Tana River mainstream.

$\delta^{18}O_{DO}$ signatures at Chania Stream ranged from +22.5 to +25.5 ‰ (Supplement Table 3), with a diurnal mean of +24.0 ± 0.8 ‰. Although the magnitude of change was small, the values decreased during day time and increased at night time (Fig. 10). $\delta^{18}O_{DO}$ signatures showed a strong negative correlation with DO saturation (Pearson correlation: $p < 0.01$, $r^2 = 0.84$; $n = 23$; Fig. 11a). Masinga Dam values (+25.4 to +28.2 ‰) showed moderate diurnal fluctuation, while the Tana River mainstream (+24.6 to +25.5 ‰) values did not exhibit clear diurnal fluctuations. The $\delta^{18}O_{DO}$ values at both the Masinga Dam and Tana River mainstream remained above the atmospheric equilibrium (+24.2 ‰) during the whole diurnal cycle, but values at Chania Stream oscillated above and below atmospheric equilibrium, though with a pronounced diurnal cycle (Fig. 10b). The maximum $\delta^{18}O_{DO}$ value (+25.5 ‰) at Chania Stream was recorded at 04:00 LT (all times mentioned are in local time) corresponding with lowest DO saturation (90.8 %). Generally, $\delta^{18}O_{DO}$ values at Chania Stream were below atmospheric equilibrium (+24.2 ‰) during daytime (07:00 to 17:00) and above atmospheric equilibrium at night (18:00 to 06:00).

The DIC values during the diurnal cycles ranged from 0.506 to 0.573 mmol L$^{-1}$, 0.603 to 0.719 mmol L$^{-1}$, and 1.152 to 1.224 mmol L$^{-1}$, whereas $\delta^{13}C_{DIC}$ ranged from −7.2 to −5.8 ‰, −10.4 to −9.6 ‰, and −8.9 to −8.0 ‰ for Chania Stream, Masinga Dam and the Tana River mainstream, respectively (Supplement Table 3). The latter two
sites showed minimal $\delta^{13}\text{DIC}$ diurnal amplitude ($\sim 0.9\%$) compared to Chania Stream (1.4%). $\delta^{13}\text{C}_{\text{DIC}}$ values at Chania Stream typically increased during daytime and decreased at night (Fig. 12a). While the amplitudes of both DIC and $\delta^{13}\text{C}_{\text{DIC}}$ at Chania Stream were small, their diurnal cycles were pronounced but showed reversed patterns (Fig. 12a). In contrast, both the Masinga Dam and Tana River mainstream did not exhibit clear diurnal patterns for either $\delta^{13}\text{C}_{\text{DIC}}$ or DIC. However, the DIC concentrations at the Tana River mainstream (1.191 ± 0.020 mmol L$^{-1}$) were significantly high (ANOVA, $p < 0.01$) compared to those of Chania Stream (0.547 ± 0.021 mmol L$^{-1}$) and Masinga Reservoir (0.666 ± 0.027 mmol L$^{-1}$).

During the diurnal cycles, $p\text{CO}_2$ values ranged from 174 to 614 ppm, 2621 to 3684 ppm, and 980 to 1131 ppm at Chania Stream, Masinga Dam and Tana River mainstream, respectively (Supplement Table 3). $p\text{CO}_2$ fluxes and $p\text{CO}_2$ showed an identical diurnal pattern at Chania Stream (Fig. 13a), but $p\text{CO}_2$ fluxes at Masinga Dam and Tana River mainstream were consistently high throughout diurnal cycle and did not exhibit clear diurnal patterns.

The gas transfer velocity ($k_{600}$) computed from the flux $p\text{CO}_2$ and the air–water $p\text{CO}_2$ gradient was on average much higher at Chania Stream (25.6 ± 13.0 cm h$^{-1}$) than at Masinga Reservoir (7.7 ± 2.4 cm h$^{-1}$) and at Tana River at Bura (1.3 ± 0.4 cm h$^{-1}$) (Supplement Table 3).
Discussion

4.1 Longitudinal patterns of dissolved inorganic carbon and $\delta^{13}$C$_{DIC}$

In the Tana River basin, DIC generally increased downstream during all three seasons, a strong indication of DIC build-up possibly contributed by rock weathering. This is particularly evident along the Tana River mainstream, where DIC strongly increased downstream during all seasons (Fig. 4a), which corresponded to an increase in suspended matter (Bouillon et al., 2009; Tamooh et al., 2012). Nyambene Hills tributaries recorded exceptionally high DIC concentrations (Fig. 4a) during all three seasons, a phenomenon we associate with a high rate of rock weathering. The lithology of the Nyambene Hills subcatchment mainly includes Quaternary volcanic rocks. The high rate of chemical weathering could result from the presence of ashes and pyroclastic rocks, as these have much higher weathering rates than basalts and trachytes (Hughes et al., 2012). HCO$_3^-$ in rivers is mainly derived from the weathering of carbonate and silicate rocks. The relative proportion of HCO$_3^-$ in rivers derived from weathering of carbonate and silicate rocks depends to a very large extent on the lithology of the drainage basin (Garrels and Mackenzie, 1971; Meybeck, 1987). In Kenya, sedimentary carbonate rocks are associated with the Mozambique Belt, south of the Tana drainage basin (Warden and Horkel, 1984). The Tana drainage basin is rich in silicate rocks as it is mainly composed of Tertiary and Quaternary volcanic rocks, mainly basalt and a broad spectrum of alkaline rocks (e.g. Baker, 1987). Also, Kenya is extremely rich in alkaline rocks and carbonates (Woolley, 2001), including calcite carbonate (Rosatelli et al., 2003). Finally, saline and sodic soils characterise the dry lands in Kenya (Dijkshoorn et al., 2010), and the latter soils are rich in carbonates.

TA is well correlated with dissolved silicate (DSi) and the sum of Ca$^{2+}$ and Mg$^{2+}$ (Fig. 14), suggesting that HCO$_3^-$ in the Tana originates from both carbonate and silicate rock weathering. The strong correlation between TA and Na$^+$ provides additional evidence for silicate weathering, which is the main source of Na$^+$ in rivers (data not shown). To discriminate the origin of HCO$_3^-$, we applied the simple stoichiometric model of Garrels and Mackenzie (1971), whereby the contribution to TA from carbonate weathering (TA$_{carb}$) is computed from Ca$^{2+}$ and Mg$^{2+}$, and the contribution to TA from silicate weathering (TA$_{sil}$) is computed from DSi according to

$$TA_{carb} = 2 \times ([Ca^{2+}] + [Mg^{2+}] - [SO_4^{2-}]),$$

$$TA_{sil} = [DSi]/2.$$
SO\textsubscript{4}\textsuperscript{2−} allows for Ca originating from the dissolution of gypsum (CaSO\textsubscript{4}) to be accounted for. However, SO\textsubscript{4}\textsuperscript{2−} data were only available for the 2009 wet season campaign, so we did not apply this correction. For the 2009 wet season campaign, SO\textsubscript{4}\textsuperscript{2−} corresponded on average to 11 % of Ca\textsuperscript{2+}. The resulting estimated relative contributions of silicate weathering to total rock weathering ranges between 7 and 78 % and averages 28 %, close to the overall average of 24 % for African rivers reported by Garrels and Mackenzie (1971). Modelled TA (TA\textsubscript{carb} + TA\textsubscript{sil}) is very close to measured TA, as most data points fall on the 1 : 1 line, with the exception of eight outliers (Fig. 15), which are mostly from the three rivers in the Nyambene Hills (Murera, Rojewero and Mutundu).

As previously reported by Hughes et al. (2012), the Nyambene Hills region is characterised by high silicate weathering, with DSi values up to 1148 µmol L\textsuperscript{−1} (Supplement Table 1), above the maximum of 958 µmol L\textsuperscript{−1} previously reported by Hughes et al. (2012). In the absence of detailed lithological information of the drainage basin in Nyambene Hills, it is difficult to explain the discrepancy between modelled and observed TA. Precipitation of hydrous sodium silicates and zeolites (Sebag et al., 2001) could explain the discrepancy because it would lead to a removal of DSi but leaving HCO\textsubscript{3}− unaffected. The stoichiometric model of Garrels and Mackenzie (1971) assumes that the weathering of silicate rocks leads to a release of HCO\textsubscript{3}− and DSi according to a 1 : 2 ratio. However, the dissolution of certain silicates such as plagioclase feldspar (NaCaAl\textsubscript{3}Si\textsubscript{5}O\textsubscript{16}) may lead to the release of HCO\textsubscript{3}− and DSi according to a 6 : 4 ratio. When we apply this ratio to the data points of Nyambene Hills rivers, the recomputed modelled TA is in better agreement with observed TA than based on the HCO\textsubscript{3}− : DSi ratio of 1 : 2 (Fig. 15).

The dynamics of riverine δ\textsuperscript{13}DIC are primarily controlled by both in-stream and watershed processes (Finlay and Kendall, 2007). The wide range of δ\textsuperscript{13}DIC values recorded in the present study (−15.0 to −2.4 ‰) suggests variable carbon sources. The δ\textsuperscript{13}DIC values of the Aberdare Range (−6.8 ± 1.6 ‰) and Mt Kenya (−6.7 ± 2.4 ‰) tributaries, and particularly those in high-altitude regions (>1700 m), were characterised by high δ\textsuperscript{13}DIC values consistent with carbonate mineral weathering. These high values may be reinforced by the fact that high-altitude tributaries are characterised by fast and turbulent flows, thus enhancing CO\textsubscript{2}
evasion to the atmosphere. Since the CO\textsubscript{2} fraction is isotopically depleted relative to HCO\textsubscript{3}\textsuperscript{-} and CO\textsubscript{3}\textsuperscript{2-}, this process leads to a gradual enrichment of the remaining DIC pool (e.g. Doctor et al., 2008). We investigated the downstream (\(<1\) km) gradients from the Satima springs, a high-altitude stream in the Aberdare Range (3600 m). We noticed a rapid increase in $\delta^{13}$C\textsubscript{DIC} values ($-20.8$ to $-8.8\%$) over the first 200 m, corresponding to a fast decrease in $p$CO\textsubscript{2} (4789 to 468 ppm) due to rapid CO\textsubscript{2} degassing into the atmosphere (Fig. 16). High $\delta^{13}$C\textsubscript{DIC} values ($-4.5 \pm 0.1\%$, $n = 4$) were recorded at Masinga Reservoir during the dry season campaign (Bouillon et al., 2009), consistent with CO\textsubscript{2} degassing combined with CO\textsubscript{2} consumption by primary production. The increase in water residence time at the reservoir also favours CO\textsubscript{2} exchange with the atmosphere. High primary production was demonstrated by high chlorophyll $a$ concentrations in the Masinga Reservoir. Similar observations have been reported in other dammed rivers (Brunet et al., 2005; Zeng et al., 2011). $\delta^{13}$C\textsubscript{DIC} values were relatively depleted in the lower Tana mainstream during the three seasons. Lower $\delta^{13}$C\textsubscript{DIC} values correlated with an increase in total suspended matter (TSM) (Bouillon et al., 2009; Tamooh et al., 2012), suggesting that degradation of laterally derived organic matter plays an important role in controlling $\delta^{13}$C\textsubscript{DIC} values along the lower Tana River mainstream. When excluding values from Nyambene Hills tributaries, the negative relationship between $\delta^{13}$C\textsubscript{DIC} values and DIC concentration (Pearson correlation: $p < 0.01$, $r^2 = 0.62$; $n = 39$) and the positive correlation between $\delta^{13}$C\textsubscript{DIC} values and DO (Pearson correlation: $p < 0.01$, $r^2 = 0.52$; $n = 39$) during the wet season campaign further suggests the contribution of organic matter degradation.

### 4.2 Factors regulating $p$CO\textsubscript{2} and CO\textsubscript{2} fluxes

As reported for other tropical rivers (Richey et al., 2002; Mayorga et al., 2005; Aufdenkampe et al., 2011), the Tana River was generally supersaturated in CO\textsubscript{2} with respect to atmospheric equilibrium. This was the case during all three seasons. The overall median $p$CO\textsubscript{2} for the Tana River (1002 ppm) is, however, relatively low compared to the global median of $\sim 3600$ and $\sim 4300$ ppm for tropical rivers and streams, respectively (Aufdenkampe et al., 2011). The Tana River tributaries at high altitudes were characterised by relatively low $p$CO\textsubscript{2} concentrations compared to other tropical high-altitude streams, e.g. 11 000 to 25 000 ppm for Amazon headwaters (Davidson et al., 2010). One possible explanation is relatively low in-stream respiration as Tana River headwater streams are characterised by low suspended sediment (Bouillon et al., 2009; Tamooh et al., 2012) and low temperatures, resulting in lower CO\textsubscript{2} production from organic matter degradation. In addition, the CO\textsubscript{2} carried by groundwater inputs rapidly degasses to the atmosphere, as observed in the Aberdare Range headwaters. The CO\textsubscript{2} from groundwater at Satima springs rapidly degassed (4789 to 468 ppm) with increasing distance from the spring source (Fig. 16), resulting in $\sim 90\%$ CO\textsubscript{2} evasion over a distance of a few hundred metres. Similar observations have been reported from both temperate (Doctor et al., 2008; Wallin et al., 2012) and tropical headwater streams (Johnson et al., 2008; Davidson et al., 2010), thus depicting small streams as hotspots for CO\textsubscript{2} evasion. Further, we observed a strong coupling between isotopic signatures of DIC and $p$CO\textsubscript{2} where $\delta^{13}$C\textsubscript{DIC} values ($-20.8$ to $-8.8\%$) became rapidly enriched downstream, thus providing further evidence of rapid degassing downstream from groundwater inputs (Fig. 16). In contrast to Aberdare and Mt Kenya streams, Nyambene Hills tributaries recorded relatively high $p$CO\textsubscript{2} values (median

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**Fig. 15.** Observed total alkalinity against modelled total alkalinity along the Tana River basin.

**Fig. 16.** A plot showing behaviour of CO\textsubscript{2} concentration and $\delta^{13}$C\textsubscript{DIC} at the Satima springs groundwater source in the headwaters of the Aberdare Range.
1377 ppm, n = 12; Fig. 6b) as well as high CO\textsubscript{2} evasion (median \( \sim 260 \text{mmol m}^{-2} \text{d}^{-1} \), Fig. 7, Supplement Table 1), notably in Murera and Mutundu tributaries during both the wet and end of wet season campaigns. However, Nyambene Hills tributaries were characterised by a cover of high organic matter on the river bed up to 17% (Tamooh et al., 2012), possibly contributing to additional generation of CO\textsubscript{2} through respiration. This way, Nyambene Hills subcatchment emerges as a major hotspot for CO\textsubscript{2} emissions within the Tana River basin.

Primary production strongly controlled pCO\textsubscript{2} in the reservoirs, as demonstrated by low pCO\textsubscript{2} values during the dry season at Masinga Reservoir (313 to 443 ppm) and Kambugu Dam during both the wet (118 ppm) and end of the wet season (305 ppm) campaigns (Supplement Table 1). Low turbidity coupled with high water residence time and temperatures (\( \sim 30^\circ \text{C} \)) in the reservoirs enhanced phytoplankton production, thus consuming CO\textsubscript{2}. In contrast, the depth profile data obtained at Masinga Dam (up to 15 m deep) recorded consistently high pCO\textsubscript{2} values (3385 \( \pm \) 212 ppm, n = 5) during the end of the wet season campaign. This suggests that the dam water was vertically mixed owing to the fact that this sampling campaign coincided with an unusual dam infilling (following high precipitation in headwaters) and excess outflow at the spillway, thus enhancing upwelling. Furthermore, a maximal pCO\textsubscript{2} value (5059 ppm) was recorded at 20 m depth (Supplement Table 1). Conversely, Kambugu Dam recorded highly stratified pCO\textsubscript{2} values during both the wet season (118 ppm at surface waters and 2556 ppm at 20 m depth) and end of the wet season campaigns (305 ppm at surface waters and 3270 ppm at 20 m depth) (Supplement Table 1). In addition, consistently high pCO\textsubscript{2} values (up to 5204 ppm) and CO\textsubscript{2} flux (600 mmol m\textsuperscript{-2} d\textsuperscript{-1}) were recorded downstream of the Masinga Reservoir exit (Masinga Bridge), a phenomenon we attribute to the release of hypolimnetic, CO\textsubscript{2}-rich water. However, the CO\textsubscript{2} degassed rapidly < 5 km downstream, where we observe \( \sim 90 \% \) of CO\textsubscript{2} being degassed before reaching the entrance of Kambugu Dam (Supplement Table 1). Thus, Masinga Reservoir contributes significantly to CO\textsubscript{2} emission downstream, consistent with observations reported in some tropical reservoirs where CO\textsubscript{2} fluxes ranging from 412 to 1494 mmol m\textsuperscript{-2} d\textsuperscript{-1} were recorded immediately downstream of the reservoirs (Guérin et al., 2006; Abril et al., 2005).

While different in-stream and watershed processes regulate riverine pCO\textsubscript{2} dynamics, organic matter respiration within the river seemed to be the major controlling process in the Tana River mainstream, as reflected by high pCO\textsubscript{2} values during the 2009 wet season campaign. Furthermore, the lower Tana River mainstream was characterised by high suspended sediment and dissolved organic carbon, thus fuelling in-stream respiration (Tamooh et al., 2012). This is further corroborated by the general negative correlation observed between \( \delta^{13}\text{C}_{\text{DIC}} \) values and pCO\textsubscript{2} values (Fig. 6a). In-stream processing and decomposition of terrestrially derived organic carbon have been reported to primarily regulate CO\textsubscript{2} evasion in most large tropical rivers (Richey et al., 2002; Mayorga et al., 2005).

### 4.3 Dynamics of aquatic metabolism

River metabolism is primarily influenced by the availability of light, nutrients, temperature and organic matter subsidies. Both primary production rates and chlorophyll \( a \) concentrations increased consistently downstream in the Tana River basin (Fig. 9), contrary to expectation, as the turbidity increased downstream (Bouillon et al., 2009; Tamooh et al., 2012). However, primary production rates here are based on incubations of surface water within the euphotic zone (\( \sim 0.5 \text{m deep} \)) and do not represent depth-integrated values (deeper in the water column light is obviously extinct). The downstream increase in primary production can be attributed to higher temperatures coupled with higher water residence time in the lower Tana River mainstream. In addition, during the wet season, the nitrate concentrations increased downstream (Pearson correlation: \( r^2 = 0.75, p < 0.01; n = 50 \), Fig. 8b), particularly along the lower Tana River mainstream, thus potentially enhancing primary production. This counterintuitive observation of high productivity in turbid rivers has also been reported for some large temperate rivers, such as the Mississippi (Kendall et al., 2001; Delong and Thorp, 2006) and Meuse (Descry and Gosselain, 1994), as well as in the White Nile (Sinada and Karim, 1984). While most headwater tributaries are characterised by low suspended matter, primary production rates were low owing to nutrient limitation, low temperature and high stream flow velocity – factors unfavourable for efficient phytoplankton production. Generally, primary production in tropical rivers has been reported to be quite variable ranging from lows of 10–200 mg C m\textsuperscript{-2} d\textsuperscript{-1} to over 1000 mg C m\textsuperscript{-2} d\textsuperscript{-1} in relatively unpolluted systems, but can exceed 6000 mg C m\textsuperscript{-2} d\textsuperscript{-1} in polluted systems (Davies et al., 2008).

The mean river depth between Kora Bridge and the Tana River mouth during the wet season ranges between 2.0 and 3.0 m, while the average respiration rate along the same stretch was 1.92 \( \pm \) 0.72 \( \mu \text{mol L}^{-1} \text{h}^{-1} \). This would translate to a depth-integrated pelagic respiration ranging between 92 and 138 mmol m\textsuperscript{-2} d\textsuperscript{-1}, while the median CO\textsubscript{2} efflux was 156 mmol C m\textsuperscript{-2} d\textsuperscript{-1} (Table 2, Fig. 7a). Assuming the net primary production is negligible; this suggests that the bulk of CO\textsubscript{2} evasion (77 to 114%) along the lower Tana mainstream can be accounted for by in-stream respiration (Table 2, Fig. 7a). These estimates are based on the assumption that the respiration quotient between O\textsubscript{2} and CO\textsubscript{2} is 1.3, as suggested by Richardson et al. (2013). Similar observations have been reported in the Amazon Basin, where in-stream respiration was highly variable but could account for up to \( \sim 100 \% \) of the CO\textsubscript{2} evasion in some rivers (Ellis et al., 2012).

In contrast, most headwater tributaries in the Tana River basin are shallow (\( \sim 0.5–1 \text{m deep} \)), with
depth-integrated respiration rates ranging between ~10 and 33 mmol m$^{-2}$ d$^{-1}$, while the median CO$_2$ fluxes ranged between 38 and 260 mmol m$^{-2}$ d$^{-1}$ (Table 2, Fig. 7a). This implies that in-stream respiration accounts for a smaller fraction of total CO$_2$ evasion (5 to 68 %) in the studied tributaries compared to the Tana River mainstream. This is particularly the case for the Nyambene Hills tributaries, whose pelagic respiration accounts for only 5 to 10 % of total CO$_2$ evasion (Table 2, Fig. 7a). This suggests the excess bulk CO$_2$ evasion must be sustained by other processes such as inputs of groundwater (highly CO$_2$ oversaturated to soil and root respiration) and/or in-stream benthic respiration. The latter hypothesis is in agreement with the fact that most Nyambene Hills tributaries were characterised by high organic matter on the bottom of the river bed possibly fuelling respiration.

4.4 Factors regulating diurnal variations in surface water biogeochemistry

Diurnal fluctuations of biogeochemical variables were substantial in the Chania headwater stream, moderate at Masinga Reservoir and not detectable within the lower Tana mainstream. Dissolved oxygen was supersaturated with respect to atmospheric equilibrium at the headwaters site (Chania Stream), reaching a maximum value of 124.0 % during daytime (11:00) and minimum saturation of 89.4 % at 06:00, shortly before sunrise. Similarly, the pH increased during daytime (indicative of CO$_2$ consumption by photosynthesis) and decreased at night (indicative of CO$_2$ release by respiration). These patterns were absent at both the Masinga Dam and lower Tana River mainstream, where biogeochemical parameters remained nearly constant during the whole diurnal cycle. Masinga Dam surface waters were particularly undersaturated in DO (63.0 to 73.6 %, Table 1), suggesting the dominance of respiration and net heterotrophy. The absence and/or small fluctuation of DO at Masinga and the Tana River mainstream may be attributed to high suspended loads (Bouillon et al., 2009; Tamooh et al., 2012), which attenuate photosynthetic light, as opposed to the headwaters site (Chania Stream), which is characterized by low suspended matter concentrations.

The $\delta^{18}$O$_{DO}$ signatures fluctuated by ~3.0, 2.8, and 0.9‰ at Chania Stream headwaters, Masinga Reservoir and Tana River mainstream, respectively. The signatures at Chania Stream were isotopically light during daytime and enriched at night, fluctuating slightly below and above the atmospheric equilibrium value (+24.2 ‰). This suggests that aquatic P dominated riverine biological activities during the day but reversed to R dominance at night. Similar observations have been reported in small streams elsewhere (Parker et al., 2005; Waldron et al., 2007; Gammons et al., 2011; Parker et al., 2010; Nimick et al., 2011). In contrast, Masinga Dam and Tana River mainstream $\delta^{18}$O$_{DO}$ signatures were consistently above atmospheric equilibrium (+24.2 ‰), suggesting the two sites were dominated by R and thus predominantly heterotrophic during the whole diurnal period.

Precipitation and dissolution of CaCO$_3$ in freshwaters can be deduced from changes in TA (e.g. McConnaughey et al., 1994). At Masinga Dam and Tana River mainstream, TA did not show a distinct pattern during the diurnal cycle. For Chania Stream, the TA values followed a regular pattern, with a decrease from 01:00 to 13:00 of about 0.035 mmol L$^{-1}$, (Fig. 13b) inconsistent with the one expected from daily cycles of precipitation/dissolution of CaCO$_3$ (increase of TA during night-time and decrease of TA during daytime; de Montety et al., 2011; Gammons et al., 2011). The changes of TA at Chania Stream followed those of conductivity, suggesting overall changes in the chemistry of flowing waters rather
than in-stream processes affecting TA. We conclude that changes in DIC can be ascribed to \( P \) and \( R \) and that precipitation/dissolution of \( \text{CaCO}_3 \) was not a major process contributing to daily changes of DIC. The DIC concentration showed a fluctuation of \( \sim 0.067, 0.116, \) and \( 0.072 \text{ mmol L}^{-1} \) at Chania Stream, Masinga Reservoir and Tana River mainstream, respectively. However, DIC concentrations and \( \delta^{13}\text{C}_{\text{DIC}} \) values showed a strong inverse relationship (Pearson correlation: \( p < 0.01, r^2 = 0.87; n = 22; \) Fig. 11c) characterised by a pronounced diurnal cycle at Chania Stream only. DIC concentrations at Chania Stream increased at night due to aquatic community respiration, and decreased during daytime due to \( \text{CO}_2 \) uptake by photosynthesis. Studies elsewhere have reported substantial diurnal changes in DIC concentrations ranging from 0.300 to 0.420 \text{ mmol L}^{-1} \) (Poulson and Sullivan, 2010; Gammons et al., 2011) with a record fluctuation of \( \sim 1.133 \text{ mmol L}^{-1} \) reported by Parker et al. (2010). However, this large diurnal change in DIC concentration is reported to have been driven by unique calcite precipitation and not in-stream \( P \) or \( R \) (Gammons et al., 2007; Nimick et al., 2011).

Similarly, \( \delta^{13}\text{C}_{\text{DIC}} \) values at Chania Stream showed a modest fluctuation of \( \sim 1.4\%_e \), with signatures increasing during daytime as \( \text{CO}_2 \) was preferentially utilised by aquatic periphyton phototrophs at a faster rate than \( ^{13}\text{CO}_2 \). At night, \( \delta^{13}\text{C}_{\text{DIC}} \) signatures were isotopically depleted as \( \text{CO}_2 \) was released as a result of aquatic community respiration. In contrast, \( \delta^{13}\text{C}_{\text{DIC}} \) at Masinga Dam and the Tana River mainstream showed minor fluctuations (\( \sim 0.8 \) and 0.9\%e, respectively) and did not exhibit systematic diurnal patterns.

Chania Stream showed strong diurnal cycles for both \( p\text{CO}_2 \) and \( \text{CO}_2 \) fluxes, with in-stream \( P \) and \( R \) as the major drivers as reflected by a strong negative correlation between \( DO \) and \( p\text{CO}_2 \) (Pearson correlation: \( p < 0.01, r^2 = 0.74; \) Fig. 11e). In contrast, \( p\text{CO}_2 \) at Masinga Dam (2621 to 3684 ppm) and Tana River mainstream (980 to 1131 ppm) was consistently above atmospheric equilibrium (385 ppm), and the two sites were sources of \( \text{CO}_2 \) to the atmosphere during the whole diurnal period (95 to 331 \text{ mmol CO}_2 \text{ m}^{-2} \text{ d}^{-1} \) at Masinga; 2 to 16 \text{ mmol CO}_2 \text{ m}^{-2} \text{ d}^{-1} \) in the lower Tana River). While the general observation is that in-stream \( P \) and \( R \) control biogeochemical processes in Chania Stream, it is worth mentioning that \( P \) is largely driven by periphyton as outlined below. We computed net community production (NCP), gross primary production (GPP) and respiration (\( R \)) at community level based on the diurnal changes of \( \text{O}_2 \), using an oxygen mass balance, the so-called Odum method (Odum, 1956). Accordingly, the hourly change of \( \text{O}_2 \) in surface waters during night-time corresponds to the hourly rate of community respiration (CR); the hourly change of \( \text{O}_2 \) in surface waters during daytime corresponds to the hourly rate of net community production. Assuming that the hourly rate of community respiration is constant through the daily cycle, the hourly rate of GPP is computed as NCP + CR. In order to account for the effect of changes of temperature on the solubility of \( \text{O}_2 \), we used apparent oxygen utilisation (AOU) night-time temporal changes to compute \( R \) during the 24 h cycle at Chania from the linear regression of AOU versus time. The change of AOU as a function of time was very linear (\( r^2 = 0.84 \)) and yielded an \( R \) value of 2.44 ± 0.33 \text{ mmol O}_2 \text{ L}^{-1} \text{ h}^{-1} \). The change of oxygen during daytime corresponds to the difference between gross primary production (GPP) and \( R \). If the rate of \( R \) during night-time is considered to be the same as during daytime, then this allows for GPP to be computed. The daytime temporal change of AOU computed by linear regression at Chania allowed for computation of a GPP value of 6.41 ± 3.92 \text{ mmol O}_2 \text{ L}^{-1} \text{ h}^{-1} \). The error on the GPP estimate is higher than on the \( R \) estimate because the AOU data showed more scatter during daytime (\( r^2 = 0.19 \)). The oxygen mass balance approach provides estimates of metabolic rates (GPP and \( R \)) at community level, integrating both planktonic and benthic compartments. Both GPP and \( R \) based on the oxygen mass balance have rates (6.41 and 2.44 \text{ mmol O}_2 \text{ L}^{-1} \text{ h}^{-1} \) that are one to three orders of magnitude higher than those derived from planktonic incubations (0.006 and 0.40 \text{ mmol O}_2 \text{ L}^{-1} \text{ h}^{-1} \). This can be due to the strong contribution to community metabolism of periphyton and benthic bacterial respiration.

The \( \text{CO}_2 \) flux measurements with floating domes in aquatic environments have been criticised in the past for a variety of possible biases (Raymond and Cole 2001), although there is evidence that these measurements provide reasonable estimates of k600 (Guérin et al., 2007; Vachon et al., 2010). The k600 values we obtained are consistent with those typically reported for streams (Wanninkhof et al., 1990), rivers (Raymond and Cole 2001) and lakes (Cole and Caraco, 1998). The higher k600 values at Chania Stream compared to Tana River fit the pattern described by Raymond et al. (2012) of k600 decreasing with increasing stream order. The higher k600 at Masinga Reservoir compared to the Tana River might be related to lower fetch limitation and more exposure leading to higher contribution of wind-induced turbulence to k600 (Borges and Abril, 2011).

5 Conclusions

While the wide range of \( \delta^{13}\text{C}_{\text{DIC}} \) values (−20.8 to −2.4\%e) in the Tana River basin suggests variable DIC sources, the primary process regulating of DIC concentrations is weathering, as clearly demonstrated by a strong correlation between TA and the sum of \( \text{Ca}^{2+} \) and \( \text{Mg}^{2+} \) concentration. However, Nyambene Hills subcatchment is lithologically distinct from the rest of the basin, as reflected by its systematically high TA during all seasons. Global syntheses suggest that streams are generally characterised by higher \( p\text{CO}_2 \) than large rivers, but our data show the opposite pattern with higher concentrations of \( \text{CO}_2 \) recorded at the Tana River mainstream compared to headwater streams. Chlorophyll \( a \) and primary production increased consistently downstream, contrary to what could be
expected from the RCC and from high TSM concentrations, due to favourable conditions for phytoplankton development such as high nutrients, temperature and water residence time. In-stream $R$ in the lower Tana River mainstream accounted for the bulk of total CO$_2$ evasion (77 to 114 %) to the atmosphere, while it could only account for 5 to 68 % of CO$_2$ evasion in headwaters, thus pointing toward greater contributions by other watershed processes or in-stream processes such as benthic respiration. Focusing on a depth-integrated primary production measurement in future is critical in order to give an accurate estimate of $P : R$ ratios for Tana River. The diurnal biogeochemical fluctuations were substantial in headwater streams, moderate at Masinga Reservoir and not detectable at Tana River mainstream Masinga Reservoir and Tana River mainstream. However, the timing for Masinga Reservoir sampling was unusual due to inflowing of the reservoir resulting in high water turbulence associated with high turbidity and vertical mixing of water, thus diurnal biogeochemical fluctuations at Masinga Dam were not representative of a typical situation. Hence, there is a need to focus on a more representative diurnal sampling in the reservoirs in future studies, particularly the during dry season, when phytoplankton production is expected to be significant.

Supplementary material related to this article is available online at http://www.biogeosciences.net/10/6911/2013/bg-10-6911-2013-supplement.pdf.

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