Coupling the chemical dynamics of carbonate and dissolved inorganic nitrogen systems in the eutrophic and turbid inner Changjiang (Yangtze River) Estuary

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Abstract

To better understand biogeochemical processes controlling CO$_2$ dynamics in those eutrophic large-river estuaries and coastal lagoons, we investigated surface water carbonate system, nutrients, and relevant hydrochemical parameters in the inner Changjiang (Yangtze River) Estuary, covering its channel-like South Branch and the lagoon-like North Branch, shortly after a spring-tide period in April 2010. In the North Branch, with a water residence time of more than 2 months, biogeochemical additions of ammonium (7.4 to 65.7 µmol kg$^{-1}$) and alkalinity (196 to 695 µmol kg$^{-1}$) were detected along with high salinity of 4.5 to 17.4. In the South Branch upper-reach, unusual salinity values of 0.20 to 0.67 were detected, indicating spillover waters from the North Branch. The spillover waters enhanced the springtime Changjiang export fluxes of nutrients, dissolved inorganic carbon, and alkalinity. And they affected the biogeochemistry in the South Branch, by lowering water-to-air CO$_2$ flux and continuing the nitrification reaction. In the North Branch, $p$CO$_2$ was measured from 930 to 1518 µatm at the salinity range between 8 and 16, which was substantially higher than the South Branch $p$CO$_2$ of 700 to 1100 µatm. Based on field data analyses and simplified stoichiometric equations, we suggest that the North Branch CO$_2$ productions were quantified by biogeochemical processes combining organic matter decomposition, nitrification, CaCO$_3$ dissolution, and acid-base reactions in the estuarine mixing zone. Although our study is subject to limited temporal and spatial coverage of sampling, we have demonstrated a procedure to quantificationally constrain net CO$_2$ productions in eutrophic estuaries and/or coastal lagoons, by coupling the chemical dynamics of carbonate and dissolved inorganic nitrogen systems.
1 Introduction

Large-river estuaries are important interfaces between continents and the oceans. They are biogeochemical hot spots since they receive large inputs of particulate matters, organic carbon, and nutrients from continents and oceans to support high rates of metabolism and/or chemical reactions. Significantly, CO$_2$ emission from estuaries, coastal lagoons, and salt marshes to the atmosphere has been proposed as an important component of the global carbon cycle (Frankignoulle et al., 1996, 1998; Cai and Wang, 1998). The global estuarine CO$_2$ emission rate has been estimated at 0.1 to 0.5 GtC yr$^{-1}$ (Borges, 2005; Borges et al., 2005; Cai, 2011; Chen et al., 2013). Also the chemical dynamics in large-river estuaries and their biogenic element fluxes have global and/or regional impacts on marine biogeochemistry (e.g., Justić et al., 1995; Bricker et al., 2008; Liu et al., 2015).

So far, mechanisms supporting the estuarine CO$_2$ emission rates need to be better understood, especially in those eutrophic large-river estuaries. Cai (2011) suggested that respiration processes may not occur in the large-river estuaries due to short water transit or residence times. However, many medium to large rivers have forked estuaries. For example, the inner estuary of the Yangtze River (Changjiang) is divided into two primary branches by the Chong-ming Island (Fig. 1). The Channel-like South Branch is subject to very low respiration rates (Zhai et al., 2007), while the lagoon-like North Branch shows substantial additions of free CO$_2$ and dissolved organic matters (Zheng et al., 2011; Guo et al., 2014). We contend that, if the estuarine areas have a sufficient water residence time (such as the North Branch of the inner Changjiang Estuary), they may function as sites of terrestrial carbon incineration.

In addition, biogeochemistry of dissolved inorganic carbon and nutrients in the North Branch are rarely studied. How the North Branch biogeochemistry sustains the estuarine CO$_2$ additions reported by Zheng et al. (2011) remains unknown. Furthermore, the North Branch is occupied by saline water rather than freshwater, especially in relatively dryer seasons (He et al., 2006; Zheng et al., 2011; Guo et al., 2014). According
to earlier results (e.g., Mao et al., 2001), the saltwater spillover from the North Branch usually occurs during spring-tide periods (with tidal ranges of more than 2.5 m) in winter and spring, with water discharge rates of lower than 25,000 m$^3$ s$^{-1}$ (from the Datong Station, ~400 km upstream of the study area). The saltwater spillover from the North Branch has received increasing attention since the occurrence of salty water in the South Branch threatens an important water source of Shanghai, the Qingcaosha Reservoir (e.g., Shen et al., 1980; Mao et al., 2001; He et al., 2006; Wu et al., 2006; Xue et al., 2009; Qiu and Zhu, 2013). However, the potential impacts of spillover signals from the North Branch (Fig. 1a) on the budgets of carbon and nitrogen in the South Branch are poorly understood.

To evaluate the impacts of the North Branch saltwater spillover on the South Branch biogeochemistry, we investigated carbonate system and nutrients in the inner Changjiang Estuary in April 2010, covering both of the two primary branches (Fig. 1), shortly after a spring tide period. Together with simultaneous dissolved oxygen (DO) and partial pressure of CO$_2$ ($p$CO$_2$) data, this dataset also provides an opportunity to quantificationally examine how coupled dynamics of nitrogen and carbon elements affect CO$_2$ degassing fluxes from this important large-river estuary to the atmosphere, which may help to understand CO$_2$ dynamics and the controlling processes in those eutrophic large-river estuaries and coastal lagoons of the world.

2 Materials and methods

2.1 Study area

Changjiang is the fourth largest river in the world by virtue of the water discharge of \( \sim 944 \times 10^9 \text{ m}^3 \text{ yr}^{-1} \) (Dai and Trenberth, 2002). It exports large amounts of nutrients, alkalinity and carbon to the adjacent East China Sea (ECS) continental shelf (e.g., Chen and Wang, 1999; Wang and Wang, 2006; Zhang et al., 2007), enhancing the coastal eutrophication and red tides there (e.g., Chen et al., 2003; Zhou et al., 2008).
Changjiang is also one of the most important solid transporting rivers (Gaillardet et al., 1999), although its downstream solid content has declined from ~ 600 mg L⁻¹ in the 1960s to < 400 mg L⁻¹ in the 2000s (Li and Zhang, 2003; Lin et al., 2007). Since its upstream water flows through the Yun–Gui Plateau and the Sichuan Basin, where the basement rocks are abundant in carbonates (Chen et al., 2002), the suspended sediment in the Changjiang River is rich of calcite (Chen et al., 2001). Also the Changjiang water is characterized by high alkalinity (Chen et al., 2002; Wu et al., 2007; Zhai et al., 2007) as compared with many other major rivers in the world.

The inner Changjiang Estuary is a large tidal estuary, complicated by bathymetry, islands, and deep channels (Xue et al., 2009). Nearly all of the river flow is transported via its South Branch into the ECS (Shen, 2001; Qu, 2010). The North Branch of the inner Changjiang Estuary is relatively shallow (water depth ~ 3 m) and open to navigation only at high tides in a day. At low tide periods, however, the North Branch is isolated from the Changjiang main stream.

In the outer Changjiang Estuary, a coastal current (i.e., Yellow Sea Coastal Current, YSCC for short) affects the hydrology and hydrochemistry during the northeast monsoon period from October to April in the following year (Fig. 1a; Chen, 2009). This current has higher dissolved inorganic carbon (DIC) and total alkalinity (TAlk) as compared with most of ECS surface waters (Zhai et al., 2014).

### 2.2 Survey design

During 2 to 7 April 2010, a sampling cruise was carried out in the inner Changjiang Estuary. Shortly before the cruise, the water discharge rates (from the Datong Station) increased from 17 600 m³ s⁻¹ in 27 March to 20 800 m³ s⁻¹ in 2 April (Fig. 2a). Therefore, this cruise represented a transitional period between the dry seasons (usually with the water discharge rate of less than 14 000 m³ s⁻¹, from December of the last year to February) and the flood seasons (usually with the water discharge rate of more than 40 000 m³ s⁻¹, from May to September). Overall, our cruise included three surveys/legs in the South Branch, one survey in the North Branch, and one survey in...
the outer Changjiang Estuary. The three South Branch surveys occupied the 3rd day, 6th day, and 7th day after the spring-tide day in 1 April (Fig. 2b). In the North Branch, two anchored stations were repeatedly sampled for 12 to 20 h against tide height and salinity variations.

2.3 Sampling and analyses

Using an underway pumping system similar to Zhai et al. (2005, 2007), surface water (at a depth of ~ 1 m) was pumped from a side intake for continuous measurements of hydrochemical parameters such as temperature, salinity, DO and \( p\text{CO}_2 \). Via a side vent of our pumping system, discrete samples for Winkler DO, pH, DIC, TAlk, nutrients and particulate matters were collected at selected sites (Fig. 1).

Surface water temperature and salinity were continuously determined with a pre-calibrated YSI® 6600 m, with the precisions of ±0.01 °C and ±0.01 salinity units. And the underway salinity data were validated by simultaneous discrete salinity data. Aqueous \( p\text{CO}_2 \) was continuously detected by a Li-Cor® non-dispersive infrared spectrometer (Li-7000) together with a continuous flow and fully sealed cylinder-type equilibrator (Zhai et al., 2007; Jiang et al., 2008). For calibration purposes, four CO\(_2\) gas standards with CO\(_2\) molecular fractions from 400 to 1510 µmol mol\(^{-1}\) were applied. The uncertainty of these standards was ~ 1%, which represents the maximum level of uncertainty during the period of extensive measuring of \( p\text{CO}_2 \) and data processing (see details in Zhai et al., 2005). The field-measured atmospheric CO\(_2\) data were corrected to survey-based barometric pressure at 10 m above the water surface and 100% humidity at water surface temperature and salinity, following the procedure described in Zhai et al. (2007).

For discrete salinity measurements, water samples were sealed in 140 mL high-density polyethylene bottles and kept at room temperature. They were determined in a week using a WTW TetraCon® 325 probe based on conductivity measurements, with a precision of ±0.1 salinity units (Yan et al., 2012). Winkler DO and NIST (National Insti-
stitute of Standards and Technology, USA)-traceable pH (at 15°C) were sampled and determined according to Zhai et al. (2007, 2012). The possible nitrite interference in the DO titration was removed by adding 0.01% NaN\textsubscript{3} during sample fixation (Wong, 2012). pH was measured on board using a precision pH meter and an Orion\textsuperscript{®} 8102BN Ross electrode, which were calibrated against three NIST-traceable buffers (pH = 4.00, 7.03, and 10.12 at 15.0°C, Thermo Fisher Scientific Inc., USA). The precisions of DO and pH data were ±0.5% DO and ±0.01 pH, respectively (Zhai et al., 2012). To express the oxygen consumption during water mixing, DO saturation was calculated from field-measured DO concentrations divided by DO concentration at equilibrium with the atmosphere. The latter was calculated as per the Benson and Krause (1984) equation.

For measurements of nutrient elements and TAlk, trinary water samples were filtered on board with 0.45 µm cellulose acetate membranes (Zhai et al., 2007; Yan et al., 2012). One of these was poisoned with 0.1% chloroform and preserved at 4°C for NH\textsubscript{4}\textsuperscript{+}–N (ammonium) and silicate determination (within a time frame of 10 days upon sampling). The second one was deep-frozen and kept at −20°C for NO\textsubscript{3}−–N (nitrate), NO\textsubscript{2}−–N (nitrite) and phosphate determination (within a time frame of 25 days upon sampling). The third one (for TAlk) was stored in a 140 mL high-density polyethylene bottle, immediately mixed with 50 µL of saturated HgCl\textsubscript{2}, and then sealed and preserved at room temperature until determination (within a time frame of 15 days upon sampling). Water samples for DIC were unfiltered but allowed to settle before measurement. They were stored in 60 mL borosilicate glass bottles (bubble free), and also preserved with 50 µL of saturated HgCl\textsubscript{2} and determined within a time frame of 7 days upon sampling.

Water samples for suspended particulate matter (SPM), particulate organic carbon (POC), and particulate inorganic carbon (PIC) were filtered on board with carbon-free 0.7 µm quartz microfiber (GFF) membranes. The SPM data were collected after drying in an oven at 50°C. And then the membranes were divided into two equal parcels. One of them was prepared for total particulate carbon measurements (without acid fuming), and another one was fumed with concentrated HCl so as to remove carbonate before POC determination. Both total particulate carbon and POC were measured according
to the JGOFS protocols (Knap et al., 1996), using a PE-2400 SERIES II CHNS/O analyzer. Finally, the PIC data were obtained from the difference between total particulate carbon and POC.

$\text{NO}_3^-$–N, $\text{NO}_2^-$–N, and phosphate were determined using an AA3 Auto-Analyzer (Bran + Luebbe Co., Germany), while $\text{NH}_4^+$–N and silicate were measured using a Tri-223 continuous Auto-Analyzer (see details in Yan et al., 2012). Briefly, $\text{NO}_3^-$–N and $\text{NO}_2^-$–N were measured by reducing $\text{NO}_3^-$ to $\text{NO}_2^-$ with a Cd column, and then determining $\text{NO}_2^-$ using the standard pink azo dye spectrophotometric method. Phosphate, silicate and $\text{NH}_4^+$–N were measured based on the standard phospo-molybdenum blue, silicon molybdenum blue and indophenol blue spectrophotometric procedures. Note that an appropriate quantity of more reagent NaOH was added during the $\text{NH}_4^+$–N measurement so as to make the final pH within the optimum range of 10.5 ± 0.1 (Pai et al., 2001; Tzollas et al., 2010).

Following Zhai et al. (2007), DIC was measured by infrared detection after acid extraction of a 0.5 mL sample with a Kloehn® digital syringe pump, while TAlk was determined by Gran acidimetric titration on a 25 mL sample with another Kloehn® digital syringe pump, using a precision pH meter and an Orion® 8102BN Ross electrode for detection. To ensure the measuring quality of DIC and TAlk, certificated reference materials from Andrew G. Dickson’s laboratory (Scripps Institute of Oceanography) were regularly checked at a precision of ±2 µmol kg$^{-1}$.

### 2.4 Flux estimation

Riverine element export fluxes were estimated based on the element concentrations at the river end, i.e. west side of the Chongming Island (Fig. 1a). Water discharge data needed were measured at the Datong Station, as released by the China Bureau of Hydrology (http://xxfb.hydroinfo.gov.cn/). The North Branch saltwater spillover induced element fluxes were estimated based on a mass balance approach. See Sects. 4.1 and 4.4 for details.
2.5 Calculating carbonate system parameters from DIC and TAlk

For the purpose of modeling calculations, aqueous $pCO_2$ and concentrations of bicarbonate and carbonate ions were calculated from the DIC, TAlk, silicate, phosphate, seawater temperature, and salinity values using the program CO2SYS.xls (Pelletier et al., 2011), an updated version of the original CO2SYS.EXE (Lewis and Wallace, 1998). The dissociation constants for carbonic acid were those determined by Millero et al. (2006), and the dissociation constant for the $HSO_4^-$ ion was determined as per Dickson (1990).

3 Results

3.1 Hydrological and particle backgrounds

During the cruise, water temperature ranged from 10.6 to 13.8°C. In the North Branch, water temperature was detected as low as 10.9±0.3°C, lower than the nearby offshore sea surface temperature observed in early spring (Zhai et al., 2014). The relatively high temperature of >13°C was only detected in the South Branch.

Salinity in the South Branch varied day by day (Fig. 3a). In the 3rd day after the spring-tide day, it ranged from 0.13 to 0.67, with a salinity peak at 121°22′ E (Fig. 3a), ~20 km downstream the west side of the Chong-ming Island. In the 6th day after the spring-tide day, it changed from 0.23 to 0.54, with a peak at ~40 km downstream the earlier salinity peak (Fig. 3a). These patterns were similar to results reported by He et al. (2006). In the 7th day after the spring-tide day, the salinity peak in the South Branch moved to downstream further, while salinity at the upstream and middle stations ranged from 0.14 to 0.16 (Fig. 3a). In the North Branch, however, very high salinity of 4.5 to 17.4 was determined in the surface water (Fig. 1b), showing the occupation of saline water (He et al., 2006; Zheng et al., 2011; Guo et al., 2014). East to the line of
122° E, surface salinity ranged from 0.23 to 18.4, showing the combined influences of Changjiang dilution and YSCC waters (Fig. 1a; Zhai et al., 2014).

The South Branch SPM ranged from 25 to 202 mg L\(^{-1}\). It was lower than earlier SPM values of 240 to 720 mg L\(^{-1}\) between 1960 and 2001 (Li and Zhang, 2003), but similar to recent SPM data of 20 to 190 mg L\(^{-1}\) between 2003 and 2005 (Lin et al., 2007). In the North Branch, however, surface water SPM was at very high levels of 390 to 455 mg L\(^{-1}\), with POC of 259 to 417 µmol L\(^{-1}\) and PIC of 253 to 293 µmol L\(^{-1}\). Particle related data are presented in the Appendix (Fig. A1) for public reference.

3.2 Distributions of hydrochemical parameters in the South Branch

In the South Branch, surface water DO saturation ranged from 86 to 93 % (Fig. 3b), similar to previous results reported by Zhai et al. (2007). In contrast, pH varied well and declined slightly from 3 April to 7 April (Fig. 3c). In 3rd day after the spring-tide day, pH showed a peak value of 8.06 at the site with the highest salinity. In 6th day after the spring-tide day, although the salinity varied from 0.23 to 0.54, pH ranged smoothly from 7.88 to 7.94. During the last survey in 7 April, pH ranged from 7.75 to 7.88.

\(\text{NH}_4^+ - \text{N}\) ranged from 5.0 to 25.2 µmol kg\(^{-1}\) (Fig. 3d), while \(\text{NO}_2^- - \text{N}\) declined from 4.05–4.36 µmol kg\(^{-1}\) at our upstream stations to 2.03 µmol kg\(^{-1}\) at the river mouth stations (Fig. 3e). As the products from chemical fertilizer application and soil erosion in the drainage basin of Changjiang, riverine \(\text{NO}_3^- - \text{N}\) ranged from 126.5 to 139.9 µmol kg\(^{-1}\) (Fig. 3f). The concentration of \(\text{NO}_3^- - \text{N}\) was one or two magnitudes higher than those of \(\text{NH}_4^+ - \text{N}\) and \(\text{NO}_2^- - \text{N}\). To sum up, the concentration of dissolved inorganic nitrogen (DIN, the sum of concentrations of \(\text{NH}_4^+ - \text{N}\), \(\text{NO}_2^- - \text{N}\), and \(\text{NO}_3^- - \text{N}\)) ranged from 142.2 to 167.3 µmol kg\(^{-1}\) (Fig. 3g). Except for a sampling site presumably influenced by a sewage outlet (Chai et al., 2006), the distributions of the two major DIN species (\(\text{NH}_4^+ - \text{N}\) and \(\text{NO}_3^- - \text{N}\)) in the surface water roughly reflected the salinity variation patterns (Fig. 3). Considering the above-mentioned salty water mass, most \(\text{NH}_4^+ - \text{N}\) data in 3 April were higher than those in 6 April (Fig. 3d), while most \(\text{NO}_3^- - \text{N}\)
data showed an inverse trend (Fig. 3f). DIN in this water mass slightly declined during the three days from 3 April to 6 April (Fig. 3g).

As a particle-concentrated element (e.g., Das et al., 2006; Leote et al., 2013), phosphate was measured at relatively low levels from 0.87 to 1.63 µmol kg\(^{-1}\) (Fig. 3h). However, silicate was at relatively high levels from 99.6 to 104.9 µmol kg\(^{-1}\) (Fig. 3i).

Except for the sampling site presumably influenced by sewage, the two carbonate system parameters (i.e. TAlk and DIC) also reflected the salinity variation patterns (Fig. 3l and m). TAlk ranged from 1495 to 1694 µmol kg\(^{-1}\), while DIC ranged from 1520 to 1654 µmol kg\(^{-1}\).

The South Branch \(p\text{CO}_2\) ranged from 628 to 1148 µatm (Fig. 3n). Its distribution patterns roughly mirrored the distributions of pH (Fig. 3). Similar to earlier results reported by Zhai et al. (2007) and Chen et al. (2008), \(p\text{CO}_2\) was always higher than the water-air equilibration level of 398 to 411 µatm (Fig. 3n). Hydrochemical parameters against salinity in the South Branch are plotted in the Appendix (Fig. A2) for public reference.

### 3.3 Water mixing behaviors of hydrochemical parameters in the North Branch and the outer estuary area

In the North Branch, surface water DO saturation ranged from 83 to 95\% (Fig. 4a). Low DO saturation values of < 86\% were mostly observed at the middle salinity from 8.4 to 12.4. Correspondingly, significant additions of \(p\text{CO}_2\) and \(\text{NH}_4^+\text{--N}\) were detected (Fig. 4). And the associated pH was measured at relatively low level from 7.90 to 7.92 (Fig. 4b). \(\text{NO}_2^-\text{--N}, \text{NO}_3^-\text{--N}\) and DIN also showed additions (Fig. 4e–g). However, silicate behaved like a conservative element (Fig. 4i), with a very tight linear relationship against salinity (\(n = 13, r = 0.9993\)).

In the outer estuary area, only two stations were sampled for nutrient analyses. Silicate data in both the outer area and the South Branch followed the water mixing line obtained from the North Branch (Fig. 4i), suggesting that water mixing in the outer area was controlled by nearly the same end-members as those in the North Branch. Therefore, we assumed conservative water mixing lines of dissolved nitrogen species in the
Changjiang Estuary (Fig. 4). They were:

\[
\begin{align*}
\text{NH}_4^+ - N_{\text{conservative}} \, (\mu \text{mol kg}^{-1}) &= -0.3041 \times \text{Salinity} + 12.5 \\
\text{NO}_2^- - N_{\text{conservative}} \, (\mu \text{mol kg}^{-1}) &= -0.1163 \times \text{Salinity} + 2.5 \\
\text{NO}_3^- - N_{\text{conservative}} \, (\mu \text{mol kg}^{-1}) &= -4.1905 \times \text{Salinity} + 135 \\
\text{DIN}_{\text{conservative}} \, (\mu \text{mol kg}^{-1}) &= -4.6667 \times \text{Salinity} + 150
\end{align*}
\]

where seawater end-members of NO$_3^-$-N (≈ 3 µmol kg$^{-1}$ at a salinity of 31.5) and DIN (≈ 10 µmol kg$^{-1}$ at a salinity of 30) referred to spring YSCC values reported by Chen et al. (2009) and Wang et al. (2003).

TAlk and DIC also showed additions in the North Branch, while they behaved like conservative elements in the outer area (Fig. 5). In the outer area, the simplified water mixing lines of TAlk and DIC, with seawater end-members of them from YSCC obtained in April 2007 (Zhai et al., 2014), were as follow:

\[
\begin{align*}
\text{TAlk}_{\text{conservative}} \, (\mu \text{mol kg}^{-1}) &= 22.068 \times \text{Salinity} + 1580 \\
\text{DIC}_{\text{conservative}} \, (\mu \text{mol kg}^{-1}) &= 14.551 \times \text{Salinity} + 1600
\end{align*}
\]

Most of field-measured TAlk and DIC values in the outer area satisfactorily followed Eqs. (5) and (6) (Fig. 5), giving confidence in the water mixing line reduction. The intercepts of Eqs. (5) and (6) were consistent with TAlk and DIC values at the river mouth (1595 to 1615 µmol kg$^{-1}$), and slightly higher than TAlk and DIC values at the river end (1495 to 1585 µmol kg$^{-1}$) (Fig. 3). However, they were ≈ 200 µmol kg$^{-1}$ lower than the freshwater end-member values of TAlk and DIC in spring of 2006 and 2007 (≈ 1800 µmol kg$^{-1}$) (Zhai et al., 2007, 2014).

The assumed conservative mixing line of $p$CO$_2$ (Fig. 4c) was calculated from conservative TAlk (Eq. 5), DIC (Eq. 6), silicate (Fig. 4i), and a fixed phosphate of 1.4 µmol kg$^{-1}$ (Fig. 4h). It was a sunken line, with an oversaturated $p$CO$_2$ value of 596 µatm at salinity 1 and undersaturated $p$CO$_2$ value of 325 µatm at salinity 20. The freshwater end-member value was similar to field-measured $p$CO$_2$ at the river mouth (Fig. 4c), while
the relatively high-salinity value was consistent with the earlier reported YSCC $pCO_2$ range of 310 to 377 µatm obtained in April 2007 (Zhai et al., 2014). To better evaluate the biogeochemical additions/removals of nitrogen and carbon elements, we calculated the conservative concentrations of DIN species and carbonate system parameters at salinity, according to Eqs. (1)–(6). And then we obtained the differences between the measured concentrations and the conservative concentrations along the mixing lines, i.e., $\Delta NH_4^+ - N$, $\Delta NO_2^- - N$, $\Delta NO_3^- - N$, $\Delta DIN$, $\Delta TAlk$ and $\Delta DIC$ in Eqs. (7)–(12), with positive values indicating biogeochemical additions and negative values indicating removals. Note that both of the additions and removals are relative to the conservative mixing between the Changjiang freshwater end-member near the river mouth and the spring YSCC surface water end-member.

$$\Delta NH_4^+ - N = [NH_4^+ - N] - [NH_4^+ - N\ \text{conservative}]$$  (7)

$$\Delta NO_2^- - N = [NO_2^- - N] - [NO_2^- - N\ \text{conservative}]$$  (8)

$$\Delta NO_3^- - N = [NO_3^- - N] - [NO_3^- - N\ \text{conservative}]$$  (9)

$$\Delta DIN = ([NH_4^+ - N] + [NO_2^- - N] + [NO_3^- - N]) - DIN\ \text{conservative}$$  (10)

$$\Delta TAlk = TAlk - TAlk\ \text{conservative}$$  (11)

$$\Delta DIC = DIC - DIC\ \text{conservative}$$  (12)

Figure 6 summarizes $\Delta NH_4^+ - N$, $\Delta NO_2^- - N$, $\Delta NO_3^- - N$, $\Delta DIN$, $\Delta DIC$ and $\Delta TAlk$ against salinity in the North Branch. During our survey, biogeochemical additions of DIN species and carbonate system parameters were estimated at 7.4 to 65.7 µmol kg$^{-1}$ ($\Delta NH_4^+ - N$), 1.24 to 16.15 µmol kg$^{-1}$ ($\Delta NO_2^- - N$), 6.3 to 26.0 µmol kg$^{-1}$ ($\Delta NO_3^- - N$), 27 to 103 µmol kg$^{-1}$ ($\Delta DIN$), 243 to 753 µmol kg$^{-1}$ ($\Delta DIC$), and 196 to 695 µmol kg$^{-1}$ ($\Delta TAlk$). Except for $\Delta NO_3^- - N$, most of those biogeochemical additions were obtained at salinity between 8 and 16 (Fig. 6), suggesting the associated biogeochemical processes mainly occurred at the middle salinity area.
4 Discussion

4.1 Residence time estimation

Estuaries are not only transport passages of terrestrial materials from the continents to the oceans, but also chemical reactors and/or buffers (Officer, 1979). Before discussing biogeochemical processes and element transport fluxes in the inner Changjiang Estuary, the residence times need to be examined. To estimate the residence times, we balanced water and salt budgets under the condition of the North Branch saltwater spillover shortly after the spring-tide period:

\[ Q_S = Q_R + Q_N \]  
\[ Q_S \times S_S = Q_R \times S_R + Q_N \times S_N \]

where \( Q_S, Q_R \) and \( Q_N \) are water discharges from the South Branch, from the upper river, and the mean water flux of the North Branch saltwater spillover during the spring-tide period in early April 2010 (Fig. 1a); \( S_S, S_R \) and \( S_N \) are mean salinity of the South Branch water, the Changjiang River water and the North Branch water. In this study, \( Q_R \) and \( S_R \) were 20 000 m\(^3\) s\(^{-1}\) and 0.14, respectively (Figs. 2a and 3a). \( S_S \) and \( S_N \) were estimated at 0.30 and 15.26, based on a compilation of our underway data obtained from 2 to 5 April (data partially reported in Figs. 1b and 3a). Therefore, \( Q_S \) and \( Q_N \) were estimated at 20 214 and 214 m\(^3\) s\(^{-1}\), respectively (Table 1).

Based on Shen (2001), Meng and Cheng (2005), the navigation map (2010 edition), and our field records, mean water depths of the South Branch (\( d_S \)) and the North Branch (\( d_N \)) were estimated at 8 and 3 m, respectively, while water areas of the South Branch (\( A_S \)) and the North Branch (\( A_N \)) were 900 and 422 km\(^2\), respectively (Table 1). Therefore, the mean residence times of the North Branch water (\( \tau_N \)) and the South
Branch water \((\tau_S)\) were estimated as bellow:

\[
\tau_N = A_N \times d_N/Q_N = 5.9 \times 10^6 \text{s} = 68 \text{day} \tag{15}
\]

\[
\tau_S = A_S \times d_S/Q_S = 0.36 \times 10^6 \text{s} = 4 \text{day} \tag{16}
\]

We must point out that Eq. (16) ignored the tidal effect. That is, effluent water particles across the interface between inner and outer estuaries can return to the inner estuary (i.e., the South Branch) during the next tide cycle, leading to a longer exposure time of water particles than the steady-state residence time (Monsen et al., 2002). Based on Fig. 3a, the North Branch saltwater spillover (presumably in 1 April) induced salinity peak in the South Branch moved from 121°22′ E in 3 April to 121°50′ E in 6 April. Even in 7 April, the salinity peak was still observable inside the river mouth (west of 122°00′ E) (Fig. 3a). Thus the South Branch water was exposed to the estuarine biogeochemical processes for \(~7\) days in spring, more than 1.5 times of the residence time estimated under the steady-state assumption via Eq. (16).

Similar to the South Branch, the exposure time of the North Branch water estimated via the steady-state Eq. (15) is also subject to uncertainties. However, Eq. (15) suggested that the North Branch water had a very long residence time of more than 2 months in spring. It provided reaction times for many biogeochemical processes to function.

### 4.2 Maintaining mechanisms of the biogeochemical additions in the North Branch

In the North Branch, DIC and TAlk were linearly correlated \((n = 13, r = 0.996)\), with a slope of 0.9888 that was very close to 1 (Fig. 7a). Since DIC is the sum of free \(\text{CO}_2\), \(\text{HCO}_3^-\) and \(\text{CO}_3^{2-}\), and \(\text{TAlk} = [\text{HCO}_3^-] + 2[\text{CO}_3^{2-}] + [B(\text{OH})_4^-] + [\text{OH}^-] - [\text{H}^+] + \text{other trace species}\), Fig. 7a suggested that both DIC and TAlk in the North Branch were mainly composed of \(\text{HCO}_3^-\) ion. Note that this pattern was much different from the usual DIC vs. TAlk plot (with a slope lower than 0.5) as driven by water mixing between the Changjiang freshwater and ECS seawaters (Zhai et al., 2007).
ΔTAlk and ΔDIN were also linearly correlated in the North Branch \( n = 13, r = 0.985 \), with a slope of 6.56 (Fig. 7b). This plot suggested that both the carbonate system and dissolved inorganic nitrogen dynamics in the North Branch might controlled by same biogeochemical processes. To examine these processes from a quantitative point of view, we analyzed the stoichiometric relationship of the Redfield respiration. Also resolved were the nitrification, the CaCO\(_3\) dissolution, and the acid–base reaction during the oxidation of organic matters.

In a saline aquatic environment with abundant oxygen and the pH of ~ 8, the Redfield respiration of biogenic organic matters is directly associated with a release of NH\(_4^+\)–N:

\[
(CH_2O)_{106}(NH_3)_{16}H_3PO_4 + 106O_2 \rightarrow 92CO_2 + 16NH_4^+ + 14HCO_3^- + HPO_4^{2-} + 92H_2O \quad (\text{R1})
\]

As compared with the decomposition of organic matters, the oxidation of NH\(_4^+\)–N (a key step of nitrification) is a time-expensive reaction (e.g., Dai et al., 2008), especially when the temperature is lower than 15°C. The nitrification occurs as below:

\[
2NH_4^+ + 3O_2 + 4HCO_3^- \rightarrow 2NO_2^- + 4CO_2 + 6H_2O \quad (\text{R2})
\]
\[
2NO_2^- + O_2 \rightarrow 2NO_3^- \quad (\text{R3})
\]

Combining Reactions (R1)–(R3), a complete Redfield respiration of biogenic organic matters is:

\[
(CH_2O)_{106}(NH_3)_{16}H_3PO_4 + 138O_2 + 18HCO_3^- \rightarrow 124CO_2 + 16NO_3^- + HPO_4^{2-} + 140H_2O \quad (\text{R4})
\]

In the North Branch, these processes were associated with the consumption of DO and the releases of free CO\(_2\) and the three dissolved inorganic nitrogen species (Fig. 4). To reveal how many NH\(_4^+\)–N released through respiration were transformed into NO\(_2^-\)–N and/or NO\(_3^-\)–N, we plotted ΔNH\(_4^+\)–N vs. ΔDIN (Fig. 7c). The North Branch ΔNH\(_4^+\)–N and ΔDIN were linearly correlated \( n = 13, r = 0.94 \), with a slope of 0.63. This slope...
suggested that Reactions (R2) and (R3) may only transform \((1 - 0.63) \times 100\% = 37\%\) of respiration-induced DIN additions into \(\text{NO}_2^- - \text{N}\) and/or \(\text{NO}_3^- - \text{N}\). Combining Reactions (R1) and (R4) based on this ratio, ignoring the incomplete nitrite oxidation, the local status of Redfield respiration in the North Branch was better characterized as below:

\[
(\text{CH}_2\text{O})_{106}(\text{NH}_3)_{16}\text{H}_3\text{PO}_4 + 118\text{O}_2 \rightarrow 104\text{CO}_2 + 10\text{NH}_4^+ + 6\text{NO}_3^- + 2\text{HCO}_3^- + \text{HPO}_4^{2-} + 110\text{H}_2\text{O} \quad (R5)
\]

Furthermore, the North Branch was an environment with abundant supplies of PIC (mostly calcite or \(\text{CaCO}_3\)) and seawater (having the \(\text{CO}_3^{2-}\) ion). Besides \(\text{CO}_2\) degassing, most of the above-mentioned estuarine \(\text{CO}_2\) productions are potentially removed by \(\text{CaCO}_3\) dissolution (Abril et al., 2003) and/or \(\text{CO}_3^{2-}\) titration:

\[
\text{CO}_2 + \text{CaCO}_3 + \text{H}_2\text{O} \rightarrow \text{Ca}^{2+} + 2\text{HCO}_3^- \quad (R6)
\]

\[
\text{CO}_2 + \text{CO}_3^{2-} + \text{H}_2\text{O} \rightarrow 2\text{HCO}_3^- \quad (R7)
\]

Except for the \(\text{CO}_3^{2-}\) titration (Reaction R7), all of respiration (Reaction R5), nitrification (Reaction R2) and \(\text{CaCO}_3\) dissolution (Reaction R6) affected the ratio of \(\Delta\text{DIC}\) to \(\Delta\text{TAlk}\) (Fig. 7d). In the North Branch, both \(\Delta\text{DIC}\) and \(\Delta\text{TAlk}\) were positive, suggesting net biogeochemical additions. \(\Delta\text{DIC} = 1.09 \times \Delta\text{TAlk} (n = 13, r = 0.992)\). The slope of 1.09 (higher than the \(\text{CaCO}_3\) dissolution induced slope of 0.5) suggested that respiration and \(\text{CaCO}_3\) dissolution controlled the North Branch carbonate system as below:

\[
(\text{CH}_2\text{O})_{106}(\text{NH}_3)_{16}\text{H}_3\text{PO}_4 + 118\text{O}_2 + 88\text{CaCO}_3(s) \rightarrow 16\text{CO}_2 + 88\text{Ca}^{2+} + 178\text{HCO}_3^- + 10\text{NH}_4^+ + 6\text{NO}_3^- + \text{HPO}_4^{2-} + 22\text{H}_2\text{O} \quad (R8)
\]

In the South Branch, however, most \(\Delta\text{DIC}\) and \(\Delta\text{TAlk}\) were negative, suggesting biogeochemical removals. \(\Delta\text{DIC} = 0.66 \times \Delta\text{TAlk} - 21 \mu\text{mol kg}^{-1}\) (Fig. 7d). The slope of 0.66 together with the negative \(\Delta\text{DIC}\) and \(\Delta\text{TAlk}\) values suggested that both \(\text{CO}_2\) degassing
and nitrification controlled the South Branch carbonate system (Fig. 7d). The later process was also evidenced by the above-mentioned dissolved nitrogen system dynamics in Fig. 3. Although CaCO$_3$ dissolution may also occur in the South Branch, as indicated by the facts that several ΔDIC and ΔTAlk plots in the South Branch were positive, and the slope of 0.66 was similar to the CaCO$_3$ dissolution induced slope of 0.5 (Fig. 7c), its effects on the South Branch carbonate system were negligible (Zhai et al., 2007).

In summary, the biogeochemical additions of NH$_4^+$–N and carbonate observed in the North Branch were driven by the biogeochemical processes combining organic matter decomposition, nitrification, and CaCO$_3$ dissolution, as described by Reaction (R8). Abundant organic matters and CaCO$_3$ particles, together with the quite long residence time and suitable environmental conditions such as temperature and oxygen, had made the North Branch into a natural reactor. This big reactor affected both element export fluxes (via its water spillover, see Sect. 4.4) and the water-to-air CO$_2$ flux in this important estuary (via increasing aqueous pCO$_2$, see Sect. 4.3).

### 4.3 Estuarine CO$_2$ dynamics in the North Branch

Reaction (R8) suggested that, if most of estuarine CO$_2$ products from Redfield respirations are removed by CaCO$_3$ dissolution, the ratio of ΔTAlk to ΔDIN should be 176/16 = 11.125. Alternatively, if none of the CO$_2$ products is removed by the CaCO$_3$ dissolution, the ratio of ΔTAlk to ΔDIN should be 2/16 = 0.125 based on Reaction (R5). Therefore, the real ratio of ΔTAlk to ΔDIN of 6.56 (Fig. 7b) suggested that only 6.56/(11.125 – 0.125) x 100% = 60% of respiration-induced free CO$_2$ was removed via CaCO$_3$ dissolution. Ignoring the minor impacts of CO$_2$ degassing fluxes (due to the much slower air-water exchanging rates as compared with the acid–base reactions), the other 40% of estuarine CO$_2$ products were potentially titrated by CO$_3^{2-}$ ion supplied by the seawater end-member, as indicated by Reaction (R7).

To discuss this effect, we calculated $pCO_2^{\text{addition}}$ in the North Branch via $pCO_2$ minus $pCO_2^{\text{conservative}}$, based on Fig. 4c. We then calculated concentrations of the additional
free CO₂, using \([\text{CO}_2^\ast]_{\text{addition}} = K_H \times p\text{CO}_2\) \(^\text{addition}\), where \(K_H\) is the solubility coefficient of CO₂, calculated via the Weiss (1974) equation. Figure 8a showed that \([\text{CO}_2^\ast]_{\text{addition}}\) ranged from 2.7 to 53 \(\mu\text{mol kg}^{-1}\) in the North Branch, usually lower than those predicted values based on Reaction (R8) and our \(\Delta\text{DIN}\) data. The differences were likely caused by acid-base titration between respiration-induced free CO₂ and seawater-introduced \(\text{CO}_2^2\) ion.

Figure 8b showed that conservative concentrations of non-carbonate alkalinity (such as borate ions) ranged from 7 \(\mu\text{mol kg}^{-1}\) at salinity 4 to 30 \(\mu\text{mol kg}^{-1}\) at salinity 16. These non-carbonate alkalinity values primarily accounted for the observed differences between \(\text{TAlk}\) and DIC in the North Branch, i.e. 22 to 29 \(\mu\text{mol kg}^{-1}\) as suggested by Fig. 7a. This comparison provided another evidence supporting the idea that seawater-introduced \([\text{CO}_3^2\text{-}]_{\text{conservative}}\) was mostly titrated by respiration-induced CO₂, and transferred into \(\text{HCO}_3^-\) ions.

Furthermore, we regarded the conservative concentrations of \(\text{CO}_3^2\text{-}\) ion (Fig. 8b) as the maximum removals via \(\text{CO}_3^2\text{-}\) titration. Figure 8a showed that the predicted values via DIN minus \([\text{CO}_3^2\text{-}]_{\text{conservative}}\) were highly consistent with the field-measured values of \([\text{CO}_2^\ast]_{\text{addition}}\) at salinity of 5 to 13, indicating that seawater-introduced \([\text{CO}_3^2\text{-}]_{\text{conservative}}\) was fully titrated by respiration-induced CO₂. However, the predicted values were lower than the real at salinity of > 13, presumably due to the overestimate of \(\text{CO}_3^2\text{-}\) titration reactions there.

In summary, we have quantificationally demonstrated that the observed aqueous \(p\text{CO}_2\) in the North Branch was determined by water mixing and several biogeochemical processes. Although 80 to 85% of estuarine CO₂ production from organic matter decomposition and nitrification had been removed by \(\text{CaCO}_3\) dissolution (~60%) and \(\text{CO}_3^2\text{-}\) titration (50 to 60% of the residuals, Fig. 8), the North Branch acted as a significant source area of the atmospheric CO₂ with higher \(p\text{CO}_2\) than the South Branch (Fig. 4c). To determine the seasonality and magnitude of the North Branch water-to-air CO₂ flux, more investigations are needed.
4.4 Spillover fluxes of TAlk, DIC, and nutrients from the North Branch into the Changjiang mainstream

The North Branch biogeochemical signals intrude to Changjiang mainstream (the South Branch) via the tide induced saltwater spillover process in dry seasons (Fig. 1a), potentially increasing element export fluxes to the ECS. Considering the steady state spillover flux of saline water (∼ 214 m³ s⁻¹, Table 1) from the North Branch into the Changjiang mainstream, the spillover flux of salt was estimated at 3265 kg s⁻¹, slightly higher than the riverine transport salt flux (Table 2). Similar to the salt flux estimation, riverine transport fluxes and spillover fluxes of TAlk, DIC, and nutrients were calculated (Table 2).

Generally, spillover fluxes of TAlk, DIC, and nutrients from the North Branch into the Changjiang mainstream were minor, as compared with the Changjiang transport fluxes. However, the springtime areal yield rates of TAlk, DIC, and DIN species from the North Branch were 34 to 133 times higher than those from the Changjiang drainage basin, as Changjiang has a drainage area of 1.8 × 10⁶ km² (Chen et al., 2002), roughly 4000 times of the North Branch water area. Note that the organic matters and CaCO₃ particles supporting the springtime TAlk, DIC, and DIN production in the North Branch were also supplied by Changjiang during its flood seasons. Therefore, the North Branch may serve as an estuarine chemical buffer, to some extent smoothing seasonal variations of Changjiang element export fluxes.

During our sampling period, the spillover of TAlk, DIC, and DIN species from the North Branch affected the biogeochemistry in the South Branch. Firstly, the spillover water transported more TAlk than DIC into the South Branch (Table 2), leading to the South Branch pCO₂ decline from ∼ 1000 µatm at salinity of ∼ 0.14 (with the water-to-air pCO₂ difference of ∼ 600 µatm) to ∼ 700 µatm at salinity of > 0.4 (with the water-to-air pCO₂ difference of ∼ 300 µatm) (Fig. 3n). If the air-water gas transfer velocity was changeless, the 50% decline of water-to-air pCO₂ difference means that the water-to-air CO₂ degassing flux had decreased by 50% due to the spillover of salty water.
Secondly, the spillover water likely contained active nitrifiers, which continued the nitrification reaction in the South Branch, as indicated by the $\text{NH}_4^+\text{-N}$ decline and the $\text{NO}_3^-\text{-N}$ increase of the salty water mass from 3 to 6 April (Fig. 3). This was also evidenced by the pH decline and the $p\text{CO}_2$ increase signals (at salinity of $>0.4$) during the same period (Fig. 3), as shown by Reaction (R2).

5  Concluding remarks

This study showed that the spillover waters from the North Branch had minor contributions to the Changjiang transport fluxes of nutrients, dissolved inorganic carbon, and alkalinity. They affected the biogeochemistry in the South Branch, by lowering water-to-air $\text{CO}_2$ fluxes and continuing the nitrification. Significantly, several primary biogeochemical processes such as organic matter decomposition, nitrification, $\text{CaCO}_3$ dissolution, and acid-base reaction occurred in the North Branch of the Changjiang Estuary, leading to the unusual high $p\text{CO}_2$ values at middle salinity and the biogeochemical additions of dissolved inorganic nitrogen species and carbonate system parameters. Similar biogeochemical processes may occur in many eutrophic estuaries and/or coastal lagoons of the world. This study demonstrated a procedure to quantitatively analyze the coupled dynamics of dissolved inorganic nitrogen and carbonate systems, which may help to better understand the combined effects of metabolic processes and chemical reactions on $\text{CO}_2$ fluxes in estuaries and/or coastal lagoons with similar physical and biogeochemical conditions.

Appendix A: Distributions of surface particulate parameters

Particulate samples were collected at selected sites during our April 2010 cruise. Their suspended particulate matter concentrations, particulate organic carbon data, and particulate inorganic carbon data were presented against longitude (Fig. A1). Usually the North Branch had more SPM, POC, and PIC concentrations than the South Branch.
Appendix B: Hydrochemical parameters vs. salinity in the south Branch

Despite the very low salinity, we plotted those ammonium, nitrate, dissolved inorganic nitrogen, partial pressure of CO$_2$, total alkalinity, and dissolved inorganic carbon data obtained in the South Branch against salinity (Fig. A2). From 3 to 6 April, we obtained a decline of ammonium and increases of nitrate and partial pressure of CO$_2$ in the same salinity range from 0.4 to 0.6, indicating nitrification.

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References


Coupling the chemical dynamics of carbonate and dissolved inorganic nitrogen systems

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Coupling the chemical dynamics of carbonate and dissolved inorganic nitrogen systems

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Table 1. Summary of hydrological characteristics in the inner Changjiang Estuary during the spring-tide period in early Apr 2010. See the text for details.

<table>
<thead>
<tr>
<th></th>
<th>Water discharge (m$^3$s$^{-1}$)</th>
<th>Salinity</th>
<th>Area (km$^2$)</th>
<th>Depth (m)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Upper river</td>
<td>$Q_R = 20000$</td>
<td>$S_R = 0.14$</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>South Branch</td>
<td>$Q_S = 20214$</td>
<td>$S_S = 0.30$</td>
<td>$A_S = 900$</td>
<td>$d_S = 8$</td>
</tr>
<tr>
<td>North Branch</td>
<td>$Q_N = 214$</td>
<td>$S_N = 15.26$</td>
<td>$A_N = 422$</td>
<td>$d_N = 3$</td>
</tr>
</tbody>
</table>
Table 2. A steady-state estimation of Changjiang transport fluxes and the North Branch (NB) spillover fluxes of salt, TAlk, DIC, and nutrients during a spring-tide period in Apr 2010. See the text for details.

<table>
<thead>
<tr>
<th>Chemical species</th>
<th>Changjiang transport</th>
<th>Spillover from the NB</th>
<th>% added by the NB</th>
</tr>
</thead>
<tbody>
<tr>
<td>Water flux (m³ s⁻¹)</td>
<td>20 000ᵃ</td>
<td>214ᵃ</td>
<td>1.1 %</td>
</tr>
<tr>
<td>Salt flux (kg s⁻¹)</td>
<td>0.14ᵃ × 20 000 = 2800</td>
<td>15.26ᵃ × 214 = 3265</td>
<td>116 %</td>
</tr>
<tr>
<td>TAlk flux (mol s⁻¹)</td>
<td>1530ᵇ × 20 000/1000 = 30 600</td>
<td>(2336ᶜ × 214)/1000 = 500</td>
<td>1.6 %</td>
</tr>
<tr>
<td>DIC flux (mol s⁻¹)</td>
<td>1550ᵇ × 20 000/1000 = 31 000</td>
<td>(2310ᶜ × 214)/1000 = 494</td>
<td>1.6 %</td>
</tr>
<tr>
<td>NH₄⁺−N flux (mol s⁻¹)</td>
<td>18ᵇ × 20 000/1000 = 360</td>
<td>(55ᶜ × 214)/1000 = 12</td>
<td>3.3 %</td>
</tr>
<tr>
<td>NO₂⁻−N flux (mol s⁻¹)</td>
<td>4.2ᵇ × 20 000/1000 = 84</td>
<td>(11ᶜ × 214)/1000 = 2.4</td>
<td>2.9 %</td>
</tr>
<tr>
<td>NO₃⁻−N flux (mol s⁻¹)</td>
<td>128ᵇ × 20 000/1000 = 2560</td>
<td>(102ᶜ × 214)/1000 = 22</td>
<td>0.9 %</td>
</tr>
<tr>
<td>DIN flux (mol s⁻¹)</td>
<td>150ᵇ × 20 000/1000 = 3000</td>
<td>(168ᶜ × 214)/1000 = 36</td>
<td>1.2 %</td>
</tr>
<tr>
<td>Phosphate flux (mol s⁻¹)</td>
<td>1.3ᵇ × 20 000/1000 = 26</td>
<td>(1.3ᶜ × 214)/1000 = 0.28</td>
<td>1.1 %</td>
</tr>
<tr>
<td>Silicate flux (mol s⁻¹)</td>
<td>102ᵇ × 20 000/1000 = 2040</td>
<td>(67ᶜ × 214)/1000 = 14</td>
<td>0.7 %</td>
</tr>
</tbody>
</table>

ᵃ Refer to Table 1.
ᵇ Element concentrations at the river end. Refer to Fig. 3.
ᶜ Averaged element concentrations in the North Branch. Refer to Figs. 4 and 5.
Figure 1. Area map (a) and surface distributions of salinity, total alkalinity and ammonia during the April 2010 cruise (b–d). Following Zhai et al. (2007), we regarded the line of 122°E as the boundary between inner sub-estuaries and the outer estuary area. ECS = East China Sea; CM = Chong-ming Island; YSCC = Yellow Sea Coastal Current. As sketched in (a), the inner Changjiang Estuary is geographically divided into three sub-estuaries by the Chong-ming Island and several sandbanks, i.e. the North Branch (I), the north channel of the South Branch (II), and the south channel of the South Branch (III). Four neighboring sampling sites in April 2007 (Zhai et al., 2014) were sketched in (a) by “+” symbols. The pentacle shows a reference station for the tidal cycle.
Figure 2. Water discharges from the Changjiang River (a) and the tidal cycles (b). Grey bars represent survey periods. The arrow denotes the highest tide time in 1 April, i.e. the spring-tide day. The reference station for the tidal cycle was at 31°27′42″ N and 121°24′42″ E, as sketched in Fig. 1a. SB denotes our South Branch surveys, while NB shows the North Branch survey. “Outer” means our outer estuary survey east to 122° E.
Figure 3. Evolution of surface water hydrochemical parameters in the South Branch of the inner Changjaing Estuary in early April 2010, with arrows showing signals likely from a sewage outlet, as noticed by Chai et al. (2006). Dashed lines in (d–m) indicate the assumed element concentrations at the river end.
Figure 4. Surface water distributions of dissolved oxygen, pH, partial pressure of CO$_2$, and dissolved inorganic nutrients against salinity during the survey. Grey line in (c) is calculated based on conservative mixing lines of TAlk, DIC, and silicate.
Figure 5. Surface water distributions of total alkalinity and dissolved inorganic carbon against salinity during the survey. Several April 2007 data obtained in several neighboring sites (Fig. 1a) by Zhai et al. (2014) were sketched by crossover symbols.
**Figure 6.** Biogeochemical additions of dissolved inorganic nitrogen species and carbonate system parameters against salinity in the North Branch.
Figure 7. DIC vs. TAlk (a), ΔTAlk vs. ΔDIN (b), ΔNH₄⁺-N vs. ΔDIN (c), and ΔDIC vs. ΔTAlk (d).
**Figure 8.** A comparison between field-measured $[\text{CO}_2^*]$ addition and the predicted values (a) and the modeled conservative concentrations of $\text{CO}_3^{2-}$ ion and non-carbonate alkalinity (b). In (a), conservative $[\text{CO}_3^{2-}]$ concentrations were regarded as the maximum removals of earlier free $\text{CO}_2$ productions. In (b), non-carbonate alkalinity was defined here via $(\text{TAlk} - [\text{HCO}_3^-] - 2[\text{CO}_3^{2-}])$, where $[\text{HCO}_3^-]$ and $[\text{CO}_3^{2-}]$ were calculated via DIC and TAlk data.
Figure A1. Surface water concentrations of suspended particulate matter (a), particulate organic carbon (b), and particulate inorganic carbon (c) in the inner Changjiang Estuary in early April 2010.
Figure A2. Surface water distributions of ammonium, nitrate, dissolved inorganic nitrogen, partial pressure of CO$_2$, total alkalinity, and dissolved inorganic carbon against salinity in the South Branch of the inner Changjaing Estuary in early April 2010.