The response of inorganic carbon distributions and dynamics to upwelling-favorable winds on the northern Gulf of Mexico during summer

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Article info
Article history:
Received 31 March 2015
Received in revised form 1 August 2015
Accepted 26 August 2015
Available online 1 September 2015

Keywords:
Gas exchange
Carbon cycle
Hypoxia
Mississippi River plume
Continental shelf
Carbon cycle

Abstract
Upwelling-favorable winds and an offshore-distributed Mississippi and Atchafalaya River plume trajectory were observed in summer 2009 in contrast to the mean conditions from 2002 to 2010 (upwelling-unfavorable winds and an alongshore river plume trajectory), a set of conditions which was also observed in summer 2007. The responses of dissolved inorganic carbon (DIC) distributions and dynamics to upwelling-favorable winds are studied by comparing the contrasting conditions between summer 2009 and summer 2007 on the northern Gulf of Mexico. Patterns of surface water partial pressure of CO2 (pCO2), DIC, δ13C in DIC, and total alkalinity (TA) determined in July 2009 and August 2007 were strongly related to river plume trajectories, and differed between the two summers.

The slope of the relationship between dissolved oxygen (DO) and DIC in summer 2007 was comparable to the Redfield O/C ratio of 1.3, which was attributed to respiration of organic matter in the bottom water. The slope of the DO and DIC relationship and δ13CDIC values in bottom waters during July 2009 were clearly affected by mixing since their salinities were <35. A three end-member mixing model was used to remove mixing effects in δ13CDIC, to estimate the organic source of respiration, and (2) in DIC concentrations, to calculate DIC removal and release. δ13CDIC results in both summers were consistent with an apparent release of DIC in hypoxic waters (DO less than 2 mg L−1) associated with respiration of surface organic matter. The area-weighted surface DIC removal (i.e., biological production) was lower in 2009 than in 2007 on the shelf, as the plume was distributed offshore. The release of DIC in bottom waters was higher over the shelf in 2009 and was surmised to be related to stronger mixing, which was favorable for the DO supply for respiration. Overall, surface waters on the continental shelf in the region of study in July 2009 acted as a weak CO2 source to the atmosphere, but a weak CO2 sink in August 2007. We contend that the inorganic carbon distribution and concentrations on the shelf were related to regional wind forcing, through its influence on the distribution of coastal currents and plume trajectories and their subsequent impact on biogeochemical processes.

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1. Introduction

Despite its relatively small global areal coverage, the coastal ocean contributes substantially to the global carbon cycle through net biological production and export of organic carbon to the open ocean (Bauer and Bianchi, 2011) as well as through air-sea carbon dioxide (CO2) gas exchange (Cai, 2011; Chen and Borges, 2009; Cai et al., 2006). The coastal ocean faces serious perturbations that can alter the carbon cycle, such as through varying river export of inorganic carbon and rising atmospheric CO2 levels (Bauer et al., 2013; Raymond et al., 2008), riverine nutrient export (Chou et al., 2013), typhoon-enhanced primary production (Chen et al., in...
anomalous weather induced air–sea CO₂ variations (Ávila-López et al., in preparation), and long-term ocean acidification (Lui et al., in preparation). In particular, surface nitrogen enrichment through increasing river nitrate flux (Medsaas et al., 2001) can lead to cultural eutrophication (surface algal blooms) and consequent bottom water hypoxia (dissolved oxygen [DO] < 2 mg L⁻¹), especially on large-river-dominated continental shelves (Rabalais et al., 2010; Rabouille et al., 2008). Physically, coastal oceans are characterized by a complex shelf circulation that is driven by variations in freshwater discharge, upwelling along the shelf edge, and local wind forcing (Lentz and Frewings, 2012). Variations in shelf circulation can alter the cross- and along-shelf distribution of river plumes and associated nutrients (Blanchi et al., 2010) and consequently the severity of eutrophication.

Outflow from the Mississippi and Atchafalaya rivers is usually distributed westward along the Louisiana shelf during non-summer months (Cochrane and Kelly, 1986; Ohlmann and Niiler, 2005; Smith and Jacobs, 2005) in response to variable wind forcing primarily out of the north and east (Wang et al., 1998). Biogeochemically this circulation pattern produces an alongshore distribution of pCO₂ (partial pressure of CO₂) on the Louisiana shelf (Lohrenz et al., 2010; Huang et al., 2013). Surface water dissolved inorganic carbon (DIC) also exhibits an alongshore distribution, increasing toward the west (Cai, 2003; Guo et al., 2012). During summer months, wind forcing out of the south and west can result in a reversal of shelf circulation leading to a pattern of river plume discharge distributed to the east and cross-shelf (Cochrane and Kelly, 1986; Ohlmann and Niiler, 2005; Smith and Jacobs, 2005; Fichot et al., 2014). This general summer pattern of Louisiana shelf circulation is greatly dependent on local wind forcing, however (Cochrane and Kelly, 1986; Ohlmann and Niiler, 2005; Smith and Jacobs, 2005). Upwelling favorable winds on continental shelves without large freshwater discharges can induce deep or bottom waters rising toward surface. On this large-river dominated continental shelf, upwelling favorable wind forcing (northerly wind) has been observed irregularly during summers on the northern Gulf of Mexico with high chlorophyll a concentrations expending toward the west (Cai, 2003; Guo et al., 2012). During summer months, wind forcing out of the south and west can result in a reversal of shelf circulation leading to a pattern of river plume discharge distributed to the east and cross-shelf (Cochrane and Kelly, 1986; Ohlmann and Niiler, 2005; Smith and Jacobs, 2005; Fichot et al., 2014). This general summer pattern of Louisiana shelf circulation is greatly dependent on local wind forcing, however (Cochrane and Kelly, 1986; Ohlmann and Niiler, 2005; Smith and Jacobs, 2005). Upwelling favorable winds on continental shelves without large freshwater discharges can induce deep or bottom waters rising toward surface. On this large-river dominated continental shelf, upwelling favorable wind forcing (northerly wind) has been observed irregularly during summers on the northern Gulf of Mexico with high chlorophyll a concentrations expending toward the west (Cai, 2003; Guo et al., 2012). During summer months, wind forcing out of the south and west can result in a reversal of shelf circulation leading to a pattern of river plume discharge distributed to the east and cross-shelf (Cochrane and Kelly, 1986; Ohlmann and Niiler, 2005; Smith and Jacobs, 2005; Fichot et al., 2014). Specifically in July 2009, continuous upwelling-favorable winds along the Louisiana coast have been characterized with cross-shelf river plume trajectories in July 2009 (Zhang et al., 2012; Xue et al., 2013) and have been suggested to be related to a small areal extent of bottom water hypoxia thereafter (Feng et al., 2014; Turner et al., 2012; Bianchi et al., 2010; Forrest et al., 2011; Evans and Scavia, 2011). In order to evaluate the role of biogeochemistry in shelf circulations and their responses to global anthropogenic changes (Rabalais et al., 2010; Doney et al., 2011), there is a need to comprehensively describe the response of inorganic carbon to the upwelling-favorable wind on this shelf, especially given the limited extent of observations during upwelling-favorable wind forcing.

In this study, we assess the circulation and carbon biogeochemical response of the Louisiana shelf to changes in predominant wind fields. We do this by identifying the average summer (July/August) circulation climatology and then comparing that to biogeochemical observations made during August 2007 and July 2009, which respectively represent upwelling-unfavorable and upwelling-favorable summer conditions. Our approach includes an examination of satellite-derived chlorophyll-a (chl-a) and modeled surface currents for a climatologically average summer using (1) multi-year averages of July and August, (2) a summer with upwelling unfavorable winds (August 2007), and (3) a summer with upwelling favorable winds (July 2009). We then describe the distribution of sea surface salinity, surface water pCO₂, DIC, and total alkalinity (TA) in the surface and bottom water in both wind conditions. Finally, we describe and discuss air–sea CO₂ flux, the spatial extent of bottom water DIC, δ¹³C in DIC (δ¹³CDIC), and the relationship between DIC and DO in the bottom water on the Louisiana shelf.

2. Methods

2.1. Sampling and external data

Cruises were conducted August 18–24, 2007 on board OSV Bold, and July 19-29, 2009 on board R/V Cape Hatteras on the Louisiana shelf and its adjacent regions. TA and DIC samples were collected from Niskin bottles in thoroughly flushed 250 ml borosilicate glass bottles and then preserved with 100-μL HgCl₂ saturated solution. DIC samples were measured shortly after the cruise by acidifying 0.5 mL of sample and quantifying the released CO₂ with an infra-red gas analyzer (LI-COR® 6252). TA samples were measured by Gran titration using 0.1 M HCl and a semi-automated titrator. Both TA and DIC measurements had a precision of 0.1% (Cai et al., 2010; Huang et al., 2012b). δ¹³CDIC samples were collected in 2 mL vials and stored at 4 °C until analysis. δ¹³CDIC was measured using liquid chromatography coupled to an isotope-ratio mass spectrometer with a precision of 0.06‰ (Brandes, 2009). DO concentrations were measured using the sensor mounted on a vertical conductivity, temperature, and depth (CTD) array. The calibration of DO concentration was verified by comparison to Winkler titration based on samples collected from the Niskin bottle.

For sea surface salinity and pCO₂ measurements, seawater was sampled from the ship flow-through whose inlet was at about one meter depth. pCO₂ was measured by a flow-through system with a shower head equilibrator plus a CO₂ analyzer (LI-COR® 7000) (Jiang et al., 2008; Huang et al., 2015). The LI-COR® 7000 was calibrated every 3.5 to 6 h using four certified gas standards, which had dry CO₂ molar fractions of 197.45, 400.57, 594.65, and 975.26 ppm referenced against standards traceable to the National Institute of Standards and Technology. Area averaged pCO₂ values were gridded to a resolution of 0.1° × 0.1° and were used to calculate the air–sea CO₂ flux using the following equation:

\[
\text{CO}_2 \text{ flux} = k \times K_0 \times (p\text{CO}_2\text{sw} - p\text{CO}_2\text{aw})
\]

where \(k\) represents the gas transfer velocity; \(K_0\) is the solubility of CO₂ (Weiss, 1974); and \(p\text{CO}_2\text{sw}\) and \(p\text{CO}_2\text{aw}\) are the pCO₂ in surface seawater and the overlying atmosphere, respectively. We adopt the equation given by Ho et al. (2006) to calculate the air–sea gas transfer coefficient. More details about the calculation of air–sea CO₂ flux were described by Huang et al. (2015). Spatial wind forcing data were from QuikSCAT (Quick Scatterometer from NOAA’s Ocean Watch Live Access Server, http://las.pfeg.noaa.gov/ocean-Watch/oceanWatch.php) ocean surface winds data with a resolution of 0.25°. The wind forcing time series record was from the JUM1 station at Louisiana coast (29.255°N, 94.663°W) and the buoy (NOAA station #42043) near Galveston Texas (28.982°N, 94.898°W). Eight-day composites of MODIS-Aqua chl-a data from 2002 to 2010 were acquired from NASA’s (The National Aeronautics and Space Administration’s) Ocean Color Web (http://ocean-color.gsfc.nasa.gov/) and were used to illustrate the synoptic trajectory of the river plume (Del Castillo et al., 2001; Jones and Wiggert, 2015; Nababan et al., 2011). Hu et al. (2003) have compared satellite-derived and in situ measurements on the northern Gulf of Mexico and have reported a percentage error less than 50% between them. We applied the Hybrid Coordinate Ocean Model (HYCOM, Naval Research Laboratory, http://hycom.
org], HYCOM + NCODA Gulf of Mexico 1/25° Analysis) derived sea surface currents to illustrate general shelf circulation from 2003 to 2010. The slope of the DO-to-DIC relationship was determined by Type II regression (York, 1966) considering uncertainties in both DO and DIC analyses.

2.2. Three end-member mixing model

We follow Huang et al. (2012a) and assume DIC and nutrient concentrations were dominated by mixing and biological activity. Conservatively, one unit of water is the sum of the fraction (f) from each end-member: the Mississippi River (subscripted M), the Atchafalaya River (subscripted A), and seawater (subscripted S) as follows:

\[1 = f_M + f_A + f_S \]  \hspace{1cm} (2)

\[\text{Sal} = \text{Sal}_M \times f_M + \text{Sal}_A \times f_A + \text{Sal}_S \times f_S \]  \hspace{1cm} (3)

where \(\text{Sal}\) is our measured salinity, \(\text{Sal}\) with abbreviations is the salinity of each end-member, respectively. Measured TA and NO3 are further affected by biological uptake.

\[\text{TA} = \text{TA}_M \times f_M + \text{TA}_A \times f_A + \text{TA}_S \times f_S + \text{NO3}_{\text{Bio}} \]  \hspace{1cm} (4)

\[\text{NO3} = \text{NO3}_M \times f_M + \text{NO3}_A \times f_A + \text{NO3}_S \times f_S - \text{NO3}_{\text{Bio}} \]  \hspace{1cm} (5)

where subscripted “Bio” represents the biological uptake of the parameter. In order to combine Eqs. (4) and (5), Eq. (5) can be reorganized as follows:

\[\text{NO3}_{\text{Bio}} = \text{NO3}_M \times f_M + \text{NO3}_A \times f_A + \text{NO3}_S \times f_S - \text{NO3} \]  \hspace{1cm} (6)

Thus, NO3 Bio in Eq. (4) can be replaced by Eq. (6). Eq. (4) is further reorganized as follows:

\[\text{TA} + \text{NO3} = (\text{TA} + \text{NO3})_M \times f_M + (\text{TA} + \text{NO3})_A \times f_A + (\text{TA} + \text{NO3})_S \times f_S \]  \hspace{1cm} (7)

As TA and NO3 are measurements, the fractions from each endmember were solved along with the end-members of salinity, TA, DIC, and NO3 (Table A in Appendix) using Eqs. (2), (3), and (7) with a non-negative solver (NNLS in Matlab®). Furthermore, we define the difference between DIC and the conservative mixing DIC value as biological removal or release (\(\Delta\text{DIC}\)):

\[\Delta\text{DIC} = \text{DIC}_M \times f_M + \text{DIC}_A \times f_A + \text{DIC}_S \times f_S - \text{DIC} \]  \hspace{1cm} (8)

We assume the formation of \(\delta^{13}\text{C}_{\text{DIC}}\) in bottom hypoxic waters is the product of respiration and mixing of river and sea end-members.

\[\delta^{13}\text{C} \times \text{DIC} = \delta^{13}\text{C}_{\text{DIC}_M} \times f_M + \delta^{13}\text{C}_{\text{DIC}_A} \times f_A + \delta^{13}\text{C}_{\text{DIC}_S} \times f_S + \delta^{13}\text{C}_{\text{resp}} \times \Delta\text{DIC} \]  \hspace{1cm} (9)

where \(\delta^{13}\text{C}_{\text{resp}}\) is assumed to be the \(\delta^{13}\text{C}\) values of organic matter source for respiration.

3. Results

3.1. Wind forcing

The average spatial distribution of wind forcing was weak and southerly or southeasterly over the northern Gulf of Mexico during July and August as demonstrated by the eight-year climatology (Fig. 1a). These average conditions were similar to those observed during our August 2007 cruise (Fig. 1b). During our July 2009 cruise, however, the wind pattern was quite different, coming from the south along the Texas coast and the southwest over the Louisiana coast. The wind forcing was upwelling-favorable for the Louisiana shelf (Fig. 1c). Based on data from buoy and land stations, the average wind direction and speed in July and August over the interval 2004 to 2011 were southerly (182 ± 94.7° and 3.2 ± 1.8 m s⁻¹ at buoy LUML1, Louisiana and 175 ± 61.7° and 4.6 ± 2.1 m s⁻¹ at Galveston, TX). Wind forcing in August 2007 was similar: 169.4 ± 94.6° and 3.1 ± 1.4 m s⁻¹ at LUML1, Louisiana and 172.2 ± 82.1° and 4.0 ± 2.1 m s⁻¹ at Galveston, TX (F-test, \(P > 0.05\)). In July 2009 however, the average wind direction and speed were significantly different from the average: 221.6 ± 55.3° and 3.4 ± 1.7 m s⁻¹ at station LUML1 (F-test, \(P < 0.05\)). In Galveston TX it was not different: 189.8 ± 46.8° and 5.0 ± 1.8 m s⁻¹ (F-test, \(P > 0.05\)).
3.2. Distributions of plume trajectories

The nine-year average (2002–2010) distribution of MODIS-Aqua chl-a concentration in the months of July and August illustrates the average trajectory of the Mississippi and Atchafalaya River outflow (Fig. 2a). High chl-a concentrations were evident around the Mississippi River delta and westward along the Louisiana shelf towards Texas. The HYCOM-simulated eight-year average (2003–2010) surface shelf circulation is similar to that implied by the distribution of chl-a with weak currents on the inner shelf and strong horizontal currents near the boundaries between offshore and open Gulf of Mexico waters between 89 and 95°W (Fig. 2a). Only a few scattered high chl-a patches extend offshore to the south in this multi-year average distribution. The chl-a distribution observed in August 2007 was similar to the long-term trend, thus confirming that this cruise was conducted during upwelling-unfavorable summer conditions (Fig. 2b). In this upwelling-unfavorable summer, salinities less than 33 were observed immediately adjacent to the Southwest pass of the Mississippi River and along the inner Louisiana shelf towards Texas (Fig. 3a). The distributions of chl-a and salinity displayed an alongshore plume trajectory in August 2007. Looking seaward (Box A in Fig. 2), we observed high chl-a concentrations on the inner and middle shelves and low chl-a concentrations on the outer shelf and open gulf for both the multi-year average and August 2007 time frames (Fig. 4). Furthermore, the cross-shelf component of surface currents in the same region (Box A) was similar between the eight-year average (0.02 m s⁻¹) and the value in August 2007 (0.01 m s⁻¹).

Plume distributions as illustrated by chl-a were quite different in July 2009, being high in the inner shelf and out to the open gulf east of 91.5°W (Fig. 2c), especially immediately south of the river delta (Fig. 2c). Inner-shelf currents were particularly strong toward Louisiana (Fig. 2c), resulting in salinities being above 35 at stations on the inner shelf to the west of 93°W (Fig. 3b). Lower salinity waters (salinity less than 33) were confined mostly to the east of 91.5°W, except for a small pocket immediately next to the Atchafalaya River and Bay (Fig. 3b), and they extended to the open gulf to the east of 91.5°W. Cross-shelf currents were observed along 28°N between 89°W and 94°W during our July 2009 sampling (Fig. 2c). The offshore component in the region of Box A was 0.09 m s⁻¹, which was faster than the 8-year average (0.02 m s⁻¹).
offshore. Such low values were only observed on the inner shelf in (Fig. 6). Although we only have limited values were observed to the west of 93°W (Fig. 6b, f). The δ13CDIC values that extended from the inner shelf to the west of 93°W during our July sampling period. High salinity seawater on the inner shelf west of 92.5°W was observed, and its source was evident from δ13CDIC, which is discussed in the following paragraphs.

3.3. Distributions of TA, pCO2, DIC, and δ13CDIC

Sea surface spatial distributions of TA, DIC, and δ13CDIC were different between August 2007 and July 2009. In August 2007, low TA and DIC values were observed on the inner shelf west of 93.5°W (Fig. 5). But high TA, pCO2, DIC, and δ13CDIC values were observed at the same area in July 2009 (Fig. 5). We also observed low TA, pCO2, DIC, and δ13CDIC values that extended from the inner to outer shelves east of 91°W in July 2009 (Fig. 5). In contrast, high values west of 93°W in July 2009 were less observed in August 2007. The isotope data, δ13CDIC, were consistent with satellite and modeled currents in July 2009, i.e., δ13CDIC values were close to 0‰ (open ocean signal) on the inner shelf to the west of 93°W and were −2‰ (river induced signal) at the offshore along 90.4°W.

There were similar gradients in TA and DIC in bottom waters that differed between 2007 and 2009. TA increased in the offshore direction while DIC was low on both inner and outer shelves and highest on the middle shelf in August 2007 (Fig. 6). For July 2009, we only focused on the bottom waters north of 28.2°N. TA and DIC still showed a cross-shelf gradient to the east of 92°W but stable values were observed to the west of 93°W in the bottom water (Fig. 6). Although we only have limited δ13CDIC data in bottom waters, data showed a pattern similar to above features: lower δ13CDIC values (< −1‰) in bottom waters along 91°W toward offshore. Such low values were only observed on the inner shelf in August 2007 (Fig. 6c).

3.4. Distributions of air–sea CO2 flux, surface DIC removal, and bottom DIC release

In comparing air–sea CO2 fluxes between the two conditions, we confined our analysis to the smaller survey area of the August 2007 cruise (generally within the 100-m isobath). Results show that the shelf acted as a weak CO2 sink to the atmosphere (−2.0 ± 0.20 mmol m−2 d−1) in August 2007, but a weak source (1.4 ± 0.13 mmol m−2 d−1) in July 2009 (Table 1). This difference in behavior was mainly due to the difference in seawater pCO2 levels on the western area between the two years (Fig. 5b,f). The ΔCO2 flux was 4.96 mmol m−2 d−1 on the western area in contrast to 2.7 mmol m−2 d−1 on the eastern area.

In the surface water, biological DIC removal was high (e.g., net biological production of organic carbon) on the inner shelf and low on the outer shelf in August 2007 (Fig. 7a). In July 2009, surface DIC removal was observed to the east of 91.5°W (generally 38–70 μM), but surface DIC release was also observed to the west of 93°W (mean ΔDIC = −11 μM with the lowest value of −77 μM on the inner shelf along 93.4°W) (Fig. 7b). Overall, the integrated surface DIC removal on the shelf in July 2009 was 81.8 μM less than in August 2007 (Table 1). This result was consistent with observations that high productivity was transported offshore and low productivity waters were observed on the shelf to the west of 92.5°W in July 2009.

In the bottom water, DIC release (net biological respiration of organic carbon) demonstrated a cross-shelf gradient in August 2007, i.e., the released DIC concentration was low on the inner and outer shelves (< 200 μM) and was high on the middle shelf (> 200 μM) (Fig. 7c). Only a small patch of DIC removal was observed on the shallow inner shelf to the west of 93°W in August 2007. In July 2009, bottom water-released DIC was generally observed over this shelf. Released DIC concentration was particularly high at a few stations on the middle to offshore to the east of 92°W (Fig. 7d). The integrated result showed that released DIC concentration was 68.3 μM more in July 2009 than in August 2007 (Table 1).

3.5. DO–DIC relationships and δ13CDIC–salinity relationships

We observed negative linear relationships between DO and DIC in Louisiana shelf waters for brackish and high salinity waters and the slopes differed between upwelling-unfavorable and upwelling-favorable summer conditions (Fig 8a). To better illustrate the DO–DIC relationships, we separately compared brackish and high salinity waters. We define inner “brackish waters” as inner shelf water with salinity < 35, water depth < 20 m, DO < 150 μM,
and DIC < 2200 μM. The biogeochemical composition is highly influenced by the river end-member (low salinity, shallow depth, and high DO). Brackish waters along the 10 and 20 m isobaths parallel the trajectory of plume water in the upwelling-unfavorable year. Bottom waters offshore of the 20 m isobath showed little salinity variation (< 0.4) and were similar in composition to the open ocean end-member (high salinity, deeper depth and high DO). We define “high salinity waters” as more offshore water with salinity > 35, depth > 20 m and DO < 150 μM. In all cases, DO and DIC demonstrated negative linear relations: for brackish waters, the slope was $-1.0 \pm 0.2$ ($R^2=0.826$); and for high salinity waters, the slope was $-1.2 \pm 0.1$ ($R^2=0.836$). Under upwelling-favorable conditions of July 2009, hypoxic conditions were less observed (Fig. 8b) and the slope of the DO–DIC relationship was more negative with brackish and high salinity waters.

The variation of $\delta^{13}$C$_{DIC}$ was indicative of an influence of mixing (river and seawater) as well as metabolism (Figs. 6 and 9). $\delta^{13}$C$_{DIC}$ values were generally higher in surface than in bottom waters, reflecting the net effect of biological production and the net effect of respiration, respectively (Fig. 9). As salinities were lower in hypoxic waters in July 2009 than August 2007, $\delta^{13}$C$_{DIC}$ values were also lower in these hypoxic waters in July 2009 (Fig. 9). This mixing effect on $\delta^{13}$C$_{DIC}$ was removed by the mixing model and the result is discussed in Section 4.3.

4. Discussion

4.1. The biogeochemical response of surface waters to upwelling-favorable winds

The characteristics of spatial patterns in surface $pCO_2$, DIC, TA,
Fig. 6. Distributions of TA (μM), DIC (μM), and δ¹³CDIC (‰) in bottom waters. TA (a), DIC (b), and δ¹³CDIC (c) values in the bottom water were lower on the inner shelf and were higher on the middle and outer shelves in August 2007. Despite the fact that such cross-shelf gradients can also be observed in the region right south to Atchafalaya Bay in July 2009, TA (d) and DIC (e) values were higher to the west of 93° W. δ¹³CDIC (f) values were lower on the inner shelf and displayed lower values than the ones in August 2007 along 91° W.

Fig. 7. Distributions of surface water-removed DIC and bottom water-released DIC concentrations (μM). Surface DIC removals were generally higher in August 2007 (a) than those in July 2009 (b) and the difference between them was higher to the east of 91.5° W (c). In the bottom water, released DIC was high alongshore around the middle shelf to the west of 91.5° W in August 2007 (c) and such high DIC release was not observed to the west of 91.5° W in July 2009 (d).
and chl-a were generally consistent with the characteristics of river discharge trajectories represented by satellite chl-a distributions, i.e., alongshore distribution in August 2007 and offshore transport in July 2009. The distribution of CO₂-undersaturated waters (relative to the atmospheric value) as reported in previous studies conducted in summer (Lohrenz et al., 2010; Lohrenz and Cai, 2006, Guo et al., 2012) is generally alongshore, as was the distribution of DIC removal in August 2007. However, in July 2009, the region of CO₂ undersaturation and strong DIC removal was restricted to the east of 91.5°W, which reflected the location of the river plume under the influence of upwelling-favorable winds at that time. As a consequence of this different pattern of CO₂ drawdown and DIC removal, the shelf was a weak source in 2009 in contrast to being a sink of atmospheric CO₂ in 2007 (Table 1). This biological carbon uptake was also evident by pigment concentrations measured on the same cruise on the open gulf during July 2009: higher chl-a concentrations in evident by pigment concentrations measured on the same cruise on the open gulf during July 2009: higher chl-a concentrations in low-to-middle salinity waters (Chakraborty and Lohrenz, 2015). Changes in Δ^{13}C_DIC might be related to processes such as riverine input of inorganic and organic materials along with mixing, production, respiration, and air–sea gas exchange. We attribute the Δ^{13}C_DIC values that are higher than the general conservative mixing relationship between river and sea end-members as having likely been elevated by net photosynthesis, while those below this mixing line were likely reduced by net respiration (Fig. 9). Changes in Δ^{13}C_O₂ in hypoxic waters will be further discussed in the next two sections.

### 4.2. Bottom-water respiration in August 2007

The relationship between DO and DIC suggests that the variations of DO and DIC concentrations in the bottom water were dominated by microbial respiration. In August 2007, the slope of the DO and DIC regression was $1.0 \pm 0.2$ ($R^2 = 0.826$) in brackish waters and $1.2 \pm 0.1$ ($R^2 = 0.836$) for high salinity waters. Both slopes are similar to Redfield stoichiometry, i.e., the O/C molar ratio is 1.30 (138/106, Redfield, 1958). Such Redfield-type

![Fig. 8. The relationships between DO and DIC in August 2007. The relationship between DO and DIC in bottom waters can be observed in brackish and high salinity waters individually (a), which were separated by an alongshore isobath zone between 10 and 20 m. The slopes of the DO-to-DIC relationship ($1.0 \pm 0.2$, $R^2 = 0.826$ for brackish waters; and $1.2 \pm 0.1$, $R^2 = 0.836$ for high salinity waters) were both close to the estimation of Redfield respiration (represented by two red arrows in b). In July 2009, the slopes of the relationship between DO and DIC were higher than those in August 2007 and also higher than Redfield-ratio predicted lines (gray dash lines). (For interpretation of the references to color in this figure legend, the reader is referred to the web version of this article.)](image-url)
respiration has been observed in hypoxic coastal waters as well (Cantoni et al., 2012; Maske et al., 2010; Hu et al., in press). In addition, we observed that strong surface DIC removal was on the inner shelf (in both surface and shallow bottom waters) while strong bottom DIC removal was on the middle shelf (Fig. 7a,d), suggesting that particulate organic carbon moved from inner shelf to outer shelf. This suggestion was consistent with Fry et al. (2015).

Those values of δ¹³CDIC below the river-to-sea mixing lines (Fig. 9) were mostly associated with middle and bottom waters and were believed to be related to depletion of δ¹³CDIC due to respiration activity. Based on our three end-member model, δ¹³Cresp displayed an average of −21.3 ± 2.5‰ in hypoxic waters in August 2007 (Table 2). This result was consistent with results in previous studies, which ranged from −22.9‰ to −22.7 ± 10.3‰ (Strauss et al., 2012; Hu et al., in press), and also in the range of δ¹³C values of −22.5 to −21.2‰ for surface sediment on the inner shelf (Bianchi et al., 2002). For the two outliers in shallow waters (−47.4‰, −57.4‰ at 10 m bottom depth), these two δ¹³CDIC values were probably affected by factors in addition to our assumption (mixing and respiration), such as methane from the marsh and air-sea gas exchange. Lower δ¹³Cresp values in 10-m depth compared to values in >10-m depth were also observed by Strauss et al. (2012) in hypoxic waters. In addition, the three end-member model generated results comparable to the salinity normalized method used by Hu et al. (in press) as the water salinity was above 35 in August 2007 and the effect of variations of river end-members on high salinity waters can be neglected. Assuming 1‰ difference in δ¹³C in river end-member DIC variations, the variation is less than 0.002‰ when salinity was higher than 35 and was smaller than the resolution of measurements (± 0.06‰).

The river end-members of δ¹³CDIC in the Mississippi River (−10.7‰ at salinity 1.36) and the Atchafalaya River (−9.7‰ at salinity 0.35) were close to the value of −10‰ reported by Dubois et al. (2010) for the lower Mississippi River waters. The δ¹³CDIC sea end-member was 0.65‰ in August 2007 and was between the two end-members used by Strauss et al. (2012) and Hu et al. (in press), i.e. 0.5‰ and 0.8‰, respectively. In addition, carbonate precipitation and dissolution can also affect δ¹³CDIC. However, this effect can be limited as the fractionation factor of carbonate precipitation is relatively small (<3‰; Romanek et al., 1992) and the contribution of coccolithophores to variations in DIC is believed to be small (Cai, unpublished data; Guo et al., 2012).

### 4.3. The impact of shifting surface biological production on bottom-water DIC release in July 2009

The area-integrated bottom water-released DIC concentration was 68.3 µM higher on the shelf in July 2009 than in August 2007 (Table 1). Please note that the standard deviations of the area-integrated concentration (± 88.2 and ± 69.2 µM) imply spatial variations in August 2007 and July 2009 and they should not be neglected. To reduce these spatial variations, we discuss DIC release in sub-regions. The carbon processing in bottom waters was observed to be similar in the overlapping plume distributions from July 2009 and August 2007 (that is, on the shelf to the east of 92.5°W) and was suggested to be different in other three regions as a consequence of the relocated plume. These include (1) the shelf (10 m depth) waters, (2) the inner shelf to the west of 92.5°W, and (3) the open gulf to the east of 92.5°W. On the shelf to the east of 92.5°W in July 2009, strong released DIC concentrations in hypoxic waters were observed at two stations along 91°W (Fig. 7d). Mixing apparently affected these hypoxic waters as their salinities (34–36) were lower than those of hypoxic waters in August 2007 (Fig. 8). The three end-member model was applied again to remove the mixing effect and to estimate the organic matter respired in hypoxic waters in July 2009. The δ¹³Cresp results ranged from −23.5‰ and −22.1‰ at bottom depths of 21.6 and 17.3 m, respectively, similar to previous results from August 2007. In contrast, the δ¹³Cresp in shallow waters (bottom depth of 10 m) was −29.4‰ and was more ¹³C enriched than the δ¹³Cresp values at the same depth in August 2007 (−47.4 and −57.4‰). As this

### Table 2

<table>
<thead>
<tr>
<th>Cruise</th>
<th>Station</th>
<th>Latitude °N</th>
<th>Longitude °W</th>
<th>Depth (m)</th>
<th>Sal.</th>
<th>DIC (µM)</th>
<th>δ¹³CDIC (%)</th>
<th>δ¹³Cresp (%)</th>
<th>DO (µM)</th>
</tr>
</thead>
<tbody>
<tr>
<td>GM0708</td>
<td>A04</td>
<td>29.133</td>
<td>89.750</td>
<td>18</td>
<td>36.16</td>
<td>2239.7</td>
<td>−0.33</td>
<td>−19.9</td>
<td>30.9</td>
</tr>
<tr>
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<td>A06</td>
<td>29.010</td>
<td>89.751</td>
<td>25</td>
<td>36.22</td>
<td>2267.1</td>
<td>−0.85</td>
<td>−24.5</td>
<td>24.7</td>
</tr>
<tr>
<td>GM0708</td>
<td>B03</td>
<td>29.047</td>
<td>90.164</td>
<td>10</td>
<td>33.81</td>
<td>2195.8</td>
<td>−1.35</td>
<td>−47.4</td>
<td>9.1</td>
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<tr>
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<td>B06</td>
<td>28.993</td>
<td>90.076</td>
<td>18</td>
<td>36.00</td>
<td>2290.7</td>
<td>−0.95</td>
<td>−23.7</td>
<td>4.4</td>
</tr>
<tr>
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<td>B09</td>
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<td>89.994</td>
<td>37</td>
<td>36.48</td>
<td>2259.7</td>
<td>−0.30</td>
<td>−17.3</td>
<td>41.9</td>
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<td>90.525</td>
<td>10</td>
<td>32.10</td>
<td>2159.0</td>
<td>−1.25</td>
<td>−57.4</td>
<td>31.3</td>
</tr>
<tr>
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<td>C06</td>
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<td>90.463</td>
<td>18</td>
<td>35.83</td>
<td>2241.7</td>
<td>−0.38</td>
<td>−19.7</td>
<td>5.3</td>
</tr>
<tr>
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<td>C08</td>
<td>28.789</td>
<td>90.277</td>
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<td>35.91</td>
<td>2247.3</td>
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<td>−22.5</td>
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</tr>
<tr>
<td>GM0708</td>
<td>D03</td>
<td>28.718</td>
<td>90.834</td>
<td>14</td>
<td>35.88</td>
<td>2260.2</td>
<td>−0.69</td>
<td>−21.6</td>
<td>8.1</td>
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<tr>
<td>GC0907</td>
<td>D2</td>
<td>28.584</td>
<td>91.024</td>
<td>21.6</td>
<td>35.70</td>
<td>2340.2</td>
<td>−1.47</td>
<td>−23.5</td>
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<tr>
<td>GC0907</td>
<td>D1</td>
<td>28.785</td>
<td>91.008</td>
<td>9.3</td>
<td>34.04</td>
<td>2305.7</td>
<td>−2.02</td>
<td>−28.9</td>
<td>4.7</td>
</tr>
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</table>
offshore transport event in July 2009 was ephemeral (Feng et al., 2014), the effect of air–sea gas exchange on δ13CDIC was expected to be less than in August 2007 when the water residence time was longer. Thus, this δ13Cresp value for 10-m bottom depth in July 2009 might reflect a signal close to the δ13C values of soil and plant material in freshwater and salt marshes in Louisiana (−26.5 to −27.9‰) (DeLaune, 1986). A better understanding of the underlying mechanisms for the observed δ13Cresp results will require further observation as this system was highly dynamic and our estimation was based simply on mixing and respiration. Furthermore, the bottom DIC release was largely reduced on the shelf to the west of 92.5°W in July 2009 but the overlying surface ΔDIC was negative (release of CO2). This DIC release might be related to benthic respiration which has already been suggested to be important in this far-field plume during upwelling-unfavorable conditions (Hetland and DiMarco, 2008). On the third region (the surface open gulf to the east of 92.5°W), elevated biological uptake (characterized by strong DIC removal and high chl-a concentrations) was observed, implying that the biological pump on this region was enhanced during this rare event in July 2009.

4.4. Complex mechanisms behind this highly dynamic system in July 2009

The surface shelf circulation on the northern Gulf of Mexico, while highly dynamic and complex in summer, was particularly unusual in summer 2009. Coastal current reversal during summer months has been associated with freshwater discharge offshore and to the east of the Mississippi delta (Wiseman et al., 1997; Nowlin et al., 2005), a pattern consistent with our observations (Fig. 3). Lentz and Frewings (2012) have theoretically described the response of shelf circulation to wind direction for an ideal coastline and topography. Chu et al. (2005) used drifters to reconstruct the shelf circulation in this study area from fall to winter and have modeled the shelf circulation by applying different wind forcing (northeasterly and southwesterly). Their findings attributed the reverse of the Texas–Louisiana shelf current from westward to eastward to the shelf-wide southwestwardly wind forcing. Feng et al. (2014) used a 3D model to simulate the reduced hypoxic areal extent in July 2009 and noted the importance of the time period when upwelling wind occurs as a factor influencing the strength of current reversals. The persistent upwelling-favorable wind altered the vertical and horizontal distribution of low salinity waters (Feng et al., 2014). Apparently, riverine nutrients followed the plume trajectory and were removed proportionally to DIC uptake. In addition, light and stratification are non-negligible controlling factors as they have been identified to be important in biological activity and phytoplankton community variability (Lohrenz et al., 1999; Lehrter et al., 2009).

River discharge also played a non-negligible role in addition to the upwelling-favorable wind. The Mississippi and Atchafalaya river discharges were increasing prior to our sampling period in July 2009, while they remained low before the August 2007 sampling effort. Huang et al. (2013) and Fichot et al. (2014) speculated that cross-shelf transport was not only related to upwelling-favorable wind but also to high river discharge in this study area. This is likely, as river discharge has been shown to be correlated with the areal coverage of the river plume and the concentration of total suspended material (Shi and Wang, 2009). In addition, a severe drought occurred in Texas during summer 2009 (NOAA, National Climatic Data Center report). These conditions would have led to low export of freshwater discharge to the shelf and less river-to-sea mixing along the Texas coastline. High salinity water, thus, had a greater probability of intruding onto the inner Louisiana shelf to the west of 93°W inner shelf, as the δ13DIC values on the inner shelf were close to the values in high-salinity offshore waters (Fig. 3b).

Gulf-wide wind forcing has been suggested to affect the Loop Current and the coastal currents along Texas and Louisiana shelves (Chang and Oey, 2010), as the 10-day surface current exhibits coherence with the establishment of a high pressure atmospheric pressure system over the Gulf of Mexico (Barron and Vastano, 1994). In a conceptual model provided by Rabalais et al. (2010), the biogeochemical processes were postulated to directly respond not only to local wind forcing, but also to regional weather variation. We suggest that wind conditions dominated the distribution of surface primary production, which subsequently influenced the biogeochemistry of subsurface waters. An atmosphere-terrestrial-ocean biogeochemical model or a large data analysis among these three datasets is needed to fully explain the biogeochemical response on the shelf to regional weather variations.

5. Summary

Under an upwelling-unfavorable summer condition as in August 2007, surface distributions of chl-a, pCO2, DIC, and TA values were observed to track the Mississippi and Atchafalaya River plume trajectory, which was alongshore and confined to the shelf. Conditions in July 2009, however, differed in that the river plume, along with changed surface distributions of chl-a, pCO2, DIC, and TA values, were distributed to the east of 91.5°W and extended to the gulf. Similarly, δ13C values normally characteristic of inner-shelf waters were observed in waters on middle-to-outer shelves, consistent with the offshore plume trajectory in July 2009. High-productivity plume waters were offshore and low-productivity seawater was on the western shelf in July 2009, resulting in less DIC removal on the shelf under this upwelling-favorable wind event than the mean conditions. Higher area-integrated bottom DIC concentrations on the shelf under this upwelling-favorable wind event compared to the mean conditions might be related to mixing, which was favorable for DO supply. An examination of surface air–sea CO2 fluxes revealed that the Louisiana shelf acted as weak source of atmospheric CO2 in July 2009 in contrast to a weak sink in August 2007. The slope of the DO and DIC relationship in August 2007 was comparable to the Redfield stoichiometry of 1.3 for the O/C ratio and was an indication of the influence of respiration in the bottom waters. The slope of the DO and DIC relationship as well as the δ13C values in bottom water were affected by mixing during upwelling-favorable wind forcing in July 2009. After removing mixing effects, patterns of δ13C also provided evidence that the source of DIC released via respiration was of marine origin in both August 2007 and July 2009. Our results provide clear evidence that wind direction and shelf circulation on the Louisiana shelf play a major role in the distribution of salinity, biological communities, community metabolism, and biogeochemical properties, such as chl-a, DO, pCO2, and DIC values. For this highly anthropogenically influenced system, we suggest that anomalous weather events such as that observed in July 2009 may become more frequent under future climatic conditions with major consequences for carbon processes.

Acknowledgements

We thank the captains and their crews of the OSV Bold and R/V Cape Hatteras for their support in field sampling. Austin Todd provided many helpful comments. This work was supported by the National Science Foundation (OCE-0752110 and OCE-0752254), the National Aeronautics and Space Administration (NNX10AU06G, NNX10AU78G, NNH14AY07I, and NNX14AO73G),
Table A
Endmembers for the three-endmember mixing model. The river end-member of δ13C in August 2007 was lacking and was replaced by the values from July 2009.

<table>
<thead>
<tr>
<th>End-member</th>
<th>Salinity TA (μ mol kg⁻¹)</th>
<th>DIC (μ mol kg⁻¹)</th>
<th>NO₃ (μ mol kg⁻¹)</th>
<th>δ¹³C (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>August 2007 cruise</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Mississippi River</td>
<td>0.1</td>
<td>2420.0</td>
<td>2450.0</td>
<td>51.4</td>
</tr>
<tr>
<td>Atchafalaya River</td>
<td>0.1</td>
<td>1620.0</td>
<td>1650.0</td>
<td>35.0</td>
</tr>
<tr>
<td>Sea</td>
<td>36.04</td>
<td>2399.3</td>
<td>2082.8</td>
<td>2.3</td>
</tr>
<tr>
<td>July 2009 cruise</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Mississippi River</td>
<td>0.1</td>
<td>2140.0</td>
<td>2170.0</td>
<td>100.7</td>
</tr>
<tr>
<td>Atchafalaya River</td>
<td>0.1</td>
<td>1580.0</td>
<td>1610.0</td>
<td>45.7</td>
</tr>
<tr>
<td>Sea</td>
<td>36.75</td>
<td>2424.7</td>
<td>2055.2</td>
<td>0.7</td>
</tr>
</tbody>
</table>

and the BP-funded Gulf of Mexico Research Initiative RFP-I award (GoMRI-020).

Appendix

Table A

References


Huang, W.-J., Wang, Y., Cai, W.-J., 2012b. Assessment of sample storage techniques and the BP-funded Gulf of Mexico Research Initiative RFP-I award (GoMRI-020).


