Spatial variability of surface-water pCO$_2$ and gas exchange in the world’s largest semi-enclosed estuarine system: St. Lawrence Estuary (Canada)

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Abstract

The incomplete spatial coverage of partial pressure of CO$_2$ (pCO$_2$) measurements across estuary types represents a significant knowledge gap in current regional- and global-scale estimates of estuarine CO$_2$ emissions. The Estuary and Gulf of St. Lawrence (EGSL) at the lower limit of the subarctic region in eastern Canada is the largest estuarine system in the world, and is characterized by an exceptional richness in environmental diversity. It is among the world’s most intensively studied estuaries, yet there are no published data on its surface-water pCO$_2$ distribution. To fill this data gap, a comprehensive dataset was compiled from direct and indirect measurements of carbonate system parameters in the surface waters of the EGSL during the spring or summer of 2003-2016. The calculated pCO$_2$ ranged from 435-765 μatm in the shallow, partially mixed Upper Estuary, 139-578 μatm in the deep, stratified Lower Estuary, and 207-478 μatm along the Laurentian Channel in the Gulf. Overall, at the time of sampling, the St. Lawrence Estuary served as a weak source of CO$_2$ to the atmosphere, with an area-averaged CO$_2$ degassing flux of 1.00 to 2.06 mmol C m$^{-2}$ d$^{-1}$ (0.37 to 0.75 mol C m$^{-2}$ yr$^{-1}$). A preliminary analysis of factors controlling the spatial variability of surface-water pCO$_2$ reveals that respiration (Upper Estuary), photosynthesis (Lower Estuary), and temperature (Gulf) are dominant controls.

Keywords:
Coastal ocean, Estuaries, CO$_2$, Carbon cycle, Air-sea CO$_2$ exchange, St. Lawrence Estuary, Gulf of St. Lawrence, Laurentian Channel

1. Introduction
Although estuaries occupy a very small fraction (0.2 %) of the global ocean surface area, their CO$_2$ emissions are disproportionately large compared with CO$_2$ exchanges between the open ocean and the atmosphere (Bauer et al., 2013). With an estimated global efflux of 0.25 ± 0.25 Pg C yr$^{-1}$ (Regnier et al., 2013), estuarine CO$_2$ degassing is thought to counterbalance CO$_2$ uptake on the continental shelves (Laruelle et al., 2010; Cai, 2011). Almost every estuary on Earth, for which data are available, is generally supersaturated with CO$_2$ with respect to the atmosphere (Cai and Wang, 1998; Frankignouelle et al., 1998; Borges, 2005; Borges et al., 2005; Borges et al., 2006; Chen and Borges, 2009; Laruelle et al., 2010; Cai, 2011; Chen et al., 2012; Bauer et al., 2013; Chen et al., 2013; Regnier et al., 2013), with values of partial pressure of CO$_2$ (pCO$_2$) ranging from 400 to 10,000 μatm (in contrast, the atmospheric pCO$_2$ in coastal zones was approximately 360–385 μatm in year 2000) (Cai, 2011). Although estuaries are generally net sources of CO$_2$, there is considerable variability and uncertainty in estimates of their CO$_2$ emissions, reflecting the limited spatial and temporal coverage of pCO$_2$ measurements in estuaries as well as their heterogeneous nature (hydrological and geomorphological differences, differences in magnitude and stoichiometry of carbon and nutrient inputs) (Bauer et al., 2013; Regnier et al., 2013).

Estuaries are geochemical reaction vessels through which continentally weathered organic matter and inorganic nutrients must pass to enter the coastal ocean (Kaul and Froelich, 1984). Horizontal transport is controlled by a set of physical attributes (tides, wind, bathymetry, basin geography, river flow) that determine the estuarine filter function (Cloern, 2001). The longer the freshwater flushing (or turnover) time of the estuary, the more opportunity there is for water-column biological activity, benthic exchanges and particle-dissolved phase interactions to influence its biogeochemistry (Statham, 2012). The high pCO$_2$ observed in estuaries results from the in situ microbial degradation of internally and externally supplied organic carbon and the lateral transport of inorganic carbon from rivers, coastal wetlands and ground waters (Bauer et al., 2013). In strongly tidal (macrotidal) systems, long water and particle residence times (on the order of weeks to months; Middelburg and Herman, 2007) allow for the extensive modification and degradation of particulate organic carbon during estuarine transport (Borges et al., 2006; Chen and Borges, 2009). In the absence of seasonal or permanent water stratification, the decoupling between production and degradation of organic matter at and below the surface, respectively,
does not occur, resulting in less efficient export of dissolved inorganic carbon (Borges, 2005). Strongly tidal estuaries also tend to exhibit lower levels of photosynthetic activity (Monbet, 1992) and have greater suspended particulate matter concentrations within their high-turbidity regions (Uncles et al., 2002; Middelburg and Herman, 2007) where suspended particles and organic-rich aggregates serve as “hot spots” of microbial recycling (Statham, 2012). Field measurements suggest that 10 % of the CO₂ emissions from the inner estuary of macrotidal systems is sustained by the ventilation of riverine CO₂, whereas 90 % is due to local net heterotrophy (Borges et al., 2006) fueled by inputs of terrestrial and riverine-algae derived (planktonic) detritus and, in populated areas, sewage (Chen and Borges, 2009).

North American estuaries rank first in terms of global estuarine surface area (41 %) but account for the lowest numerically averaged CO₂ flux per unit area (12 %) among all continents (Chen et al., 2013). These estimates are subject to large uncertainties due to data paucity. A recent synthesis by Regnier et al. (2013) highlights the meagre spatial coverage of estuarine pCO₂ measurements, particularly along the Canadian eastern seaboard. Ironically, the Estuary and Gulf of St. Lawrence (EGSL) in eastern Canada is the largest semi-enclosed estuarine system in the world, and is among the world’s most intensively studied estuaries (El-Sabh and Silverberg, 1990), but was left unmentioned in recent global (Cai, 2011; Chen et al., 2012; Chen et al., 2013) and regional (Laruelle et al., 2015) data compilations. Furthermore, the majority of previous estuarine CO₂ studies have focused on small-scale river-dominated estuaries, whereas there has been limited research on CO₂ dynamics in large estuaries and bay systems (Joesoef et al., 2015) including the areas of mixing at sea (outer estuaries or river plumes) (Borges et al., 2005). A comparative study by Jiang et al. (2008) reveals large differences in CO₂ degassing between marine-dominated and river-dominated estuaries while several authors note that outer estuaries (plumes) can differ substantially from inner (inland) estuaries in terms of CO₂ emissions (Frankignoulette et al., 1998; Borges, 2005; Chen et al., 2012). On the U.S. east coast, the ratio of non-riverine (flushed by tidal action and receiving minimum freshwater inputs) to river-dominated estuaries is nearly 1:1, demonstrating the geographic importance of coastal estuaries/bays on the eastern seaboard of North America (Cai, 2011).
The large-scale (width often considerably greater than the internal Rossby radius; Cyr et al., 2015), macrotidal (mean tidal range greater than 2-4 m; Monbet, 1992) St. Lawrence Estuary is an excellent analogue of marine-dominated systems. Throughout its length, the full spectrum of oceanic variability can be found (Mertz and Gratton, 1990). Moreover, the basin characteristics and water transport time scales of the St. Lawrence Estuary provide an almost ideal natural laboratory for geochemical studies. Its surface waters have a renewal time of several months while its bottom waters take several years to replenish, allowing for the comparison of spatial-temporal variations in physical and chemical properties (El-Sabh and Silverberg, 1990). Given its bimodal bathymetry, the St. Lawrence Estuary also permits the investigation of biogeochemical processes in two types of estuary: (1) the shallow, partially mixed Upper Estuary where physical mixing and abiotic processes dominate, and (2) the deep, stratified Lower Estuary where biological cycling and oceanic processes prevail (Yeats, 1990). As yet, no systematic study of the surface-water pCO₂ distribution in the St. Lawrence has been published and, hence, the present study provides the first comprehensive description of its mixed-layer inorganic carbon chemistry, including (1) a multi-year compilation of springtime and summertime pCO₂ calculated from field measurements of alkalinity and pH; (2) an area-averaged estimate of the air-sea CO₂ gas flux; and (3) an analysis of the relative importance of thermodynamic (temperature) and biological (photosynthesis, respiration) processes in controlling the spatial variability of estuarine pCO₂.

2. Materials and methods

2.1. Study area–St. Lawrence Estuary and Gulf

The greater St. Lawrence system (Fig. 1) connects the chain of Great Lakes, the second largest terrestrial freshwater reservoir in the world, to the Atlantic Ocean (Yang et al., 1996). With a drainage basin of approximately 1.32 million km², the St. Lawrence River channels the second largest freshwater discharge (11,900 m³ s⁻¹) on the North American continent, second only to that of the Mississippi (El-Sabh and Silverberg, 1990). The catchment area geology is dominated by silicate rocks of the Precambrian Shield and carbonates of the Paleozoic lowlands, whose components influence the downstream evolution of river chemistry (Yang et al., 1996). The erosion of the
carbonate rocks of the drainage basin is practically constant, as the quantity of bicarbonate ions carried by the river varies little from season to season (Pelletier and Lebel, 1979). On a yearly basis, between 15-20 % of the outflow of dissolved inorganic carbon from the St. Lawrence River into its estuary originates from the system’s tributaries (e.g., Ottawa River, Mascouche River) while 80-85 % is from the Great Lakes (Hélie et al., 2002). The St. Lawrence Estuary (SLE) begins at the landward limit of the salt water intrusion near Île d’Orléans (5 km downstream of Québec City) and stretches 400 km seaward to Pointe-des-Monts where it widens into the Gulf of St. Lawrence (GSL) (El-Sabh and Silverberg, 1990), a semi-enclosed sea with an area of approximately 240,000 km² (Dufour and Ouellet, 2007) connected to the Atlantic Ocean through Cabot Strait and the Strait of Belle Isle (Coote and Yeats, 1979).

Traditionally, the SLE is divided into two segments based on its bathymetry and hydrographical features (Ingram and El-Sabh, 1990). The Upper St. Lawrence Estuary (USLE), from Île d’Orléans, just downstream of Québec City, to Tadoussac, near the mouth of the Saguenay Fjord, covers an area of 3,470 km². It is relatively narrow (2 to 24 km wide) and mostly shallow (depths less than 30 m; d’Anglejan, 1990), and features an uneven, fairly complex bottom topography characterized by several disconnected channels and troughs separated by ridges and islands (El-Sabh and Murty, 1990). Topographically modified motions give rise to very large tidal ranges and currents (up to 10 m and 3 m s⁻¹, respectively; Mertz and Gratton, 1990). In this tidally energetic region, wind mixing is one to two orders of magnitude smaller than tidal mixing (Painchaud et al., 1995). Owing to the resuspension of bottom sediments (tide, wind or wave generated) and the net non-tidal estuarine circulation (d’Anglejan and Smith, 1973), a well-developed turbidity maximum stretches between Île d’Orléans and Île-aux-Coudres (Painchaud and Therriault, 1989) where suspended particulate matter concentrations vary from 10 to more than 200 mg l⁻¹ (Silverberg and Sundby, 1979). The sources of particulate organic matter (POM) in the estuary are still debated (Gearing and Pocklington, 1990). Carbon isotope studies indicate that less than half of the POM is derived from terrestrial sources (Pocklington and Leonard, 1979) and is quite refractory to biodegradation (Lucotte et al., 1991), whereas the major contributor to POM is believed to be “fresh” organic matter, i.e., living or recently living material, of river-borne origin (Tan and Strain, 1983; Hélie and Hillaire-Marcel, 2006). During the spring freshet in April-May, when freshwater discharge delivers 40 % of the annual
solid inputs to the estuary, the input of terrigenous POM is equivalent to the average
POM kept in suspension in the turbidity maximum (Lucotte, 1989).

The Lower St. Lawrence Estuary (LSLE) is fairly unique in that its character is more
oceanic than most estuaries due to its grand size in all three dimensions and
unimpeded connection with Labrador and Slope waters from the Atlantic Ocean (El-
Sabgh and Silverberg, 1990). Relative to the USLE, the LSLE is much larger (9,350 km²;
d’Anglejan, 1990), wider (30 to 50 km) and deeper (~300 m), and displays a smoother,
less variable bottom topography. Tidal currents are weaker (on the order of 30 cm s⁻¹
or less; Mertz and Gratton, 1990) and, under these less turbulent conditions, the Lower
Estuary is the major sink of continental inputs to the St. Lawrence system. Most (~75 %)
of the terrigenous POM carried by the St. Lawrence River is deposited on the LSLE floor
(Lucotte et al., 1991). The dominant bathymetric feature is the Laurentian Channel (or
Trough), a deep, central, U-shaped glaciated valley that extends 1,240 km from the
eastern Canadian continental shelf break through the GSL and into the LSLE
(d’Anglejan, 1990). The termination (head) of the Laurentian Channel at an abrupt and
shallow sill near Tadoussac, at the mouth of the Saguenay Fjord, marks the region of
transition between the Upper and Lower Estuary and is an area of complex tidal
phenomena (Gratton et al., 1988). Due to rapid shoaling, tidal movements, including
internal tides and strong flows over the steep sill, locally generate significant mixing of
near-surface waters with deeper saline waters, resulting in an important feeding habitat
for several large marine mammals (Dufour and Ouellet, 2007) as well as a nutrient-rich
surface layer that flows continuously through the LSLE (Coote and Yeats, 1979; Saucier
and Chassé, 2000). The Lower Estuary’s seaward outflow, together with the Gaspé
Current, a rapidly moving coastal jet, are a major input of nutrients and zooplankton to
the near-surface waters of the GSL (Coote and Yeats, 1979; Plourde and Runge, 1993).
Mesoscale features such as coastal jets, internal Kelvin waves, baroclinic eddies and
unstable waves are all possible due to strong Coriolis effects (Ingram and El-Sabh,
1990).

The Lower St. Lawrence Estuary is one of the most laterally stratified estuaries in
the world (Larouche et al., 1987). During summertime, the SLE can be described as a
three-layer system on the basis of thermal stratification (Gratton et al., 1988). Each
spring, a new surface layer flow is initiated by the freshwater runoff from the St.
Lawrence River, Saguenay Fjord and rivers on the north shore of the estuary (Dufour and Ouellet, 2007; see Fig. 1). Discharge from the St. Lawrence River (mean annual discharge of 10,000 m$^3$ s$^{-1}$, peaking at 15,000 m$^3$ s$^{-1}$ during the spring freshet; Painchaud and Therriault, 1989) provides about 80% of the total freshwater input to the estuary (Ingram and El-Sabh, 1990), whereas the combined runoff from the Saguenay and Manicouagan rivers accounts for most of the remainder (Tee, 1990). The warm and relatively fresh surface layer (0 to 30 m) overlies the cold intermediate layer or CIL (30-150 m deep; $S_p = 32.0$ to $32.6$) that is formed by the advection of the wintertime surface mixed layer from the GSL (Galbraith, 2006). Below the CIL, a warmer (2 to 6 °C) and saltier ($S_p = 33$ to 35) bottom layer (>150 m deep), originating from the mixing of western-central Atlantic and Labrador shelf waters that intrude at depth primarily through Cabot Strait, flows sluggishly landward (~0.5 cm s$^{-1}$; Bugden, 1988) toward the head region of the Laurentian Channel (Saucier et al., 2003; Gilbert et al., 2005).

### 2.2. Water-column sampling and analytical procedures

Water samples were collected on board the RV Coriolis II during ten research cruises within the St. Lawrence Estuary and Gulf in the spring or summer season between 2003 and 2016. The sampling locations are shown in Fig. 2. Samples were taken at ~3m depth and throughout the water column, along the central axis of the estuary and Laurentian Channel, and in the freshwaters of the St. Lawrence River near Québec City, to obtain freshwater and seawater end-member values. A comprehensive dataset describing the inorganic carbon chemistry in surface mixed layer waters was compiled from field measurements of temperature (T), practical salinity ($S_p$), pH$_{NBS}$ and/or pH$_T$, total alkalinity (TALK), soluble reactive phosphate (SRP), and dissolved silicate (DSi), nitrate (NO$_3$) and oxygen (DO).

$T$ and $S_p$ were determined in situ using the conductivity-temperature-depth (CTD) probe (SeaBird SBE 911) mounted on the sampling rosette. The temperature probe was calibrated by the manufacturer, whereas the conductivity sensor was calibrated by the manufacturer and recalibrated using discrete salinity samples collected throughout the water column and analyzed on a Guildline Autosal 8400 salinometer calibrated with IAPSO standard seawater. Water samples destined for pH
and TAlk measurements were transferred directly from the 12L Niskin bottles mounted on the CTD-rosette system to, respectively, 125mL plastic bottles without headspace and 250mL glass bottles as soon as the rosette was secured onboard. In the latter case, a few crystals of HgCl$_2$ were added before the bottle was sealed with a ground-glass stopper and Apiezon® Type-M high-vacuum grease.

pH was determined onboard at 25 °C, potentiometrically on the NBS/NIST scale (infinite dilution convention, pH$_{\text{NBS}}$) for low salinity waters (S$_P$ < 5) and potentiometrically and/or colorimetrically on the total hydrogen ion concentration scale (constant ionic medium convention, pH$_T$) for higher salinity waters. Potentiometric pH measurements were carried out using a Radiometer Analytical® GK2401C combination glass electrode connected to a Radiometer Analytical® PHM84 pH/millivolt-meter. Prior to each measurement, the electrode was calibrated against three NIST-traceable buffer solutions: pH-4.00, pH-7.00 and pH-10.00 at 25°C. The electrode response to these buffers was then least-squares fitted to obtain the Nernstian slope. For S$_P$ > 5, pH measurements were converted to the pH$_T$ scale using TRIS buffer solutions prepared at S$_P$ = 5, 15, 25, or 35 for which the pH$_T$ was assigned at 25°C (Millero, 1986). Colorimetric pH measurements were carried out using a Hewlett-Packard UV-Visible diode array spectrophotometer (HP-8453A) and a 5cm quartz cell after thermal equilibration of the sampling bottles in a constant temperature bath at 25.0 ± 0.1°C. Phenol red (Robert-Baldo et al., 1985) and m-cresol purple (Clayton and Byrne, 1993) were used as color indicators. The pH$_T$ of the water samples and buffer solutions were calculated according to the equation of Byrne (1987). The reproducibility of the pH measurements was typically better than ± 0.003.

TAlk was measured at McGill University using an automated Radiometer (TitraLab865®) potentiometric titrator and a Red Rod® combination pH electrode (pHC2001). The dilute HCl titrant was calibrated prior, during, and after each titration session using certified reference materials provided by Andrew Dickson (Scripps Institute of Oceanography). Raw titration data were processed with a proprietary algorithm specifically designed for shallow end-point detection. The reproducibility of the method was better than 0.5 %.
DO concentrations were determined by Winkler titration (Grasshoff et al., 1999) on distinct water samples recovered directly from the Niskin bottles. The relative standard deviation, based on replicate analyses of samples recovered from the same Niskin bottle, was better than 1%. These measurements further served to calibrate the SBE-43 oxygen probe mounted on the rosette. For the determination of nutrient concentrations, aliquots of the water samples taken from the Niskin bottles were syringe filtered through a 0.45 µm Millipore polycarbonate (MA) filter. DSI was measured onboard on the same day of sampling using the method described in Grasshoff et al. (1999). Water samples destined for NO₃ and SRP measurements were transferred, respectively, into acid-washed 15ml polyethylene and borosilicate tubes, quickly frozen and stored at -20 °C. Their concentrations were determined using standard colorimetric methods adapted from Grasshoff et al. (1999) with a SEAL Autoanalyzer III at the Institut des Sciences de la Mer de Rimouski. The analytical detection limit was 0.04 µM for NO₃, 0.05 µM for SRP and 0.1 µM for DSI. Based on replicate analyses of the standards, the reproducibility of these measurements was typically 1%.

The in situ pressure and density of the samples were calculated from the Thermodynamic Equation of Seawater - 2010 (TEOS-10) using the Gibbs Seawater (GSW) Oceanographic Toolbox (MATLAB-version 3.05; McDougall and Barker, 2011). All field measurements reported in µmol L⁻¹ were converted to µmol kg⁻¹ using the in situ density data.

2.3. Calculation of aqueous pCO₂

2.3.1. pCO₂ in surface mixed layer waters

Aqueous pCO₂ (pCO₂(water)) is defined as the partial pressure of carbon dioxide in wet (100% water-saturated) air that is in equilibrium with the water sample. Values of pCO₂(water) (µatm) and total dissolved inorganic carbon, or DIC (µmol kg⁻¹), were calculated from the measured pH (total or NBS scale) and TA(κ) (µmol kg⁻¹), at in situ temperature (°C), salinity (Sₛ) and pressure (dbar), using the program CO2SYS (MATLAB-version 1.1; van Heuven et al., 2011) and the carbonic acid dissociation constants, K₁ and K₂, of Cai and Wang (1998) for estuarine waters. Wherever data were
available, the contributions to TAlk from phosphate and silicate were included.

Although the K₁ and K₂ formulations from Lueker et al. (2000) are recommended for best practices by Dickson et al. (2007), they are not suitable for the low-salinity conditions found in estuaries (S_P < 19) (Orr et al., 2015). The revised equations for K₁ and K₂ from Cai and Wang (1998) are applicable over a larger range of salinities (0 to 40) and, thus, were used to examine the carbonate system in the estuarine waters of our study area.

This study focuses on the inorganic carbon dynamics of near-surface waters. To obtain individual data points of surface-water pCO₂ at each sampling location, the pCO₂(water) data in the surface mixed layer (SML) were averaged. The SML is the site of active air-sea interaction where heat and gases are exchanged directly with the atmosphere and within which physical (temperature, salinity, density) and chemical (dissolved gases) properties are vertically homogeneous due to turbulent mixing (Sprintall and Tomczak, 1992). The lower limit to air-sea interaction, i.e., the mixed layer depth (MLD), is demarcated by the pycnocline, a sharp density gradient that generally coincides with both a temperature (thermocline) and salinity (halocline) gradient. Here, we determine the thickness of the SML using a density-based criterion, in which the MLD is defined as the depth at which a threshold difference of 0.03 kg m⁻³ from the sea surface occurs (de Boyer Montégut et al., 2004). In the following sections, surface-water pCO₂ will be taken to mean the SML-averaged pCO₂.

2.3.2. Sources of error in pCO₂ calculation

The importance of using appropriate formulations of K₁ and K₂ in estuarine waters is shown by the discrepancies in calculated values of pCO₂ (pH/TAlk) at low salinities (Fig. 3). The percent difference between pCO₂ values calculated from the dissociation constants of Cai and Wang (1998) and those from Lueker et al. (2000) was on average 2.12%. Calculations differed by 1.13-18.5% at low salinities (S_P < 19), whereas, at S_P > 19, the calculated pCO₂ were much closer (only ~1.49% difference).

The K₁ and K₂ formulations of Millero (2010), i.e., the most recent alternative proposed for estuarine waters (S_P = 1 to 50), yielded pCO₂ values that differed substantially from those of Cai and Wang (1998) at S_P < 19, with the largest divergence reaching 34.5% (average difference of 13.7%). At S_P = 0, the pCO₂ values calculated from Cai and
Wang (1998) compared very well with those given by the Millero (1979) constants for freshwater (difference of only ~0.08 %), whereas the pCO$_2$ values calculated from Millero (2010) showed very poor agreement with the freshwater results (differing by ~34.3 %). These discrepancies highlight the need for new or revised measurements of the carbonic acid dissociation constants under estuarine conditions (in brackish waters) especially at $S_F < 5$. Studies which use the best-practices formulations of $K_1$ and $K_2$ to calculate estuarine pCO$_2$ may underestimate CO$_2$ emissions at low salinities, whereas those that implement the Millero (2010) formulations may produce overestimates.

Another potential source of error in the calculation of pCO$_2$ (pH/TAlk) in low-salinity estuarine waters is the contribution of organic species to the total alkalinity. While the contributions of borate, phosphate and silicate species are taken into consideration by the program CO2SYS, the magnitude of organic alkalinity (Org-Alk) is usually assumed to be small or negligible, and is simply ignored when using TAlk to calculate pCO$_2$ in open ocean waters. In riverine and coastal waters, however, the contribution of organic species to the TAlk can be significant (Yang et al., 2015). Rivers draining organic-rich soils and non-carbonate rocks have low DIC concentrations (a few hundred μmol L$^{-1}$) that are often exceeded by dissolved organic carbon (DOC) concentrations (Abril et al., 2015). As discussed by Hunt et al. (2011), a significant contribution of Org-Alk (the organic acid anions in DOC) leads to an overestimation of calculated pCO$_2$ using any program that accounts only for the inorganic species contributing to TAlk. A comparison of the calculated TAlk (DIC/pH) (direct DIC measurements were carried out in 2014) and the measured TAlk reveals that Org-Alk is on the order of ~20 μmol kg$^{-1}$ for the St. Lawrence River end-member, whereas it is as high as ~120 μmol kg$^{-1}$ for the Saguenay River end-member (A. Mucci, pers. comm.). Given that TAlk exceeds ~1000 μmol kg$^{-1}$ throughout our study area and the Saguenay River contribution to the surface waters of the SLE is limited (at most ~6 % at the head of the estuary; Mucci et al., submitted), consideration of the Org-Alk in the calculation of pCO$_2$ (pH/TAlk) yielded values that were at most 1.9 % different from those uncorrected for Org-Alk. Bearing in mind the uncertainties in the $K_1$ and $K_2$ formulations as well as the measurement uncertainties, the influence of Org-Alk on the calculation of pCO$_2$ (pH/TAlk) did not represent a significant source of error.

2.4. Temperature normalization of pCO$_2$
The effect of temperature on aqueous pCO$_2$ is primarily the manifestation of changes in the solubility of CO$_2$ gas in water (Takahashi et al., 1993). The temperature dependence of pCO$_2$ in seawater, i.e., $\partial \ln(pCO_2) / \partial T = 0.0423 \, ^\circ C^{-1}$, was determined experimentally by Takahashi et al. (1993) on a single North Atlantic surface water sample with $S_P = 35.380$ under isochemical conditions. As this oft-used approximation for thermally induced changes in pCO$_2$ was derived from direct measurements in seawater, we use a different approach to remove the temperature effect on the estuarine pCO$_2$ in our study area. The in situ pCO$_2$ were normalized to the average surface-water temperature (SST = 7.82 °C) using the temperature normalization method of Jiang et al. (2008) in which pCO$_2$ are re-calculated from the TAlk and DIC data at a common temperature. The results yielded a temperature coefficient of $\partial \ln(pCO_2) / \partial T = 0.0404 \, ^\circ C^{-1}$ ($R^2 = 0.99$), in excellent agreement with that of Takahashi et al. (1993). The pCO$_2$ changes due to temperature deviations from the mean SST were calculated as:

$$\Delta pCO_2(\text{thermal}) = pCO_2(\text{in-situ}) - pCO_2(\text{temp-norm})$$  \hspace{1cm} (1)

where $\Delta pCO_2(\text{thermal})$ is the pCO$_2$ change due to temperature effects, pCO$_2(\text{in-situ})$ is the in situ pCO$_2$ and pCO$_2(\text{temp-norm})$ is the temperature-normalized pCO$_2$. Since changes in pCO$_2$ at a constant temperature primarily reflect changes in DIC, the spatial variations in temperature-normalized pCO$_2$ were attributed to the combined influences of non-thermal processes, i.e., water mass mixing and biological activity. Theoretically, pCO$_2(\text{temp-norm})$ can be further partitioned into the pCO$_2$ change due to biology and pCO$_2$ due to mixing through an analysis of the water mass structure, e.g., an optimum multiparameter (OMP) water mass analysis. Results from its application will be presented in a subsequent study.

### 2.5. Air-sea CO$_2$ flux estimation

Air-sea CO$_2$ gas exchange ($F$, mmol C m$^{-2}$ d$^{-1}$) at each sampling location was estimated as follows:

$$F = k \cdot K_0 \cdot (pCO_2(\text{water}) - pCO_2(\text{air}))$$  \hspace{1cm} (2)
where \( k \) (cm h\(^{-1}\)) is the gas transfer velocity of CO\(_2\), \( K_0 \) (mol kg\(^{-1}\) atm\(^{-1}\)) is the solubility coefficient of CO\(_2\) at in situ surface-water temperature and salinity (Weiss, 1974), and \( p_{CO_2}^{(\text{water})} \) and \( p_{CO_2}^{(\text{air})} \) (µatm) are the partial pressures of CO\(_2\) in the water and the air, respectively. The difference between \( p_{CO_2}^{(\text{water})} \) and \( p_{CO_2}^{(\text{air})} \) (\( \Delta p_{CO_2} \)) determines the direction of gas exchange across the air-sea interface. Positive values of \( F \) indicate release of CO\(_2\) by the surface water, whereas negative values indicate CO\(_2\) uptake by the surface water. Conversion factors were applied to express the final \( F \) with the aforementioned units.

Atmospheric \( p_{CO_2} \) (\( p_{CO_2}^{(\text{air})} \)) was calculated using the monthly averages of the measured mole fraction of CO\(_2\) in dry air (\( x_{CO_2} \), at the greenhouse gas observational station in Fraserdale, Ontario) obtained from the Climate Research Division at Environment and Climate Change Canada. The mean \( p_{CO_2}^{(\text{air})} \) in the sampling month was computed using the relationship (Takahashi et al., 2002):

\[
p_{CO_2}^{(\text{air})} = x_{CO_2} \cdot (P_b - P_w)
\]

where \( x_{CO_2} \) is in ppm, \( P_b \) (atm) is the atmospheric (or barometric) pressure at the sea surface, and \( P_w \) (atm) is the equilibrium (or saturation) water vapor pressure at in situ surface-water temperature and salinity (Weiss and Price, 1980). One-month averaged barometric pressures were calculated using the hourly station pressure data from Environment Canada at the following weather observing stations: Québec/Jean Lesage International Airport (Upper Estuary), Mont-Joli Airport (Lower Estuary), and Gaspé Airport (Gulf of St. Lawrence). The \( P_b \) at station elevation was converted to mean sea level pressure using the formula of Tim Brice and Todd Hall (NOAA’s National Weather Service, http://www.weather.gov/epz/wxcalc_stationpressure).

The formulation of the gas transfer velocity, \( k \), is the largest source of error in the computation of air-sea CO\(_2\) fluxes (Borges et al., 2004a,b). Properly constraining values of \( k \) in estuaries is problematic (Raymond and Cole, 2001) due to their hydrodynamic and geomorphologic complexity (Abril et al., 2000). Gas transfer is thought to be regulated by turbulence at the air-water interface (Wanninkhof, 1992). Wind stress plays a key role in the generation of turbulence at the ocean surface through the
transfer of momentum to waves and currents (Ho et al., 2011), whereas, in estuarine environments and especially macrotidal estuaries, surface turbulence can be created by interactions of wind forcing, tidal currents and boundary friction (Zappa et al., 2003, Borges et al., 2004a,b; Zappa et al., 2007) and, in turbid estuaries, attenuated by suspended material (Abril et al., 2009). The turbulence generated from bottom stress varies with water depth and tidal velocity (Raymond et al., 2000), and is important only in shallower estuaries with high current speeds (Cerco, 1989). Raymond and Cole (2001) have shown that wind stress controls turbulence at the air-water interface for all systems with depths greater than 10 meters (at depths < 10 m, either wind or bottom stress may dominate).

Several different predictive relationships between wind speed and gas transfer have been proposed based on laboratory and field studies. Here, we estimate the gas transfer velocity from short-term (or steady) wind speed measurements using the equations of Wanninkhof (1992) revised by Wanninkhof (2014) and Raymond and Cole (2001):

\[ k_{W-14} = 0.251 u^2 (Sc / 660)^{-0.5} \]  \hspace{1cm} (4)

\[ k_{R&C-01} = 1.91 e^{0.35u} (Sc / 660)^{-0.5} \]  \hspace{1cm} (5)

where \( u \) is the wind speed (m s\(^{-1}\)) and \( Sc \) is the Schmidt number (\( Sc = \mu/D \), where \( \mu \) is the kinematic viscosity of the water and \( D \) is the diffusion coefficient) for CO\(_2\) gas in solution. The Schmidt number for CO\(_2\) in seawater at 20 °C is 660 and was adjusted to \( Sc=600 \) for freshwater. Hourly wind speed data were obtained from Environment Canada at the aforementioned weather observing stations, and averaged over the sampling month to obtain short-term wind speeds. The correction to a common Schmidt number was performed using the equations of Wanninkhof (1992) for the temperature dependence of \( Sc \) for CO\(_2\) gas in seawater (\( Sp=35 \)) and freshwater, respectively, and assuming that \( k \) is proportional to \( Sc^{-0.5} \).
Because of increased turbulence, one would expect k values calculated from estuarine parameterizations to be higher than those predicted from oceanic parameterizations at equivalent wind speeds (Abril et al., 2000). Within the confines of the SLE, estimates of k using the Wanninkhof (2014) relationship ranged from 1.6 to 4.5 cm h\(^{-1}\) whereas those calculated from Raymond and Cole (2001) were between 3.8 and 8.1 cm h\(^{-1}\). Hence, we take the air-sea CO\(_2\) flux values calculated with \(k_{W-14}\) to be the theoretical lower limit of gas exchange (\(F_{W-14}\)), whereas those computed from \(k_{R&C-01}\) represent the upper limit of gas exchange (\(F_{R&C-01}\)).

In order to estimate the area-averaged CO\(_2\) flux in the SLE, the estuary proper was divided into five segments, with each section containing at least one sampling location. The fluxes for each segment were normalized to the sectional surface area and then summed to obtain a spatially integrated CO\(_2\) flux (\(F_{area-avg}\)) for air-sea gas exchange in the whole estuary as follows (Jiang et al., 2008):

\[
F_{area-avg} = \frac{\sum F_i S_i}{\sum S_i}
\]  

(6)

where \(F_i\) is the average of all the fluxes within segment i, and \(S_i\) is the surface area of segment i. Sectional surface areas were tabulated in MATLAB using the land mask of eastern Canada obtained from Fisheries and Oceans Canada. An area-averaged CO\(_2\) flux was obtained for both the upper and lower limits of gas exchange in the SLE. These two final estimates are assumed to bracket the real areal CO\(_2\) flux.

2.6. Conceptual framework for the analysis of variations in biogenic gas concentrations

A comparison of the distribution of biologically reactive dissolved gases, e.g., CO\(_2\) and O\(_2\), can provide useful information about the physical (thermal) and biological processes controlling their concentrations (Richey et al., 1988). Temperature-related gas solubility effects occur in the same direction for CO\(_2\) and O\(_2\), whereas biological production and respiration affect CO\(_2\) and O\(_2\) in opposite directions. Following the approach of Carrillo et al. (2004), the saturation states (or % saturation) of pCO\(_2\) and DO, with respect to the atmosphere, were compared in order to determine the relative importance of temperature changes (heating or cooling) and biological activity.
(photosynthesis or respiration) in the surface waters at each sampling location. The pCO$_2$ percent saturation ($\%$pCO$_2$$_{(sat)}$) was calculated as follows:

$$\%pCO_2_{(sat)} = \left( \frac{pCO_2_{(water)}}{pCO_2_{(air)}} \right) \cdot 100$$  \hspace{1cm} (7)

The DO percent saturation ($\%$DO$_{(sat)}$) was calculated as:

$$\%DO_{(sat)} = \left( \frac{DO}{DO^*} \right) \cdot 100$$  \hspace{1cm} (8)

where DO$^*$ is the equilibrium DO concentration ($\mu$mol kg$^{-1}$) at in situ surface-water temperature and salinity (Benson and Krause, 1984).

According to the method of Carrillo et al. (2004), data points fall into one of four quadrants on a graph of $\%$DO$_{(sat)}$ versus $\%$pCO$_2$$_{(sat)}$, with the origin at 100 % saturation for both gases. Quadrant I (upper left; supersaturated DO, undersaturated pCO$_2$) suggests net photosynthesis, Quadrant II (upper right; supersaturated DO and pCO$_2$) indicates the effects of heating, Quadrant III (lower right; undersaturated DO, supersaturated pCO$_2$) implies net respiration, and Quadrant IV (lower left; undersaturated DO and pCO$_2$) represents the effects of cooling. Although general patterns become apparent, we urge caution in the interpretation of these results as significant limitations apply. Surface-water CO$_2$ and O$_2$ may be acted upon by other forcings such as air-sea gas exchange. The net transfer of CO$_2$ and O$_2$ gases occurs across the air-sea interface whenever their partial pressures in the surface water differ from those in the atmosphere. Because of their differential gas exchange rates (i.e., O$_2$ exchanges ~19 times faster than CO$_2$; Peng et al., 1987), CO$_2$ and O$_2$ dynamics may be decoupled in surface waters, causing an asymmetry in the observed CO$_2$:O$_2$ relationship (Carrillo et al., 2004).

3. Results and discussion

3.1. Spatial variability of surface-water pCO$_2$

Data were compiled from all ten cruises to describe the inorganic carbon chemistry in the SML waters of the St. Lawrence River, Upper Estuary, Lower Estuary
and Gulf (Table 1). Large spatial variations in surface-water pCO₂ were observed in the EGSL system, with values ranging from 139 to 765 µatm (453 ± 133 µatm) during the spring/summer sampling periods. The pCO₂ were higher in the USLE (571 ± 72 µatm) than in the LSLE (394 ± 117 µatm) and GSL (354 ± 82 µatm), whereas the atmospheric pCO₂ showed less variability, ranging from 372 to 405 µatm, during the sampling months. From Fig. 4, it can be seen that the USLE was always a CO₂ source (pCO₂(water)) above atmospheric equilibrium) while the LSLE and GSL were generally either a CO₂ sink or nearly neutral (pCO₂(water) below or close to atmospheric equilibrium).

Within the confines of the SLE, the surface-water pCO₂ generally decreased with increasing distance from the head of the estuary (Île d’Orléans) and along the salinity gradient (Fig. 4 and 5). The highest values of pCO₂ were observed near the landward limit of the salt water intrusion in the SLE’s upper reaches, in the vicinity of the Cap Tourmente intertidal flats and marshes. This area (3 x 10⁶ m²) is located along the core of the estuary’s maximum turbidity zone (MTZ) (Lucotte and d’Anglejan, 1986). The lowest surface-water pCO₂ were found downstream of the MTZ in the lower reaches of the SLE near Pointe-des-Monts, where the channel widens into the gulf. Due to favorable environmental conditions (nutrients, light, stratification), phytoplankton blooms typically occur in late spring or early summer in the LSLE (Zakardjian et al., 2000), with maximal biological production occurring in its downstream portion due to the mixing of cold, nutrient-rich waters, upwelled at the head of the Laurentian Channel, with warmer freshwaters flowing in from the north shore rivers (Savenkoff et al., 1994). Seaward from the estuary-gulf boundary, the pCO₂ gradually increased from 207 to 478 µatm, coinciding with a large increase in surface-water temperature (T = 3.9 to 13.7 °C).

The spatial variability of surface-water pCO₂ due to temperature variations was removed by normalizing the pCO₂ data to a common temperature (T = 7.82 °C). A comparison of the in situ and corresponding temperature-normalized pCO₂ shows that spatial changes in water temperature lower or raise the pCO₂ by -170 to 181 µatm. The maximum (minimum) values of ΔpCO₂(thermal), expressed as a percent change, were 38 % (-14 %) in the USLE, 24 % (-31 %) in the LSLE, and 29 % (-17 %) in the GSL. Temperature normalization, however, removed only a small part of the overall spatial variability of surface-water pCO₂ (Fig. 6). Given that the spread of the temperature-
normalized pCO$_2$ data remained large (153–668 µatm; 449 ± 133 µatm), most of the spatial heterogeneity in surface-water pCO$_2$ can be explained by non-thermal processes (water mass mixing and/or biological activity).

### 3.2. Air-sea CO$_2$ flux and spatial integration

Large spatial variability of the air-sea CO$_2$ flux was observed in the EGSL system during the spring/summer sampling periods, with fluxes ranging from -21.9 to 28.4 mmol m$^{-2}$ d$^{-1}$ (Fig. 7). Values of F were always positive in the USLE (2.0 to 28.4 mmol m$^{-2}$ d$^{-1}$) and either negative or positive in the LSLE (-21.9 to 15.1 mmol m$^{-2}$ d$^{-1}$) and GSL (-8.4 to 3.6 mmol m$^{-2}$ d$^{-1}$). As expected, $F_{R&C:01}$ (estuarine parameterization of k) were larger than $F_{W:14}$ (oceanic parameterization of k) due to the inherently greater surface turbulence in estuarine systems. The average difference between fluxes calculated from Wanninkhof (2014) and those from Raymond and Cole (2001) was 71.5%. Details of the k and F values given by each parameterization of the gas transfer velocity are shown in Table 2. Irrespective of the parameterization, the calculated CO$_2$ fluxes were more positive in the USLE (9.2 ± 5.3 mmol m$^{-2}$ d$^{-1}$) than in the LSLE (0.8 ± 7.1 mmol m$^{-2}$ d$^{-1}$) and GSL (-1.1 ± 3.0 mmol m$^{-2}$ d$^{-1}$).

To obtain an area-averaged CO$_2$ flux for the whole estuary, the SLE was divided into five segments, with each segment containing at least one sampling location. The data used to calculate the area-averaged CO$_2$ flux are listed in Table 3. Overall, the SLE served as a weak source of CO$_2$ to the atmosphere at the time of sampling, with an area-averaged CO$_2$ degassing flux of 1.00 to 2.06 mmol C m$^{-2}$ d$^{-1}$ (0.37 to 0.75 mol C m$^{-2}$ yr$^{-1}$). This efflux compares favorably with that of the Delaware Estuary (2.4 ± 4.8 mol C m$^{-2}$ yr$^{-1}$; Joesoef et al., 2015), another large estuarine system with a long water residence time, but is significantly lower than estimates in the marine-dominated Sapelo and Doboy Sound estuaries (10.5 to 10.7 mol C m$^{-2}$ yr$^{-1}$; Jiang et al., 2008).

From a compilation of 165 estuaries worldwide, almost all systems, with the exception of those in the Arctic (-1.1 mol C m$^{-2}$ yr$^{-1}$), serve as sources of CO$_2$ to the atmosphere (Chen et al., 2013). Chen et al. (2013) concluded that the world’s upper estuaries ($S_F < 2$) are strong sources (39.0 ± 55.7 mol C m$^{-2}$ yr$^{-1}$), mid estuaries ($2 < S_F < 25$) are moderate sources (17.5 ± 34.2 mol C m$^{-2}$ yr$^{-1}$), and lower estuaries ($S_F > 25$) are weak sources (8.4 ± 14.3 mol C m$^{-2}$ yr$^{-1}$). Predictably, with its maritime region occupying
almost three-fourths of the total surface area, the SLE behaves like an outer estuary with only small CO₂ evasion. The lack of temporal coverage of surface-water pCO₂ data, however, prevents us from reliably synthesizing an annual air-sea CO₂ flux.

3.3. Major controlling factors of estuarine pCO₂

The pCO₂ in the surface mixed layer is a function of its temperature (T), salinity (S_p), dissolved inorganic carbon (DIC) and total alkalinity (TAlk), as described by the following relationship (Takahashi et al., 1993):

\[
dpCO_2 = (\partial pCO_2/\partial T) dT + (\partial pCO_2/\partial S_p) dS_p + (\partial pCO_2/\partial DIC) dDIC + (\partial pCO_2/\partial TAlk) dTAlk \tag{9}
\]

Through changes in T, S_p, DIC and TAlk, variations in surface-water pCO₂ are mainly controlled by dynamic processes (water mass mixing), thermodynamic processes (temperature and salinity changes), air-sea gas exchange and biological processes (photosynthesis, respiration) (Poisson et al., 1993). Among these, the effects of temperature and DIC, i.e., the DIC inputs from internal (biological activity) and external (physical mixing) sources, are generally the most important drivers of estuarine pCO₂ variability. Whereas the variations in pCO₂ due to water mass mixing and those due to in situ biological activity will be separated in a future study through a modified OMP water mass analysis, here, we evaluate the relative importance of thermal and biological processes in controlling the spatial distribution of pCO₂ in our study area.

To disentangle the biological and temperature effects on pCO₂ variability, the %DO_{(sat)} were plotted against the %pCO₂_{(sat)}, with the origin at 100 % saturation for both gases. This simple approach uses the four possible combinations of %pCO₂_{(sat)}/%DO_{(sat)} as integrated measures of thermally and biologically induced changes. From Fig. 8, it can be seen that microbial respiration was the major driver of pCO₂ variability in the USLE, whereas the effects of photosynthesis and temperature predominated in the LSLE and GSL. In the strongly stratified Lower Estuary, the biological drawdown of pCO₂ surpassed the increasing effect of sea surface warming (Fig. 9). Values of surface-water pCO₂ were mostly below atmospheric level despite increases in temperature (T = -0.32 to 12.6 °C). This pattern is consistent with the
finding that, in spring/summer, the increasing effect of warming on pCO₂ is counteracted by the photosynthetic utilization of CO₂, particularly in a strongly stratified shallow mixed layer (Takahashi et al., 1993). In the Gulf of St. Lawrence, the temperature dependence of pCO₂ exerted a major influence, causing values of surface-water pCO₂ to increase (207 to 478 µatm) concomitantly with temperature (T = 3.9 to 13.7 °C) (Fig. 9).

4. Conclusions

Because of its large physical dimensions and unimpeded connection to the Atlantic Ocean, the St. Lawrence Estuary encompasses both a river-dominated inner estuary, where physical mixing and abiotic processes dominate, and a marine-dominated outer estuary, where biological cycling and oceanic processes prevail. The physical and biogeochemical processes of these contrasting environments are reflected in the spatial distribution of surface-water pCO₂ (139-765 µatm). The shallow, partially mixed Upper Estuary, with a turbidity maximum controlled by tide- and wind-induced turbulence, was, during our sampling period, a source of CO₂ to the atmosphere due to microbial respiration (low biological productivity), whereas the deep, stratified Lower Estuary, with its stable, summertime three-layer vertical structure, was generally a sink of atmospheric CO₂ due to the enhanced biological drawdown of pCO₂ (light availability, nutrient supply, strong stratification).

Overall, the large subarctic St. Lawrence Estuary was a weak source of CO₂ to the atmosphere, with an area-averaged CO₂ degassing flux of 1.00 to 2.06 mmol C m⁻² d⁻¹ (0.37 to 0.75 mol C m⁻² yr⁻¹). This efflux is somewhat smaller than the numerically averaged CO₂ flux per unit area (2.19 mol C m⁻² yr⁻¹) reported from North American estuaries by Chen et al. (2013), highlighting their relatively small contribution (~12 %) to global estuarine CO₂ emissions. The pronounced shift in source/sink dynamics in the St. Lawrence Estuary, between its river-dominated (9.2 ± 5.3 mmol m⁻² d⁻¹) and marine-dominated (0.8 ± 7.1 mmol m⁻² d⁻¹) regions, is consistent with the conclusions of the comparative study carried out by Jiang et al. (2008) that showed large differences in CO₂ degassing between riverine (inner) and maritime (outer) estuaries. Given the limited research on CO₂ dynamics in large estuaries and bay systems, which cover approximately one-half of the estuarine surface area on the U.S. east coast, as
well as the large uncertainties in the calculation of pCO₂ (carbonic acid dissociation constants, organic alkalinity contribution), current global-scale estimates of estuarine CO₂ degassing may be overestimated. To better constrain the role of large estuaries/bays in the estuarine carbon cycle, more extensive spatial and temporal coverage of pCO₂ measurements is needed.

Data availability
Data presented in this paper (Figures 4 and 8) are available upon request from one of the authors (alfonso.mucci@mcgill.ca).

Author contribution
A.D. and A.M. conceived the project. A.M. acquired and processed the data prior to 2016. A.D. conducted the data analysis and wrote the first draft of the paper whereas A.M. provided editorial and scientific recommendations.

Competing interests
The authors declare that they have no conflict of interest.

Acknowledgements
We wish to thank the Captains and crews of the RV Coriolis II for their unwavering help over the years. We also wish to acknowledge Gilles Desmeules and Michel Rousseau for their dedicated electronic and field sampling support as well as Constance Guignard for her help in cruise preparation and field data acquisition. Most of the data acquisition was carried out opportunistically on research cruises funded by grants to A.M. or Canadian colleagues by the Natural Sciences and Engineering Research Council of Canada (NSERC) whereas the work was funded by a Regroupement Stratégique grant from the Fonds Québécois de Recherche Nature et Technologies (FQRNT) to GEOTOP as well as NSERC Discovery and MEOPAR grants to A.M. A.D. wishes to thank the Department of Earth and Planetary Sciences at McGill for financial support in the form of scholarships and assistantships.

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Table 1. Mean, standard deviation and range of the surface-water temperature (T), practical salinity (S_P), dissolved inorganic carbon (DIC), total alkalinity (TAlk) and in situ partial pressure of CO₂ (pCO₂) in the St. Lawrence River (near Québec City), Upper Estuary (Île d’Orléans to Tadoussac), Lower Estuary (Tadoussac to Pointe-des-Monts) and Gulf (Pointe-des-Monts to Cabot Strait) during all sampling months.

<table>
<thead>
<tr>
<th></th>
<th>T (°C)</th>
<th>S_P</th>
<th>DIC (µmol kg⁻¹)</th>
<th>TAlk (µmol kg⁻¹)</th>
<th>pCO₂ (µatm)</th>
</tr>
</thead>
<tbody>
<tr>
<td>River</td>
<td>14.2 ± 3.9</td>
<td>0.03 ± 0.05</td>
<td>1242 ± 132</td>
<td>1204 ± 99</td>
<td>604 ± 76</td>
</tr>
<tr>
<td>(N=3)</td>
<td>(9.8-17.2)</td>
<td>(0-0.09)</td>
<td>(1148-1335)</td>
<td>(1124-1314)</td>
<td>(550-658)</td>
</tr>
<tr>
<td>Upper</td>
<td>9.6 ± 3.6</td>
<td>10.9 ± 8.0</td>
<td>1514 ± 242</td>
<td>1492 ± 272</td>
<td>571 ± 72</td>
</tr>
<tr>
<td>(N=46)</td>
<td>(4.2-17.4)</td>
<td>(0-24.5)</td>
<td>(1081-2005)</td>
<td>(969-2030)</td>
<td>(435-765)</td>
</tr>
<tr>
<td>Lower</td>
<td>5.6 ± 2.8</td>
<td>26.8 ± 2.7</td>
<td>1843 ± 89</td>
<td>1962 ± 87</td>
<td>394 ± 117</td>
</tr>
<tr>
<td>(N=60)</td>
<td>(-0.32-12.6)</td>
<td>(21.2-32.7)</td>
<td>(1634-2083)</td>
<td>(1752-2185)</td>
<td>(139-578)</td>
</tr>
<tr>
<td>Gulf</td>
<td>8.8 ± 3.1</td>
<td>30.1 ± 1.5</td>
<td>1936 ± 64</td>
<td>2094 ± 61</td>
<td>354 ± 82</td>
</tr>
<tr>
<td>(N=30)</td>
<td>(3.9-13.7)</td>
<td>(25.5-31.5)</td>
<td>(1761-2032)</td>
<td>(1921-2175)</td>
<td>(207-478)</td>
</tr>
</tbody>
</table>
Table 2. Mean, standard deviation and range of ΔpCO\textsubscript{2}, k\textsubscript{W-14}, k\textsubscript{R&C-01}, F\textsubscript{W-14} and F\textsubscript{R&C-01} in the surface waters of the St. Lawrence River, Upper Estuary, Lower Estuary and Gulf during all sampling months. k\textsubscript{W-14} are the gas transfer velocities given by the Wanninkhof (2014) relationship, whereas k\textsubscript{R&C-01} are those given by Raymond and Cole (2001). Values of F\textsubscript{W-14} are taken to be the theoretical lower limit of air-sea gas exchange, whereas values of F\textsubscript{R&C-01} are the upper limit; the extreme F data points are shown in bold.

<table>
<thead>
<tr>
<th></th>
<th>ΔpCO\textsubscript{2} (µatm)</th>
<th>k\textsubscript{W-14} (cm h\textsuperscript{-1})</th>
<th>k\textsubscript{R&amp;C-01} (cm h\textsuperscript{-1})</th>
<th>F\textsubscript{W-14} (mmol m\textsuperscript{-2} d\textsuperscript{-1})</th>
<th>F\textsubscript{R&amp;C-01} (mmol m\textsuperscript{-2} d\textsuperscript{-1})</th>
</tr>
</thead>
<tbody>
<tr>
<td>River</td>
<td>217 ± 99</td>
<td>3.0 ± 1.4</td>
<td>6.1 ± 2.0</td>
<td>5.8 ± 3.2</td>
<td>12.7 ± 6.4</td>
</tr>
<tr>
<td>(N=3)</td>
<td>(147-287)</td>
<td>(1.9-4.5)</td>
<td>(4.3-8.2)</td>
<td>(3.5-8.0)</td>
<td>(8.2-17.3)</td>
</tr>
<tr>
<td>Upper</td>
<td>184 ± 72</td>
<td>2.8 ± 0.8</td>
<td>5.6 ± 1.1</td>
<td>6.1 ± 3.0</td>
<td>12.3 ± 5.4</td>
</tr>
<tr>
<td>(N=46)</td>
<td>(43-386)</td>
<td>(1.6-4.5)</td>
<td>(3.8-8.1)</td>
<td>(2.0-14.7)</td>
<td>(3.6-28.4)</td>
</tr>
<tr>
<td>Lower</td>
<td>8.7 ± 115</td>
<td>3.2 ± 0.5</td>
<td>5.9 ± 0.6</td>
<td>0.6 ± 4.8</td>
<td>1.0 ± 8.9</td>
</tr>
<tr>
<td>(N=60)</td>
<td>(-266-188)</td>
<td>(2.0-3.8)</td>
<td>(4.3-6.9)</td>
<td>(-12.1-8.3)</td>
<td>(-21.9-15.1)</td>
</tr>
<tr>
<td>Gulf</td>
<td>-28.4 ± 91</td>
<td>1.2 ± 0.3</td>
<td>3.4 ± 0.3</td>
<td>-0.7 ± 1.5</td>
<td>-1.6 ± 3.9</td>
</tr>
<tr>
<td>(N=30)</td>
<td>(-178-107)</td>
<td>(0.8-1.7)</td>
<td>(2.8-4.1)</td>
<td>(-3.6-1.1)</td>
<td>(-8.4-3.6)</td>
</tr>
</tbody>
</table>
Table 3. Sectional and area-averaged air-sea CO$_2$ fluxes (mmol C m$^{-2}$ d$^{-1}$) in the St. Lawrence Estuary during all sampling months. To obtain the area-averaged CO$_2$ flux, the SLE was divided into five segments at equal intervals. The first row of the table shows the surface area (km$^2$) of each segment. The flux data in each segment were numerically averaged to obtain sectional fluxes, which were then area weighted and summed to obtain the spatially integrated whole-estuary flux (in bold). The F$_{W.14}$ and F$_{R&C.01}$ data provide the lower and upper estimates, respectively.

<table>
<thead>
<tr>
<th></th>
<th>Seg 1 (N=17)</th>
<th>Seg 2 (N=23)</th>
<th>Seg 3 (N=21)</th>
<th>Seg 4 (N=17)</th>
<th>Seg 5 (N=8)</th>
<th>Whole estuary</th>
</tr>
</thead>
<tbody>
<tr>
<td>Surface area (km$^2$)</td>
<td>1,098</td>
<td>2,250</td>
<td>2,726</td>
<td>3,404</td>
<td>3,303</td>
<td>12,781</td>
</tr>
<tr>
<td>F$_{W.14}$ (mmol m$^{-2}$ d$^{-1}$)</td>
<td>7.2</td>
<td>5.4</td>
<td>4.3</td>
<td>-1.7</td>
<td>-4.0</td>
<td><strong>1.00</strong></td>
</tr>
<tr>
<td>F$_{R&amp;C.01}$ (mmol m$^{-2}$ d$^{-1}$)</td>
<td>14.5</td>
<td>11.0</td>
<td>7.9</td>
<td>-3.3</td>
<td>-7.4</td>
<td><strong>2.06</strong></td>
</tr>
</tbody>
</table>
Figure 1. Map of the greater St. Lawrence system, including the chain of Great Lakes, the St. Lawrence River, the Upper St. Lawrence Estuary (USLE), the Lower St. Lawrence Estuary (LSLE), and the Gulf of St. Lawrence (GSL). From: Overview of the State of the St. Lawrence 2014 (with permission).
Figure 2. Map showing the four principal subdivisions of the study area and the sampling locations (markers). Water samples were collected during ten spring/summer research cruises: July 2003, June 2006, May 2007, July 2007, June 2009, July 2009, July 2010, May 2011, June 2013 and May 2016. The estuary, from the landward limit of the salt water intrusion near Île d’Orléans (5 km downstream of Québec City) to Pointe-des-Monts, where the coastline diverges, extends ~400 km and covers a total surface area of ~12,820 km$^2$. The solid gold line follows the 200m isobath of the Laurentian Channel.
Figure 3. Differences of pCO$_2$ ($\Delta$pCO$_2$) calculated from measured pH and TAlk with different published formulations of $K_1$ and $K_2$ including Cai and Wang (1998) [C&W], Lueker et al. (2000) [Luek], Roy et al. (1993) [Roy], Millero (2010) [Mill], and Millero (1979) for pure water only ($S_P = 0$) [Fresh]. All calculations were carried out at 15 °C (pCO$_2$ @ 15 °C). The constants of Lueker et al. (2000) are recommended for best practices by Dickson et al. (2007), whereas those of Roy et al. (1993) ($S_P = 5$ to 45) are recommended by Dickson and Goyet (1994). Both the constants of Cai and Wang (1998) and Millero (2010) have been proposed as more appropriate for the study of the carbonate system in estuarine waters.
Figure 4. Spatial distributions of surface-water pCO$_2$ (circles, squares) and practical salinity (x symbols) in the St. Lawrence River, Estuary and Gulf during spring/summer cruises. Horizontal lines show the mean pCO$_2$(air) in the sampling months. The pCO$_2$ data points above atmospheric level are sources of CO$_2$ to the atmosphere whereas those below atmospheric level are sinks of atmospheric CO$_2$. 
Figure 5. Spatial distribution of surface-water pCO$_2$ (µatm) in the St. Lawrence Estuary and Gulf during all spring/summer cruises. Linear interpolation was used between the sampling locations (+ symbols). The mean atmospheric pCO$_2$ during the sampling months ranged from 372 to 405 µatm.
Figure 6. The surface-water in situ pCO$_2$, temperature-normalized pCO$_2$, and $\Delta$pCO$_2$(thermal) averaged over salinity bins of 3. The open circles show the average temperature for each salinity bin. To correct for the increasing/decreasing effect of temperature on surface-water pCO$_2$, the in situ pCO$_2$ were normalized to the average surface-water temperature of the study area (T = 7.82 °C). The $\Delta$pCO$_2$(thermal) are due to temperature deviations from T = 7.82 °C, whereas variations in temperature-normalized pCO$_2$ are due to water mass mixing and/or biological activity.
Figure 7. Box plot showing the variability of air-sea CO$_2$ fluxes in the four principal subdivisions of the study area (St. Lawrence River, Upper Estuary, Lower Estuary and Gulf). The box spans the interquartile range (25–75 percentiles), the dashed line is the median, and the whiskers extend to the extreme data points. The F$_{W-14}$ data were combined with the F$_{R&C-01}$ data from all spring/summer sampling months to depict the upper and lower limits of gas exchange.
Figure 8. Comparison of the saturation states (or % saturation) of pCO$_2$ and DO in the surface mixed layer waters of the St. Lawrence River, Estuary and Gulf. Dashed lines delineate the 100 % saturation levels for both gases. Surface-water samples (markers) fall into one of four quadrants representing the dominant controls on CO$_2$/O$_2$ dynamics. Quadrants I and III indicate the effects of photosynthesis/respiration, whereas Quadrants II and IV indicate the effects of heating/cooling.
Figure 9. Spatial distributions of surface-water pCO$_2$ (circles) and temperature (diamonds) in the St. Lawrence River, Estuary and Gulf during spring/summer cruises. Temperatures ranged from 4.2 to 17.4 °C (generally decreasing) in the USLE, -0.32 to 12.6 °C (generally increasing) in the LSLE, and 3.9 to 13.7 °C (generally increasing) in the GSL. The horizontal line shows the mean atmospheric pCO$_2$, pCO$_2$(air), during all sampling months.