Variability of carbon monoxide and carbon dioxide apparent quantum yield spectra in three coastal estuaries of the South Atlantic Bight

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Abstract. The photochemical oxidation of oceanic dissolved organic carbon (DOC) to carbon monoxide (CO) and carbon dioxide (CO2) has been estimated to be a significant process with global photoproduction transforming petagrams of DOC to inorganic carbon annually. To further quantify the importance of these two photoproducts in coastal DOC cycling, 38 paired apparent quantum yield (AQY) spectra for CO and CO2 were determined at three locations along the coast of Georgia, USA over the course of one year. The AQY spectra for CO2 were considerably more varied than CO. CO AQY spectra exhibited a seasonal shift in spectrally integrated (260 nm–490 nm) AQY from higher efficiencies in the autumn to less efficient photoproduction in the summer. While full-spectrum photoproduction rates for both products showed positive correlation with pre-irradiation UV-B sample absorption (i.e. chromophoric dissolved organic matter, CDOM) as expected, we found no correlation between AQY and CDOM for either product at any site. Molecular size, approximated with pre-irradiation spectral slope coefficients, and aromatic content, approximated by the specific ultraviolet absorption of the pre-irradiated samples, were also not correlated with AQY in either data set. The ratios of CO2 to CO photoproduction determined using both an AQY model and direct production comparisons were 23.2 ± 12.5 and 22.5 ± 9.0, respectively. Combined, both products represent a loss of 2.9 to 3.2 % of the DOC delivered to the estuaries and inner shelf of the South Atlantic Bight yearly, and 6.4 to 7.3 % of the total annual degassing of CO2 to the atmosphere. This result suggests that direct photochemical production of CO and CO2 is a small, yet significant contributor to both DOC cycling and CO2 gas exchange in this coastal system.

1 Introduction

The ocean represents one of the largest and most dynamic reservoirs for reduced carbon on Earth with a pool of dissolved organic carbon (DOC) on the order of 600 Pg/C (Hansell et al., 2009; Canadell et al., 2007). Coastal systems are a dynamic subset of this DOC pool with carbon inputs from terrestrial sources adding to in-situ production. Rivers contribute up to 0.2 Pg/C DOC to coastal systems per year with most having had little exposure to sunlight before its arrival in coastal waters (Ludwig et al., 1996). Carbon of terrestrial origin is generally refractory to microbial oxidation and strongly absorbs solar radiation in blue and ultraviolet (UV) wavelengths (Bricaud et al., 1981; Carlson, 2002). Regardless of source, the optically active fraction of DOC is referred to as chromophoric dissolved organic matter (CDOM) and absorbs much of the energetic UV radiation entering the water column. This both shields the marine biological community from damage and captures the energy required to initiate most photochemical reactions in the surface ocean.

Despite the continuous input of CDOM to the coastal ocean via continental sources, ocean water as a whole is not highly coloured, due largely to photochemical transformations that are known to alter the absorptive characteristics of coloured components within the DOC pool (Andrews et al., 2000; Del Vecchio and Blough, 2002; Goldstone et al.,
The absorption of solar radiation by CDOM can lead to a whole host of chemical reactions, including formation of reactive oxygen species such as hydroxyl radical (OH-) and superoxide (O$_2^-$) (Micienski et al., 1993; Moffett and Zafiriou, 1993; Zika et al., 1985; Blough and Zepp, 1995), breakdown of large organic molecules into lower molecular weight carbon compounds (Wetzel et al., 1995; Kieber et al., 1989, 1990), alteration of the redox state of biologically important metals (White et al., 2003; Barbeau, 2006; Barbeau et al., 2003), and formation of oxidized inorganic carbon species, such as CO and CO$_2$ (measured as DIC) (Clark et al., 2004; Johannessen and Miller, 2001; Miller and Zepp, 1995; White et al., 2010; Ziołkowski and Miller, 2007; Wang et al., 2009; Xie et al., 2004; Zafiriou et al., 2003). On an annual global basis, photoproduction of these two carbon compounds may be almost equal to oceanic new production ($\sim$10$^{15}$ moles C yr$^{-1}$; Johannessen, 2000; Wang et al., 2009).

Photochemical production is the major source of CO in the surface ocean and, as a relatively insoluble gas, it can be transferred to the lower atmosphere where it competes with methane as a major sink for hydroxyl radical, thus indirectly contributing to changes in greenhouse gas concentrations (Shindell et al., 2009). CO in seawater can also be used as a substrate for marine bacteria (Tolli et al., 2006; Kitić et al., 2011), efficiently competing with gas exchange to lessen its transfer to the atmosphere. Even with these two removal pathways, CO is supersaturated with respect to the atmosphere in the surface ocean, having concentrations from $\sim$2 nM in the open ocean to $\sim$12 nM in the coastal ocean (Zafiriou et al., 2008, 2003). As is the case with other photoproducts, oceanic CO concentrations show a distinct diurnal signal with a late afternoon peak. This may reflect the photoinhibition of CO uptake by bacteria, with higher bacterial consumption rates when solar UV radiation is less intense and substrate concentrations have been elevated by photochemistry (Tolli and Taylor, 2005). Photochemical production of CO in the oceans has been well studied recently (Day and Faloona, 2009; Miller and Moran, 1997; Stubbins et al., 2008, 2006; White et al., 2010; Xie et al., 2009; Zafiriou et al., 2003, 2008; Ziołkowski and Miller, 2007; Kitić et al., 2011), and global annual production of CO in the worlds oceans is currently estimated at between 30 and 84 Tg C yr$^{-1}$ (Zafiriou et al., 2003; Stubbins et al., 2006; Fichot and Miller, 2010).

Increased CO$_2$ concentrations in the surface ocean from rising atmospheric CO$_2$ levels are creating a more acidic ocean, raising concerns for the health of calcareous organisms critical to carbonate balance in the oceans (Moy et al., 2009; Fabry et al., 2008). Overall, the ocean is a sink for atmospheric CO$_2$; however, some coastal systems, such as the terrestrial influenced South Atlantic Bight (USA) and the Pearl River estuary (China), have been reported as seasonal sources of CO$_2$ to the atmosphere (Guo et al., 2009; Jiang et al., 2008b; Wang et al., 2005), with the inner shelf and estuaries being the strongest sources (Cai, 2011; Jiang et al., 2008a) presumably due to heterotrophic production in these regions. Photochemical production of dissolved inorganic carbon (DIC) in the form of CO$_2$ from the oxidation of DOC has the potential to add to the source strength in these areas with high organic carbon content.

The photochemical production of CO$_2$ (generally measured as total DIC but discussed as CO$_2$) has been studied in both freshwater (Anesio and Granelli, 2004, 2003; Bertilsson and Tranvik, 2000; Granelli et al., 1998; Salonen and Vahatalo, 1994) and marine systems (Belanger et al., 2006; Gao and Zepp, 1998; Johannessen and Miller, 2001; Miller and Zepp, 1995), though the extent of coverage in marine systems is considerably less. Several noteworthy studies have pointed towards a strong global photochemical signal of DIC production in the ocean (White et al., 2010; Johannessen and Miller, 2001; Miller and Zepp, 1995).

The analytical constraints on measuring the photochemical production of CO$_2$ in seawater containing an inorganic carbon pool over 100× more concentrated than changes created by photochemical production are significant. Consequently, there has been some focus on establishing a valid ratio relating CO$_2$ and CO photoproduction that would allow use of the more prevalent CO data to assess the magnitude of photochemical CO$_2$ production in the oceans. Early estimates of the CO$_2$:CO photochemical production ratio were 15–20 (Miller and Zepp, 1995; Mopper and Kiefer, 2000), but more recent studies have shown that this ratio is much more variable, ranging from ~2 to 98 in some cases (White et al., 2010). This ratio may be dependent on the source material comprising the CDOM. Photochemical efficiencies for both CO and CO$_2$ (i.e. apparent quantum yield spectra, AQY) have been reported to vary depending on carbon source (Ziołkowski and Miller, 2007; Johannessen and Miller, 2001) with CO AQY spectra appearing to be more constant than those for CO$_2$.

Together, CO and CO$_2$ constitute the largest analytically identifiable carbon photoproducst in the ocean and, as such, have potential to affect the cycling of DOC in the oceans. The following study provides new data that will better constrain the natural variability of CO and CO$_2$ photoproduction and its significance to DOC cycling in the coastal ocean. Thirty-eight (38) samples were collected from three sites along the coast of Georgia, USA, over the course of a year for paired determinations of CO and CO$_2$ AQY spectra together with measurements of potential environmental parameters that may provide insight into these photochemical processes. The results presented here provide the most extensive set of paired CO and CO$_2$ AQY and photoproduction data collected to date, allowing a robust consideration of their relative production rates and photochemical efficiency.
2 Methods

2.1 Sample collection

Samples were collected for both CO and CO$_2$ photochemical experiments, monthly during spring high tide and quarterly during spring low tide. Three sites were chosen within the Georgia Coastal Ecosystems Long Term Ecological Research (GCE LTER) area, meant to represent three variants of coastal estuarine systems (Fig. 1). Sapelo Sound (31.537779° N, 81.176860° W) is a coastal marine dominated site, with little freshwater input over the year. Altamaha Sound (31.314000° N, 81.265333° W) receives outflow from the Altamaha River, which drains the largest watershed in the state of Georgia, and is a mixed riverine and marsh site. Doby Sound (31.376373° N, 81.281718° W) is primarily a coastal marine dominated site but receives significant freshwater input from the Altamaha River during periods of high flow. Samples were collected in concert with the GCE LTER monthly mini-cruise program. Sample characteristics can be found in Table 1. Samples were collected in acid-washed polycarbonate bottles (Nalgene, 2l) and stored on ice until returned to the laboratory (<3 h). Samples were then immediately filtered using an acid-cleaned 0.2 µm nylon cartridge filter (Whatman Polycap AS75) and were then stored in the dark at 4°C until the photochemical evaluation (0–6 months). Previous evaluations in our lab (unpublished) and work such as Stedmon and Markager (2001) and Swan et al. (2012) have shown that storage had little effect on CDOM absorption, and thus storage was determined not to be an issue.

2.2 Sample and labware preparation

To analyze the photoproduction of DIC (i.e. CO$_2$) in micromolar amounts in seawater, the ambient DIC in all samples (i.e. CO$_2$, HCO$_3^-$, CO$_3^{2-}$) was removed from the sample prior to irradiation following the methods of Johannessen and Miller (2001). Briefly, 11 of sample was transferred to a UV-C sterilized glass kettle, acidified to pH 2–3 using a nominal amount of concentrated HCl to minimize dilution (Fisher Scientific), and bubbled overnight (~8 h) under positive pressure with CO$_2$-free (soda lime column, 12″×1.5″, Fisher Scientific, indicating grade) room air to ensure complete removal of DIC from the solution. Successful treatment was confirmed via direct analysis using a Shimadzu TOC CPN in IC mode (see Sect. 2.3).

Quartz spectrophotometric cells (Spectrocell Inc, 10 cm pathlength) for use in DIC photochemical experiments were sterilized with a UV-C lamp at close range (less than 30 cm) for an hour before filling to ensure that CO$_2$ produced during the course of an irradiation was due to photochemical production rather than biological respiration. Once DIC was removed from a sample, it was rebuffered to its initial pH (~7–8 pH) using crystalline sodium borate (Fisher Scientific, ACS grade). Immediately after raising the pH, the spectrophotometric cells were filled using a gas-tight glass dispenser while keeping the sample under positive pressure with CO$_2$-free air. Cells were then sealed headspace-free, using 13 mm diameter, 1-mm-thick Teflon-faced butyl rubber septa, placed in a chilled aluminum block (held at 15°C), and mounted within a Suntest solar simulator (Suntest CPS) for radiation exposure.

Samples for measurement of CO photochemical production did not require the removal of DIC and were not subject to contamination from microbial respiration. Consequently, these were simply filtered a second time (0.2 µm Whatman Polycap AS75) just prior to placing them directly into acid-washed and oven-dried spectrophotometric cells, sealed without a headspace, and exposed in the solar simulator in the same manner as was done in the CO$_2$ experiments.

2.3 Photochemical irradiations

Photochemical irradiations were performed following the multispectral methods of Johannessen and Miller (2001). A tightly fitted 2.4-cm-thick black plastic lid with 1.6 cm diameter holes drilled directly above each spectrophotometric cell served as effective Gershun tubes for each cell to minimize off-axis light, thus minimizing reflection at the air-quartz surface of the cells. To create distinct irradiation
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Table 1. Sample characteristics and AQY fitting parameters for all samples.

<table>
<thead>
<tr>
<th>Date</th>
<th>Site</th>
<th>Tide</th>
<th>Salinity</th>
<th>DOC (µmol l⁻¹)</th>
<th>ag320 (m⁻¹)</th>
<th>CO</th>
<th>CO₂</th>
</tr>
</thead>
<tbody>
<tr>
<td>11/11/08</td>
<td>ALT</td>
<td>HT</td>
<td>32.2</td>
<td>226</td>
<td>4.101</td>
<td>7.996</td>
<td>0.039</td>
</tr>
<tr>
<td>11/11/08</td>
<td>ALT</td>
<td>LT</td>
<td>22.1</td>
<td>432</td>
<td>13.775</td>
<td>8.310</td>
<td>0.026</td>
</tr>
<tr>
<td>11/12/08</td>
<td>SAP</td>
<td>HT</td>
<td>33.2</td>
<td>287</td>
<td>3.263</td>
<td>7.700</td>
<td>0.060</td>
</tr>
<tr>
<td>11/12/08</td>
<td>SAP</td>
<td>LT</td>
<td>32.6</td>
<td>275</td>
<td>6.155</td>
<td>8.012</td>
<td>0.037</td>
</tr>
<tr>
<td>11/13/08</td>
<td>DOB</td>
<td>HT</td>
<td>33</td>
<td>213</td>
<td>4.113</td>
<td>7.933</td>
<td>0.038</td>
</tr>
<tr>
<td>11/13/08</td>
<td>DOB</td>
<td>LT</td>
<td>30.8</td>
<td>335</td>
<td>7.438</td>
<td>8.156</td>
<td>0.032</td>
</tr>
<tr>
<td>12/10/08</td>
<td>DOB</td>
<td>HT</td>
<td>30.2</td>
<td>226</td>
<td>5.453</td>
<td>8.147</td>
<td>0.034</td>
</tr>
<tr>
<td>12/10/08</td>
<td>DOB</td>
<td>LT</td>
<td>30.8</td>
<td>335</td>
<td>9.711</td>
<td>8.361</td>
<td>0.028</td>
</tr>
</tbody>
</table>

conditions for the samples in each of 14 spectrophotometric cells, duplicate Schott-glass long-pass cutoff filters (WG280, WG295, WG305, WG320, GG385, GG420, GG475) were placed directly over the lid, between the quartz cells and the light source. An opaque disk was used above the 15th cell in the chilled Al block to provide a dark control. The spectral irradiance under the lid and under each filter was measured at 1 nm resolution using a 2-inch integrating sphere attached by a 60 cm fiber optical cable to an Optronic model 756 spectroradiometer that was calibrated using a NIST-certified tungsten halogen standard lamp powered by an Optronic model OL752-10 power supply. Exposures lasted either 4 or 6 h for CO₂ experiments and 3 h for CO experiments. This delivered a total photon dose under the 305 nm (WG305) cutoff filter that most closely simulates the solar spectrum in our system, of 0.93, 1.23, and 1.85 millimoles photons cm⁻² for 3, 4, and 6 h respectively. The UV-visible absorbance of each sample was measured directly in the spectrophotometric exposure cells on a Perkin-Elmer Lambda-40 spectrophotometer before and after each irradiation. The absorbance spectrum was converted to absorption for use in apparent quantum yield calculations using the equation

\[
\alpha(\lambda) = \frac{A(\lambda) \cdot 2.303}{\ell}
\]

(1)
where \( a \) is the Napierian absorption coefficient (m\(^{-1}\)), \( A \) is the absorbance measured by the spectrophotometer and is unitless, \( \ell \) is the pathlength of the spectrophotometric cell (m), and 2.303 converts from log 10 scale to natural log scale (i.e. Napierian). In all experiments, photochemical fading was less than 1% at all wavelengths. This small change in \( a(\lambda) \) was accounted for in AQY calculations by assuming linear fading and using the average at each wavelength between the starting and ending absorption spectra.

Photochemically produced CO\(_2\) was measured with a Shimadzu TOC V-CPN analyzer configured for IC analysis. It should be noted here that, like almost all studies of CO\(_2\) photochemical production in natural waters, this method quantifies CO\(_2\) as dissolved inorganic carbon (DIC) and that discussion of CO\(_2\) production in any measured sample assumes that changes in DIC are from CO\(_2\) to dissolved inorganic carbon (DIC) and that dis-

The AQY is expressed as a unitless ratio and all quantities are spectral. Thus, in order to describe the efficiency of these photochemical reactions, both the amount of product formed and the total amount of photons absorbed by the CDOM in the sample over the course of the experiment must be known. Products are measured by the analytical techniques described above, and the number of photons absorbed over the course of an experiment for all samples is calculated using the following equation from Hu et al. (2002):

\[
Q_a(\lambda) = E_0(\lambda) \times (a_g(\lambda)/a(\lambda)) \times S \times [1 - \exp(-a(\lambda) \times \ell)],
\]

where \( Q_a \) is the moles of photons absorbed/second by the sample, \( E_0(\lambda) \) is the scalar irradiance entering the top of the cell (mols photons m\(^{-2}\) s\(^{-1}\)), \( S \) is the irradiated surface area of the spectrophotometric cell (m\(^2\)), \( a_g \) and \( a \) are the absorption coefficients for CDOM and for the total solution, respectively (m\(^{-1}\)), and \( \ell \) is the pathlength of the spectrophotometric cell (m). Making the assumption that, in filtered samples, essentially all of the absorption in the cell occurs due to CDOM, (i.e. \( a_g \approx a \)) this equation can be simplified to

\[
Q_a(\lambda) = E_0(\lambda) \times S \times [1 - \exp(-a(\lambda) \times \ell)].
\]

This rigorous treatment of \( Q_a \) is necessary to avoid overestimating the absorption of photons by samples that experience varying light fields over the length of the spectrophotometric cell due to inner filter effects (i.e. self shading) (Hu et al., 2002).

Due to the multispectral nature of these irradations, an iterative, non-linear fitting routine is used to determine the AQY spectrum (Johannessen and Miller, 2001; Ziolkowski and Miller, 2007; White et al., 2010; Xie et al., 2009). The experimental photochemical production under each cutoff filter is calculated according to the equation

\[
\frac{dP}{dt}(\lambda) = AQY(\lambda) \cdot Q_a(\lambda)
\]

where \( \frac{dP}{dt} \) is the production rate of the product in question (CO or CO\(_2\)) over the course of the experiment. The spectral equation for the AQY is defined a priori as

\[
AQY(\lambda) = e^{-(m_1 + m_2(\lambda - m_3))}
\]

and initial estimates for the fitting coefficients, \( m_1, m_2, \) and \( m_3 \) are taken from literature values (Johannessen and Miller, 2001; Ziolkowski and Miller, 2007). The MATLAB \textit{nlinfit} routine is used to iteratively find the best fit to the defined AQY equation, adjusting the \( m \) coefficients and calculating the single AQY spectrum that best describes all of the different photochemical production values measured in samples

\[\text{www.biogeosciences.net/9/4279/2012/}
\]

\[\text{Biogeosciences, 9, 4279–4294, 2012}\]
distributed under the varied spectral irradiation environments created by Schott cutoff filters. Previous published work on CO and CO₂ AQY spectra has shown that this multispectral approach compares well with traditional monochromatic studies, particularly in the important UV region of the spectrum, and both do a credible job of predicting measured full spectrum photochemical production rates (White et al., 2010; Ziolkowski and Miller, 2007).

2.5 Determination of CDOM-normalized production rates

As mentioned above, the irradiance spectrum under the 305 nm cutoff filter most closely matches the spectral distribution of sunlight. Consequently, photoproduction in cells under the 305 nm cutoff filter can be used as a proxy for photoproduction in sunlight when total photon dose is matched. For some comparisons, we have used photoproduction under the 305 nm filter (referred to as “measured production”) to provide useful insight about the variation of photoproduction in the system. As can be seen clearly in Eqs. (4) and (5) above, it should be noted that measured production is intrinsically linked to both AQY and the absorption coefficient of CDOM in the system with samples having higher absorption generally exhibiting higher measured production without necessarily reflecting differences in photochemical efficiency.

To assess variations in photoproduction due to changes only in the AQY spectrum independently from absorption in the original sample, CDOM normalized production can be calculated, using the photoproduction Eq. (5) where the change in concentration of product per unit time (e.g. mol s⁻¹) is a function of the scalar irradiance, total number of photons absorbed (Eq. 4), and the AQY(λ), with each of these quantities being spectrally defined. By employing both a standard absorption spectrum (for this paper we use a₂g from the November 2008 Altamaha high tide sample) and a constant, defined scalar irradiance spectrum, we can compare the consequences of variations in the AQY spectrum on photoproduction by integrating Eq. (5) over photochemically active wavelengths (260 nm–490 nm). This allows an evaluation of both the spectral slope coefficient of each AQY spectrum together with its relative magnitude. For consistency, the spectral irradiance used was the same as in the experimental setup, leading to units of production in the spectrophotometric cell of nmol product s⁻¹ cell⁻¹ (~30 ml). Errors are calculated from the root-mean-square error of the AQY fit, as determined by the nlinfit routine, and propagated through the calculation of CDOM normalized production. It should be noted that these rates are not true environmental rates, but were solely calculated for the purpose of eliminating the variation due to CDOM and irradiance in the photochemical reactions that would naturally occur in an environmentally rigorous calculation of photoproduction. The purpose of this calculation was to specifically isolate only the effect of changing AQY spectra on photoproduction, and we do not want to mislead the reader by suggesting that these rates are estimates of those found in the coastal waters sampled. For a discussion of the environmentally relevant rates, please refer to Sect. 2.7.

2.6 Determination of sample optical proxies

Helms et al. (2008) have shown that the spectral slope coefficient for the exponential best fit of CDOM spectra over the wavelength range 275 nm–295 nm (S₂75–295) correlates with the molecular mass of the DOM in a seawater sample. For this CDOM spectral range, a shallow spectral “slope” in the UV indicated larger molecular size while a steep S₂75–295 indicated smaller molecular size. Spectral slope coefficients were determined for our samples using a non-linear fit to the CDOM absorption spectrum over the prescribed wavelengths, 275–295 nm. Using MATLAB’s nlinfit routine, the CDOM spectra were fit to the equation

\[ a_g(\lambda) = B \cdot e^{-S \lambda} \]  

(7)

where \( a_g \) is absorption in m⁻¹, \( B \) is a fitting constant, \( \lambda \) is wavelength in nm, and \( S \) is the spectral slope coefficient.

Specific ultraviolet absorption (SUA) is the absorption coefficient of CDOM at a specific UV wavelength divided by the total DOC concentration of the same sample. When highly absorbing moieties such as those containing aromatic structures make up a larger fraction of the total DOC pool, SUVA is expected to be elevated over those samples without aromatics. Weishaar et al. (2003) have confirmed this strong positive correlation between DOC % aromatic content and SUVA at 254 nm (i.e. SUVA₂⁵₄) in a variety of naturally occurring water samples.

The GCE-LTER nutrient monitoring program collected samples for DOC analysis simultaneously with our photochemical samples (data obtained via http://www.gce-lter.marisci.uga.edu and displayed in Table 1). The absorption spectra of samples prior to irradiation were used to calculate SUVA₂⁵₄ using the equation

\[ SUVA_{254} = \frac{a_g(254)}{DOC} \]  

(8)

where \( a_g(254) \) is the absorption coefficient at 254 nm and DOC is the concentration of dissolved organic carbon in mg C 1⁻¹ reported by the GCE-LTER.

2.7 Annual photochemical production

To use our CO and CO₂ AQY data set for an estimate of the total annual photoproduction of these two oxidation products in the South Atlantic Bight (USA), we employed a photoproduction equation adapted from Fichot and Miller (2010) and Stubbins et al. (2006):

\[ \sum_{\text{area}} P(\text{total}) = \int_{490}^{290} E_d(\lambda) \cdot R(\lambda) \cdot AQY(\lambda) \cdot 12, \]  

(9)
where $P(\text{total})$ is the annual photoproduction rate (g C yr$^{-1}$ m$^{-2}$) summed over the area of interest, $E_{d0−}(\lambda)$ is the downwelling spectral irradiance for 30° N latitude (mol photons yr$^{-1}$ m$^{-2}$) calculated using the STAR model (System for Transfer of Atmospheric Radiation, University of Munich, Ruggaber et al., 1994) following the methodology described in Fichot and Miller (2010), AQY is the “mean” apparent quantum yield spectrum for the chemical species in question (unitless ratio), 12 is the molecular mass of carbon (g mol$^{-1}$), $\lambda$ is wavelength (nm) and $R(\lambda)$ is the ratio of CDOM absorption to total light attenuation (unitless ratio), as derived from a large coastal and oceanic data set (Fichot, 2004). The ratio follows a linear relationship spectrally:

$$R(\lambda) = \frac{a_g(\lambda)}{K_D(\lambda)} = -0.0028(\lambda) + 1.575,$$

(10)

where $a_g$ is absorption (m$^{-1}$) and $K_D$ is the downwelling attenuation coefficient (m$^{-1}$). The “mean” apparent quantum yield spectra were determined by fitting all data from all experiments involving a particular product to the AQY equation (Eq. 6).

### 3 Results

#### 3.1 Apparent quantum yield spectra and CDOM-normalized photoproduction rates

Examining our entire AQY data set together shows that results for both CO and CO$_2$ fall within the range of literature values from previous studies, as shown in Fig. 2. Also, as expected from previous studies, the overall efficiency of CO production is lower than that seen for CO$_2$ production over the UV wavelength range (280–400 nm). Additionally, the AQY spectra describing CO efficiency have a tighter distribution than those observed for CO$_2$. Table 1 shows the individual AQY fitting parameters for every sample shown in Fig. 2 together with its associated field data.

CDOM-normalized CO and CO$_2$ photoproduction rates calculated from our AQY data set for each sample can be seen in Fig. 3. Neither product shows obvious patterns between sites, though there appears to be somewhat more variation among the terrestrially influenced sites (i.e. Doboy Sound and Altamaha Sound) for CO$_2$. CO rates show a seasonal signal at all three sites, with lower production efficiency in the spring and summer months and higher production efficiency in the autumn and winter months for both high and low tide conditions. Considering all sites and tides, the difference between summer and autumn (August and November) is 21.7% for CO, and is statistically significant (1-way ANOVA, $p < 0.01$).

#### 3.2 CO$_2$ to CO production ratios

For comparison with previous studies and for potential use in regional coastal photochemical production estimates, the

ratio between photochemical production of CO$_2$ and CO was examined in two ways. First, the ratio was calculated using the measured production values (CO$_2$:CO$_{\text{measured}}$) for each sample. Second, the ratio was calculated from the CDOM-normalized production values for each sample (CO$_2$:CO$_{\text{normalized}}$) generated from specific AQY spectra and Eq. (7). Values for both ratio methods are presented in Table 2, both showing a similar large ratio range and having nearly identical mean values (22.5 and 23.2).

### 3.3 Photochemical production relative to salinity and optical properties

Figure 4 shows the relationships between photochemical production and salinity. Measured production appears to decrease with increasing salinity for both photochemical products seen in Fig. 4a, but this is likely due to a negative correlation between CDOM and salinity. When normalized for CDOM absorbance (Fig. 4b), there is no relationship between salinity and the efficiency of photochemical production of either product.

Because the absorption of a photon by CDOM must occur for a photochemical reaction to occur, a relationship between CDOM “concentration” and measured production should exist (see Eqs. 4 and 5). Using the absorption coefficient of the sample at 320 nm ($a_g$320, m$^{-1}$), Fig. 5a shows that samples with higher $a_g$320 absorb more photons and exhibit a correspondingly higher production of both photochemical products. The relationship between $a_g$320 and CDOM normalized production, shown in Fig. 5b, shows no such trend for either product with $a_g$320, indicating that efficiency is relatively constant for all CDOM “concentrations” in this study.

The measured production of both products shows a slight decreasing trend with relation to CDOM spectral slope, but the scatter around this trend is quite large (Fig. 6a). Once production is normalized for CDOM variations (Fig. 6b), there is less scatter than seen with measured production alone, but no trend exists with respect to spectral slope.

Examination of the measured production of both products compared to carbon normalized optical properties using SUVA$_{254}$ (Fig. 7a) also showed a large amount of scatter. The slight positive correlation seen in this relationship is not statistically significant. Again, using CDOM normalized production compared to SUVA$_{254}$ shows less scatter than measured production, but no trend for either CO or CO$_2$ was observed, suggesting that, within the range of CDOM and SUVA$_{254}$ values seen in this study, aromatic content does not seem to affect the efficiency for producing CO and CO$_2$ through photochemical processes (Fig. 7b).
3.4 Estimated photochemical production

Table 3 shows annual photochemical production estimates for the estuaries and inner shelf of the South Atlantic Bight (SABi) using the methods described in Sect. 2.7 and the areal extent values ($37.7 \times 10^9$ m$^2$) that Cai (2011) used to calculate inorganic carbon cycling in this coastal region. Photochemical production of CO$_2$ was calculated in two ways. First we used the more easily defined “mean” AQY spectrum for CO photoproduction from our data to calculate the annual photoproduction of CO in the SABi as $7.13 \times 10^9$ g C yr$^{-1}$. Then using the mean ratio of CO$_2$ to CO production from our data (22.5), we calculated the annual SABi photoproduction of CO$_2$ to be $1.60 \times 10^{11}$ g C yr$^{-1}$. The second method used to estimate photoproduction of CO$_2$ in the SABi employed the “mean” AQY spectrum obtained directly from our CO$_2$ data set, which gave an annual SABi photoproduction of $1.38 \times 10^{11}$ g C yr$^{-1}$.

4 Discussion

4.1 Seasonal changes in photoproduction efficiency

The two sites that experience significant freshwater flow during the year (Doboy and Altamaha sounds) exhibit considerable seasonal variability in their photochemical efficiencies as can be seen by examining their CDOM-normalized production (Fig. 3 top panel) for CO$_2$. The Sapelo Sound samples show considerably less variability. This is likely due to
the homogeneity of its carbon source material (CDOM) since it is a tidally dominated system with a DOC source dominated by tidal exchange with coastal water and input from surrounding marshes. This is unlike the mixed influence of marsh, riverine, and coastal DOC experienced by Doboy and Altamaha sounds where the Altamaha River delivers upland (i.e. nonmarsh-derived) DOC with a seasonal signal, thus creating a more complex and dynamic DOC system.

Fluctuations in riverine discharge and coastal hydrology may also add to the observed variability of photochemical reactivity in our samples. Due to higher sediment loads and reduced residence times during high-flow events in the Altamaha River, CDOM reaching our sampling sites during these periods would have minimal pre-exposure to sunlight and could influence its photochemical reactivity relative to CDOM delivered during low-flow conditions with a longer pre-exposure to sunlight. It is also documented that estuarine iron is usually positively correlated with river flow and particulate loading (e.g. Lippiatt et al., 2010) and that its complexation with DOM can greatly affect photochemical reactions in estuarine systems (Gao and Zepp, 1998; White et al., 2003). These potential variations in the chemical nature of CDOM would certainly add to the variation of AQY in coastal systems and may not be directly related to salinity.

CDOM normalized photoproduction of CO shows less variability than does CO₂ across all samples, suggesting that the molecular source that dominates CO production is more homogenous throughout these coastal and estuarine samples.
than that responsible for CO$_2$ photoproduction. Another possible source of this larger CO$_2$ variation is that the chromophores within CDOM that initiate CO photoproduction vary more directly with overall CDOM absorbance than do the chromophores responsible for CO$_2$ photoproduction. In such a case, using CDOM to normalize photoproduction would create a larger variation in normalized CO$_2$ production than in CO. This would be compatible with reported positive correlations between CO photoproduction and aromatic content (Stubbins et al., 2008), a natural chromophore thought to be a very strong contributor to the overall CDOM absorbance.

All three sampling sites showed a decrease in photochemical production efficiency for CO in the spring and summer with an increase in the autumn and winter months. While the total variation is small (21.7% between November and August over all sites and tides), it is a statistically significant change. In Georgia, the marshes are dominated by the smooth cordgrass *Spartina alterniflora* that is highly productive in the spring and summer months and undergoes senescence beginning in September and October. Autumn, therefore, represents a period of increased organic matter input to the ocean from the marshes, a situation similar to Gardner et al. (2005) in the Neponset River estuary. This increased flux of organic carbon into the system may help to explain the higher efficiency of CO photoproduction in the autumn and winter months. Interestingly, this trend is seen at all three sites for CO, suggesting that marsh-derived DOC may be the dominant control on the seasonal variability of AQY for CO photoproduction. Additionally, this seasonal trend with lower efficiencies seen in the summer months could be driven by increased solar irradiation and pre-exposure of the CDOM to extensive sunlight, leading to photobleaching and potentially lowering the efficiency of the photochemical reaction. CDOM released in the autumn and winter months would likely receive more solar irradiation prior to sample collection, giving mean ratios of 22.5 (measured production) and 23.2 (CDOM normalized) (see Table 2). While the overall range of CO$_2$:CO ratios was large (4–73 (measured production), 12–62 (CDOM normalized)), it agrees well with the ratios reported in Miller and Zepp (1995) (12–65) with a mean of 20. Miller and Zepp (1995) found that pre-faded samples showed smaller CO$_2$ to CO production ratios. Pre-exposure of the CDOM to sunlight could explain the variation of CO$_2$ to CO production ratios in the present study since samples from riverine sources, presumably having had less sunlight-exposure, showed higher ratios (see Fig. 3) than samples from the more coastal Sapelo Sound site, which likely received more solar irradiation prior to sample collection. Regardless of the cause, it is clear from our spatialtemporal study that CO$_2$ photoproduction is more variable than CO photoproduction (Figs. 2, 3). This is consistent with both CO$_2$ AQY spectra being more affected by pre-exposure to sunlight than CO AQY spectra, and differing molecular

### Table 3. Annual photoproduction calculated for the estuaries and inner shelf of the South Atlantic Bight (SABI) for CO and CO$_2$.

<table>
<thead>
<tr>
<th>Photoproduct</th>
<th>Annual SAB production (g C yr$^{-1}$)</th>
<th>Method used</th>
</tr>
</thead>
<tbody>
<tr>
<td>CO</td>
<td>$7.13 \times 10^9$</td>
<td>“Mean” AQY</td>
</tr>
<tr>
<td>CO$_2$</td>
<td>$1.36 \times 10^{11}$</td>
<td>“Mean” AQY</td>
</tr>
<tr>
<td>CO$_2$</td>
<td>$1.60 \times 10^{11}$</td>
<td>CO$_2$:CO measured production ratio</td>
</tr>
</tbody>
</table>

While some estimates of regional and global photochemical production of CO$_2$ have been made (Mopper and Kieber, 2000; Miller and Zepp, 1995), these are generally based on ratios between CO$_2$ and CO photoproduction, taking advantage of comparatively well-defined CO photoproduction estimates. The first study to define a ratio between CO$_2$ and CO photoproduction reported values between 15 and 20 (CO$_2$:CO) (Miller and Zepp, 1995), suggesting that CO$_2$ is a dominant carbon photoproduc resulting from the irradiation of CDOM. To date, this has been the only study where CO and CO$_2$ were measured from the same experimental exposure cell at the same time. Consequently, the question remains as to what effect the absence of the carbonate system, required for the CO$_2$ analysis, and small differences in pH may have on measured photoproduction of CO. More recent studies have shown that the ratio between CO$_2$ and CO photoproduction is more variable and can range from 2 to 98 depending on the water type (White et al., 2010). In this study, the ratio of CO$_2$ to CO was calculated in two different ways, giving mean ratios of 22.5 (measured production) and 23.2 (CDOM normalized) (see Table 2). While the overall range of CO$_2$:CO ratios was large (4–73 (measured production), 12–62 (CDOM normalized)), it agrees well with the ratios reported in Miller and Zepp (1995) (12–65) with a mean of 20. Miller and Zepp (1995) found that pre-faded samples showed smaller CO$_2$ to CO production ratios. Pre-exposure of the CDOM to sunlight could explain the variation of CO$_2$ to CO production ratios in the present study since samples from riverine sources, presumably having had less sunlight-exposure, showed higher ratios (see Fig. 3) than samples from the more coastal Sapelo Sound site, which likely received more solar irradiation prior to sample collection. Regardless of the cause, it is clear from our spatialtemporal study that CO$_2$ photoproduction is more variable than CO photoproduction (Figs. 2, 3). This is consistent with both CO$_2$ AQY spectra being more affected by pre-exposure to sunlight than CO AQY spectra, and differing molecular
compositions of the source material for the two different photoproducts as discussed above in Sect. 4.1.

4.3 Water sample characteristics and their relationship to photoproduction efficiencies

Salinity has been suggested as a potential tracer for AQY variations in studies such as Xie et al. (2009) in the Beaufort Sea and Zhang et al. (2006) in the St. Lawrence River estuary. Each have shown a decrease in the photochemical efficiency of CO production with decreasing salinity. Stubbins et al. (2011) also found an inverse trend between CO AQYs and salinity in the Tyne River estuary. This was not explained by simple conservative mixing and suggests that this relationship is more strongly tied to CDOM. Conversely, this trend was not seen by White et al. (2010) in the Delaware River estuary where they reported nearly constant photochemical efficiencies (AQY at 330 nm) for both CO₂ and CO throughout the estuary, with the exception of a strong increase for both products at a salinity of zero in the Delaware River. The current study shows no indication of varying photochemical efficiencies with salinity for either product. This may reflect the limited salinity range that describes the vast majority of our samples, with most falling between 20 and 33. Without larger and more evenly distributed salinity data such as that examined by Zhang et al. (2006) and Xie et al. (2009) (salinities from 0 to 35) as well as White et al. (2010) (salinities from 0 to 23), any salinity related AQY variation present in our samples would be very difficult to define.

The clear increase in photoproduction with increasing CDOM absorption is expected (see Eq. 7) and could correlate with changes in CO and CO₂ production efficiency. However, the CDOM normalized production data show no such trend for either product, indicating that CDOM in darker waters is neither more nor less efficient at producing CO and CO₂ than CDOM in less absorbive samples (Fig. 5). This is in contrast to other studies in the St. Lawrence River estuary (Zhang et al., 2006), the Beaufort Sea (Xie and Zafiriou, 2009) and the Tyne River estuary (Stubbins et al., 2011). These studies did not address CO₂ photochemistry, but all three found an increase in CO AQY values with increasing CDOM absorption coefficients using transect data from high absorption (upper estuary) to low absorption (coastal) waters. In contrast, results from our coastal study do not represent transect data from upper estuaries to the open ocean taken over a short time period, but rather repeated sampling of specific estuarine locations over the course of a year. The constant efficiency of CO photoproduction over all samples seen in our yearlong investigation is consistent with a relatively homogeneous influence of marsh-derived carbon at all three sites.

While the measured production in this study shows a slight decrease for both CO and CO₂ with increasing spectral slope, there is no such trend with CDOM normalized production for either photoproduct (Fig. 6). If, in fact, spectral slope values reflect molecular size variations over the sample range collected in this study, the lack of trend with CDOM normalized photoproduction suggests that it does not affect photochemical efficiency for either photoproduct. The E₂:E₃ ratio (i.e. the ratio between absorption at 254 nm and 365 nm) is similar to spectral slope, and has been shown to be an indicator of the charge-transfer character of CDOM (Del Vecchio and Blough, 2004). Del Vecchio and Blough (2004) have shown that increased absorption at long wavelengths is an indication of higher charge-transfer character. This charge-transfer character has been shown to be a potential predictor of the photochemical activity of reactive oxygen species, such as \(^{1}\text{O}_2\) and \(\text{H}_2\text{O}_2\) AQYs (Dalrymple et al., 2010; Sharpless, 2012). We investigated the relationship between our CO and CO₂ reaction efficiencies and E₂:E₃, but found that there was no significant relationship between them (data not shown). As with the spectral slope, it is possible that the limited range of E₂:E₃ ratios present in our data set (4.95–7.24) is not enough to show a definitive relationship.

Stubbins et al. (2008) showed that the aromaticity of a sample was positively correlated with increased photoproduction of carbon monoxide. While aromatic content was not directly measured in this study, Weishaar et al. (2003) demonstrated a strong positive linear relationship between SUVA\(\text{254}\) and aromatic content of natural water samples. Examining this relationship between SUVA\(\text{254}\) (as a proxy for aromatic content) and measured CO and CO₂ photoproduction showed a slight positive correlation as expected based on the work by Stubbins et al. (2008). However, due to the small range of SUVA\(\text{254}\) values measured in our sampling regime, this relationship did not prove to be statistically significant. The photochemical production efficiency, as indicated by CDOM-normalized photoproduction for both CO and CO₂, showed no statistically significant relationship with aromaticity, although a slight indication of a negative correlation can be seen in Fig. 7 for CO. Zhang et al. (2006) showed that SUVA\(\text{254}\) was a good indicator of the efficiency of CO photoproduction in the St. Lawrence River estuary, showing a significant positive correlation. The SUVA\(\text{254}\) values obtained by Zhang et al. (2006), however, were higher (4–7) and spanned a larger range than those from our study of the Georgia coast where most SUVA\(\text{254}\) values were concentrated in the 2–3 range. This smaller range of SUVA\(\text{254}\) and variability in our data set makes it less likely that robust trends can be defined.

4.4 Contribution of photochemistry to the coastal system

Based on previous estimates (White et al., 2010; Johannessen and Miller, 2001; Miller and Zepp, 1995), there is a strong potential for photochemistry to significantly impact the coastal and global cycling of DOC by formation of inorganic carbon in the form of CO and CO₂. In this study, photochemistry was found to contribute to the direct oxidation of DOC in the estuaries and inner shelf for the South
Atlantic Bight (SABi) by conversion of $7.13 \times 10^9$ g C yr$^{-1}$ in the form of CO, along with $1.60 \times 10^{11}$ g C yr$^{-1}$ in the form of CO$_2$. Calculations using our “mean” AQY give a slightly smaller estimate for the contribution of photochemistry in the form of CO$_2$. The ratio method for calculating CO$_2$ photoproduction in the ocean is often used for open ocean and global calculations due to the lack of reliable AQY data for CO$_2$ in blue water marine systems. Adding CO and CO$_2$ photoproduction together, the contribution of photochemistry to the formation of inorganic carbon products is between $1.46 \times 10^{11}$ g C yr$^{-1}$ and $1.67 \times 10^{11}$ g C yr$^{-1}$ in the SABi. Cai (2011) and Jiang et al. (2008a), using field measurements of in situ CO$_2$ saturation state, found this same area to be a source of CO$_2$ to the atmosphere, degassing $2.29 \times 10^{12}$ g C yr$^{-1}$ to the atmosphere. This result is supported by estimates of remineralization rates in a heterotrophic ecosystem (Cai, 2011). Using Cai’s (2011) values for comparison, photochemical production of inorganic carbon products in this system could contribute between 6.4 % to 7.3 % of the total inorganic carbon signal annually.

Based on numbers from Hopkinson (1988), Alberts and Takacs (1999); Moran et al. (1999), the total DOC input to the entire South Atlantic Bight annually is $5.1 \times 10^{12}$ g C yr$^{-1}$ (Cai, 2011), with the majority (~86%) of this carbon being marsh-derived DOC. A significant portion of this organic carbon is transformed, presumably by heterotrophic bacteria, to inorganic carbon during its transit from marsh and river to the open ocean (Cauwet, 2002). Comparing the magnitude of DOC input to photochemical estimates, it is possible that between 2.9 % and 3.3 % of this carbon annually may be directly removed from the DOC pool by photochemical oxidation to CO and CO$_2$ in the SABi.

5 Conclusions

The extensive nature of this AQY data set contributes improved constraints for the spatiotemporal variability of both CO and CO$_2$ photochemical production in the subtropical coastal ocean. It represents a longer-term examination of coastal CO photochemistry than the much more common transect studies published to date. The photochemical efficiency of CO production over all Georgia coastal estuarine environments studied varied annually within 21.7 % with a seasonal pattern. The variation observed in our CO$_2$ data set is overall larger than that for CO. The Sapelo Sound samples that should be most representative of offshore coastal waters, however, showed a CO$_2$ photoproduction constrained to within 9.6 % year-round. The average CO$_2$ to CO photoproduction ratio for this study was ~23. Together, these findings allow for better constraints on estimates of the direct influence of photochemistry in coastal organic carbon models. Our calculations indicate the potential for direct mineralization of DOC by photochemical oxidation to inorganic gases in the SABi from 2.9 % to 3.3 %. This suggests that photochemical oxidation in estuarine and coastal environments is a small but potentially significant contributor to coastal carbon cycling.

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