Interactive comment on "Sources, fluxes, and behaviors of fluorescent dissolved organic matter (FDOM) in an estuarine mixing zone: Results from the Nakdong-River Estuary, Korea" by Shin-Ah

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Thank you for sending us the minor comments. We made necessary corrections as below. We hope that our manuscript is acceptable for publication in *Biogeosciences* at this time.

Minor comments

Lines 142-145: I could not understand this description. What was compared between uncorrected and corrected values for the inner filter effect? Did the authors compare fluorescence intensity at a specific pair of Ex/Em wavelengths (e.g., Ex/Em=350/450)? Please clarify it in the revised manuscript.

=> We compared the peak values for the three components that were used in this study after scanning all wavelengths for five samples. The values are compared for the three peaks between the "uncorrected" values without inner filer correction and the "corrected" values with inner filter correction. This is clarified in the revised version (lines 142-144).

Lines 165 and 181: I think "Fig. 1" should be "Fig. 2". Please check these.

=> corrected.

Lines 197-204: A description regarding with C4 is missing, even though the authors described that "Four components were identified in the water samples from the EEMs dataset". The description should be added.

=> We identified four components using PARAFAC model but we used the three peaks which were validated for the known FDOM components. Thus, we used only three components in this study. In order to clarify this, we changed this sentence to "Three components were identified in the water samples from the EEMs dataset".

Line 242: I think "leached" is better than "extracted" here. Please consider it.

=> corrected.

1	Sources, fluxes, and behaviors of fluorescent dissolved organic matter (FDOM) in the
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20 Abstract

We monitored seasonal variations of dissolved organic carbon (DOC), stable carbon isotope of 21 DOC (δ^{13} C-DOC), and fluorescent dissolved organic matter (FDOM) in water samples from a 22 fixed station in the Nakdong-River Estuary, Korea. Sampling was performed every hour during 23 24 spring tide once a month from October 2014 to August 2015. The concentrations of DOC and humic-like FDOM showed significant negative correlations against salinity ($r^2=0.42-0.98$, p < 25 0.0001), indicating that the river-originated DOM components are the major source and behave 26 conservatively in the estuarine mixing zone. The extrapolated δ^{13} C-DOC values (-27.5 – 27 24.5‰) in fresh water confirm that both components are mainly of terrestrial origin. The slopes 28 of humic-like FDOM against salinity were 60-80% higher in the summer and fall, due to higher 29 terrestrial production of humic-like FDOM. The slopes of protein-like FDOM against salinity, 30 however, were 70-80% higher in spring, due to higher biological production in river water. Our 31 32 results suggest that there are large seasonal changes in riverine fluxes of humic and proteinlike FDOM to the ocean. 33

35 **1. Introduction**

The global annual flux of dissolved organic carbon (DOC) via rivers is approximately 0.17 - 0.36×10^{15} g (Meybeck, 1982; Ludwig et al., 1996; Dai et al., 2012). The DOC delivered from riverine discharges as well as *in situ* production through biological activities significantly affects carbon and biogeochemical cycles in coastal waters (Hedges, 1992; Bianchi et al., 2004; Bauer et al., 2013; Moyer et al., 2015).

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42 Generally, DOC includes fluorescent dissolved organic matter (FDOM), which emits fluorescent light due to its chemical characteristics. As FDOM accounts for 20 - 70% of the 43 DOC in coastal waters (Coble, 2007) and controls the penetration of harmful UV radiation in 44 the euphotic zone, it plays a critical role in carbon cycles as well as biological production. In 45 addition, FDOM is known as a powerful indicator of humic and protein-like substances (Coble, 46 2007) in coastal waters. River discharge is generally the main source of humic-like FDOM in 47 coastal waters, although it is also produced through in situ microbial activity (Romera-Castillo 48 49 et al., 2011). In contrast, protein-like FDOM is known to be from biological production as well as anthropogenic sources (Baker and Spencer, 2004). Terrestrial humic substances behave 50 conservatively in coastal areas due to their refractory characteristics (Del Castillo et al., 2000), 51 52 whereas protein substances behave non-conservatively in many estuaries due to their relatively rapid production and degradation (Vignudelli et al., 2004). 53

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55 The magnitudes of DOC and FDOM fluxes from rivers are generally dependent on 56 rainfall, discharge, and temperatures (Maie et al., 2006; Jaffé et al., 2004; Huang and Chen, 57 2009). In the estuarine mixing zone, intensive biogeochemical processes occur through photo-58 oxidation, microbial degradation, or physicochemical transformations (i.e., flocculation, 59 sedimentation) (Bauer and Bianchi, 2011; Moran et al., 1991; Benner and Opsahl, 2001; 60 Raymond and Bauer, 2001). Recent studies have demonstrated large seasonal variations as high 61 as 40%, in DOC export from rivers to the ocean (Burns et al., 2008; Bianchi et al., 2004; Dai 62 et al., 2012). However, the seasonal variations in sources, fluxes, and behaviors of DOC and 63 FDOM in the estuarine mixing zone are still poorly understood.

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In this study, we analyzed DOC, δ^{13} C-DOC, and FDOM in estuarine water samples 65 collected monthly from the Nakdong-River estuary. Sampling was conducted at a fixed 66 platform, which has been utilized for monitoring various environmental parameters. This 67 sampling station is advantageous because we can collect water samples for a wide range of 68 69 salinities throughout tidal fluctuations. Using the data obtained from this unique station, we were able to determine (1) the behaviors of DOM in the estuarine mixing zone, (2) the fluxes 70 of DOM from rivers based on the slopes between salinities and DOM components, and (3) the 71 changes in DOM sources using δ^{13} C-DOC in the estuarine samples. The slope measurement in 72 73 the mixing zone may represent the endmember of DOM components in rivers better than sitespecific measurements in the river, by integrating larger spaces and times. 74

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76 2. Materials and methods

77 2.1 Study site

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The Nakdong-River Estuary, which is the estuary of the longest river in Korea, is a

major source of water supplying the needs for drinking, agriculture, and industry. The main channel of Nakdong River is approximately 510 km in length with a watershed area of approximately 23,380 km². It faces the south-eastern coastal area of the Korean peninsula, passing through Busan which is the second largest city in Korea. The mean annual precipitation is 1150 mm, and most precipitation (60-70%) occurs during the summer monsoon and typhoon seasons (Jeong et al., 2007). To manage water supply and saltwater intrusion, estuary dams were constructed in the mouth of the river in 1987.

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87 2.2 Sampling

Water samples were collected at the sampling site which is located 560 m downstream 88 from the dam (Fig. 1). The sampling period was from October 2014 to August 2015. The 2-L 89 water sampling was conducted every hour for 24 hours during spring tide using an auto-sampler 90 91 (RoboChemTMAutosampler, Model S3-1224N, Centennial Technology, Korea), with a depth of the water intake 1 m below the surface. After samples were collected in acid-cleaned 92 polyethylene bottles, they were moved to the laboratory within 24 hours. All water samples 93 94 were filtered using pre-combusted GF/F filters. The FDOM samples were stored in precombusted amber glass vials and kept below 4°C in a refrigerator before analysis. The DOC 95 and δ^{13} C-DOC samples were acidified to pH ~2 using 6 M HCl to avoid bacterial activities and 96 stored in pre-combusted glass ampoules. Ampoules were fire-sealed to prevent the samples 97 from any contaminations. The samples were analyzed for DOC and CDOM within a week. 98 99 Salinity was measured using a YSI Pro Series conductivity probe sensor in the laboratory. The real-time and compulsory discharge volume data from the dam are available at 100 http://www.water.or.kr, provided by K-Water. The monitoring program at this station is 101

maintained by Korea Environment Management Corporation (KOEM). The water temperature
data are recorded automatically at the site. The data are available at https://www.koem.or.kr.

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105 *2.3 Analytical methods*

The concentrations of DOC were determined by a high-temperature catalytic oxidation 106 (HTCO) method using a TOC-VCPH analyzer (Shimadzu, Japan). Standardization was 107 108 performed based on the calibration curve of acetanilide in ultra-pure water. The acidified samples were purged with carbon dioxide (CO_2) free carrier gas for 2 min to remove inorganic 109 carbon. The samples were then injected into a combustion column packed with Pt-coated 110 alumina beads and heated to 720°C. The CO₂ evolving from combusted organic carbon was 111 detected by a non-dispersive infrared detector (NDIR). Our DOC method was verified with 112 Deep Seawater Reference (DSR) samples for DOC (44-46 µmol L⁻¹) produced by the 113 University of Miami, USA. 114

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The values of δ^{13} C-DOC were measured using a TOC–IR–MS instrument consisting 116 of an IR-MS instrument coupled with a Vario TOC cube (Isoprime, Elementar, Germany). The 117 118 TOC instrument uses a common high-temperature catalytic combustion method (Kirkels et al., 2014). The analytical method is fully described in Kim et al. (2015). Briefly, 10 mL of filtered 119 samples was purged with O₂ gas for 20–30 min to completely remove DIC after the samples 120 121 were acidified to pH ~2. Then, 1 mL of the sample was injected into Pt-impregnated catalyst in a quartz tube. In this tube, the DOC was converted completely to CO₂ at 750°C, which was 122 then fed through a water trap followed by a halogen trap. After DOC was detected by an NDIR 123

detector, the CO₂ gas was entered the TOC-IR-MS interface by the O₂ carrier gas. In the 124 interface, the CO₂ was transferred to the IR-MS instrument following the removal of any 125 126 interfering gasses. The δ^{13} C-DOC value of blank was measured using the Low Carbon Water (LCW) from Hansell lab (University of Miami), which contains less than 2 µM DOC. Certified 127 IAEA-CH6 sucrose (International Atomic Energy Agency, $-10.45 \pm 0.03\%$) prepared with the 128 129 low carbon water was used as a standard solution. A standard sample was analyzed for every sample queue (once before or after ten samples) to check a drifting effect during the 130 131 measurements. The blank correction was performed using a method previously described in De Troyer et al. (2010) and Panetta et al. (2008). Our measurement result of δ^{13} C-DOC for the 132 DSR (University of Miami) was $-21.5 \pm 0.1\%$, which is consistent with the results reported 133 134 by Panetta et al. (2008) and Lang et al. (2007). The reproducibility of TOC-IR-MS was ~0.3‰.

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FDOM fluorescence was determined in a scan mode using a spectrofluorometer 136 (SCINCO FluoroMate FS-2) within two days after sampling. Emission (Em) spectra were 137 collected from 250 to 600 nm at 2 nm intervals at excitation (Ex) wavelengths from 250 nm to 138 500 nm at 5 nm intervals. Backgrounds were subtracted for fresh distilled water prepared daily 139 from the sample data to eliminate Raman Scatter peaks (Zepp et al., 2004). All data were 140 obtained in counts per second (cps) and converted to a ppb quinine sulfate standard solution in 141 0.1 N sulfuric acid at Ex/Em of 350/450 nm. The inner filter effect was negligible for these 142 estuarine water samples since the correlation between the uncorrected and corrected values for 143 the inner filter effect was very significant for the three identified peaks (r²=1, n=5). EEMs-144 PARAFAC analysis was performed using a MATLAB R2013a program with a DOMFluor 145 146 toolbox.

148 **3. Results and Discussion**

Salinities ranged from 0.1 to 28.5 over the sampling period of a year. Salinities in the
sampling location were dependent primarily on the volume of river-water discharge from the
dam. The volumes of river discharge were relatively larger in October, April, July, and May.
The mean annual surface water temperature was 16°C, with the lowest temperature (avg. 8°C)
in December and the highest temperature in August (avg. 26°C).

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155 *3.1 Behaviors and sources of DOC in the estuarine mixing zone*

The concentrations of DOC ranged from 100 to 300 μ M, with the highest concentrations in July (avg. 243 μ M) and the lowest concentrations in February (avg. 115 μ M), consistent with the typical DOC concentration ranges in coastal waters (Wang et al., 2004; Raymond and Bauer, 2001). The concentrations of DOC correlated significantly with salinities ($r^2 = 0.59-0.92$, p < 0.0001), indicating that DOC behaves conservatively in the mixing zone of this estuary (Fig. 2A), which is commonly observed in estuarine mixing zones (Laane, 1980; Mantoura and Woodward, 1983; Del Castillo et al., 2000; Clark et al., 2002; Jaffé et al., 2004).

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If the high salinity periods are excluded, both the slope and y-intercept of DOC concentrations versus salinities were highest in July (Fig. 2), which could be due to a higher terrestrial DOC loading in the summer period, as observed in Horsens Fjord, Denmark (Markager et al., 2011). For this comparison, we excluded the high-salinity periods (>20), including December, January, February, and June, since they showed a narrow and low DOC 169 concentration range (103-163 μ M), resulting in large uncertainties by extrapolating them to the 170 fresh water.

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The carbon isotope values in the Nakdong-River Estuary ranged from -28.2 ‰ to -173 17.6. In order to determine the source of DOC in fresh water, we plotted δ^{13} C-DOC values 174 against salinities (Fig. 2B). The conservative mixing curve of δ^{13} C values can be obtained using 175 the two endmember mixing equation (Spiker, 1980; Raymond and Bauer, 2001):

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$$\delta^{13} C_s = \frac{F_r \delta^{13} C_r [DOC]_r + (1 - F_r) \times \delta^{13} C_m [DOC]_m}{[DOC]_s}$$
(1)

where $\delta^{13}C_s$, $\delta^{13}C_r$ and $\delta^{13}C_m$ are the $\delta^{13}C$ -DOC values at a given sample salinity, river endmember salinity, and marine endmember salinity, respectively; F_r is the riverine freshwater fraction calculated from the measured salinities; [DOC]_s and [DOC]_m are the DOC concentrations at a given salinity and marine endmember salinity, respectively; [DOC]_r is the endmember DOC value for the river water (Fig. 2).

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183 The riverine DOC endmember values (S=0‰) ranged from 174 to 284 μ M. The marine endmember value (S=29‰) of DOC is 100 μ M with the δ^{13} C-DOC value of -19‰. If these 184 values from each month are applied, the δ^{13} C-DOC endmember values for the river water 185 extrapolated to be from -27.5 to -24.5‰ (average: -26.2‰). Overall, the carbon isotope values 186 of our samples are fitted well into the conservative mixing curve of the overall trend, with a 187 slight change using different endmember values for different months (Fig. 2B). In general, 188 δ^{13} C-DOC values range from -22 to -18‰ for marine phytoplankton, from -34‰ to -23‰ 189 for terrestrial C3 plants, and from -16‰ to -10‰ for terrestrial C4 plants (Gearing 1988; Clark 190

and Fritz, 1997). Carbon isotope values in our study confirm that the main source of DOC in
the estuarine mixing zone is dominantly from terrestrial C3 plants over all seasons. However,
the value was heavier at lower salinity ranges (S<10) in March and April samples, perhaps in
association with the higher biological production in the river.

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196 *3.2 Behaviors and sources of FDOM in the estuarine mixing zone*

197 Three components were identified in the water samples from the EEMs dataset. Based on the excitation-emission peak location, Component 1 (FDOM_H, Ex/Em = 320/418 nm) is 198 found to be a terrestrial humic-like component (C peak) shown by Coble (2007). Component 199 2 (FDOM_P, Ex/Em = 280/328 nm) is found to be a tryptophan-like component (T peak), which 200 is produced by microbial processes. Component 3 (Ex/Em = 300,325/364 nm) is found to be a 201 marine humic-like component (M peak). Since Component 3 values were significantly 202 correlated with Component 1 ($r^2=0.95$) values, we simply focused on Component 1 (FDOM_H) 203 and Component 2 (FDOM_P) for data interpretations. 204

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The concentrations of FDOM_H ranged from 2.4 to 19.7 quinine sulfate unit (QSU), with the highest concentration in July (avg. 17.6 QSU) and the lowest concentration in June (avg. 3.4 QSU) (Fig. 2C). The concentrations of FDOM_P ranged from 0.6 to 22.4 QSU, with the highest concentration in March (avg. 15.1 QSU) followed by October (avg. 13.6 QSU) (Fig. 2D).

The concentrations of both FDOM components were significantly correlated with 212 salinities ($r^2 = 0.42-0.98$, p<0.0001 for FDOM_H and $r^2 = 0.27-0.96$, p<0.0001 for FDOM_P), 213 214 indicating that they are conservative in the mixing zone (Fig. 2). The slopes of FDOM_H and FDOM_P for each month ranged from -0.15 to -0.59 and -0.15 to -0.71, respectively. The higher 215 FDOM_H slopes in July and October were similar to the trend of DOC (Fig. 2C), which could 216 be due to higher terrestrial FDOM production. However, the seasons (March and April) in 217 which higher FDOM_P slopes occurred differ from those of DOC and FDOM_H, indicating that 218 219 both FDOM components have different source inputs (Fig. 2D).

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221 Although there are large differences in scattering of FDOM components against salinities, it is very difficult to compare scatterings for different seasons in order to discuss the 222 different behaviors of DOM since the scattering is generally larger for the narrow salinity 223 224 ranges. If the winter data are excluded, in March, during the highest biological production period in the river, the correlation coefficient against salinities was the highest for FDOM_P and 225 lowest for FDOM_H. In contrast, in June, during the highest fluvial DOM discharge period, the 226 correlation coefficient against salinities was the highest for FDOM_H and lowest for FDOM_P. 227 This suggests that the biological production and removal, together with other generally known 228 factors such as photo-degradation and sedimentary inputs, may affect the scattering of these 229 FDOM components in the estuarine mixing zone. 230

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As such, there was a significant positive correlation between FDOM_H and DOC concentrations throughout all sampling periods ($r^2 = 0.93$, p<0.0001) (Fig. 3A), suggesting that the main source of FDOM_H and DOC is terrestrial based on δ^{13} C-DOC values. Since FDOM 235 does not usually contribute to a major portion of DOC, a positive correlation between FDOM and DOC has only been observed in specific areas, such as river-estuarine systems (Del 236 237 Vecchio and Blough, 2004; Coble, 2007). Stedmon et al., (2006) demonstrated that stronger correlations were observed between DOC and FDOM as humic substances derived from 238 terrestrial DOM are more colored than DOM produced in situ. In general, terrestrial DOM 239 240 occurring in rivers originates mainly from plant decomposition and leaf litter in the form of humic substances (Huang and Chen, 2009). As such, Gueguen et al., (2006) showed that humic 241 242 materials are more effectively leached from soils during August and September under high temperatures. Thus, higher FDOM_H slopes in August, October, and November, relative to the 243 other periods, could be associated with higher terrestrial inputs of degradation products of soil 244 245 organic matter (Dowell, 1985; Qualls et al., 1991).

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247 In the study region, FDOM_P concentrations were poorly correlated with DOC concentrations ($r^2=0.11$) (Fig. 3B). The slopes of FDOM_P concentrations against DOC 248 concentrations varied significantly over different seasons, with steeper gradients in the spring 249 (March and April) and fall (October). In general, FDOM_P is known to be produced efficiently 250 by biological production in water (Coble, 1996; Belzile et al., 2002; Steinberg et al., 2004; 251 Zhao et al., 2017). Thus, higher FDOM_P concentrations, relative to DOC concentrations, in the 252 spring and fall seems to be associated with the spring and fall phytoplankton blooms in river 253 waters (Mayer et al., 1999; Zhang et al., 2009). 254

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256 *3.3 Fluxes of DOC and FDOM in the estuarine mixing zone*

The fluxes of DOC and FDOM from rivers to the ocean are calculated using the endmember values (C) of these components in rivers multiplied by the river discharge volumes (Q) for each month (Fig. 4). For this estimation, we assumed that (1) the endmember values are the same as the intercepts of the DOC, FDOM_H, and FDOM_P versus salinity plots, and (2) the endmember values measured in the spring tides represent the concentrations of these components for each month.

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River discharge was highest in April and July following heavy precipitation, and the 264 largest discharge volume was about five-fold higher than that of winter discharges (Fig. 4A). 265 However, the monthly variations of DOC endmember (y-intercept) values were quite constant, 266 ranging from 174 - 284 µM. This indicates that the concentrations of DOC in the river are 267 independent of river discharge volumes (Fig. 4B). The DOC endmember values were highest 268 269 in December, followed by July and June (Fig. 4B). The monthly variation trend of FDOM_H endmember values was similar to that of DOC, except for the December value. Excluding the 270 December values, the FDOM_P endmember values were highest in March, February, and 271 October. These endmember trends are consistent with the slope variations explained in the 272 previous section. Although there are large uncertainties in fresh water endmember values of 273 DOC and FDOM in winter owing to narrow, high salinity ranges, we used the endmember 274 values for the flux comparisons since the contribution of the uncertainties may be relatively 275 small due to smaller river discharge volumes in winter. 276

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The riverine DOC flux ranged from 1.6×10^6 mol day⁻¹ (February) to 12.3×10^6 mol day⁻¹ (July), indicating that there are large variations of DOC fluxes to the ocean. The riverine

flux of FDOM_H and FDOM_P ranged from 1.4×10^9 QSU m³ day⁻¹ (December) to 23.1×10^9 QSU m³ day⁻¹ (July) and from 1.6×10^9 QSU m³ day⁻¹ (June) to 16.4×10^9 QSU m³ day⁻¹ (March), respectively. The seasonal variation trend of FDOM_H was similar to that of DOC. The fluxes of FDOM_P in December and March were twofold higher than those of FDOM_H whereas the flux of FDOM_H in July was 2-3 folds higher than that of FDOM_P. This shows that the fluxes of both components of FDOM differ significantly by seasons owing to the different source inputs even though their magnitudes are controlled mainly by river discharges

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It is well known that the single sampling event is not enough to capture the full range 288 289 of natural variability in DOM abundance over all seasons (Stedmon et al., 2006; Huang and Chen, 2009; Markager et al., 2011; Dai et al., 2012; Moyer et al., 2015). Overall, our results 290 show that monthly variations are significant. This implies that our understanding of DOC 291 292 fluxes from large rivers is largely biased, depending on sampling resolution, methods, and hydrogeological settings of a specific river. For example, if summer data are extrapolated to 293 annual river water discharge, the DOC and FDOM_H fluxes can be overestimated up to three 294 times for the Nakdong River. 295

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297 **4. Conclusions**

The concentrations of FDOM_H and DOC showed significant negative correlations against salinities throughout all sampling periods, indicating that they behave conservatively in this estuarine mixing zone. The slopes of both DOC and FDOM_H concentrations versus salinities were highest in July, due to the largest terrestrial DOC loadings. The carbon isotope values showed that the main source of DOC in the estuarine mixing zone is terrestrial C3 plants

303	over all seasons. The slopes of $FDOM_P$ versus salinity were relatively higher in March and
304	April in association with the spring phytoplankton blooms in river and estuarine waters. The
305	monthly fluxes of DOC, $FDOM_H$, and $FDOM_P$ showed large seasonal variations (5-10 folds),
306	suggesting that the estimation of annual riverine fluxes of DOC, $FDOM_H$, and $FDOM_P$ requires
307	careful considerations of seasonal changes in rivers.
308	
309	Competing interests
310	The authors declare that they have no conflict of interest.
311	
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Figure 1. Map of the Nakdong-River Estuary. The square indicates a fixed monitoring site,located 560 m downstream from the dam.



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Figure 2. Salinities versus the concentrations of (A) DOC, (B) δ^{13} C-DOC, (C) FDOM_H, and (D) FDOM_P. The values for the regression lines are excluded for high-salinity periods (>20), including December, January, February, and June, which have large uncertainties in extrapolation. The solid curve (B) is the average conservative mixing line for the two endmember mixing equation. The dotted lines represent the monthly changes in mixing lines for the different monthly endmember values.



453 Figure 3. The plots of the concentrations of DOC versus the concentrations of (A) FDOM_H and
454 (B) FDOM_P.



Figure 4. Temporal variations in discharge volumes, the endmember values of DOC, FDOM_H,
and FDOM_P, and riverine fluxes of DOC, FDOM_H, and FDOM_P in the Nakdong-River Estuary
from October 2014 to August 2015.