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1 **Carbon leaks from flooded land: do we need to re-plumb the inland water active**  
2 **pipe?**

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15 **ABSTRACT**

16 At the global scale, inland waters are a significant source of atmospheric carbon (C),  
17 particularly in the tropics. The active pipe concept predicts that C emissions from  
18 streams, lakes and rivers are largely fuelled by terrestrial ecosystems. The traditionally  
19 recognized C transfer mechanisms from terrestrial to aquatic systems are surface runoff  
20 and groundwater drainage. We present here a series of arguments that support the idea  
21 that land flooding is an additional significant process that fuels inland waters with C at  
22 the global scale. Whether the majority of CO<sub>2</sub> emitted by rivers comes from floodable  
23 land (approximately 10% of the continents) or from well-drained land is a fundamental  
24 question that impacts our capacity to predict how these C fluxes might change in the  
25 future. Using classical concepts in ecology, we propose, as a necessary step forward, an  
26 update of the active pipe concept that differentiates floodable land from drained land.  
27 Contrarily to well-drained land, wetlands combine strong hydrological connectivity with  
28 inland waters, high productivity assimilating CO<sub>2</sub> from the atmosphere, direct transfer of  
29 litter and exudation products to water and waterlogged soils, a generally dominant  
30 allocation of ecosystem respiration below the water surface and a slow gas exchange  
31 rate at the water-air interface. These properties force plants to pump atmospheric C to  
32 wetland waters and, when hydrology is favourable, to inland waters as organic C and  
33 dissolved CO<sub>2</sub>. This wetland CO<sub>2</sub> pump may contribute disproportionately to CO<sub>2</sub>  
34 emissions from inland waters, particularly in the tropics, and consequently at the global  
35 scale. In future studies, more care must be taken in the way that vertical and horizontal  
36 C fluxes are conceptualized along watersheds and 2D-models that adequately account  
37 for the hydrological export of all C species are necessary. In wetland ecosystems,  
38 significant effort should be dedicated to quantifying the components of primary  
39 production and respiration in air, water and waterlogged soils, and these metabolic



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40 rates should be used in coupled hydrological-biogeochemical models. The construction

41 of a global typology of wetlands also appears necessary to adequately integrate

42 continental C fluxes at the global scale.

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45 **1. INTRODUCTION**

46 Continental surfaces play a major role on the present and past climates, in particular  
47 through the exchange of greenhouse gases (GHGs) such as carbon dioxide (CO<sub>2</sub>) and  
48 methane (CH<sub>4</sub>) with the atmosphere (Ciais et al. 2013). Conversely, the global climate  
49 affects the continental carbon (C) budget, as biological productivity and the capacity of  
50 ecosystems to store C are influenced by temperature, rainfall and other climatic  
51 variables (Heimann and Reichstein 2008; Reichstein et al. 2013). The continental C  
52 budget is in addition affected by direct human alterations such as  
53 deforestation/reforestation and other land use changes. On continents, the C cycle is  
54 tightly coupled to the water cycle, and CO<sub>2</sub> and CH<sub>4</sub> budgets strongly depend on how and  
55 how much water circulates through the plants, soil, groundwater, and surface waters to  
56 the coastal ocean. Biogeochemical processes and fluxes in the critical zone, the  
57 permeable layer of the continents from the vegetation top to the aquifer bottom (Lin  
58 2010), have varied drastically at geological time scales (Knoll and James 1987).  
59 Emissions of GHGs from continental ecosystems are expected to be highly sensitive to  
60 precipitation and hydrology in the future. Water is necessary for plant photosynthesis;  
61 moisture strongly controls respiration in soils; the presence of water promotes  
62 anaerobic conditions and CH<sub>4</sub> production in wetlands, while soil desiccation promotes  
63 soil CH<sub>4</sub> oxidation. Water also considerably contributes to continental C budgets because  
64 rivers transport C laterally; C being later trapped in sediments, emitted as CO<sub>2</sub> and CH<sub>4</sub>  
65 to the atmosphere, or exported to the ocean (Garrels and Mackenzie 1971; Meybeck  
66 1982; Cole et al. 2007).

67

68 In terms of CO<sub>2</sub> and CH<sub>4</sub> fluxes, continental landscapes act as a heterogeneous mosaic,  
69 and some ecosystems store or emit more atmospheric C than others. Some small



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70 surfaces can behave as hotspots and disproportionately contribute to the total C mass  
71 balance at the regional, continental and global scales. Surface waters are recognized  
72 hotspots for CO<sub>2</sub> and CH<sub>4</sub> fluxes (Cole et al. 1994; Cole and Caraco 2001; Bastviken et al.  
73 2011; Raymond et al. 2013; Holgerson and Raymond 2016). Natural surface waters  
74 include the open waters of streams, lakes, rivers and estuaries (approximately 3.5% of  
75 the continents) as well as intermittently flooded land, where a canopy of vegetation is  
76 active above the water and/or when water is temporarily absent: swamps, marshes and  
77 floodplains, also called wetlands, that occupy approximately 10% of the continents  
78 (Downing 2009). In general, inland waters and wetlands show higher atmospheric C  
79 exchange rates per surface area than the surrounding land: Wetlands are recognized for  
80 their high productivity, sedimentary organic carbon (OC) burial and CH<sub>4</sub> emissions  
81 (Mitsch et al 2013). Inland waters (rivers, streams, lakes and reservoirs) act as a very  
82 significant source of atmospheric CO<sub>2</sub> at the global scale (Raymond et al. 2013).

83

84 Although the magnitude of CO<sub>2</sub> outgassing from inland surface waters at the global scale  
85 is still subject to large uncertainties, there is consensus that the quantity of C exported  
86 from land to freshwaters (1.9-3.2 PgC yr<sup>-1</sup>) was larger than the C flux ultimately reaching  
87 the ocean (0.9 PgC yr<sup>-1</sup>, Fig. 1b). Cole et al. (2007) have conceptualized inland waters as  
88 an active pipe (Fig. 1b), receiving, processing, emitting, and storing terrestrial C during  
89 its travel from land to the ocean, as opposed to a passive pipe that simply transports  
90 terrestrial C conservatively to the ocean (Fig. 1a), as generally assumed in earlier  
91 literature from the 1970's and 1980's (Garrels and Mackenzie 1971; Meybeck 1982).  
92 Since this definition, it has been assumed that most of the C emitted by inland waters  
93 was initially fixed upland by terrestrial vegetation, then transported from soils to  
94 aquatic systems with runoff and drainage, and finally emitted downstream as CO<sub>2</sub> to the



95 atmosphere. Because no satisfactory methods are available yet to estimate directly the  
96 flux of C across the land-water boundary (e.g., Deirmendjian et al. 2018), this flux is  
97 calculated as the sum of outgassing from inland waters, burial in freshwater and  
98 estuarine sediments, and export to the coastal ocean (Cole et al. 2007). However, the  
99 processes controlling C fluxes at the land-water interface are poorly understood and  
100 some potential inconsistencies could arise when comparing C budget derived from  
101 terrestrial studies with those derived from aquatic studies. Here, we provide some  
102 additional evidence demonstrating that the transfer of terrestrial C to rivers could occur  
103 preferentially from floodable ecosystems. We suggest that wetlands behave not only as a  
104 significant source of atmospheric CH<sub>4</sub> and a long-term C sink in soils (Mitsch et al. 2013)  
105 but also as an efficient CO<sub>2</sub> pump that exports dissolved and particulate C to inland  
106 waters. Using classical concepts in ecology, we analyse qualitatively and quantitatively  
107 how ecosystem production and respiration affect C export from drained-land and from  
108 wetlands. We stress that our current understanding of processes and our ability to  
109 measure and quantify C metabolic and hydrological fluxes must be considerably  
110 improved to understand the origin of carbon in inland waters and predict future  
111 continental GHG budgets in the mosaic of continental ecosystems.

112

## 113 **2. CONCEPTUALIZING AND FORMULATING C FLUXES**

114 Fluxes of C through the boundaries of an ecosystem, *i.e.*, vertical exchange with the  
115 atmosphere and burial in soils and sediments on the one hand, and horizontal exchange  
116 between lands, wetlands and aquatic ecosystems on the other hand, are driven by  
117 metabolic processes in each ecosystem and physical processes that transport C such as  
118 hydrology, wind, turbulent mixing, sediment deposition/resuspension, etc. Following



119 the conventions of Chapin et al. (2006), the net CO<sub>2</sub> exchange of an ecosystem with the  
120 atmosphere is partitioned into several forms of C fluxes (Fig. 2):

$$121 \quad -NEE = NECB + F_{\text{other}} + E \text{ (Eq. 1)}$$

122 where NEE is net ecosystem exchange (the net CO<sub>2</sub> flux from the ecosystem to the  
123 atmosphere), NECB is the net ecosystem carbon balance (the net C accumulation in the  
124 ecosystem), F<sub>other</sub> is the sum of vertical fluxes of volatile forms of C other than CO<sub>2</sub> (CH<sub>4</sub>,  
125 carbon monoxide, volatile organic carbon) from the ecosystem to the atmosphere and E  
126 is horizontal C export by hydrological transport, trading of food, feed and wood (Ciais et  
127 al. 2008). Among the components of E, only hydrological horizontal transport of C will  
128 be discussed in this paper. Note that, by convention, NEE is opposite in sign to NECB  
129 because NEE is defined by atmospheric scientists as a C input to the atmosphere,  
130 whereas NECB is defined by ecologists as a C input to ecosystems (Chapin et al. 2006).

131

132 Regarding metabolic fluxes, net ecosystem production (NEP) is defined as:

$$133 \quad NEP = GPP - ER \text{ (Eq. 2)}$$

134 where GPP is gross primary production and ER is ecosystem respiration. For conceptual  
135 and methodological reasons, it is necessary to consider separately the autotrophic and  
136 heterotrophic components of ER as:

$$137 \quad NEP = GPP - AR - HR \text{ (Eq. 3),}$$

$$138 \quad NPP = GPP - AR \text{ (Eq. 4), and:}$$

$$139 \quad NEP = NPP - HR \text{ (Eq. 5)}$$

140 where AR and HR are, respectively, the autotrophic and the heterotrophic components  
141 of ER and NPP is net primary production. A positive NEP (Eq. 2) reduces the  
142 concentration of CO<sub>2</sub> and/or dissolved inorganic carbon (DIC) inside the ecosystem and  
143 generates a gradient that causes atmospheric CO<sub>2</sub> to enter the ecosystem. One process



144 that makes -NEE diverge from NEP and NECB is when significant amounts of inorganic C  
145 enter or leave the ecosystem as DIC in the aquatic phase with horizontal hydrological  
146 transport rather than through atmospheric exchange (Chapin et al. 2006). In addition to  
147 this divergence between -NEE and NEP, NECB deviates from NEP when C enters or  
148 leaves the ecosystem in forms others than CO<sub>2</sub> or DIC (Eq. 1). This includes horizontal  
149 transport of particulate and dissolved OC by hydrological processes, as well as vertical  
150 CH<sub>4</sub> fluxes, a secondary C flux that is significant for the active pipe concept, as well as for  
151 climate regulation.

152

153 An adequate conceptualization of atmospheric C fluxes along watersheds implies first  
154 the definition of functional boundaries inside the boundless C cycle (Battin et al. 2009),  
155 at least between three types of ecosystems that have fundamentally different properties  
156 with respect to atmospheric CO<sub>2</sub> (Fig. 2): (1) the terrestrial, never flooded land and its  
157 biosphere (forest, crops, shrub, grassland and their well-drained soils and  
158 groundwater); (2) the floodable land and its mosaics of wetlands with extremely  
159 variable ecological and hydrological properties; (3) the open waters of streams, lakes  
160 and rivers. Some estimations of CO<sub>2</sub> outgassing from inland waters have included  
161 wetland surface areas (Richey et al. 2002; Aufdenkampe et al. 2011; Sawakuchi et al.  
162 2017), while some others have not (Cole et al. 2007; Tranvik et al. 2009; Raymond et al.  
163 2013). However, wetlands are functionally different from inland waters because their  
164 canopy of vegetation can alter the direction of atmospheric CO<sub>2</sub> exchange (Raymond et  
165 al. 2013; Abril et al. 2014). Assuming that the CO<sub>2</sub> flux at the water-air interface  
166 equals -NEE in wetlands (Richey et al. 2002) implicitly supposes that GPP and the aerial  
167 compartment of AR (Fig. 2b) are null or exactly balanced, which is incorrect. A functional  
168 definition of wetlands with respect to C cycling could be the *intermittent and/or*





169 *vegetated flooded land*, in contrast with the well-drained land which is never flooded and  
170 whose topsoil is never waterlogged, and with the permanent and open waters of lakes  
171 and rivers without emerged or floating vegetation. This definition allows clear  
172 delineation of the three sub-systems using remote sensing (e.g., Melack and Hess 2010)  
173 and is also functional with respect to the conceptualization and quantification of C  
174 cycling (Fig. 2).

175

176 Second, our conceptual model should be two-dimensional (vertical and up-downriver),  
177 and should consider the hydrological net export term  $E$  in Eq. 1 as a potentially  
178 significant component of  $-NEE$  and  $NECB$  (Fig. 2), in accordance with the active pipe  
179 concept. In well-drained terrestrial ecosystems, surface runoff and drainage export  $C$  to  
180 inland water, and  $E$  is necessarily always positive. In inland water and wetlands,  $E$  must  
181 be conceptualized and quantified as the net balance between hydrological import to and  
182 export from the ecosystems and, depending on each case,  $E$  can be positive or negative.  
183 In fact,  $C$  fluxes along watersheds must be seen as a cascade from one sub-system  
184 upstream to another sub-system downstream. Several chemical forms of  $C$  are involved  
185 in the  $E$  term, which can be written as the sum of the export of four terms:

$$186 \quad E = E_{POC} + E_{DOC} + E_{CO_2} + E_{CH_4} \text{ (Eq. 6)}$$

187 Particulate and dissolved organic C (POC and DOC) are derived from NPP; DIC is in part  
188 the result of ER, that release dissolved  $CO_2$  (as well as  $CH_4$ ) to waters and in part the  
189 result of chemical weathering that generates alkalinity. Weathering of carbonate and  
190 silicate rocks is mediated by soil  $CO_2$  derived from respiration, so that weathering is also  
191 a component of ER. Because chemical weathering is assumed to occur mostly upland,  
192 alkalinity is considered as a relatively conservative chemical form of river C, although  
193 some exceptions have been reported in floodplains of tropical rivers (Boucher et al.



194 2012; Geeraert et al. 2017). Here, we will discuss only the fraction of DIC that occurs as  
195 excess  $\text{CO}_2$ , that is, the DIC that is potentially lost after complete water-air equilibration  
196 (Abril et al. 2000). Concerning dissolved  $\text{CH}_4$ , the role of wetlands was identified in the  
197 literature for sustaining  $\text{CH}_4$  emissions in adjacent rivers (Borges et al. 2015b) and lakes  
198 (Juutinen et al. 2003). However, owing to its low solubility and the fact that emissions  
199 from wetlands occur mostly as ebullition or through plants (contributing to the  $F_{\text{other}}$   
200 term in Fig. 2B), the contribution of  $E_{\text{CH}_4}$  to E is small (few percent) in most ecosystems.  
201  
202 NEE is generally negative in forests (Luyssaert et al. 2010; Ciais et al. 2013) and  
203 wetlands (Morison et al. 2000; Saunders et al. 2007; Lu et al. 2016) but positive in lakes  
204 and rivers (Cole et al. 1994; 2007; Raymond et al. 2013) (Fig. 3). Compared to NEE,  
205 exchange of  $\text{CH}_4$  with the atmosphere ( $F_{\text{other}}$  in Eq. 1) is significant in wetlands but not in  
206 forests (Ciais et al. 2013; Saunois et al. 2016) and probably not in inland waters. Indeed,  
207 budgets of  $\text{CH}_4$  emissions from inland waters strongly depend on whether wetland areas  
208 were included or not and, in general, open waters of rivers and lakes emit  $\text{CH}_4$  at rates  
209 approximately 100 times lower than  $\text{CO}_2$  (Melack et al. 2004; Bastviken et al 2011;  
210 Borges et al. 2015a). The occurrence of a horizontal transport of C by streams and rivers  
211 implies a positive E term in terrestrial ecosystems, where -NEE should exceed NECB.  
212 The E is probably also large in wetlands, where -NEE likely exceeds net storage in soils  
213 plus  $\text{CH}_4$  emissions (Eq. 1; Fig. 1c). In contrast, in aquatic systems NECB exceeds -NEE  
214 and E is negative (Cole and Caraco 2001; Battin et al. 2008) because these ecosystems  
215 receive in general more C from upstream than they export downstream. In addition, the  
216 fact that part of E occurs as OC implies that NEP exceeds NECB in terrestrial systems and  
217 wetlands that export OC, whereas NECB will exceed NEP for instance in lakes or  
218 estuaries that receive and store large amounts of allochthonous OC in their sediments



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219 (Lovett et al. 2006; Cole et al. 2007; Tranvik et al. 2009). In general, C fluxes at the  
220 boundaries of ecosystems and metabolic fluxes inside the ecosystems suggest that the  
221 magnitude of the export term E in Eq. 1 and Fig. 2 and the deviation of -NEE from NECB  
222 and from NEP, will strongly depend on their hydrological connectivity, together with the  
223 allocation of GPP and ER in air and water.

224

225

226 **3. THE INLAND WATER PERSPECTIVE**

227 Global estimates of CO<sub>2</sub> emissions from inland waters (Cole et al. 1994; Raymond et al.  
228 2013; Lauerwald et al. 2015) are derived from CO<sub>2</sub> flux intensities computed from the  
229 water-air gradient of the partial pressure of CO<sub>2</sub> (pCO<sub>2</sub>) and the gas transfer velocity at  
230 the water-air interface and scaled to the surface area of lakes and rivers. Each of the  
231 three terms suffers for uncertainties and generally poor data coverage. Cole et al. (1994)  
232 provided the first quantification of the CO<sub>2</sub> emission to the atmosphere from lakes (0.1  
233 PgC yr<sup>-1</sup>), which was later confirmed by an updated calculation by Sobek et al. (2005).  
234 Cole and Caraco (2001) estimated global CO<sub>2</sub> degassing for rivers and streams, which  
235 has been recently re-evaluated by Raymond et al. (2013) and Lauerwald et al. (2015).  
236 The two latter studies are based on pCO<sub>2</sub> computed from pH and alkalinity from the  
237 same database (GLORICH, Hartmann et al. 2014) but with different data selection  
238 criteria and scaling approaches. Raymond et al. (2013) extrapolated discrete pCO<sub>2</sub>  
239 values per COSCATS catchment aggregated units (Meybeck et al. 2006) and obtained a  
240 global CO<sub>2</sub> emission to the atmosphere of 0.3 PgC yr<sup>-1</sup> from lakes and 1.8 PgC yr<sup>-1</sup> from  
241 rivers and streams. A potential problem in this estimation comes from the calculation of  
242 pCO<sub>2</sub> from pH and alkalinity, which greatly overestimates pCO<sub>2</sub> (up to several hundred  
243 percent) in many acidic organic rich “black” waters such as those found in the tropics  
244 and the boreal zone (Abril et al. 2015). Lauerwald et al. (2015) computed river pCO<sub>2</sub>  
245 values on a regular grid (1°x1°), using a multiple regression model based on the  
246 GLORICH pCO<sub>2</sub> data and modelled terrestrial NPP on the catchment, population density,  
247 air temperature and slope; this method provided a lower estimate of global CO<sub>2</sub>  
248 emission for rivers of 0.7 PgC yr<sup>-1</sup>. The strong divergence of global CO<sub>2</sub> emission  
249 estimates in these two studies most likely reflects the low data coverage in tropics that  
250 account for nearly 80% of the modelled global emission, although in the GLORICH



251 database nearly all of the data in the tropics are from the Amazon. Recent direct pCO<sub>2</sub>  
252 measurements in several African rivers (Borges et al. 2015a), and in the Amazon (Abril  
253 et al. 2014) scaled to the tropics with wetland coverage (Borges et al. 2015b) provide a  
254 value of 1.8±0.4 PgC yr<sup>-1</sup> of CO<sub>2</sub> outgassing from tropical rivers alone (latitude < 25°),  
255 and thus support the higher estimate of Raymond et al. (2013). A larger estimate of the  
256 global river CO<sub>2</sub> outgassing of 3.9 PgC yr<sup>-1</sup> has been published recently (Sawakuchi et al.  
257 2017). However, we choose not to consider this number in our analysis because it is  
258 based on observations in the Amazon River that include the floodplain areas with a  
259 canopy of vegetation above the water.

260

261 According to the active pipe concept (Fig. 1b), the emission of CO<sub>2</sub> to the atmosphere  
262 from inland waters is attributed to terrestrial C fixed by plants on the catchment. The  
263 transfer occurs as (1) an input of dissolved CO<sub>2</sub> (and CH<sub>4</sub>) originating from soil  
264 respiration, that will be further degassed from waters (E<sub>CO2</sub> and E<sub>CH4</sub> in Eq. 6); (2) an  
265 input of particulate and dissolved organic C (E<sub>DOC</sub> and E<sub>POC</sub>) followed by heterotrophic  
266 degradation to CO<sub>2</sub> and CH<sub>4</sub> in the aquatic system (Del Giorgio et al. 1999; Prairie et al.  
267 2002; Cole et al. 2000; Battin et al. 2008; Hotchkiss et al. 2015). Inland waters,  
268 particularly lakes, also store significant quantities of OC mainly of terrestrial origin in  
269 their sediments (Cole et al. 2007; Tranvik et al. 2009). In aquatic systems, all the GPP  
270 and ER occur in water and sediments (Fig. 2c) and can be quantified with *in vitro* or *in*  
271 *situ* incubations. In addition, the CO<sub>2</sub> outgassing flux measured with floating chambers in  
272 open waters give a direct estimate of -NEE (although this method may create artefacts at  
273 the water-air interface), and diurnal changes in water pCO<sub>2</sub> (or oxygen concentration)  
274 can provide an estimate of GPP and ER. Battin et al. (2008) made a global synthesis of  
275 aquatic metabolism rate measurements and confirmed that stream, river and estuarine



276 ecosystems are overall net heterotrophic and respire a total flux of 0.3 PgC yr<sup>-1</sup>. The fact  
277 that net heterotrophy (negative NEP) is in general lower than CO<sub>2</sub> outgassing in inland  
278 waters, led Hotchkiss et al. (2015) to differentiate “internal CO<sub>2</sub>” (from -NEP) from  
279 “external CO<sub>2</sub>” coming from groundwater inputs of DIC. Indeed, inputs of groundwater  
280 DIC are acknowledged as sustaining a significant fraction of the CO<sub>2</sub> emissions from  
281 lakes (Butman and Raymond 2011; McDonald et al. 2013) and from rivers, especially  
282 headwaters (Johnson et al. 2008; Hotchkiss et al. 2015; Deirmendjian and Abril 2018).  
283 Horizontal transfer of respiration-derived DIC from terrestrial and wetland ecosystems  
284 to aquatic ecosystems explain why aquatic NEE (CO<sub>2</sub> outgassing) greatly exceeds -NEP  
285 (negative NEP, net heterotrophic ecosystems) in rivers (Abril et al. 2014; Hotchkiss et al.  
286 2015; Borges et al. 2015a). Conversely, this outgassing flux from aquatic systems implies  
287 that in terrestrial ecosystems and wetlands that release DIC laterally, NEP exceeds -NEE.  
288



289

290 **4. THE TERRESTRIAL PERSPECTIVE**

291 Hydrological C export as a significant loss term for terrestrial ecosystems has been  
292 considered in more detail only relatively recently (e.g., Ciais et al. 2008) and is included  
293 in only a very limited number of global terrestrial models (Krinner et al. 2005).  
294 Terrestrial C budgets at the plot and the continental scales are based on different  
295 methods not consistent and precise enough to estimate hydrological C export as a  
296 residual flux. In addition, no direct standardized experimental method is available yet to  
297 directly estimate the flux of C across the boundary between land and water, and the E  
298 term in Eq. 1 for terrestrial systems is almost always calculated from a C mass balance in  
299 inland waters (Fig. 1b; Ciais et al. 2013). Terrestrial -NEE calculated as the difference  
300 between land use change and net land C flux is estimated at 2.6 PgC yr<sup>-1</sup> for the years  
301 2000s (Ciais et al. 2013). In a conceptual model that ignores the different functionalities  
302 between floodable and drained land (Fig. 1b), depending on what estimates are used for  
303 the outgassing term (Raymond et al. 2013; Lauerwald et al. 2015) and for the sediment  
304 burial term (Cole et al. 2007; Tranvik et al. 2009), the hydrological export necessary to  
305 balance the inland water C budget is 1.9-3.2 PgC yr<sup>-1</sup>, which corresponds to 75-125% of  
306 the present net atmosphere-land C flux (Fig. 1b). The atmosphere-land net C flux of 2.6  
307 PgC yr<sup>-1</sup> is derived from multiple approaches including atmospheric CO<sub>2</sub> inversion,  
308 terrestrial ecosystem models and forest inventories (Ciais et al. 2013). The atmospheric  
309 CO<sub>2</sub> inversion method integrates large continental areas that include inland waters.  
310 Thus, the global -NEE calculated from continental-scale inversion models accounts for  
311 CO<sub>2</sub> outgassing from inland waters. Intriguingly, the results of inversion methods are  
312 relatively consistent with forest inventories and process-based models that do not  
313 necessarily account for hydrological export (Ciais et al. 2013). However, when a



314 comparison is made at the plot scale with eddy-covariance data, model performance is  
315 generally poor (Schwalm et al. 2010), and for instance modelled GPP can be  
316 overestimated by more than 100% in tropical forests (Stöckli et al., 2008). If a -NEE  
317 from atmospheric inversion is assumed close to NECB from inventories and process-  
318 based models, then the E term (Eq. 1) is expected to be small, within the error of flux  
319 estimates from the terrestrial perspective. If outgassing of CO<sub>2</sub> from freshwater is  
320 already included in -NEE calculated by atmospheric inversion methods, and if this -NEE  
321 value (2.0-3.0 PgC yr<sup>-1</sup>) is very close to that of NECB (1.8-2.3 PgC yr<sup>-1</sup>), then terrestrial  
322 ecosystems cannot export the 0.6-1.0 PgC yr<sup>-1</sup> of recalcitrant OC that is buried in inland  
323 waters (0.2-0.6 PgC yr<sup>-1</sup>) and exported to the ocean (0.4 PgC yr<sup>-1</sup>).

324

325 Spatially, global forest carbon accumulation occurs in boreal and temperate regions,  
326 whereas tropical forests were found to be near neutral, with net emissions from land  
327 use change being compensated by sinks in preserved tropical forests (Pan et al. 2011).  
328 In contrast, Lauerwald et al. (2015) estimated that 78% of global CO<sub>2</sub> outgassing by  
329 rivers occurred at a latitude lower than 25°. Such latitudinal uncoupling between CO<sub>2</sub>  
330 uptake by forests and CO<sub>2</sub> outgassing from rivers and lakes is intriguing and merits an  
331 explanation. Indeed, it would imply that different climatic and/or anthropogenic forces  
332 are driving these continental fluxes, in contradiction with the positive spatial correlation  
333 between river pCO<sub>2</sub>, air temperature and terrestrial NPP at the global scale (Lauerwald  
334 et al. 2015). It should not be forgotten, however, that these correlations could be  
335 indirect. Indeed, field pCO<sub>2</sub> data in the Amazon and in African rivers including the Congo,  
336 reveal a strong positive influence of flooding and the presence of wetlands on water  
337 pCO<sub>2</sub> (Abril et al. 2014; Borges et al. 2015a,b).

338





339 In terrestrial systems, few local studies at the plot scale compare -NEE or NECB  
340 measurements with E derived from groundwater, spring and/or stream sampling. These  
341 studies lead to very different conclusions from those of global modelling studies. In  
342 remnant mature forests of Para, Brazil, Davidson et al. (2010) estimated the export of  
343 dissolved CO<sub>2</sub> from soil and groundwater to streams at a value of 2-3 orders of  
344 magnitude lower than the soil respiration and NPP. In temperate climate, Kindler et al.  
345 (2011) quantified C leaching by combining a soil-water model and dissolved C analysis  
346 in soil water; these authors reported significant E flux in croplands (25% of NECB),  
347 grasslands (22%) but not in forests (less than 3%). In a temperate, forested and well-  
348 drained watershed, Deirmendjian et al. (2018) monitored dissolved C concentrations in  
349 groundwater and streams and estimated a total export E of 2% of -NEE as measured by  
350 eddy-covariance at the same site. These modest export rates from forests in this limited  
351 number of studies appear contradictory with the necessity of a large E term from  
352 terrestrial ecosystems (1.9-3.2 PgC yr<sup>-1</sup> in Fig. 1b) to fuel inland waters at the global  
353 scale (Cole et al. 2007; Ciais et al. 2013).

354

355 From an ecological point of view, a modest hydrological C export from well-drained  
356 lands is also supported by the nature of their NEP components and more specifically by  
357 the allocation of GPP and ER between air and water (Fig. 2,3). In terrestrial systems, GPP  
358 assimilates atmospheric CO<sub>2</sub>, and AR releases CO<sub>2</sub> partly in air (ARa), as foliar  
359 respiration, woody tissue respiration, and partly in soil (ARs), as root respiration. HR  
360 occurs almost entirely in soils (HRs). In forests, belowground respiration generally  
361 accounts for 30-80% of ER, and aboveground respiration accounts for the remaining  
362 fraction of ER (Davidson et al. 2006). Belowground respiration in soils (ARs and HR)  
363 produces CO<sub>2</sub> mainly in superficial well-drained soils, where root density is highest and



364 which are enriched in biodegradable organic matter by litter fall and root exudation  
365 (Ryan and Law 2005). When the land is well-drained, this CO<sub>2</sub> is released in the  
366 unsaturated zone of the soil and mostly returns to the atmosphere across the soil-air  
367 interface. In a tallgrass prairie, downward transfer of soil CO<sub>2</sub> to groundwater was only  
368 approximately 1% of the soil-air CO<sub>2</sub> efflux (Tsy-pin and Macpherson 2012). For this  
369 reason, CO<sub>2</sub> efflux from soils as measured with static chambers (Fig. 3) is commonly  
370 used as an integrative measure of soil respiration (Ryan and Law 2005; Davidson et al.  
371 2006) and until now, by considering the loss of CO<sub>2</sub> that dissolves in groundwater as  
372 negligible or within the error of estimation of metabolic flux at the ecosystem scale.  
373

374 The transfer of C from well-drained terrestrial ecosystems to aquatic systems (Fig. 3)  
375 occurs through mechanical erosion of superficial soil by runoff that mobilizes POC  
376 including young litter, more refractory mineral-bound OC, as well as dissolved humic  
377 OC, and percolation of rainwater through soils that dissolves gaseous CO<sub>2</sub> and soil OC  
378 and liberates DIC and DOC in groundwater, which is further drained to streams and  
379 rivers. The fraction of HR that occurs in groundwater is probably modest in well-drained  
380 ecosystems, as the deepest water-saturated soil horizons contain much less  
381 biodegradable organic matter than the superficial soil (Ryan and Law 2005;  
382 Deirmendjian et al. 2018). A modest export rate from forests is thus consistent with the  
383 allocation of forest metabolism (in particular ER) mainly above the water table (Fig. 2a),  
384 and with only few percent of -NEE ultimately reaching the aquatic system in non-  
385 flooding conditions (Fig. 3).

386

387



388 **5. THE WETLAND PERSPECTIVE**

389 Even though wetlands cover an area of only approximately 10% of land surface  
390 (Downing 2009), they act as hotspots of productivity and CH<sub>4</sub> emissions (Saunio et al.  
391 2016). In addition, wetlands have strong hydrological connections with streams, rivers  
392 and lakes. Ecologists formulated the hypothesis of wetlands as efficient C-exporters long  
393 ago. Mulholland and Kuenzler (1979) reported several-fold higher DOC export from  
394 swamps than from the surrounding landscape in North Carolina (US). Junk (1985)  
395 described floodplain wetlands as a source of POC for the Amazon River; Wetzel (1992)  
396 named littoral wetlands of lakes has “metabolic gates” for nutrients and organic C  
397 between terrestrial and aquatic ecosystems. More recently, using a landscape ecological  
398 approach, Jenerette and Lal (2005) commented on the determinant influence of  
399 hydrology on wetland C fluxes, including downstream export to open waters.  
400 Consequently, hydrological variation (the second dimension of the conceptual 2D-  
401 Model) was identified as a factor of large uncertainty in wetland C cycling (Jenerette and  
402 Lal 2005). Indeed, current available quantitative information on the C export flux (Eq. 6)  
403 is particularly scarce. In wetlands, the quantification of metabolic C fluxes, and the  
404 understanding of biogeochemical processes regulating -NEE, NEP, ER, and NECB have a  
405 high degree of uncertainty. The partitioning of wetland community metabolism between  
406 air, water and sediment, and the complex biological and physical processes that transfer  
407 C in gaseous, dissolved, and particulate forms between these three sub-compartments  
408 are only partially understood (e.g., Hamilton et al. 1995); they are also highly variable in  
409 time and space, and difficult to measure in practice.

410

411 The few estimates of wetland C fluxes at the global scale strongly vary depending first on  
412 the surface area considered for upscaling (Fig. 1c). Lenher and Döll (2004) calculated a



413 wetland surface area of 9-11  $10^6$  km<sup>2</sup>, Mitsch et al. (2013) have used a value of 7  $10^6$   
414 km<sup>2</sup>, and Downing (2009) re-evaluated the total wetland area including smaller systems  
415 to 13-16  $10^6$  km<sup>2</sup>. Based on remote sensing data, Papa et al. (2010) provide a mean total  
416 surface area of 3.4  $10^6$  km<sup>2</sup>, with 56% located in the tropics, in agreement with previous  
417 estimates by Pringent et al. (2001; 2007). More recently, Lu et al. (2016) use a larger but  
418 probably unrealistic value of 33  $10^6$  km<sup>2</sup>. Global wetland C fluxes consist in three major  
419 terms in Eq. 1: (1) -NEE obtained from eddy-covariance measurements was up-scaled to  
420 a value of 3.2 PgC yr<sup>-1</sup> (Lu et al. 2016), an estimate that needs to be corrected to 1.3 PgC  
421 yr<sup>-1</sup> when applying the surface area re-evaluated by Downing (2009); in addition, the  
422 arithmetic mean of available eddy covariance data (Lu et al. 2016) is probably not the  
423 most appropriate way to upscale -NEE at the global scale, and a more precise typology of  
424 wetland -NEE is necessary, based for instance on the classification of Lehner and Döll  
425 (2004). (2) NECB is assumed as equal to organic C sequestration in soils and estimated  
426 from <sup>210</sup>Pb and <sup>137</sup>Cs core dating (Mitsch et al. 2013), a method that ignores slow decay  
427 in the soil C pool and can result in unrealistically high soil C sequestration rates  
428 (Bridgham et al 2014); Indeed, Mitsch et al. (2013) proposed a global C sequestration  
429 value of 0.8 PgC yr<sup>-1</sup>, whereas Bridgham et al. (2014) re-evaluated this value to less than  
430 0.1 PgC yr<sup>-1</sup>. (3) The  $F_{\text{other}}$  term for wetlands is mainly composed of CH<sub>4</sub> emissions and  
431 estimated from bottom-up approaches using static chambers and process-based models  
432 (Mitsch et al. 2013; Saunois et al. 2016), and top-down inversion models based on  
433 atmospheric data (Saunois et al. 2016). Recent published estimates for the global  
434 wetland CH<sub>4</sub> flux range between 0.2 PgC yr<sup>-1</sup> (Saunois et al. 2016) and 0.6 PgC yr<sup>-1</sup>  
435 (Mitsch et al. 2013). Wetland C sources and sinks are thus subject to large uncertainties  
436 but still support the possibility of a residual C flux able to contribute significantly to  
437 river C budgets at the global scale (Fig. 1c.).



438

439 Eddy covariance reveals strong negative NEE (CO<sub>2</sub> sink) in most wetlands (Morison et al.  
440 2000; Jones and Humphries 2002; Saunders et al. 2007; Lu et al. 2016). However, if  
441 wetland E is ignored but significant, GPP, NPP, NEP, and NECB deduced from the diurnal  
442 changes of eddy CO<sub>2</sub> fluxes (Lu et al. 2016) would be overestimated and, inversely, ER  
443 would be underestimated (Eqs.1-6). This point is particularly crucial because in  
444 wetlands the aerial compartment contains most of the photosynthetic parts of the  
445 ecosystem (GPP, NPP) fixing CO<sub>2</sub> directly from the atmosphere, whereas the aquatic  
446 compartment contains the respiratory parts of the ecosystem (ER, HR and a large  
447 fraction of AR) releasing CO<sub>2</sub> to waters but only part of it back to the atmosphere  
448 because of gas-exchange limitation at the water-air interface (Fig. 3). Wetland 1D mass-  
449 balance budgets also include an estimation of NPP, based on biomass inventories  
450 (Mitsch et al. 2013; Sjögersten et al. 2014). One problem with NPP data is not accounting  
451 for all the C transferred by the plant from the atmosphere to soil and waters. As the sum  
452 of NEP and HR (Eq. 5), NPP does not include the fraction of GPP that is recycled by AR,  
453 and most importantly, the root respiration in sediment and water, which is highly  
454 significant below floating plant meadows (Bedford et al. 1991; Hamilton et al. 1995) and  
455 in flooded forest (Piedade et al. 2010). Total AR should be divided into three  
456 components according to:

$$457 \quad \text{AR} = \text{AR}_a + \text{AR}_w + \text{AR}_s \quad (\text{Eq. 7})$$

458 where AR<sub>a</sub>, AR<sub>w</sub> and AR<sub>s</sub> are the fraction of AR occurring in air, water and soils,  
459 respectively (Fig. 3). In wetlands, a canopy of vegetation protects the water-air interface  
460 from wind stress and the gas transfer velocity is lower compared to surrounding open  
461 waters (Foster-Martinez and Variano 2016; Ho et al. 2018). Consequently, only a limited  
462 fraction of AR<sub>w</sub> and AR<sub>s</sub> will contribute to the CO<sub>2</sub> fluxes measured with static chambers



463 in wetlands. This is a second reason why wetland mass balances are incomplete and  
464 may artificially shift wetlands to atmospheric C sources or sinks (Sjögersten et al. 2014).  
465  
466 The allocation of C stocks and metabolism above and below water is fundamentally  
467 different in wetlands compared to well-drained land, and this considerably modifies  
468 their ecological functionalities (Fig. 2 and 3). Although some wetland plants also use DIC  
469 from water for photosynthesis, a large majority of wetland GPP is made by the emerged  
470 part of plants that fix atmospheric CO<sub>2</sub> during the emersion periods, and/or during the  
471 flooding because of their emerged or floating canopies (Piedade et al. 1994; Parolin et al.  
472 2001; Engle et al. 2008). A large fraction (excluding wood) of the wetland biomass  
473 produced annually is transferred directly to water and sediment as litter fall and fine  
474 root production, where it fuels HR, including methanogenesis. Albeit important for CH<sub>4</sub>  
475 oxidation (Segarra et al. 2015), this leads to a  $F_{\text{other}}$  (Eq. 1) as CH<sub>4</sub> fluxes more  
476 significantly in wetlands than in well-drained terrestrial ecosystems (Ciais et al. 2013;  
477 Saunio et al. 2016). In addition, because of anaerobic conditions in their soils, water-  
478 tolerant plants can develop morphological aeration strategies (Haase and Rättsch 2010)  
479 that actively transport oxygen to the root zone and enhance respiration and the release  
480 of dissolved CO<sub>2</sub>, CH<sub>4</sub> and other fermentative organic compounds such as ethanol to  
481 waters and pore waters (Bedford et al. 1991; Hamilton et al. 1995; Piedade et al. 2010).  
482 Plants also transport CH<sub>4</sub> directly from sediments to the atmosphere (Byrnes et al.  
483 1995). This is why wetland water below plant canopies is generally hypoxic and highly  
484 supersaturated in CO<sub>2</sub> (Bedford et al. 1991; Abril et al. 2014) and CH<sub>4</sub> (Hamilton et al.  
485 1995; Borges et al. 2015b). Because the water-air interface behaves as a strong physical  
486 barrier for gas diffusion, depending on hydrological features, dissolved CO<sub>2</sub> from



487 swamps, marshes and floodplains waters can be transported downriver for long  
488 distance before being emitted to the atmosphere (Abril et al. 2014; Borges et al. 2015b).

489

490 The process that captures atmospheric CO<sub>2</sub> and exports organic and inorganic C to  
491 rivers and lakes can be called the *wetland CO<sub>2</sub> pump*. This biological pump is also  
492 consistent with chamber measurements that generally identify CO<sub>2</sub> sinks in vegetated  
493 flooded areas and CO<sub>2</sub> sources in adjacent open waters (Pierobon et al. 2011; Ribaud et  
494 al. 2012; Peixoto et al. 2016). It is worth noting that little is known on how wetland -NEE  
495 is affected by hydrology. For instance, a swamp of papyrus (*Cyperus papyrus*) on a  
496 sheltered shore of Lake Naivasha, Kenya, was a CO<sub>2</sub> sink during immersion but a CO<sub>2</sub>  
497 source during emersion, when large amounts of plant detritus accumulated in soils were  
498 exposed to air (Jones and Humphries 2002). In contrast, in the more hydrologically  
499 dynamic Amazon floodplain, Brazil, a stand of *Echinochloa polystachya*, another C<sub>4</sub> plant,  
500 was a CO<sub>2</sub> sink during both immersion and emersion (Morison et al. 2000). This suggests  
501 that a more efficient hydrological export of C in Amazon floodplains compared to Lake  
502 Naivasha could have promoted an annual negative NEE (Eq. 1). Such competition  
503 between C export and burial is also consistent with the more efficient C sequestration in  
504 low flow-through wetlands (Mitsch et al. 2014).

505

506 Concerning wetland metabolic C balance, the fraction of OC produced by NPP that is not  
507 respired *in situ* or buried in the wetland soil is exported to rivers systems as OC (Fig. 3),  
508 according to:

$$509 \quad \text{NPP} = \text{B} + \text{HR} + \text{E}_{\text{POC}} + \text{E}_{\text{DOC}} \quad (\text{Eq. 8})$$

$$510 \quad \text{NEP} = \text{B} + \text{E}_{\text{POC}} + \text{E}_{\text{DOC}} \quad (\text{Eq. 9})$$



511 where B is the OC burial in the wetland soil. Thus, the export of POC and DOC from  
512 wetlands is expressed as:

$$513 \quad E_{\text{POC}} + E_{\text{DOC}} = \text{NEP} - B = \text{NPP} - \text{HR} - B \quad (\text{Eq. 10})$$

514 Downstream, this organic material will undergo intense degradation in inland water  
515 (negative NEP), contributing to CO<sub>2</sub> outgassing through the OC detrital pathway (Cole  
516 and Caraco 2001; Battin et al. 2008).

517 Plants and microbes respiring in water, sediments, and the root zone (AR<sub>w</sub> and AR<sub>s</sub> and  
518 HR) release dissolved CO<sub>2</sub> in wetland water. AR<sub>a</sub> is the only component of ER not  
519 contributing to E<sub>CO<sub>2</sub></sub>. The fraction  $\alpha$  of wetland ER occurring in water and sediment  
520 (AR<sub>w</sub> and AR<sub>s</sub>) and almost all of the microbial HR (Eq. 11), release dissolved CO<sub>2</sub> (and  
521 CH<sub>4</sub>) to waters:

$$522 \quad \alpha \text{ER} = \text{AR}_w + \text{AR}_s + \text{HR} \quad \text{with } (0 < \alpha < 1) \quad (\text{Eq. 11})$$

523 part of these dissolved gases are emitted to the atmosphere, and another part is  
524 exported by the water flow:

$$525 \quad \alpha \text{ER} = \text{FCO}_2 + \text{FCH}_4 + \text{E}_{\text{CO}_2} + \text{E}_{\text{CH}_4} \quad (\text{Eq. 12})$$

$$526 \quad \text{with } \text{E}_{\text{CO}_2} = \alpha \beta \text{ER} \text{ and } \text{FCO}_2 = \alpha(1 - \beta) \text{ER} \text{ and } (0 < \beta < 1) \quad (\text{Eq. 13})$$

527 For simplification, we do not include E<sub>CH<sub>4</sub></sub> in this last equation because this term is  
528 assumed to be modest (few %) compared to E<sub>CO<sub>2</sub></sub>. Indeed, the  $\beta$  term might be much  
529 smaller for CH<sub>4</sub> than for CO<sub>2</sub> due to preferential CH<sub>4</sub> ebullition and transport through  
530 plants in wetlands (Chanton and Whiting 1995). For CO<sub>2</sub>, the fraction  $\beta$  depends on  
531 hydrological and geomorphological parameters such as water depth, velocity and gas  
532 exchange in the wetland. Using a simple model of lateral dissolved gas transport (Abril  
533 et al. 2014), typical values of 1 cm s<sup>-1</sup> for the gas transfer velocity (Foster-Martinez and  
534 Variano 2016; Ho et al. 2018) and 5000 ppmv for water pCO<sub>2</sub>, we calculated a  $\beta$  value of





535 0.93 for a water column of 1 m-depth flowing at a velocity of 10 cm s<sup>-1</sup> in a 100 m-long  
536 wetland. When the water depth is established at 0.1 m instead of 1 m or the water  
537 velocity is established at 1 cm s<sup>-1</sup> instead of 10 cm s<sup>-1</sup>,  $\beta$  decreases to 0.53. Consequently,  
538 a large majority of the CO<sub>2</sub> produced by wetland below-water respiration is outgassed to  
539 the atmosphere outside of the wetland. Finally, accounting for all terms in Eq. 6 in  
540 wetlands leads to total export expressed as:

$$541 \quad E = (E_{\text{DOC}} + E_{\text{POC}}) + (E_{\text{CO}_2} + E_{\text{CH}_4}) = (\text{NPP} - \text{HR} - \text{B}) + (\beta \alpha \text{ER} - \text{FCO}_2 - \text{FCH}_4) \quad (\text{Eq. 14})$$

$$542 \quad E = (E_{\text{DOC}} + E_{\text{POC}}) + (E_{\text{CO}_2} + E_{\text{CH}_4}) = (\text{NPP} - \text{HR} - \text{B}) + (\beta (\text{AR}_w + \text{AR}_s + \text{HR}) - \text{FCO}_2 - \text{FCH}_4) \quad (\text{Eq. 14})$$

$$543 \quad E = \text{NPP} - \text{B} + \beta \text{AR}_w + \beta \text{AR}_s + (\beta - 1) \text{HR} - \text{FCO}_2 - \text{FCH}_4 \quad (\text{Eq. 15})$$

544 The correct 2D wetland mass balance budget in wetlands is also calculated as:

$$545 \quad \text{NPP} + \beta \text{AR}_w + \beta \text{AR}_s - (1 - \beta) \text{HR} = \text{B} + \text{FCO}_2 + \text{FCH}_4 + E \quad (\text{Eq. 16}).$$

546 The three terms AR<sub>w</sub> and AR<sub>s</sub> and HR together with the E term, are generally neglected  
547 in wetland C budgets (Mitsch et al. 2013; Sjögersten et al. 2014).

548

549 WHAT TOOLS DO PLUMBERS NEED?

550 Quantifying hydrological C export from wetlands at the community, ecosystem, regional,  
551 and global scales would require information that to date is still missing or incomplete.

552 General recommendations include more systematic field observations of C fluxes across  
553 the boundaries of wetlands with the atmosphere, the upland and the river. Eddy

554 covariance data is still lacking in some remote wetlands where logistics are complicated  
555 (Lu et al. 2016), for example in floodplains of large tropical rivers, which host highly

556 productive flooded forests and floating macrophytes (Piedade et al 1994; Morison et al.

557 2000), and largely contribute to riverine global CO<sub>2</sub> and CH<sub>4</sub> emissions (Richey et al.

558 2002; Engle et al. 2008; Bloom et al. 2010; Abril et al. 2014, Borges et al. 2015a). Eddy



559 covariance measurements should also be more systematically coupled at the same site  
560 with chamber measurements, hydrological C fluxes and C sequestration studies but  
561 accounting for the longer time-scale of the sequestration rates based on core dating.

562

563 The quantification in the field of the amount of C that enters or leaves wetland  
564 ecosystems horizontally with water flow is challenging because many wetlands have  
565 complex morphologies and multiple pathways of hydrological transport that should be  
566 apprehended using hydrodynamical modelling. In addition to hydrological complexity,  
567 the C forms may largely change when water crosses the wetland and for instance,  
568 terrestrial mineral-bound POC can be trapped and replaced by wetland POC, DOC and  
569 dissolved CO<sub>2</sub>. Isotopic and molecular tracers can help in differentiating terrestrial from  
570 wetland OC, when the signatures of the two sources are well separated, for instance, in  
571 watersheds dominated by C3 forests, the contribution of wetland C4 macrophytes can  
572 be tracked in riverine POC, DOC and DIC (Quay et al. 1992; Mortillaro et al. 2011; Albéric  
573 et al. 2018). In contrast, OC from flooded forests is more difficult to differentiate from  
574 that coming from *terra firme* forests (Ward et al. 2013) when many tree species are  
575 common to both ecosystems (Junk et al. 2010). Radiocarbon age in rivers can be  
576 interpreted as the time spent by C in soils and, when young C predominates, they  
577 suggest a rapid transfer from plants to waters (Mayorga et al. 2005), as in wetlands  
578 ecosystems. However, some wetlands such as peats can also export old dissolved C to  
579 streams (Billet et al. 2007).

580

581 Original experimental work in mesocosms that simulate flooding, as well as wetland  
582 ecosystem manipulations are necessary to characterize and quantify hydrological C  
583 export per flooded area, as well as the fraction of ecosystem respiration occurring below



584 water; methods must be developed to estimate HR, ARw and ARs (Eq. 11-13). Soil core  
585 incubations or submerged static chambers for instance, provide an estimate of HRs plus  
586 a fraction of ARs in some wetlands with small plants; in the absence of phytoplankton,  
587 dark bottle incubations measure HRw but miss ARw released by the submerged part of  
588 plants. Special mesocosms adapted to the metabolism of semi-aquatic plants are  
589 necessary. Data of metabolic rates are still missing but would be necessary to build  
590 coupled hydrological-biogeochemical models for wetlands and inland waters. Process-  
591 based biogeochemical models are indeed promising approaches for quantifying C  
592 exports from flooded lands (e.g., Sharifi et al. 2013; Lauerwald et al. 2017). Ideally, these  
593 models could simulate the most important biological processes in the wetland: GPP,  
594 NPP, litter fall, and the different components of ER in air, water and soil, together with  
595 hydrological transport and gas emission. Few modelling studies account for DOC export  
596 (Sharifi et al. 2013), most miss the DIC export as dissolved CO<sub>2</sub> and do not correctly  
597 account for the autotrophic respiration terms (ARw and ARs), or the heterotrophic  
598 microbial processes in the root zone (HRs) (Fig. 2). Recently, Lauerwald et al. (2017)  
599 developed a new type of model of C cycling in large rivers that mimics the most  
600 important physical and biological processes, including land flooding; when applied to  
601 the Amazon River, the model calculated a total CO<sub>2</sub> outgassing flux close to that upscaled  
602 from field measurements (Richey et al. 2002); in addition, the computed relative  
603 contributions to the total dissolved C inputs of surface runoff (14%), drainage (28%)  
604 and flooding (57%) were consistent with recent field evidence that wetlands  
605 predominantly fuel CO<sub>2</sub> outgassing from the Amazon River (Abril et al. 2014).  
606  
607 Finally, a precise upscaling of wetland and inland waters global C budgets requires an  
608 adequate typology of C cycles that accounts for the different hydrological and



609 biogeochemical functioning of peats, swamps, marshes and floodplains, and their spatial  
610 distributions along climatic zones (Lehner and Döll 2004). Ideally, process-based  
611 models should be built and validated in individual wetland types, and then aggregated to  
612 a global model able to quantify C fluxes between drained land, floodable land, rivers and  
613 lakes and the atmosphere at the continental scale. Such modelling tools will also be  
614 highly valuable to predict the impacts of climate and land use changes on these  
615 continental C fluxes. Knowing the relative contribution of well-drained land and wetland  
616 to inland water CO<sub>2</sub> emissions is crucial for quantifying the continental greenhouse gas  
617 budget (Fig. 1) and to predict its sensitivity and feedback on climate warming. For  
618 instance, the intensification of floods and droughts or river damming have the potential  
619 to drastically modify C fluxes at the land-water-atmosphere interface and alter or  
620 enhance the hotspot character of wetlands in the continental C cycle. Such evolution  
621 must be monitored in the field, better understood, conceptualized, and modelled in  
622 order to guide environmental conservation strategies in the next decades.

623

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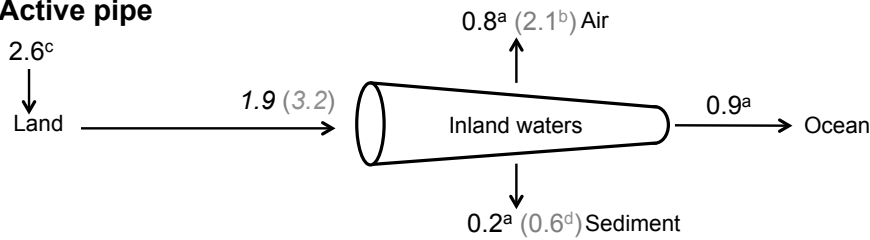
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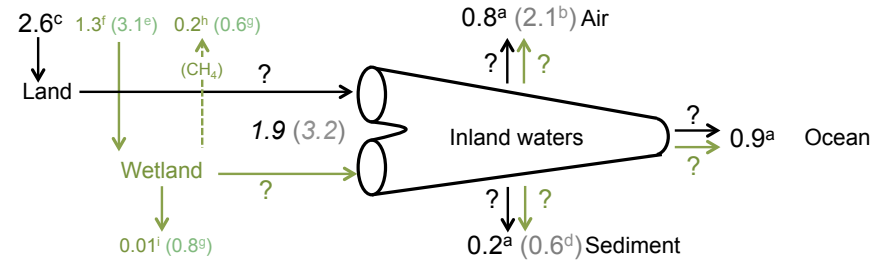
**a. Passive pipe**



**b. Active pipe**



**c. Re-plumbed active pipe**

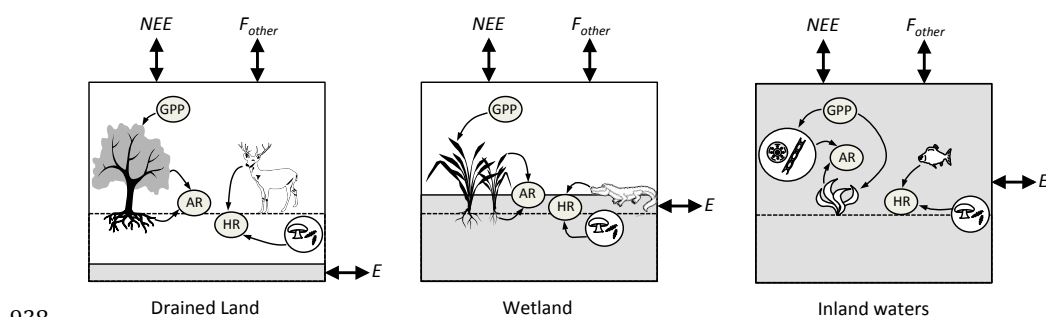


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927 Fig. 1. An update of the active pipe concept, including wetlands in the C budget of inland  
 928 waters. <sup>a</sup> from Cole et al. (2007); <sup>b</sup> from Raymond et al. (2013) (note that the estimate of  
 929 global CO<sub>2</sub> outgassing from Cole et al. (2007) is similar to that of Lauerwald et al. 2015);  
 930 <sup>c</sup> calculated as the difference between land use change and net land flux in Ciais et al.  
 931 (2013); <sup>d</sup> from Tranvik et al. (2009); <sup>e</sup> from Lu et al. (2016); <sup>f</sup> from Lu et al. (2016)  
 932 corrected for a global wetland surface area of Downing et al. (2009); <sup>g</sup> from Mitsch et al.  
 933 (2013); <sup>h</sup> from Saunois et al. (2016); <sup>i</sup> corrected from Mitsch et al. (2013), according to  
 934 Bridgham et al. (2014). Numbers in italics are calculated as the sum of all others fluxes  
 935 and include a high (grey) and a low (black) estimate. Black arrows represent C  
 936 originating from well-drained, terrestrial ecosystems, and green arrows represent  
 937 wetland C.



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939 Fig. 2. Relationship among the carbon (C) fluxes (in italics) that determine net  
 940 ecosystem carbon balance (NECB) (the net of all C imports to and exports from the  
 941 ecosystem), and the metabolic fluxes (inside grey oval) that determine net ecosystem  
 942 production (NEP). (Adapted from Chapin et al. 2006 to include aquatic compartments).  
 943 The boxes represent the ecosystems (drained land, wetland, inland waters). Fluxes  
 944 contributing to NECB are (i) net ecosystem exchange ( $NEE$ ) with the atmosphere  
 945 (emissions to or uptake from the atmosphere of carbon dioxide,  $CO_2$ ); (ii) fluxes of  
 946 carbon forms other than  $CO_2$  ( $F_{other}$ ), which include methane ( $CH_4$ ), carbon monoxide  
 947 ( $CO$ ), and volatile organic C (VOC); (iii) lateral export ( $E$ ) or import of dissolved organic  
 948 and inorganic C and particulate organic C by hydrological transport and other processes  
 949 such as animal movement, wind deposition and erosion, and anthropogenic transport or  
 950 harvest. In this study, we consider  $F_{other}$  as the flux of  $CH_4$  from the ecosystem to the  
 951 atmosphere, and  $E$  as hydrological export from the ecosystem as POC, DOC, dissolved  
 952  $CO_2$  and dissolved  $CH_4$ . Fluxes contributing to NEP are gross primary production ( $GPP$ )  
 953 and ecosystem respiration (ER). ER includes autotrophic respiration (AR) by the  
 954 different components of vegetation (leaves, wood, roots and photosynthetic microbes)  
 955 and heterotrophic respiration (HR) by prokaryotes, fungi and animals. The shaded  
 956 volume in each box indicates the part of the ecosystem occupied by water.  $GPP$  and ER  
 957 occur mostly above the water table in well-drained ecosystems, partly above and below

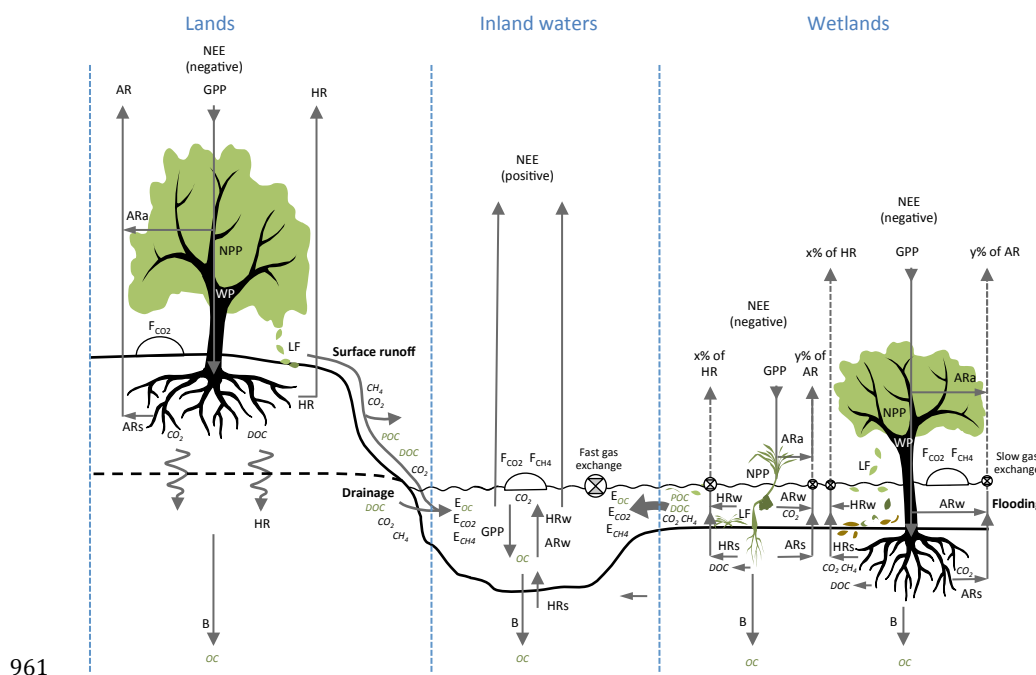


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958 the water table in wetland ecosystems, and exclusively in water and sediments in

959 aquatic ecosystems.

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962 Fig. 3 Functional differences of carbon metabolism and hydrological export in well-  
 963 drained and flooded land. NEE: net ecosystem exchange; GPP: gross primary production;  
 964 NPP: net primary production; WP: wood production; LF: litter fall; AR: autotrophic  
 965 respiration; ARa: autotrophic respiration in air; ARw: autotrophic respiration in water;  
 966 ARs: autotrophic respiration in soils and sediments; HR: heterotrophic respiration; HRw  
 967 heterotrophic respiration in water; HRs heterotrophic respiration in sediments; B: long-  
 968 term burial in soils and sediments. POC: particulate organic C; DOC: dissolved organic C;  
 969  $E_{OC}$ : export of organic carbon (sum of DOC and POC);  $E_{CO_2}$ : export of dissolved  $CO_2$ ;  $E_{CH_4}$ :  
 970 export of dissolved  $CH_4$ ;  $F_{CO_2}$  and  $F_{CH_4}$ : fluxes of  $CO_2$  and  $CH_4$  at the soil-air or water-air  
 971 interface (as determined with static chambers). Note that, by convention, NEE is  
 972 opposite in sign to GPP and NPP because NEE is defined by atmospheric scientists as a C  
 973 input to the atmosphere, whereas GPP and NPP are defined by ecologists as C inputs to  
 974 ecosystems (Chapin et al. 2006). C export to river systems results from the interactions  
 975 between metabolic processes and C transport processes between air, plants, soils,



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976 sediments and waters and are fairly different in wetlands ecosystems (right) and  
977 terrestrial, never-flooded, ecosystems (left). In terrestrial systems, carbon export occurs  
978 as surface runoff and drainage and includes a small fraction of LF, root exudation, ARs,  
979 and HR. In contrast, in flooded wetlands (right), almost all LF and root exudation (that  
980 releases DOC), as well as a substantial fraction of ecosystem respiration  
981 (ARw+ARs+HRw+HRs) are transferring C to the aquatic system as OC and dissolved  
982 gases; in addition, slow gas exchange (low gas transfer velocity) in protected wetlands  
983 favours lateral export of dissolved CO<sub>2</sub> and CH<sub>4</sub>. These lateral C fluxes are enhanced in  
984 wetlands compared to drained systems and should generate strong discrepancies  
985 between ecosystem metabolic fluxes (GPP, NPP, ER, and NECB) and vertical C fluxes  
986 measured in the field with static chambers (F<sub>CO2</sub> and F<sub>CH4</sub>), and eddy covariance towers  
987 (NEE).  
988  
989