Climate-driven Variations in Nitrogen Retention from a Riverine Submerged Aquatic Vegetation Meadow

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Abstract

Large rivers can retain a substantial amount of nitrogen (N), particularly in submerged aquatic vegetation (SAV) meadows that may act as disproportionate control points for N retention in rivers. However, the temporal variation of N retention remains unknown since past measurements were snapshots in time. Using high frequency measurements over the summers 2012-2017, we investigated how climate variation influenced N retention in a SAV meadow at the confluence zone of two agricultural tributaries entering the St. Lawrence River. Distinctive combinations of water temperature and level were recorded between years, ranging from extreme hot-low (2012) and cold-high (2017) summers (2 @C and 1.4 m interannual range). Using an indicator of SAV biomass, we found that these extreme hot-low and cold-high years had reduced biomass compared to hot summers with intermediate levels. In addition, change in main stem water levels were asynchronous with the tributary discharges that controlled NO3- inputs at the confluence. We estimated daily N uptake rates from a moored NO3- sensor, and partitioned these into assimilatory and dissimilatory pathways. Measured rates were variable but among the highest reported in rivers (median 576 mg N m-2 d-1; range 60 - 3893 mg N m-2 d-1) and SAV biomass promoted greater proportional retention and permanent N loss through denitrification. We estimated that the SAV meadow could retain up to 0.8 kt N per year and 87% of N inputs, but this valuable ecosystem service is contingent on how climate variations modulate both N loads and SAV biomass.

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1	Climate-driven Variations in Nitrogen Retention from a Riverine Submerged
2	Aquatic Vegetation Meadow
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16	Key Points:
17	• Nitrogen retention and biomass were measured at high resolution over six summers in a
18	submerged aquatic vegetation meadow of a large river
19	• Among the highest riverine nitrate uptake rates were recorded and 47-87% of loads were
20	retained with plants favoring denitrification
21	• Interannual climate variation influenced nitrate retention by altering water levels,
22	temperature, plant biomass and tributary nitrate load
23	

24 Abstract

25 Large rivers can retain a substantial amount of nitrogen (N), particularly in submerged 26 aquatic vegetation (SAV) meadows that may act as disproportionate control points for N 27 retention in rivers. However, the temporal variation of N retention remains unknown since past 28 measurements were snapshots in time. Using high frequency measurements over the summers 29 2012-2017, we investigated how climate variation influenced N retention in a SAV meadow at 30 the confluence zone of two agricultural tributaries entering the St. Lawrence River. Distinctive 31 combinations of water temperature and level were recorded between years, ranging from extreme 32 hot-low (2012) and cold-high (2017) summers (2 °C and 1.4 m interannual range). Using an 33 indicator of SAV biomass, we found that these extreme hot-low and cold-high years had reduced 34 biomass compared to hot summers with intermediate levels. In addition, change in main stem 35 water levels were asynchronous with the tributary discharges that controlled NO₃⁻ inputs at the 36 confluence. We estimated daily N uptake rates from a moored NO₃⁻ sensor, and partitioned these 37 into assimilatory and dissimilatory pathways. Measured rates were variable but among the highest reported in rivers (median 576 mg N m⁻² d⁻¹, range 60 - 3893 mg N m⁻² d⁻¹) and SAV 38 39 biomass promoted greater proportional retention and permanent N loss through denitrification. 40 We estimated that the SAV meadow could retain up to 0.8 kt N per year and 87% of N inputs, 41 but this valuable ecosystem service is contingent on how climate variations modulate both N 42 loads and SAV biomass.

43

Plain Language Summary

44 Large rivers remove significant amounts of nitrogen pollution generated by humans in 45 waste waters and from fertilizers applied to agricultural lands. Underwater meadows of aquatic plants remove nitrogen particularly well. To keep the river clean, plants use the nitrogen 46 47 themselves, and promote conditions where bacteria can convert this pollution to a gas typically 48 found in air. Measuring nitrogen removal in rivers is really difficult, and we don't know how 49 climate conditions influences this removal or plant abundance. We successfully measured 50 nitrogen pollution removal from an underwater plant meadow in a large river over six summers. We found that plant abundance and river nitrogen inputs were critical to determine how much 51 52 pollution was removed, and that these were controlled by climatic conditions. Plant abundance 53 was controlled by both water temperatures and levels. When water was warm and levels were

- 54 neither too high nor too low, conditions were perfect for lots of plants to grow who mainly
- 55 stimulated bacteria that removed nitrogen. We showed that the amount of nitrogen pollution
- 56 removed over the summer by the meadow changes with climatic conditions but in general
- 57 represents the amount produced by a city of half a million people.

59 1 Introduction

60 Human activities on land has led to increased delivery of nitrogen (N) to aquatic 61 ecosystems, resulting in the degradation of receiving waters (Carpenter et al., 1998; Galloway et 62 al., 2003). During the transfer from land to sea, these impacts are modulated by river networks 63 that retain a considerable amount of N, either through temporary biotic uptake in the water 64 column or permanent removal by denitrification in anoxic sediments (Hall et al., 2009; 65 Seitzinger et al., 2006). Within hydrographic networks, modelling efforts suggest that large 66 rivers have a substantial influence on basin-wide N retention (Seitzinger et al., 2002; Wollheim 67 et al., 2006; Ye et al., 2017). This influence is explained by their broader reaches that increase 68 water residence time and contact rate with reactive surfaces, combined with higher N loads due 69 to their downstream position. However, N retention in rivers is highly heterogeneous (Piña-70 Ochoa & Álvarez-Cobelas, 2006) and specific locations, like submerged aquatic vegetation 71 (SAV) meadows, can display a disproportionate retention, thus acting as control points 72 (Bernhardt et al., 2017, e.g. Pinardi et al., 2009; Preiner et al., 2020).

73 This disproportionate role of SAV meadows can be explained by a suite of positive 74 feedbacks enhancing plant growth and N uptake. For example, SAV reduce flow velocities and 75 increase water transparency through sediment deposition (Hilt, 2015; Scheffer et al., 1993). The 76 role of SAV in N dynamics might be amplified by their location within rivers. For example, high 77 N retention has been observed in a meadow located in a confluence zone of one river (Hudon & 78 Carignan, 2008), but near undetectable denitrification rates were reported at one located in the 79 main channel of another (Tall et al., 2011). This might be due to the specific geomorphologies, 80 like deltas, at confluence zones that increase water residence time and favors N retention in 81 comparison to the higher velocities in main channels. Greater N retention at confluence zones 82 can also be explained by their typically high productivity caused by the increased nutrient supply 83 from incoming tributaries (Benda et al., 2004; Rice et al., 2008). Nevertheless, information on 84 the effect of SAV on N dynamics in rivers remain scarce, with only a handful of studies 85 addressing the topic in rivers with discharges greater than 100 m³ s⁻¹ (Diamond et al., 2021; 86 Hudon & Carignan, 2008; Hudon et al., 2017; Tall et al., 2011) and a few in rivers with 87 discharge of less than 20 m³ s⁻¹ (Audet et al., 2021; Desmet et al., 2011; Heffernan & Cohen, 2010; Pinardi et al., 2009; Preiner et al., 2020). As a result, we have some understanding, albeit 88 89 fragmented, of N dynamics in SAV meadows across rivers. Yet, past evidences are snapshots

over a day, a season, or a year at most, and do not capture the temporal variability in N retention
imposed by climate.

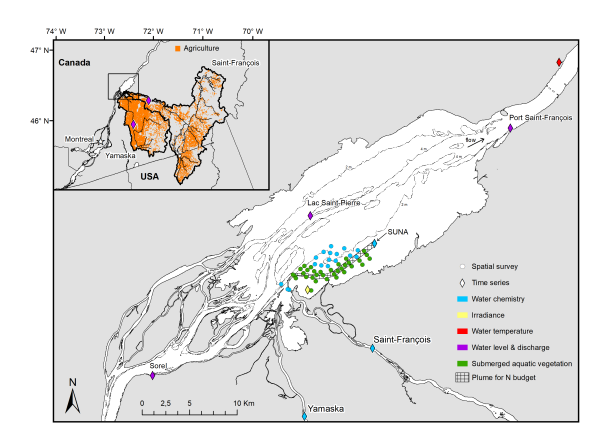
92 Indeed, N retention fluctuates among years as a function of climate variations as shown 93 by previous studies relating net human watershed N input to riverine exports (Ballard et al., 94 2019; Goyette et al., 2019; Howarth et al., 2012). In temperate regions, water flow (levels or 95 discharge) and temperature tend to be negatively correlated because of higher evapotranspiration 96 of land plants with increased temperatures (e.g. Hudon et al., 2010). As a result, colder than 97 average growing seasons should exhibit higher runoff watershed losses, faster within-river flow, 98 and consequently increased N export. In contrast, warmer growing seasons should result in lower 99 discharge and higher N retention through enhanced plant and microbial activity, potentially 100 compounded by lower precipitation and higher water residence times. However, the effect of 101 climate and temperature variation on riverine SAV dynamics and overall N retention remains 102 unknown. Rising temperatures can affect SAV biomass either by directly increasing growth rate 103 (Brown et al., 2004; Farquhar et al., 1980) or by improving light availability through water level 104 decline. During extreme drought, water levels can go below a deleterious threshold after which 105 air exposure can lead to the dry-out of above ground biomass (Ersoy et al., 2020; Hudon et al., 106 2010). Growth rates could also increase to a temperature optimum where SAV thereafter could 107 be outcompeted by periphyton or phytoplankton (Short et al., 2016), which may result in a 108 decrease in N retention through reduced reactive surfaces and higher flow velocities. As SAV 109 meadows in rivers represent a potential control point for N loss, characterizing how climate 110 variations influence this critical ecosystem service is of high importance.

111 Accurate estimates of N loss in large rivers are particularly challenging because methods 112 for measuring SAV biomass and N retention throughout or over multiple growing seasons are 113 restricted and time consuming. Capturing changes in SAV biomass over a growing season 114 requires the processing of many plant samples, or in typically turbid waters, conducting 115 numerous remote sensing surveys using a sonar (Botrel et al., 2022). One potential solution to 116 track temporal SAV biomass is to use the difference in slope in surface water level between 117 gauging stations around large riverine meadows. Variations in water level slope tend to follow 118 plant development because as SAV accrues biomass, flow is obstructed by the biomass in large 119 meadows resulting in an increase surface water elevation upstream (Boudreau et al., 1994). This 120 approach has been used successfully to estimate biomass in a large river (Giacomazzo et al.,

121 2020; Vis et al., 2007). As for N uptake measurements, few reliable methods exist in large rivers. 122 The typical isotopic or nutrient release experiments used to measure N uptake rates in streams 123 are impractical in large rivers because of the enormous quantities of injected N needed to detect 124 and track a downstream signal in a large water mass (e.g. Mulholland et al., 2002; Newbold et 125 al., 1981; Peterson et al., 2001). Consequently, past N uptake estimates in large rivers are very 126 coarse as they were either derived from large scale mass balance (Alexander et al., 2000; 127 Howarth et al., 1996) or from models that upscaled measurements taken in small streams (Hall et 128 al., 2013; Wollheim et al., 2006; Ye et al., 2017). However, the development of in situ high 129 frequency NO₃⁻ sensors provides a unique opportunity to directly measure variation in N uptake rates, particularly in rivers subjected to human derived N inputs where NO3 is typically the 130 131 dominant N form (Caraco & Cole, 1999). Using these sensors, N uptake can be estimated daily 132 by a passive mass balance between two stations along a river reach. Furthermore, when 133 autochthonous primary production is high, NO_3^{-1} concentrations tend to display a diel variation 134 driven by daily change in sunlight and gross primary productivity (GPP) that can be used to 135 estimate autotrophic NO₃⁻ uptake (Hall et al., 2009; Heffernan & Cohen, 2010). As a result, high 136 frequency NO₃⁻ signals allow for the partitioning of total N uptake rates into assimilatory and 137 dissimilatory pathways, providing additional information on N fate.

138 Given the lack of knowledge on temporal N retention dynamics in riverine SAV bed, our 139 objective is to characterize how interannual climate variation affects N retention in a large 140 riverine SAV meadow located at a confluence zone. We hypothesized that retention would be a 141 function of NO_3^{-1} loads and of vegetation biomass, both of which are related to climate-driven 142 variations. To do so, we estimated SAV biomass and environmental conditions over six summers 143 in a natural widening of the St. Lawrence River (SLR), a large temperate river with seasonal ice 144 cover. Using a passive mass balance approach with an in situ sensor, we contrasted interannual 145 variations of N retention within the flow path of the SAV meadow and partitioned retention into 146 autotrophic assimilation and denitrification. N retention was then upscaled to the entire SAV bed 147 using a simple regression model.

148 **2 Materials and Methods**



149

Figure 1. Map of the study area and site locations in Lake Saint-Pierre. The limits of Yamaska and Saint-François river watersheds (thick outline) and farmlands (orange) are delineated and the frame indicates the location of the enlarged area (inset). The dotted lines in the lake are the 2 m and 4 m isobaths. Notice that tributaries gauging stations are only shown in inset.

154 2.1 Study Site

We studied N retention in a large (~10 km²) SAV meadow receiving waters at the confluence of two agriculturally impacted rivers (Goyette et al., 2016), the Saint-François (10.18 x 10³ km², 2012-2017 median summer discharge = 74 m³s⁻¹) and the Yamaska (4.45 x 10³ km², 20 m³s⁻¹). At the same location, these two tributaries join the fluvial Lake Saint-Pierre (LSP), a ~ 400 km² shallow widening (mean depth of 3 m) of the St. Lawrence River (SLR, 8th order, 9640 m³/s), Québec (Canada, Figure 1). Annual loads of total N from the Yamaska (7.03 x 10³ tons yr⁻¹) far exceeded those of the Saint-François River (4.45 x 10³ tons yr⁻¹), in spite of its much

- 163 drainage basin was also shown by its high conductivity and high flashiness after local rain
- 164 events. Conductivity in the tributaries is distinct (median, $med_{Saint-François} = 370 \ \mu S \ cm^{-1}$,
- 165 $med_{Yamaska} = 206 \ \mu S \ cm^{-1}$), allowing for its use as a conservative tracer of water mass.

166 Located at the mouth of their confluence with the SLR, the SAV meadow under study 167 was well positioned to intercept and filter the nutrient-rich waters from both tributaries, as was 168 previously suggested by a nutrient mass balance study in LSP (Hudon & Carignan, 2008). Lake 169 Saint-Pierre supports about half of the remaining SLR wetlands (Hudon et al., 2018; Jean et al., 170 2002), and has been designated as both a Ramsar site and a UNESCO biosphere reserve. 171 Although its natural capital has been recognized, LSP hydrodynamics have been highly modified 172 by human activities. Discharge from the Great Lakes to the SLR is regulated at the Moses-173 Saunders dam (Cornwall, ON, Massena, NY), which has resulted in an overall reduction in water 174 level extremes in LSP of 0.7 m on average between 1912 and 1994 (Hudon, 1997; Hudon et al., 175 2006). The effects of flow regulation have been compounded by the dredging of a 11.3 m-deep 176 navigation channel that focuses water flow from the Great Lakes to the central part of the lake 177 (0.5-1 m s⁻¹, representing 55-88 % of total discharge (Hudon & Carignan, 2008; Morin & Côté, 178 2003, Figure 1). As a result, flow is markedly slower ($< 0.5 \text{ m s}^{-1}$) in the lateral shallow zones 179 particularly during summer, when mixing of incoming tributaries with this central water mass is 180 limited (Vis et al., 2007). Reduction in flow velocity in the lateral zones is also explained by the 181 presence of extensive SAV present up to a maximum depth of 3 m (Hudon, 1997).

182 While the bulk of SLR flow is controlled anthropogenically, water levels nevertheless 183 follow interannual fluctuations as a function of climate variation. Flow regulation has a 184 stabilizing effect on waters originating from the Great Lakes (772 x 10^3 km²), which account for 185 approximately 70 % of the SLR waters entering LSP (Hudon et al., 2017). The climate effect is 186 thus primarily the result of tributary inputs, the largest of which is the Ottawa River (146.3 x 10^3 187 km^2), a North shore tributary accounting for ~20% of water inputs in LSP. The much larger sizes 188 and areas covered by the SLR watershed compared to its individual tributaries further induce a 189 potentially complex influence of climate at their confluence zones. The SLR main stem climate 190 variation is an aggregate of the basin-wide Great Lakes and overall tributaries water inputs, 191 while individual tributaries discharges and NO_3^- concentrations are independent of these large-192 scale variations and represent localized precipitation patterns.

193 2.2 Spatial Survey of SAV Biomass and Water Chemistry

194 Field surveys were carried out during maximum SAV biomass accumulation, around 195 early August from 2012 to 2017, and once during the early growing season in June 2012. We 196 measured SAV biomass at approximately 35 sites. Water temperature and conductivity were 197 measured using a YSI 556 MPS or a YSI 600XL at each site as well as at ~17 additional sites 198 within and around the meadow (Figure 2). SAV was collected every summer using the rake 199 technique, with additional diver-collected quadrats in 2016. For the rake technique, a 0.35 m-200 wide double-headed rake was lowered in the water and dragged toward the boat over a length of 201 approximately 1 m. In 2016, divers harvested all plant material within a 0.30 m by 0.30 m PVC 202 frame placed on the lake bottom. At each site, 3 to 5 replicates were collected, and plant material 203 was rinsed on-boat, stored in plastic bags, and frozen once on shore. Upon return to the 204 laboratory, plant samples were thawed, wrung out manually, sorted by species, separated from 205 filamentous algae, dried to a constant mass at 60°C, and weighed (0.001g). The sum of species drv biomass in g m⁻² was calculated per sample and averaged at each site (3-5 replicated) SAV 206 207 species composition was dominated by Vallisneria americana and Potamogeton richardsonii, 208 but also included Heteranthera dubia, Stuckenia pectinata, Elodea spp, Myriophyllum spp. and 209 Chara spp. Because rake biomass collections are underestimated due to loss of plant material 210 during sampling, rake estimates were converted to quadrat equivalency using a previously 211 determined general relationship (Botrel et al., 2022).

- 212 2.3 Time Series Data
- 213 2.3.1 Temporal SAV Biomass Estimates

214 Daily SAV biomass was estimated between June and November using the slope in 215 surface water level between gauging stations (Figure 1). Slopes were calculated as water surface 216 elevation upstream minus downstream divided by distance between successive stations (23 km, 217 Supporting information, SI, Text S1). As variations in water level slope are also affected by 218 signals other than SAV growth, such as tides and winds (Vis, 2004), we extracted the SAV 219 signal with the Hilbert-Huang transform method using the hht and EMD packages in R (Bowman 220 & Lees, 2013; Huang et al., 1998; Kim & Oh, 2018; Wu & Huang, 2009). We chose this signal 221 extraction method because it is adaptative and intrinsic, and is entirely based on data without

imposing an a priori hypothesis on the shape of the SAV growth curve (Wu et al., 2007). We
analyzed water level slopes only during the growing season (June 21 to September 22), therefore
SAV signal corresponded to the EMD residuals or the seasonal trend. To reduced signals mixing
at different time scales (i.e. mode mixing), we used a noise-assisted version of EMD, ensemble
empirical mode decomposition (EEMD, Bekka & Berrouche, 2013). Confidence levels were
estimated to assess between year differences using the bootstrapping approach of Ezer & Corlett
(2012, SI Text S1).

229 To reconstruct SAV growth using changes in water level slopes as an indicator of daily 230 biomass, we assessed the optimal gauging station representative of SAV biomass. For this 231 validation, we used the Pearson correlation between the slope signal to the overall biomass 232 measured during spatial surveys (SI Text S1 and Figure S1). The slope calculated between the 233 downstream gauging stations (Port St-François) to the middle LSP station yielded a higher 234 correlation (r = 0.98) than the slope between upstream Sorel station and the same middle station 235 (r = 0.33). Using this former slope value and to provide estimates using a common metric, we 236 calculated maximum summer SAV biomass using a simple linear regression between rake 237 estimated biomass and slopes ($r^2 = 0.97$, p < 0.0001). For visualization of SAV growth patterns 238 and statistical analysis, we directly used the slope between the Port St-François and LSP station 239 as an indicator of temporal SAV biomass changes.

240 2.3.2 Nutrient Inputs

241 To estimate daily NO₃⁻, total phosphorus (TP) and turbidity in the tributaries, we used the composite approach from the R package loadflex that includes regression modelling with 242 243 discharge as a predictor and a residual correction (Appling et al., 2015). The best regression 244 models were selected using Akaike information criterion (AIC) from among 9 models fitted for 245 data between 2009 and 2018 in the rloadest package (Johnson & Omland, 2004; Runkel & De 246 Cicco, 2017). Weekly water chemistry data from both provincial and national agencies collected 247 in the Saint-François and Yamaska rivers (Table S1) were used to derive models. To avoid 248 autocorrelation when fitting models, a subset of data at time intervals greater than 7 days was 249 used; the complete dataset was used for the residual correction. Daily discharge data was 250 acquired at upstream stations in Drummondville for Saint-François and Saint-Hyacinthe for 251 Yamaska (Table S1). Missing data (3 days) from Saint-François were filled using data available

from a station 1 km upstream, Hemming Falls (r = 0.93), while missing data from the Yamaska

253 dataset (22 days) were filled using the linear regression between daily discharge at Saint-

Hyacinthe and the sum of discharges at the two main upstream sub-watershed stations (Farnham and Noire, $R^2 = 0.96$).

256 Water chemistry monitoring stations were located near the mouth of the tributaries, but 257 discharge stations were located 47 km and 63.5 km upstream for Saint-François and Yamaska 258 respectively. Discharge was corrected for additional water inflow using the pro-rate technique 259 and was multiplied by the proportional increase in drainage area at the respective water 260 chemistry stations (1.05 and 1.34 for Saint-François and Yamaska, Morse, 1990). We also 261 corrected daily discharge according to transit time estimated with Saint-Venant equations which 262 included information on channel geometry and friction coefficients (P. Fortin ECCC pers. 263 comm.). During summer (from June 21 to September 22), transit time was longer than 24h for 264 4% and 65% of dates for Saint-François and Yamaska respectively. Longer transit times in 265 Yamaska were probably due to its smaller watershed size and the higher distance between the 266 gauging station to the river mouth. To describe total tributary inputs during the SAV growing 267 season, discharges are reported as the sum of both tributaries. Summertime concentrations 268 received by the SAV bed were weighted by the proportion of individual tributary (i) discharge to the total discharge ([NO₃⁻] (mg L⁻¹) = $\sum_{i}^{2} ([NO_{3}^{-}]_{i} \times \frac{Q_{i}}{Q_{total}}))$. 269

270

2.3.3 Downstream NO₃-, Temperature and Light Estimates

271 To estimate NO₃⁻ concentrations downstream of the SAV meadow, a SUNA V1 sensor 272 (Satlantic) using in situ ultraviolet absorption spectroscopy (200-400 nm spectra) was moored 273 from the end of June to end of September in 2012 to 2016 (Figure 2). The sensor, equipped with 274 a battery pack, flow cell, and a Sea-Bird electronic pump, was mounted on an aluminum 275 stepladder which kept the sensor at 75 cm above the sediment. Before and after moorings, 276 calibration was verified using distilled water and known NO₃⁻ concentrations. To limit the 277 interference of large particles on the ultraviolet lamp, a pre-filter with a mesh size of 5 mm was 278 installed at the water intake. Each hour, water was pumped, and four measures were recorded. 279 Only measurements within manufacturer-specified accuracy (+- $0.028 \text{ mg N L}^{-1}$) were deemed 280 acceptable and median values were kept for further analysis. A sensor to measure hourly

dissolved oxygen concentrations (D-Opto) was also deployed in 2016, with punctual

282 measurements over other years.

283 Continuous local water temperature was taken from the SUNA sensor, and validated with D-Opto measurements in 2016 (r = 0.99). Additional water temperature time series were 284 285 acquired at Trois-Rivières station (Table S1) and correlated with daily water temperature 286 measured by the SUNA and mean water temperatures measured during spatial surveys (r = 0.72) 287 and r = 0.75, respectively). Hourly short-wave radiation was acquired from NASA datarod at a 288 location close to the SAV bed (46.1247 N, 72.9018 W, Teng et al., 2016, 289 apps.hydroshare.org/apps/data-rods-explorer). Short-wave radiation flux (300-2000 nm in Wm⁻²) 290 was converted to photosynthetically active radiation (PAR, 400-700 nm) in photon flux density 291 units (mol m⁻² h⁻¹) using a conversion ratio of 2.114 (Britton & Dodd, 1976). To describe SAV 292 growing season, we calculated the accumulated degree days using a reference temperature of 293 7°C, which corresponds to the known sprouting temperature of Vallisneria americana (Lacoul & 294 Freedman, 2006), a common species at our study site. Similarly, cumulative PAR was calculated 295 as the sum of daytime hourly irradiance.

296 $2.3.4 \text{ NO}_3^-$ Budget and Fate

297 We calculated a nitrate (NO₃⁻) budget for a plume spanning from the mouths of the 298 tributaries (input) to the downstream edge of the SAV meadow where the SUNA was located 299 (output, Figure 1). NO₃⁻ was the dominant N form in these tributaries (60 % on average) and is a 300 significant agricultural pollutant. A complete summary of equations used to derive budgets is 301 presented in Table 1. The shape of the plume was drawn from trajectories of drifters deployed in 302 2005, 2006 and 2012 (SI Figure S1). A plume width of 500 m was used for calculations, 303 corresponding to the distance between stations of the spatial survey and was chosen based on 304 conductivity measurements as a conservative tracer of water masses. When choosing the location 305 for the outflow station in 2012, the objective was to measure NO₃⁻ from the Saint-Francois water 306 mass flowing through the SAV bed. Since 2012 was an extremely low water level year, the 307 sensor was placed at the offshore limit of this water mass to ensure water was deep enough to 308 allow measurements throughout the summer. However, the study area is a complex zone at a 309 confluence of the Saint-François and Yamaska rivers. Based on delineation using optical 310 properties from satellite observations (pers. comm. P. Massicotte) and conductivity, we observed

311 that in subsequent higher water level years, the same outflow location was most likely exposed to 312 a mixture of waters from both rivers. Daily upstream NO₃⁻ concentrations were thus estimated 313 from the NO₃⁻ loadflex predictions multiplied by the contribution of Saint-François and Yamaska 314 to the water flowing at the downstream site (Table 1, equation 1). To estimate contributions, we 315 used a two end member mixing model with conductivity as the conservative tracer and applied 316 an analytical solution (Table 1, equation 2). Contributions were estimated once per summer 317 using matching monthly conductivity at the nearest sampling station to the SUNA and 318 conductivity measurements in the tributaries from this study and the provincial monitoring 319 station (Table S1).

320 Daily discharge at the outflow site was computed as the area of the plume cross-sectional 321 transect multiplied by flow velocity (Table 1, equation 3). The cross-sectional area was estimated 322 from the sum of cell depths measured for the outflow transect on a bathymetric raster map 323 (Hudon & Carignan, 2008), corrected for daily water level height, and multiplied by the 324 bathymetric cell width. Daily velocity was estimated from Delft3D simulations, a tridimensional 325 hydrodynamic model established for our study area (Bulat et al., 2019). Model simulations were 326 run for each year using median, minimum, and maximum input discharges at Saint-François over 327 the SUNA mooring period, and yearly spatial polygons of SAV height established from SAV 328 echosounding surveys (Botrel et al., 2022). Mean depth average velocity was computed on the 329 resulting spatial polygon grid of 70 m resolution at the outflow transect (7 polygons). Mean 330 residence time was also estimated from depth-averaged velocity assuming a travelling distance 331 of 11.61 km, but the mean was computed over the complete plume surface area. To estimate both 332 daily velocity and daily residence time, interpolations were computed using yearly linear 333 regressions between daily Saint-François discharge and the mean depth average velocities or the 334 mean residence time (i.e. Delft3D inputs and outputs).

Daily total areal uptake (Ut) was considered to be the sum of daily autotrophic assimilation (Ua), denitrification (Ud), and heterotrophic assimilation (Uh, Table 1, equation 4). Ut was calculated as the difference in daily median outflow NO_3^- concentration between upstream input and output (downstream) while taking residence time into account, multiplied by the ratio of outflow discharge divided by the plume area (Table 1, equation 5). Daily proportional uptake (R) was calculated as the median NO_3^- concentration at the outflow minus upstream inputs, divided by those inputs (Table 1, equation 6). Ua was calculated from diel NO_3^-

342 variation using equation 7, Table 1, and U_d as the difference between U_t and U_a . Since hourly 343 NO_3^{-1} data at the downstream station was a composite of multiple frequency signals, we extracted 344 diel variations using Hilbert-Huang transform, similar to the SAV time series and to the protocol 345 of Chamberlin et al. (2021). This method is best suited for nonstationary data, a common feature 346 of hydrological time series (Lloyd et al., 2014). Days for which more than five consecutive 347 hourly measurements were missing were excluded from the data series (5 days in 2012 and 2013, 348 1 day in 2014), otherwise missing values were linearly interpolated. For our five NO₃- time 349 series (2012-2016), IMF4 always corresponded to the diel signal with mean instantaneous 350 periods between 21 and 23 hours. When a NO₃⁻ spike introduced oscillations on the diel IMFs 351 (Stallone et al., 2020), those sections were removed from the analysis. We then filtered the 352 resulting U_a to obtain time intervals between daily maximums of 20 to 28 hours. 353 U_a estimated from NO₃⁻ time series were validated to autotrophic assimilation estimates 354 based on gross primary production (U_{a-GPP}) calculated from the oxygen time series in 2016 355 (equation 8). For this U_{a-GPP} calculation, we assumed a photosynthetic coefficient of 1, an 356 autotrophic respiratory coefficient of 0.5, and used a mean C : N ratio measured for both SAV 357 and macroalgae in LSP (molar ratio of 13:1). Daily gross primary production (GPP) and 358 ecosystem respiration (ER) were calculated using the bookkeeping method in the R package 359 LakeMetabolizer (Winslow et al. 2016). For gas exchange estimation, the depth of the surface 360 mixed layer was assumed to be the depth at the outflow site, and oxygen saturation was 361 calculated using in situ water temperature, and atmospheric pressure estimates at the Nicolet 362 weather station. The coefficient of gas exchange was computed using wind speed (Cole and 363 Caraco (1998) estimates from LSP weather station or, when data were missing, from Nicolet 364 (Table S1). Nighttime and daytime periods were estimated using the R package suncalc 365 (Thieurmel & Elmarhraoui, 2019). Validation of the estimates showed that U_a and U_{a-GPP} were 366 comparable, as most values fell along the 1:1 line and overall means were similar (SI Figure S3). 367 Ecosystemic respiration estimates also allowed for calculation of U_h for 2016, assuming a 368 heterotrophic growth efficiency of 0.2 and molar C : N ratio of 20 : 1 (Table 1, equation 9, Hall 369 & Tank, 2003; Heffernan & Cohen, 2010). As U_h represented less than 0.05% of U_t, this term 370 was considered negligible and was not considered in denitrification estimates. Finally, to isolate 371 the effect of biotic activity on retention (Wollheim et al., 2006; Stream Solute Workshop, 1990),

372 we calculated the uptake velocity (V_f), or the mass transfer coefficient from water to the benthos

373 (Table 1, equation 10). We calculated both total (V_{f-t}) and process-specific velocity (for 374 autotrophic assimilation V_{f-a} and denitrification V_{f-d}).

375 2.4 Statistical Analysis

376 We used linear discriminant analysis (LDA) to assess the occurrence of among-year 377 differences in environmental variables and N budget during summertime, and to identify which 378 variables generated those differences. Given the smaller number of observations for process-379 specific rates (*n* total uptake = 292, *n* process-specific = 99), interannual differences were 380 assessed using nonparametric Kruskall-Wallis tests. A principal component analysis (PCA) was 381 used to describe correlations between NO_3^- budget terms and environmental variables. We then 382 determined predictors of Ut using a regression approach. As our data were nested per year and 383 displayed temporal autocorrelation, we conducted regressions as linear mixed models. To find 384 optimal model structures, we applied the protocol of Zuur et al. (2009). We first removed 385 collinear variables and kept the variables that were most correlated to our response variable, U_t . 386 We then selected random effects using a fixed structure of all possible covariates with restricted 387 maximum likelihood estimation (REML). The best model was selected to minimize the sample-388 corrected Akaike Information Criterion (AICc). Using this optimal random structure, fixed 389 effects were similarly selected using AICc but with maximum likelihood estimation (ML). The 390 final model was fitted using REML and validated for normality and homogeneity by visual 391 inspections of standardized residuals against fitted values and explanatory variables. Using the 392 fixed component of the mixed model, daily retention was predicted from the weighted discharge 393 estimates of NO₃⁻ concentrations inputs as the explanatory variable. Total summertime retention 394 was estimated by multiplying these predictions with the SAV meadow area (10 km²) and 395 summing these daily values. Similarly, we estimated total NO_3^- inputs by summing daily loads 396 from the two tributaries. All statistical analyses were conducted in R (RCoreTeam, 2020) using 397 the MASS package for LDA, vegan for PCA and nlme for linear models (Oksanen et al., 2020; 398 Pinheiro et al., 2020; Venables & Ripley, 2002). Prior to analysis, data were transformed when 399 necessary to satisfy normality and homoscedasticity assumptions.

- 401 **Table 1.** List of Equations Used for Nitrate Budget Calculations. No. equation number. In the
- 402 'Abbreviated description' column, abbreviations that are used in subsequent equations are only
- 403 described once in sequential order.

No	Equations	Abbreviated description Nitrate input ([NO3-] _{in}): StF Saint-François River, Yam, Yamaska River, f fraction where $f_{Stf} + f_{Yam} = 1$		
1	$[NO_{3}^{-}]_{in} (mg L^{-1}) = [NO_{3}^{-}]_{StF} \times f_{StF} + [NO_{3}^{-}]_{Yam} \times f_{Yam}$			
2 <i>ª</i>	$\mathbf{f}_{\mathbf{StF}} = \frac{(C_{\text{out}} - C_{\text{Yam}})}{(C_{\text{StF}} - C_{\text{Yam}})}$	Fraction of Saint-François River water mass (fstf) : out outflow, C conductivity		
3	$\mathbf{Q_{out}} (m^3 s^{-1}) = (\sum_{i=1}^{n} h_{out,i} \times w_i) \times u_{out}$	Discharge at outflow (Q_{out}) : h depth, i ith bathymetric raster cell, w width, u depth average velocity		
4	$\mathbf{U}_{t} (mgN m^{-2}d^{-1}) = U_{a} + U_{d} + U_{h}$	Total nitrate uptake rate (Ut) : Ua autotrophic assimilation, Ud denitrificatic Uh heterotrophic assimilation		
5	$\mathbf{U}_{t} (\text{mgN m}^{-2} \text{d}^{-1}) = \frac{Q_{\text{out,d}}}{A} ([\text{NO}_{3}^{-}]_{\text{in}-\tau} - [\text{NO}_{3}^{-}]_{\text{out,med}})$	Total nitrate uptake rate (Ut) : A area of the plume, τ residence time in days, med median		
6	$\mathbf{R} \text{ (unitless)} = 1 - \frac{[\text{NO}_3^-]_{\text{out,med}}}{[\text{NO}_3^-]_{\text{in}-\tau}}$	Proportional retention (R)		
7 <i>b</i>	$\mathbf{U}_{\mathbf{a}} (\text{mgN m}^{-2} \text{d}^{-1}) = \frac{Q_{\text{out,d}}}{A} \sum_{t=hmax0}^{hmax1} ([\text{NO}_3^-]_{max (hmax0)} - [\text{NO}_3^-]_h)$	Autotrophic assimilation (Ua) : t time of day, hmax0 hour of first daily NO ₃ - maximum, hmax1 hour of last daily NO ₃ - maximum, h hour		
8 <i>b,c</i>	$\mathbf{U}_{\mathbf{a}-\mathbf{GPP}} (\mathrm{mgN} \ \mathrm{m}^{-2}\mathrm{d}^{-1}) = \frac{(\mathrm{GPP} \times \mathrm{h} \times \mathrm{p} \times \mathrm{r}_{\mathrm{a}})}{(\mathrm{C} : \mathrm{N})}$	Autrotrophic assimilation from GPP (Ua- GPP) : GPP gross primary productivity, p photosynthetic coefficient, ra autotrophic respiratory coefficient, C : N carbon to nitrogen ratio		
9 <i>b,c</i>	$\mathbf{U_h} (\text{mgN m}^{-2}\text{d}^{-1}) = \frac{((\text{ER} - (r_a \times \text{GPP})) \times \text{HGE})}{(\text{C} : \text{N})}$	Hetetrophic assimilation (Uh): ER ecosystem respiration, HGE heterotrophi growth efficiency		
10 ^d	$V_{f} (m d^{-1}) = \frac{U}{[NO_{3}^{-}]_{out,med}}$	Uptake velocity (V _f): U uptake rate		

406

404

407 **3 Results**

408

3.1 Interannual Variation in Environmental Variables and Temporal SAV Biomass

409 There were strong and significant interannual variations in environmental conditions 410 during the SAV growing seasons (MANOVA, p < 0.0001) caused by several variables that 411 varied differently among summers. Variables that contributed to among-year differences were 412 mainly water level, water temperature, tributary discharge, and NO₃⁻ input concentrations (LDA, 413 SI Table S2). These conditions created a sharp gradient among summers, ranging from hot 414 temperatures with low water levels to cold summers with high water levels. Mean summer water levels and temperatures spanned 1.4 m and 2°C, respectively coinciding with historical extremes 415 416 in 2012 and 2017 (Figure 2). Exceptionally low water levels and high temperatures were 417 observed in 2012, whereas 2017 coincided with major flooding from the Ottawa River during the 418 spring freshet and sustained high discharge throughout the season. The other four years, either 419 tended to be hot with low water levels (2016) or had intermediate water levels (0.69 to 0.77 m 420 CD) but varying mean water temperatures (2013, 2014, 2015). In these latter years, 2014 was 421 colder, both in mean temperature (21°C) and accumulated degree day (Figure 2b). For the 422 remaining two years (2013, 2015), the mean temperatures were similar. However, the greater 423 variability in daily temperatures with sporadic colder days in 2013 ($min_{2013} = 17^{\circ}C$, $min_{2015} =$ 424 18.8°C, SI Figure S4) resulted in a lower accumulated degree day that year compared to 2015.

Hotter years (2012, 2015, 2016), with both higher accumulated degree days and mean 425 426 water temperature, tended to receive more PAR than colder years, but this was not always the 427 case. For example, the second hottest summer (2016) had similar irradiance levels to the two 428 coldest ones (2014, 2017). Whereas mean water temperature and water levels followed a pattern 429 of negative correlation in the SLR, there was no correlation between SLR temperatures and 430 tributary variables (Figure 2e-g). This suggests that tributary loadings were influenced 431 asynchronously from flow patterns in the mainstem. Tributary discharge and weighted NO₃⁻ had 432 similar patterns albeit NO₃⁻ concentrations were markedly lower in 2012 as compared to other 433 years (med₂₀₁₂ = 0.26 mg N L^{-1} , med_{other} = $0.46-0.72 \text{ mg N L}^{-1}$). This was a function of more 434 variable NO₃⁻ and considerably higher concentrations in the Yamaska as compared to the Saint-435 François since its proportion to total discharge stayed relatively stable throughout the different summers (med_{all} = 0.20, med_{year} 0.27 to 0.13, SI Figure S5). NO₃⁻ concentrations from Yamaska 436

- 437 were particularly high during the intermediate temperature years ($med_{2015} = 1.75 mg L^{-1}$, med_{2013} 438 = 1.77 mg L⁻¹, $med_{other} = 0.08$ to 0.82 mg L⁻¹), up to 5.5 mg L⁻¹. In contrast, TP and turbidity did 439 not show any obvious differences among years, but TP was lower in 2015 ($med_{2015} = 30.2 \mu g P$ 440 L⁻¹, $med_{other} = 33.6$ to 44.0 $\mu g P L^{-1}$, Figure 2g) and turbidity higher in 2012 ($med_{2012} = 17 NTU$, 441 $med_{other} = 10$ to 15 NTU, SI Figure S4) as compared to other years. 442 From our comparison with field biomass estimates, we deemed the downstream LSP 443 water level slope a good indicator of SAV biomass. Looking at the estimated temporal SAV
- 444 biomass changes throughout the six summers, two patterns of seasonal progression were
- 445 revealed (Figure 3). The first pattern reflected a reduced SAV biomass reaching its peak earlier
- 446 during the season (late July to late August). This pattern was observed over both the colder years
- 447 with high water levels (2013, 2014, 2017) and the hottest year with very low water levels (2012),
- 448 suggesting that extreme environmental conditions can restrict SAV growth. In addition, three of
- 449 these four years (all but 2013) experienced lower NO_3^- inputs from tributaries. The second
- 450 pattern occurred over years of highest SAV biomass, peaking later during the season
- 451 (September). SAV growth was favored by warm summers (2015, 2016) with intermediate water
- 452 levels and high NO₃⁻ inputs. Overall, these observations suggest that maximum SAV biomass
- 453 could vary following a dome-shaped curve where the optimum coincided with summers of warm
- 454 temperature, average water levels and abundant nitrate.
- 455

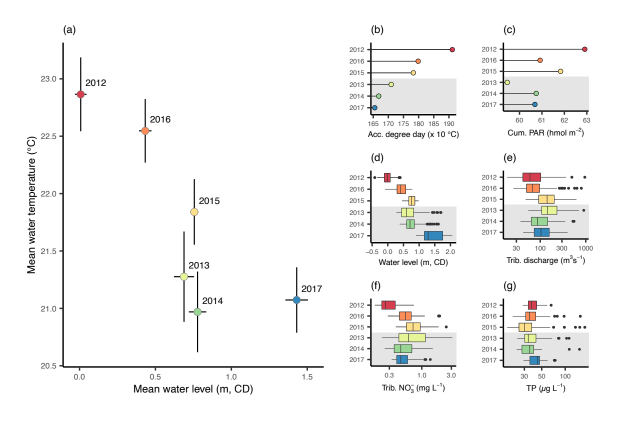




Figure 2. Interannual variation in environmental variables during summers from 2012 to 2017 457 458 (June 21 to September 22) for the main stem (a-d) and tributaries (e-g). Error bars are 95% 459 confidence intervals around the mean. Dot and boxplots are ordered by decreasing water 460 temperature as accumulated degree day using a warm to cold color gradient; cold summers are indicated by the light grey shaded background. Vertical bars within boxes indicate median value, 461 462 box boundaries represent 25th and 75th percentile and whiskers range from 10th to 90th 463 percentiles. Cum. cumulative, PAR photosynthetic active radiation, Acc. accumulated, CD Chart Datum, Trib. tributaries. Cumulative summer PAR and degree-days > 7°C were calculated from 464 May 1st to September 22. Water levels are from LSP station and SLR temperature from Trois-465 Rivières. Tributaries discharge correspond to the sum of Yamaska and Saint-François discharges. 466 467 Nitrate and total phosphorus are flow-weighted concentration from the two tributaries.

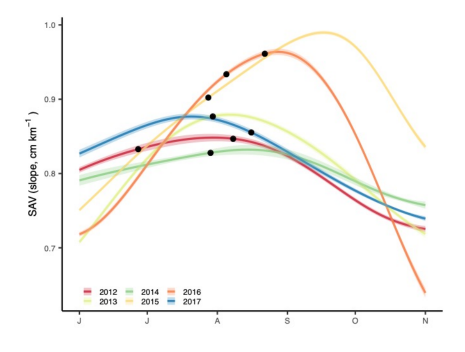




Figure 3. Interannual variation in temporal SAV biomass changes estimated daily from water level slope. Light-colored bands are 95% confidence intervals, but are not always visible given their small sizes. For each year, a date of formal biomass measurement allowed for estimated SAV biomass to be validated (black points). Slopes are the trend extracted using EEMD from raw surface water elevation, which corresponded to the SAV flow obstruction effect. SAV submerged aquatic vegetation, LSP Lac Saint-Pierre. Lines are color-coded by water temperature using a warm to cold color gradient.

477 3.2 NO_3^- Budget

To estimate the NO_3^- budget for the plume flowing across the SAV meadow (Figure 1), we first compared the modelled NO_3^- input to the output of the in situ sensor measurements over five summers (Figure 4). The time series for the sensor were complete for 2012 and 2015 (88 and 93 days) and truncated by about 29 days for the other years due to instrument failure, with 2014 having the shortest time series (24 days). The comparison revealed that the NO_3^- output followed the same temporal pattern as the estimated inputs from the tributaries, but at markedly lower concentrations, regardless of the contribution of each tributary to NO_3^- inputs.

485 The contribution of each tributary to the incoming NO_3^- loads, estimated from 486 conductivity, varied among years. Over the hot summers of 2012 and 2016, NO_3^- inputs

487	originated solely from Saint-François River, whereas over the remaining years (2013-2015), both
488	rivers contributed approximately equally. We also observed that the daily NO3 ⁻ concentration
489	range measured at the outflow, which reflected daily retention by autotrophic assimilation, was
490	most often considerably below estimated inputs, suggesting substantial removal by
491	denitrification. The sensor occasionally exhibited abrupt changes and elevated daily
492	concentrations, sometimes above estimated inputs, particularly in years when the Yamaska was
493	contributing to the inputs. These changes coincided with both high NO3 ⁻ concentrations from the
494	Yamaska River combined with short-term high discharge events that increased its proportion to
495	the total water inputs (SI Figure S5). We thus evaluated that on those days, tributary
496	contributions to the outflowing water mass were poorly estimated. As such, we excluded these
497	dates from budgets ($n_{excluded} = 31$, $n_{budget} = 292$) and assumed that the remaining NO ₃ ⁻ inputs were
498	accurately estimated.
499	The NO ₃ ⁻ budget in the plume flowing across the SAV meadow also displayed high

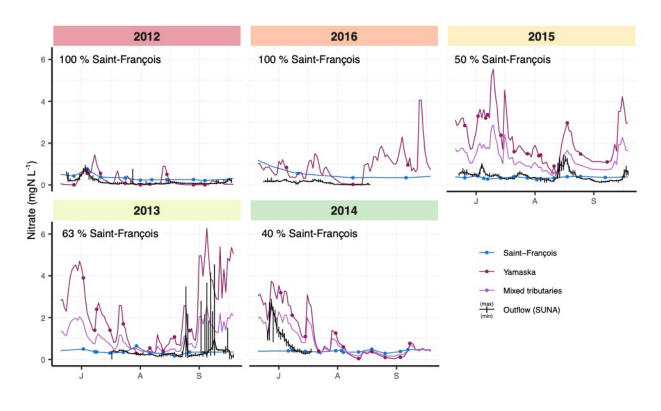
199 500 interannual variability (MANOVA p < 0.001) similar to the observed variation in temporal SAV 501 biomass, and NO₃⁻ inputs and outputs for the complete SAV meadow (Figure 5). All variables 502 contributed to among-year differences in budgets (LDA SI Table S3 and for process-specific 503 rates and velocities p < 0.03). Using PCA, we observed that the uptake rates (particularly for U_t, 504 U_a and U_d) were highly correlated to NO₃⁻ concentrations and tributary discharge (r > 0.53, 505 Figure 5a). Variables describing proportional uptake (R and U_d/U_t) and V_f , both total and 506 process-specific, where orthogonal to the uptake rates and were related to temperature and SAV 507 abundance, both of which were markedly higher in August 2016 (up to day 228 of the year, last 508 observation that year). In contrast, overall uptake rates tended to be differentiated by more 509 generalized annual conditions where hotter and lower water levels years resulted in lower NO₃⁻ 510 inputs and total retention than colder, higher water level years.

As such, this pattern of higher uptake measured in the plume was explained by the high NO₃⁻ loads in the colder years where both discharge and NO₃⁻ concentrations from tributaries (inputs) increased simultaneously (r = 0.82) and were strongly correlated to U_t ($r_{NO3} = 0.95$, $r_Q =$ 0.84). Interannual variations in NO₃⁻ loads entering the meadow induced sharp differences in measured areal total uptake rates among years with median differences of up to 1.6 g m⁻² d⁻¹ (range: med₂₀₁₂ = 0.1 g m⁻² d⁻¹, med₂₀₁₄ = 1.7 g m⁻² d⁻¹, Figure 5b). Similarly to total NO₃⁻ inputs to the overall SAV meadow (Figure 2), high inflowing NO₃⁻ concentrations from the tributary

518 plume reflected the greater contribution of the heavily enriched and flashy Yamaska River 519 waters. Inflowing NO₃⁻ concentrations to the plume were generally similar to those for the 520 complete SAV meadow (Figure 2), except for 2013 and 2014 (Figure 5c). These discrepancies 521 were due to the shortened periods when sensor data were available those years, with midsummer 522 2013 and early June 2014 having lower and higher NO₃⁻ inputs, respectively, than the rest of the 523 summer (Figure 4). Both U_a and U_d followed a somewhat similar pattern as NO_3^- inputs (SI 524 Figure S6), but 2013 displayed a distinctive pattern of high U_a as compared to its low inputs in 525 NO₃⁻. This is in part explained by lower U_d values, likely resulting from higher discharge and 526 lower plant biomass. Apart from that year, U_d tended to be higher and had a wider range than U_a, reaching its maximum at 2.6 g m⁻² d⁻¹ (in 2015) compared to the maximum U_a of 1.9 g m⁻² d⁻¹ (in 527

528 2014).

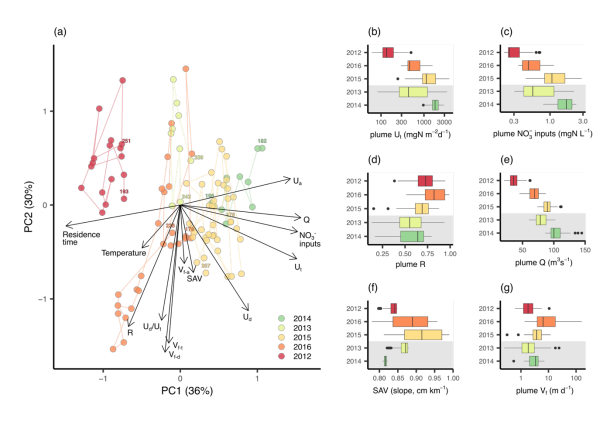
529 In contrast to uptake rates, R was highest during hotter summers with a median increase 530 of 5 to 27% as compared to colder years (med_{hot} = 69 to 82 %, med_{cold} = 55 to 64 %, Figure 5d). 531 The low R measured over the two hottest years (2012 and 2016) could be explained by the up to 532 three-fold reduction in discharge in comparison with colder years (med_{2012,2016} = 36-69 m³ s⁻¹, 533 $med_{2013, 2014} = 78-100 \text{ m}^3 \text{ s}^{-1}$). The low tributary discharge in 2012 and 2016 also nearly doubled 534 water residence time in the plume (med_{2012, 2016} = 50-60 h) compared to other summers (med₂₀₁₃₋ 535 $_{2015}$ = 35-38 h, Figure S6). However, in 2015 the residence time and discharge were similar to the 536 colder summers (med₂₀₁₅ = 35h, med_{2013, 2014} = 35-38 h). In this case, the high R could be 537 attributed to the higher SAV biomass of that year compared to colder ones; this factor likely also 538 contributed to the high R in 2016. The higher total $V_f(V_{f,t})$ in the high biomass years (2015, 539 2016) also suggested that the increased biotic activity resulted in higher overall retention and 540 uptake rates (Figure 5e-g). Additionally, the summers of 2015 and 2016 tended to have 541 proportion of denitrification from total uptake (U_d/U_t) 11 to 37 % higher than others, suggesting 542 an interplay between climate and SAV biomass on N fate (SI Figure S6). To further investigate 543 whether the changing SAV biomass affected the partitioning of N retention pathways, we 544 compared yearly medians of process-specific (V_{f-a}, V_{f-d}) to total uptake velocity (V_{f-t}, Figure 6). The higher V_{f-t} in the high biomass years could be attributed to either generally higher uptake 545 546 velocity due to denitrification (V_{f-d}, in 2015) or higher daily values that increased variation 547 around the estimate (2016). By comparison, when SAV biomass was reduced, V_{f-d} was lower 548 (2013) or similar (2012, 2014) to uptake velocity due to autotrophic assimilation (V_{f-a}).





550 Figure 4. Daily NO₃⁻ inputs from individual tributaries as well as their estimated mixing 551 concentrations in the plume compared to the median NO₃⁻ measured at the SAV bed outflow by 552 the in situ sensor. Black vertical bars represent daily minimum and maximum NO3-553 concentration at the outflow. For tributaries, lines are fitted concentration values from the 554 loadflex composite approach and points are measurements. Estimated input contributions from 555 the Saint-François River are indicated on each panel with the remaining proportion coming from 556 the Yamaska River. Mixed tributary lines for 2012 and 2016 are not presented since estimated 557 inputs are solely from the Saint-François River. Panels are ordered by summer water temperature 558 and water level (from hot-low level on top to cold-high on the bottom).

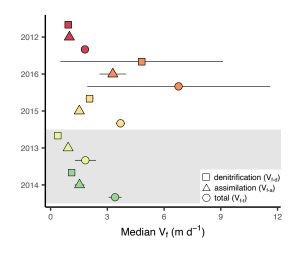
559



562 Figure 5. Correlative relationships and interannual variation in NO₃⁻ budget terms and their 563 potential predictors in the plume flowing across the SAV meadow. A) principal component 564 analysis (PCA) using scaling 2 where angles between arrows approximate correlation between 565 variables. B) to G) show interannual distributions with vertical bars within boxes indicating median value, box boundaries representing 25th and 75th percentile and whiskers ranging from 566 567 10th to 90th percentiles. PC principal components with explained variance in parentheses, U 568 uptake rate, R proportion of NO3⁻ retained, Q discharge, SAV submerged aquatic vegetation, V_f 569 uptake velocity, t total, a autotrophic assimilation, d denitrification. In a), colored numbers 570 correspond to the first and last day of the year of estimation, and lines connect observations in 571 temporal order. Years are ordered by decreasing water temperature using a warm to cold color 572 gradient; cold summers are indicated by light grey shaded background.

573





575

576 Figure 6. Comparison of median uptake velocity (V_f) from denitrification and autotrophic 577 assimilation to total uptake velocity per summer. Error bars are standard error. Years are ordered 578 by decreasing water temperature using a warm to cold color gradient; cold summers are indicated 579 by the light grey shaded background.

580 3.3 Prediction of Total NO₃⁻ Retention by the SAV Meadow

581 To estimate summertime NO_3^- retention in the complete SAV meadow, we built 582 regression models using our NO₃⁻ budget in the plume and its relationship with environmental 583 variables. The model to predict U_t with the highest AIC included both NO_3^- inputs and the 584 temporal SAV biomass estimate (SI Table S4) but, this model was not selected because the AIC 585 improvement was minor (< 2) compared to the simplest model with only one predictor. Given 586 the strong relationship between retention and inputs, U_t could be predicted solely from NO₃⁻ 587 concentration of those inputs (marginal $R^2 = 0.88$, SI Table S5). The intercepts of the NO₃⁻ and Ut relationship differed among years (conditional $R^2 = 0.92$). Although the AIC difference was 588 589 minor with the model without random structure, variable intercept per year was selected because 590 it homogenized the residual variance among years (SI, Table S6). Most importantly, the model 591 was substantially improved by adding an autocorrelation structure per date (autoregressive 592 model, AR-1), which considers that consecutive observations were highly correlated with each 593 other (r = 0.82). Using the fixed effect of the model with only NO₃⁻ as a predictor (SI Table S5) 594 and weighted concentrations of tributaries derived from the plume, we predicted Ut for the

- 595 complete SAV bed and summer period and derived R from yearly NO₃⁻ load (Table 2). Retention
- 596 was highly variable among years, showing a fourfold difference between extremes. The
- 597 interannual pattern in retention was similar to patterns in inputs, albeit the inputs exhibited a
- 598 much higher interannual variation than the outputs, with almost a sixfold difference between
- 599 extreme N loads. As a result, R tended to be somewhat higher (63-87%) in the warmer years
- 600 (2012, 2016, 2015) than in colder years (47-62%). A nearly complete retention (87%) was
- observed in 2016, concomitant with lower NO_3^- inputs. When calculating U_d from the overall
- 602 SAV bed U_t using the median plume U_d/U_t, these differences in retention with climate were more
- 603 important. Year with higher SAV biomass or lower NO₃⁻ inputs had 30 to 58% of inputs
- 604 permanently removed through denitrification compared to 18-22% during colder years with
- 605 reduced SAV biomass and higher NO_3^- loads.

606	Table 2. Whole-SAV meadow summertime NO_3^- inputs, total and denitrification uptake (U_t, U_d) ,
607	and proportional retention (R) modelled for six summers (June 21 to September 22). Nitrate
608	inputs were estimated from the sum of daily tributary loads. Ut was modelled from daily inputs,
609	using NO_3^- concentrations as the predictive variable, while U_d was estimated by multiplying U_t
610	by the plume yearly median proportion of U_d on U_t . Maximum SAV biomass was estimated from
611	the linear regression between rake estimated biomass and water level slope. CI confidence
612	intervals, denit. denitrification. Summers are ordered by decreasing water temperature.

Year	Total NO ₃ -	Ut	Ut 95% CI	Ud	R	R	Maximum SAV
	inputs				total	denit.	biomass
	(tonnes)	(tonnes)	(tonnes)	(tonnes)	(%)	(%)	(g m ⁻²)
2012	304	192	[173, 214]	91	63	30	31
2016	533	466	[392, 555]	311	87	58	109
2015	1188	751	[610, 927]	441	63	37	127
2013	1811	846	[670, 1073]	250	47	14	52
2014	726	444	[373, 529]	158	61	22	20
2017	622	384	[329, 448]	113	62	18	50

614 **4 Discussion**

615 In this study, we provide high frequency measures of N retention and its partitioning into 616 autotrophic and heterotrophic pathways across multiple years from a riverine SAV meadow, in 617 one of the largest rivers to date. We hypothesized that retention, measured as either uptake rates 618 (U) and proportion retained from inputs (R), would be a function of NO₃⁻ loads and of vegetation 619 biomass, both of which expected to be subject to climate-driven variations. We found that N 620 retention by the meadow in the main stem significantly attenuated the incoming loads from two 621 enriched agriculturally impacted tributaries. Estimated rates were among the highest reported so far in rivers (U_t median 576 mg N m⁻² d⁻¹, range 60 - 3893 mg N m⁻² d⁻¹), showing both annual 622 623 and interannual variations of this ecosystem service as a function of climate conditions. This 624 variation highlights the importance of measuring N retention rates multiple times within a year 625 and across several years. NO₃⁻ uptake rates in the SAV meadow were a function of NO₃⁻ loads, 626 which varied largely as a function of climate driven changes in discharge from the tributaries. 627 However, the climate effect on loads were not in synchrony with the water level changes in the 628 main stem. This was probably due to differential effect of precipitation patterns on basin-wide 629 water inputs and aspects related to regulated flow upstream from the study site versus localized 630 water inputs at the confluence. Proportional retention was higher in the hottest years and 631 coincided with either higher water residence time or higher SAV biomass in the meadow or both. 632 Maximum summer SAV biomass appeared to vary along a dome-shaped curve, and optimal 633 conditions were observed in summers with warm water temperatures, average water levels, and 634 loadings of abundant NO₃. Higher SAV biomass also influenced the partitioning of N retention 635 pathways by promoting permanent removal through higher denitrification. Our work indicates 636 that this riverine SAV meadow provides an important ecosystem service by reducing N export to 637 downstream waters by 47 to 87% and permanently removing between 14 and 58% of N inputs. 638 Thus, management actions should aim at preserving or restoring such critical riverine N removal 639 control points.

640

4.1 Nitrogen Retention in Riverine SAV Meadows

641 Vegetated sites in rivers are recognized for performing multiple aquatic ecosystem

642 services including N removal (Caraco et al., 2006). However, measuring seasonal and

643 interannual changes in SAV biomass and how it affects N processing in large meadows remains

644 a challenge due to the restricted ability to assess biomass changes underwater and the difficulty 645 to measure N retention in large river. We successfully followed SAV biomass daily using the 646 indirect measure of water level slope and N retention using a high frequency sensor. Using this 647 information, we found that only part of our hypothesis was supported as there was no conclusive 648 evidence of a strong relationship between uptake rates and SAV biomass. This was likely due to 649 the overriding role of NO_3^- inputs in determining total uptake rates (U_t) of the meadow. The 650 strong relationship between inputs and Ut is not suprising as N retention is typically modelled 651 using first-order kinetics, where process rates are a linear (log-log) function of substrate 652 concentrations (e.g. Seitzinger et al., 2006; Wollheim et al., 2006). For a maximal functioning, N 653 removal sites thus have to be spatially connected to high N loading locations, as is the case for 654 the confluence zone of this study (Cheng et al., 2020; Mitsch et al., 2001). 655 The strong positive effect of SAV biomass on N uptake rates has previously been 656 reported when comparing river reaches with contrasting plant cover (85% vs 45%, SI Table S7, 657 Preiner et al., 2020). We did not observe a clear pattern with plant biomass probably because 658 SAV cover was always high across the N budget plume, thus always favoring high N uptake. 659 Rather, the interannual differences in SAV was one of plant density, or biomass, which likely 660 changed SAV surface area as these two measures of SAV are well correlated (Armstrong et al., 661 2003). Greater SAV surface area obstructs flow, which reduces velocity and increases water 662 residence time (Madsen et al., 2001). This latter increase should also increase the proportion of N 663 retained since a prolonged contact between water column NO₃⁻ with plants and sediments can 664 lead to the exhaustion of this substrate pool. Indeed, and as typically observed (Seitzinger et al., 665 2006), there was a general inversed interannual pattern between proportion retained and 666 discharge (Figure 5d-e). However, high biomass years (2015, 2016) had higher than expected retention (1st and 3rd highest) given their fast flow and high discharge (4nd and 2nd highest). 667

Higher biomass years therefore probably had added plant surface area facilitating N exchangebetween plants, their epiphytes, sediments, and water column.

Additionally, our results suggest that changing SAV biomass affected N fate. When SAV
was abundant, denitrification rates in the plume were high and accounted for 59 to 67% of total
uptake in contrast to 30 to 48% during years when biomass was reduced. Similarly, when using
V_f, a metric that is independent of concentrations and hydrologic characteristics (Ensign &
Doyle, 2006; Wollheim et al., 2006), the biotic demand of denitrification (V_{f-d}) was higher than

675 that of assimilation during high biomass years, and vice versa. At the scale of the SAV bed, these 676 differences translated into an up to four times higher rate of permanent removal in high biomass 677 years compared to lower ones. These results are in agreement with studies that considered both assimilatory and dissimilatory pathways (Heffernan & Cohen, 2010; James, 2010; Pinardi et al., 678 679 2009; Preiner et al., 2020, summarized in SI Table S7), while reports focusing solely on N 680 assimilation by SAV and their epiphytes generally found that overall uptake of water column 681 NO₃⁻ concentrations were typically lower (Desmet et al., 2011; Diamond et al., 2021). The 682 dominance of denitrification in a higher biomass year can be explained by the structural 683 complexity of SAV and increased surface area that supports biofilms where denitrification can 684 occur (Eriksson & Weisner, 1996). SAV also favors denitrification in sediments, by promoting 685 organic matter particle settling, which increases sediment respiration rates and the anoxic 686 conditions needed for denitrification (Cornwell etal. 1999). Furthermore, SAV provide labile 687 dissolved organic matter directly to denitrifiers within the sediments through the release of 688 exudates from their roots (Karjalainen et al., 2001). Overall, in advection-dominated systems 689 such as rivers where NO_3^- is continuously supplied from the watershed (Seitzinger et al., 2006), 690 the dominance of denitrification on total uptake in SAV beds is likely a function of how plants 691 amplify reactive surface areas, enrich sediments, and increase contact rates through a reduction 692 in flow creating optimal environmental conditions for denitrifying microbes.

693 Given the optimal conditions for N processing at our SAV control point, both total N 694 uptake and denitrification rates measured in this study ($U_t = 576 [60-3893 \text{ mg N m}^2 \text{ d}^{-1}]$, $U_d =$ 338 mg N m⁻² d⁻¹ [1-2624 mg N m⁻² d⁻¹] SI Table S7) are among the highest reported, but remain 695 696 within the range of previous estimates derived from similar methods in SAV meadows [up to 697 840 mg Nm⁻² d⁻¹], Heffernan & Cohen, 2010; Preiner et al., 2020). Our estimates are also higher 698 than those previously reported in a meta-analysis of denitrification in rivers worldwide (up to 699 1143 mg m⁻² d⁻¹, Piña-Ochoa & Álvarez-Cobelas, 2006). This may be due to differences across 700 methods, temporal and spatial scales of inquiry, and the presence, size and location of the SAV 701 bed under study. The meta-analysis involved short-term laboratory (cores) or in situ (chambers) 702 incubations providing discrete denitrification rates in time. We report daily and interannual net 703 uptake rates at the reach-scale that reflect the balance between processes that produce 704 (nitrification) and consume NO₃⁻ (assimilation, denitrification), and its variability throughout 705 multiple growing seasons. Our estimates may be higher because of a greater sensitivity of the

two-station mass-balance approach to increase in NO₃⁻ concentration and reach length (von 706 707 Schiller et al., 2011). Still, the uptake rates from this method are generally in agreement with multiple site mass-balance and incubation approaches (Piña-Ochoa & Álvarez-Cobelas, 2006; 708 von Schiller et al., 2011). The importance of the size and location of vegetated areas on flow 709 710 modification and N attenuation is highlighted by the similar denitrification rates measured in a 711 dense floating bed of *Trapa natans* in a sheltered bay of the Hudson River (1.5 km², 518-994 mg 712 N m⁻² d⁻¹), in contrast to the undetectable rates in a small (0.6 km²) Vallisneria meadow in a fast-713 flowing channel from the same river (Tall et al., 2011). The floating meadow acted in a similar 714 way to this study large SAV meadow (~10 km²) that both intercepted nutrient-rich waters and 715 reduced velocities (Bulat et al., 2019), thus promoting high nitrate removal. Another line of 716 evidence supporting the robustness of our estimates is that our measured uptake velocities (V_f, median 3.0 m d⁻¹) were similar to those reported for the Upper Snake River (WY, USA, 12 m³ s⁻ 717 718 ¹, 0.6 to 13.0 m d⁻¹, Tank et al., 2008). However, our estimates were much more variable both 719 within and among seasons (range $0.2 - 160.8 \text{ m d}^{-1}$), highlighting the considerable heterogeneity 720 of biotic activity, load and NO₃⁻ retention capacity in time at this dynamic confluence zone.

721 4.2 Complex Effects of Climate at a Confluence Zone

722 By measuring environmental conditions over 6 summers, we showed that different 723 climatic variables drove complex variations in SAV biomass and N retention in a large riverine 724 SAV meadow. The predominant influence of climate on SAV biomass was perceptible through 725 the large range of water levels and temperatures among years: our observations spanned the 726 lowest (2012) and highest (2017) mean summer water levels recorded at LSP gauging station for 727 more than a century (data available since 1914, meds-sdmm.dfo-mpo.gc.ca). Our observations 728 also included the warmest (2012) and the third coolest (2014) mean summer water temperature 729 over the last 20 years (data available since 2000, ogsl.ca). These climatic variations translated 730 into a \approx 6-fold range in maximum SAV biomass within the study area during the 2012-2017 731 period (Table 2).

The range in climate variation captured on such a short time scale reflect the sensitivity of rivers to these changes and impacts that are likely already taking place (Gudmundsson et al., 2021; Nijssen et al., 2001). Among these impacts, we could expect that increased precipitation should favor N transport, as is generally seen in anthropogenically altered river networks

736 (Goyette et al., 2019; Howarth et al., 2012). Inflowing NO_3^- concentrations to the SAV meadow 737 were correlated to the discharge of incoming tributaries, but concentrations changes were mostly 738 a function the relative contribution of each tributary to the total at the confluence site, 739 particularly from the smaller, more NO₃⁻ rich, and flashier Yamaska River. This river displayed 740 the typical seasonality of agricultural watershed where discharge and NO₃⁻ concentrations vary in 741 synchrony (Van Meter et al., 2020). Indeed, in smaller catchments like the Yamaska, riverine 742 nutrient loads are typically more variable than in larger catchments, like the Saint-François (2.1 743 times larger than Yamaska) and the SLR main stem, which integrate loads from many sub-744 watersheds that dampens individual basin signals (Burt & Pinay, 2005; Chezik et al., 2017). 745 However, the variation in NO₃⁻ inputs was disconnected from temperature and water level 746 variation in the main stem that was more reflective of water inputs at the scale of the entire SLR 747 watershed. These asynchronies might become more common as more pronounced climate-driven 748 changes and reduced summer discharge are expected over the next decade in southern SLR 749 tributaries, in contrast to maintained or amplified discharge in northern SLR tributaries (Boyer et 750 al., 2010; Ouranos, 2015). Regardless, the highest SAV biomass were recorded at the confluence 751 zone over years of intermediate water level and temperature conditions.

752 Past observations suggest that optimal conditions for SAV biomass are met when water levels are high in LSP, and are limited when levels are low, favoring reed bed expansion (Hudon, 753 754 1997). We observed the latter in the extremely low-level year of 2012. SAV biomass was among 755 the lowest recorded during this study and can be explained by the large areas of riverbed being 756 exposed to air that year, potentially leading to SAV loss through the dry-out of above ground 757 biomass. The elevated water temperatures might also have favored phytoplankton and 758 metaphyton growth that year through slow water transit time, high water column illumination 759 and increased nutrient release from the sediments. Indeed, low water levels in LSP are associated 760 with increased filamentous green algae biomass (Cattaneo et al., 2013). Overall highest TP (~39 761 μ g L⁻¹), chlorophyll *a* (~3.6 mg L⁻¹) and lowest secchi depths (~67 cm) were also measured 762 within the bed in 2012 despite average TP inputs (Figure 2g). In contrast with Hudon (1997), 763 however, our result suggests that SAV biomass is also constrained at extremely high water level 764 years when temperatures are cold. The effect of high waters can be explained by SAV light 765 limitations through the typical positive relationship between maximum SAV colonization depth 766 and water transparency (e.g. Chambers & Kalff, 1985). Incoming turbidity did not display a

coherent pattern as a function of water level changes (Figure S4), and increased water height in
itself likely created deep areas where light did not reach the bottom, thus restricting plant
colonization. In the LSP fluvial lake, the deeper waters also correspond to an increased exposure
to fast currents, further limiting SAV growth (Hudon et al., 2000).

When water depth and transparency allowed light penetration to the sediments and thus 771 772 plants were not limited by light, increases in temperature probably directly favored SAV growth 773 and biomass accumulation. This has previously been shown in nearby shallow lakes where sharp 774 differences in SAV biomass for the same lakes were observed in a cold and a hot year (Rooney 775 & Kalff, 2000). The direct effect of temperature is likely a function of increased photosynthetic 776 rates that drive plant growth (Brown et al., 2004; Farguhar et al., 1980; Riis et al., 2012). 777 Experimental studies support this potential direct effect in mesotrophic conditions, when light 778 availability does not limit plant growth (Barko et al., 1982; Ersoy et al., 2020). These conditions 779 were met in intermediate water level and hotter years (2015-2016) where TP within the meadow 780 ranged from $\sim 19-26 \mu g P L^{-1}$. During those years, SLR temperature was near the previously 781 determined growth optimum of \sim 25-28°C, below deleterious temperatures (28-30 °C, Barko et 782 al., 1982; Riis et al., 2012). Indeed, during our field survey, water temperatures were always 783 below 27.2 °C and generally around 24 °C during 2015 and 2016. The intermediate water level of those years likely also increased light availability to a more extensive area for SAV growth 784 785 (Beklioglu & Altinayar, 2006). SAV growth during those years might also have been facilitated 786 by the time lag that allowed for plant bed recovery following the 2012 drawdown that potentially 787 damaged winter buds and reduced biomass the subsequent years. We argue that this is unlikely 788 as SAV bed can regenerate from seed banks (Kimber et al., 1995), and the 2012 event still left 789 large meadow areas covered with water. Additionally, the highest biomass year (2016) was 790 followed by a low biomass year, suggesting that autocorrelation between consecutive years was 791 not a dominant driver of SAV biomass during our study period. Our results rather indicate that 792 water levels combined with water temperature seems critical in determining biomass as for 793 example in 2014 when water levels were similar to the high biomass year of 2015 and 2016, 794 temperatures were coldest and SAV biomass was lowest. As such, when areal light to sediment 795 is apparently equal, temperatures in hot years directly stimulated SAV growth in LSP.

796 **5** Conclusions

797 In this study, we showed the complex influence of climate on N retention in a SAV 798 meadow in the SLR and report some of the highest measured estimates in a large river from this 799 control point. This influence includes modulating N loads to the meadow combined with SAV 800 biomass and water residence time within it, mitigating the total amount and the proportion of N 801 retained, as well as N fate. Our results suggest that SAV biomass is restricted by extreme dry and 802 wet years, which may increase in frequency with climate change, thus suggesting a diminishing 803 role of SAV on N retention in the SLR. Indeed, SLR water levels have been decreasing since the 804 1970s (Hudon et al., 2018), and climate projections of increasing summer air temperatures will 805 further reduce levels through higher evaporative loss. However, higher precipitations are 806 projected with climate change, particularly in the North (Ouranos, 2015), which might also lead 807 to more floods as observed in 2017 and sustained high water levels throughout summers. There 808 was no direct correlation between SLR water levels and tributary NO₃⁻ loads at our confluence 809 site, probably due to asynchronous precipitation patterns across sub-watersheds in the SLR basin. 810 During years with overall higher tributary discharge, short-term, heavy precipitation patterns 811 favored the flashiness of the Yamaska River, yielding high inflow of turbid, nutrient-rich waters. 812 Our sensor time series displayed more frequent NO₃⁻ spikes following summer rain, potentially 813 reflecting higher N export during these events due to reduced water residence time. As extreme 814 precipitation events are expected to become more common with climate change, these moments might increasingly contribute to N exports (Goyette et al., 2019; Lu et al., 2020). However, we 815 816 could not calculate N budgets during these events because of the changing tributary contribution 817 to waters flowing through the SAV meadow. Future studies should investigate how such events 818 contribute to overall N exports and assess whether SAV lose their capacity to dampen hydraulic 819 flows and retain N under such circumstances.

Although wetlands in watersheds are considered to have a disproportionate role in N retention on the landscape (Cheng et al., 2020), we argue that shallow-water underwater plant meadows should also be considered in conservation assessments. Even when SAV biomass was low, the meadow provided an important ecosystem service by retaining large amounts of N, thus decreasing impacts of overall N delivery from agricultural rivers to the SLR estuary. The proportion retained and permanently removed was highest during hotter years with high SAV biomass, while the absolute amount retained was related to N loads which reflected localized

827 precipitation patterns from the two tributary watersheds. Although the effect of SAV is seasonal, 828 it coincides with the period when N export has the highest potential impact in the SLR estuary by 829 contributing to the reported eutrophication, which partially explains growing hypoxia in its deep 830 waters (Gilbert et al., 2005; Lehmann et al., 2009). We estimate that during the SAV growing 831 season, between 2 to 9 t N per day could be retained, which is higher than previous estimates of 832 1.5 t N per day from this same location using a coarser mass-balance approach (Hudon et al., 833 2017). Assuming a 1.1 kg N per summer based on an annual per capita excretion of 4.4 kg N yr⁻¹ (Howarth et al., 1996), this would be the equivalent of retaining the N generated by a city of 0.17 834 835 to 0.77 million people over the SAV growing season. Although we illustrated the value of SAV 836 meadows from a N perspective, other cultural and provisioning services are supplied by SAV in 837 general (Hilt et al., 2017) and in particular at our site, commercial and recreational fisheries 838 (Giacomazzo et al., 2020; He et al., 2016). Therefore, management action should aim at 839 preserving or restoring such critical sites within the shallow-water areas of large rivers.

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852 **Open Research**

B53 Data used for figure creation and statistical analysis in this study are all openly available

at the Zenodo repository via <u>https://doi.org/10.5281/zenodo.6413187</u> with license under a

855 Creative Commons Attribution 4.0 International.

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