Physical and stoichiometric controls on stream respiration in a 1 headwater stream 2

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4 Jancoba Dorley¹, Joel Singley^{2,3}, Tim Covino^{4,5}, Kamini Singha⁶, Michael Gooseff^{7,8}, David Van 5 Horn⁹, Ricardo González-Pinzón¹

6 Correspondence to: Ricardo González-Pinzón (gonzaric@unm.edu)

7 ¹Civil, Construction and Environmental Engineering, University of New Mexico, Albuquerque, NM USA

89 ²Environmental Studies Program, University of Colorado, Boulder, CO USA

³Biology, Marine Biology, and Environmental Science, Roger Williams University, Bristol, RI USA

10 ⁴Ecosystem Science and Sustainability, Colorado State University, Fort Collins, CO USA

11 ⁵Department of Land Resources and Environmental Sciences, Montana State University, Bozeman, MT USA

12 ⁶Geology and Geological Engineering, Hydrologic Science and Engineering Program, Colorado School of Mines, 13 Golden, CO USA

14 ⁷Civil, Environmental and Architectural Engineering, University of Colorado, Boulder, CO USA

15 ⁸Institute of Arctic and Alpine Research, University of Colorado, Boulder, CO USA

16 ⁹Department of Biology, University of New Mexico, Albuquerque, NM USA 17

18 Abstract. Many studies in ecohydrology focusing on hydrologic transport argue that longer residence times across a

19 stream ecosystem should consistently result in higher biological uptake of carbon, nutrients, and oxygen. This

20 consideration does not incorporate the potential for biologically mediated reactions to be limited by stoichiometric

21 imbalances. Based on the relevance and co-dependences between hydrologic exchange, stoichiometry, and

22 biological uptake, and acknowledging the limited amount of field studies available to determine their net effects on

23 the retention and export of resources, we quantified how microbial respiration is controlled by the interactions and

24 supply of essential nutrients (C, N, P) in a headwater stream in Colorado, USA. For this, we conducted two rounds

25 of nutrient experiments, each consisting of four sets of continuous injections of Cl- as a conservative tracer,

26 resazurin as a proxy for aerobic respiration, and one of the following nutrient treatments: a) N, b) N+C, c) N+P, and

27 d) C+N+P. Nutrient treatments were considered as known system modifications to alter metabolism, and statistical

28 tests helped identify the relationships between hydrologic transport and respiration metrics. We found that as

29 discharge changed significantly between rounds and across stoichiometric treatments, a) transient storage mainly

30 occurred in pools lateral to the main channel and was proportional to discharge, and b) microbial respiration

31 remained similar between rounds and across stoichiometric treatments. Our results contradict the notion that

32 hydrologic transport alone is a dominant control on biogeochemical processing and suggest that complex

33 interactions between hydrology, resource supply, and biological community function are responsible for driving in-

34 stream respiration.

36 1 Introduction

37 High biochemical processing rates in streams and rivers occur at locations and times where the dynamic 38 interconnections among hydrologic exchange, residence time, nutrient supply, and microbial biomass combine to 39 form optimum conditions for metabolic activity (i.e., the transformation of nutrients, carbon, and oxygen or another 40 electron acceptor into energy and biomass). The exchange of water between the main channel and transient storage 41 zones, where most microbes exist, is the primary mechanism supplying carbon, nutrients, and oxygen to 42 metabolically active zones (Gooseff et al. 2004; Covino et al. 2010b, 2011; Knapp et al. 2017; Gootman et al. 43 2020). The extent of water exchange controls the residence time of solutes (Drummond et al., 2012; Gomez et al., 44 2012; Patil et al., 2013), their chemical signatures (Covino and McGlynn 2007), as well as the microbial 45 composition and their metabolic functioning (Blume et al. 2002; Navel et al. 2011; Li et al. 2020). Exchange 46 patterns are influenced by geomorphologic conditions (Kasahara and Wondzell 2003; Cardenas et al. 2004; Gooseff 47 et al. 2005; Emanuelson et al. 2022), hydrologic conditions (i.e., discharge and surrounding water table 48 configuration) (Gooseff et al. 2005; Wondzell 2006; Ward et al. 2013; Ward and Packman 2019), and biofilm 49 growth (Battin et al. 2003; Wen and Li 2018). The spatiotemporal variability in exchange processes and resource 50 availability (e.g., seasonal variations in nutrient loads) create heterogeneous hydrologic and biogeochemical 51 gradients across space and time, within which ecosystem metabolism occurs (Mulholland et al., 1985; Mulholland & 52 Hill, 1997). 53 To date, studies with a focus on hydrologic transport argue that longer residence times across a stream

54 ecosystem should consistently result in higher biological demand for carbon, nutrients, and oxygen (Valett et al.

55 1996; Gooseff et al. 2005; Wondzell 2006; Gomez et al. 2012; Zarnetske et al. 2012; Ward et al. 2013; Li et al.

56 2021), not fully incorporating the potential for biologically mediated reactions to be limited by stoichiometric

57 imbalances. Ecological stoichiometry is the notion that biota balance the consumption of nutrients with energy

58 requirements. Redfield (1934) noted that marine phytoplankton generally contained a ratio of C:N:P of 106:16:1 in

59 their biomass, and these ratios are similar to those available in their environment. This "Redfield ratio" suggests that

60 an ecosystem requires an optimal ratio of available nutrients to flourish and has been used as a guide for many other

61 environmental stoichiometry studies. In a study of streams across eight biomes, Dodds et al. (2004) noted that N

62 consumption depends in part on the C:N ratio of organic matter in streams and suggested that shifts in these state

63 ratios likely influence N retention.

64 The net effect of supply and demand of resources can be explored with the non-dimensional Damköhler 65 number, *Da* (Harvey et al. 2013; Pinay et al. 2015; Krause et al. 2017; Ocampo et al. 2020), which quantifies the 66 ratio of transport (i.e., supply) to biological uptake (i.e., demand) timescales along flow paths (Oldham et al. 2013;

67 Liu et al. 2022). Similar to any other non-dimensional number, *Da* offers simplicity and objectivity for inter-site and

68 intra-site comparisons. Da has been used to provide insight into the factors limiting the supply and demand of

69 resources (Harvey et al. 2005), as values of $Da \sim 1$ define a balance between transport and uptake time scales, which

70 theoretically result in maximal resource retention. Accordingly, where or when $Da \ll 1$, i.e., the uptake timescale is

71 much greater than the transport timescale, uptake is suboptimal, and it is referred to as reaction limited because even

though resources became available through hydrologic exchange, they were not fully taken up (i.e., assimilated).

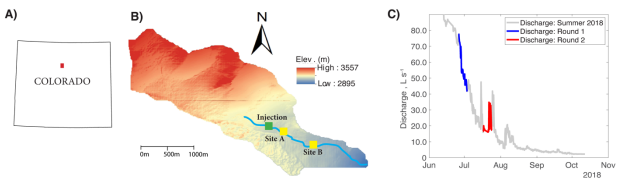
73 Conversely, where or when $Da \gg 1$, i.e., the transport timescale is much greater than the uptake timescale, resources

- 74 become scarce or transport-limited, and biologically inactive subregions start to develop (González-Pinzón and
- Haggerty 2013; Harvey et al. 2013; Gootman et al. 2020). While *Da* captures essential components of the potential
- 76 interactions between the supply and demand of ecologically relevant resources, it does not explicitly capture the role
- of stoichiometric limitations on the supply (i.e., C:N:P ratios in water fluxes) and demand (C:N:P biomass
- 78 composition and needs) of resources (Tromboni et al. 2018). This is because *Da* numbers are estimated from solute-
- 79 specific mass balances, which inform transport and reaction timescales for one resource at a time (e.g., only N), in
- 80 isolation of other stoichiometrically relevant resources that can become limiting factors (e.g., C and P).
- 81 Based on the relevance and co-dependences between hydrologic exchange, stoichiometry, and biological
- 82 uptake, and the limited amount of field studies available to determine their net effects on the retention and export of
- 83 resources, we sought to quantify how metabolic activity is controlled by the interactions and supply of essential
- 84 nutrients (C, N, P). More specifically, we tested if variations in stoichiometric conditions can induce metabolic
- 85 limitations at which residence time alone becomes a weak predictor of stream respiration. We addressed the
- 86 following research question: *How is microbial respiration controlled by hydrologic exchange vs. stoichiometric*
- 87 *conditions (i.e., supply of C, N, and P)*? We hypothesized that aerobic respiration would be maximized when
- 88 nutrient supply and demand were nearly balanced for a given hydrologic condition. To test this, we conducted a
- 89 repeated set of stream tracer injections in Como Creek, a mountain stream in Colorado, USA, varying stream C
- 90 (acetate; sensu Baker et al., 1999), N (NaNO₃), and P (KH₂PO₄) concentrations to manipulate stoichiometry and
- 91 nutrient supply. We repeated experiments under different flow conditions to quantify the tradeoffs between supply
- 92 (transport and delivery of nutrients), and demand (microbial respiration). We tested for statistical relationships
- 93 between hydrologic transport metrics and respiration metrics using the resazurin-resorufin tracer system (González-
- 94 Pinzón et al., 2012; Knapp et al., 2018) and contextualized our findings within the framework of the Damköhler
- 95 number.

96 2 Methods

97 2.1 Site Description

98 Our research experiments were conducted in Como Creek, a forested pool and riffle stream in Colorado, 99 USA. Como Creek is a tributary to Boulder Creek, with land cover consisting of approximately 20% alpine 100 meadow-tundra and 80% conifer forest. The study reach drains a 5.4 km² catchment, with elevations ranging from 101 2895-3557 m and a mean average precipitation of 883 mm/y (Ries III et al. 2017; Emanuelson et al. 2022). Como 102 Creek has a snowmelt-driven hydrograph with stream discharges ranging from 1-98 L/s and features short-lived 103 increases in discharge during the monsoon season between July and August (Figure 1). The study reach is a multi-104 thread channel with substrate ranging from small gravel to bedrock. Additionally, the channel has an average width-105 to-depth ratio of 11.5, a sinuosity of 1.1, and an average longitudinal slope of 21% (Natural Resources Conservation 106 Service).



1082018109Figure 1: A) Location of Como Creek watershed in Colorado, B) detailed map of the watershed where Sites A and B are11050 and 350 m downstream from the injection location, and C) hydrograph and timing of experimental work; each round111of experiments consisted of four treatments featuring N, N+C, N+P, and C+N+P nutrient additions.112

113 2.2 Stream tracer injection experiments

114 We conducted two rounds of experiments, each consisting of four sets of continuous injections (lasting \sim 4-115 7 h) of Cl⁻ as a conservative tracer, resazurin (referred to as Raz hereafter) as a proxy for aerobic respiration, and one 116 of the following nutrient treatments: a) N, b) N+C, c) N+P, and d) C+N+P. In our study, the nutrient treatments are 117 treated as known system modifications (control variables) to alter metabolism. Also, we use the transformation of 118 Raz, which occurred at the same spatiotemporal scales of the nutrient additions, to calculate how changes in 119 stoichiometric conditions and discharge affect respiration. Briefly, the reactive tracer Raz (blue in color) is 120 irreversibly reduced to resorufin (Rru, red) under aerobic respiration, and the relationship between Raz 121 transformation and oxygen consumption is linear (González-Pinzón et al. 2012, 2014, 2016; Knapp et al. 2018; 122 Dallan et al. 2020). 123 Before each tracer injection, we used the Tracer Injection Planning Tool (TIPT) (González-Pinzón et al. 124 2022) to estimate the amount of tracer mass needed to reach steady state conditions at the downstream site and to 125 estimate the duration of the tracer breakthrough curves. From our field sampling, ambient concentrations of nitrate 126 averaged 0.035 (± 0.002) mg/L. We corroborated this value with a study by (Smith et al. 2003), who generated 127 estimates of background total nitrogen (TN) and total phosphorous (TP) yield and concentrations throughout the 128 stream-river network in 14 ecoregions of the conterminous US. That study found 75th % quartile TN= 0.21 (±0.05) 129 mg/L and TP= 0.02 (±0.005), which indicates relatively low nutrient concentrations compared to agricultural 130 streams in the US Midwest featuring ambient concentrations of up to two orders of magnitude higher. Based on 131 estimated discharges and reach lengths, we targeted a maximum concentration of 2 mg/L for Cl, and 100 µg/L at the 132 most downstream locations. The concentrations for nitrogen, phosphorus, and carbon were based on the expected 133 detection limit of phosphate (i.e., 0.1 mg/L) for common ion chromatographs. From that minimum phosphate 134 concentration expected, we scaled the masses of nitrogen and carbon using the 106C:16N:1P Redfield ratio 135 (Redfield, 1934). Table 1 shows the masses injected and the discharges observed during the studies. Note that we 136 allowed the stream to return to ambient concentrations for one day after each set of injections. 137

Table 1: Tracer injection data for each round of experiments at Como Creek.

Date	Treatment	Discharge (L/s)	Start time	End time	NaCl (g)	KNO3 (g)	KPO4 (g)	Sodium Acetate (g)	Raz (g)
Round 1									
6/26/18	Ν	74	11:30	17:00	32653	502	-	-	150
6/28/18	N+C	61	10:08	14:10	32680	500	-	2000	150
6/30/18	N+P	53	10:00	17:00	32680	500	400	-	150
7/2/18	C+N+P	49	9:59	14:00	32680	500	400	2000	150
Round 2									
7/17/18	Ν	20	10:30	14:35	10000	100	-	-	30
7/19/18	N+C	17	10:00	13:59	10000	100	-	400	30
7/21/18	N+P	17	10:00	14:06	10000	100	80	-	30
7/23/18	C+N+P	25	9:30	13:35	10000	100	80	400	30

140

141 We collected 20 mL aliquots in each tracer injection 50m and 350m downstream of the injection site 142 (labeled Sites A and B, Figure 1) to generate tracer breakthrough curves (BTCs) for Raz. All samples were filtered 143 immediately after being collected using a 0.7 µm GF/F filter (Sigma-Aldrich) and kept on dry ice during transport 144 until they were frozen at -4°C for laboratory analysis for Raz concentrations. All analyses took place within a week 145 after the end of each round of injections. At the laboratory, each sample was buffered to a pH of 8.5 (1:10 buffer-to-146 sample) following Knapp et al. (2018). The fluorescence signals were measured with a Cary Eclipse Fluorescence 147 Spectrophotometer (Agilent Technologies) using excitation/emission wavelengths of 602/632 nm for Raz and 148 571/584 nm for Rru and converted to concentrations based on an 8-point calibration curve (R²=0.99). 149 We monitored specific conductivity (SC) and temperature using Campbell Scientific CS547A sensors 150 connected to Campbell Scientific CR 1000 dataloggers, which recorded and stored those measurements every 10 151 minutes. From the grab samples, we measured chloride using a Dionex ICS-1000 Ion Chromatograph with 152 AS23/AG23 analytical and guard columns. Cl data were augmented with background-corrected SC data to model 153 conservative transport. 154 We monitored changes in stream stage every 10 minutes at the end of the study reach using pressure 155 transducers (Campbell Scientific CS420) connected to a datalogger (Campbell Scientific CR 1000). We used 156 established stage-discharge relationships specific for the study site, as provided by the site managers. The discharge

157 values reported in Table 1 represent mean values observed during a given experiment.

158

2.2 Conservative transport modelling and metrics

We calibrated the conservative transport parameters of the transient storage model presented in Equations 1 and 2 using Cl⁻ and streamwater electrical conductivity data observed at Sites A and B. For this, we used the Matlab (The Mathworks Inc., Natick, Massachusetts) script from Knapp et al. (2018), which features a joint calibration of

162 conservative and reactive solutes through a non-linear, least squares optimization routine.

$$163 \qquad \frac{\partial c}{\partial t} = -u\frac{\partial c}{\partial x} + D\frac{\partial^2 c}{\partial x^2} - \frac{A_s}{A}\frac{\partial c_{ts}}{\partial t} + q_{in}c - \lambda_{mc}c \tag{1}$$

$$164 \qquad \frac{\partial c_{ts}}{\partial t} = k(c - c_{ts}) - \lambda_{ts} c_{ts} \tag{2}$$

- 166 where c [ML⁻³] and, c_{ts} [ML⁻³] are the concentrations in the main channel and aggregate transient storage zone; x
- 167 [L] is the distance of the study reach; t [T] is time; u [LT⁻¹] and D [L²T⁻¹] are parameters representing advective
- 168 flow velocity and dispersion coefficient, respectively; q_{in} [T⁻¹] is a volumetric flux parameter accounting for lateral
- 169 inputs; $k[T^{-1}]$ is the first-order mass transfer rate coefficient parameter between the main channel and the aggregate
- 170 transient storage zone; $A_s/A[-]$ is the capacity ratio parameter representing the relative contribution of transient
- 171 storage-dominated to advection-dominated compartments in the stream, represented as areas along the reach; and
- 172 λ_{mc} and λ_{ts} [T⁻¹] are processing-rate coefficients in the main channel and transient storage zones (equaling zero for
- a conservative tracer).
- 174 We completed the parameter estimation using the Differential Evolution Adaptive Metropolis (DREAM
- 175 [ZS]) algorithm (Vrugt et al. 2009). We jointly fit Cl- and Raz data in a first step of 100,000 model generations. We
- 176 assessed model convergence using Gelman and Rubin \hat{R} statistics (Gelman and Rubin 1992). The goodness of fit
- 177 between measured and simulated BTCs was quantified through the calculation of the residual sum of squares,
- 178 (nRSS) (-), normalized by the squared theoretical peak tracer concentrations of each tracer BTC of the respective
- 179 tracer at the given location. The medians of the best 1,000 model simulations were used to assess the agreement
- 180 between our final model fits and a subset of possible curve fits. The details on the model calibration procedure that
- 181 we use in this work were presented in the supporting information of Gootman et al. (2020). Examples of observed
- and fitted breakthrough curves can be found in Figures S1-S3.
- 183 We estimated conservative transport timescales from the transport parameters to describe the transient 184 storage timescale, $\tau_{sz} = 1/k$ [T], and the mean travel time between sites A and B, τ [T], which was computed as: 185 $\tau = \frac{m_{1,cl}}{m_{0,cl}}$ (3)

186
$$m_n = \sum_{i=1}^r \left(\frac{t_i + t_{i+1}}{2}\right)^n \left(\frac{c_i + c_{i+1}}{2}\right) (t_{i+1} - t_i)$$
(4)

187 where $m_{0,cl}$ and $m_{1,cl}$ are the zeroth and first-centralized temporal moments of the Cl⁻ BTCs from each sampling 188 site, *i* is a time index, *r* is the total number of samples available in a BTC.

189 **2.3** Estimating the transformation of Raz as a proxy for microbial respiration:

190 We used the net transformation rate coefficients of Raz, λ_{Raz} [T⁻¹], as a proxy for microbial respiration, 191 λ_{Raz} incorporates transformation in the main channel and in transient storage zones, and was estimated and 192 estimated them-following the work by González-Pinzón and Haggerty (2013), which who derived algebraic 193 relationships with analytical solutions to calculate processing rate coefficients from the transient storage model 194 presented in Equations 1 and 2:

$$195 \qquad \lambda_{Raz} = \frac{\ln\left(m_{0,Raz}^{inj}/m_{0,Raz}^{BTC}\right)}{\tau} \left(1 + \frac{\underbrace{\frac{dispersion term, \Phi}{\ln\left(m_{0,Raz}^{inj}/m_{0,Raz}^{BTC}\right)}}{Pe}\right) \tag{5}$$

196 where $m_{0,Raz}^{inj} = M_{Raz}/Q$ is the zeroth temporal moment of Raz at the injection site [M L⁻³ T⁻¹], M_{Raz} is the mass of 197 Raz added to the injectate, Q is the stream discharge [L³T⁻¹]; $m_{0,Raz}^{BTC}$ is the dilution-corrected zeroth temporal

- 198 moment of Raz estimated with BTC data from a sampling site; and Pe = L u/D is the Peclet number [-], which
- describes the relative importance of advection and dispersion in the system. As noted by González-Pinzón and
- Haggerty (2013), when Pe>>10, which is the case in advection-dominated systems such as open channel flow, the
- 201 <u>dispersion term</u> Φ is negligible and $\lambda_{Raz} \approx \ln (m_{0,Raz}^{inj}/m_{0,Raz}^{BTC})/\tau$.
- 202 Since we can only get one processing<u>transformation</u>-rate coefficient from every observed BTC available 203 from Equation (5), or from the direct calibration of the transient storage model, we expanded the work by González-204 Pinzón and Haggerty (2013) to incorporate the conceptual principles proposed in the<u>used the</u> Tracer Addition for 205 Spiraling Curve Characterization (TASCC) framework (Covino et al. 2010b), where multiple rate coefficients can 206 be estimated from an equivalent version of Equation 5.
- 207 Briefly, TASCC uses the dynamic range of solute concentrations sampled in BTCs to characterize uptake 208 kinetics over the range of experimental concentrations observed from ambient to saturation concentrations. In 209 TASCC, the ratio of reactive to conservative solute concentrations for every independent sample across the tracer 210 BTCs is compared to the ratio of the concentrations of the injection solution to determine uptake metrics. If the 211 added solutes are non-reactive, they will transport conservatively, and the ratio of the reactive to conservative solute 212 concentrations will remain constant. Alternatively, if the added solutes are limiting, co-limiting or reactive, they will 213 not transport conservatively, and the ratio of the reactive to conservative solute concentrations will change over time 214 as a function of reactivity.
- To incorporate the TASCC framework into the algebraic equation developed by González-Pinzón and Haggerty (2013) and estimate transformation rate coefficients for Raz from each pair of conservative (i.e., $C_{cons.}$) and reactive tracer concentrations (i.e., C_{Raz}), we need to replace m_0 with $C_{Raz}/C_{cons.}$ This guarantees that the mean value of all the processing rate coefficients is equal to the processing rate coefficient estimated from the zeroth temporal moment analysis of model derived simulations from Equations (1) and (2).TASCC-based transformation rate coefficients for Raz were estimated using Accordingly:

221
$$\lambda_{Raz,sample} = \frac{\ln\left[\frac{C_{Raz}}{C_{cons.}}\right]_{inj} - \ln\left[\frac{C_{Raz}}{C_{cons.}}\right]_{BTC}}{\frac{fx}{u}} \left(1 + \frac{\frac{dispersion term,\Phi}{\ln\left[\frac{C_{Raz}}{C_{cons.}}\right]_{inj}} - \ln\left[\frac{C_{Raz}}{C_{cons.}}\right]_{BTC}}{Pe}\right)$$

(6)

222

223 Equation 6 directly links different transport mechanisms used to explain the transport and fate of solutes 224 (i.e., advection, dispersion, transient storage, and reactivity) with TASCC, an algorithm vielding higher information 225 content from experimental work. We note here that alternative forms of Equation 6 can be derived for solute 226 transport models, including additional reactions such as sorption and production. Therefore, similar new equations 227 could be derived to provide mechanistic explanations to TASCC related findings noticing hysteresis behavior in 228 nutrient uptake between the rising and falling limbs of experimental BTCs (Gibson et al. 2015; Trentman et al. 229 2015; Rodríguez-Cardona et al. 2016; Brooks et al. 2017; Day and Hall 2017). Finally, fFrom each transformation rate coefficient λ_{Raz} and $\lambda_{Raz,sample}$, we also estimated an uptake (or mass transfer) velocity of Raz, $V_{f_{Raz}} = \lambda_{Raz}h$ 230 <u>or</u> $V_{f_{Raz,sample}} = \lambda_{Raz,sample} - h$, where h is the mean depth of the stream. Following Ensign and Doyle (2006), 231

- 232 uptake velocities represent the vertical velocity of solute molecules through the water column towards the benthos
- 233 and are typically used in stream ecology to normalize processing-rate coefficients by the influence from contrasting
- 234 discharge magnitudes to facilitate the comparison of results from small streams and large rivers. As demonstrated in

235 <u>Covino et al. (2010b), the range of $\lambda_{Raz,sample}$ and $V_{f_{Raz,sample}}$ values encompass the λ_{Raz} and $V_{f_{Raz}}$ values</u>

- 236 obtained from processing rates derived from temporal moments analyses (e.g., Equation (5)).
- 237 Finally, reach-scale Damköhler numbers, *Da* [-], were calculated using the following equation: $Da = \frac{\text{transient storage timescale}}{\text{transformation timescale}} = \tau_{sz} \lambda_{Raz}.$ 238 (7)

239 2.4 Statistical tests

240 We calculated standard deviations (std) based on repeated measures of the distribution of the transport 241 parameters of Equations 1 and 2 to create upper and lower boundaries of the uncertainties in our measurements (i.e., 242 mean \pm std). Because our data were not normally distributed, we used the Mann-Whitney U nonparametric statistical 243 test to determine if there were statistically significant differences between nutrient treatments across rounds (e.g., N 244 vs. N in rounds 1 and 2), following a similar procedure in Ensign and Doyle (2006). For the Mann-Whitney U test, 245 we set our significance level (α , alpha) equal to 0.05. 246 We explored the Pearson correlation coefficient (r) matrix between the transport parameters of Equations 1

- 247 and 2, and associated metrics, to establish direct (r > 0.1), inverse (r < -0.1), and non-existent correlations (-0.1 < r) 248 <0.1) (Bowley 2008). We classified the strength of the correlations as uncorrelated ($0 \le r \le |0.1|$), weakly correlated 249 (|0.1| < r < |0.5|), moderately correlated (|0.5| < r < |0.8|), strongly correlated (|0.8| < r < |1.0|), and included p-values for
- 250 each correlation.

251 Lastly, we tested differences in mean values of the transport parameters of Equations 1 and 2, and

- 252 associated metrics, between nutrient treatments within each experimental round (e.g., N vs. N+C vs. N+P vs.
- 253 C+N+P in round 1) using the Student's t-test based on deviation from the group's mean value (Blair et al. 1980).

254 **3** Results and Discussion

255 **3.1** Conservative transport and metrics of physical controls

256 Between experimental rounds 1 and 2, stream depth (h) and discharge (Q) decreased, causing significant 257 differences in stream velocity (u), dispersion (D), mass-transfer rate coefficients (k), transient storage time scales 258 (τ_{TS}) and mean travel times (τ) (Figure 2). The only parameter that did not show significant differences was the 259 relative contribution of the main channel to storage zone areas, A_s/A .

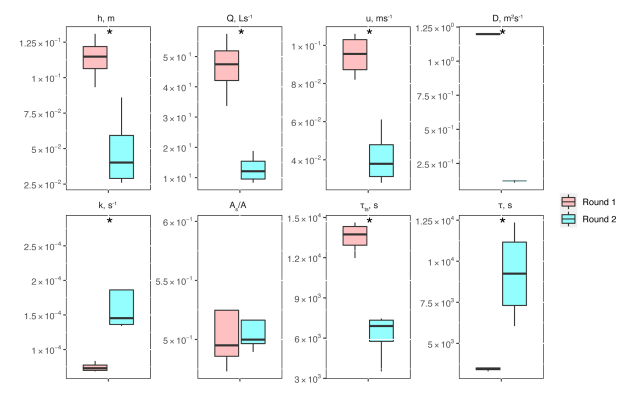
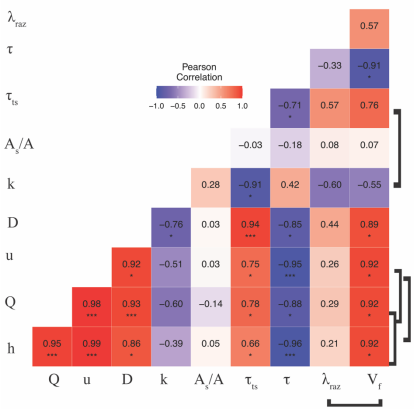
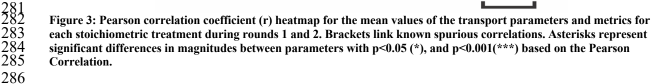


Figure 2: Conservative transport parameters and metrics of physical controls estimated for the two experimental rounds: stream depth (*h*), stream velocity (*u*), dispersion (*D*), mass transfer rate coefficients (*k*), the ratio of transient storagedominated to advection-dominated compartments (A_s/A), transient storage time scales (τ_{TS}) and mean travel times (τ). Asterisks represent statistical differences in magnitudes for rounds 1 and 2 with p<0.05 (*) based on the Mann-Whitney U nonparametric statistical test.

268 The correlation matrix between parameters and metrics (Figure 3) shows that Q (and interrelated quantities 269 h and u), D, and τ_{ts} were all directly correlated (from moderately to strongly). Mean travel times between sites, τ , 270 were directly and weakly correlated with k and the ratio A_s/A , and inversely correlated (from weakly to strongly) 271 with the rest of the conservative transport parameters and metrics. Finally, the ratio A_s/A was generally uncorrelated 272 or weakly correlated with other quantities. Even though the correlations of some interdependent quantities are 273 known to be spurious, e.g., Q vs. u and λ_{Raz} vs. $V_{f_{Raz}}$ (González-Pinzón et al. 2015), we included all relevant 274 measured and modeled quantities in Figure 3 to allow readers to explore different data pairs. For clarity, we 275 differentiate with brackets all known spurious correlations. Note that we did not flag the correlation between A_s/A 276 and Q (and their interrelated quantities h and u) as spurious because the ratio of areas is an indicator of the relative 277 volume-based contribution from advection-dominated to transient storage-dominated compartments, instead of 278 actual estimates of cross-sectional areas (Kelleher et al. 2013; González-Pinzón et al. 2013; Knapp and Kelleher 279 2020).





287 One of the metrics of interest in stream reactive-transport modeling is the transient storage timescale (τ_{ts} = 288 1/k, which quantifies the exposure that solutes have to biological communities in metabolically active transient 289 storage zones. In our study site, τ_{ts} decreased one order of magnitude from round 1 to round 2, and were comparable 290 to the range of values observed in other studies involving forested mountain streams (Valett et al. 1996; Hall et al. 291 2002). Due to the geomorphology of the stream, which is characterized by pool and riffle sequences, but steep 292 longitudinal and valley slopes and shallow bedrock, transient storage was expected to occur mainly in the main 293 channel (Fields and Dethier 2019; Barnhart et al. 2021; Emanuelson et al. 2022). As flow receded from round 1 to 294 round 2, we observed the disconnection of in-stream pools contributing to transient storage, which explains the 295 direct correlation between discharge and transient storage timescales. Another indication of the dominant 296 contribution of in-stream pools to total transient storage is the lack of change of A_s/A with discharge. Since A is 297 expected to vary proportional with discharge (i.e., $Q = A \cdot u$), a constant A_s/A suggests that the contribution of 298 transient storage-dominated (i.e., A_s) compartments (i.e., A) also varied proportionally with discharge.

299 **3.2** Raz transformation (a proxy for respiration) as a function of physical controls

300 Our results indicate that the mean values of the transformation rate coefficient of Raz (λ_{Raz}) were directly 301 and moderately correlated with the transient storage timescale (τ_{ts}), as other studies on reactive transport have

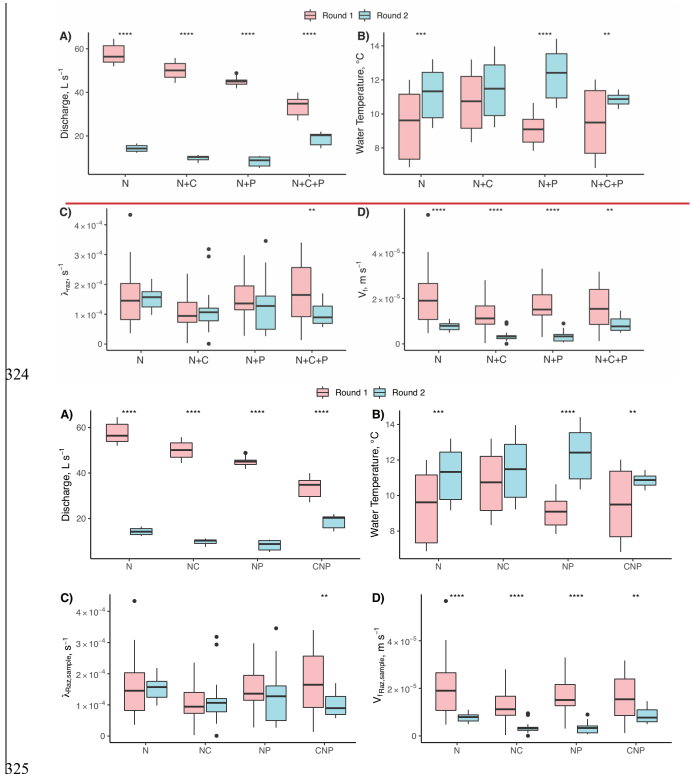
- 302 shown (Valett et al. 1996; Hall et al. 2002; Gomez et al. 2012; Zarnetske et al. 2012; Kiel and Bayani Cardenas
- 303 2014; Gootman et al. 2020). Mean λ_{Raz} values were directly and weakly correlated with discharge (Q) (also depths
- 304 *h* and velocities *u*) and dispersion (*D*), and directly and moderately correlated with τ_{ts} . Mean λ_{Raz} values were
- inversely and weakly correlated with mean travel times (τ), and inversely and moderately correlated with mass-
- 306 transfer rate coefficients (k) (Figure 3). Raz uptake velocities $(V_{f_{Raz}})$ showed spurious, direct and strong
- 307 correlations with discharge (Q) (also h and u), strong correlations with dispersion (D) and transient storage
- 308 timescales (τ_{ts}), and strong indirect correlations with mean travel times (τ) and k (moderate). Finally, both λ_{Raz} and
- 309 $V_{f_{Raz}}$ were uncorrelated with A_s/A . Unlike studies where an increased transient storage timescale (τ_{ts}) is mainly
- 310 associated with slower hyporheic flows due to lower discharges (Q) (Zarnetske et al. 2007; Schmid et al. 2010), τ_{ts}
- 311 in our study site increased with *Q* because the geomorphology of the channel and the valley favored in-stream
- transient storage in lateral pools (Jackson et al. 2012, 2013, 2015). Similar declines in transient storage with falling
- 313 discharge have been observed in other streams with comparable geomorphic characteristics (Covino et al. 2010a;
- Emanuelson et al. 2022), however, the absence of concurrent declines in respiration suggest biological control by
- 315 some other mechanism.

316 **3.3 Raz transformation (a proxy for respiration) as a function of physical and stoichiometric controls**

317 Our results suggest no significant changes in respiration despite significant differences in discharge (Q),

temperature, and nutrient treatments. Between experimental rounds, the mean values of Q (and h and u by

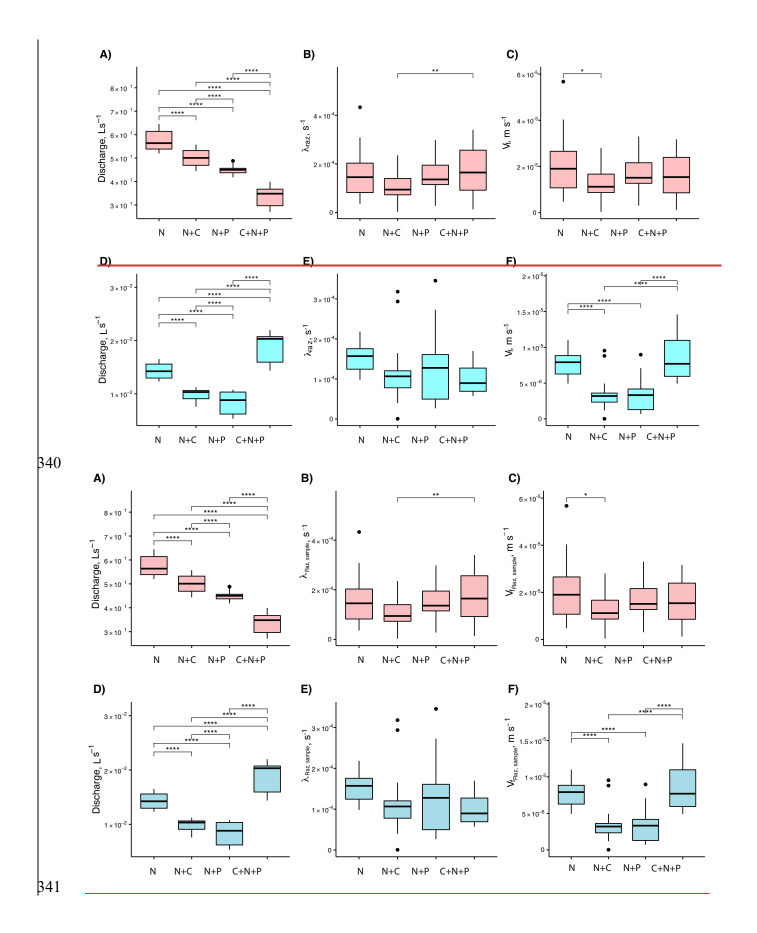
- 319 extension) and temperature (except for N+C) were statistically different for each treatment comparison (Figure 4A).
- B20 For $\lambda_{Raz,sample}$, we only found statistical differences between rounds for the C+N+P treatments (Figure 4C). Due to
- 321 the large influence of Q on the uptake velocity of Raz $(V_{f_{Raz,sample}})$ through stream depth (h), the statistical
- differences between rounds seen for Q were also seen for $V_{f_{Raz,sample}}$ (Figure 4D).
- 323





325 326 327 328 329 330 Figure 4: Comparison of A) stream discharge values recorded at the gaging station, B) stream water temperatures, C) transformation rate coefficients of resazurin ($\lambda_{Raz,sample}$) resulting from Equation 6, and associated D) uptake velocities of resazurin ($V_{f_{Raz,sample}} = \lambda_{Raz,sample} h$) estimated for each experimental nutrient treatment addition during rounds 1 and 2. Due to the large influence of Q on the uptake velocity of Raz ($V_{f_{Raz,sample}}$) through stream depth (h), most of the statistical differences between rounds seen for Q were also seen for $V_{f_{Raz,sample}}$. Asterisks represent significant

differences in magnitudes between rounds with p<0.01(**), and p~0 (****) based on the Mann-Whitney U nonparametric statistical test. When looking at the data collected from each round, we found that mean Q values were statistically different across nutrient treatments (Figures 5A and 5D). For mean $\lambda_{Raz,sample}$ values, the only treatments with statistical differences were the N+C and C+N+P from round 1 (Figures 5B and 5E). Finally, $V_{f_{Raz,sample}}$ mean values were only statistically different for the N vs N+C treatments for round 1, and for all but the N+C vs N+P and N vs C+N+P treatments for round 2 (Figures 5C and 5F).



- Figure 5: Comparison of stream discharges (A and D), transformation rate coefficients of resazurin ($\lambda_{Raz,sample}$) resulting from Equation 6 (B and E), and associated uptake velocities of resazurin ($V_{f_{Raz,sample}}$) (C and F) across treatments for round 1(top row) and 2 (bottom row). Due to the large influence of Q on the uptake velocity of Raz ($V_{f_{Raz,sample}}$) through stream depth (h), most of the statistical differences between rounds seen for Q were also seen for $V_{f_{Raz,sample}}$. Asterisks represent significant differences in magnitudes for treatments N, N+C, N+P, and C+N+P with p<0.05 (*), p<0.01(**), and p~0 (****) based on the Mann-Whitney U nonparametric statistical test.
- 348 349

For each of the eight nutrient injections, we related the mean transient storage timescales, τ_{ts} , which

- 350 indicate exposure times between solutes and microbial communities, and the mean transformation timescales of Raz,
- $1/\lambda_{Raz}$, which indicate respiration (Figure 6). This Damköhler-based analysis allows us to visualize the interplay
- between physical, biological, and stoichiometric controls in the stream. We found that the range of variation of the
- 353 mean transient storage timescales was three times greater than that of the mean transformation timescales. In round
- 1, all the stoichiometric treatments resulted in transport-limited conditions due to the high values of τ_{ts} , i.e., the
- 355 average particle of Raz that entered a metabolically active compartment underwent transformation and more Raz
- 356 could have been transformed if it had been available. Thus, in round 1, respiration was high relative to the supply of
- 357 solutes to the metabolically active transient storage zones. In round 2, all stoichiometric treatments, except N,
- 358 resulted in reaction-limited conditions, i.e., the average particle of Raz entering a metabolically active compartment
- 359 left it without undergoing transformation. Thus, in round 2, respiration was slow relative to the exposure of solutes
- to microbial communities.
- 361
- 362

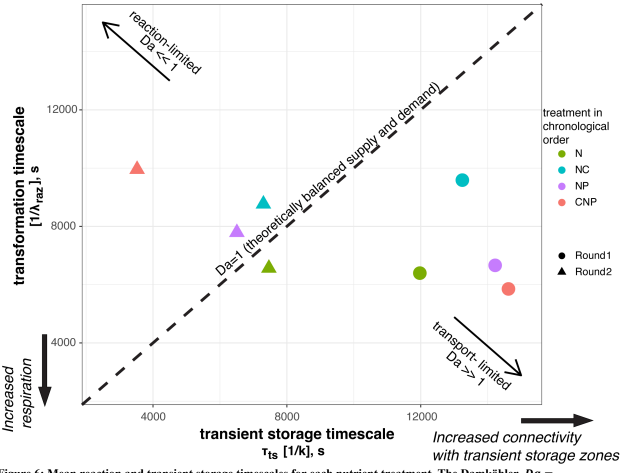


Figure 6: Mean reaction and transient storage timescales for each nutrient treatment. The Damköhler, *Da* =
 transient storage timescale/ transformation timescale, indicates reaction-limited and transport-limited conditions.

367 3.4 How is microbial respiration controlled by hydrologic exchange vs. stoichiometric conditions (i.e., supply 368 of C, N, and P)?

369 We characterized microbial respiration with the transformation timescale of Raz, $1/\lambda_{Raz}$; the extent of 370 hydrologic exchanges with the transient storage timescale, τ_{TS} , and the relative size of the main channel and 371 transient storage areas, A_s/A ; and stoichiometric conditions with our controlled nutrient additions (i.e., N, N+C, 372 N+P, and C+N+P treatments). The most salient findings indicate that a) discharge (Q) changed significantly 373 between rounds (Figure 4a) and across stoichiometric treatments (Figure 5a, 5d), and was directly and moderately 374 correlated with τ_{TS} and uncorrelated with A_s/A (Figure 3), suggesting that most transient storage occurred in lateral 375 pools in the channel, which increased in quantity and extent proportionally with Q, and b) the respiration activity 376 indicated by λ_{Raz} remained similar between rounds with significantly different Q (Figure 4b), and across controlled 377 stoichiometric treatments also featuring different Q (Figure 5b, 5e). Thus, we observed that respiration remained 378 largely unchanged or constant with varying physical and stoichiometric conditions.

379 Several hypotheses may explain the invariant respiration observed between experimental rounds and
 380 treatments. First, tradeoffs in metabolic rates may have occurred as the stream shifted from high to low flows. At

381 high flows during late June and early July, lateral pools in the main channel were inundated, and transient storage

- 382 timescales likely associated with these pools were high. Under these conditions, the observed respiration was
- 383 probably supported by low levels of processing in the hyporheic zone due to the prevalence of bedrock substrate and
- relatively low respiration from benthic biomass due to scour from high flows (Francoeur and Biggs 2006; Katz et al.
- 385 2018). However, the combination of longer transient storage timescales and an expanded total surface area resulted
- in moderate total respiration. In contrast, during the low flows seen in the second round of injections, surface area,
- 387 and transient storage timescales were decreased due to the contraction of the channel. Under these conditions,
- biomass increased likely due to decreased scour and increased stability (Francoeur and Biggs 2006; Katz et al. 2018;

Cargill et al. 2021), increased water temperatures (Perkins et al. 2012), and increased processing of autochthonous

- 390 carbon (Wagner et al. 2017) (Figure S4). This may have supported elevated areal metabolic rates in benthic biofilms
- 391 (Battin et al. 2016), maintaining relatively constant respiration levels with respect to the first round of injections.
- 392 An alternative hypothesis to explain the consistency of the observed respiration values is that some other 393 factor constraints respiration values within a narrow range. For example, the limitation of a key nutrient or metabolic 394 resource may constrain respiration. While we designed the experiments to relieve stoichiometric constraints, it is 395 possible that the quantities of C, N, and P in the injectate we were logistically able to introduce to the stream were 396 insufficient to overcome demand. Also, the form of the resources may not have been readily available to 397 communities adapted to these locals, as stream microbial communities most efficiently process the forms and 398 diversity of dissolved organic matter found in their native habitats, and they express extracellular enzymes in ratios 399 appropriate to acquire limiting nutrients (Hill et al. 2012; Lane et al. 2012; Wilhelm et al. 2015; Logue et al. 2016). 400 In previous studies, transient storage and nutrient uptake have presented contradictory relationships, which
- 401 we summarize below.

- 402 Inconclusive relationships: Martí et al. (1997) did not find correlations between NH₃ uptake length and 403 A_s/A in a desert stream using data from eight tracer injections. Webster et al. (2003) did not find statistically 404 significant relationships between NH4 uptake and As/A using the 11-stream LINX-I dataset that included arctic to 405 tropical streams. From thirty seven injections conducted in thirteen streams at Hubbard Brook Experimental Forest 406 (HBEF), Hall et al. (2002) found weak correlations (R²=0.14-0.35) between transient storage parameters and NH₄ 407 demand. Using data from seven streams in New Zealand, Niyogi et al. (2004) did not find significant correlations 408 between soluble reactive phosphorous (P-SRP), NO₃ uptake velocities, and A_s/A. Bukaveckas (2007) reported an 409 indefinite relationship between transient storage and NO3 and P-SRP retention efficiencies from tracer injections in a 410 reference (N=13 injections), a channelized (N=14 injections), and a restored (N=17 injections) stream reach in the 411 midwestern US. Lastly, the LINX-II dataset from ¹⁵N-NO₃ injections in 72 streams located in eight regions of the 412 US showed no relationship between NO₃ uptake and the fraction of median travel time due to transient storage 413 (F_{med}^{200}) (Hall et al. 2009).
- 414 *Weak to moderate relationships:* Thomas et al. (2003) showed that transient storage accounted for 44% to 415 49% of NO₃ retention measured by ¹⁵N in a small headwater stream in North Carolina. Mulholland et al. (1997) 416 found larger PO₄ uptake rates in a stream with higher transient storage, when they compared two forested streams. 417 Ensign and Doyle (2005) found an increase in A_s/A and the uptake velocities for NH₄ and PO₄ after the addition of

418 flow baffles to two streams. Lautz and Siegel (2007) found a modest correlation (R²=0.44) between NO₃ retention

- 419 efficiency and transient storage in the Red Canyon Creek watershed, WY.
- 420 *Strong relationships:* Valett et al. (1996) found a strong correlation ($R^2=0.77$) between transient storage
- 421 and NO₃ retention in three first-order streams in New Mexico. From nine tracer injections in two urban streams in
- 422 the eastern US, Ryan et al. (2007) found strong relationships between P-SRP retention and transient storage metrics
- 423 $(k, A_s/A; \mathbb{R}^2 > 0.84)$ when the variables were measured in different seasons. Sheibley et al. (2014) observed that the
- 424 retention of NO₃ in seven agricultural streams in the US was positively correlated with A_s/A and the average water
- 425 flux through the storage zone per unit length of stream ($q_s = kA$), and negatively correlated with the transient
- 426 storage timescale (τ_{ts}). However, they found no significant correlation between NH₄⁺ and SRP retention and 427 transient storage metrics.
- 428 The studies referenced above were performed in streams with contrasting physical, chemical, and 429 biological conditions. Together, they offer a broader perspective on the inconsistent relationship between transient 430 storage metrics and metabolic processing. Those studies do not feature co-injections of C, N, and P macronutrients 431 (e.g., N+C, N+P, N+C+P), even while some tracked ambient processing rates of more than one nutrient. Therefore, 432 they generally represent solute-specific analyses, where the uptake of one nutrient at a time was analyzed and, thus, 433 did not account for stoichiometric controls on nutrient uptake (however, see Tromboni et al. (2018) for an example 434 of recent trend changes in this research area). By combining both transport and stoichiometric analyses, our study 435 offers evidence that stoichiometric controls have an ambiguous relationship to reach-scale metabolic activities, and 436 that further investigations should be conducted using greater quantities and types of resources.

437 4 Conclusions

438 We conducted two rounds of four stoichiometric treatments (i.e., N, C+N, N+P, and C+N+P) in a 439 headwater stream in Colorado to quantify changes to stream respiration during flow recession and answer the 440 question: How is respiration controlled by hydrologic exchange vs. stoichiometric conditions (i.e., supply of C, N, 441 and P)? We found that discharge changed significantly between rounds and across stoichiometric treatments, and 442 that it was directly and moderately correlated with transient storage timescales but uncorrelated with the ratio of 443 contributions from advection-dominated to transient storage-dominated compartments (i.e., A_s/A). This suggests 444 that most transient storage occurred in lateral pools within the main channel, which increased in quantity and extent 445 proportionally with discharge. We also found that respiration remained similar despite significant changes in 446 discharge and stoichiometric treatments. Our results contradict the notion that hydrologic transport alone is a 447 dominant control on biogeochemical processing, and suggest that complex interactions between hydrology, resource 448 supply, and biological community function are responsible for driving in-stream respiration.

449 Author contribution: RGP, TC, KS, and MG secured the funding for this research. All co-authors designed carried
 450 out the experiments. JD and RGP processed Raz samples, performed solute transport simulations, statistical analyses,

- 451 and prepared the manuscript with input from all co-authors. DVH supported the contextualization of hydrological and
- 452 ecological interactions. All co-authors approved the final version of the manuscript.
- 453 **Competing interests:** The authors declare no competing interests.

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458 **Data availability:** The data used in his article can be found in the CUAHSI HydroShare repository. Gonzalez-Pinzon,
459 R. (2022). Resazurin tracer data from experiments in Colorado (2018) and Iowa (2019),
460 HydroShare, http://www.hydroshare.org/resource/50ae3c59bebe4cb383e31408a0c10012

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