Contrasting Raz–Rru stream metabolism and nutrient uptake downstream of urban wastewater effluent sites

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Abstract: Understanding how key stream ecosystem functions respond to wastewater treatment plant effluent is critical for assessing the ability of stream ecosystems to ameliorate anthropogenic nutrient loading and to effectively manage and restore impacted systems. We evaluated instream metabolism, reactive solute transport, and nutrient uptake along two 1.5 to 2-km-long reaches of a 2nd-order stream in the urbanized suburbs of Philadelphia, Pennsylvania, USA, each directly downstream of a wastewater treatment plant outfall. We compared metabolism based on resazurin–resorufin (Raz–Rru) with nutrient uptake and dissolved oxygen (DO) metabolism calculations. Plateau co-injections of the Raz–Rru metabolic tracer system and fluorescein provided integrated stream metabolism measurement. We sampled tracer concentrations hourly along longitudinal profiles and recorded them continuously, along with DO, at 2 discrete locations. The smaller reach 1, characterized by higher nutrient concentrations and canopy cover, had higher short-term transient storage and Raz uptake velocity. In contrast reach 2, with lower nutrient concentrations and less canopy cover, had higher nitrate and phosphorus uptake along with higher rates of gross primary productivity (GPP) and ecosystem respiration (ER). Temporal analysis indicated nitrate uptake increased over the afternoon at reach 2, whereas Raz uptake declined at both reaches. Our results suggest that nutrient uptake and GPP are sensitive to excessive nutrient concentrations and light in our system. In contrast, lower light and higher transient storage are likely driving larger, reach-scale spatial differences in Raz-based ER. Increasing nitrate uptake at reach 2, which lags behind diel DO concentrations, is likely the result of assimilatory N uptake coupled to GPP moderated by nitrification and denitrification, whereas decreasing rates of Raz transformation are likely related to diel variation in heterotrophic uptake. Our ability to resolve sub-daily changes in ER illustrates one of the key advantages of the Raz–Rru tracer system. However, our results also show the need for further investigation into the drivers of sub-daily ecosystem metabolism in streams as well as the mechanistic differences between DO- and Raz-based estimates of ER. Contrasting results from different measures of metabolic activity between reaches and over time highlight the complexity of metabolic processes in high-nutrient systems.

Key words: instream metabolism, wastewater treatment plant, nutrient uptake, nitrate, total dissolved phosphorus, urban stream, resazurin

Anthropogenic inputs of nutrients to streams, including effluent from wastewater treatment plants (WWTPs), are a major driver of excessive nutrient loading and impaired water quality in streams worldwide. Increased inputs of nutrients to streams lead to increased algae growth, decreased dissolved oxygen (DO), and fish mortality (Dodds and Welch 2000, Smith 2003, Carey and Migliaccio 2009). In the United States (US), hypoxic zones in receiving waters continue to grow despite efforts throughout the country to curb nutrient inputs (van Meter et al. 2018). Although understanding the impacts of both point and non-point sources is required to fully quantify nutrient budgets, understanding WWTP effects...
is particularly important because they contribute to large nutrient loads in receiving streams. Despite improvements in treatment technology, US wastewater effluent that has gone through biological-nutrient-removal tertiary treatment typically still contains 3 to 8 mg/L of total nitrogen (TN) and 1 to 2 mg/L of total phosphorus (TP), and upgrading plants to further remove nutrients before discharge is cost-prohibitive (Carey and Migliaccio 2009). The fate of WWTP-derived nutrients in streams and their influence on stream ecosystem function require further study.

The addition of nutrients and organic carbon, as well as changes in nutrient uptake rates, in WWTP-impacted streams can alter magnitude and patterns of ecosystem respiration (ER) and gross primary production (GPP). Gücker et al. (2006) found ER rates as high as ~59 g O₂ m⁻² d⁻¹ downstream of WWTPs in German streams. Aristi et al. (2015) saw increases in ER downstream of WWTPs, whereas Wassenaar et al. (2010) found that the ER response downstream from WWTPs varied with season. GPP responses to effluent are more variable than ER responses, with observed rates as high as 59 g O₂ m⁻² d⁻¹ downstream of WWTPs (Gücker et al. 2006). This variation in response to effluent may also be driven by changes in other environmental conditions besides nutrient concentrations, similar to what has been seen in higher-nutrient urban streams without effluent. In those cases, change in riparian shading was an important driver in longitudinal GPP change (Ledford et al. 2017, Reisinger et al. 2019).

Although WWTP effluent increases nutrient concentrations in streams, the efficiency of nutrient uptake (i.e., concentration relative to flux) typically declines, driven by saturation kinetics (Dodds et al. 2002, Mulholland et al. 2008). Uptake lengths of N can increase downstream of WWTPs (i.e., less uptake), with some studies finding almost ½ of the studied streams showing no net nutrient retention (Martí et al. 2004, Gibson and Meyer 2007, Figueroa-Nieves et al. 2016). Another study found that nitrate uptake efficiencies did not have a uniform response to discharge and increased nutrient loads from 2 WWTPs (Gücker et al. 2006). Downstream of 1 plant, uptake rate and uptake velocity did not change, whereas the other plant showed large increases in both variables. Nutrients from WWTP effluent can be taken up via assimilatory and dissimilatory pathways. Assimilatory pathways result in the incorporation of nutrients into biomass, whereas dissimilatory pathways, including denitrification, include the use of nutrients as electron acceptors, changing the molecular form of the nutrient (Burgin and Hamilton 2007, Burgin et al. 2011).

Many metabolic processes, including assimilatory and dissimilatory uptake, vary with daily frequency, although the timing of these processes, and of resulting changes in stream solute concentrations, are not always predictable. For example, daytime autotrophic nitrate uptake, which is coupled to GPP, can lead to lower nitrate concentrations during the day relative to a nighttime peak. In contrast, assimilatory P uptake and denitrification do not have consistent diel cycles (Heffernan and Cohen 2010, Nimick et al. 2011 and references therein, Cohen et al. 2013), and we know very little about diel variations in heterotrophic activity or uptake. Resolving diel signals and process dynamics in streams receiving WWTP effluent is further confounded by the occurrence of diel shifts in effluent quantity and quality and diel variations in nutrient demand and uptake (Gammons et al. 2011). Rahm et al. (2016) measured rapid uptake of nutrients downstream of a WWTP in central New York, USA, and although they could not determine the pathway, molecular assays indicated that denitrifying communities were being released in the effluent. The molecular form of N in the effluent understandably plays a large role in its fate.

Understanding how ecosystem functioning responds to WWTP effluent inputs (enhanced or inhibited) and what controls spatial and temporal variability in response (nutrient loading, effluent magnitude, hydrologic variability, metabolically active transient storage) is critical to assessing the capacity of stream ecosystems to ameliorate anthropogenic nutrient loading. These processes are all theoretically linked, and previous studies have compared stream metabolism (DO or resazurin–resorufin [Raz–Rru] based) and nutrient cycling in WWTP-impacted systems (e.g., Gücker and Pusch 2006, Aristi et al. 2015, Ribot et al. 2019) and DO-based stream metabolism and Raz transformation (e.g., González-Pinzón et al. 2012, 2014, 2016, Kurz et al. 2017, Knapp and Cirpka 2018). Raz-transformation rates have not been reported previously in a WWTP-impacted stream system, although Raz has been applied to study the metabolic activity of WWTP-derived sludge (Strotmann et al. 1993, McNicholl et al. 2007) and the metabolic response of WWTP-impacted flume biofilms to labile carbon (Ribot et al. 2019).

The complexity of metabolic processes in time and space suggests that applying methods that resolve different processes over different scales is important to unravelling the key drivers of coupled metabolic processes in WWTP-impacted systems. Our goals in this study were: 1) to compare metabolic processing of effluent inputs from 2 WWTPs on the same stream, 2) to assess if hourly variations in metabolic activity occur in response to nutrient or temperature variation over the course of an afternoon (~5 h), and 3) to compare the results of 3 methods measuring rates of whole-stream metabolic processes (Raz–Rru transformation, nutrient uptake, and DO-based GPP and ER). We hypothesized that: 1) the reach with a higher portion of WWTP input (and higher nutrient concentrations) would have lower rates of metabolic activity because of nutrient saturation; 2) metabolic rates would increase through the afternoon as nutrient concentrations and temperature rose as part of a diurnal effluent cycle; and 3) spatial and temporal patterns in metabolic parameters, including nutrient uptake, aerobic respiration, and whole-stream ER and GPP, would vary consistently between reaches as measured through
the 3 methods of metabolism evaluation (Raz–Rru transformation, nutrient uptake, and DO-based GPP and ER).

**METHODS**

**Site description**

Wissahickon Creek is a 3rd-order stream at its outlet with the Schuylkill River and flows through Montgomery and Philadelphia counties, Pennsylvania, USA (Fig. 1A). It is a 166-km² suburban-to-urban watershed with a population of 222,000 (CSC 2014). Four WWTPs are permitted to discharge into the stream: 2 on the mainstem and 2 on Sandy Run Tributary (Fig. 1A). The stream is listed as impaired because of nutrients, nuisance algae, and sediments (USEPA 2003). Although 51% of the watershed has impervious or low-pervious land cover, the corridor along the Wissahickon has benefited from open space protection. About 75% of the upper watershed has a forested riparian buffer on 1 or both sides, with that number increasing to 80% for the entire watershed because of forest cover in Fairmont Park, Philadelphia (Heritage Conservancy 2012).

This paper focuses on the upper half of the mainstem of the creek where the uppermost 2 WWTPs discharge. We chose these 2 sites because they have differing % of flow coming from the WWTP, resulting in differing nutrient concentrations downstream of their respective outfalls. In the headwaters, Upper Gwynedd WWTP effluent contributes ~70 to 80% of discharge during baseflow downstream of this plant (reach 1; Fig. 1B). Grab samples show the effluent typically has 20 to 30 mg N/L as nitrate and 0.25 to 0.35 mg total dissolved phosphorus (TDP)/L. Effluent discharge generally exhibits a daily pattern of a morning rise, daytime plateau, and overnight fall. Previous longitudinal grab samples showed a downstream decrease in nitrate for 10 km downstream of the plant but did not show much change in TDP concentration over the same distance, potentially indicating different processing controls for the 2 nutrients. Reach 1 is ~2 km in length, has an average wetted width of 8.4 m, and an average depth of 22 cm (Table 1). This width combined with mature trees in the riparian zone results in a mean riparian canopy cover of 65% along this reach (Fig. S1A). Similar high nutrient loads and longitudinal changes in nutrients have been observed in the downstream reaches below 2 of the other WWTPs in the watershed (Ledford and Toran 2020). In contrast, neither nitrate nor phosphorus show a decrease in concentration downstream of Ambler WWTP, which is ~10 km downstream.
of the Upper Gwynedd WWTP (Fig. 1C). The Ambler WWTP effluent contributes ~40 to 50% of the total baseflow downstream of the plant (reach 2). Effluent nutrient concentrations range from 15 to 25 mg N/L of nitrate, and TDP concentrations are highly variable, ranging from 0.2 to 1.5 mg TDP/L. Effluent discharge typically peaks in the morning and evening, with falling discharge during the early afternoon and overnight. Reach 2 is ~1.5 km in length leading to a mean 55% canopy cover (Fig. S1B).

### Experimental design

To measure the impact of WWTP effluent on stream ecosystem function, we used 3 independent measures of whole-stream metabolic processing at various spatial and temporal resolutions in the 2 study reaches. First, we used plateau injections of the reactive tracer Raz and calculated longitudinal estimates of aerobic respiration and the relative influence of metabolically active transient storage to whole-stream metabolism at sub-daily timescales. Raz undergoes an irreversible transformation to Rru under the mildly reducing conditions produced by respiring cells (O’Brien et al. 2000, Haggerty et al. 2008, Knapp et al. 2018). Thus, the rate of Raz–Rru transformation represents an integrated measure of (near-instantaneous) whole-system aerobic ER. The Raz–Rru system has been used broadly to assess the reactivity of stream reaches and to identify the portion of transient storage that is metabolically active (Argerich et al. 2011, González-Pinzón et al. 2012, 2014, Blaen et al. 2018, Ward et al. 2019). Second, we calculated nitrate and TDP retention downstream of the WWTP inputs at sub-daily timescales based on longitudinal declines in concentration (Martí et al. 2004). Third, we calculated daily rates of whole-stream GPP and ER at the midpoint of each study reach using the 1-station diel DO method (Hall and Hotchkiss 2017).

### Stream solute tracer injections

#### Tracer injections

To calculate reach- and sub-reach-scale transient storage and aerobic respiration we used plateau injection tests of conservative and metabolically reactive tracers combined with the nitrate and TDP inputs from the WWTP effluent (Martí et al. 2004). We conducted an instream constant-rate injection of the conservative tracer fluorescein and reactive tracer Raz downstream of both WWTP effluent input points targeting a 4 to 5-h daytime plateau of ~30 ppb fluorescein and ~50 ppb Raz. At reach 1, we mixed 400 g fluorescein and 646 g Raz powder with 100 L of tap water in an opaque mixing drum and injected across the stream width downstream of the WWTP input at a rate of 2.7 to 2.8 mL/s for ~9.5 h (0436–1406h, 26 September 2017; Fig. 1B). Similarly, at reach 2, we mixed 400 g fluorescein and 626 g Raz powder with 100 L of tap water in an opaque mixing drum and injected across the stream width 150 m downstream of the WWTP input at a rate of 3.1 mL/s for a duration of 8 h 33 min (0743–1616h, 12 September 2017; Fig. 1C). We measured the injection flow rate into the stream periodically throughout both injections to account for potential variation, which was negligible.

We evaluated tracer concentrations with both grab samples and data logging sensors. We measured concentration breakthrough curves (BTC) of fluorescein, Raz, and Rru at 10-s resolution with 2 calibrated in-situ GGUN FL30 fluorometers (Albillia, Neuchâtel, Switzerland; Lemke et al. 2014).

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### Table 1. Major structural and transport parameters compared between sites. Bolded values indicate observed higher relative values between reaches. $\mu_2$ (h$^2$) represents the 2nd central temporal moment (temporal variance or symmetrical spreading). $\mu_3$ (h$^3$) represents the 3rd central temporal moment (temporal extent of late-time tailing). $D_{app}$ represents apparent dispersion. CV = coefficient of variation. Ranges that are temporal are indicated with $\text{S}$ and ranges that represent spatial change are indicated with $\text{S}$.

| Parameter                  | Reach 1          | Reach 2          |
|----------------------------|------------------|------------------|
| Discharge (m$^3$/s) (±SE)  | 0.09 (±8%)       | 0.34 (±7.2%)     |
| Velocity (m/s)             | 0.06             | 0.12             |
| Specific discharge (m$^3$/s)| 0.011            | 0.019            |
| Mean depth (m; range)      | 0.22 (0.12–0.28)$^S$ | 0.21 (0.10–0.31)$^S$ |
| Mean width (m; range)      | 8.4 (6.6–11.6)$^S$ | 19.0 (12.1–26.3)$^S$ |
| Reach $\mu_2$ (h$^2$)     | 0.38             | 0.17             |
| Reach $\mu_3$ (h$^3$)     | 0.37             | 0.08             |
| Reach CV                   | 0.20             | 0.22             |
| Reach skewness, $\gamma$  | 1.61             | 1.13             |
| $D_{app}$ (x10$^4$ m$^2$/h)| 3.8              | 1.5              |
| pH                         | 7.73–8.55$^T$    | 8.44–8.66$^T$    |
| Canopy cover (average % open; range) | 35.3 (23.9–48.9)$^S$ | 44.8 (35.1–67.6)$^S$ |
located at 1494 and 2894 m downstream of the reach 1 effluent input and 650 and 1250 m downstream of the reach 2 effluent input (Fig. 1B, C). A 3rd fluorometer measured only fluorescein at 694 and 1750 m in reaches 1 and 2, respectively. We collected longitudinal grab samples for analysis of major anions, cations, nutrients, and the concentrations of fluorescein, Raz, and Rru at hourly resolution during the tracer plateau. We collected hourly samples from 1215 to 1615h (with a final sample at 1700h) at reach 1 every 200 m along the study reach from 494 m downstream of the effluent input (400 m downstream of the tracer injection point) to 2094 m downstream. We collected hourly samples from 1230 to 1630h at reach 2 every 100 m along the study reach from 550 m downstream of the effluent input (400 m downstream of the tracer injection point) to 1650 m downstream. Differences in longitudinal spacing were because the number of sample sites was the same for both tests but the 2 reaches had different distances to the nearest major hydrologic feature (a low-head dam at reach 1 and a tributary at reach 2). We collected additional grab samples at the fluorometer locations pre- and post-plateau for analysis of tracer concentrations to cross-check the calibrated fluorometer BTC concentrations. We field filtered all Raz, Rru, and fluorescein grab samples through a 0.22-µm-glass-fiber filter and stored them at 4°C in amber glass vials until analysis. Within 48 h of collection, we analyzed samples on a Quantamaster™ QM-4-CW lab spectrofluorometer (Photon Technology International, Birmingham, New Jersey). We created manual calibration curves at 490 nm excitation and 503 nm emission for fluorescein (Siejak and Frackowiak 2005), 570 nm excitation and 573 nm emission for resorufin, and 602 nm excitation and 621 nm emission for resazurin (Haggerty et al. 2008). We selected excitation peaks from literature values and chose emission peaks by selecting the peak emission from scans of standards of each dye run at the peak excitation wavelength reported in the literature.

**Transient storage metrics** Transfer functions are representative of sub-reach travel time distribution and reflect both instream and hyporheic transport and associated advective, dispersive, and short-term storage processes. To determine transfer functions for the sub-reaches between fluorometers, we used the nonparametric deconvolution approach described in Cirpka et al. (2007), and the associated MATLAB® code (The MathWorks®, Natick, Massachusetts), to analyze injection input signal and conservative tracer BTCs. To characterize and compare this transport and storage behavior between study reaches, we calculated a series of transport and transient storage metrics for each sub-reach transfer function, following the equations in Ward et al. (2016, 2019) and Schmadel et al. (2016). Metrics included the 1st temporal moment ($M_1$, h), representing the mean travel time; the 2nd central temporal moment ($\mu_2$, h$^2$), representing the temporal variance or symmetrical spread-
We then used the sub-reach depths to calculate the length-weighted mean stream depth of the 2 study reaches.

**Raz–Rru transformation rates** To estimate Raz–Rru transformation rates, we used the longitudinal profiles of Raz and Rru tracer concentrations from the grab samples collected during the tracer plateau, as in Kurz et al. (2017), using the analytical solution:

\[
k_{t,ad} = \left[ \ln\left( \frac{C_{Rru}}{C_{Raz}} + P \right) - \ln\left( \frac{C_{Rru,0}}{C_{Raz,0}} + P \right) \right] t_{ad}^{-1}
\]

(Eq. 1),

where \( k_{t,ad} \) (h) is the apparent Raz–Rru transformation rate coefficient, assuming transformation is a 1st-order process (Haggerty et al. 2008). \( C_{Rru} \) and \( C_{Raz} \) are the concentrations of Rru and Raz (ppb), \( C_{Rru,0} \) and \( C_{Raz,0} \) are the input concentrations of Rru and Raz, \( P \) is the production–decay ratio (unitless), and \( t_{ad} \) is the advective travel time (h), such that \( k_{t,ad} \) is the slope of the regression between travel time and the Raz–Rru transformation rate. By using the Rru:Raz ratio in our calculations we eliminate the need to correct for conservative mass loss or dilution. We assume irreversible sorption of both tracers is minimal given the pH remained above 7.7 at both sites (Table 1) and that any mass losses of Raz or Rru other than the decay of Raz to Rru are minimal or equal between the 2 tracers, therefore, \( P = 1 \). The concentration of Rru at the injection site is theoretically 0, however, when solving for \( k_{t,ad} \) as the slope of \( \ln(C_{Rru} / C_{Raz}) + 1 \) and \( t_{ad} \) do we not enforce an intercept of 0. We interpolated the advective travel time to each sampling station from the downstream distance of each sampling site and the between-fluorometer sub-reach travel times calculated from the fluorometer BTCs. We used the fitlm function in MATLAB to test the slope of each sampling-period regression. \( k_{t,ad} \) is reported as 0 for sampling periods with a p-value above 0.06 (Table S1).

Raz to Rru transformation rates have been shown to be proportional to changes in oxygen, which is a common measurement of respiration rates (González-Pinzón et al. 2012). Therefore, estimated \( k_{t,ad} \) can be used as a proxy of instantaneous rates of ER. To compare Raz-transformation values among reaches with different hydrologic conditions, we used the uptake velocity of Raz (\( v_{t-Raz} \), mm/min), calculated as:

\[
v_{t-Raz} = k_{t,ad} \times d \quad (\text{Eq. 2}),
\]

where \( d \) (m) is the average water depth. \( v_{t-Raz} \) can be thought of as the effective velocity of the transformation of Raz into Rru (Haggerty et al. 2014). We used the fitlm function in MATLAB to test temporal trends in all Raz transformations. To compare Raz transformation between sites, when there was strong evidence of uptake, we used the test2function in MATLAB to perform a Welch’s t-test with unequal variance.

**Nutrient uptake**

For our 2nd whole-stream metabolism method of calculating nitrate and TDP retention downstream of WWTP inputs, we analyzed the longitudinal grab samples for these nutrients. We field filtered 60-mL samples for nitrate and chloride to 0.22 μm and froze them until analysis. We measured nitrate and chloride, along with other anions, on an ICS-1000 ion chromatography system (Dionex™, Sunnyvale, California). We collected 125 mL of sample in pre-cleaned bottles for TDP, field filtered samples to 0.22 μm, and acidified them with 3 drops of trace-metal-grade nitric acid. We stored samples at 4°C until analysis on an iCAP™ 7000 ICP-OES system (Thermo Fisher Scientific, Waltham, Massachusetts) for TDP and other cations. Following methods of Martí et al. (2004), we treated the WWTP effluent as a nutrient injection, with nitrate and TDP concentrations normalized to chloride at each sampling location and then used to calculate uptake parameters. We calculated uptake (\( k \)) as the negative slope of the regression between advective travel time (\( t_{ad} \)) from the effluent input and the natural log of dilution-corrected nutrients. Linear regressions fit these relationships best, indicating pseudo-1st-order decay (Hensley et al. 2014), and we used the fitlm function in MATLAB to test the slopes of these regressions (Table S1). We considered \( k \) to be detectable when the slope of the regression had a p-value <0.06. Choosing this alpha level allowed us to include 2 sampling periods at reach 1 with \( k \) that appeared less important even though not as strongly detected as other sampling periods. Time periods that did not have detectable uptake or showed nutrient release are reported as 0.

For the time periods with detectable \( k \), we calculated uptake velocity, uptake length, and areal uptake rates. Uptake velocity (or mass transfer coefficient) considers differences in stream velocity, resulting in the rate of movement from the stream to the bed. We calculated uptake velocity (\( v_t \)) as:

\[
v_t = kd \quad (\text{Eq. 3}).
\]

Nutrient uptake length is conceptualized as the average distance a nutrient molecule travels downstream in inorganic form before being assimilated. We calculated uptake length (\( S_u \)) as:

\[
S_u = Q / (v_t w)
\]

(Eq. 4),

where \( Q \) is mean stream discharge in m³/s and \( w \) is mean wetted width in m. Finally, the areal uptake rate considers different levels of ambient nutrient concentrations, resulting in the mass assimilated/area streambed/unit time (Stream Solute Workshop 1990). Areal uptake rate (\( U_i \)) was calculated as:

\[
U_i = v_C C
\]

(Eq. 5),

where \( C \) is the mean nutrient concentration of the reach during each sampling time (Stream Solute Workshop...
1990). We identified temporal trends in nutrient uptake in the same manner as Raz by using the fitlm function in MATLAB. When there was positive, non-0 uptake, we compared uptake parameters \((k, v_f, \text{ and } U_f)\) between the reaches with a Welch’s \(t\)-test with unequal variance (ttest2 function in MATLAB).

**Daily whole-stream metabolism**

To obtain our 3rd measure of whole-stream metabolic processing, we used the 1-station diel DO method (Hall and Hotchkiss 2017). We used in-situ sensors (EXO™ multiparameter sonde and HOBO U20 and Pendant loggers) to measure DO, temperature, water depth, and light every 15 min at the midpoint of the 2 study reaches: 1494 m downstream of the effluent input at reach 1 (Fig. 1B) and 650 m downstream of the effluent input at reach 2 (Fig. 1C). These records are part of a larger spatial and temporal data set of DO measurements within the Creek; 7 to 8 d of data are used here, from 22 to 29 September 2017 for reach 1 and 12 to 18 September 2017 for reach 2. Solar noon varied from 1257h EDT on 12 September to 1251h EDT on 29 September 2017. We used the StreamMetabolizer package (Appling et al. 2018) in R (version 3.6.1; R Project for Statistical Computing, Vienna, Austria) to calculate daily estimates of whole-stream GPP, ER, and net ecosystem productivity. We used the Bayesian approach and did not pool \(K_{600}\) values, with \(K_{600}\) values calculated by the model for this period as \(2.8 \pm 0.5/d\) at reach 1 and \(5.1 \pm 1.2/d\) at reach 2. \(R^2\) for the fit between modeled and observed DO for these time periods averaged 92% at reach 1 and 99% for reach 2. We did a Welch’s \(t\)-test with unequal variance to compare GPP and ER over the period modeled to test for differences.

**RESULTS**

**Stream solute tracer injections**

**Solute transport**  Transient storage, one driver of metabolism rates, showed different storage processing between reaches. We observed lower discharge and longer advective travel times in reach 1 relative to reach 2 (Table 1). Both central moments were higher in reach 1, consistent with the longer reach 1 advective transport timescales that allowed time for non-advective processes, including dispersion and transient storage, to act on the tracer signal. The CV, skewness, and \(D_{app}\) metrics, all normalized by this advective time scale, provide a more accurate comparison of transient storage between study reaches. CV was similar across both reaches, whereas \(D_{app}\) and skewness were higher in reach 1. This result indicates that both dispersion and short-term storage processes were higher in reach 1 relative to reach 2 even when accounting for the longer advective timescales in reach 1.

**Reach characterization**  Discharge during the tests averaged 0.09 m\(^3\)/s at reach 1 and 0.34 m\(^3\)/s at reach 2 (Table 1). Depths recorded by loggers indicate minimal change in discharge over the course of sampling, with <1 cm of change at reach 1 sites and 1 to 2 cm of change at reach 2 sites (Fig. 2B, F). Temperature increased \(~1.8\) to \(1.9^\circ\text{C}\) over the sampling period in reach 1 and \(~0.6\) to \(1.8^\circ\text{C}\) in reach 2 (Fig. 2A, E). One location was monitored for specific conductivity in reach 1, showing a plateau at the beginning of sampling followed by an increase over the afternoon (Fig. 2G). Specific conductivity increased over the afternoon at the reach 2 WWTP, but this change was attenuated with distance (Fig. 2G). Canopy cover at reach 1 averaged 35.3% open channel, and reach 2 had more open channel (44.8%; \(p = 0.01, \text{df} = 19.4; \text{Fig. S1}\)).

**Raz-based metabolism**  All sampling times at reach 1 and 4 of 6 sampling times at reach 2 showed detectable transformation of Raz to Rru (all tests of regressions \(p < 0.06\) except 1615 and 1700h at reach 2; Table S1). Cumulative Raz transformation to Rru increased with longitudinal distance in both study reaches (Fig. S2A, B). The intercept of this relationship suggests an initial concentration of 10% Rru in the injection, which is consistent with the purity of Raz injected. Estimates of retardation factors from our tests are 1.08 for Rru and 0 for Raz, lower than those reported in Lemke et al. (2014; 1.36 and 1.22, respectively) (Fig. S3A, B). These low retardation factors indicate there was negligible sorption of Raz and Rru at both sites, consistent with the high pH values at the sites (Table 1). Raz to Rru transformation rates, \(k_{trans}\) and the uptake velocity of Raz, \(v_{RC_{Raz}}\), varied by an order of magnitude between the 2 study reaches (Table 2), with higher uptake velocities being observed in reach 1 than reach 2 (\(p = 3.5E^{-4}, \text{df} = 4.4; \text{Fig. 3A, B}\)). In both reaches, \(k_{trans}\) and \(v_{RC_{Raz}}\) declined over the course of the afternoon observation period, \(v_{RC_{Raz}}\) from 0.13 to 0.08 mm/min (\(-33%; p = 0.02, \text{df} = 3; \text{Fig. 3A, B}\)) in reach 1 and from 0.02 mm/min to below detection in reach 2 (\(p = 0.007, \text{df} = 4; \text{Fig. 3A, B}\)).

**Nutrient uptake**

Our 2nd method of ecosystem metabolism evaluation, measuring nutrient retention downstream of WWTPs, showed differences between reaches. Nitrate concentrations in reach 1 were nearly twice as high as those in reach 2 (\(p = 4.1E^{-9}, \text{df} = 7.5\)); however, nitrate fluxes in reach 1 were <½ the fluxes measured reach 2 (Table 2). Concentrations from each individual grab sample site generally increased over the afternoon, although mean concentration across the entire reach during the test period only increased at reach 2 (\(p = 0.03, \text{df} = 4; \text{Table 2}\)). Nitrate concentrations decreased with distance downstream of the
plant at both sites during most sampling periods. All samplings at reach 1 showed detectible uptake, and 3 time periods at reach 2 had detectible uptake of N (1515, 1615, and 1700h), 1 had marginal uptake (1415h), and 2 had negative uptake (1215 and 1315h; Table S1, Fig. S4A, C). Nitrate uptake length at reach 1 ranged from 7.9 to 16.9 km, whereas reach 2 had late-afternoon (1515, 1615, and 1700h) uptake with lengths from 6.7 to 9.0 km (Table 2). In addition, areal rates of NO₃⁻ uptake showed indications of saturation kinetics, with very high areal uptake rates (988–2554 mg N m⁻² d⁻¹) but low uptake velocities (0.038–0.160 mm/min) between sites at both reaches during periods with uptake. When considering only sample periods with detectible positive uptake, uptake velocities were lower in reach 1 than reach 2 (p = 0.02, df = 2.3), as was areal uptake (p = 0.03, df = 2.8). There was no measurable change in nitrate uptake rates over the afternoon at reach 1, whereas both \( v_t \) (p = 0.009, df = 4) and \( U \) (p = 0.005, df = 4) increased over the afternoon at reach 2 (Fig. 3A, B).

Phosphorus concentrations and trends also differed between the 2 reaches. TDP concentrations were ~20% higher in reach 1 than reach 2 (p = 3.6E−4, df = 5.8; Table 2). TDP increased over the afternoon at all sites in reach 1, but was more variable at reach 2, plateauing at many sites in the later afternoon. Average whole-reach TDP concentrations increased over the afternoon at both reaches (reach 1: p = 0.002, df = 4; reach 2: p = 0.04, df = 5) (Table 2). At reach 1, 3 time periods had detectible uptake for TDP (1430, 1530, and 1630h; Table S1), 1 time period showed marginal uptake (1300h), and 1 time period had a negative uptake (1230h). At reach 2, only 1 time period had detectible uptake for TDP at reach 2 (1215h), 2 time periods were marginal (1315 and 1700h), and the remaining 3 time periods had negative uptake (1415–1615h) (Table S1, Fig. S4B, D).
Table 2. Major metabolic parameters compared between sites. Bolded values indicate higher values between reaches, as identified with the analysis of variance, with p-values < 0.05. Sampling intervals with undetectable uptake or release are reported as 0 for uptake rates and velocities or ∞ for uptake lengths. Up and down arrows indicate increasing or decreasing temporal trends during the afternoon study period with p-values reported. No arrow means there was no temporal trend in that variable over the afternoon. All ranges are temporal. GPP = gross primary production, ER = ecosystem respiration, $k_{sad}$ = Raz–Rru transformation rate coefficient, $S_w$ = uptake length, $v_l$ = uptake velocity, TDP = total dissolved phosphorous, $U$ = areal uptake rate.

| Parameters | Reach 1 | Reach 2 |
|------------|---------|---------|
| NO$_3^-$ (mg N/L) | 18.2–19.0 | 10.2–11.1 (↑ p = 0.03) |
| Average NO$_3^-$ flux (g/s) | 1.7 | 3.9 |
| TDP (mg/L) | 0.37–0.47 (↑ p = 0.002) | 0.27–0.36 (↑ p = 0.04) |
| Average TDP flux (mg/s) | 37 | 120 |
| GPP (g O$_2$ m$^{-2}$ d$^{-1}$) | 2.7 (0.9–4.9) | 7.0 (4.3–10.8) |
| ER (g O$_2$ m$^{-2}$ d$^{-1}$) | -7.4 (-5.6–-8.9) | -10.1 (-7.2–-13.3) |
| # significant Raz transformation | 5 (of 5) | 4 (of 6) |
| Raz $k_{sad}$ (x10$^{-4}$ min$^{-1}$) | 3.74–5.77 (↑ p = 0.02) | 0–0.98 (↓ p = 0.007) |
| Raz $v_l$ (mm/min) | 0.082–0.127 (↑ p = 0.02) | 0–0.021 (↓ p = 0.007) |
| # significant N uptake | 5 (of 5) | 3 (of 6) |
| NO$_3^-$ $S_w$ (km) | 7.9–16.9 | 6.7–∞ |
| NO$_3^-$ $v_l$ (mm/min) | 0.038–0.081 | 0–0.160 (↑ p = 0.009) |
| NO$_3^-$ $U$ (mg N m$^{-2}$ d$^{-1}$) | 988–2235 | 0–2554 (↑ p = 0.008) |
| # significant TDP uptake | 3 (of 5) | 1 (of 6) |
| TDP $S_w$ (km) | 9.0–∞ | 6.1–∞ |
| TDP $v_l$ (mm/min) | 0–0.071 | 0–0.176 |
| TDP $U$ (mg m$^{-2}$ d$^{-1}$) | 0–42 | 0–80 |

Neither reach had a temporal trend in TDP uptake. Because of the small number of samples with detectible uptake, spatial statistics for TDP could not be calculated.

DO-based metabolism

The 3rd method of metabolism evaluation, the 1-station diel DO method, also showed differences between the 2 reaches. On the day of the injection, DO varied from 1.86 to 9.59 mg/L at reach 1, with the minimum at 0545h and the peak at 1645h (Fig. 2D), and temperature ranged from 22 to 26°C (Fig. 2A). At reach 2, DO ranged from 7.44 to 12.10 mg/L, with the minimum at 0045h and maximum at 1130h (Fig. 2H), and temperature ranged from 16 to 20°C (Fig. 2E). Using 8 d of DO measurements, both sites were always net heterotrophic, although reach 1 was more heterotrophic (i.e., had more negative net ecosystem production). At reach 1, mean GPP was 2.7 g O$_2$ m$^{-2}$ d$^{-1}$ (±1.6 SD), and mean ER was -7.4 g O$_2$ m$^{-2}$ d$^{-1}$ (±1.1). In contrast, mean GPP at reach 2 was 7.0 g O$_2$ m$^{-2}$ d$^{-1}$ (±2.2), and mean ER was -10.1 g O$_2$ m$^{-2}$ d$^{-1}$ (±2.3). Instream productivity was higher in reach 2 than reach 1 (GPP: p = 0.002, df = 10.9; ER: p = 0.022, df = 8.2) (Fig. S5A, B).

DISCUSSION

This study examined spatial and temporal patterns of 3 different metabolic indicators downstream of 2 WWTPs, reporting results from the Raz–Rru tracer system in a WWTP-impacted system for the first time. One of the key advantages of the Raz–Rru tracer system over traditional DO-based daily estimates is the ability to resolve sub-daily changes in ER and provide daytime heterotrophic metabolism estimates. The Raz–Rru tracer confirmed low metabolic activity, potentially due to nutrient saturation, along with low DO-based GPP and ER rates. Spatial differences were observed downstream from 2 WWTPs only 10 km apart, but these differences apparently were not related to differing effluent contributions as expected. Furthermore, temporal variations were not consistent between the reaches or between the different metabolic indicators. In this section we discuss the drivers for these different responses and illustrate the need for multiple indicators to investigate ecosystem metabolism in streams.

Wastewater impacts on metabolic activity

Previous research has shown that downstream of WWTPs, effluent inputs saturate nutrient demand and reduce nutrient removal efficacy (Martí et al. 2004, Covino et al. 2010, Hensley et al. 2014), with concurrent effects on ER and GPP (Gücker et al. 2006), which our results support. Our observed rates for DO-based GPP and ER and for nutrient uptake all fell within the ranges seen at other
WWTP-impacted sites, and other studies that used DO to measure productivity rates in WWTP-impacted streams reported rates that were both similar to ours (Aristi et al. 2015) and higher (Gücker et al. 2006). Along with similar GPP and ER rates, long uptake lengths have been measured for nitrate in most WWTP-impacted systems (Haggard et al. 2001, Martí et al. 2004, Gücker et al. 2006; Fig. 4), and our system showed the same pattern. Our high nitrate uptake rates were driven by extremely high nitrate concentrations in our streams, as expected downstream of WWTPs. Nitrate uptake velocities were low in our system, as seen in other eutrophic streams (Ensign and Doyle 2006, Gücker and Pusch 2006, Hall et al. 2009, Hensley et al. 2014). Low uptake velocities combined with long uptake lengths indicate that demand for nitrate was lower than the load. Phosphorus uptake was also on par with other WWTP removal results (Haggard et al. 2001, Martí et al. 2004, Gücker and Pusch 2006). Haggard et al. (2005), Gücker and Pusch (2006), and Gücker et al. (2006) all reported a large range in TDP areal uptake rates, from ~14 up to 2506 mg P m$^{-2}$ d$^{-1}$, with many higher than the maximum of 80 mg P m$^{-2}$ d$^{-1}$ measured across our reaches. We infer that high nitrate and TDP concentrations do not limit rates of metabolic processing.

Raz-transformation rates in a WWTP-impacted stream system have not been reported previously, but our Raz results are consistent with DO-based studies reporting reduced ER in such systems. The observed rates of Raz transformation are on the low end of values previously reported for natural and artificial stream systems not affected by WWTP. When compared with streams of similar size, the higher values observed in reach 1 are within the low end of the range of other reported values (Blaen et al. 2018, Knapp and Cirpka 2018), whereas those in reach 2 are an order of magnitude lower than previously observed values. These low Raz-transformation rates illustrate the sensitivity of the method, showing distinctly different rates downstream of nearby WWTPs, and confirm the impact of nutrient saturation on stream metabolism.

**Drivers of spatial patterns in metabolic processes**

Metabolic activity clearly differed between the 2 study reaches; however, this spatial difference was not consistent among the various metabolic metrics measured. Nitrate uptake rates and DO-based GPP and ER were all lower in reach 1, which had higher nutrient concentrations than reach 2, in support of our hypothesis 1. Reach 2 may also be less effective at P retention, given that more sampling times had no detectible P retention. In contrast, Raz transformation was higher in reach 1. This difference suggests that, counter to our hypothesis 3 and that proposed by Haggerty et al. (2009), the primary drivers of the spatial patterns in these metabolic processes were decoupled in our system.

Variable environmental conditions between reaches may have influenced nutrient dynamics and metabolism in our study. The corresponding differences in nutrient uptake and DO-based metabolism between reaches, and their inverse relationships with nutrient concentrations and canopy...
cover, suggest that excessive nutrient concentrations, shading, or both, inhibit both nutrient uptake and GPP and ER at reach 1 relative to reach 2. Such clear coupling between instream metabolism and nutrient uptake velocities is well supported in the literature (Heffernan and Cohen 2010, Hensley and Cohen 2016). The fact that our GPP rates are similar to urban streams with lower nutrient loads (Clapcott et al. 2016, Alberts et al. 2017) indicates that reduced light, more so than excessive nutrients, may limit GPP in reach 1 (Bernhardt et al. 2018, Reisinger et al. 2019). Although both reaches have a similar depth, reach 2 is more than twice as wide as reach 1 (Table 1) with a more open canopy (Fig. S1), suggesting similar light penetration to the benthic surface at both reaches but a greater and more highly lit surface area for benthic microalgae growth in reach 2. These differences indicate that photoautotrophic productivity and nutrient uptake may be more important than heterotrophic and dissimilatory activity in reach 2 relative to reach 1, at least in the afternoon. The relative predominance of photoautotrophic activity in reach 2 is also consistent with its lower relative transient storage (Table 1), potentially limiting the transport of nutrients from the water column to the subsurface and, thus, limiting denitrification. The spatial differences in nutrient uptake lengths between sites are inversely related to differences in specific discharge, which is the opposite of the scaling trend typically observed in streams (Hall et al. 2013). This inverse trend suggests that local reach conditions, including site-specific differences in nutrient uptake and GPP driven by light availability, nutrient concentrations, and benthic contact, rather than stream discharge, are the predominant drivers of the observed spatial differences in nutrient uptake.

Higher Raz-transformation rates and uptake velocities were observed in reach 1 than reach 2. The higher rates were associated with higher nutrient concentrations (Table 2), dispersion ($D_{app}$), and short-term storage processes (skewness), as well as lower relative stream size (as measured by discharge and specific discharge) and light availability (as proxied by canopy cover) (Table 1). Unlike nutrient uptake, the higher nutrient concentrations in reach 1 do not appear to inhibit relative Raz transformation, and, in fact, there is some evidence that higher nutrient concentrations can stimulate microbial respiration, at least in lower-nutrient systems (e.g., Ramírez et al. 2003, Halvorson et al. 2016). Thus, the higher Raz-transformation rates in reach 1 are most likely the result of enhanced short-term transient storage processes, including hyporheic exchange, in-channel storage, or both, promoting higher heterotrophic activity.
relative to reach 2. Reduced light availability and higher nutrient concentrations in reach 1 result in lower relative photoautotrophic activity relative to reach 2.

Drivers of temporal patterns in metabolic processes

Temporal differences in environmental conditions, and not WWTP effluent magnitude, appear to be driving differences between the reaches. Nutrient concentrations, nitrate uptake, water temperature, and Raz-transformation rates were all observed to change over the course of the afternoon tracer tests (~1200–1700h) in 1 or both study reaches. Although we did not directly measure light across the entire reaches, it presumably also changed (solar noon was at 1254h ± 3 min). In contrast, neither the magnitude of WWTP effluent inputs nor stream discharge (as approximated by stream depth; Fig. 2B, F) showed a temporal change over the sampling period as we had expected.

The presence of temporal patterns in nutrient concentrations and uptake differed between the 2 reaches. Neither nutrient removal efficiency nor nitrate concentrations changed over the afternoon in reach 1, contrary to our hypothesis 2 that these parameters would be time varying along with effluent magnitude, although TDP concentrations increased. However, at reach 2, nitrate uptake, along with both nitrate and TDP concentrations, increased over the afternoon (Fig. 3B). Although this reach 2 trend was in line with our hypothesis 2 and paralleled the temporal trend in temperature, it was not perfectly in-phase with rates of instantaneous GPP, which would be expected to mirror the reach 2 afternoon plateau in DO concentration (Fig. 2H). This offset in phase suggests that assimilation, which is typically tightly coupled to the timing of GPP (Heffernan and Cohen 2010, Kurz et al. 2013) was not the sole driver of the increasing rates of nitrate removal observed in reach 2. Higher daytime nitrification, enhanced by light, DO, and temperature, has been shown to promote denitrification (Laursen and Seitzinger 2004), which could have been the case in this study. However, unlike many other WWTP-impacted systems (Marti et al. 2004, Figueroa-Nieves et al. 2016), ammonium concentrations are reasonably low in our WWTP effluent (<0.1 mg/L as reported by WWTPs), and nitrification of effluent is not likely to be a major factor influencing N cycling in our system. Further, unlike for other well-lit systems, our observations do not support inhibition of daytime denitrification in the subsurface by benthic microalgae productivity (Nimick et al. 2011 and references therein). The lack of other temporal trends in nutrient uptake highlights the complexity of disentangling diel nutrient processing dynamics, especially given the short length of our observation window. In addition, the time-varying response to nitrate and TDP uptake indicates complex interactions between nutrient concentrations, loads, and the timing of N and P demand by microbes (Heffernan and Cohen 2010, Cohen et al. 2013). We did not expect the variation in temporal change of nutrient uptake between reaches (including the lack of temporal change in reach 1), and this variation indicates that nutrient uptake at these sites may be decoupled from other temporal factors, including nutrient concentration, discharge, and temperature, at the timescale sampled.

In contrast to nutrient uptake, the observed temporal declines in $k_{\text{rad}}$ and $V_{\text{F-Raz}}$ at both reaches over the afternoon (Fig. 3A, B) illustrate the ability of the Raz–Rru tracer system to resolve sub-daily changes in reach-scale ER but are opposite in direction from our expectation (hypothesis 2). Although it is well known that autotrophic processes, including increased rates of daytime photoautotrophic nitrate uptake and nighttime autotrophic P uptake, can vary with sub-daily frequency (Heffernan and Cohen 2010, Nimick et al. 2011 and references therein, Cohen et al. 2013), comparatively little is known about diel variations in heterotrophic activity or uptake. ER in streams is often assumed to be either diurnally stable, as is the case for the diel DO method for estimating stream metabolism (Odum 1956, Owens 1974), or to increase predictably with temperature, such as in the van’t Hoff-Arrhenius equation. Testing these assumptions, González-Pinzón et al. (2016) observed no significant differences between daytime and nighttime respiration rates estimated from Raz transformation. They suggested this lack of difference was likely because most stream respiration takes place in the hyporheic zone where diurnal fluctuations in stream temperature and light are considerably attenuated. In contrast, Tobias et al. (2007) and Hotchkiss and Hall (2014) found evidence using stable oxygen isotopes that daytime ER was higher than nighttime. However, their estimates were subject to uncertainty resulting from estimated parameters, which limited their ability to further interrogate potential drivers of this daytime increase.

Previous research has reported increased biochemical reaction rates with temperature, and corresponding spatial and temporal coupling between temperature and ER in streams (Sinsabaugh 1997, Perkins et al. 2012, Beaulieu et al. 2013, Hotchkiss and Hall 2014). However, in our system surface water temperatures increased over the afternoon (Fig. 2A, E), reflecting an inverse relationship with $k_{\text{rad}}$ and $V_{\text{F-Raz}}$. Even if Raz transformations primarily occur in the subsurface, the temperature there is also expected to increase some, even accounting for a potential lag between surface and subsurface patterns. Likewise, temporal variation in nutrient concentrations does not appear to influence temporal patterns in $k_{\text{rad}}$ and $V_{\text{F-Raz}}$. Both nitrate (reach 2 only) and TDP concentrations increased during the observation window, inverse with Raz transformation, but there is no evidence that nutrients were limiting in our system. Also, comparison of the 2 reaches does not appear to support nutrient concentration inhibition of Raz transformation.

Light is the major driver of GPP in streams, which was illustrated in our study by the lower shading and higher...
GPP rates in reach 2. In turn, daily estimates of ER are often, but not always, predicted by GPP (e.g., Beaulieu et al. 2013), in part because of the availability of new or more labile carbon to drive ER (e.g., Kaplan and Bott 1982, De Lange et al. 2003, Heffernan and Cohen 2010). Tobias et al. (2007) and Hotchkiss and Hall (2014), suggest this relationship between light and ecosystem metabolism could downscale to sub-daily linkages. Further, daytime increases in benthic algae productivity have also been shown to increase oxygen penetration in the subsurface (Nielsen et al. 1990, Lorenzen et al. 1998), which would expand the aerobic zone for subsurface Raz transformation during the day. Our observed pattern of decreasing $V_{f-Raz}$ over the afternoon does not appear to clearly support either of these light-driven mechanisms. However, we note that our ability to fully resolve the drivers of the observed temporal change in Raz-transformation rates is confounded by the short (4–5 h) observation window during our tracer tests and the timing of this window ($\sim$1200–1700h) that roughly spanned the period between the solar peak and the lagged peak in DO. This decline in $V_{f-Raz}$ appears to be in-phase with the diel pattern of daily solar radiation (declining after $\sim$1254h). However, the timing of most light-driven metabolic processes, including GPP and assimilatory nitrate removal, lag behind the solar peak (Heffernan and Cohen 2010) and would, therefore, still have been increasing during the afternoon observation window. For instance, this lag was seen in DO concentrations, especially in reach 1 (Fig. 2D). The differences in temporal responses of the metabolic indicators illustrates the need for further investigation into the drivers of ecosystem metabolism in streams.

**Contrasting metabolic indicators**

The 3 metabolic indicators, Raz transformation, nutrient uptake, and DO-based GPP and ER, differed in the patterns between reaches. Unlike our expectations (hypothesis 3), the trend in Raz transformation did not mirror nitrate uptake or ER. Conversely, spatial nitrate uptake patterns were similar to those of DO-based GPP and ER, all which had higher rates in reach 2. The patterns also differed over the afternoon, with a decrease in uptake velocity for Raz in both reaches, an increase in nitrate uptake at reach 2, and no change for all other nutrient uptake velocities.

The relationship between Raz-based and DO-based measurements of respiration is dependent on the assumptions of each method. A lack of relationship between Raz-based and DO-based estimates of ER have been reported before within stream systems (Kurz et al. 2017, Knapp and Cirpka 2018) even though Raz transformation has been shown to be well correlated with oxygen turnover both in batch experiments and broadly across stream systems (González-Pinzón et al. 2012, 2014, 2016, Knapp and Cirpka 2018). However, this apparent contradiction could be attributed to methodological differences and assumptions. Raz-transformation rates measured during the observation window of a tracer test are not necessarily representative of the full 24-h integral of the diel DO method. In our case, however, this explanation is not sufficient to account for the spatial difference in diel DO- and Raz-based respiration measurements. That is, respiration would have had to have been extremely variable over time and, in the case of reach 2, unrealistically high during the unsampled periods of the day. Additionally, there could be losses of Raz or Rru through processes other than the decay of Raz to Rru, which would only matter if the relative magnitude of Raz vs Rru loss varied over space, time, or both. Sorption of both tracers was negligible, but both Raz and Rru are somewhat sensitive to photodecay, with the time-scale of Rru photodecay being an order of magnitude faster than that of Raz (10s of h vs 100s of h, respectively; Haggerty et al. 2008). Although relative differences in Raz vs Rru photodecay cannot explain the directionality in observed temporal patterns, they could contribute to the spatial differences between the 2 sites. Enhanced photodecay in the more open canopy of reach 2 would reduce the apparent $k_{Rru}$ values, but an unrealistic increase in photodecay of $>$20-fold would be needed to account for the entire spatial difference observed.

More plausibly, we attribute the difference in DO- and Raz-based ER results to a lack of understanding of the physical or biological processes that drive Raz transformation in stream systems. It is possible that the tracer-based approach does not sample the same physical compartments of the ecosystem as the diel DO method, despite both being metrics of whole-stream oxygen consumption. Only flowpaths falling within the time period of the tracer test are reflected in the tracer observations, although plateau injections such as ours more completely sample longer flowpaths than are sampled by slug injections. Likewise, the differing results between methods may reflect a lack of biological process understanding regarding Raz transformation specifically or ecosystem metabolism more broadly, including the roles of heterotrophs, autotrophs, or oxygen-cycling reactions. Notably, González-Pinzón et al. (2012) demonstrated that the magnitude of Raz transformation relative to oxygen loss differed between pure cell cultures, suggesting that microbial community structure could strongly affect the degree of Raz transformation observed relative to oxygen loss. This dependency likely holds for Raz transformation relative to other microbially mediated functions, such as nutrient cycling.

The inverse spatial patterns observed in the uptake velocities of nutrients and Raz suggest the drivers of nitrate uptake and respiration are decoupled in our system. Although Haggerty et al. (2009) predicted Raz transformations and nutrient uptake rates could be correlated if
transient storage and nutrient retention are related, our findings indicate the limitations of this assumption. It is understood that metabolic processes, while coupled, are not always synchronous in their timing. For example, the timing of assimilation varies for different elements at different times of day because of biota processing along different pathways (Heffernan and Cohen 2010, Cohen et al. 2013, Appling and Heffernan 2014, Hensley and Cohen 2016, Bernhardt et al. 2018), and rates of denitrification have been shown to be driven by the previous day’s GPP (Heffernan and Cohen 2010). Such lags could explain the observed temporal trend in sub-daily Raz transformation, but our limited observation window and differences in temporal trends in nutrient uptake (or lack thereof) limit our ability to resolve these nuances further.

Collectively, our results demonstrate that identifying the drivers of different measures of metabolic rates in systems with complex physical and chemical inputs, such as stressed urban streams, can be challenging. In addition, the 2 reaches, which are only 10 km apart, show large spatial and temporal differences in metabolic processing within each measurement method, highlighting the complexity of metabolic responses to small changes in hydrology, geochemistry, and light availability. The extremely high nutrient flux at both sites supports a system that is energy, not nutrient, limited (Reisinger et al. 2019). In the wider, more open canopy and lower nutrient reach 2, photoautotrophic productivity and nutrient uptake appear to be more important than heterotrophic and dissimilatory removal. In contrast, heterotrophic activity appears to be more important in reach 1, where higher transient storage potentially facilitated the transport of nutrients from the water column to the subsurface. Temporal trends in nitrate uptake, only discerned in reach 2, could be the result of time-varying rates of primary productivity and coupled nitrification–denitrification.

Different observed outcomes among the multiple methods of metabolism evaluation suggests a measure of uncertainty should be incorporated into single-method stream metabolism estimates in WWTP-impacted systems. The Raz–Rru tracer system provides a unique opportunity to advance our understanding of the drivers of heterotrophic activity, especially at sub-daily timescales. Combining and comparing multiple measures of metabolic activity allows for more complete inference of the factors driving temporal and spatial patterns in stream metabolic processes. Further research is needed to support a more mechanistic understanding of the drivers of Raz transformation relative to often contrasting DO-based estimates of ER and other metrics of ecosystem metabolism in streams.

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