

1 **Seasonal dynamics of zooplankton nutrient recycling in a hypereutrophic reservoir**

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18

19 **ABSTRACT** (up to 200 words)

20 Consumer-driven nutrient cycling influences aquatic ecosystem functioning by altering
21 energy flow as well as the movement and transformation of nutrients. In hypereutrophic lakes,
22 zooplankton nutrient recycling has largely been considered negligible due to the high
23 concentrations of available nutrients. A recent comparative analysis found that zooplankton
24 community composition may influence nutrient availability, particularly phosphorus availability,
25 in hypereutrophic lakes. However, the magnitude and timing of the effect of zooplankton
26 nutrient recycling and grazing on phytoplankton in hypereutrophic lakes remains unclear. We
27 quantified zooplankton, phytoplankton, and nutrient concentration dynamics during the summer
28 of 2019 in a temperate, hypereutrophic reservoir. We found that zooplankton excretion
29 contributed a substantial proportion (23-46%) to the inorganic phosphorus (P) pool in early
30 summer when P concentrations were low and limiting phytoplankton growth. Zooplankton
31 excretion of P also significantly ($p=0.003$) contributed to variation in phytoplankton community
32 composition in early summer. Further, we found evidence that zooplankton affected
33 phytoplankton size distributions through selective grazing of smaller phytoplankton cells.
34 Overall, our results demonstrate the important role of zooplankton in hypereutrophic lakes as
35 excretion helped drive springtime phytoplankton dynamics through nutrient recycling while
36 grazing influenced phytoplankton size distributions later in the summer.

37

38 **INTRODUCTION**

39 Animal consumers drive nutrient cycling in aquatic ecosystems by controlling the
40 movement and transformation of nutrients over time and across space (Atkinson *et al.*, 2017).
41 Aquatic consumers such as zooplankton ingest primary producers then excrete and egest
42 metabolized and unassimilated materials as waste, recycling nutrients back into the ecosystem
43 (Vanni, 2002). In turn, these excreted, recycled nutrients can be taken up by phytoplankton to
44 produce new biomass (Sarnelle and Knapp, 2005). Mismatches between consumer demand for
45 and assimilation efficiency of elements and the elemental composition of primary producers
46 drives the stoichiometry of nutrients recycled back into the ecosystem (Elser and Hassett, 1994;
47 Sterner, 1990). Elemental mismatches can then lead to greater nutrient recycling of a particular
48 element that may result in changes to ecosystem nutrient limitation and alter trophic interactions
49 between consumers and their resource (Elser *et al.*, 2000; Dobberfuhl and Elser, 2000).

50 The community composition of both phytoplankton and zooplankton can influence the
51 stoichiometry of recycled nutrients and generate strong differences in nitrogen (N) and
52 phosphorus (P) recycling (Balseiro *et al.*, 1997). For example, copepods and small cladocerans
53 generally retain more N whereas *Daphnia* generally retain more P (Elser and Urabe, 1999).
54 Differences in N and P retention between zooplankton taxa can result in copepod and small
55 cladoceran dominated communities retaining more N and recycling more P into the water
56 column, thereby driving phytoplankton to N-limitation (Elser *et al.*, 2000, 1988). Phytoplankton
57 community composition varies with lake trophic state and nutrient limitation as different
58 phytoplankton genera prefer different nutrient forms (Andersen *et al.*, 2020). Cyanobacteria-
59 dominated phytoplankton communities, which often arise in nutrient enriched ecosystems, are
60 uniquely resistant to zooplankton grazing due to the ability of many genera to form colonies or
61 filaments, as well as their poor nutritional quality, toxin production and rapid growth rates
62 (Moustaka-gouni and Sommer, 2020). This results in a loss of nutrient and energy transfer to
63 higher trophic levels due to poor assimilation efficiency (Karpowicz *et al.*, 2021, 2020). In
64 combination, zooplankton-phytoplankton interactions affect nutrient recycling in aquatic
65 ecosystems; however, the effects may vary depending on the severity of nutrient enrichment.

66 Much of our understanding regarding zooplankton nutrient recycling comes from
67 oligotrophic and eutrophic ecosystems (Elser *et al.*, 2000; Moegenburg and Vanni, 1991), though
68 many temperate lakes are increasingly becoming hypereutrophic due to continued land use
69 conversion and climate change (Stoddard *et al.*, 2016). The extremely high nutrient
70 concentrations in hypereutrophic lakes can produce unique conditions compared to less enriched
71 waterbodies such as large seasonal variability in nutrient limitation of phytoplankton growth
72 (Andersen *et al.*, 2020), substantial internal phosphorus loading under oxic conditions (Song and
73 Burgin, 2017), and a more complex mix of top-down and bottom-up forces affecting
74 phytoplankton communities (Matsuzaki *et al.*, 2018). However, the contribution of zooplankton
75 nutrient recycling in hypereutrophic ecosystems is often considered negligible as high
76 concentrations of inorganic nutrients within the water column can dampen the influence of
77 nutrient inputs from consumer excretion (Spooner *et al.*, 2013). Despite this, zooplankton may
78 still influence nutrient availability in hypereutrophic lakes as nutrient limitation and zooplankton
79 biomass shift throughout the growing season. Additionally, selective feeding on small
80 phytoplankton by small-bodied zooplankton increases the dominance of large phytoplankton

81 species, including filamentous and colonial cyanobacteria (Erdoğan *et al.*, 2021). This may
82 influence nutrient availability as cyanobacteria have the capacity for luxury nutrient uptake,
83 subsequent storage of excess nutrients, and the ability to use N forms inaccessible to other
84 phytoplankton species, such as diazotrophs that can fix atmospheric nitrogen (Cottingham *et al.*,
85 2015). As hypereutrophic lakes are dominated by smaller-bodied zooplankton including
86 microzooplankton and ciliates, selective grazing pressure on the phytoplankton community may
87 indirectly influence nutrient availability.

88 A recent analysis of mesozooplankton (i.e., copepods, cladocerans, and rotifers; hereafter
89 zooplankton) stoichiometric traits found that the community N:P ratio shifted towards N-rich
90 species with increasing eutrophication (Moody and Wilkinson, 2019). As such, in hypereutrophic
91 lakes, zooplankton may be increasing P recycling. This hypothesis was supported by the fact that
92 the seston N:P in hypereutrophic lakes was lower compared to less enriched lakes concurrent
93 with more N-rich zooplankton communities that exhibit greater P recycling. This analysis
94 suggests that the unique functioning of hypereutrophic lakes, even compared to eutrophic lakes,
95 was due in part to the consumers inhabiting them. However, this was a comparative study among
96 lakes based on a single sampling point in summer. It is well established that zooplankton and
97 phytoplankton communities are dynamic and undergo an annual succession over the course of a
98 summer growing season, which can vary depending on lake trophic state and other variables
99 (Sommer *et al.*, 2012). Furthermore, the balance of top-down and bottom-up forces in lakes
100 varies with nutrient ratios and concentrations across a season (Rogers *et al.*, 2020). In the scope
101 of this comparative study (Moody and Wilkinson, 2019), the seasonal variability within
102 zooplankton, phytoplankton, and nutrient dynamics was not captured. As such, it remains unclear
103 how hypereutrophic lakes are influenced by zooplankton nutrient recycling throughout the
104 summer growing season.

105 We investigated the role of zooplankton nutrient recycling in a hypereutrophic waterbody
106 by quantifying zooplankton, phytoplankton, and water column nutrient dynamics across the
107 summer growing season. We used effect traits, which directly link individual activity to
108 ecosystem processes (Hébert *et al.*, 2017, 2016b), to estimate storage and flux of nutrients driven
109 by zooplankton consumers. Effect traits like size (Litchman *et al.*, 2013) or elemental
110 composition (Sterner and Elser, 2002) can be used to assess nutrient cycling by quantifying
111 zooplankton body stoichiometry and excretion rate, as well as to infer growth and predation

112 based on community body size structure. With this approach we asked: (1) do zooplankton in
113 hypereutrophic lakes alter P availability, and how does that vary over the course of a summer
114 growing season? And (2) do zooplankton contribute to variation in phytoplankton community
115 composition amidst other environmental variables? We hypothesized that zooplankton nutrient
116 recycling would contribute to P availability before the onset of high internal P loading in
117 midsummer. Furthermore, we expected zooplankton excretion would contribute to variation in
118 phytoplankton community composition due to body stoichiometry shifting nutrient availability.
119 Finally, we hypothesized that zooplankton grazing would affect phytoplankton size structure
120 throughout the summer due to selective grazing on smaller phytoplankton.

121

122 **METHODS**

123 *Study Lake*

124 Green Valley Lake (46°06'02" N, 94°23'05" W) is a hypereutrophic reservoir built in 1952 as an
125 impoundment of the Platte River in southwestern Iowa (USA). The maximum depth is 7.3 m,
126 with an average depth of 3.2 m and a surface area of 156 ha. The fish community is dominated
127 by crappie (*Pomoxis spp.*), bluegill (*Lepomis macrochirus*), and largemouth bass (*Micropterus*
128 *salmoides*). Additionally, there is a small population of common carp (*Cyprinus carpio*) and
129 channel catfish (*Ictalurus punctatus*). The watershed is dominated by row crop agriculture
130 (68.4% corn/soybean rotation). Consequently, Green Valley Lake is enriched with nutrients and
131 beset by annual phytoplankton blooms dominated by cyanobacteria (Supplementary Figure S1).
132 To characterize zooplankton nutrient recycling in Green Valley Lake, zooplankton,
133 phytoplankton, and water samples for nutrient analysis were taken weekly at the deepest point in
134 the reservoir during the summer of 2019. Samples were taken from early May (day of year; DOY
135 143) to early September (DOY 251). An additional sampling event occurred on DOY 273, but
136 only zooplankton and nutrient samples were collected at that time. Additionally, a YSI EXO3
137 sonde (Yellow Springs Instruments, Yellow Springs, Ohio, USA) was deployed at 0.5 m at the
138 deepest point in the reservoir and collected temperature, total dissolved solids, and pH
139 measurements every 15 minutes. Daily averages of each variable were used in our analyses.

140

141 *Plankton Analysis*

142 Zooplankton and phytoplankton biomass and community composition were quantified for
143 each sampling event during the summer growing season. Zooplankton were sampled via a
144 vertical tow of a Wisconsin net (63 μm mesh) from 6 m depth. Samples were preserved with a
145 formaldehyde solution (5% concentration after sample addition) in the field and later transferred
146 to 70% ethanol. Phytoplankton samples were a composite sample over depth. Water was
147 collected in a 4 L Van Dorn sampler from 0.25, 1, 2, 3, and 4 m depths, then mixed in a 20 L
148 carboy in the field. Samples were taken to 4 m depth which was the average thermocline depth at
149 the sampling point (4.0 ± 1.23 m; s.d.). A 1 L sample was then taken from the carboy following
150 thorough mixing and preserved with Lugol's solution in the field.

151 Zooplankton samples were identified and enumerated with a Leica MZ8
152 stereomicroscope connected to Motic Images software. For each sample a 1 mL subsample was
153 taken where a minimum of 60 individual zooplankton were identified and enumerated. If less
154 than 60 organisms were in the subsample a second 1 mL subsample was counted. Individuals
155 were identified to genus for cladocerans and rotifers, order for copepods, and class for ostracods.
156 Copepod nauplii could not be identified to order and were simply identified as nauplii. We
157 measured zooplankton lengths to calculate dry mass per liter using length-weight regressions
158 (Dumont *et al.*, 1975; McCauley, 1984). For visual display of the zooplankton data, they were
159 separated into nine taxonomic groups: *Daphnia*, *Simocephalus*, *Ceriodaphnia*, *Bosmina*,
160 *Chydorus*, rotifers, calanoids, cyclopoids, nauplii, and ostracods (Supplementary Table S1).
161 *Simocephalus* contributed only 7% of total community biomass at its peak biomass and so were
162 grouped with *Daphnia* for later analyses.

163 The 1 L phytoplankton samples were transferred to a graduated cylinder and allowed to
164 settle in a dark environment for 8 days before the supernatant was removed with a vacuum
165 pump, leaving 50 mL of concentrated sample. A subsample was then removed from the
166 concentrated sample and identified and enumerated using a modified Palmer-Maloney chamber.
167 Phytoplankton were identified to genus and measured using a calibrated ocular reticle on a Leitz
168 DM IL inverted microscope at 400x magnification. A minimum of 300 natural units across 8
169 fields were measured for each sample. Biovolume per liter was calculated based on
170 phytoplankton shape and then converted to wet biomass per liter assuming a 1:1 ratio between
171 wet mass and biovolume (Hillebrand *et al.*, 1999; Sournia, 1978). Like zooplankton,
172 phytoplankton genera were separated into six groups for visual display: bacillariophytes,

173 chlorophytes, chryso- and cryptophytes, *Aphanothece*, *Microcystis*, and other cyanophytes
174 (Supplementary Table S2). Both *Aphanothece* and *Microcystis* were the dominant genera of
175 cyanobacteria, contributing the majority of phytoplankton biomass ($88 \pm 18\%$; s.d.) and therefore
176 were visualized separately.

177

178 ***Nutrient Analysis***

179 To quantify the concentration and forms of nutrients in Green Valley Lake, surface water
180 samples were collected at a depth of 0.25 m at the deep point. A subset of the water was first
181 filtered through Whatman glass fiber filters (pore size = 0.45 μm) in the field, preserved with
182 concentrated sulfuric acid to a pH of 2, and stored at 4 °C until analysis for soluble reactive
183 phosphorus (SRP) and nitrate + nitrite (NO_x). Unfiltered sample water was preserved with
184 concentrated sulfuric acid to a pH of 2 and stored at 4 °C until analysis for total phosphorus (TP)
185 and total nitrogen (TN). The ascorbic acid method was used to quantify P concentrations with
186 filtered water for SRP and unfiltered water that had undergone persulfate digestion for TP.
187 Second-derivative ultraviolet spectroscopy was used to quantify NO_x concentrations in filtered
188 samples and TN concentrations following persulfate digestion. N species were analyzed using an
189 Agilent Cary 8454 UV-VIS spectrophotometer (Agilent Technologies Inc, Santa Clara, CA,
190 USA) and P species were analyzed using a Seal Analytical AQ2 Discrete Analyzer (Seal
191 Analytical Inc. Mequon, WI, USA). For data analysis, nutrient concentrations below detection
192 were replaced with the instrument long-term method detection limit.

193 To investigate zooplankton nutrient recycling between different nutrient pools we quantified
194 the available concentrations of nitrogen (N) and phosphorus (P) as well as calculated total and
195 inorganic N and P. Nutrient limitation of phytoplankton was determined by the N:P ratio of total
196 N and P with N:P > 20 indicating P limitation (Guildford and Hecky, 2000). A decade of
197 summertime measurements of NH_x in Green Valley Lakes indicates that it is rarely detectable
198 (Supplementary Material).

199

200 ***Zooplankton Stoichiometry and Excretion***

201 To assess the contribution of zooplankton to nutrient availability we determined
202 zooplankton elemental composition, nutrient storage, and excretion rate. We estimated elemental
203 composition and total nutrient storage by zooplankton ($\text{L}^{-1} \text{d}^{-1}$) following methods described

204 previously (Moody and Wilkinson, 2019). Briefly, taxa-specific %N and %P collected from the
205 literature (Hamre, 2016; Hébert *et al.*, 2016a; Hessen *et al.*, 2007) were used to estimate total
206 nutrient storage by multiplying %N and %P by the biomass of each taxa and summing across the
207 community on each sampling date. Although we are using trait data from largely oligotrophic
208 lakes, zooplankton express fairly strong stoichiometric homeostasis (Persson *et al.*, 2010) as well
209 as low intraspecific stoichiometric variation between lakes (Prater *et al.*, 2017) and variable food
210 quality (Teurlinckx *et al.*, 2017). Thus, it is unlikely that intraspecific variation in %N and %P
211 values in our study lake will have a large influence.

212 Excretion rates of N and P by zooplankton were estimated using published allometric
213 equations (Supplementary Material). The equations relate zooplankton body size to N and P
214 excretion rates derived from a compiled dataset of marine and freshwater zooplankton species
215 (Hébert *et al.*, 2016b, 2016a). Briefly, for each sampling event we input the average dry mass per
216 zooplankton taxon into the allometric equations to determine individual N and P excretion rates
217 ($\mu\text{M N or P individual}^{-1} \text{ hour}^{-1}$) then converted the hourly excretion rate to a daily rate (day^{-1}).
218 Next, we multiplied the daily excretion rates by the density of each taxon ($\text{individuals L}^{-1}$), then
219 summed the excretion rates across taxa to calculate a zooplankton community excretion on that
220 sampling date ($\mu\text{M N or P day}^{-1}$). We calculated uncertainty in the excretion estimates by
221 propagating the variation in the slope and intercept from the allometric equations presented in
222 Hébert *et al.*, (2016b). We also calculated zooplankton excretion using other published
223 allometric equations from Wen and Peters (1994) that were derived from different underlying
224 datasets. The overall pattern of zooplankton excretion did not differ between the two methods;
225 however, the Wen and Peters (1994) based estimates of excretion were slightly higher
226 (Supplementary Table S3). We chose to use the more conservative estimate of zooplankton
227 excretion rates based on Herbert *et al.* (2016) in our analysis as the method also allowed us to
228 estimate uncertainty.

229

230 ***Data Analysis***

231 To assess the magnitude of zooplankton N and P excretion in Green Valley Lake we
232 compared the estimated concentration of excreted N and P over the course of a day to the
233 measured surface water concentrations of inorganic N and P for each sampling event. This was
234 expressed as a percent of the inorganic nutrient pool:

$$\left(\frac{\mu\text{M } N \text{ or } P \text{ excreted by zooplankton community in a day}}{\mu\text{M of inorganic } N \text{ or } P \text{ present in the surface waters}} \right) * 100 \quad (1)$$

235

236 To assess how zooplankton affected the rate of nutrient cycling over the course of the
 237 growing season we calculated the zooplankton nutrient turnover time of the inorganic P pool.
 238 Turnover, in this case, is the number of days it would take for zooplankton excretion of P,
 239 estimated for a specific sampling date to reach the concentration SRP measured on the same
 240 sampling date, assuming no uptake from phytoplankton (SRP/excretion rate). The turnover time
 241 varies depending on the rate of zooplankton excretion and concentration of SRP in the surface
 242 waters. We did not have a weekly measurement of ammonium as it was likely undetectable
 243 based on previous monitoring, so we could not calculate turnover of the soluble N pool. Short
 244 turnover times of inorganic P indicate zooplankton are significantly contributing to the inorganic
 245 P pool in Green Valley Lake. Long nutrient turnover times indicate factors other than
 246 zooplankton excretion are driving nutrient availability.

247 To assess the drivers of phytoplankton community composition across the growing season
 248 we performed a distance based-redundancy analysis (db-RDA). In order to ascertain whether
 249 zooplankton nutrient recycling partially explained phytoplankton community composition we
 250 included zooplankton excretion and body stoichiometry with other potentially important
 251 environmental variables including inorganic nutrient concentration (Filstrup and Downing,
 252 2017), temperature (Striebel *et al.*, 2016), pH (Rönicke *et al.*, 2010), and total dissolved solids
 253 (Ahmed and Wanganeo, 2015). Phytoplankton species biomass data were Hellinger transformed
 254 and species that only occurred once in the full summer dataset and contributed <1% of total
 255 biomass were removed to decrease the weight of rare species. Environmental variables were z-
 256 scored in order to correct for differences in scale and magnitude (Legendre and Legendre, 1998).
 257 The db-RDA was performed using a Bray-Curtis distance matrix using the square roots of
 258 dissimilarities to avoid negative eigenvalues (Legendre and Anderson, 1999). Missing or lost
 259 samples were removed from the final analysis. The best model was determined using forward
 260 and backward stepwise regression with model significance determined using a Monte Carlo
 261 permutation test (999 permutations, *p*-value < 0.05).

262 To investigate the importance of zooplankton top-down control we calculated the
 263 percentage of zooplankton biomass to phytoplankton biomass. The percentage of zooplankton

264 biomass in comparison to phytoplankton biomass has been used as a proxy for describing the
265 importance of top-down v. bottom-up control in lakes, with a high percentage (~40-50%)
266 indicating strong top-down control and a low percentage (~10%) indicating weak top-down
267 control (Leroux and Loreau, 2015; Havens and Beaver, 2013). Additionally, we compared the
268 size distributions of zooplankton and phytoplankton communities and individual zooplankton
269 mass, as well as their changes over time. Zooplankton length data were compiled for up to 25
270 individuals per taxa per sampling date then used to estimate body mass with taxa-specific length-
271 weight regressions (Dumont *et al.*, 1975; McCauley, 1984). When more than 25 individuals of a
272 species were present in a sample, the total number was counted, but length was only measured
273 for the first 25 individuals. The greatest axial linear dimension (GALD) of phytoplankton was
274 measured as the greatest distance across an individual cell, colony, or filament (i.e., natural unit),
275 such as would be encountered by a zooplankton grazer. Distributions of zooplankton length and
276 body mass were compared to the distribution of phytoplankton GALD for each sampling date to
277 investigate the size distribution dynamics over time in the two trophic levels. Additionally, we
278 performed a linear regression of mean phytoplankton GALD versus mean zooplankton size. All
279 analyses were performed using the statistical software R version 4.0.4 (R Core Team, 2021) with
280 the, *magrittr*, and *vegan* packages (Bach and Wickham, 2020; Oksanen *et al.*, 2020).

281

282 **RESULTS**

283 First, we assessed zooplankton and phytoplankton community composition and biomass
284 during the summer growing season. Zooplankton biomass peaked ($249 \mu\text{g L}^{-1}$) in late May and
285 early June (DOY 150-164), rapidly decreased to its minimum ($\sim 2 \mu\text{g L}^{-1}$) in mid-July to late
286 August (DOY 192 – DOY 234), then returned to early summer concentrations through
287 September (Figure 1A). The early summer zooplankton community was dominated by *Daphnia*
288 and calanoid copepods which transitioned in early July (DOY 199) to a community dominated
289 by *Chydorus* and cyclopoid copepods, then transitioning back to *Daphnia* in late August (Figure
290 1A). Zooplankton top-down control was very low (<10%) over the course of the growing season
291 with the percentage of zooplankton biomass to phytoplankton biomass peaking on DOY 164 at
292 6.9%. Similar to zooplankton biomass, phytoplankton biomass was initially high in the spring,
293 mainly composed of bacillariophytes, before rapidly decreasing during the clear-water period
294 between DOY 150 – 164 when zooplankton biomass was at its highest (Figure 1B). Following

295 DOY 172, the phytoplankton community was overwhelmingly composed of cyanophytes, mainly
296 *Microcystis*, with phytoplankton reaching peak biomass on DOY 213 (~329 mg L⁻¹). The other
297 dominant cyanophyte was the diazotroph *Aphanothece*, which was present from DOY 192 – 228.

298 Nutrient concentrations and limitation were dynamic throughout the summer (Figure 2).
299 Inorganic N concentrations were highest in the spring and decreased by 80% to low or
300 undetectable concentrations after DOY 178 (Figure 2A). At the same time, there was a rapid
301 increase in inorganic P of 394% from DOY 172 to 178 and a 937% increase from DOY 178 to
302 DOY 206 (Figure 2B). Total N:P declined rapidly in mid-July (DOY 192), transitioning the
303 ecosystem from P- to intermittent N-limitation. There was also a shift in inorganic N:P to N-
304 limitation in mid-July that was persistent for the remainder of the summer (Figure 2C).
305 Zooplankton community body N:P was the highest in mid-summer, coincident with the rapid
306 transition between P- and N-limitation, then steadily declined throughout the rest of the sampling
307 period (Figure 2D). The storage of N in zooplankton biomass was negligible over the course of
308 the growing season (Figure 2C), however P storage nearly equaled inorganic P concentrations in
309 the water column early in the summer when zooplankton biomass peaked (Figure 2D).

310 Supporting our hypothesis, zooplankton excretion contributed substantially to the
311 inorganic P pool from late May to late June (DOY 143-172), with daily excretion ranging
312 between 23-46% of the inorganic P standing stock (Figure 3A). Following DOY 172, the
313 contribution of zooplankton excretion to the inorganic P pool dropped below 1% for the
314 remainder of the sampling period. Furthermore, zooplankton excretion was contributing to a
315 rapid turnover of the inorganic P pool in early summer with turnover times ranging between 3 –
316 4 days but increased well beyond 365 days as inorganic P concentrations increased in late June
317 (Supplementary Table S4). Estimated zooplankton N excretion was never more than 3.3% of the
318 inorganic N pool over the course of the growing season (Figure 3B).

319 In support of our second hypothesis, the daily mass of zooplankton excretion
320 significantly explained variance in phytoplankton community composition during early summer
321 (DOY 143 – 164, Figure 4, Table 1). We found that the db-RDA discriminated the
322 phytoplankton community into distinct communities defined by pre- and post-cyanobacterial
323 dominance ($F=2.44$, $p=0.003$). A suite of environmental variables explained 41.9% of the
324 variation in phytoplankton community composition including inorganic P ($p=0.003$) and N
325 ($p=0.020$) concentrations, zooplankton P excretion ($p=0.004$), temperature ($p=0.032$), and total

326 dissolved solids ($p=0.046$). Phytoplankton community composition was correlated with total
327 dissolved solids, zooplankton N and P excretion, and inorganic N concentrations in early
328 summer prior to the cyanobacteria bloom beginning on DOY 172 where phytoplankton
329 community composition became more correlated with inorganic P concentrations and
330 temperature. However, the significant axes did not explain a large proportion of the variation
331 with the first axis explaining 28.4% and the second axis explaining only 13.4% of variation.

332 Furthermore, there was evidence that zooplankton were influencing the size distribution
333 of phytoplankton GALD in mid- to late summer. There was a higher average GALD when mean
334 zooplankton length was at its lowest (Figure 5A), and the density of smaller zooplankton
335 individuals began increasing relative to the total size distribution in early June (DOY 164). Small
336 individual zooplankton dominated the zooplankton size distribution from late June to early
337 August (DOY 178 – 213). This was concurrent with a period in which larger phytoplankton
338 dominated the GALD distribution (Figure 5A). Phytoplankton average GALD was greatest in
339 July (mean = $32.5 \pm 19.6 \mu\text{m}$; s.d.) concurrent with the period of the summer where zooplankton
340 average length was at its lowest (mean = $171 \pm 102 \mu\text{m}$; s.d.). In late July through August the
341 difference in zooplankton length and phytoplankton GALD steadily increased, surpassing the
342 mean differences observed in early summer (Figure 5B). A similar pattern was observed between
343 phytoplankton GALD and zooplankton dry mass (Supplementary Figure S2). There was a weak
344 negative relationship between GALD and zooplankton length ($p=0.0119$, $R^2=0.42$;
345 Supplementary Figure S3A), and zooplankton body mass ($p=0.0306$, $R^2=0.33$; Supplementary
346 Figure S3B).

347

348 **DISCUSSION**

349 *Effect of zooplankton excretion on nutrient availability*

350 We found that zooplankton excretion contributed substantially to the inorganic P pool in
351 Green Valley Lake, but only during the early summer (DOY 143 – 178). It was during this
352 period that inorganic P was at low concentrations in the water column and phytoplankton growth
353 was likely P-limited, indicating that zooplankton-mediated recycling contributed to meeting
354 nutrient demand by phytoplankton during this time. This early-season P availability, facilitated
355 by zooplankton recycling, may have helped initialize the cyanotoxin-producing cyanobacteria
356 bloom that flourished later in the season and persisted until late summer (Isles and Pomati,

357 2021). The large contribution of zooplankton to inorganic P availability is consistent with the
358 hypothesis from Moody and Wilkinson (2019) that N-rich zooplankton communities,
359 characteristic of hypereutrophic lakes, contribute to increased P availability within nutrient-rich
360 ecosystems. Our study in Green Valley Lake, however, revealed that this zooplankton-mediated
361 flux of P is mainly confined to the early part of the growing season.

362 Beginning on DOY 178, the concentration of inorganic P in Green Valley Lake increased
363 substantially, diminishing the importance of zooplankton-recycled P, and driving the ecosystem
364 to co-limitation or N-limitation for the rest of the season. The transition between P and N-
365 limitation or co-limitation is a dynamic that has been reported in other eutrophic and
366 hypereutrophic ecosystems (Andersen *et al.*, 2020; Wang *et al.*, 2019). The transition is likely a
367 result of increased internal P loading (Albright and Wilkinson, 2022) and differences in N uptake
368 strategies and preferences in the phytoplankton community, particularly when cyanobacteria
369 begin to dominate (Li *et al.*, 2020; Glibert *et al.*, 2016). Furthermore, our estimates of P turnover
370 by zooplankton indicated rapid turnover of inorganic pools during early summer, but drastically
371 slowed once inorganic P concentrations rose. These results support our conclusions that
372 zooplankton nutrient recycling was an important P flux during the early summer growing season,
373 but not an important flux once internal loading increased P availability. Additionally, it is
374 unlikely zooplankton had much influence over the turnover of total P and N across the entire
375 summer, likely due to the lack of top-down control by the zooplankton community on
376 phytoplankton throughout the summer growing season.

377 Overall, the contribution of zooplankton-recycling to the inorganic N pool in Green
378 Valley Lake was never greater than 5%. However, the uptake of ammonium from zooplankton
379 excretion by phytoplankton may have been too fast to result in a measurable concentration,
380 masking the contribution of zooplankton excretion to N availability. Alternatively, we may be
381 underestimating N excretion given that our estimates of zooplankton excretion were not taxon-
382 specific but instead were based on a consolidated dataset of both cladocerans and copepods. This
383 is particularly true when Cladocera dominate in the early and late-summer periods, which could
384 increase community N excretion as Cladocera retain more P than N due largely to their body
385 stoichiometry (Elser *et al.*, 1988). Overall, our estimates of zooplankton excretion were low
386 relative to the concentrations of inorganic nutrients in the ecosystem across the summer;

387 however, they were comparable with other studies using similar allometric equations (Conroy *et*
388 *al.*, 2005) or direct measurement (den Oude and Gulati, 1988) in eutrophic ecosystems.

389 In addition to zooplankton, other consumers can play a key role in nutrient recycling in
390 eutrophic ecosystems, particularly detritivores and planktivores such as gizzard shad (Sharitt *et*
391 *al.*, 2021; Vanni *et al.*, 2006) and mussels (Arnott and Vanni, 1996). However, neither gizzard
392 shad nor zebra mussels have been reported in Green Valley Lake. While we did not quantify the
393 contribution of nutrient recycling by other consumers to availability in Green Valley Lake, these
394 organisms certainly contributed. There is a common carp (*Cyprinus carpio*) population in Green
395 Valley Lake which can influence nutrient cycling through bioturbation and excretion (Weber and
396 Brown, 2009); however, the population is small. We hypothesize that the contributions of fish
397 and other organisms would have a similar seasonality given the large contribution of internal P in
398 the latter half of the season.

399

400 ***Role of zooplankton excretion and grazing on phytoplankton community structure***

401 The redundancy analysis of phytoplankton community composition showed that the daily
402 estimated rate of zooplankton P excretion was related to variation in phytoplankton community
403 composition prior to the cyanobacteria bloom later in summer. This suggests that zooplankton P
404 recycling, in part, influenced phytoplankton biomass and composition. The phytoplankton
405 community was dominated by bacillariophytes and chlorophytes until DOY 172 when
406 cyanophytes dominated the community. This transition is reflected in the discrimination of the
407 phytoplankton community between pre- and post-cyanobacterial dominance in the redundancy
408 analysis (db-RDA). The early summer phytoplankton community was also significantly related
409 to the concentration of inorganic N and total dissolved solids. This corresponds with the seasonal
410 dynamic of nutrient limitation as both chlorophytes and bacillariophytes perform well under P-
411 limitation (Berg *et al.*, 2003). Furthermore, the inorganic N pool was highest in early summer
412 and predominantly composed of nitrate which can be taken up and used by bacillariophytes
413 (Andersen *et al.*, 2020). The higher concentrations of total dissolved solids in the spring are
414 reflective of the input of ions from the watershed with spring melt, including nutrients such as
415 nitrate. These inputs combined with zooplankton excretion drove the community composition in
416 the spring.

417 Beginning on DOY 172, the phytoplankton community was overwhelmingly dominated
418 by *Microcystis* leading to very stable community composition during mid- and late summer. The
419 mid- to late-summer phytoplankton community was significantly related to temperature and
420 inorganic P concentrations, consistent with other studies describing increasing temperature and
421 N-limitation as key drivers of cyanobacteria dominance (Bogard *et al.*, 2020; Hayes *et al.*, 2020).
422 In fact, diazotrophic cyanobacteria (specifically, *Aphanothece spp.*) did not appear in the Green
423 Valley phytoplankton community until the onset of N-limitation. Other environmental factors
424 were likely influencing the phytoplankton community as the db-RDA described only 41.9% of
425 variation in the phytoplankton community composition. Phytoplankton community turnover is a
426 complex phenomenon driven by a multitude of environmental factors (Wentzky *et al.*, 2020;
427 Sommer *et al.*, 2012), including nutrient and light availability, the latter of which we did not
428 measure. Given the high biomass of phytoplankton, light limitation through self-shading likely
429 played a significant role in phytoplankton dynamics.

430 While we did not observe significant top-down control of zooplankton on phytoplankton
431 growth based on the Z:P ratios, we did find evidence that zooplankton may have influenced the
432 size structure of the phytoplankton community. The negative relationship between zooplankton
433 length and phytoplankton GALD is consistent with other studies in hypereutrophic lakes
434 indicating that small-bodied zooplankton preferentially graze on small-sized phytoplankton,
435 favoring growth of large filamentous and colonial phytoplankton (Bairagi *et al.*, 2019; Onandia
436 *et al.*, 2015). This is evidenced by the large bloom of *Microcystis* colonies midsummer that drove
437 the increase in phytoplankton GALD we observed in July through early August. It is likely that
438 smaller-bodied zooplankton were contributing, in part, to the dominance of *Microcystis* colonies
439 and higher phytoplankton GALD. By grazing on smaller sized phytoplankton cells or colonies,
440 zooplankton can eliminate smaller phytoplankters leaving a greater proportion of individuals
441 with large GALD to dominate the overall size distribution. The size structure of communities is
442 closely tied to food web structure and energy flow (Brose *et al.*, 2017), indicating that the
443 influence of zooplankton on phytoplankton size structure was influential for the transfer, uptake,
444 and recycling of nutrients by phytoplankton. However, it is unlikely zooplankton were the sole
445 cause of increased phytoplankton GALD as the drawdown of inorganic N we observed
446 midsummer coincided with the bloom of *Microcystis* beginning on DOY 172. Inorganic N is

447 known to promote *Microcystis* growth and was likely influencing the proliferation of *Microcystis*
448 colonies (Chen *et al.*, 2019).

449 It is also likely that microzooplankton and ciliates played an important role grazing on
450 small phytoplankton species; however, we did not quantify these communities in this study.
451 Furthermore, our phytoplankton counting methods were unable to facilitate the identification of
452 nano- or picophytoplankton species in the water column. Microzooplankton, nano- and
453 picophytoplankton are increasingly recognized as key components of the plankton food web and
454 contribute a significant percentage of grazing pressure on phytoplankton in highly productive
455 ecosystems (Agasild *et al.*, 2007; Zingel *et al.*, 2007). Future studies should examine their
456 seasonal dynamics and potential contribution to ecosystem processes more thoroughly as they
457 can be key components of zooplankton-phytoplankton interactions in nutrient-rich lakes.

458

459 **CONCLUSIONS**

460 While the importance of consumer-driven nutrient recycling has been demonstrated in
461 less eutrophic waterbodies, the role that zooplankton consumers play in nutrient availability and
462 phytoplankton dynamics in hypereutrophic lakes remained unclear. Our results support a
463 previous comparative study indicating that zooplankton community composition may influence
464 nutrient availability in hypereutrophic ecosystems, as well extend our understanding of the
465 temporal dynamics of zooplankton and phytoplankton interactions. We found evidence of the
466 importance of zooplankton nutrient cycling in a hypereutrophic reservoir with zooplankton
467 excretion providing a large portion of the available P early in the summer, prior to the onset of
468 the cyanobacteria-dominated bloom later in the season. Additionally, zooplankton influenced the
469 early summer phytoplankton community composition through excretion as well as phytoplankton
470 size structure, particularly later in the summer when cyanobacteria were blooming. As
471 demonstrated here, the role of zooplankton nutrient recycling in hypereutrophic lakes is an
472 important component of phytoplankton dynamics and ecosystem function that should be
473 considered in greater detail. Unlike previous assumptions that zooplankton do not contribute
474 substantially to nutrient cycling and phytoplankton dynamics, our results suggest that
475 zooplankton do in fact do contribute to those dynamics, predominantly for a short period early in
476 the summer. Future work should investigate the dynamics of zooplankton nutrient recycling

477 across different climate contexts and over longer time periods, including dynamics through
478 winter and autumn.

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484

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492

493 **DATA ARCHIVING**

494 The data for this study will be archived using the Environmental Data Initiative repository and
495 given a unique digital object identifier. Data will be uploaded in a comma delimited file format
496 with metadata composed of contact information, detailed variable descriptions, complete
497 taxonomic information, temporal resolution, and descriptions for a given variable when
498 appropriate. Metadata will follow the ecological metadata language and be published under a
499 creative commons license. Scripts for data analysis and figure generation will be available and
500 maintained online through GitHub (<https://github.com/tjbutts/hyper-plankton>) and will
501 eventually be published in Zenodo for long-term storage.

502

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660

661 **TABLE & FIGURE LEGENDS**

662 **Table 1.** Statistics for the distance based-Redundancy Analysis (db-RDA) of phytoplankton
663 community composition in Green Valley Lake from May to September 2019.

664
665 **Figure 1.** (A) Zooplankton biomass and community composition and (B) phytoplankton biomass
666 and community composition over the course of the growing season in Green Valley Lake, IA.

667
668 **Figure 2.** (A) Surface water nitrogen and (B) phosphorus concentrations split between total,
669 inorganic, and zooplankton body storage over the course of the growing season. Additionally,
670 (C) surface water molar nitrogen: phosphorus (N:P) ratios split between total and inorganic pools
671 as well as (D) molar N:P ratios of the zooplankton community.

672
673 **Figure 3.** The estimated concentration of total zooplankton community excretion produced over
674 a day compared with the surface water inorganic phosphorus and nitrogen concentrations
675 measured the same day as a percentage. Estimates of zooplankton excretion were derived from
676 published allometric equations of zooplankton body size and excretion rate (Hébert, *et al.*, 2016).
677 The dark lines represent the estimated excretion of either phosphorus or nitrogen, and the shaded
678 area represents the error associated with the estimate for each sampling day.

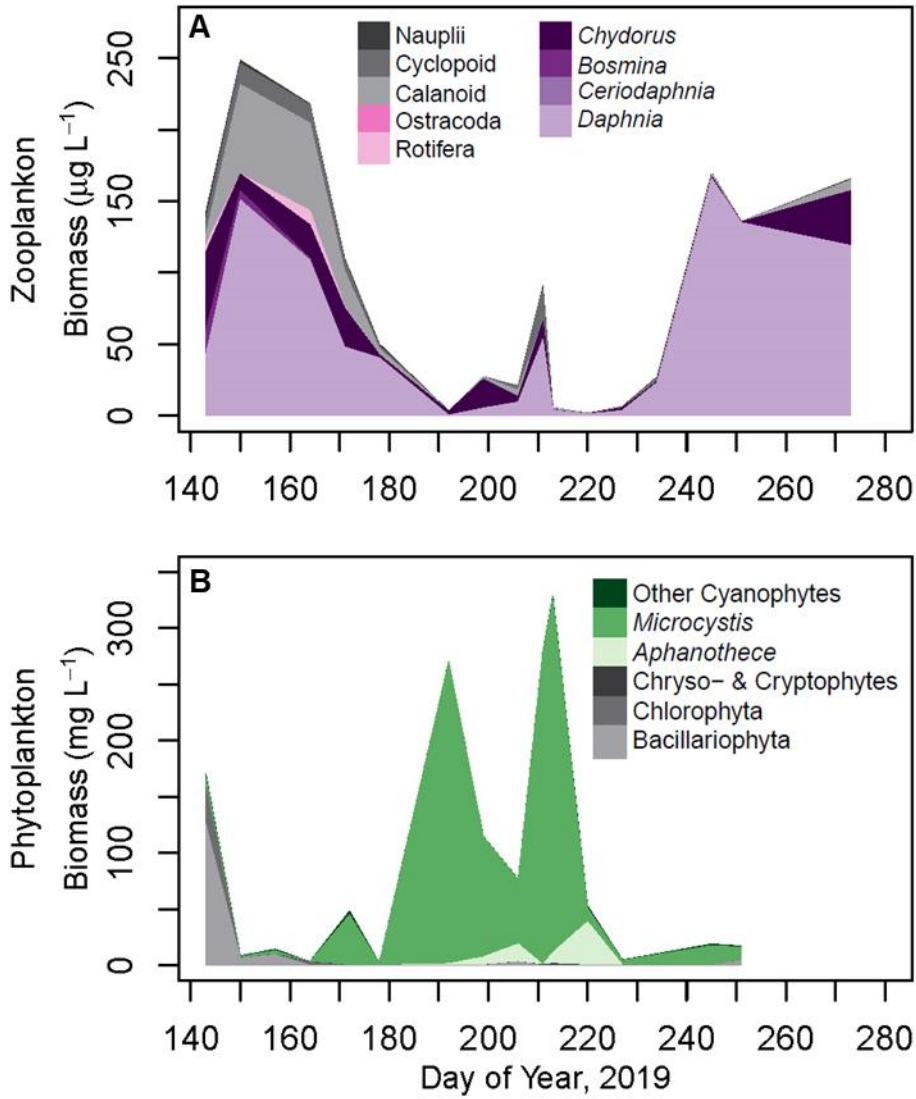
679
680 **Figure 4.** Distance based-Redundancy Analysis (db-RDA) of the phytoplankton community in
681 Green Valley Lake from May to September 2019. Dots represent sampling points, and the
682 numbers 1-14 are DOY 143, 150, 164, 172, 178, 192, 199, 206, 211, 213, 220, 227, 245, 251,
683 respectively. DOY 245 (13) was omitted from the diagram as there were no available data for
684 inorganic N and P thus the data were omitted from the analysis. The explanatory variables are
685 represented by arrows with significance denoted by an asterisk.

686
687 **Figure 5.** (A) Density ridgeline plots of phytoplankton greatest axial distance (GALD, μm) and
688 zooplankton body size (μm) over the course of the growing season in Green Valley Lake, IA.
689 The black vertical line within each distribution represents the mean. (B) Mean difference
690 between zooplankton length and phytoplankton GALD. DOYs that are missing either
691 phytoplankton GALD or zooplankton length are the result of sample loss or no available data.

692 **TABLES**

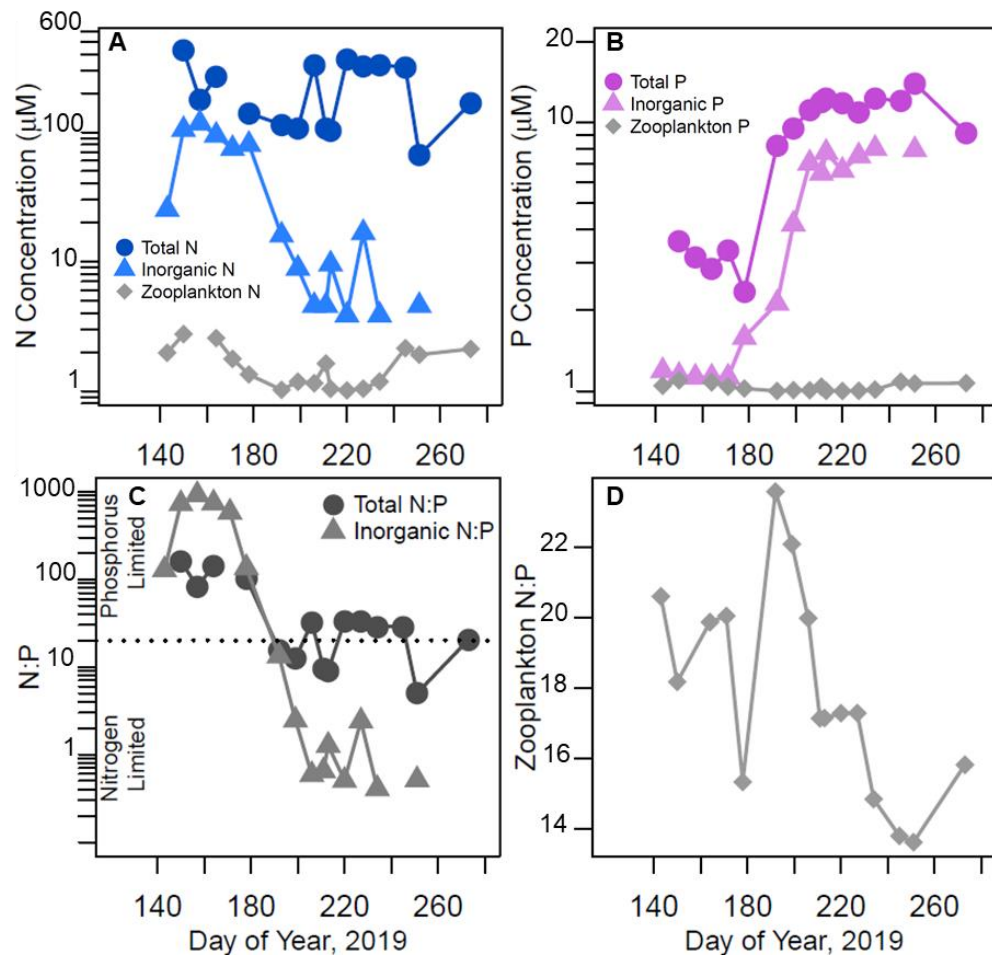
693 Table 1.

| Permutation test variable | Sums of Squares | pseudo-<i>F</i> | <i>p</i>-value |
|----------------------------------|------------------------|------------------------|-----------------------|
| Full model | 2.58 | 2.44 | 0.003 |
| First axis | 1.03 | 5.87 | 0.002 |
| Second axis | 0.49 | 2.76 | 0.030 |
| Inorganic P | 0.54 | 3.07 | 0.003 |
| Inorganic N | 0.41 | 2.32 | 0.020 |
| Zooplankton P excretion | 0.58 | 3.27 | 0.004 |
| Zooplankton N excretion | 0.28 | 1.59 | 0.099 |
| Temperature (°C) | 0.37 | 2.12 | 0.032 |
| Total Dissolved Solids | 0.40 | 2.28 | 0.046 |
| Residual | 1.06 | | |



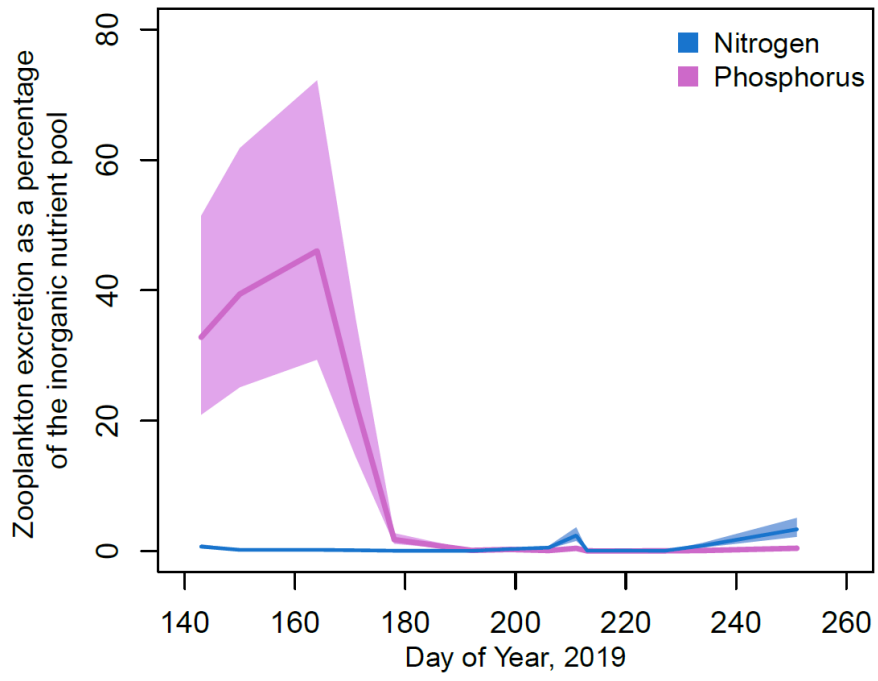
695

696 Figure 1.



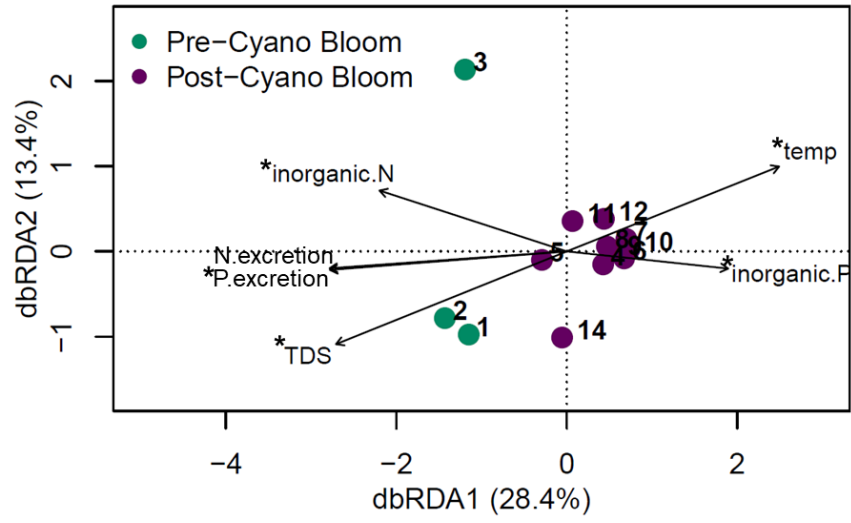
697

698 Figure 2.



699

700 Figure 3.

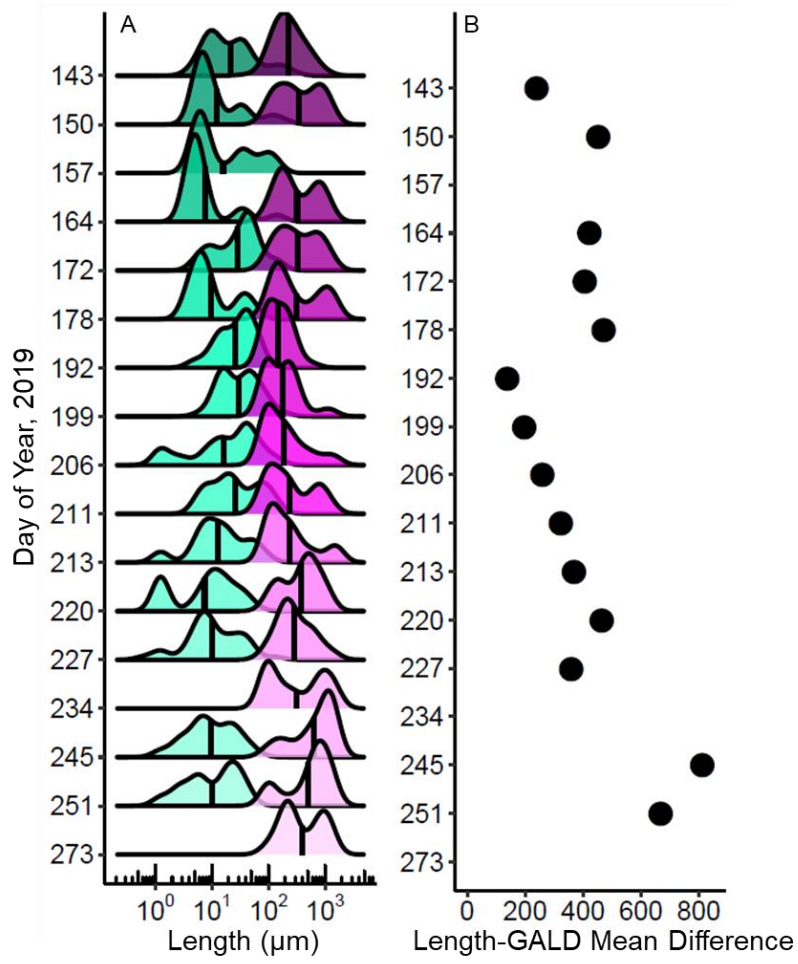


701

702 Figure 4.

703

■ Phytoplankton GALD ■ Zooplankton Length



705

706 Figure 5.

707 **Supplementary Material**

708 *Nutrient concentrations and speciation*

709 The following equations describe how we defined the major fractions of nitrogen (N) and
710 phosphorus (P) in Green Valley Lake. Total N in freshwater is composed organic and inorganic
711 fractions:

$$TN = orgN + DIN \quad (1)$$

712 where TN is total N, $orgN$ is organic N in both the particulate (organisms and detritus) and
713 dissolved (urea) form, and DIN is dissolved inorganic N composed of NOx and NHx representing
714 nitrate + nitrite and ammonium + ammonia, respectively. Previous data from the last decade in
715 Green Valley Lake indicated NHx were extremely low or undetectable in the surface waters
716 during the summer months. If we assume that NHx is undetectable (1) simplifies to:

$$TN = orgN + NOx \quad (2)$$

717 allowing calculation of $orgN$ by rearranging (2):

$$orgN = TN - NOx \quad (3)$$

718 Thus, we could characterize N pools as total (TN) representing dissolved and particulate forms of
719 N, organic ($orgN$) representing dissolved organic N (urea) and seston, and inorganic N (NOx)
720 representing DIN in the surface waters. For our analyses we focused on the TN and DIN pools.

721 Similarly, P is composed of organic and inorganic fractions in reservoir surface waters:

$$TP = POP + PIP + DIP + DOP \quad (4)$$

722 where TP is total P, POP is particulate organic P, PIP is particulate inorganic P, DIP is dissolved
723 inorganic P, and DOP is dissolved organic P. DIP and PIP were both present within the water
724 column, but our focus for this study was on DIP which is far more bioavailable to phytoplankton
725 than PIP (Zhou *et al.*, 2005) and thus more influential to nutrient cycling via zooplankton-
726 phytoplankton interactions. Previous data from the last decade in Green Valley Lake indicated
727 PIP was extremely low or undetectable in the surface waters during the summer months. Thus,
728 (4) can be simplified by combining DOP and POP to one organic pool ($orgP$) and using SRP as
729 a measure of DIP over the course of the growing season:

$$TP = orgP + SRP \quad (5)$$

730 Therefore, we could characterize P pools as total (*TP*) representing dissolved and particulate
 731 forms of P, organic (*orgP*) representing dissolved organic P and seston, and inorganic (*SRP*)
 732 representing *DIP* in the surface waters. For our analyses we focused on the TP and SRP pools.

733 Ammonium + ammonia (NH_x) (EPA method 103-A v6) and inorganic suspended solids
 734 were measured at the same location in the lake three times during the summer by the Iowa
 735 Ambient Lakes Monitoring program (IDNR 2021). Ammonium was analyzed through the
 736 alkaline phenate method on a Seal Analytical AQ2 Discrete Analyzer and inorganic particulates
 737 were determined via difference between total and volatile suspended solids (USGS method I-
 738 3765-85).

739

740 *Zooplankton excretion equations*

741 Individual zooplankton excretion of P was determined using the following equation from Hébert
 742 *et al.*, (2016):

$$\ln(P_{exc,h}) = 2.50 + (0.84\ln(Z_{BS})) \quad (6)$$

743 where $P_{exc,h}$ is excreted P (nM of P individual⁻¹ hour⁻¹) and Z_{BS} is the dry mass of an individual
 744 zooplankter (mg). Zooplankton excretion of N was determined in a similar manner:

$$\ln(P_{exc,h}) = 0.56 + (0.70\ln(Z_{BS})) \quad (7)$$

745 where $N_{exc,h}$ is excreted N (nM of N individual⁻¹ hour⁻¹).

746 Data were then converted to μM of N or P per day using the following conversions:

$$\frac{nmol\ N\ or\ P}{individual \cdot hour} \cdot \frac{24\ hours}{1\ day} \cdot \frac{individuals}{L} \cdot \frac{1\ \mu mol}{1000\ nmol} = \frac{\mu M\ N\ or\ P}{day} \quad (8)$$

747 The allometric equations were derived from a combined dataset of marine and freshwater
 748 zooplankton. Using only the freshwater data did not significantly change the slope, nor was the
 749 relationship between excretion and body size significant due to the much smaller sample size.

750 Thus, we only present the combined freshwater and marine model as presented in Hébert *et al.*
 751 (2016). Additionally, we used zooplankton excretion equations from Wen and Peters (1994).

752 Specifically, we used their multivariate regression equations for crustacean zooplankton which
 753 corrected for temperature (K) and experimental duration (h) in their estimates of excretion. As

754 our data did not have an experimental duration, we dropped the experimental duration correction
755 resulting in the following equations:

$$\text{Log}_{10}(P_{exc,wp}) = -5.28 + (0.61 * \text{log}_{10}(Z_{BS})) + (0.01 * T) \quad (9)$$

756 Where $P_{exc,wp}$ is excreted P ($\mu\text{g d}^{-1}$), Z_{BS} is the body size of an individual zooplankter (μg), and T
757 is water temperature (K). Similarly, for N excretion:

$$\text{Log}_{10}(N_{exc,wp}) = -3.47 + (0.74 * \text{log}_{10}(Z_{BS})) + (0.00002 * T^2) \quad (10)$$

758 Where $N_{exc,wp}$ is excreted N ($\mu\text{g d}^{-1}$), Z_{BS} is the body size of an individual zooplankter (μg), and T
759 is water temperature (K). The pattern of zooplankton excretion was consistent between the two
760 methods; however, the magnitude of excretion was different (Supplementary Table S3).

761

762 SUPPLEMENTARY REFERENCES

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766 Assessment Section. AQuIA [database].

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768 rates by zooplankton. *Limnol. Oceanogr.*, **39**, 1669–1679.

769 Zhou, A. *et al.* (2005) Phosphorus adsorption on natural sediments: Modeling and effects of pH
770 and sediment composition. *Water Res.*, **39**, 1245–1254.

771

772 *Tables*

773 **Table S1.** Zooplankton genera, order, or class identified over the course of the growing season in
774 Green Valley Lake.

| Taxonomic Group | Taxa identified in Green Valley Lake included in grouping |
|------------------------|---|
| Large Cladocera | <i>Daphnia</i> <i>Simnocephalus</i> <i>Ceriodaphnia</i> |
| Small Cladocera | <i>Bosmina</i> <i>Chydorus</i> |
| Ostracod | Ostracoda |
| Calanoids | Calanoida |
| Cyclopoids | Cyclopoida |
| Nauplii | Copepod nauplii |
| Rotifers | <i>Asplanchna</i> <i>Keratella cochlearis</i> <i>Keratella quadrata</i> <i>Pompholyx</i> <i>Trichocerca</i> <i>Filinia</i> |

775

776 **Table S2.** Phytoplankton genera identified over the course of the growing season in Green
 777 Valley Lake.

| Taxonomic Group | Taxa identified in Green Valley Lake included in grouping |
|------------------------------------|--|
| Bacillariophyta | <i>Asterionella</i> |
| | <i>Fragilaria</i> |
| | <i>Stephanodiscus</i> |
| | <i>Unknown pennate bacillariophyte</i> |
| | <i>Unknown centric bacillariophyte</i> |
| Chlorophyta | <i>Chalmydomonas</i> |
| | <i>Coelastrum</i> |
| | <i>Cosmarium</i> |
| | <i>Desmodesmus</i> |
| | <i>Elakatothrix</i> |
| | <i>Eudorina</i> |
| | <i>Monoraphidium</i> |
| | <i>Oocystis</i> |
| | <i>Pediastrum</i> |
| | <i>Schroederia</i> |
| | <i>Staurastrum</i> |
| Unknown chlorophyte | |
| Chyrso - & Cryptophytes | <i>Mallomonas</i> |
| | <i>Cryptomonas</i> |
| | <i>Komma</i> |
| <i>Aphanothece</i> (Cyanophyte) | <i>Aphanothece</i> |
| <i>Microcystis</i> (Cyanophyte) | <i>Microcystis</i> |
| | <i>Microcystis (Single-celled)</i> |
| Other Cyanophytes | <i>Aphanizomenon</i> |
| | <i>Aphanocapsa</i> |
| | <i>Merismopedia</i> |

Planktolyngbya

Pseudanabaena

Snowella

Woronichinia

Dolichospermum

779 **Table S3.** Estimated zooplankton excretion of N and P ($\mu\text{M d}^{-1}$) using different published
780 allometric equations from Hébert *et al.* (2016) and Wen and Peters (1994). Uncertainty estimates
781 derived from the allometric equation parameters in Hébert *et al.* (2016) are presented in
782 parentheses.

| Zooplankton Excretion ($\mu\text{M N or P day}^{-1}$) | | | | |
|---|----------------------|--------------|----------------------|--------------|
| DOY | Nitrogen Excretion | | Phosphorus Excretion | |
| | Hébert | Wen & Peters | Hébert | Wen & Peters |
| 143 | 0.159 (0.143- 0.242) | 0.073 | 0.062 (0.040-0.100) | 0.080 |
| 150 | 0.177 (0.116-0.270) | 0.082 | 0.056 (0.036-0.088) | 0.072 |
| 164 | 0.167 (0.110-0.255) | 0.083 | 0.058 (0.037-0.091) | 0.081 |
| 171 | 0.087 (0.057-0.133) | 0.039 | 0.029 (0.018-0.045) | 0.036 |
| 178 | 0.034 (0.022-0.051) | 0.014 | 0.010 (0.007-0.016) | 0.012 |
| 192 | 0.003 (0.002-0.004) | 0.002 | 0.001 (0.001-0.002) | 0.002 |
| 199 | 0.022 (0.014-0.033) | 0.012 | 0.008 (0.005-0.012) | 0.011 |
| 206 | 0.015 (0.010-0.022) | 0.007 | 0.005 (0.003-0.007) | 0.006 |
| 211 | 0.068 (0.045-0.104) | 0.035 | 0.023 (0.014-0.035) | 0.032 |
| 213 | 0.004 (0.002-0.005) | 0.002 | 0.001 (0.001-0.007) | 0.001 |
| 220 | 0.001 (0.001-0.002) | 0.001 | 0.000 (0.000-0.002) | 0.001 |
| 227 | 0.005 (0.003-0.007) | 0.002 | 0.002 (0.001-0.003) | 0.002 |
| 234 | 0.018 (0.012-0.027) | 0.008 | 0.005 (0.003-0.008) | 0.007 |
| 245 | 0.109 (0.072-0.167) | 0.046 | 0.031 (0.020-0.049) | 0.037 |
| 251 | 0.095 (0.062-0.145) | 0.042 | 0.029 (0.019-0.046) | 0.036 |
| 273 | 0.120 (0.079-0.183) | 0.051 | 0.039 (0.025-0.061) | 0.046 |

783

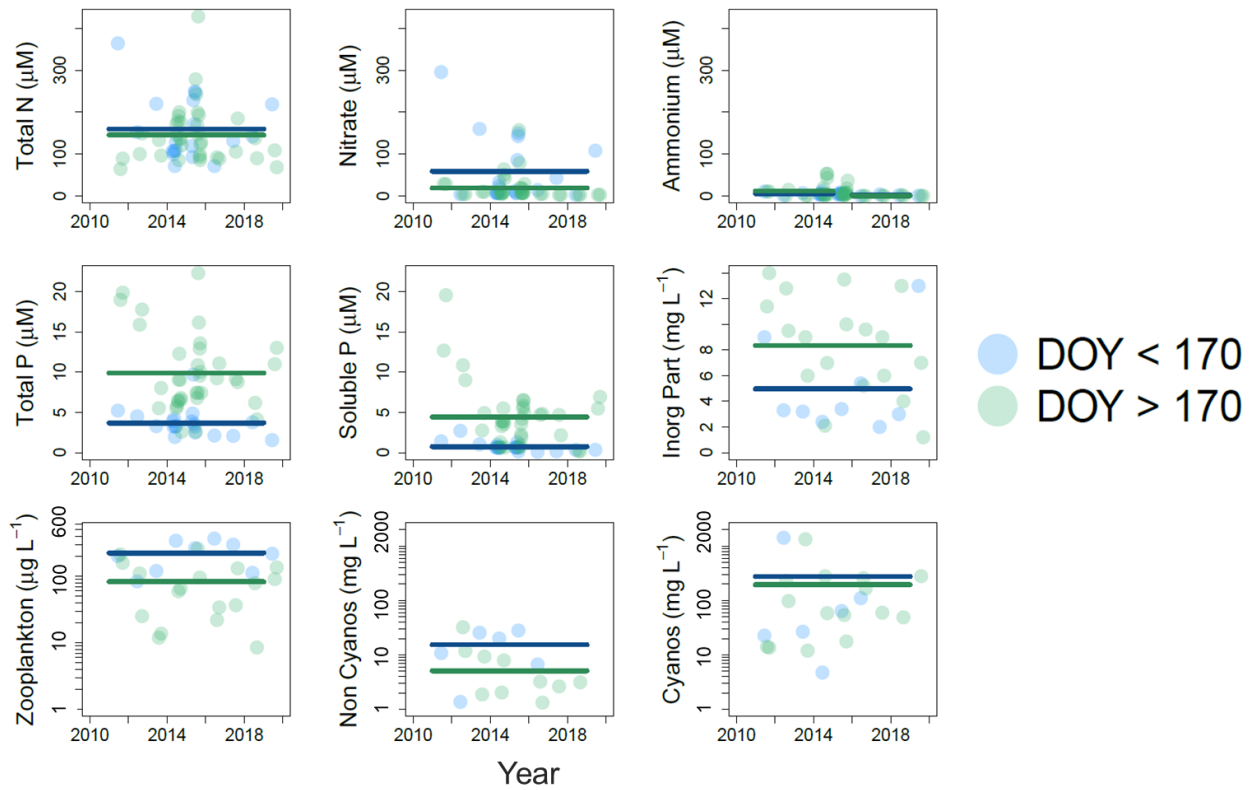
784

785 **Table S3.** Potential zooplankton nutrient turnover of various nutrient pools in Green Valley Lake
 786 representing the number of days it would take zooplankton excretion alone to meet the water
 787 column concentration of total phosphorus, total nitrogen, or inorganic phosphorus on a given
 788 sampling day. Missing values were the result of sample loss or the lack of available data and are
 789 denoted by NA.

| <i>Nutrient Pool</i> | <i>DOY</i> <i>143</i> | <i>DOY</i> <i>150</i> | <i>DOY</i> <i>164</i> | <i>DOY</i> <i>172</i> | <i>DOY</i> <i>178</i> | <i>DOY</i> <i>192 - 273</i> |
|----------------------|--------------------------|--------------------------|--------------------------|--------------------------|--------------------------|--------------------------------|
| Soluble Phosphorus | 3 | 3 | 2 | 4 | 57.3 | >365 |

790

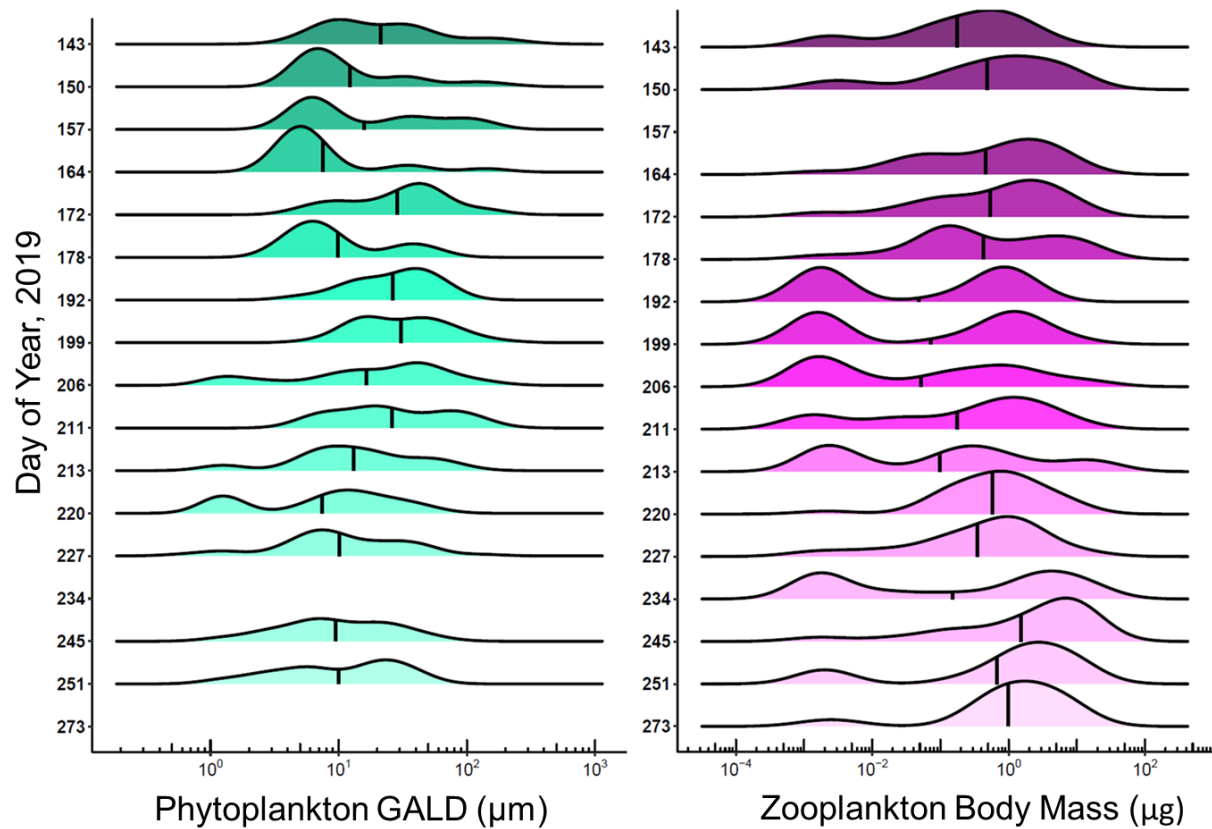
791 **FIGURES**



792

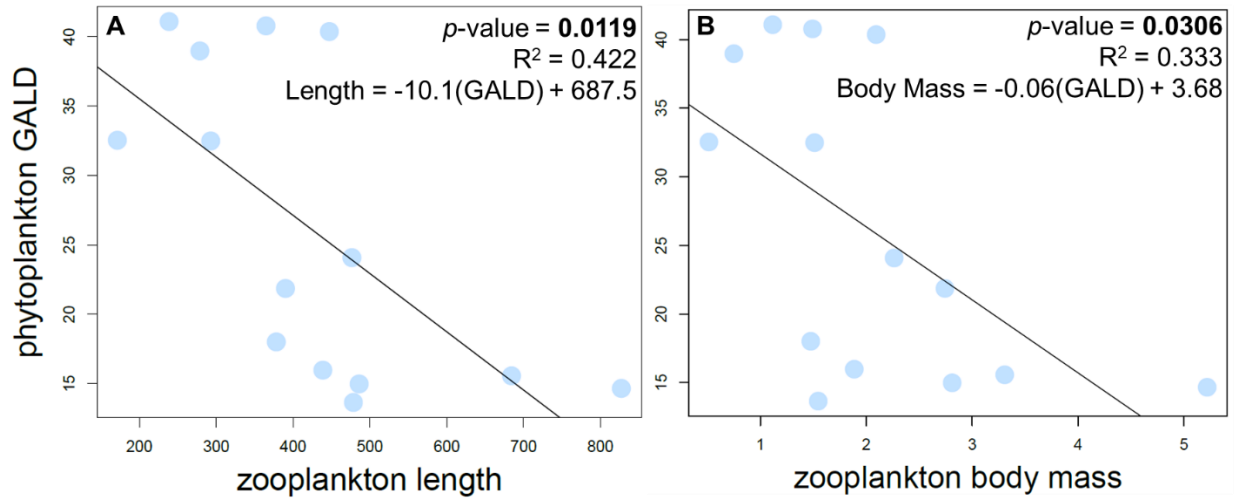
793 **Figure S1.** Historical water quality and plankton data for Green Valley Lake. The solid
 794 horizontal line is the mean value for the period 2011 – 2019 split between before or after the
 795 clear-water period which we determined was around DOY 170 using a breakpoint analysis. From
 796 left to right, top to bottom the variables represented are total nitrogen, nitrate, ammonium, total
 797 phosphorus, soluble reactive phosphorus, inorganic particulates, zooplankton biomass, non-
 798 Cyanophyta biomass, and Cyanophyta biomass. Data were collated from the Ambient Lakes
 799 Monitoring program in the state of Iowa (IDNR, 2021). Ammonium concentrations became
 800 extremely low or undetectable past 2015 and thus the mean value was split between pre- and
 801 post-2015.

802



803

804 **Figure S2.** Density ridgeline plots of phytoplankton greatest axial distance (GALD, μm) and
 805 zooplankton body mass (μg) over the course of the growing season in Green Valley Lake, IA.
 806 The black vertical line within each distribution represents the mean. DOYs that are missing
 807 either phytoplankton GALD or zooplankton length are the result of sample loss or no available
 808 data.



809

810 **Figure S3.** Linear regression of (A) zooplankton body length (μm) and (B) zooplankton body
 811 mass (μg) by phytoplankton greatest axial linear distance (GALD, μm).