

Warming alters food web-driven changes in the CO₂ flux of experimental pond ecosystems

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21	Evidence shows the important role biota play in the carbon cycle, and strategic
22	management of plant and animal populations could enhance CO ₂ uptake in aquatic
23	ecosystems. However, it is currently unknown how management-driven changes to
24	community structure may interact with climate warming and other anthropogenic
25	perturbations to alter CO ₂ fluxes. Here we show that under ambient water temperatures,
26	predators (Three-Spined Stickleback) and nutrient enrichment synergistically increased
27	primary producer biomass, resulting in increased CO2 uptake by mesocosms in early
28	dawn. However, a 3°C increase in water temperatures counteracted positive effects of
29	predators and nutrients leading to reduced primary producer biomass, and a switch from
30	CO ₂ influx to efflux. This confounding effect of temperature demonstrates that climate
31	scenarios must be accounted for when undertaking ecosystem management actions to
32	increase biosequestration.

33

34 Keywords: trophic cascades, biosequestration, carbon cycling, climate change

35 1. INTRODUCTION

In freshwater ecosystems, projected increases in water temperatures (3-5°C) [1] are likely to interact with bottom-up and top-down processes to modify community structure [2] and CO₂ dynamics [3,4]. Heterotrophs contribute to the net carbon balance of ecosystems by consuming organic matter and respiring it as CO₂. Predators and herbivores can further influence carbon balance by directly or indirectly (via trophic cascades) shifting the balance between heterotrophic respiration and photosynthesis. However, warming and eutrophication can modify food web structure by increasing species extinctions, especially at higher trophic levels [5], altering species

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interactions. Phytoplankton blooms caused by eutrophication and trophic cascades can only
enhance long-term carbon storage if plant matter escapes mineralization and is buried in
sediments; however, higher water temperatures increase metabolism and remineralization rates.
Because temperature modifies food web structure with consequences for CO₂ assimilation and
remineralization [6], alterations to animal and plant populations could cause complex climate
feedbacks in a warmer world.

49 We tested two hypotheses of how elevated water temperatures could alter the effects of top-down and bottom-up manipulations on the CO₂ flux of freshwater pond mesocosms. 50 51 Mesocosm food webs contained phytoplankton, periphyton, zooplankton, benthic 52 macroinvertebrates and in treatments containing fish, Gasterosteus aculeatus (Three-Spined Stickleback; Table S1). First, warming would increase the strength of trophic cascades and 53 54 decrease mesocosm CO₂ emissions. We predicted that warming would increase indirect positive effects of predators on primary production [7,8] leading to higher CO_2 influx in treatments 55 containing predominantly odd-numbered food chains with top predators. Second, warming 56 would alter interactive effects between eutrophication and predators on primary producers and 57 CO_2 flux. In a previous study we described the temporal response of phytoplankton to the same 58 three variables, showing that warming reduced the positive effects of nutrients on primary 59 producers [7]. We predicted that this interaction between nutrients and warming would lead to 60 increased CO₂ emissions [9]. However, because our previous study also showed that warming 61 enhanced top-down control [7], it is difficult *a priori* to predict the combined effects of all three 62 stressors on CO₂ flux. 63

64 2. MATERIALS AND METHODS

65 (a) *Experimental set-up*

66	We manipulated water temperature, nutrients, and the presence of Stickleback in a 2x2x2 design
67	using 40, open-air, well mixed, 1,136 L Rubbermaid plastic mesocosms (0.6 m deep, 1.5 m in
68	diameter) to test their independent and interactive effects on consumer biomass, producer
69	biomass and CO ₂ flux. Each treatment was replicated 5 times. Water temperatures were either
70	ambient or $3.04^{\circ} \pm 0.05^{\circ}$ C (mean \pm s.e.) above ambient. Mesocosms had either ambient nutrient
71	levels or were meso-eutrophic (N:P ratio of 22) with monthly additions of both nitrogen and
72	phosphorus (264 μ g of nitrogen/L as NaNO ₃ and 27 μ g of phosphorus/L as KH ₂ PO ₄). Finally,
73	mesocosms contained either five Stickleback (54.4 ± 0.05 mm standard body length) or no fish.
74	Detailed experimental methods can be found in Supplementary Material.
75	(b) Organism sampling and CO ₂ measurements
76	Primary producer biomass, consumer biomass (not including fish), and water CO ₂ concentrations
77	were collected one year (May) and 16 months (October) following the start of the study.
78	Phytoplankton and periphyton biomasses and benthic and pelagic consumer biomasses were
79	combined for total primary producer and total consumer biomasses, respectively.
80	Water samples for dissolved CO ₂ were extracted at dawn using 50-mL gas-tight syringes
81	and headspace equilibrium analysis. CO2 concentrations were measured on a 5890 Series II gas
82	chromatograph within 24 h.
83	CO_2 flux (mg C m ⁻² d ⁻¹) to the atmosphere was calculated as follows:
84	$CO_{2 flux} = (CO_{2 water} - CO_{2 sat}) k$
85	Here CO_{2water} is the temperature corrected CO_2 concentration of the water, CO_{2sat} is the
86	concentration the water would have if it were at equilibrium with the atmosphere (390 ppm), k is

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the CO₂ exchange velocity coefficient (0.63 m d⁻¹), which was estimated using literature values for our study sites average wind speed (2.8 ± 0.09 m s⁻¹)[10].

89 (c) Statistical analyses

Treatment effects on primary producer biomass, consumer biomass, and CO₂ flux were tested 90 using linear mixed-effects models (α = 0.05) in R 3.1.1 (R Development Core Team, 2014). 91 Individual mesocosms and date were treated as random factors. Date was included as a random 92 factor to account for non-independence between dates and because our sampling design did not 93 have sufficient power to describe temporal differences. In order to understand how treatment-94 mediated changes to food web structure influenced CO₂ flux, it was imperative to analyse only 95 dates where pelagic and benthic organisms and CO₂ were sampled at the same time. Because of 96 the highly destructive nature of benthic sampling on the community (see supplementary for 97 detailed methods) we limited our collections to only two occasions. 98

99 3. RESULTS & DISCUSSION

Under ambient water temperatures, we detected independent and interactive effects of nutrients 100 and predators on total consumer and producer biomass that led to increased CO₂ influx (Table 1, 101 Fig. 1). Increased CO₂ influx was likely the result of predator- (in predator only) and nutrient-102 103 induced (in nutrient addition treatments only) increases in primary production [9,11]. In the 104 absence of predators, nutrient additions doubled consumer and plant biomass, which would have increased both primary production and respiration (Tables S2). However, because increases in 105 CO₂ influx were still observed in these treatments, the effect of nutrients on CO₂ assimilation 106 appears greater than their effect on respiration. Fish reduced consumer biomass by 70% and 107 increased primary producer biomass by 32% (Table S2). Trophic cascades were enhanced by 108

nutrient enrichment, with a 27-fold increase in producer biomass (Table S2). Similar to Cole et
al. [10], mesocosms with both nutrient additions and top predators had the greatest CO₂ influx.

In general, warming had strong negative effects on trophic cascade strength and the 111 influence of nutrients on primary producers, generating top-heavy food webs with reductions in 112 CO₂ influx (Table 1; Fig. 1). Higher emissions in warmed mesocosms likely resulted from 113 114 positive effects of warming on respiration rates [3] and observed negative effects of warming on primary producers. Contrary to our prediction, warming only strengthened trophic cascades in 115 non-fertilized mesocosms. In fertilized mesocosms, warming dampened consumer effects 116 117 resulting in similar CO₂ fluxes as mesocosms without predators (Fig. 1, Table S2). The negative 118 effect of warming on primary producers was strongest in mesocosms with nutrient additions and especially large in mesocosms with nutrients and predators. Warmed mesocosms containing both 119 120 added nutrients and predators had 90% less primary producer biomass compared to ambient 121 temperature mesocosms (Table S2). The negative effect of warming on primary producer 122 biomass in our study and others [3,12] may have been the result of increased consumption by 123 herbivores under warmer temperatures [6,8], or the replacement of productive phytoplankton taxa by stress tolerant, but less productive taxa. Although further studies are needed to 124 understand the mechanisms by which increased temperatures alter CO₂ cycling, our results 125 suggest that future increases in temperatures could reduce biosequestration by aquatic 126 ecosystems. 127

Mesocosms are well suited to explore complex interactions between global change drivers that are otherwise difficult to study in natural systems. Nevertheless, the use of mesocosms and our context-dependent results should be considered. First, our mesocosms did not receive large terrestrial subsidies, which can make up > 50% of the dissolved organic C pool

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132 in natural ecosystems [13]. Thus, respiration in natural lentic ecosystems is not constrained by *in* 133 *situ* primary production, as was the case in our mesocosms. Because respiration rates have a stronger temperature-dependence than photosynthetic rates [3], the positive effects of warming 134 on CO₂ efflux seen in our study may be conservative. Second, the effects seen on communities 135 and ecosystem process in this study may be only short-term, transient responses to perturbations 136 that could greatly differ from long-term ones. Future studies should investigate whether the 137 effects seen in our study are consistent across diurnal cycles and longer time scales within a 138 myriad of aquatic ecosystems. Finally, the direction and magnitude of predator effects on food 139 webs and carbon cycling may depend on predator identity, food chain length, and species 140 diversity [8]. Despite these limitations our study is the first to our knowledge to unravel the 141 cumulative effects of eutrophication, warming, and alterations to top predators on the CO_2 flux 142 143 of an ecosystem.

Our findings support other studies [9,14] which suggest that maintaining viable predator 144 populations in odd-numbered food webs and/or adding nutrients could increase natural 145 146 biosequestration. However, our results reveal how previously unknown interactions between warming, nutrients, and changes to top predators could create positive climate feedbacks by 147 reducing the capacity of top-down and bottom-up forces to reduce the production of *in situ* CO₂. 148 The number of freshwater ecosystems experiencing negative or undesirable anthropogenic 149 impacts is likely to increase in the future due to a combination of a 2.4-2.7 fold increase in 150 eutrophication [15] and increased rates of trophic downgrading [16]. As most freshwater 151 ecosystems are sources of CO₂, our results suggest that CO₂ emissions from aquatic systems 152 could increase as temperatures rise. 153

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203						
203	Fthic	s. This work was conducted with the approval by the University of British Columbia's				
205	Animal Ethics Committee (animal ethics permit #A08-0220)					
206	06 Data accessibility Raw data are available from the Dryad digital repository					
207	(http://dx.doi.org/10.5061/dryad.1f090)					
208	Author contributions T.B.A., E.H., H.S.G., P.K, J.B.S., J.S.R contributed to study design and					
209	writing of the manuscript. Data were collected by T.B.A., E.H., H.S.G. and P.K. All authors gave					
210	final approval for publication, and agreed to be accountable for all aspects of the work.					
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- 216 **Competing interests statement** We have no competing interests.

Figure Legend 218

- Figure 1 Interaction plots illustrating the impacts of warming, nutrient additions, and predators 219
- on community biomass and CO₂ flux of mesocosms. Effects of nutrients and warming on 220
- consumer biomass (a), primary producer biomass (c), and CO₂ flux (e) in food webs where fish 221
- 222 are absent. Effects of nutrients and warming on consumer biomass (b), primary producer
- 223 biomass (d), and CO₂ flux (f) in food webs containing fish. Dotted lines in graphs (e) and (f)
- represent CO₂ source/sink boundaries. Means \pm 95 % C.I.s represent fixed effects and were 224
- ne()" fu approximated using the "predictSE.lme()" function in the "AICcmodavg" package in R. 225

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227 Tables

228	Table 1 Summary statistics of linear mixed-effects models for individual and interactive
229	effects of warming (W), nutrient additions (N), and predators (P) on consumer biomass,
230	primary producer biomass, and CO ₂ flux of mesocosms. <i>P</i> -values in bold are statistically
231	significant.

	Consumer biomass		Primary producer biomass		CO ₂ flux	
	F _{1,32}	Р	F _{1,32}	Р	F _{1,32}	Р
Ν	55.796	< 0.001	148.673	< 0.001	31.950	<0.001
Р	26.756	< 0.001	28.483	< 0.001	44.814	<0.001
W	3.218	0.081	32.620	< 0.001	77.174	< 0.001
N:P	1.775	0.191	3.815	0.060	0.061	0.807
W:N	0.102	0.751	18.839	< 0.001	6.712	0.014
W:P	0.035	0.852	2.378	0.133	13.024	0.001
W:P:N	0.072	0.790	11.901	0.002	11.744	0.002

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