On the sensitivity of food webs to multiple

stressors

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Data accessibility statement: Empirical food web data for the St. Lawrence System used for this study are openly available from the primary sources in which they were published (Morissette *et al.* 2003; Savenkoff *et al.* 2004; Savenkoff 2012).

39 Proposal letter

Instructions: Unsolicited proposals, which will be evaluated by the Ideas and Perspectives or Reviews and Syntheses Editors, in consultation with the Editorial Board and Editor-inChief, prior to a full submission. Proposals should be no more than 300 words long, describe the nature and novelty of the work, the contribution of the proposed article to the discipline, and the qualifications of the author(s) who will write the manuscript. Proposals should be sent to the Editorial Office (ecolets@cefe.cnrs.fr).

Notes: Contribution portion is unconvincing and letter is 42 words over the limit.

As global change and the human footprint keeps expanding worldwide, properly evaluating the impacts of multiple stressors on natural systems is becoming an increasingly vital 48 issue. Even after decades of research on the impacts of multiple stressors, a general lack of theoretical understanding of their impacts remains. In particular, the role of species and their interactions in mediating the effects of multiple stressors, although widely ac-51 knowledged, has yet to be formally explored. We propose, to our knowledge, the first such 52 exploration by conceptualizing the impacts of multiple stressors on complex food webs and by exhaustively simulating the many pathways through which the dynamics of the most 54 common 3-species motifs in empirical food webs are impacted by stressors using generalized Lotka Volterra models. We uncover that interactions, their configuration and a species 56 trophic position greatly influence a species sensitivity to and amplification of the impacts of stressors. Species in omnivory and tri-trophic food chains in particular are susceptible 58 to the impacts of stressors and to non-additive impacts, which arise chiefly from impacts to species interactions. We then use simulation results to infer species sensitivity to stressors 60 in empirical food webs of the St. Lawrence System in eastern Canada. We find that, in the mid-1980s, species were trophically predisposed to the dramatic shifts in trophic structure 62 that unfolded following the groudfish stock collapse of the early 1990s. This contribution 63 is significant and timely for two main reasons. First, from a theoretical stand point, this is the first exhaustive exploration of the impacts of stressors on food web motifs. Second, we provide a clear link between theory and management by illustrating how our framework can be applied to detect most sensitive species in a real food web exposed to various stressors. 67 Our group of co-authors is singularly positioned to write this paper. D. Beauchesne and P. Archambault both have strong backgrounds in environmental assessments in general and cumulative impacts in particular. K. Cazelles and D. Gravel are both experts in theoretical ecology with strong backgrounds and recognition in the field.

72 Abstract

- Instructions: The abstract page should contain a short summary not exceeding 150 words for Letters, and 200 words for Ideas and Perspectives and Reviews and Syntheses.
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As global change and the human footprint expands worldwide, evaluating the impacts of multiple stressors is becoming increasingly vital. Even after decades of research on the impacts of multiple stressors, a general lack of theoretical and empirical understanding of their impacts remains. In particular, the role of species and their interactions in mediating the impacts of stressors, although widely acknowledged, has yet to be formally explored. Here, we conceptualize the impacts of multiple stressors on complex food webs and exhaustively explore how stressors may impact the most common 3-species motifs in empirical food webs using generalized Lotka-Volterra models. We uncover that interactions, their configuration and a species trophic position greatly influence a species sensitivity to and amplification of the impacts of stressors. Species in omnivory and tri-trophic food chains in particular are susceptible to stressors and to non-additive impacts, which arise chiefly from disruptions to species interactions. We then infer species sensitivities in empirical food webs of the 87 St. Lawrence System. We find that, in the mid-1980s, species were trophically predisposed 88 to the dramatic shifts in trophic structure that unfolded following the groudfish stock collapses of the early 1990s. In conceptualizing and exploring the impacts of multiple stressors on food webs, we provide a clear link between theory and management through a famework 91 that clearly demonstrates that an holistic understanding of complex systems is key to proper management.

$_{94}$ Glossary

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- Stressor: externality that arises from natural or human-induced biophysical processes or from anthropogenic activities and that impacts environmental processes and disturbs natural systems;
- Direct effect: effect of a species on another without the involvement of a third species;
- Indirect effect: effects that link at least two direct effects through an intermediary species;
- Pathway of effect: composed of impacted biological processes (e.g. mortality rates),
 it is the pathway through which stressors directly and indirectly impact ecological
 communities. A unitary pathway of effect is an indvidual impacted biological process,
 whereas an integrative pathway of effect is the set of all unitary pathways of effect that
 combine to collectively impact an ecological community;
 - Trophic sensitivity: difference in species equilibrium abundance before and after the permanent appearance of stressors;
 - Weak entry point: highly susceptible species to a pathway of effect, whether to its benefit and strongly positive trophic sensitivity (i.e. positive weak entry point) or to its detriment and strongly negative trophic sensitivity (i.e. negative weak entry point);
 - Non-additive impact: trophic sensitivity to an integrative pathway of effect that is greater (i.e. synergistic impact) or lower (i.e. antagonistic impact) than the sum of trophic sensitivities to unitary pathways of effect;
- Trophic amplification: difference between a species trophic sensitivity to an integrative pathway of effect and the sum of its trophic sensitivity to unitary pathways of effect;
 - Biotic buffer: species on which an integrative pathway of effect has a lower impact (i.e. antagonism) than the summed impact of unitary pathways of effect;
 - Biotic amplifier: species on which an integrative pathway of effect has a greater

impact (i.e. synergism) than the summed impact of unitary pathways of effect.

Ecosystems worldwide are increasingly affected by a vast array of environmental pressures,

1 Introduction

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commonly referred to as stressors (Boonstra et al. 2015; Halpern et al. 2019). Stressors 123 are driven by natural or human-induced biophysical processes such as ocean acidification 124 and warming, or from anthropogenic activities such as fisheries. Stressors can be defined 125 as externalities that affect environmental processes and disturb natural systems. Individually, stressors impact all levels of biological organization and cause dramatic changes to whole ecosystems. For example, ocean acidification reduces coral and mollusk calcification, 128 metabolic, growth and mortality rates, and has been linked to altered carbon fluxes in ecological networks (Fabry et al. 2008; Kroeker et al. 2013; Bove et al. 2019). Fisheries induced 130 the collapse of commercial species, decreased the mean trophic level in coastal and oceanic 131 environments by targeting large predators and impaired a variety of essential ecosystem 132 services (Pauly et al. 1998; Myers & Worm 2003; Worm et al. 2006). 133 Stressors rarely occur individually and their interactions are extremely alarming because 134 they are dreadfully impactful, pervasive (e.q. Crain et al. 2008; Piggott et al. 2015; Jackson 135 et al. 2016) and unpredictable (Darling & Côté 2008; Côté et al. 2016). Global changes and 136 local human activities result in increasingly intricate environmental stress exposure regimes, 137 compounding the pathways through which natural systems can be altered (Halpern et al. 138 2019). For example, coral reefs contend with a suite of pressures including fishing, warming 139 temperatures, ocean acidification and water pollution (McClanahan et al. 2014; Harborne et al. 2017). Moreover, striking contrasts exist between the impacts of individual and interacting stressors. For instance, Wiedenmann et al. (2013) showed that the susceptibility of corals to temperature-induced bleaching increases with the concentration of dissolved 143 inorganic nitrogen, and a recent study by Lapointe et al. (2019) showed that the synergy

between nutrient enrichment and global warming is already causing major coral reef loss in the Florida Keys. Other dramatic examples include up to 100-fold increases in the sensitivity of certain organisms to toxicants when exposed to other environmental stressors (Liess et al. 2016) and the reversal of the positive effects of acidification on primary producer biomass 148 by warmer waters in boreal lakes (Christensen et al. 2006). 149 Beyond their obvious direct impacts, stressors ripple through ecological communities by 150 way of the interactions structuring the complex network in which species are embedded 151 (Wootton 2002; Bascompte 2009; Montoya et al. 2009; O'Gorman & Emmerson 2009; 152 O'Gorman et al. 2012). Surprising observations arise from complex networks, such as a 153 predator positively affecting its own prey (Abrams 1992). Ample empirical evidence exist of 154

such trophically-mediated effects across ecosystems globally (Paine 1980; Estes et al. 2011).

Classic examples include sea otters (Enhydra lutris) indirectly shielding kelp forests from

browsing by sea urchins (Strongylocentrotus sp.; Estes & Palmisano 1974) and the release

of cottonwood (Populus spp.) and willows (Salix spp.) from elk (Cervus elaphus) browsing

following the reintroduction of wolves (Canis lupus) in Yellowstone National Park (Ripple

& Beschta 2003). A species's susceptibility to trophically-mediated effects is influenced by

are more resilient than species with specialized diets (*i.e.* specialists; Montoya *et al.* 2009; Clavel *et al.* 2011), and apex predators are generally more vulnerable to trophically-mediated

its trophic role and position. For example, species with diversified diets (i.e. generalists)

effects (Ripple et al. 2015; Stier et al. 2016).

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How ecological networks are structured, *i.e.* the number, configuration and strength of interactions between species, also influences the propagation of stressors and the stability and resilience of whole systems (Wootton 2002; Montoya *et al.* 2009; Bartley *et al.* 2019). Stressors can modify these structural properties and rewire entire food webs (Blanchard 2015; Kortsch *et al.* 2015; Bartley *et al.* 2019). Links can be added or removed (*i.e.* topological rewiring; Bartley *et al.* 2019) through primary and secondary species extinctions (*e.g.* Allesina *et al.* 2006; Eklöf & Ebenman 2006), climate-related distributional shifts (*e.g.*

Kortsch et al. 2015; Bartley et al. 2019) or invasive species introductions (e.g. Vander Zanden et al. 1999; David et al. 2017). Alteration to the flow of energy also arise when consumer modify their space and resource use (i.e. interaction strength rewiring; Bartley et al. 2019).

In spite of all this knowledge, single-stressor and single-species assessments remain over-176 whelmingly employed (O'Brien et al. 2019), and most large-scale multiple stressors studies 177 remain focused on impacts to habitats (e.g. Ban et al. 2010; Halpern et al. 2019). This 178 conveniently circumscribes assessments to sets of tractable units, but it ignores systems com-179 plexity and aggregates biological processes across levels of organization (Halpern & Fujita 180 2013; Giakoumi et al. 2015). Furthermore, methodologies tend to assume that impacts of 181 multiple stressors are additive (e.g. Halpern et al. 2019), or rely null models providing little 182 insights into the ecological mechanisms governing impacts of multiple stressors (Griffen et 183 al. 2016; Jackson et al. 2016; De Laender 2018; Schäfer & Piggott 2018). 184

Confronted with the challenge of managing and preserving complex systems, we clearly need 185 holistic frameworks that consider the complexities of multiple stressors in ways that are 186 informative to management. Such frameworks should be grounded in theory, scaled for gains 187 in ecological understanding rather than convenience, and allow for dynamic identification of 188 conservation targets. It should also be flexible and data-oriented so that it can make use 189 of the expanding corpus of openly available environmental knowledge and computational 190 capabilities. This need to unite practice and theory is reflected in the recent upsurge in 191 theory-driven modelling approaches to evaluate the impacts of multiple stressors on complex 192 systems (e.q. Hodgson et al. 2017; De Laender 2018; Galic et al. 2018; Thompson et al. 193 2018). 194

Here, our objective is to theoretically and holistically investigate the role of species and their interactions in driving species sensitivity to the impacts of multiple stressors. In doing so, we seek to answer questions of particular significance to management and the application

of holistic environmental approaches: 1) should species interactions be considered in impact assessments, 2) should the effects of stressors be evaluated separately or in combination, 199 and 3) which species are most sensitive to stressors based on their trophic position? The 200 paper is divided in two parts. In the first part, we conceptualize how multiple stressors 201 permeate complex ecological communities. We then simulate the impacts of stressors on 202 the equilibrium dynamics of the most frequent three-species motifs in food webs (i.e. food 203 chain, omnivory, exploitative competition, and apparent competition) to explore the many 204 pathways through which species can be impacted in complex communities. In the second 205 part, we illustrate our framework by inferring the sensitivity of species in the St. Lawrence 206 System, in Eastern Canada, using data from three empirical food webs describing different 207 regions of the St. Lawrence and exposed to up to eight different sources of stress. 208

209 2 Of food webs and multiple stressors

2.1 A gateway through ecological communities

We begin by conceptualizing how multiple stressors impact species embedded in complex 211 ecological communities. To illustrate this, we use a simplified 6-species food web composed 212 of populations of beluga (Delphinapterus leucas) and humpback (Megaptera novaeangliae) 213 whales, Atlantic cod (Gadus morhua), capelin (Mallotus villosus), krill (Euphausiacea) and 214 copepods (Copepoda; Figure 1A). The dynamic ecological communities is driven by biological 215 processes operating at the scale of individual populations (e.g. reproduction and mortality) 216 and of the whole community (e.q. consumer-resource interactions). Through interactions, species influence the dynamics of other species both directly and indirectly. Direct effects arise when a species affects another without the involvement of a third species (Abrams et al. 219 1996; Wootton 2002). For example, cod consumes capelin in our system, directly affecting 220 its prey and weaving the dynamics of both populations together (Figure 1A). 221

Indirect effects are effects that link at least two direct effects through an intermediary species, which results in an interaction chain also known as a density-mediated indirect effect (Wootton 1993, 2002; Abrams et al. 1996). In our system, krill is indirectly affected by cod through their respective interaction with capelin (Figure 1A). Indirect effects can be as important of, 225 and propagate faster than, direct effects (Wootton 1993, 2002; Menge 1995; Yodzis 2000). 226 Trophic cascades, i.e. the propagation of impacts by consumers downward through whole 227 food webs (e.g. cod-capelin-krill; Figure 1A) and apparent competition, i.e. alternate prey 228 species of a generalist predator (e.g. krill-capelin-copepod; Figure 1A) are well-documented 229 and common types of indirect effects in empirical food webs (e.g. Holt 1977; Paine 1980; 230 Menge 1995; Estes et al. 2011). In complex food webs, the net effect of a single or of multiple 231 species on another is the integration of all direct and indirect effects propagating through 232 individual pathways to a focal species (Abrams et al. 1996). For example, the net effect of 233 cod on beluga depends on the direct effect linking both species and the indirect effect of cod 234 on beluga through capelin (Figure 1A). 235 Now imagine that 3 distinct sources of stress appear in the system: climate change-induced temperature anomalies, commercial shipping and trawl fishing (Figure 1A). Stressors impact 237 populations and whole communities by disrupting the biological processes that govern their 238 dynamics (e.g. Galic et al. 2018; Guiden et al. 2019; Hodgson & Halpern 2019; Hodgson 239 et al. 2019). Stressors can disrupt multiple processes and multiple process can be disrupted 240 by a single stressor. Species-specific sensitivities influence the resulting impacts of stressors. 241 For example, species may be well adapted to (e.g. northern shrimp Pandalus borealis and 242 Greenland halibut Reinhardtius hippoglossoides; Pillet et al. 2016), avoid (e.g. Atlantic 243 cod; Chabot & Claireaux 2008), or be killed by (e.g. sessile benthic invertebrates; Eby et 244 al. 2005; Belley et al. 2010) hypoxic environments. To address our objective, however, we 245 focus on the net impact of stressors, rather than how impacts manifest in the first place. 246 Identifying and quantifying species-specific sensitivities is best addressed through in situ 247

sampling and targeted experimental investigations. These have limited applicability for

complex communities influenced by many stressors, and are thus beyond the scope of this manuscript.

Let us assume that temperature anomalies impact the reproductive capabilities of copepods 251 (i.e. growth rate) and the effectiveness of their predators to assimilate them (i.e. conversion 252 rate), that shipping alters the feeding behaviour of whales (i.e. attack rate), and that fish-253 eries impact the mortality of cod and capelin (Figure 1A, B). Stressors permeate ecological 254 communities through the indirect pathways of effects provided by species interactions (Woot-255 ton 2002; Bascompte 2009; Montoya et al. 2009; O'Gorman & Emmerson 2009; O'Gorman 256 et al. 2012). For example, the direct impact of shipping on whale behaviour indirectly prop-257 agates to their prey (Figure 1A). By extension to net effects in food webs, the net impact of a 258 single or of multiple stressors on a species must integrate all direct and trophically-mediated 259 impacts propagating through individual pathways to a focal species. As such, the net impact of shipping on whale also depends on the indirect impact propagating to capelin, cod and krill, and cycling back to whales (Figure 1A).

2.2 Handling food web complexity

The number and complexity of pathways through which a species may affect or be affected by other species – and through which impacts may permeate communities – increases ex-265 ponentially with the number of species and interactions in a network (Menge 1995). To 266 illustrate this, let us imagine that community dynamic is governed by resource growth, mor-267 tality, attack and conversion rates (Figure 1B). Our six-species system would then be driven 268 by 21 distinct biological processes, offering over 2 000 000 unique pathways through which 269 the system could be disrupted. Such complexity has proven prohibitive to investigations 270 into community dynamics and the impacts of disturbances on food webs (Wootton 2002; 271 Montoya et al. 2009). 272

273 Studying smaller subgraphs – community motifs or modules – emerged as an alternative

to gather insights into the dynamics and stability of complex ecological communities (Holt 1997; Holt & Hochberg 2001). Motifs are collections of n-species that, when put together, construct whole food webs (Milo et al. 2002; Stouffer et al. 2007). They form the backbone 276 of food webs and provide a mesoscale characterization of the structural properties of commu-277 nities (Bascompte & Melián 2005; Stouffer et al. 2007; Stouffer & Bascompte 2010, 2011). 278 Investigations into three-species motifs are particularly common in the literature (e.q. Menge 279 1995; Milo et al. 2002; Stouffer et al. 2007, 2012). On average, 95% of three-species motifs 280 in empirical food webs are composed of tri-trophic food chain (e.g. cod-capelin-krill; Figure 281 1A), omnivory or intraguild predation (e.g. beluga-cod-capelin; Figure 1A), exploitative 282 competition (e.q. humpback whale-capelin-beluga; Figure 1A) and apparent competition 283 (e.g. krill-capelin-copepod; Figure 1A) motifs (Camacho et al. 2007; Stouffer & Bascompte 284 2010). Focusing on motifs rather than whole food webs restricts the complexity we must 285 contend with to better understand the role of species and their interactions in mediating the 286 impacts of multiple stressors. For example, impacts to omnivory interactions are possible 287 through 9 biological processes and 511 unique pathways (Figure 1B). We now shift our focus 288 to the dynamics of those four motifs particularly relevant to the structural properties of 289 empirical food webs.

2.3 Simplified dynamics of multiple stressors

We explore the dynamic impacts of multiple stressors with the omnivory interaction connecting cod, beluga and capelin in our system (Figure 1C). Net effects are typically measured as
variations in species abundances or densities in food webs, which integrate all trophicallymediated effects operating on the system collectively (Wootton 2002; Montoya et al. 2009).
Likewise, we evaluate how pre-stressor abundances at equilibrium shift after the permanent
appearance of stressors in a system as a measure of net impact.

Impacts to single biological processes travel through unitary pathways of effect (k), such

as an increase in cod mortality ($k = \{m_y\}$; Figure 1B). Unitary pathways of effect that can induce contrasting population trajectories. The impact of fishing on capelin mortality 300 $(k = \{r_x\})$ favours cod and results in reduced abundances for capelin and beluga (Figure 301 1C-1). In this scenario, cod are likely released from beluga predation due to their drop in 302 numbers (i.e. mesopredator release; Ritchie & Johnson 2009). This trophically-mediated 303 effect could ultimately exacerbate the impact of fishing on capelin by favouring one of its 304 predators. Meanwhile, impacting cod mortality $(k = \{m_y\})$ results in the growth of the 305 capelin and beluga populations (Figure 1C-2). Surprisingly, the cod population remains 306 relatively unchanged (Figure 1C-2), likely because the increase in prey availability offsets 307 the impact of fishing (i.e. compensatory dynamics; Gonzalez & Loreau 2009). Finally, the 308 beluga population appears insensitive to the impact of shipping $(k = \{\alpha_{xz}\})$ and $k = \{\alpha_{yz}\}$; 309 yet shipping likely disrupts the top-down control of beluga on cod to the benefit of cod and 310 to the detriment of capelin (Figure 1C-3). 311 Unless a single biological process is impacted, unitary pathways of effect combine to form 312 an integrative pathway of effect (K). Shipping and fishing collectively impact our system 313 through an integrative pathway of effect $(K = \{r_x, m_y, \alpha_{xz}, \alpha_{yz}\})$ that benefits cod and 314

$$S_{m,K} = \frac{a_{m,K} - a_m}{a_m},\tag{1}$$

where a_m and $a_{m,K}$ are the pre- and post-stressors abundances of species m, respectively. In the remainder of the text, the term pathway of effect without a qualifier (*i.e.* integrative or unitary) refers to integrative pathways of effect. Note that by definition $S_{m,K}$ is bounded negatively to -1, as species abundances cannot be negative. We refer to species that are highly susceptible to impacts – whether positively or negatively – as weak entry points and

reduces capelin and beluga (Figure 1B4). We define a species (m) trophic sensitivity $(S_{m,K})$

as the net impact -i.e. the pre- and post-stressors variation in abundance - resulting from

an integrative pathway of effect K (Figure 1B):

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distinguish between negative and positive weak entry points.

In multi-species systems, where many direct and indirect trophic effects are operating si-324 multaneously, impacts of stressors can be amplified or dampened through biotic interactions 325 (Ives 1995; Wootton 2002; Thompson et al. 2018). Uncovering synergies and antagonisms 326 has been a hallmark of investigations into the impacts of multiple stressors (e.g. Crain et 327 al. 2008; Darling & Côté 2008; Côté et al. 2016; Galic et al. 2018; Thompson et al. 2018). 328 These so-called non-additive impacts arise when the net impact of an integrative pathway 329 of effect is greater (i.e. synergy) or lower (i.e. antagonism) than the sum of the impacts 330 of unitary pathways of effect. We define a species (m) trophic amplification $(A_{m,K})$ as the 331 difference between its trophic sensitivity to an integrative pathway of effect (K) and the sum 332 of its trophic sensitivities to the unitary pathways of effect forming K ($k \in K$; Figure 1C-4):

$$A_{m,K} = \sum_{k \in K} S_{m,k} - \frac{1}{|K|} S_{m,K}, \tag{2}$$

where |K| is the number of unitary pathways of effect k forming the integrative pathway of effect K. Synergisms and antagonisms are identified by positive and negative trophic 335 amplifications, respectively. From this definition of non-additive impacts, a single stressor 336 can elicit non-additive impacts by disrupting multiple biological processes. In constrast, non-337 additive impacts are usually defined as arising from more than one stressor. However, we 338 argue that, at the scale of communities, a stressor could indeed elicit non-additive impacts on itw own. In our system, shipping and fishing elicit synergistic impacts on capelin and beluga, and a slightly antagonistic impact on cod. Similarly, we refer to species as biotic amplifiers or biotic buffers if their trophic amplification is positive or negative, respectively. 342 Hence, capelin and beluga are biotic amplifiers, whereas cod is a biotic buffer (Figure 1C-4). 343 A species' trophic sensitivity – or lack thereof – can also arise from different mechanisms. 344 Unitary pathways of effect may reinforce each other, whereas others may cancel each other 345 out (Wootton 2002; Montoya et al. 2009). For example, the positive impact of cod mortality 346

on capelin (Figure 1C-2) is offset by the negative impacts of capelin mortality and altered beluga behaviour (Figure 1C-1, C-3, C-4). Comparing the effective and expected impacts of a unitary pathway of effect – i.e. the average impact of an integerative pathways of effect – provides a measure of variance associated to trophic sensitivity to an integrative pathway of effect (K):

$$V_{m,K} = \sum_{k \in K} \left(S_{m,k} - \frac{1}{|K|} S_{m,K} \right)^2.$$
 (3)

Low variance arise from sets of unitary pathways of effect whose individual impacts are relatively similar, whereas high variance identify sets of contrasting unitary pathways of effect. In our system, beluga ($V_{beluga,K} = 0.22$) and capelin ($V_{capelin,K} = 0.18$) are exposed to unitary pathways of effect that tend cancel each other out, whereas cod ($V_{cod,K} = 0.09$) is exposed to unitary pathways of effect that reinforce each other.

$_{57}$ 2.4 Holistic impacts of stressors

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volved in an omnivory interaction and exposed to a specific pathway of effect (Figure 1C-4). 359 There are far more potential pathways of effect. Restricting impacts of stressors to resource 360 growth, mortality, conversion and attack rates, there are 7 biological processes and 127 dis-361 tinct pathways of effect for the tri-trophic food chain, competitive exploitation and apparent 362 competition motifs, and 9 biological processes and 511 pathways of effect for the omnivory 363 motif. We now model the dynamics of those motifs using generalized Lotka-Volterra equation systems with Type 1 functional response (Table S1) to explore all possible pathways of effect. 366 Two additional motifs are included as controls to test the importance of species interac-367 tions in mediating the impacts of stressors: a partially connected motif with a disconnected 368

We have thus far explored the trophic sensitivity, amplification and variance of species in-

species and a predator-prey interaction resulting in 31 distinct pathways of effects, and a disconnected motif with three fully independent species resulting in 7 pathways of effect.

Resources were modeled using equations of the form:

$$\frac{dX_i}{dt} = X_i(r_i - \alpha_{ii}X_i - \sum_j \alpha_{ij}X_j),$$

where X_i denotes species i, r_i is the intrinsic resource growth rate, α_{ii} is the densitydependent effect of the resource on itself and α_{ij} is the rate at which consumer j affects
resource i, i.e. the attack rate.

³⁷⁵ Consumers were modeled using equations of the form:

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$$\frac{dX_j}{dt} = X_j(-m_j + \sum_i e_{ij}\alpha_{ij}X_i - \alpha_{jk}X_k),$$

where m_j is the mortality rate of species j and e_{ij} is the rate at which resource i biomass is

transformed into consumer j biomass, i.e. the conversion rate, and is a scaling parameter of

the attack rate which cannot exceed 1. 378 We studied the equilibrium dynamics of species constrained to coexistence, i.e. the abun-379 dances of all species had to be positive before and after the appearance of stressors in the 380 system. See Stouffer & Bascompte (2010) and Stouffer et al. (2012) for investigations into 381 the role of motifs and species to community persistence and stability. As no equilibrium ex-382 ists for the exploitative competition motif, competitive parameters between consumers were 383 included to the consumer models to constrain growth and limit competitive exclusion. Refer 384 to table S1 for the detailed equation system of each motif considered. Equation systems 385 were solved using SageMath (Developers 2019). All other analyses were performed using R (Team 2019).

Initial parameter values for intrinsic growth (r) and resource density-dependence (α_{ii}) were

fixed to 1 and 0.001, respectively, to bound all resource solutions. Competitive parameters

for the exploitative competition motif were also fixed to 0.001, and conversion rates (e) were fixed to 0.5. A set of 100 initial values for mortality $(m \in [0.01, 0.5])$ and attack rates 391 $(\alpha_{ij} \in [0.0001, 0.01])$ was sampled using a simulated annealing algorithm. The algorithm 392 optimized parameter solutions for consumer abundance, i.e. the algorithm was more likely 393 to select solutions if the abundance of both consumers in a particular motif increased at 394 each iteration. Complete sets of parameters were only retained if initial abundances and 395 abundances resulting from all possible pathways of effect were positive (explained below). 396 All possible pathways of effect were simulated by modifying the equilibria equation parameter 397 values by 1%. Parameters were modified to simulate negative impacts only, e.g. by increasing 398 the mortality rate of the predator in the food chain motif by 1%. Simulated impacts were limited to resource growth rates (r) mortality rates (m), conversion rates (e) and attack rates (α_{ij}) . Trophic sensitivity $(S_{i,K})$ and trophic amplification (A_i) for all unique motif positions (i) were then computed (see equations 1 and 2 above). The overall trophic sensitivity 402 and amplification of unique motif positions was computed as the average sensitivity and 403 amplification over all possible pathways of effect. We refer to those quantities as a position 404 sensitivity and a position amplification. 405 Arbitrary thresholds were used to identify negative $(S_{i,K} < 1)$ and positive $(S_{i,K} > 1)$ 406 weak entry points, biotic buffers $(A_{i,K} < 0.02)$ and biotic amplifiers $(A_{i,K} > 0.02)$. These 407 thresholds are used solely as references for discussion purposes to identify species that are 408 more or less sensitive and prone to non-additive impacts. 400

⁴¹⁰ 2.5 Impacts on motifs

Unsurprisingly, greater trophic sensitivities arise from integrative than unitary pathways of effect for types interation types (Figure 2). Interactions (food chain, omnivory, exploitative and apparent competition) also lead to greater trophic sensitivities than controls (partially connected and disconnected; Figures 2, 3). Among interaction types, omnivory and food chains are generally more sensitive and prone to amplification than competitive interactions (Figures 2, 3).

Weak entry points are prevalent in all interactions (Figures 2, 3). Most notably, predators 417 and resources in omnivory and food chains are predominantly negative weak entry points, 418 whereas mesopredators are positive weak entry points (Figures 2, 3). Likewise, consumers 419 in exploitative competition and resources in apparent competition are generally negative 420 and positive weak entry points, respectively (Figures 2, 3). Resources and consumers in 421 exploitative and apparent competition are the sole insentive positions among interactions 422 (Figures 2, 3). Interestingly, this insensitivity arises from negligible impacts rather than 423 unitary pathways of effect cancelling each other out (see variance in Figure 2). 424

Non-additive impacts are largely exclusive to species in omnivory interactions and to the predator in food chains; integrative pathways of effect consistently lead to synergistic or antagonistic impacts and unitary pathways of effect are highly contrasted (Figure 2). Meticulous knowledge on realized pathways of effect is therefore necessary to properly typecast these species as either biotic buffers or amplifiers, even though predators and consumers are, on average, biotic amplifiers (Figure 3). We can, however, typecast these positions are acutely susceptible to non-additive impacts.

Unsurprisingly, impacts to mutliple types of biological processes lead to greater trophic sensitivities and amplifications (Figure 4). Impacts to species mortality or growth rates in particular influence trophic sensitivity (Figure 4). Yet it is worth noting that impacts to mortality or growth rates do not affect trophic amplification; instead, non-additive impacts are driven by impacts on species interactions, *i.e.* attack (α_{ij}) and conversion (e) rates (Figure 4).

3 Scaling back to complex food webs

Thus far, we simplified food web complexity using motifs, conceptualized the role of species and their interactions in mediating the impacts of multiple stressors in ecological communities, and evaluated how interaction type influences a species trophic sensitivity and amplification. We now scale back to species in complex food webs. As a food web can be deconstructed into n-species motifs, it can be pieced back together to 443 assess the structural roles of species and their interactions in complex food webs (Stouffer et 444 al. 2012; Cirtwill & Stouffer 2015). All the positions a species holds in n-species interactions 445 in a food web forms its motif profile. In our system, cod is twice a predator in food chains, 446 once a consumer in an omnivory interaction and once a consumer in exploitative competition 447 (Figure 1D). Each three-species interaction is impacted through a specific pathway of effect 448 from which we can evaluate trophic sensitivity and amplification (Figure 1D, E). For example, 449 cod is negatively impacted through the food chains and competitive exploitation interactions, 450 whereas it benefits from impacts through the omnivory interaction (Figure 1E). Impacts to 451 cod are also amplified through the food chain with capelin and copepod, yet buffered through 452 the omnivory and exploitative competition interactions (Figure 1E). We summarize trophic sensitivities (S_m) and amplifications (A_m) across a species motif profile (M) by summing 454 individual trophic sensitivities and amplifications (Figure 1E):

$$S_m = \sum_{i \in M} S_{i,K^i}; \tag{4}$$

$$A_m = \sum_{i \in M} A_{i,K^i},\tag{5}$$

where K^i is the pathway of effect impacting species through motif i. Summarizing by adding individual trophic sensitivities and amplifications allows for individual pathways of effect to reinforce and cancel each other out. For instance, we expect pathways of effect to negatively and synergistically impact cod in our system (Figure 1E).

$_{ t 460}$ 3.1 An empirical illustration: the St. Lawrence System

We illustrate a species trophic sensitivity and amplification in complex food webs using 461 empirical food webs from the St. Lawrence System in eastern Canada. The St. Lawrence 462 System is formed by one of the largest estuaries in the world and a vast interior sea. Variable 463 environmental and oceanographic processes make it suitable for the establishment of diverse 464 and productive ecological communities (El-Sabh & Silverberg 1990; Savenkoff et al. 2000). 465 The St. Lawrence System provides a wealth of ecosystem services that benefit the Canadian 466 economy. It sustains rich commercial fisheries, grants access to one of the most densely 467 populated regions in North-America through more than 40 ports, is home to an expanding 468 aquaculture production, and has an expanding tourism industry (Beauchesne et al. 2016; 469 Archambault et al. 2017; Schloss et al. 2017). These human-induced stressors blend with 470 climate related stressors to result in intricate stress exposure regimes (Beauchesne 2019). 471

472 **3.1.1** Food webs

We use empirical data on food webs in the three main regions of the St. Lawrence for 473 different periods: the Northern Gulf of St. Lawrence (mid-1980s; Morissette et al. 2003), 474 the Southern Gulf of St. Lawrence (mid-1980s; Savenkoff et al. 2004), and the St. Lawrence 475 Estuary (early 2000s; Savenkoff 2012). The total number of species and functional groups 476 differs between food webs ($n_{SSL} = 30$; $n_{NSL} = 32$; $n_{ESL} = 41$;), yet there is significant overlap ($n_{all} = 21$). Food web resolution is biased towards commercial fish for all food webs. Interactions were identified as a species or functional group's diet composition in percent. 479 Only diet percent > 0.1 were considered as interactions. Refer to Morissette et al. (2003), Savenkoff et al. (2004) and Savenkoff (2012) for more details on the food webs. Note that 481 detailed results are only presented and discussed for the Northern Gulf of St. Lawrence food web in the manuscript. Results for the Estuary and Southern Gulf are nonetheless available as supplementary material (Figures S1, S2).

485 3.1.2 Stressor impacts

- The most prevalent sources of offshore human- and climate-induced stress in the St. Lawrence
 System are shipping, trawl, trap and pelagic fisheries, ocean acidification, hypoxia, and
 bottom- and surface-water temperature anomalies (Beauchesne 2019). We inferred the impacts of individual sources of stress on the biological processes governing these food webs
 (Figure 5) through broad trait-matching rules guiding from expert knowledge and the scientific litterature to simplify the number of assumptions required.
- 1) Position in the water column *i.e.* deep or surface-dwelling species determined a species exposure to stressors. Acidification, hypoxia and bottom temperature anomalies are widespread in the deep layers of the St. Lawrence, whereas surface temperature anomalies and shipping are prevalent in the surface layer (Beauchesne 2019);
- Mobility established a species vulnerability to hypoxia and temperature anomalies.

 Hypoxia and temperature anomalies were considered as impactful to the physiology of
 species with low mobility, whereas the behaviour of mobile species was considered as
 impacted by hypoxia only;
- 3) Ocean acidification was considered impactful to the physiology of carbonate-secreting organisms (e.g. mollusks and crustaceans; Kroeker et al. 2013);
- 502 4) Shipping was considered as impactful to whale behaviour (Christiansen *et al.* 2013;
 503 Lesage *et al.* 2017);
- 5) Fisheries were considered impactful to a species mortality. The catch data provided in
 the food web descriptions provided a list of targeted species (Morissette *et al.* 2003;
 Savenkoff *et al.* 2004; Savenkoff 2012). The gear types used to capture targeted species
 were identified with landing data from the Department of Fisheries and Ocean's Canada

(DFO 2016). Reported whale bycatch and seals and seabird hunting were considered as impacts to mortality (Morissette et al. 2003; Savenkoff et al. 2004; Savenkoff 2012).

3.1.3 Pathways of effect

- The motif profile of each species was evaluated from the structure of each food web (Figure 5).

 For each three-species interaction in which a species was identified, the realized pathway of

 effect was identified as a function of biological processes impacted (Figure 5). The following

 rules were applied to identify realized pathways of effect:
- 1) impacts on mortality disrupt resource growth rates (r) and consumer mortality rates (m),
- 2) impacts on behaviour disrupt consumer attack rates (α_{ij}) , and
- $_{518}$ 3) impacts on physiology disrupt consumer conversion rates (e).
- Simulation results from the holistic exploration of the impacts of stressors on motifs were then used as heuristics to infer the trophic sensitivity and amplication of species to specific pathways of effect in the food webs.

3.2 Sensitivity and amplification in the St. Lawrence System

A species trophic position – as described by its motif profile – informs us on expected trophic sensitivities and amplifications. The most trophically sensitive species in the St. Lawrence are most frequently positioned as predators in food chain, omnivory and exploitative competition interactions (Figure 5); these generally lead to negative impacts (Figure 3). The least sensitive species, meanwhile, generally occupy positions that benefit from (e.g. consumersin omnivory interactions) or are less sensitive to (e.g. resource in exploitative competition) to stressors (Figures 3, 5). Interestingly, trophic sensitivities and amplifications are not correlated; in fact, there are very few biotic buffers and most species are biotic amplifiers to

some degree (Figure 5). Furthermore, the number of stressors impacting a species does not necessarily translate to higher trophic sensitivities or amplifications. For example, shrimp trophic sensitivity was very low even though its mortality, physiology and behaviour were all potentially impacted; marine mammal and sea birds, in contrast, were negative weak entry points and biotic amplifiers even without stressors (Figure 5).

We can largely summarize results for the mid-1980s Northern St. Lawrence food web with 536 two major observations. Large apex predators (e.g. Atlantic cod, Greenland halibut and 537 large demersals) and marine mammals were both negative weak entry points and biotic 538 amplifiers (Figure 5). Smaller forage species, meanwhile, were either positive weak entry 539 points and biotic amplifiers (e.g. capelin and crustaceans), or largely insensitive to stressors 540 (e.q. shrimp; Figure 5). On the whole, these observations foreshadow the ensuing groundfish 541 stock collapse of the early to mid-1990s in the St. Lawrence (Savenkoff et al. 2007; Morissette et al. 2009) and elsewhere in the Northern Atlantic Ocean (Worm & Myers 2003; Frank et al. 2005). Triggered by overfishing and poor environmental conditions (Dempsey et al. 2018), the groundfish stock collapse resulted in dramatic shifts in trophic structure that saw the fall of piscivorous groundfish and the rise of small pelagics and benthic crustaceans (Savenkoff et al. 2007; Morissette et al. 2009) that mostly endure 30 years later (Bourdages et al. 547 2018). Marine mammals, meanwhile, shifted their resource use and their biomass increased 548 in the St. Lawrence (Morissette et al. 2009; Gavrilchuk et al. 2014). Trophic interpretations 549 explaining the shifts are plentiful (e.g. Jackson et al. 2001; Worm & Myers 2003; Frank et al. 550 2005); it now appears that species were also trophically predisposed more or less susceptible 551 to stressors in the St. Lawrence. 552

Still, trophic sensitivities and amplifications are context-specific rather than species-specific.

Indeed, species like cod, shrimp and large crustaceans have variable trophic sensitivities and
amplifications in the food webs analysed: cod was more susceptible to impacts in the North
than in other regions of the St. Lawrence, shrimp benefited more from impacts in the Estuary,
and large crustaceans benefited in the Gulf, yet were negatively impacted in the Estuary

(Figure 6). Similarly, stressor type alters a species trophic sensitivity and amplification. For instance, fisheries and climate combine to increase and decrease sensitivity of cod and shrimp, respectively (Figure 6). Stressors can also strengthen or weaken their respective impacts. For instance, fisheries weaken the impact of climate stressors on shrimp, although it greatly increases trophic amplification (Figure 6). All these observations bolster our assessment that detailed knowledge of realized pathways of effect and of the structure of food webs is necessary to properly capture the impacts of multiple stressors.

565 4 What it all means

From cells to ecosystems, stressors are increasingly impacting natural environments. Here, we show how stressors impacting individual species ripple through entire ecological communities. 567 In conceptualizing the impacts of multiple stressors on food webs, we sought to develop a 568 theory-grounded framework to tackle the challenge of incorporating the complexities of real-569 world systems to impact assessments, management and conservation. We hypothesized that 570 interactions, their configuration and species trophic positions greatly influence a species 571 sensitivity to and amplification of the impacts of stressors. We numerically explored the 572 many pathways of effect through which the dynamics of the four most common 3-species 573 motifs (i.e. tri-trophic food chain, omnivory, competitive and apparent competition) can 574 be impacted using generalized Lotka-Volterra models. We then used simulation results as 575 heuristics to infer species sensitivity to and amplification of multiple stressors in the food 576 webs of the St. Lawrence System in eastern Canada. In doing so, we searched for theory-577 grounded answers to questions of particular relevance to management. 578

579 Should species interactions be considered in impact assessments?

We find, as anticipated, that species interactions play a crucial role in mediating the impacts
of multiple stressors through food webs and that considering species in isolation can grossly

underestimate the impacts of stressors. Our result support longstanding evidence on the importance of interactions in spreading and amplifying the effects of disturbances through food webs (e.g. Menge 1995; ???; Wootton 1993, 2002; Yodzis 2000; Montoya et al. 2009; O'Gorman & Emmerson 2009) and extend this conclusion to the impacts of multiple stressors (see Thompson et al. 2018).

In the context of multiple stressors, our results further establish that impacts to multiple
types of biological processes – e.g. mortality and conversion rates – greatly increase a
species's sensitivity to stressors and that non-additive impacts emerge chiefly from impacts
to species interactions (i.e. attack and conversion rates), rather than impacts to their growth
or mortality rates. These results suggest that using a single indicator of population health
such as reproduction or mortality downplays the impacts of stressors; they also imply that, for
complex food webs, cataloguing and quantifying species interactions is necessary to properly
evaluate the often suprising impacts of multiple stressors.

We further uncover that the type of interaction greatly influences the susceptibility of species 595 to the impacts of multiple stressors. Omnivory and food chain interactions in particular lead 596 to heightened trophic sensitivities for all species involved; they also capture almost all non-597 additive impacts arising through species interactions in our simulations. Food chains are 598 generally overrepresented (i.e. observed more often than randomly expected) in empirical 590 food webs, whereas omnivory interactions are often used to differentiate food webs based 600 on their structural properties (Bascompte & Melián 2005; Stouffer et al. 2007; Monteiro & 601 Faria 2016; Klaise & Johnson 2017). While the role of omnivory in stabilizing food webs is 602 debated in the literature (e.g. McCann et al. 1998; Bascompte & Melián 2005; Stouffer et 603 al. 2007; Monteiro & Faria 2016; Klaise & Johnson 2017), our result suggest that food webs 604 overrepresented with omnivory motifs may be more susceptible to multiple stressors. 605

606 Should impacts of stressors be evaluated jointly?

We find that trophic amplification almost exclusively affects species in food chain and om-607 nivory interactions. For these, non-additive impacts are so prevalent that evaluating impacts 608 on a stressor by stressor basis will almost always result in over- or under-estimations of net 609 impacts of stressors. Moreover, the type of non-additive impacts is so variable that dis-610 tinguishing between synergistic and antagonistic impacts is only possible through detailed 611 knowledge on the pathways of effect through which stressors are impacting a system. Consid-612 ering that food chain and omnivory interactions are important building blocks of empirical 613 food web structure (e.g. Bascompte & Melián 2005; Stouffer et al. 2007; Monteiro & Faria 614 2016; Klaise & Johnson 2017), non-additive impacts are likely common. Based on this, we 615 conclude that stressors should be evaluated jointly. 616

Which species are most susceptible to stressors?

- At the motif and food web scales, predators are generally most susceptible to stressors, acting
 as negative weak entry points and either biotic amplifiers or buffers as a function of specific
 pathways of effect. This is expected, as apex predators are both preferentially targeted for
 hunting and fishing, and more vulnerable to trophically-mediated effects (Pauly et al. 1998;
 Estes et al. 2011; Ripple et al. 2015; Stier et al. 2016).
- Our results further show that direct impacts are unnecessary for a species to be highly susceptible to stressors in a food web; conversely, a species can be largely insensitive or favoured even if directly impacted by multiple stressors.
- A species susceptibility is context-dependent
- We did this at equilibrium. These hold for food webs at equilibrium, meaning that the topology of the food web stays the same. If the topology changes, then the dynamics would also change.

5 The way forward

Here, we clearly demonstrate that management and conservation need to move away from 631 single species and single stressor environmental assessments and conservation in favour of holistic approaches that explicitly consider the intricacies of ecological networks. By doing so, we join others in advocating for and providing evidence in support of the application of holistic environmental approaches and the conservation of ecological networks (e.q. McCann 2007; Tylianakis et al. 2008, 2010; O'Gorman et al. 2012; Thompson et al. 2018). Such a 636 transition will require infrastructure and investments in broader monitoring program target-637 ing non-commercial and unthreatened species, which is something we tend to avoid. It will 638 also require detailing the structure of networks, which is challenging (Paine 1992). Emerg-639 ing approaches such as isotopic analyses (???), environmental DNA (???), and machine and 640 deep learning (???) could be leveraged to assist and optimize the characteriation of food 641 web structure. It will also require embracing system's complexity rather than trying to distill 642 it. That is not say that simplifications are unwarranted; rather, say something clever here. 643 Conceptually, we favoured simplified model formulations to tackle the complexities of eval-644 uating the impacts of multiple stressors on food webs. Ample opportunities remain to glean deeper insights into the dynamics of ecological systems impacted by multiple systems. First, our results hint at the importance of interaction strengths to properly understand the impacts of stressors (see also e.g. Paine 1992; McCann et al. 1998; Montoya et al. 2009; 648 O'Gorman & Emmerson 2009; Gellner & McCann 2016) without fully exploring them. 649

650 Scaling up to food webs

We also considered interactions and stressors as binary events, when that is obviously not the
case. Our approach could however easily be adapted to consider the probabilities of interactions and exposure to stressors. This would however require very detailed environmental
data that we generally lack at the moment.

655 6 Figures

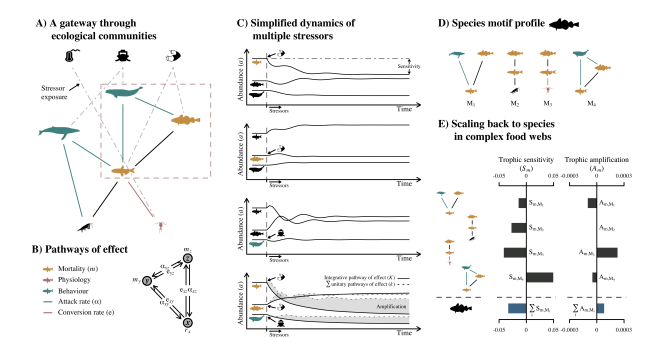


Figure 1: A) Conceptualization of the impacts of multiple stressors on complex food webs using a simplified 6-species food web composed of populations of beluga (Delphinapterus leucas) and humpback (Megaptera novaeangliae) whales, Atlantic cod (Gadus morhua), capelin (Mallotus villosus), krill (Euphausiacea) and copepods (Copepoda), and impacted by climate change-induced temperature anomalies, commercial shipping and trawl fishing. B) Stressors impact food webs by disrupting biological processes such as mortality (m), attack (α) and conversion (e) rates. Species are either directly or indirectly impacted by stressors through species interactions. C) Disrupting biological processes impacts community dynamics and results in variations in species abundances (trophic sensitivity; S_m). Food web complexity is simplified by focusing on subsets of species interactions – such as the omnivory interaction linking beluga, cod and capelin – called motifs. Impacts to individual biological processes arise through unitary pathways of effects (k) and result in contrasting population trajectories (1-3). Unitary pathways of effect combine to form an integrative pathway of effect (K)and collectively impact species in a system (4). The difference between the sum of trophic sensitivities to unitary pathways of effect $(k \in K)$ and trophic sensitivity to the integrative pathway of effect identifies synergistic and antagonistic impacts (trophic amplification; A_m). D) A species motif profile (M) is composed of all the position it holds in n-species interactions in a food web. E) A pathway of effect and resulting trophic sensitivities and amplification can be evaluated across a species motif profile, the sum of which summarizes that species overall trophic sensitivity and amplification.

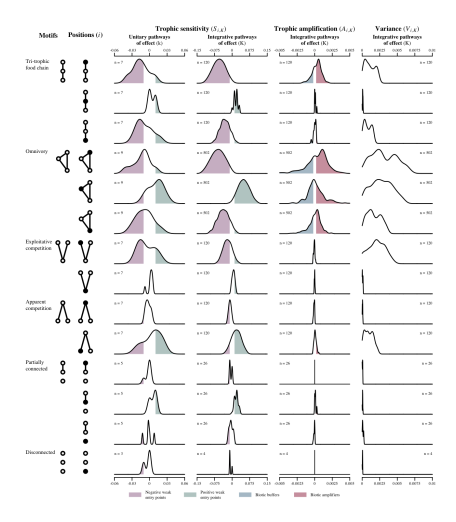


Figure 2: Density plots of the trophic sensitivity $(S_{i,K})$, amplification $(A_{i,K})$ and variance $(V_{i,K})$ of 13 unique motif positions (i) resulting from all unitary (k) and integrative (K) pathways of effect simulated on the dynamics of 3-species motifs (i.e. tri-trophic food chain, omnivory, exploitative competition, apparent competition, partially connected and disconnected). Impacts of stressors on individual biological processes – i.e. mortality (m), attack (α) and conversion (e) rates – form unitary pathways of effect that collectively impact food webs through integrative pathways of effect. A species trophic sensitivity is difference in equilibrium abundance before and after the permanent appearance of stressors; a species trophic amplification is the difference between its trophic sensitivity to an integrative pathway of effect and the sum of its trophic sensitivity to unitary pathways of effect. Pathways of effect that lead to a position being a weak entry point (i.e. highly sensitive to disturbances), a biotic buffer (i.e. synergistically impacted by stressors) or a biotic amplifier (i.e. antagonistically impacted by stressors) are identified as colored areas under the density curves.

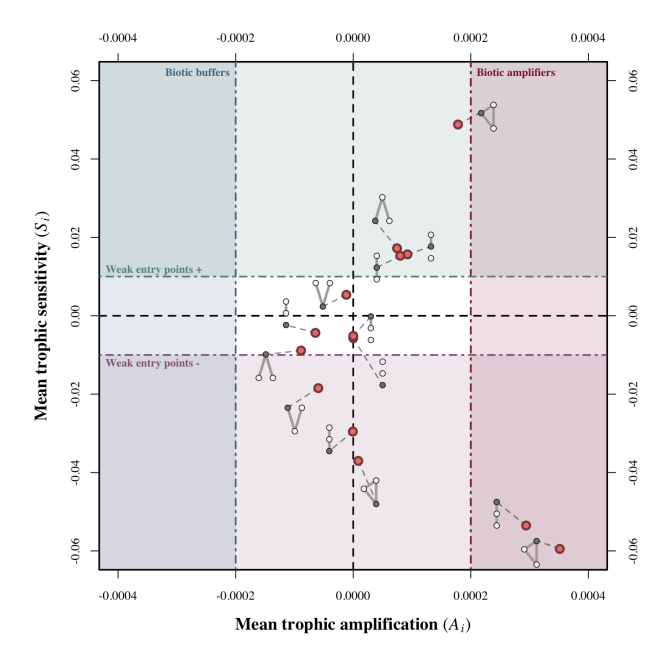


Figure 3: Mean trophic sensitivity (S_i) as a function of mean trophic amplification (A_i) to all possible pathways of effect (K) for the 13 unique 3-species motif positions explored, *i.e.* tri-trophic food chain, omnivory, exploitative competition, apparent competition, partially connected and disconnected. Motif positions identified as weak entry points (i.e. highly sensitive to disturbances), biotic buffers (i.e. synergistically impacted by stressors) and biotic amplifiers (i.e. antagonistically impacted by stressors) are identified as colored areas on the graph.

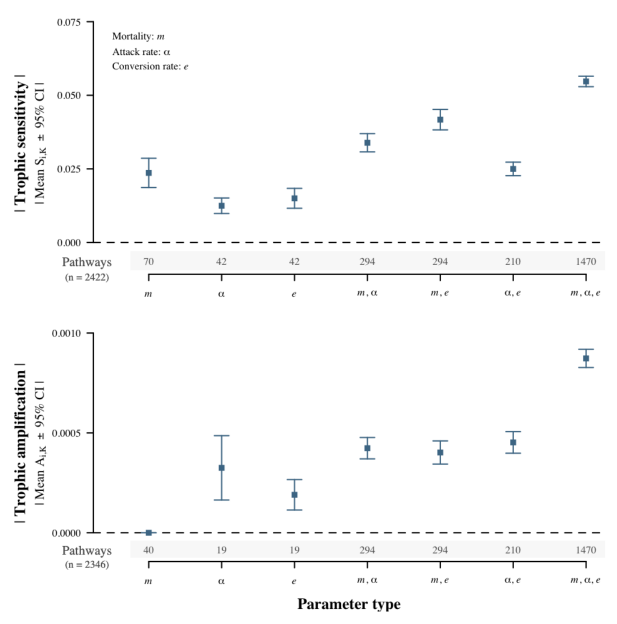


Figure 4: Mean absolute trophic sensitivities $(S_{i,K})$ and amplifications $(A_{i,K})$ as a function of the types of parameters forming an integrative pathway of effect (K). All pathways of effect impacting mortality (m), attack (α) and conversion (e), were grouped to evaluate the respective and combined importance of the types of parameters in driving trophic sensitivity and amplification.

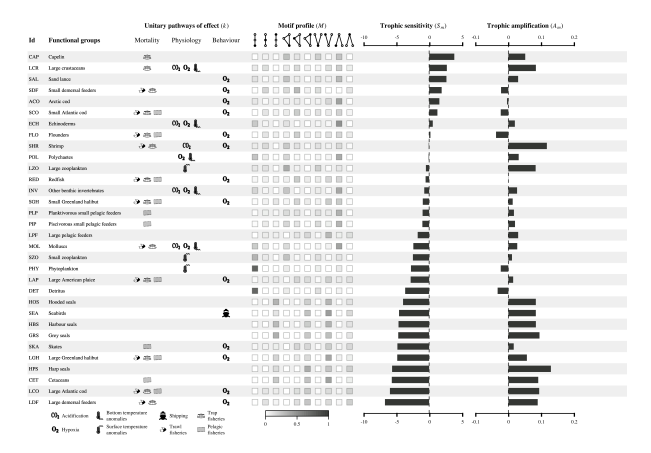


Figure 5: Trophic sensitivities (S_m) and amplifications (A_m) of species and function groups of the Northern Gulf of St. Lawrence in the mid-1980s (Morissette et al. 2003). A species trophic sensitivity and amplification is summarized using the sum of simulated trophic sensitivities $(S_{i,K})$ and amplifications $(A_{i,K})$ to pathways of effect (K; Figure 2) across a species motif profile (M). The left-hand side of the figure presents species and functional groups, unitary pathways of effects (k) arising from individual stressors and their potential impacts on population level mortality, physiology and behaviour, and the motif profile (M) of species and functional groups measured as the frequency of times they hold unique positions in tritrophic food chain, omnivory, exploitative and apparent competition interactions structuring of the food web. Main stressors in the Northern Gulf of St. Lawrence were fisheries (i.e. demersal destructive, demersal non-destructive high-bycatch and pelagic high-bycatch), climate change (i.e. ocean acidification, hypoxia and bottom and surface temperature anomalies), and shipping (Beauchesne et al. in review). Impacts of stressors on individual biological processes form unitary pathways of effect (k) that collectively impact food webs through integrative pathways of effect (K). The right-hand side of the figure presents trophic sensitivities and amplifications of species and functional groups. Negative or positive trophic sensitivities denote expected decreases or increases in species abundance as a response to pathways of effect. Species or functional groups with lowest or highest trophic sensitivities are positive or negative weak entry points (i.e. highly sensitive to disturbances), respectively. Negative or positive trophic amplifications identify species or functional groups expected to be impacted synergistically (i.e. biotic amplifiers) or antagonistically (i.e. biotic buffers) by stressors.

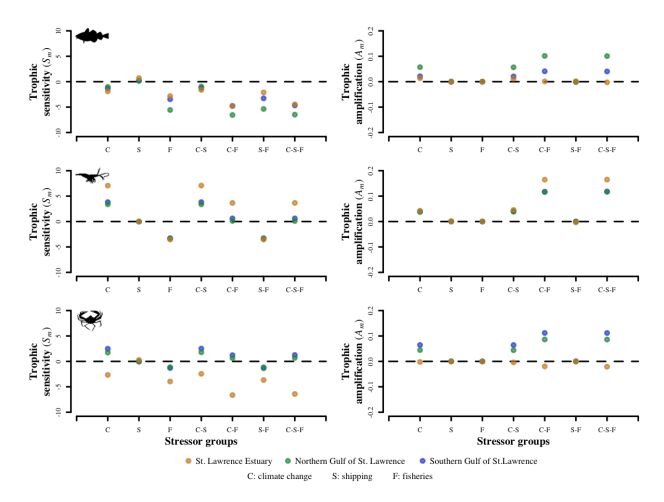


Figure 6: Comparison of the trophic sensitivities (S_m) and amplifications (A_m) to the impacts of different groups of stressors (Figure 5) for Atlantic cod ($Gadus\ morhua$), shrimp ($Pandalus\ borealis$) and large crustaceans (Crustacea) between the food webs of the Southern and Northern Gulf of St. Lawrence in the mid-1980s (Morissette $et\ al.\ 2003$; Savenkoff $et\ al.\ 2004$) and the St. Lawrence Estuary in the early 2010s (Savenkoff 2012).

7 References

- ⁶⁵⁷ Abrams, P.A. (1992). Predators that Benefit Prey and Prey that Harm Predators: Unusual
- Effects of Interacting Foraging Adaptation. The American Naturalist, 140, 573–600.
- Abrams, P.A., Menge, B.A., Mittelbach, G.G., Spiller, D.A. & Yodzis, P. (1996). The Role
- of Indirect Effects in Food Webs. In: Food Webs: Integration of Patterns & Dynamics (eds.
- Polis, G.A. & Winemiller, K.O.). Springer US, Boston, MA, pp. 371–395.

- Allesina, S., Bodini, A. & Bondavalli, C. (2006). Secondary extinctions in ecological net-
- works: Bottlenecks unveiled. Ecological Modelling, Special Issue on the Fourth European
- 664 Conference on Ecological Modelling, 194, 150–161.
- Archambault, P., Schloss, I.R., Grant, C. & Plante, S. (Eds.). (2017). Les hydrocarbures
- dans le golfe du Saint-Laurent Enjeux sociaux, Économiques et environnementaux. Notre
- 667 Golfe, Rimouski, Qc, Canada.
- Ban, N.C., Alidina, H.M. & Ardron, J.A. (2010). Cumulative impact mapping: Advances,
- relevance and limitations to marine management and conservation, using Canada's Pacific
- waters as a case study. Marine Policy, 34, 876–886.
- Bartley, T.J., McCann, K.S., Bieg, C., Cazelles, K., Granados, M. & Guzzo, M.M. et al.
- 672 (2019). Food web rewiring in a changing world. Nature Ecology & Evolution, 3, 345–354.
- Bascompte, J. (2009). Disentangling the Web of Life. Science, 325, 416–419.
- Bascompte, J. & Melián, C.J. (2005). Simple Trophic Modules for Complex Food Webs.
- 675 Ecology, 86, 2868–2873.
- ⁶⁷⁶ Beauchesne, D. (2019). Next Generation Planning Structuring and Sharing Environmental
- ⁶⁷⁷ Drivers Data for the St. Lawrence. Frontiers in Marine Science, 1, 1,
- Beauchesne, D., Grant, C., Gravel, D. & Archambault, P. (2016). L'évaluation des impacts
- cumulés dans l'estuaire et le golfe du Saint-Laurent : vers une planification systémique de
- l'exploitation des ressources. Le Naturaliste Canadien, 140, 45–55.
- Belley, R., Archambault, P., Sundby, B., Gilbert, F. & Gagnon, J.-M. (2010). Effects of
- bypoxia on benthic macrofauna and bioturbation in the Estuary and Gulf of St. Lawrence,
- 683 Canada. Continental Shelf Research, 30, 1302–1313.
- 684 Blanchard, J.L. (2015). A rewired food web. *Nature*, 527, 173–174.
- Boonstra, W.J., Ottosen, K.M., Ferreira, A.S.A., Richter, A., Rogers, L.A. & Pedersen,
- 686 M.W. et al. (2015). What are the major global threats and impacts in marine environments?

- Investigating the contours of a shared perception among marine scientists from the bottom-
- ⁶⁸⁸ up. Marine Policy, 60, 197–201.
- Bourdages, H., Marquis, M.-C., Nozères, C. & Ouellette-Plante, J. (2018). Assessment of
- 690 northern shrimp stocks in the Estuary and Gulf of St. Lawrence in 2017: Data from the
- research survey. DFO Can. Sci. Advis. Sec. Res. Doc. 2018/057. Iv + 67 p. (No.
- $692 \quad 2018/057$).
- Bove, C.B., Ries, J.B., Davies, S.W., Westfield, I.T., Umbanhowar, J. & Castillo, K.D.
- 694 (2019). Common Caribbean corals exhibit highly variable responses to future acidification
- and warming. Proceedings of the Royal Society B: Biological Sciences, 286, 20182840.
- 696 Camacho, J., Stouffer, D.B. & Amaral, L.A.N. (2007). Quantitative analysis of the local
- structure of food webs. Journal of Theoretical Biology, 246, 260–268.
- ⁶⁹⁸ Chabot, D. & Claireaux, G. (2008). Environmental hypoxia as a metabolic constraint on
- fish: The case of Atlantic cod, Gadus morhua. Marine Pollution Bulletin, 5th International
- Conference on Marine Pollution and Ecotoxicology, 57, 287–294.
- Christensen, M.R., Graham, M.D., Vinebrooke, R.D., Findlay, D.L., Paterson, M.J. &
- Turner, M.A. (2006). Multiple anthropogenic stressors cause ecological surprises in boreal
- lakes. Global Change Biology, 12, 2316–2322.
- Christiansen, F., Rasmussen, M. & Lusseau, D. (2013). Whale watching disrupts feeding
- activities of minke whales on a feeding ground. Marine Ecology Progress Series, 478, 239–
- 706 251.
- Cirtwill, A.R. & Stouffer, D.B. (2015). Concomitant predation on parasites is highly variable
- but constrains the ways in which parasites contribute to food web structure. The Journal
- 709 of Animal Ecology, 84, 734–744.
- 710 Clavel, J., Julliard, R. & Devictor, V. (2011). Worldwide decline of specialist species: Toward
- a global functional homogenization? Frontiers in Ecology and the Environment, 9, 222–228.

- Côté, I.M., Darling, E.S. & Brown, C.J. (2016). Interactions among ecosystem stressors and
- their importance in conservation. Proceedings of the Royal Society B: Biological Sciences,
- 714 283, 20152592.
- Crain, C.M., Kroeker, K. & Halpern, B.S. (2008). Interactive and cumulative effects of
- multiple human stressors in marine systems. *Ecology Letters*, 11, 1304–1315.
- Darling, E.S. & Côté, I.M. (2008). Quantifying the evidence for ecological synergies. *Ecology*
- 718 Letters, 11, 1278–1286.
- David, P., Thébault, E., Anneville, O., Duyck, P.F., Chapuis, E. & Loeuille, N. (2017).
- 720 Chapter One Impacts of Invasive Species on Food Webs: A Review of Empirical Data.
- In: Advances in Ecological Research, Networks of Invasion: A Synthesis of Concepts (eds.
- Bohan, D.A., Dumbrell, A.J. & Massol, F.). Academic Press, pp. 1–60.
- De Laender, F. (2018). Community-and ecosystem-level effects of multiple environmental
- change drivers: Beyond null model testing. Global change biology, 24, 5021–5030.
- Dempsey, D.P., Gentleman, W.C., Pepin, P. & Koen-Alonso, M. (2018). Explanatory Power
- of Human and Environmental Pressures on the Fish Community of the Grand Bank before
- and after the Biomass Collapse. Frontiers in Marine Science, 5.
- Developers, T.S. (2019). SageMath, the Sage Mathematics Software System Version 8.8.
- DFO. (2016). Zonal Interchange File Format (ZIFF) data. A compilation of landing data
- 730 from logbook data between 2010 and 2015. Gestion des données, Institut Maurice Lam-
- ontagne, Department of Fisheries and Oceans (DFO) Mont-Joli, Canada. Department of
- Fisheries and Oceans.
- Eby, L.A., Crowder, L.B., McClellan, C.M., Peterson, C.H. & Powers, M.J. (2005). Habitat
- degradation from intermittent hypoxia: Impacts on demersal fishes. Marine Ecology Progress
- 735 Series, 291, 249–262.
- 736 Eklöf, A. & Ebenman, B. (2006). Species loss and secondary extinctions in simple and

- complex model communities. Journal of Animal Ecology, 75, 239–246.
- El-Sabh, M.I. & Silverberg, N. (1990). Oceanography of a Large-Scale Estuarine System.
- 739 Springer New York.
- Estes, J.A. & Palmisano, J.F. (1974). Sea Otters: Their Role in Structuring Nearshore
- 741 Communities. Science, 185, 1058–1060.
- Estes, J.A., Terborgh, J., Brashares, J.S., Power, M.E., Berger, J. & Bond, W.J. et al.
- 743 (2011). Trophic Downgrading of Planet Earth. Science, New Series, 333, 301–306.
- Fabry, V.J., Seibel, B.A., Feely, R.A. & Orr, J.C. (2008). Impacts of ocean acidification on
- marine fauna and ecosystem processes. ICES Journal of Marine Science, 65, 414–432.
- Frank, K.T., Petrie, B., Choi, J.S. & Leggett, W.C. (2005). Trophic Cascades in a Formerly
- Cod-Dominated Ecosystem. Science, 308, 1621–1623.
- Galic, N., Sullivan, L.L., Grimm, V. & Forbes, V.E. (2018). When things don't add up:
- Quantifying impacts of multiple stressors from individual metabolism to ecosystem process-
- ₇₅₀ ing. Ecology Letters, 21, 568–577.
- Gavrilchuk, K., Lesage, V., Ramp, C., Sears, R., Bérubé, M. & Bearhop, S. et al. (2014).
- Trophic niche partitioning among sympatric baleen whale species following the collapse of
- groundfish stocks in the Northwest Atlantic. Marine Ecology Progress Series, 497, 285–301.
- Gellner, G. & McCann, K.S. (2016). Consistent role of weak and strong interactions in high-
- and low-diversity trophic food webs. *Nature Communications*, 7, 11180.
- Giakoumi, S., Halpern, B.S., Michel, L.N., Gobert, S., Sini, M. & Boudouresque, C.-F. et al.
- 757 (2015). Towards a framework for assessment and management of cumulative human impacts
- on marine food webs. Conservation Biology, 29, 1228–1234.
- Gonzalez, A. & Loreau, M. (2009). The Causes and Consequences of Compensatory Dy-
- namics in Ecological Communities. Annual Review of Ecology, Evolution, and Systematics,
- 761 40, 393–414.

- Griffen, B.D., Belgrad, B.A., Cannizzo, Z.J., Knotts, E.R. & Hancock, E.R. (2016). Re-
- thinking our approach to multiple stressor studies in marine environments. Marine Ecology
- 764 Progress Series, 543, 273–281.
- Guiden, P.W., Bartel, S.L., Byer, N.W., Shipley, A.A. & Orrock, J.L. (2019). Predator-
- Prey Interactions in the Anthropocene: Reconciling Multiple Aspects of Novelty. Trends in
- 767 Ecology & Evolution, 34, 616–627.
- Halpern, B.S., Frazier, M., Afflerbach, J., Lowndes, J.S., Micheli, F. & O'Hara, C. et al.
- (2019). Recent pace of change in human impact on the world's ocean. Scientific Reports, 9,
- 770 11609.
- Halpern, B.S. & Fujita, R. (2013). Assumptions, challenges, and future directions in cumu-
- lative impact analysis. *Ecosphere*, 4, art131.
- Harborne, A.R., Rogers, A., Bozec, Y.-M. & Mumby, P.J. (2017). Multiple Stressors and
- the Functioning of Coral Reefs. Annual Review of Marine Science, 9, 445–468.
- Hodgson, E.E., Essington, T.E. & Halpern, B.S. (2017). Density dependence governs when
- population responses to multiple stressors are magnified or mitigated. Ecology, 98, 2673–
- 777 2683.
- Hodgson, E.E. & Halpern, B.S. (2019). Investigating cumulative effects across ecological
- scales. Conservation Biology, 33, 22–32.
- Hodgson, E.E., Halpern, B.S. & Essington, T.E. (2019). Moving Beyond Silos in Cumulative
- ⁷⁸¹ Effects Assessment. Frontiers in Ecology and Evolution, 7.
- Holt, R.D. (1977). Predation, apparent competition, and the structure of prev communities.
- Theoretical population biology, 12, 197–229.
- Holt, R.D. (1997). Community modules. In: Multitrophic Interactions in Terrestrial Ecosys-
- tems, 36th Symposium of the British Ecological Society (eds. Grange, A.C. & Brown, V.K.).
- Blackwell Science, Oxford, pp. 333–349.

- Holt, R.D. & Hochberg, M.E. (2001). Indirect interactions, community modules and bio-
- logical control: A theoretical perspective. Evaluating indirect ecological effects of biological
- 789 control, 13–37.
- ⁷⁹⁰ Ives, A.R. (1995). Measuring Resilience in Stochastic Systems. *Ecological Monographs*, 65,
- 791 217-233.
- Jackson, J.B.C., Kirby, M.X., Berger, W.H., Bjorndal, K.A., Botsford, L.W. & Bourque,
- ⁷⁹³ B.J. et al. (2001). Historical Overfishing and the Recent Collapse of Coastal Ecosystems.
- 794 Science, 293, 629–637.
- Jackson, M.C., Loewen, C.J.G., Vinebrooke, R.D. & Chimimba, C.T. (2016). Net effects
- of multiple stressors in freshwater ecosystems: A meta-analysis. Global Change Biology, 22,
- 797 180–189.
- ⁷⁹⁸ Klaise, J. & Johnson, S. (2017). The origin of motif families in food webs. *Scientific Reports*,
- 799 7, 16197.
- Kortsch, S., Primicerio, R., Fossheim, M., Dolgov, A.V. & Aschan, M. (2015). Climate
- change alters the structure of arctic marine food webs due to poleward shifts of boreal
- generalists. Proceedings of the Royal Society B: Biological Sciences, 282, 20151546.
- Kroeker, K.J., Kordas, R.L., Crim, R., Hendriks, I.E., Ramajo, L. & Singh, G.S. et al.
- 804 (2013). Impacts of ocean acidification on marine organisms: Quantifying sensitivities and
- interaction with warming. Global Change Biology, 19, 1884–1896.
- Lapointe, B.E., Brewton, R.A., Herren, L.W., Porter, J.W. & Hu, C. (2019). Nitrogen
- enrichment, altered stoichiometry, and coral reef decline at Looe Key, Florida Keys, USA:
- A 3-decade study. Marine Biology, 166, 108.
- Lesage, V., Omrane, A., Doniol-Valcroze, T. & Mosnier, A. (2017). Increased proximity of
- vessels reduces feeding opportunities of blue whales in the St. Lawrence Estuary, Canada.
- 811 Endangered Species Research, 32, 351–361.

- Liess, M., Foit, K., Knillmann, S., Schäfer, R.B. & Liess, H.-D. (2016). Predicting the
- synergy of multiple stress effects. Scientific Reports, 6, 32965.
- McCann, K. (2007). Protecting biostructure. Nature, 446, 29–29.
- McCann, K., Hastings, A. & Huxel, G.R. (1998). Weak trophic interactions and the balance
- 816 of nature. *Nature*, 395, 794.
- McClanahan, T.R., Graham, N.A. & Darling, E.S. (2014). Coral reefs in a crystal ball:
- Predicting the future from the vulnerability of corals and reef fishes to multiple stressors.
- 819 Current Opinion in Environmental Sustainability, Environmental change issues, 7, 59–64.
- Menge, B.A. (1995). Indirect Effects in Marine Rocky Intertidal Interaction Webs: Patterns
- and Importance. Ecological Monographs, 65, 21–74.
- Milo, R., Shen-Orr, S., Itzkovitz, S., Kashtan, N., Chklovskii, D. & Alon, U. (2002). Network
- Motifs: Simple Building Blocks of Complex Networks. Science, 298, 824–827.
- Monteiro, A.B. & Faria, L.D.B. (2016). The interplay between population stability and food-
- web topology predicts the occurrence of motifs in complex food-webs. Journal of theoretical
- 826 biology, 409, 165–171.
- Montoya, J., Woodward, G., Emmerson, M.C. & Solé, R.V. (2009). Press perturbations and
- indirect effects in real food webs. *Ecology*, 90, 2426–2433.
- Morissette, L., Castonguay, M., Savenkoff, C., Swain, D.P., Chabot, D. & Bourdages, H. et al.
- (2009). Contrasting changes between the northern and southern Gulf of St. Lawrence ecosys-
- tems associated with the collapse of groundfish stocks. Deep Sea Research Part II: Topical
- studies in Oceanography, The Proceedings of the ECONORTH Symposium on Ecosystem
- B33 Dynamics in the Norwegian Sea and Barents Sea, 56, 2117–2131.
- Morissette, L., Despatie, S.-P., Savenkoff, C., Hammill, M.O., Bourdages, H. & Chabot, D.
- 2003). Data gathering and input parameters to construct ecosystem models for the northern
- Gulf of St. Lawrence (Mid-1980 s). Can. Tech. Rep. Fish. Aquat. Sci./Rapp. Tech. Can.

- 837 Sci. Halieut. Aquat., 100.
- Myers, R.A. & Worm, B. (2003). Rapid worldwide depletion of predatory fish communities.
- 839 Nature, 423, 280.
- O'Brien, A.L., Dafforn, K.A., Chariton, A.A., Johnston, E.L. & Mayer-Pinto, M. (2019).
- After decades of stressor research in urban estuarine ecosystems the focus is still on single
- stressors: A systematic literature review and meta-analysis. Science of The Total Environ-
- 843 ment.
- O'Gorman, E.J. & Emmerson, M.C. (2009). Perturbations to trophic interactions and the
- stability of complex food webs. Proceedings of the National Academy of Sciences, 106,
- 846 13393-13398.
- O'Gorman, E.J., Fitch, J.E. & Crowe, T.P. (2012). Multiple anthropogenic stressors and
- the structural properties of food webs. *Ecology*, 93, 441–448.
- Paine, R.T. (1980). Food Webs: Linkage, Interaction Strength and Community Infrastruc-
- ture. Journal of Animal Ecology, 49, 667–685.
- Paine, R.T. (1992). Food-web analysis through field measurement of per capita interaction
- 852 strength. *Nature*, 355, 73.
- Pauly, D., Christensen, V., Dalsgaard, J., Froese, R. & Torres, F. (1998). Fishing Down
- 854 Marine Food Webs. *Science*, 279, 860–863.
- Piggott, J.J., Townsend, C.R. & Matthaei, C.D. (2015). Reconceptualizing synergism and
- antagonism among multiple stressors. Ecology and Evolution, 5, 1538–1547.
- Pillet, M., Dupont-Prinet, A., Chabot, D., Tremblay, R. & Audet, C. (2016). Effects of
- exposure to hypoxia on metabolic pathways in northern shrimp (Pandalus borealis) and
- Greenland halibut (Reinhardtius hippoglossoides). Journal of Experimental Marine Biology
- and Ecology, 483, 88–96.
- Ripple, W.J. & Beschta, R.L. (2003). Wolf reintroduction, predation risk, and cottonwood

- recovery in Yellowstone National Park. Forest Ecology and Management, 184, 299–313.
- Ripple, W.J., Newsome, T.M., Wolf, C., Dirzo, R., Everatt, K.T. & Galetti, M. et al. (2015).
- Collapse of the world's largest herbivores. Science Advances, 1, e1400103.
- Ritchie, E.G. & Johnson, C.N. (2009). Predator interactions, mesopredator release and
- biodiversity conservation. Ecology Letters, 12, 982–998.
- Savenkoff, C. (2012). Input data and parameter estimates for ecosystem models of the lower
- 868 St. Lawrence Estuary (20082010). Canadian Technical Report of Fisheries and Aquatic
- Sciences 2999, Department of Fisheries and Oceans, Mont-Joli, Québec, Canada.
- 870 Savenkoff, C., Bourdages, H., Swain, D.P., Despatie, S.-P., Hanson, J.M. & Méthot, R. et
- al. (2004). Input data and parameter estimates for ecosystem models of the southern Gulf
- of St. Lawrence (mid-1980s and mid-1990s). Canadian Technical Report of Fisheries and
- Aquatic Sciences 2529, Department of Fisheries and Oceans, Mont-Joli, Québec, Canada.
- 874 Savenkoff, C., Swain, D.P., Hanson, J.M., Castonguay, M., Hammill, M.O. & Bourdages, H.
- et al. (2007). Effects of fishing and predation in a heavily exploited ecosystem: Comparing
- periods before and after the collapse of groundfish in the southern Gulf of St. Lawrence
- 877 (Canada). Ecological Modelling, 204, 115–128.
- 878 Savenkoff, C., Vézina, A.F., Roy, S., Klein, B., Lovejoy, C. & Therriault, J.C. et al. (2000).
- Export of biogenic carbon and structure and dynamics of the pelagic food web in the Gulf
- of St. Lawrence Part 1. Seasonal variations. Deep Sea Research Part II: Topical Studies in
- 881 Oceanography, 47, 585–607.
- 882 Schäfer, R.B. & Piggott, J.J. (2018). Advancing understanding and prediction in multiple
- stressor research through a mechanistic basis for null models. Global Change Biology, 24,
- 884 1817–1826.
- Schloss, I.R., Archambault, P., Beauchesne, D., Cusson, M., Ferreyra, G. & Levasseur, M.
- et al. (2017). Cumulative potential impacts of the stress factors associated with human ac-

- tivities on the St. Lawrence marine ecosystem. In: Hydrocarbon in the Gulf of St. Lawrence
- Social, economic and environmental issues (eds. Archambault, P., Schloss, I.R., Grant, C.
- & Plante, S.). Notre Golfe, Rimouski, Qc, Canada, pp. 133–165.
- 890 Stier, A.C., Samhouri, J.F., Novak, M., Marshall, K.N., Ward, E.J. & Holt, R.D. et al.
- 891 (2016). Ecosystem context and historical contingency in apex predator recoveries. Science
- 892 Advances, 2, e1501769.
- 893 Stouffer, D.B. & Bascompte, J. (2010). Understanding food-web persistence from local to
- global scales. Ecology Letters, 13, 154–161.
- 895 Stouffer, D.B. & Bascompte, J. (2011). Compartmentalization increases food-web persis-
- tence. Proceedings of the National Academy of Sciences, 108, 3648–3652.
- 897 Stouffer, D.B., Camacho, J., Jiang, W. & Nunes Amaral, L.A. (2007). Evidence for the
- existence of a robust pattern of prey selection in food webs. Proceedings of the Royal Society
- 899 B: Biological Sciences, 274, 1931–1940.
- Stouffer, D.B., Sales-Pardo, M., Sirer, M.I. & Bascompte, J. (2012). Evolutionary Conser-
- vation of Species' Roles in Food Webs. Science, 335, 1489–1492.
- ⁹⁰² Team, R.C. (2019). R: A Language and Environment for Statistical Computing.
- Thompson, P.L., MacLennan, M.M. & Vinebrooke, R.D. (2018). Species interactions cause
- non-additive effects of multiple environmental stressors on communities. Ecosphere, 9,
- e02518.
- Tylianakis, J.M., Didham, R.K., Bascompte, J. & Wardle, D.A. (2008). Global change and
- species interactions in terrestrial ecosystems. Ecology letters, 11, 1351–1363.
- Tylianakis, J.M., Laliberte, E., Nielsen, A. & Bascompte, J. (2010). Conservation of species
- one interaction networks. Biological Conservation, 143, 2270–2279.
- Vander Zanden, M.J., Casselman, J.M. & Rasmussen, J.B. (1999). Stable isotope evidence
- for the food web consequences of species invasions in lakes. Nature, 401, 464–467.

- Wiedenmann, J., D'Angelo, C., Smith, E.G., Hunt, A.N., Legiret, F.-E. & Postle, A.D. et
- 913 al. (2013). Nutrient enrichment can increase the susceptibility of reef corals to bleaching.
- 914 Nature Climate Change, 3, 160–164.
- 915 Wootton, J.T. (1993). Indirect Effects and Habitat Use in an Intertidal Community: Inter-
- action Chains and Interaction Modifications. The American Naturalist, 141, 71–89.
- Wootton, J.T. (2002). Indirect effects in complex ecosystems: Recent progress and future
- challenges. Journal of Sea Research, Structuring Factors of Shallow Marine Coastal Com-
- 919 munities, Part I, 48, 157–172.
- Worm, B., Barbier, E.B., Beaumont, N., Duffy, J.E., Folke, C. & Halpern, B.S. et al. (2006).
- ⁹²¹ Impacts of Biodiversity Loss on Ocean Ecosystem Services. Science, 314, 787–790.
- Worm, B. & Myers, R.A. (2003). Meta-Analysis of Cod-Shrimp Interactions Reveals Top-
- Down Control in Oceanic Food Webs. *Ecology*, 84, 162–173.
- Yodzis, P. (2000). Diffuse Effects in Food Webs. *Ecology*, 81, 261–266.