

~~Feeding on multiple prey at a time:~~ In defense of the original Type I functional response:

The frequency and population-dynamic effects of ~~functional response linearity~~ feeding on multiple prey at a time

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RUNNING TITLE: Multi-prey functional response

Code and data availability

The FoRAGE compilation is available from the *Knowledge Network for Biocomplexity* (DeLong & Uiterwaal, 2018). All code and data are available at https://github.com/marknovak/FR_n-prey-at-a-time and [FigShare_url_posted_after_acceptance](#).

Author contributions

MN conceived of the study, performed the analyses, and wrote the first draft. JPD compiled functional response datasets. KEC and JPD discussed the analyses and edited the manuscript.

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Conflict of interest disclosure

We declare to have no conflict of interest relating to the content of this manuscript.

1 Abstract

2 Ecologists differ in the degree to which they consider the linear Type I functional response to
3 be an unrealistic versus sufficient representation of predator feeding rates. Empiricists tend to
4 consider it unsuitably non-mechanistic and theoreticians tend to consider it necessarily simple.
5 Holling’s original rectilinear ~~model-Type I response~~ is dismissed by satisfying neither desire,
6 with most compromising on the smoothly saturating Type II response for which searching and
7 handling are assumed to be mutually exclusive activities. We derive a “multiple-prey-at-a-time”
8 ~~functional~~-response and a generalization that includes the Type III to reflect predators that can
9 continue to search when handling an arbitrary number of already-captured prey. The multi-
10 prey model clarifies the empirical relevance of ~~Holling’s-the~~ linear and rectilinear models and
11 the conditions under which linearity can be a mechanistically-reasoned description of predator
12 feeding rates, even when handling times are long. We find support for ~~the-presence-of~~ linearity in
13 35% of 2,591 compiled empirical datasets ~~;-and-find-evidence-and support for the hypothesis~~ that
14 larger predator-prey body-mass ratios permit predators to search while handling greater numbers
15 of prey. Incorporating the multi-prey response into the Rosenzweig-MacArthur population-
16 dynamics model reveals that a non-exclusivity of searching and handling can lead to coexistence
17 states and dynamics that are not anticipated by theory built on ~~Holling’s-traditional-the Type~~
18 ~~I, II, or III response~~ models. In particular, it can lead to bistable fixed-point and limit-cycle
19 dynamics with long-term crawl-by transients between them under conditions where abundance
20 ratios reflect top-heavy food webs and the functional response is linear. We conclude that
21 functional response linearity should not be considered empirically unrealistic but also that ~~that~~
22 ~~more-bounded-conclusions-more cautious inferences~~ should be drawn in theory presuming the
23 ~~linear~~ Type I to be appropriate.

24 KEYWORDS: ~~type-0 functional response~~, *generalized Holling model, predator-prey body-mass ra-*
25 *tio, consumer-resource cycles, long transients, alternative states, top-heavy food webs,* ~~digestion,~~

27 Introduction

28 The way that predator feeding rates respond to changes in prey abundance, their functional
29 response, is key to determining how species affect each other’s populations (Murdoch & Oaten,
30 1975). The challenge of empirically understanding and appropriately modeling functional re-
31 sponses is therefore central to myriad lines of ecological research that extend even to the pro-
32 jection of Earth’s rapidly changing climate (DeLong, 2021; Rohr *et al.*, 2023).

33 The simplest functional response model, the ~~Holling~~-Type I response, describes feeding rates
34 as increasing linearly with prey abundance. Interpreted to represent an analytically-tractable
35 first-order approximation to all other prey-dependent forms (Lotka, 1925; Volterra, 1926), its
36 simplicity has caused the Type I to become foundational to theory across Ecology’s many sub-
37 disciplines. Nonetheless, there is a common and persistent belief among empirically-minded
38 ecologists that the Type I response is unrealistic and artifactual. Indeed, it is typically dismissed
39 *a priori* from both empirical and theoretical efforts to “mechanistically” characterize predator
40 feeding rates (e.g., Baudrot *et al.*, 2016; Kalinkat *et al.*, 2023). This dismissal is similarly levied at
41 the piecewise rectilinear ~~(a.k.a. Type 0) model, originally depicted by Holling (1959b)~~response
42 (e.g., Koen-Alonso, 2007), originally referred to by Holling (1959a) as the Type I ~~(Denny, 2014)~~
43 response (Denny, 2014; Holling, 1965), in which feeding rates increase linearly with prey abun-
44 dance to ~~an abrupt maximum(e.g., Koen-Alonso, 2007)~~a relatively abrupt maximum. Support
45 comes from syntheses concluding functional response linearity to be rare, with feeding rates
46 more consistent with smoothly saturating Type II responses being by far the more frequently

47 inferred (Dunn & Hovel, 2020; Jeschke *et al.*, 2004).

48 Countering justifications for the continued use of the linear Type I response in theory relate
49 to the challenge of extrapolating the inferences of mostly small-scale experiments to natural field
50 conditions (DeLong, 2021; Griffen, 2021; Jeschke *et al.*, 2004; Li *et al.*, 2018; Novak & Stouffer,
51 2021b; Novak *et al.*, 2017; Uiterwaal *et al.*, 2018). For example, prey abundances in the field may
52 vary relatively little over relevant scales, making linearity a sufficiently good approximation for
53 how species affect each other (Wootton & Emmerson, 2005). Further, prey abundances in nature
54 are typically much lower than those used in experiments to elicit predator saturation (Coblentz
55 *et al.*, 2023), which may consequently be rare in nature (but see Jeschke, 2007). Functional
56 responses could therefore be approximately linear even for predator-prey interactions having
57 very long handling times (e.g., Novak, 2010).

58 Here, our goal is to offer a further way of resolving ecologists' views on the linear and
59 rectilinear models by considering a reason for feeding rates to exhibit linear prey dependence
60 over a large range of prey abundances. This reason is not one of experimental design or variation
61 in prey abundances per se, but rather is attributable to the mechanics of predator-prey biology:
62 the ability of predator individuals to handle and search for more than just one prey individual
63 at a time (i.e. the non-exclusivity of handling and searching). Although it is straightforward
64 to show how the linear Type I can emerge when handling times are assumed to be entirely
65 inconsequential, and although functional response forms that could result from a non-exclusivity
66 of handling and searching have been considered before (Jeschke *et al.*, 2002; 2004; Mills, 1982;

67 Sjöberg, 1980; Stouffer & Novak, 2021), we contend that the empirical relevance and potential
68 prevalence of such “multiple-prey-at-a-time” feeding (henceforth multi-prey feeding) are not
69 sufficiently understood due to an inappropriately literal interpretation of the “handling time”
70 parameter of functional response models (see *Discussion* and DeLong, 2021; Jeschke *et al.*, 2002;
71 2004). Likewise, the potential implications of multi-prey feeding for predator-prey coexistence
72 and population dynamics have not, to our knowledge, been assessed.

73 We begin by providing a derivation of a simple multi-prey functional response model for
74 a single predator population feeding on a single prey species that relaxes the assumption of
75 searching and handling being exclusive activities. This derivation helps clarify the empirical
76 relevance of ~~Holling’s~~ the linear and rectilinear models and the conditions under which these
77 can be good descriptions of feeding rates (Jeschke *et al.*, 2004). We then further generalize the
78 multi-prey model to include the Holling-Real Type III response and fit all models to a large
79 number of datasets assembled in a new version of the FoRAGE compilation (Uiterwaal *et al.*,
80 2022). This allows us to quantify the potential prevalence of multi-prey feeding and to test the
81 hypothesis that larger predator-prey body-mass ratios permit predators to handle and search for
82 more prey at a time. We also assess the predicted association between larger body-mass ratios
83 and more pronounced Type III responses. Finally, we incorporate the multi-prey response into
84 the Rosenzweig & MacArthur (1963) “paradox of enrichment” population-dynamic model to
85 assess its potential influence on predator-prey coexistence and dynamics.

86 With our statistical analyses demonstrating that many datasets are indeed consistent with

87 multi-prey feeding and that larger predator-prey body-mass ratios are indeed more conducive
88 to multi-prey feeding (and more pronounced Type III responses), our mathematical analyses
89 demonstrate that even small increases in the number of prey that a predator can handle at a
90 time can lead to dynamics that are not anticipated by theory assuming ~~Holling's traditional~~
91 ~~functional response forms~~ Type I, II, or III response models.

92 **A functional response for multi-prey feeding**

93 **Holling's Type II response**

94 The multi-prey model may be understood most easily by a contrast to Holling's Type II model
95 ~~(a.k.a. the disc equation).~~ (a.k.a. the disc equation, Holling, 1959b). There are several ways
96 to derive the Type II (Garay, 2019), but the most common approach takes the perspective of a
97 single predator individual that can either be searching or "handling" a single prey individual at
98 any point in time: In the time T_S that a predator spends searching it will encounter prey at a
99 rate proportional to their abundance N , thus the number of prey eaten is $N_e = aNT_S$ where a
100 is the attack rate. Rearranging we have $T_S = N_e/aN$. With a handling time h for each prey,
101 the length of time spent handling all eaten prey will be $T_H = hN_e$. Given the presumed mutual
102 exclusivity of the two activities, $T_S = T - T_H$ where T is the total time available. Substituting
103 the second and third equations into the fourth, it follows that $N_e = aNT/(1 + ahN)$. We arrive
104 at the predator individual's feeding rate by dividing by T , presuming steady-state predator
105 behavior and constant prey abundances.

106 An alternative derivation on which we build to derive the multi-prey model considers a tem-

107 poral snapshot of a predator population composed of many identical and independent individuals
 108 (see also Real (1977) and the *Supplementary Materials*). Assuming constant prey abundance
 109 and steady-state conditions, the rate at which searching individuals P_S become handling indi-
 110 viduals P_H must equal the rate at which handling individuals become searching individuals such
 111 that $aNP_S = \frac{1}{h}P_H$, visually represented as

$$\begin{array}{ccc}
 & N & \\
 & \swarrow & \\
 P_S & \xrightarrow{a} & P_H \\
 & \searrow & \\
 & N_e & \\
 \hline
 & N & \\
 & \swarrow & \\
 P_S & \xrightarrow{a} & P_H \\
 & \searrow & \\
 & N_e & \\
 \hline
 \end{array} \tag{1}$$

115 Given the mutual exclusivity of searching and handling, $P_S = P - P_H$, where P is the total
 116 number of predators. Substituting this second equation into the first, it follows that the total
 117 number of handling predators $P_H = ahNP/(1 + ahN)$. Eaten prey are generated at rate $\frac{1}{h}P_H$
 118 by all these predators as they revert back to searching. We thus obtain Holling's Type II
 119 (per-predator) model by multiplying the proportion of handling predators, P_H/P , by $\frac{1}{h}$.

120 The multi-prey response

121 The derivation of the multi-prey response follows the same logic but assumes that searching
 122 and handling are not mutually exclusive activities until an arbitrary count of n prey individuals
 123 are being handled (see the *Supplementary Materials* for a more explicit derivation); handling
 124 need not reflect literal handling but rather could also reflect a process of digestion and stomach
 125 fullness.

126 With constant prey abundance and steady-state conditions as before, we assume that preda-
 127 tors continue to handle each prey with handling time h and that predators handling less than
 128 n prey continue to search for and encounter prey at rate aN . The rate at which searching
 129 individuals P_S become P_{H_1} individuals handling one prey is then equal to the rate at which they
 130 revert back to being searching individuals with no prey, thus $P_{H_1} = ahNP_S$. Likewise, the rate
 131 at which P_{H_1} individuals become P_{H_2} individuals handling two prey must equal the rate these
 132 revert back to handling just one prey, thus $P_{H_2} = ahNP_{H_1} = (ahN)^2P_S$. That is,

$$\begin{array}{c}
 133 \quad P_S \xrightarrow[\frac{N}{N_e}]{\frac{a}{\sqrt{1/h}}} P_{H_1} \xrightarrow[\frac{N}{N_e}]{\frac{a}{\sqrt{1/h}}} P_{H_2} \xrightarrow[\frac{N}{N_e}]{\frac{a}{\sqrt{1/h}}} \dots \xrightarrow[\frac{N}{N_e}]{\frac{a}{\sqrt{1/h}}} P_{H_n} \cdot \\
 \hline
 134 \quad P_S \xrightarrow[\frac{N}{N_e}]{\frac{a}{\sqrt{1/h}}} P_{H_1} \xrightarrow[\frac{N}{N_e}]{\frac{a}{\sqrt{1/h}}} P_{H_2} \xrightarrow[\frac{N}{N_e}]{\frac{a}{\sqrt{1/h}}} \dots \xrightarrow[\frac{N}{N_e}]{\frac{a}{\sqrt{1/h}}} P_{H_n} \cdot \quad (2) \\
 135 \quad \text{~~~~~}
 \end{array}$$

136 Generalizing by induction, the number of predators P_{H_i} handling i prey will be $(ahN)^iP_S$ for
 137 $i \in \{1, 2, 3, \dots, n\}$. The proportion of predators handling i prey at any point in time will then
 138 be

$$139 \quad \frac{P_{H_i}}{P} = \frac{(ahN)^iP_S}{P_S + P_{H_1} + \dots + P_{H_n}} = \frac{(ahN)^i}{1 + \sum_{i=1}^n (ahN)^i} \quad (3)$$

140 (Fig. S.1). With each of these groups generating eaten prey at rate $\frac{1}{h}P_{H_i}$, the per predator
 141 feeding rate of the population is obtained by a summation across all groups, giving

$$142 \quad f(N) = \frac{\frac{1}{h} \sum_{i=1}^n (ahN)^i}{1 + \sum_{i=1}^n (ahN)^i} \quad (4)$$

143 (Fig. 1). This is the multi-prey model for integer values of n . However, because the geometric
 144 series $\sum_{i=1}^n x^i = x(1 - x^n)/(1 - x)$ for $x \neq 1$, we can also write the model more generally for

145 arbitrary values of n as

$$146 \quad f(N) = \frac{aN(1 - (ahN)^n)}{1 - (ahN)^{n+1}} \quad (5)$$

147 to reflect predator populations capable of searching while handling a non-integer (e.g., average)
148 number of prey individuals.

149 We note that Sjöberg (1980) derived equivalent formulations in Michaelis-Menten enzyme-
150 kinematics form with parameters having correspondingly different statistical properties (Novak
151 & Stouffer, 2021a; Rohr *et al.*, 2022). We also note that despite the appearance of two sum-
152 mations in eqn. 4 and the unusual appearance of subtractions in eqn. 5 (see *Supplementary*
153 *Materials*), the model has only three parameters and thus has a parametric complexity no
154 greater than that of the Holling-Real Type III model and many others (see Table 1 of Novak
155 & Stouffer, 2021a). In fact, for subsequent model-fitting, we will combine the multi-prey and
156 Holling-Real models to a four-parameter generalization,

$$157 \quad f(N) = \frac{aN^\phi(1 - (ahN^\phi)^n)}{1 - (ahN^\phi)^{n+1}}, \quad (6)$$

158 which can be simplified to the other models when $\phi = 1$. Parameter ϕ (a.k.a. the Hill exponent)
159 can be interpreted as the number of prey encounters a predator must experience before its
160 feeding efficiency is maximized (Real, 1977).

161 **Relevance of the Type I response**

162 The conditions under which ~~Holling's the~~ linear, rectilinear, and Type II models can be good
163 descriptions of predator feeding rates are clarified by observing that the multi-prey response

164 simplifies to the Type II when $n = 1$ and approaches the rectilinear model as n increases
 165 (Fig. 1). Further, the linear Type I is obtained when $n = \infty$ (Fig. 1) because the infinite
 166 power series $\sum_{i=1}^{\infty} x^i = x/(1 - x)$ for $|x| < 1$. Incorporating this infinite power series into
 167 eqn. 3 shows that the expected proportion of predators handling prey at any given time will
 168 be ahN under the Type I. Importantly, this proportion differs from the expectation of zero
 169 that would be inferred to emerge by letting $h \rightarrow 0$ in the way the Type I is typically derived
 170 ~~(e.g., Rohr *et al.*, 2022)~~(e.g., Holling, 1965; Rohr *et al.*, 2022). In other words, the multi-prey
 171 model shows that handling times need not be inconsequential for the functional response to
 172 exhibit linear density dependence (Jeschke *et al.*, 2004). Rather, even the Type I can be a
 173 very good approximation of feeding rates when n is high and less than 100% of predators are
 174 handling prey (i.e. $ahN < 1$), which requires that prey abundances remain less than $1/ah$.
 175 (For comparison, note that under the Type II the quantity $1/ah$ reflects the prey abundance
 176 at which 50% of predators will be handling prey (i.e. the per predator feeding rate is at half
 177 its maximum of $1/h$) ~~;-it-which~~ is equivalent to the half-saturation constant of the Michaelis-
 178 Menten formulation. ~~)-Of futher note is that under the multi-prey model, $1/ah$ is also the~~
 179 prey abundance at which the proportions of predators handling 1,2,...,n prey are all equal
 180 (Fig. S.1).

181 Empirical support for multi-prey feeding

182 The multi-prey model shows that a spectrum of functional response forms can exist between the
 183 extremes of the Type I and Type II when handling and searching are not assumed to be mutually

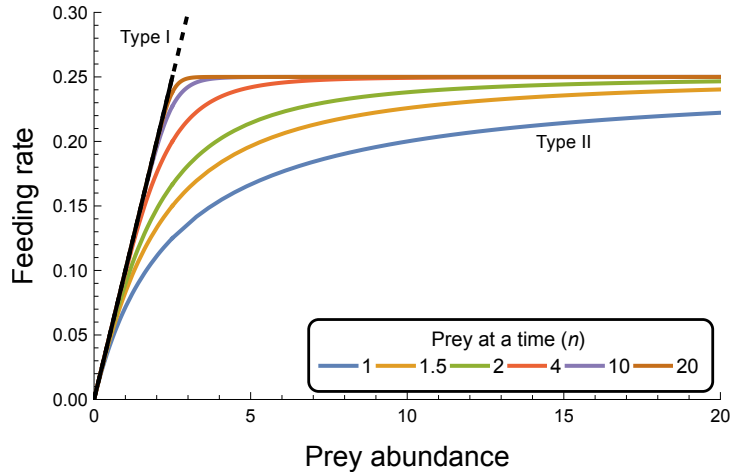


Figure 1: The potential forms of the multi-prey response. The multi-prey model diverges from the Type II (for which $n = 1$) and approaches the rectilinear model as the number n of prey individuals that a predator can handle while continuing to search increases. When $n = \infty$ it reduces to the linear Type I which can remain a biologically appropriate description of predator feeding rates so long as $ahN < 1$ (indicated by non-dashed region of the black line). *Parameter values*: attack rate $a = 0.1$ and handling time $h = 4$.

184 exclusive (Fig. 1). This motivated us to test two main hypotheses using the large number of
 185 empirical functional response studies that exist in the literature. The first hypothesis was that
 186 prior syntheses indicating the Type I response to be rare (Dunn & Hovel, 2020; Jeschke *et al.*,
 187 2004) were biased against the Type I despite its potential empirical appropriateness. That is,
 188 feeding rates may have had response shapes between the Type II and rectilinear model (close
 189 to the Type I for prey abundances $< 1/ah$) but were classified as Type II due to the lack of a
 190 sufficiently simple rectilinear-approaching model in prior analyses. The second hypothesis was
 191 due to Sjöberg (1980) who motivated parameter n by considering it to be a measure of food
 192 particle size relative to a zooplankter's gut capacity, with low n reflecting capacity for few large
 193 prey and high n reflecting capacity for many small prey. We thus expected predator-prey pairs
 194 with larger body-mass ratios to exhibit larger estimates of n when their functional responses

195 were assumed to follow the multi-prey model. For generality and to safeguard against potential
196 statistical model-comparison issues (see below), we included the Type I, II, III, multi-prey, and
197 the generalized (eqn. 6) model in our comparisons. We were thus also able to test an additional
198 hypothesis, due to Hassell *et al.* (1977), that larger body-mass ratios are associated with more
199 pronounced Type III responses (i.e. larger values of ϕ).

200 We used the FoRAGE database of published functional response datasets to assess these
201 hypotheses (Uiterwaal *et al.*, 2022). Our v4 update contains 3013 different datasets representing
202 1015 unique consumer-resource pairs (i.e. not just predator and prey species, though we continue
203 to refer to them as such for simplicity). For our analyses, we excluded datasets having a sample
204 size less than 15 observations as well as structured experimental studies that implemented less
205 than 4 different treatment levels of prey abundance (see the *Supplemental Materials* for addi-
206 tional details). Our model-fitting procedure followed the approach used by Stouffer & Novak
207 (2021) and Novak & Stouffer (2021b), assuming one of two statistical models for each dataset:
208 a Poisson likelihood for observational (field) studies and when eaten prey were replaced during
209 the course of the experiment, and a binomial likelihood when eaten prey were not replaced.
210 Experimental data available in the form of treatment-specific means and uncertainties were an-
211 alyzed by a parametric bootstrapping procedure in which new datasets were created assuming
212 either a treatment-specific Poisson or binomial process as dictated by the study’s replacement
213 of prey. In cases where measures of the uncertainty around non-zero means were not available,
214 we interpolated them based on the global log-log-linear relationship between means and stan-

215 dard errors across all datasets following Uiterwaal *et al.* (2018); for zero means, we interpolated
216 missing uncertainty values assuming a linear within-dataset relationship. Unlike in Stouffer &
217 Novak (2021) and Novak & Stouffer (2021b), we added a penalty to the likelihoods to discourage
218 exceptionally large estimates of n and ϕ (see the *Supplementary Materials*) and bootstrapped
219 data available in non-summarized form as well, using a non-parametric resampling procedure
220 that maintained within-treatment sample sizes for treatment-structured datasets. Both replace-
221 ment and non-replacement data were bootstrapped ~~50~~100 times which was enough to obtain
222 sufficient precision on the parameter point estimates.

223 **Frequency of multi-prey feeding**

224 We used the Bayesian Information Criterion (BIC) to test our first hypothesis, counting the
225 number of datasets whose bootstrapped mean BIC score supported a given model over the other
226 models by more than two units ($\Delta\text{BIC} > 2$). Our choice to use BIC was motivated both by
227 its purpose of selecting the generative model (rather than the best out-of-sample predictive
228 model, as per AIC) and by its generally stronger penalization of parametrically-complex models
229 (thereby favoring simpler models, relative to AIC). Conclusions regarding evidence in support of
230 the multi-prey model were thereby made more conservative, with our inclusion of models having
231 equal or greater parametric complexity helping to guard against an inappropriate reliance on
232 the asymptotic nature of BIC's consistency property.

233 The result of this first analysis was that, overall, ~~912~~(35925 (36%)) of all 2,591 datasets
234 provided support for functional response linearity (i.e. the Type I and multi-prey models), with

235 ~~990~~ 998 (38%) of all datasets providing support for multi-prey feeding more generally (i.e. the
236 Type I, multi-prey, and generalized eqn. 6 models). When considering only those datasets
237 that could differentiate among all five of the models, 7 (5.3%) of 132 replacement datasets and
238 ~~153~~ ~~(9.7~~ 143 (9.1%) of 1575 non-replacement datasets identified the multi-prey model (eqn. 5)
239 as the sole best-performing model (Fig. 2a-2b). An additional ~~36~~ ~~(27~~ 37 (28%) replacement
240 and ~~433~~ ~~(18~~ 451 (29%) non-replacement datasets identified the multi-prey model as performing
241 equivalently well to their best-ranked model(s). Although the Type I and the generalized model
242 were the least frequently sole-supported models, they were supported by datasets representing
243 all four of the most common predator taxonomic groups that constituted 90% of all datasets in
244 FoRAGE (insects, arachnids, crustaceans, and fishes; Fig. S.2).

245 **Effects of predator-prey body-mass ratio on n and ϕ**

246 To test the second and third hypotheses, we excluded datasets for which the Type I had alone
247 performed best and regressed the remaining datasets' bootstrapped median point estimates of n
248 and ϕ against their study's predator-prey body-mass ratio ($ppmr$), these having been compiled
249 in FoRAGE for most datasets. ~~(Datasets for which all other models performed better or equally~~
250 ~~well could be included because for them n and ϕ could equal 1.)~~ Although roughly 90% of these
251 datasets had estimates of $n \leq 8$ and $\phi \leq 2$ (Figs. S.3 and S.4), all three variables exhibited
252 substantial variation in magnitude. We therefore performed linear least-squares regression using
253 $\log_2(n)$ and $\log_2(\phi)$ versus $\log_{10}(ppmr)$.

254 Our analysis supported the hypothesis that predator-prey pairs with larger body-mass ratios

255 tend to exhibit larger estimates of n (Fig. 2c; $\log_2(n) = 0.55 + 0.15 \cdot \log_{10}(ppmr)$, $p < 0.01$, Table
256 S.1), but the predictive utility of this relationship was extremely poor (~~$R^2 = 0.03$~~). $R^2 = 0.02$.
257 We also found support for the hypothesis that larger body-mass ratios are associated with
258 larger values of ϕ , although the magnitude of this effect was weaker than it was for n (Fig. S.5;
259 ~~$\log_2(\phi) = 0.27 + 0.06 \cdot \log_{10}(ppmr)$~~) $\log_2(\phi) = 0.26 + 0.06 \cdot \log_{10}(ppmr)$, $p < 0.01$, Table S.2) and
260 was of similarly poor predictive utility ($R^2 = 0.02$).

261 To assess the sensitivity of our result for n to variation among datasets, we performed
262 additional regressions that restricted the considered datasets to (i) those having estimates of $n >$
263 1 (Fig. 2c, Table S.1), (ii) those with sample sizes exceeding the median sample size of all datasets
264 (Fig. S.6, Table S.3), and (iii) the four most common predator taxonomic groups (insects,
265 arachnids, crustaceans, and fishes), including for this last regression a two-way interaction term
266 between predator group identity and predator-prey body-mass ratio (Fig. 2d, Table S.4). These
267 analyses evidenced statistically clear, albeit predictively poor, positive relationships between n
268 and predator-prey body-mass ratios for all predators in general and for each predator group
269 individually as well.

270 Population-dynamic effects of multi-prey feeding

271 Given the empirical evidence that multi-prey feeding may indeed be common and a viable way to
272 describe functional responses, we next investigated its potential consequences for predator-prey
273 dynamics. Our goal was to understand how assuming either a Type I or Type II response could
274 lead to incorrect conclusions regarding these dynamics. We used the well-studied Rosenzweig &

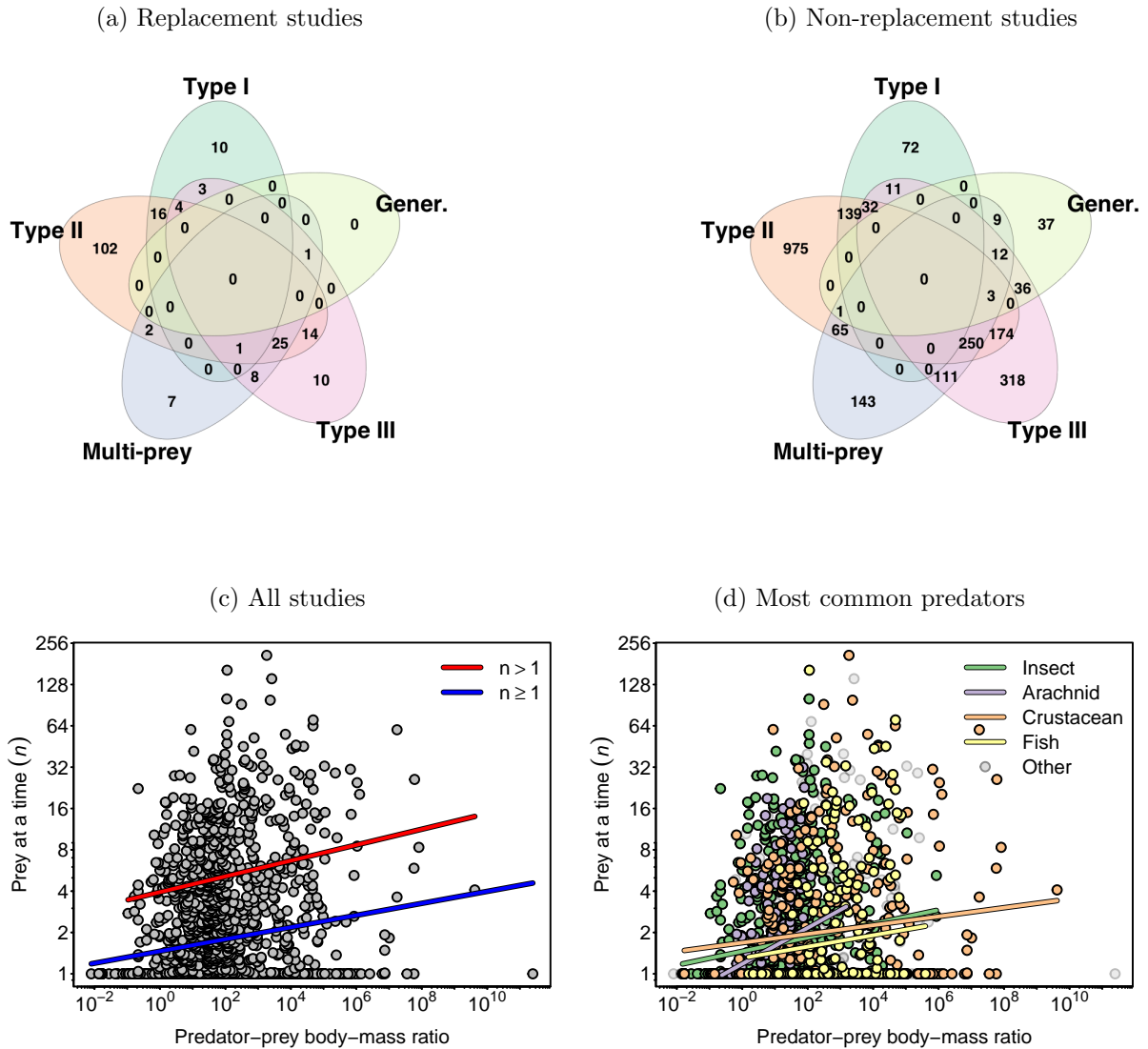


Figure 2: Empirical support for multi-prey feeding. Figs. 2a and 2b depict Venn diagrams categorizing the datasets of FoRAGE by their support for one or more of the five models as evaluated using a cut-off of 2 BIC units. Figs. 2c and 2d depict the observed relationship between estimates of n and the body-mass ratio of the studies' predator-prey pairs, excluding datasets for which the Type I model alone performed best. Regression lines in Fig. 2c reflect all considered datasets or only those with estimates of $n > 1$ (Table S.1). Regression lines in Fig. 2d reflect the identity of the four most common predator groups ($n \geq 1$, Table S.4).

275 MacArthur (1963) model to achieve this goal, employing both graphical (i.e. isocline) analysis
276 and simulations.

277 The model describes the growth rates of the prey N and predator P populations as

$$\frac{dN}{dt} = rN \left(1 - \frac{N}{K} \right) - f(N)P \quad (7a)$$

$$\frac{dP}{dt} = ef(N)P - mP, \quad (7b)$$

278 where r and K are the prey's intrinsic growth rate and carrying capacity, $f(N)$ is the functional
279 response, and e and m are the predator's conversion efficiency and mortality rate. Logistic
280 prey growth and Holling's Type II response have become the component parts of the canonical
281 Rosenzweig-MacArthur model for which enrichment in the form of an increasing carrying ca-
282 pacity causes the populations' dynamics to transition from a regime of monotonically-damped
283 stable coexistence to damped oscillations to sustained limit cycles (Rosenzweig, 1971). Other
284 prey growth and Type II-like functional response forms affect a similar destabilization sequence
285 (e.g., Freedman, 1976; May, 1972; Rosenzweig, 1971; Seo & Wolkowicz, 2018). The location
286 of the Hopf bifurcation between asymptotic stability and limit cycles is visually discerned in
287 the model's P vs. N phase plane (Fig. 3) as the point where the vertical N^* predator iso-
288 cline intersects the parabolic P^* prey isocline at its maximum, half-way between $-1/ah$ and
289 K (Rosenzweig, 1969; Rosenzweig & MacArthur, 1963). That is, the coexistence steady state
290 entails a globally-stable fixed point when the isoclines intersect to the right of the maximum
291 and entails a locally-unstable fixed point with a globally-stable limit cycle when they intersect
292 to the left of the maximum (Seo & Wolkowicz, 2018). Graphically, increasing K destabilizes

293 dynamics by stretching the prey isocline, moving its maximum to the right while the position of
 294 the vertical predator isocline remains unchanged. In contrast, when logistic growth and a Type
 295 I are assumed, the prey isocline is a linearly-decreasing function of prey abundance (Fig. 3) and
 296 predator-prey coexistence entails a globally-stable fixed point for all levels of enrichment.

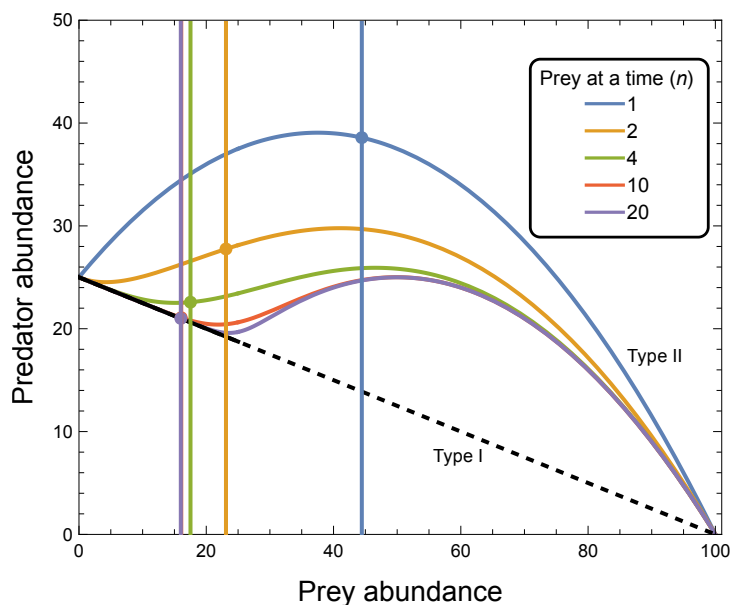


Figure 3: Predator and prey isoclines of the Rosenzweig-MacArthur model modified to include the multi-prey response correspond to those observed with the Type I and Type II responses when $n = \infty$ and $n = 1$ respectively. As the number n of prey that a predator can handling while searching increases, the prey abundance at which the predator’s growth rate is zero (i.e. the vertical predator isocline, N^*) decreases from its value under the Type II response ($m/a(e - mh)$) and converges rapidly on the value expected under the Type I response (m/ae). In contrast, predator abundances at which the prey’s growth rate is zero, P^* , converge on those expected under the Type I response only at low prey abundances to affect a second region of asymptotically stable dynamics; the “hump” does not flatten as it would if the handling time were presumed to be inconsequential (i.e. $h = 0$). Limit cycles occur when the predator and prey isoclines intersect on the left flank of the hump. With increasing n , the inflection point between the low-prey region of stability and limit cycles approaches the prey abundance where all predators are busy handling predators under the rectilinear model, $1/ah$ (indicated by non-dashed region of the black prey isocline). *Other parameter values:* attack rate $a = 0.02$, handling time $h = 2$, prey growth rate $r = 0.5$, prey carrying capacity $K = 100$, conversion efficiency $e = 0.25$, predator mortality rate $m = 0.08$.

297 **Graphical analysis**

298 For our analysis we insert the multi-prey response (eqn. 5) for $f(N)$ in eqn. 7. Solving $dP/dt = 0$
 299 for the N^* predator isocline then requires solving

$$300 \quad \frac{m}{e} = f(N^*) \implies N^* = \frac{m(1 - (ahN^*)^{n+1})}{ae(1 - (ahN^*)^n)}. \quad (8)$$

301 This leads to a solution for N^* that is independent of the predator's abundance (i.e. remains
 302 vertical in the P vs. N phase plane) but is unwieldy for $n > 2$ (see *Supplementary Materials*).
 303 Nonetheless, it represents a generalization of the predator isocline obtained for the Rosenzweig-
 304 MacArthur model with $n = 1$, $N^* = \frac{m}{a(e-mh)}$, and converges on $N^* = m/ae$ as $n \rightarrow \infty$ when
 305 $ahN^* < 1$, just as obtained assuming the Type I. In fact, N^* transitions smoothly from the
 306 former to the latter as n increases (Fig. 3) because eqn. 8 is a monotonically declining function
 307 of n for $ahN^* < 1$.

308 Solving $dN/dt = 0$ for the P^* prey isocline leads to the solution

$$309 \quad P^* = \frac{rN}{f(N)} \left(1 - \frac{N}{K}\right) = \frac{r(K - N)(1 - (ahN)^{n+1})}{aK(1 - (ahN)^n)}. \quad (9)$$

310 This too represents a generalization of the Rosenzweig-MacArthur model's prey isocline, $P^* =$
 311 $r(K - N)(1 + ahN)/aK$, which is itself a generalization of the isocline $P^* = r(K - N)/aK$
 312 obtained with the Type I as $n \rightarrow \infty$. Between these the prey isocline under the multi-prey
 313 response transitions from a parabolic dependence on the prey's abundance to having a second
 314 region within which it is a declining function of prey abundance (Fig. 3). This second region
 315 has a slope of $-r/aK$ at its origin regardless of n and is limited to low prey abundances of

316 $N < 1/ah$; as n increases, the region's upper extent approaches the prey abundance at which
317 all predators are busy handling prey under the rectilinear model. That is, for $1 < n < \infty$ the
318 "hump" shape of P^* does not flatten out as it does when one assumes handling times to become
319 negligible. Rather, similar to what can occur for the Type III response (Uszko *et al.*, 2015), the
320 prey isocline exhibits two regions of negative prey dependence (where $\frac{dP^*}{dN} < 0$) that flank an
321 intermediate region of positive prey dependence (where $\frac{dP^*}{dN} > 0$).

322 **Implications for coexistence and dynamics**

323 The emergence of a second prey abundance region where the slope of the prey isocline is neg-
324 ative means that a second asymptotically-stable coexistence equilibrium — one having a high
325 predator-to-prey abundance ratio — is possible should the two isoclines intersect within it. The
326 fact that this may occur is discerned by noting that N^* (eqn. 8) is independent of r and K ,
327 and that P^* (eqn. 9) is independent of m and e ; the positions of the two isoclines are thus
328 independent except via the functional response parameters a , h , and n . In fact, because N^*
329 decreases while the upper limit of the low prey abundance region of P^* increases towards $1/ah$
330 as n increases, it is readily possible — conditional on the values of the other parameters — to
331 observe a stable state at $n = 1$ to first transition to limit cycles and then return to fixed-point
332 stability as n alone is increased. This is illustrated by Fig. 4 in the context of enrichment for
333 values of K between approximately 75 and 115. Multi-prey feeding may thus be seen as another
334 mechanism contributing to stability at high productivity (Roy & Chattopadhyay, 2007). In-
335 deed, in addition to rescuing predators from deterministic extinction at low levels of enrichment

336 where a single-prey-at-a-time predator could not persist ($20 < K < 40$ in Fig. 4), sufficiently
 337 large values of n can preclude the occurrence of limit cycles altogether ($n > 9$ in Fig. 4).

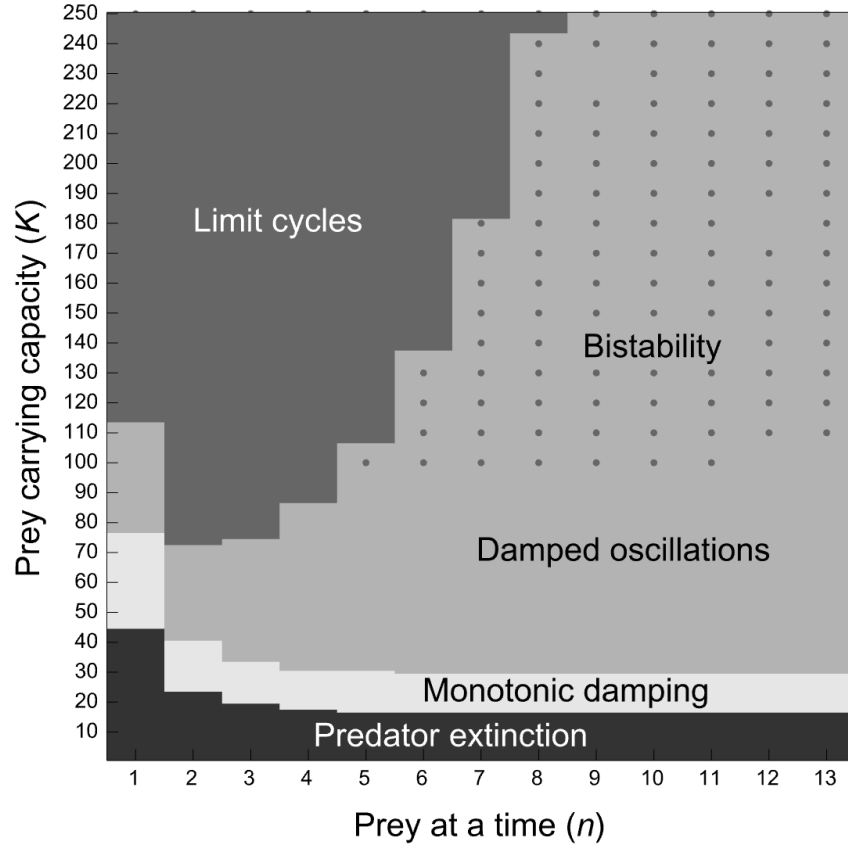


Figure 4: The destabilization with enrichment that is seen under the classic Rosenzweig-MacArthur model (where $n = 1$) is altered when predators can search for and handle multiple prey at a time ($n > 1$). At low prey carrying capacities ($K < 40$), multi-prey feeding rescues predators from deterministic extinction. At intermediate carrying capacities ($40 < K < 110$), low levels of multi-prey feeding destabilize dynamics by causing perturbation responses to transition from a transient regime of monotonic damping to one of damped oscillations or from damped oscillations to a persistent limit cycle regime. Further increases in multi-prey feeding can have a qualitatively stabilizing influence on dynamics, with sufficiently high n precluding a transition to limit cycles altogether so long as perturbations are sufficiently small. Large perturbations, on the other hand, will cause a transition to an alternative stable state consisting of limit cycle dynamics (see Fig. 5). *Other parameter values* as in Fig. 3.

338 Notably, however, the just-described high-predator low-prey steady state is only a locally

339 stable fixed point and coexists with a stable limit cycle that surrounds it (Figs. 4 and 5). The
340 high-predator low-prey state thus exhibits bi-stability. The consequences of this bi-stability are
341 that predator-prey interactions with multi-prey feeding are destined to exhibit (i) transitions to
342 persistent limit cycles when subjected to large perturbations that send abundances beyond the
343 domain of attraction of the fixed-point steady state (Fig. 5*a,c*), and (ii) transient dynamics that
344 are prone to damped oscillations (rather than monotonic damping) in response to small per-
345 turbations within the domain of attraction. These transient oscillations occur for substantially
346 lower levels of enrichment than is the case for single prey-at-a-time predators (Fig. 4). Moreover,
347 their temporal duration can be exceedingly long (Fig. 5*b*) because the limit cycle acts akin to a
348 crawl-by attractor (Hastings *et al.*, 2018) that impinges upon the steady state’s local resilience.
349 Thus, when subjected to continual perturbations in an explicitly stochastic setting (Barraquand
350 *et al.*, 2017), the system can readily transition between the stable fixed-point attractor and the
351 stable limit cycle attractor that surrounds it (Fig. 6), resulting in dynamical epochs of irregular
352 duration that are characteristic of many empirical time-series (Blasius *et al.*, 2020; Rubin *et al.*,
353 2023). Therefore, multi-prey feeding does not provide a robust mechanism against instability
354 at high productivity but rather leads to a richer range of population dynamics and coexistence
355 states than can result from Type I, II, or III responses alone.

356 Discussion

357 Our study was motivated by the apparent disconnect that exists between the way that many
358 empirically-minded ecologists perceive ~~the Type I model~~ functional response linearity and the

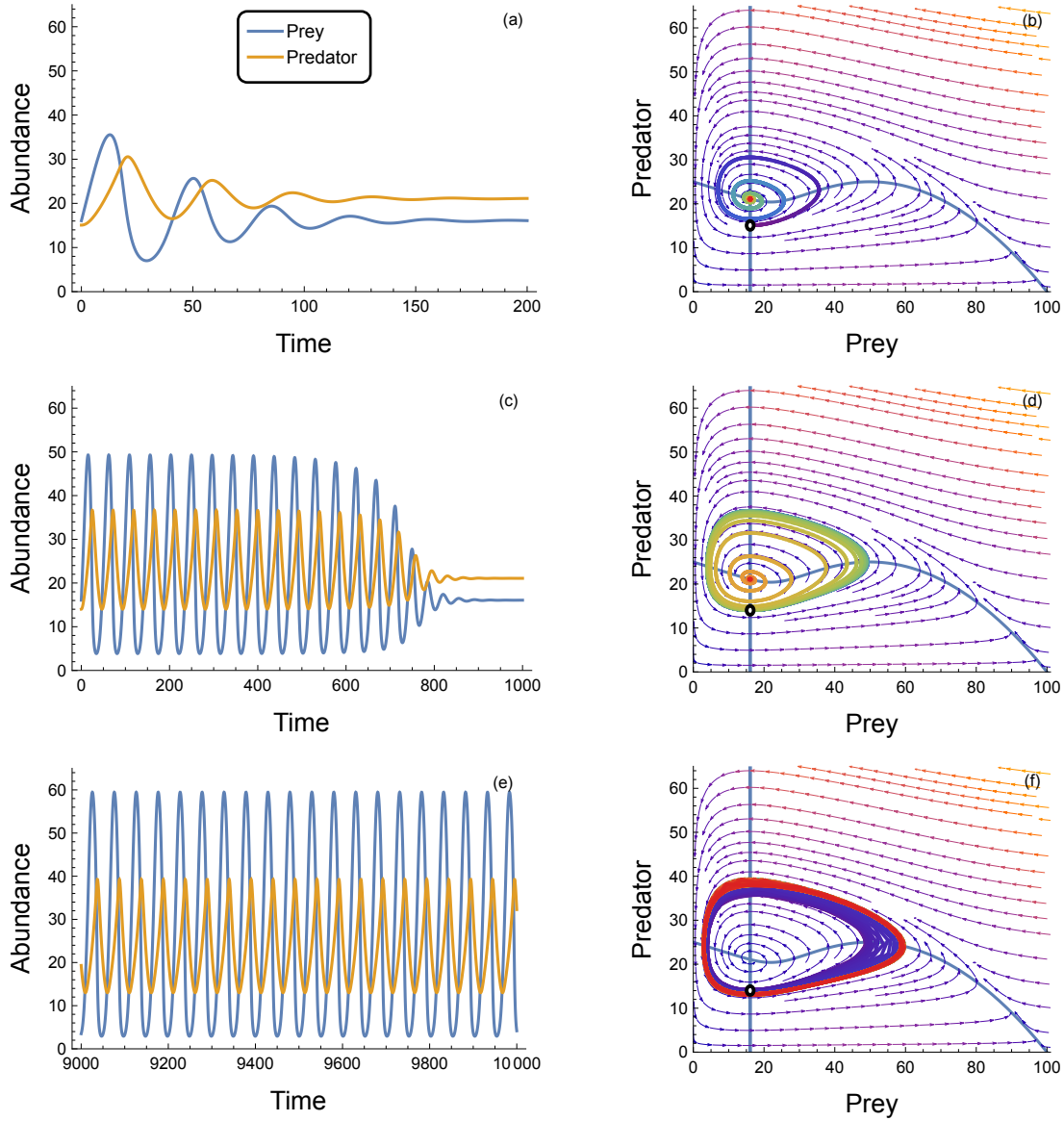


Figure 5: Because of the system's bi-stability at high predator-to-prey abundance ratios, even small differences in the size of a perturbation to the steady state can affect a large change in the duration of the system's transient response (compare panels *a* and *b* with *c* and *d*) and can even cause the system to become entrained in a stable limit cycle (illustrated in panels *e* and *f*). The only difference between each of the above panel rows is that the predator's initial population size $P(0)$ is perturbed away from its P^* steady state as: (*a*, *b*) $P(0) = P^* - 6$; (*c*, *d*) $P(0) = P^* - 7.0645$; and (*e*, *f*) $P(0) = P^* - 7.065$. For all cases $N(0) = N^*$. Parameter values as Fig. 3 with $n = 10$.

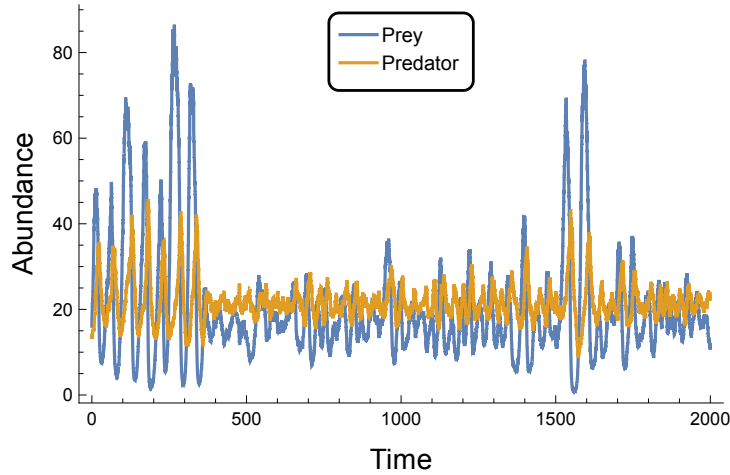


Figure 6: When subjected to continually-occurring stochastic perturbations, the high-predator low-prey coexistence state can exhibit time periods during which its dynamics are influenced primarily by the stable fixed-point attractor and time periods during which dynamics are primarily influenced by the alternative stable limit cycle attractor, switching between these on an irregular basis. Simulation implemented using an Itô ~~integral process~~ [stochastic differential equations](#) as $dN = rN(1 - N/K) - f(N)P dt + \sigma NdW$ and $dP = ef(N) - mP dt - \sigma PdW$, with $f(N)$ as in eqn. 5 and Gaussian white environmental noise $dW(t)$ of volatility $\sigma = 0.04$ (*cf.* Barraquand, 2023). *Other parameter values* and initial population sizes as in Fig. 5c-d.

359 way that many modelers and theory-minded ecologists justify its use in their representations
360 of consumer-resource interactions. While the former are prone to dismiss the Type I as being
361 overly simplistic and hence unsuitable for describing predator feeding rates, the latter are prone
362 to rely on and justify its sufficiency for the sake of computational ease and analytically-tractable
363 insight. Since the potential for predators to feed on multiple prey at a time (i.e. the non-
364 exclusivity of handling and searching activities) has been little considered by either group, we
365 set out to address three aspects of this disconnect: (i) deriving a multiple-prey-at-a-time model
366 that mechanistically connects the linear and rectilinear models to the more empirically palatable
367 Type II model, (ii) assessing the extent to which published datasets provide support for multi-

368 prey feeding, and (iii) investigating how multi-prey feeding and the linear density dependence it
369 can impose on feeding rates can alter our understanding of predator-prey coexistence. Because
370 they bear insight with which to elaborate on the circumstances under which linearity may be
371 empirically relevant, we structure the discussion of our work by considering the latter two aspects
372 first.

373 **Empirical support**

374 Our statistical analysis of the datasets compiled in FoRAGE demonstrates that both the Type
375 I and multi-prey models are viable descriptions (*sensu* Skalski & Gilliam, 2001) of the feeding
376 rates that predators have exhibited in many single-prey experiments (Figs. 2a-2b). This result
377 is consistent with handling and searching being non-exclusive activities for a substantial number
378 of predator-prey pairs. Although ~~this~~ our result contrasts with the prior syntheses of Jeschke
379 *et al.* (2004) and Dunn & Hovel (2020), these (*i*) did not consider models capable of response
380 forms in between the strictly linear Type I and Type II forms and (*ii*) either relied on the
381 conclusions reached by each studies' original authors (who used varied model-fitting and com-
382 parison approaches) or visually assessed functional response forms from plotted data. One might
383 argue that many of the datasets providing sole support to the Type I in our analysis came from
384 experiments using prey abundances that were insufficient to elicit saturation (see also Coblenz
385 *et al.*, 2023), but the point can be made that, from an information-theoretic perspective, the
386 Type I performed best across the range of prey abundances that the original authors consid-
387 ered empirically reasonable (and logistically feasible). The even greater number of datasets that

388 provided sole support to the multi-prey model, along with the result that many of the point
389 estimates for parameter n (the maximum number of prey eaten at a time) were sufficiently large
390 to affect a response approaching a rectilinear response (~~cf.~~ Figs. 1 and 2c), indicates that feeding
391 rates exhibited a ~~significant~~-region of linearity for many predator-prey interactions having long
392 handling times as well. Moreover, the statistically-clear positive relationships we observed in our
393 subsequent regression analyses of n and predator-prey body-mass ratios (Figs. 2c-2d) ~~confirm~~
394 support Sjöberg's hypothesis regarding a proximate reason for this linearity; ~~it~~ it being more
395 likely to occur for larger predators feeding on small prey because handling is less preclusive of
396 searching.

397 Unfortunately, the amount of variation in n that was explained by body-mass ratio alone was
398 extremely low, making the relationship of little predictive utility relative to several other body-
399 mass relationships (e.g., Brose *et al.*, 2006; Coblenz *et al.*, 2023; Hatton *et al.*, 2015; Rall *et al.*,
400 2012). That said, the relationship's low explanatory power is not unsurprising given that none
401 of the experiments in FoRAGE was designed with the multi-prey model in mind. In particular,
402 and although most estimates of n were of a seemingly reasonable magnitude (Fig. S.3), we
403 caution against giving too much credence to the very large-valued estimates we observed. This
404 is for two primary reasons. First, given that a given dataset's ability to distinguish between
405 possible values of n diminishes rapidly as n increases (Fig. 1), datasets exhibiting saturation at
406 high prey abundances but having few or no observations near the inflection point of $1/ah$ will
407 have been sensitive to issues of parameter identifiability. Low identifiability will have caused an

408 inflation of estimates despite our effort to guard against it by removing datasets with fewer than
409 4 prey abundance levels. Second, given that initiating experiments with predator individuals
410 having empty guts is a common protocol (Griffen, 2021; Li *et al.*, 2018), many experiments
411 will have strictly violated the assumption of predator behavior being at steady state. This will
412 also have inflated estimates of n by causing transient rates of prey ingestion to exceed rates
413 of handling completion (i.e. $aN > 1/h$) to affect faster-than-steady-state feeding, especially at
414 prey abundances below $1/ah$. We therefore suggest that the very large estimates of n observed
415 in our analyses be better interpreted as qualitative (rather than quantitative) support for the
416 non-exclusivity of searching and handling and encourage future experiments and analyses with
417 additional covariate predictors to better understand the biological sources of variation in n .
418 (Similar issues pertain to the estimation and interpretation of ϕ .)

419 **Mechanistic approximations**

420 The multi-prey model may be considered a mechanistic model in that its derivation and each
421 of its parameters has at least one biologically-specific interpretation. However, it is also rather
422 phenomenological in that it encodes only an essence of the biologically possible non-exclusivity of
423 searching and handling processes. For example, the model's derivation assumes that the attack
424 rate and handling time remain constant and independent of the number of prey that predators
425 are already handling (below the maximum number n). Although this assumption may result in
426 a very good approximation to feeding rates, it is unlikely to reflect biological reality particularly
427 as the number of prey being handled by a given predator approaches n . In such circumstances

428 either or both searching and handling process rates are likely to become dependent on the feeding
429 rate and thereby on prey abundance (see also Okuyama, 2010; Stouffer & Novak, 2021).

430 Functional responses where such dependence is important may be better and more mech-
431 anistically described by more flexible models (see also Novak & Stouffer, 2021a). Prominent
432 among these is the extended Steady State Saturation model (SSS¹) of Jeschke *et al.* (2004) in
433 which handling and digestion are explicitly distinguished (see *Supplementary Materials*). In
434 this four-parameter model, searching and handling are mutually exclusive, but searching and
435 digestion are not because the predator’s search effort depends on its gut fullness (i.e. hunger
436 level) and is thus dictated by the digestion rate. A phenomenological shape parameter controls
437 the non-linearity of the search-effort hunger-level relationship. For high values of this shape pa-
438 rameter (reflecting predators that search at their maximum rate even when their guts are quite
439 full) and inconsequential handling times, the model approaches the rectilinear model, just like
440 the multi-prey model at high n , while for consequential handling times it retains a saturating
441 curvature at low prey abundances (see Figs. A1 and A2 of Jeschke *et al.*, 2004).

442 **Population-dynamic effects**

443 The population-dynamic consequences of the extended SSS model remain unstudied, but our
444 analysis of the simpler multi-prey model reveals the relevance of it and other models for un-
445 derstanding how the linearity of multi-prey feeding can impact predator-prey dynamics. These
446 other models are the arctangent and hyperbolic tangent models because for these it has been

¹We would be remiss not to point out that all functional response models of which we are aware assume steady state conditions at the behavioral foraging scale. The SSS model’s name does not, therefore, reflect a limitation that is unique to it.

447 more rigorously shown that two limit cycles — one stable and the other unstable — can co-occur
448 with a locally-stable fixed point at low prey abundances (Seo & Kot, 2008; Seo & Wolkowicz,
449 2015; 2018), just as we observed for the multi-prey model (see also Freedman, 1980). The key
450 feature common to all three models is that they affect a prey isocline that *decreases* from a
451 *finite*-valued origin at zero prey abundance. This differs from the Type II and other functional
452 responses that ~~exhibit saturating curvature~~ are concave and increasing with prey density at low
453 prey abundance. For these the prey isocline *increases* from a finite-valued origin, the low-prey
454 fixed point is unstable, and only the stable limit cycle is thus of relevance under logistic prey
455 growth. It also differs from functional responses that accelerate at low prey abundances (e.g., the
456 Type III) and from consumer-resource models more generally in which, for example, prey have
457 a physical refuge, exhibit sublinear density-dependence, or experience density-independent im-
458 migration. For these the prey isocline decreases from an origin that approaches infinity and the
459 low prey steady state is a stable fixed point around which limit cycles do not occur (e.g., Case,
460 2000; Uszko *et al.*, 2015). We surmise that the linearity brought about by the non-exclusivity of
461 searching and handling in the multi-prey model is (i) replicated by the more phenomenological
462 arctangent and hyperbolic tangent models, and that (ii) it is the cause of the greater range of
463 dynamical outcomes that these functional responses affect as compared to responses exhibiting
464 nonlinearity at low prey abundances.

465 ~~The broader implication of our analysis is that population-dynamic theory that relies on the~~
466 ~~Type I may not be as globally relevant from a biological perspective as its mathematics would~~

467 suggest. In particular, it shows that the stabilization which the Type I contributes to dynamics
468 is dependent on perturbation magnitude. More specifically, the relevance of theory that relies
469 on the Type I is limited to perturbations that are small enough to preclude the influence of the
470 attracting stable limit cycle that will exist when the functional response is described as having
471 a potentially unobserved maximum feeding rate.

472 Our consideration of enrichment effects illustrates a specific example of this. When the
473 functional response is assumed to be Type I, the fixed point is globally stable and perturbations
474 to it decay monotonically. In contrast, when the functional response is linear only at low
475 prey abundances, as when multi-prey feeding occurs, the fixed point is only locally stable and
476 perturbations can elicit cycles that may persist for many generations or even indefinitely. In
477 fact, as indicated by Rubin *et al.* (2023) in their analysis of a stochastic implementation of
478 the Rosenzweig-MacArthur model, the dynamics will additionally be influenced by the crawl-by
479 inducing origin (dual extinction) and prey-only (carrying capacity) steady states that will extend
480 the lifetime of long-term transients even further. This influence, too, will not be observed when
481 a Type I is assumed because these unstable steady states will rarely if ever be approached.

482 **Relevance revisited**

483 As discussed above (see *Relevance of Type I response*), the multi-prey model shows that handling
484 times need not be inconsequential to observe linear prey dependence when the number of prey
485 that a predator individual can handle at a time is relatively high and the maximum proportion of
486 individuals in a predator population that are simultaneously handling prey remains sufficiently

487 low. This is not to say that other factors and processes cannot cause functional responses to be
488 very nonlinear, but within the confines of our work’s assumptions the latter condition can be
489 satisfied as long as prey abundances remain less than $1/ah$.

490 Our statistical and mathematical analyses add insight into when the conditions for linearity
491 are more likely to be met. Specifically, functional responses are more likely to exhibit linearity
492 when predator-to-prey body-mass ratios are high (Fig. 2c), when predator-to-prey abundance
493 ratios are high (Fig. 3), and thus, we predict, in top-heavy systems with high predator-to-prey
494 biomass ratios. Top-heavy interactions and food webs more generally occur in all ecosystem
495 types (McCauley *et al.*, 2018), but are more likely for ectothermic and invertebrate consumers,
496 in aquatic habitats, among higher trophic levels, and in ecosystems of low total biomass (Brose
497 *et al.*, 2006; Hatton *et al.*, 2015; Perkins *et al.*, 2022). The development of methods for gaug-
498 ing the nonlinearity of functional responses in diverse field settings (e.g., Novak *et al.*, 2017;
499 Uiterwaal & DeLong, 2024) will be useful for directly testing our prediction that these same
500 systems should also exhibit more linear functional responses. New methods that make use of
501 the greater information content associated with counts of the numbers of prey being handled
502 (Fig. S.1) should be particularly useful.

503 Importantly, our work also shows that predator-prey dynamics need not be destabilized
504 by food web top-heaviness. Rather, paralleling theory assuming Type III responses (Kalinkat
505 *et al.*, 2013; Uszko *et al.*, 2015), increases in top-heaviness can lead to greater food web stability
506 — be it stable coexistence potential or perturbation resilience (Fig. S.7) — when multi-prey

507 feeding occurs, provided that perturbations are small enough for population abundances to
508 remain well within the local attractor of the stable fixed point (Fig. 5). This contrasts with
509 existing theory on top-heavy food webs that has largely assumed Type II responses (McCauley
510 *et al.*, 2018). Indeed, our analyses show that even small departures from mutual exclusivity
511 can lead to qualitatively different coexistence states and dynamics than predicted by existing
512 theory, including the possibility of long-term transients and the just-mentioned bi-stability of
513 fixed-point and limit-cycle dynamics. Food web models that incorporate multi-prey feeding
514 and how its prevalence may change with species- and system-level attributes will be useful for
515 understanding just how much multi-prey feeding must occur within food webs as a whole to alter
516 ~~their~~ community structure and dynamics. A first step towards such food web models will be
517 to extend the multi-prey model to multi-species formulations appropriate for generalist rather
518 than single-prey-species predators.

519 **Conclusions for bridging theory and empirical insight**

520 Natural history observations show that diverse types of predators are capable of (literally) han-
521 dling and searching for prey simultaneously: sea otters capture several snails on a dive; crabs
522 process mussels with their mouthparts while picking up more with their claws; spiders capture
523 insects in their webs while processing others for later ingestion. Many more situations relevant
524 to multi-prey feeding become apparent and potentially relevant to the context of functional re-
525 sponses when it is recognized that the “handling time” parameter of most models represents not
526 just the literal manipulation of prey (e.g., that may be seen by an observer of the interaction)

527 but rather reflects the feeding process that limits a predator's maximum feeding rate, including
528 possible limits to stomach fullness and digestion (DeLong, 2021; Jeschke *et al.*, 2002; 2004).
529 Sculpin fishes, for example, have been observed with over 300 identifiable mayflies in their stom-
530 achs (Preston *et al.*, 2018), the majority of which could not have been captured simultaneously
531 and for which literal handling must therefore have been inconsequential relative to digestion.

532 The degree to which searching and (general) handling actually represent mutually exclu-
533 sive activities, and the degree to which each of the many processes potentially encapsulated
534 by a handling time parameter measurably contributes to a predator's functional response, is
535 nonetheless poorly discerned from observation alone. Knowing that handling times are short or
536 long, or that searching and literal handling do or do not overlap, is neither sufficient to dismiss
537 or assume a given functional response model on *a priori* grounds. This is because all models
538 are phenomenological approximations of biological process at some level. This applies as much
539 to predator-prey interactions studied in controlled experiments as it does to those studied in
540 natural settings, and is particularly true in the context of building understanding and theory
541 when extrapolating the former to the latter across Ecology's wide-ranging scales.

542 ~~We thus~~ In this context we draw two overarching conclusions ~~:-from our analyses:~~ that
543 functional response linearity should not be dismissed by empiricists as an irrelevant description of
544 predator feeding rates, and that modelers and theoreticians should be more cautious in reaching
545 empirical conclusions of system dynamics when presuming the linear Type I response to be
546 appropriate. ~~In even broader terms, our research demonstrates how disagreements between~~

547 ~~different perspectives can be addressed by identifying and easing the fundamental assumptions~~
548 ~~that underpin them, and how improved communication between empiricists and theoreticians~~
549 ~~will benefit Ecology as a whole (Grainger *et al.*, 2022).~~

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Supplementary Materials for
~~Feeding on multiple prey at a time:~~In defense of the original
Type I functional response:
The frequency and population-dynamic effects of ~~functional~~
~~response linearity~~feeding on multiple prey at a time

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Multi-prey functional response model

Derivations

More explicit derivations of the Type II and multi-prey models are as follows.

Holling Type II model

Assuming a predator population P of fixed size that is composed of only P_S searching and P_H handling sub-populations, let the rate of change in abundance of the two sub-populations be described by

$$\frac{dP_S}{dt} = -aN P_S + \frac{1}{h} P_H \quad (\text{S.1a})$$

$$\frac{dP_H}{dt} = aN P_S - \frac{1}{h} P_H. \quad (\text{S.1b})$$

Correspondingly, the rate at which eaten prey N_e are generated is

$$\frac{dN_e}{dt} = \frac{1}{h} P_H. \quad (\text{S.2})$$

As in the main text, a is the per capita attack rate, h the handling time, and N the prey's abundance (which is also assumed fixed at the behavioral time scale we are considering).

Setting $\frac{dP_H}{dt} = 0$ (i.e. assuming steady state conditions), we substitute $(P - P_H)$ for P_S and rearrange to determine the proportion of the whole population that is busy handling:

$$aN(P - P_H) = \frac{1}{h} P_H \quad (\text{S.3a})$$

$$\implies aNP = aNP_H + \frac{1}{h} P_H \quad (\text{S.3b})$$

$$= (aN + \frac{1}{h}) P_H \quad (\text{S.3c})$$

$$\implies \frac{P_H}{P} = \frac{aN}{aN + \frac{1}{h}} \quad (\text{S.3d})$$

$$= \frac{ahN}{1 + ahN}. \quad (\text{S.3e})$$

The total number of handling predators is thus

$$P_H = \frac{ahNP}{1 + ahN}. \quad (\text{S.4})$$

Since the rate at which each these P_H predators finishes handling its prey is $\frac{1}{h}$, it follows that the rate at which eaten prey are “generated” by the whole predator population is

$$\frac{dN_e}{dt} = \frac{1}{h} P_H = \frac{aN P}{1 + ahN} \quad (\text{S.5})$$

and thus that the *per predator* feeding rate (the functional response) is

$$f(N) = \frac{1}{P} \frac{dN_e}{dt} = \frac{1}{h} \frac{P_H}{P} = \frac{aN}{1 + ahN}. \quad (\text{S.6})$$

Multi-prey model

Again assume a predator population P of fixed size that is composed of P_S searching and handling sub-populations, but now split handling predators into those capable of searching while handling less than n prey individuals at any moment time. We therefore have that

$$P = P_S + P_{H_1} + P_{H_2} + \dots + P_{H_n} \quad (\text{S.7})$$

and describe the rate of change for each sub-populations by

$$\frac{dP_S}{dt} = -aN P_S + \frac{1}{h} P_{H_1} \quad (\text{S.8a})$$

$$\frac{dP_{H_1}}{dt} = aN P_S - \frac{1}{h} P_{H_1} \quad (\text{S.8b})$$

$$\frac{dP_{H_2}}{dt} = aN P_{H_1} - \frac{1}{h} P_{H_2} \quad (\text{S.8c})$$

\vdots

$$\frac{dP_{H_n}}{dt} = aN P_{H_{(n-1)}} - \frac{1}{h} P_{H_n} . \quad (\text{S.8d})$$

Correspondingly, the rate at which eaten prey N_e are generated is now

$$\frac{dN_e}{dt} = \frac{1}{h} \sum_{i=1}^n P_{H_i} . \quad (\text{S.9})$$

By setting $\frac{dP_{H_i}}{dt} = 0$ for all sub-populations, rearranging, and iteratively substituting, we have that

$$aN P_S = \frac{1}{h} P_{H_1} \implies P_{H_1} = ahN P_S \quad (\text{S.10a})$$

$$aN P_{H_1} = \frac{1}{h} P_{H_2} \implies P_{H_2} = ahN P_{H_1} \quad (\text{S.10b})$$

$$= ahN (ahN P_S) \quad (\text{S.10c})$$

$$= (ahN)^2 P_S \quad (\text{S.10d})$$

$$aN P_{H_2} = \frac{1}{h} P_{H_3} \implies P_{H_3} = ahN P_{H_2} \quad (\text{S.10e})$$

$$= ahN ((ahN)^2 P_S) \quad (\text{S.10f})$$

$$= (ahN)^3 P_S \quad (\text{S.10g})$$

\vdots

$$aN P_{H_{(n-1)}} = \frac{1}{h} P_{H_n} \implies P_{H_n} = ahN P_{H_{(n-1)}} \quad (\text{S.10h})$$

$$= ahN ((ahN)^{n-1} P_S) \quad (\text{S.10i})$$

$$= (ahN)^n P_S , \quad (\text{S.10j})$$

with the last lines for P_{H_n} inferred by induction. The proportional abundance of each i th sub-population is thus

$$\frac{P_{H_i}}{P} = \frac{(ahN)^i P_S}{P} \quad (\text{S.11a})$$

$$= \frac{(ahN)^i P_S}{P_S + P_{H_1} + P_{H_2} + \dots + P_{H_n}} \quad (\text{S.11b})$$

$$= \frac{(ahN)^i P_S}{P_S + ahNP_S + \dots + (ahN)^n P_S} \quad (\text{S.11c})$$

$$= \frac{(ahN)^i}{1 + ahN + \dots + (ahN)^n} \quad (\text{S.11d})$$

$$= \frac{(ahN)^i}{1 + \sum_{k=1}^n (ahN)^k}. \quad (\text{S.11e})$$

Each of the sub-populations generates eaten prey at rate $\frac{1}{h}$, thus the rate at which eaten prey are generated by the whole population is

$$\frac{dN_e}{dt} = \frac{1}{h} \sum_{i=1}^n P_{H_i} \quad (\text{S.12a})$$

$$= \frac{1}{h} \sum_{i=1}^n \frac{P_{H_i}}{P} P \quad (\text{S.12b})$$

$$= \frac{1}{h} \sum_{i=1}^n \frac{(ahN)^i}{1 + \sum_{k=1}^n (ahN)^k} P \quad (\text{S.12c})$$

$$= \frac{\frac{1}{h} \sum_{i=1}^n (ahN)^i}{1 + \sum_{i=1}^n (ahN)^i} P. \quad (\text{S.12d})$$

The *per predator* feeding rate is therefore

$$f(N) = \frac{1}{P} \frac{dN_e}{dt} = \frac{\frac{1}{h} \sum_{i=1}^n (ahN)^i}{1 + \sum_{i=1}^n (ahN)^i} \quad (\text{S.13})$$

as given in eqn. 4 of the main text.

Proportion of predators feeding on 1 to n prey

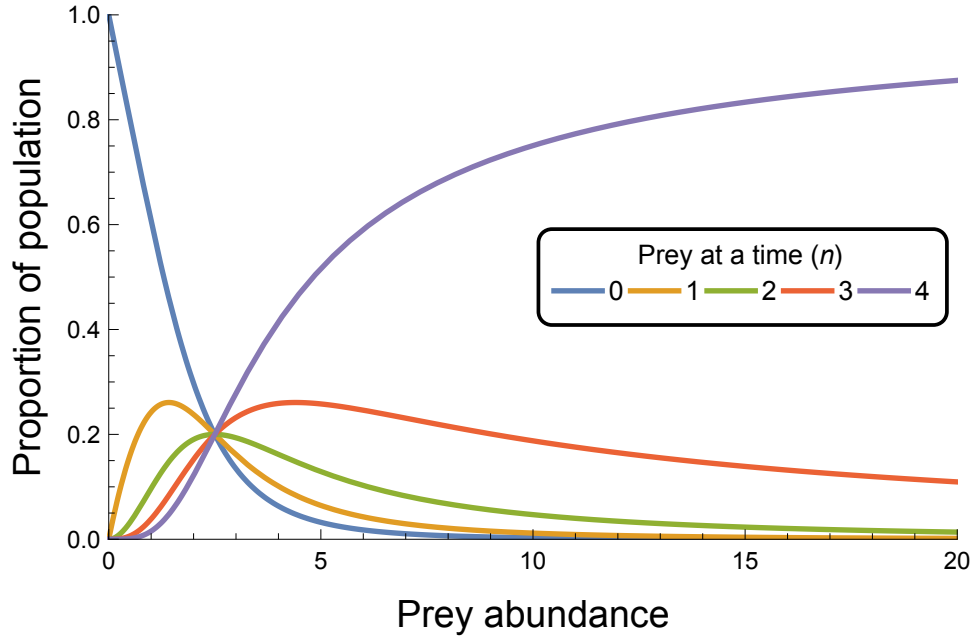


Figure S.1: The expected proportions of predator individuals that will be observed not feeding or handling $i = 1, 2, 3$ or 4 prey changes with prey abundance (here visualized for a predator population whose individuals can handle up to $n = 4$ prey at a time). Individuals from each of the handling groups consumes prey at rate $1/h$, thus the predator population's (i.e. the average individual's) functional response is the product of $1/h$ and the sum of these handling-predator proportions. The prey abundance at which the expected proportions of individuals handling $0, 1, 2, 3$ or 4 prey are all equal occurs at prey abundance $1/ah$. *Parameter values:* the attack rate is $a = 0.1$ and the handling time is $h = 4$.

Equivalence of eqns. 4 and 5 for integer values of n

Letting $n = 1$, we have

$$\begin{aligned} f(N) &= \frac{aN(1 - (ahN)^n)}{1 - (ahN)^{n+1}} = \frac{aN(1 - (ahN))}{1 - (ahN)^2} = \frac{aN(1 - ahN)}{1^2 - (ahN)^2} \\ &= \frac{aN(1 - ahN)}{(1 + ahN)(1 - ahN)} \\ &= \frac{aN}{1 + ahN}. \end{aligned}$$

Letting $n = 2$, we have

$$\begin{aligned} f(N) &= \frac{aN(1 - (ahN)^n)}{1 - (ahN)^{n+1}} = \frac{aN(1 - (ahN)^2)}{1 - (ahN)^3} = \frac{aN(1 + ahN)(1 - ahN)}{(1 + ahN + (ahN)^2)(1 - ahN)} \\ &= \frac{aN(1 + ahN)}{1 + ahN + (ahN)^2} \\ &= \frac{\frac{1}{h} \sum_{i=1}^2 (ahN)^i}{1 + \sum_{i=1}^2 (ahN)^i}. \end{aligned}$$

Letting $n = 3$, we have

$$\begin{aligned} f(N) &= \frac{aN(1 - (ahN)^n)}{1 - (ahN)^{n+1}} = \frac{aN(1 - (ahN)^3)}{1 - (ahN)^4} = \frac{aN(1 + ahN + (ahN)^2)(1 - ahN)}{(1 + ahN + (ahN)^2 + (ahN)^3)(1 - ahN)} \\ &= \frac{aN(1 + ahN + (ahN)^2)}{1 + ahN + (ahN)^2 + (ahN)^3} \\ &= \frac{\frac{1}{h} \sum_{i=1}^3 (ahN)^i}{1 + \sum_{i=1}^3 (ahN)^i}. \end{aligned}$$

Their equivalence for higher integer values of n follows similarly.

Analysis of FoRAGE datasets

Data exclusions and re-scaling

The most recent version of FoRAGE (v.4) contains a total of 3013 datasets from which we excluded 422 for our analyses. Most of these were excluded because they entailed less than 4 prey-abundance treatment levels or because they had fewer than 15 data points (i.e. replicates) overall, but we also excluded several datasets because they provided prey abundances as densities for treatments that were implemented using arenas of varying size without specifying what those arena sizes were; entailed feeding rates of a variable but unspecified number of predators known to exhibit predator-dependent feeding rates; and/or entailed feeding rates of variable but unspecified experimental duration. Nine datasets were excluded because our models failed to reach convergence.

Our analyses required integer counts of prey abundance and eaten prey because we assumed binomial and Poisson likelihood functions to accommodate the increasing variance that accompanies an increase in the expected number of eaten prey (Novak & Stouffer, 2021b). For most datasets in which prey abundances were expressed as prey densities and/or predation was expressed as feeding rates, integer counts of prey abundance and prey eaten could be calculated using provided information on the area size(s) used (area or volume), the number of predators per treatment, and experimental duration(s). For raw-data datasets where this information was not provided, as well as datasets expressing densities and feeding rates on a mass basis (e.g., micro-grams of prey available or eaten), we (i) multiplied prey densities by the minimum scalar value necessary to obtain integer values across all prey densities (which we then used as prey abundance counts), and (ii) multiplied prey feeding rates by the minimum scalar value necessary to obtain integer values across all feeding rates (which we then used as counts of prey eaten). We multiplied prey abundances by an additional minimum scalar value for non-replacement datasets (reported as raw-data or as means) where the units in which densities and feeding rates were measured caused there to be more prey eaten than were seemingly available. Although these procedures will have altered the interpretation of the attack rate and handling time parameters (i.e. our estimates of a and h are not comparable across datasets), neither procedure will have affected our estimates of n for the multi-prey model (because it is dimensionless) except, potentially, through an influence on the variance of the likelihood models (larger counts of prey eaten being permitted a higher variance than low counts of prey eaten). Although we did not observe any relationship between estimates of n and the magnitudes of re-scaling across our datasets, its potential influence is worthy of future analytical study.

Penalized likelihood

Many datasets were not sufficiently informative to constrain estimates of n and ϕ . We therefore implemented a penalized likelihood approach, augmenting the two aforementioned likelihood functions with a penalty term proportional to the values of n and ϕ to discourage large values of n and ϕ . More specifically, we performed model fitting using

$$\mathcal{L}_p = \mathcal{L} + \lambda \cdot \ln(n) + \lambda \cdot \ln(\phi) \tag{S.14}$$

as the loss function, where \mathcal{L} is the negative log-likelihood and λ determines the strength of the penalty for values of n and ϕ . Although it is possible to treat λ as a free parameter that is estimated for each dataset, we set $\lambda = 1/\ln(20)$. A value of n or ϕ equal to 20 therefore penalized the negative log-likelihood by 1 unit (equivalent to 1/2 the penalty associated with each parameter of a model under AIC).

Supplementary figures and statistical tables

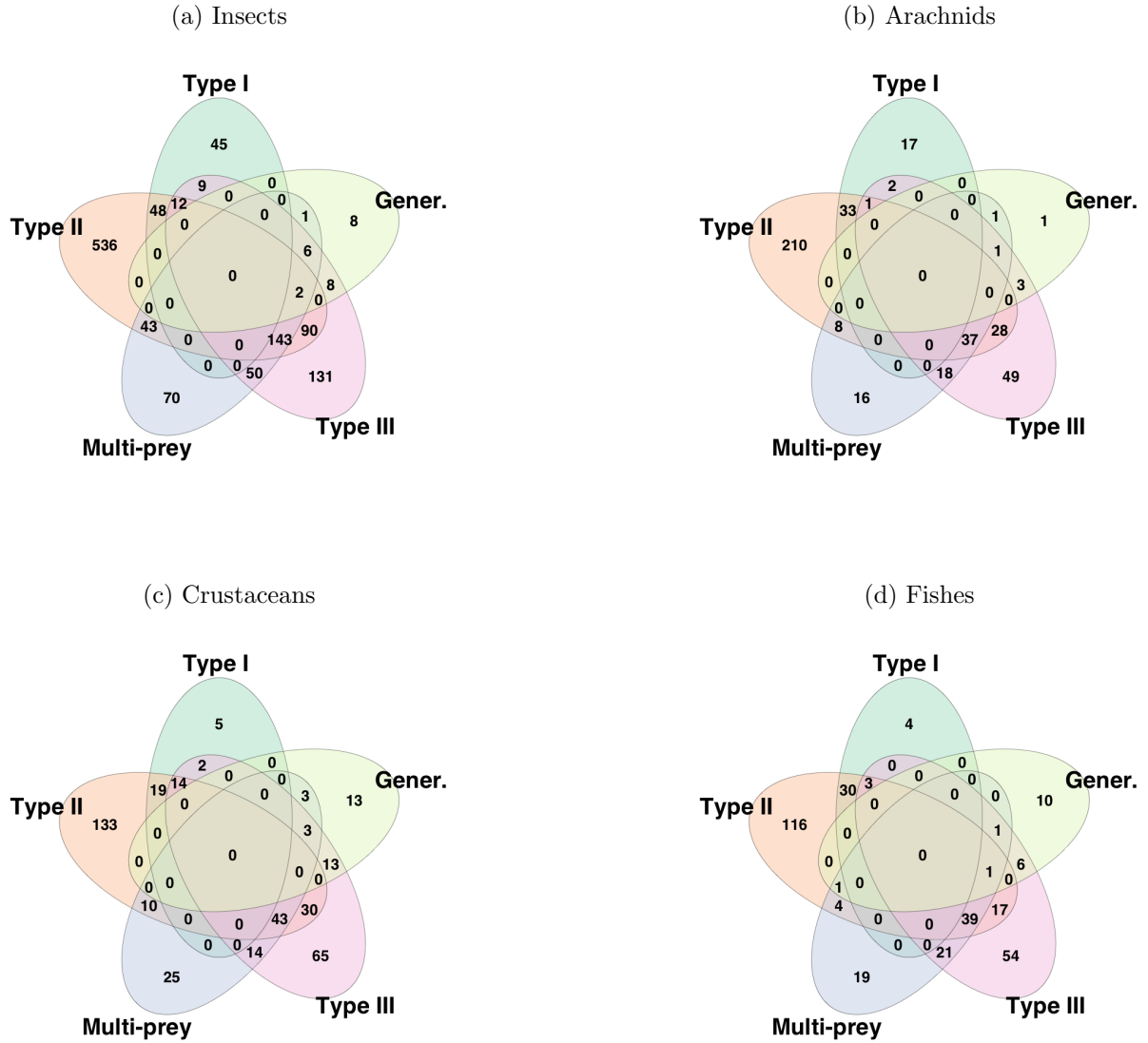


Figure S.2: Venn diagrams categorizing the datasets of the four most common predator groups by their support for one or more of the considered models based on their BIC scores.

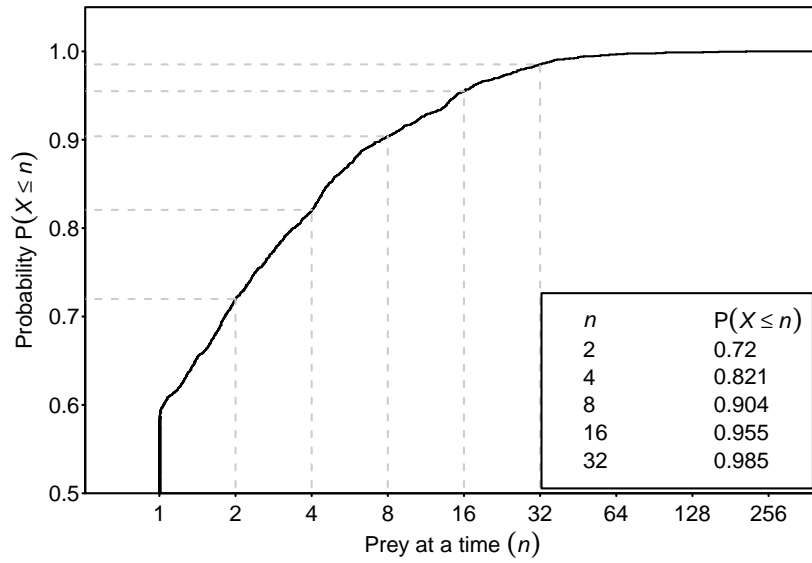


Figure S.3: Cumulative probability distribution of the estimates of n (assuming the multi-prey model) from across all datasets excluding those for which the Type I model alone performed best.

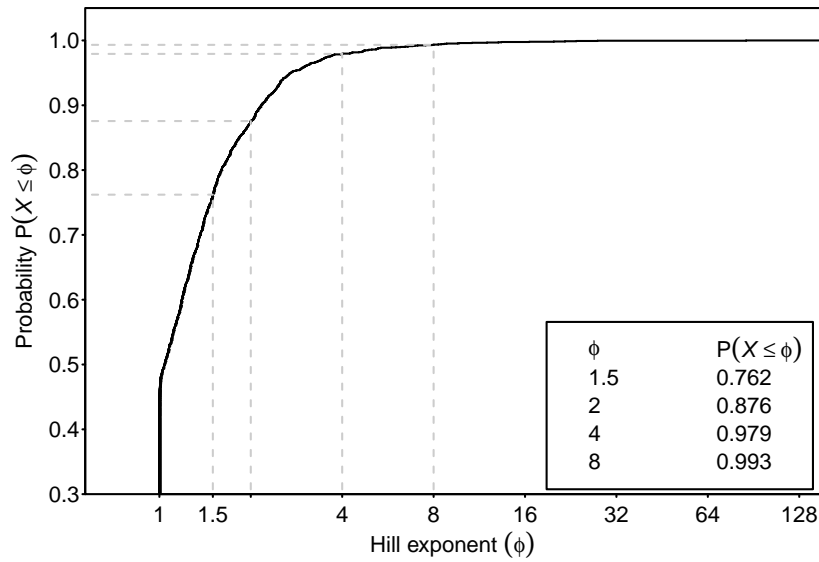


Figure S.4: Cumulative probability distribution of the estimates of ϕ (assuming the Holling-Real Type III model) from across all datasets excluding those for which the Type I model alone performed best.

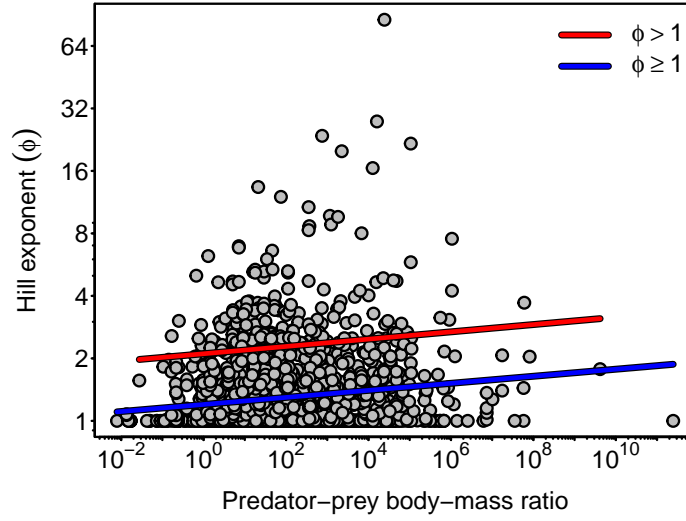


Figure S.5: The relationship between $\log_2(\phi)$ and $\log_{10}(\text{PPMR})$ assuming the Holling-Real model excluding datasets for which the Type I model alone performed best (Table S.2).

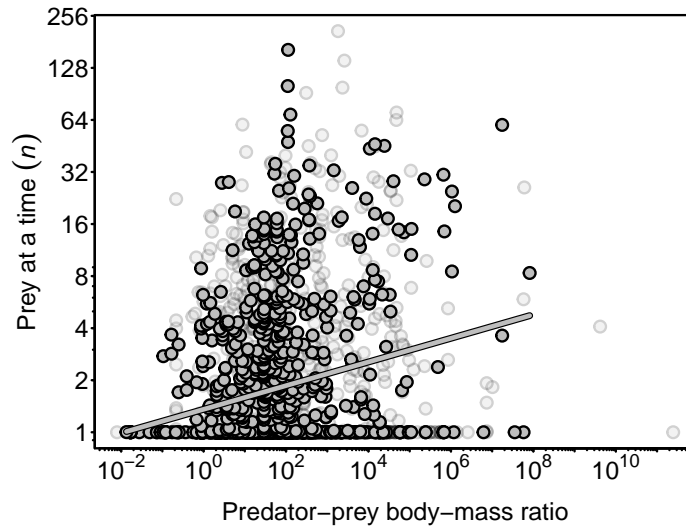


Figure S.6: The relationship between $\log_2(n)$ and $\log_{10}(\text{PPMR})$ assuming the multi-prey model when considering only those datasets having a sample size greater than the median sample size of all datasets excluding those for which the Type I model alone performed best (Table S.3).

Table S.1: Summary statistics (with 95% confidence intervals) for the least-squares linear regressions of $\log_2(n)$ of the multi-prey model on $\log_{10}(\text{PPMR})$ when considering all studies ($n \geq 1$) or only those studies for which $n > 1$.

	Estimates	
	$n \geq 1$	$n > 1$
Intercept	0.546*** (0.455, 0.638)	1.976*** (1.806, 2.147)
$\log_{10}(\text{PPMR})$	0.145*** (0.106, 0.184)	0.190*** (0.122, 0.258)
Observations	2,137	715
R^2	0.024	0.041
Adjusted R^2	0.024	0.039
Residual Std. Error	1.342 (df = 2135)	1.334 (df = 713)
F Statistic	53.006*** (df = 1; 2135)	30.186*** (df = 1; 713)

*p<0.1; **p<0.05; ***p<0.01

Table S.2: Summary statistics (with 95% confidence intervals) for the least-squares linear regressions of $\log_2(\phi)$ of the Holling-Real Type III on $\log_{10}(\text{PPMR})$ when considering all studies ($\phi \geq 1$) or only those studies for which $\phi > 1$.

	Estimates	
	$\phi \geq 1$	$\phi > 1$
Intercept	0.262*** (0.222, 0.302)	1.074*** (0.974, 1.173)
$\log_{10}(\text{PPMR})$	0.056*** (0.039, 0.073)	0.058*** (0.020, 0.097)
Observations	2,137	511
R^2	0.020	0.017
Adjusted R^2	0.019	0.015
Residual Std. Error	0.583 (df = 2135)	0.667 (df = 509)
F Statistic	42.597*** (df = 1; 2135)	8.810*** (df = 1; 509)

*p<0.1; **p<0.05; ***p<0.01

Table S.3: Summary statistics (with 95% confidence intervals) for the least-squares linear regression of $\log_2(n)$ of the multi-prey model on $\log_{10}(\text{PPMR})$ when considering only those studies having a sample size greater than the median sample size of all studies.

	Sample size >36
Intercept	0.440*** (0.309, 0.571)
$\log_{10}(\text{PPMR})$	0.228*** (0.167, 0.289)
Observations	981
R ²	0.052
Adjusted R ²	0.051
Residual Std. Error	1.289 (df = 979)
F Statistic	53.442*** (df = 1; 979)

*p<0.1; **p<0.05; ***p<0.01

Table S.4: Summary statistics (with 95% confidence intervals) for the multiple least-squares linear regression of $\log_2(n)$ of the multi-prey model on $\log_{10}(\text{PPMR}) \times$ predator group for the four most common predator taxonomic groups.

	Focal predators
Intercept (Insect)	0.544*** (0.409, 0.678)
$\log_{10}(\text{PPMR})$	0.167*** (0.090, 0.244)
Arachnid	-0.305** (-0.580, -0.029)
Crustacean	0.208 (-0.063, 0.479)
Fish	-0.164 (-0.680, 0.352)
$\log_{10}(\text{PPMR})$:Arachnid	0.269** (0.050, 0.488)
$\log_{10}(\text{PPMR})$:Crustacean	-0.061 (-0.165, 0.044)
$\log_{10}(\text{PPMR})$:Fish	-0.029 (-0.201, 0.144)
Observations	1,917
R ²	0.032
Adjusted R ²	0.029
Residual Std. Error	1.318 (df = 1909)
F Statistic	9.150*** (df = 7; 1909)

*p<0.1; **p<0.05; ***p<0.01

Population-dynamic effects

Supplementary figures

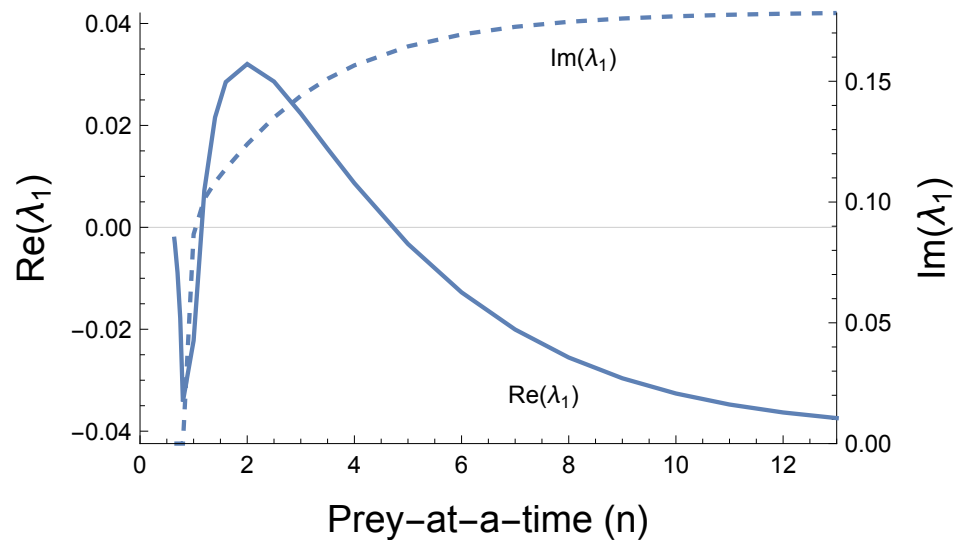


Figure S.7: The coexistence state is asymptotically stable when the real part of the dominant eigenvalue $\text{Re}(\lambda_1)$ is negative. This occurs for $n \approx 1$ where it is globally stable and for $n > 5$ where it is only locally stable. Post-perturbation dynamics towards the stable equilibrium exhibit monotonic damping when the imaginary part $\text{Im}(\lambda_1)$ is zero as occurs for $n \approx 1$, but exhibit damped oscillations when $\text{Im}(\lambda > 0)$ as occurs for higher n . *Other parameter values as in Fig. 3.*

A reformulation of the extended Steady State Saturation model

Jeschke *et al.* (2004) introduced a functional response model that, like the multi-prey model, is capable of exhibiting a continuum of shapes between the Type I and Type II response forms. In its original formulation, their model is written as

$$\frac{e(1 + aN(b + c)) - \sqrt{e(4acN + e(1 + aN(b - c))^2)}}{2c(e(1 + abN) - 1)}, \quad (\text{S.15})$$

where N is the prey's abundance, a is the attack rate, b is the handling time, c is the digestion time, and e is a dimensionless shape parameter interpreted as affecting the trade-off between search effort and hunger level (i.e. gut fullness). The model approaches the rectilinear model as $e \rightarrow \infty$ when $b = 0$ (see Fig. A2 of Jeschke *et al.*, 2004). For $e = 1$ it reduces to the ‘‘Steady State Saturation’’ (SSS) model of Jeschke *et al.* (2002), written in its original formulation as

$$\frac{1 + aN(b + c) - \sqrt{1 + aN(2(b + c) + aN(b - c)^2)}}{2abcN}. \quad (\text{S.16})$$

Both models may be expressed in a formulation more similar to the Holling form that eases a comparison to other functional response models. This may be done by deriving them using the citardauq formula. The SSS may thereby be rewritten as

$$\frac{2aN}{1 + aN(b + c) + \sqrt{1 + aN(2(b + c) + aN(b - c)^2)}}. \quad (\text{S.17})$$

(Note that the equation presented in the original version of Novak & Stouffer (2021a) is incorrect but has subsequently been corrected (Novak & Stouffer, 2024).) The extended SSS with parameter e may be rewritten as

$$\frac{2aN}{1 + aN(b + c) + \frac{1}{e}\sqrt{e(4acN + e(1 + aN(b - c))^2)}}. \quad (\text{S.18})$$

With four parameters, the extended SSS model is capable of exhibiting more variation in shape than the three-parameter multi-prey model. In particular, with sufficiently high e and appropriately chosen non-zero values of b and c , it exhibits curvature at the low prey abundances where the multi-prey model with high n is effectively linear (see Figs. A1 and A2 of Jeschke *et al.*, 2004).