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Bivariate stochastic modelling of functional response with natural mortality

M.J. FADDY, J.S. FENLON and D.J. SKIRVIN

A correction due to Abbott (1925) is the standard method of dealing with control mortality in insect bioassay to estimate the mortality of an insect conditional on control mortality not having occurred. In this paper a bivariate stochastic process for overall mortality is developed in which natural mortality and predation are jointly modelled to take account of the competing-risks associated with prey loss. The total mortality estimate from this model is essentially identical with that from more classical modelling. However, when predation loss is estimated in the absence of control mortality the results are somewhat different, with the estimate from the bivariate model being lower than that from using Abbott's formula in conjunction with the classical model. It is argued that over-dispersion in observed mortality data corresponds to correlated outcomes (death or survival) for the prey initially present, while Abbott's correction relies implicitly on independence.

Keywords: Stochastic modelling; Competing risks; Estimation of predator mortality; Over-dispersion.

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1 INTRODUCTION

It is often the case in bioassays that mortality of insects can occur for reasons other than those due to the substance under investigation. Morgan (1992) makes the distinction between ‘natural’ and ‘control’ mortality, citing Hoekstra (1987) and Preisler (1989). Natural mortality may occur as well as mortality due to handling of the insects, or mortality may be due to the fact that the insects are not in their natural habitat. In order to assess the degree of ‘control mortality’, groups of insects are often kept under conditions that are the same as those for the treated groups, except for the absence of the treatment. Abbott’s formula (Abbott, 1925) is then commonly used to estimate treatment mortality in the absence of control mortality for dose response assays. For a given dose d , if the expected proportion responding is p then this is not directly observable because of contamination through natural mortality; so if the expected overall proportion dead is p^* , then p^* is made up of the proportion of insects φ that die naturally, and a proportion p of the remainder $(1 - \varphi)$ that die due to the treatment, so that $p^* = \varphi + (1 - \varphi)p$. The actual probability of dying because of the dose is therefore given by $p = (p^* - \varphi)/(1 - \varphi)$. Hewlett and Plackett (1979) remark that it is rarely possible to test the assumptions on which this formula is based, and cite Kuenen (1957). They make the rather surprising comment that “control response should be avoided if possible”.

A particular type of assay in predator-prey studies, known as a functional response assay, involves experiments in which different numbers of prey are made available to a single predator (see, *e.g.* Fenlon and Faddy, 2006). The data presented here derive from a predator-prey study in which the assay arena has been extended to simulate small plants. The purpose of the original study was to derive realistic estimates of predation rates for input into larger

simulation models to examine practical biological control scenarios (Skirvin *et al.* 2002). However, in moving to larger arenas it was not possible to guarantee recovery of non-predated prey, so that a simultaneous assay procedure was used with separate arenas for control and predator groups. Classical models with beta-binomial distributed residuals about a mean function, as described in Fenlon and Faddy (2006), are first fitted to the data, with predator mortality estimated using Abbott's formula. The main contribution of the paper is the development of a bivariate stochastic process model for both predation loss and natural mortality as an extension of the univariate stochastic model in Fenlon and Faddy (2006), and the fitting of this model to the data. This bivariate modelling owes its genesis to a recent paper by Faddy and Smith (2005). Estimates of predator mortality in the absence of control mortality from the classical and stochastic modelling approaches are then compared, and the paper concludes with some speculations on the reasons for the apparent differences between these estimates.

2 DATA

Details of the experimental procedure are given in Skirvin and de Courcy Williams (1999) and Skirvin and Fenlon (2001). The assay system comprised cut stems with four tri-lobate leaves of *Choisya ternata* (a popular garden ornamental sometimes called Mexican orange blossom). Mini-plants were 'primed' with webbing of the spider mite *Tetranychus urticae*, a pest of many garden plants, before elimination of all life stages of the spider mite.

Different levels of pest load were then transferred to individual stems using a fine paintbrush, and a single *Phytoseiulus persimilis* (predator) was introduced onto each stem before the stems were placed in individual perspex cylinders which were sealed with ventilated lids. The number of assays at each availability level (2, 5, 10, 20 and 40 prey

initially) with the predator present varied (average 19), and there were 25 control assays with 40 prey initially, in which no predator was present, to monitor natural loss(es).

For each individual assay the number of surviving prey was counted after 24 hours, and the number predated or lost determined by difference. Data from assays in which the predator was not found, or was dead, were excluded. Similarly, control assays in which a predator was found were also removed from the analysis. The data with some summary statistics are presented in Table 1 and plotted in Figure 1. With the exception of availability level 2, it appears that all these data show over-dispersion relative to the binomial distribution, with the over-dispersion tending to increase with prey availability (the variance ratio statistics in the final row of Table 1 bear this out).

3 CLASSICAL MODELLING

3.1 BETA-BINOMIAL MODEL

In the summaries of the data it is quite clear that simple binomial modelling is really of no value since over-dispersion relative to the binomial distribution is significant. So, a beta-binomial (residual) distribution was used:

$$P(n \text{ prey lost}) = \binom{N}{n} \frac{B\left(n + \frac{\mu}{N\theta}, N - n + \frac{N-\mu}{N\theta}\right)}{B\left(\frac{\mu}{N\theta}, \frac{N-\mu}{N\theta}\right)}$$

where $B(\cdot, \cdot)$ is the beta function. The mean of this distribution is μ and the variance

$\mu\left(1 - \frac{\mu}{N}\right)\left[1 + \left(\frac{\theta}{\theta+1}\right)(N-1)\right]$ where θ is an over-dispersion parameter. The mean response of

the compound data (predation with natural mortality) was modelled as a function of the availability N , with some adjustment being made for natural (or control) mortality using Abbott's formula. The underlying model then takes the form:

$$\mu_{control} = N\varphi \quad (1)$$

and
$$\mu_{compound} = N\varphi + (1 - \varphi)\mu_{predator} \quad (2)$$

where $\mu_{control}$, $\mu_{compound}$ and $\mu_{predator}$ correspond to the means for the respective subscript groups. Only data relating to the control and compound means are observable, with the predator mean being inferred. As in Fenlon and Faddy (2006) $\mu_{predator}$ is modelled using a Gompertz function constrained to pass through the origin:

$$\mu_{predator} = a\{\exp[-b\exp(-cN)] - \exp[-b]\} \quad (3)$$

which corresponds to a Type III functional response model (Holling, 1959). Some additional modelling of the over-dispersion parameter θ as a function of N was indicated by the data – the details are deferred to the next sub-section.

3.2 RESULTS

A saturated beta-binomial model was first fitted to the control and compound data by maximum likelihood; this consisted of individual two-parameter beta-binomials for each prey availability level, N . It was noted that estimates of the beta-binomial over-dispersion parameter, θ , generally declined with prey availability, so several forms were tried, with means from equations (1), (2) and (3), to describe the control and compound data over all availability levels:

- (a) $\theta = d$,
- (b) $\theta = d_0$ for control data, and $\theta = d_1$ for the compound data,
- (c) $\theta = d + e/N$ for all data,

(d) $\theta = d_0$ for control data, and $\theta = d_1 + e/N$ for the compound data.

All fits resulted in large estimated a 's and small b 's in the form of $\mu_{predator}$ given by (3), corresponding to the limiting ($a \rightarrow \infty$ and $b \rightarrow 0$ with $ab = a'$) form:

$$\mu_{predator} = a' [1 - \exp(-cN)], \quad (4)$$

with a' and $c > 0$; *i.e.*, type II behaviour without an inflexion point (Holling, 1959). The respective log-likelihoods from the model fits were -249.0 , -248.5 , -248.3 and -248.2 , and the corresponding AIC values ($-2 \times \{\log\text{-likelihood} - \text{no. of parameters}\}$) 505.9 , 507.0 , 506.6 and 508.4 , which all suggest that staying with the simplest model (a) is not unreasonable.

However, the corresponding generalised Pearson statistics (sums of squared standardised residuals), which give measures of goodness of fit, were 125.8 , 123.9 , 118.9 and 119.4 with 115 , 114 , 114 and 113 d.f., respectively. These show quite a marked reduction when θ is made dependent on N , and suggest that a model based on (c) might be preferred, with the generalised Pearson statistic indicating a better fit. Such a decline in θ with N does moderate the dispersion for larger N , more in accord with the values in Table 1. This model also compares favourably with the saturated model (deviance = 5.3 on 7 d.f.). The fit of the model to the compound (predator + control) mortality data is illustrated in Figure 1 where the estimated mean from (2) and (4) is plotted with \pm one standard deviation limits. The fit to the control data gave estimated mean 6.8 and variance 16.2 , compared with the observed values of 6.9 and 17.8 shown in Table 1. The most extreme observation is the count of 19 in this control data sub-set, and corresponds to a tail probability of just under 0.01 , which would not be unexpected in a data-set of size 119 in total. In Figure 2 is the (Abbott's formula corrected) mean predator mortality derived from the estimated mean compound and control mortality given by (1) and (2); the \pm one standard deviation limits here have been

estimated under the assumption that a beta-binomial distribution applies with the same over-dispersion parameter as that associated with the compound mortality.

4 STOCHASTIC PROCESS MODELLING

4.1 BIVARIATE STOCHASTIC MODEL

In Fenlon and Faddy (2006) probability distributions on $0, 1, \dots, N$ for the number of prey lost were constructed from a univariate Markov process $\{X(t); t \geq 0\}$ with $X(0) = 0$ and rate parameters $\lambda_0, \lambda_1, \dots, \lambda_{N-1}, \lambda_N$ (with $\lambda_N = 0$) where:

$$P\{X(t + \delta t) = n + 1 | X(t) = n\} = \lambda_n \delta t.$$

The modelling of prey loss is here extended to a bivariate Markov process

$\{X(t), Y(t); t \geq 0\}$, with one component $X(t)$ for natural mortality and the other $Y(t)$ for predator mortality, with $0 \leq X(t) + Y(t) \leq N$ and $X(0) = Y(0) = 0$. So that two transitions now have to be considered, with infinitesimal probabilities:

and

$$\begin{aligned} P\{X(t + \delta t) = x + 1, Y(t + \delta t) = y | X(t) = x, Y(t) = y\} &= \lambda_{xy}^{(1)} \delta t \\ P\{X(t + \delta t) = x, Y(t + \delta t) = y + 1 | X(t) = x, Y(t) = y\} &= \lambda_{xy}^{(2)} \delta t \end{aligned} \quad (5)$$

representing, respectively, an increase in the natural mortality of the prey, and an increase in predator-induced mortality. Whilst $X(t)$ can be observed $Y(t)$ cannot; what is observed, rather, is the total mortality $X(t) + Y(t)$. However, since the processes may not be independent, the rates $\lambda_{xy}^{(1)}$ of natural mortality and $\lambda_{xy}^{(2)}$ of predator mortality cannot simply be added together to describe the total mortality.

The solution for the probabilities $p(x, y) = P\{X(1) = x, Y(1) = y\}$ where the process of prey loss is taken, without loss of generality, to run for one unit of time can be expressed in terms of a matrix, \mathbf{Q} , of transition rates from (5). The rows and columns of this matrix are indexed by (x, y) taking the values $(0,0), (0,1), \dots, (0,N); (1,0), (1,1), \dots, (1,N-1); \dots; (N,0)$ with the three non-zero elements of \mathbf{Q} in the row corresponding to (x, y) being:

$$-(\lambda_{xy}^{(1)} + \lambda_{xy}^{(2)}) \text{ in the column corresponding to } (x, y),$$

$$\lambda_{xy}^{(1)} \text{ in the column corresponding to } (x+1, y), \text{ and}$$

$$\lambda_{xy}^{(2)} \text{ in the column corresponding to } (x, y+1),$$

except when $x+y=N$ in which case all the entries are zero. The probability $p(x, y)$ is then the appropriate element in the row vector (*cf.* Fenlon and Faddy, 2006):

$$\begin{aligned} [p(0,0) \ p(0,1) \ \dots \ p(0,N) \ p(1,0) \ p(1,1) \ \dots \ p(1,N-1) \ \dots \ p(N,0)] \\ = [1 \ 0 \ \dots \ 0] \exp(\mathbf{Q}), \end{aligned} \quad (6)$$

with the probability of a total of n prey lost then given by the convolution $\sum_{x=0}^n p(x, n-x)$, for $n=0,1,\dots,N$.

A reasonable description of control mortality would be in terms of the total numbers lost, *i.e.* $\lambda_{xy}^{(1)} = f(x+y)$. For predation mortality, following Fenlon and Faddy (2006), a product form is used where $\lambda_{xy}^{(2)} = g(y)h(x+y)$; here $g(y)$ can be thought of as a predator behaviour component [*e.g.* $g(y)$ constant would correspond to constant predator activity regardless of the number of prey consumed] and $h(x+y)$ as relating to the prey [*e.g.* $h(x+y) = N-x-y$ would correspond to the remaining prey being exchangeable (Faddy and Fenlon, 1999)]. The expression:

$$\lambda_{xy}^{(1)} = -\log(1-\varphi)(N-(x+y)) \quad (7)$$

corresponds to a simple binomial distribution with probability φ for control mortality in the absence of predator mortality. And, following Fenlon and Faddy (2006), the form

$$\lambda_{x,y}^{(2)} = \exp(\alpha + \beta y) \frac{\left[1 - \exp\left\{-\delta(N - (x + y))^\varepsilon\right\}\right]}{\delta} \quad (8)$$

was used, which, for $\beta > 0$, corresponds to greater predator activity with increasing prey consumed and, for $\delta > 0$ and/or $\varepsilon \neq 1$, non-exchangeable prey.

4.2 RESULTS

The bivariate stochastic process model for both control and predator mortality described by the transition rates in equations (5) and the probabilities from (6) with the expression (7) for $\lambda^{(1)}$, as has already been remarked, corresponds to a simple binomial distribution with probability φ for control mortality in the absence of predator mortality. However, the analysis of section 3.2 showed that the control mortality exhibited over-dispersion relative to the binomial distribution, and that a beta-binomial distribution was more appropriate. Beta-binomially distributed control mortality can be incorporated into the bivariate stochastic process with the probability of x prey lost naturally and y lost by predation being given by:

$$\int_0^1 p(x, y | \varphi) \frac{\varphi^{r-1} (1-\varphi)^{s-1}}{B(r, s)} d\varphi, \quad (9)$$

where $p(x, y | \varphi)$ is the probability of x prey lost naturally and y prey lost by predation for a given value of φ , determined from equations (6), (7) and (8), and the other component in the integrand of (9) is the probability density function of a beta distribution.

Use of probabilities given by equation (9) is very costly in computing time, as numerical computation of the integral would involve many matrix exponential evaluations of quite

large matrices (6); to expedite these computations a discrete binomial mixture distribution for control mortality was used as an alternative. A discrete mixed binomial model uses probabilities $\sum_{i=1}^k w_i b(N, \varphi_i, x)$, where $0 \leq w_i \leq 1$, $\sum_{i=1}^k w_i = 1$ and $0 \leq \varphi_i \leq 1$ for $i = 1, 2, \dots, k$,

k being the number of components in the mix, and $b(N, \varphi_i, x) = \binom{N}{x} \varphi_i^x (1 - \varphi_i)^{N-x}$,

$0 \leq x \leq N$, are binomial probabilities. Table 2 shows details of maximum likelihood fitting of these distributions to the control data: a two component binomial mixture model would be selected according to the AIC values, and it shows a fit to the data that is comparable to the beta-binomial.

Fitting the bivariate stochastic process model with a two component binomial mixture distribution for the control mortality simply involves probabilities $p(x, y | \varphi_1)$ and $p(x, y | \varphi_2)$ created from only two matrices Q_1 and Q_2 derived from equations (6), (7) and (8) with probabilities φ_1 and φ_2 , respectively, and using the discrete version of expression (9):

$$w_1 p(x, y | \varphi_1) + (1 - w_1) p(x, y | \varphi_2).$$

Parameter estimates and their corresponding standard errors from fitting this model to the control and compound mortality data are given in the second and third columns of Table 3. The log-likelihood under this model is -245.4 , compared to -248.3 from the previously fitted beta-binomial model. However, these two models have different sub-models (two-component mixed binomial and beta-binomial, respectively) for the control data, and the likelihoods for the compound data only are more comparable: -179.4 and -180.5 , respectively.

The results in Table 3 suggest that setting $\beta = 0$ in equation (8), corresponding to constant predator activity, might be acceptable, and indeed, fitting such a reduced model results in only a small decline in the log-likelihood from -245.4 to -246.0 . However, the overall generalised Pearson statistic increases from 114.6 on 112 d.f. to 125.1 on 113 d.f. when $\beta = 0$, indicating a poorer fit, and the latter model seems less convincing for the data corresponding to $N = 5$ and 20. Exceedance probabilities of apparent outliers for the full ($\beta > 0$) model correspond to one with tail probability about 0.01 and three with tail probabilities between 0.01 and 0.05, which would seem unremarkable given the size of the data-set.

Shown in Figure 1 is the estimated mean total mortality from the full ($\beta > 0$) model (along with the data and estimated means from the previously fitted beta-binomial model); the figure also shows the spread of \pm one standard deviation about the mean for both models. These two fits are clearly very similar. The estimated mean control mortality was 6.8, similar to the earlier fitted beta-binomial value, but the estimated variance was slightly larger at 19.0 (*cf.* 16.2) so that the 2-component mixed binomial distribution used here is somewhat better able to accommodate the extreme count of 19 in the control data sub-set.

The estimated predation when natural mortality is not present can be determined from the distribution of $Y(1)$ corresponding to $\lambda^{(1)} \equiv 0$ in (5), and is shown in Figure 2 (along with that from the beta-binomial model); the \pm one standard deviation limits here have been estimated from this reduced model with no further assumptions necessary. Of note is the smaller magnitude of the estimate from the stochastic process model compared with that from the classical beta-binomial model.

5 DISCUSSION

The main contribution of this paper has been the construction of a bivariate stochastic process model for natural and predator mortality, and the comparison of the results from fitting this model to a data-set with those from a more classical approach. The differentiation methods used to estimate net predation for the two models in the absence of natural mortality gave rather different results: the stochastic process model used the univariate process with the natural mortality transition rates set to zero, whereas the beta-binomial modelling used Abbott's formula. This resulted in rather different estimates, with those from the stochastic model being lower than those from the beta-binomial (Figure 2), even though the estimates of total mortality were virtually identical (Figure 1). While both mean responses and the standard deviation from the stochastic process model have stabilised by $N = 40$, the standard deviation from the beta-binomial model continues to increase; however, the assumption under which this has been estimated, *i.e.* that the level of over-dispersion is the same as that for total mortality, is clearly unverifiable. An aesthetic argument can be made for the stochastic process modelling which has a consistency insofar as the 'tension' between the control and compound mortality (with the former sometimes outrunning the latter for large N) leads to an early horizontal asymptote of the compound mortality (effectively satiation) within the experimental range. Furthermore, the stochastic modelling attempts to model the actual temporal process of predation, and offers a possible explanation of how the predator behaves in relation to changing prey availability, even though the data are observed only at a single time point. The problem associated with over-dispersion in the control data has been addressed in the stochastic modelling by the use of a discrete binomial mixture model to obviate the need for costly numerical integration involving many matrix exponential evaluations associated with using a beta-binomial distribution.

Abbott's formula essentially estimates for an individual prey the *conditional* probability of loss due to the predator *given* that natural death has not occurred, where predator and natural mortality can be considered as competing risks. A more appropriate estimate is that of the *unconditional* probability of loss due to the predator. These probabilities will be the same if the risks of predator and natural mortality are assumed to apply *independently* to the prey. A further assumption that loss (either natural death or death from predation) occurs independently between prey when there are several available initially will lead to a binomial distribution of the number of prey lost during the course of an experiment or study, with Abbott's formula giving an estimate of the unconditional mean number of prey lost to the predator.

However, if the observed number of prey lost shows residual variation in excess of that corresponding to binomial variation then *both* the above assumptions of independence are called into question. Extra-binomial variation in numbers lost can be explained either by a residual distribution that shows over-dispersion (such as the beta-binomial) or a stochastic process running during the course of the experiment with rates of loss of individual prey that are not constant but a function of the accumulating number of prey lost. In both of these models, the over-dispersion corresponds to correlated outcomes (death or survival) for the initial number of prey. And there is an equivalence between these models in that any distribution showing over-dispersion relative to the binomial has a representation in terms of a stochastic process with varying rates of individual prey loss (Faddy, 1997) – some increase in these rates with the accumulating number of prey lost gives rise to the over-dispersion. Such a stochastic representation of predator mortality will result in the two processes, predator loss and loss due to natural mortality, not being independent of one another. This is because individual prey that might die naturally later in the experiment

would be exposed to higher rates of predation than those that might die earlier, since more prey are likely to be consumed by the predator in a longer period of time. So Abbott's formula, which (generally) leads to an estimate of the mean number of prey lost to the predator conditional on natural death not occurring, will *not* give an estimate of the unconditional mean number of prey lost to the predator because of this dependence between natural and predator mortality. However, the *bivariate* stochastic model of predator and natural mortality will give an estimate of this unconditional mean number of prey lost to the predator if the natural mortality component of the model is set to zero.

There is no doubt that there are challenges in modelling these data, and deriving estimates of predator mortality. Further models may be developed, but what has been presented here does offer a novel approach to the problem and a very plausible estimate of net predation, with interpretable dynamics. The use of stochastic process modelling ultimately provides an appropriate temporal structure giving a reasonable explanation for observed data on functional response, as has been previously argued in Fenlon and Faddy (2006).

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Table 1: observed numbers and summary statistics of *T. urticae* adults consumed or lost (with predator) and control c (without predator) for groups of assays with the same prey availability; ρ is the ratio of sample to binomial variance (a measure of over-dispersion).

assay no.	prey availability					
	2	5	10	20	40	40(c)
1	2	5	10	17	22	19
2	2	5	9	15	19	14
3	2	5	9	14	18	13
4	2	5	8	14	16	13
5	2	5	8	12	13	10
6	2	5	8	12	13	10
7	2	5	8	12	11	9
8	2	4	7	11	11	7
9	2	4	7	9	11	7
10	2	4	7	8	10	7
11	2	3	7	8	10	7
12	2	3	6	8	9	6
13	2	3	6	7	9	6
14	2	3	6	6	7	5
15	2	3	5	5	7	5
16	1	3	5	3	5	5
17	1	3	5			4
18	1	2	5			4
19	1	1	5			4
20		1	5			4
21			5			4
22			3			3
23			2			3
24						2
25						2
mean	1.8	3.6	6.3	10.1	11.9	6.9
variance	0.2	1.7	3.8	15.4	22.1	17.8
proportion	0.89	0.72	0.63	0.50	0.30	0.17
ρ	0.9	1.7	1.6	3.1	2.6	3.1

Table 2: log-likelihood, generalised Pearson and AIC statistics for mixed binomial models in comparison to the beta-binomial model for the control data.

Model	log-lik.	gen. Pearson	d.f.	AIC
simple binomial	-77.9	74.8	24	157.7
2-component mix	-65.9	25.4	22	137.8
3-component mix	-65.7	24.2	20	141.5
beta-binomial	-67.8	28.0	23	139.6

Table 3: parameter estimates of the bivariate stochastic process model of natural mortality and predator mortality.

Parameter	full model		model with $\beta = 0$	
	estimate	s.e.	estimate	s.e.
$\hat{\phi}_1$	0.12	0.014	0.12	0.014
$\hat{\phi}_2$	0.34	0.041	0.35	0.039
\hat{w}_1	0.77	0.094	0.74	0.084
$\hat{\alpha}$	0.86	0.28	0.93	0.28
$\hat{\beta}$	0.057	0.041	–	–
$\hat{\delta}$	0.44	0.14	0.41	0.13
$\hat{\varepsilon}$	0.77	0.23	0.79	0.24

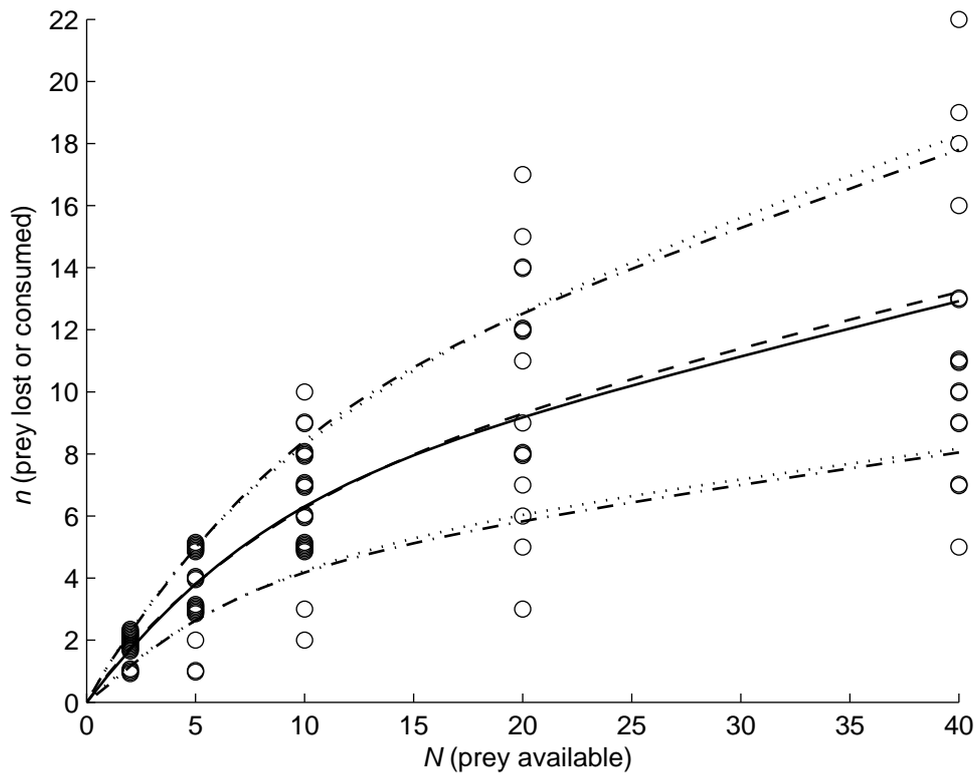


Figure 1: estimates of mean total prey lost or consumed from the bivariate stochastic process model (—) and the beta binomial model (---); \pm one standard deviation limits are shown by (- · -) and (·····), respectively, for the two models. Data are denoted by open circles, jittered slightly to indicate multiple data-points.

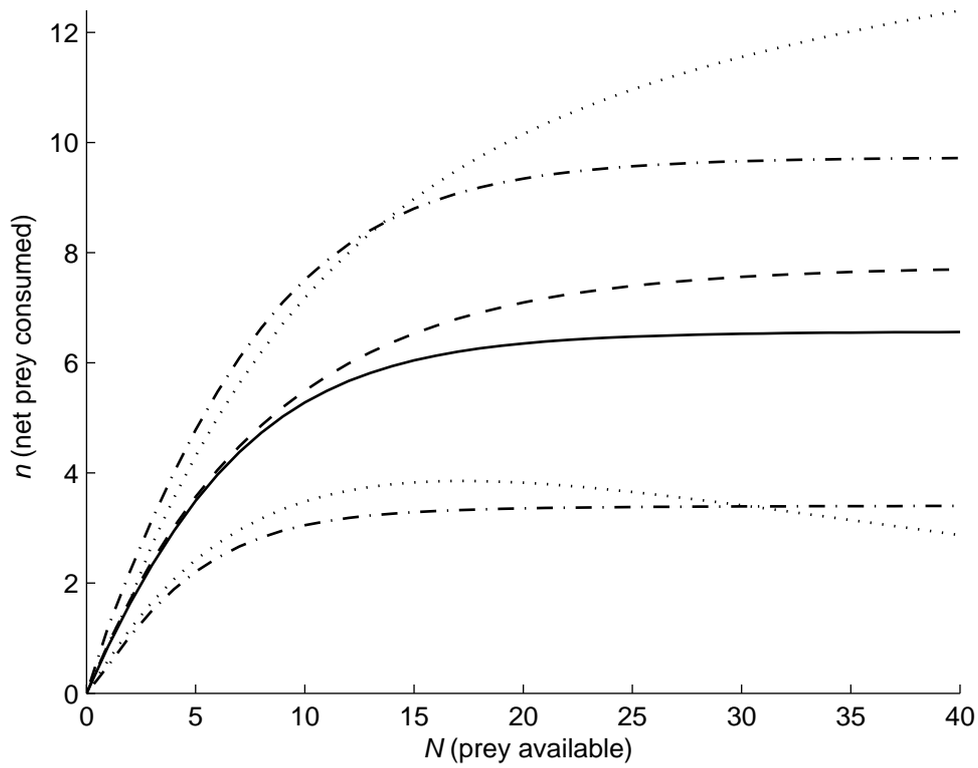


Figure 2: estimates of mean net prey consumed by the predator from the bivariate stochastic process model (—) and the beta binomial model (---); \pm one standard deviation limits are shown by (- · -) and (·····), respectively, for the two models.