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[Re] Modeling Insect Phenology Using Ordinal Regression and Continuation Ratio Models

Philipp H. Boersch-Supan^{1,2, ID}

¹British Trust for Ornithology, Thetford, United Kingdom – ²University of Florida, Gainesville, FL, USA

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Introduction

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Phenology, the timing of seasonal biological phenomena, is a key aspect of plant and animal life. It defines the timing and duration of growth and reproduction and thereby determines the ability to capture seasonally variable resources [1]. Phenological analyses often focus on the timing of particular events, such as the dates of peak plant flowering [2]. However, for many biological phenomena exact dates of particular events are more difficult to observe than the state of the system itself. For example, repeated but sparse survey visits may record whether a plant is in bud, flowering, or setting fruit, but not the exact dates when each of those stages was reached. Such observations can be used to categorize an organism's state into discrete classes which usually follow natural ordering, e.g. from least to most developed. The resulting data can be described using ordinal regression models [3, 4].

I here replicate a number of ordinal regression models that were developed by Dennis, Kemp, and Beckwith⁵ and Candy⁶ to describe insect phenology.

2 Data

The models replicated in this study are fitted to a data set on the phenology of the western spruce budworm *Choristoneura freemani* (Lepidoptera: Tortricidae), a defoliating moth that is widespread in western North America [7]. This data set was originally published in [5] and is a subset of a larger budworm survey data set analysed in [8]. The data consist of 12 sampling occassions at which counts of individual budworms in each of seven development stages (five larval instars, pupae, and adults) were recorded. The only available covariate is a measure of seasonal progression, the accumulated degree days calculated using a threshold of 5.5°C. Candy⁶ noted an inconsistency in these data, namely that the reported total number of individuals did not correspond to the sum across the seven development stages for two of the sampling occasions. I therefore use the data set as it was republished in [9], where numbers in each stage have been assumed correct and the totals for each sampling occasion were adjusted accordingly.

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The authors have declared that no competing interests exist.
Code is available at https://github.com/pboesu/replication_candy_1991..
Data is available at https://github.com/pboesu/replication_candy_1991.

3 Methods

The statistical models replicated here are different types of ordinal regression models [4] all with the aim of predicting the proportion of an insect population in a particular development stage at any given given time. In particular, they represent three different parametrisations of the so-called cumulative model and one version of the so-called sequential model. A recent summary of the theory underlying these models is provided in [10].

The models generally assume that the development of an insect follows an unobservable stochastic process S(t) consisting of accumulated increments of development over time t. As the amount of S(t) increases, the insect passes through successive stages, delimited by moults, with the jth moult occurring when the development threshold a_i is reached:

 $\begin{array}{lll} \operatorname{stage} 1: & S(t) \leq a_1 \\ \operatorname{stage} 2: & a_1 < S(t) \leq a_2 \\ & \vdots & \vdots \\ \operatorname{stage} r - 1: & a_{r-2} < S(t) \leq a_{r-1} \\ \operatorname{stage} r: & a_{r-1} < S(t) \end{array}$

The a_i values are typically unknown and must be estimated from the data.

3.1 Cumulative model with constant variance

If the cumulative number of individuals observed in stages 1 to j is given by $m_{ij} = \sum_{k=1}^{j} n_{ik}$ then the ordinal regression model [3] is specified by

$$E(m_{ij}) = N_i Pr(S(t) < \alpha_j), \qquad j = 1, \dots, r$$

$$= N_i G(\alpha_i + \beta z_i)$$
(2)

where G is the cumulative probability density function of S(t), α_j are ordered thresholds or cut-point parameters, β is a vector of regression parameters and z_i is a vector of predictor variables. If the probability of an individual being in stage j or earlier at time t_i is

$$\mu_{ij} = \mathbf{E}(m_{ij})/N_i$$

one can define ${\cal G}^{-1}$ as the link function of a generalised linear model with the linear predictor

$$\eta_{ij} = \alpha + \beta z_i$$

. This ordinal regression model is commonly known as the cumulative model, and is applied to the budworm data in [6] using the logit and complementary log-log link functions. In both cases the parameterisation results in a constant variance for S(t). Candy⁶ reexpresses the model in terms of stage-specific counts n_{ij}

$$\mathbf{E}(n_{ij}) = N_i \{ G(\alpha_j + \beta z_i) - G(\alpha_{j-1} + \beta z_i) \}$$
(3)

and fits it using a Poisson likelihood [11]. No code or inital values for the likelihood optimisation are provided for this estimation procedure in the original paper. I therefore created an R version of the estimation procedure which directly optimizes a Poisson log-likelihood for (3) using the optim function. The cumulative model is also implemented in various R packages, including in the vglm function in VGAM [12] and the clm function in ordinal [13] and I here make use of these to fit the model to the budworm data.

3.2 Cumulative model with proportional variance

Dennis, Kemp, and Beckwith⁵ proposed a different parameterisation of the ordinal model is based on assuming a logistic distribution for S(t), such that the probability that an insect's development at time t has not exceeded s amounts to

$$Pr[S(t) \le s] = 1 / \left\{ 1 + \exp\left[-\left(\frac{s-t}{\sqrt{b^2 t}}\right)\right] \right\}$$
 (4)

where b^2 is a positive constant which also must be estimated from the data. This distribution has a mean of t and a variance of $(\pi^2/3)b^2t$. At any fixed time t the thresholds a_j segment the porbaility distribution function in to r parts and the area under the curve between a_{i-1} and a_i gives the probability that the insect will be in stage i at time t. This modelling approach is applied to a dataset consisting of samples that record the number of insects x_{ij} in stage j at times t_1, t_2, \ldots, t_q and the x_{ij} are assumed to be random samples from a multinomial distribution with corresponding multinomial probabilities p_{ij}

$$p_{ij} = Pr[a_{j-1} < S(t_i) \le a_j] \tag{5}$$

$$= 1 / \left\{ 1 + \exp\left[-\left(\frac{a_j - t_i}{\sqrt{b^2 t_i}}\right)\right] \right\} - 1 / \left\{ 1 + \exp\left[-\left(\frac{a_{j-1} - t_i}{\sqrt{b^2 t_i}}\right)\right] \right\}$$
 (6)

To fulfill the constraint that $\sum_{j=1}^{r} p_{ij} = 1$ it is further assumed that $a_0 = -\infty$ and $a_r = +\infty$. The model has r unknown parameters a_1, \ldots, a_{r-1} and b^2 which can be found by maximising the corresponding log-likelihood function which takes the form

$$\ell = \log C + \sum_{j=1}^{r} \sum_{i=1}^{q} x_{ij} \log p_{ij}$$
 (7)

where C is a combinatorial constant that is independent of the parameter values. Dennis, Kemp, and Beckwith⁵ provided SAS code and initial values to estimate the parameters under this likelihood using an interatively reweighted non-linear least squares approach based on PROC NLIN. This was updated to run in a contemporary version of SAS (SAS 9.4) and is provided in the article code repository. However, since SAS is a proprietary software package, I created an R version of the estimation procedure which directly optimizes the log-likelihood (7) using the optim function and initial values provided in [5].

Candy⁶ re-expresses (6) to match the form of (3), which results in the following reparameterisation $\alpha_j = a_j/b$, $\beta = -1/b$, and $z_i = \sqrt{t_i}$, and uses the poisson likelihood approach described above for parameter estimation. No code is given for the estimation of the version of the model in [6], however an earlier manuscript by the same author [9] provides a set of example macros for the software package GLIM [14] which is no longer actively developed or distributed. Initial values from the GLIM code were used in the estimation with R.

3.3 Sequential model

A different class of ordinal regression models, the sequential model, can be derived by treating the observations as the result of a strictly ordinal counting process, in the sense that to achieve a stage j, all lower stages $1,\ldots,j-1$ have to be achieved. The general form of this model is known as the sequential model, and rather than assuming a single latent process S(t) as in the cumulative model there is a latent continuous variable S_j for each category j. As in the cumulative model this can be framed as a GLM

$$S_i = \eta + \epsilon_i \tag{8}$$

with a linear predictor η and an error term ϵ_j which has mean zero and is distributed following some distribution G. This leads to a model of the form

$$Pr[S = j | S \ge j, \eta] = G(a_j - \eta) \tag{9}$$

When G is the logistic distribution this model is also known as the continuation ratio model [15]. Confusingly, there are two common versions of the model in the literature both using this name. The one outlined above describing the probability of the sequential process *stopping* at stage j, and the other describing the probability of the process *continuing* beyond stage j, i.e. $Pr(Y > j|Y \ge j)$ [10, 12]. The paper replicated here [6] used the stopping parametrisation of this model. In their notation this leads to an expected value for the stage-specific counts n_{ij}

$$E(n_{ij}) = N_i G(\beta_{01} + \beta_{11} t_i), j = 1$$

$$= N_{ij}^* G(\beta_{0j} + \beta_{1j} t_i), j = 2, \dots, r - 1$$
(10)

with

$$N_{ij}^* = \left(N_i - \sum_{k=1}^{j-1} n_{ik}\right)$$

and conditional probabilities

$$p_{ij}^* = G(\beta_{0j} + \beta_{1j}t_i), \qquad j = 1, \dots, r - 1$$
 (11)

Candy⁶ uses GLM estimation routines in GLIM to fit these models assuming the n_{ij} are binomially distributed conditional on N_i for stage 1 and conditional on N_{ij}^* for stages $2,\ldots,r-1$. No code is provided for this estimation procedure in the original paper, but the model is straightforward to fit using the glm function in R with a model formula of the form cbind(count,total - N_star) ~ stage + stage:time - 1. The sequential model with stopping parameterisation is also implemented in VGAM::vglm [12] and I here make use of it to fit the model to the budworm data.

4 Results

4.1 Cumulative model with constant variance

No code was provided in [6] for the parameter estimation of the cumulative model. Parameters were estimated using a direct optimisation of the Poisson likelihood for (3), as well as with the R functions VGAM::vglm and ordinal::clm. The cloglog link model

Table 1. Parameter estimates for the cumulative model with constant variance. This table replicates results presented in the first two rows of Table 2 of [6]. Note that ordinal::clm uses a parameterisation $\alpha_j - \beta z_i$ for the linear predictor yielding a parameter estimate for β with the opposite sign than the other methods. The cloglog link model failed to fit using VGAM::vglm.

α_1	α_2	α_3	α_4	α_5	α_6	β	Link	Method
5.49	9.39	12.23	15.70	21.26	27.25	-0.05	logit	Original [6]
5.53	9.48	12.35	15.86	21.46	27.52	-0.05	logit	Roptim
5.47	9.36	12.21	15.67	21.22	27.19	0.05	logit	Rclm
5.47	9.36	12.21	15.67	21.22	27.19	-0.05	logit	Rvglm
3.32	5.85	7.72	10.03	13.46	17.52	-0.03	cloglog	Original [6]
3.48	6.10	8.05	10.45	14.01	18.23	-0.03	cloglog	Roptim
3.32	5.85	7.71	10.02	13.46	17.52	0.03	cloglog	Rclm
NA	NA	NA	NA	NA	NA	NA	cloglog	R∨glm

Table 2. Parameter estimates for the cumulative logit model with proportional variance. This table replicates results presented in the first row of Table 1 of [8] and the last row of Table 2 of [6].

a_1	a_2	a_3	a_4	a_5	a_6	b^2	Method	Eqn.
121.080	204.360	264.410	342.473	465.620	599.570	1.559	Original[8]	6
120.000	204.700	264.600	341.300	464.500	595.700	1.412	SAS NLIN	6
120.033	204.659	264.586	341.285	464.480	595.690	1.412	R optim	6
α_1	α_2	α_3	α_4	α_5	α_6	β	Method	Eqn.
101.000	172.200	222.700	287.200	390.900	501.300	-0.842	Original [6]	3
100.990	172.181	222.598	287.134	390.771	501.157	-0.841	R optim	3

Table 3. Parameter estimates for the sequential model with stopping ratios. This table replicates results presented in Table 3 of [6]. The cloglog link model failed to fit using VGAM::vglm.

Parameter	eta_{-1}	eta_{-2}	β_{-3}	β_{-4}	$eta_{ extsf{-}5}$	β_{-6}	Link	Method
$\begin{array}{c} \beta_{0j} \\ \beta_{0j} \\ \beta_{0j} \end{array}$	10.410	12.960	12.020	11.160	17.700	33.730	logit	Original [6]
	10.410	12.959	12.020	11.165	17.698	33.726	logit	R glm
	10.410	12.959	12.020	11.165	17.698	33.726	logit	R vglm
$eta_{1j} \ eta_{1j} \ eta_{1j} \ eta_{1j}$	-0.085	-0.062	-0.046	-0.033	-0.038	-0.056	logit	Original [6]
	-0.085	-0.062	-0.046	-0.033	-0.038	-0.056	logit	R glm
	-0.085	-0.062	-0.046	-0.033	-0.038	-0.056	logit	R vglm
$eta_{0j} \ eta_{0j} \ eta_{0j} \ eta_{0j}$	7.350	8.530	9.120	8.440	10.090	16.300	cloglog	Original [6]
	7.347	8.537	9.124	8.442	10.087	16.298	cloglog	R glm
	NA	NA	NA	NA	NA	NA	cloglog	R vglm
$eta_{1j} \ eta_{1j} \ eta_{1j} \ eta_{1j}$	-0.065	-0.044	-0.037	-0.026	-0.023	-0.029	cloglog	Original [6]
	-0.065	-0.044	-0.037	-0.026	-0.023	-0.029	cloglog	R glm
	NA	NA	NA	NA	NA	NA	cloglog	R vglm

failed due to numerical errors when using the vglm function. Parameter estimates were close to those of the original study for the two R packages, and differed slightly for the optim method (Table 1), the latter exhibiting a noticeable sensitivity to the choice of starting values.

4.2 Cumulative model with proportional variance

The original SAS code provided in [5] required minimal updates to run in a contemporary version of SAS (SAS 9.4). Translating the model code to R was straightforward once I took the decision to not reimplement the IRNLS approach, but instead implemented a direct minimsation of the negative log likelihood with optim. Parameter estimates from SAS NLIN and R optim (Table. 2) were virtually identical, but differed slightly from the parameter estimates presented in [8], which was assumed to be the original source for the parameter estimates, as no actual parameter estimates were presented in [5]. Based on these three sets of parameter estimates it was also possible to redraw two figures from [5]. Figure 1 and 2, respectively, show that despite minor parameter differences there is an overall good agreement between the original results and the replication.

4.3 Sequential model

No code was provided in [6] for the parameter estimation of the sequential model. Parameters were estimated using the base R glmfunction, as well as with the VGAM::vglmfunction. The cloglog link model failed due to numerical errors when using the vglmfunction. Parameter estimates were identical to the original study (Table 3).

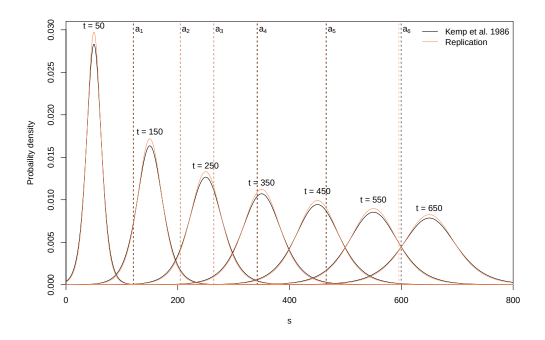


Figure 1. Logistic PDF of the Dennis, Kemp, and Beckwith⁵ model plotted for seven fixed values of t. Area under the PDF between a_{j-1} and a_j gives the expected proportion of insects in stage j at time t. Values of a_j and b^2 used in the graph are the estimates given in Table 1 of [8] (black lines) and the estimates from the replication (red lines). This figure replicates Figure 2 in [5].

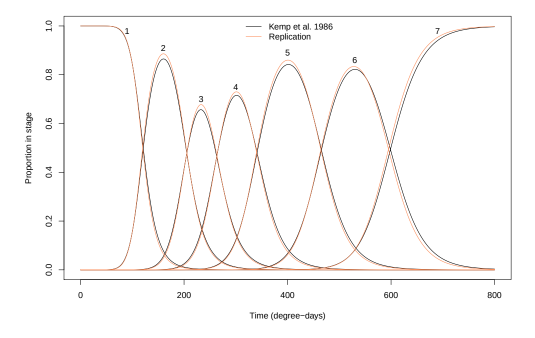


Figure 2. Expected proportion of insects in stages 1-7 plotted as functions of time t. Values of a_j and b^2 used in the graph are the estimates given in Table 1 of [8] (black lines) and the estimates from the replication (red lines). This figure replicates Figure 3 in [5].

5 Discussion

Overall the results from both [5] and [6] could be replicated closely.

The SAS code provided in [5] required only minimal updates to run in a contemporary version of SAS (SAS 9.4) and produced virtually identical estimates as the R reimplementation. These estimates, however, differed slightly from the parameter values reported in [8]. Given that the same initial values were used in all implementations, I believe that this disagreement is most likely caused by the inconsistencies in the published data set described in Section 2. The corrections applied to the data by [6] result in a data set that is internally consistent but potentially different to that on which the estimates in [8] are based.

No code was provided in [6]. However, the mathematical and verbal descriptions of the models were detailed enough to re-implement the estimation procedure for all models in R. GLIM code of the cumulative model with proportional variance was available from an earlier manuscript[9]. This allowed me to use the same initial values as the original study for this model. Initial values for the model with constant variance had to be guessed. The direct optimisation of the likelihood is sensitive to the choice of initial values, and this may explain the difference in the parameter estimates between the different methods. Another factor may be slight differences in the numerical implementation of the inverse link functions. Naive implementations of both the inverse logit and inverse complementary log-log function suffer from numerical underflow and/or overflow. The GLIM code from [9] uses multiple thresholding steps during the calculation of the linear predictor to mitigate against this, whereas the R implementation makes use of a single thresholding step in the inverse link function (gtools:inv.logit and VGAM::clogloglink, respectively). The numerical instability of the complementary log-log link may have contributed to the failure to fit either of the two complementary log-log models using the VGAM::vglmfunction. However, despite several troubleshooting attempts the exact reason for the numerical issues leading to this failure could not be established.

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