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A Computer Model for Simulating Population Development of the Indianmeal Moth (Lepidoptera: Pyralidae) in Stored Corn

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ABSTRACT  The Indianmeal moth, Plodia interpunctella (Hübner) (Lepidoptera: Pyralidae), is a common pest of stored corn, Zea mays L. We developed a computer model to simulate population development of the Indianmeal moth in stored corn by using previously published data describing immature developmental times and survivorship, and adult longevity and fecundity. The model accurately simulated population development of Indianmeal moths in corn stored during fall and into winter of three separate storage seasons in South Carolina. This is the period when the Indianmeal moth is a pest in stored corn in South Carolina. The model predicted that populations would increase after winter as grain temperatures rose, but observed populations in the grain bins never increased after winter. Despite this, the model should be useful from a management perspective because the corn is being sold off or used up after winter, and the observed Indianmeal moth populations never reached damaging levels after winter.

KEY WORDS  Plodia interpunctella, simulation model, stored-product insect

The Indianmeal moth, Plodia interpunctella (Hübner) (Lepidoptera: Pyralidae), is a common pest of stored corn, Zea mays L. (Arbogast and Throne 1997). Adults are short-lived, and a female can lay several hundred eggs (Mohandass et al. 2007). Larvae feed on kernels near the surface of the grain mass, and silken webbing produced by larvae can interfere with grain handling procedures when there are large infestations. Last-instar larvae move out of the grain to find a place to pupate, often on the surface of the grain.

Computer models can be useful for optimizing control strategies for pests of stored grain or for making real-time pest management decisions (Throne 1995). For example, Arthur et al. (2001) used a model for the maize weevil, Sitophilus zeamais Motschulsky (Coleoptera: Curculionidae), to determine feasibility of aeration (blowing cool air through grain to lower the temperature) for control. Models for simulating population development of insect pests of stored wheat have been incorporated into an expert system that inputs real-time insect sampling data and grain abiotic conditions, uses the models to predict insect population growth, and then uses a rule base to make management recommendations (Flinn et al. 2007).

Several types of simulation models have been developed for the Indianmeal moth. Briggs et al. (2000) developed a model for investigating basic ecological phenomena in the Indianmeal moth, but the model was not intended for pest management uses and does not include abiotic inputs such as grain temperature and moisture. Na et al. (2000) developed a phenology model for predicting peaks of moth emergence. Kaliyan et al. (2007) developed a model for predicting mortality of Indianmeal moths at cold temperatures during winter.

The objective of our study was to develop a computer model that could be used to simulate population development of the Indianmeal moth in stored corn from the initiation of storage (usually around September in the southeastern United States) to winter. This is the period when the Indianmeal moth is a pest in stored corn in the southeastern United States. Populations usually are at low levels after winter (Arbogast and Chini 2005), so the Indianmeal moth is not a pest after winter. The eventual plan is that this model, along with models for other pests of stored corn (e.g., Maier et al. 1996), will be incorporated into an expert system, such as that developed for stored wheat, Triticum aestivum L. (Flinn and Hagstrum 1990), to aid in decision making for management of insect pests in stored corn.

Materials and Methods

Model Development. Methods used for model development were as in Throne (1989). We used data from Arbogast (2007a) for a wild strain of Indianmeal moth from stored corn to develop equations describ-
ing the relationship between temperature and egg-to-
adult development (Table 1). The data were collected
at 20–35°C, and we extrapolated them to 10°C. Above
35°C and below 10°C, development rate was set to that
at 35 and 10°C, respectively. Sex did not affect develop-
ment rate, and we did not include moisture content
in the equation because there was no apparent pattern
to changes in developmental rate with moisture con-
tent (Arbogast 2007a). Survivorship from egg to adult
did not differ with temperature or moisture content at
20–30°C and averaged 54% (Arbogast 2007a). Above
30°C, we used linear regression to predict out to no
survivorship at 40°C [survivorship = 3.36 – (0.084 ×
temperature [°C])]. Below 20°C, we used linear re-
gression to predict out to no survivorship at 0°C (sur-
vivorship = 0.042 × temperature [°C]).

No published data are available describing longevity
of adult males. Brower (1976) showed no differ-
ce in longevity of males and females at 27°C and 60%
RH. Although not compared statistically, longevity of
males and females did not seem to differ at 28°C and
65% RH in a study by Huang and Subramanyam
(2003). So, we used female longevity data from Ar-
bohydrate (2007b) to describe longevity of males. The
relationship was linear from 20 to 35°C (Table 1), and
we extrapolated the equation to 10 and 40°C. Above
40°C, we used the same longevity as at 40°C; below
10°C, longevity was set to 20 d.

The female stage was split into young and old fe-
males, based on data from the longevity study by
Arbogast (2007b). Young females were considered to
be those up to the age where 75% of eggs are laid.
There are no data to indicate that moisture content has
an effect on longevity, so we used a linear equation to
predict young female longevity (Table 1). Above
40°C, longevity was set as at 40°C; below
15°C, longevity was set to 7 d. Longevity of old adult
females was set to 2.7 d.

Fecundity data are from Arbogast (2007b). A qua-
Dratic equation was used to describe the data (Table
1), which are extrapolated from 10 to 40°C. Effects of
relative humidity on fecundity are simulated based on
Mbata (1985) using linear regression to reduce fecun-
dity when relative humidity is below 75% (% total
fecundity = 0.01262 × relative humidity).

Inputs for the model are the starting and ending
days of the simulation, with day 1 corresponding to 1
January and day 365 corresponding to 31 December of
nonleap years; numbers of eggs, adult males, young
adult females, and old adult females; and hourly tem-
peratures and relative humidities for the simulation
period. During simulation, development rate of each
of the above-mentioned stages was calculated based
on temperature and relative humidity. Individuals en-
ter a stage simultaneously at the start of a simulation,
but passage to the next stage is distributed over time
using a time-varying distributed delay (Manetsch
1976). The shape of the curve describing the number
of individuals leaving a stage over time was approxi-
mated by a member of the Erlang family of density
functions. The particular curve for each stage was
designated by the parameter K, which is calculated as
DEL²/s², where DEL and s² are the mean time
to development at a given temperature and the associ-
ated variance. K was calculated for a stage at each
temperature and then an average K for a stage across
all temperatures was used in the model, but K was not
allowed to exceed 12 times the shortest developmental
time or else the delay becomes unstable (Abkin and
Wolf 1976). Emerging adults were distributed at a 1:1
sex ratio (Arbogast 2007a). Number of eggs laid was
divided evenly over the young or old female stages at
a constant temperature.

Model Validation. We used data from Arbogast and
Chini (2005) to test the validity of the model. They
collected wandering-phase larvae in cardboard rolls
on the surface of stored corn weekly throughout the
storage season during 3 yr on two sites in South Caro-
olina. They also recorded hourly temperatures and
weekly moisture contents at nine locations on the
surface of the grain in each bin, and we used the mean
of those nine hourly temperatures or moisture con-
tents, after conversion to hourly relative humidity
(using the equation in Throne 1994), as abiotic inputs
for the model. Although our intent was to simulate
population development in the fall, we continued sim-
ulations throughout the storage season to determine
whether temperatures were responsible for limiting
population growth in spring. The amount of corn in
the bins was constant during the studies. The valida-
tion studies were terminated when the farmer wanted
to start removing corn from the bins.

We used data only from the Bamberg Co. site be-
cause there were too many missing data and some
anomalies in the insect data for the Barnwell Co. site.
There were occasionally missing temperature or mois-
ture content data for the Bamberg Co. site, and we
interpolated between adjacent values to compensate
for the missing data. We initiated the model with
young adult females. We determined the number of
young adult females to initiate simulations by trying
different initial numbers and then using the number

<table>
<thead>
<tr>
<th>Process</th>
<th>a</th>
<th>b</th>
<th>c</th>
<th>n</th>
<th>R²</th>
</tr>
</thead>
<tbody>
<tr>
<td>Egg-to-adult development (d)</td>
<td>355.5 ± 24.1</td>
<td>−21.52 ± 1.81</td>
<td>0.3557 ± 0.0328</td>
<td>24</td>
<td>0.93</td>
</tr>
<tr>
<td>Male longevity (d)</td>
<td>23.04 ± 0.759</td>
<td>−0.4860 ± 0.027</td>
<td>4</td>
<td>0.99</td>
<td></td>
</tr>
<tr>
<td>Young female longevity (d)</td>
<td>18.00 ± 0.000</td>
<td>−0.4000 ± 0.000</td>
<td>4</td>
<td>1.0</td>
<td></td>
</tr>
<tr>
<td>Lifetime fecundity</td>
<td>−1,639 ± 219</td>
<td>144.2 ± 16.4</td>
<td>−2.766 ± 0.298</td>
<td>4</td>
<td>0.99</td>
</tr>
</tbody>
</table>

All equations are of the form $Y = a + bX + cX^2$, where $X$ is temperature (°C) and $Y$ is the process being modeled.
that resulted in populations that best matched the observed data because we did not know the initial size or infestation dates of the infesting populations in the bins. We determined validity of the model by determining whether the simulated numbers of adults were within the 95% CL on mean number of pupae found in cardboard rolls. This introduces some error, but numbers of pupae in cardboard rolls were determined in the field study, whereas our model predicts egg to adult development because effects of temperature and moisture content on development of individual Indianmeal moth stages has not been determined.

Results

Simulated numbers of adults emerging between 31 October 1990 and 20 February 1991 were within observed 95% CL except on the first sampling date when simulated number of adults emerging was lower than the observed lower confidence limit by less than one adult (Fig. 1). After 20 February (data not shown), grain temperature began to rise and the simulated numbers of adults emerging began to increase, but the population in the bin never increased through the rest of study which ran into summer.

Simulated numbers of adults emerging between 20 September 1992 and 21 February 1993 were within observed 95% CL except on 6 December when simulated number of adults emerging was higher than the observed higher confidence limit by less than one adult (Fig. 3). After 21 February (data not shown), grain temperature began to rise and the simulated number of adults emerging began to increase, but the population in the bin never increased through the rest of study which ran into summer.

Discussion

The model fit the data well through the early growth phase of the population in fall and into winter when Indianmeal moths are a problem in corn storages in South Carolina. The fit was best during the first and third storage years, with simulated numbers differing from observed numbers on only one date and by less...
than one individual. During the second storage year, the fit was not as good early in the season with the model overpredicting population growth. We assumed in the simulations that female moths invaded the bins on the first day of storage, and this may not have always been the case. This may account for differences between simulated and observed population levels early in the second storage year.

The observed populations never increased in the bins in the spring despite favorable abiotic grain conditions (Arbogast and Chini 2005). Arbogast and Chini (2005) hypothesized that this may have been due to adversely low temperatures in winter killing the insects, adversely high headspace temperatures in the spring killing insects on the grain surface, and natural enemies. There was no evidence for diapause in the South Carolina populations (Arbogast 2005), although diapause is known in Indianmeal moths (Kikukawa et al. 2009). Simulating populations during spring and early summer is less of a concern from a management perspective because the corn is being sold off or used up during this period, and the observed Indianmeal moth populations never reached damaging levels during this time. However, further studies to clarify what happens to limit growth of the Indianmeal moth populations during this time would be interesting. Indianmeal moth populations in a botanical warehouse followed the same pattern in North Carolina, with large populations in fall, populations declining in winter, and the populations not increasing in size in spring (Arbogast et al. 2002).

Although we had to make some assumptions about affects of extreme temperatures on population growth because of lack of data at these temperatures, that the simulated data fit the observed data well indicates that our assumptions were reasonable or that population development at these temperatures contributes little to population dynamics during fall and into winter. The apparent validity of the model will make it useful for helping to optimize insect pest management in stored corn. As we develop models for other insect pests of stored corn, these models will be used to develop an expert system to aid in decision making for management of insect pests in stored corn.

Acknowledgment

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