Mummy Berry Fruit Rot and Shoot Blight Incidence in Blueberry: Prediction, Ranking, and Stability in a Long-term Study

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Abstract. Mummy berry (Monilinia vaccinii-corymbosi) is an important disease of cultivated blueberry (Vaccinium spp.). The disease has two distinct phases: a blighting phase initiated by ascospores and a fruit infection stage initiated by conidia during bloom. In this study, we investigated, in a nursery setting, blueberry cultivar resistance to both phases of the disease and, using multiple “standards” with a range of susceptibilities, examined, over 9 to 12 years, factors affecting disease incidence in controlled inoculations. The analyses of our data show that a minimum of 8 years of testing is necessary to obtain stable rankings of cultivar susceptibility for the fruit infection phase of the disease. Insufficient years of data were available to estimate this for the blight phase. Eight years are necessary largely as a result of uncertainty arising from the large environment × genotype interaction, estimated to be more than double any other source of observed variation, other than that resulting from sampling/individual plants. For individual cultivars, temperature and the amount and frequency of precipitation in January to March (when neither plant nor pathogen were presumed active and when both were in cold frames somewhat protected from environmental conditions) were predictive of later disease incidence. For most cultivars, the same weather variables at the same time period were found to be predictive for independently modeled cultivars. Additional cultivars, with only a few years’ data, were grouped with the standard with which they shared similar environmental (year) responses and possibly similar disease predictive models.

Mummy berry is one of the most important diseases of blueberry in North America (Eck and Childers, 1966). The disease affects all cultivated species, including highbush (Vaccinium corymbosum L.), lowbush (V. angustifolium Aiton), and rabbiteye (V. ashei Reade = V. virgatum Aiton) (Hildebrand et al., 1995). Wild species of blueberry such as V. myrillioides Michaux and V. elliottii Chapman are also susceptible (Hildebrand et al., 1995). The disease is caused by the fungus Monilinia vaccinii-corymbosi (Reade) Honey. There are two distinct infection phases, each of which must be completed for the disease to persist. The primary phase results in blighting of shoots and sometimes flower clusters. This phase is initiated by germination of pseudosclerotia that form apothecia (Batra, 1983). The ascospores produced by the apothecia are wind-dispersed to young vegetative blueberry tissues during and shortly after bud break (Cox and Scherrn, 2001). Conidia, produced on the blighted tissues, are dispersed to the stigma and surface of open flowers (Batra and Batra, 1985). The conidia germinate and grow down the stylar canal and colonize the ovaries. The fungal thallus that grows within the infected fruit forms pseudosclerotia that fall to the ground and serve as the overwintering stage.

Several early studies documented levels of resistance to M. vaccinii-corymbosi in limited collections of highbush blueberry cultivars (Nelson and Bittenbender, 1971; Pepin and Toms, 1969; Varney and Stretch, 1966). The most methodical screening for blight resistance (primary phase) was an evaluation of 52 highbush blueberry cultivars using concentrated inoculum in a nursery-type setting (Stretch et al., 1995) that identified several cultivars with good horticultural quality and high levels of resistance. In a later study, Ehlenfeldt et al. (1997) manipulated shoot elongation of resistant and susceptible cultivars to evaluate the nature of resistance. In susceptible cultivars, susceptibility was highest at short shoot lengths (i.e., young shoots) and decreased as shoot length increased (more mature shoots). Among resistant cultivars, there were indications of resistance resulting from avoidance (through delayed bud break) and biochemical or structural factors. Lehman and Oudeman (1997b) similarly demonstrated that susceptibility decreased as shoots elongated, suggesting the development of ontogenetic resistance. Ehlenfeldt et al. (1996) evaluated cultivar differences in 48-cultivar subset of those evaluated by Stretch et al. (1995) and found a significant positive correlation (r = 0.74) between blight incidence and average shoot length during the first week of ascospore release. Fruit susceptibility in the mummy berry disease cycle was also investigated by Stretch and Ehlenfeldt (2000). No correlation was found between blight resistance and stress, frost damage, and other environmental factors, which could affect disease outcome.

In this study, we compiled data from many years of screening blueberry cultivars for resistance to both phases of mummy berry disease. In developing resistance rankings, we found that even the cultivars included as standards varied in response from year to year and occasionally shifted ranks. To determine the number of years necessary for stable rankings, we decided to more fully analyze the response of the standards and better quantify the components of disease resistance beyond simply plant genotype. We applied variance decomposition methods to assess the contributions of: cultivar (genetic factors), year (weather and other environmental factors), and their interaction. As part of year and year × cultivar interaction, we investigated if weather variables such as temperature, rainfall, and humidity were useful predictors of disease incidence. The year × cultivar interaction is typically considered to be an “unpredictable” part of the cultivar × environment interaction (Lin and Binns, 1988). However, we found that weather variables from earlier in the year, in part, could explain year-to-year ranking shifts among cultivars. Thus, unique predictive equations were used for average disease incidence, based on weather variables and the cultivar’s long-term mean resistance, may be derived for each cultivar.

Materials and Methods

Blight screening materials

Based on data from earlier screenings (Ehlenfeldt and Stretch, 2000; Stretch and Ehlenfeldt, 2000; Stretch et al., 1995), the following cultivars were selected as shoot blight standards to represent both highbush (HB) and rabbiteye (RE) genotypes and a broad range of responses from low to high. The cultivars that were included across 1996.
to 2007 were: ‘Bluejay’ (HB), ‘Brightwell’ (RE), ‘Callaway’ (RE), ‘Coastal’ (RE), ‘Coville’ (HB), ‘Northblue’ (HH, a half-high type, i.e., a lowbush × highbush hybrid), ‘Sunrise’ (HB), and ‘Toro’ (HB). ‘Climax’ (RE) was included from 1996 to 2006. An additional 36 cultivars were coscreened for 3-year periods and analyzed against the shoot blight standards (Table 1).

Fruit infection screening materials
Similarly, six cultivars were selected as fruit infection standards based on data from earlier screenings (Stretch and Ehlenfeldt, 2000) and because they represented a range of responses from low to high. The cultivars that were included across 1995 to 2007 included: ‘Atlantic’ (HB), ‘Bluejay’ (HB), ‘Blueray’ (HB), ‘Northsky’ (HH), ‘Rancocas’ (HB), and ‘Sierra’ (HB). An additional 32 cultivars were coscreened for 3-year periods and analyzed against the fruit infection standards (Table 2). Detailed background and pedigree information on the cultivars used in either screening test may be obtained by request from the authors. Additional information on some cultivars is available in the Germplasm Resources Information Network online database (GRIN; http://www.ars-grin.gov/npgs).

Blight screening methods
The tests were conducted outdoors in randomized complete block designs with five replicates (potted plants) per cultivar. Pots were arranged in a grid pattern in each replicate with the specific dimensions of the grid determined by the number of entries (cultivars) and the space allocated each year. The space between plants averaged 0.45 m and blocks were laid out linearly. The plants were potted in 2.84-L pots in a sand:peat mix (1:1). Plants for blight evaluation were typically 30 to 60 cm and were 5- to 6–year-old plants that had been cut back and allowed to regrow several times. The M. vaccinii-corymbosi pseudoclerotia (mummies) used in each test were collected from the P.E. Marucci Center for Blueberry and Cranberry Research, Chatsworth, NJ. Pseudoclerotia were collected in September and October from the field, transplanted into soil-filled (1:1 sand:peat mix) 0.92-L plastic pots, and overwintered in plastic-covered cold frames. At the start of each test (early April), pots containing pseudoclerotia with emerging apothecia were removed from the covered cold frames and placed at evenly spaced intervals within and around the periphery of the experiment area to provide an even distribution of inoculum. To enhance the infection process, the entire plot received supplemental misting for 1 h every evening during ascospore release (applied volume ≈5 mm/d).

Trials were typically initiated the first week of April with the timing determined by initiation of shoot development in the earliest emerging cultivars. Until the start of each test, the plants were sheltered in well-ventilated, plastic-covered cold frames. During the test, plants were examined for blighted shoots on a weekly basis between early May, when symptoms became visible, and late May/early June, when blighting was essentially complete. At each inspection, blighted shoots were tabulated and removed. At the end of the blight period, remaining healthy shoots were counted on each plant so that the percentage of blighted shoots could be calculated. Cultivars to be evaluated were included for 3 years and sometimes as many as 4 to 6 years if widely variable results were observed.

Fruit infection screening methods
The plants were maintained in 2.84-L pots in a sand:peat (1:1) planting mix. The test was typically conducted in a cold frame covered with shadecloth (55% transmission) as a randomized complete block design of five replicates (potted plants) per cultivar. Plants for fruit infection evaluation were typically 50 to 80 cm and were also 5- to 6–year-old plants that had been cut back and allowed to regrow several times. The space between plants averaged 0.45 m between rows and 0.36 m between plants within rows.

In all years, blighted plants of ‘Bluehaven’ and ‘Bluejay’ (also in 2.84-L pots) with surrounding tissue were evenly distributed between every two rows of plants to serve as inoculum sources located within 0.5 m or less of the plants to be tested. In previous studies, ‘Bluehaven’ and ‘Bluejay’ had averaged 63.9% and 35.6% blight, respectively, and had ranked as two of 47 and 11 of 47 with percentages of blight, respectively, obtained from the Oregon Plant Pathology database (P. Huttel et al., 1996). Additional

| Table 1. Means and sds of the blighted proportion (disease incidence) of shoots for cultivar standards and cultivars with a similar yearly disease incidence pattern. |
|-----------------|-----------------|-----------------|
| Cultivar standards for shoot blight | Blighted proportion of shoots mean and (so) | Cultivars with similar yearly responses |
| Bluejay (HB) 0.20 (0.23) | Bluebelle, Chanticleer, Clara, Cumberland, Echota, Pearl River, Putte, Sampson, Scammell, Tibblue |
| Toro (HB) 0.22 (0.24) | Clara, Goldtraube, Jubilee, Olympia, O’Neal, Triblue, Wolcott |
| Coastal (RE) 0.36 (0.27) | Bluebelle, Collins, Triblue |
| Callaway (RE) 0.37 (0.32) | Ascorba, Bluegem, Bluebelle, Chandler, Clara, Cooper, Olympia, Pender, Santa Fe, Wolcott |
| Sunrise (HB) 0.45 (0.32) | Ascorba, Bluegem, Britolblue, Chandler, Ethel, Friendship, Misty, O’Neal, Ozarkblue, Pender, Santa Fe, Triblue |
| Coville (HB) 0.58 (0.35) | Black Giant, Denise Blue, Goldtraube, O’Neal, Powderblue, Sunshine Blue |
| Baldwin (RE) 0.60 (0.38) | Clara, Friendship, Goldtraube, Misty, O’Neal, Putte, Sampson, Suwannee, Triblue |
| Brightwell (RE) 0.62 (0.35) | Clara, Collins, Denise Blue, Goldtraube, Premier, Suwannee, Triblue |
| Northblue (HH) 0.77 (0.24) | Concord, Jewel, Reka, Triblue |

Table 2. Means and sds of the infected proportion (disease incidence) of berries for cultivar standards and cultivars with a similar yearly disease incidence pattern.

<table>
<thead>
<tr>
<th>Cultivar standards for fruit infection</th>
<th>Infected proportion of berries—mean and (so)</th>
<th>Cultivars with similar yearly responses</th>
</tr>
</thead>
<tbody>
<tr>
<td>Northsky (HH) 0.02 (0.03)</td>
<td>Collins, Croatian, Harrison, June, Morrow, Murphy, Northblue, Windsor</td>
<td></td>
</tr>
<tr>
<td>Bluejay (HB) 0.10 (0.11)</td>
<td>Cabot, Cape Fear, Croatian, Friendship, Nelson, Northblue, Oranblue, Polaris, Reka, Sunrise, St. Cloud</td>
<td></td>
</tr>
<tr>
<td>Rancocas (HB) 0.22 (0.22)</td>
<td>Bluetta, Blue Rose, Bonifacy, Echota, Gila, Goldtraube, Northblue, Zuckertraube</td>
<td></td>
</tr>
<tr>
<td>Blueray (HB) 0.35 (0.30)</td>
<td>Elizabeth, Elliott, Grover, Ivanhoe, Katherine, Legacy, Wareham</td>
<td></td>
</tr>
<tr>
<td>Sierra (HB) 0.41 (0.29)</td>
<td>Black Giant, Denise Blue, Goldtraube, O’Neal, Powderblue, Sunshine Blue</td>
<td></td>
</tr>
<tr>
<td>Atlantic (HH) 0.53 (0.26)</td>
<td>Clara, Collins, Denise Blue, Goldtraube, Premier, Suwannee, Triblue</td>
<td></td>
</tr>
</tbody>
</table>

Standards for fruit infection incidence have 11 to 12 years of data. Standards are ordered from most resistant to least resistant. Similarity does not imply similar levels of disease incidence, but rather a high correlation of response for years in common. Some of the standards were similar to each other, so cultivars may be similar to more than one standard. RE = rabbiteye; HB = highbush; HH = half-high.
plants of ‘Bluehaven’ and ‘Bluray’ were added over time, as necessary, to maintain inoculum levels. To facilitate transfer of conidia from the blighted tissue to the stigmata, two hives of honeybees were placed in the netted cold frame, one at each end of the experimental area. To ensure that the resultant fruit developed only under the imposed conditions of the experiment, open flowers were removed from the test plants before setting the inoculum plants and bees in the cold frame. The experimental area was sprinkler-irrigated (22 mm/d) every evening during conidia production. Fruit were picked while still green, 4 to 6 weeks after flowering, and stored at 2 °C until evaluated. To evaluate for infection, each fruit was sliced in half crosswise and observed for the distinctive white fungal growth pattern of the mummy berry pathogen. For purposes of analysis, the incidence of mummy berry fruit infection was expressed as an uninfected proportion of berries. As with blight screening, cultivars were typically evaluated for 3 years, but sometimes longer if widely divergent results were observed within a cultivar.

**Statistical methods**

**Minimum estimate for reliable resistance ratings.** To determine the number of years of testing necessary to obtain reliable estimates of percent infection, we had data available from various cultivars per year over an 11- to 12-year period for all data for that cultivar) and the average of years in the model decreased. As a measure of change, we calculated the mean square error (variance + bias squared) for the mean (the logit of the proportion of uninfected observations). The bias is the difference between the estimate for all years (the mean for all data points) and the average estimate from models with 1 or more years of data dropped. To determine whether the variance estimate for total years differed significantly from model-based estimates from resampling, we asked “where in the distribution of model-based variance estimates from resampling do the total years estimate lie?” i.e., was it in the bottom 5% or above this value? Recall that the average model-based variance estimate from resampled data sets was always greater than the total years estimate.

**Weather variables.** Several weather variables were evaluated as a likely source of year-to-year variation in infection incidence. Available weather data included: daily minimum-, maximum-, and mean- temperature, daily precipitation, and relative humidity (U.S. National Oceanic and Atmospheric Administration). Weather variables were assessed over the period of 1 Jan. to 30 May (ordinal Days 1 to 150). Growing degree-days [average of the daily maximum and minimum temperatures compared with a 7.2 °C (45 °F) base temperature] were calculated from these data and evaluated along with the weather variables.

Variables were grouped by 10-d increments (i.e., mean maximum temperature for ordinal Days 1 to 10, ordinal Days 11 to 20, and so on) because this produced the best compromise between resolution and a meaningful average (i.e., gave the highest proportion of significant correlations). Five-d and 20-d increments were also initially evaluated. For each of the cultivar standards, the correlation between the mean disease incidence (on both the logit and arcsine square root-transformed scales) and each of the weather variables for each 10-d increment was calculated. If at least two cultivar standards had significant correlations with the weather variable for that time period, we investigated the relationship further by looking at scatterplots of disease incidence versus weather variable to verify that the correlation was not the result of one or two outliers and that the relationship between the weather variable and other (nonsignificant) cultivars was not grossly different. This analysis provided us with a pool of candidate regressors.

We then performed stepwise regression separately for each cultivar (evaluating both the logit and arcsine square root-transformed scales). The resulting model was inspected and occasionally reduced to a border-line significant weather variables whose influence on the growth model was not necessarily similar in average disease incidence over years for the standards. For those with at least 3 years of data in common with one of the standards, we tested the correlation between the two.

We accepted any cultivar as being similar to a standard in yearly pattern (although not necessarily similar in average disease incidence) if the correlation coefficient between the two was statistically significant (P < 0.05) and positive.

**Results**

**Disease incidence in the standards.** We first analyzed the fruit infection results because this data set had more years of data available. Preliminary line plots of fruit disease incidence over years for the standards were roughly parallel across some periods of years, but shifted ranks in others, indicating existence of a genotype-by-environment interaction (Fig. 1). To determine the number of years necessary to obtain reliable estimates of percent disease, we used the data for six cultivars over an 11- to 12-year period for assessing incidence of fruit infection. The analysis was conducted separately for each cultivar. We present the results for ‘Atlantic’ (ATL) and ‘Northsky’ (NOS) as examples (Fig. 2A–B).

For ATL, the 11-year estimate (all estimates are on the logit scale; the logit scale is log (1 – p), where p is the estimate of...
The variance of the mean is \( y = 0.13 \) (horizontal dotted line on graph). This is the model “population” estimate. The solid line gives the mean square error (MSE). The figure also shows the sum of the squared bias and the variance of estimates of the mean from resampling. The contribution of the squared bias is small. For each number of years resampled, we also plotted the average estimate of the variance of the mean from modeling (each time we resampled the data, we modeled it and estimated the variance of the mean, then averaged those estimates) referred to subsequently as model estimates. Note that the MSE from resampling for 4 or more years is smaller than the model estimates for partial years, as would be expected, because the former is based on resampling a “population” of 11 years, whereas the latter is attempting to estimate this parameter for an infinite population of years.

Several important results are shown in Figure 2A. The curve of the MSE does not flatten appreciably until \( \approx 8 \) years of data are available. With only 2 years of data, the variance of the mean (often reported as its square root, i.e., the SE of the mean) estimated by the model is considerably smaller than the estimate from resampling. Thus, with only 2 years of data, the SE of the mean from a model estimate would be misleadingly small, which would result in differences between cultivars being assigned far too liberally. This was also true for 3-year estimates for all standards, except ATL.

The average of the model estimates for partial years for ATL does not show any indication that it has reached an asymptote at 10 years, so we assume an asymptote is still several years beyond this. At 8 to 10 years, the 11-year estimate was above the bottom 5% of the model estimates from resampling (data not shown). At 7 years, the 11-year estimate crossed into the bottom 5% (data not shown), suggesting that 8 years is the minimum number of years that can be used for a reliable resistance estimate for this cultivar. For less than 7 years, the distribution of the model estimates of the variance from resampling widens such that the 11-year estimate is no longer found in the tail of the distribution.

The patterns for fruit infection in the remaining five standards were very similar to ATL, except for NOS, in which the resampling variances for 2 or 3 years of data were very large (Fig. 2B). For this cultivar, only the variance estimates from resampling are depicted for 2 and 3 years of data. The model estimates are not shown; they were so large as to be judged unreasonable (that is, anyone analyzing these data would reject those modeling results). This occurred because NOS is a resistant cultivar and there were many infection-free plants on the logit scale, this value (one) gets pushed toward infinity, so variance estimates of the mean are inflated. For ‘Blueray’, the 12-year variance estimate crossed into the tail of the model estimates from resampling at 7 years (suggesting a minimum of 8 years is necessary). For ‘Rancocas’, the 12-year variance estimate crossed into the 5% tail at 5 years (but was below the 10% tail starting at 8 years). For the other two cultivars (Bluejay and Sierra), the distribution of the resampling variance estimates never became sufficiently narrow for the total years variance estimate to fall into the tail.

Results for blighting were similar (not shown) for the nine standards with 8 years of data. Overall, variances tended to be larger.
with the same gross underestimates of the model-based variance of the mean with only 2 years of data. The larger variances showed that resistance to shoot blight was more variable than years means was fruit infection. In only one cultivar, Sunrise, did the 8-year variance estimate fall into a 10% tail for 5 and fewer years (i.e., this would suggest that for ‘Sunrise’, 6 years of evaluation are sufficient to estimate a variance based on 8 years of data). The distributions of the resampling variance estimates were generally wider for these data than for the fruit disease incidence data, so statistical power was low.

Predictive weather variables. Weather variables were evaluated for possible correlations with disease incidence for both shoot blighting and fruit infection in each standard cultivar. We give results for these models in Tables 3 and 4, respectively. These models show that disease incidence in the spring can be well-predicted based on weather conditions from earlier that same year, although there are differences among the cultivar models.

For shoot blighting (Table 3), the most important predictors were precipitation weather variables, either rain frequency, \( P_t \) (on ordinal Days 91 to 100, i.e., end of March), or average amount of precipitation, \( P_{avg} \) (on ordinal Days 21 to 30, i.e., end of January) based on arcticise square root-transformed data. These models have high \( R^2 \) values ranging from 0.50 for ‘Toro’ to 0.95 for ‘Baldwin’ and ‘Sunrise’ and therefore should do a good job predicting average disease incidence. These values show a general trend of \( R^2 \) being higher for more susceptible cultivars, suggesting better predictability of disease incidence in such cultivars. For mummy berry fruit infection (Table 4), the common predictive variable for all models is the average maximum temperature across ordinal Days 51 to 60 (late February). For fruit infection, \( R^2 \) values had less predictive value than for shoot blight, ranging from 0.31 for ‘Atlantic’ to 0.76 for ‘Bluejay’, and did not show any trend with disease incidence ranging.

Variance decomposition. The sources of variation in the model were cultivar (genotype), year (environment), cultivar \( \times \) year (genotype \( \times \) environment interaction), and plant-to-plant variation + residual (sampling error and any other source of variation not previously considered) (Table 5). For each cultivar, the year-to-year variation can be further decomposed into that part resulting from weather and that part resulting from other year-to-year effects, because the \( R^2 \) values in Tables 3 and 4 for each standard estimates the ratio of variation explained by the model (weather variables) to the total year-to-year variation.

The variance component estimates for the two disease phases were similar. The largest variance component was that resulting from the sum of plant-to-plant variation, sampling error, and other unknown sources. Although we did not separate the sampling error variance from other sources of variation for this component, sampling error is readily calculated for each sample, because it depends only on \( n \), the size of the sample (number of berries or shoots), and the parameter, \( p \), estimating disease incidence. It is small when \( n > 100 \) and is likely responsible for less than 10% of this variance component for these data, because \( n \) was typically greater than 50 (on the ascemic transformation scale, sampling error variance is \( \approx 0.005 \) for \( n = 50 \) with a disease incidence of 50%). The cultivar main effect is relatively small and, for both phases of the disease, the cultivar \( \times \) year interaction was larger than either of the two main effects (cultivar and year).

Ranking of standard cultivars for disease incidence. The standards, with many years of data, are ranked in Tables 1 and 2. Although these rankings are based on ascemic square root-transformed data, they are reported in the table as percentages. \( \hat{p} \) are slightly larger for shoot blight data, because they are based on fewer years than the fruit infection. Because there were only a few years of data on all other cultivars not used as standards, a ranking of them would not be stable. Cultivars with yearly patterns similar to one of the standards are given in Tables 1 and 2.

### Discussion

Our analysis of fruit disease incidence in the standard cultivars over 11 years indicated significant and large genotype-by-environmental interactions. This means that rankings of resistance (fruit infection) are not stable from year to year and that the 2 or 3 years of data typically used for publication are insufficient to reliably estimate disease resistance. Because most cultivars are only tested for 2 to 3 years, this could lead to erroneous rankings depending on which years were sampled. There is a large body of literature on the stability of crop yield, both across locations and over years, and various definitions of stability (Lin and Binns, 1991), and our finding of a large cultivar-by-year interaction is not uncommon, although we measured disease incidence rather than yield.
Our question differs from that commonly asked in yield stability literature, because we wanted to know how many years of data would be necessary to obtain a good measure of year-to-year variability rather than which cultivars tend to produce similar yields over various locations and years.

Our data show the curve of the MSE takes several years to begin flattening, and even with 11 years of data, the curves did not flatten completely (this was true for all standards). The results suggest that for cultivars that do not vary excessively from year to year (i.e., the distribution of the model variance estimates from resampling is not too wide), 8 years of data are necessary for a resistance estimate that is roughly as good as an 11- or 12-year estimate and that fewer years may produce unreliable estimates. This assumes that the 8 years are a fair representation of all the years sampled in the inference period. Our blight disease incidence results showed larger variances over similar time ranges; hence, it is likely that to reliably evaluate resistance, more years of data would be necessary than were used for fruit infection estimates.

Because accumulating 8 or more years of infection data on large numbers of cultivars is not generally feasible, we sought to determine if weather data before and during manifestation of the disease could be used to help predict disease incidence occurring that same year. We found the most important predictor of blight to be either average amount of precipitation at the end of January or rain frequency at the end of March. The most important predictor for fruit infection was found to be the average maximum temperature in late February.

We have shown that one or more weather variables can be statistically linked to disease incidence. Our models have high $R^2$ values, suggesting they should do a good job predicting average incidence of the disease, but it was unclear how these factors are actually linked to disease incidence. Weather variables in the context of our screenings (except perhaps temperature) were not directly causal. Our plants and mummies were in covered cold frames until early April so that precipitation could not have had a direct impact. However, precipitation often is accompanied by decreased light, increased humidity, and increased soil moisture, and any of these factors, not directly measured, could influence some aspect of plant or fungus physiology and the subsequent disease incidence. Thus, the weather variables found useful in the modeling can be thought of as proxies for the true variables that affect infection.

For blighting, genetic effects among pseudoclerotia affect the infection process. Lehman and Oudemans (1997a) demonstrated mummy berry populations from localized areas were influenced by natural selection to adapt to better match the phenology of their host plant population. Similarly, fruit infection results from the interaction of many factors. Lehman et al. (2007) suggest differences in pollinator activity, attractiveness of cultivars to pollinators, and differences in cultivar phenology may all account for environmental differences between years.

Variance decomposition indicated that, for both phases of the disease, the cultivar × year interaction was larger than either main effect. Thus, not only will overall disease incidence (or pressure) and cultivar response differ from year to year based on prevailing weather conditions and other factors, but they will not vary in a consistent manner.

Despite year-to-year variability, it is possible to group cultivars that respond similarly to the established standards with the idea that, by using the weather variable models derived for each standard cultivar, we might now be able to better understand the year-to-year responses of these “similar cultivars.” Similarity of response does not imply similarity of resistance, but does imply that within years used for the comparison, these cultivars ranked consistently above or below the standard with which they are grouped (i.e., had a similar cultivar × year interaction). Thus, by defining groups that share similar responses to environmental factors (e.g., possibly weather variables), we may be able to improve predictions of disease incidence for individual cultivars within these groups.

Literature Cited


