Molecular Polymorphism and Phenotypic Diversity in the Eutypa Dieback Pathogen Eutypa lata

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ABSTRACT


Pathogen adaptation to different hosts can lead to specialization and, when coupled with reproductive isolation, genome-wide differentiation and ecological speciation. We tested the hypothesis of host specialization among California populations of Eutypa lata (causal fungus of Eutypa dieback of grapevine and apricot), which is reported from >90 species. Genetic analyses of nine microsatellite loci in 182 isolates from three hosts (grapevine, apricot, and willow) at three locations were complemented by cross-inoculations on cultivated hosts grapevine and apricot to reveal patterns of host specialization. The cultivated hosts are likely more important sources of inoculum than the wild host willow, based on our findings of higher pathogen prevalence and allelic richness in grapevine and apricot. High levels of gene flow among all three hosts and locations, and no grouping by clustering analyses, suggest neither host nor geographic differentiation. Cross-inoculations revealed diversified phenotypes harboring various performance levels in grapevine and apricot, with no apparent correlation with their host of origin. Such phenotypic diversity may enable this pathogen to persist and reproduce as a generalist. Regular genetic reshuffling through sexual recombination, frequent immigration among hosts, and the lack of habitat choice in this passively dispersed fungus may prevent fixation of alleles controlling host specialization.

In agricultural landscapes, i.e., a mosaic of crops exerting differential selective pressures on pathogen populations, the distribution of genetic variation (i.e., population genetic structure) in pathogen populations often coincides with the spatial heterogeneity of hosts (20). Host heterogeneity impacts two evolutionary forces: selection and gene flow. Frequently, populations exploiting different resources are genetically differentiated. For fungal plant pathogens, this pattern usually results from the divergent selection pressures exerted by distinct host plants (20), whereas active habitat choice, which limits migration among environments, is far less relevant for passively dispersed organisms such as fungi.

The ascomycete fungus Eutypa lata (Pers:Fr.) Tul. & C. Tul. (=Eutypa armeniacae Hansf. & Carter) (Xylariales, Diatrypaceae) is considered a wide host range pathogen, based on the fact that it is reported to infect and sporulate on a worldwide total of 70 genera (37 families) of host plants (15). However, reports of hosts that suffer from Eutypa dieback (i.e., severe disease symptoms) are restricted to cultivated hosts, namely grapevine and apricot (8). E. lata is found in Europe, North America, Australia, and South Africa. Recent investigation of its worldwide population genetic structure suggests a European origin of the fungus, with its current global distribution likely resulting from multiple introductions of genetically diverse genotypes into new areas (North America, Australia, and South Africa) via transport of infected plant material (45).

During the dormant (rainy) season, wind-dispersed ascospores infect the woody stems of the host through pruning wounds (8), and establish a localized infection (wood canker) that kills distal shoots (dieback). The mycelium decays the wood, in part through cation of phytotoxic fungal metabolites, via the vascular system, production of cell-wall–degrading enzymes (41). The translo-

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The relative importance of specific hosts in the spread of Eutypa dieback to healthy vineyards is unclear, in part based on conflicting evidence of host specialization. For example, Trouillas and Gubler (46) found no evidence of host specificity; isolates originating from different hosts were all able to infect grapevine and there was no apparent phylogenetic grouping according to either host or geographic origins. As such, all hosts might be thought to be sources of inoculum. In contrast, a previous study suggested that grapevine is susceptible to a wide range of isolates, of which only a few are also infectious on hosts other than grapevine (in this case, pear). In other words, the pathogenic ability to infect particular hosts may be a unique feature of particular isolates.

We evaluated the host specificity of *E. lata* in California, based on population genetics analyses of the pathogen from three hosts (grapevine, apricot, willow). We tested the hypothesis that pathogen populations from different hosts or locations are genetically differentiated, which would support the more important role of host adaptation (selection) than gene flow in shaping population genetic structure. Previous findings revealing that *E. lata* isolates from different hosts were all infectious in grapevine (46) do not allow testing for a higher performance on the host of origin, which would suggest host specialization. Accordingly, we conducted cross-inoculations of the two most economically and historically important hosts of *E. lata*, grapevine and apricot, to test for host specialization. Evidence for host-based genetic differentiation and/or host specialization would help focus management strategies on the most significant source of inoculum.

### MATERIALS AND METHODS

**Experimental design.** From October 2009 to June 2011, hierarchical sampling was carried out in three locations of California (Napa, San Benito, and Solano Counties). In each location, three host plants, grapevine (*Vitis vinifera*), apricot (*Prunus armeniacae*), and willow (*Salix sp.*), were sampled, resulting in nine sampling sites (Table 1). Willow was considered the most important source of inoculum among native hosts by Trouillas and Gubler (46), based on the relatively high incidence of perithecia (sexual fruiting bodies) observed in nature. A total of 182 *E. lata* isolates were collected. Spatial distances among the three locations ranged from 60 to 350 km (Fig. 1). Within each location, spatial distances among the vineyard, apricot orchard, and riparian area did not exceed 6 km. This sampling scheme was designed to determine whether isolates from one host, across locations, were genetically more similar than isolates from different hosts in the same location.

At each location, vineyards, apricot orchards, and riparian areas were surveyed for plants with general dieback symptoms that are typical of all trunk diseases of grape (i.e., dead spur positions, shoot dieback, poor growth) and for the more specific, characteristic foliar symptoms of Eutypa dieback (in grapevine: stunting of shoots, shortening of internodes, chlorosis, necrosis and cupping of leaves; in apricot: intense gummosis, branch death, and wilting of leaves) (8). At each sampling site, between 47 and 118 plants were sampled for fungal isolation (Table 1). Mono-sporous isolates were recovered from perithecia and mass-hyphal isolates from necrotic wood, as previously described (46). Mass-hyphal isolates were further purified through hyphal-tip subcultures. Disease incidence was estimated as the proportion of plants from which *E. lata* was successfully isolated. To test equality of disease incidence between host plants, confidence intervals for incidence differences (CI) were calculated as

\[
CI = p_a - p_b \pm 1.96 \times \sqrt{\left(\frac{p_a \times q_a}{n_a} - 1\right) + \left(\frac{p_b \times q_b}{n_b} - 1\right)}
\]

where \(p_a\) and \(p_b\) are disease incidences in each host; \(q_a\) and \(q_b\) = 1 - \(p\); \(n_a\) and \(n_b\) are the number of samples from each host; and 1.96 corresponds to the 95% confidence level.

**Population genetic analyses.** All 182 isolates were genotyped with nine polymorphic microsatellite markers that have been suitable for *E. lata* population studies (6,45), using published polymerase chain reaction (PCR) conditions (45). Briefly, cultures were incubated on potato dextrose agar ([PDA], Difco, Detroit, MI) at 25°C in dark for 14 days. DNA was extracted from aerial mycelia (DNaseq Plant kit, Qiagen). Species identity was confirmed by PCR amplification and sequencing of the rDNA internal transcribed spacer region (ITS) (ITS1 and ITS4 [47]). Based on phylogenetic analyses of ITS sequences, these 182 isolates clustered with type specimens in maximum likelihood analyses and were identified as *E. lata* (data not shown). Allele assignments were performed on an ABI PRISM 3100xl sequencer (GENEMARKER v.1.75, SoftGenetics LLC). All 182 isolates were genotyped twice, to ensure repeatability. The allelic data set contained no missing data.

The number of repeated multilocus haplotypes, within and across sampling sites, was determined using GENALEX 6 (33), and this number was used to estimate, at each sampling site, the

**TABLE 1. Hosts of origin, geographic locations, and genetic diversity of Eutypa lata populations from three host plants in California (182 total isolates)**

<table>
<thead>
<tr>
<th>Host of origin</th>
<th>Code*</th>
<th>Location</th>
<th>Preponderant crop</th>
<th>Number of isolates</th>
<th>Number of samples*</th>
<th>Incidence*</th>
<th>Number of haplotypes</th>
<th>Clonal fraction</th>
<th>(R^2)</th>
<th>(H^p)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Grapevine</td>
<td>SBenGr</td>
<td>San Benito</td>
<td>Apricot</td>
<td>21</td>
<td>48</td>
<td>0.44</td>
<td>21</td>
<td>0.00</td>
<td>3.34 (1.86)</td>
<td>0.56 (0.14)</td>
</tr>
<tr>
<td></td>
<td>NapaGr</td>
<td>Napa</td>
<td>Grapevine</td>
<td>29</td>
<td>70</td>
<td>0.41</td>
<td>29</td>
<td>0.00</td>
<td>4.36 (2.70)</td>
<td>0.63 (0.15)</td>
</tr>
<tr>
<td></td>
<td>SolaGr</td>
<td>Solano</td>
<td>Apricot</td>
<td>42</td>
<td>118</td>
<td>0.36</td>
<td>42</td>
<td>0.00</td>
<td>4.33 (2.57)</td>
<td>0.60 (0.14)</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td><strong>Average</strong></td>
<td>0.39 a</td>
<td></td>
<td></td>
<td>3.13 a</td>
<td>0.60 a</td>
</tr>
<tr>
<td>Apricot</td>
<td>SBenAp</td>
<td>San Benito</td>
<td>Apricot</td>
<td>17</td>
<td>47</td>
<td>0.36</td>
<td>17</td>
<td>0.00</td>
<td>3.60 (1.85)</td>
<td>0.58 (0.13)</td>
</tr>
<tr>
<td></td>
<td>NapaAp</td>
<td>Napa</td>
<td>Grapevine</td>
<td>15</td>
<td>48</td>
<td>0.31</td>
<td>14</td>
<td>0.07</td>
<td>3.44 (1.33)</td>
<td>0.58 (0.16)</td>
</tr>
<tr>
<td></td>
<td>SolaAp</td>
<td>Solano</td>
<td>Apricot</td>
<td>34</td>
<td>74</td>
<td>0.46</td>
<td>34</td>
<td>0.00</td>
<td>3.83 (2.17)</td>
<td>0.58 (0.14)</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td><strong>Average</strong></td>
<td>0.39 a</td>
<td></td>
<td></td>
<td>2.94 ab</td>
<td>0.58 a</td>
</tr>
<tr>
<td>Willow</td>
<td>SBenWi</td>
<td>San Benito</td>
<td>Apricot</td>
<td>6</td>
<td>54</td>
<td>0.11</td>
<td>6</td>
<td>0.00</td>
<td>NA</td>
<td>0.56 (0.17)</td>
</tr>
<tr>
<td></td>
<td>NapaWi</td>
<td>Napa</td>
<td>Grapevine</td>
<td>13</td>
<td>53</td>
<td>0.25</td>
<td>13</td>
<td>0.00</td>
<td>3.44 (1.67)</td>
<td>0.55 (0.15)</td>
</tr>
<tr>
<td></td>
<td>SolaWi</td>
<td>Solano</td>
<td>Apricot</td>
<td>5</td>
<td>114</td>
<td>0.04</td>
<td>5</td>
<td>0.00</td>
<td>NA</td>
<td>0.56 (0.33)</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td><strong>Average</strong></td>
<td>0.11 b</td>
<td></td>
<td></td>
<td>2.58 b</td>
<td>0.55 a</td>
</tr>
</tbody>
</table>

*Population code for each host at each location.

*Number of plants from which a symptomatic wood sample was collected for fungal isolations.

*Proportion of plants from which *E. lata* was cultured. Values followed by different letters are significantly different, based on lack of overlap of 95% confidence intervals.

*Mean allelic richness (standard deviation in parentheses) per host plant, averaged across loci and sampling sites (37). Values followed by different letters are significantly different (assessed after 5,000 random permutations of haplotypes across hosts; \(P > 0.05\)). NA = not applicable; Number of haplotypes < 10.

*Mean gene diversity (standard deviation in parentheses) per host plant, averaged across loci and sampling sites (32). Values followed by different letters are significantly different (assessed after 5,000 random permutations of haplotypes across hosts; \(P > 0.05\)).
estimates of variance components and squared-distances among all pairs of haplotypes used to produce framework incorporates haplotype divergence into a matrix of hierarchical levels (12). Two distinct analyses were performed to quantify the partitioning of molecular variation at different hierarchical levels (12). Two distinct analyses were performed: samples were grouped either according to their host of origin or their location (i.e., county) of origin. The AMOVA framework incorporates haplotype divergence into a matrix of squared-distances among all pairs of haplotypes used to produce estimates of variance components and F-statistic analogs, designated as \( \Phi \)-statistics (12). GENALEX was used to compute and test the statistical significance of \( \Phi \)-statistics based on 999 random permutations of haplotypes.

To examine population subdivision, we used a Bayesian method of assignment implemented in the program STRUCTURE 2.3.1 (39). This program uses a Markov Chain Monte Carlo algorithm to assign multilocus haplotypes to a genetic cluster, assuming Hardy-Weinberg equilibrium and minimizing linkage disequilibrium among loci within clusters. Analyses were performed both using uniform priors and using the location of origin or host of origin of isolates as prior information to assist clustering (model LocPrior [24]). The likelihood of the posterior probability distributions was computed for each number of clusters \( K \) from 1 to 12. To check for consistency of likelihood values for each \( K \) value between runs, each \( K \) was simulated six times, with a run length of 300,000 iterations after the specified burn-in (300,000 iterations), using the admixture model of genetic ancestry and the correlated model of allele frequency (14). We estimated the number of clusters as \( \Delta K \), which is based on the second-order rate of change of the likelihood of posterior probability of the data \( L(K) \) between successive \( K \) values (11).

**Cross-pathogenicity tests.** A total of 24 isolates was selected for cross-inoculations, with 12 isolates from San Benito and 12 isolates from Napa, in February 2012. From each location, six isolates originated from grapevine and six from apricot. Symptom development of Eutypa dieback in grapevine woody tissue requires an incubation period ≥12 months. As such, this was a 14-month experiment encompassing two greenhouses, in which we propagated the perennial hosts from dormant plant material. Inoculum consisted of a 3-mm-diameter agar plug colonized by actively growing mycelium (PDA, Difco Laboratories). The 24 isolates were inoculated to grapevine (‘Thompson Seedless’) and apricot (‘Patterson’), and noninoculated controls were mock-inoculated with sterile PDA to give a total of 50 isolate \( \times \) host combinations. Two replicate experiments were conducted 2 weeks apart in two separate greenhouses (\( n = \) six plants per treatment, 600 total plants). In each greenhouse, plants were arranged in a completely randomized design.

Grapevines were propagated from 1-year-old dormant cuttings of ‘Thompson Seedless.’ Cuttings were cut to uniform length (=30 cm) containing three nodes, surface-sterilized in 1% sodium hypochlorite (Clorox, Oakland, CA) for 15 min, rinsed in water overnight, and then callused in a mixture of perlite and vermiculite (1:1, vol/vol), at 30°C and 100% RH for 3 weeks. Once root and shoot initials emerged from the callus tissue, the stem of each cutting was wounded at approximately 3 cm below the uppermost node with a 3-mm-cork borer, and a 3-mm mycelial plug from a 10-day-old PDA culture was inserted into the wound and sealed with Vaseline and Parafilm. Noninoculated controls were ‘mock-inoculated’ with a sterile PDA plug. After inoculation, cuttings were submerged in melted paraffin wax (Gulf Wax, Royal Oak Enterprises, Atlanta, GA) within 4 cm of the roots and potted in UC mix (5) amended with slow-release fertilizer (Osmocote Pro 24-4-9, Scotts, Marysville, OH).

Apricots were propagated from dormant, bare root trees of ‘Patterson’ (=1 cm diameter) grafted onto Nemaguard rootstock. On the day of inoculations, apricot trees were taken out of cold storage (2°C), and the inoculation and potting procedures were identical to those for the grapevines, except with no waxing step. Grapevines and apricots were grown in the greenhouse at the University of California Experimental Station in Davis from February 2012 to April 2013 (natural sunlight photoperiod, 25 ± 1°C [day], 18 ± 3°C [night]). In early November, plants were pruned and the temperature was decreased (10 ± 2°C [day], 4 ± 2°C [night]) until the end of January. Plants were watered twice per week for 15 min using a drip-irrigation system (0.5 l h\(^{-1}\)).

The performance of *E. lata* isolates was based on two distinct measures: the incidence of foliar symptoms and the extent of wood lesions in inoculated stems. Following bud break in the second growing season (mid-February 2013), plants were monitored for foliar symptoms twice per week for 6 weeks. Incidence of foliar symptoms was calculated as the proportion of plants per treatment expressing the following characteristic symptoms of Eutypa dieback: necrotic areas on some leaves, dwarfed leaves with necrosis, reduction in leaf and shoot size, rapid dieback of stunted shoots, and/or no shoot growth. Characteristic foliar symptoms on apricot included a rapid dieback of the shoots above the inoculation site during the growing season or no shoot growth in the second growth cycle. Wood symptoms were evaluated 14 months after inoculation (April 2013). Plants were removed from the soil; roots and shoots were excised; and bark was scraped from the woody stem. Stems were surface-sterilized in 1% sodium hypochlorite solution for 2 min, rinsed with tap water,
and cut longitudinally. The length of wood lesions extending above and below the point of inoculation was measured with a caliper.

There are two ways to test for host specialization in a “common garden” experiment, as conducted here: (i) by comparing the performance of a pathogen population from a given host in different hosts (the “home versus away” criterion), or (ii) by comparing the performance of a pathogen population in its native habitat (i.e., host of origin) with that of another pathogen population from a different habitat (the “local vs. foreign” criterion) (26). In both cases, the relevant statistic is the significance of the interaction between the pathogen population (from the host of origin) and test host (host inoculated). The incidence of foliar symptoms (pathogenicity) was compared among treatments using a generalized linear model with a binomial error distribution and a logit link function (GLIMMIX procedure of SAS version 9.2; SAS Institute, Cary, NC). When testing for a host specialization pattern (higher average incidence of foliar symptoms on host of sampling), the host of origin, test host, and their two-way interactions were considered fixed effects. In the model, replicate plants and fungal isolates were considered random effects; they were nested within test host and host of origin, respectively. Contrasts allowed testing the “home vs. away” criterion (comparisons between the two hosts for one pathogen population) and the “local vs. foreign” criterion (comparisons between the two pathogen populations on one host). Severity of wood symptoms (i.e., wood lesion length; aggressiveness) was compared among treatments using an analysis of variance (ANOVA; MIXED procedure of SAS); host specialization was tested as previously described for incidence of foliar symptoms.

RESULTS

Disease incidence. General dieback symptoms were observed in all three hosts and in all three locations. Characteristic Eutypa dieback symptoms (intense gummosis, branch death, and wilting of leaves) were observed in some apricot trees at each location. Similarly, diagnostic symptoms were observed in several grapevines at locations surveyed during spring months. Perithecia of the fungus, which were embedded in a black stroma located with-in the bark, were observed on each host species. Based on recovery of the fungus in culture, the incidence of E. lata in vineyards and apricot orchards ranged from 31 to 46% (Table 1). In contrast, incidence in willow trees was lower, ranging from 4 to 25%. Across locations, the average incidence in grapevine and apricot was 39% and did not differ significantly (95% CI of –0.10 to 0.10, P = 0.05, 401 df). Disease incidence averaged across locations was significantly lower in willow (11%) than in grapevine (95% CI of 0.21 to 0.35, P = 0.05, 453 df). Similarly, disease incidence was significantly lower in willow than in apricot (95% CI of 0.20 to 0.36, P = 0.05, 368 df).

Genetic diversity. Only two repeated multilocus haplotypes were identified from the same sampling site (apricot in Napa, NapaAp). The clonal fraction was null in all but one sampling site (NapaAp, Table 1). In addition, two pairs of repeated haplotypes were identified from distinct sampling sites: two identical haplo-types were identified from apricot in Napa and Solano, and two identical haplotypes were identified from apricot in Solano and San Benito. Gene diversity (H) was homogenous across sampling sites and varied between 0.55 and 0.63. Differences in allelic richness (R) were revealed across sampling sites: the mean number of alleles per locus, estimated for 13 haplotypes (the smallest sample size in the data set), ranged from 3.34 (SBenGr) to 4.36 (NapaGr). When H and R, averaged across sampling sites per host, were compared among hosts (grapevine, apricot, willow), significant differences in allelic richness were revealed (P = 0.048). Consistently for these two measures of genetic diversity, sampling sites from grapevine were the most genetically diverse (Table 1).

The hierarchical AMOVA, first using location as a grouping factor, revealed that 99% of the genetic variation was found within-in sampling sites, and differences among locations did not contribute to total genetic variation (Table 2). Genetic differences among sampling sites within a location contributed to 1% of genetic variance. When host was used as a grouping factor, results were similar and 99% of the genetic variation was found within sampling sites. The results of the AMOVA analyses, therefore, indicated that there were no significant differences in pathogen populations among sampling sites from different locations or from different hosts. The low genetic differentiation among samples from different hosts was illustrated by the negative and nonsignificant estimator of genetic differentiation (ΦST = –0.004; P = 0.67; Table 2), suggesting high levels of gene flow among the three hosts.

Clustering analysis. The Bayesian method of assignment of multilocus haplotypes detected population subdivision, as evidenced by the estimated modal values of the statistic δK (Fig. 2). Analyses using sampling location as prior information suggested the presence of three genetic clusters, whereas analyses using host of sampling as prior information did not reveal any clear population structure (Fig. 2). Analyses that made no use of prior population information revealed that either three or five genetic clusters of multilocus haplotypes best described the genetic variation. According to these results, we presented membership coefficients to each cluster for K = 3 and K = 5 (Fig. 3). Bar plots for K = 3, using either no prior population information or sampling location as prior information to assist clustering, revealed a high proportion of multilocus haplotypes with admixed ancestry (<50% probability of membership to any cluster). In contrast, bar plots representing the membership coefficients to each cluster for K = 5 (no use of prior population information) revealed a clearer pattern of population subdivision, while the proportion of multilocus haplotypes with admixed ancestry was low (16 out of 182 haplotypes). Assessment of membership probabilities relative to the host of origin of each haplotype did

<table>
<thead>
<tr>
<th>Source of variation</th>
<th>df</th>
<th>Variance components</th>
<th>% variation</th>
<th>Φ-statistics</th>
<th>P values</th>
</tr>
</thead>
<tbody>
<tr>
<td>Among locations</td>
<td>2</td>
<td>0.00</td>
<td>0</td>
<td>ΦRT = –0.004</td>
<td>0.61</td>
</tr>
<tr>
<td>Among sampling sites within locations</td>
<td>6</td>
<td>3.75</td>
<td>1</td>
<td>ΦRT = 0.006</td>
<td>0.34</td>
</tr>
<tr>
<td>Within sampling sites</td>
<td>173</td>
<td>674.53</td>
<td>99</td>
<td>ΦRT = 0.002</td>
<td>0.39</td>
</tr>
<tr>
<td>Among hosts</td>
<td>2</td>
<td>0.00</td>
<td>0</td>
<td>ΦRT = –0.004</td>
<td>0.67</td>
</tr>
<tr>
<td>Among sampling sites within hosts</td>
<td>6</td>
<td>3.76</td>
<td>1</td>
<td>ΦRT = 0.006</td>
<td>0.35</td>
</tr>
<tr>
<td>Within sampling sites</td>
<td>173</td>
<td>674.53</td>
<td>99</td>
<td>ΦRT = 0.001</td>
<td>0.46</td>
</tr>
</tbody>
</table>

* Sampling site is the location x host combination (e.g., grapevines in San Benito or SBenGr).
* Φ-statistics are analogues of F-statistics, estimated through AMOVA (12). ΦRT = proportion of variance among regions, relative to total variance. ΦPR = proportion of variance among populations within regions, relative to variance among and within populations. ΦPT = proportion of variance among populations, relative to total variance.
* Probability that a statistic obtained from 999 random permutations of haplotypes across the data set is greater than the value observed.
not reveal any genetic clustering by host plant. Indeed, most
genetic clusters consisted of haplotypes from all hosts and loca-
tions. For example, cluster 4 included haplotypes from each of the
nine sampling sites (Fig. 4). The five genetic clusters were not
supported by any reasonable biological interpretation, such as
geography or host of origin. The lack of substantive biological
reasons to support the outputs of these analyses led us to consider
this identified genetic structure as marginal.

Cross-pathogenicity tests. Inoculations conducted in the
greenhouse on grapevine and apricot plants revealed a large vari-
ation in both the severity of wood symptoms and the incidence of
folic symptoms among the 24 isolates (Fig. 5). When inoculated
to grapevines, there was a smaller range in length of wood lesions
(55 to 145 mm) than on apricot (50 to 360 mm). Ranges of folic
symptom incidence were similar in both hosts (approximately 20
to 100%). Some isolates caused large wood lesions and frequent
folic symptoms on grapevine and apricot (e.g., SBen204, SBen218). By contrast, some isolates caused small wood lesions and a low incidence of folic symptoms on both hosts (e.g.,
SBen015). In a few cases, some isolates seemed better adapted to
one host than the other; for example, isolate SBen020 caused
larger lesions and a higher incidence of folic symptoms on
apricot than on grapevine (Fig. 5).

Host specialization. A significant difference in E. lata patho-
genicity was found between isolates from the two hosts of origin,
with higher average incidence of folic symptoms for isolates
from apricot than isolates from grapevine (P = 0.037; Table 3).
Contrasts revealed greater incidence of folic symptoms of
grapevine isolates when inoculated to grapevine than to apricot
(“home versus away” comparison, P = 0.024; Table 3, Fig. 6A),
suggesting a better performance on the host of origin. Nonethe-
less, this pattern of host specialization was not observed for
apricot isolates; they caused greater incidence of folic symptoms
on grapevine than on apricot. The absence of a significant host of
origin × test host interaction (P = 0.072; Table 3) and the absence
of crossing reaction norms (Fig. 6A) revealed no clear host-
specialization pattern for E. lata based on incidence of folic
symptoms.

Host of origin, test host, and their interaction had significant
effects on the length of wood lesions, indicating a pattern of host
specialization based on this life-history trait (Table 4). Indeed,
isolates from apricot caused larger wood lesions than isolates
from grapevine when inoculated to apricot (“local vs. foreign”
comparison, P = 0.0006; Table 4, Fig. 6B). The two significant
“home vs. away” comparisons revealed an overall higher severity
of wood lesions of isolates from both hosts when inoculated to
apricot (Table 4 and Fig. 6B). For example, grapevine isolates
caused larger wood lesions in apricot than in grapevine (P =
0.0002; Table 4). These results indicated an overall greater level
of aggressiveness in (or susceptibility of) apricot, independently
of the host in which these isolates originated.

DISCUSSION

We tested the hypothesis of host specialization in the wood-
canker pathogen E. lata, and our results did not provide evidence
for host-associated genetic differentiation or preferential host
exploitation in this pathogen. The pathogen population infecting
multiple hosts in California was genetically homogenous, based
on our finding of no genetic structuring according to either host or
geographic origins. The assessment of host specialization to
grapevine and apricot for two life-history traits, the pathogen’s
ability to cause folic symptoms and to cause large wood lesions,
did not support a consistent pattern of host adaptation. These
results support an overall greater role of gene flow than selection
in structuring the pathogen population in California. The three
host plants examined may not exert strong divergent selection
pressures on the pathogen population, and free migration among
hosts may be facilitated by the large population sizes and high
fecundity that characterize fungal pathogens.

As concerns about “pathogen pollution” and emerging infec-
tious diseases of plants increases (4), a thorough understanding of
population dynamics of pathogens that infect multiple host
species is required (10). The effect of host species diversity on
disease risk in a host of interest can have several outcomes de-
pending on the relative properties of each host (27). Different
hosts usually have different “reservoir potential,” corresponding
to the product of the proportion of plants infected by the pathogen
and the probability that this host will serve as a source of inoculum (29). With respect to E. lata, the fungus produces sexual
fruiting bodies on any infected and dying hosts (46). Hence, once
infected, grapevine, apricot, and willow constitute a source of
infectious propagules. As a first approximation, we can thus
consider that the reservoir potential of each host depends only on
the pathogen incidence (i.e., prevalence) in each host. Our find-
ings illustrate the contrasted reservoir potential of the three host
plants examined; grapevine and apricot had similar incidences of
the pathogen (39%), whereas willow had a significantly lower
reservoir potential, with an incidence of 11%, averaged across the
three locations. These results support the hypothesis that culti-
vated crops, such as grapevine and apricot, are more important
sources of inoculum than are wild plants, such as willow, present
in riparian areas (13,38).

Our finding of higher genetic diversity of E. lata populations on
grapevine and apricot further support this hypothesis. When
measured in terms of allelic richness, the genetic diversity of the
E. lata populations on willow was significantly lower than those
of populations on grapevine or apricot. This lower genetic
diversity of populations on willow is in agreement with a smaller
effective population size of E. lata populations in riparian areas.
As previously discussed (45), E. lata is a wound pathogen, and
grapevine and apricot are regularly and severely “wounded” when
they are pruned each winter, which is a cultural practice used to
maintain high crop productivity. In contrast, willow is a wild host;
it is not pruned and thus has fewer infection courts. This may

fig. 2. Change in the statistic deltaK relative to the number of clusters K (1 to
12) tested on the 182 Eutypa lata multilocus haplotypes with the program
STRUCTURE. Estimates of deltaK are based on the second order rate of change
of the likelihood of posterior probability of the data between successive K values.
explain the low genetic diversity of *E. lata* in willow, relative to the higher incidence, effective population size, and genetic diversity observed in *E. lata* populations in apricot orchards and vineyards.

In populations of plant-pathogenic fungi, the spatial scale of pathogen dispersal usually affects the spatial scale of genetic differentiation, which in turn may affect the scale of local adaptation (i.e., higher performance on sympatric than on allopatric genotypes of a given host [44]). In the present study, we anticipated that a pattern of adaptation of *E. lata* to a particular host may not be detectable at a local spatial scale (i.e., between an adjacent vineyard and orchard) because of high dispersal levels among hosts that may “swamp” locally adapted genotypes. Instead we hypothesized that such a pattern may be more likely detectable at a regional spatial scale (i.e., San Benito versus Napa), because these two locations are separated by 360 km, which may restrict habitat connectivity. However, our finding of no significant genetic differentiation of *E. lata* populations suggests that gene flow among hosts and locations at both local and regional scales prevents genetic differentiation in this pathogen. High levels of gene flow were previously shown among *E. lata* populations from different continents (36,45). Gene flow at local spatial scales is likely a result of passive ascospore dispersal by wind, whereas gene flow at continental scales is more likely a result of human-mediated transport of infected plant material (45). A high level of gene flow among local *E. lata* populations in California may promote the mixing of locally adapted genotypes and migrant, maladapted genotypes in each local pathogen population.

As for the homogenizing effect of gene flow that hinders the differentiating effect of selection, sexual recombination also tends to break up locally adapted allelic combinations (linkage dis-

![Fig. 3. Posterior probability membership of each individual to each genetic cluster for K = 3 and K = 5. Posterior probability memberships were estimated both using uniform priors and using the location of origin as prior information to assist clustering. Each color represents a genetic cluster, and each individual is represented by a vertical line partitioned into colored segments whose lengths represent the admixture proportions from K clusters.](image-url)
equilibrium created by selection), except when genes controlling habitat specialization and mate choice (i.e., assortative mating) are identical or physically linked (19). Although we did not test for multilocus or pairwise linkage disequilibrium in the present study, previous genetic analyses of E. lata populations with the same set of microsatellite markers (45) and different markers (35, 36) support the hypothesis that E. lata is a random mating species. Indeed, repeated haplotypes are rare. It has been suggested that life history traits of many plant-pathogenic fungi, especially when mating takes place within the host plant, favor the fixation of alleles controlling host specialization (21). Host adaptation induces pleiotropically assortative mating, which can eventually lead to reproductive isolation and ecological speciation (20). Hence, if California is a recent geographic area occupied by this pathogen (45), a short evolutionary timescale may have prevented host adaptation; host range expansion may be too recent to generate divergences within the pathogen population. Also, we cannot reject the hypothesis that our neutral genetic markers are not suited to detect differentiation among host-adapted populations; instead, examining genes under positive selection may allow the detection of host specialization (3). Alternatively, if host adaptation evolved in source populations of the pathogen before introduction to California, patterns of host adaptation should still be detectable because mating occurs within host plants, thereby facilitating the fixation of alleles controlling host adaptation in local pathogen populations (21). Our results suggest that such a pattern of host adaptation was absent in source populations of this pathogen.

The characterization of 24 E. lata isolates for foliar symptom incidence and wood lesion length provided further insights into the phenotypic diversity in populations of this fungus. The population examined in California can be considered polymorphic, based on the variation in these two life-history traits. Such polymorphism can be considered adaptive when natural selection promotes phenotypic diversity (31), as is the case when heterogeneous environments (e.g., distinct hosts) favor frequency-dependent selection (40). However, our findings do not support such adaptive polymorphism. For instance, the absence of a pattern of
host specialization for the incidence of foliar symptoms does not suggest that the host plants exert a strong selection pressure for this trait. Moreover, some isolates caused severe foliar symptoms on both hosts, regardless of their host of origin. Considering the large genotypic polymorphism detected among the isolates studied and the apparent lack of adaptive phenotypic diversity, such genetic variation could be considered adaptive if it allows the pathogen to perform efficiently in an increased range of niches, relative to less genetically diverse pathogen populations; this hypothesis remains to be tested. The large phenotypic variation observed among the 24 isolates for severity of wood symptoms also precluded the detection of a pattern of host specialization. Some isolates could be considered adapted to grapevine and maladapted to apricot (e.g., SBen022); other isolates could be considered adapted to apricot but not to grapevine (e.g., Napa215); and isolates could be maladapted to both hosts (e.g., SBen015) or adapted to both hosts (e.g., SBen207). Overall, our finding of such diversified phenotypes, representing variable performance levels of isolates in different hosts, suggests a generalist strategy in *E. lata* that may enable this species to persist and reproduce on a large range of hosts.

The broad host range of *E. lata* in California (46) and our findings supporting a generalist strategy in this wood-decay fungus suggest that “preadapted” genotypes were present in the

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**Fig. 5.** Length of wood lesions and incidence of foliar symptoms caused by 24 *E. lata* isolates on **A**, grapevine, and **B**, apricot plants. Isolates from Napa and San Benito are labeled Napa and SBen, respectively. Isolates recovered from grapevine are labeled from 001 to 023, whereas isolates recovered from apricot are labeled 201 to 218. Incidence of foliar symptoms was assessed at the beginning of the second growth cycle, at 12 months after inoculation in the greenhouse. Severity of wood symptoms was expressed as wood lesion length (mm) and was measured in the inoculated stems at 14 months after inoculation.
pathogen population at the time of introduction to California, and such genotypes were able to use novel resources for survival and reproduction (i.e., phenotypic plasticity). Organisms have a range of environments in which they can survive and reproduce, and this range is larger than the sole environments in which they have evolved (2). This corresponds to the process of ecological fitting, implying that the pathogen traits necessary to infect a plant (i.e., for the plant to serve as a host) did not result from a long co-evolutionary history with that host, but rather that such traits evolved under different environmental conditions (2,25). We hypothesize that the introduced population of *E. lata* in California evolved as a soft-rot pathogen in its native range, which allowed subsequent host-range expansion in novel areas where new hosts with similar physiology and defense responses were present (i.e., host resource tracking). Assuming that defense responses among these related hosts were conserved, the pathogen may have been able to infect hosts it had not previously encountered (1). In conclusion, *E. lata* and possibly other wood-decay fungi infecting exposed wood may possess the characteristics that would allow them to infect new hosts they have not coevolved with in the past through ecological fitting. Future studies on adaptive patterns in wood-decay fungi might consider ecological fitting as a cause of host-range expansion.

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