A continuum of genetic divergence from sympatric host races to species in the pea aphid complex

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Sympatric populations of insects adapted to different host plants, i.e., host races, are good models to investigate how natural selection can promote speciation in the face of ongoing gene flow. However, host races are documented in very few model systems and their gradual evolution into good species, as assumed under a Darwinian view of species formation, lacks strong empirical support. We aim at resolving this uncertainty by investigating host specialization and gene flow among populations of the pea aphid complex, *Acyrthosiphon pisum*. Genetic markers and tests of host plant specificity indicate the existence of at least 11 well-distinguished sympatric populations associated with different host plants in Western Europe. Population assignment tests show variable migration and hybridization rates among sympatric populations, delineating 8 host races and 3 possible species. Notably, hybridization correlates negatively with genetic differentiation, forming a continuum of population divergence toward virtually complete speciation. The pea aphid complex thus illustrates how ecological divergence can be sustained among many hybridizing populations and how insect host races blend into species by gradual reduction of gene flow.

The beginning of this century has seen the reconciliation of Darwin’s view that species gradually emerge by the primary action of natural selection, with the modern evolutionary synthesis defining speciation as the evolution of reproductive isolation (1–4). Adaptations to different habitats and resources may induce reproductive isolation (reviewed in refs. 1, 5, and 6) if morphology or niche selection determine mate choice and if hybrids between ecologically divergent taxa are unfit in the parental environments. Theoretical models and recent empirical studies (reviewed in refs. 7 and 8) have shown that these ecological reproductive barriers may evolve without the geographical separation of populations, an extrinsic barrier to hybridization (gene flow) that was long thought to be a requisite for speciation in animals (2). Divergence in sympatry (i.e., within the same region) implies a more gradual reduction of gene flow that should reflect increasing reproductive isolation and the continuum of speciation from polymorphic populations to good species. Such evolution is not observable in nature at reasonable timescales, but it could be inferred from the multiple stages of sympatric divergence observed within a variety of organisms (9).

The pea aphid complex encompasses sympatric populations showing differential preference and fitness on specific host plants (16–20); hereafter, we refer to such populations as “biotypes,” a word that is useful for cases in which genetic bases of variation have not yet been established. Among these morphologically similar populations, only those feeding on a few crops have been characterized genetically. In North America, populations specialized on red clover and alfalfa are genetically distinct (21), and genetic trade-offs prevent the optimal use of both host species by pea aphids (22). In Europe, 2 host-adapted populations coexist on the same plant species with another one specialized on pea and broad bean, which exhibits a higher genetic divergence (23, 24). The precise levels of reproductive isolation between these host-specialized populations remain nonetheless unresolved. Other populations feeding on wild plants in Western Europe have been recognized at the rank of subspecies or species on the basis of subjective morphological or ecological criteria (15, 25). The pea aphid constitutes therefore an ideal model for studying a range of stages of speciation in sympatry.

Results and Discussion

Multiple Host-Specialized Sympatric Populations. We examined western European pea aphids on 19 widely distributed, sympatric host species, mostly wild plants, using microsatellite DNA markers and laboratory tests of host plant specificity. The Bayesian assignment method implemented in the program Structure (26–28), with no a priori information on the origin of individuals, identified 11 well-distinquished genetic clusters, or populations, associated with different host plants and not with sampling locations (Fig. L4). This sharp population genetic structure is mirrored by clear differences in performance (survival and growth) of aphid lineages tested on representative plants in the laboratory (Fig. 1B). Consistent with expectations, pea aphid lineages performed better on the test plant from which individuals of their genetic population were usually collected (supporting information [SI] Fig. S1). Phenotypic data thus demonstrate that genetic clusters correspond to 11 host-specialized populations, which we hereafter call biotypes.

Three of these biotypes, noted “E” on clovers (*Trifolium,* “G”

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on pea (Pisum) and annual vetches (Vicia faba, hirsuta, and sativa), and “K” on alfalfa (Medicago sativa), correspond to populations that were previously characterized genetically on narrower host ranges (23, 24). The existence of biotypes “A” on Cytisus scoparius, “B” on Ononis sp., “D” on Lotus pedunculatus, and “I” on Lathyrus pratensis was expected, given earlier phenotypic measures of host specialization on these plants (17, 18).

The 4 remaining biotypes “C, H, J, and F” were not identified before the present study, to our knowledge. Notably, the first 3 are specialized on plant species belonging to genera harboring other biotypes: L. corniculatus, V. cracca, and M. lupulina, respectively, with biotype F being specialized on 2 species of Melilotus. Given that many other legume genera harbor pea aphids (25), the discovery of these 4 biotypes strongly suggests the existence of many other biotypes in the pea aphid complex.

### Gene Flow and Genetic Differentiation Among Pea Aphid Biotypes.

Because pea aphids grow and mate on their host plants (25, 29), the association between biotype and plant species informs us on the potential for gene flow. The Bayesian method implemented in the program Geneclass (30) revealed the presence of “migrant” aphids, which correspond to individuals collected on a plant species that is not typical of their biotype (Table 1). Host specialization tests on laboratory lineages initiated by some of these migrant individuals were consistent with their genetic assignment, as they showed similar patterns of host performance to other lineages of the same genetic cluster (Fig. S1). Immigration rates are averaged at 9.4% across host species and indicate potential gene flow among biotypes. To quantify the levels of actual gene flow, we looked for first-generation hybrids, i.e., individuals sharing ancestries with 2 biotypes, using the programs Structure and NewHybrids (31). We also measured the genetic differentiation at microsatellite loci between biotypes and in sympatry, using hierarchical analyses of molecular variance (32, 33). The frequency of hybrids per biotype pair appeared moderate (Fig. 2, circles). However, because of the various parental origins for hybrids within this complex of 11 biotypes (Table S1), their cumulative proportion per biotype could reach 9% (Fig. 2, diamonds) and negatively correlates with their level of genetic differentiation. This correlation can be explained if the proportion of hybrids partly reflects reproductive isolation in pea aphid biotypes. Reproductive isolation determines effective gene flow in sympatric populations and hence their genetic differentiation at equilibrium between gene flow and drift (34). If equilibrium is not reached, genetic differentiation may mostly reflect divergence time, rather than effective gene flow. Likewise, its correlation with hybrid frequency could be explained by increasing reproductive isolation during divergence, as expected under the process of speciation. Variable spatial distance separating host species at all sampling sites may also account for the correlation between hybrid frequency and genetic divergence in pea aphid biotypes. However, the genetic structure of these biotypes appears moderately affected by distances of several hundreds of kilometers, because they constitute homogeneous genetic clusters at the scale of western Europe (Fig. 1A). Therefore, the distance separating host species at sampling sites, which was of hundreds of meters at most, would not explain the various levels of differentiation observed here (Fig. 2). The negative correlation between population differentiation and the proportion of hybrids among the 11 biotypes would thus portray various stages of speciation. Specifically, the 3 most differentiated biotypes A, B, and I, on C. scoparius, Ononis species, and L. pratensis, can be considered to be nearing complete speciation because no hybrid could be detected with any other sympatric biotype (Fig. 2). By contrast, the 8 other biotypes that presented various proportions of hybrids would constitute host-specialized races (11). They likely belong to the same species, given the diverse origins of

![Fig. 1. Genetic structure and host specialization in western European pea aphids. (A) Population structure: horizontal segments represent 1,090 individuals (14-microsatellite genotypes), and colors represent their proportions of ancestry to inferred populations coded from A to K (see B). Genotypes are sorted by host plants and then by locations, as indicated by the vertical bar on the left. White, eastern France; black, northwestern France; gray, eastern Germany. (B) A phenogram of 45 aphid laboratory lineages assigned to 10 populations and based on their performance (survival and growth) on 10 representative plant species (photographs and names in boldface type in A). Branch lengths thus represent differences in host specialization on these plants. Italicized letters indicate biotypes that were previously characterized genetically (23, 24) and letters in boldface type indicate biotypes whose existence was suspected from earlier host specialization tests (17, 18).](http://www.pnas.org/cgi/doi/10.1073/pnas.081117106)

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their hybrids and lower pairwise genetic differentiation (Table S1). This hypothesis is supported by very low sequence polymorphism at noncoding nuclear makers, in comparison to interspecific divergence with a related species *A. kondoi* at the same loci (Fig. S2).

**Implications for Ecological Divergence and Reproductive Isolation.**

Gene flow between populations often constrains their adaptive divergence (35). In the pea aphid, pairwise differences in host performance between biotypes do not correlate with their genetic differentiation (Fig. 3) and do not provide evidence that current hybridization hinders ecological divergence. Strong selection by host plants (ref. 19 and Fig. S1) may explain this observation, genetic differentiation being likely higher at loci controlling host adaptation compared to the average genetic divergence at our microsatellites (36). Also, western European races remain highly differentiated at many microsatellite markers (usually >50%, Fig. 2) despite appreciable hybridization. This observation strongly suggests that hybrids contribute only weakly to subsequent generations, in other words, postzygotic isolation. In North American host races, postzygotic isolation is the by-product of host plant adaptation, as trade-offs in host performance prevent the optimal use of parental plants by hybrid lineages (22). Postzygotic isolation may have an ecological basis in European biotypes, which present pronounced host specialization (ref. 19 and Fig. S1). They also produce hybrids

### Table 1. Distribution of sampled pea aphids belonging to 11 biotypes on the surveyed plants

<table>
<thead>
<tr>
<th>Collection plant</th>
<th>A</th>
<th>B</th>
<th>C</th>
<th>D</th>
<th>E</th>
<th>F</th>
<th>G</th>
<th>H</th>
<th>I</th>
<th>J</th>
<th>K</th>
<th>Total</th>
<th>Migrants (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td><em>Cytisus scoparius</em></td>
<td>156</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>156</td>
<td>0.0</td>
</tr>
<tr>
<td><em>Ononis repens</em></td>
<td>48</td>
<td>1</td>
<td>1</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>50</td>
<td>4.0</td>
</tr>
<tr>
<td><em>O. spinosa</em></td>
<td>77</td>
<td>2</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>79</td>
<td>2.5</td>
</tr>
<tr>
<td><em>Lotus corniculatus</em></td>
<td>56</td>
<td>2</td>
<td>1</td>
<td>4</td>
<td>9</td>
<td>2</td>
<td>4</td>
<td>78</td>
<td>28.2</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td><em>L. pedunculatus</em></td>
<td>5</td>
<td>68</td>
<td>1</td>
<td>2</td>
<td></td>
<td></td>
<td></td>
<td>76</td>
<td>10.5</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td><em>Trifolium campestrum</em></td>
<td>22</td>
<td>3</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>25</td>
<td>12.0</td>
</tr>
<tr>
<td><em>T. dubium</em></td>
<td>12</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>12</td>
<td>0.0</td>
</tr>
<tr>
<td><em>T. repens</em></td>
<td>66</td>
<td>4</td>
<td>2</td>
<td>4</td>
<td>1</td>
<td>77</td>
<td>14.3</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td><em>T. pratense</em></td>
<td>1</td>
<td>1</td>
<td>202</td>
<td>3</td>
<td>3</td>
<td>210</td>
<td>3.8</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td><em>Melilotus albus</em></td>
<td>52</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>55</td>
<td>5.5</td>
</tr>
<tr>
<td><em>M. officinalis</em></td>
<td>54</td>
<td>2</td>
<td></td>
<td>1</td>
<td>57</td>
<td>5</td>
<td>53</td>
<td>5.3</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td><em>Pisum sativum</em></td>
<td>139</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>139</td>
<td>0.0</td>
</tr>
<tr>
<td><em>Vicia faba</em></td>
<td>2</td>
<td>48</td>
<td></td>
<td>4</td>
<td>54</td>
<td>11.1</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td><em>V. sativa</em></td>
<td>1</td>
<td>30</td>
<td>5</td>
<td></td>
<td></td>
<td>36</td>
<td>16.7</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td><em>V. hirsuta</em></td>
<td>6</td>
<td>37</td>
<td>1</td>
<td>2</td>
<td>4</td>
<td>50</td>
<td>26.0</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td><em>V. cracca</em></td>
<td>2</td>
<td>1</td>
<td>102</td>
<td>1</td>
<td>1</td>
<td>114</td>
<td>10.5</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td><em>Lathyrus pratensis</em></td>
<td>1</td>
<td>16</td>
<td>97</td>
<td>2</td>
<td></td>
<td>116</td>
<td>16.4</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td><em>Medicago lupulina</em></td>
<td>1</td>
<td>1</td>
<td>2</td>
<td>1</td>
<td>113</td>
<td>4</td>
<td>122</td>
<td>7.4</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td><em>M. sativa</em></td>
<td>2</td>
<td></td>
<td></td>
<td>5</td>
<td>182</td>
<td>190</td>
<td>4.2</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Total</td>
<td>159</td>
<td>125</td>
<td>67</td>
<td>70</td>
<td>316</td>
<td>107</td>
<td>279</td>
<td>138</td>
<td>99</td>
<td>133</td>
<td>203</td>
<td>1696</td>
<td>9.4</td>
</tr>
</tbody>
</table>

Biotypes are denoted by the same letters as in Fig. 1. Numbers in boldface type indicate “resident” individuals, as opposed to “migrants” found on plants typical of other biotypes. To give a more accurate picture of migration patterns between plants, this distribution considers all aphids that could be clearly assigned to a biotype (thus excluding hybrids), counting individuals of the same parthenogenetic lineage (see Methods). Because all aphids were wingless and sampled on separate plants, these individuals represent different settlements.

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**Fig. 2.** Correlation of the proportion of hybrids with the genetic differentiation between biotypes of the pea aphid. The proportions of hybrids both per pair of biotypes (circles) and per biotype (diamonds, with 95% confidence intervals) correlate negatively with genetic differentiation (67.8% variance explained, \(P < 0.001\), Mantel test). Letters denote biotypes as in Fig. 1. Genetic differentiation between 2 biotypes at microsatellite markers (standardized \(F_{SC}\)) was estimated within location by hierarchical analyses of molecular variance. For a given biotype (diamond), \(F_{SC}\) was averaged over the 10 pairwise comparisons involving this biotype.
western Europe and most of Eurasia (references in International Legume Database & Information Service; http://www.ildis.org). They were sampled in grasslands and perturbed areas, where they usually grow in mixtures, or in adjacent monocultures of *Pisum, V. faba*, and some *M. sativa*. The main sampling location measured 40 km in diameter and was centered on Lanteny, eastern France (46°03′ N, 5°32′ E). This area was surveyed in August 2006 and in June and July 2007. Complementary sampling locations were the regions surrounding the towns of Le Rheu, northwestern France (48°06′ N, 1°47′ W) surveyed in June 2006 and Jena, located in eastern Germany (50°55′ N, 11°35′ E), surveyed in July 2006. In each location, wingless aphids from each plant species were sampled in at least 2 sites separated by a distance of several kilometers. No more than 1 individual per plant was sampled and then stored in 95% ethanol.

**Genotyping and Sequencing.** Aphids were genotyped with 2 multiplexes of 7 microsatellite loci (43–46) amplified and analyzed as in ref. 47. *Table S2* details the loci used, the redesign of some primers, and specific conditions of PCR amplification.

In any sample set from the same host species and location, the probability that 2 individuals produced by sexual recombination had the same genotype at all loci, *P*(null) (49), computed with the program GIMLET (48), was <0.1%. Given that aphids reproduce asexually from spring to autumn, this low probability indicated that identical 14-locus genotypes had resulted from parthenogenesis and not from sexual reproduction, which occurs during the fall. Including several individuals of the same genotype in the analyses would be equivalent to using the same individuals multiple times when analyzing strictly sexual populations. To circumvent this problem, we retained only 1 individual per 14-locus genotype (50) in the sample set where the corresponding genotype was most frequent.

Null alleles were suspected at 3 loci (*Table S2*) showing heterozygote deficit and a lack of amplification in rare individuals. These loci were discarded for analyses, except for assignment tests, which are only weakly affected by null alleles (51) and improve with the number of loci used (52).

We amplified and sequenced the noncoding, flanking regions of 3 microsatellite loci (*Table S2*) for each allele found in at least 1 homogenous individual (detected by the genotyping procedure) and in an outgroup, *A. kondoi*. We used a standard protocol described in ref. 47. Three loci were discarded after comprehensive assessment of the genetic diversity found in the pea aphid at these loci, while direct sequencing avoided the risk of cloning PCR products bearing in vitro mutations. Sequences were aligned with the program BioEdit (53), coding contiguous missing nucleotides as single indels. Maximum-likelihood trees (*Fig. S2*) were built by heuristic searches in PAUP* 4.0b10 (54), using default settings and the most likely substitution model determined by Akaika information criteria in Modeltest 3.7 (55).

**Detection of Host-Associated Populations.** Using Structure 2.2 (26–28), we assigned individuals to inferred populations, assuming genetic admixture and correlated allele frequencies between them. We varied parameter *k* (the number of assumed populations) from 2 to 19. Twenty simulations were run for each value of *k* for 2 × 10⁶ Markov Chain Monte Carlo (MCMC) steps, the first 10⁵ being discarded. The posterior probability of the different values of *k* (56) did not outline a more likely number of populations (*Fig. S3*).

We however retained the structuring solution of the run showing higher posterior probability at *k* = 11, because no other host-associated population was detected at higher values of *k* or when separately analyzing the structure of clusters B, E, F, and G, which are associated with several host species.

**Detection of Hybrids.** The parental origins of individuals were inferred one generation backward, with the “use PopInfo” option in Structure, which specifies a prior on their source population. Here, we used the collection plant of an individual as a prior and defined 11 putative populations by their host ranges, on the basis of results from the Structure analysis with admixture (*Fig. 1A*). We assumed prior migration rates of 10% between populations, as previously estimated between North American host races (39) (parameter “MIGRPRIOR”). Other parameters were the same as specified in the admixture model at *k* = 11. Assignment tests on generated individuals of known parental origins presented low error rates (<3%) under these settings, especially for highly differentiated biotypes (*SI Methods* and *Fig. S4*).

We then used the program NewHybrids (31) to identify hybrids in pairs of genetic clusters determined by Structure. We defined only 1 class of hybrids, excluding *F₂* and backcrosses, and used Jeffrey’s priors (31) as these settings proved the most accurate for simulated individuals. The accurate detection of *F₂* and backcrosses usually requires many more loci (52). An individual was considered as a hybrid between 2 biotypes if it was assigned as such with a posterior probability ≥95% in NewHybrids and had intermediate ancestry to

differentiation (standardized Fsc)

[Fig. 3.](#) Adaptive divergence between biotypes of the pea aphid does not correlate with their genetic differentiation. Adaptive divergence is based on host-specialization tests. It represents the Euclidian distance between 2 biotypes from their performance (produced biomass in milligrams) on their 2 representative host plants (see Fig. 1).
these biontbes in Structure. The parental origin of all individuals bearing different genotypes is shown in Table S1.

**Assignment of Migrants.** To genetically assign migrants with confidence, it was necessary to identify individuals potentially belonging to other undetected biontbes specialized on host plants that were not inspected. To detect these outliers, we computed the likelihoods of assignment of all individuals to 11 reference populations with Baudoin and Lebrun's Bayesian criterion (57) implemented in the program Geneclass 2 (30). On the basis of the output of Structure under the Popinfo model (see above), reference populations were defined as genetic clusters comprising only resident aphids (see Table 1), as these were less likely to be outliers. Migrants had likelihood values that were in the range of those of residents and thus showed no evidence that they belonged to undetected biontbes, with the exception of 3 individuals (Fig. S5). These presented clearly lower likelihoods of belonging to their reference population, and they were excluded from Table 1 and from further analyses. This analysis allowed us to delineate biontbes as groups of individuals assigned to a given host-associated population, i.e., to exclude outliers and F1 hybrids.

**Genetic Differentiation.** Genetic differentiation (Figs. 2 and 3) was computed by hierarchical analyses of molecular variance (33) under Arlequin v. 3.11 (32). We structured the data set by location and then by biontte. Considering 2 biontbes at a single time, we computed the within-group (here, within-location) component of molecular variance, FST, which represents the mean genetic differentiation between these biontbes in sympatry. To correct for variations in genetic diversity at our markers, which may result from different mutation rates or effective population sizes among biontbes, FST's were standardized according to Meirmans (58). FST's values for all pairs of biontbes are shown in Table S1.

**Host-Specialization Tests.** Seventy individuals sampled from various plants were transferred to the laboratory and reared on separate broad bean plants (V. faba), a suitable host for all known biontbes of the pea aphid (18), to generate paraphenogenetic lineages. None of them rejected this new host, but 3 of the aphids failed to develop lineages as they were parasitized by hymenoptera.

For performance tests, we selected a subsample of 45 aphid lineages assigned to 10 biontbes and bearing different genotypes, maximizing their diversity in terms of collection plant and location (Fig. S1). Lineages from the same host plant and location were chosen at random. No hybrid lineage was identified in our live collection for testing. Biontte C (on L. corniculatus) was not included because we initially expected the same biontte as that associated with L. pedunculatus. Hence, no plant of L. corniculatus was grown for performance tests.

Chosen lineages were maintained on broad bean plants at low density for at least 3 generations. Using a protocol modified slightly from ref. 47, we installed 10 first instar nymphs of each lineage on a potted plant (measuring at least 20 cm in height) in a climatic chamber (17 °C, 16-h photophase). After 9 days, surviving aphids, which had typically reached early adult stage, were weighed with a precision of 10⁻⁶ g. Three replications per lineage per plant species were carried out in a climatic chamber.

As a proxy for performance, we used the total biomass produced by a lineage i on a given plant species a (hereafter termed Mia), combining its survival and growth after 9 days, and averaged over the 3 replicates. By doing so, lineages were each described by 10 performance variables (on each test plant). A principal component analysis (PCA) was computed on this data set. The hierarchical ascendant classification (HAC) of lineages (Fig. 1B) was based on the first 4 components of the PCA. The adjusted Ward's criterion of aggregation (59), which consists of merging iteratively the 2 lineages or groups of lineages with the lowest increase in intragroup variance and the lowest decrease in intergroup variance of the partition. This process yields a binary segmentation tree reflecting the hierarchy of similarities between lineages and wherein branch lengths represent the decrease of variance involved in the creation of a node. This analysis was done using SPAD 6 (http://spadssoftware.com).

The Euclidian distance between 2 lineages based on their performances when reared on their 2 typical host species was computed as

\[ ((M_{ia} - M_{ja})^2 + (M_{ib} - M_{jb})^2 \]

whereby lineages i and j belong to different biontbes that are specialized on host plants a and b, respectively. The Euclidian distance between 2 biontbes (Fig. 3) was averaged over all pairwise comparisons between lineages.

The correlation between ecological and genetic differentiation (FST, previously computed in Arlequin) was tested using program FSTAT v. 2.9.3.2 (60).

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