

Molecular Differentiation at Nuclear Loci in French Host Races of the European Corn Borer (*Ostrinia nubilalis*)

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ABSTRACT

French populations of the European corn borer consist of two sympatric and genetically differentiated host races. As such, they are well suited to study processes that could be involved in sympatric speciation, but the initial conditions of host-race divergence need to be elucidated. Gene genealogies can provide insight into the processes involved in speciation. We used DNA sequences of four nuclear genes to (1) document the genetic structure of the two French host races previously delineated with allozyme markers, (2) find genes directly or indirectly involved in reproductive isolation between host races, and (3) estimate the time since divergence of the two taxa and see whether this estimate is compatible with this divergence being the result of a host shift onto maize after its introduction into Europe ~500 years ago. Gene genealogies revealed extensive shared polymorphism, but confirmed the previously observed genetic differentiation between the two host races. Significant departures from the predictions of neutral molecular evolution models were detected at three loci but were apparently unrelated to reproductive isolation between host races. Estimates of time since divergence between French host races varied from ~75,000 to ~150,000 years, suggesting that the two taxa diverged recently but probably long before the introduction of maize into Europe.

THE European corn borer (ECB), *Ostrinia nubilalis* Hübner (Lepidoptera: Crambidae), exists as a number of sympatric, genetically differentiated but partially interfertile taxa across its geographical range in northern Eurasia and northern America (HUDON *et al.* 1989). As such, it is a well-suited biological model to study processes that could be involved in the early steps of speciation, especially sympatric and ecological speciation (SCHLUTER 2001; VIA 2001; BERLOCHER and FEDER 2002; GAVRILETS 2003; RUNDLE and NOSIL 2005).

The ECB’s area of origin is thought to be central Asia (HUDON *et al.* 1989). Its larvae feed on a large number of wild and cultivated host plants (CAFFREY and WORTHLEY 1927; HODGSON 1928; PONSARD *et al.* 2004), but it is

commonly found on maize (*Zea mays* L.). ECB populations collected as larvae on maize stalks in certain locations in Europe (PEÑA *et al.* 1988) and in North America (KLUN 1975; KOCHANSKY *et al.* 1975; GLOVER *et al.* 1990) are polymorphic with respect to sex pheromone communication. On the basis of this polymorphism, two pheromonal races can be distinguished: the Z-race using a 97:3 blend of Z:E isomers of the 11-tetradecenyl acetate (11-14:OAc), and the E-race using blends of Z11-14:OAc and E11-14:OAc in a 1:99 to 4:96 ratio (KLUN 1975). The percentage of hybrid moths is lower than expected under random mating between E- and Z-race moths, suggesting partial reproductive isolation between the two pheromonal races (GLOVER *et al.* 1990).

In contrast to other locations in Europe and North America, ECB pheromonal races in France can be distinguished according to the host plant(s) they use (BOURGUET *et al.* 2000; MARTEL *et al.* 2003). Indeed, in France, but not in the Balkans, Italy, and North America,

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maize is infested exclusively by moths using the Z sex pheromone blend (PÉLOZUELO *et al.* 2004), whereas hop (*Humulus lupulus* L.) and mugwort (*Artemisia vulgaris* L.) are infested exclusively by moths using the E pheromone blend (THOMAS *et al.* 2003; PÉLOZUELO *et al.* 2004). These host races have thus been referred to as the "maize race" and the "mugwort–hop race," respectively. The percentage of hybrids between the two host races is lower than those observed in the United States between the two pheromonal races (MALAUSA *et al.* 2005), suggesting that the pheromone difference is probably not the only factor serving to isolate the maize race from the mugwort–hop race.

Given that maize was introduced into Eurasia only ~500 years ago and has become a major host plant of many ECB populations worldwide, it is reasonable to assume that populations of the maize race have adapted recently to this host plant (THOMAS *et al.* 2003). It is also tempting to hypothesize that the substantial reproductive isolation and genetic divergence between the two French host races is the result of a host shift onto maize followed by host-plant specialization. Indeed, a set of ecological features that might constitute host-plant adaptations, such as emergence timing (THOMAS *et al.* 2003) and oviposition preferences (BETHENOD *et al.* 2005; MALAUSA *et al.* 2007), have been found to distinguish both host races in France. However, as with other well-documented host-race models [*e.g.*, *Rhagoletis pomonella* (FEDER *et al.* 2003), *Zeiraphera diniana* (EMELIANOV *et al.* 2004), *Acyrtosiphon pisum* (VIA 1999), and *Enchenopa binotata* (WOOD *et al.* 1999)] the exact circumstances of the divergence between ECB races remain elusive.

The genetic divergence between the two pheromonal races on the one hand and the two host races on the other hand was first investigated by means of allozyme markers. American surveys were performed on >30 loci (HARRISON and VAWTER 1977; CARDÉ *et al.* 1978; CIANCHI *et al.* 1980) while 6 loci were analyzed in France (BOURGUET *et al.* 2000; MARTEL *et al.* 2003; THOMAS *et al.* 2003; BONTEMPS *et al.* 2004). Major differences in allelic frequencies were detected between the American pheromonal races at only 1 locus, *Tpi* (triose phosphate isomerase), and between French host races at 2 loci: *Tpi* and *Mpi* (mannose phosphate isomerase). More recently, the divergence between North American ECB pheromonal races was investigated using a gene genealogical approach based on DNA sequences at 4 nuclear loci, including *Tpi*, and at the mitochondrial *COI* locus (DOPMAN *et al.* 2005). Only *Tpi* showed clear differentiation of the two pheromonal strains: all haplotypes of the E-race, although sampled from two quite distant locations, formed a very homogeneous cluster that did not include any Z-race haplotype. For the mitochondrial *COI* gene, the results from DOPMAN *et al.* (2005) for American populations are in agreement with those of MARTEL *et al.* (2003), who did not find any differentia-

tion within or between French ECB populations at this locus.

No other genetic data exist on ECB populations collected in France. As a consequence, the delineation of both host races remains incomplete and the hypothesis of divergence triggered by a host shift of some populations onto maize has never been tested. Moreover, no gene has been detected to be involved or located in a chromosomal region involved in host-plant adaptation and reproductive isolation of French host races. This work aims at examining nuclear DNA sequence polymorphism in French ECB populations collected on maize, mugwort, hop, pepper, cocklebur, sunflower, and sorghum. Specifically, our goal is to determine whether the distribution of polymorphisms (1) confirms the host-race delineation found with allozyme markers and possibly even reveals additional genetically differentiated taxa, (2) reveals patterns of monophyly in the host races or selective effects acting at some loci, and/or (3) provides an estimate of divergence time compatible with the scenario of host-race formation by host shift onto maize after its introduction into Europe ~500 years ago.

MATERIALS AND METHODS

Insect samples: Samples of ECB populations were collected over the course of 2002 and 2003 as fifth larval instars from a total of seven plant species and 27 populations throughout France (Figure 1). In three cases (Muret, Pierrelatte, Grignon), populations from different plants were collected in the same sites. The sex of each individual was determined. Only females were used, as female Lepidoptera are heterogametic (ZW), *i.e.*, haploid at all loci on the Z chromosome, while males are diploid (ZZ). We aimed at collecting ~10 females for each population. Samples included populations collected from maize ($N = 93$, 10 populations), mugwort ($N = 52$, 5 populations), hop ($N = 43$, 4 populations), sorghum *Sorghum* sp. ($N = 22$, 2 populations), cocklebur *Xanthium* sp. ($N = 20$, 2 populations), pepper *Capsicum frutescens* L. ($N = 37$, 3 populations) and amaranth *Amaranthus* sp. ($N = 11$, 1 population). Populations collected from mugwort could be collected only in the northern part of France, as this plant species does not appear to be infested in the southern part of this country (MARTEL *et al.* 2003). The geographic distances between all pairs of sampling sites were estimated as linear distances, estimated in kilometers, between points plotted on a map of France (using Google Earth at <http://earth.google.com/>).

Nuclear loci: Sequence data were obtained from four nuclear loci: *Mpi*, *Tpi*, *Pbp*, and *Ket*. *Mpi* and *Tpi* are both enzymes of carbohydrate metabolism. *Pbp* encodes a pheromone binding protein, which displays affinity to certain pheromone components in insect species (DU and PRESTWICH 1995). *Ket* encodes kettin, a high molecular weight protein of the insect flight muscle (KOLMERER *et al.* 2000). *Tpi* and *Ket* loci are not thought to have relevance for host selection or pheromone production and response, but they are located on the Z sex chromosome that also carries genes affecting male behavioral response to female sex pheromones (*Resp*) and the major factor of postdiapause development time (*Pdd*) (DOPMAN *et al.* 2004, 2005).

DNA extraction, amplification, and sequencing: DNA extraction followed the CTAB protocol (DOYLE and DOYLE

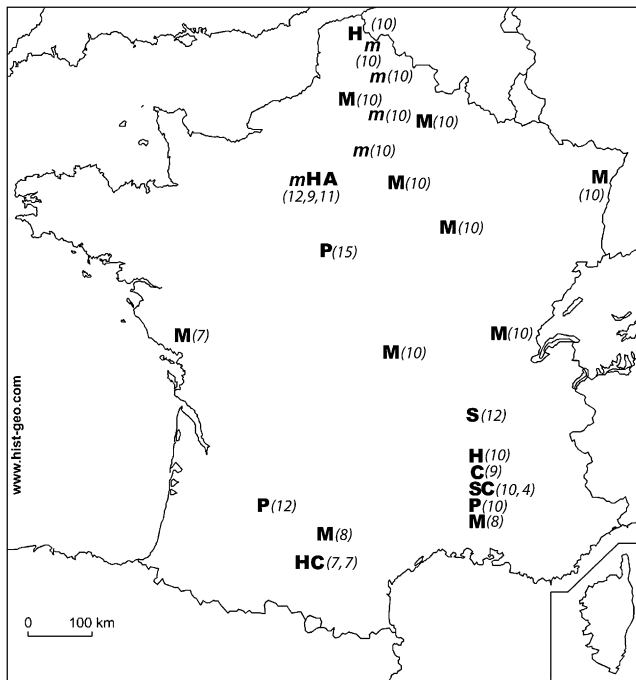


FIGURE 1.—Location of sampling sites. Host plants: M, maize; m, mugwort; P, pepper; H, hop; C, cocklebur; S, sorghum; A, amaranth. The number of females sampled in each population is indicated in parentheses. Letters grouped on the same line indicate sites where several plants, located <10 km from each other, were sampled.

1987). An ~1600-bp DNA fragment including the coding region (5 exons of a total length of 732 bp) and introns (~870 bp) of *Tpi* was amplified using primers ECBtpi_for1A and ECBtpi_rev5 (5'-ATAGTTTACGAATTACGAGTT-3') (DOPMAN *et al.* 2005) and PCR profile: 1.5 min at 95°, 35 cycles of 1 min at 95°, 30 sec at 55°, and 1 min at 72°, followed by 3 min at 72°. An ~1400-bp fragment of *Ket*, including a 232-bp exon, was amplified using primers ECBketF and ECBketR (DOPMAN *et al.* 2005) and PCR profile: 1.5 min at 95°, 35 cycles of 1 min at 95°, 30 sec at 58°, and 1 min at 72°, followed by 3 min at 72°. An ~400-bp fragment of *Mpi*, including two exons of 60 and 77 bp, was amplified with primers Mpi-15 and Mpi-16 (LENIAUD *et al.* 2006) and PCR profile: 1.5 min at 95°, 35 cycles of 30 sec at 95°, 30 sec at 60°, and 30 sec at 72°, followed by 3 min at 72°. Finally, an ~1650-bp DNA fragment, including *Pbp* and its three exons of 126, 181, and 182 bp, was amplified with primers ECEP5 and ECPA (WILLETT and HARRISON 1999) and PCR profile: 1.5 min at 95°, 35 cycles of 30 sec at 95°, 30 sec at 60°, and 1 min at 72°, followed by 3 min at 72°. Raw sequences of all loci were submitted to GenBank under the accession nos. EF396347–EF396478.

Because *Tpi* and *Ket* are located on the Z chromosome and because we used only female ECB in our study, PCR products for these two genes were sequenced directly. *Tpi* PCR products were sequenced by Genome Express S.A. (Meylan, France). *Ket* PCR products were purified with 2 μ l of Exosap (USB, Cleveland) solution for 5 μ l of PCR product and sequenced with a General Electrics Healthcare automatic sequencer using the BigDye terminator kit 3.1 (General Electrics Healthcare, Little Chalfont, UK). Sequencing primers were ECBtpi_for1A and ECBtpi_rev5 for *Tpi* and ECBketF for *Ket*. Of 278 individual samples, we obtained 222 sequences unambiguously readable over the full length of the fragment for *Tpi* and 219 such sequences for *Ket*.

Because *Mpi* and *Pbp* are located on autosomes, PCR products containing parts of these two genes were cloned before sequencing. A total of 94 individuals were chosen among those sequenced for *Tpi* and *Ket* by randomly picking 2–3 individuals from each population. PCR products obtained from these individuals were sent to Genoscreen (Lille, France) for cloning using the TOPO TA cloning kit (Invitrogen, Carlsbad, CA). PCR products were ligated in 96-well plates following the TA cloning kit protocol (2004) and transformed with the one-shot Top10 competent cells provided with the TA cloning kit, in 96-well plates (Millipore plasmid miniprepkit LSK P09624, Millipore, Billerica, MA). The transformation products were spread on an Luria Burani (LB)-ampicillin (100 μ g \cdot μ l⁻¹) solidified solution contained in petri dishes (AES–AEB521279). Individual colonies were then each transferred into a well of a 96-well plate (Millipore plasmid miniprepkit LSK P09624) and maintained in a liquid LB-ampicillin solution (100 μ g \cdot μ l⁻¹). Plasmids were extracted with a miniprepkit (Millipore) and sequenced with an Applied Biosystems 3730 XL automatic sequencer. Sequence reactions used the BigDye 3.1 kit (Applied Biosystems, Foster City, CA). *Mpi* clones were sequenced using the two primers provided in the TOPO TA cloning kit (Invitrogen). *Pbp* clones were sequenced using the forward 5'-GACAAAC TAAATTGTTAGGCA-3' and the reverse 5'-GATTCCAACCTCC ATCGCAT-3' primers, which were designed by Genoscreen to improve the quality of final sequences by increasing the length of the overlapping zone between forward and reverse sequences (giving a final complete sequence of ~1300–1400 bp).

For both *Mpi* and *Pbp*, four clones were initially sequenced for each individual, with the hope of unambiguously determining at least one allele per locus. However, for ~30% of the individuals, the sequencing of these four clones yielded more than two different haplotypes, a problem also reported for *Pbp* by WILLETT and HARRISON (1999). In a diploid organism, this can be due to (1) multiple copies of the gene in the genome, (2) Taq DNA polymerase errors, or (3) *in vitro* recombination during the PCR. Taq errors and *in vitro* recombination are known to occur [“jumping PCR,” see PAABO *et al.* (1990); BRADLEY and HILLIS (1997)]. We directly sequenced PCR products of individuals for which cloning had yielded more than two haplotypes and for which no length polymorphism among clone sequences was detected. We considered that Taq errors were the most likely explanation when substitutions observed in clone sequences were absent from direct PCR product sequences of the same individual. For all but five (when considering both *Pbp* and *Mpi*) of the remaining individuals, manual inspection of the sequences showed that the set of haplotype sequences obtained was compatible with a scenario of *in vitro* recombination, *i.e.*, we found that the additional haplotypes found in PCR products at low frequencies consisted of a combination of successive fragments of the two sequences. In the five remaining individuals, we could not assess the compatibility of sequences with an *in vitro* recombination scenario, due to too large a number of Taq errors (including errors at polymorphic sites). For *Pbp*, we concluded that Taq errors occurred in ~20% of the clones and *in vitro* recombination in ~10% of the clones. For *Mpi*, these percentages were ~10 and ~5%, respectively.

To choose which haplotypes to keep for subsequent analyses, we proceeded as follows. When the initial four to eight clone sequences revealed only one or two haplotypes for a given individual, we used the most common sequence provided it was observed at least three times. When this number was not reached, we either sequenced additional clones when length polymorphism was observed or directly sequenced the PCR products when no length polymorphism was detected. For all individuals showing more than two haplotypes and no insertion/deletion (indel) in the region responsible for the

existence of these apparently more than two haplotypes, we directly sequenced the PCR products and kept the clone sequence(s) that unambiguously matched the PCR product sequence. For individuals with fragment length polymorphisms for which direct sequencing did not help to distinguish true haplotype(s) from the artifact(s), we sequenced up to 12 additional clones and used only the sequence found in the majority of clones, provided this majority consisted of three or more clones. All other sequences were discarded.

This procedure resulted in a much smaller data set than we were initially hoping to gather: of the 94 individuals (*i.e.*, 188 alleles) initially included in the cloning protocol, 79 and 83 individuals finally yielded one usable sequence each for *Pbp* and *Mpi*, respectively.

Data analysis: Sequences were aligned using CLUSTALW 1.83 (THOMPSON *et al.* 1994) and adjusted manually using BIOEDIT 7.01 (HALL 1999). The correspondence between sequences and chromatograms was checked manually using CHROMAS 2.3 (Technelysium Pty, Tewantin, Queensland, Australia). All sequences were blasted (March 6, 2006) using the BlastN and BlastX functions of the NCBI-BLAST website (<http://www.ncbi.nlm.nih.gov/BLAST/>), which confirmed that they all corresponded to the expected locus.

For *Tpi*, *Ket*, and *Pbp*, several complex sets of indels were observed, corresponding to substitutions and length polymorphism within the indels. As the sequences were long and the number of informative sites was high, we excluded indels from our analysis. For *Mpi*, we failed to align the entire data set because a region of ~200 bp (of 300–400 bp, depending on the length of the indels) proved extremely variable. Hence, we restricted our analysis to the ~160-bp region for which the sequences could be aligned. Delineation of coding and non-coding regions as well as positions of variable sites in the final alignments is shown for each gene in the supplemental online material (<http://www.genetics.org/supplemental/>).

We used DNASP 4.01 (ROZAS *et al.* 2003) to calculate the following summary statistics describing our data set: number of haplotypes (H), haplotype diversity (H_d), nucleotide diversity estimated by π (NEI 1987) and θ (WATTERSON 1975), and minimum number of recombination events (R_{m}). These descriptive statistics were estimated for the entire data set and for data sets restricted to populations collected on the same host-plant species.

Mutation models (Jukes and Cantor, Kimura 2P, Tajima and Nei, Tamura, Tamura and Nei) available in ARLEQUIN 3.01 (EXCOFFIER *et al.* 2005) were fitted to the data and their likelihood was tested for each locus using the MODELTEST 3.7 program (POSADA and CRANDALL 1998). Tamura and Nei's model was retained for *Tpi* (uniform substitution rates among sites), *Ket* (uniform substitution rates), and *Pbp* (different substitution rates, $\gamma = 0.77$), and Kimura's 2P was retained for *Mpi* (different substitution rates, $\gamma = 0.253$). To look for possible clustering by host plant, neighbor joining (NJ) trees (inferences tested by a bootstrap procedure of 1000 replications) were constructed with MEGA 3.1 software (KUMAR *et al.* 2004) for each locus, using the most appropriate mutation model as determined by MODELTEST 3.7.

The level of genetic differentiation between groups of populations collected on the same host-plant species was assessed by calculating among- and within-group fixation indices (F_{CT} and F_{SC}) in an analysis of molecular variance (AMOVA) using ARLEQUIN 3.01. Again, the most appropriate mutation model to be used in the AMOVA was chosen for each locus according to prior tests using MODELTEST 3.7. The P -values associated with among-group fixation indices were obtained by permutation tests performed by ARLEQUIN 3.01 (10,000 permutations). As the number of available sequences varied among loci, the mean F_{CT} value across all four loci could not be

obtained using the standard procedure of ARLEQUIN 3.01. Instead, we calculated the mean F_{CT} value over the four loci and combined the corresponding P -values using Fisher's method (FISHER 1932).

We also tested for a correlation between the matrices of pairwise genetic and geographic distances with Mantel tests performed with GENEPOP 3.4 (RAYMOND and ROUSSET 1995). Pairwise genetic distances were calculated for each locus by computing pairwise F_{ST} in the "Population Comparisons" section of ARLEQUIN 3.01 (the haplotype distance matrices were computed using mutational models chosen according to the goodness of fit assessed with MODELTEST 3.7). The average of this pairwise genetic distance was calculated as the average of the four distance values estimated at the four loci for the pair of populations considered. This analysis was conducted using data on populations sampled from (1) maize, (2) mugwort, or (3) all host-plant species. Populations collected from plants other than maize and mugwort were not numerous enough to test for isolation by distance within each data subset.

Possible departures from the standard neutral model of molecular evolution—potentially revealing demographic events or the existence of selective effects at certain loci—were tested for each locus by Tajima's D -test, Fu and Li's D^* -test, and Fu's F_s -test using DNASP 4.01. Neutrality tests were performed on the entire data set and on data subsets each consisting of populations collected on the same host-plant species, provided at least three populations collected on this host plant were available. Thus, tests on separate host-related groups were conducted only for maize, pepper, mugwort, and hop. P -values associated with these tests were obtained from coalescent simulations performed by DNASP 4.01 taking into account estimates of the intragenic recombination parameter R . Following Fu (1997), the P -value limit for significance in Fu's F_s tests was set to $P = 0.02$ instead of 0.05. The signature of selective constraints on coding regions was tested using the CODEML program of the PAML package (YANG, 1997) by comparing likelihood of models where the ratio of synonymous and nonsynonymous substitutions ($d_{\text{N}}/d_{\text{S}}$) was either fixed to 1 (neutral) or let as a free parameter.

The time elapsed since the initial divergence between populations was estimated using the updated version of the Isolation Model (IM) program released July 31, 2006 (see HEY and NIELSEN 2004 for details on the initial release) on the subset of sequences obtained for larvae collected on maize and mugwort. Again, only sequences obtained for individuals collected on one of these two host plants were used because the individuals collected on the other host plants could not be assigned to one of the host races with complete certainty. *Tpi* and *Mpi* were excluded from this analysis as neutrality tests revealed a significant departure from neutral evolution at these loci in at least one population (see RESULTS). Since intragenic recombination was detected, we followed HEY and NIELSEN's (2004) advice and used, for any given locus, only the largest region of the sequence that was not subject to recombination. The presence of recombination between sites was checked using the HUDSON and KAPLAN algorithm (1985) included in DNASP 4.01. We analyzed two data sets: both included the longest portion (511 bp) of *Ket* (where intragenic recombination was rare), but they differed in the region of *Pbp* that was used (158 bp in one data set and 194 bp in the other). Summary descriptive statistics for these two data sets are detailed in supplemental online material (<http://www.genetics.org/supplemental/>). Although IM did not reject the infinite site mutation model for the two loci, results of MODELTEST 3.7 clearly indicated that the HKY model (Hasegawa–Kishino–Yano; HASEGAWA *et al.* 1985) provided a better fit to the data for both *Ket* and *Pbp*. Thus, we decided to use this latter mutation model for IM runs. First runs used parameter values recommended

TABLE 1

Descriptive statistics of polymorphism found at each of the four nuclear loci, for the entire data set

Locus	<i>N</i>	<i>H</i>	<i>H_d</i>	π	θ	<i>P_s</i>	<i>R_m</i>
<i>Tpi</i>	222	28	0.740 (0.023)	0.00166 (0.00017)	0.00358 (0.00062)	33	2
<i>Ket</i>	219	23	0.893 (0.009)	0.00407 (0.00017)	0.00650 (0.00130)	25	4
<i>Pbp</i>	79	38	0.940 (0.016)	0.02101 (0.00086)	0.02164 (0.00184)	139	14
<i>Mpi</i>	83	17	0.902 (0.013)	0.04935 (0.00110)	0.03151 (0.00630)	25	2

Standard deviation of estimates is in parentheses. *N*, number of sequences; *H*, number of haplotypes; *H_d*, haplotype diversity; π , pi nucleotide diversity; θ , theta nucleotide diversity; *P_s*, number of polymorphic sites; *R_m*, minimum number of recombination events.

by HEY and NIELSEN (2004) for priors of upper bounds of population size theta, migration rate *m*, and divergence time *t*. Following the advice of the IM manual (version of July 31, 2006), convergence and mixing of runs were evaluated by examining (1) the level of autocorrelation between final and initial parameter values, (2) the estimated value of the effective sample size (ESS), (3) the parameter trends plots, and (4) the consistency of results over different runs using the same model (three runs were performed for each data set). Data were not informative enough to estimate all parameters of complex models including different values of theta, *m*, and population size fluctuations. Runs with such models did not converge and/or resulted in flat posterior distribution for *m* and *t*. We thus used the simplest possible model, consisting of one ancestral population splitting into two populations of same size and experiencing symmetric gene flow. In final runs, priors were set to 8 for theta, 10 for *t*, and 75 for *m*. Optimal chain swapping rates and ESS numbers were obtained with 20 chains and the following heating scheme: “-f g -g1 0.7 -g2 0.9.” The burn-in period was set to 500,000 iterations and convergence of runs was reached after a duration composed of between ~7 and ~12 million iterations. Finally, the “-p5” option was used to have a closer look at gene flow between populations by estimating the maximum likelihood number of migration events and the mean time at which they occurred in genealogies that had at least one migration event. Estimates of the mean mutation rate per gene per generation and of the number of generations per year are required to convert results into demographic units. Mutation rates per site in nuclear genes are generally expected to be ~10⁻⁹ per nucleotide site per generation, or 10⁻⁶ per (entire) gene per year (FUTUYMA 1998; RIDLEY 2004). To be conservative we used a mutation rate of 10⁻⁸ per site per generation, which corresponded to a value of ~2.8 × 10⁻⁶ and ~3.1 × 10⁻⁶ per gene per generation for data set A and data set B, respectively. Again, to be conservative, as the ECB can achieve two generations per year, we chose to use a generation time (*g*) of 0.5 year. This resulted in the mutation rate per gene per year (μ) being ~5.6 × 10⁻⁶ and ~6.2 × 10⁻⁶ for data sets A and B, respectively.

RESULTS

All loci were polymorphic, but the extent of polymorphism varied among loci (Table 1). Haplotype diversity was comparable at *Ket*, *Pbp*, and *Mpi* (~0.9, Table 1) and markedly lower at *Tpi* (~0.7, Table 1). Nucleotide diversity, whether estimated by π or θ , was an order of magnitude lower at *Tpi* and *Ket* (10⁻³) than at *Pbp* and *Mpi* (10⁻²). Intra-genic recombination (*R_m*, HUDSON and KAPLAN 1985) was detected at all four loci, although at

quite different frequencies (Table 1). Descriptive statistics within populations collected from each plant are available as supplemental online material (<http://www.genetics.org/supplemental/>).

Differentiation between populations: Over all populations and all loci, individuals did not clearly cluster according to their host plant in NJ trees (Figure 2). Rather, populations collected from any pair of plants typically shared several haplotypes. The only exception was one *Pbp* haplotype that was shared by 43% of the individuals collected from mugwort but occurred much less frequently, if at all, in populations collected from other host-plant species.

Fixation indices (Table 2) essentially showed that the population structure detected in French populations with allozyme markers (BOURGUET *et al.* 2000; MARTEL *et al.* 2003; LENIAUD *et al.* 2006) was also detectable at DNA sequence level. Significant genetic divergence was found between populations collected on mugwort and populations collected on any of the other host plants except hop. Conversely, populations collected on maize did not show any significant divergence from populations collected on any other host plant except mugwort. Moreover, divergences from populations collected on mugwort (estimated by *F_{CT}* over all genes) were of similar magnitude for populations collected on maize, cocklebur, pepper, and sorghum. No significant divergence was detected in the present study between populations collected on hop and those collected on maize. Restricting the AMOVA to northern France (Table 2) did not change the conclusion about populations feeding on maize and mugwort being significantly divergent and about populations feeding on mugwort and hop not being significantly divergent. However, in this region, significant divergence between populations from maize and hop was detected.

Mantel tests revealed no correlation between matrices of pairwise genetic and geographic distances, be it with populations collected from mugwort (*P* = 0.141), from maize (*P* = 0.438), or from all host-plant species (*P* = 0.477).

Selective effects: Neutrality tests performed on the entire data set showed that *Tpi* and *Mpi* were not at mutation-drift equilibrium. Indeed, except for the Fu

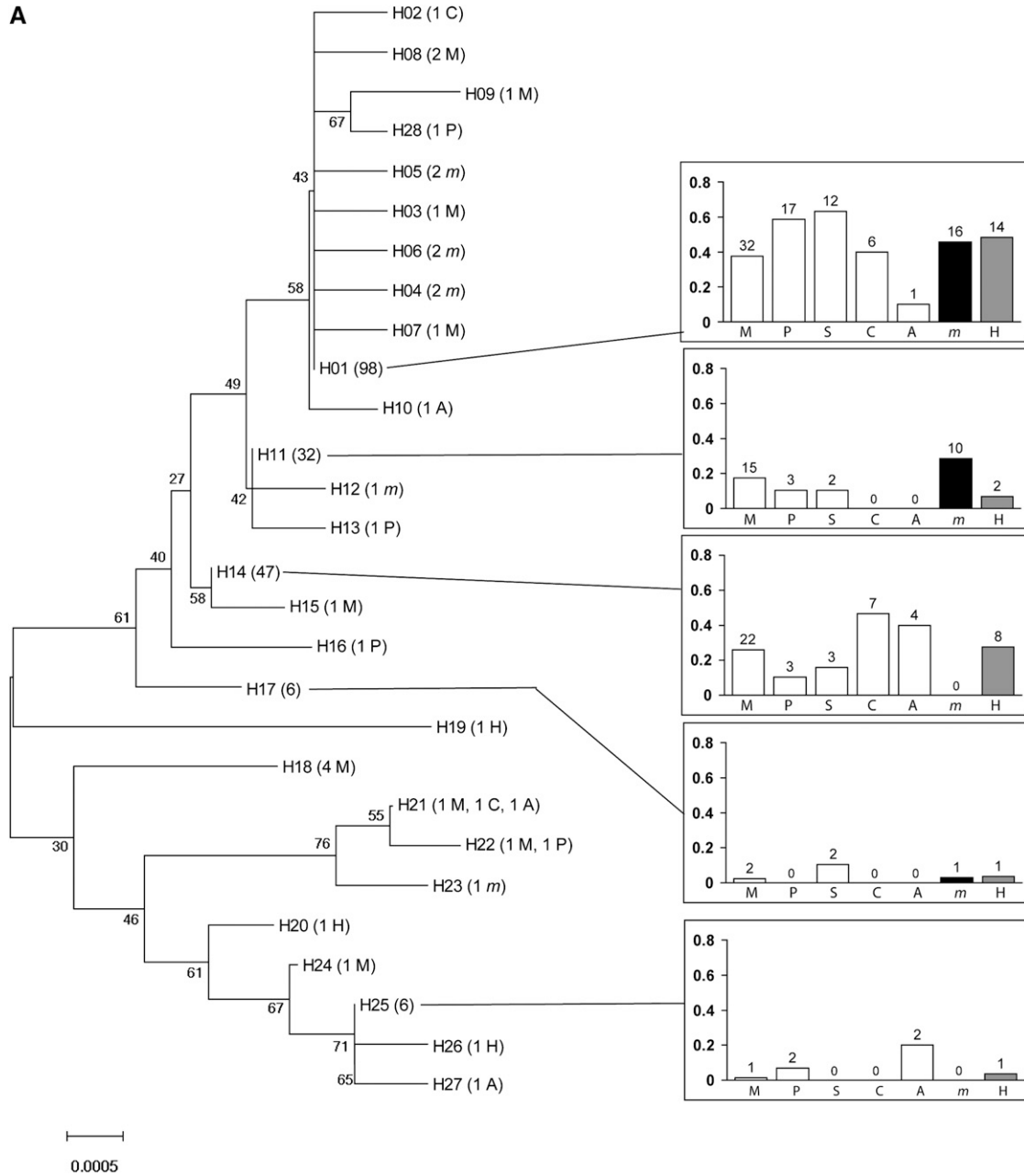


FIGURE 2.—Neighbor joining trees for the four loci: (A) *Tpi*, (B) *Ket*, (C) *Pbp*, (D) *Mpi*. H01–H30: haplotype number, followed by the number of individuals carrying each haplotype (in parentheses) and the host plant(s) from which these individuals were collected (names are coded as in Figure 1). For haplotypes shared by more than four individuals sampled from more than four different plants, a diagram (right) represents (i) among all individuals collected from a given plant, the proportion sharing the haplotype (bars) and (ii) the corresponding number of individuals from this plant (above bars). When sequences were available, *Ostrinia furnacalis* (the Asian corn borer, ACB) was used as an outgroup.

and Li's D^* for *Mpi*, values of the three statistics were all significantly different from zero for *Tpi* and *Mpi* (Table 3). *Tpi* was characterized by negative values, whereas *Mpi* displayed positive values. For *Ket*, the values of the three statistics were negative but none of them were significantly different from zero. For *Pbp*, D^* , D , and F_s values were very close to zero with no particular trend. Results of the analyses performed on each host-plant-specific data set were consistent with these global trends, except

in hop populations for *Ket* (Table 3). Finally and noteworthy, the three different test statistics used in this analysis lead to consistent results (Tajima's D revealed five while the two other tests revealed four significant departures from neutral evolution) (Table 3). d_N/d_S was significantly different from one in the coding regions of *Tpi* ($d_N/d_S = 0.22$) and *Pbp* ($d_N/d_S = 0.11$) but not in those of *Mpi* ($d_N/d_S = 0.44$) and *Ket* ($d_N/d_S = 0$; only three sites were variable).

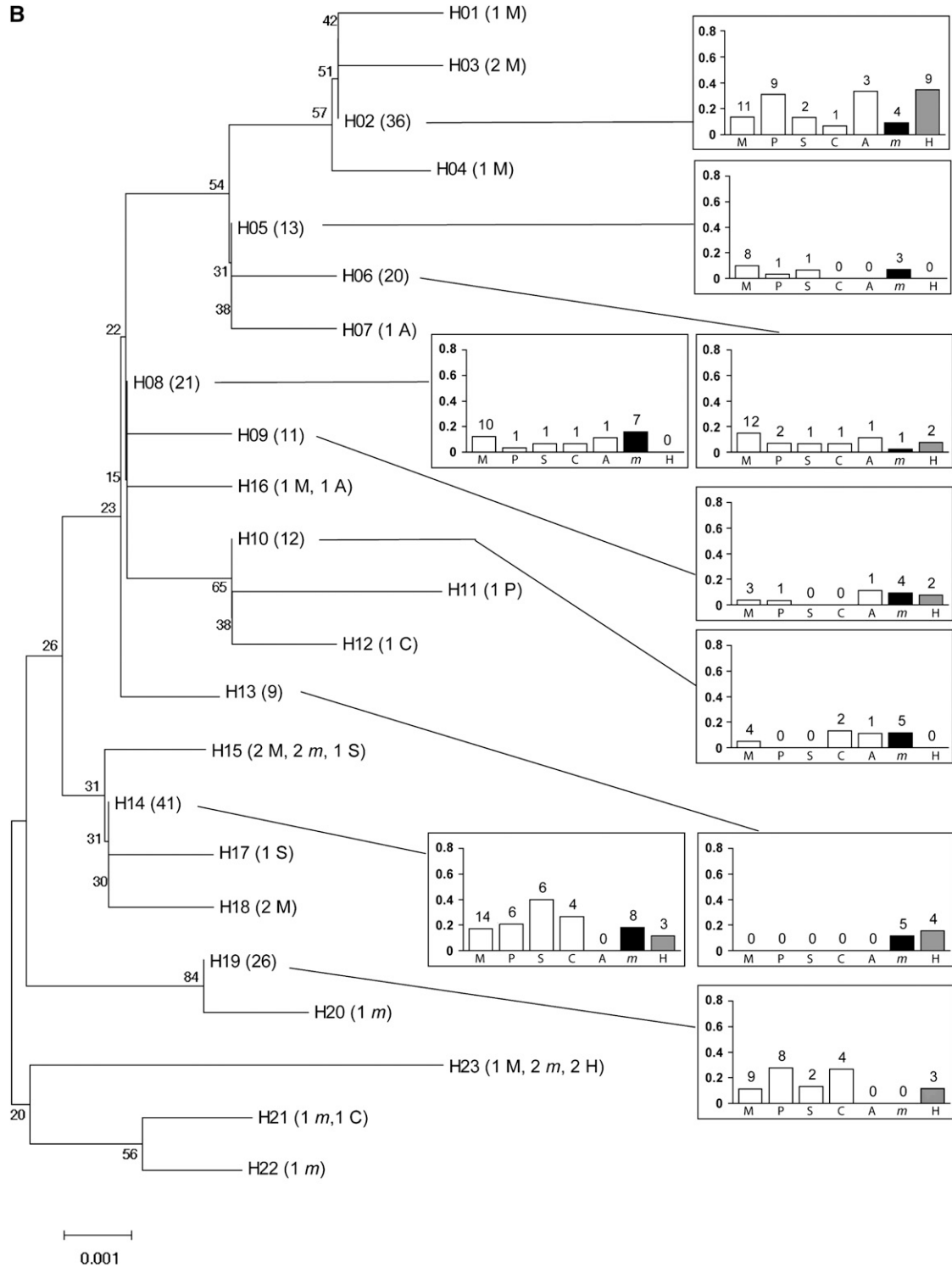


FIGURE 2.—Continued.

Divergence time between populations: Overall, results were consistent over data sets and replicates (Figure 3, Table 4), albeit quite imprecise, as posterior distributions were long-tailed for t and m . The effective population size theta estimate with the highest maximum likelihood was consistently $\sim 150,000$ – $200,000$ individuals (see Table 4 for estimates of the 90% highest posterior density in-

terval, the HPD90% interval, resulting from the different runs). The estimated divergence time t varied between $\sim 75,000$ and $\sim 150,000$ years. Despite a large HPD90% interval, in no instance did its lower bound include values compatible with a divergence time of 500 years. In all cases, the likelihood values attributed to this divergence time were extremely low. An unrealistically high ($>10^{-4}$

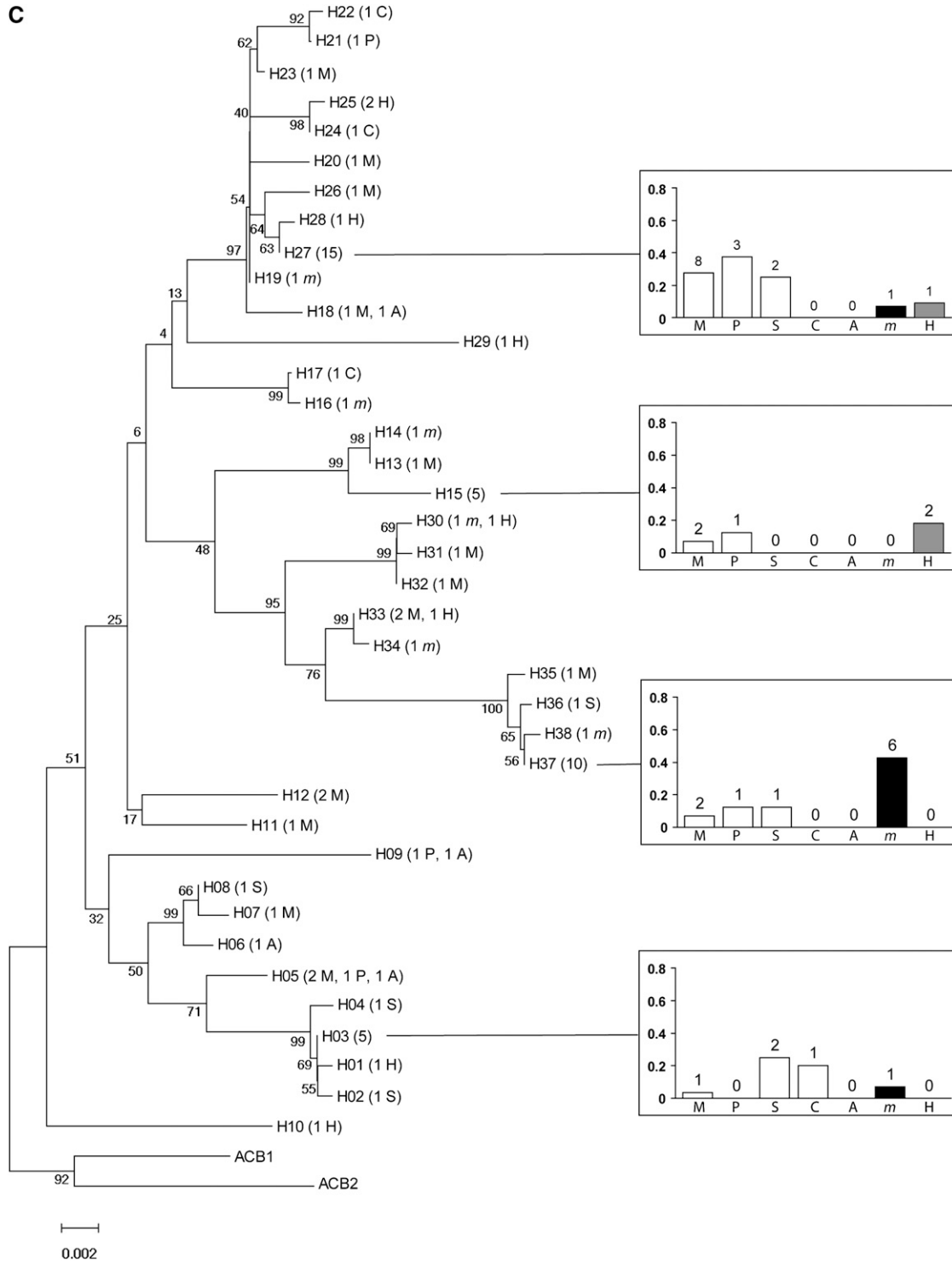


FIGURE 2.—Continued.

and $>5 \times 10^{-5}$ for data sets A and B, respectively) mutation rate per gene per year would be necessary for the lower bound of the HPD90% interval of the divergence time to include the value of 500 years. Estimates for the migration rate m ranged from 7 to 11 and HPD90% intervals were large (Table 4). Finally, estimates for $2Nm$ between the two taxa were relatively high, *i.e.*, ~ 10 .

DISCUSSION

The genetic structure at the four nuclear loci we studied (*i.e.*, *Tpi*, *Ket*, *Mpi*, and *Pbp*) was significantly influenced by the host plants on which the ECB populations were collected (Table 2). ECB populations collected on mugwort were significantly differentiated

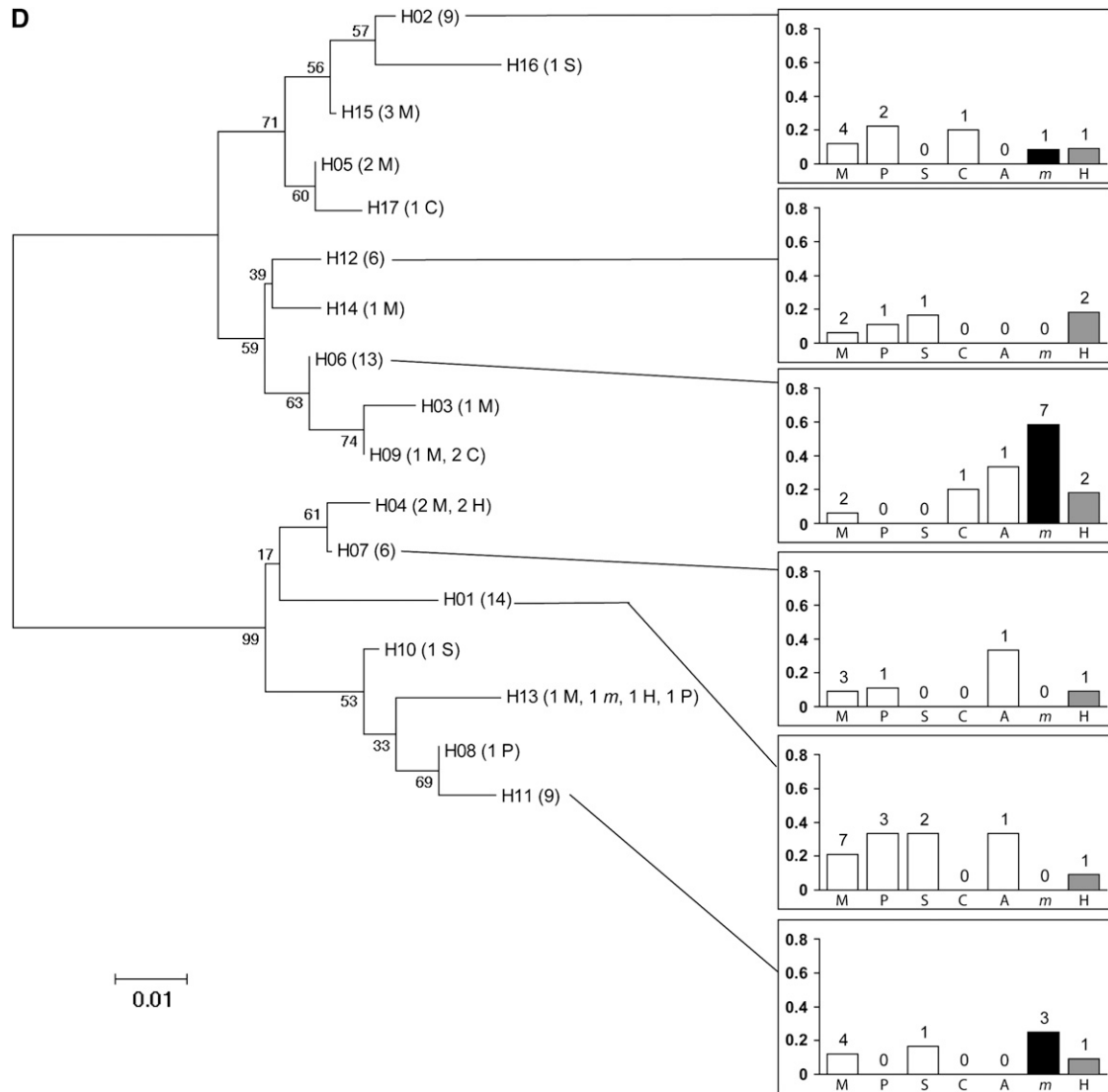


FIGURE 2.—Continued.

from those collected on maize. In addition, populations collected on pepper, sorghum, and cocklebur were genetically more similar to those feeding on maize than to those feeding on mugwort. Hence, our former conclusion based exclusively on allozyme markers also holds when investigating polymorphism at the DNA sequence level: in France, ECB populations are divided into at least two host races (BOURGUET *et al.* 2000; MARTEL *et al.* 2003; THOMAS *et al.* 2003). One, referred to as the maize race, is feeding mostly on maize but occasionally on pepper, sorghum, cocklebur, and sunflower (LENIAUD *et al.* 2006). The second, referred to as the mugwort–hop race, feeds on mugwort and hop.

Although in general our results fit quite well with the host-race delineation based on allozyme data, we did find one discrepancy. When populations from all over France were included in the analysis, we failed to detect any differentiation between populations collected on maize and on hop. Such differentiation could be detected only when

our analysis was restricted to populations from northern France. In southern France, mugwort stands are apparently free of ECB (MARTEL *et al.* 2003), whereas maize fields are often substantially infested by this pest (A. WEISSENBARGER, unpublished data), and hop is restricted to wild stands along rivers (T. MALAUSA, D. BOURGUET and S. PONSARD, personal observations). Hence, in this geographical area, adult moths of the mugwort–hop race are probably at low density and strongly outnumbered by adults of the maize race. As the two host races still hybridize in field conditions (MALAUSA *et al.* 2005), populations of the mugwort–hop race may thus receive higher gene flow from populations feeding on maize in southern than in northern France.

Overall, the level of divergence between populations was low, even when comparing populations collected on maize with populations collected from mugwort. These two groups of populations displayed extensive shared polymorphisms (Table 1), and consequently

TABLE 2
Fixation indices between European corn borer populations collected from maize, mugwort, and hop, and between maize, mugwort, and other plants

Comparison		<i>Tpi</i>		<i>Ket</i>		<i>Pbp</i>		<i>Mpi</i>		All genes	
Plant 1	Plant 2	F_{CT}	F_{SC}	F_{CT}	F_{SC}	F_{CT}	F_{SC}	F_{CT}	F_{SC}	F_{CT}	F_{SC}
Overall France											
Maize	Mugwort	0.05**	-0.01	0.03*	0.06**	0.14**	-0.03	0.00	0.12	0.06***	0.05*
	Hop	0.00	0.04	0.01	0.09**	-0.05	0.04	-0.06	0.17*	0.00	0.08**
	Pepper	0.01	-0.01	-0.01	0.19***	-0.06	0.01	-0.07	0.07	0.00	0.05**
	Sorghum	0.01	-0.01	0.01	0.05	0.05	-0.03	-0.09	0.09	0.02	0.03
	Xanthium	-0.02	-0.01	0.05	0.05*	0.04	-0.01	0.18	0.13	0.07	0.05
Mugwort	Hop	0.04	0.11*	-0.02	0.13**	0.12*	-0.08	-0.06	0.16	0.04	0.10**
	Pepper	0.01	0.01	0.05	0.16***	0.15*	-0.14	0.06	-0.13	0.07*	0.04**
	Sorghum	0.00	0.03	0.05	0.06*	0.17***	-0.23	0.03	-0.09	0.06***	0.02
	Xanthium	0.09	0.04	0.04	0.06*	0.31**	-0.23	0.08	0.03	0.13***	0.03
Northern France only											
Maize	Mugwort	0.05	0.01	0.06**	0.03	0.25***	-0.16	-0.10	0.31*	0.09***	0.09
	Hop	0.00	0.10	0.11***	0.08	0.05	-0.14	-0.14	0.42**	0.04***	0.15**
Mugwort	Hop	-0.05	0.13	-0.05	0.14***	0.12*	-0.20	-0.23	0.21	0.03	0.12**

F_{CT} is relative to among-group variance and F_{SC} to within-group variance. Within-population variance is not shown. P -values testing whether indices are significantly >0 were obtained by permutation tests with 10,000 permutations (SCHNEIDER *et al.* 2000). * $P < 0.05$, ** $P < 0.01$, *** $P < 0.001$.

none of the trees based on any of the four markers exhibits any clustering of the two ECB host races (Figure 2). This suggests that reproductive isolation between the maize-Z and mugwort-hop-E races is recent and/or incomplete. More generally, reproductive barriers between diverging or recently diverged species are often semipermeable, which leads to patterns of shared alleles at most loci. Shared polymorphism between closely related species has been observed in other animals. For example, MACHADO *et al.* (2002) observed shared polymorphism at 12 out of 14 nuclear loci between *Drosophila pseudoobscura* and *D. persimilis*. In field crickets, BROUGHTON and HARRISON (2003) did not find any fixed differences at 4 nuclear loci between *Gryllus firmus* and *G. pennsylvanicus*, two species that are estimated to have diverged $\sim 10^3$ - 10^4 generations ago. In *R. pomonella*, a model species thought to have experienced a recent host shift, the sequences of three nuclear loci did not cluster according to the host plant from which the individuals were collected (FEDER *et al.* 2003). Similarly,

recently diverged cichlid fish species also exhibit a pattern of shared polymorphisms (HEY *et al.* 2004). Our results thus tend to confirm that the two French *Ostrinia* taxa constitute two recently diverged host races or sibling species, rather than two distant species in which divergence would be strong at most loci, including nuclear genes (BLUM *et al.* 2003; BARR and MCPHERON 2006; see review in HARE 2001).

However, our estimates of divergence time between taxa suggest that the abundant shared polymorphism between both *Ostrinia* host races is likely due to a high level of gene flow having occurred since the divergence of taxa, rather than to a short time since divergence. Using mutation rates usually observed at nuclear genes (FUTUYMA 1998; RIDLEY 2004) in IM simulations, we estimated the divergence time to be between $\sim 75,000$ and 150,000 years and $2Nm$ to have values ~ 10 . Although these estimates are rough, they seem compatible either with an allopatric divergence followed by secondary contact between both taxa or with a sympatric

TABLE 3
Tests for neutral evolution of molecular polymorphism

	<i>Tpi</i>				<i>Ket</i>				<i>Pbp</i>				<i>Mpi</i>			
	N	D	D^*	F_s	N	D	D^*	F_s	N	D	D^*	F_s	N	D	D^*	F_s
All populations	222	-1.64*	-3.86**	-12.66**	219	-1.03	-1.25	-7.04	79	-0.19	-0.38	-0.01	83	1.23*	0.23	1.52*
Maize	86	-1.24	-0.74	-2.90	81	-0.72	-0.37	-3.93	29	-0.12	0.04	1.67	35	1.03	0.62	0.76
Pepper	29	-1.03	-0.41	-0.40	29	0.19	-0.73	-0.35	8	-0.46	-0.25	3.10	10	0.83	0.57	1.64
Mugwort	35	-1.88**	-2.45*	-1.90	44	-1.11	-0.10	-4.07	14	0.14	-0.25	3.67	12	1.07	1.06	5.29*
Hop	28	-0.72	-0.27	0.20	26	0.21	1.47*	0.94	11	-0.15	-0.02	1.46	11	0.28	0.05	-0.28

N , number of sequences; D , Tajima's D ; D^* , Fu and Li's D^* ; F_s , Fu's F_s . * $P < 0.05$ ($P < 0.02$ for Fu's F_s); ** $P < 0.01$.

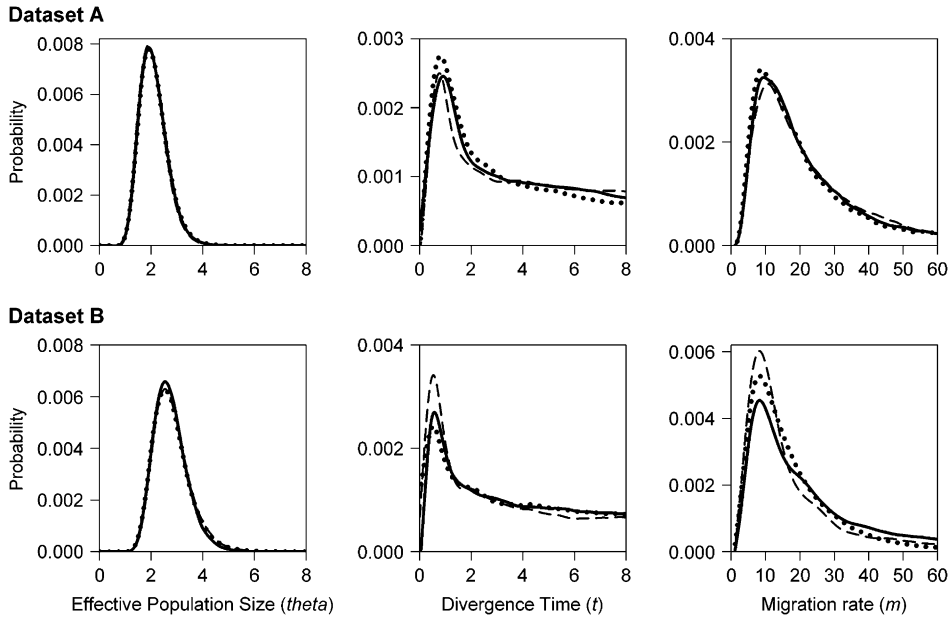


FIGURE 3.—Posterior distribution produced by the IM program for the effective population sizes θ of maize and mugwort populations, their divergence times t , and the migration rates m for each of the three runs using data set A and data set B (containing a common set of *Ket* sequences and two different parts of *Pbp* sequences; see supplemental material for details at <http://www.genetics.org/supplemental/>).

divergence, but they do not support a scenario of recent divergence triggered by a shift onto maize ~500 years ago. Indeed, one would have to assume a mutation rate $>5 \times 10^{-5}$ substitutions per gene per year to obtain divergence time estimates whose confidence intervals include 500 years. Although little is known about rates of evolution at the two loci *Ket* and *Pbp*, such a value seems unrealistically high. An earlier divergence may thus have occurred due to previous host shift (*e.g.*,

onto *Sorghum* sp. or *Panicum* sp.). This pattern of slow divergence under a regime of high gene flow strengthens the conclusions of EMELIANOV *et al.* (2004), who concluded from several studies on the larch budmoth *Z. diniana* that a long and relatively stable sympatric phase of genetic divergence in the presence of gene flow is often a feature of speciation.

Finally, we did not find evidence that any of the genes studied marked a genome region involved in

TABLE 4
Results of isolation model simulations

Data set	Replicate no.	Effective population size (θ)		Divergence time (t)		Migration rate (m)	
		HiPt	HPD90%	HiPt	HPD90%	HiPt	HPD90%
A	1	1.835	1.235–2.925	0.955	0.195–8.795	9.863	2.888–45.038
	2	1.875	1.205–2.885	0.835	0.265–9.995 ^a	10.613	3.038–48.188
	3	1.785	1.225–2.935	0.765	0.155–9.085	9.338	2.663–47.963
Parameters converted into demographical units (individuals for θ or years for t)							
	1	161,450	108,660–257,352	168,049	34,314–1,547,629	2Nm =	9.05
	2	164,969	106,020–253,833	146,933	46,632–1,758,790 ^a		9.95
	3	157,051	107,780–258,232	134,615	27,275–1,598,659		8.33
B	1	2.495	1.705–3.745	0.455	0.205–9.965	7.750	2.250–56.450
	2	2.585	1.705–3.895	0.545	0.025–8.945	7.750	1.750–47.250
	3	2.535	1.655–3.845	0.495	0.025–9.355	7.650	2.050–38.050
Parameters converted into demographical units (individuals for θ or years for t)							
	1	198,107	135,380–274,859	72,256	32,555–1,582,472	2Nm =	9.67
	2	205,253	135,380–309,269	86,548	3,970–1,420,493		10.02
	3	201,283	131,410–305,299	78,607	3,970–1,485,602		9.70

Model parameters estimates (θ , t , m) for data set A and B. Parameters were converted into demographic units using the following formulas: θ (individuals) = $\theta/(4 \times \mu)$; T (years) = t/μ . The migration parameter $2Nm$ was calculated by multiplying $(0.5 \times \theta)$ and m . HiPt, bin with the maximum residence time; HPD90%, 90% highest posterior density interval.

^a Due to the long-tailed posterior distribution, the HPD90% upper bound was higher than the upper bound of the prior distribution and could not be estimated.

reproductive isolation between the two *Ostrinia* host races. This result contrasts with the findings of DOPMAN *et al.* (2005) who documented a clear clustering of the North American E pheromonal race at *Tpi* and a departure of its polymorphism from neutral evolution restricted to this same E-race. In France, differentiation at *Tpi* is significant but low (Table 4) and there is only little evidence that its departure from a neutral molecular evolution was restricted to populations of the E-race (Table 3). Further, this departure can be explained, as in the United States, by one or several selective sweep(s), but it remains unclear whether these sweeps were caused by the selective constraints detected on the coding region of *Tpi* or by selection acting on a gene located near *Tpi*. This latter hypothesis was supported by DOPMAN *et al.* (2005) because *Pdd*, a locus involved in the determinism of the postdiapause-development duration (a trait probably under strong selection in most populations), maps at the same chromosomal location as *Tpi*. However, neither *Tpi* nor *Pdd* seems to contribute to reproductive isolation between French host races: divergence at *Tpi* was not greater than that observed at other loci and was negligible when compared to that estimated between *Ostrinia* American pheromonal races (DOPMAN *et al.* 2005) or between host races of *Z. diniana* in genomic regions thought to be involved in reproductive isolation (EMELIANOV *et al.* 2004).

Conclusion: Although French host races of the ECB have been found to be reproductively isolated (MALAUSA *et al.* 2005), analysis of molecular polymorphisms at four nuclear loci did not reveal a clear pattern of monophyly for each race. Instead, extensive haplotype sharing was observed for all genes, even at the *Tpi* locus in which a clear separation of E and Z populations was previously observed in North America (DOPMAN *et al.* 2005). However, the results of this study confirmed the significant genetic differentiation observed in previous work on the French ECB populations using allozyme markers (BOURGUET *et al.* 2000; MARTEL *et al.* 2003; THOMAS *et al.* 2003; MALAUSA *et al.* 2005; LENIAUD *et al.* 2006). IM coalescent-based simulations allowed estimation of the time since divergence of the two host races between 75,000 and 150,000 years. Although additional data from mitochondrial or microsatellite loci will be needed to confirm this first estimation, this suggests that a scenario of speciation by host shift onto maize, after the introduction of this plant into Europe ~500 years ago, is not likely. Finally, although neutrality tests revealed several departures from the mutation-drift equilibrium for the molecular polymorphism of two genes, we did not identify any gene involved in the reproductive isolation between the two French host races.

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