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## A complex invasion story underlies the fast spread of the invasive box tree moth (*Cydalima perspectalis*) across Europe

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1 A complex invasion story underlies the fast spread of the invasive box tree moth (*Cydalima*  
2 *perspectalis*) across Europe

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16

## 17 **Abstract**

18 Many recently established non-native insect species appear to be spreading across Europe significantly  
19 faster than before. The box tree moth (*Cydalima perspectalis*), a native to Asia, is illustrative of this  
20 trend. First recorded in 2007 in Germany, the moth has then colonized in less than 10 years more than  
21 30 countries in Europe and Asia Minor, causing significant damage to wild and ornamental *Buxus* trees.  
22 It has been hypothesized that the trade of ornamental box trees between China and Europe was  
23 responsible for the moth introduction while plant trade among European countries may have caused  
24 its rapid spread. To clarify the pest invasion history, we analyzed the genetic diversity and structure of  
25 its populations in the native and invaded ranges, using a 1495-bp fragment of the mitochondrial  
26 cytochrome oxidase I and II genes. Moth genetic diversity in Asia compared to the one observed in the  
27 invaded Europe and Asia Minor suggested that the invasive populations probably originated from  
28 eastern China. Furthermore, the high genetic diversity coupled with the spatial genetic structure in the  
29 invaded range suggested the occurrence of several introduction events, probably directly from China.  
30 Moreover, the spatial genetic structure in Europe and Asia Minor may also reflect secondary invasions  
31 within invaded range because of ornamental plant trade among European countries.

32 **Keywords:** *Cydalima perspectalis*, *Buxus*, invasion, insect, ornamental plant trade, multiple  
33 introductions

34

35 Key Message:

36 The invasive moth, *Cydalima perspectalis*, has spread rapidly across Europe and Asia Minor, causing  
37 significant damage to both wild and ornamental *Buxus* trees.

38 Genetic analyses suggested China, and mainly eastern China, as the source of the populations invasive  
39 in Europe but also that multiple introduction events likely occurred.

40 The fast spread of the moth across Europe may result from a combination between these multiple  
41 introductions and human-mediated long-distance transportations of infested *Buxus* trees with  
42 ornamental plant trade among European countries.

43

44 Author Contribution Statement

45 ABr, JR, AR, MAAR conceived the research. ABr, DA, MK, HM, GV, ABe, AR performed sampling. ABr,  
46 DA, CC performed experiments. ABr, DA analyzed data. ABr, DA, GV, JR, AR, MAAR wrote the paper.  
47 All authors approved the paper.

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66

67

## 68 Introduction

69 Biological invasions are continuing to occur worldwide despite an increasing awareness of how  
70 globalization is resulting in human-mediated introductions of species with major ecological, economic,  
71 and sociological effects (Roques 2010a; Simberloff et al. 2013; Meurisse et al. 2018). Moreover, the  
72 arrival of new non-native species is not showing any signs of plateauing (Seebens et al. 2017). In  
73 Europe, most of the insect species that have arrived during the recent decades are first-time invaders,  
74 which have never been categorized as invasive elsewhere (Seebens et al. 2018). Most of newly insect  
75 invaders are originating from Asia (Roques 2010a), with an ever-increasing proportion of  
76 phytophagous species (Roques et al. 2016). The ornamental plant trade is considered as the major  
77 pathway by which these phytophagous insects have been accidentally introduced into Europe (Kenis  
78 et al. 2007; Roques 2010a; Eschen et al. 2017), in parallel with a significant increase in the importation  
79 of live plants to the continent since 1995 (Van Kleunen et al. 2018). Quite simultaneously, faster rates  
80 of spread following establishment were observed in many of these recently-introduced insects  
81 (Roques et al. 2016). Unregulated trade, especially of ornamental plants, as well as the progressive  
82 liberalization of trade and travel during the 1990s may have facilitated such rapid expansions of non-  
83 native species across Europe (Roques et al. 2016).

84 The box tree moth, *Cydalima perspectalis* (Walker, 1859) (Lepidoptera: Crambidae), a native to  
85 Asia, is representative of the non-native species having spread rapidly across Europe. Its native range  
86 includes China, Korea, and Japan (Maruyama and Shinkaji 1987; Xiao et al. 2011; Kim and Park 2013),  
87 where the insect is known to develop on several *Buxus* species (Buxaceae) (Wan et al. 2014). In Europe,  
88 the moth larvae only feed on leaves and shoots of *Buxus* species (Leuthardt and Baur 2013; Matošević  
89 et al. 2017), eventually causing plant death (Kenis et al. 2013; Wan et al. 2014). *C. perspectalis* was first  
90 observed on ornamental box trees in urban areas but it has now spread to natural forests in some  
91 countries, causing severe defoliation in native box stands, such as for *B. sempervirens* and *B. colchica*  
92 (Kenis et al. 2013; John and Schumacher 2013; Gninenko et al. 2014; Mitchell et al. 2018). *C.*  
93 *perspectalis* was first recorded in Europe in early 2007, at two different sites in Germany (Krüger 2008).  
94 Later that same year, it was observed in Switzerland and the Netherlands (Leuthardt et al. 2010; Van  
95 der Straten and Muus 2010) (Figure 1). Then, over a period of less than 10 years, the insect spread  
96 across the whole of Europe and into Asia Minor, and it is at present observed in more than 30 countries,  
97 ranging from the United Kingdom (Salisbury et al. 2012) to Iran (Mitchell et al. 2018).

98 It has been hypothesized that the moth was accidentally introduced primarily via the trade of  
99 ornamental box trees between China and Europe (Leuthardt et al. 2010; Casteels et al. 2011; Nacambo  
100 et al. 2014), and that subsequent trade among European countries led to its fast spread (EPPO 2012;  
101 Kenis et al. 2013; Matošević 2013). Indeed, China shipped large quantities of *Buxus* trees to several  
102 European countries between 2006 and 2010 (EPPO 2012). As a result, the moth could have been  
103 introduced several times in different countries after its initial appearance in 2007 in Germany.  
104 However, few interception records are available from the European plant health and quarantine  
105 services (EPPO 2012) because the moth was included on the EPPO alert list only from 2007 to 2011  
106 (Strachinis et al. 2015). During this period, *C. perspectalis* was intercepted once in 2008 in the  
107 Netherlands (EPPO 2012), a country that has been the largest importer of ornamental plants to Europe

108 over the recent years (Eschen et al. 2017). It is also noticeable that in the Netherlands, Belgium, and  
109 England, the moth was first recorded from nurseries (Van der Straten and Muus 2010; Casteels et al.  
110 2011; Salisbury et al. 2012) whilst the first mention in Russia was on box trees imported for the Winter  
111 Olympics in Sochi (Gninenko et al. 2014). Actually, box trees, especially *Buxus sempervirens*, are very  
112 popular ornamental plants (Matošević 2013; Mitchell et al. 2018) and drive significant commercial  
113 trade within Europe and adjacent countries (EPPO 2012; Dehnen-Schmutz et al. 2010).

114 Human-mediated introductions often involve complex invasion pathways (Garnas et al. 2016;  
115 Meurisse et al. 2018), for which historical information may be missing or misleading. Hence, molecular  
116 data can often supplement what is known about a species' invasion history, helping to clarify the likely  
117 invasive pathways by revealing the presence of e.g. genetic bottlenecks, multiple introduction events,  
118 or admixture (Estoup and Guillemaud 2010; Lawson Handley et al. 2011; Estoup et al., 2016; Fraimout  
119 et al. 2017). The sequencing of mitochondrial DNA (mtDNA) is an efficient first step for disentangling  
120 the pathways followed by non-native species. It can be used to identify source populations, founding  
121 events, and the occurrence of multiple introductions (Muirhead et al. 2008; Estoup and Guillemaud  
122 2010; Cristescu 2015). Thus, this approach has been employed in a number of non-native species (e.g.,  
123 Auger-Rozenberg et al. 2012; Garipey et al. 2014; Javal et al. 2017; Lesieur et al. 2018). Identifying  
124 source populations and clarifying invasive pathways help to complement management strategies and  
125 increases our understanding of how non-native species spread (Muirhead et al. 2008; Lawson Handley  
126 et al. 2011).

127 The objectives of this study were to clarify the geographical origin(s) and colonization history of  
128 the populations having invaded Europe. To this end, we compared the genetic diversity and structure  
129 of the populations of the box tree moth in the native Asian range and in most of the invaded countries  
130 of Europe and Asia Minor, using mtDNA cytochrome oxidase I and II (COI - COII) genes.

131

## 132 Materials and methods

### 133 *Sampling*

134 Moth populations were sampled throughout their native and invaded ranges (Table 1;  
135 Supplementary Materials Table S1, Figures S1 and S2).

136 In the native Asian range, we attempted to collect specimens from 2012 to 2017 in both natural  
137 stands and ornamental plantations of *Buxus* trees growing in the putative distribution range of the  
138 moth (Figure 2a). Pheromone traps were used for adult moths and hand sampling for larvae and pupae  
139 (Suppl. Mat. Table S1). However, surveys and trappings were unsuccessful in Japan even though *C.*  
140 *perspectalis* has been recorded there (Maruyama and Shinkaji 1987; Kawazu et al. 2007). In South  
141 Korea, we sampled one population from an urban area in Seoul (Table 1; N20). In China, 19 populations  
142 were obtained, covering most of the putative range from north-eastern China (Liaoning province),  
143 northern China (Nei Mongol and Beijing provinces), and eastern China (Shandong, Anhui, Jiangsu,  
144 Shanghai, Zhejiang, and Fujian provinces), to south-central China (Henan province) and south-western

145 China (Yunnan and Guizhou provinces). Only the southern provinces of Guangdong and Guangxi could  
146 not be sampled. However, all but one of these Chinese populations were collected in urban areas  
147 despite our efforts in natural stands of *Buxus* where moth density appeared to be very low. Only in  
148 Fuyang, Zhejiang province (Table 1; N17), larvae could be collected outside urban areas, in a sentinel  
149 plant nursery established at the boundary between a natural forest and agricultural lands (Kenis et al.  
150 2018).

151 In the invaded range, samples could be collected in 23 different countries (Table 1) from  
152 regions that were colonized by the moth between 2007 and 2016 (Figures 1 and S2). Ideally, we  
153 intended to collect larvae via hand sampling. When that was not possible, adults were captured using  
154 pheromone traps. Moths were mostly collected in urban areas except in the following sites: Bzyb Valley  
155 (Table 1; I27) and Mtirala Park in Georgia (Table 1; I28); all sites in Russia (Table 1; I58-I60); Si Sangan  
156 National Park in Iran (Table 1; I41); and Roquefort-sur-Garonne (Table 1; I24) and Marcillac-Vallon in  
157 France (Table 1; I23).

158 Upon collection, all specimens were placed in 96% alcohol and stored at -21°C to preserve their  
159 DNA until the analyses could take place.

160

#### 161 *DNA extraction, amplification, and sequencing*

162 The larvae and adults were dissected. DNA was extracted from the thoracic muscles of the  
163 larvae or the legs of the adults using the DNeasy® Blood and Tissue Kit (Qiagen, Hilden, Germany). We  
164 amplified a section of the mitochondrial genome that included part of the COI-COII genes. We  
165 employed a pair of primers developed for a related species, *Diaphania* (= *Glyphodes*) *pyloalis* (Zhu et  
166 al. 2013). The primers were renamed LeCyd-F2 (5' TGGAGCAGGAACAGGATGAAC 3') and Cynna-R2 (5'  
167 GAGACCANTACTTGCTTTCAG 3'). Amplification was carried out in a total PCR volume of 25 µL, which  
168 contained 1 µL of DNA, 15.8 µL of ultrapure water, 2.5 µL of 10X DreamTaq Green Buffer, 2.5 µL of  
169 dNTP (10mM), 0.5 µL of MgCl<sub>2</sub> (2.5mM), 1 µL of each primer (10µM), 0.5 µL of betaine solution (5 M),  
170 and 0.2 µL of DreamTaq DNA polymerase (5 units/µL). Thermocycling was performed using a Veriti®  
171 96-well Fast Thermal Cycler (Applied Biosystems, Foster City, CA, USA) and the following procedure:  
172 an initial 5-min denaturation step took place at 95°C and was followed by 25 amplification cycles (94°C  
173 for 35 s, 60°C for 45 s, and 72°C for 3 min). PCR products were analyzed by gel electrophoresis in a  
174 1.5% agarose gel to check for successful amplification. Those of approximately 2000 bp in length were  
175 purified using the NucleoFast® 96 PCR Clean-up Kit (Macherey-Nagel, Düren, Germany). A fragment of  
176 around 1500 bp of purified DNA that included COI, ARNt L2, and COII was then sequenced using Cynna-  
177 R2 and the internal primer Jerry-F 5' CAACATTTATTTGATTTTTGG 3' because the PCR products were  
178 too long to be sequenced directly. Sequencing was carried out using the Big Dye Terminator Cycle  
179 Sequencing Kit (v. 3.0, Applied Biosystems, Foster City, CA, USA) and an ABI Prism 3500 Genetic  
180 Analyzer. The two sequenced strands were then aligned and the absence of double peaks on  
181 electropherograms was manually verified using CodonCode Aligner v. 3.7.1 (CodonCode Corporation,  
182 Centerville, MA, USA).

183



184 *Genetic Analyses*

185 We successfully sequenced mtDNA from 132 and 305 individuals from the native and invaded  
186 ranges, respectively. Sequence data from Matošević et al. (2017) was also included in the dataset for  
187 the invaded range. Sequences were aligned using Clustal W (Thompson et al. 1994), which was  
188 implemented in BioEdit v. 7.1. The final alignment was obtained without any insertions or deletions.  
189 All sequences were truncated at the same length (1495 bp). The presence of stop codons were checked  
190 using MEGA v. 6 (Kumar et al. 2008). We compared our sequences with sequences in GenBank and  
191 BOLD using BLAST to confirm that individuals had been properly identified as *C. perspectalis* and to  
192 check for possible contamination.

193 A phylogenetic analysis to investigate the relationships among mtDNA haplotypes was  
194 performed by maximum-likelihood (ML) method computed using the software MEGA v. 6. Evaluation  
195 of statistical confidence in nodes was based on 10,000 bootstrap replicates. The distance between DNA  
196 sequences were calculated based on Kimura's two-parameter method (Kimura 1980). Two related  
197 species, *D. pyralis* (Genbank Accession No. KM576860) and *Glyphodes quadrimaculalis* (Genbank  
198 Accession No. KF234079), were used as outgroup taxa. A statistical parsimony network with a 95%  
199 confidence level was constructed using TCS v. 1.21 (Clement et al. 2000). We determined haplotype  
200 number (H), haplotype diversity ( $h$ ), the average number of nucleotide differences ( $k$ ), and nucleotide  
201 diversity ( $n$ ) using DNAsp v. 5 (Librado and Rozas 2009). Haplotype distribution and frequency were  
202 projected onto maps of Asia, Europe, and Asia Minor using ArcGis v. 10.6 (ESRI, Redlands, CA, USA). To  
203 characterize the moth's native range, spatial genetic structure was first assessed by testing if  $G_{ST}$  (the  
204 coefficient of genetic variation over all the populations) was significantly smaller than  $N_{ST}$  (the  
205 coefficient taking into account similarities among haplotypes)—10,000 permutations were  
206 implemented in Permut (Pons and Petit 1996).

207 Analysis of molecular variance (AMOVA) was performed to look for evidence of genetic  
208 structure in the moth's native and invaded ranges using Arlequin v. 3.5 (Excoffier and Lischer 2010).  
209 To carry out the analysis, we grouped the populations according to different criteria. First, we took  
210 into account the populations' geographical locations. Second, we took into account information  
211 related to the ornamental plant trade, such as production areas and volume of imported and/or  
212 exported plants. The objective was to estimate the impacts on genetic structure in both the native and  
213 invaded ranges simultaneously. In the native range, sequences were grouped into five clusters based  
214 on population geographical location and known *Buxus* tree production in Chinese provinces (René  
215 Eschen pers. comm.) (Figures 2a and S1): (1) north-eastern province of Liaoning grouped with South  
216 Korea (N8, N20); (2) northern province of Nei Mongol and Beijing area (N2, N9); (3) eastern provinces  
217 of Fujian, Henan, Shandong, and Shanghai area (N3, N4, N6, N10, N11, N12, N13, N14); (4) south-  
218 eastern provinces of Anhui, Jiangsu and Zhejiang (N1, N7, N17, N18, N19); and (5) south-western  
219 provinces of Guizhou and Yunnan (N5, N15, N16). In the invaded range, information on ornamentals  
220 was country-specific (Dehnen-Schmutz et al. 2010; EPPO, 2012; Eschen et al. 2017), and therefore  
221 sequence data were first grouped per country. Then, countries were grouped into four clusters based  
222 on geographical location and the commercial value of ornamental plant imports/exports (Dehnen-  
223 Schmutz et al. 2010; Eschen et al. 2017). The groups were as follows: (1) Germany, Netherlands,

224 France, Italy, Belgium; (2) Switzerland, Spain, Portugal, Luxembourg; (3) Austria, Hungary, Czech  
225 Republic, Slovenia, Slovakia, Croatia, Greece, Serbia; and (4) Romania, Turkey, Russia, Bulgaria,  
226 Georgia, Iran.

227

## 228 Results

229 Twelve haplotypes were identified based on the sequences of the 437 specimens collected  
230 across the species' native and invaded ranges (HTA1-HTC1, Table 1; GenBank accession numbers:  
231 **MK611945-MK611956**). These haplotypes comprised 21 single nucleotide polymorphisms. There was  
232 no evidence of contamination nor of nuclear copies of mitochondrial DNA (numts). The BLAST search  
233 confirmed specimen identification, based on the few samples of *C. perspectalis* present in GenBank  
234 and BOLD.

235 The haplotypes formed three haplogroups (A, B, and C), which were separated by at least seven  
236 mutation steps (Figure 2b). The sixteen intermediate haplotypes were not present in our sample pool.  
237 The topology of the phylogenetic tree was similar to the haplotype network (Suppl. Mat. Figures S3).  
238 Two haplotypes, HTA1 and HTB1, were prevalent—they were displayed by approximately 36% of the  
239 individuals sequenced. Two other haplotypes, HTA2 and HTB2, had prevalence of 13.5% and 8.9%,  
240 respectively. These four haplotypes were found in both the native and invaded range. Overall,  
241 haplotype diversity ( $h$ ) was 0.709 ( $\pm 0.012$ ), the average number of nucleotide differences ( $k$ ) was 4.10,  
242 and nucleotide diversity ( $n$ ) was 0.00274 ( $\pm 0.0000$ ).

243

### 244 *Genetic diversity and structure in the native range*

245 Twelve haplotypes were observed among the sequences of the 132 specimens collected in  
246 China (19 sampling locations) and South Korea (1 sampling location) (Table 1, Figures 2a and 2b). The  
247 average pairwise sequence difference between haplotypes was 0.004, and ranged from 0.1% to 0.7%,  
248 which is consistent with intraspecific distances. The three haplogroups were represented. Haplogroups  
249 A and B were observed across the range of Chinese populations, but haplogroup C was only found in  
250 the Fuyang population (N17, Zhejiang province). Haplogroups A and B comprised 5 and 6 haplotypes,  
251 respectively, while haplogroup C was made up of only one. In most populations (65%), haplotypes  
252 belonging to haplogroups A and B co-occurred (e.g., Beijing, N2; Ordos, N9; Tai'an, N11). The values of  
253 the diversity indices are provided in Table 1.

254 Only haplotype HTB1 was common to both South Korea and China (Figure 2a), where it was  
255 shared by 32.6% of the individuals sequenced. HTB3 was only observed in South Korea (N20: 46%).  
256 HTA1 and HTB2 were prevalent across China (30.3% and 22.0%, respectively). HTA1, HTB1, and HTB2  
257 occurred in 14 populations. With the exception of HTA2 and HTA4, which were each found in two  
258 populations in different provinces (HTA2: N9, Nei Mongol province, and N11, Shandong province;  
259 HTA4: N12, Shandong province, and N14, Shanghai province), all the other haplotypes were observed  
260 at single locations in China. Nine populations contained three haplotypes, whereas three populations  
261 contained a single haplotype. Among these three populations, only in the Wendeng population (N13,



262 Shandong province) was more than one individual sampled. In native populations, haplotype number  
263 (H) ranged from 1 to 4, haplotype diversity ( $h$ ) ranged from 0 to 1, the average number of nucleotide  
264 differences ( $k$ ) ranged from 0.40 to 5.98, and nucleotide diversity ( $n$ ) ranged from 0 to 0.004.

265 Pronounced genetic structure was observed:  $N_{ST}$  (0.277) was significantly higher than  $G_{ST}$   
266 (0.181;  $p$  value < 0.01). The AMOVA results also supported the existence of genetic structure as all the  
267 fixation indices were significant ( $p$  value < 0.05), including  $F_{CT}$  (Table 2). Genetic differences among  
268 groups and among populations within groups nevertheless accounted for a small percentage of the  
269 genetic variance (8.4% and 9.4% respectively). The largest amount of genetic differentiation (82.2%)  
270 was found within populations (highly significant fixation index;  $p$  value < 0.001).

271

### 272 *Genetic diversity and structure in the invaded range*

273 Only five haplotypes (HTA1, HTB1, HTA2, HTA4, HTB2) emerged from the sequences of the 305  
274 specimens obtained from the 72 locations sampled in Europe and Asia Minor. All of these haplotypes  
275 were also present in China (Table 1). The values of the diversity indices in the invaded range were lower  
276 than those in the native range ( $h=0.668$ ,  $k=3.79$ , and  $n=0.0027$ ).

277 HTA1 and HTB1 were observed across the invasive populations (Figure 3a and 3b). HTA1 was  
278 found in 47 populations and 39.3% of the individuals sequenced, while HTB1 was found in 45  
279 populations and 38.0% of the individuals sequenced. HTA2 was seen in 29 populations. HTA4 and HTB2  
280 had restricted distributions—they occurred in populations associated with the first records of *C.*  
281 *perspectalis* in Europe. HTA4 was only found in the Weil-am-Rhein population (I34) in Germany and in  
282 the Saint Louis population (I18) in France, which is located near the border with Germany. HTB2 was  
283 found mostly in populations in Kehl (I33) and Strasbourg (I19), two nearby locations. Twenty-eight  
284 populations contained just one haplotype, whereas 34 populations contained two haplotypes, and 9  
285 populations contained 3 haplotypes. In invasive populations, haplotype number (H) ranged from 1 to  
286 4, haplotype diversity ( $h$ ) ranged from 0 to 1, the average number of nucleotide differences ( $k$ ) ranged  
287 from 0.4 to 8, and nucleotide diversity ( $n$ ) ranged from 0 to 0.00535. We did not observe any genetic  
288 diversity in Switzerland, Austria, and Greece. Germany and France had the highest genetic diversity:  
289 four and five haplotypes, respectively.

290 The AMOVA results revealed the presence of genetic structure within the invasive populations  
291 (Table 2). The values of all the fixation indices were highly significant ( $p$  value < 0.01). The largest  
292 amount of genetic differentiation (76.09%) was found within populations. Genetic differences among  
293 populations within groups and among groups accounted for a smaller percentage of genetic variation  
294 (12.30% and 11.61%, respectively).

295

## 296 Discussion

297 Based on mtDNA sequence diversity, three major findings resulted from this study. First, the  
298 box tree moth displayed a complex genetic structure with a mix of deeply differentiated haplogroups  
299 in its native Asian range, probably as a result of anthropogenic activities within this region. Second, the  
300 comparison of the genetic diversity patterns between Asia and the invaded Europe indicated eastern

301 China as the likely source of the European populations. Finally, historical records coupled to the genetic  
302 diversity and structure observed in the invaded range suggested that multiple introduction events may  
303 have occurred.

#### 304 *Genetic structure of the box tree moth in its native range*

305 *Cydalima perspectalis* displayed a weak but significant spatial genetic structure across its  
306 native range. From our analyses, three main distribution regions can be delimited based on: (1) the  
307 occurrence of a private haplotype (HTB3) in South Korea, (2) the co-occurrence of haplogroups A and  
308 B in northern and eastern China; and (3) the high prevalence of haplogroup B in southern China. Similar  
309 phylogeographic pattern was observed in some lepidopterans native to Asia. The Asiatic rice borer,  
310 *Chilo suppressalis*, thus showed three genetically diverse and geographically localized clades in China  
311 corresponding to north-eastern, central and southern China (Meng et al. 2008). These same major  
312 lineages were also defined for a swallowtail, *Papilio bianor* (Zhu et al. 2011). Actually, these regions  
313 have been pointed out as Glacial refugia for the last two species (Meng et al. 2008; Zhu et al. 2011).  
314 Since *Buxus* fossils were found in southern China (Ma et al. 2015; Huang et al. 2018), this region may  
315 also have served as refugia for some populations of *C. perspectalis*.

316 Only specimens collected from urban areas could be analyzed because we were unable to  
317 obtain samples from natural stands of *Buxus* despite intensive efforts. Indeed, the moth's putative  
318 distribution in Asia appeared to be solely based on records from urban areas (Kawazu et al. 2007; Kim  
319 and Park 2013; Nacambo et al. 2014), which makes it difficult to define its natural distribution. A large  
320 part of the natural stands of *Buxus* species are found in southern China (Figure 2a) (Fang et al. 2011).  
321 Thus, the proximity to forests may explain the significant genetic diversity that we observed in our  
322 data, even if our samplings in this region were carried out on box trees planted in towns. Similarly, the  
323 proximity of the sentinel plant nursery in Fuyang (south-eastern China) to natural stands may also  
324 account for the unique occurrence of haplogroup C at that site.

325 The mtDNA diversity of *C. perspectalis* populations in Asia corresponds to three distinct  
326 haplogroups, two of which being widely distributed. The spatial co-occurrence of divergent  
327 haplogroups suggests that this moth has a complex history. Indeed, the existence of divergent  
328 haplogroups may reflect ancient phylogenetic differentiation (Avice et al. 1987), whereas their co-  
329 occurrence may have resulted from secondary contact, produced by moth movements and/or by  
330 human-mediated dispersal of infested plants. Haplotype co-occurrence has also been observed in the  
331 native Chinese range of the Asian long-horned beetle, *Anoplophora glabripennis*, leading to suggest  
332 that beetles have been moved by man to northern and eastern China with plants used for reforestation  
333 (Carter et al. 2009; Javal et al. 2017). The oriental fruit moth, *Grapholita molesta*, displayed the same  
334 genetic pattern, which likely reflect recent dispersal through human activities (Song et al. 2018).

335 More generally, human-mediated dispersal can indeed reshape the genetic structure of insect  
336 populations and largely modify their primary natural phylogeographic pattern (Stone et al. 2007; Song  
337 et al. 2018). In recent years, *Buxus* trees have increasingly been planted as ornamentals in China,  
338 especially in the northern part of the country. Because no box trees grow in the wild in northern China,  
339 it can be assumed that *C. perspectalis* has been introduced into the cities of this region (e.g., Beijing)  
340 as a result of ornamental plantations (Nacambo et al. 2014). As our samplings were essentially carried

341 out in urban areas, the observed genetic structure is likely to represent a combination between the  
342 moth phylogeographic history and its human-mediated dispersal with ornamental plant trade.  
343 Additional samplings in natural stands, especially from other provinces of south-central China and  
344 Japan would be required to precise the evolutionary history of *C. perspectalis* in Asia and understand  
345 better its present spatial genetic structure.

#### 346 *A Chinese origin for the populations invasive in Europe and Asia Minor*

347 Five haplotypes were observed in *C. perspectalis* invaded range. It corresponded to a  
348 significant fraction (41.7%) of the genetic variability observed in the native Asian populations. This  
349 pattern was commonly noticed in biological invasions (Dlugosch and Parker 2008; Estoup and  
350 Guillemaud 2010; Lawson Handley et al. 2011; Cristescu 2015). For example, a high genetic diversity in  
351 the invaded range was also observed for the micromoth *Phyllonorycter issikii* (Kirichenko et al. 2017),  
352 and the brown marmorated stink bug, *Halyomorpha halys* (Gariepy et al. 2014, 2015), two invaders  
353 that came to Europe from Asia.

354 For *C. perspectalis*, the South Korean haplotype was not observed in the invasive populations  
355 whereas all the haplotypes found in the invaded range were also observed in China, including two of  
356 the three most common haplotypes. This finding strongly suggested that these invasive populations  
357 have a Chinese origin. It could be coherent with the data on *Buxus* tree imports (EPPO 2012), which  
358 also pointed out China as the most probable source of invasion (Leuthardt et al. 2010; Casteels et al.  
359 2011; Nacambo et al. 2014). Over recent decades, China has effectively emerged as a key exporter of  
360 ornamental plants (Dehnen-Schmutz et al. 2010; Kenis et al. 2018). For example, between 2005 and  
361 2010, the Netherlands obtained more than 80% of its imported ornamental plants (i.e., import volume)  
362 from eastern Asia, and especially from China (van Valkenburg et al., 2014). Moreover, China was the  
363 greatest supplier of *Buxus* trees to other European countries during that same period (EPPO 2012;  
364 Kenis et al. 2018).

365 Both the spatial genetic structure of the moth populations within China, the distribution of the  
366 production areas of ornamental *Buxus* in this country, and the higher economic development of coastal  
367 Chinese provinces (Roques 2010b) suggested eastern China as the location of the invasive source. All  
368 the haplotypes observed in Europe were also found in the Shandong province, and four were observed  
369 around Shanghai. The coastal provinces in eastern China are more economically developed than are  
370 other Chinese provinces, and Shanghai is one of China's key economic centers, from whence large  
371 quantities of products are exported throughout the world (Roques 2010b; Lu et al. 2018). Moreover,  
372 half of the areas in which *Buxus* trees are produced for export are found in eastern China, notably  
373 around Shanghai (René Eschen pers. comm.). These places could potentially be the sources of invasive  
374 *C. perspectalis* populations. However, correctly identifying source populations or invasion pathways is  
375 not always straightforward. The native range of an invader can be too large to be exhaustively  
376 characterized (e.g., Orlova-Bienkowskaja et al. 2015; Orlova-Bienkowskaja and Volkovitsh 2018). The  
377 fine-scale reconstruction of an invasion history is possible when there is significant genetic structure  
378 in the native range (Lombaert et al. 2011; Cristescu 2015). Here, it was challenging to define the source  
379 of the invasive moth populations because there was a significant genetic diversity in the invaded range

380 coupled to a low level of genetic variation among the analyzed Chinese populations, and a lack of  
381 samples from populations from other native areas in southern China and Japan.

382

### 383 *A complex invasion process*

384 The populations sampled where *C. perspectalis* was first recorded in 2007 in Europe (Krüger  
385 2008; Van der Straten and Muus 2010; I34 and I33 in Germany and I52 and I53 in the Netherlands;)   
386 differed genetically. German populations presented two haplotypes, which were missing in the  
387 Netherlands, whereas the Dutch populations showed one haplotype not found in Germany. Moreover,  
388 three of these four populations shared only one haplotype. This pattern could result from (i) a single  
389 founder event with a significant genetic diversity introduced before 2007 in one of these localities,  
390 followed by a secondary spread. In the invasion processes, it is common knowledge that insects can  
391 be present before their first observations (Allendorf and Lundquist 2003; Crooks 2005). Furthermore,  
392 this genetic diversity could reflect the introduction of insects belonging to populations from eastern  
393 China, in which we observed often signs of admixture. Such a pattern was already observed for  
394 *Ambrosia artemisiifolia*, the annual weed which recently invaded Europe (van Boheemen et al. 2017).

395 The genetic pattern of *C. perspectalis* can also be explained by (ii) multiple introductions  
396 arriving directly from China to several places at the same period. *Buxus* trees were commercially  
397 imported from China by different countries of the EU, at least the Netherlands and Italy, between 2006  
398 and 2010 (EPPO 2012). No information is available for Germany, but Weil am Rhein where the insect  
399 was first observed, regularly received large shipments of Chinese imports (Casteels et al. 2011). This  
400 trade information coupled with our genetic data may suggest more than one introduction directly from  
401 China. These events of multiple introductions are a common feature in biological invasions (Estoup  
402 and Guillemaud 2010; Lawson Handley et al. 2011; Cristescu 2015). It has been observed for many non-  
403 native insects across the world (e.g., *Diabrotica virgifera virgifera*, Ciosi et al. 2008; *Hyalopterus pruni*,  
404 Lozier et al. 2009; *Cactoblastis cactorum*, Marsico et al. 2011; *Leptoglossus occidentalis*, Lesieur et al.  
405 2018), and recently, for non-native species originating from China (*Harmonia axyridis*, Lombaert et al.  
406 2010; *H. halys*, Garipey et al. 2015; *A. glabripennis*, Javal et al. 2017; *Drosophila sukuzii*, Fraimout et al.  
407 2017).

408 In the same way, multiple introductions may increase the probability of spread of non-native  
409 species inside the invaded range (Cristescu 2015). Indeed, anthropogenic activity is known to promote  
410 subsequent introduction events, leading to dispersal and range expansion (Estoup and Guillemaud  
411 2010). It is assumed that *C. perspectalis* has a natural dispersal rate of around 10 km per year (Van der  
412 Straten and Muus 2010; Casteels et al. 2011), which is far too limited to explain its fast colonization of  
413 Europe and Asia Minor. The spatial genetic structure suggested that different groups exist. This pattern  
414 likely resulted from several complex introduction events, some possibly directly from China and/or  
415 some within the invaded range. The speed of the invasion can be explained by those different  
416 assumptions, even if the important ornamental plant trade inside Europe (Dehnen-Schmutz et al. 2010;  
417 Eschen et al. 2015) may have clearly played a role in the moth dispersal. For example, in 2008, the  
418 moth was intercepted in the Netherlands in a *Buxus* shipment of unknown provenance for exportation

419 inside Europe (EPPO 2012). Besides, Gninenko et al. (2014) supposed that the insect was introduced  
420 from Italy in 2012 in Caucasian forests due to the import of infested box trees for the Winter Olympics  
421 in Sochi. If considering Hungary, Turkey and Romania, where the pest was recorded in 2011 (Sáfián  
422 and Horváth 2011; Hizal et al. 2012; Gutue et al. 2014), the geographical distances separating the  
423 localities are greater than 400 km (Figure 1), which suggests simultaneous introductions but from  
424 unknown origins.

425 Our study provides a new example of a pest species with a complex invasion history. We have  
426 highlighted the role played by the ornamental plant trade in its introduction and dispersal, even if more  
427 detailed data on *Buxus* tree imports to Europe are needed to decipher the number of introduction  
428 events. However, to better pinpoint the moth's region of origin in China and to disentangle the effects  
429 of anthropogenic activity on its current distribution patterns, we need to carry out broader sampling  
430 efforts and acquire more information about the moth's natural distribution in its native range  
431 (Muirhead et al. 2008). Additionally, more powerful tools such as microsatellite or SNP markers (Estoup  
432 and Guillemaud 2010; Cristescu 2015; Estoup et al. 2016) could help flesh out *C. perspectalis* invasion  
433 scenarios and clarify how the species spread across Europe and Asia Minor in less than ten years.

434

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439 Ethical approval: All applicable international, national, and/or institutional guidelines for the care and  
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441

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663

664 **Legends**

665

666 Table 1. Genetic diversity statistics of native and invasive populations of *Cydalima perspectalis* based  
667 on the current study and Matošević et al. (2017).

668

N°	Country	Populations	N	HTA1	HTA2	HTA3	HTA4	HTA5	HTB1	HTB2	HTB3	HTB4	HTB5	HTB6	HTC1	H	<i>h</i> (±SD)	<i>k</i>	<i>n</i> (±SD)
<b>Native Range</b>			132	40	3	1	2	1	43	29	6	1	1	1	4	12	0.755 (0.018)	4.25	0.0028 (0.0001)
N1	China	Huaibei	5						4	1						2	0.400 (0.237)	0.40	0.0003 (0.0002)
N2		Beijing	7	5					1	1						3	0.524 (0.209)	3.62	0.0024 (0.0008)
N3		Fuzhou	1	1												1	-	-	-
N4		Youxi	5	1					3	1						3	0.700 (0.218)	3.20	0.0021 (0.0011)
N5		Guiyang	8	1					4	3						3	0.679 (0.122)	2.29	0.0015 (0.0008)
N6		Xinyang	1						1							1	-	-	-
N7		Nanjing	5	3					1	1						3	0.700 (0.218)	4.60	0.0031 (0.0009)
N8		Shenyang	7	2					4	1						3	0.667 (0.160)	3.62	0.0024 (0.0008)
N9		Ordos	5	2	2									1		3	0.800 (0.164)	3.80	0.0025 (0.0013)
N10		Jinan	5	2		1			2							3	0.800 (0.164)	4.60	0.0031 (0.0008)
N11		Tai'an	8	2	1				3	2						4	0.821 (0.101)	4.43	0.0030 (0.0006)
N12		Dongying	3	1			1		1							3	1 (0.272)	5.33	0.0036 (0.0015)
N13		Wendeng	2							2						1	0	-	0
N14		Shanghai	14	5			1	1		7						4	0.659 (0.090)	4.59	0.0031 (0.0003)
N15		Kunming	8						4	2		1	1			4	0.750 (0.139)	0.93	0.0006 (0.0002)
N16		Lijiang	9						4	5						2	0.556 (0.090)	0.56	0.0004 (0.0001)
N17		Fuyang	12	4					3	1				4		4	0.773 (0.069)	5.98	0.0040 (0.0003)
N18		Hangzhou	8	6					1	1						3	0.464 (0.200)	3.25	0.0022 (0.0009)
N19		Lishui	6	5						1						2	0.333 (0.215)	2.67	0.0018 (0.0012)
N20	South Korea	Seoul	13						7		6					2	0.538 (0.060)	1.08	0.0007 (0.0001)
<b>Invaded range</b>			305	120	56		3		116	10						5	0.668 (0.012)	3.79	0.0027 (0.0001)
I1	Austria	Vienna	6						6							1	0	-	0
I2		Rankweil	2						2							1	0	-	0
I3	Belgium	Ghent	5	5												1	0	-	0
I4		Mechlin	1						1							1	-	-	-
I5		Vremde	3	1					2							2	0.667 (0.314)	4.67	0.0031 (0.0015)
I6	Bulgaria	Plovdiv	4	1	3											2	0.500 (0.265)	0.50	0.0003 (0.0002)
I7		Sofia	5	1					4							2	0.400 (0.237)	2.80	0.0019 (0.0011)
I8	Croatia	Vinica	6		1				5							2	0.333 (0.215)	2.67	0.0018 (0.0012)
I9		Osor	1						1							1	-	-	-
I10		Artatore	1	1												1	-	-	-
I11		Zagreb	5	3					2							2	0.600 (0.175)	4.20	0.0028 (0.0008)
I12		Višnjevac	5						5							1	0	-	0



N°	Country	Populations	N	HTA1	HTA2	HTA3	HTA4	HTA5	HTB1	HTB2	HTB3	HTB4	HTB5	HTB6	HTC1	H	<i>h</i> (±SD)	<i>k</i>	<i>n</i> (±SD)
I13	Czech Rep.	Brno	5	2	2				1							3	0.800 (0.164)	3.40	0.0023 (0.0011)
I14	France	Grenoble	3	3												1	0	-	0
I15		Orléans	5	2					3							2	0.600 (0.175)	4.20	0.0028 (0.0008)
I16		Tours	7	4	1				2							3	0.667 (0.160)	3.62	0.0024 (0.0008)
I17		Bastia	4	2	2											2	0.667 (0.204)	0.67	0.0005 (0.0001)
I18		Saint Louis	9	6			1			2						3	0.556 (0.165)	3.33	0.0022 (0.0009)
I19		Strasbourg	4							4						1	0	-	0
I20		Paris	5	5												1	0	-	0
I21		Bordeaux	4		2				2							2	0.667 (0.204)	5.33	0.0036 (0.0011)
I22		La Rochelle	5	1	4											2	0.400 (0.237)	0.40	0.0003 (0.0002)
I23		Marcillac Vallon	2		1				1							2	1 (0.500)	8.00	0.0054 (0.0027)
I24		Roquefort-sur-Garonne	4	2	1				1							3	0.833 (0.222)	4.00	0.0027 (0.0012)
I25		Nantes	5	3	1				1							3	0.700 (0.218)	3.20	0.0021 (0.0011)
I26		Lagnes	11	7	2				1	1						4	0.600 (0.154)	2.80	0.0019 (0.0008)
I27	Georgia	Bzyb Valley	5	3	1				1							3	0.700 (0.218)	3.20	0.0021 (0.0011)
I28		Mtirala National Park	5	2	3											2	0.600 (0.175)	0.60	0.0004 (0.0001)
I29		Zugdidi	4		3				1							2	0.500 (0.265)	4.00	0.0027 (0.0014)
I30		Tbilisi	5		2				3							2	0.600 (0.175)	4.80	0.0032 (0.0009)
I31	Germany	Lorsch	5	3					2							2	0.600 (0.175)	4.20	0.0028 (0.0008)
I32		Kandern	4	4												1	0	-	0
I33		Kehl	3							3						1	0	-	0
I34		Weil Am Rhein	4	2			2									2	0.667 (0.204)	0.67	0.0005 (0.0001)
I35	Greece	Filiria	2						2							1	0	-	0
I36		Katerini	7						7							1	0	-	0
I37	Hungary	Harkány	5						5							1	0	-	0
I38		Hódmezővásárhely	5	3					2							2	0.600 (0.175)	4.20	0.0028 (0.0008)
I39		Budaörs	5	3					2							2	0.600 (0.175)	4.20	0.0028 (0.0008)
I40		Kőszeg	5	1					4							2	0.400 (0.237)	2.80	0.0019 (0.0011)
I41	Iran	Si Sangan National Park	3	1					2							2	0.667 (0.314)	4.67	0.0031 (0.0015)
I42	Italy	Bologna	3		3											1	0	-	0
I43		Ruta	3	1	1				1							3	1 (0.272)	5.33	0.0036 (0.0015)
I44		Legnaro	5						5							1	0	-	0
I45		Tregnago	4						4							1	0	-	0
I46		Cesa	2	1	1											2	1 (0.500)	1.00	0.0007 (0.0003)

N°	Country	Populations	N	HTA1	HTA2	HTA3	HTA4	HTA5	HTB1	HTB2	HTB3	HTB4	HTB5	HTB6	HTC1	H	<i>h</i> (±SD)	<i>k</i>	<i>n</i> (±SD)
I47		Florence	4		3				1							2	0.500 (0.265)	4.00	0.0027 (0.0014)
I48		Lucca	4	1	2				1							3	0.833 (0.222)	4.17	0.0028 (0.0013)
I49		Turin	4	2	2											2	0.667 (0.204)	0.67	0.0005 (0.0001)
I50		Viterbo	4	2	1				1							3	0.833 (0.222)	4.00	0.0027 (0.0012)
I51	Luxembourg	Luxembourg	2	1					1							2	1 (0.5)	7.00	0.0047 (0.0023)
I52	Netherlands	Giessen	9	1					8							2	0.222 (0.166)	1.56	0.0010 (0.0008)
I53		Boskoop	3	3												1	0	-	0
I54		Rotterdam	5	3					2							2	0.600 (0.175)	4.20	0.0028 (0.0008)
I55		Rhederbrug	3						3							1	0	-	0
I56	Portugal	Vila Nova de Cerveira	5	2	3											2	0.600 (0.175)	0.60	0.0004 (0.0001)
I57	Romania	Timișoara	3		2				1							2	0.667 (0.314)	5.33	0.0036 (0.0017)
I58	Russia	Solokhaul	4	3	1											2	0.500 (0.265)	0.50	0.0003 (0.0002)
I59		Komsomolsk	2	2												1	0	-	0
I60		Krasnodar	5	4	1											2	0.400 (0.237)	0.40	0.0003 (0.0002)
I61	Serbia	Belgrade	5	2					3							2	0.400 (0.237)	4.20	0.0028 (0.0008)
I62	Slovakia	Zvolen	5		5											1	0	-	0
I63		Bratislava	4						4							1	0	-	0
I64	Slovenia	Dobrovnic	2						2							1	0	-	0
I65		Nova Gorica	6	2					4							2	0.533 (0.172)	3.73	0.0025 (0.0008)
I66		Sečovlje	2						2							1	0	-	0
I67	Spain	Besalú	4	3	1											2	0.500 (0.265)	0.50	0.0003 (0.0002)
I68	Switzerland	Delémont	6	6												1	0	-	0
I69		Liestal	3	3												1	0	-	0
I70		Monteggio	3	3												1	0	-	0
I71	Turkey	Istanbul	5	3					2							2	0.600 (0.175)	4.20	0.0028 (0.0008)
I72		Yalova	2	1	1											2	1 (0.5)	1.00	0.0007 (0.0003)

669

670 The information provided for each location is population number (N°); number of individuals (N); haplotype number (H); haplotype diversity (*h*, with  
671 standard deviation SD); average number of nucleotide differences (*k*); and nucleotide diversity (*n*, with standard deviation SD).

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674 Table 2. Results of the hierarchical AMOVA of COI-COII sequence data obtained from *Cydalima*  
 675 *perspectalis* in its (a) native range and (b) invaded range.

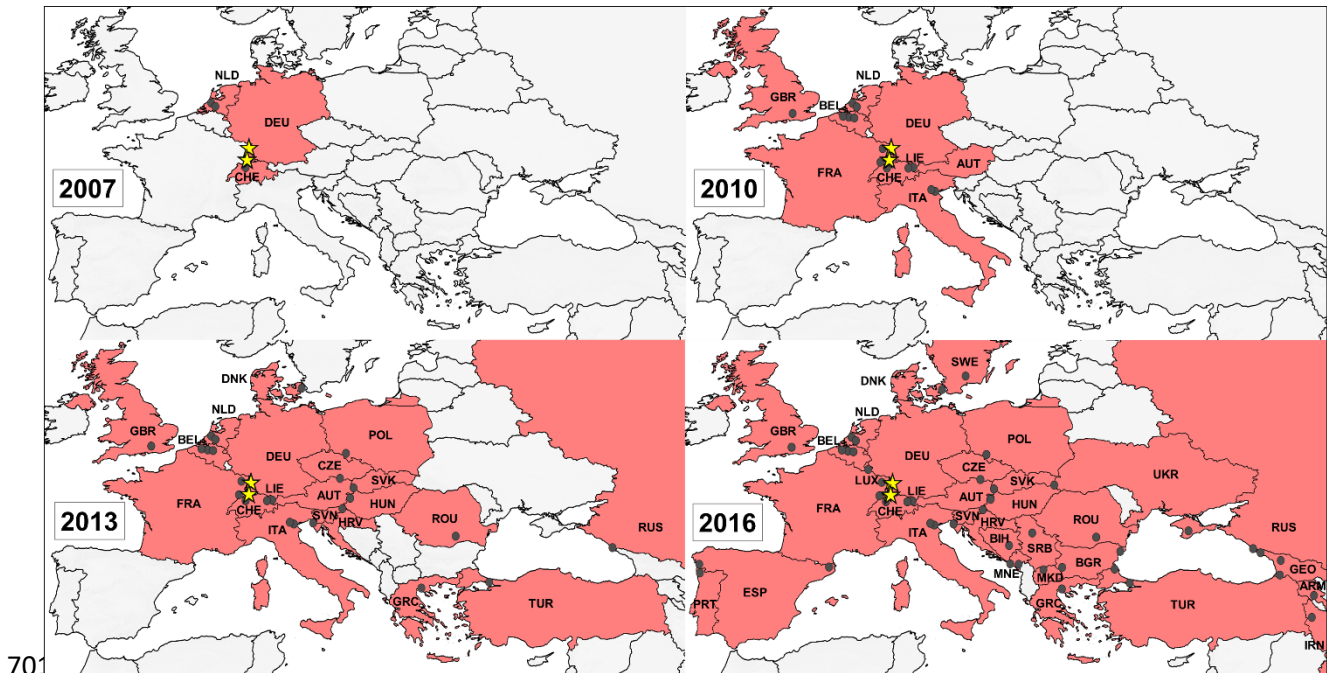
Source of variation	df	Sum of squares	% of variation	Fixation index
(a) Native range				
Among groups	4	6.067	9.37	$F_{CT} = 0.093^*$
Among populations within groups	15	7.725	8.43	$F_{SC} = 0.094^*$
Within populations	112	35.693	82.20	$F_{ST} = 0.178^{***}$
(b) Invaded range				
Among groups	3	11.091	11.61	$F_{CT} = 0.116^{**}$
Among populations within groups	20	15.291	12.30	$F_{SC} = 0.139^{***}$
Within populations	281	75.132	76.09	$F_{ST} = 0.239^{***}$

676 Statistical probabilities were derived from 50,175 permutations; \* $p$  value < 0.05, \*\* $p$  value < 0.01, \*\*\* $p$   
 677 value < 0.001. (a) In the native range, five groups were defined: (1) N8, N20; (2) N2, N9; (3) N3, N4, N6,  
 678 N10, N11, N12, N13, N14; (4) N1, N7, N17, N18, N19; and (5) N5, N15, N16 (see Table 1 for population  
 679 numbers). (b) In the invaded range, four groups were defined: (1) Germany, Netherlands, France, Italy,  
 680 Belgium; (2) Switzerland, Spain, Portugal, Luxembourg; (3) Austria, Hungary, Czech Republic, Slovenia,  
 681 Slovakia, Croatia, Greece, Serbia; and (4) Romania, Turkey, Russia, Bulgaria, Georgia, Iran.

682

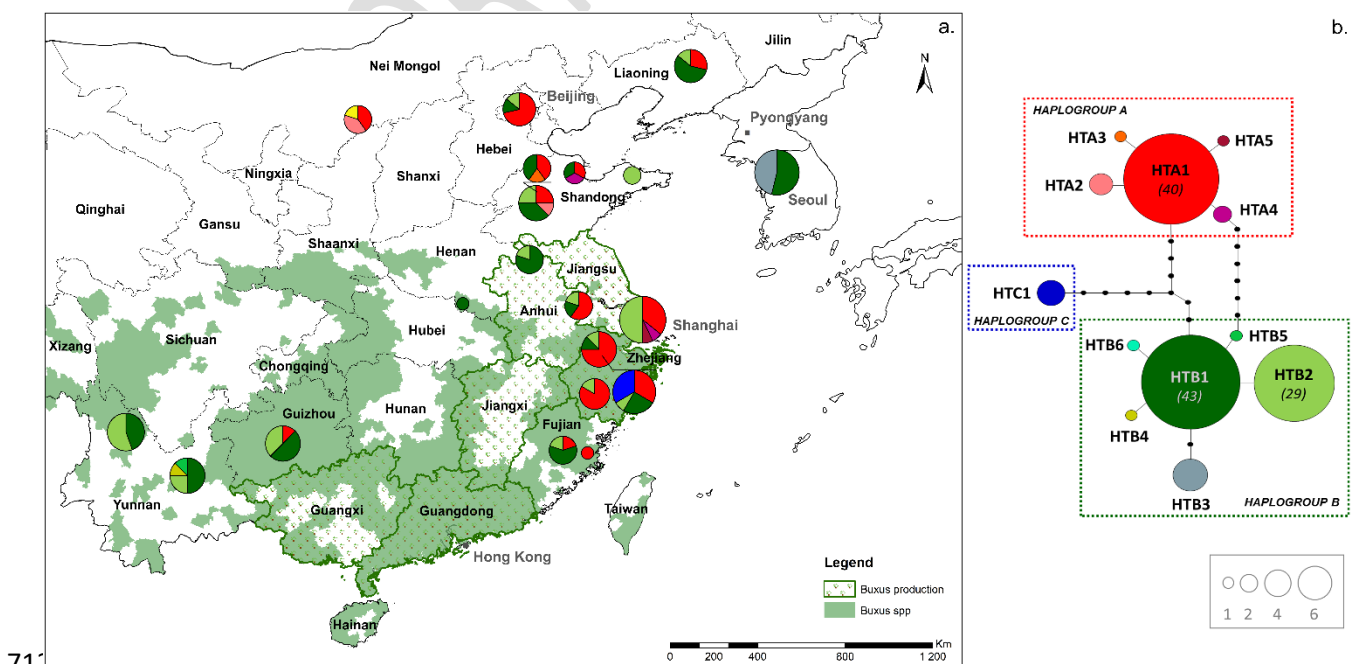
683 Figure 1. Spread of *Cydalima perspectalis* across Europe and Asia Minor between 2007 (the year the  
 684 species was first observed) and 2016. The two yellow stars indicate the first places that *C. perspectalis*  
 685 was detected—in Germany (DEU) in 2007 (Krüger 2008). The gray dots represent the first  
 686 observation(s) of the moth in each country, which were determined based on a literature review.  
 687 Country abbreviations are as follows (listed by year of first moth observation): CHE: Switzerland  
 688 (Leuthardt et al. 2010); NLD: Netherlands (Van der Straten and Muus 2010); FRA: France (Feldtrauer  
 689 et al. 2009); GBR: United Kingdom (Salisbury et al. 2012); AUT: Austria; LIE: Liechtenstein; DNK:  
 690 Denmark; ITA: Italy (Bella 2013); BEL: Belgium (Casteels et al. 2011); ROU: Romania (Gutue et al. 2014);  
 691 TUR: Turkey (Hizal et al. 2012); HUN: Hungary (Sáfián and Horváth 2011); CZE: Czech Republic (Bella  
 692 2013); SVN: Slovenia (Seljak 2012); HRV: Croatia (Koren and Crne 2012); POL: Poland (Blaik et al. 2016);  
 693 RUS: Russia (Gninenko et al. 2014); SVK: Slovakia (Bella 2013); GRC: Greece (Strachinis et al. 2015);  
 694 ESP: Spain (Pérez-Otero et al. 2015); BGR: Bulgaria (Beshkov et al. 2015); SRB: Serbia (Vajgand 2016);  
 695 MNE: Montenegro (Hrnčić et al. 2017); BIH: Bosnia and Herzegovina (Ostojić et al. 2015); MKD:  
 696 Macedonia (Načeski et al. 2018); GEO: Georgia and Abkhazia (Matsiakh et al. 2018); UKR: Ukraine  
 697 (Nagy et al. 2017); LUX: Luxembourg (Ries et al. 2017); PRT: Portugal (Maria da Conceição de Lemos  
 698 Viana Boavida pers. comm.); ARM: Armenia (Shiroma Sathyapala pers. comm.); IRN: Iran (Mitchell et  
 699 al. 2018); and SWE: Sweden (Bengtsson 2017).

700



702 Figure 2a. Spatial distribution of *Cydalima perspectalis* COI-COII haplotypes in the moth's native range.  
 703 The color codes indicate the color used in the haplotype network (see Figure 2b). The putative natural  
 704 range of *C. perspectalis* was characterized based on records in the literature and moths collected as  
 705 part of this study. The distribution of *Buxus* species was estimated based on Fang et al. (2011), and  
 706 *Buxus* production in Chinese provinces was estimated using unpublished data provided by René  
 707 Eschen.

708 2b. Network of *Cydalima perspectalis* COI-COII haplotypes based on the haplotype frequencies  
 709 observed in the moth's native range. Each circle represents a haplotype (HTA1 to HTC1) and is labeled  
 710 using a specific color. Circle size is proportional to the number of individuals. Each line between circles  
 711 corresponds to a mutational step, and the small black circles are missing intermediate haplotypes.

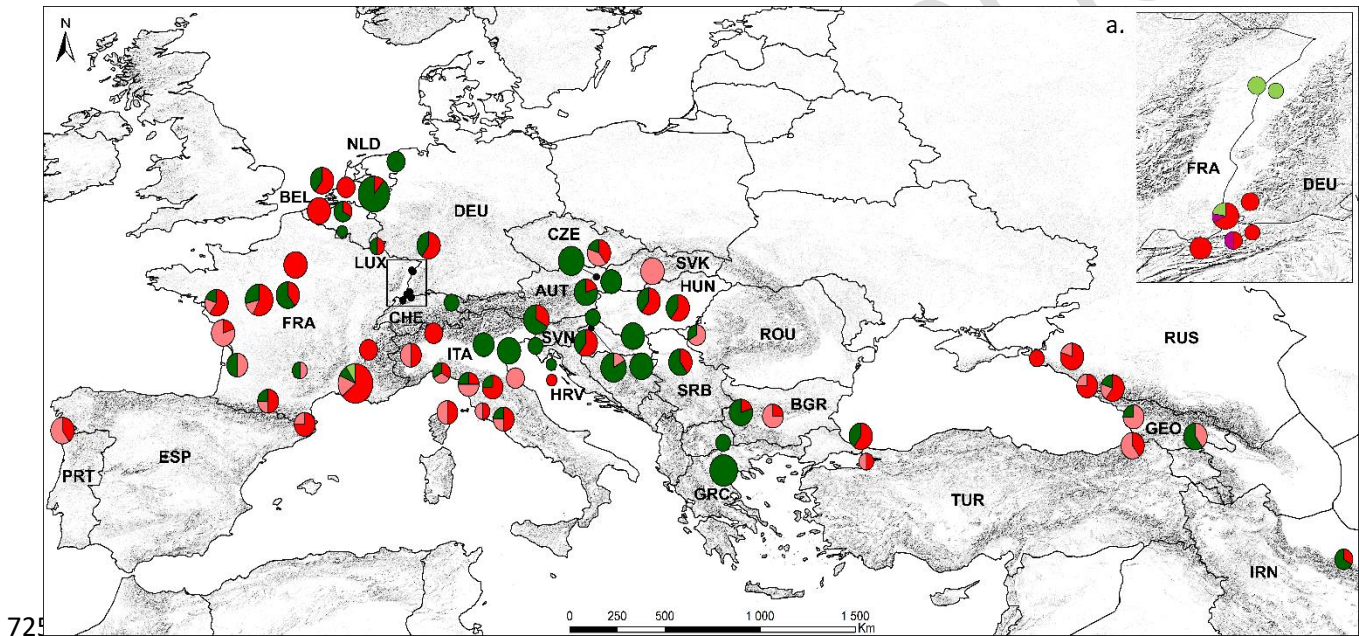


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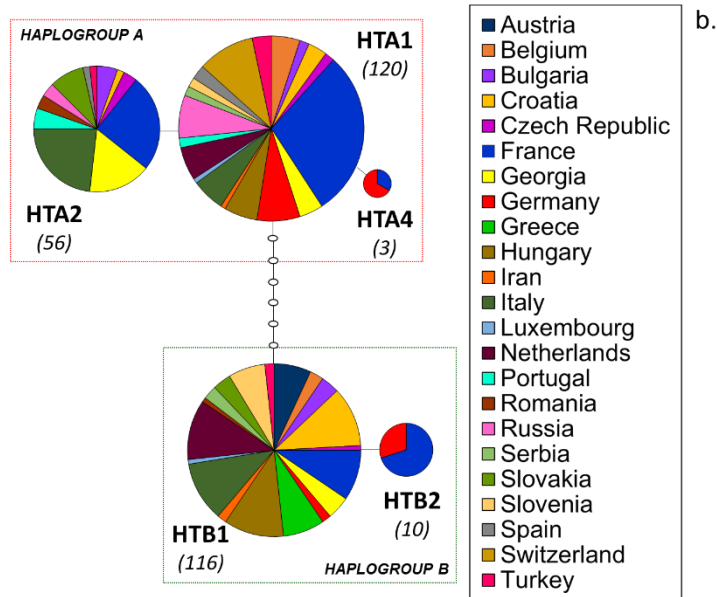
23

713 Figure 3a. Geographical distribution of *Cydalima perspectalis* COI-COII haplotypes in the moth's  
 714 invaded range based on the results of the current study and Matošević et al. (2017). Circle size is  
 715 proportional to the number of individuals. The color codes indicate the color used in the haplotype  
 716 network (see Figure 2b). Country abbreviations are as follows: DEU: Germany; CHE: Switzerland; NLD:  
 717 Netherlands; FRA: France; AUT: Austria; ITA: Italy; BEL: Belgium; ROU: Romania; TUR: Turkey; HUN:  
 718 Hungary; CZE: Czech Republic; SVN: Slovenia; HRV: Croatia; RUS: Russia; SVK: Slovakia; GRC: Greece;  
 719 ESP: Spain; BGR: Bulgaria; SRB: Serbia; GEO: Georgia; LUX: Luxembourg; PRT: Portugal; and IRN: Iran.

720 3b. Network of *Cydalima perspectalis* COI-COII haplotypes based on the haplotype frequencies  
 721 observed in the moth's invaded range. Each circle represents a haplotype. Different colors represent  
 722 sampled invaded countries by *Cydalima perspectalis*. Circle size is proportional to the number of  
 723 individuals. Each line between circles corresponds to a mutational step, and the small empty circles  
 724 are missing intermediate haplotypes.







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