Phylogeography of the soil-borne vector nematode *Xiphinema index* highly suggests Eastern origin and dissemination with domesticated grapevine

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The soil-borne nematode *Xiphinema index* is closely linked to its main host, the grapevine, and presents a major threat to vineyards worldwide due to its ability to transmit *Grapevine fanleaf virus* (GFLV). The phylogeography of *X. index* has been studied using mitochondrial and microsatellite markers in samples from most regions of its worldwide distribution to reveal its genetic diversity. We first used the mitochondrial marker *CytB* and illustrated the low intraspecific divergence of this mainly meiotic parthenogenetic species. To generate a higher polymorphism level, we then concatenated the sequences of *CytB* and three mitochondrial markers, *ATP6*, *CO1* and *ND4*, to obtain a 3044-bp fragment. We differentiated two clades, which each contained two well-supported subclades. Samples from the eastern Mediterranean and the Near and Middle East were grouped into three of these subclades, whereas the samples from the western Mediterranean, Europe and the Americas all belonged to the fourth subclade. The highest polymorphism level was found in the samples of one of the Middle and Near East subclades, strongly suggesting that this region contained the native area of the nematode. An east-to-west nematode dissemination hypothesis appeared to match the routes of the domesticated grapevine during Antiquity, presumably mainly dispersed by the Greeks and the Romans. Surprisingly, the samples of the western subclade comprised only two highly similar mitochondrial haplotypes. The first haplotype, from southern Iberian Peninsula, Bordeaux and Provence vineyards, exhibited a high microsatellite polymorphism level that suggests introductions dating from Antiquity. The second haplotype contained a highly predominant microsatellite genotype widespread in distant western countries that may be a consequence of the massive grapevine replanting following the 19th-century phylloxera crisis. Finally, our study enabled us to draw a first scaffold of *X. index* diversity at the global scale.

To date, more than 4,100 species of plant-parasitic nematodes have been described1 worldwide. They represent a threat to agriculture estimated at approximately $US 80 billion per year2. Among the species attacking crops, the dagger nematode *Xiphinema index* has a high economic impact in vineyards worldwide. During its feeding on the plant, *X. index* may transmit *Grapevine fanleaf virus* (GFLV)3–5 to grapevines. GFLV is known as the most severe viral disease of grapevines. It is responsible for fanleaf degeneration6, which occurs in temperate regions of vine cultivation7,8. The vector *X. index* is an ectoparasitic nematode present in Mediterranean environments and temperate climates where grapevine grows. It has a limited host range, and domesticated grapevine (*Vitis vinifera* subsp. *vinifera* or *sativa*)9 is by far its major host10,11. Therefore, its detection on other hosts occurs only when

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the location has a previous history of grapevine cultivation. Notably, its presence has not been reported in native forests or climax vegetation. An exception is in the Middle East, where its presence has been reported in Iranian natural woodland (Sturhan in Weischer), in the forests lying along the Caspian Sea where wild grapevines (V. vinifera subsp. sylvestris) may be common.

Because X. index is the vector of GFLV, estimating its variability is of interest to the sustainable management of this viral disease. In grapevine, alternatives to the use of highly toxic chemical nematicides, such as the use of nematode-resistant rootstocks or plants with an antagonistic effect on X. index, are promising leads for nematode and subsequently virus control. The genetic variability of this nematode is expected to depend on its reproductive mode. Females of X. index reproduce by meiotic parthenogenesis, but occasional sexual reproduction may also occur, although males are rare.

The native area of a nematode species is another key point to consider for variability studies, i.e., where it should show its highest genetic diversity. For X. index, this information will help to trace its routes of introduction and will provide a better understanding of its dispersal. It has been hypothesized that GFLV, together with its vector, was introduced to western vineyards from the Middle East with domesticated grapevine. Nevertheless, there is still no conclusive argument available today to support the hypothesis that the nematode originates from the Middle East. The objective of this study was to address the genetic diversity of X. index in relation to its geographic distribution.

We report hereafter a diversity study using mitochondrial (maternal coding DNA) and microsatellite (nuclear noncoding DNA) markers. These markers are complementary and characterize different evolutionary time scales. Mitochondrial markers are robust and commonly used for phylogeographic studies in species belonging to diverse zoota, such as nematodes, insects, fishes or mammals. By contrast, microsatellite markers, the preferred tool for population genetics, are suitable to the level of the individual and have allowed the identification of rare hybridization events in X. index. In the model nematode Pristionchus pacificus, microsatellite markers have shown much higher mutation rates (approx. 10-fold) than mitochondrial markers.

In our study, we first considered the mitochondrial gene coding for Cytochrome B (CytB) in X. index. Because of the low resolution of this single marker, we then described the nematode phylogeographical patterns using CytB concatenated with three other mitochondrial genes, ATP synthase subunit 6 (ATP6), Cytochrome c oxidase subunit 1 (CO1) and NADH dehydrogenase subunit 4 (ND4). We then relied on eight microsatellite markers specific to X. index to refine our results.

Our data set for X. index contributed to (i) establishing the patterns of genetic variability observed in the sampling regions that we have covered, (ii) proposing a putative native area in the eastern Mediterranean and (iii) drawing hypotheses about its dissemination routes and the correlated dispersal itineraries of its host, domesticated grapevine, from very early antiquity. Finally, our work enabled us to draw a first scaffold of X. index diversity at the global scale.

**Material and Methods**

**Origins of nematode samples.** We designated individuals from the same origin as ‘samples’ and not as ‘populations’ because of their different sampling procedures and histories, as reported hereafter. The term ‘sample’ was also used because of the low nematode numbers available in many of them. Eighty-two total samples of X. index, all originating from grapevine plots, were used (Table S1). Within a vineyard plot, a sample generally corresponded to one soil lump obtained from a single or a few clods dug in a viral disease focus showing GFLV aerial symptoms. The total soil weight ranged from 0.25 to 2.0 kg. Out of the 82 samples of X. index, 33 originated from greenhouse rearing of an initial field sample. Rearing was performed in pots on grapevine or fig hosts grown in a collection initiated in 1993 at INRA-ISA, Sophia-Antipolis, France (Table S1). Approximately half of the other X. index samples originated from soils received by express mail and directly used on their date of reception for nematode extraction and storage at −80 °C. The remaining half of the samples consisted of individuals extracted from soil by their provider, fixed in 70% ethanol solution and sent to INRA-ISA for direct processing or storage at −80 °C. In our study, the samples of X. index covered a geographic area ranging from the Middle and Near East and the eastern Mediterranean to Europe and North and South America.

**Selection of samples and individuals for genotyping.** Three individuals per sample were initially used for genotyping. For an overall evaluation of the diversity of X. index, our study was first conducted using the CytB marker alone from all samples (Table S1). Then, the study focused on a subset of samples that contained each of the CytB haplotypes obtained and covered the worldwide distribution area of the nematode. In particular, all the samples from the eastern Mediterranean and Near and Middle Eastern locations were included together with samples from distant locations, such as Chile, California and Hungary, and representative samples from Italy, France, Spain and Portugal. The three individuals of these samples were genotyped for the ND4, CO1 and ATP6 markers, and the new haplotypes were also retained. When several haplotypes were obtained within the three individuals of the same sample, three additional individuals, when available, were genotyped to confirm previous results and/or detect putative new variants that were also included. This procedure allowed us to compose a final subset of 35 samples grouping 43 total individuals (last column of Table S1) that we considered representative of the mitochondrial diversity. The individuals of the subset were then genotyped for their multilocus genotypes (MLGs) using eight microsatellite loci. As for mitochondrial haplotypes, when several MLGs were obtained within the three individuals of a sample, three additional individuals, when available, were genotyped to confirm the results and/or detect putative new variants.

**Preparation of DNA templates from single individuals.** DNA from a single nematode individual was isolated by a simplified procedure: an adult or juvenile was hand-picked and placed in a 0.5 ml PCR tube containing 50 μl lysis buffer (KCl 50 mM, Gelatin 0.05%, Tris pH 8.2 10 mM, Tween 20 0.45%, Proteinase K 60 μg/
ml and MgCl₂ 2.5 mM). Then, Eppendorf tubes were alternatively moved from liquid nitrogen to a 55 °C water bath 10 times to facilitate breakdown of the nematode body. This step was followed by an incubation at 60 °C for 90 min (tubes were vortexed at least once during incubation to help break up the tissues) and by heating at 95 °C for 10 min to inhibit the reaction of Proteinase K. Finally, individuals were cooled at 4 °C, vortexed briefly (2–3 sec) and centrifuged shortly at 6,000 rpm for 30 sec. DNAs were stored at −20 °C until use for PCR or further experiments.

**Primer design for CytB, ATP6, ND4 and CO1 mitochondrial genes.** We performed BLAST alignments between each of the CytB, ND4, CO1 and ATP6 sequences of *X. americanum* obtained by He et al.34 (accession number NC_005928) and the EST database from *X. index* on NemaBLAST (http://nematode.net/NN3_front-page.cgi). We found putative partial sequences of the four homologous mitochondrial genes in *X. index* (contigs XI01293 for CytB, XI01255 for ATP6, XI01185 for ND4 and XI01295 for CO1). For each gene, an alignment was performed between the *X. americanum* sequence and each of these partial sequences using Clustal W in Mega software version 6.0.35. Then, specific primers were designed from these alignments for each gene and used to amplify specific fragments from *X. index* (CytB-852 bp, ND4-644 bp, CO1-998 bp and ATP6-550 bp) (Table S2).

**PCR amplification, purification and sequencing of mitochondrial genes.** The primers designed enabled the amplification of partial sequences of all four mitochondrial genes from *X. index*. All amplifications were carried out in a 50 μL reaction mixture containing 3 μL of DNA template, 5 μL of reaction buffer, 2 μL of each primer at 10 μM and 0.4 units of Taq polymerase (AmpliTaq, Applied Biosystems/Perkin Elmer). Amplifications were performed on a Hybaid thermocycler with the following steps: (i) 95 °C for 3 min; (ii) 35 cycles of 30 sec at 94 °C, 1 min at 59 °C (52 °C for ATP6) and 1 min 30 sec at 72 °C; and (iii) 72 °C for 5 min. The presence of the expected PCR fragments was checked by running the PCR products on 1% agarose gel in 0.5X TAE. DNA sequences were then obtained from the purified PCR products. Occasionally, samples gave nonspecific bands (blurred bands) for the markers CytB and CO1. In such cases, 40 μL of PCR product was loaded onto a 1.5% agarose gel in TAE. The band was recovered from the gel with a sterile scalpel, purified using the MinElute Gel Extraction Kit protocol (Qiagen) and sequenced. DNA sequences obtained from purified PCR products or purified bands were aligned visually. Because all the sequences are protein coding and have no introns or gaps, alignment was straightforward.

**Microsatellite genotyping.** We used a set of eight out of the nine primer pairs (Table S3) designed by Villate et al.36 for which the forward primers carried a fluorescent tag. One primer pair (Xi08) was omitted because its poor polymorphism within the samples did not yield significant additional information.39 PCR was carried out in 10 μL simple reactions containing 2 μL of DNA extract, 2 μL of 5 X Taq reaction buffer, 0.15 units of Taq polymerase, 5 U/μL Taq DNA polymerase (AmpliTaq, Applied Biosystems/Perkin Elmer), 0.3 μL of each primer (forward primers were fluorolabeled with a FAM, PET, NED or VIC dye at the 5′ end) at 10 μM. For each reaction, we mixed four pairs of primers in one tube called mix 1 (microsatellites Xi29, Xi04, Xi16 and Xi13) or mix 2 (microsatellites Xi24, Xi22, Xi32 and Xi27) (Table S3). Reactions were carried out in a PTC-100 thermocycler (MJ Research) with the following amplification conditions: 95 °C for 3 min; 30 cycles of 30 sec at 94 °C, 1 min 30 sec at 57 °C and 1 min 30 sec at 72 °C; and 72 °C for 5 min. The presence of the expected PCR fragments was checked by running the PCR products on 1% agarose gel in 0.5X TAE. The band was recovered from the gel with a sterile scalpel, purified using the MinElute Gel Extraction Kit protocol (Qiagen) and sequenced. DNA sequences obtained from purified PCR products or purified bands were aligned visually. Because all the sequences are protein coding and have no introns or gaps, alignment was straightforward.

**Data analysis.** For the mitochondrial genes, we computed the mean distances between sequences with Mega software version 6.035 using the Kimura-2-parameters model and the maximum likelihood (ML) method to build phylogenetic trees.34,37 We also completed the bootstrap values of the ML method by the Bayesian posterior probability.38 The sequence of *X. americanum* was included as an outgroup in our phylogenetic analysis.34 For the microsatellite data, we first analyzed the results with the GeneMarker program version 1.75 (Applied Biosystems). Then, we used the POPULATION software version 1.2.3.1 (available at http://bioinformatics.org/populations/) to compute the distances between individuals by Cavalli-Sforza and Edwards distance (CSE)39 and draw neighbor-joining (NJ) trees.

**Results**

**Overall diversity in *X. index* using the CytB gene.** Six CytB haplotypes were obtained from our samples overall. We estimated the evolutionary intraspecific divergence between the CytB sequences using representatives of CytB and other haplotypes obtained hereafter from other mitochondrial genes (Table S4). This divergence was low and reached a maximal value of 0.012. All individuals grouped into two well-supported clades (Fig. S1). The first clade contained only seven individuals and was divided into two subclades holding, respectively, four individuals from Israel and Palestine (a single haplotype) and three individuals from Samos and South Italy (two haplotypes). The second clade included all the individuals from western Europe, North and South America (a single haplotype in 69 individuals) and the rest of the eastern samples (two haplotypes in 13 individuals).

**Phylogeographic pattern in *X. index* inferred from single and concatenated sequences of mitochondrial genes.** We generated a phylogenetic tree for each of the mitochondrial gene markers ATP6, CO1 and ND4 using the subset of 35 samples (43 *X. index* individuals) retained to represent the worldwide variability of the nematode. The ML trees (Fig. S2A–C) produced for each gene revealed the same general topology as for CytB (Fig. S1). Several new haplotypes were detected for ATP6 (e.g., in the Tabriz and Sharekord Iranian samples and in the Alasehir Turkish sample) and for CO1 (i.e., a particular haplotype common to South American samples and some samples from the southern Iberian Peninsula and France) (Fig. 1). Among the four genes, ATP6 showed the highest polymorphism (Table S5).
We next generated a consensus ML tree using the 3044-bp fragment obtained from the concatenated sequences of the four markers to further support these topologies (Figs 1 and 2A). The X. index individuals were distributed into two clades, each clade being separated into two well-supported subclades. In clade I, a first subclade designated ‘Western’ (W) gathered all individuals from Western Europe, North and South America (two haplotypes). Meanwhile, a second subclade designated ‘Near and Middle East’ (NME) grouped individuals encompassing, from west to east, Crete, Cyprus, Turkey, Egypt and Iran. In clade II, a first subclade designated ‘Israel and Palestine’ (IP) grouped the three samples from Israel and Cis-Jordan that belonged to a single haplotype, whereas a second subclade designated ‘Samos and South Italy’ (SSI) grouped three samples showing two haplotypes.

The Western subclade was divided into two groups, designated Western 1 (W1) and Western 2 (W2), due to a single substitution (CO1-438) (Figs 1 and 2A). W1 comprised 14 samples with the same haplotype ranging from the island of Crete and Pecs in Hungary in its eastern geographic expansion to Northern California in its western expansion. By contrast, W2 comprised 12 samples located in three distant regions: the southern Iberian Peninsula (Spain and Portugal), Bordeaux (France) and Provence (France). The nine samples from NME showed five total haplotypes. NME revealed two specific haplotypes in the sample from Tabriz (Iran) and a single specific haplotype in each of the samples from Sharekord (Iran) and Alasehir (Turkey) (Figs 1 and 2A).

Subclades NME, IP and SSI, from the Central and Eastern Mediterranean, totaled eight haplotypes, while the widely spread subclade W totaled only two haplotypes (Figs 1 and 2A). These data strongly suggest that the samples from the Central and Eastern Mediterranean possess a higher diversity than those from western countries.

**Diversity of X. index inferred from microsatellite markers.** Several limiting factors, such as the low number of samples, the low number of individuals available per sample and the highly heterogeneous sampling procedures, prevented us from performing reliable formal population genetics statistics, such as comparison of allelic and genotypic richness or mean heterozygosity. Nevertheless, we were able to refine the mitochondrial classification using the MLGs from the subset of 35 samples. While the concatenated mitochondrial sequences exhibited only 10 total haplotypes (Fig. 1), microsatellite data revealed 35 MLGs within these samples, which was consistent with their higher mutation rate (Fig. 3). Despite its low bootstrap values (data not shown), the unrooted microsatellite NJ tree that we constructed (Fig. 4) showed a better resolution than the mitochondrial markers in the different subclades (Figs 1 and 2A).
We first analyzed the samples from the Western subclade (W) that had been distributed into only two mitochondrial haplotypes (groups W1 and W2 differing by a single substitution). Within the individuals from the widespread haplotype W1 (samples mainly from Europe and North America), microsatellite polymorphism was very limited due to a highly prevalent MLG, and we designated these individuals as the poorly polymorphic subgroup W1-PP (Figs 3 and 4). This W1-PP subgroup showed only four variant genotypes, of which two were monophyletic and differed by a single mutation from the dominant genotype and two were highly divergent (5 and 6 mutations) (Figs 3 and 4). These latter genotypes were distant from the cluster of the other genotypes (Fig. 4). Figure 3 also shows the genotypes of 10 additional samples (not included in the subset) that were classified in the W mitochondrial group according to their CytB haplotype. All 10 samples, despite their diverse origins (France, USA, Portugal, Spain and Italy), belonged to the predominant MLG and confirmed its wide and frequent distribution. Considering the 35 representative samples, the individuals from haplotype W2 clustered into two contrasting microsatellite subgroups. The first grouped the polymorphic samples (W2-P) that we detected in three regions (southern Iberian Peninsula, Bordeaux and Provence). This W2-P subgroup was very diverse, with 13 total MLGs, of which 10 were monophyletic and three were paraphyletic (Figs 3 and 4). The second microsatellite subgroup contained three samples from Chile that corresponded to a single MLG (Figs 3 and 4). Six additional samples from five other locations in Chile and one location in Argentina (Fig. 3) also belonged to this genotype, and we designated this monomorphic South American subgroup as W2-CA.

Analysis of the diversity of the NME subclade revealed nine total MLGs. Its individuals were dispersed in the tree, but the four most extreme variants belonged to the Iranian samples (3) and the Turkish sample Alasehir (1). The four samples of subclade IP showed four MLGs that were gathered in sister groups, as expected from their geographical proximity (Figs 3 and 4). By contrast, the three samples from subclade SSI were split into several clusters that were not sister groups (Figs 3 and 4).
Discussion

Genetic diversity of mitochondrial sequences. Our study first allowed us to evaluate the overall diversity of *X. index*, a species reproducing mainly by meiotic parthenogenesis. Using the common *CytB* gene with a wide range of nematode geographical origins, we revealed its low intraspecific divergence. As expected from a species with this reproduction mode, divergence (0 to 0.012) was lower than within the amphimictic sister species *X. diversicaudatum* (0.018 to 0.074)\(^4\), another important virus vector nematode of grapevine. Our study then reported polymorphisms from three additional genes. Among them, *ATP6* was identified for the first time in *X. index* and turned out to be the most polymorphic of the four mitochondrial sequences tested. The low diversity between those genes (ranging from 1.40% in *ND4* to 1.82% in *ATP6*) confirmed the previous data obtained with the *CO1* gene\(^4\). Similar results have been obtained for the mitotic parthenogenetic plant-parasitic nematodes *Meloidogyne* spp.\(^4\). Nevertheless, diversity values are much lower than in the amphimictic species *G. pallida*, which showed 12% *CytB* divergence among native Peruvian clades associated with relatively high levels of diversity and gene flow\(^4\).

The phylogeography of *X. index* might be closely linked to grapevine. The nematode is ectoparasitic and able to survive for many years in soil. Its long-distance dispersal is linked with grapevine dissemination by man through the transport of rooted plants together with their substrate\(^4\). The oldest domesticated grape seeds that have been discovered suggest an origin of grape cultivation in approximately 6000 BCE in Georgia and northeastern Turkey\(^4\). Additionally, the earliest evidence of grape production dates back to 5400–5000 BCE and has been found in the Northern Zagros mountains in northwestern Iran\(^4\). A Middle Eastern grape-growing and wine-making area was presumably located in Northern Mesopotamia and in the Turkish mountains from eastern Taurus\(^4\). From there, grape cultivars have presumably been transplanted into the Central and Southern Zagros Mountains (3000 BCE) in eastern Mesopotamia and the Jordan Valley to Egypt (circa 4000–3000 BCE) in...
western Mesopotamia\textsuperscript{50}. The available data on grapevine nuclear microsatellite diversity corroborate these putative ancestral domestication events occurring in the Caucasus and the Fertile Crescent\textsuperscript{51}.

Mitochondrial phylogeography suggests that the Near and Middle East contains the native area of \textit{X. index}. Interestingly, even though sampling has been limited, detection of \textit{X. index} has not been reported in native forests or climax vegetation\textsuperscript{12}, except in the Middle East. In this region, its presence has been reported in Iranian natural woodland, e.g., in the forests along the Caspian Sea, where wild grapevines may be common (Sturhan in Weischer)\textsuperscript{13}. This explains why \textit{X. index} occurs frequently in the cultivated grapevines of northern and western Iran\textsuperscript{14}. In our results, the highest mitochondrial diversity was observed in the Near and Middle East subclade. In particular, the evidence of three different and unique haplotypes in the Tabriz and Sharekord Iranian samples suggests that this region is the closest to the native area of \textit{X. index}. The microsatellite analysis also revealed a higher diversity in the NME subclade. All these results are in line with a scenario in which the NME area would contain or be the closest to the cradle of \textit{X. index} prior to its dispersal by man through grapevine domestication.

Putative dissemination of eastern samples belonging to the Near and Middle East subclade.

In the hypothesis of an expansion of domesticated grapevine from Southern Caucasus regions, it appears credible that \textit{X. index} has been spread with its host by two different routes. A first route to the west may have produced the NME subclade (from which the W subclade has then emerged), whereas a second route in the southern direction (through southern Mesopotamia) may have given birth to the subclades IP and SSI (Fig. 2B). This is in agreement with grapevine historical data\textsuperscript{8,49,52} and chloroplast DNA and microsatellite information\textsuperscript{50,51}, which show a diffusion of viticulture from the Near East through two routes around the Mediterranean Basin. A northern route was traced from eastern to Western Europe (Hittite, Phrygian, Greek and Roman people in the Antiquity)\textsuperscript{49} and a southern route went through Egypt and the Maghreb up to Gibraltar and the Iberian Peninsula (via Phoenicians and Romans in Antiquity and Arabs from the 7th century ACE).

In the northern route, domesticated grapevines appeared in Asia Minor, southern Greece, Crete and Cyprus in 3000–2500 BCE and in the southern Balkans\textsuperscript{80} in 2000–1500 BCE. They finally reached southern Italy, southern

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**Figure 4.** Unrooted tree of individuals of \textit{X. index} based on microsatellite multilocus genotypes. Data are from the subset of 35 representative samples. The NJ tree based on distance CSE was performed using 2000 iterations. The colors (pink: W1; orange: W2; blue: NME; green: IP; grey: SSI) correspond to the mitochondrial topology (see Figs 1 and 2A). For sample codes, see Table S1.
France, Spain and Portugal in the first millennium BCE. For *X. index*, the NME subclade matches the area of the Greek civilization (western Turkey, Cyprus, Crete and Egypt) and might represent the northern Mediterranean route of grapevine varieties. Nevertheless, the detection of an Egyptian sample that marks the southern part of this putative Greek dissemination area suggests an introduction of the nematode there during late Antiquity.

**Putative dissemination of eastern samples belonging to the IP and SSI subclades.** With the historical expansion of grapevine in mind, we may hypothesize that the samples from Israel and Palestine (subclade IP) are initial steps along the southern route. Nevertheless, an overlap of the NME and IP subclades has certainly occurred, at least in Lebanon. Indeed, within *CytB* from single individuals recovered in two Lebanese locations, we obtained two different haplotypes. The first individual (Xi-Le-Le; from Kefraya) had the specific IP haplotype, while the second individual (Xi-Le-Le; from an unspecified location) belonged to the most common NME haplotype (e.g., the same as Xi-Cy-Cy) (Fig. S1).

Samples from Samos and South Italy (subclade SSI) might originate from the same region, as they are closely related to the IP subclade (Fig. 2B). From our microsatellite data (Fig. 4), the MLGs Xi-Gr-Sa-A and Xi-Gr-Sa-B from Samos on the one hand and the MLGs Xi-It-Pa and Xi-It-PC from South Italy on the other hand do not belong to sister groups. This result argues for an ancient divergence between them. In Antiquity (2000–500 BCE), the Lebanon area was under the authority of the Phoenician kingdom, which had also settled many trade harbors along the southern Mediterranean coast to the southern Iberian Peninsula (Malaga, Cadiz and Lisbon). Phoenicians had a great influence on the spread of grapevine across the Mediterranean Basin, but it is unlikely that they had trade harbors in the northern Mediterranean region. Consequently, nematode introduction from the SSI subclade may result from Greek and/or Phoenician trading in the last two millennia (1500–200 BCE) or later, during the Roman Empire.

**Putative origin and dissemination of nematodes of the Western subclade.** The western samples of *X. index* are closely linked to the NME subclade. We may hypothesize that they originated from it. The nematode might have been introduced by the Romans after the final unification of Mediterranean territories into the Roman Empire. An alternative hypothesis is that the Greeks had introduced *X. index* even earlier via their distant trading in northern Mediterranean harbors in southern Italy and southern France (1500–200 BCE) (Fig. 2B). Both W1-PP and W2-P might originate from the same Near or Middle Eastern location.

**Putative dissemination of nematodes of the Western 2 group (W2-P and W2-CA).** The high diversity of MLGs in the W2-P samples suggests that repeated introductions from the same mitochondrial haplotype have occurred during the Roman Empire epoch (100 BCE–200 ACE), after the unification of the western Mediterranean territories. Fréjus (in Provence) was the capital of the new Roman Narbonensis Province in 22 BCE. It was a place of rapid development of viticulture. Almost simultaneously, in Bordeaux, archeological data show that cultivated grapevine was intensively developed from 50 to 250 ACE. The clustering of Fréjus, Bordeaux and southern Iberian Peninsula samples (for both mitochondrial and microsatellite data) argues for dissemination events that would have occurred on a limited time scale and possibly from a common origin. The long and peaceful Roman period in the western Mediterranean may have promoted a wide-scale diffusion of grapevine together with the nematode at that time.

In the Iberian Peninsula (and a fortiori in its southern part), chloroplast and microsatellite diversities support the hypothesis of an earlier introduction of grapevine, probably by the Phoenicians, making this region a secondary center of domestication. If we consider that the SSI subclade is linked with Phoenician grapevine dissemination, we should have detected SSI-linked individuals within the samples from the southern Iberian Peninsula. As we did not, our current results are not in line with the hypothesis of another nematode route of Phoenician origin into this region. However, it is also plausible that we failed to track the putative *X. index* to the Iberian Peninsula because the early-introduced Phoenician grapevines were nematode-free.

In America, cultivated grapevine has been imported from Europe since the 16th century by Christian missionaries. Introduction of *X. index* into the New World certainly occurred through the Spanish colonization of South America. The monomorphism of these samples (W2-CA) (Figs 3 and 4) suggests a single introduction event into Argentina and Chile that may have occurred directly from Spain or indirectly from an intermediate region such as the Canary Islands. Interestingly, few grapevine varieties from the Old World were detected in Chile. This low diversity suggests that the plant material originated from few introduction events and/or from a limited area of Spain, which is in line with the absence of nematode polymorphism in our Southern American samples.

**Putative origin and dissemination of the Western 1 group (W1-PP).** The samples from the second Western haplotype (W1-PP) are almost monomorphic and, within them, an identical genotype ranges from Hungary and Crete eastward to western and southwestern Europe and to Madeira and California westward (Fig. 3). This suggests a recent spread of this haplotype, presumably from a geographically limited monomorphic focus. We hypothesize that this recent dissemination followed the massive replanting of vineyards in a large part of the worldwide grapevine distribution area after the 19th-century phylloxera crisis. The invariant W1 MLG may originate from a location initially contaminated by a single introduction event and that has provided nematode-infected rootstocks for the stepwise replanting of highly distant vineyards. Interestingly, the W1-PP samples exhibit four variant genotypes, two of which are highly different and originate from Apulia (Terlizzi) and North Crete, respectively, and two of which are less variable but originate from the same Apulian sample and from Sardinia. Detecting such a polymorphism in southern Greece and southern Italy is in line with the hypothesis of a Greco-Roman origin of some or all of the samples of the Western subclade (Fig. 2B).
Unlike the putative nematode introduction event(s) from the 16th century in South America, our data suggest that the introduction in California occurred later in the 19th–20th centuries with vines grafted on European phylloxera-resistant rootstocks. Interestingly, several locations from the southern Iberian Peninsula, Bordeaux and Provence harbor both W1-PP and W2-P individuals, which illustrates their dual historical infection.

The global picture of X. index diversity is an ongoing study. Although our study has benefited from a worldwide range of samples covering 16 countries over four continents, many other relevant areas need to be explored to confirm and refine the portrait of dissemination that we have drawn.

The eastern Mediterranean area shows the highest diversity, but our few geographical sampling points provide only preliminary insight into this region. For instance, other countries such as Syria, Iraq, Armenia and Georgia will have to be considered. Future efforts will also aim to decipher the Phoenician heritage and its putative routes from Lebanon. We lack data from Northern Africa that could show whether a specific dissemination route by the Phoenicians and/or later by the Arabs exists in southern Mediterranean countries. Xiphinema index is present in Maghreb countries60 in grapevine plots planted by the French in the 19th–20th centuries after European colonization, and consequently, at least individuals from the W1 widespread genotype should be found there.

In South America and, in particular, in Andean countries, a survey for new samples will bring additional information on the marked bottleneck effect that we revealed in Chile and Argentina. In the Southern Hemisphere, other vineyards installed by Western colonizers exist in South Africa and Australia10,11 and should be sampled.

The diversity of this nematode and its history of introduction in those distant areas will be worth investigating. Another effort should be dedicated to the putative dispersal of nematode-contaminated plant material after the phylloxera crisis at the end of the 19th century. Because of their recent occurrence, these events should be easier to document. Revealing the putative scenarios explaining the spread of a single genotype over such a vast geographical area would be a challenge.

Finally, X. index appears to be a marker of the dispersion of grapevine, and future data may improve our knowledge of the history of this fascinating crop as a witness to human history.

Data Availability
EBI accession numbers of the CyTB, ATP6, COI, and ND4 mitochondrial gene sequences (LT996601 to LT996818) are reported in Table S1. Sequences have been uploaded at the: http://www.ebi.ac.uk/ena/data/view/PRJEB26007

They will be available upon manuscript acceptance.

References
1. Decraemer, W. & Hunt, D. J. In Plant nematology (eds Perry, R. N. & Moens, M.) 3–32 (CABI International, 2006).
2. Nicol, J. M. et al. In Genomics and molecular genetics of plant-nematode interactions (eds Jones, J., Ghesyen, G. & Fenoll, C.) 21–43 (Springer Netherlands, 2011).
3. Hewitt, W. B., Raski, D. J. & Goheen, A. C. Nematode vector of soil-borne fanleaf virus of grapevines. Phytopathology 48, 586–595 (1958).
4. Taylor, C. E. & Brown, D. J. E. Nematode vectors of plant viruses. (CAB International, 1997).
5. Jones, J. T. et al. Top plant-parasitic nematodes in molecular plant pathology. Mol. Plant Pathol. 14, 946–961 (2013).
6. Raski, D. J., Goheen, A. C., Lider, L. A. & Meredith, C. P. Strategies against Grapevine fanleaf virus and its nematode vector. Plant Dis. 67, 335–338 (1983).
7. Martelli, G. P. & Savino, V. In Compendium of grape diseases (eds Pearson, R. & Goheen, A.) 48–49 (APS Press, 1990).
8. Andrè-Lint, P. et al. Grapevine fanleaf virus: Still a major threat to the grapevine industry. J. Plant Pathol. 86, 183–195 (2004).
9. This, P., Lacombe, T. & Thomas, M. R. Historical origins and genetic diversity of wine grapes. Trends Genet. 22, 511–519 (2006).
10. Siddiqui, M. R. In C.I.H Descriptions of plant parasitic nematodes Set 3, No 45. (Commonwealth Institute of Parasitology, C.A.B. International, 1986).
11. Van Zyl, S., Vivier, M. A. & Walker, M. A. Xiphinema index and its relationship to grapevines: A review. S. Afr. J. Enol. Vitic. 33, 21–32 (2012).
12. Esenjaud, D. In Raveurages de la vigne (eds Esenjaud, D. et al.) 18–45 (Éditions Ferté, Bordeaux, 2008).
13. Weisker, B. In Nematode vectors of plant viruses (eds Lamberti, F., Taylor, C. & Seinhorst, J.) 291–307 (NATO-Plenum Press, 1975).
14. Mofajahed, H., Sturhan, D., Akhiani, A. & Barouti, S. Xiphinema species in Iranian vineyards. Nematol. Medit. 8, 165–170 (1980).
15. Esenjaud, D., Bouquet, A., Demangeat, G., Van Helden, M. & Ollat, N. Nematode-resistant rootstocks as a major component of the management alternative for Grapevine fanleaf virus control in grape. Acta Hortic. 904, 111–115 (2011).
16. Villate, L., Morin, E., Demangeat, G., Van Helden, M. & Esenjaud, D. Control of Xiphinema index populations by fallow plants under greenhouse and field conditions. Phytopathology 102, 627–634 (2012).
17. McDonald, B. A. & Linde, C. Pathogen population genetics, evolutionary potential, and durable resistance. Annu. Rev. Phytopathol. 40, 349–379 (2002).
18. Dalmasso, A. Gametogenesis of genus Xiphinema and Longidorus (Nematoda: Dorylaimida). Comptes Rendus Acad. Sci. Serie D: Sciences naturelles 6, 824–827 (1970).
19. Villat, L., Esenjaud, D., Van Helden, M., Stoeckel, S. & Plantard, O. Genetic signature of amphimixis allows for the detection and fine scale localization of sexual reproduction events in a mainly parthenogenetic nematode. Mol. Ecol. 19, 856–873 (2010).
20. Allendorf, F. W. & Lundquist, L. L. Introduction: population biology, evolution, and control of invasive species. Conserv. Biol. 17, 24–30 (2003).
21. Hewitt, W. B. Virus and virus diseases of the grapevine. Rev. Appl. Mycol. 47, 433–455 (1968).
22. Martelli, G. P. & Taylor, C. E. In Advances in disease vector research. 151–189 (Springer, 1990).
23. Brown, D. J. F., Taylor, C. E., Choleva, B. & Romanenko, N. D. The occurrence of Longidoridae (Nematoda: Dorylaimida) in western USSR with further comments on longidorid nematodes in Europe and the Mediterranean Basin. Nematol. Medit. 18, 199–207 (1990).
24. Robbins, R. T. & Brown, D. J. F. Comments on the taxonomy, occurrence and distribution of Longidoridae (Nematoda) in North America. Nematologica 37, 395–419 (1991).
25. Doucet, M. E., Ferraz, L. C. C. B., Magunacelaya, J. C. & Brown, D. J. F. The occurrence and distribution of longidorid nematodes in Latin America. Rev. J. Nematol. 6, 111–128 (1998).
26. Plantard, O. et al. Origin and genetic diversity of Western European populations of the potato cyst nematode (Globodera pallida) inferred from mitochondrial sequences and microsatellite loci. Mol. Ecol. 17, 2208–2218 (2008).
27. Lilja, T., Troell, K., Kirik, H. & Lindström, A. A distinct group of north European Aedes vexans as determined by mitochondrial and nuclear markers. Med. Vet. Entomol. 32, 282–289 (2018).
28. Viret, A. et al. Absence of spatial genetic structure in common dentex (Dentex dentex Linnaeus, 1758) in the Mediterranean Sea as evidenced by nuclear and mitochondrial molecular markers. PLoS one 13, e019866 (2018).
29. Kawamura, K., Kaieda, S., Kato, M. & Kobayashi, S. Invasion genetics of nutria (Myocastor coypus) in Okayama, Japan, inferred from mitochondrial and microsatellite markers. Eur. J. Wildlife Res. 64, 1–13 (2018).
30. Villate, L., Esbenjaud, D., Coedel, S. & Plantard, O. Development of nine polymorphic microsatellite markers for the phytoparasitic nematode Xiphinema index, the vector of the grapevine fanleaf virus. Mol. Ecol. Res. 9, 229–232 (2009).
31. Molnar, R. I., Witte, H., Dinkelacker, I., Villate, L. & Sommer, R. J. Tandem-repeat patterns and mutation rates in microsatellites of the nematode model organism Pristionchus pacificus. G3 (Bethesda) 2, 1023–1034 (2012).
32. Molnar, R. I., Bartelmes, G., Dinkelacker, I., Witte, H. & Sommer, R. J. Mutation rates and intraspecific divergence of the mitochondrial genome of Pristionchus pacificus. Mol. Biol. Evol. 28, 2317–2326 (2011).
33. Villate, L. et al. Spatial distribution of the dagger nematode Xiphinema index and its associated Grapevine fanleaf virus in French vineyard. Phytopathology 98, 942–948 (2008).
34. He, Y., Jones, J., Armstrong, M., Lamberti, F. & Moens, M. The mitochondrial genome of Pristionchus americanum sensu stricto (Nematoda: Enoplea): Considerable ecozonization in the length and structural features of encoded genes. J. Mol. Evol. 61, 819–833 (2005).
35. Tamura, K., Stecher, G., Peterson, D., Filipski, A. & Kumar, S. MEGA6: molecular evolutionary genetics analysis version 6.0. Mol. Biol. Evol. 30, 2725–2729 (2013).
36. Kishino, H. & Hasegawa, M. Evaluation of the maximum likelihood estimate of the evolutionary tree topologies from DNA sequence data, and the branching order in Homininae. J. Mol. Evol. 29, 170–179 (1989).
37. Felsenstein, J. & Churchill, G. A. A Hidden Markov Model approach to variation among sites in rate of evolution. Mol. Biol. Evol. 13, 93–104 (1996).
38. Pagel, M. & Meade, A. A phylogenetic mixture model for detecting pattern-heterogeneity in gene sequence or character-state data. Syst. Biol. 53, 571–581 (2004).
39. Cavalli-Sforza, L. L. & Edwards, A. W. F. Phylogenetic analysis: models and estimation procedures. Evolution 21, 550–570 (1967).
40. Nguyen, V. C. Genetic diversity of the grapevine vector nematode Xiphinema index and application to optimize the resistance strategy. PhD thesis. Doctoral School ‘Sciences de la Vie et de la Santé – Université Côte d’Azur, France, 179 pp (2018).
41. Gutiérrez-Gutiérrez, C. et al. Genetic structure of Xiphinema pachtaicum and X. index populations based on mitochondrial DNA variation. Phytopathology 101, 1168–1175 (2011).
42. Hugall, A., Moritz, C., Stanton, J. & Wolstenholme, D. R. Low, but strongly structured mitochondrial DNA diversity in root knot nematodes (Meloidogyne). Genetics 136, 903–912 (1994).
43. Picard, D., Sempere, T. & Plantard, O. A northward colonisation of the Andes by the potato cyst nematode during geological times suggests multiple host-shifts from wild to cultivated potatoes. Mol. Phylogenet. Evol. 42, 308–316 (2007).
44. Berkowitz, M. World’s earliest wine. Archaeology 49, 26 (1996).
45. Blasco, J. L. In L’Histoire du vin, une histoire de rites (ed. O. I. V.) 31–41 (Salomon, Paris, 1997).
46. Bottero, J. In L’Histoire du vin, une histoire de rites (ed. O. I. V.) 13–24 (Salomon, Paris, 1997).
47. McGovern, P. E., Glusker, D. L. & Exner, L. J. Neolithic resinated wine. J. Am. Soc. Sci. 93, 91–100 (1991).
48. Balmelle, C. Progress in Old World palaeoethnobotany. J. Roy. Anth. Soc. Touraine 11, 93–100 (1991).
49. Sechrist, R. World’s earliest wine. Aust. J. Grape Wine Res. 19, 299–310 (2013).
50. Pougnet, R. In La vigne et le vin 15–20 (La Manufacture et la Cite des sciences et de l’industrie, Graficas, Paris, 1988).
51. Sechrist, R. Planet of the grapes: a geography of wine. (Preager ABC-CLIO, 2017).
52. Arroyo-García, R. et al. Multiple origins of cultivated grapevine (Vitis vinifera l. ssp. sativa) based on chloroplast DNA polymorphisms. Mol. Ecol. 15, 3707–3714 (2006).
53. Bacilleri, R. et al. Genetic structure in cultivated grapevines is linked to geography and human selection. BMC Plant Biol. 13, 25 (2013).
54. Zohary, D. In The origins and ancient history of wine (eds Mc Govern, P., Fleming, S. & Katz, S.) 23–30 (Gordon and Breach, 1996).
55. Zohary, D. In Progress in Old World palaeoethnobotany (eds Van Zeist, W., Wasylikowa, K. & Behre, K.) 161–177 (Balkema, Rotterdam, 1991).
56. Hopf, M. In Progress in Old World palaeoethnobotany (eds Van Zeist, W., Wasylikowa, K. & Behre, K.) 241–277 (Balkema, Rotterdam, 1991).
57. Balmelle, C. et al. La viticulture antique en Aquitaine. Gallia 129–164 (2001).
58. Tapia, A. M. et al. Determining the Spanish origin of representative American grapevine varieties. Am. J. Enol. Vitic. 58, 242–251 (2007).
59. Millia-Tapia, A. et al. Naturalised grapevines collected from arid regions in Northern Chile exhibit a high level of genetic diversity. Aust. J. Grape Wine Res. 19, 299–310 (2013).
60. Pougnet, R. In La vigne et le vin 15–20 (La Manufacture et la Cité des sciences et de l’industrie, Graficas, Paris, 1988).
61. Pougnet, R. Le Phylloxera et les maladies de la vigne. La lutte victorieuse des savants et des vignerons français (1850–1900) (édilivre, St Denis, France, 2015).
62. Dalmasso, A. & Cuany, A. Importance of the transmission par le sol de la dégénérescence infectieuse de la vigne dans l’Algérois. FAO Plant Prot. Bull. 17, 58–60 (1969).

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Author Contributions
D.E., V.C.N. and L.V. designed the study; D.E., V.C.N., L.V., P.C. and C.G.G. collected the samples; V.C.N. and D.E. conducted the literature review; V.C.N., D.E., C.V.G. and L.V. analyzed the data; V.C.N. and D.E. wrote the paper with input from L.V. and O.P.

Additional Information
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