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Sixty years from the first disease description, a novel badnavirus associated with chestnut mosaic disease

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2 chestnut mosaic disease

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- 22 The nucleotide sequences reported here have been deposited in GenBank under the accession
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ABSTRACT

Although the chestnut mosaic disease (ChMD) was described several decades ago, its etiology is still 26 not elucidated. Here, using classical approaches in combination with high throughput sequencing 27 (HTS) techniques, we identify a novel *Badnavirus* that is a strong etiological candidate for ChMD. 28 Two disease sources from Italy and France were submitted to HTS-based viral indexing. Total RNAs 29 30 were extracted, ribodepleted and sequenced on an Illumina NextSeq500 (2x150 or 2x 75 nt). In each source, we identified a single contig of about 7.2 kilobases that corresponds to a complete circular 31 32 viral genome and shares homologies with various badnaviruses. The genomes of the two isolates have an average nucleotide identity of 90.5% with a typical badnaviral genome organization 33 comprising three open reading frames. Phylogenetic analyses and sequence comparisons show that 34 35 this virus is a novel species for which we propose the name Chestnut mosaic virus (ChMV). Using a 36 newly developed molecular detection test, we systematically detected the virus in symptomatic graft-inoculated indicator plants (chestnut and American oak), as well in chestnut trees presenting 37 typical ChMD symptoms in the field (100% and 87% in France and Italy surveys, respectively). 38 39 Datamining of publicly available chestnut SRA transcriptomic data allowed the reconstruction of two additional complete ChMV genomes from two Castanea mollissima sources from the USA, as well 40 41 as ChMV detection in *C. dentata* from the USA. Preliminary epidemiological studies, performed in France and in Central Eastern Italy, showed that ChMV has a high incidence in some commercial 42 43 orchards, with a low within-orchard genetic diversity.

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46 European chestnut (Castanea sativa Mill.) has a long-standing tradition of cultivation in many European countries. It is an important species, both economically as a source of timber and fruit 47 and ecologically through the multiple ecosystemic services it provides. In Europe, chestnut covers 48 about 2.5 million hectares, mainly concentrated in France, Italy, Spain, Portugal, Switzerland, the 49 Balkan regions, and Southern England (Conedera et al. 2016). Chestnut (Castanea spp.) can be 50 51 heavily affected by various pathogens. The most detrimental are caused by fungal-like organisms (Oomycetes) and fungi such as Phytophthora cambivora Petri and P. cinnamomic Rands., the agents 52 of ink disease, or Cryphonectria parasitica, the causal agent of chestnut blight, which all provoke 53 disorders that can lead to tree mortality (Prospero et al. 2012; Rigling and Prospero, 2018). In Italy, 54 Gualaccini (1958) described a chestnut disease associated with viral symptoms (mosaic, shoots with 55 asymmetric leaf blade deformation) which was again reported in Campania during the 80s, 56 (Ragozzino and Lahoz 1986), and in the Marche region (central eastern Italy) in 2000 (Antonaroli 57 and Perna; 2000). In France, the disease was first identified circa 1987 on cultivars of C. sativa x C. 58 crenata hybrids from commercial orchards located in the south-west of the country. Desvignes 59 60 (1999b) made a more detailed description of the symptoms, which are characterized by necrotic lesions in the bark and wood that turns into cankers, chlorotic lesions and yellow stripes on leaf 61 62 veins and partial limb atrophy, and called this disease Chestnut Mosaic Disease (ChMD). This disease can heavily affect the production of both young and secular trees (Antonaroli and Perna, 2000). It 63 64 has also been reported in Japan and Hungary (Shimada, 1962; Horvath et al. 1975). Even if its etiology has long remained unknown, researchers hypothesized that the causal agent of ChMD 65 could be a virus, introduced in Europe between 1940 and 1960 when a number of C. crenata 66 67 cultivars were imported from Japan for breeding purposes. Investigations in France and Italy 68 established that the causal agent can be eliminated by thermotherapy, is aphid-transmissible, and 69 is graft-transmissible to Castanea and Quercus species in which it may elicit symptoms (Desvignes

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70 and Lecocq, 1995; Desvignes, 1999b; Vettraino et al. 2005). The susceptibility to the ChMD agent of Castanea species/cultivars has been evaluated in several studies (Desvignes, 1992; 1999b; 71 72 Desvignes and Lecoq, 1995). Three categories of cultivars could thus be defined from tolerant to moderately and fully susceptible. Graft incompatibility was also observed when cultivars of different 73 susceptibilities are assembled by grafting. Most of the C. sativa cultivars and hybrids are tolerant to 74 75 ChMD, although some well-known French hybrids like 'Maraval' (Ca 74) are fully susceptible, and used for indexing purposes to detect the ChMD agent in tolerant cultivars (Desvignes and Lecoq, 76 1995). 77

In the last decade, a number of studies have highlighted the potential of non-targeted molecular 78 diagnostics based on high-throughput sequencing (HTS) to elucidate the etiology of viral plant 79 80 diseases and to provide viral sequence data from which rapid diagnostic molecular assays can be developed (Martin et al. 2016; Villamor et al. 2019). Since 2009, HTS combined with bioinformatics 81 have been used for the discovery, characterization, and *de novo* assembly of the genome of known 82 and novel plant viruses and viroids (Rott et al. 2017; Kreuze et al. 2009). This has accelerated the 83 application of HTS technologies in the field diagnostic of diseases (Massart et al. 2014), and in 84 quarantine regulations (Martin et al. 2016; Massart et al. 2017). 85

86 Badnaviruses are plant pararetroviruses belonging to the family Caulimoviridae that have 87 emerged as serious pathogens causing severe yield losses in a wide range of economically important 88 crops all over the world (Bhat et al. 2016). The genome of badnaviruses is composed of a noncovalently closed, circular double-stranded DNA (ranging from 7.2 to 9.2 kbp) and is encapsidated 89 in bacilliform virions. This genome typically harbors three open reading frames (ORFs) encoding, 90 91 respectively, a protein of unknown function, the virion-associated protein (VAP), and a polyprotein 92 containing functional and structural domains [movement protein (MP), coat protein (CP), aspartic 93 protease (AP), reverse-transcriptase (RT) and RNase H)] (Hohn and Rothnie, 2013; Bhat et al. 2016).

Badnaviruses can also be present as integrated sequences in some host plant genomes (endogenous
badnaviruses) (Staginnus et al. 2009; Bhat et al. 2016). The contribution of these integrated
sequences to host and virus evolution is still poorly understood (Geering et al. 2014).

Given the very limited knowledge on the etiology of ChMD, and based on previously published 97 studies (Desvignes, 1992; 1999a, 1999b; Desvignes and Lecoq, 1995; Desvignes and Cornaggia, 98 99 1996), we investigated the hypothesis that a virus might be involved in this disease. Combining HTSbased viral indexing and classical approaches, we report here the complete genome sequence of a 100 101 novel badnavirus species for which the name Chestnut mosaic virus (ChMV) is proposed. We further show that there is a strict correlation between the presence of the virus and the appearance of 102 typical ChMD symptoms in various graft-inoculated indicator plants. Preliminary epidemiological 103 studies carried out in Italy and in France reveal that the virus can have high incidence in some 104 105 orchards, and, as expected, can be associated with symptomatic or asymptomatic infections.

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MATERIALS AND METHODS

Plant samples and virus isolates. Virus isolates included in this study are listed in Supplementary 108 Table S1. Isolate LC1224H is originated from a red oak (*Quercus rubra*) artificially inoculated in 1992 109 110 with a chestnut mosaic source from a hybrid Castanea sativa x Castanea crenata included in a French breeding program. Leaves of grafted oaks displayed typical symptoms characterized by 111 112 chlorotic mottle, yellow veins, and mosaic (Desvignes and Lecoq 1995) (Figure 1A). Isolate FRIc1224A was derived from the same source, and is the result of a back-inoculation by grafting of 113 LC1224H to the natural chestnut hybrid Maraval (Ca 74; C. crenata x C. sativa) indicator (Desvignes 114 115 et al. 1992). Isolate LC1224F originated from a Maraval indicator inoculated by aphid transmission 116 from an initial ChMD source in a C. crenata x C. sativa French hybrid (Desvignes and Cornaggia, 117 1996). The LCA552 and LCA584 isolates were collected from C. sativa trees in France in 2009 and

118	2018, while the T32018 disease source was isolated from a French hybrid C. crenata x C. sativa in
119	2018. All of these isolates have been held and propagated on 'Maraval' indicator plants at the CTIFL
120	virology laboratory (Lanxade, France).

121 In the framework of a survey carried out in Italian chestnut orchards to monitor chestnut blight 122 (Acquasanta Terme (AP), locality Umito, Italy) (Murolo et al. 2018), typical leaf symptoms of ChMD 123 were recorded in 2016. Six symptomatic plants were collected, pooled (10 -15 symptomatic shoots) 124 and included in the HTS analysis (ITumito39 source).

In order to evaluate the incidence of ChMV, chestnut trees from INRAE chestnut biological 125 resource center (https://www6.bordeaux-aquitaine.inrae.fr/biogeco/Ressources) were sampled. 126 This orchard is located on the Villenave d'Ornon INRAE center (France) with trees distributed in 127 three plots (A, E, or Port, Table S1). A total of 43 C. sativa, 14 C. mollissima, six C. crenata and 32 128 129 hybrid chestnut trees were sampled, corresponding to a total of 38 symptomatic trees with typical ChMD symptoms, 47 asymptomatic trees, and 10 trees with dubious or atypical symptoms. In 130 addition, in the Central eastern Italy Marche region, leaves from 60 symptomatic and from 10 131 asymptomatic grafted C. sativa cv. Marrone trees of different ages were collected in a commercial 132 chestnut orchard (Plot I, Table S1). 133

Isolates FRIc1224A and ITumito39 were used for the HTS analysis, whereas all other samples were
 included either in the incidence analysis or in the causal relationship analysis (Table S1).

Total RNA extraction and RNA-Seq analysis. Symptomatic leaves from a 'Maraval' indicator (FRIc1224A) were collected and used to extract total RNAs according to the protocol described by Reid et al. (2006). For the Italian material, total RNAs were extracted from symptomatic leaves according to the protocol described by Gambino et al. (2008). Total RNAs were then submitted to a DNAse treatment following the manufacturer's recommendations (Fisher Scientific, Illkirch, France). Ribosomal RNAs were removed using a RiboMinus Plant Kit for RNA-Seq (Invitrogen, Fisher Scientific, Illkirch, France) before cDNA library synthesis with the Illumina TruSeq Stranded RNA
library Prep kit (Illumina, Inc., San Diego, CA) and sequenced on an Illumina NextSeq500 (2x150 nt
or 2x75 nt) in a multiplexed format (GIGA-Genomics facility, Université de Liège, Belgium).

Bioinformatic analysis. Primary quality analyses were performed using Geneious Prime 145 2019.2.1 Software (https://www.geneious.com). *De novo* assemblies of quality filtered reads were 146 performed using Velvet (Zerbino and Birney, 2008), Geneious R 11 (https://www.geneious.com), 147 Spades (Bankevich et al. 2012), or using the CLC genomics workbench 8.0 148 and http://www.clcbio.com). Contigs were annotated by BlastN and BlastX comparisons with nucleotide 149 and non-redundant protein GenBank databases, respectively. Blast results were screened using e-150 value thresholds of 10⁻⁶ and 10⁻⁴ for BlastN and BlastX, respectively. Publicly available chestnut RNA-151 Seq transcriptomic data were retrieved from the NCBI Sequence Read Archive (SRA) and 152 downloaded reads were mapped against the sequence of the FRIc1224A isolate using CLC Genomics 153 Workbench 11.0. When needed, de novo assembly and contig annotations were also performed as 154 described above. 155

Total DNA extraction and PCR confirmation of genome completeness and circularity. In order 156 to verify both the completeness of the assembled genome sequences and genome circularity, pairs 157 158 of specific outward-facing primers were designed for each isolate (Ch-Bad-6976F/ Ch-Bad-252R for the isolate FRIc1224A and Bad-Ch-6481F/Bad-Ch-325R for the isolate ITumito39, Table S2). Leaf 159 160 tissues (0.5 g) were pulverized in liquid nitrogen and total DNA was extracted in CTAB buffer (2% cetyl trimethylammonium bromide, 100 mM Tris-HCl, 1.4 M NaCl, 20 mM EDTA), adding 3% 161 polyvinyl pyrrolidone 40, and 0.5% sodium metabisulfite (Doyle and Doyle, 1990). Finally, the DNA 162 163 pellets were resuspended in 50 µl sterile water. PCRs were performed in a 50 µl reaction volume 164 containing 10 mM Tris-HCl (pH 8.5), 2 mM MgCl₂, 50 mM KCl, 0.2 mM dNTPs, forward and reverse 165 primers at 1µM each, and 1.25U of Dream Taq (ThermoFisher) using 50 ng of the template. After an

initial denaturation step at 95°C for 4 min, 40 or 35 cycles, respectively, were set at 94°C for 30 sec,
60°C (Ch-Bad-6976F/ Ch-Bad-252R) or 55°C (Bad-Ch-6481F/325R) for 30 sec, and 72°C for 90 sec,
followed by a final extension step of 10 min at 72°C. PCR amplification products were sequenced on
both strands (GATC, Eurofins, Ebersberg, Germany).

ChMV molecular detection and variant analysis by PCR. For the molecular detection of ChMV, 170 two sets of primers were designed in conserved regions of ORF3 designed using the sequences of 171 isolates FRIc1224A and ITumito39. One primer pair (Ch-Bad-1466F/Ch-Bad-1800R, Table S2) allows 172 the amplification of a genomic region (335 nt) in the MP domain (Figure 2), whereas the second pair 173 174 (Ch-Bad-5860F/Ch-Bad-6109R, Table S2) amplifies a 232-nt fragment in the RH domain (Figure 2). 175 An aliquot of 25 ng of total DNA was used for the PCR assays in a 50 µl volume containing 10 mM Tris-HCl (pH 8.5), 2 mM MgCl₂, 50 mM KCl, 0.2 mM dNTPs, forward and reverse primers at 1 μ M 176 each, and either 1.25U of DreamTag or 1U of GoTag. After an initial denaturation step at 95°C for 4 177 min, 35 cycles were set at 94°C for 30 sec, 56°C for 30 sec, and 72°C for 90 sec, followed by a final 178 extension step of 10 min at 72°C. Amplicons were analyzed by electrophoresis on 1.5% agarose gel 179 180 and were directly sequenced on both strands (GATC).

Possible phytoplasma infection was evaluated using primer pair P1/P7 (Deng and Hiruki. 1991;
Smart et al. 1996) and, in nested PCR, primers R16F2n/R2 (Gundersen and Lee 1996).

Sequence and phylogenetic analyses. The full-length genomes were analyzed by ORF Finder (http://www.ncbi.nlm.nih.gov/projects/gorf/) to identify putative ORFs in the viral genome. Deduced amino acid (aa) sequences were analyzed for conserved protein domains gathered in Conserved Domains Database (CDD) (http://www.ncbi.nlm.nih.gov/structure/cdd.shtml) and theoretical molecular weights were calculated using ExPASy (http://web.expasy.org/compute_pi/). Multiple alignments of nucleotide (nt) or amino acid (aa) sequences were performed using the ClustalW program (Thompson et al. 1994) implemented in MEGA version 7.0 (Kumar et al. 2016).

190	Genetic distances (p-distances using strict nt or aa identity) were calculated using MEGA 7.0.
191	Phylogenetic trees were reconstructed using the neighbor-joining method implemented in MEGA
192	7.0 and robustness of nodes was assessed from 1,000 bootstrap resamplings.

193

RESULTS

Determination of the complete genome sequence of a novel badnavirus from two chestnut 194 disease sources. Two ChMD sources were included in the HTS analysis. The French source 195 (FRIc1224A) showed typical ChMD symptoms, with leaf deformation, yellow veins and chlorotic 196 diffuse mottling (Figure 1B) and the Italian source (ITumito39) is a mixture of six plants showing 197 intensive vein banding and leaf blade deformation (Figure 1C). High-throughput sequencing of 198 199 ribodepleted RNAs extracted from the sources FRIc1224A and ITumito39 yielded a total of 200 10,737,052 reads and 4,135,330 reads, respectively. *De novo* assembly and Blast annotation allowed 201 for the identification of a single long contig with significant homology with badnaviruses. These contigs were respectively 7,264 and 7,214 bp long and showed short terminal redundancies, 202 203 consistent with the structure of the long RNA transcript involved in the replication of badnaviruses (Teycheney et al. 2020) and suggesting they represented the full coverage of a circular badnaviral 204 genome. A total of 39,657 reads were integrated in the FRIc1224A contig, representing 0.37% of 205 206 total reads, with a mean coverage depth of 795X, whereas 611 reads (0.015% of total reads) were 207 integrated in the ITumito39 contig, with a mean coverage depth of 14.4X. The circularity and completion of the DNA genome sequence of each isolate were validated by a PCR on purified DNA 208 209 extracted from the host plants and using specific outward-facing primers designed from the contig 210 sequences. The respective 436 nt- and 1,007 nt fragments were amplified and sequenced, confirming DNA genome completeness and circularity (data not shown). The assembled sequences 211

have been deposited in GenBank under accession numbers MT269853 and MT261366, respectively.

213 No other plant virus was detected in the two datasets during the Blast annotation of contigs.

Genome organization of chestnut mosaic virus and determination of its phylogenetic 214 215 relationships. The badnaviral genomes characterized independently from the French and Italian ChMD sources are respectively 7,160 bp and 7,161 bp long, within the range of badnavirus genome 216 sizes (Teycheney et al. 2020). The genomic organization is the same for both isolates, comprising 217 three open reading frames (ORFs) encoded on the positive strand (Figure 2), and is typical for 218 219 badnaviruses (Teycheney et al. 2020). The ORF1 (nt 245-751, numbering according to the isolate FRIc1224A sequence) encodes a protein of 169 aa (19.8 kDa), the ORF2 (nt 751-1161) encodes a 220 221 137-aa protein (15 kDa), and the third ORF (nt 1,163-6,721) encodes a polyprotein of 1,853 aa (211.7 kDa) with five conserved protein domains (Figure 2): a viral movement protein (MP, cl03100), a zinc-222 binding motif (ZnF, pfam00098), a retroviral aspartyl protease domain (RVP, pfam00077), a reverse 223 224 transcriptase domain (RT, cd01647) and a ribonuclease H domain (RH, cl14782). The two "Cys" 225 motives (C-X₂-C-X₄-H-X₄-C, and C-X₂-C-X₁₁-C-X₂-C-X₄-C-X₂-C) usually found in the coat protein of 226 badnaviruses (Bath et al. 2016) were also detected in the ORF3-deduced protein, between aa positions 777-790 and 902-928. 227

Both isolates are closely related, with an overall 90.5% nt identity. The three indels observed between the two sequences are located in the intergenic region, the isolate ITumito39 ended up being one nucleotide longer. The three ORFs have the same sizes, are strictly colinear and the encoded proteins share respectively 95.2% (ORF1), 95.5% (ORF2) and 94.8% (ORF3) aa identity.

To characterize the phylogenetic relationships and taxonomic position of the chestnut badnavirus, a phylogenetic tree was reconstructed using an alignment of full genome nucleotide sequences of *Badnavirus* genus members, with the rice tungro bacilliform virus used as an outgroup (Figure 3). Both isolates cluster in group 3 defined by Wang et al. (2014), together with gooseberry

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236 vein banding virus (GVBV), rubus yellow net virus (RYNV), grapevine vein clearing virus (GVCV), birch leafroll-associated virus (BLRaV), wisteria badnavirus 1 (WBV1), and pagoda yellow mosaic-237 associated virus (PYMaV) (Figure 3). Nevertheless, they are clearly distant from all of these species, 238 defining a novel branch, supported by a 99% bootstrap value (Figure 3). Tree topology was similar 239 when using an alignment of representative badnaviral ORF3 protein sequences (Figure S1). To 240 241 confirm these analyses, pairwise comparisons of genome sequences showed that the isolate FRIc1224A has only weak identity levels with representative members of the genus Badnavirus, 242 comprised between 42.1% nt identity (sugarcane bacilliform IM virus; 42.5% for the isolate 243 ITumito39) and 50.9% (WBV1; 50.8% for the isolate ITumito39). The same tendency is observed 244 when considering the genome proteins. The ORF1-encoded protein shows only weak homology with 245 246 the corresponding proteins of WBV1 (27.8% aa identity) and PYMAV (26.1% aa identity), and the ORF2-encoded protein shares only 33.1% aa identity with the corresponding protein of the most 247 closely related virus, WBV1. The polyprotein encoded by ORF3 shares 49.5% aa identity with the 248 corresponding protein of the closest relative, PYMAV. Using the ORF3 region (RT and RH domain) 249 250 used for taxonomical discrimination in the family Caulimoviridae (Teycheney et al. 2020), the FRIc1224A isolate shows between 64% (with GVBV) and 68.4% (with BLRaV) nt identity (Table 1), 251 252 which is below the 80% nt identity value used as the species demarcation threshold in the family. Therefore, this virus represents a novel species in the family *Caulimoviridae*, for which we propose 253 254 the name *Chestnut mosaic virus* (ChMV). In the same taxonomically informative region, the isolates FRIc1224A and ITumito39 share 91.9% nt identity (97.8% aa identity), indicating that they belong to 255 the same viral species (Table 1). 256

Identification of ChMV in publicly available chestnut HTS data. The datamining of chestnut
 HTS data from various chestnut sources publicly available at GenBank [EST sequences, whole
 genome assembly, RNA-Seq and Genotyping-by-sequencing (GBS) reads available as Sequence Read

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260 Archives (SRA)] allowed the identification of ChMV in several of those datasets (Table S3). In particular, two complete genomes were obtained from datasets involving C. mollissima cv. Vanuxem 261 in the USA, one from the whole genome assembly (JRKL01079565) and the other by de novo 262 assembly of RNA-Seq data (SRX4015368) with 99.2% and 97.4% nt identity, respectively, with the 263 FRIc1224A isolate over the whole genome (Figure 4). In addition, partial ChMV genome assemblies 264 265 of >3 kbp could be obtained from a range of other datasets generated in the USA or in China from C. mollissima (Table S3), all of which showed significant relatedness with the FRIc1224A sequence 266 267 as shown by a phylogenetic tree reconstructed using nucleotide alignments of concatenated ChMV sequences retrieved from the various datasets (Figure 4). In addition, partial ChMV genomes could 268 also be reconstructed from two datasets obtained from *C. dentata* in the USA. Interestingly, one of 269 270 these two C. dentata isolate sequences shows closest relationship with ITumito39 sequence (Figure 4) with only 89.2% nt identity with the isolate FRIc1224A as compared to 93.9% nt identity with 271 ITumito39. The second isolate of C. dentata appears to be equally related to the FRIc1224A and 272 ITumito39 isolates, with 90.9% and 90.6% nt identity, respectively. 273

274 Incidence and genetic variability of ChMV in France and Italy. The incidence and genetic variability of ChMV were investigated by analyzing two genomic regions of ORF3, one 335-nt long 275 276 located in the MP domain amplified using primer pair Ch-Bad-1466F/Ch-Bad-1800R and the other 277 232-nt long in the RNase H domain and amplified with primer pair Ch-Bad-5860F/Ch-Bad-6109R 278 (Table S2, and Figure 2). The two primer pairs were designed to be able to detect both isolates FRIc1224A and ITumito39. In Italy, a total of 70 C. sativa cv Marrone samples were collected in the 279 same location, while in France, 95 chestnut accessions belonging to three different *Castanea* species 280 281 or hybrids were sampled in three plots (A, Port, E). Both symptomatic and asymptomatic samples 282 were collected, as well as some samples with atypical or dubious symptoms. Globally, ChMV was 283 frequent in the surveyed plots, with 57/70 (81.5%) infected C. sativa samples in Italy and 65/95 trees

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(68%) in France (Table 2). In the Italian orchard, half of the asymptomatic trees were found to be infected by ChMV, compared to 87% of the symptomatic ones (Table 2). None of the analyzed samples were found positive using a phytoplasma-specific PCR assay. In the French collection, the virus was detected in 100% (38/38) of the trees showing typical ChMD symptoms, and in 49% (23/47) of the asymptomatic trees, including two trees that were symptomless but showed strong symptoms on rootstock off-shoots (Figure S2). ChMV was also detected in four out of the 10 trees showing atypical/doubtful symptoms.

The genetic variability of ChMV was evaluated by analyzing the sequences of the two PCR 291 amplicons generated for the incidence survey. Considering the relative homogeneity of the origin 292 of the Italian samples, the number of samples included in this analysis was limited to 13 (four from 293 asymptomatic trees, and nine symptomatic ones) (Table S1). In total, the final dataset consisted in 294 53 isolates for which the sequence of the two genomic regions were available (49 from the incidence 295 survey and four from independent ChMD sources held in collection at CTIFL, see below). As 296 illustrated by the unrooted neighbor-joining tree reconstructed from the alignments of RT-RnaseH 297 298 domain nucleotide sequences (Figure S3A), ChMV diversity is structured into two clusters, defined by the geographical origin of the samples (Italy and France). The sequences determined from the 299 300 four independent French disease sources (FRIc1224A, T30218, LCA552, LCA584) belong to the same «French» cluster. Overall, the level of genetic diversity is very low in this genomic region, with an 301 302 average pairwise nt divergence (diversity) of 2.2% +/- 0.5%. This value is even lower when considering the intra-group diversity, as 0.2% +/- 0.1% within the French cluster and 0.1% +/- 0.1% 303 304 within the Italian ones. In contrast, the inter-group diversity reaches 6.3% +/- 1.5%, confirming the 305 existence of two geographical clusters. The same trends are observed when analyzing the genomic 306 region located in the MP domain (Figure S3B). The same geographical clustering could be observed, 307 with the exceptions of three French isolates that seem to be more closely related to the Italian

308	cluster (Figure S3B). Another French isolate, 20971-E remains isolated and does not fit in either
309	group. The average nt divergence in this region is slightly higher than in the RT-RNaseH region, (5.8%
310	+/- 0.7%), and the inter-group diversity reaches a value of 13.4% +/- 1.8%, as compared to the 6.3%
311	value for the other region.

312

DISCUSSION

Since the seminal work of Desvignes and collaborators in the 1990s, it has been acknowledged 313 that the agent responsible for ChMD is most likely a thermosensitive, graft-transmissible virus that 314 315 can be transmitted experimentally and probably naturally by the aphid Myzocallis castanicola 316 Desvignes, 1992; 1999a; 1999b; Desvignes and Lecoq, 1995; Desvignes and Cornaggia, 1996). Depending on the chestnut genotype, this infection can either be asymptomatic or can result in the 317 expression of severe and conspicuous ChMD symptoms. In chestnut orchards in the Marche region 318 (Italy), both young and mature plants were shown to be affected, significantly decreasing chestnut 319 320 production. Symptoms have also been observed in some Quercus species following experimental 321 graft inoculation. To date, however, the causal agent remains to be identified.

Here, by using HTS-based viral indexing, we were able to identify and characterize, in two independent ChMD sources, two isolates of the same novel virus. Phylogenetic and sequence analyses showed that this virus belongs to the genus *Badnavirus*, in the family *Caulimoviridae*, and could be considered as a new species, for which the name *Chestnut mosaic virus* (ChMV) is proposed. Interestingly, this new virus clusters with a group of badnaviruses that includes RYNV, GVBaV, and GVCV.

There is unambiguous evidence that ChMV as reported here is an episomal virus. It was detected in graft-inoculated indicators, and not in non-inoculated control plants of the same variety, demonstrating its graft-transmissibility, a property of episomal viruses. This line of evidence is further reinforced by the detection of ChMV in symptomatic, graft-inoculated indicator *Quercus* 332 plants and, again, not in the corresponding control plants. In parallel, the HTS detection of ChMV from DNAse-treated RNAs, the failure to detect ChMV in a range of the surveyed chestnut trees and 333 the sequence diversity identified in ChMV all rule out a scenario in which an endogenous ChMV 334 genome, integrated in the chestnut genome could be responsible for the HTS and PCR results 335 reported here. There was in fact no indication of ChMV in the chestnut genome assembly 336 337 (JRKL01079565) since no integration borders could be identified and a single contig, representing a complete unintegrated viral genome transcript, was identified. Integration of ChMV as an 338 339 endogenous viral element (EVE, Bhat et al. 2016) therefore does not appear to be a general genomic feature of chestnut. 340

According to the simplified hierarchical approach proposed by Fox (2020) for assessing causal 341 relationships in plant virology, ChMV appears as a good candidate, if not as the causative agent of 342 343 ChMD. There are several arguments and experimental evidence supporting this idea. Following HTS analyses, ChMV was the sole virus detected in the French source FRIc1224A, coming from a ChMD 344 source initially involving a C. sativa x C. crenata hybrid. It was also the sole virus detected in the 345 346 Italian ChMD source analyzed by HTS. Using molecular detection tests developed in this work, the virus was consistently found in other symptomatic accessions derived from the same diseased 347 348 source (LC1224H, a Q. rubra artificially inoculated and LC1224F, an indicator plant inoculated by aphid transmission; Figure 5). In addition, three other independent chestnut sources shown by 349 350 biological indexing on the 'Maraval' indicator to be affected by ChMD were found to be infected by ChMV (LCA552, LCA584 and T32018 in Figure 5). There is therefore a correlation between the 351 352 appearance of ChMD symptoms and the presence of ChMV in the graft-inoculated indicators, 353 supporting the hypothesis of a causal relationship between ChMV infection and ChMD. In total, five 354 independent ChMD sources collected between 1990 and 2018 in two countries (Italy and France)

were ChMV positive, satisfying the Bradford-Hill's experimental and consistency criteria (Bradford
Hill, 1965; Fox, 2020).

Preliminary studies indicate that ChMV is highly prevalent in the analyzed orchards in France 357 and Italy, confirming the earlier results of Desvignes (1999a). In parallel, the identification of ChMV 358 sequences in publicly available HTS data provides a strong indication of the presence of ChMV in C. 359 360 mollissima in the USA and in China and in C. dentata in the USA. In the surveys, ChMV was not systematically associated with symptomatic infections, although its frequency was systematically 361 362 higher in symptomatic plants. This result was expected since previous grafting experiments had demonstrated that not all chestnut varieties/species are susceptible to ChMD and develop 363 symptomatic infections (Desvignes and Lecoq, 1995; Desvignes, 1992; 1999b). Biological indexing 364 on the susceptible 'Maraval' indicator has in particular identified latent ChMV infections in many 365 366 symptomless C. sativa varieties or C. sativa x C crenata hybrids (Desvignes and Lecoq, 1995; Desvignes, 1992; 1999b). On the other hand, all surveyed symptomatic plants in France were found 367 to harbor the virus, while it was detected in 52/60 (87%) of tested symptomatic Italian trees. The 368 369 failure to detect ChMV in eight symptomatic Italian trees might reflect sequence variability and an incomplete inclusiveness of the PCR primers used or low or uneven virus accumulation. Indeed, 370 371 using biological indexing, Desvignes et al. have previously found an uneven distribution of the ChMD agent in infected trees leading to a failure to detect it in parts of some infected trees (Desvignes and 372 373 Lecoq, 1995; Desvignes et al. 1999b).

Taken together, and even though Koch's postulates were not fully verified, the experiments reported here make a very strong case for a role of ChMV as the causal agent of the chestnut mosaic disease. The low ChMV diversity observed in France and Italy are consistent with the scenario of its recent introduction in Europe (Desvignes and Lecoq, 1995), while the genetic separation of the Italian and French clusters is suggestive of separate introduction events. These results and the

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associated development of molecular tools for the detection of ChMV will help speed up the selection of virus-free mother plants and mitigate the virus spread in new chestnut orchards and layerings. However, many questions remain regarding the variability of symptom intensity in relationship to cultivar susceptibility, ChMV-induced graft incompatibility, the impact of pedoclimatic conditions and of synergic and competitive interferences with other chestnut pathogens, and silvicultural management.

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 Bruijn graphs. Genome Res. 18:821-829.

489 TABLE 1. Percentage of identity between the ORF3 region encoding the Reverse Transcriptase -

490 Rnase H^a of chestnut mosaic virus (ChMV) isolate FRIc1224A and the corresponding genomic regions

491 of the isolate ITumito39 and of the most closely related members of the genus *Badnavirus*

Virus ^b	Nucleotide identity (%)	Amino acid identity (%)
ChMV ITumito39	91.9%	97.8%
RYNV	65.1%	71.6%
GVBV	64%	68.9%
GVCV	66.8%	72.6%
BLRaV	68.4 %	72.6%
WBV1	68.2%	71.6%
PYMaV	67.7%	71.9%

^a This region is the one typical used for taxonomic discrimination in the family *Caulimoviridae*(Teycheney et al. 2020)

^b Acronyms used: RYNV, rubus yellow net virus; GVBV, gooseberry vein banding virus; GVCV,
 grapevine vein-clearing virus; BLRaV, birch leafroll-associated virus; WBV1, wisteria badnavirus 1;

496 PYMaV, pagoda yellow mosaic-associated virus.

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- 499 **TABLE 2.** Number and percentage of chestnut mosaic virus-infected plants regarding the plot, the
- 500 *Castanea* species sampled, and the symptomatology

Origin of the	Infected/total	Infected/asympto	Infected/sympto	Infected/plants
sampled plants	plants (%)	matic plants (%)	matic plants (%)	with atypical
				symptoms (%)
France (overall)	65/95 (68.4%)	23/47 (48.9%)	38/38 (100%)	4/10 (40%)
Plot A	13/23 (56.5%)	6/15 (40%)	6/6 (100%)	1/2 (50%)
Plot E	48/66 (72.7%)	15/28(53.6%)	30/30 (100%)	3/8 (37.5%)
Plot Port	4/6 (66.6%)	2/4 (50%)	2/2 (100%)	na ^b
C. crenata	5/6 (83%)	1/1 (100%)	4/4 (100%)	0/1
C. sativa	30/43 (70%)	11/20 (55%)	18/18 (100%)	1/5 (20%)
C. mollissima	13/14 (93%)	1/2 (50%)	10/10 (100%)	2/2 (100%)
Hybrid ^a	17/32 (53.1%)	10/24 (41.6%)	6/6 (100%)	1/2 (50%)
Italy Castanea	57/70 (81.5%)	5/10 (50%)	52/60 (87%)	na ^b
sativa				

- ^a Interspecific hybrids between *C. crenata, C. mollisima* and *C. sativa*
- ^b not applicable
- 503

504 CAPTIONS FOR FIGURES

Fig. 1. Symptoms of chestnut mosaic disease on various hosts. (A) Isolate LC1224H: Red oak (*Quercus rubra*) graft-inoculated with a diseased source; (B) Isolate FRIc1224A: 'Maraval' Ca 74 graft-inoculated with LC1224H; (C) Isolate ITumito39: symptomatic leaves from cv Marrone grafted onto *Castanea sativa*; (D) non-inoculated *Q. rubra*; (E) non-inoculated 'Maraval' Ca 74; (F) Asymptomatic leaves from cv Marrone grafted onto *C. sativa*.

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Fig. 2: Schematic representation of the genomic organization of the chestnut mosaic virus. The tRNA binding site is indicated and defines the position 1 on the genome. The three open reading frames (ORFs) are shown as grey arrows, as well as their position in parentheses. Five conserved motifs are identified in the ORF3 polyprotein: MP, Viral movement protein (pfam01107); ZnF, Zinc finger (pfam00098); RVP, Retroviral aspartyl protease (pfam00077); RT, Reverse transcriptase (cd01647); RH, Ribonuclease H (cd09274)

517

Fig. 3. Phylogenetic tree reconstructed using the complete genome sequences of badnavirus members. Virus names as well as GenBank accession numbers are indicated. The tree was reconstructed using the neighbor-joining method, and randomized bootstrapping was performed to evaluate the statistical significance of branches (1,000 replicates). Bootstrap values above 70% are shown. The scale bar represents 5% nucleotide divergence between sequences. The groups as defined by Wang et al. (2014) are indicated. Chestnut mosaic virus isolates determined in this work are indicated by black triangles. Rice tungro bacilliform virus was used as outgroup.

525

Fig. 4. Unrooted neighbor-joining phylogenetic tree reconstructed from the alignment of concatenated nucleotide sequences related to chestnut mosaic virus detected by datamining of publicly available transcriptomic chestnut data. Randomized bootstrapping was performed to evaluate the statistical significance of branches (1,000 replicates). Bootstrap values above 70% are shown. The scale bar represents 10% nucleotide divergence between sequences.

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Fig. 5. Detection of chestnut mosaic virus in various samples by PCR using primers pairs Ch-Bad1466F/1800R (A) and Ch-Bad5860F/6109R (B). Lane 1: LC1224F; Lane 2: LC1224H; Lane 3: FRIc1224A; Lane 4: T32018; Lane 5: LCA552; Lane 6: LCA584; Lane 7: 'Maraval' Ca 74 non-inoculated plant; Lane 8: *Quercus rubra* non-inoculated plant; Lane 9: no template; L: molecular weight marker. Horizontal bars on the left of the figure indicate the size of the amplification products. The isolates are listed in Table S1.

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539 e-EXTRA FIGURE CAPTIONS AND TABLE TITLES

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Supplementary Fig. S1. Phylogenetic tree reconstructed using the ORF3-deduced amino acid
sequences of badnavirus members. Virus names as well as GenBank accession numbers are
indicated. The tree was reconstructed using the neighbor-joining method, and randomized
bootstrapping was performed to evaluate the statistical significance of branches (1,000 replicates).
Bootstrap values above 70% are shown. The scale bar represents 10% amino acid divergence
between sequences. Chestnut mosaic virus isolates determined in this work are indicated by black
triangles. Rice tungro bacilliform virus was used as outgroup.

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549 Supplementary Fig. S2. Leaves of the sample 20893. (A) rootstock leaves (*Castanea sativa*); (B) 550 cultivar leaves

551

552 Supplemental Fig. S3. Unrooted neighbor-joining phylogenetic trees reconstructed from the 553 alignment of nucleotide sequences of the PCR fragments targeted partial RT-RnaseH domain (A) 554 and partial MP domain (B) obtained from a range of chestnut mosaic virus isolates (listed in Table 555 S1). Randomized bootstrapping was performed to evaluate the statistical significance of branches 556 (1,000 replicates). Bootstrap values above 70% are shown. The scale bar represents 5% (A) or 10% 557 (B) nucleotide divergence between sequences.

558

Supplemental Table S1. List of chestnut samples used in the present study together with relevant
 ChMV accession numbers

561

562 Supplemental Table S2. Primers used for genome circularity confirmation and for molecular 563 detection of chestnut mosaic virus by PCR

564

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- 565 Supplemental Table S3. Datamining of publicly available chestnut HTS data for chestnut mosaic
- 566 virus sequences





Fig. 2: Schematic representation of the genomic organization of the chestnut mosaic virus. The tRNA binding site is indicated and defines the position 1 on the genome. The three open reading frames (ORFs) are shown as grey arrows, as well as their position in parentheses. Five conserved motifs are identified in the ORF3 polyprotein: MP, Viral movement protein (pfam01107); ZnF, Zinc finger (pfam00098); RVP, Retroviral aspartyl protease (pfam00077); RT, Reverse transcriptase (cd01647); RH, Ribonuclease H (cd09274)

338x190mm (96 x 96 DPI)



Fig. 3. Phylogenetic tree reconstructed using the complete genome sequences of badnavirus members. Virus names as well as GenBank accession numbers are indicated. The tree was reconstructed using the neighbor-joining method, and randomized bootstrapping was performed to evaluate the statistical significance of branches (1,000 replicates). Bootstrap values above 70% are shown. The scale bar represents 5% nucleotide divergence between sequences. The groups as defined by Wang et al. (2014) are indicated. Chestnut mosaic virus isolates determined in this work are indicated by black triangles. Rice tungro bacilliform virus was used as outgroup.

190x338mm (96 x 96 DPI)





Fig. 5. Detection of chestnut mosaic virus in various samples by PCR using primers pairs Ch-Bad1466F/1800R (A) and Ch-Bad5860F/6109R (B). Lane 1: LC1224F; Lane 2: LC1224H; Lane 3: FRIc1224A; Lane 4: T32018; Lane 5: LCA552; Lane 6: LCA584; Lane 7: 'Maraval' Ca 74 non-inoculated plant; Lane 8: Quercus rubra non-inoculated plant; Lane 9: no template; L: molecular weight marker. Horizontal bars on the left of the figure indicate the size of the amplification products. The isolates are listed in Table S1.

338x190mm (96 x 96 DPI)





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Fig. S2



Fig. S3A



Fig. S3B



1 Supplementary Table S1. List of chestnut samples used in the present study together with relevant ChMV

2 accession numbers

Isolate ID ^a	Chestnut Accession ID	Chestnut Accession name	Species ^b	Rootstock	Country	Plot ^{.c}	Accession numbers ^d	Use ^e
20885	Ca04x Ca03	-	C. crenata	Not graft	France	А	MT339547; MT339503	Incidence
20887	Ca 715	'Merle'	C. sativa	G1. Ca15	France	A	MT339548; MT339504	Incidence
20890	Ca 663	'Trigueira'	C. sativa	unknown	France	Port	MT339549; MT339505	Incidence
20891	Ca 664	'Longal Special'	C. sativa	Ca 74	France	Port	MT339550; MT339506	Incidence
<u>20895</u>	Ca 43	'Vignols'	C. sativa x C. crenata	Ca 116	France	A	MT339551; MT339507	Incidence
20899	Ca 599	'Ibuki'	C. crenata	G1.Ca 02	France	A	MT339552; MT339508	Incidence
20901	Ca 598	'Rihei'	C. crenata x C. mollissima	G1. Ca102	France	A	MT339553; MT339509	Incidence
20902	Ca 75	'Fertil'	C. mollissima	Ca 07	France	A	MT339554; MT339510	Incidence
20904	Tree- A68Ks ⁱ	-	C. mollissima	Not graft	France	A	MT339555; MT339511	Incidence
20907	Ca 118	'Marlhac'	C. sativa x C. crenata	Not graft	France	A	MT339556; MT339512	Incidence
20908	Ca 124	'Maridonne'	C. sativa x C. crenata	G1.Ca116	France	A	MT339557; MT339513	Incidence
20909	Ca 125	'Bouche de Bétizac'	C. sativa x C. crenata	Ca 74	France	A	MT339558; MT339514	Incidence
20910	Ca 860	-	hybrid SC	unknown	France	E	MT339559; MT339515	Incidence
20911	Ca 860	-	hybrid SC	unknown	France	E	MT339560; MT339516	Incidence
20913	Ca 844 ⁱ	-	C. mollissima	Not graft	France	E	MT339561; MT339517	Incidence
20917	Ca 837 ^h	-	C. mollissima	Not graft	France	E	MT339562; MT339518	Incidence
20923	Ca 846 ^h	-	C. mollissima	Not graft	France	E	MT339563; MT339519	Incidence
20925	Ca 741	'Dauphine'	C. sativa	Ca 07	France	E	MT339564; MT339520	Incidence
20926	Ca 665	'Longal'	C. sativa	Ca 07	France	E	MT339565; MT339521	Incidence
20930	Ca 860	-	hybrid SC	unknown	France	E	MT339566; MT339522	Incidence
20932	Ca 564	'Ipharra 16'	C. crenata	Ca 07	France	E	MT339567; MT339523	Incidence
<u>20935</u>	Ca 576	'Sardonne'	C. sativa	Ca 07	France	E	MT339568; MT339524	Incidence
20937	Ca 138	'Marron de Redon'	C. sativa	Ca 07	France	E	MT339569; MT339525	Incidence
20943	unknown	-	C. sativa	Ca 74	France	E	MT339570; MT339526	Incidence
20944	Ca 520	'Montagne'	C. sativa	Ca 07	France	E	MT339571; MT339527	Incidence
20945	Ca 106	'Marron Comballe'	C. sativa	Ca 07	France	E	MT339572; MT339528	Incidence

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20947	Ca 111	'Marron de Lyon'	C. sativa	Ca 07	France	E	MT339573; MT339529	Incidence
20951	Ca 126	-	C. sativa x C. crenata	Ca 07	France	E	MT339574; MT339530	Incidence
20956	Ca 105	'Sardonne'	C. sativa	Ca 07	France	E	MT339575; MT339531	Incidence
20959	Ca 03	-	C. crenata	Ca 74	France	E	MT339576; MT339532	Incidence
20960	Ca 127	-	(C. crenata x C. sativa) x C. mollisima	Ca 07	France	E	MT339577; MT339533	Incidence
20961	Ca 127	-	(C. crenata x C. sativa) x C. mollisima	Ca 07	France	E	MT339578; MT339534	Incidence
20964	Ca 112	'Bournette'	C. crenata x C. sativa	Ca 07	France	E	MT339579; MT339535	Incidence
20965	Ca 501	-	C. sativa	Ca 07	France	E	MT339580; MT339536	Incidence
20966	Ca 501	-	C. sativa	Ca 07	France	E	MT339581; MT339537	Incidence
20967	Ca 665	'Longal'	C. sativa	Ca 07	France	E	MT339582; MT339538	Incidence
20968	Ca 48	'Précoce Migoule'	(C. crenata x C. sativa)	Ca 07	France	E	MT339583; MT339539	Incidence
20971	Ca 151	'Bouche Rouge'	C. sativa	Ca 07	France	E	MT339584; MT339540	Incidence
20975	unknown	-	C. sativa x C. crenata	Ca 07	France	E	MT339585; MT339541	Incidence
20889	unknown	-	C. sativa	unknown	France	Port	na	Incidence
20893	Ca 106	'Marron Comballe'	C. sativa	unknown	France	Port	na	Incidence
20898	Ca 105	'Sardonne'	C. sativa	G1. Ca394	France	А	na	Incidence
20900	Ca 501	-	C. sativa	G1.Ca486	France	А	na	Incidence
20903	Ca744	'Quing Zha'	C. mollissima	G1.Moll	France	А	na	Incidence
20912	Ca 845 ^h	-	C. mollissima	Not graft	France	E	na	Incidence
20914	Ca 843 ^h	-	C. mollissima	Not graft	France	E	na	Incidence
20915	Ca 842 ^h	-	C. mollissima	Not graft	France	E	na	Incidence
<u>20918</u>	Ca 838 ^h	-	C. mollissima	Not graft	France	E	na	Incidence
<u>20919</u>	Ca 839 "	-	C. mollissima	Not graft	France	E	na	Incidence
20920	Ca 840 "	-	C. mollissima	Not graft	France	E	na	Incidence
20921	Ca 841 "	-	C. mollissima	Not graft	France	E	na	Incidence
20922	Ca 74	'Maraval'	C. crenata x C. sativa	Not graft	France	E	na	Incidence
20924	Ca 825	Besse'	C. sativa	Ca 07	France	E	na	Incidence
20940	Ca 393	Chevanceaux'	C. sativa	Ca 07	France	E	na	Incidence
20941	Ca 460	-	(C. crenata x C. sativa) x (C. crenata x C. sativa)	Ca 74	France	E	na	Incidence
29942	Ca 511	'Marrone di Chiusa Pesio'	C. sativa	Ca 74	France	E	na	Incidence
20948	Ca 116	-	C. sativa x C. crenata	Ca 74	France	E	na	Incidence
20949	Ca 135	'Précoce des Vans'	C. sativa	Ca 74	France	E	na	Incidence
20052	Ca 521	-	C. crenata	Ca 07	France	E	na	Incidence

20955	Ca 399	'Comballe'	C. sativa	Ca 07	France	E	na	Incidence
20962	Ca 639	-	C. sativa	Ca 07	France	E	na	Incidence
20963	Ca 730	'Sauvage Marron'	C. sativa	Ca 07	France	E	na	Incidence
20969	Ca 381	-	C. sativa	Ca 07	France	E	na	Incidence
20970	Ca 381	-	C. sativa	Ca 07	France	Е	na	Incidence
20972	Ca 151	'Bouche Rouge'	C. sativa	Ca 07	France	E	na	Incidence
FRIC1224A	Ca 74	'Maraval'	C. crenata x C. sativa	na	France	Ctifl	MT269853 (complete genome)	HTS
LC1224H	na	na	Q. rubra	na	France	Ctifl	MT339586; MT339542	CRA
T32018	Ca 74	Maraval	C. crenata x C. sativa	na	France	Ctifl	MT339587; MT339543	CRA
LCA552	Ca 74	Maraval	C. sativa	na	France	Ctifl	MT339588; MT339544	CRA
LCA584	Ca 74	Maraval	C. sativa	na	France	Ctifl	MT339589; MT339545	CRA
LC1224F	Ca 74	Maraval	C. crenata x C. sativa	na	France	Ctifl	MT339590; MT339546	CRA
ITUmito39	unknown	Marrone	C. sativa	na	Italy	I	MT261366 (complete genome)	HTS
ITCh2	unknown	Marrone	C. sativa	na	Italy	I	MT270674; MT270683	Incidence
ITCh3	unknown	Marrone	C. sativa	na	Italy	I	MT270667 ^f	Incidence
ITCh4	unknown	Marrone	C. sativa	na	Italy	I	MT270668; MT270684	Incidence
ITCh8	unknown	Marrone	C. sativa	na	Italy	I	MT270685 ^g	Incidence
ITCh10	unknown	Marrone	C. sativa	na	Italy	I	MT270669; MT270678	Incidence
ITCh11	unknown	Marrone	C. sativa	na	Italy	I	MT270664; MT270679	Incidence
ITCh12	unknown	Marrone	C. sativa	na	Italy	I	MT270665; MT270680	Incidence
ITCh14	unknown	Marrone	C. sativa	na	Italy	I	MT270666; MT270681	Incidence
ITCh20	unknown	Marrone	C. sativa	na	Italy	I	MT270670; MT270675	Incidence
ITChAs3	unknown	Marrone	C. sativa	na	Italy	I	MT270671; MT270676	Incidence
ITChAs4	unknown	Marrone	C. sativa	na	Italy	I	MT270672 ^f	Incidence
ITChAs5	unknown	Marrone	C. sativa	na	Italy	I	MT270677 ^g	Incidence
ITChAs8	unknown	Marrone	C. sativa	na	Italy	I	MT270673; MT270682	Incidence
ITChAs9	unknown	Marrone	C. sativa	na	Italy	I	na	Incidence
ITCh1	unknown	Marrone	C. sativa	na	Italy	I	na	Incidence
ITCh5	unknown	Marrone	C. sativa	na	Italy	I	na	Incidence
ITCh6	unknown	Marrone	C. sativa	na	Italy	I	na	Incidence
ITCh7	unknown	Marrone	C. sativa	na	Italy	I	na	Incidence
ITCh9	unknown	Marrone	C. sativa	na	Italy	I	na	Incidence
ITCh13	unknown	Marrone	C. sativa	na	Italy	I	na	Incidence
ITCh14	unknown	Marrone	C. sativa	na	Italy	I	na	Incidence
ITCh15	unknown	Marrone	C. sativa	na	Italy	I	na	Incidence
ITCh20	unknown	Marrone	C. sativa	na	Italy	I	na	Incidence
ITCh21	unknown	Marrone	C. sativa	na	Italy	1	na	Incidence

ITCh22	unknown	Marrone	C. sativa	na	Italy	I	na	Incidence
ITCh23	unknown	Marrone	C. sativa	na	Italy	I	na	Incidence
ITCh24	unknown	Marrone	C. sativa	na	Italy	I	na	Incidence
ITCh25	unknown	Marrone	C. sativa	na	Italy	I	na	Incidence
ITCh27	unknown	Marrone	C. sativa	na	Italy	I	na	Incidence
ITCh28	unknown	Marrone	C. sativa	na	Italy	I	na	Incidence
ITCh29	unknown	Marrone	C. sativa	na	Italy	I	na	Incidence
ITCh30	unknown	Marrone	C. sativa	na	Italy	I	na	Incidence
ITCh31	unknown	Marrone	C. sativa	na	Italy	I	na	Incidence
ITCh32	unknown	Marrone	C. sativa	na	Italy	I	na	Incidence
ITCh33	unknown	Marrone	C. sativa	na	Italy	I	na	Incidence
ITCh35	unknown	Marrone	C. sativa	na	Italy	I	na	Incidence
ITCh36	unknown	Marrone	C. sativa	na	Italy	I	na	Incidence
ITCh37	unknown	Marrone	C. sativa	na	Italy	I	na	Incidence
ITCh38	unknown	Marrone	C. sativa	na	Italy	I	na	Incidence
ITCh39	unknown	Marrone	C. sativa	na	Italy	I	na	Incidence
ITCh41	unknown	Marrone	C. sativa	na	Italy	I	na	Incidence
ITCh42	unknown	Marrone	C. sativa	na	Italy	I	na	Incidence
ITCh43	unknown	Marrone	C. sativa	na	Italy	I	na	Incidence
ITCh44	unknown	Marrone	C. sativa	na	Italy	I	na	Incidence
ITCh46	unknown	Marrone	C. sativa	na	Italy	I	na	Incidence
ITCh47	unknown	Marrone	C. sativa	na	Italy	I	na	Incidence
ITCh48	unknown	Marrone	C. sativa	na	Italy	I	na	Incidence
ITCh49	unknown	Marrone	C. sativa	na	Italy	I	na	Incidence
ITCh50	unknown	Marrone	C. sativa	na	Italy	I	na	Incidence
ITCh51	unknown	Marrone	C. sativa	na	Italy	I	na	Incidence
ITCh52	unknown	Marrone	C. sativa	na	Italy	I	na	Incidence
ITCh54	unknown	Marrone	C. sativa	na	Italy	I	na	Incidence
ITCh55	unknown	Marrone	C. sativa	na	Italy	I	na	Incidence
ITCh56	unknown	Marrone	C. sativa	na	Italy	I	na	Incidence
ITCh58	unknown	Marrone	C. sativa	na	Italy	I	na	Incidence
ITCh59	unknown	Marrone	C. sativa	na	Italy	I	na	Incidence
ITCh60	unknown	Marrone	C. sativa	na	Italy	I	na	Incidence

^a Isolates from asymptomatic trees are indicated in bold; isolates with doubtful symptoms are underlined; isolates from asymptomatic cultivars with typical symptoms visible on rootstock regrowths are indicated in bold italic.

5 ^b; hybrid SC : hybrid between *C. sativa* and *C. crenata*, regardless the knowledge about which are the female and male parents.

6 ^c: Five plots have been sampled (A, Port, E, I, Ctifl)

^d: Accession numbers are relative to the sequences obtained with both PCR detection assays (Ch-Bad-1466F/Ch-Bad-1800R and
 Ch-Bad-5860F/Ch-Bad-6109R)

9 e: Isolates were included either in the HTS analysis, or in the incidence analysis, or in the causal relationship analysis (CRA)

10 ^f: Only the fragment amplified with Ch-Bad-1466F/Ch-Bad-1800R was sequenced

11 ^g: Only the fragment amplified with Ch-Bad-5860F/Ch-Bad-6109R was sequenced

12 ^h 'Mengshankui' (*C. mollissima* cultivar) seedling

13 ⁱ C. mollissima seedling

14 15

3 4

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Supplemental TABLE S2. Primers used for genome circularity confirmation and for molecular	
detection of chestnut mosaic virus by PCR	

Primer	Nucleotide	Genome	Amplicon size	Isolates detected
	sequence (5' to 3')	coordinates		
Ch-Bad-6976F	CCCGAGCCATTTAC	6,976-6,999	436 nt	FRIc1224A
	ACTTCACAAC			
Ch-Bad-252R	TCACTCATCTACCTC	252-230		
	ACACGCTC			
Ch-Bad-6481F	GAAATTGAATTGGA	6,481-6,501	1,007 nt	ITumito39
	AGGAAGA			
Ch-Bad-325R	TCAGATCAGCAAAC	344-325-		
	TCGAAC			
Ch-Bad-1466F	TATCAGCACTACAG	1,466-1,488	335 nt	
	TGAACAACC			nolvvalent
Ch-Bad-1800R	GTCATGACGCAAAC	1,800-1,781		poryvalent
	TTGGAA			
Ch-Bad-5860F	AGTATGTAAATGG	5,857-5,878	232 nt	
	GCACCGTTC			nolyvalent
Ch-Bad-6109R	GTTGATCCATCGCA	6,109-6,090		poryvarent
	CTCTTG			

1 **Supplemental TABLE S3.** Datamining of publicly available chestnut HTS data for chestnut mosaic

2 virus sequences

Type of	Dataset	Castanea	Country	Length of	% mapped	% nt identity	
dataª		species/cultivar		assembled	reads ^b	with FRIc1224A	
				sequence			
				(nt)			
EST	GO917001	C. mollissima, BX3,	USA	436	na	99%	
		clone Vanuxum					
WGA	JRKL01079565	C. mollissima, cv	USA	7,164 (full	na	99%	
		Vanuxem		lenght)			
RNA-Seq	SRX4015368	C. mollissima	USA	7,161 (full	0.004%	97.4%	
				lenght)			
RNA-Seq	SRX001805	C. mollissima, cv	USA	5,889	0.04%	95-100%	
		Vanuxem		(scaffold)			
GBS	SRX5144434	C. mollissima	China	5,933	0.06%	98.7% (average)	
				(scaffold)			
GSB	SRX5144449	C. mollissima,	China	6,396	0.04%	$0.7 \Gamma_{0}^{0}$	
		clone Vanuxem		(scaffold)		97.5% (average)	
GBS	SRX5145051	C. mollissima, cv	China	5,873	0.02%	0.0% (average)	
		Vanuxem		(scaffold)		99% (average)	
GBS	SRX5145050	C. mollissima	China	6,680	0.01%		
				(scaffold)		99% (average)	
GBS	SRX51445044	C. mollissima	China	6,183	0.01%	97.4% (avorago)	
				(scaffold)		97.4% (average)	
GBS	SRX51444621	C. mollissima	China	5,875	0.02%	$0E E^{(2)}$	
				(scaffold)		95.6% (average)	
GBS	SRX51444448	C. mollissima	China	2,948	0.04%	00% (average)	
				(scaffold)		22% (average)	
GBS	SRX51444446	C. mollissima	China	5,873	0.04%	90% (average)	
				(scaffold)		99% (average)	

GBS	SRX5825095	C. dentata	USA	1,857	0.0036%	20,20 (every set)	
				(scaffold)		89.2% (average)	
GBS	SRX5145077	C. dentata	USA	6,068	0.01%	90.9% (average)	
				(scaffold)			

3 ^a EST: expressed sequence tags; WGA: whole genome assembly; GBS: genotyping by sequencing

4 ^b Reads mapped to the genomic sequence of chestnut mosaic virus (French isolate)

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