Role of fresh dead wood in the epidemiology and the biological control of the chestnut blight fungus

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Abstract

The invasive fungus Cryphonectria parasitica, the causal agent of chestnut blight, is able to survive and sporulate on the bark of fresh dead Castanea sativa wood for at least two years. Here, we experimentally investigated the role of fresh dead wood in the epidemiology of chestnut blight, specifically in the spread of the hyperparasitic virus CHV1, which acts as biocontrol agent of C. parasitica. A total of 152 artificially initiated, virulent bark cankers in four chestnut stands were treated with virus-infected asexual spores originating either from sporulating dead wood or from a spore suspension. Molecular markers for both the virus and the fungal carrier were used to examine the spread of the applied biocontrol virus. Fourteen
months after treatment, 42-76% of the conidial spray-treated cankers and 50-60% of the cankers exposed to a sporulating dead stem had been virus-infected by the applied hypovirulent conidia in all four study sites. Virus infection reduced canker expansion and promoted canker healing (callusing). Thus, fresh chestnut dead wood may play an important role in supporting the successful spread of natural hypovirulence in chestnut forests. Further, combined with the application of virus-infected conidial suspensions, it may help promote the establishment of artificially released hypoviruses in chestnut stands to control chestnut blight.

Introduction

The ascomycete fungus *Cryphonectria parasitica* (Murr) Barr., the causal agent of chestnut blight, is a necrotrophic pathogen causing perennial bark lesions (cankers) on the stem and branches of susceptible host trees, particularly species of the genus *Castanea* (Rigling and Prospero 2018). Parts of the tree distal to the infection point may wilt and die. Originating from eastern Asia, *C. parasitica* was accidentally introduced into Europe in the 1930s, where it became invasive and affected the European chestnut (*Castanea sativa* Mill.). Following an initially high tree mortality, beginning from the 1950s many chestnut stands started to recover because of the spontaneous spread of a hyperparasitic mycovirus (Heiniger and Rigling 1994).

*Cryphonectria hypovirus* 1 (CHV1) is a RNA virus located in the cytoplasm of *C. parasitica* (Shapira et al. 1991). Its presence reduces virulence and sporulation capacity of the infected fungal strain, causing hypovirulence (Rigling and Prospero 2018). Cankers induced by virus-infected *C. parasitica* strains typically stop expanding and do not kill the affected trees. To date, natural hypovirulence successfully controls chestnut blight in many European regions, but there are certain areas (e.g. in parts of Southern and Northern Europe) where this is still not the case.
CHV1 does not occur outside its fungal host and is transmitted either horizontally or vertically (Rigling and Prospero 2018). Horizontal transmission by cytoplasmic exchange can occur after hyphal anastomosis between *C. parasitica* strains belonging to the same vegetative compatibility (vc) type. Vertical transmission enables spread of the virus by asexual spores (conidia). In fact, a variable proportion of conidia produced by virus-infected *C. parasitica* strains is itself virus-infected (e.g. Peever et al. 2000). On the contrary, sexual ascospores never carry the virus (Carbone et al. 2004).

*Cryphonectria parasitica* may sporulate on the surface of the bark cankers on living trees, in particular on actively growing, virus-free cankers (Rigling and Prospero 2018). Sporulation on non-expanding, virus-infected cankers is rare or absent, questioning their role in the spread of CHV1 (Peever et al. 2000; Prospero et al. 2006). Prospero et al. (2006) showed that both virus-free and virus-infected strains were able to survive for more than one year in preexisting cankers on stacked dead wood stems. Moreover, *C. parasitica* stromata abundantly developed on the surface of former cankers and adjacent bark areas. Sporulation was also observed on healthy stems that were cut and stacked indicating that these were purely saprophytically colonized by *C. parasitica* (Prospero et al., 2006).

The importance of the saprotrophic activity of *C. parasitica* (Figure 1) for the epidemiology of hypovirulence is still poorly understood. In this study, we conducted a field experiment to determine if virus-infected spores produced on dead chestnut wood can transmit the virus to bark cankers on living trees. To simulate sporulating chestnut wood, we inoculated freshly cut stems in the laboratory with a virus-infected *C. parasitica* strain and incubated them until complete colonization and stromata production was observed. In addition, we produced aqueous solutions of virus-infected conidia. Both the sporulating chestnut stems and the conidial suspensions were applied to virus-free cankers, and the spread of both the released virus strain as well as the fungal carrier strain was followed using
molecular markers. So far, only either the viral or the fungal haplotype spread was studied in
field experiments and not both together. Specifically, we determined if the virus-infected
conidia are able to infect preexisting virulent cankers on living trees, thereby reducing canker
growth and activity. Finally, we tested if the spread of the viral strains in vivo can be
forecasted by assessing the biocontrol potential of the fungal strain in vitro.

Materials and Methods

Chestnut stands

The experiment was conducted in four pure chestnut (C. sativa) coppice stands in
Switzerland (Table 1). Two sites were located in the Chablais region, specifically in Bex-
Montet (MON) and Bex-Creux Boyon (CRE), whereas the other two were situated in the La
Côte region, one in Vinzel (VIN) and one in Villars-sous-Yens (VSY). In Chablais, C.
parasitica genotype CpMG30 (vc type EU-1) is dominant, and in La Côte genotype CpMG15
(vc type EU-2) (Prospero and Rigling 2012). In all four stands, hypovirulence was artificially
introduced by treating chestnut blight cankers with virus-infected C. parasitica strains (Table
1). Natural hypovirulence only occurs in VSY, where the virus CHV1-M4740 has been found
since 2005 (Prospero and Rigling 2016).

Inoculation of artificial cankers

In June 2014, 50 cankers were artificially initiated in each of the sites MON, VIN and VSY,
and 37 cankers in CRE where fewer chestnut trees were available, resulting in a total of 187
cankers. To induce these cankers, a small agar plug taken from the margin of a growing
virus-free C. parasitica culture was placed into a hole made in the bark of a healthy chestnut
sprout using a cork borer (diameter 5 mm). Before each hole was made, the bark was
disinfecte by wiping with 70% ethanol. For inoculation, an isolate of the most common vc
type and microsatellite genotype of the region was used (Prospero and Rigling 2012; Chablais: CpMG30 of EU-1; La Côte: CpMG15 of EU-2). After inoculation, the wounds were sealed with tape. About three months after canker inoculation, the tape was removed and the cankers were treated as stated below.

## Treatments

At each experimental site, 20 cankers were treated with sporulating dead wood stems previously inoculated with a virus-infected *C. parasitica* strain, 20 cankers were sprayed with a suspension of virus-infected conidia, and 10 cankers were left untreated as a negative control (16, 16 and 5 cankers, respectively, in CRE).

**Dead Wood (DW) treatment.** For dead wood production, healthy *C. sativa* stems (6–8 cm in diameter) from a coppice stand in southern Switzerland were cut into 30 cm-long segments and autoclaved for 15 min at 121°C. Thereafter, segments were sealed at both ends with melted paraffin wax. Small agar disks (5 mm diameter) from a growing virus-infected *C. parasitica* culture were inserted into 15 holes made in the bark of the dead wood stems with a cork borer and sealed with tape. The stems for Chablais were inoculated with the *C. parasitica* haplotype CpMG8 (EU-1; fungal carrier M1709) infected by the virus CHV1-M3623, and those for La Côte with CpMG47 (EU-2; fungal carrier M4000) containing the same virus (Table 2). This virus differs in its sequence from the haplotypes used previously for treatments in the stands (Table S1). The inoculated dead wood stems were placed onto plastic supports in boxes (57 x 37 x 13 cm) filled with 4 liters demineralized water. The boxes were sealed with a transparent plastic sheet and incubated for six weeks at temperatures between 15 and 25°C. After this period, the inoculated *C. parasitica* haplotype had completely colonized the stems and produced numerous asexual fruiting bodies containing virus-infected spores. In
September 2014, the sporulating dead wood stems were attached to trees in the field 50 cm above the cankers using cable ties.

**Conidial Spray (SP) treatment.** The same *C. parasitica* and virus haplotypes as for the DW treatment were used (i.e. CpMG8 for Chablais and CpMG47 for La Côte). Hypovirulent *C. parasitica* isolates were grown for two weeks on potato dextrose agar (PDA; 39 gl⁻¹; Difco™ Laboratories, Detroit, USA) in Petri dishes at 25°C under light conditions (3330 lx, 14 h photoperiod, MA Illuminance Meters, Minolta, Japan). For spore harvesting, 15 ml sterile distilled water was poured on each culture and recollected using a pipette. A 10⁷ conidia/ml solution was prepared in 1% methylcellulose (Sigma-Aldrich®, St. Louis, USA) (Larena et al. 2010) to protect conidia from desiccation and increase the stickiness of the solution. Before treatment, the virus presence in conidia was verified by cultivating a bulk of them on PDA and assessing the culture type (orange: virus-free; white: virus-infected; Bissegger et al. 1997). In the field, in September 2014 the conidial solution (ca. 2 × 10⁷ conidia per canker) was sprayed on the cankers and distributed evenly with a brush. The treated cankers were sealed with plastic foil to prevent conidia from drying out or being washed away by rain. As conidia were sprayed directly on the canker surface and only had to penetrate into it for virus transmission, this treatment represented a simplified version of the DW treatment where conidia first had to reach the canker.

**Sampling and scoring the cankers and dead wood stems**

To isolate *C. parasitica* from the cankers and from the dead wood stems, bark pieces were removed using a bone marrow biopsy needle (diameter 2 mm; Jamshidi gauge, BD, Franklin Lakes, USA) (Meyer et al. 2015). The biopsy needle was sterilized before each bark piece was removed. The sampling holes in the cankers were sealed with LacBalsam® (Frunol Delicia® GmbH, Delitzsch, Germany) to close a potential entry point for *C. parasitica* spores.
C. parasitica was isolated from the bark samples as described by Bryner et al. (2014). The presence of the virus in the C. parasitica isolates was assessed based on the characteristic white culture morphology induced by a CHV1 infection (Bissegger et al. 1997). If at least one C. parasitica isolate per canker showed white culture morphology, the canker was scored as virus-infected, whereas if all isolates recovered were orange, the canker was considered as virus-free.

To determine the effect of a virus-infection on canker development and activity, the cankers were measured (length and width) and assessed for activity (active or callused; Bryner et al. 2013) at each sampling date (March 2015, July 2015, and November 2015). During the first two sampling dates (March and July 2015), only half of the cankers and dead wood stems were sampled, but all cankers were measured and scored. At the third sampling event (November 2015), all cankers and dead wood stems were sampled and scored.

**Sampling and scoring of natural cankers**

In June 2016, a total of 88 natural cankers were sampled (Chablais: 21 cankers in CRE and 22 in MON; La Côte: 24 in VIN and 21 in VSY) to determine possible infection with the released virus haplotype CHV1-M3623. Due to differences in the distribution of sprout clusters, in Chablais natural cankers were mostly sampled from the same sprout clusters as the experimental cankers, whereas in La Côte samples were taken from different sprouts.

**Fungal DNA and viral RNA extraction**

RNA was extracted from all virus-infected isolates originating from a canker, but only from one virus infected isolate per dead wood stem. The extraction was performed with the Plant/Fungi Total RNA Purification Kit (Norgen Biotek Corp., Thorold, Canada), excluding the DNA digestion step to allow further determination of the fungal haplotype by microsatellite
genotyping (see below). For cankers and dead wood stems from which only virus-free isolates were recovered, one of the three isolates was randomly selected for DNA extraction and microsatellite analysis. DNA extraction was performed using the KingFisher™ 96 Flex (Thermo Fisher) according to the protocol of the manufacturer.

**Virus haplotypes**

For virus haplotype determination, the complementary DNA synthesis and subsequent amplification/sequencing of a 693 bp region of the open reading frame (ORF) A were performed as described by Prospero and Rigling (2016). ORF A sequences of the virus haplotypes were analyzed with CLC Main Workbench (v. 7.6.2) and aligned in MEGA (v. 7.0.14). For phylogenetic analysis, the sequences were then separated into two different alignments according to the two regions Chablais (CRE, MON) and La Côte (VIN, VSY). The single alignments were processed with Gblocks (v. 0.91b, http://www.phylogeny.fr/, accessed 11.04.2016; Dereeper et al. 2008), trimming all sequences to 568 bp. Statistical selection of best-fit models and building of phylogenetic trees were carried out as described by Meyer et al. (2017), but only maximum likelihood (ML) analyses were performed. The trees were rooted with the reference sequence (CHV-I Euro7; Chen and Nuss 1999) and nodes were collapsed by support at a 50% threshold. Based on the phylogenetic tree, the virus sequences were assigned to different groups associated with either the applied virus haplotype or virus haplotypes already present in the region (Table 2).

**Fungal haplotype**

*Cryphonectria parasitica* isolates were genotyped at ten polymorphic microsatellite loci, as described by Prospero and Rigling (2012). Allele patterns were then compared with those of the multilocus genotypes used in the experiment and, if no match was found, with those of
genotypes occurring in the specific populations or previously identified in Switzerland (Prospero and Rigling 2012).

Biocontrol potential of the *C. parasitica* strains

The performance of the fungal biocontrol strains (Table 2) was estimated by their conidial production (sporulation) and the vertical virus transmission into conidia. Mycelial plugs were punched out from the edge of actively growing 7-day-old cultures with a sterile borer. The plugs were placed upside down in the center of PDA dishes which were wrapped with Parafilm and incubated at 24°C under a light intensity of 3330 lx and 14 h photoperiod. Five replicated plates were prepared for each fungal strain. After 32 days, conidia were washed from the surface of the cultures using 15 ml sterile water. Serial dilutions were made and the conidia concentration determined using a hemocytometer as described in Bryner and Rigling (2011). Conidial suspensions (100 µl) were then spread on PDA and incubated two days at room temperature. For each fungal strain, 28–96 germinated conidia were selected under a dissecting microscope and transferred onto new PDA plates using a sterile needle. The presence of the virus was determined by assessing the culture morphology (orange or white).

Data analysis

For each treated canker, the following features were assessed: (i) infection by a CHV1 virus, if at least one of the three isolates (upper, middle, lower part) showed a virus-infected phenotype; (ii) infection by the virus haplotype CHV1-M3623, if at least one of the three isolates (upper, middle, lower part) was infected by this specific haplotype used for the experiment; (iii) canker size calculated as an ellipse from the measured width and length of each canker; canker size was log-transformed for analysis, as it was not normally distributed; and (iv) canker morphology (active or healed). For each feature, we ran a generalized and
linear mixed model using treatment as a fixed effect and nested random effects to represent
the experimental sites nested within regions that corresponded to the two studied C. 
parasitica strains (CpMG8 for Chablais and CpMG47 for La Côte). Virus infection and canker
morphology were considered as binomial variables, and the logarithm of canker size was
included as a Gaussian variable. We assessed if there was a significant site effect by running
the model using site and treatment as fixed factors and region as a random effect.

The dynamics of treatment effects over time were assessed using generalized linear
mixed models with the following response variables: infection by any CHV1 virus, infection by
CHV1-M3623, and canker size (log-transformed). Treatment and sampling date were
included as fixed factors. Sampled individual and sampling site were included as nested
random factors. In the linear model linking canker size to treatment and date, an interactive
effect between the two fixed factors was included. Because of the relatively small sample
size, it was not possible to test for interactive effects in the models for infection by any virus
and infection by CHV1-M3623.

The in vitro biocontrol potential of each hypovirulent C. parasitica isolate was
evaluated as follows. We used a linear mixed model for the number of spores (log-
transformed) as a function of fungal strain and a generalized mixed model for the percentage
of virus-infected spores as a function of fungal strain (using a binomial error term). Both
models included the region (Chablais or La Côte) where the fungal strain was sampled as a
random effect.

All graphics and statistical analysis were produced in R (3.4.3). Linear mixed effect
models were run using the package nlme (Pinheiro et al. 2018) and generalized mixed effect
models using the package lme4 (Bates et al. 2015).

**Results**
**Experimental cankers**

**Canker infection by CHV1**

In November 2015, 92.5% of the 187 inoculated cankers yielded at least one *C. parasitica* isolate and 60.7% were virus-infected. At all four sites, the prevalence of CHV1 was significantly higher in treated cankers (DW: 50–84%; SP: 68–95%) than in control (untreated) cankers (0–55%) (Figure 2A, Table S2). Sequence analysis revealed that 81% of the virus-infected cankers were infected by the applied CHV1-M3623. The cankers treated with DW (42–74% of them were virus-infected across sites) and SP (50–60%) were significantly more likely to be infected with this haplotype than the control cankers (0–40%) (Figure 2B, Table S2). No statistical difference was found between the two types of treatment (DW versus SP) in CHV1-M3623 infection probability.

Although different virus-carrying fungal haplotypes were used in the two study regions (CpMG8 for Chablais and CpMG47 for La Côte), the factor region did not significantly affect canker infection by CHV1-M3623 (La Côte: 67% of the cankers; Chablais: 54%). A site influence was observed for total virus infection but not for CHV1-M3623 infection, with a significantly higher incidence of virus-infected cankers in VIN (81%, \( P < 0.001 \)) than in the other sites (CRE: 51%, MON: 56%, VSY: 52%) after controlling for the effect of the treatment.

**Canker morphology**

The presence of CHV1-M3623 significantly reduced canker size, with a significantly stronger effect observed for the SP treatment (Figure 2C, Table S2). Non-treated cankers were on average 26.7 cm\(^2\) in size, whereas SP treated cankers were 8.8 cm\(^2\) and DW treated cankers were 18.5 cm\(^2\). No differences in the size of CHV1-M3623-infected cankers were found between regions or among sites. Our analysis showed that the DW treatment led, on average, to a 25% reduction in canker size and the SP treatment led to a stronger reduction of 62.4%.
The incidence of healed cankers in November 2015 differed between the treatments: significantly more SP treated cankers (10–70%) were healed (Figure 2D, Table S2) compared with DW treated (5–35%) and non-treated cankers (0–20%). Among study sites, the incidence of healed cankers across treatments was significantly higher (46%) in VSY than in the other three sites (CRE: 24%, MON: 14%, VIN: 14%).

**Treatment effect over time**

Overall CHV1 and CHV1-M3623 infection of the cankers increased for all canker groups (SP treated, DW treated and control) between the first and second sampling date (March and July 2015; Figure 3A–D, Table S3). From July to November 2015, however, no significant changes in the incidence of infected cankers were observed anymore. Size increased over time for all cankers (Figure 3E–F). However, as early as July, the growth of SP treated cankers was significantly reduced compared with control cankers, whereas DW treated cankers only showed a significant difference in November (Table S3).

**Cryphonectria parasitica genotypes**

Microsatellite profiles were obtained from 170 out of the 173 *C. parasitica* isolates genotyped. In Chablais, experimental cankers were initiated with CpMG30 and CpMG8 used as virus carrier. In November 2015, CpMG30 was still present in 95% of the 75 experimental cankers (Figure 4). Only four cankers in CRE harbored other haplotypes. In La Côte, the experimental cankers were initiated with CpMG15 and CpMG47 used as the virus carrier. In November 2015, CpMG15 was present in 89% of the cankers and 10 cankers were found with other fungal haplotypes (Figure 4).

**Virus haplotypes**
Overall, 85 experimental cankers were infected by virus haplotypes, with sequences identical or similar (1–3 SNP differences) to the applied CHV1-M3623 (Figures 4, S1 and S2). Other viruses clustered with CHV1-M3625, except for three virus sequences from MON (in a control canker), newly classified as CHV1-MON. In La Côte, viruses not originating from CHV1-M3623 clustered either with CHV1-M4762 or CHV1-M4740. Only one CHV1-M4762 haplotype was isolated in VSY, whereas all others were found in VIN. The CHV1-M4740 group was only present in VSY in two cankers.

Survival of *C. parasitica* on dead wood stems

In November 2015 (14 months after treatments), in La Côte the virus-carrying CpMG47, could still be isolated from 35% (VSY) and 65% (VIN) of the dead wood stems, and in Chablais CpMG8 from 44% (MON) and 70% (CRE). All *C. parasitica* cultures isolated from dead wood stems were virus-infected.

Natural cankers

*Cryphonectria parasitica* genotypes

Microsatellite profiles were obtained for 82 isolates out of 88 natural cankers. In Chablais, nine *C. parasitica* haplotypes were present (Figure 4), but only two of them were frequent, i.e. CpMG33 and CpMG44. A similar situation was observed in La Côte, with one single haplotype (CpMG15) predominant. Haplotypes CpMG79, CpMG80, CpMG81, CpMG82, CpMG83 and CpMG84 were found for the first time in Switzerland.

Virus haplotypes

A total of 51 CHV1 sequences (26 in Chablais, 25 in La Côte) were obtained from 27 naturally occurring cankers. The released CHV1-M3623 was detected in 10 natural cankers in
Chablais, mainly in sprout clusters with a treated experimental canker (Figure 4, Table 3) and in one case without treated canker. Other haplotypes belonged to CHV1-M3624 or CHV1-M3625, or to the new haplotype CHV1-MON. In La Côte, none of the natural cankers sampled was infected by CHV1-M3623 (Table 3). The virus-infected natural cankers contained CHV1-M4762, CHV1-M4740, or the new haplotype CHV1-VSY. The latter differed by nine SNPs from CHV1-M4762 and 12 SNPs from CHV1-M4740 (Figure S2).

**Sporulation and vertical virus transmission in vitro**

Sporulation on PDA at 24 °C was significantly affected by fungal haplotype (Figure 5A, Table S4) and virus transmission into the conidia by the fungus-virus combination (Figure 5B, Table S4). The highest vertical transmission rate (91% infected conidia) was observed for M4762 (La Côte), whereas the lowest rate (13%) was observed for M3624 (Chablais). Virus-infected haplotypes applied in La Côte produced, on average, more virus-infected spores than haplotypes used in Chablais (75% vs. 24%), even after controlling for haplotype. Vertical transmission rate positively correlated with the amount of spores produced (Figure 5C, Table S4).

**Discussion**

Although *Cryphonectria parasitica* is mainly known for being an aggressive primary pathogen on American (*Castanea dentata*) and European (*C. sativa*) chestnut, it also has the ability to survive and sporulate on the bark of moribund or recently dead chestnut branches or stems for more than one year (Prospero et al. 2006). The relevance of dead chestnut wood in forests affected by chestnut blight is, however, still controversial for the epidemiology of the disease, especially regarding the occurrence of hypovirulence. Previously, Prospero et al. (2006) showed that recently dead chestnut wood supports asexual sporulation of CHV1-
infected *C. parasitica* strains and, therefore, might promote the natural CHV1 dissemination. However, the role of dead chestnut wood in biocontrol of chestnut blight remained unclear. Here, we used molecular markers in a field experiment and demonstrated that virulent *C. parasitica* cankers become virus-infected by hypovirulent conidia produced on fresh dead wood. As expected, infection by the virus significantly reduced canker growth and promoted canker healing. Fresh dead chestnut wood can, thus, act as reservoir of virus-infected *C. parasitica* inoculum and be beneficial for the natural biological control of the disease.

Fourteen months after DW and SP treatment most experimental cankers had been infected by the applied hypovirulent conidia in all four study sites. Most likely conidia germinated, hyphal fusion with the canker isolate occurred and virus was transmitted into the canker isolate. Since the conidial suspension was sprayed directly on the canker surface, whereas the sporulating stem was set 50 cm above the canker, one might have expected a higher canker infection rate by SP treatment. The results suggest that the amount of virus-infected conidia coming in contact with a virulent canker might not be decisive for virus transmission to the canker. However, more detailed experimental analyses (e.g. treatment with different concentrations of conidia) should be conducted to confirm this hypothesis. The conidia produced on the dead wood stems were most likely washed down to the target cankers by rain water or transported by insects moving along the tree stems. Unlike the virus, the fungal strain of the conidia rarely became established in the target cankers and only acted as virus carrier, as shown before by Hoegger et al. (2003). Virus infection significantly reduced canker expansion in both treatments. The trend over time reveals a faster effect of the SP treatment in comparison to the DW treatment. A similar trend was observed for the healing process of virus-infected cankers. These differences between the two treatments may be due to the larger amount of virus inoculum provided by the SP treatment than with the DW treatment.
Our findings may have implications for chestnut forest management, as they suggest a possible beneficial role of fresh dead wood for the biocontrol of chestnut blight in forest stands where hypovirulence is already present. Previous recommendations were to remove dead chestnut wood from forests with established natural hypovirulence because dead wood fosters sexual reproduction of virus-free *C. parasitica* strains (Bryner et al. 2014; Heiniger and Rigling 1994). An increased diversity of vc types through sexual reproduction might disrupt natural hypovirulence by reducing virus spread. In chestnut stands with no hypovirulence (e.g. at the disease front), removing dead chestnut wood is meaningful as it reduces the amount of substrate on which virulent fungal strains can sporulate. Since bark is the main substrate for *C. parasitica* growth and sporulation, fresh dead wood with intact bark is particularly critical to be removed. In stands where *C. parasitica* is already controlled by CHV1, this sanitation practice may, however, not be that important. For instance, field investigations in Italy evidenced limited tree mortality in unmanaged plots (Turchetti and Maresi 2008). Our study shows that virus-infected conidia produced on fresh dead wood are able to transmit the virus to cankers on living trees. Nevertheless, to definitively conclude on the role of fresh dead chestnut wood in the biocontrol of *C. parasitica*, experimental studies comparing the natural spread of CHV1 in managed (i.e. without dead wood) versus unmanaged (i.e. with dead wood) would be needed. Particularly helpful for forest managers would be to determine whether there is an incidence threshold for CHV1 in a *C. parasitica* population above which dead wood is beneficial for the spread of hypovirulence and, thus, should be left in the stand.

DW and SP treatments could be used for promoting the establishment of CHV1 in chestnut stands. Currently, canker treatment is performed by applying hypovirulent mycelial slurries to the canker surface or in holes cut in the bark around the margin of a canker (Rigling and Prospero 2018). This method is particularly suitable and effective for young trees with smooth and thin bark (Heiniger and Rigling 2009; Milgroom and Cortesi 2004). However,
it has some limitations for the treatment of large and old trees with thick bark, as it is difficult
to bore holes and cankers are often diffuse. Moreover, such treatments can be extremely time
consuming and expensive, especially if cankers are located in the crown of the trees. Finally,
as virus-infected cankers produce a small amount of virus-infected conidia or even do not
sporulate at all (Peever et al. 2000; Prospero et al. 2006), the treated cankers rarely
contribute to virus dispersal. Sporulating dead wood stems could, during the same amount of
time, target specific cankers and increase the amount of virus-infected inoculum in the forest.

The SP treatment could be an interesting approach to the therapeutic treatment of
cankers, e.g. on grafted orchard trees. The effect on canker morphology was considerably
greater for SP than DW treatment, as the sprayed cankers were much smaller and healed
faster. Covering the cankers with a plastic sheet for three months, to prevent the applied
spores from drying out and being washed away, might have created favorable conditions
(humidity, temperature) for germination of the conidia. SP treatment for the biocontrol of
chestnut blight was already tested a few decades ago in North America (Scibilia and Shain
1989). Although the treated cankers were converted, the method was not further developed,
mainly because of the high susceptibility of C. dentata to C. parasitica infection (i.e. new
virulent cankers appeared on the same tree and killed it). Since the European chestnut is
slightly less susceptible to the pathogen than the American chestnut, this should not be a
major problem in Europe. Even if such cankers were to appear, there should be enough time
to treat them before they would become lethal. DW and SP treatments could be combined in
a forest stand by (i) applying the spray on cankers on living trees, and (ii) either applying the
spray on fresh dead wood stems present in the stand or placing sporulating dead wood stems
in strategic positions, e.g. the crown of blighted trees. As the virus disperses mainly by
nearest neighbor contact (Liu et al. 2000), its transmission depends on the distance between
affected chestnut trees. In our experiment, CHV1-M3623 was able to infect natural cankers
that were present either on the same tree or in the same sprout cluster as a treated experimental canker and in one case in a cluster located 2.3 m away. Therefore, by placing sporulating dead wood stems in the crown of blighted trees, the virus-infected conidia may reach and infect existing cankers. Chestnut stems releasing virus-infected conidia can be produced in the laboratory under controlled conditions taking advantage of the competitive ability of *C. parasitica* in colonizing the bark of freshly cut chestnut stems.

Before application in the field, several fungal-viral combinations should be tested for their potential effectiveness in the biocontrol of *C. parasitica*. As the present study and previous studies (Melzer et al. 1992) showed, significant differences exist between hypovirulent haplotypes in sporulation ability and vertical transmission rate of the virus. In our case, the viral population was more dynamic in La Côte than in Chablais, i.e. the viral haplotypes already present in the stands infected the experimental cankers more often. Given that chestnut stands are more widespread in Chablais and hypovirulence was introduced there a few years earlier (2003) than in La Côte (2007) (Heiniger and Rigling 2009), we expected to observe the opposite pattern. Different performances of the hypovirulent *C. parasitica* strains may account for this unexpected result. The biocontrol strains released in Chablais produced significantly fewer virus-infected spores *in vitro* than the three strains present in La Côte. We found that the incidence of virus-infected spores correlated positively with the number of spores produced. Thus, evaluating spore production might be important for selecting the best biocontrol agent *in vitro*; this approach should, however, be tested with more hypovirulent fungal strains. Although different vc types are dominant in Chablais and in La Côte, their incidence in the local *C. parasitica* populations is similar (S. Prospero, *unpublished data*). It is, thus, unlikely that vc type diversity has strongly affected the spread of the released viruses.
**Acknowledgements**

We thank Esther Jung and Hélène Blauenstein for support in the laboratory, Melissa Dawes for English revision, and the Federal Office for the Environment FOEN and the Swiss cantons of Valais and Vaud for financial support. We are also grateful to a reviewer for valuable comments on the manuscript.

**Literature Cited**


### Table 1. Main characteristics of the four experimental sites where this study was conducted.

<table>
<thead>
<tr>
<th>Characteristics</th>
<th>Chablais</th>
<th>La Côte</th>
</tr>
</thead>
<tbody>
<tr>
<td>Altitude (m a.s.l.)</td>
<td>630</td>
<td>475</td>
</tr>
<tr>
<td></td>
<td>480</td>
<td>655</td>
</tr>
<tr>
<td>Forest type</td>
<td>Coppice</td>
<td>Coppice</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>First observation of</td>
<td>1997</td>
<td>2005</td>
</tr>
<tr>
<td>chestnut blight</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Dominant vc types(^a)</td>
<td>EU-1, EU-5</td>
<td>EU-1, EU-5</td>
</tr>
<tr>
<td></td>
<td></td>
<td>EU-2</td>
</tr>
<tr>
<td>Dominant fungal genotypes(^b)</td>
<td>CpMG30</td>
<td>CpMG15</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Natural hypovirulence</td>
<td>-</td>
<td>CHV1-M4740</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>First biocontrol treatment(^c)</td>
<td>2003</td>
<td>2005</td>
</tr>
<tr>
<td></td>
<td>(EU-1 : CHV1-M3624)</td>
<td>(EU-2 : CHV1-M4762)</td>
</tr>
</tbody>
</table>

\(^a\)Prospero and Rigling (2016).

\(^b\)According to Prospero and Rigling (2012).

\(^c\)Previous biocontrol treatments were conducted as described by Rigling and Prospero (2018).
Table 2. Characteristics of the virus-infected *Cryphonectria parasitica* strains used in the *in vitro* fitness test.

<table>
<thead>
<tr>
<th>Region</th>
<th>Fungal strain</th>
<th>Culture type</th>
<th>Viral strain (CHV1-MXXX)</th>
<th>Fungal strain SSR</th>
<th>Vc type</th>
<th>Host tissue</th>
<th>Year/Location of isolation in Switzerland</th>
<th>Treatment</th>
</tr>
</thead>
<tbody>
<tr>
<td>La Côte</td>
<td>M4000 w</td>
<td>CHV1-M4000</td>
<td>CPMG47</td>
<td>EU-2</td>
<td>Canker</td>
<td>1992/Lattecaldo</td>
<td></td>
<td>Deadwood/Spray</td>
</tr>
<tr>
<td></td>
<td>M4762 w</td>
<td>CHV1-M4762</td>
<td>CPMG15</td>
<td>EU-2</td>
<td>Canker</td>
<td>2005/Villars-sous-Yens</td>
<td></td>
<td>Previous treatment</td>
</tr>
<tr>
<td></td>
<td>M4740 w</td>
<td>CHV1-M4740</td>
<td>CPMG15</td>
<td>EU-2</td>
<td>Canker</td>
<td>2005/Villars-sous-Yens</td>
<td></td>
<td>Natural virus</td>
</tr>
<tr>
<td>Chablais</td>
<td>M1709 w</td>
<td>CHV1-M1709</td>
<td>CPMG8</td>
<td>EU-1</td>
<td>Canker</td>
<td>1992/Novaggio</td>
<td></td>
<td>Deadwood/Spray</td>
</tr>
<tr>
<td></td>
<td>M3624 w</td>
<td>CHV1-M3624</td>
<td>CPMG33</td>
<td>EU-1</td>
<td>Canker</td>
<td>1999/Bex</td>
<td></td>
<td>Previous treatment</td>
</tr>
</tbody>
</table>

- MXXX = fungal strain number in the culture collection of WSL;
- w = white (virus-infected; Bissegger et al. 1997);
- CHV1 = *Cryphonectria* hypovirus 1 infected isolates;
- SSR = microsatellite genotype (Prospero and Rigling 2012);
- Vc type = vegetative compatibility type;
- Previous biocontrol treatments were conducted as described by Rigling and Prospero (2018).
Table 3. Location and virus infection of the naturally occurring *Cryphonectria parasitica* bark cankers in the four study sites.

<table>
<thead>
<tr>
<th>Stand</th>
<th>Region</th>
<th>Cankers (N)</th>
<th>CHV1-M3623 Cankers</th>
<th>Distance range&lt;sup&gt;a&lt;/sup&gt; (average)</th>
<th>CHV1-M3623 infected (N)</th>
<th>Cankers (N)</th>
<th>Distance range&lt;sup&gt;a&lt;/sup&gt; (average)</th>
<th>CHV1-M3623 infected (N)</th>
</tr>
</thead>
<tbody>
<tr>
<td>CRE</td>
<td>Chablais</td>
<td>4</td>
<td>1</td>
<td>15</td>
<td>0.2–1.4 (0.6)</td>
<td>5</td>
<td>2</td>
<td>2.4–3.3 (2.8)</td>
</tr>
<tr>
<td>MON</td>
<td>Chablais</td>
<td>4</td>
<td>2</td>
<td>14</td>
<td>0.3–2.3 (0.8)</td>
<td>1</td>
<td>4</td>
<td>2.9–8.3 (5.5)</td>
</tr>
<tr>
<td>VIN</td>
<td>La Côte</td>
<td>2</td>
<td>0</td>
<td>5</td>
<td>0.5–5.0 (2.0)</td>
<td>0</td>
<td>14</td>
<td>1.6–7.3 (3.5)</td>
</tr>
<tr>
<td>VSY</td>
<td>La Côte</td>
<td>2</td>
<td>0</td>
<td>7</td>
<td>0.3–1.3 (0.6)</td>
<td>0</td>
<td>11</td>
<td>1.4–12 (4.5)</td>
</tr>
<tr>
<td>Total</td>
<td>-</td>
<td>12</td>
<td>3</td>
<td>41</td>
<td>-</td>
<td>6</td>
<td>31</td>
<td>-</td>
</tr>
</tbody>
</table>

<sup>a</sup>Distance (in m) between the sampled natural cankers and the closest experimental canker that was treated with either dead wood or conidial spray.
Figure 1. Stromata of *Cryphonectria parasitica* produced on dead chestnut wood with intact bark. (A) Standing dead chestnut sprout; (B) Cut chestnut stem left in the forest. Photos: Phytopathology, WSL.

Figure 2. Presence of the virus in the experimental cankers and effects induced by viral infection. (A) Incidence of CHV1-infected cankers; (B) Incidence of cankers infected by the released haplotype CHV1-M3623; (C) Canker sizes, and (D) Incidence of healed cankers.

Figure 3. Development over time (March – November 2015) of the incidence of CHV1-infected experimental cankers (A, B), of the incidence of experimental cankers infected by the released haplotype CHV1-M3623 (C, D), and of experimental canker size (E, F) in the two treatments (dead wood and conidial spray), and in the controls (non-treated cankers).

Figure 4. *Cryphonectria parasitica* haplotypes and CHV1 haplotypes identified in the experimental cankers non-treated (CO), treated with dead wood (DW) and treated with conidial spray (SP), and in naturally occurring cankers (Natural cankers) at the four study sites.

Figure 5. Biocontrol potential of the virus-infected *Cryphonectria parasitica* haplotypes from Chablais (white) and from La Côte (grey) (Table 2). (A) Number of conidia produced on PDA at 24 °C; (B) Incidence of virus-infected conidia; (C) Correlation between the number of produced conidia and their virus infection rate.
### Figure S1

Rooted maximum likelihood tree of a partial sequence (568 nt) from the ORF A of the CHV1 strains from Creux Boyon (CRE) and Montet (MON) in the Chablais region. Viral strains were recovered from the upper margin (o), the center (m) and the lower margin (u) of the cankers from deadwood (Dw) and conidial spray (Sp) treated or untreated (control, C) cankers. The other CHV1 strains originate from naturally occurring cankers. 1000 bootstraps were performed. Only bootstrap values higher than 65% are shown. Scale bar = substitution per site. Euro7 subtype I reference sequence was used to root the tree. CHV1-M3624 and CHV1-M3625 are virus haplotypes from former treatments. CHV1-M3623 is the haplotype applied with the dead wood stems and the conidial spray. CHV1-MON is a new viral haplotype.

### Figure S2

Rooted maximum likelihood tree of a partial sequence (568 nt) from the ORF A of the CHV1 isolates from Vinzel (VIN) and Villars-Sous Yens (VSY) in the La Côte region. CHV1 strains were recovered from the upper margin (o), the center (m) and the lower margin (u) of the cankers from deadwood (Dw) and conidial spray (Sp) treated or untreated (control, C) cankers. The other CHV1 strains originate from naturally occurring cankers. 1000 bootstraps were performed. Only bootstrap values higher than 65% are shown. Scale bar = substitution per site. Euro7 subtype I reference sequence was used to root the tree. CHV1-M4762 is a virus haplotype from a former treatment, CHV1-M4740 a naturally occurring haplotype. CHV1-M3623 is the haplotype applied with the dead wood stems and the conidial spray. CHV1-VSY is a new viral haplotype.
Stromata of *Cryphonectria parasitica* produced on dead chestnut wood with intact bark. (A) Standing dead chestnut sprout; (B) Cut chestnut stem left in the forest. Photos: Phytopathology, WSL.

207x276mm (300 x 300 DPI)
Stromata of *Cryphonectria parasitica* produced on dead chestnut wood with intact bark. (A) Standing dead chestnut sprout; (B) Cut chestnut stem left in the forest. Photos: Phytopathology, WSL.
Presence of the virus in the experimental cankers and effects induced by viral infection. (A) Incidence of CHV1-infected cankers; (B) Incidence of cankers infected by the released haplotype CHV1-M3623; (C) Canker sizes, and (D) Incidence of healed cankers.

423x317mm (72 x 72 DPI)
Development over time (March – November 2015) of the incidence of CHV1-infected experimental cankers (A, B), of the incidence of experimental cankers infected by the released haplotype CHV1-M3623 (C, D), and of experimental canker size (E, F) in the two treatments (dead wood and conidial spray), and in the controls (non-treated cankers).
Cryphonectria parasitica haplotypes and CHV1 haplotypes identified in the experimental cankers non-treated (CO), treated with dead wood (DW) and treated with conidial spray (SP), and in naturally occurring cankers (Natural cankers) at the four study sites.

317x246mm (72 x 72 DPI)
Biocontrol potential of the virus-infected *Cryphonectria parasitica* haplotypes from Chablais (white) and from La Côte (grey) (Table 2). (A) Number of conidia produced on PDA at 24 °C; (B) Incidence of virus-infected conidia; (C) Correlation between the number of produced conidia and their virus infection rate.
Supplementary materials for:

Role of fresh dead wood in the epidemiology and the biological control of the chestnut blight fungus

Joana Beatrice Meyer, Loïc Chalmandrier, Fabio Fässler, Christopher Schefer, Daniel Rigling, Simone Prospero

Table S1. Single nucleotide polymorphisms between the CHV-1 haplotype that was released in this study (CHV1-M3623) and the haplotypes that were already present in the stands (La Côte: CHV1-M4762 and CHV1-M4740; Chablais: CHV1-M3624 and CHV1-M3625).

<table>
<thead>
<tr>
<th>Nucleotide position in the sequence</th>
<th>CHV1-M3623</th>
<th>CHV1-M4762&lt;sup&gt;a&lt;/sup&gt;</th>
<th>CHV1-M4740&lt;sup&gt;b&lt;/sup&gt;</th>
<th>CHV1-M3624&lt;sup&gt;a&lt;/sup&gt;</th>
<th>CHV1-M3625&lt;sup&gt;b&lt;/sup&gt;</th>
</tr>
</thead>
<tbody>
<tr>
<td>35</td>
<td>A</td>
<td>G</td>
<td>C</td>
<td>C</td>
<td>G</td>
</tr>
<tr>
<td>41</td>
<td>T</td>
<td>C</td>
<td>C</td>
<td>G</td>
<td>C</td>
</tr>
<tr>
<td>121</td>
<td>T</td>
<td>T</td>
<td>G</td>
<td>T</td>
<td>T</td>
</tr>
<tr>
<td>163</td>
<td>A</td>
<td>C</td>
<td>G</td>
<td>G</td>
<td>A</td>
</tr>
<tr>
<td>186</td>
<td>C</td>
<td>C</td>
<td>T</td>
<td>C</td>
<td>G</td>
</tr>
<tr>
<td>187</td>
<td>A</td>
<td>G</td>
<td>C</td>
<td>G</td>
<td>A</td>
</tr>
<tr>
<td>209</td>
<td>T</td>
<td>G</td>
<td>T</td>
<td>G</td>
<td>A</td>
</tr>
<tr>
<td>249</td>
<td>C</td>
<td>C</td>
<td>T</td>
<td>C</td>
<td>G</td>
</tr>
<tr>
<td>286</td>
<td>T</td>
<td>G</td>
<td>T</td>
<td>C</td>
<td>G</td>
</tr>
<tr>
<td>313</td>
<td>C</td>
<td>C</td>
<td>C</td>
<td>G</td>
<td>A</td>
</tr>
<tr>
<td>390</td>
<td>A</td>
<td>G</td>
<td>A</td>
<td>A</td>
<td>A</td>
</tr>
<tr>
<td>447</td>
<td>T</td>
<td>G</td>
<td>A</td>
<td>G</td>
<td>A</td>
</tr>
<tr>
<td>489</td>
<td>T</td>
<td>C</td>
<td>A</td>
<td>G</td>
<td>A</td>
</tr>
<tr>
<td>547</td>
<td>G</td>
<td>A</td>
<td>G</td>
<td>A</td>
<td>A</td>
</tr>
</tbody>
</table>

<sup>a</sup>Virus used for former treatments (Prospero and Rigling 2016).

<sup>b</sup>Naturally occurring virus in VSY.
**Table S2.** Estimates of linear and generalized linear model characterizing the link between cankers features and applied treatments.

<table>
<thead>
<tr>
<th>Response variable</th>
<th>Virus-infection (CHV1-M3623)</th>
<th>Virus infection (all viruses)</th>
<th>Size (log)</th>
<th>Size of control cankers (log)</th>
<th>Healed</th>
</tr>
</thead>
<tbody>
<tr>
<td>Model type</td>
<td>Binomial</td>
<td>Binomial</td>
<td>Gaussian</td>
<td>Gaussian</td>
<td>Binomial</td>
</tr>
<tr>
<td>Number of observations</td>
<td>173</td>
<td>173</td>
<td>185</td>
<td>87</td>
<td>187</td>
</tr>
<tr>
<td>Intercept</td>
<td>-1.945</td>
<td>-1.070</td>
<td>5.287 (t = 42.27, p &lt; 0.01)</td>
<td>5.38 (t = 34.86, p &lt; 0.001)</td>
<td>-2.58 (z = -3.622, p &lt; 0.001)</td>
</tr>
<tr>
<td>Treatment</td>
<td>DW</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>2.122 (z = 3.614, p &lt; 0.001)</td>
<td>1.717 (z = 3.421, p &lt; 0.001)</td>
<td>-0.2969 (t = -1.973, p = 0.05)</td>
<td>-0.371 (t = -1.74, p = 0.0855)</td>
<td>1.04 (z = 1.52, p = 0.13)</td>
</tr>
<tr>
<td></td>
<td>SP</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>2.363 (z = 4.035, p &lt; 0.001)</td>
<td>2.051 (z = 4.050, p &lt; 0.001)</td>
<td>-0.977 (t = -6.48, p &lt; 0.001)</td>
<td>-1.22 (t = -5.591, p &lt; 0.001)</td>
<td>1.99 (z = 2.97, p = 0.003)</td>
</tr>
<tr>
<td>Random effect std. deviation</td>
<td>Fungi strain</td>
<td>0.00</td>
<td>0.00</td>
<td>4.14 x 10^{-5}</td>
<td>2.384 x 10^{-5}</td>
</tr>
<tr>
<td></td>
<td>Site within Fungi strain</td>
<td>0.00</td>
<td>0.6439</td>
<td>2.80 x 10^{-5}</td>
<td>2.72 x 10^{-6}</td>
</tr>
</tbody>
</table>
Table S3. Estimates of the linear model linking canker size to sampling date and treatment.

<table>
<thead>
<tr>
<th>Response variable</th>
<th>Virus-infection (CHV1-M3623)</th>
<th>Virus-infection (all viruses)</th>
<th>Size (log)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Model type</td>
<td>Binomial</td>
<td>Binomial</td>
<td>Gaussian</td>
</tr>
<tr>
<td>Number of observations</td>
<td>310</td>
<td>325</td>
<td>252</td>
</tr>
<tr>
<td>Intercept</td>
<td>-4.35 (z = -3.86, p &lt; 0.001)</td>
<td>-4.59 (z = -4.86, p &lt; 0.001)</td>
<td>3.613 (t = 17.56, p &lt; 0.001)</td>
</tr>
<tr>
<td>Date</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>July</td>
<td>2.54 (z = 4.01, p &lt; 0.001)</td>
<td>2.57 (z = 4.13, p &lt; 0.001)</td>
<td>1.078 (t = 7.867, p &lt; 0.001)</td>
</tr>
<tr>
<td>November</td>
<td>2.42 (z = 4.03, p &lt; 0.001)</td>
<td>2.12 (z = 3.76, p &lt; 0.001)</td>
<td>1.726 (t = 12.59, p &lt; 0.001)</td>
</tr>
<tr>
<td>Treatment</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>DW</td>
<td>3.28 (z = 3.88, p &lt; 0.001)</td>
<td>3.03 (z = 4.25, p &lt; 0.001)</td>
<td>0.3435 (t = 1.979, p = 0.0514)</td>
</tr>
<tr>
<td>SP</td>
<td>3.22 (z = 3.65, p &lt; 0.001)</td>
<td>2.98 (z = 4.05, p &lt; 0.001)</td>
<td>-0.09348 (t = -0.5461, p = 0.59)</td>
</tr>
<tr>
<td>Date x Treatment</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>DW - July</td>
<td>/</td>
<td>/</td>
<td>-0.2756 (t = -1.668, p = 0.0973)</td>
</tr>
<tr>
<td>DW - November</td>
<td>/</td>
<td>/</td>
<td>-0.5137 (t = -3.108, p = 0.002)</td>
</tr>
<tr>
<td>SP - July</td>
<td>/</td>
<td>/</td>
<td>-0.6481 (t = -3.972, p &lt; 0.001)</td>
</tr>
<tr>
<td>SP - November</td>
<td>/</td>
<td>/</td>
<td>-0.9246 (t = -5.668, p &lt; 0.001)</td>
</tr>
<tr>
<td>Random effect</td>
<td>Site</td>
<td>1.28 x 10-14</td>
<td></td>
</tr>
</tbody>
</table>
Std. deviation Individual within sites

3.351

2.10
Table S4. Linear model estimate linking sporulation and rate of vertical CHV1 transmission of CHV1-M3623-infected *Cryphonectria parasitica* strains in *in vitro* experiments to fungal strain and virus-infection.

<table>
<thead>
<tr>
<th>Response variable</th>
<th>Sporulation (24°C) – log transformed</th>
<th>Spore infection rate</th>
</tr>
</thead>
<tbody>
<tr>
<td>Number of observations</td>
<td>49</td>
<td>72</td>
</tr>
<tr>
<td>Intercept</td>
<td>14.9 (t = 69.5, p &lt; 0.001)</td>
<td>-1.84 (t = -11.3, p &lt; 0.001)</td>
</tr>
<tr>
<td>Strain</td>
<td>M1709</td>
<td>1.41 (t = 6.99, p &lt; 0.001)</td>
</tr>
<tr>
<td></td>
<td>M4000</td>
<td>2.21 (t = 9.92, p &lt; 0.001)</td>
</tr>
<tr>
<td></td>
<td>M4762</td>
<td>4.23 (t = 13.8, p &lt; 0.001)</td>
</tr>
<tr>
<td></td>
<td>M4740</td>
<td>2.77 (t = 11.5, p &lt; 0.001)</td>
</tr>
</tbody>
</table>
Rooted maximum likelihood tree of a partial sequence (568 nt) from the ORF A of the CHV1 strains from Creux Boyon (CRE) and Montet (MON) in the Chablais region. Viral strains were recovered from the upper margin (o), the center (m) and the lower margin (u) of the cankers from deadwood (Dw) and conidial spray (Sp) treated or untreated (control, C) cankers. The other CHV1 strains originate from naturally occurring cankers. 1000 bootstraps were performed. Only bootstrap values higher than 65% are shown. Scale bar = substitution per site. Euro7 subtype I reference sequence was used to root the tree. CHV1-M3624 and CHV1-M3625 are virus haplotypes from former treatments. CHV1-M3623 is the haplotype applied with the dead wood stems and the conidial spray. CHV1-MON is a new viral haplotype.
Rooted maximum likelihood tree of a partial sequence (568 nt) from the ORF A of the CHV1 isolates from Vinzel (VIN) and Villars-Sous Yens (VSY) in the La Côte region. CHV1 strains were recovered from the upper margin (o), the center (m) and the lower margin (u) of the cankers from deadwood (Dw) and conidial spray (Sp) treated or untreated (control, C) cankers. The other CHV1 strains originate from naturally occurring cankers. 1000 bootstraps were performed. Only bootstrap values higher than 65 % are shown. Scale bar = substitution per site. Euro7 subtype I reference sequence was used to root the tree. CHV1-M4762 is a virus haplotype from a former treatment, CHV1-M4740 a naturally occurring haplotype. CHV1-M3623 is the haplotype applied with the dead wood stems and the conidial spray. CHV1-VSY is a new viral haplotype.