

Besides stem canker severity, oilseed rape host genotype matters for the production of Leptosphaeria maculans fruit bodies

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1 Besides stem canker severity, oilseed rape host genotype matters for the production of 2 Leptosphaeria maculans fruit bodies 3 4 Lydia Bousset*, Patrick Vallée, Régine Delourme, Nicolas Parisey, Marcellino Palerme, Melen 5 Leclerc 6 7 INRAE, UMR1349 IGEPP, F-35653 Le Rheu, France 8 *Email: lydia.bousset @inrae.fr, Phone: +33 2 23 48 51 85, Fax: +33 2 23 48 51 50 9 10 Total word count: main body of the text: 4875 words 11 12 Introduction: 995 13 Materials and Methods: 1590 14 Results: 722 15 Discussion: 1596 16 17 With 8 Figures and 3 Tables (2 with colours) 18 Supporting information with 14 Figures and 13 Tables (2 with colours) 19 20 **Abstract** 21 22 For fungal cyclic epidemics on annual crops, the pathogen carry-over is an important but poorly 23 documented step. Plant resistance affects the pathogen development within the epidemics but we lack 24 data on the inter-annual transmission of inoculum. For Leptosphaeria maculans on 15 oilseed rape 25 genotypes in field during 4 growing seasons, stem canker severity was visually scored at harvest. The

number of fruit bodies produced on incubated stubble was quantified using an automated image

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analysis framework. Our results confirm that fruit body production increases with disease severity and is significantly affected by host genotype and nitrogen supply. Tracking individual stems through incubation, we confirm for the first time that the oilseed rape genotype has a direct effect on inoculum production, not only disease severity. This major effect of genotype on inoculum carry-over should be taken into account in models of varietal deployment strategies.

Keywords: blackleg, disease phenotyping, epidemiology, image analysis, inoculum potential, transmission

Introduction

A key development in disease control has been the breeding and deployment of crop varieties with genetically controlled resistance to pathogens, but most pathogens have repeatedly evolved and overcome qualitative resistance genes (Burdon *et al.* 2016). Therefore, a critical challenge is to design and implement durable crop protection strategies against rapidly evolving pathogens (Cowger & Brown 2019). Any mitigation strategy that reduces the transfer of inoculum between seasons, therefore the effective size of pathogen populations, should limit the evolutionary potential of pathogens, and increase resistance durability (Bousset & Chèvre 2013; Zhan *et al.* 2015). In experiments, combining quantitative with qualitative resistance (Brun *et al.* 2010: Delourme *et al.* 2014; Lasserre-Zuber *et al.* 2018), burying stubble to reduce inoculum transmission (Daverdin *et al.* 2012), or removing leaf litter in combination with reduced fungicide application (Didelot *et al.* 2016) has delayed adaptation. In models and experiments, the spatio-temporal distribution of host resistance in cultivated landscapes was optimised to mitigate disease transmission (Lô-Pelzer *et al.* 2010; Bousset *et al.* 2018; Papaïx *et al.* 2018; Rimbaud *et al.* 2018; Watkinson-Powell *et al.* 2019). However, model validation and accurate predictions rely on the availability of data that are difficult and costly to collect at these large scales.

On annual crops, many plant diseases have cyclic epidemics (Zadoks & Schein 1979; Bousset & Chèvre 2013; Fig. 1A). Their dynamics are highly influenced by both temporal and spatial discontinuities, induced by the climate (e.g. seasonality) or by human actions (e.g. sowing and harvesting). At the beginning of the cropping season, the primary inoculum level is a product of carry-over from the prior crop(s) and/or dispersal. Mitigation strategies can thus target this carry-over of the pathogen using resistant varieties (Marcroft *et al.* 2004a), biocontrol agents (Bailey *et al.* 2004), or preventive management of crop residues (Wherrett *et al.* 2003). However, inoculum survival and transmission are difficult to estimate in field and predict (Bailey *et al.* 2004; Bousset *et al.* 2015).

Host resistance impacts pathogen life history traits during the epidemic phase (e.g. infection efficiency, latent period or sporulation) (Bruns *et al.* 2012; Delmas *et al.* 2016; Dumartinet *et al.* 2020; Leclerc *et al.* 2019; Bove & Rossi 2020). Qualitative host resistance reduces the number of compatible infections whereas quantitative host resistance reduces the amplification during the epidemics (Fig. 1B). In addition, the selective deployment of resistant host genotypes can reduce landscape connectivity following the intercrop (Lô-Pelzer *et al.* 2010; Bousset & Chèvre 2013; Papaïx *et al.* 2018; Rimbaud *et al.* 2018; Watkinson-Powell *et al.* 2019). Little is known about the direct effect of quantitative host resistance on initial amount of inoculum at the beginning of the next cropping season (i.e. inter-annual transmission, Fig. 1B). Quantitative resistance affects transmission in *Phytophthora infestans* through a trade-off between crop infection and survival on tubers (Pasco *et al.* 2016). A trade off was detected in *Zymoseptoria tritici*, but varietal effects were not investigated (Suffert *et al.* 2018a). In downy mildew, the effect of quantitative resistance was assessed on oospore production (inoculum for inter-annual transmission; Delbac *et al.* 2019).

Inoculum transmission between seasons was described in *Leptosphaeria maculans* on winter oilseed rape sown in autumn. This fungal pathogen initiates epidemics early in the cropping season (in autumn) with stubble-borne ascospores that can spread between fields across the landscape (West *et al.* 2001). It produces Phoma leaf spots on host leaves that are observed between autumn and early spring. Then, stem cankers develop from spring to summer, up to the time of harvest, following

systemic growth of fungal hyphae from leaf spots to the leaf petiole through xylem vessels, and subsequently to the stem base. The fungus can survive as hyphae in crop stubble, more specifically in stems around crown level, forming two kinds of fruit bodies: pycnidia and pseudothecia. Pseudothecia can only be formed following sexual reproduction if isolates of opposite mating types co-occur in the same oilseed rape stem. Infected stubble ensures the carry-over of the fungus from one cropping season to the next and serves as the main source of inoculum.

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For L. maculans, several studies have shown that the number of produced pseudothecia, that release initial inoculum ascospores, was correlated with Phoma stem canker severity (McGee & Emmett 1977; Marcroft et al. 2004a; Lô-Pelzer et al. 2009) and that some quantitative host resistances tends to decrease severity at harvest (Marcroft et al. 2004b; Lô-Pelzer et al. 2009). Host genotype can significantly affect both the visual density of pseudothecia, and the number of released ascospores (Marcroft et al. 2004b). However, properly disentangling the effects of host genotype on stem canker severity and fruit body production remains to be achieved. Therefore, the question of a direct genetic effect on the production of fruit bodies, and thus temporal transmission of inoculum between seasons, is still unknown for this pathosystem. The development of high-precision and high-throughput phenotyping methods could improve experimental quantification of plant pathogens, alleviate the tedious manual quantification of fruit bodies on many genotypes, and improve knowledge and data on pathogen carry-over. Imaging and processing algorithms to segment and count regions, circumvented the long and tedious manual counting (Stewart et al. 2016; Karisto et al. 2018; Yang & Hong 2018) and were applied to the automated quantification of fruit bodies of *L. maculans* (Bousset et al. 2019). This offers a means to overcome the experimental bottleneck and assess the effect of host quantitative resistance on pathogen carry-over.

In this study, we address the effect of host resistance on the carry-over of inoculum between consecutive years, to understand how host genotype might affect fruit body production and the effects of a common cropping practice, i.e. nitrogen fertilization, and disease severity at harvest in the preceding growing season (year). We consider *L. maculans* on oilseed rape as an example

pathosystem and use an imaged-based phenotyping method for quantifying fruit bodies that appeared on diseased stems collected in experimental field plots, after an incubation period. We analyse the main drivers of this poorly known life-history trait of the pathogen.

Materials and methods

Experimental fields and contamination of field plots

This experiment was run on the INRAE UE La Motte experimental station located in Le Rheu (48.1°N, 1.5°W), in Brittany, France. Winter *B. napus* genotypes were grown in 5-row plots (2.5 m²) and assessed for stem canker (details are given in Table S1.1). The cropping season extends from sowing in autumn to harvest the following summer and was repeated in the 4 years named 12-13, 13-14, 14-15 and 15-16. We followed the same 15 genotypes in years 12-13, 13-14 and 14-15, (Table 1; S1.2) and six genotypes were added in year 15-16 (Table 1). The varieties were chosen to represent a wide range of winter oilseed rape diversity and derived from different breeding programs.

In the four years, nitrogen supply was normal (Nh) according to usual practices in conventional cultivation to achieve yield potential of modern winter oilseed rape varieties (Bouchet et al. 2014) (Table S1.3). In years 14-15 and 15-16, the panel was replicated with low (Nl) nitrogen availability. On these Nl plots, no organic matter was spread on the fields for 3 years before the trials and the previous crops were grown under a low input management system. This limited the amount of mineral N in soil in the experimental plots, avoiding organic matter mineralisation into nitrogen later available to the plants.

Homogeneous disease infection was achieved with contaminated stems collected from the previous year on the susceptible Bristol and miscellaneous varieties. In the experimental area and around Phoma stem cankered stubble, only *L. maculans* leaf spots were observed which supported the hypothesis of the very small prevalence of the other causal agent *L. biglobosa* in this area (Bousset et al. 2019). Stems were scattered throughout the plots in each cropping season at a density of two stems per m² when the crop was at the two to three leaf growth stage. Each variety was grown on

three replicated plots, from which stems were pooled for the current analysis. Sprinklers were run twice per day (4 mm per day) from plant emergence to the appearance of first leaf spots to promote fruit body maturation and spore release. This kept the time from spreading the contaminated stems to the appearance of leaf spots similar over years (Fig. S1E). No fungicides were applied throughout the season. Phoma stem canker severity was assessed 2-3 weeks before crop maturity on a 1 to 6 scale as follows: S1 = no disease, S2a = 1-5%, S2b - 6-25%, S3 = 26-50%, S4 = 51-75%, S5 = 76-99%, S6= 100% of crown cross section with Phoma stem canker symptoms. Severity classes 2a and 2b were distinct for the analysis of fruit body formation on stems, but pooled before the calculation of plot canker severity. The G2 aggregated disease index was calculated as follows (Aubertot et al. 2004):

 $G2 = (0 \times n_1 + 1 \times n_2 + 3 \times n_3 + 5 \times n_4 + 7 \times n_5 + 9 \times n_6)/N$

with n_i being the number of stems in severity class S_i , respectively and N the total number of stems.

The climate of the experimental area is oceanic. Meteorological data were obtained from the INRAE CLIMATIK database, for Le Rheu weather station, on an hourly basis. Cumulative temperature (Fig. S1ab), rainfall (Fig. S1cd), and number of days that are favourable for pseudothecia maturation (Fig. S1e) were calculated from start (shortly after harvest in June; Table S1.1) to end of incubation when stems were retrieved from the field (at the end of the following winter; Table S1.1). A day was considered favourable if the mean temperature was between 2 and 20°C and if the cumulative rainfall over the previous 11 days before (including the day in question) exceeded 4 mm (Aubertot *et al.* 2006; Lô-Pelzer *et al.* 2009). Given these parameter values, 64 favourable days were required for 50% of pseudothecia to reach maturation. We were able to compare the number of calendar days needed to reach maturation in different years (Fig. S1E).

Stubble incubation to promote fruit body formation

Phoma stem canker severity was assessed at harvest, and then tracked for each individual stem piece comprising the crown and upper 10 cm (Bousset *et al.* 2019). The sampling was constrained by the availability of diseased stems across severity classes. After pooling the three replicated plots of

each variety, 30 stems in each of the 6 severity classes were randomly selected. If fewer than 30 stems (per class) were available, then all were kept. Within each variety and severity class, groups of 5 stem pieces attached together on BBQ sticks were labelled with a barcode (Bousset *et al.* 2019).

The selected stem pieces were incubated in field conditions at INRAE Le Rheu. Over summer, they were placed on a 1:1:1 mix of sand, peat and compost spread on the ground to avoid weed emergence, then transferred in autumn in experimental plots of winter oilseed rape to mimic incubation under the following season's crop and to allow detection of released spores. Complete maturation was indicated by the end of the appearance of new leaf spots on susceptible plants around the incubating stem pieces. The stem pieces were recollected, washed, dried and stored (Table 1). All stem pieces from a given year were incubated in the same time and location, starting in the summer following harvest.

Image acquisition, processing and post-processing

The fruit bodies appeared on diseased stem pieces were quantified using image-based phenotyping settings described in Bousset *et al.* (2019). In short, a picture of each group of 5 dry stems was taken with the barcoded label, placed on a glass plate, 16 cm above a blue background (PVC sheet Lastolite Colormatt electric blue). Two FotoQuantum LightPro 50 × 70cm softboxes were placed on both sides of the stem pieces with 4 daylight bulbs each (5400K, 30W) within the lower 45° angle. Pictures were taken with a Nikon D5200 with an AF-S DX Micro Nikkor 40mm 1:2.8G lens, on a Kaiser Repro stand, with a wired remote control. Aperture was set at F22 for maximal depth of field, iso 125, daylight white balance. Pictures were saved as RGB images with a resolution of 6000 × 4000 pixels.

Pictures were pre-processed by reading the barcode to rename the files, splitting each digital image in several new images, each containing only one stem segmented with an unsupervised method. Finally, each new image was cropped to keep only the 5 cm portion on the crown end of the stem, i.e. where pseudothecia are mainly located.

Then, stem pixels were classified either in state F (fruit bodies) or in state S (stem) using a supervised machine learning algorithm. We did not re-train the algorithm for processing our images, because learning and testing data used by Bousset et al. (2019) already included some images collected for this study. Predicted images were post processed through a computer-assisted expert curation using a graphical user interface. Each processed image was assigned a post-processed state, i.e. "correct" or "incorrect", by visually evaluating the predicted image compared to the original one (Bousset *et al.* 2019).

Statistical analyses

We analysed the effects of genotype, year and nitrogen fertilization on stem canker severity using a proportional odds model. We used an ordered logit regression model that handles ordinal data. The design of the experimental data only allowed us to estimate and test the first order effects and the interaction between genotype and nitrogen fertilization.

The density of fruit bodies detected was analysed on the post-processed images validated by the curator (state "correct"). For each image i we considered the number of pixels in states F (fruit bodies) and S (stem), i.e. n_{Ei} and $n_{S.i}$, and used a likelihood function based on $n_{Ei} \sim B[(n_{Ei} + n_{S.i}), p]$ with a logit link function to build Generalised Linear Models and analyse the effects of selected factors with Wald tests. We first considered the complete unbalanced dataset that only enables the identification and test of first order effects (i.e. year, severity prior incubation, plant genotype and nitrogen fertilization) and the interactions between genotype and nitrogen fertilization, and stem severity with nitrogen fertilization (Table 1). Second, in order to go further and investigate second and third order effects (i.e. interactions) we split the 4 years data into 3 datasets: Data14-16 (2 years 14-15 and 15-16, 2 nitrogen fertilization levels Nh and Nl, 15 genotypes), Data12-16 (4 years 12-13, 13-14, 14-15 and 15-16, 15 genotypes, all on high level of fertilization Nh), and Data15-16 (year 15-16, 21 genotypes and 2 nitrogen fertilization levels Nh and Nl). Note that for all these subdatasets the severity prior incubation (7 levels) was also included as a categorical variable in the regression models.

The whole image processing was developed in Python (Van Rossum & Drake 1995) whereas statistical analyses were performed using R (R Core Team 2019).

Prediction of inoculum potential

We simulated the production of inoculum in a population of infected host plants by coupling two stochastic processes. We first considered 10 experimental plots of 100 plants, whose severity drawn from a Multinomial distribution with event probabilities corresponding to those estimated with the ordered logistic regression model (Tables S5.1 and S5.2). Second, we assumed that the size of each individual stem is 700 000 pixels, i.e. twice the average observed number of pixels per processed image, and for each stem we drew the number of fruit body pixels using a Binomial distribution, with probability parameter previously estimated in the GLM model (Tables S5.3 and S5.3). To illustrate the importance of host genotype on fruit body production, we compared the output of the simulation with versus without host genotype.

Results

Severity data at harvest

The observed distribution of canker severities was assessed at harvest, before the sampling of stems selected for fruit body formation assays (See Materials and Methods). Model of the complete ordinal dataset indicated effects (p-values <0.05) of the genotype, the year, the fertilisation and an interaction between genotype and fertilisation on canker severity at harvest (Aggregated G2 index, Table 2; Fig. S2.1; Fig. S2.2).

Fruit body formation was strongly influenced by year, genotype and disease severity

Across datasets, fruit body formation was affected by host genotype, year, and disease severity at harvest. The year was the most influential parameter, with about 20% of the explained deviance for all analyses (Table 3). Host genotype explained between 6 and 17% of the explained deviance of fruit body formation. Finally, severity of infected stems before incubation explained between 5 and 10 %

of the total deviance. Raw data as well as pairwise comparisons of modalities with least-squares means indicated differences between genotypes and years when stems were collected. The production of fruit bodies was lowest in year 15-16 and highest in year 13-14 (Fig. 3; Supplementary Information S3).

The amount of fruit bodies produced increased with stem canker severity as already shown by Bousset et al. (2019). Stem canker severity classes were ranked from the lowest to highest, except for classes S6 and S5 that were in reversed order (Fig. 4a; Supplementary Information Fig. S3.1; 3.4; 3.7). However, for each severity class, the fruit bodies produced depended on the genotype (Fig. 4; 5; Supplementary Information Fig. S4.1; S4.2). Some genotypes like Al and Fa produced greater numbers of fruit bodies at each severity class than genotypes like As and Av. Noteworthy, here the stem pieces have been individually tracked, so the production of a genotype can be compared at each severity class. One can also note that a severely infected genotype like Yu (having no stem in classes S1 and S2a) is not the one producing the greatest amount of fruit bodies at a given severity class. In contrast, a mildly infected genotype like Da (having no stem in classes S5 and S6) produced more fruit bodies at a given severity class than e.g. genotypes As or Av.

Because several genotypes did not occur either in the first two (S1-S2) or the last two (S5-S6) severity classes, it was not possible to estimate all the coefficients of the genotype × severity interaction on the full dataset. However, when considering the genotypes for which this interaction was identifiable, we found a significant genotype × severity interaction, suggesting that the amount of produced fruit bodies by each severity class may change with host genotypes.

Nitrogen fertilization of the crop has a significant but minor effect on fruit body formation

We detected a significant fertilization \times host genotype interaction, indicating that fertilization has varying effects on disease symptoms of different host genotypes. (Fig. 2). The fertilization \times disease severity and fertilization \times year interactions were also significant. However, as shown by the little deviance explained by these first, second and third order factors, nitrogen fertilization had a

minor influence on fruit body production (Table 3; Fig. 6). In fact, nitrogen explained below 1% of the deviance in the Data15-16 dataset, and only the nitrogen × genotype interaction explained 5% of the deviance (Table 3).

Simulated fruit body formation

The stochastic simulation of fruit body production based on fitted event probabilities (proportional odds and generalized linear models) allowed us to predict the potential level of inoculum produced by field plots for each genotype x year x nitrogen fertilization case (Table S5.1; S5.2; S5.3; S5.4). Simulations outputs illustrate the variability of fruit body formation and confirm the importance of host genotype on this life-history trait of the fungal pathogen (Fig. 7; Fig. S5.1).

When the simulated fruit body production was plotted over canker severity at harvest using the aggregated G2 index, among the genotypes, either susceptible (high canker severity) or with higher levels of quantitative resistance (low canker severity), the pixels of fruit bodies are variable (Fig. 8). A very susceptible genotype is not always the one producing the greatest amount of fruit bodies for the following year. Moreover, genotypes with higher levels of quantitative resistance have low canker severity at harvest but are still diverse with regard to fruit body production (Fig. 8).

Discussion

With field experiments and image-based automated quantification of fruit bodies on individual stems, we disentangled the role of the year, the genotype, the disease severity and the nitrogen fertilization. We confirmed for the first time that the oilseed rape genotype has a direct effect, not only through disease severity, on a seldom-informed trait of the fungal pathogen life cycle. The effect of host genotype on both the visual density of pseudothecia and the numbers of released ascospores had already been observed by Marcroft *et al.* (2004b). However, given the experimental design used, the authors could not clearly distinguish the effect of stem canker severity from the effect of host

genotype. Lô-Pelzer *et al.* (2009) followed by Bousset et al. (2019) detected the effect of host quantitative resistance and field on disease severity and on fruit body production. Yet, those studies were unable to disentangle the role of host genotype, cropping practices as well as the between-season variability. In our study, because all genotypes were planted in the same field and later incubated at the same place, we could exclude confusion with an effect of the environment. Furthermore, as stems were tracked individually, we were able to separate the effect of disease severity from the effect of genotype. One way forward would be assessing canker severity as a quantitative measure for each stem, without losing part of the information into severity classes, for instance by using image-based phenotyping. Then, it would be possible to assess the quantitative relationship between severity and fruit body production among genotypes.

The number of ascospore per pseudothecium ranged from 1000 to 10 000 (Lô-Pelzer *et al.* 2009) and 1000 to 14 000 (Schneider 2005). Estimates were based either on calculating the minimal and maximal volumes of pseudothecia considered as a sphere and asci considered as a spheroid, or on crushing 50 pseurothecia in water with bleach and counting the ascospores with a haemocytometer (Lô-Pelzer *et al.* 2009). Therefore, the numbers of ascospores available for release in cropping areas are huge. Nevertheless, clear gradients of decrease in leaf spot numbers on farmer's oilseed rape fields were observed with increasing distance from emitting spore sources (previous years' oilseed rape fields) (Bousset *et al.* 2015). Further, how far a source would spill ascospores depends on the number of emitted ascospores. This indicates that however large the number of ascospores is, decreasing their number by the means of genotype will contribute to decreasing disease transmission between successive years. In the panel of genotypes considered in the study, the host modulates the production of fruit bodies rather than suppressing it. Therefore, plant genotypes cannot prevent epidemics but may help to mitigate them. As stated later in the discussion, our findings can help identify conditions (i.e. arrangements of cultivars and fields) within a landscape allowing a durable control of the pathogen, which is a current challenge.

Differences in host phenology may contribute to the observed variability among host genotypes. From the biotrophic and asymptomatic presence of the fungus in the stem, visible cankers appear progressively when the crop matures. Moreover, the delay between infection and symptom appearance (i.e. asymptomatic phase) is known to vary between host-plants in a population, and the distribution of the incubation period could also change with some variables like the host-age when the infection occurs (Leclerc *et al.* 2014). When collecting disease data from genotypes differing in phenology, the sampling can occur at different stages in these processes, and thus introduce more variability in the relationship between the visual Phoma stem canker severity and the resulting fruit bodies produced. In oilseed rape stems, pathogen load, i.e. the amount of fungal mycelium, remains low throughout winter and starts to increase as *L. maculans* changes from biotrophic to necrotrophic after host flowering (Gervais *et al.* 2017). Nevertheless, in our study neither the time of flowering nor the time between flowering and harvest appeared as a major determinant of the genotype effect (Table S1.2). A more precise quantification of mycelium, by qPCR or by the visualisation of cankers through non-destructive imaging like MRI, could be helpful to better understand the mechanisms involved in the genotype effect.

Genotype effect could be due to a difference in the amount of nutrients available for the fungus. Indeed, in *P. infestans*, the most severely affected plants during the cropping season were the least prone to the intercrop survival of the pathogen on tubers. As quantitative host resistance reduces disease severity during the cropping season, it thus seems to promote intercrop survival (Pasco *et al.* 2016). However, in our study some genotypes may be severely infected but produce a moderate quantity of fruit bodies (Yu) while others may produce more fruit bodies (Da) than expected based on their Phoma stem canker severity class. Quantitative host resistance does affect inoculum production by decreasing disease severity, as number of pseudothecia on stems increase with Phoma stem canker severity (McGee & Emmett 1977; Marcroft *et al.* 2004a; Lô-Pelzer *et al.* 2009; Bousset *et al.* 2019). In our study, the genotypes Av, Cb, Da, Gr, Jn, Ko had higher than average levels of quantitative resistance on stem canker severity (Table S1.2; Fopa-Fomeju *et al.* 2015; Kumar *et al.*

2018), but they did not show any specific trend regarding fruit body production at a given severity class (Fig. 4; 5). Further, the nitrogen supply affected both disease severity at harvest and fruit body production, though the magnitude of the effect on fruit bodies was relatively small. It could be worth considering this for breeding cultivars for low-input systems. Genetic analyses are needed to identify the determinants underlying fruit body production in the genotypes. So far, plant resistance is still only characterised during the epidemics, e.g. in grapevine (Bove & Rossi 2020). However, pathogen life-cycle stages related to intercrop transmission, like the production of oospores in *Plasmopara viticola*, are starting to be evaluated (Delbac *et al.* 2019). In pathosystems such as downy mildew on grapevine or stem canker on oilseed rape, as host resistance affects pathogen transmission and survival, a low production of fruit bodies could be worth selecting in breeding schemes. At least, discarding the most fruit body-prone genetic backgrounds would be a step forward.

The highlighted effect of host genotype on inoculum carry-over may also be explained by a bias in mating of the fungal pathogen. Because the fungus is heterothallic, mating only occurs when the two mating types are present. A small canker with mycelia of both mating types is thus suitable for mating. In contrast, if a large canker is caused by only one individual, then no pseudothecia could be produced. The occurrence of Allee effects in fungi, i.e. reduced success of mating at low population density, has started to be investigated under controlled conditions for *Zymoseptoria tritici* (Suffert *et al.* 2018b) but should deserve further investigations. In the particular case of *L. maculans*, mycelial growth in the petiole has started to be investigated (Huang *et al.* 2019), but the precise location of the fungus in stem and its consequences on mating remains unknown. Further studies would be interesting and could rely on the use of GFP-transformed strains in controlled conditions or sequencing methods combined with population genetics analyses in natural monitored epidemics.

In agreement with previous findings (Lô-Pelzer *et al.* 2009; Bousset *et al.* 2019), our study confirmed the importance of between-year variability that may be related to variable environmental conditions. The highest fruit bodies numbers (Fig. 2) and rainfall during incubation (Fig. S1d) were

observed in year 13-14. In contrast, the lowest fruit bodies numbers and rainfall during incubation were observed in year 15-16. While the effects of climatic variables on maturation of the fruit bodies have been modelled (Aubertot *et al.* 2006), their influences on the pathogen during the cropping season and during the intercrop still deserve further investigations. In particular, besides influencing the development of the host plant, some main climatic variables such as temperature may drive within-host pathogen growth, influence pathogen load at stem base, thus influence mating of the fungus and therefore fruit body production. Linking environmental conditions with host and pathogen development is a current challenge for most pathosystems to improve predictions of epidemics and yields. In the particular case of oilseed rape stem canker, further studies are needed to address the influence of crop growth in real situations as well as intercrop practices of the production of fruit bodies (McCredden *et al.* 2017).

As plant genotype appeared to be an important driver of the production of inoculum, this should be taken into account in models used to compare strategies for the deployment of varieties in the landscape (Lô-Pelzer *et al.* 2010; Papaïx *et al.* 2018; Rimbaud *et al.* 2018; Watkinson-Powell *et al.* 2019). So far, the effect of qualitative host resistance on infection and some effects of quantitative host resistance on pathogen development are considered. Our results suggest that it would be worth considering the effect of host quantitative resistance on the pathogen during the intercrop. One striking result of our stochastic simulations of fruit body production is that the ranking of the genotypes (Fig. 7) appeared to be very different from the ranking obtained when considering the disease severity at harvest (Fig. 2). For example, the genotypes Ja, Pr and Yu with a high disease severity at harvest (Fig.2) produce fewer fruit bodies than others (Fig.7). In contrast, the genotypes Fa, Fr and Po have a moderate disease severity at harvest (Fig.2) but later produce higher amounts of fruit bodies that others (Fig.7). Recognition of this trade-off in pathogen resistance properties may help breeding varieties less prone to inoculum production and guide further breeding schemes.

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Authors' contributions

LB, PV and RD carried out experiments, ML carried out statistical analyses, LB, ML, NP and MP analysed pictures. LB conceived and designed the study and prepared the manuscript, read and approved by all authors. The authors declare the absence of conflict of interest.

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542 Zadoks JC, Schein RD, 1979. Epidemiology and Plant Disease Management, Oxford University Press, New York, NY, USA. 543 Zhan J, Thrall PH, Papaïx J, Xie L, Burdon JJ, 2015. Playing on a pathogen's weakness: Using 544 545 evolution to guide sustainable plant disease control strategies. Annual Review of 546 Phytopathology **53**: 19–43. 547 548 549 **Figure Legends** 550 Fig. 1. Schematic representation of cyclic epidemics and their control by plant resistance. (A) Cyclic 551 epidemics are characterized over years by the disease being present in the landscape with a stable 552 long term dynamics (black dashed line), but short term dynamics alternating between epidemic phase 553 (red lines) when the crop is present and decrease (blue lines) during the intercrop (Ic.). (B) Plant 554 genetic resistance can contribute to the control of cyclic epidemics during the crop with qualitative resistance reducing infection and quantitative resistance reducing the increase. During the intercrop, 555 556 the selective deployment of genotypes reduces landscape connectivity, and the plant genotype 557 influence the amount of inoculum produced (this study). 558 559 Fig. 2. Canker severity (G2 index) on field plots depending on the nitrogen level and the genotype. 560 The model was adjusted to the data of 12-13 to 15-16 years. Full names of genotypes and flowering 561 dates are available (Supplementary Information Table S1.2). 562 Fig. 3. Fraction of fruit body pixels depending on year of sampling (12-13, 13-14, 14-15 and 15-16) 563 for the fifteen genotypes in the Data12-16 dataset. Full names of genotypes are available 564 565 (Supplementary Information Table S1.2). 566

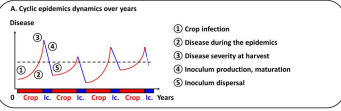
567 Fig. 4. Fraction of fruit body pixels depending for the S1 to S6 severity classes in the Data14-16 dataset. (A) For all genotypes; (B) to (H) per genotype for each of the six severity classes. Full 568 569 names of genotypes are available (Supplementary Information Table S1.2). 570 571 Fig. 5. Fraction of fruit body pixels depending on the severity before incubation exemplified for the 572 fifteen genotypes in the Data14-16 dataset. This result was observed for all datasets (Supplementary 573 Information Fig. S4.1; S4.2). Full names of genotypes are available (Supplementary Information 574 Table S1.2). 575 576 Fig. 6. Predicted fraction of fruit body pixels depending on the twenty one genotypes in the Data15-577 16 dataset for (A) normal (Nh) or (B) reduced (Nl) nitrogen fertilization of the crop. Full names of 578 genotypes are available (Supplementary Information Table S1.2). 579 Fig. 7 Boxplot of simulated numbers of fruit body pixels depending on the normal (Nh) or reduced 580 581 (NI) nitrogen level and on the genotype (21 genotypes). The effect of genotype was taken into account both for the distribution of stems in canker severity classes and on the prediction of fruit body pixels, 582 583 adjusting the model on observed data. Overall are values without the genotype effect. Data are means 584 of 10 simulated field plots with 100 plants of 700 000 pixels. Full names of genotypes are available 585 (Supplementary Information Table S1.2). 586 Fig. 8 Average simulated numbers of fruit body pixels depending on the canker severity at harvest 587 588 (G2 index calculated for the ensemble of stems) for the fifteen genotypes over all years. In 589 simulations, the effect of genotype was taken into account both for the distribution of stems in canker 590 severity classes and on the prediction of fruit body pixels, adjusting the model on observed data. Data 591 are means of 10 simulated field plots with 100 plants of 700 000 pixels. Among the genotypes, either

susceptible (high canker severity) or with higher levels of quantitative resistance (low canker

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- severity), the pixels of fruit bodies are variable. A very susceptible genotype (Yu) is not the one
- leaving the greatest amount of fruit bodies.
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- 596

Fig 1



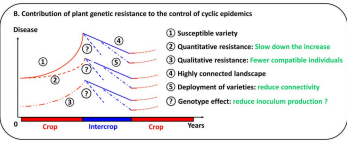
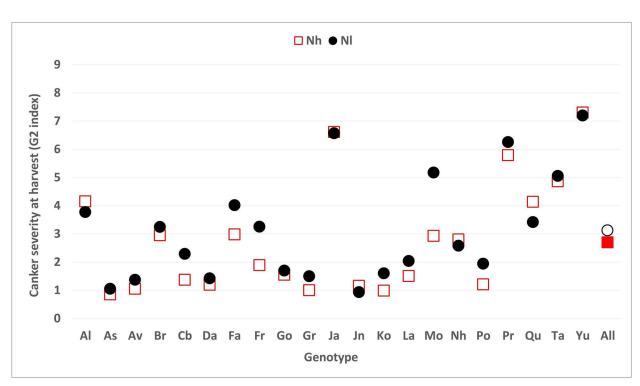
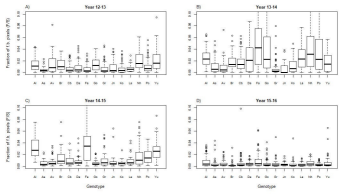
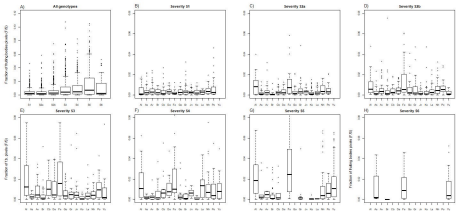
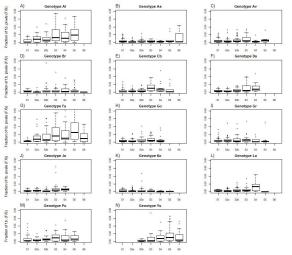


Fig 2









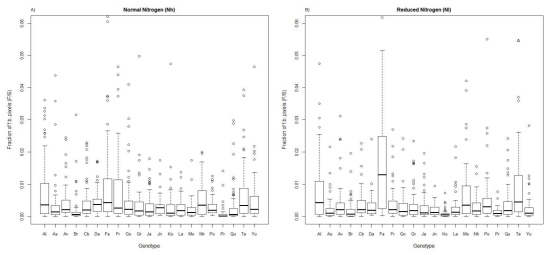


Fig 7

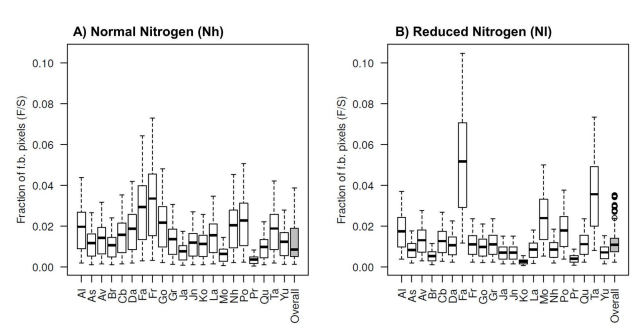


Fig 8

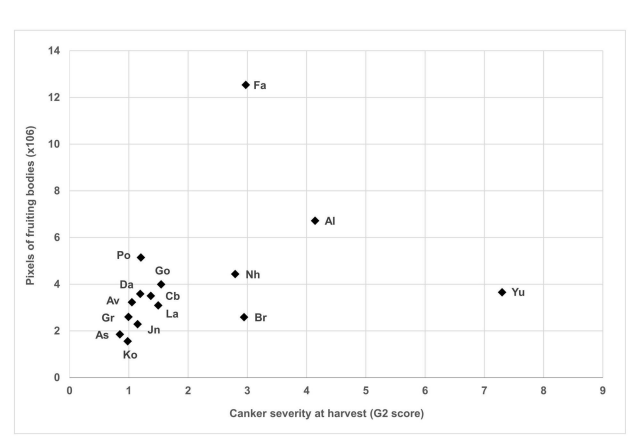


Table 1. Numbers of stems for the complete and the three sub datasets (Data14-16, Data15-16) given overall (italicised), but also by year and nitrogen fertilization treatment (Nl and Nh) and by genotype.

Dataset	Year	Nitrogen	Post process			Genoty	pe																			
			Before	Discard	After	Al	As	Av	Br	Cb	Da	Fa	Go	Gr	Jn	Ко	La	Nh	Po	Yu	Fr	Ja	Мо	Pr	Qu	Та
Complete	All		6758	18.3%	5525	390	318	366	325	356	286	338	290	348	247	291	320	363	291	273	114	133	137	76	181	82
	12-13	Nh	938	36	902	96	50	70	58	67	64	56	49	54	60	58	44	77	60	39	-	-	-	-	-	-
	13-14	Nh	919	48	871	49	59	75	52	88	48	73	42	65	12	31	80	86	70	41	-	-	-	-	-	-
	14-15	Nh	826	222	604	72	41	37	15	37	40	57	40	40	43	27	29	57	35	34	-	-	-	-	-	-
	14-15	NI	789	318	471	38	31	30	47	23	25	39	23	32	24	27	36	24	23	49	-	-	-	-	-	-
	15-16	Nh	1626	240	1386	67	78	77	77	73	57	61	66	71	44	78	71	76	47	59	48	68	91	35	96	46
	15-16	NI	1660	369	1291	68	59	77	76	68	52	52	70	86	64	70	60	43	56	51	66	65	46	41	85	36
Data1416	All		4045	10.1%	3029	245	209	221	215	201	174	209	199	229	175	202	196	200	161	193	-	-	-	-	-	-
	14-15	Nh	826	222	604	72	41	37	15	37	40	57	40	40	43	27	29	57	35	34	-	-	-	-	-	-
	14-15	NI	789	318	471	38	31	30	47	23	25	39	23	32	24	27	36	24	23	49	-	-	-	-	-	-
	15-16	Nh	1208	206	1002	67	78	77	77	73	57	61	66	71	44	78	71	76	47	59	-	-	-	-	-	-
	15-16	NI	1222	270	952	68	59	77	76	68	52	52	70	86	64	70	60	43	56	51	-	-	-	-	-	-
Data1216	All		3685	3.1%	3379	284	228	259	202	265	209	247	197	230	159	194	224	296	212	173	-	-	-	-	-	-
	12-13	Nh	938	36	902	96	50	70	58	67	64	56	49	54	60	58	44	77	60	39	-	-	-	-	-	-
	13-14	Nh	919	48	871	49	59	75	52	88	48	73	42	65	12	31	80	86	70	41	-	-	-	-	-	-
	14-15	Nh	826	222	604	72	41	37	15	37	40	57	40	40	43	27	29	57	35	34	-	-	-	-	-	-
	15-16	Nh	1002	206	1002	67	78	77	77	73	57	61	66	71	44	78	71	76	47	59	-	-	-	-	-	-
Data1516	All		3286	6.1%	2677	135	137	154	153	141	109	113	136	157	108	148	131	119	103	110	114	133	137	76	181	82
	15-16	Nh	1626	240	1386	67	78	77	77	73	57	61	66	71	44	78	71	76	47	59	48	68	91	35	96	46
	15-16	NI	1660	369	1291	68	59	77	76	68	52	52	70	86	64	70	60	43	56	51	66	65	46	41	85	36

 Table 2. Analysis of deviance for the observed distribution of canker severities at harvest

Factor	Df	χ2	P-value
Year	3	7742	< 2.2e-16
Nitrogen	1	211	< 2.2e-16
Genotype	20	8828	< 2.2e-16
Nitrogen × Genotype	20	1971	< 2.2e-16

Table 3: Analysis of deviance for the three datasets, with percentage of deviance explained by each factor and by the model

	Analysis of deviance				Model
Dataset	Factor	Df	P-value	Explained deviance	R ²
All	Year	3	<0.05	0.20	0.34
	Genotype	20	<0.05	0.061	
	Severity	6	<0.05	0.069	
	Nitrogen	1	<0.05	<0.001	
	Nitrogen X Genotype	20	<0.05	0.014	
	Nitrogen X Severity	6	<0.05	<0.001	
Data14-16	Nitrogen	1	<0.05	0.01	0.51
	Year	1	<0.05	0.20	
	Genotype	14	<0.05	0.17	
	Severity	6	<0.05	0.08	
	Nitrogen X Year	1	<0.05	0.0002	
	Nitrogen X Genotype	14	<0.05	0.003	
	Year X Genotype	14	<0.05	0.01	
	Year X Severity	6	<0.05	0.01	
	Nitrogen X Severity	6	<0.05	0.01	
	Nitrogen X Year X Genotype	14	<0.05	0.02	
	Nitrogen X Year X Severity	6	<0.05	0.002	
Data12-16	Year	3	<0.05	0.235	0.52
	Genotype	14	<0.05	0.115	
	Severity	6	<0.05	0.096	
	Year X Genotype	42	<0.05	0.062	
	Year X Severity	18	<0.05	0.011	
Data15-16	Nitrogen	1	<0.05	0.0003	0.28
	Genotype	20	<0.05	0.17	
	Severity	6	<0.05	0.05	
	Nitrogen X Genotype	20	<0.05	0.05	
	Nitrogen X Severity	6	<0.05	0.009	