Influence of Airborne and Seed Inoculum in the Initiation of Leaf, Stem, and Root Infection by Systemic *Botrytis cinerea* in Lettuce (*Lactuca sativa*)

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**Abstract**

Studies showed that both airborne seed infection of *B. cinerea* have been implied as major sources of primary inoculum in different part of lettuce plant. Results from quantitative real-time PCR and visual observations showed that seed infection was the main source of inoculum in the field trial conducted in this study. However, glasshouse studies established that the pathogen can be transmitted from infected seeds into roots, shoots and leaves without showing symptoms. Plants in the field trial remained free of infection for approximately four months before symptoms were observed in the crop. Covering the crop during part of the growing season was found to prevent the growth of the pathogen, despite the use of infected seed, which indicate that changes in the physiological condition of the plant and/or environmental conditions most especially over watering may trigger disease development. However, once the disease symptoms appeared in the field it quickly spread throughout the plants. Level of inoculum equivalent to spore concentrations between 30 and 60 spores per m$^2$ of air was only detected on three occasions during the growing season. The temporal pattern and level of detection of *B. cinerea* DNA in spore tape samples indicated that airborne inoculum was limited and most likely represented rain-splashed conidia rather than putative ascospores.

**Keywords:** *B. cinerea*; Lettuce plant; Real-time PCR; Seed infection; Spore trap

**Introduction**

Lettuce *Lactuca sativa* (Asteraceae; Compositae) is one of the World’s most important salad crops and horticulturists regard it as one of the most profitable to grow [1]. Leaf, stem, root and seed infections caused by systemic *B. cinerea* are common in lettuce grown in the whole of the UK [2], Infection of lettuce plants by *B. cinerea* is favoured by cool moist weather conditions [3,4]. However, the levels of disease in crops can alter from year to year, with the cost of losses in winter lettuce estimated at £257 million due to *B. cinerea* infection alone in fungicide-treated crops [5]. Initial infection can result from infected lettuce stubble and/or crop debris on the soil [6,7] and most commonly occurs when successive lettuce crops have been grown in the same field. Symptom development in the fields with no previous record of lettuce cropping can arise from the sowing of infected seed or by infection from air-dispersed spores. Once infection is established in a crop, the infected lettuce plants show abundant white, cottony mycelial growth and hard, black sclerotic bodies on the underside of the lower leaves and the basal part of the stem [4]. The head and stem of infected plants often disintegrates into a soft watery mass covered with sclerotia [8].

It has been reported by [1,2,9] that *B. cinerea* infection was seed-borne, and it is thought that seed infection could be extremely important for the spread of *B. cinerea* into new areas with no record of lettuce cultivation [10,11]. In another study Shafia [12] found that the efficacy of disease transmission on seeds was dependent on the rate of seed germination and temperature, [1,10,13] showed that root, stem, leaf infection could develop from symptomless seeds. The symptoms of *B. cinerea* infection generally occur and can be identified by the presence of dense gray or light brown airborne spores (ascospores or conidia), which may require high humidity and free water for infection to occur [8,14,15]. Recent work using DNA-based markers has confirmed that *B. cinerea* populations are highly genetically variable, with the majority of variation distributed within individual fields [16,17]. These findings agree with the results of Ma and Michailides, Martin [18] who reported moderate genetic variation within *B. cinerea* isolates collected from fig plants in the same location in California however, differences between different hosts. Similarly, this was similarly Newton [19] reported that population of *B. cinerea* from different host’s species (Strawberry, dandelion, black current, primula) sampled from different locations of the UK were genetically variable and most importantly highly variable within host plants. The objective of this study therefore, was to determine the role of seeds and airborne inoculum in the initiation of *B. cinerea* infection in leaf, stem, root and seed infection of lettuce plant in a field experiment where crop debris was excluded as a potential inoculum source. Covering the crop until disease symptoms appeared in the remaining plots was to test the hypothesis that conidia and ascospore release from plant material in autumn and winter could initiate leaf and stem infection in lettuce plant.

**Materials and Methods**

**Field trial**

The area designated for the research has been cropped with lettuce for the last five years, having been fallow for the last two years. The Nearest lettuce stubble field was at least 0.8 km away and the plots were about 500 m from grassland. The whole field measured 33 × 14 m and contained 12 small plots (3 × 3 m). The lettuce plots were either sown with ‘clean’ seed from Fothergills Seeds, New market, UK or from ‘infected’ seed harvested in the previous experiment from untreated plots at School of Biological Science, Reading. The certified lettuce seeds supplied by Fothergills Seeds, New market had been assessed

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experimentally and were found to be free of disease symptoms, whereas most of the infected seeds showed symptoms of *B. cinerea* both on the surface and interior of the seeds. All seeds were sown without the use of seed treatments on 2 January 2010.

Specially designed tents made from aluminium tubing and acetate clear plastics were placed in the centre of selected plots to exclude airborne inoculum [1]. By covering the crop, the effect of excluding conidia and airborne spores as an inoculum source, which is an issue in other closely related species such as light leaf spot of oilseed rape (*Pyrenopeziza brassicae*), could be tested in the period of the growing season when ascospores release is anticipated. The covers were later removed to determine whether the clean lettuce plants which had been excluded from airborne inoculum could subsequently become infected. The use of *Arabidopsis* plants between the plots was to help minimise the spread of inoculum from plot to plot via rain splash. The mini tents were secured in the soil to a depth of 250 mm, covering all the plants in the centre square metre of each plot in a ‘miniature greenhouse’. All the tents were thoroughly and regularly checked and any hole noticed was repaired using parcel tape until the development of disease symptoms, thereafter all the tents were removed from the trial plots. None of the plots in the field experiment received any fungicide treatments thereafter all the tents were removed from the trial plots. None of the plots in the field experiment received any fungicide treatments throughout the growing season, however, all other inputs were made according to standard farm practice with a spring application of 167 kg ha one of ammoniacal nitrogen 18.4%, nitric N 11.6% and water-soluble sulphur (SO3) 19.0% (Sulphur Gold, TERRA Nitrogen UK Ltd).

### Field trial sampling and visual assessments

Assessments in the field were carried out every week. The study used conventional visual plant pathological techniques, in combination with spore trapping and quantitative real-time PCR which was developed previously [20] to check for the development of disease symptoms, until the time that they first appeared in multiple plots, symptoms were visible in all uncovered plots sown with infected seeds. All tents were then removed and plants were sampled from each of the nine, 1 m² subplots, contained in all 12 of the main plots. The visual assessments for *B. cinerea* in leaves and stems were carried out by counting the number of lesions on whole plants, and the plant growth stages were also recorded for each subset of 10 plants sampled. After assessment of the disease, all the plants were discarded except for the 10 plant samples from the central 1 m² subplot. Plants from this subplot were then further dissected, were each of the plant was separated into roots, white stems (colourless and located below the surface of the soil), green stems (found above the surface of the soil) and leaves (10 of each for each plant section, from three replicates). These were then frozen for future DNA extraction and PCR testing. The same procedure was used for the samples taken on 24 April 2010. Also visual assessments were performed on 10 plants per plot each month throughout the growing season.

### Trapping of airborne inoculum

A seven-day recording volumetric spore trap Burkard (Burkard Manufacturing Co. Ltd.) was set up in the central strip of the field immediately after the crop was sown *Arabidopsis*. A 3° radius directly surrounding the spore trap was kept clear to decrease rain splash-dispersed spores being sucked into the trap. The spores were collected on Melinex polyester film tape (Burkard Manufacturing) coated with a mixture of paraffin wax and petroleum jelly. The tape was replaced every seven days. For analysis, the tape was cut into 48 mm sections, each representing a 24 h period.

DNA extraction and quantification of *B. cinerea* DNA using real-time PCR

DNA was extracted from all sampled plant and fungal material using the method of [1,21] except that the DNA extraction buffer was amended with 5 mM 1,10-phenanthroline monohydrate and 2% (wt/ vol) polyvinylpyrrolidone K30 (Sigma-Aldrich Chemie GmbH) to clean the DNA [22]. The DNA was quantified using the fluorescent dye thiazole orange (Sigma-Aldrich) as described previously [21]. Spore tape DNA extractions were done using Ballotini beads (Jencons Ltd.) in a Fast Prep machine (Savant Instruments) as described by Calderon et al. [23]. DNA was extracted from half of each 24-h tape sample and 1 lL from 200 lL final DNA solution was used in each PCR assay. Quantitative real-time PCR measurements were carried out in a Strata gene Mx3000P real-time PCR machine (Stratagene, La Jolla, California, USA) with the cytochrome b Locked Nucleic Acid (LNA) probe assay recently developed [20].

### Determination of *B. cinerea* seed infection and its transmission in controlled environment

Seeds from the two seed batches used in the field experiment were visually assessed and tested by real-time PCR to establish differences in seed infection levels and to determine whether symptomless seeds were infected with *B. cinerea*. Five seeds with and without symptoms were randomly selected on the grain surface of each seed batch and analysed. For the controlled environment study, five hundred seeds from each seed batch were sampled. Each seed was thoroughly examined with a hand lens and were separated into seed lots with and without symptoms. Symptomless seeds from the ‘clean’ seed batch and seeds with symptoms from the infected seed batch were sown into autoclaved compost (John Innes, No 2) in propagator trays (50 × 20 cm) and incubated at separate controlled environment cabinets (Fisons 600H). The light intensity used was 200 l Einsteinm)2 s)1 PAR. After germination, the seedlings in the trays were watered daily, so that the soil remained damp regardless of the incubation temperature. Seedlings were sampled at growth stage 8–9 (coleoptiles breaking through soil surface) and at 14–15 (4 or 5 leaves unfolded). The samples were collected on different sampling dates, as the growth of the seedlings differ.

The basic aluminium frame of the tent manufactured from 2.5 cm diameter tubing. Ten seedlings were sampled at each growth stage using ethanol-cleaned forceps. Seedlings were dissected into lower root, seed, upper root and above soil stem/leaf sections and *B. cinerea* infection levels determined using real-time PCR. Microscopy was carried out according to the methods developed [1,19] to examine the plant material for fungal structures of *B. cinerea*.

### Data analysis

The visual observations and quantitative real-time PCR data sets were transformed using natural logarithms, ln (×+0.5) for visual data and ln (×+0.1) for PCR data, to normalize the residuals from fitted models. Results was examined with a 2-way analysis of variance (ANOVA) using Genstat, 14th edition, (VSN International, Ltd), with plant, plot and block as random factors. The weather data for the 2009-2010 growing season were obtained from the Meteorological Database.

### Results

#### Effect of seed source and covering of plants on leaf infection

The first symptom of leaf blotch, a single lesion in one of the plots drilled with infected seed, was seen on 3 February 2010. The tents were removed two weeks later when all uncovered plots sown with farm-
saved infected seed showed symptoms. Plants from farm-saved seed had an average of 5.6 lesions per plant compared to the symptomless clean seed, which had only 0.18 lesions per plant (Table 1). However, covering the crop significantly reduced the formation of symptoms; at the time the tents were removed, there was no symptoms in any of the covered areas, whether drilled with symptomless infected or uninfected seed. Disease was recorded in seedlings surrounding the tents in the same plots (data not shown). However, after removal of the tents, disease levels in these previously covered areas remained lower than in plots that had not been covered. While in the covered plots, disease levels in plots drilled with uninfected clean seed remained much lower than in those drilled with farm-saved infected seed, and the same effect was also seen for the plots that were not covered. Results obtained from the ANOVA on all plots found that tenting greatly reduced symptoms of B. cinerea (P < 0.001) and plots sown with uninfected clean seed had less disease than plots sown with infected seed (P < 0.001). Statistically, the interaction was significant (P < 0.001) because the covered plants were free of symptoms regardless of seed source. The difference between the seed types and the effect of covering the crop with tents both remained highly significant (P < 0.001) for up to two months after the removal of the tents. However, four months after removal of the tents disease severity levels were relatively closer. The main effect differences were highly significant (P = 0.006 for seed source, P < 0.001 for tenting) and the interaction was not significant (P = 0.4).

Detection of B. cinerea in plant samples using real-time PCR

All plant materials roots, stem and leaves sampled from plots which were not covered, had detectable amounts of B. cinerea DNA in real-time PCR throughout the trial. However, plant samples from uncovered plots sown with an infected seeds had higher levels of B. cinerea than those collected from uncovered plots sown with uninfected clean seeds. The roots sampled from uncovered plots sown with an infected seeds contained around 6 pg B. cinerea DNA, compared to around 0.6 pg for the uninfected seeds. The PCR results from the February samples showed that no B. cinerea DNA could be detected in samples from plots sown with either the infected seed (plots 4, 7 and 11) or symptomless uninfected seeds when grown undercover. Nearly all plant parts sampled from all plots contained B. cinerea DNA April 2010. The highest levels of B. cinerea DNA were mostly found in the leaves and decreased towards the roots. B. cinerea DNA levels were lower in plots sown with uninfected seed (P < 0.001) and in covered plots (P < 0.001). The interaction between seed and cover was significant, showing that covering prevented disease development even when the seed was heavily infected with B. cinerea (P < 0.001). The differences in B. cinerea DNA levels between the different plant parts (roots, seeds, stems and leaves) were also significant (P < 0.001) as was the interaction between plant parts and cover (P < 0.001). Patterns and levels of significance were similar on both sampling dates (Table 1).

Seed infection and transmission of disease in controlled environment

Seeds with symptoms had approximately twice as much B. cinerea DNA as randomly-selected seeds from the same batch (Table 2). Most seeds without symptoms tested positive for presence of B. cinerea in real-time PCR. However, the average infection level of seeds sampled from the symptomless clean seed batch was approximately 10-fold lower than seeds sampled from the farm-saved infected seed batch, irrespective of the sampling method.

Most parts of plants produced from symptomless uninfected seed tested negative for B. cinerea DNA at all stages of growth and temperatures tested (data not shown). As expected, more samples were positive for B. cinerea infection when infected seeds were sown 6). The highest levels of pathogen DNA was detected after growth at16. C. At this temperature, DNA of B. cinerea was detected in all plant parts sampled at growth stage 8–9 and 14–15. None of the plants grown in the controlled environment showed any symptoms of leaf infection at growth stage 14–15. No spores or hyphal / mycelium structures resembling B. cinerea was seen in any of the samples examined by light microscopy.

Discussion

The emergence of the lettuce seedling under the covers was quicker

| Seed          | Tent cover | Leaf gray mold1 | Leaf gray mold2 |
|---------------|------------|-----------------|-----------------|
| Uninfected seed | open       | 0.19            | 53              |
| Infected seed | Tented     | 0.95            |                 |
| cv of mean 3  | Tented     | 0.51            | 13%             |
| cv of mean 1  | Tented     | 0.34            | 22%             |

1Disease was expressed as number of lesions per plant. It was analysed on log scale: main effects and interaction significant (P < 0.001, df 1, 8). Block and plot variances not significantly larger than plant variance. B. cinerea.

2Disease was expressed as number of lesions per plant. It was analysed on log scale: main effects of seed source, P < 0.001 (1.8 df), tenting P < 0.001. Interaction P = 0.4. Blocks did not differ more than expected from plot-plot variation, but this was significantly more than plant-plant variation (P < 0.001; 6108 df).

3The coefficient of variation from sem on log scale; the variability applies to any mean in the column closer, the main effect differences were highly significant (P < 0.006 for seed source, P < 0.001 for tenting) and the interaction was no longer significant (P = 0.4).

| Seed lot | sampling method1 | visual symptom2 | pathogen DNA3 (Pg) per sample |
|----------|------------------|-----------------|-----------------------------|
| Uninfected seed | At Random        | 0               | 0.18                        |
|           | Selective        | 1.2             | 0.26                        |
| Infected seed | At Random        | 0.6             | 1.53                        |
|           | Selective        | 1.2             | 3.0                         |
| SED or Proportionate error In each mean | 0.26 | 34% |

1Seeds that were randomly selected or selected for symptoms of B. cinerea infection.

2Symptom visual scores: 0; no symptoms; 1; flecking symptoms; 2; lesions visible. Main effect of sampling method, P < 0.001; seed lot and interaction P > 0.1, not significant.

3Back-transformed mean; analysis on log scale (equivalent to the measurement scale). ANOVA: main effect of seed lot, P < 0.001; sampling method and interaction P > 0.2, not significant.

Table 2: Level of B. cinerea in clean (symptomless) or infected B. cinerea seed of various origins, determined by visual assessments and quantitative real-time PCR.
by about one week because of the warmer climate and thus the plants grow at a faster rate throughout the growing season. On the day that the tents were removed (growth stage 23–24), there were clear effects of both covering and seed batches on the epidemic development of *B. cinerea* (Table 1). Plants in symptomless open plots sown with uninfected seeds had significantly less disease than open plots sown with infected seed. These results were supported by the real-time PCR data. All plant parts (roots, stems and leaves) sampled from plots sown with infected seeds showed higher levels of *B. cinerea* DNA than equivalent samples taken from plots sown with uninfected seeds. Detection of *B. cinerea* in all of the plant parts sampled indicates a systemic infection spreading from the seed. However, the greater infection levels were always found in infected uncovered seed and these caused greater disease levels throughout the growing season when compared to all the other seed types and conditions. *Botrytis cinerea* was initially not detected in plants sampled from covered subplots, irrespective of seed source for reasons which are unclear. Suppression of disease development in the covered crop may have been due to the differences in soil water potential which could have influenced rhizosphere bacterial communities involved in disease suppression. Another possible explanation may be the environmental conditions for the covered crop may not have been conducive for the fungal growth as it may have been too warm and dry, but ideal for greater plant growth enabling disease escape. Also Shafia [12] demonstrated that the rate of germination of a seedling could prevent the infection occurring at increased temperature higher than the optimal 16_C by causing the coleoptiles to grow away more rapidly from the infected area of the seed. Winter lettuce is now drilled as early as possible in the UK, when the soil temperature will be higher and therefore this should reduce the effective transmission of *B. cinerea* from seed to seedlings.

However, during this experiment no data for the environmental impact of the covers were taken so it is not possible to determine what factors caused the lack of both visual disease and positive PCR results for samples taken from under covers in February from both infected and uninfected seed. Four weeks after the removal of the tents, *B. cinerea* DNA was detected in plant samples from all plots and this was eventually followed by leaf gray mold symptoms (Table 1). This may be due to a delayed spread of inoculum from the seed following a change in the environment, because highest levels of pathogen DNA were still recorded for plant parts (including roots) sampled from plots sown with the infected seed. However, it is also possible that infection may have developed from the outside plots following the removal of the covers and these plants could have been more susceptible to symptom development due to their earlier 'softer' growth environment containing elevated temperatures and humidity and reduced amounts of other outside factors such as wind and frost. The development of a similar disease pattern following the removal of the covers is probably the results of conidia spreading across all the plots with the exception of the infected uncovered seed plots which has initially higher pathogen levels because of the role of seed infection detection threshold were detected in spore tape samples from the end of January until the end of July 2010 with the occasional higher level of up to 1.9 pg of pathogen DNA. This pattern is not typical of other ascospore-producing fungi present in other arable crops such as *Mycosphaerella graminicola*, *Pyrenopeziza brassicae* and *Oculimacula yallundae*, which all tend to produce ascospores more abundantly from senesced plants and/or stubble during late summer to early winter. Moreover, these pathogens also produce some spores during late spring to summer [1,23,24]. The period in which positive tape samples were detected coincided largely with the period that symptoms were visible on the leaves and not during late summer to early winter. Because of the low level of detection in comparison with other ascospore producing pathogens, it is likely that the *B. cinerea* DNA detected originated from ascospores, rather than from ascospores. Therefore, it was suggested that sexual recombination events may be common in populations of *B. cinerea* and this may therefore, be the major reason for its epidemiology and adaptation to the environment [17,25,26]. Asexual spores of *B. cinerea* are relatively small in comparison with ascospores of other fungi and therefore during humid conditions, water droplets containing spores could have been sucked directly into the spore trap. It was not possible to identify ascospores of *B. cinerea* by examining the spore tapes by microscopy since the morphology of the telephor is unknown. In order to identify ascospores in positive samples, work now needs to be carried out to develop an immunofluorescence microscopy assay using specific antibodies or in situ PCR to positively identify the source of fungal material that is causing a positive result for spore tape DNA samples.

In the controlled environment study however, no root, stem or leaf symptoms were seen throughout the experiment. But, *B. cinerea* DNA was detected in seedlings from both infected and uninfected seeds at all temperatures used. Out of the temperatures tested, 16_C was optimal for *B. cinerea*, confirming earlier work [12]. However, [11] using competitive PCR demonstrated that visual assessments poorly corroborate with levels of *B. cinerea* DNA from the same seed.

Plant roots were colonized by *B. cinerea* in both the controlled environment and the field experiments. There was as previous report of *B. cinerea* infection on the root system, and affected plants had reduced root length [27]. In 2003 [28] isolated *B. cinerea* from the roots of Primula plant, and recent work [29] has suggested that soil borne inoculum and root infection could be important in the development of rice blast disease, caused by *Magnaporthe grisea*. This fungus had previously been regarded as a foliar pathogen however, [29] were able to show that root colonization can lead to systemic invasion and development of disease on aerial parts of the plant. Unfortunately, no infection structure of *B. cinerea* was seen by light microscopy. Understanding which stages/structures of *B. cinerea* are important for the infection process and its life cycle could be investigated by green fluorescent protein technology to visualize by confocal microscopy, fungal development in plant [30]. The role of seed infection as primary inoculum for leaf and stem gray mold disease of *B. cinerea* in winter lettuce crops in the UK has probably been over looked because of the long latency of the disease. It is clear that *B. cinerea* can be transmitted from seeds, but may produce no symptoms in the plant for several months (latent infection), and currently almost all of the commercial seed treatments used by growers are not active against *B. cinerea* infection contained in the seed. But changes in the physiological condition of the plant and/or environmental conditions may trigger disease development. Seed borne infection may also contribute to the genetic variation of *B. cinerea* populations as batches of seed are often combined and travel for long distances before being sown. In addition to using resistant lettuce varieties and foliar fungicide applications, the testing of seed batches for *B. cinerea* by PCR, with rejection of severely contaminated batches, or additional seed treatments with fungicides might improve disease control. In addition to this more work is needed to establish what levels of seed infection can be tolerated before rejection should occur as most of the clean symptomless seeds used in this study were later found to be infected. Seed batches can also be tested for the presence of alleles conferring resistance to fungicides. Mutations in cytochrome b and b-tubulin encoding genes, targets for the quinone outside inhibitors (QoI) and methyl benzimidazole carbamate (MBC) fungicides, respectively, have been identified in populations of *B. cinerea* [31]. Seed infection may also be a reason behind the
introduction of populations of B. cinerea, resistant to compounds such as QoI fungicides. The quinone outside inhibitors fungicide resistance that was developed in France during the summer of 2008 [32] may pose a greater risk to the UK growers if seed is imported from an infected area. Therefore, on the final note routine monitoring of seed and other source of B. cinerea inoculum using PCR should become vital for the long term viability of lettuce cropping in the UK.

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