ABSTRACT: The cultivated area of sunflower in Brazil is expanding considerably over the last years as the result of the great interest in the biodiesel production derived from that crop. One consequence of that expansion is the occurrence of several diseases some of devastating nature in part of the growing areas with sunflower. This study evaluated the effects of the association of sunflower seeds infected by *Sclerotinia sclerotiorum* and the pathogen transmission rates from these infected seeds, considering some factors which may interfere in that interaction. Four inoculum potentials, two isolates of the cited fungus, two sunflower cultivars, 'Helio 250' and 'Helio 253' and two environmental temperatures, 20°C and 25°C, were used for the purpose in this work. Seed germination and health, emergence speed index, and plant populations were variables analyzed. From the results, increased levels of inoculum potentials led to gradual reduction of the mean values of germination, emergence rate index and plant populations, regardless the genotype, fungal isolates and temperatures. Transmission rates were higher at the highest levels of inoculum potential, the maximum reaching 80%. These results show the significance of *S. sclerotiorum* inoculum associated with sunflower seeds both in the establishment of plants in fields and also in spreading the pathogen inoculum between crop fields.

KEYWORDS: *Helianthus annuus*. Physiologic conditioning. Transmission. White mold.
MATERIAL AND METHODS

Obtaining and multiplication of fungal isolates
Two isolates of *S. sclerotiorum* used in this study were obtained from sunflower seeds, collected in Lavras, State of Minas Gerais MG (CMLAPS 250) and Montividiu – State of Goiás (CMLAPS 423), both belonging to the mycological collection of the Seed Pathology Laboratory of the Federal University of Lavras. The isolates were grown initially on PDA media plates and incubated at 20 ± 2°C, with a photoperiod of 12 hours for five days.

Seed inoculation procedures
Seeds of two sunflower cultivars, 'Helio 250' (C1) and 'Helio 253' (C2), were disinfested with sodium hypochlorite 1% for 30 seconds, rinsed in distilled water and then left to dry for 48 hours in a laminar flow hood, on germitest paper. The seed inoculation was done by the osmotic technique developed by Costa et al. (2003) and Machado et al. (2012). By that technique, PDA substrate modified osmotically by the solute mannitol, at water potential of -1.4 MPa adjusted by software MPPS (MICHEL; RADCLIFFE, 1995) was poured to Petri dishes of 15 cm diameter. Mycelial plugs of *S. sclerotiorum* isolates were placed on the media and then kept in an incubation chamber at 20°C and photoperiod of 12 hours for five days. On the top of the fungal colonies disinfested seeds were distributed in a single layer, making sure all seeds were placed in contact with the fungal mycelium. The dishes were then transferred to incubation chamber at 20°C for the periods: 24h (P1), 48h (P2), 72h (P3) and 96h (P4) hours, each period standing for one inoculum potential of the pathogen. A control was prepared for each incubation period using the same seed substrate with mannitol in the absence of *S. sclerotiorum*.

Assessing the quality of sunflower seeds infected by *Sclerotinia sclerotiorum* in different levels of inoculum potential
The health condition and physiological performance of the infected and non infected seeds were evaluated according to protocols described in Brazil (2009b). The seed health test consisted in incubation of seeds on a semi selective-bromophenol agar substrate (NEON), as widely described in literature (NASSER et al., 1999; PERES et al., 2002; NAPOLEÃO et al., 2006), using a total of 200 seeds of control and inoculated seeds for each inoculum potential. The plates were distributed in incubation chamber at 20°C. At the third and fifth days of incubation, the seeds were examined for the formation of yellow-red halos around seeds, which is an indication of the presence of *S. sclerotiorum* in them. The plates were also analyzed in stereo microscope for mycelium growth and presence of sclerotia of that fungus (BRAZIL, 2009b).

Standard germination test was run by the roll paper method (BRAZIL, 2009b) in which 200 seeds per treatment were used with incubation of seeds in germination cabinet with temperature adjusted to 25 ± 2°C. Evaluation was made on the fifth and ninth day, according to Brazil, 2009a.

To evaluate the emergence speed index, initial and final stands, seeds were sown in 300 mL plastic cups containing a 2:1 mixture of sand and substrate (Tropstrato HA Hortaliças). For each cultivar, 100 inoculated seeds and 100 of non inoculated seeds were sown, each seed being planted in single plastic cups, of 300 mL which were placed together in polyethylene boxes for a total of four boxes per cultivar and inoculums potential. The experiment was conducted in a growth rooms at 20 and 25°C. The seedling/plant emergence rate was determined by daily counts of seedlings until the stabilization of plant population. The ratio of emergence rate was calculated according to formula described by Maguire (1962).

Determination of potential transmission rate of *Sclerotinia sclerotiorum* from sunflower seeds to plants
The total transmission rate was estimated by the sum of the number of seeds/seedlings killed in pre-emergence as the result of the pathogen action plus the number of emerged plants with symptomatic infection and asymptomatic disease (Baker and Smith, 1966; Bergstrom and Shah, 2000). Plants were considered symptomatic when presenting wilt followed by necrosis or damping-off. In V3 growth stage (15 days after sowing – DAS), one of the cotyledons of each plant was collected and, at 25 DAS, each plant was sectioned in three parts (cotyledon/stem insertion, apex and root crown). The evaluation was carried out by the destructive method, collecting the four fragments, 1.5 cm size, of each plant. All sections were disinfested in 70% alcohol sodium hypochlorite and 1% sterile distilled water for 1 min. After drying on paper towels, fragments of the same plant were distributed on semi-selective agar bromophenol (Neon) medium in Petri dishes and kept in an incubation at 20°C for seven days. After this period, the occurrence of mycelial growth from each plant was observed by the color change of the substrate and, when necessary, fungal structures were confirmed with the aid of light microscopy at 80X.
magnification. The transmission of the pathogen to the emerged plant and dead non emerged seedlings was considered positive when at least one fragment of the symptomatic or asymptomatic plant presented mycelial growth of *S. sclerotiorum*.

**Data analysis**

For germination and incidence were used 50 seeds for each cultivar, isolate and inoculum potential, in a total of 200 seeds per treatment. For emergence speed index, initial and final stands 25 seeds for each cultivar, isolate, mannitol and inoculum potential in a total of 100 seeds per treatment were used. The experimental design was complete randomized with four replications. Statistical analysis of variance (ANOVA) was performed by R statistical software v 2.15.1. Data were analyzed using Tukey test and P-value less than 0.05 was considered significant. Analysis was conducted in double factorial among cultivars and isolates (I 250, I 423 and mannitol) x inoculum potential (P1, P2, P3 and P4) in each temperature (20 and 25°C). The graphics were prepared using the Sigma Plot 12.1 program (Systat Software Inc.).

**RESULTS**

**Relationship between inoculum potential and quality of sunflower seeds infected by *S. sclerotiorum***

The initial germination percentages of sunflower seeds of the cultivars used in this study, 'Helio 250' and 'Helio 253' were 98% and 93%, respectively. From the seed health pre-test presence natural incidence of *S. sclerotiorum* was not found in the seed lots used.

Infected seeds by the two isolates of *S. sclerotiorum* presented in this work reductions in the variable germination according to the inoculum potentials used (Figure 1A). For one cultivar, 'Helio 250', the reduction was of 25% and 76.5% in relation to both isolates, CMLAPS 423 and CMLAPS 250, respectively. The reduction of that variable was even higher (73%) for the cultivar 'Helio 253', for both isolates compared to the control.

Regarding the effect of the water restrictor used to inhibit seed germination during the inoculation of seeds in agar substrate in the absence of the pathogen, there was also a decline in the percentage of seed germination, proportional to the increase of the periods of time of contact between seeds and water restrictor in agar substrate. But the declining slope was not significant in comparison with the reduction observed when *S. sclerotiorum* was present in the agar substrate.

The incidence of *S. sclerotiorum* in sunflower seeds, as provided by the health testing, was correlated with inoculum potential of the fungus used for both cultivars and fungal isolates (Figure 1B). In relation to cultivars, it was observed a similar reaction to *S. sclerotiorum* isolates with a gradual increase in the incidence of infection, reaching 100% at the highest level of inoculum potential (P4).

![Figure 1. A) Percentage seed germination in sunflower infected by *S. sclerotiorum* in different inoculum potentials (P1, P2, P3, P4) of the fungus and B) incidence of *S. sclerotiorum* in sunflower seeds infected at different inoculum potentials. Bars represent Tukey 0.05 mean separation test; overlapping bars indicate lack of significant difference.](image-url)

For the variables: emergence rate index, initial and final stands, variations were observed in the factors analyzed at different levels of seed infection (Figures 2, 3 and 4). The highest values of...
vigor (emergence rate index) were observed on the potential P1, this value decreasing with increasing inoculum potential, with similar trends regarding cultivars, isolates and temperature (Figure 2 – A and B). For sunflower seeds submitted to different periods of contact with the water restrictor in the absence of the pathogen, reductions were observed in the seed performance, but in smaller proportions compared to the treatments evaluated in the presence of the fungus.

Figure 2. Emergence rate index of sunflower seeds, cultivars: 'Helio 250' and 'Helio 253' inoculated with two isolates of S. sclerotiorum (CMLAPS 250 and CMLAPS 423) in contact with the water restrictor (manitol) under temperatures of 20°C (A) and 25°C (B) in a growth room with different levels of inoculum potential. Bars represent Tukey 0.05 mean separation test; overlapping bars indicate lack of significant difference between treatments.

For the initial stand, at both temperatures of 20 and 25°C a drastic reduction was observed in the percentage of the plant population for both cultivars and isolates of S. sclerotiorum. The average values of initial reductions in the stands were approximately 95% on the higher inoculum potential compared to the lowest level. The cultivar 'Helio 250' infected with the isolate CMLAPS 423, for two temperatures 20 and 25°C (Figure 3 - A and B) showed a decline less severe in initial stand and emergence rate index in potential inoculum tested when compared with the other results, with the exception of potential P4, in which there was no difference among cultivars. These results show that the presence of S. sclerotiorum in association with sunflower seeds may cause seedling collapse in pre and post-emergence, reducing the plant population in crop fields.

Figure 3. Initial Stand (%) of sunflower seeds, cultivars: 'Helio 250' and 'Helio 253' inoculated with two isolates of S. sclerotiorum (CMLAPS 250 and CMLAPS 423) in contact with the water restrictor (manitol) under temperatures of 20°C (A) and 25°C (B) in a growth room with different levels of inoculum potential. Bars represent Tukey 0.05 mean separation test; overlapping bars indicate lack of significant difference.
In relation to final stand, evaluated at 25 days after sowing, there was little variation among cultivars for each isolate at both temperatures used (Figure 4 – A and B). Under those temperatures the plant establishment was reduced largely, reaching up to 95% in the higher inoculum potential for both cultivars and isolates of \textit{S. sclerotiorum}.

The reduction of the final stand was also observed in the treatment without the presence of \textit{S. sclerotiorum}, but with the addition of mannitol to the agar medium. In this case there was an average reduction of 20% in the sunflower population under both temperatures (20 and 25°C).

From Figure 4 it is clear that the reduction in final stand was higher with an increase in inoculum potential of \textit{S. sclerotiorum} in sunflower seeds, which can be better visualized in Figure 5, showing the effect of the fungus on cultivars and at distinct temperatures used in this work.
**Potential transmissibility of S. sclerotiorum by sunflower infected seeds under controlled conditions**

The transmission process based on assessment of white mold infections in symptomatic sunflower plants was demonstrated in this study, being the transmission rates directly proportional to inoculum potentials of the pathogen in infected seeds. However, the rates varied according to the cultivar and environment temperature in the early stage of plant development (Figure 6 – A and B). For the Cultivar ‘Helio 250’ the highest average transmission rate (% symptomatic plants) was observed when the seeds were infected by the isolate CMLAPS 423, with increments close to 6% to 31% and 12% to 37% at a temperature of 20°C (A) and 25°C (B), respectively, proportional to the increase of the inoculum potential. In seeds of this same cultivar infected by isolate CMLAPS 250, these rates varied in the range of nil to 31% and nil to 18% at temperatures of 20°C (A) and 25°C (B), respectively, depending on the inoculum potential. Similar results were observed for cultivar ‘Helio 253’, in which seeds were inoculated with isolate CMLAPS 423 and had the highest rates of transmission with symptomatic infection for both temperatures.

![Figure 6](image-url)

**Figure 6.** Values of transmission rate (%) of isolates of *S. sclerotiorum* (CMLAPS 250 and CMLAPS 423) from sunflower seeds (cultivars Helio 250 and Helio 253) to plants with symptomatic infection cultivated in growth chamber with controlled temperature (20°C - (A) and 25°C - (B)). Bars represent Tukey 0.05 mean separation test; overlapping bars indicate lack of significant difference.

The transmission of the pathogen to asymptomatic plants was not observed in this study as demonstrated by the normal isolation procedures and confirmed by incubation of plant fragments on semi-selective agar bromophenol (Neon), demonstrating the absence of the fungus in tissues of sunflower plants without disease symptoms.

Regarding the total transmission rate of *S. sclerotiorum* (Figure 7 – A and B), the results varied according to temperature, fungal isolates and sunflower cultivars. In general, the highest rates were observed for the isolate CMLAPS 250, independent of cultivar and temperature, the mean values being 52.2% to 77.2%. For the isolate CMLAPS 423, the mean values of transmission rates were 48.5% to 71.7% at both temperatures. The lower rates of total transmission of *S. sclerotiorum* were observed in the lowest inoculum potentials.
DISCUSSION

The association of *S. sclerotiorum* with sunflower seeds is known well from literature but the intensity of this interaction is little investigated and understood. From epidemiology view the occurrence of that pathogen in seeds of that species may lead to various consequences, starting with low plant establishment in the field and then causing reduction in yields and being an efficient mean of disease dissemination. The internal presence of *S. sclerotiorum* in sunflower seeds was able to cause serious damages to the early development of this crop. In general, the higher inoculum potentials caused the higher seed germination reductions as well as plant development after emergence. Although *S. sclerotiorum* may be considered a soil-borne organism it was clear that its association to sunflower seeds is also an important factor affecting seed quality for planting.

An interesting factor observed in this study was the effects of the mycelium infection of sunflower seeds in causing drastic damages to seed quality and then representing a serious risk for growers using infected seeds. From the results of different evaluation tests, it was clear that internal inoculum of *S. sclerotiorum* is able to cause serious reduction in the germination rates and in plant vigour in the early stage development of the infected plants. At the lowest level of inoculum potential of *S. sclerotiorum* the mean values of germination of sunflower seeds were 86% and the emergence speed index of 1.1, under the most favorable temperature, 20°C, for white mold development. The most severe effects of the pathogen on seed germination and plant vigour occurred at the highest inoculum potential, P4. The reduction of the mean value of seed germination was 44.3% in relation to the control (non inoculated seeds). In relation to vigour, the reduction in the mean value was of 52%, reaching 94% at the highest inoculum potential of the pathogen. In literature, no quantification has been found as to the effect of *S. sclerotiorum* inoculum from sunflower seeds. That may be explained by the difficulty to find appropriate methodology to perform that evaluation. In that respect the use of the osmotic technique as described by Machado et al. (2012) that measurement is made possible.

In relation to the influence of mycelium inoculum of *S. sclerotiorum* on the plant establishment in this study it was clear the strong action of the pathogen on that variable, which was evaluated at two stages of plant development. In general the impact on that variable followed the same pattern as observed for seed germination and plant vigour. At the highest inoculum potentials the effect of *S. sclerotiorum* was more severe. There was a correlation between inoculum potentials and reductions of mean values of stands evaluated on twelve and fifty-five days after sowing.

With regard the reaction of both sunflower cultivars to both *S. sclerotiorum* isolates, no marked difference was detected between them, meaning that those factors may not play a similar role as temperatures and inoculum potentials on that interaction although further investigation should be conducted in this research line, in which a higher number of sunflower cultivars and *Sclerotinia* isolates should be used.
Potential effects of other important pathogenic fungi in seeds of important crops in Brazil, such as *Fusarium oxysporum* f. sp. *phaseoli* in bean seeds, *Colletotrichum gossypii* var. *cephalosporioides* and *Fusarium oxysporum* f. sp. *vasinfectum* in cotton seeds, *Stenocarpella maydis* and *S. macrospora* in maize seeds have been reported in literature and in general inoculum potential in those cases is also responsible for the determination of variable levels of damage to seed performance and establishment of plant population in fields (COSTA et al., 2003; ARAUJO et al., 2006; SOUSA et al., 2008).

The study on seed transmissibility of *S. sclerotiorum* in sunflower provides a clear understanding about the close interaction of this pathogen and seeds of the two sunflower cultivars used in this research. The transmission rates of the pathogen varied according to inoculum potential and temperature, the higher rates being observed at the highest inoculum potentials and under lower environment temperature. The higher proportion of dead seedlings in pre-emergence stage show the great potentiality of mycelium infection by *S. sclerotiorum* in causing drastic damages to seed performance. Because its ability to survive in soil for many years, as dormant sclerotia, dead seedlings from seed infection represent an important source of inoculum of white mold in practice. From the epidemiological view such transmission pattern means an alternative way to guarantee the disease in the field with serious implications for growers. Investigation conducted by Tu (1988) on white mold in beans had shown that *S. sclerotiorum* was able to survive in infected seeds as dormant mycelium in testa and cotyledons, and the rate of survival varied from 85% to 89% and did not change appreciably over a 3-year period.

Interesting to note also in this investigation that even at low level inoculum potential, *S. sclerotiorum* was able to be transmitted to seedling and emerged plants at high proportion. At potential P1, which consisted in exposing seeds to the fungal colony for 24 hours, the total transmission rate was 50.4%; whereas at the highest potential (P4) transmission rate was 74.3%. Although similar transmission rates were observed for the two sunflower cultivars and the two isolates of the pathogen used in this work, caution should be taken in making any general conclusion on the effect of those variables. A more representative number of cultivars of sunflower and of isolates of *S. sclerotiorum* should be considered for a better understanding of that interaction.

From the examination of *S. sclerotiorum* infection in emerged plants, as part of the determination of transmission rates of this pathogen, no infection was detected in asymptomatic plants. In terms of diagnosis and management of this disease, that fact is quite relevant as asymptomatic plants in other pathosystems may be infected by their pathogens and that may be the cause of serious consequences for growers (Oren et al., 2003; Vallad et al., 2005). In that direction, further investigation should be conducted for confirmation.

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