

Effects of Sub-Labeled Rates of Dazomet and Metam-Sodium Applied under Low-Permeability Films on *Calonectria* Microsclerotia Survival

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Abstract

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Infested soil is the primary inoculum source for *Calonectria* species for initiating disease in ornamental and forestry crops. The effects of dazomet and metam-sodium on survival of microsclerotia of 28 isolates belonging to 19 *Calonectria* species were evaluated in this study under nursery conditions. Two experiments with exotic *Calonectria* species in plastic containers in a greenhouse and three trials with endemic species in field plots were performed during different seasons. The containers and plots were artificially infested with *Calonectria* microsclerotia differentiated on carnation leaf tissues. Basamid (dazomet) was applied at 100, 160, 200, 400, and 500 kg/ha, while Divapan (metam-sodium) was applied at 250, 350, 400, 700, and 1,000 l/ha in the

both the containers and plots. The fumigants were applied under virtually and totally impermeable films. Fungal survival was evaluated after 21 d using leaf tissues collected from treated soil and plated on potato dextrose agar, and the ability of microsclerotia to cause infection was tested on red clover. The survival of *Calonectria* inocula and microsclerotia decreased with increasing fumigant rates. In the greenhouse trials, where Basamid was applied at 200, 400 and 500 kg/ha, and Divapan at 400, 700 and 1,000 l/ha, no viable microsclerotia were recovered for 14 exotic *Calonectria* species, whereas viable inocula of *C. hongkongensis*, *C. naviculata* and *C. sulawesiensis* were retrieved from the fumigated plots. Low rates of Basamid (100 and 160 kg/ha) and Divapan (250 and 350 l/ha) were less effective at reducing *Calonectria* viability and, for these treatments, the rate of microsclerotia survival was highly variable among the different isolates and species. Furthermore, totally impermeable film significantly enhanced fumigant performance. Relative to endemic *Calonectria* species, all of the treatments killed microsclerotia of *C. polizzii* and *C. pauciramosa* independent from fumigant, rate, and film. This research demonstrated the possibility of reducing the application rates by up to 160 kg/ha for Basamid and 400 l/ha for Divapan under low-permeability films (VIF or TIF) for eradicating/reducing the primary inoculum of *Calonectria* species in soil.

Calonectria species (also known as *Cylindrocladium*, asexual morph) are widely distributed in tropical and subtropical climates and are pathogens of a broad range of ornamental and forestry crops, especially in nursery settings (Alfenas et al. 2013; Crous 2002; Crous et al. 1991; Lombard et al. 2009, 2010a, b, 2011; Polizzi and Crous 1999; Vitale et al. 2013b). Disease symptoms associated with *Calonectria* spp. include damping-off, blight, leaf spots, crown, and collar and root rot (Crous 2002; Henricot and Culham 2002; Lombard et al. 2010a; Polizzi et al. 2006a, b, 2007a, b, 2009, 2012; Vitale et al. 2008, 2009a; Vitale and Polizzi 2008). The management of infections in nurseries should involve the development of integrated strategies aimed at reducing both the level of the primary inoculum and the rate of infection in contaminated substrate/soil used for cultivation.

Chemical control is the most widespread approach for managing *Calonectria* diseases in nurseries (Aiello et al. 2013; Cinquerrui et al. 2017; La Mondia 2014, 2015). However, the use of some fungicides, such as benzimidazoles and sterol demethylation inhibitors should be limited because they induce a high-selective pressure for resistant isolates (Guarnaccia et al. 2014; Vitale et al. 2009b). Some biological control agents are effective against *Calonectria* infections, but their efficacies are variable depending on the application modes and timing, as well as the target species and isolates (Daughtrey and Benson 2005; Harman 2000; Vitale et al. 2012).

A primary inoculum of *Calonectria* spp. consists of microsclerotia which can survive in soil for 15 years or more (Thies and Patton 1970; Phipps and Beute 1979). In nurseries, especially those with replants, the use of infected substrate/soil is the typical source primary *Calonectria* inoculum. Making matters worse, nurseries have recently begun using containers made from recycled and composted organic substrates, such as peat, bark, wood fibers, and green waste (Chong 2005; Walters 2009), which can further increase the risk of *Calonectria* spp. infections (Noble and Roberts 2004). Once substrates are infested, the eradication of these pathogens can be very difficult. Soil solarization treatments to reduce the potential inoculum have been successfully reported against *Calonectria* spp. (Vitale et al. 2013a). However, the efficacy of soil solarization is strictly dependent on exposure time, the temperature reached in the upper 0–30-cm layer of soil, and the specific heat sensitivity of the targeted isolate (Vitale et al. 2013a). Soil fumigation is a more widespread practice for reducing inoculum in intensive cultivation systems; however, only limited studies have addressed *Calonectria* spp. infesting peanuts and forest tree nurseries (Crous 2002). According to recent European directives only metam-sodium (MS), metam-potassium and dazomet (DZ) are authorized for the disinfection of soils/substrates in agriculture. Moreover, the European Parliament and of the Council of 21 October 2009's Directive 2009/128/EC ("Sustainable Use of Pesticide") imposes severe restrictions regarding application methods and rates, including that the fumigants should be applied at reduced rates and under mulching films that decrease gas emissions and environmental damage (Dir. 2009/128/EC; Commission Implementing Reg. (UE) n. 359/2012).

As a result, our current research has focused on the more advanced multi-layer films, such as virtually impermeable film (VIF) and totally impermeable film (TIF), that are less permeable to chemicals than the low- and high-density polyethylene films (Fennimore and Ajwa 2011).

A preliminary study recently demonstrated the potential use of labeled rates of DZ and MS applied under VIF for the suppression of *Calonectria* microsclerotia in nurseries of the Mediterranean Basin (Polizzi et al. 2014). However, no data are available on the effects of sub-labeled fumigant rates and TIF performances against epidemic *Calonectria polizzii* and *Calonectria pauciramosa* and exotic species from forest crops. These information could improve the environmentally sustainability of these fumigants.

Thus, we investigated the effects of DZ and MS applied at labeled and sub-labeled rates under TIF and VIF films on 17 exotic *Calonectria* species in greenhouse trials and the effects of the fumigants applied at sub-label rates on well-established *C. polizzii* and *C. pauciramosa* species in the Mediterranean Basin in open fields under nursery conditions.

Materials and methods

Pathogen isolates. In total, 28 isolates belonging to 19 *Calonectria* species were used in this study, including two species well-established in Italy, *C. polizzii* ITEM 14877 and *C. pauciramosa* ITEM 14884, and 17 exotic species previously identified by a multi-gene sequence analysis but not currently present in Italy (Table 1). All of the isolates were grown on potato dextrose agar (PDA, Oxoid, Basingstoke, UK) prior to transfer to carnation leaf agar (CLA; Fisher et al. 1982) for microsclerotia production.

Effects of DZ and MS on exotic *Calonectria* microsclerotia. Because the exotic *Calonectria* species are not present in Italy, experiments to test fungal survival were performed in plastic containers (40 × 60 cm). Two experiments (I and II) were performed in November 2014 and May 2015, respectively, in a greenhouse to determine the effects of (DZ) and (MS) on the viability of microsclerotia of 17 *Calonectria* species (26 isolates) buried in a soil substrate. The commercial fumigants evaluated were Basamid Granulat, (99% DZ, water-dispersible granules, Kanesho Soil

Treatment, Brussels, Belgium) and Divapan, (51% MS, suspension concentrate, Taminco Italia S.r.l., Milano, Italia).

Microsclerotia were produced on CLA after 2 weeks of incubation at $25 \pm 1^\circ\text{C}$, and two carnation leaf segments (6-cm long) colonized by pathogen microsclerotia were removed from Petri dishes and placed in a nylon mesh bag. Three bags for each isolate at each fumigant rate were buried (15-cm depth) in plastic containers filled with a loamy-sand substrate.

Basamid (DZ) was applied at rates of 100, 160, 200, 400, and 500 kg/ha and mixed uniformly into the soil before the bags were buried, whereas Divapan (MS) was applied as soil drench with a hand sprinkler at rates of 250, 350, 400, 700, and 1,000 l/ha. Following fumigation, the containers were brought to water field capacity, covered, and hermetically closed with VIF, (Ecobrom[®], AgriPlast S.r.l., Vittoria, Ragusa, Italy) or TIF, (EvalKuroray, USA) by fixing their edges at the container margins with adhesive tape. Subsequently, containers were placed in a greenhouse. The same number of bags and infected carnation leaf pieces buried in untreated substrate in plastic containers served as the controls. Three replicates (three plastic containers) were used for each isolate, rate, and tested film and arranged in a randomized complete block design applied to each fumigant separately. The nylon bags were sampled 21 d after treatment. Nine leaf pieces, obtained by cutting each leaf segment, were washed with sterile distilled water (SDW), placed onto PDA plates and maintained for 1 week at $25 \pm 1^\circ\text{C}$. The percentage of leaf pieces from which pathogen colonies developed was used to determine the survival of single *Calonectria* isolates after recovery from soil.

Effects of reduced fumigant rates on the survival of *C. polizzii* and *C. pauciramosa* microsclerotia. Experiments III, IV and V were performed in November 2013, and May and July 2014 in open fields to determine the effects of sub-labeled rates of DZ and MS on the viability of microsclerotia of *C. polizzii* and *C. pauciramosa* isolates. A 40-cm layer of soil was placed on a cement bed. The experimental plots, each 2.5×5.0 m, were arranged in a randomized complete block design applied to each of both species separately with three replicates for fumigant rate and

tested film. Microsclerotia were produced on CLA as described above, and two carnation leaf segments (6-cm long) colonized by pathogen microsclerotia were removed from Petri dishes and placed in a nylon mesh bag. Three bags for each isolate at each fumigant rate were buried (15-cm depth) in the soil, and filled with loamy-sand soil. Basamid (DZ) was applied at 100, 160 and 200 kg/ha and mixed uniformly into the soil before the bags were buried, while Divapan (MS) was applied as a soil drench by a hand sprinkler at 250, 350, and 400 l/ha. After fumigation, the soil in each plot was irrigated to field capacity and covered with sheet of VIF or TIF. The film sheets were laid on the soil surface, and their edges were buried 20-cm deep. The same number of carnation leaf pieces buried in the untreated soil plot served as the control. At 21 d after treatment, nine leaf pieces, obtained by cutting each leaf segment, were washed with SDW and placed onto PDA plates and maintained for 1 week at 25 ± 1 °C. The percentage of leaf pieces from which pathogen colonies developed was used to determine the survival of *Calonectria* species after recovery from the soil.

Evaluation of the viability of *C. polizzii* and *C. pauciramosa* microsclerotia on red clover following fumigation treatments. After a 21-d exposure time to fumigation, carnation leaf samples were retrieved from both untreated and fumigated plots, and microsclerotia viability was assayed on red clover (*Trifolium pratense* L.) seedlings according to a method reported in recent papers (Waipara et al. 1996; Vitale et al. 2012). Additional bags containing two infected carnation leaves were previously buried in each plot (replicate) for all treatments. The retrieved carnation leaf samples were cut into smaller pieces (18 pieces per replicate) and mixed with peat substrate in an aluminum tray. The aluminum trays were placed in a growth chamber and brought to water field capacity before seeding the red clover (three replicate aluminum trays, each containing up to 70 seeds). The percentage of red clover seedlings showing crown and root rot symptoms of the total number of examined seedlings was recorded 8–10 d after clover seeding.

Statistical analyses. Data from independent experiments (I to V) were analyzed separately by using the Statistica package software (Version 10, Statsoft Inc., Analytical Software for Windows).

An analysis of variance was performed for each fumigant separately to evaluate the viability among *Calonectria* isolates exposed to single rates and to compare the performances of VIF and TIF in greenhouse experiments I and II (Table 2). For both experiments results were analyzed in independent experiments based on year (season) and tested fumigant. Subsequently, analyses were conducted by calculating *F*-values and significances (*P*) associated with the experimental factors (isolate, rate, and film) and whether there were significant interactions among these factors within each independent experiment. The efficacy of the fumigants and rates on *C. polizzii* ITEM 14877 and *C. pauciramosa* ITEM 14884 microsclerotia viability and the relative VIF and TIF performances at each fumigant rate in experiments III, IV and V in the soil under open-field conditions were also examined using an analysis of variance. The mean separation of the viability reduction percentages compared with relative controls in plastic containers among isolates, the fumigant effects on *C. polizzii* and *C. pauciramosa* survival rates in soil under open-field conditions, and the relative infectivity levels (disease incidence) in red clover were assessed using Fisher's least significance difference test at $\alpha = 0.05$.

Results

Effects of DZ and MS on exotic *Calonectria* microsclerotia. Isolate, rate, film, and their interactions always significantly impacted the number of viable microsclerotia buried in the fumigated plots (Table 2). TIF significantly enhanced the performance of fumigation when compared with VIF over independent experiments (Table 2). Data from experiments I and II are reported in Tables 3-6.

Because all of the interactions were significant (Table 2), the responses to fumigation among the isolates were analyzed and shown for single fumigation rates under the same film (Tables 3-6).

In experiment I, Basamid (DZ) at 400 and 500 kg/ha resulted in the elimination of microsclerotia viability for all of the *Calonectria* species, whereas at 200 kg/ha viability was eliminated in all of the species, except *C. naviculata*. When this fumigant was applied at 160 kg/ha, no viable

microsclerotia were retrieved from fumigated containers for any species, except for *C. variabilis* LPF220, *C. hongkongensis*, and *C. naviculata* isolates. This fumigant was less effective when applied at 100 kg/ha, resulting in the elimination of microsclerotia viability for 16 of 26 and 21 of 26 isolates under VIF and TIF, respectively. The highest variability of response to DZ fumigation was detected at 100 kg/ha of the commercial formulate (Basamid; Table 3). Divapan (MS) at 1,000 l/ha eliminated the microsclerotia viability of all species/isolates, while at 700 and 400 l/ha viable inocula were recovered only from *C. sulawesiensis* and *C. hongkongensis*. The commercial formulate (Divapan) at 250 and 350 l/ha showed weaker capabilities to reduce the microsclerotia viability of *Calonectria* species/isolates, and these concentrations resulted in the greatest variability in response to fumigation among the tested isolates (Table 4).

In experiment II, the highest rates of Basamid (DZ) showed excellent efficacies at 400 and 500 kg/ha in killing the microsclerotia of all *Calonectria* species. This fumigant applied at 160 and 200 kg/ha totally reduced the microsclerotia viability of all species, except for *C. naviculata*, *C. variabilis* LPF220 and *C. hongkongensis* whereas at 100 kg/ha it was, on average, less effective compared with the data of experiment I. At this fumigant rate the elimination of microsclerotia viability was detected for only 6 of 26 and 14 of 26 isolates under VIF and TIF, respectively (Table 5).

Divapan (MS) at 400, 700, and 1,000 l/ha eliminated *Calonectria* microsclerotia viability, whereas the 350 l/ha rate killed the most *Calonectria* isolates under TIF (21 of 26) and half of the isolates under VIF. The commercial formulate applied at 250 l/ha was the least efficient in reducing the viability of *Calonectria* inocula. As in the previous experiment, a high rate of reduction in viability among *Calonectria* isolates was detected at the low fumigation rates (Table 6).

Effects of reduced fumigant rates on the survival of *C. polizzii* and *C. pauciramosa* microsclerotia. Data from experiments III, IV, and V, performed in open fields, are reported in Table 7. In experiment III, all of the treatments killed microsclerotia of *C. polizzii* and *C. pauciramosa* independent of fumigant, rate, and film. In experiment IV, all of the treatments were

effective in eliminating the inocula of *C. polizzii* and *C. pauciramosa*, except the lowest rates of Basamid (DZ) (under both films) and Divapan (MS) (under VIF), which significantly reduced the viable inocula of each species compared with control. At these rates, TIF increased the efficacy of fumigant treatments compared to VIF. In the last experiment (V), only Basamid (DZ) failed to kill all of the viable microsclerotia of these two *Calonectria* species when applied at 100 kg/ha and of *C. polizzii* at 160 kg/ha under TIF. Significant differences related to fumigant rate and film were observed for *Calonectria* survival, but TIF did not improve the efficacies of treatments when compared with VIF.

Evaluation of the viability of *C. polizzii* and *C. pauciramosa* microsclerotia on red clover following fumigation treatments. Microsclerotia infectivity rates on young red clover seedlings in experiments III, IV, and V are reported in Table 8. In experiment III, no *Calonectria* infections were observed on pre-infected carnation debris after retrieval from the fumigated plots. This corroborated the pathogen survival rate of zero. The disease incidence was strongly correlated to the survival of *Calonectria* microsclerotia. Thus, in experiment IV, *Calonectria* infections were only found on infected debris recovered from plots treated with Basamid (DZ) under VIF or TIF at a rate of 100 kg/ha and Divapan (MS) under VIF at 250 l/ha. *Calonectria* infections on red clover were only obtained from microsclerotia exposed to fumigation at 160 kg/ha rate under TIF and at 100 kg/ha under both films. The pairwise combinations of the significant differences between treatments were similar to those observed for the survival experiments.

Discussion

This paper provides initial information on the efficacies of DZ and MS fumigants applied at reduced rates under low-permeability films under nursery conditions against 19 *Calonectria* species, representative of a worldwide population.

In the greenhouse experiments, both DZ and MS were effective in reducing exotic *Calonectria* microsclerotia viability, although some differences in their performances were observed between

experiments. Both Basamid (DZ) and Divapan (MS), when applied at 160 kg/ha and 400 l/ha concentration and greater, respectively, under either film were effective in eradicating *Calonectria* microsclerotia in soil, except for *C. hongkongensis* (viable up to 700 l/ha of Divapan and 200 kg/ha of Basamid), *C. sulawesiensis* (viable up to 400 l/ha of Divapan) and *C. variabilis* LPF220 and *C. naviculata* (viable up to 160 and 200 kg/ha of Basamid, respectively). At lower rates, there was greater variability in microsclerotia variability among most *Calonectria* isolates in all of the experiments. In general, TIF enhanced fumigant performance. Because high inter- and intraspecific variability of responses to fumigants was detected among *Calonectria* species/isolates, the correct identification of target *Calonectria* species could play an important role in the choice of fumigant and application rate.

The field data confirmed the efficacy of the highest rates of both fumigants against *C. polizzii* and *C. pauciramosa* as similarly reported by Polizzi et al. (2014). In comparison with a previous study that utilized higher labeled rates (400, 700, and 1,000 l/ha, and 200, 400, and 500 kg/ha of Divapan and Basamid respectively), the reduced rates (defined as less than the labeled rate) applied under VIF and TIF were also effective in reducing both endemic *Calonectria* in soil. In addition, the efficacy of the fumigant treatment appeared to be strongly influenced by the season and, to a lesser extent, by the film used. Indeed, no viable inocula were retrieved from any of the fumigated plots in the July experiment. In the remaining experiments (in November and May), the lowest rates of fumigants were less effective and did not eliminate viable inocula of either pathogen. Moreover, TIF did not always improve the efficacy of the treatment when compared with VIF. The highest *Calonectria* infections rates were obtained from inocula retrieved from untreated plots and, to a lesser extent, from plots treated at the lowest fumigant rates.

The current labeled rates are 300 l/ha in open fields and 700–1200 l/ha in greenhouses for Divapan 51 while the rates of Basamid range from 300–500 kg/ha (at a 15–20 cm depth) for some pathogens to 500–700 kg/ha (at a 25–30 cm depth) for other pathogens. Our data clearly showed the possibility of reducing the application rates by up to 160 kg/ha for Basamid (DZ) and 400 l/ha for

Divapan (MS) against all *Calonectria* inocula in soil/substrate used for ornamental and forestry crops when using low-permeability films (VIF or TIF). High rates of MS should be applied only for *C. hongkongensis*, *C. sulawesensis*, *C. variabilis*, and *C. naviculata*. The use of Basamid (DZ) applied at 100 kg/ha and Divapan (MS) at 250 l/ha in July could be encouraged for the eradication of endemic *C. polizzii* and *C. pauciramosa* under nursery conditions in the Mediterranean Basin. DZ and MS have been previously reported as effective against other soilborne pathogen inocula but at higher rates than those reported in this paper (Wang et al. 2006; Weiland 2011, 2013)

TIF is less permeable than VIF, and it allows a decrease in fumigant application rates by 40% to 50% without sacrificing fumigant efficacy (Fennimore and Ajwa 2011; McAvoy and Freeman 2013; Gao et al. 2013). However, the fumigant efficacy in our study was not always significantly greater under TIF compared with VIF. Nevertheless, TIF could be particularly attractive because it decrease gas emissions. High permeability films (low-density and high-density polyethylene) should be replaced by low-permeability films to reduce or kill *Calonectria* inocula as well as it already was reported for other soilborne fungal pathogens (Cabrera et al. 2015; Chamorro et al. 2016) because they are more environmentally friendly (Gao et al. 2011) and allow for the use of lower fumigant rates.

This investigation indicated that fumigation at reduced rates under low-permeability films is an appropriate option to reduce or kill soilborne *Calonectria* before growing seedlings, propagating plants, or replanting pot-grown plants in nurseries. Because the most widely used potting substrate is derived from mixture of local (volcanic) soil with commercial peat and mineral constituents, the sustainable disinfection of soil/substrate should be encouraged on a large-scale as an additional agronomic practice in nursery production.

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Table 1. Fungal isolates collected from ornamental and forest crops used in this study.

Isolate code ^a	Location	Host	Other collections (nr.) ^b	Species	Species complex	References
LPF130	Suzano - BA	<i>Eucalyptus</i> sp.		<i>Calonectria brachiatica</i>	<i>C. brassicae</i>	Lombard et al. 2009
LPF195	Suzano - BA	Native forest		<i>Calonectria brachiatica</i>	<i>C. brassicae</i>	Lombard et al. 2009
LPF034	Jari	<i>Eucalyptus</i> sp.		<i>Calonectria brassicae</i>	<i>C. brassicae</i>	Lombard et al. 2009
LPF290	Suzano - BA	<i>Eucalyptus</i> sp.		<i>Calonectria brassicae</i>	<i>C. brassicae</i>	Lombard et al. 2009
LPF452	Jari	<i>Eucalyptus</i> sp.		<i>Calonectria ecuadoriae</i>	<i>C. brassicae</i>	Crous et al. 2006
LPF300	Jari	<i>Eucalyptus</i> sp.		<i>Calonectria orientalis</i>	<i>C. brassicae</i>	Lombard et al. 2010c
LPF301	Jari	<i>Eucalyptus</i> sp.		<i>Calonectria orientalis</i>	<i>C. brassicae</i>	Lombard et al. 2010c
LPF031	Jari	<i>Eucalyptus</i> sp.		<i>Calonectria pini</i>	<i>C. brassicae</i>	Lombard et al. 2010c
LPF253	Amcel	<i>Eucalyptus</i> sp.		<i>Calonectria pini</i>	<i>C. brassicae</i>	Lombard et al. 2010c
LPF388	Suzano - MA	<i>Eucalyptus</i> sp.		<i>Calonectria brasiliensis</i>	<i>C. cylindrospora</i>	Lombard et al. 2010c
LPF244	Viçosa - MG	<i>Piptadenia gonoacantha</i>	CBS133608	<i>Calonectria hodgesii</i>	<i>C. cylindrospora</i>	Alfenas et al. 2013
LPF245	Viçosa - MG	<i>Anadenanthera peregrina</i>	CBS133609	<i>Calonectria hodgesii</i>	<i>C. cylindrospora</i>	Alfenas et al. 2013
LPF389	Suzano - MA	<i>Eucalyptus</i> sp.		<i>Calonectria sulawesiensis</i>	<i>C. cylindrospora</i>	Lombard et al. 2010c
LPF007	Jari	<i>Eucalyptus</i> sp.	CPC18775	<i>Calonectria variabilis</i>	<i>C. cylindrospora</i>	Crous et al. 1993a
LPF220	Amcel	<i>Eucalyptus</i> sp.		<i>Calonectria variabilis</i>	<i>C. cylindrospora</i>	Crous et al. 1993a
LPF174	Maranhão	<i>Eucalyptus</i> sp.		<i>Calonectria ovata</i>	<i>C. pteridis</i>	Victor et al. 1997
LPF257	Amcel	<i>Eucalyptus</i> sp.		<i>Calonectria ovata</i>	<i>C. pteridis</i>	Victor et al. 1997
LPF002	Jari	<i>Eucalyptus</i> sp.	CPC18771	<i>Calonectria pteridis</i>	<i>C. pteridis</i>	Crous et al. 1993b
LPF004	Jari	<i>Eucalyptus</i> sp.	CPC18773	<i>Calonectria pteridis</i>	<i>C. pteridis</i>	Crous et al. 1993b
LPF075	Araponga - MG	Native forest	CPC18745	<i>Calonectria colombiana</i>	<i>C. candelabra</i>	Lombard et al. 2010c
DISTEF173	Lamezia - Catanzaro	<i>Callistemon citrinus</i>	ITEM14884	<i>Calonectria pauciramosa</i>	<i>C. candelabra</i>	Schoch et al. 1999
DISTEF87	San Marco - ME	<i>Acacia retinodes</i>	ITEM14877	<i>Calonectria polizzii</i>	<i>C. candelabra</i>	Lombard et al. 2010b
LPF066	Araponga - MG	Native Forest	CPC18736	<i>Calonectria spathulata</i>	<i>C. candelabra</i>	Crous et al. 1994
LPF061	Suzano - BA	<i>Eucalyptus</i> sp.		<i>Calonectria pseudoscoparia</i>	<i>C. candelabra</i>	Lombard et al. 2010c
LPF212	Alagoas	<i>Eucalyptus</i> sp.		<i>Calonectria pseudoscoparia</i>	<i>C. candelabra</i>	Lombard et al. 2010c
LPF219	Suzano - BA	Native forest		<i>Calonectria zuluensis</i>	<i>C. candelabra</i>	Lombard et al. 2010b
LPF311	Jari	<i>Eucalyptus</i> sp.		<i>Calonectria naviculata</i>	Sphaero-naviculate Group	Crous et al. 1994
LPF111	Unknown	Unknown		<i>Calonectria hongkongensis</i>	Sphaero-naviculate Group	Crous et al. 2004

^a LPF: Laboratório de Patologia Florestal, Universidade Federal de Viçosa, Viçosa, Brazil, DISTEF: Dipartimento di Agricoltura, Alimentazione e Ambiente, Catania, Italy (Vitale et al., 2012).

^b CBS: CBS-KNAW Fungal Biodiversity Centre, Utrecht, The Netherlands, CPC: Pedro W. Crous working collection housed at CBS, ITEM: numbers as reported by Vitale et al. (2013).

Table 2. ANOVA effects of involved factors and their relative interactions on *Calonectria* spp. survival (%) in plastic containers from independent experiments.

Year (season)	Fumigant	Source of variation	DF ^a	MS ^a	F-value	Significance ^b
2014 (autumn)	Dazomet	Isolate	25	1376	32.8	***
		Rate	4	4859	115.9	***
		Film	1	375	8.9	**
		Isolate × rate	100	852	20.3	***
		Isolate × film	25	305	7.3	***
		Isolate × rate × film	100	211	5.0	***
	Metam-Na	Isolate	25	7294.9	25.6	***
		Rate	4	263479.8	924.7	***
		Film	1	11708.0	41.1	***
		Isolate × rate	100	2411.6	8.5	***
		Isolate × film	25	1511.7	5.3	***
		Isolate × rate × film	100	1178.3	4.1	***
2015 (spring)	Dazomet	Isolate	25	5860	33.7	***
		Rate	4	52853	304.4	***
		Film	1	7729	44.5	***
		Isolate × rate	100	2306	13.3	***
		Isolate × film	25	551	3.2	***
		Isolate × rate × film	100	468	2.7	***
	Metam-Na	Isolate	25	1920.0	11.9	***
		Rate	4	97912.5	608.2	***
		Film	1	19560.9	121.5	***
		Isolate × rate	100	1159.3	7.2	***
		Isolate × film	25	1003.0	6.2	***
		Isolate × rate × film	100	637.2	3.9	***

^a DF = degrees of freedom; MS = Mean square

^b **, *** = significant at $0.001 < p < 0.01$ and $p < 0.001$, respectively.

Table 3. Comparison of the effects of DZ (Basamid) on microsclerotia viability reductions (%) when applied to different *Calonectria* species in plastic containers under VIF or TIF in autumn 2014 (Experiment I).

Isolate	100 kg/ha ^a		160 kg/ha		200 kg/ha	
	VIF	TIF	VIF	TIF	VIF	TIF
<i>C. brachiatica</i> LPF130	100 a	100 a	100 a	100 a	100 a	100
<i>C. brachiatica</i> LPF195	100 a	100 a	100 a	100 a	100 a	100
<i>C. brassicae</i> LPF034	100 a	100 a	100 a	100 a	100 a	100
<i>C. pini</i> LPF253	100 a	100 a	100 a	100 a	100 a	100
<i>C. orientalis</i> LPF300	100 a	100 a	100 a	100 a	100 a	100
<i>C. orientalis</i> LPF301	100 a	100 a	100 a	100 a	100 a	100
<i>C. ecuadoriae</i> LPF452	100 a	100 a	100 a	100 a	100 a	100
<i>C. pteridis</i> LPF004	100 a	100 a	100 a	100 a	100 a	100
<i>C. ovata</i> LPF174	100 a	100 a	100 a	100 a	100 a	100
<i>C. ovata</i> LPF257	100 a	100 a	100 a	100 a	100 a	100
<i>C. sulawesiensis</i> LPF389	100 a	86.4 b	100 a	100 a	100 a	100
<i>C. pseudoscoparia</i> LPF061	100 a	75.3 c	100 a	100 a	100 a	100
<i>C. hodgesii</i> LPF244	100 a	100 a	100 a	100 a	100 a	100
<i>C. hodgesii</i> LPF245	100 a	100 a	100 a	100 a	100 a	100
<i>C. zuluensis</i> LPF219	100 a	100 a	100 a	100 a	100 a	100
<i>C. colombiana</i> LPF075	100 a	100 a	100 a	100 a	100 a	100
<i>C. spathulata</i> LPF066	98.8 ab	100 a	100 a	100 a	100 a	100
<i>C. variabilis</i> LPF220	96.3 ab	100 a	87.5 a	100 a	100 a	100
<i>C. brasiliensis</i> LPF388	93.8 ab	100 a	100 a	100 a	100 a	100
<i>C. brassicae</i> LPF290	92.6 ab	100 a	100 a	100 a	100 a	100
<i>C. pseudoscoparia</i> LPF212	92.6 ab	85.2 bc	100 a	100 a	100 a	100
<i>C. pini</i> LPF031	91.4 ab	100 a	100 a	100 a	100 a	100
<i>C. variabilis</i> LPF007	85.2 b	100 a	100 a	100 a	100 a	100
<i>C. pteridis</i> LPF002	66.7 c	100 a	100 a	100 a	100 a	100
<i>C. hongkongensis</i> LPF111	33.3 d	53.1 d	98.8 a	74.1 b	100 a	100
<i>C. naviculata</i> LPF311	0 e	56.8 d	66.7 b	100 a	66.7 b	100

^a Each value represents the mean of three replicates, each constituted by 54 infected carnation pieces. Recovery values followed by different letters within each column are

significantly different according to Fisher's least significant difference test ($\alpha = 0.05$)

Table 4. Comparison of the effects of MS (Divapan) on microsclerotia viability reductions (%) when applied to different *Calonectria* species in plastic containers under VIF or TIF in autumn 2014 (Experiment I).

Isolate	250 liters/ha ^a		350 liters/ha		400 liters/ha		700 liters/ha	
	VIF	TIF	VIF	TIF	VIF	TIF	VIF	TIF
<i>C. brassicae</i> LPF290	100 a	75.3 abc	100 a	79 bc	100 a	100 a	100 a	100 a
<i>C. hodgei</i> LPF244	76.6 b	100 a	98.8 ab	100 a	100 a	100 a	100 a	100 a
<i>C. hodgei</i> LPF245	58 bcd	51.9 b-f	100 a	81.5 ab	100 a	100 a	100 a	100 a
<i>C. colombiana</i> LPF075	58 bc	42 c-g	69.1 c-f	84 ab	100 a	100 a	100 a	100 a
<i>C. brassicae</i> LPF034	48.2 cde	66.7 a-e	100 a	92.6 ab	100 a	100 a	100 a	100 a
<i>C. pteridis</i> LPF004	44.4 c-f	46.9 b-f	55.6 fg	100 a	100 a	100 a	100 a	100 a
<i>C. zuluensis</i> LPF219	44.4 cde	43.2 b-f	77.8 a-f	100 a	100 a	100 a	100 a	100 a
<i>C. spathulata</i> LPF066	43.2 c-f	28.4 f-i	33.3 ghi	70.4 bc	100 a	100 a	100 a	100 a
<i>C. pini</i> LPF253	42 cde	35.8 e-h	59.3 efg	70.4 bcd	100 a	100 a	100 a	100 a
<i>C. sulawesiensis</i> LPF389	35.8 c-f	18.5 f-i	93.8 abc	76.6 bc	100 a	91.4 b	100 a	100 a
<i>C. pini</i> LPF031	34.6 d-g	34.6 e-h	67.9 def	100 a	100 a	100 a	100 a	100 a
<i>C. orientalis</i> LPF300	32.1 def	55.6 b-f	84 a-d	84 ab	100 a	100 a	100 a	100 a
<i>C. ovata</i> LPF257	27.2 e-h	70.4 a-d	54.3 efg	100 a	100 a	100 a	100 a	100 a
<i>C. pseudoscoparia</i> LPF061	25.9 e-i	33.3 e-i	80.3 a-e	40.7 e	100 a	100 a	100 a	100 a
<i>C. orientalis</i> LPF301	24.7 e-h	50.6 b-f	86.4 a-d	58 cde	100 a	100 a	100 a	100 a
<i>C. ovata</i> LPF174	23.5 f-i	77.8 ab	75.2 a-d	100 a	100 a	100 a	100 a	100 a
<i>C. brachiatica</i> LPF130	21 e-i	81.5 abc	100 a	100 a	100 a	100 a	100 a	100 a
<i>C. variabilis</i> LPF220	9.9 ghi	100 a	30.9 gh	100 a	100 a	100 a	100 a	100 a
<i>C. variabilis</i> LPF007	3.7 hi	66.7 a-e	72.8 a-f	70.4 bc	100 a	100 a	100 a	100 a
<i>C. pseudoscoparia</i> LPF212	0 i	56.8 b-f	77.8 a-f	69.1 bcd	100 a	100 a	100 a	100 a
<i>C. brasiliensis</i> LPF388	0 i	51.9 b-f	69.1 b-f	50.6 de	100 a	100 a	100 a	100 a
<i>C. ecuadoriae</i> LPF452	0 i	42 c-h	100 a	92.6 ab	100 a	100 a	100 a	100 a
<i>C. brachiatica</i> LPF195	0 i	40.7 deh	14.8 hi	53.1 cde	100 a	100 a	100 a	100 a
<i>C. pteridis</i> LPF002	0 i	7.4 ghi	64.2 def	33.3 ef	100 a	100 a	100 a	100 a
<i>C. naviculata</i> LPF311	0 i	7.4 ghi	18.5 hi	11.1 f	100 a	100 a	100 a	100 a
<i>C. hongkongensis</i> LPF111	0 i	0 i	4.9 i	11.1 f	61.7 b	13.6 c	93.8 b	96.3 b

^a Each value represents the mean of three replicates, each constituted by 54 infected carnation pieces. Recovery values followed by different letters within each column are

significantly different according to Fisher's least significant difference test ($\alpha = 0.05$)

Table 5. Comparison of the effects of DZ (Basamid) on microsclerotia viability reductions (%) when applied to different *Calonectria* species in plastic containers under VIF or TIF in spring 2015 (Experiment II).

Isolate	Rate	100 kg/ha ^a		160 kg/ha		200 kg/ha	
		VIF	TIF	VIF	TIF	VIF	TIF
<i>C. brachiatica</i> LPF130		100 a	100 a	100 a	100 a	100 a	100 a
<i>C. pteridis</i> LPF004		100 a	100 a	100 a	100 a	100 a	100 a
<i>C. ovata</i> LPF174		100 a	100 a	100 a	100 a	100 a	100 a
<i>C. zuluensis</i> LPF219		100 a	100 a	100 a	100 a	100 a	100 a
<i>C. colombiana</i> LPF075		100 a	100 a	100 a	100 a	100 a	100 a
<i>C. pseudoscoparia</i> LPF212		100 a	100 a	100 a	100 a	100 a	100 a
<i>C. variabilis</i> LPF007		95.1 ab	100 a	100 a	100 a	100 a	100 a
<i>C. hodgesii</i> LPF245		90.1 abc	100 a	100 a	100 a	100 a	100 a
<i>C. hodgesii</i> LPF244		87.7 abc	88.9 ab	100 a	100 a	100 a	100 a
<i>C. ovata</i> LPF257		86.4 abc	19.8 ab	100 a	100 a	100 a	100 a
<i>C. brassicae</i> LPF290		84 abc	100 a	100 a	100 a	100 a	100 a
<i>C. ecuadoriae</i> LPF452		72.8 bcd	100 a	100 a	100 a	100 a	100 a
<i>C. brassicae</i> LPF034		71.6 cd	77.8 ab	100 a	100 a	100 a	100 a
<i>C. spathulata</i> LPF066		66.7 cd	100 a	100 a	100 a	100 a	100 a
<i>C. pseudoscoparia</i> LPF061		66.7 cd	66.7 bc	100 a	100 a	100 a	100 a
<i>C. brachiatica</i> LPF195		58 de	100 a	100 a	100 a	100 a	100 a
<i>C. pini</i> LPF031		49.4 de	61.7 bc	100 a	100 a	100 a	100 a
<i>C. pini</i> LPF253		48.2 de	49.4 cd	100 a	100 a	100 a	100 a
<i>C. brasiliensis</i> LPF388		35.8 ef	33.3 de	100 a	100 a	100 a	100 a
<i>C. pteridis</i> LPF002		34.6 ef	100 a	100 a	100 a	100 a	100 a
<i>C. sulawesiensis</i> LPF389		33.3 ef	66.7 bc	100 a	100 a	100 a	100 a
<i>C. hongkongensis</i> LPF111		21 f	32.1 de	33.3 c	33.3c	66.7 b	66.7 b
<i>C. naviculata</i> LPF311		16 f	11.1 e	33.3 c	46.9 b	56.8 b	100 a
<i>C. orientalis</i> LPF300		11.1 f	69.1 bc	100 a	100 a	100 a	100 a
<i>C. orientalis</i> LPF301		11.1 f	48.2 cd	100 a	100 a	100 a	100 a
<i>C. variabilis</i> LPF220		7.4 f	100 a	76.5 b	100 a	100 a	100 a

^a Each value represents the mean of three replicates, each constituted by 54 infected carnation pieces. Recovery values followed by different letters within each column are

significantly different according to Fisher's least significant difference test ($\alpha = 0.05$)

Table 6. Comparison of the effects of MS (Divapan) on microsclerotia viability reductions (%) when applied to different *Calonectria* species in plastic containers under VIF or TIF in spring 2015 (experiment II).

Isolate	Rate	250 liters/ha ^a		350 liters/ha	
		VIF	TIF	VIF	TIF
<i>C. brassicae</i> LPF290		100 a	79.2 abc	100 a	83.3 c
<i>C. pseudoscoparia</i> LPF212		80.3 abc	100 a	100 a	100 a
<i>C. sulawesiensis</i> LPF389		75.3 a-d	70.4 cde	100 a	90.1 b
<i>C. hodgesii</i> LPF244		72.7 ab	100 a	100 a	100 a
<i>C. orientalis</i> LPF300		70.4 b-e	80.3 bcd	100 a	100 a
<i>C. spathulata</i> LPF066		69.2 b-e	85.2 abc	65.4 d-g	100 a
<i>C. ovata</i> LPF257		66.7 b-f	100 a	100 a	100 a
<i>C. colombiana</i> LPF075		64.2 b-f	70.4 cde	100 a	100 a
<i>C. naviculata</i> LPF311		63 b-f	33.3 fgh	86.4 bcd	58 d
<i>C. hodgesii</i> LPF245		55.6 b-g	86.4 abc	100 a	100 a
<i>C. pini</i> LPF253		51.9 c-g	63 c-f	100 a	100 a
<i>C. ovata</i> LPF174		50.6 c-g	85.2 abc	58.1 efg	100 a
<i>C. pteridis</i> LPF004		49.4 e-h	84 abc	92.6 abc	100 a
<i>C. brachiatica</i> LPF130		49.4 c-g	63 c-f	100 a	100 a
<i>C. brachiatica</i> LPF195		45.7 c-g	95.1 ab	40.7 efg	100 a
<i>C. zuluensis</i> LPF219		44.5 e-h	88.9 ab	100 a	100 a
<i>C. brassicae</i> LPF034		44.4 d-h	100 a	100 a	100 a
<i>C. brasiliensis</i> LPF388		42 e-h	54.3 d-g	86.4 abc	98.8 ab
<i>C. orientalis</i> LPF301		42 e-h	44.5 e-h	87.7 abc	100 a
<i>C. variabilis</i> LPF220		35.8 fgh*	100 a*	74.1 c-f*	100 a*
<i>C. variabilis</i> LPF007		34.6 gh*	98.8 ab*	88.9 abc	100 a
<i>C. pini</i> LPF031		33.3 gh*	81.5 abc*	51.9 fg*	100 a*
<i>C. ecuadoriae</i> LPF452		32.1 fgh	48.1 efg	100 a	100 a
<i>C. hongkongensis</i> LPF111		32.1 gh	30.9 h	51.9 g	38.3 e
<i>C. pseudoscoparia</i> LPF061		22.2 hi*	100 a*	97.5 ab	100 a
<i>C. pteridis</i> LPF002		0 i	33.3 gh	79 cde*	100 a*

^a Each value represents the mean of three replicates, each constituted by 54 infected carnation pieces. Recovery values followed by different letters within each column are significantly different according to Fisher's least significant difference test ($\alpha = 0.05$)

Table 7. Effectiveness of reduced rates of DZ (Basamid) and MS (Divapan) applied to loamy-sand soil in affecting survival (%) of *Calonectria polizzii* and *Calonectria pauciramosa* from infected carnation leaves after exposure to fumigation treatment under VIF and TIF soil mulchings in the nursery in three experiments.

Treatment & rate (liters or kg/ha)	Experiment III (summer 2013) ^a				Experiment IV (autumn 2013) ^a				Experiment V (spring 2014) ^a			
	<i>Calonectria polizzii</i>		<i>Calonectria pauciramosa</i>		<i>Calonectria polizzii</i>		<i>Calonectria pauciramosa</i>		<i>Calonectria polizzii</i>		<i>Calonectria pauciramosa</i>	
	VIF	TIF	VIF	TIF	VIF	TIF	VIF	TIF	VIF	TIF	VIF	TIF
Divapan (400)	0	0	0	0	0 a	0 a	0 a	0 a	0 a	0 a	0 a	0 a
Divapan (350)	0	0	0	0	0 a	0 a	0 a	0 a	0 a	0 a	0 a	0 a
Divapan (250)	0	0	0	0	9.9 b*	0 a*	24.7 b*	0 a*	0 a	0 a	0 a	0 a
Basamid (200)	0	0	0	0	0 a	0 a	0 a	0 a	0 a	0 a	0 a	0 a
Basamid (160)	0	0	0	0	0 a	0 a	0 a	0 a	0 a*	19.1 b*	0 a	0 a
Basamid (100)	0	0	0	0	13 b ^{ns}	17.9 b ^{ns}	36.4 c*	9.3 b*	1.2 a*	20.4 b*	18.5 b ^{ns}	16.7 b ^{ns}
Control	100	100	100	100	100 c	100 c	100 d	100 c	100 b	100 c	100 c	100 c

^a Each value represents the mean of three replicates, each constituted by 54 infected carnation pieces. Recovery values followed by different letters within each column or by * within each row are significantly different according to Fisher's least significant difference test ($\alpha = 0.05$)

Table 8. Disease incidence (%) of crown and root rot caused by *Calonectria pauciramosa* and *Calonectria polizzii* on red clover seedlings grown in peat substrate mixed with carnation leaf pieces colonized by pathogen microsclerotia retrieved from both fumigated and untreated plots.

Treatment & rate (liters or kg/ha)	Infectivity of <i>Calonectria polizzii</i> ^a						Infectivity of <i>Calonectria pauciramosa</i> ^a					
	Experiment III		Experiment IV		Experiment V		Experiment III		Experiment IV		Experiment V	
	VIF ^b	TIF	VIF	TIF	VIF	TIF	VIF	TIF	VIF	TIF	VIF	TIF
Divapan (400)	0.0	0.0	0.0 a	0.0 a	0.0 a	0.0 a	0.0	0.0	0.0 a	0.0 a	0.0 a	0.0 a
Divapan (350)	0.0	0.0	0.0 a	0.0 a	0.0 a	0.0 a	0.0	0.0	0.0 a	0.0 a	0.0 a	0.0 a
Divapan (250)	0.0	0.0	15.8 b*	0.0 a*	0.0 a	0.0 a	0.0	0.0	28.6 b*	0.0 a*	0.0 a	0.0 a
Basamid (200)	0.0	0.0	0.0 a	0.0 a	0.0 a	0.0 a	0.0	0.0	0.0 a	0.0 a	0.0 a	0.0 a
Basamid (160)	0.0	0.0	0.0 a	0.0 a	0.0 a*	28.1 b*	0.0	0.0	0.0 a	0.0 a	0.0 a	0.0 a
Basamid (100)	0.0	0.0	17.2 b	20.5 b	4.2 a*	27.6 b*	0.0	0.0	40.5 c*	13.6 b*	21.2 b	23.4 b
Control	88.5	89.6	90.3 c	85.7 c	93.8 b	89.2 c	90.3	87.4	88.4 d	84.6 c	92.8 c	87.2 c

^a Data presented are means of three replications (each consisting of 42 to 55 young red clover seedlings). Arcsine (\sin^{-1} square root x) transformation was used on percentage data prior to analysis; untransformed data are presented. Disease incidence values followed by different letters within each column or by * within each row are significantly different according to Fisher's least significant difference test ($\alpha = 0.05$). Infectivity levels of fumigated pathogen microsclerotia are always compared to those of relative controls.