Methods to detect fungi in seeds of the *Austronium graveolens*, *Bauhinia forficata* and *Cordia trichotoma* (Vell.)

Métodos de detecção de fungos em sementes de *Austronium graveolens*, *Bauhinia forficata* e *Cordia trichotoma* (Vell.)

Métodos de detección de hongos en semillas de *Austronium graveolens*, *Bauhinia forficata* y *Cordia trichotoma* (Vell.)

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Abstract

The importance of native forest seeds, free of phytopathogens, is a strategy for the implantation of quality forest plantations. The scarcity of information about the health conditions of seeds of native forest species causes the spread of diseases. The objective of this work was to determine a method of sanitary analysis to identify pathogens in seeds of the species of *Austronium graveolens*, *Bauhinia forficata* and *Cordia trichotoma* (Vell.). The treatments consisted of filter paper methods with and without disinfection in the periods of seven and 15 days, 15 days with water restriction, with BDA medium by seven and 15 days
and BDA medium with water restriction during 15 days. For each test, it was used 400 seeds. The results showed that for each forest specie seeds the best detection method was in a Petri dish with culture medium and seed disinfection, the *Austronium graveolens* and *Bauhinia forficata* seeds need an incubation period of 15 days with water restriction and *Cordia trichotoma* (Vell.) seeds of seven days.

**Keywords:** Pathogens; Health; Analizes.

**Resumo**

A importância de sementes florestais nativas, isenta de fitopatógenos, é uma estratégia para a implantação de plantios florestais com qualidade. A escassez de informações acerca das condições sanitárias das sementes de espécies florestais nativas faz com que ocorra disseminação de doenças. Diante disso, o objetivo deste trabalho foi determinar um método de análise sanitária para identificação de patógenos em sementes das espécies de Guaritá, Pata de Vaca e Louro pardo. Os tratamentos consistiram nos métodos de papel filtro com e sem desinfestação, nos períodos de sete e quinze dias, e quinze dias com restrição hídrica, e meio BDA com 7 e 15 dias e meio BDA com restrição hídrica, por 15 dias. Para cada teste utilizou-se um lote de 400 sementes. Os resultados demonstraram que o melhor método de detecção para as espécies foi Placa de petri com meio de cultura e desinfestação das sementes, sendo Guaritá e Pata de Vaca, necessário um período de incubação de quinze dias, com restrição hídrica e Louro pardo, sete dias.

**Palavras-chave:** Patógenos; Sanidade; Análises.

**Resumen**

La importancia de las semillas forestales nativas, libres de fitopatógenos, es una estrategia para plantaciones forestales de calidad. La escasez de información sobre las condiciones sanitarias de las semillas de especies forestales nativas provoca la diseminación de enfermedades. El objetivo de este trabajo fue determinar un método de análisis sanitaria para identificar patógenos en semillas de las especies de *Austronium graveolens*, *Bauhinia forficata* y *Cordia trichotoma* (Vell.). Los tratamientos consistieron en métodos de papel de filtro con y sin desinfección en los períodos de siete y quince días, quince días con restricción del agua, con medio BDA a los siete y quince días y medio BDA con restricción de agua durante quince días. Para cada prueba se utilizaron 400 semillas. Los resultados mostraron que para cada semilla de especie forestal el mejor método de detección fue en placa Petri con medio de cultivo y con uso de la desinfección de semillas. Para las semillas de
Austronium graveolens y Bauhinia forficata es necesario período de incubación de quince días con restricción del agua y para Cordia trichotoma (Vell.) se debe incubar por siete días.

**Palabras clave:** Patógenos; Sanidad; Análises.

### 1. Introduction

The demand for seeds of native species is the result of the need to conserve tropical forests, since they are essential in ecosystem recovery and conservation programs (Carvalho, et al., 2006).

Forest species such as the Austronium graveolens, Bauhinia forficata and Cordia trichotoma (Vell.) Arráb. ex Steud are native and they present wide geographical distribution, with various uses, such as medicinal attributes, civil construction, furniture quality and ornamentation (Cassimiro, et al., 2015, Araújo, et al., 2014, Martins, et al., 2013, Silva, et al., 2016).

In order to obtain quality seedlings is necessary that the seeds have a good genetic and phytosanitary characteristic. The genetic factor needs of evaluations to select, but the phytosanitary characteristic is dependent of the cultural practices. The phytosanitary aspect causes preoccupation because the seed present ability to mobilize over great distances, what it can become a disseminator, mainly of fungi, in places where there is no occurrence of diseases (Schultz, et al., 2015). In addition, the sanitary quality of seeds is one of the most important factors that affect the development of forest crops, due to the microorganisms can cause deterioration in the seeds, as well as subsequent lesions and abnormalities in seedlings, what later difficulty in the implantation of forest stands (Piveta, et al., 2010).

Studies regarding the correct identification of fungi and other pathogens in seeds of native forest species are scarce, but they are necessary. Because of this, this work aimed to obtain the best methods to detect fungi in the Austronium graveolens, Bauhinia forficata and Cordia trichotoma (Vell.) seeds.

### 2. Methodology

The study was carried out at the Laboratório de Fitopatologia, da Universidade Tecnológica Federal do Paraná – Câmpus Dois Vizinhos, Paraná State, Brazil, in the first semester of 2019. The seeds were collected in the year 2018 in forest arrays located at Campo Mourão city- Paraná State, Brazil.
The seeds were collected according to the physiological maturation characteristics of the species with basis in the higher percentage of germination and vigor. Seeds were collected and after they were stored in a cold chamber for 20 days.

Stock were divided, with groups of 400 seeds for submitted the detection methods and incubation time (Table 1).

**Table 1** - Detection methods, times and pre-treatments used in the tests.

| Detection methods | Blotter test | BDA |
|-------------------|-------------|-----|
| Incubation time * |             |     |
| Without disinfection | 7 | 15 | 7 | 15 |
| With disinfection | 7 | 15 |     |     |
| Water restriction |     | 15 | 15 |     |

* In days. Source: Authors (2020).

For the blotter test without disinfection, in the two lots (7 and 15 days of incubation) the seeds were placed in Gerbox boxes with lid. Each box consisted of a repetition with 25 seeds, it being organized with three sheets of filter paper moistened with 7 mL of sterile water, each (Brasil, 2009). In the same method, but with disinfection, in the two lots (7 and 15 days of incubation), a pre-treatment of the seeds was used before the assembly of the test, which consisted of immersing them in a sodium hypochlorite solution at 1 % for one minute, afterwards, placed on absorbent paper to dry during 12 hours.

Subsequently, the seeds were placed in Gerbox boxes with a lid, where the batch of 400 seeds was divided into 16 boxes with twenty-five seeds, each. After their disposal, the paper was weighed, and the total in grams was multiplied by 2.5, obtaining the total in milliliters of sterile water, used to moisten the paper.

For the method of sanitation in culture medium, in both tests (7 and 15 days), the seeds were previously treated with sodium hypochlorite at 1% for one minute and placed on absorbent paper to dry. After treatment, the seeds were placed in Petri dishes with BDA medium (Potato - Dextrose - Agar).

Finally, in tests with water restriction, the seeds were distributed in two tests, blotter test and BDA medium to the level of -1Mpa NaCl (1.72g 150mL-1) (Rey, et al., 2005). For the blotter test, the filter paper used in the Gerbox boxes was weighed and moistened the paper with same rule as in the previous test was used. For the BDA medium, the NaCl
concentration was homogenized with the culture medium during preparation (Rey et al., 2008).

After, the treatments were stored in a BOD-type germinator for specific periods (7 and 15 days), under temperature conditions at 25°C and a photoperiod of 12 hours. Subsequently, the seeds were evaluated with the aid of a stereoscopic microscope, in order to identify the pathogens present, with basis on the evaluation of their development and reproduction structures (mycelium and spores). The entire microflora incident in the tests was evaluated by observation of morphology of their structures and characteristic aspects of their colonies (Bernardi & Rey, 2020, Pereira et al., 2018).

The data obtained were submitted to statistical analysis with the Kruskal-Wallis non-parametric test (p <0.05), to verify whether there were statistical differences between the groups evaluated - fungi incidence.

The statistical test justifies its used, since the data in this study did not have the assumptions of the mathematical model in the parametric field of the completely randomized experimental design (normality of the data and/or homogeneity of the variance of the treatments), even when submitted to transformations by the Sasm-Agri program.

3. Results and Discussion

The fungus present highest incidence in the detection methods tested with the species of Austronium graveolens (Table 2), Bauhinia forficata (Table 3) and Cordia trichotoma (Vell.) (Table 4) was Fusarium sp. This pathogen causes damage in the germination of forest seeds, both in pre and post-emergence (Machado, 1988).

For the Austronium graveolens (Table 2), the genus Rhizopus presented a higher incidence than the other fungi among all tests used, with an average of 83% of infestation. The Bauhinia forficata showed a similar incidence of fungi Fusarium sp. (18%), Penicillium sp. (15%) and Rhizopus sp. (14.5%) in the evaluated seeds (Table 3). The Cordia trichotoma (Vell.) showed 100% of the seeds infested with Rhizopus sp., Fusarium sp. and Chaetomium sp (Table 4).
Table 2. Fungi incidence (%) for A. graveolens seeds according detection method.

| Fungi                    | Detection Methods | Blotter test | Dish Petri® |
|--------------------------|-------------------|--------------|-------------|
|                          | 7 days S/D*       | 15 days C/D**| 15 days C/D| 15 days RH***| 7 days C/D | 15 days C/D | 15 days RH |
| **Alternaria sp.**       | -                 | -            | -           | 25 a         | -          | -           | -          |
| **Aspergillus sp.**      | 18 ab             | 2 ab         | 10 ab       | 12 ab        | 3 a        | 7 a         | 6 ab       |
| **Aspergillus niger**    | -                 | -            | 1 a         | 1 a          | 1 a        | -           | 2 ab       |
| **Botrytis sp.**         | -                 | -            | -           | 1 a          | -          | -           | 1 a        |
| **Cylindrocladium sp.**  | -                 | -            | -           | -            | 9 a        | 1 a         | 5 ab       |
| **Cladosporium sp.**     | -                 | 2 ab         | 3 a         | 3 ab         | -          | -           | -          |
| **Colletotrichum sp.**   | -                 | -            | -           | -            | 3 a        | 34 ab       | 46 bc      |
| **Fusarium sp.**         | 55,5 bc           | 51 cd        | 23 b        | 23 bc        | 71 ab      | 16 a        | 9 a        | 33 abc     |
| **Mucor sp.**            | -                 | 3 abc        | -           | -            | -          | -           | -          |
| **Penicillium sp.**      | 18,5 ab           | -            | 5 ab        | 5 ab         | -          | -           | -          |
| **Phomopsis sp.**        | -                 | -            | -           | -            | -          | -           | -          |
| **Rhizoctonia sp.**      | -                 | 30 bcd       | -           | 1 a          | 31 ab      | 3 a         | 2 ab       |
| **Rhizopus sp.**         | 100 c             | 70 d         | 35 ab       | 59,5 c       | 100 b      | 100 b       | 99 b       | 100c       |
| **Stemphylium sp.**      | -                 | -            | -           | -            | -          | 5 a         | -          |
| **Trichoderma sp.**      | 0,5 a             | -            | -           | -            | 1 a        | 14 a        | -          |

Percentage of incidence followed by lowercase letters in the column differ from each other, at the level of 5% probability, by the Kruskal-Wallis test, at the level of 5% probability. S/D* = No disinfection, C/D** = With 1% NaOCl disinfection, R.H*** = NaCl Water Restriction. Source: Authors (2020).

The Table 2 showed the incubation methods used in this study and the fungi detected in each one, with its respective percentage of incidence. It was noted that there are differences between the methods used, according to the incidence of each fungus.

In all species evaluated, the genus Fusarium was detected. According to Cicarelli Netto et al. (2003), evaluated the physiological and health quality of native seeds, observed that the incidence of *Fusarium* sp. can cause damage in the seedlings produced.

Fantinel et al. (2013) reported that *Fusarium* sp. is a fungus commonly found in forest seeds. This microorganism, in the vast majority of its species is responsible for seed deterioration (Maciel et al., 2012) and damping-off (Lazarotto, et. al. 2012), requiring pathogenicity tests to verify transmission and severity of the same plants (Benetti et al. 2009). The pathogen is easily identified during the tests, due to its high rate of mycelial growth and easily identified characteristics, such as densified hyphae, which may have a white to violet color, facilitating its detection in the tests (Leslie & Summerell, 2006).

The other microorganisms detected, had an incidence below 8%, when compared to all
detection tests (Tables 2, 3 and 4).

### Table 3. Fungi incidence for *B. forficata* seeds according detection method.

| Fungi             | Blotter test | Detection Methods | Dish Petri® |         |
|-------------------|--------------|-------------------|-------------|---------|
|                   | 7 days S/D*  | 15 days S/D       | 7 days C/D**| 15 days C/D | 15 days RH*** | 7 days C/D | 15 days C/D | 15 days RH |
| Alternaria sp.    | 2,5 ab       | -                 | 7 ab        | 4 a      | -             | -          | -          |
| Aspergillus sp.   | 0,5 a        | -                 | 5 ab        | 8 ab     | 1 a           | 4 a        | 3 ab       |
| Aspergillus niger | -            | -                 | 1 a         | -        | -             | 10 ab      |            |
| Bipolaris sp.     | -            | -                 | -           | -        | -             | 0,5 a      | 3 ab       |
| Botrytis sp.      | -            | -                 | 1,5 a       | 1 a      | 2 a           | 7 ab       | -          | 5 ab       |
| Chaetomium sp.    | -            | 0,5 a             | -           | 4 ab     | 1 a           | -          | -          |
| Cylindrocladium sp| -            | -                 | -           | -        | -             | 4 ab       |            |
| Cladosporium sp.  | 2 ab         | 4 ab              | 3 a         | -        | 2 a           | -          | 2 a        | 9 ab       |
| Colletotrichum sp.| -            | -                 | -           | -        | -             | -          | -          |
| Fusarium sp.      | 12,5 b       | 3,5 a             | 17 b        | 17 ab    | 72 b          | 3 ab       | 7,5 a      | 13 ab      |
| Gibberella sp.    | -            | -                 | 1 a         | -        | -             | -          | -          |
| Penicillium sp.   | 2 a          | 41,5 b            | 8 ab        | 26 b     | 2 a           | 19 b       | 9 a        | 13 ab      |
| Pestalotia sp.    | -            | -                 | -           | -        | -             | -          | -          |
| Phomopsis sp.     | -            | -                 | -           | -        | 1 a           | 3 a        | -          | -          |
| Rhizoctonia sp.   | -            | 8 ab              | -           | 1 a      | 13 ab         | 11 ab      | 5 a        | 1 a        |
| Rhizopus sp.      | 15 ab        | 3 a               | 8 a         | 33 b     | 27 ab         | -          | -          | 30 ab      |
| Stemphylium sp.   | -            | 1 a               | -           | -        | -             | -          | -          |
| Verticillium sp.  | -            | -                 | -           | -        | 2 ab          | -          | -          |

Percentage of incidence followed by lowercase letters in the column differ from each other, at the level of 5% probability, by the Kruskal-Wallis test, at the level of 5% probability. S / D * = No disinfection, C / D ** = With 1% NaOCl disinfection, R.H *** = NaCl Water Restriction. Source: Authors (2020).

In Tables 3 and 4 is important to observe the increase in the number of genera of fungi incident on the seeds of *B. forficata* and *C. trichotoma* with use of different detection methods.
Table 4. Fungi incidence for *C. trichotoma* seeds according detection method.

| Detection Methods | 7 day | 15 day | 7 day | 15 day | 15 day |
|-------------------|-------|--------|-------|--------|--------|
|                   | S/D*  | S/D**  | C/D   | C/D**  | RH***  |
| *Alternaria* sp.  | 45 cd | 100 b  | 40 bc | 100 b  | 100 c  |
| *Aspergillus* sp. | 1 a   | -      | 9 ab  | 6 a    | 21 ab  |
| *Aspergillus niger* | -    | -      | -     | -      | 1 a    |
| *Bipolaris* sp.   | -     | -      | -     | -      | -      |
| *Botrytis* sp.    | 20 bcd| 1 a    | 15 ab | 33 ab  | 11,5 ab|
| *Chaetomium* sp.  | -     | -      | -     | -      | 100 c  |
| *Cylindrocladium* sp. | -    | -      | -     | -      | -      |
| *Cladosporium* sp. | 6 abc| 1 a    | -     | -      | -      |
| *Colletotrichum* sp. | -   | -      | -     | -      | -      |
| *Fusarium* sp.    | 100 d | 100 b  | 100 c | 100 b  | 100 b  |
| *Penicillium* sp. | 5 ab  | -      | 17 abc| 4 a    | 2,5 a  |
| *Rhizoctonia* sp. | -     | -      | -     | -      | 1 a    |
| *Rhizopus* sp.    | -     | -      | 1 a   | -      | 20 a   |
| *Stemphylium* sp. | -     | -      | -     | -      | 15 ab  |
| *Verticillium* sp. | -    | -      | -     | -      | 4 a    |

The method that proved to be inappropriate for the detection of fungi in all species was the blotter test with 7 days and without disinfestation, because it was observed that it easily detected fast-growing fungi like *Rhizopus* sp. and *Fusarium* sp. and without satisfactory results in the detection of fungi with slower vegetative development and in most phytopathogens.

In this case, the lack of pre-treatment in the seeds may have facilitated the development of hyphae that prevented the incidence of other fungi. However, Mendes et al. (2011) evaluated seeds of *Leucaena leucocephala* (Lam.) with and without disinfestation concluded that the use of previous disinfestation in the health test effectively reduced the detection of fungi, due to some of the detected microorganisms to be saprophytes and to stay on the seed surface, what it was different of results obtained in this work.

This observation was corroborated by Rey et al. (2008), where with the water restriction technique does not alter the existing microflora and incident fungi in the seeds of *Phaseolus vulgaris*. In addition, when a non-toxic restrictor was used, which delays seed germination and maintain the viability, such as NaCl, the secondary contamination was
prevented. The same, it still facilitates the evaluation of the fungi present in the test, it does not affect the seeds, it does not lead to senescence (Parisi, et al., 2019).

Regarding the total microflora, which infested or infected the seeds (data not presented), in the *Austronium graveolens* seeds, in the petri dish with culture medium method, during two incubation times, there were occurrence of 8 genera. The petri dish method with 15 days and water restriction favored the incidence of 13 genera in the *Bauhinia forficata* seeds, and the same method, however with 15 days and with disinfestation, detected 10 genera in the *Cordia trichotoma* (Vell.) seeds.

In this study, when the averages of incidence were submitted to the Kruskal-Wallis test for comparison with each other, in the species of seeds studied, there was no statistical difference among the different species of fungi evaluated. However, the results showed an ease of some methods in detecting certain phytopathogenic fungi. In the *Austronium graveolens* seeds, when the petri dish method with BDA medium was used at different times, the fungus *Cylindrocladium* sp., responsible for leaf spots and necrosis in forest species (Schultz et al., 2015), was detected. The same occurred for the *Bauhinia forficata* and *Cordia trichotoma* seeds, what demonstrated the ease of incidence of the pathogen in the method. Using the same test, the fungus *Colletotrichum* sp., agent of anthracnosi in several forest species (Carmo, et al., 2017), was observed in the *A. graveolens* and *B. forficata* seeds. These fungi are essentially dependent on high humidity (Alfenas et al., 2009), due to this, their preference for this method, where the seeds are in direct contact with the culture medium. It was noted that the use of the water restrictor in the blotter test prevented the occurrence of fungi in the test. Due to the presence of NaCl, there was a decrease in the absorption of available water in the paper (substrate) used, delaying germination, however, preventing the development of some fungi (Oliveira et al., 2011). However, when the same restrictor was added to the culture medium in a petri dish, with disinfestation and incubation for 15 days, there were the occurrence of genera potentially phytopathogenic, as Cylindrocladium (*Austronium graveolens*), Colletotrichum (*Austronium graveolens* and *Bauhinia forficata*), Bipolaris (*Bauhinia forficata*).

In addition to the occurrence of potentially pathogenic fungi such as *Fusarium* sp., (Leslie & Summerell, 2006), the genus *Rhizopus* sp. was detected with medium to high incidence in the seeds. It is considered as storage pathogen and it is associated with the deterioration of seeds, with action dependent on the physical and physiological conditions (Filho, et al., 2004). Martinelli-Seneme et al. (2006) detected fungi *Aspergillus* sp., *Cladosporium* sp., *Colletotrichum* sp., *Fusarium moliniforme*, *Penicillium* sp., *Rhizoctonia*
solani and Rhizopus stolonifer in the Bauhinia forficata seeds through blotter test with and without disinfestation by 2% NaOCl during 5 min, 10 min and 20 min, what it corroborated with the results of this work.

In tests for the detection of Cordia trichotoma (Vell.) seeds, the fungus Fusarium sp. was in 100% of seeds, what it demonstrated efficiency with the blotter test method for this native species. Similar results were described by Silva & Muniz (2003), evaluated seeds of the species collected in Rio Grande do Sul State, with a high incidence of Fusarium sp.

According to Dhingra (1985), when phytopathogenic fungi are already established in the dormant mycelium inside the seeds, they can develop, even if the amount of water is less than that required for normal seed metabolism. However, easily identified characteristics of this imperfect fungus, such as densified hyphae, which it can have a white to violet color, and a high mycelial growth rate, facilitate their detection in tests (Leslie & Summerell, 2006).

In this work, it was seen that the methods of detecting pathogens in seeds, together with the incubation time, vary among forest species. It was also noted, in agreement with data already published (Parisi, et al., 2016, Walker, et al., 2013), that fungi that are on the surface of the seeds may not cause disease and be present, only infesting them. However, they can reduce the germination or lost the viability (Costa, et al., 2016, Bernardi, et al. 2018). It was obtained that among the evaluated seeds, the method that most detected fungi, both saprophytic and phytopathogenic, was the medium in a petri dish with 15 days and water restriction (Table 2).

The interference of pathogens related to seeds can promote losses in production due to the reduction of the plant population, with increase of diseases in areas, until then, healthy. Thus, the study with fungi detected in forest seeds, frequency of incidence is important since they may provide subsidies for the production of quality seedlings.

4. Conclusion

The best method of detecting pathogens in seeds of each specie was a petri dish using B.D.A. Regarding the incubation time Austronium graveolens and Bauhinia forficata seeds needed 15 days with the addition of water restriction, while Cordia trichotoma (Vell.) seeds had a higher incidence at seven days, without using a water restrictor, both with disinfestation of the seeds.

The study demonstrated the importance of different methodologies used for the detection of phytopathogenic fungi in the forest pathology. The detection of fungi for each
Specie has an easy incidence in different methods used, which it implies with transmission of diseases and consequently problems in forest nurseries.

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