

GERMINATION *IN VITRO* DE JABUTICABEIRA *Myrciaria jaboticaba* (VELL.) BERG

ABSTRACT

Aims: The present work aimed to determine the influence of antibiotic use on seed germination and development of jaboticabeira (*Myrciaria jaboticaba*) seedlings grown *in vitro*.

Study design: The experiment was conducted in a completely randomized design, where the treatments were composed of two types of culture medium and three forms of antibiotic use.

Place and Study duration: The experiment was carried out at the Laboratory of Cell Biology and Culture of Vegetable Tissues (LABCULTIVE), at the Department of Biological Sciences (DCB), at the Agricultural Sciences Center (CCA), Federal University of Paraíba (UFPB), from November 2016 until May 2017.

Methodology: The fruits of jaboticabeira were harvested from a matrix plant and the seeds were removed manually, with subsequent elimination of the pulp and removal of the tegument. They underwent a disinfestation procedure in 70% alcohol and sodium hypochlorite and grown in culture medium.

Results: The highest germination average was obtained when the seeds were soaked for 24 hours in autoclaved water + antibiotic and when placed in liquid medium. In all analyzed variables the liquid medium provided better means. There was no statistical difference in any of the variables analyzed in relation to the use of the antibiotic in the imbibition and the non-use of the antibiotic.

Conclusion: The seeds of *Myrciaria jaboticaba* have greater germination and better development in the liquid culture medium; the presence of the antibiotic in the culture medium probably caused phytotoxicity, thus compromising the germination.

Keywords: Antibiotic; endogenous contamination; polyembryony; recalcitrance

1. INTRODUCTION

The jaboticabeira (*Myrciaria jaboticaba*) originates from the Atlantic Forest, more precisely from the Center-South of Brazil, belongs to the family Myrtaceae and to the genus *Myrciaria* [1]. One of the forms of multiplication of the jaboticabeira is via seminiferous, however, it can also be propagated by grafting [2] and air layering [3], yet, both are less used methods due to the difficulty of rooting [2].

The jaboticaba seeds besides initiating their germination late, also present uneven germination, causing a setback in the species' perpetuation, damaging mainly the production of seedlings. According to [4], the jaboticabeira seeds germination can begin from 10 to 40 days after sowing, depending on the conditions in which they are found.

In vitro culture is a technique that has been used in large scale in the production of seedlings of several fruit species. By using this technique, the seedlings develop in aseptic conditions, free of pathogens, being therefore a market that has been growing exponentially, since the producers look for seedlings that do not compromise the good formation of the orchard.

One of the most widely used tissue culture techniques is micropropagation, since it allows large-scale rapid multiplication of plants with superior agronomic characteristics [5], however, it is necessary to avoid microbial contamination through preventive measures, so that success in *in vitro* propagation. In some cases, there is a need to add antibiotics to the

29 culture medium for the microorganisms control [6];[7], since competition between the
30 explants and microorganisms occurs by the components of the culture medium, which can
31 lead to the plant material death [8].
32 The darkening of the explants has been related to the release of phenolic compounds during
33 the excision of the plant, which may inhibit its development and lead to death. [9] In addition,
34 some plant materials, usually those with woody characteristics, have a common problem that
35 is the oxidation [10]. Direct contact with the culture medium can affect the development of
36 the explant, so it is used "bridges" that act as a link between the explant and the liquid
37 culture medium, that is without addition of gelling agents, such as agar and the phytagel,
38 which also provides the decrease in the production costs of the culture media. The present
39 work aimed to determine the influence of antibiotic use and consistency of the culture
40 medium on the germination and development of jaboticabeira (*M. jaboticaba*) cultivated *in*
41 *vitro*.

42 43 **2. MATERIAL AND METHODS**

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45 The experiment was developed in the Laboratory of Cell Biology and Culture of Vegetable
46 Tissues (LABCULTIVE), in the Department of Biological Sciences (DCB), Center for
47 Agrarian Sciences (CCA), Federal University of Paraiba (UFPB).

48 49 **2.1. *Myrciaria jaboticaba* Seed Preparation**

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51 The jaboticabeira's fruits were harvested from a matrix plant located in the Macacos site,
52 located in the rural area of the city of Areia - PB. They were washed in running water to
53 remove excessive impurities, leaving only those with adequate phytosanitary characteristics
54 and no physical damage.

55 The seeds were manually removed from the fruits and the pulp was removed by washing
56 them with running water, with subsequent drying of the seeds at room temperature in the
57 shade. After two days the tegument was removed and the seeds underwent a disinfection
58 procedure, washed three times with autoclaved distilled water, then immersed in 70%
59 alcohol shaking for 30 seconds, and then washed three times in autoclaved distilled water,
60 followed by immersion in 0.63% sodium hypochlorite solution, in the latter, there was
61 mechanical agitation for 20 minutes and finally they were washed three more times with
62 autoclaved distilled water.

63 64 **2.2. Culture Media Preparation**

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66 The culture medium used was the ½MS [11]. The culture medium pH was adjusted to 5.8
67 before inclusion of 2.0 g L⁻¹ of activated carbon and 7.0 g L⁻¹ of Sigma® agar, the latter has
68 been used only in treatments 2, 4 and 6. The culture media were then autoclaved at 120 °C
69 and 1.5 atm for 20 minutes.

70 71 **2.3. Treatments**

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73 The treatments were arranged as follows:

74 Treatment 1 - After asepsis, the seeds were put to soak for 24 hours in autoclaved distilled
75 water. Afterwards, they went through the disinfection process again, following the
76 methodology described in item 2.1, and in this case, after mechanical agitation for 20
77 minutes, the washing with autoclaved distilled water occurred in the laminar flow chamber,
78 as well as the seeds transfer to tubes (Vinyl Polychlorite) with filter paper and 5 ml of liquid
79 culture medium;

80 Treatment 2 - The methodology used was identical to the previous treatment, but the seeds
81 were transferred to test tubes containing semi-solid culture medium with 5 mL;

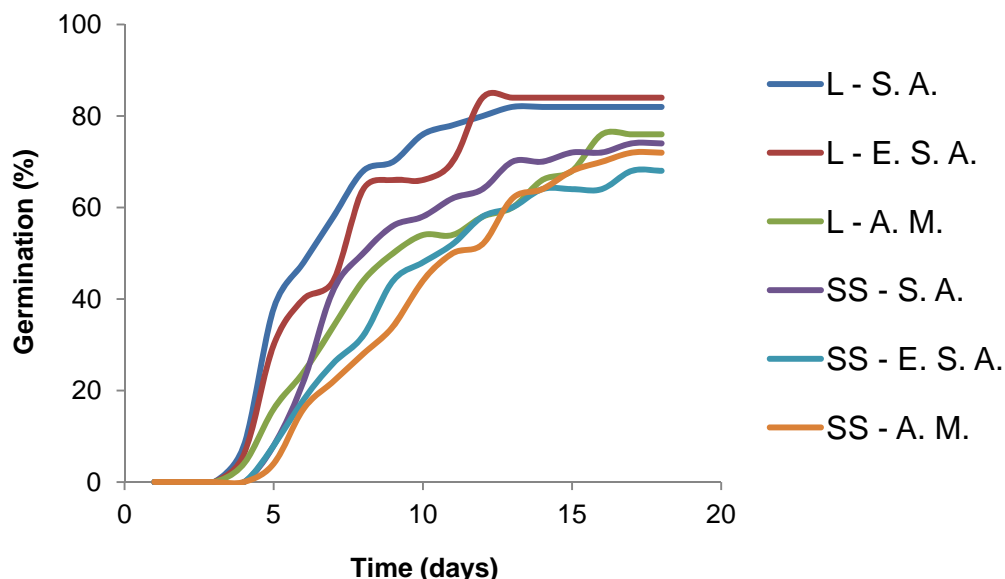
82 Treatment 3 - After asepsis, the seeds were put to soak for 24 hours in autoclaved distilled
83 water using an antibiotic capsule amoxicillin 500 mg L⁻¹ in imbibition. Then, the seeds
84 passed again through the disinfestation process, according to treatment 1;
85 Treatment 4 - The methodology used in this treatment was identical to the previous
86 treatment, but the seeds were transferred to test tubes containing 5 mL of semi-solid culture
87 medium;
88 Treatment 5 - After asepsis, the seeds were put to soak for 24 hours in autoclaved distilled
89 water. Afterwards, they went through the disinfestation process according to treatment 1, in
90 which case the liquid culture medium contained an amoxicillin 500 mg L⁻¹ antibiotic capsule;
91 Treatment 6 - The methodology used in this treatment was identical to the previous
92 treatment, but the seeds were transferred to test tubes containing 5 mL of semi-solid culture
93 medium.
94 All cultures were kept in the growth room in the presence of light with photoperiod of 16
95 hours and temperature of 25 ± 2 °C.
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97 **2.4. Experimental Design and Evaluations**

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99 The experiment was conducted in a completely randomized design, in a factorial scheme
100 2x3 (Culture media x Antibiotic conditions), totaling 6 treatments with 5 replicates. Each
101 repetition consisted of the average of 10 tubes.
102 Evaluations were carried out daily, where the percentage of germination that was obtained
103 after the beginning of the test installation was evaluated, by calculating the number of normal
104 seedlings obtained according to the Rules for Seed Analysis [12], the percentage of
105 oxidation, polyembryony and contamination.
106 For seed vigor analysis, the germination speed index (IVG) was evaluated and calculated
107 according to the formula proposed by [13] where: $IVG = G1 / D1 + G2 / D2 + \dots Gn / Dn$.
108 When the seedlings were 5 to 13 cm in length, the length of the largest root, shoot length,
109 number of leaves and the presence of stem branching were evaluated. The data were
110 submitted to analysis of variance and the means were compared by the Tukey test at 5%
111 probability using the statistical software SAS University 3.4.
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113 **3. RESULTS AND DISCUSSION**

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115 The germination of Sabará jaboticabeira seeds cultivated *in vitro* was initiated on the fourth
116 and fifth day after sowing using the liquid and semi-solid culture medium, respectively
117 (Figure 1), which is a very expressive precocity when compared to the studies on the *ex vitro*
118 germination as found by [14], when germination of the Sabará jaboticaba seeds occurred in
119 20 days after sowing in substrate composed of vegetable soil + vermiculite. In the work done
120 by [15], germination of Sabará and Cambinho jaboticaba seeds with a diameter of less than
121 6mm using Plantmax® substrate started at 25 and 27 days, respectively. According to [16]
122 the germination of Paulista and Cabinho jaboticaba seeds placed in individual Petri dishes
123 containing Germitest paper started seven days after sowing when exposed to 24 and 32 ° C
124 and when treated with fungicide solution (Benlate 500 - 15 g L⁻¹). [17] evaluating the effect of
125 maturation stage and substrate in *ex vitro* conditions on Sabará jaboticabeira, observed that
126 the germination started 18 days after sowing. The anticipation of the germinative process is
127 related to the removal of the integument, since this structure involves the embryo and it must
128 break the integument to start the germinative process, however, as this structure was
129 removed from the seeds, germination occurred more quickly.
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131 **Fig. 1. Germination curve of the six treatments. L - liquid medium; SS - semi-solid**
 132 **medium; S.A. - without use of the antibiotic; E.S.A. - imbibition of seeds in the**
 133 **antibiotic; A.M. - antibiotic in culture medium.**
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 136 There was no statistical difference regarding the use or not of the antibiotic in the semi-solid
 137 medium, but the same did not occur in the liquid medium, since when using antibiotic in the
 138 culture medium, the germination average was lower when compared to other means (Table
 139 1). This fact probably occurred due to the fact that, when coming in contact with the medium,
 140 some explants may have their development affected, due to the restrictions in the absorption
 141 rate of the nutrients in the semi-solid medium, as well as the antibiotic may have provided
 142 phytotoxicity, inhibiting the seed development.

143 The highest germination average (84%) was obtained by using liquid culture medium and
 144 seeds imbibed in antibiotics. In the work done by [18] with *sucupira-branca*, which is also a
 145 woody species, using liquid culture medium, the germination obtained was 95% when the
 146 tegument was removed from the seed and 80% when the tegument was sectioned.

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 155 **Table 1. Germination of the seeds of jaboticaba (*Myrciaria jaboticaba*) according to**
 156 **the type of culture and use of antibiotic or not. S.A without use of the antibiotic;**
 157 **E.S.A: seed embedding in the antibiotic; A.M.: antibiotic in culture medium.**

Type of Medium	Germination (%)		
	S.A.	E.S.A.	A.M.
Liquid	82 aA	84 aA	62 aB
Semi-Sólido	74 aA	68 bA	74 aA
CV(%)	15,60		

158 Averages followed by distinct letters, uppercase in the row and lowercase in the column
159 differ from each other by the Tukey test at 5% probability.

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161 All variables presented higher averages when using liquid medium, since the use of
162 "bridges" prevented the direct contact of the explant with the culture medium and
163 consequently the explant had a better development (Table 2). In relation to the type of
164 medium, there was significance for length of the largest root, with averages of 3.82 and 3.55
165 cm in the liquid and semi-solid medium, respectively, whereas for the shoot length there was
166 significance only when liquid medium was used. [15] using Sabará seeds with 6 mm and
167 larger than 8 mm in diameter, seeded in Plantmax substrate, obtained plant height averages
168 of 2.18 cm and 2.73 cm, respectively. [19] obtained the size of the seedlings of gabirobeira
169 grown *in vitro* for 60 days of 17.67 mm.

170 Regarding the number of roots and number of leaves, there was no statistical difference in
171 relation to the type of medium. The germination speed index (IVG) for seeds in liquid culture
172 medium was significant, with an average of 0.568.

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174 **Table 2. Number of roots and leaves, length of the largest root and shoot, germination**
175 **velocity index (*Myrciaria jaboticaba*) as a function of the two medium types.**

Type of Medium	Number of roots	Number of leaves	Length of largest root (cm)	Length of shoot (cm)	IVG
Liquid	0,80 a	5,60 a	3,82 a	5,69 a	0,568 a
Semi-Sólido	0,65 a	4,72 a	3,55 a	4,25 b	0,475 b
CV(%)	33,23	36,55	38,60	36,79	23,45

176 Averages followed by distinct letters differ from each other by the Tukey test at 5%
177 probability.

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179 The number of leaves was significant when the antibiotic was not used, obtaining an
180 average of 6.58, and when the seeds were soaked in the antibiotic, with an average of 5.64
181 (Table 3). In a study carried out by [15] Sabará jaboticaba seeds classified between 6-8 mm,
182 they obtained leaf number 1.78 after 46 days of cultivation. [2] when using Sabará
183 jaboticabeira stem as explant, they obtained the number of leaves of 4.2. [19] obtained the
184 number of leaves of gabirobeira grown *in vitro* for 60 days of 1.4. [20] observed that the
185 addition of rifampicin in the culture medium was phytotoxic at concentrations of 0.5 and 1.0 g
186 L⁻¹.

187 According to [21] high concentrations of antibiotic added to the culture medium can cause
188 phytotoxicity and may be a limiting factor for the development of the explants. Phytotoxic
189 action generally occurs due to disturbances of protein synthesis and inhibitory action in the
190 synthesis of RNAs and ATPs, with interference, in the energy systems of the plant [22]. In
191 relation to the number of roots and length of the aerial part, there was no significant
192 difference as a function of the antibiotic use or not.

193 The largest mean of the IVG and length of the largest root of *M. jaboticaba* was obtained
194 when the antibiotic was absent in the medium (S.A.), however, it did not differentiate when
195 the seeds imbibition was done with the said product (E.S.A.). [23] sowed totally cleaned
196 jaboticaba seeds on a substrate composed of Florestal Plantmax[®] (50% v/v) + sieved
197 organic compound (30% v/v) + vermiculite of medium granulometry (20% v/v), and they
198 obtained IVG of 1.12, when the seeds were with the attached endocarp, they obtained the
199 IVG of 0.98. [14] using semi-solid culture medium in the germination of Sabará jaboticaba
200 seeds they obtained IVG of 0.32. It is worth mentioning that the higher the IVG, the higher
201 the daily germination speed.

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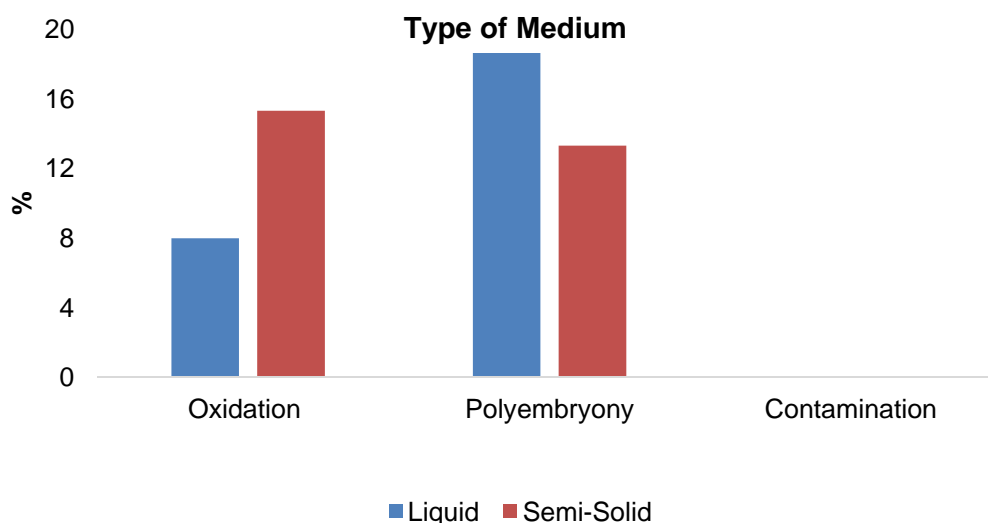
203 **Table 3. Number of roots and leaves, length of the largest root and shoot, germination**
204 **velocity index (IVG) of (*Myrciaria jaboticaba*) due to antibiotic use or not. S.A. = No**

205 use of antibiotic; E.S.A. = Soaking seeds in the antibiotic; A.M. = Antibiotic in the
 206 culture medium.

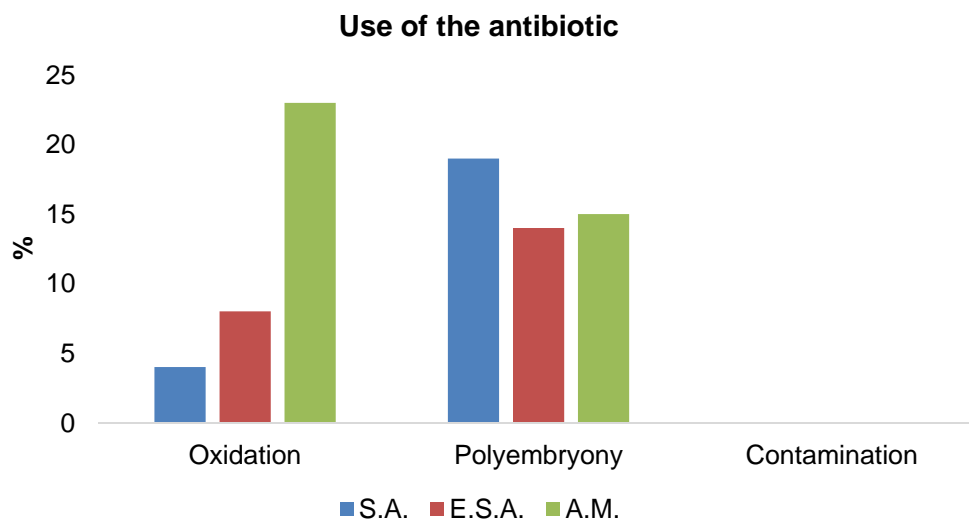
Use of the antibiotic	Number of roots	Number of leaves	Length of largest root (cm)	Length of shoot (cm)	IVG
S.A.	0,84 a	6,58 a	4,51 a	5,58 a	0,606 a
E.S.A.	0,71 a	5,64 a	3,84 ab	5,63 a	0,530 ab
A.M.	0,63 a	3,27 b	2,69 b	3,71 a	0,429 b

207 Averages followed by distinct letters differ from each other by the Tukey test at 5%
 208 probability.
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210 Regarding the type of medium, the percentage of oxidation was higher when semi-solid
 211 medium was used, and the percentage of polyembryony was higher when the liquid medium
 212 was used (Figure 2). The development of the explant may be influenced by the type of
 213 nutrient medium, explant type [24], but also by the addition of some components such as
 214 antibiotic and fungicide. As regards the use of the antibiotic, the percentage of oxidation was
 215 higher when it was used in the culture medium, possibly causing toxic effect and modifying
 216 the morphogenetic characteristics of the explant (Figure 3). The percentage of polyembryony
 217 was higher when the antibiotic was not used, and there was no contamination in any of the
 218 treatments. According to [21] one of the factors limiting the use of antibiotics in the medium
 219 is the phytotoxicity of these substances, mainly due to the common use of high
 220 concentrations.
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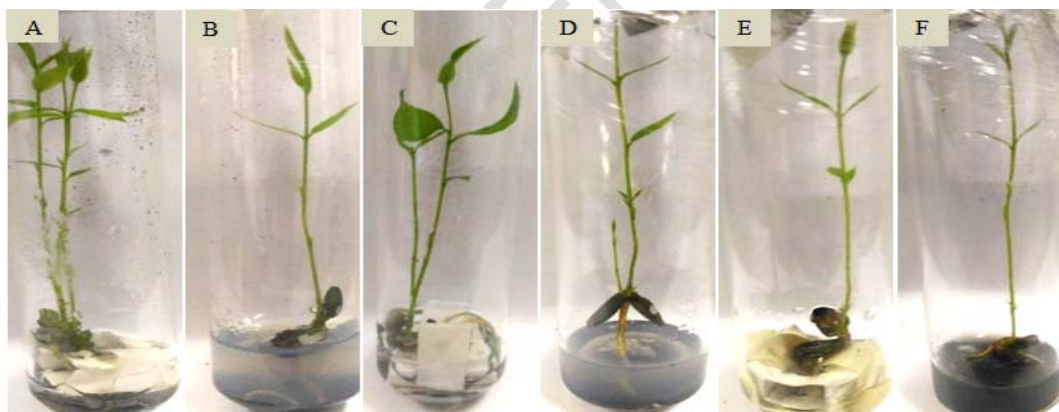
222 Fig. 2. Percentage of oxidation, polyembryony and contamination of (*Myrciaria*
 223 *jaboticaba*) as a function of the two types of medium.
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Fig. 3. Percentage of oxidation, polyembryony and contamination of (*Myrciaria jaboticaba*) due to antibiotic use or not. S.A. = No use of antibiotic; E.S.A. = Soaking seeds in the antibiotic; A.M. = Antibiotic in the culture medium.

Figure 4 shows seedlings of the six treatments at 45 days after seed inoculation, where germination and seedling development occurred uniformly in all treatments, obtaining normal and healthy seedlings.



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Fig. 4. *Myrciaria jaboticaba* seedlings obtained *in vitro* culture. Liquid medium without antibiotic (A); Semi-solid medium without antibiotic (B); Liquid medium, seeds soaked in the antibiotic (C); Semi-solid medium, seeds embedded in the antibiotic (D); Liquid medium with antibiotic in the medium (E); Semi-solid medium, with antibiotic in medium (F).

4. CONCLUSION

The seeds of *Myrciaria jaboticaba* have greater germination and better development in the liquid culture medium. The germination of *M. jaboticaba* occurred on the fourth and fifth day after sowing in the liquid and semi-solid, respectively. The presence of the antibiotic in the culture medium probably causes phytotoxicity, thus compromising the germination and development of *M. jaboticaba* seedlings.

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COMPETING INTERESTS

Authors have stated that there are no competing interests.

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